MoU with The Swiss Federal Research Institute WSL Switzerland

Activity done under MOU between HNBGU, Srinagar Garhwal and WSL, Switzerland

MOU between two institutions (i.e. HNBGU & WSL) came into existence on 20 July 2020 after signing by the Heads and Group Leaders of the both institutions. The main objectives of the MOU were to encourage visits by faculty for the purpose of engaging in research and educational activities, to support the exchange of Master and Doctoral students, to foster the exchange of academic publications and scholarly information and to develop joint research activities & to promote other academic activities, which enhance the above mentioned goals. With this brief background, the following progress has been made:

1. Online meetings and discussion on glacier studies

Soon after the MOU, we held several virtual meetings regarding the course of action on the objectives. However due to COVID-19, it was impossible for WLS Scientists to visit Indian Himalayan glaciers for the field work in the year 2020 and even in 2021.

2. Joint field work on one of the Indian Himalayan glaciers

A group of four European Research Scholars/Scientists lead by Dr. Marin Kneib (WSL) conducted fieldwork on Satopanth Glacier, Chamoli District Uttarakhand, along with HNBGU Glaciology research team during September 2022. During this field trip, various instruments (weather station, time lapse camera and pressure sensors) were setup and valuable scientific datasets including ice melting, air temperature, pressure and wind speed were observed at an altitude of 4300 m amsl over Satopanth Glacier. Some glimpse of the glacier field work is shown below.



Photograph: 1- Swiss team at Geology Department, HNBGU Srinagar; 2-Team at Mana (Badrinath) before going to Satopanth Glacier; 3- Temperature sensor; 4- Time lapse camera; and 5- Researchers working on Satopanth Glacier.

3. Setting up new proposal and funding acquisition

During the period of MOU with WSL, we have formulated a joint research proposal to study the behaviour of Himalayan and Swiss glaciers under changing climate in response to a call of the Ministry of Earth Science, Govt. of India, and the Swiss National Science Foundation (SNSF), Switzerland. The project is funded now entitled "Understanding and modeling the interactions between Debris and glacier Ice in a changing ClimatE (D-ICE)".

The above research proposal under the "Cryopsheric Modelling" theme is led by Prof. H C Nainwal (as Indian PI) and Prof. Andreas Vieli, University of Zurich, Switzerland (as Swiss PI). The other project partners of the projects are Dr. Argha Banjeree & Dr. Arjun Datta (Indian Institute of Science Education and Research, Pune), Dr. Francesca Pellicciotti (Swiss Federal Research Institute WSL, Switzerland), Dr. Atanu Bhattacharya (JIS University, Kolkatta) Dr. Bharath Shekar (Indian Institute of Technology, Bombay), Dr. Madhu Sudan Sati & Dr. Alok Sagar Gautam (HNBGU Srinagar), Prof. Ramachandran Shankar (The Institute of Mathematical Sciences, Chennai), Dr. Guillaume Jouvet (University of Zurich, Switzerland) and Dr. Tobias Bolch (University of St. Andrews, Great Britain and Northern Ireland).

MoU with University of Copenhagen, Denmark

Department of Zoology, HNB Garhwal University

MoU with University of Copenhagen, Denmark (11.10.2019)

- 1. Under Indian-Danish Network project a Workshop on "Arctic and Alpine Aquatic Science-implications of climate change on proglacial freshwater ecosystems functioning and services" (October 11-22, 2019).
- 2. Field visit to Lake Satopanth; Sampling in lake and streams in the Alaknanda Valley at Badrinath.
- 3. Due to Covid-19 Pandemic further activities under the MoU could not be undertaken as the grant was terminated.

MoU with National Bureau of Fish Genetic Resources (NBFGR), Lucknow.

- Ph.D. Thesis entitled "Evaluation of season specific differential gene expression in the Snow trout (*Schizothoraxplagiostomus*) Testis by Next Generation Sequencing (NGS)" submitted by Ms. Shriya Purohit (LZ-17196); Supervisor: Dr. Indrashis Bhattacharya, Assistant Professor, Department of Zoology; Co-Supervisor: Dr. Mahender Singh, Principal Scientist, NBFGR, Lucknow
- Mr. Rakesh Kumar (LZ-19070) registered for Ph.D. on the topic "A study of the bacterial pathogens in some Schizothoracine fish species inhabiting the Gangariver system in Uttarakhand"; Supervisor: Prof. O.P. Gusain, Department of Zoology; Co-Supervisor:Dr. Gaurav Rathore, Principal Scientist & Head, Fish Health ManagementDivision, NBFGR, Lucknow.
- 3. Ms. Yasmeen Kousar (LZ-20101) registered for Ph.D. on the topic "", Supervisor: Prof. Deepak Singh, Department of Zoology; Co-Supervisor: Dr. Mahender Singh, Principal Scientist, NBFGR, Lucknow

Research Publications

- Purohit, S., Sharma, P., Bhatt, G., Kothiyal, S., Singh, M., Nautiyal, P., & Bhattacharya, I. (2022) Evaluation of seasonal cyclicity of testicular development in adult Himalayan snow trout, *Schizothoraxplagiostomus*. Zoology Aquaculture Reports 27: 101333, 2352-5134https://www.sciencedirect.com/science/article/pii/S2352513422003295
- Purohit, S., Sharma, P., Kothiyal, S., Singh, U., Nautiyal, P., Singh, M., Bhattacharya, I. (2023). Resolving the phylogenetic relationship of Himalayan snow trout Schizothoraxplagiostomus with other species of Schizothoracine using mitochondrial CO-I and Cyt b genes. Molecular Biology Reports50(4):3927-3933<u>https://doi.org/10.1007/s11033-023-08274-y</u>

wati Nandan Bahuguna Garhwal University, Srinagar (Garhwal), Uttarakhand-246174 वती नन्दन बहुगुणा गढ़वाल विश्वविद्यालय श्रीनगर (गढ़वाल), उत्तराखण्ड– 246 174 Telephone :- 01346-251057-252143 Fax:- 01346-252247 कः / शेक्षणिक / २ ० २ जे 32 5 सेवामें

> Shriya Purohit D/O Shri Pankaj Purohit Phase II, Kargi Chawk, Dehradun

पाठयकम समिति जन्तु विज्ञान विभाग की बैठक दिनांक 27 फरवरी 2020 की मद संख्या 17.06 में लिये निर्णयानुसार एवं कुलपति महोदया के अनुमोदनोपरान्त डा0 महेन्द्र सिंह मुख्य वैज्ञानिक एन0 बीo एफ0 जीo आर0 लखनऊ को आपका सह–शोध निदेशक नियुक्त किया जाता है।

अतः आप नव नियुवल सह शोध निदेशक के निर्देशन में कार्य करना सुनिश्चित करे।

भवदीय, अनुभाग री शैक्षणिक

प्रतिलिपि :

- डा० इंद्रेश भट्टाचार्य, जूलॉजी विभाग, चौरास परिसर, ।
- डा० महेन्द्र सिंह वैज्ञानिक एन० बी० एफ० जी० आर० लखनऊ।
 ३.विभागाध्यक्ष, जूलॉजी विभाग, चौरास परिसर, ।

अनुभाग अधिकारी,शैक्षणिक

Department of Zoology & Biotemology, HNOB Granhural University, 2019 Uttarakeand. the fol Zoologe & Biologologie , FIRST worksite poll 11-moet NAME Institute Name Emgil Mone No. Sognature. Mushaver land HNBGU mushanggue Ognil. Com . 7006599736 \$ Tanuja Bartual HNBUNU tanujabartual 29 0 gmail.com 7895262897 NEE tika Sharma HNBORL neetikal 850 gmail lom @ 9675401657 Poriganka Thaken HNBGW pairjonka082914970 grail com baix MANNIGET swamp 1+NBGU namet. Impgulo & gmail. com, 2439711540 MS O.P. Gusan " opgysam 1964@ genal com RSHANKAR IMSC Shankare Imsc. res. m 944402403 Brakash Nantiyal HNDGU. ADR. pralsash. nautigel @ A. outlook.com A.R. Naulizat HAPPRC arnaulizal comail. com. d' M.C. Nathard HAPPRC Mis mental and com Meny Prakan Gruson + AMBGU mappgusbin & yahoo.co.in 9456791224 mag Sonagor 9412947733 Kanta Rala Govt. P.G. College de Kavitahala Ogmail. con Tal Deeperte lingh HNBGU bhomdasidsdeepie segnt - ty Vander Kumar: Sh. D Scholar a fanderfkge a gnrail com F Pragya Topal - Ire-PhD pragyatopal @ gmail.com Priyanka Rang Pre-PhD. Hanapinka. 97@ gmoul.com a Devendrog Singh 1) rale sal devusquat. do agmail. Com Sana Fatima P.h.D. Scholar sanafatima 1025 Dgmail.com Jahima Ph.P. Scholar HNBGU pretibiotech 88 agmail. com Preet Singh Shrikant Mishing - N. Sc. Zordog 3 sem Kant Zoology MSc. Zoology MSc. 3rd sem akashqisi 16150 Akapt Gin Akash biri Rojer v lochon 88 @ Conail. Con · Kylez Roteen Anipli Patel HMB-PhiDscholar apaga19053@gmail.com Jukesh Saloch Rapern MNBGK - Phild scholard rksalochogmail. com ranajitendra Ki@ gmail. com Sait HNBSU - " Tilenderg S. Rang ALW abhmal (aum dk FOD Abridatik nk & bio. hu. dh dur Nicls knod Univ of Copenhagen brij. Ciwsa agmail lon. A Bayl CIWSA Japan BRIJGODAL Echnstofferse e bro-hu-dk (MD) UNIV. OF COPENHAUR KURSTEN CULLISTUFFICIOSEN anoopkdobsigel @ rodiffmal. Com Pauri Campus Prof A K Dobrigal prof maingle yaho com N Sing Bula campus N. Singh

Name Institute name Name Institute name emailid Ph. No OLE GEERIZ-HALSEN, GREENKAND INSTITUTE OF NATURAL RESOURCES Signature Ph.No. Clackert Ha DEAN JACOBSEN DEPT. BIOLOGY, FBS Djacobsen@bio.Ko.dk form Jan Treetibiolecher grander Prestory Pseiti Sugh Botany HNBGU Sanafaling 102 Sagmand. Sana Patima Botany KNBQU pahang Rohul Negi Microbiology Rahul2071@ grail. com Ridel Anand Kumes High Aquitative Bio liventy akumer. ags @ gmail. cm -Arably -Caurar Shatt Zoology, HNBGU grubhatt 23 10 89 Email. Com CB Zoology Shrikant Mani Partigya Sharma Zoology Partigyasharma 18@ gmail. com Jauling Salanya Joshi Biotechnology jestis asanya @ gmail. com Zoology Sonelithali 14@gmail.com Au Soneli 'khali Shinya Purohit Zoology purchitshnipazit & quaition shope Biotechnology SACHINSINGH Sachin Sachinsingh 123321@gmail- Com Perican Ka Tha la ar Zoology pringanka 08291997 @ gmail. com Raub A kash biri akashqiri 1615@gmail.com Zoology Akapp him Zoology HNBGU. Somila Rayal Boules tree Pro programment (mail- and 16 p. delatas saya alimetass algorial PLA Challen Million Million and abertinities Alia william (200 what will 201 vertor angelight 30 ganger 1 com rplanse and - April R Lest A select - Phil controlation ramafilion to attice pradicion and all alland is al Consultance have a part of the ready and and the 10) la militada antida da unio

हेमवती नन्दन बहुगुणा गढवाल विश्वविद्यालय श्रीनगर गढ्वाल --246174 (उत्तराखण्ड) (केन्द्रीय विश्वविद्यालय)

Hernvati Nandan Bahuguna Garhwal University Srinagar (Garhwal), Uttarakhand- 246174-

(A Central university)



Tel: 01346-25225224 Email:Registrar.hnbgu@gmail.com Website: www.hnbgu.ac.in

पत्रांकः शैक्षणिक / 2022/1056

दिनाक 28/5/2022

सेवा में,

Yasmeen Kousar D/O Mohd Azam Ward No. 13, Nowshera Dist- Rajouri, J&K- 185151

पाठयकम समिति जन्तु विज्ञान विमाग की बैठक दिनॉक 29.04.2022 की मद संख्या 22.03 मे लिये निर्णयानुसार एवं कुलपति महोदया के अनुमोदनोपरान्त Dr. Mahender Singh, Principal Scientist, NBFGR को आपका सह शोध निर्देशक नियुक्त किया जाता है।

अतः आप नव नियुक्त शोध निर्देशक के शोध निर्देशन में कार्य करना सुनिष्टिचल करें।

भवदीय, अनुमाग अधिकारी, शैक्षणिक

प्रतिलिपि : 1. डा0 दीपक सिंह, जूलॉजी विभाग, बिडला परिसर श्रीनगर गढवाल। 2. Dr. Mahender Singh, Principal Scientist, NBFGR 3.विभागाच्यक्ष जूलॉजी विभाग,बिडला परिसर, श्रीनगर गढवाल।

अनुभाग अधिकारी, शैक्षणिक

HEMVATI NANDAN BAHUGUNA GARHWAL UNIVERSITY, SRINAGAR GARHWAL (A CENTRAL UNIVERSITY)

NO: Acad/2021/ 337

Dated: 17.09.2021

To,

Rakesh Kumar S/O Shri Hari Saran C/o Shri Datta Ram Bahuguna, Madhi Chauras Kirtinagar, Tehri Garhwal – 249161

Sub: Registration for Ph.D. Programme of the HNBGU.

In response to your application for Registration to the Ph.D. Programme of the University, the competent authority of the University is pleased to approve your candidature for registration to the Ph.D. Programme of the University as per followings:

TOPIC of Research:"A study of the bacterial pathogens in some Schizothoracine fish species inhabiting the Ganga river system in Uttarakhand"

Decision of the Committee held on 18.02.2021: The Title, synopsis, centre & supervisor is approved.

Name of the Supervisor: Prof. O.P. Gusain

Name of the Co-Supervisor (if any) : Dr. Gaurav Rathore

Subject: Zoology

Centre: Birla Campus, Srinagar

You have been registered since 09.08.2019 and your registration No. is LZ- 19070

By Order Registrar

Copy to:

- 1 Supervisor Prof. O.P. Gusain, Dept. of Zoology, Birla Campus, Srinagar Garhwal.
- 2 Co-Supervisor (if any) Dr. Gaurav Rathore, Head, Fish Health Management & Exotics Division, ICAR NBFGR, Lucknow.
- 3 Head of the Department Prof. P. Nautiyal, Dept. of Zoology, Birla Campus, Srinagar Garhwal.
- 4 Dean of School Prof. A.K. Dobriyal, School of Life Science, HNBGU, Srinagar Garhwal.

Deputy Registrar (Academic)

Instructions:

Your registration to Ph.D. Programme is to be governed as per rules, regulations and Ph.D. ordinances of the University. You are required to submit half yearly (Six months) reports to the Board of Studies, through the supervisor, on the work done by you and the work you proposes to do in the ensuing academic year.

Ph.D. programme shall be for a minimum duration of **three** years, including course work and a maximum of **six** years. The candidate shall be required to complete his/her research work and submit the thesis within a period of **six** years reckoned from the date of his/her enrolment, **the date of taking admission (submitting fees) for the pre-PhD. course.**

Provided that the School Board may, after considering the recommendation of the Board of Studies, in a very special case and for reasons to be recorded, grant further extension, of not more than one year.

Provided further that in case the candidate fails to submit the thesis within the period permitted for the submission of the thesis, including the periods of the extension thereof his/her admission to the Ph.D. programme shall be liable to be terminated and he/she shall, upon such termination, forfeit all the fees and other dues paid by him/her for and during such admission.

The woman candidates and persons with disability (more than 40% disability) may be allowed a relaxation of two year for Ph.D. in the maximum duration.



















MoU with Dr. Ambedkar Foundation, Ministry of Social Justice & Empowerment, GOI



पत्रांक : हे.न.ब.ग.वि.वि. / DALE/2022/0/

दिनांक : 20 /05/2022

NOTIFICATION (/2022)

In compliance to the directions issued by the Ministry of Social Justice and Empowerment, Government of India, New Delhi regading **Dr. Ambedkar Centre of Excellence (DACE)**, Prof. M.M. Semwal, Department of Political Science, HNB Garhwal University is hereby nominated/appointed as "**Programme Coordinator**" of Dr. Ambedkar Centre of Excellence (DACE) in the University. His contact details are as under:

- 1. Mobile No. 9412079266
- 2. E-mail Id. <u>mmsemwal@gmail.com</u>

This issues with the approval of the competent authority.

20/05/20 Registrar

Copy for information and necessary action to:-

- 1. Prof. M.M. Semwal, Department of Political Science, HNB Garhwal University.
- Shri Vikas Trivedi, Director, Dr. Ambedkar International Centre, Ministry of Social Justice and Empowerment, Government of India, New Delhi – 110001.
- 3. Pro Vice Chancellor for kind information.
- 4. All Deans/Hods.
- 5. Campus Directors (Tehri/Pauri/Chauras)/Director IQAC/FDC.
- 6. Finance Officer/Controller of Examination.
- 7. Joint Registrar/All Deputy Registrars.
- 8. In charge System Manager for uploading on the University Website.
- 9. PS to VC, for kind information of the Hon'ble Vice Chancellor.
- 10.Guard file.

Jostan Registrar



Information Brochure

Dr. Ambedkar Centre of Excellence (DACE) H. N. B. Garhwal University, Srinagar Garhwal (A Central University) (Under the Ministry of Social Justice and Empowerment, Govt. of India)

General Guidelines for admission in DACE (2023-24)

Important Information:

- 1. This brochure is only for general guidance for the candidates. The Common Entrance Test (CET) and admission to the DACE program shall be governed by the relevant provisions of the Ministry of Social Justice & Empowerment, Govt. of India.
- 2. The admission to this scheme is suggestive of the fact that the terms and conditions laid down in this brochure are acceptable to the candidate and his/her guardian.
- 3. Benefits under the Scheme (DACE) can be availed by a student not more than twice, irrespective of the number of chances he/she may be entitled to take in a particular competitive examination. The student will be required to submit an affidavit in this regard.
- DACE program is <u>ONLY for the Scheduled Caste and Other Backward Classes (OBC)</u> <u>Students</u> to offer them 'Free Coaching' for the Civil Services/Allied Services Examinations conducted by UPSC/State Public Service Commission and Staff Selection Commission (SSC).
- Out of 100 sanctioned seats, 70% of seats are reserved for SC and 30% of seats are reserved for Other Backward Classes (OBC) Students. As per directions from the ministry 33% of seats in each category shall be filled by eligible female candidates.
- 6. There is No actual fee for applying, admission and coaching (the course fee paid is 100% refundable) under the DACE scheme. Course fees paid at the time of admission will be refunded by the Ministry in Aadhar linked bank account of the student after submission of a receipt. The process of refund of the course fee details will be shared on the University website in due course of time as per directions from the ministry.
- 7. A stipend of Rs. 4000/ (Rupee four thousand) per month will be provided to all the students by the ministry.

- 8. An incentive of Rs 15,000/ (Fifteen thousand) shall be provided to all the successful students to prepare for the interview after being successful in the mains stage of Central Civil Services / State Civil Services Exams for Class 1 and Class 2 posts.
- 9. The candidates enrolled under this scheme shall have to attend all the classes. If any student remains absent (for more than 04 days) without acceptable reasons like medical or household emergencies, his/her candidature for the scholarship shall be cancelled. Biometric attendance based on Aadhar of the students will be recorded and shared with the ministry on a monthly basis. Leaving the coaching midway without prior approval of the competent authority, the expenditure incurred on the candidate will be recovered from the candidate concerned.
- 10. Duration of the course will be one year in case of UPSC/State Civil Services applicants and 6-9 months for SSC candidates.

11. Eligibility Criteria for Admission:

- 1. Only students belonging to SC and OBC categories, having a total family income of Rs. 8.00 lakh or less per annum from all sources will be eligible for benefits under the Scheme.
- 2. SC/OBC candidates belonging to a Minority community are not eligible under this scheme.
- 3. Income Certificate: The income declaration of self-employed parents/guardians should be in the form of a certificate issued by a Revenue Officer, not below the rank of Tehsildar. Employed parents/guardians are required to obtain an income certificate from their employer and submit a consolidated certificate from the Revenue Officer including any other additional source of income.
- 4. The minimum marks in the Graduation degree should be 50% in aggregate.
- 5. The applicants appearing in the last year/ Semester of Graduation are also eligible to apply, but Graduation degree is mandatory at the time of final admission in this programme.
- 6. Only those candidates who have passed the Common Entrance Test (CET), and have graduation degree, would be eligible for admission in DACE. The CET will be conducted by the University as per the directions provided by the Ministry of Social Justice & Empowerment, Govt. of India.
- The Candidates appearing in the final semester/ year of Graduation will not be admitted to the scheme only on the basis of qualifying for the Common Entrance Test (CET). Graduation Degree/mark sheet shall be mandatory at the time of admission.
- Final selection of the candidates for admission to the program will be strictly on the basis of CET merit. Preference may be given to the students of the last batch (2022-23) admitted under the DACE scheme subject to receiving detailed direction from the ministry.

Important Note:

- 1. It shall be the duty of the applicant to produce the prescribed/desired documents for admission. In case of non-production of the required documents, his/her claim for admission shall automatically stand cancelled.
- 2. The applicant is advised to remain vigilant in collecting information regarding the Common Entrance Test (CET), its results and other details, published at the University website and other relevant sources. The University shall not be responsible if the applicant fails to collect such information.
- 3. The information regarding the Common Entrance Test (CET) and admission process will be published on the portal of the University website.
- 4. The information supplied by the applicant in his/her application (Online) shall be final. Any subsequent change in the Online Registration Form will not be allowed. If any information provided by the applicant is found to be false or forged at any stage, his/her admission shall be cancelled.
- 5. The applicant is advised to take the print-out of Online Registration Form and keep it with him/her for future correspondence and reference.
- 6. Any additional information will be published from time to time on the University website.

Admission Procedure:

Admission will be through the Common Entrance Test (CET) conducted by the University. Candidates have to apply online for CET. Date of the entrance test (tentatively third or fourth week of September) shall be notified in due course of time on university website.

Common Entrance Test (CET) Syllabus:

The Admission Test will be of Two Hours duration. The candidate has to answer 100 questions of multiple choice (MCQ) nature. For every correct answer, 02 marks will be awarded. There will a negative marking for attempting wrong answer and 0.25 marks will be deducted for every wrong answer.

The question paper for CET would comprise of following sections, and the questions will be of intermediate standard.

- 1. General English & General Hindi
- 2. General studies related to history, polity, economy, culture, geography etc.
- 3. General Science and Environment
- 4. Reasoning and Mental ability

Counselling and Admission Process:

- 1. Admission will start in the month of the last week of September 2023 on the merit basis of CET and by the process of 'Counselling'.
- 2. The details of information regarding CET and counselling will be available on the University website: <u>www.hnbgu.ac.in</u>.
- 3. The purpose of Counselling is to provide information regarding the availability of category-wise seats on the basis of the merit of CET.

- 4. The candidate will be required to present themselves in person for Counselling and admission.
- 5. If the candidate does not turn up on such date & time along with the required documents, his/her claim for the admission will be automatically cancelled, and the candidate next in merit will be accommodated against such seat.
- 6. All the required documents, mentioned below, are to be submitted at the time of admission, failing which the applicant's claim will not be entertained. The list of documents required is as follows:
 - a. High School or equivalent examination mark sheet and certificate in original along with a photocopy thereof.
 - b. Intermediate or equivalent examination mark sheet and Certificate in original along with a photocopy thereof.
 - c. The graduation Mark sheet/Degree in original along with a photocopy thereof.
 - d. Scheduled Caste/Other backward classes Category Certificate.
 - e. Admit Card and Score card of CET/Qualifying Proof published by the University.
 - f. Proof/Details of earlier enrolment (session 2022-23) to any DACE centres in India, if any.
 - g. Income certificate issued as per provision of the scheme (issued after 31 March 2023).

Examination Centers:

There will be three centres of Examination, within the state of Uttarakhand, for appearing in CET.

- 1. Srinagar Garhwal
- 2. Dehradun
- 3. Roorkee

The University reserves all rights to change the examination centre in case of less number of applicants for any centre.



हेमवती नन्दन बहुगुणा गढ़वाल विश्वविद्यालय Hemvati Nandan Bahuguna Garhwal University श्रीनगर गढ़वाल (उत्तराखण्ड)–246174 Srinagar Garhwal (Uttarakhand) - 246174 (केन्द्रीय विश्वविद्यालय) (A Central University)

पत्रांकः हे.न.ब.ग.वि.वि. / PACE(9) 2022 / 60

दिनांक : 3 /09/ 2022

--:अल्पकालीन संविदा शिक्षक भर्ती अधिसूचनाः--

हेमवती नन्दन बहुगुणा गढ़वाल विश्वविद्यालय में सामाजिक अधिकारिता मंत्रालय भारत सरकार द्वारा वित्तपोषित **डॉo अम्बेडकर सेंटर ऑफ एक्सीलेंस (DACE**) के अन्तर्गत अनुसूचित जाति श्रेणी के छात्र / छात्राओं को संघ लोक सेवा आयोग (UPSC) सिविल सर्विसेज परीक्षाओं की कोचिंग प्रदान करने के लिये अल्पकालीन संविदा पर 03 शिक्षको (02 सामाजिक विज्ञान एवं 01 विज्ञान) की आवश्यकता है। संविदा शिक्षको को एक मुश्त रू0 1,15,000 / – वेतन प्रतिमाह की दर से भुगतान किया जायेगा। विस्तृत दिशा–निर्देशों व सूचना हेतु विश्वविद्यालय वेबसाइट www.hnbgu.ac.in देखें।

ऑनलाइन आवेदन फार्म भरने की प्रारम्भ तिथि – ऑनलाइन आवेदन फार्म भरने की अन्तिम तिथि – 05 सितम्बर 2022 (10.00AM) 25 सितम्बर 2022 (05.00 PM)

कलसचिव

प्रतिलिपि– निम्न को सूचनार्थ एवं आवश्यक कार्यवाही हेतु प्रेषित।

- कोर्डिनेटर, डॉ० अम्बेडकर सेंटर ऑफ एक्सीलेंस (DACE) को सूचनार्थ।
- 2. समस्त संकायाध्यक्ष / विभागाध्यक्ष / निदेशक / टिहरी, पौड़ी परिसर ।
- 3. समस्त प्राचार्य / निदेशक / सम्बद्ध महाविद्यालय / संस्थान ।
- 4. समस्त उप कुलसचिव/सहायक कुलसचिव।
- वित्त अधिकारी / परीक्षा नियंत्रक / कोर्डिनेटर, प्रवेश परीक्षा।
- जनसम्पर्क अधिकारी, को इस आशय से प्रेषित कि उक्त विज्ञप्ति समाचार पत्रों में निःशुल्क प्रसारित करवाने का कष्ट करें।
- सिस्टम मैनेजर, को इस आश्य से प्रेषित कि उक्त विज्ञप्ति को विश्वविद्यालय की वेबसाइट पर अपलोड़ करने का कष्ट करें।
- कुलसचिव, समस्त केन्द्रीय विश्वविद्यालय / राज्य विश्वविद्यालय को सूचनार्थ।
- 9. मा० प्रति कुलपति, महोदय को सादर सूचनार्थ।
- 10. निजी सचिव, कुलपति, माननीय कुलपति महोदया को सादर सूचनार्थ।
- 11. गार्ड फाईल।

कुलसचिव



Dr. Ambedkar Centre of Excellence (DACE)

Hemvati Nandan Bahuguna Garhwal University, Srinagar Garhwal

(A Central University)

Ref. No. HNBGU/DACE/2022/ 60

Date: 03/09 /2022

Contractual Teacher Recruitment Notification

Dr. Ambedkar Centre of Excellence of HNB Garhwal University, Srinagar (Garhwal) invites online applications from eligible candidates (eminent and professional scholars in the subjects of Social Sciences and Sciences) for **performance based** three (03) short-term positions of **Contractual Teachers** on consolidated fix salary to provide specialized coaching to the Scheduled Caste (SC) students for the Civil Services examination conducted by the UPSC as per following details:

S. N.	Positions	Essential Qualification/Eligibility	Desirable Qualification
1	Social Sciences Two (02) Position	 PhD/NET in the concerned/allied/relevant disciplines. PG- Political Science/History/Geography/ Economics/Sociology A minimum of 55% marks (or an equivalent grade in a point-scale, wherever the grading system is followed) at the Master's level UG- Social Sciences Applicant must possess a good academic record <u>Publications</u>- at least 3 research publications in peer reviewed/UGC CARE list journals 	 Experience of coaching for UPSC Civil Services Exams Qualified UPSC Preliminary exam in past Ability to teach History, Polity, Economics, Geography and other disciplines/topics for Preliminary and Mains exam of UPSC Civil Services. Working knowledge of computer and other ICT mediums.
2	Sciences One (01) Position	 PhD/NET in the concerned/allied/relevant disciplines. PG- Physics/Maths/Chemistry A minimum of 55% marks (or an equivalent grade in a point-scale, wherever the grading system is followed) at the Master's level UG- Sciences 	 Experience of coaching for UPSC Civil Services Exams Qualified UPSC Preliminary exam in past Ability to teach Logical reasoning, Aptitude reasoning and other Sciences related disciplines/topics for Preliminary and Mains exam of UPSC Civil Services. Working knowledge of computer and other ICT mediums.

Note:

- Contractual Teacher shall be paid a lump-sum monthly salary of Rs. 1,15,000/- per month only*.
 *(Subject to release of fund by DAF to DACE Centre as per Scheme).
- 2. In addition to teaching assignments with course module preparation, the contractual teacher shall perform all such other official duties which are essential for smooth functioning of the Centre.
- 3. The posts are purely contractual in nature and no claim for regularity at any stage will be entertained by the University.
- 4. Preference will be given to candidates who have prior experience of appearing in UPSC Mains examination or Interview.

Important Dates:

S.No.	Category	Fee
1.	UR/OBC/EWS Category	1000
2.	SC/ST Category and Women applicants	500

Fees once paid will not be refunded under any circumstances.

Payment should be made online only, through credit/debit card/Net Banking.

Important Instructions:

- 1. Applicants are required to apply online (Link: online.hnbgu.ac.in/dace_rec). The online link will be available live from 05/09/2022 (10:00 AM.) and will be closed on 25/09/2022 (5:00 PM).
- 2. Before applying for the post, applicants are advised to go through the Essential/ Desirable Qualifications and other general instructions carefully and satisfy themselves with their eligibility and candidature. Later on, no inquiry in this regard will be entertained.
- 3. Only completely filled forms with supporting documents will be considered for screening and incomplete applications in any respect shall be summarily rejected.
- 4. Applications must be submitted online only. The application form will be entertained through online mode only, however, applicants are also required to take print out of duly filled online application form and submit self-attested hard copy along with the requisite/uploaded certificates/documents through Registered post/ Speed post/ Courier/ by hand on the following University address, latest by 30/09/2022 (Friday) 5:00 pm. At the top of the envelope "Application for the post of(subject)" need to be mentioned clearly.
- 5. Address for Correspondence: Coordinator, Dr. Ambedkar Centre of Excellence, Department of Political Science, School of Humanities and Social Sciences, Hemvati Nandan Bahuguna Garhwal Central University, Birla Campus, Srinagar, District Pauri, Uttarakhand, India. 246174. The University shall not be responsible for any postal delays and documents received after last date of submission i.e. 30/09/2022 (Friday) on or before 5:00 pm.
- Any communication including call for interview regarding this recruitment process will be made through e-mail provided by the applicant in on-line form only, therefore, applicant must ensure providing correct valid e-mail ID. No separate communication shall be entertained by any other medium.
- 7. The university shall process applications entirely on the basis of information/documents submitted by the candidates. In case, any information/document is found false or incorrect at any stage, the responsibility and liability shall lie solely with the candidates. Therefore, the applicants are advised to fill out the application form carefully.
- 8. The prescribed essential/desirable qualifications are the minimum and the mere possession of the same does not entitle candidates to be called for the interview.
- 9. If the number of applications received in response to the advertisement will be large and it will not feasible to interview all the candidates, the university at its discretion may restrict the number of candidates to a reasonable limit on the basis of qualifications/experience higher than the minimum prescribed for the post. The university, however, will prefer, candidates possessing higher qualifications and experience.
- 10. The process of selection may include a presentation/teaching demo at the time of the interview.
- 11. Candidates are advised to visit the university website at regular intervals for any update.
- 12. Advocacy or canvassing in any form will result in disqualification.
- 13. The university reserves the right to revise/reschedule/cancel/suspend the recruitment process at any stage without assigning any reason. The decision of the university shall be final and no appeal in this regard shall be entertained.
- 14. In case of any inadvertent mistake in the process of selection that may be detected at any stage after issuing an appointment letter, the university reserves the right to modify/withdraw/cancel any communication made to the applicant in this regard.
- 15. If the services of selected candidate are not found satisfactory at any stage, his/her services may be terminated forthwith without assigning any reason.

Application Form

Personal Details			
Post Applied for		Subject applied for	Social SciencesSciences
Full Name		Marital status	
Category	-	Gender	
Date of Birth	11 ¹¹ 14	Nationality	
Father's Name		Whether you have	
Mother's Name		qualified the Civil Services Preliminary	
Aadhar No.		examination? mention your roll no. of	
Mobile No.		Pre/Main Examination (Attach proof)	
E-mail ID			
Address of Correspondence		Permanent Address	

Examination	Subject(s)	Division	Marks/Total Marks	Year	Board/University
10 th	i.				
12 th					
Bachelor's		-			
Master's					
Others					

Ph.D. de	gree details	N	
S.No.	Subject(s)	Thesis title	Date of Award

National/State Level Examination Qual	ified		
Examination (NET/SLET/SET)	Subject(s)	Year	
a			

Experiences (Research/Tead	ching)					
Organization	Designation	Status	Pay Scale/Consolidated	Nature of Duty	From	То	Total Time

Organization	Designation	Status	Pay Scale/Consolidated	Nature of Duty	From
	1.5	at .			

S.No.	Title of the Paper	Journal Name	Year	Volume	Page No	ISSN	SCI Impact Factor	Authorship Author/Co- Author	UGC listed	Scopus	Publishe

S.No.	Title of the Paper	Journal Name	Year	Volume	Page No	ISSN	SCI Impact Factor	Authorship Author/Co- Author	UGC listed	Scopus	Publishe
			4	· ·							

Publication: others (Book/Book Chapter/Conference Proceeding)								
S.No.	Publication Type	Title	ISSN/ISBN	Authorship Author/Co-Author	Publisher	Year	National/International	
		~						

Lecture Delivered							
S.No.	Title	Date	Seminar/Conference/Workshop	Organizer	State/National/International	Year	

CNIC	Name of the award	Name of the awarding	Date	Level
S.No.	Name of the award	body	Date	Level
				(National/International

Referees Details

S. No.	Name	Designation	Organization	Email	Mobile	

Discuss in about 150 words, your suitability for the post that you have applied for:

Candidate Declaration

I have read the applicable guidelines. I do hereby solemnly declare that the information given, the statements made and documents uploaded with this application form are correct and true to the best of my knowledge and belief. If any information given by me in this application is found to be false or misleading, my candidature is liable to be cancelled and I may be subjected to legal/disciplinary proceedings.

Candidate's Signature



हेमवती नन्दन बहुगुणा गढ़वाल विश्वविद्यालय Hemvati Nandan Bahuguna Garhwal University श्रीनगर गढ़वाल (उत्तराखण्ड)–246174 Srinagar Garhwal (Uttarakhand) - 246174 (केन्द्रीय विश्वविद्यालय) (A Central University)

पत्रांकः हे.न.ब.ग.वि.वि. / प्र०परी० / 2022 / 19 8

दिनांक : 29/06/2022

(कार्यालयादेश / 122)

विश्वविद्यालय द्वारा पूर्व में निर्गत अधिसूचना सं०– हे.न.ब.ग.वि.वि. / प्र0परी0 / 2022 / 121 दिनांक 30. 05.2022 के क्रम में एतदद्वारा सूचित किया जाता है, कि हेमवती नन्दन बहुगुणा गढ़वाल विश्वविद्यालय श्रीनगर गढ़वाल के द्वारा संचालित डॉ0 अम्बेडकर सेंटर ऑफ एक्सीलेंस (DACE) में प्रवेश परीक्षा के माध्यम से यूपीएससी (UPSC) परीक्षाओं के लिये निःशुल्क कोंचिंग कार्यक्रम में प्रवेश के लिए ऑनलाइन आवेदन फार्म भरने की अन्तिम तिथि दिनांक 10 जुलाई 2022 तक विस्तारित की जाती है।

कुलसचिव

प्रतिलिपि– निम्न को सूचनार्थ एवं आवश्यक कार्यवाही हेतु प्रेषित।

- कोर्डिनेटर, डॉ० अम्बेडकर सेंटर ऑफ एक्सीलेंस (DACE) को सूचनार्थ।
- समस्त संकायाध्यक्ष / विभागाध्यक्ष / निदेशक / टिहरी, पौड़ी परिसर।
- 3. समस्त प्राचार्य/निदेशक/सम्बद्ध महाविद्यालय/संस्थान।
- वित्त अधिकारी / परीक्षा नियंत्रक / कोर्डिनेटर, प्रवेश परीक्षा ।
- 5. समस्त उप कुलसचिव/सहायक कुलसचिव।
- जनसम्पर्क अधिकारी, को इस आशय से प्रेषित कि उक्त विज्ञप्ति समाचार पत्रों में निःशुल्क प्रसारित करवाने का कष्ट करें।
- सिस्टम मैनेजर, को इस आश्य से प्रेषित कि उक्त विज्ञप्ति को विश्वविद्यालय की वेबसाइट पर अपलोड़ करने का कष्ट करें।
- 8. निजी सचिव, कुलसचिव, कुलसचिव महोदय को सूचनार्थ।
- 9. मा० प्रति कुलपति, महोदय को सादर सूचनार्थ।
- 10. निजी सचिव, कुलपति, माननीय कुलपति महोदया को सादर सूचनार्थ।
- 11. गार्ड फाईल।

कुलसचिव



हेमवती नन्दन बहुगुणा गढ़वाल विश्वविद्यालय Hemvati Nandan Bahuguna Garhwal University श्रीनगर गढ़वाल (उत्तराखण्ड)–246174 Srinagar Garhwal (Uttarakhand) - 246174 (केन्द्रीय विश्वविद्यालय) (A Central University)

पत्रांकः हे.न.ब.ग.वि.वि. / प्र०परी० / २०२२ / 7 3

दिनांक : 3 / 10 / 2022

प्रेस विज्ञप्ति

एतद्द्वारा सूचित किया जाता है कि हे०न०ब० गढ़वाल विश्वविद्यालय श्रीनगर की डॉ० अम्बेडकर उत्कृष्टता केन्द्र (DACE) की प्रवेश परीक्षा, दिनांक ३० अक्टूबर २०२२ को अयोजित की गई थी। इस प्रवेश परीक्षा की उत्तर कुँजी (Answer Key) से सम्बन्धित छात्रों से प्राप्त आपत्तियों का निस्तारण करते हुये आज दिनांक 31 अक्टूबर २०२२ को परीक्षाफल घोषित किया जा चुका है। परीक्षाफल विश्वविद्यालय की वेबसाइट https://online.hnbgu.ac.in/Ambedkar_hnb/ पर उपलब्ध है।

(प्रो0 अनिल कुमार नौटियाल) कोर्डिनेटर, प्रवेश परीक्षा

प्रतिलिपि– निम्न को सूचनार्थ एवं आवश्यक कार्यवाही हेतु प्रेषित।

- कोर्डिनेटर, डॉ० अम्बेडकर सेंटर ऑफ एक्सीलेंस (DACE) को सूचनार्थ।
- समस्त संकायाध्यक्ष / विभागाध्यक्ष / निदेशक / टिहरी, पौड़ी परिसर।
- 3. समस्त प्राचार्य / निदेशक / सम्बद्ध महाविद्यालय / संस्थान ।
- 4. वित्त अधिकारी / परीक्षा नियंत्रक।
- 5. समस्त उप कुलसचिव/सहायक कुलसचिव।
- जनसम्पर्क अधिकारी, को इस आशय से प्रेषित कि उक्त विज्ञप्ति समाचार पत्रों में निःशुल्क प्रसारित करवाने का कष्ट करें।
- सिस्टम मैनेजर, को इस आश्य से प्रेषित कि उक्त विज्ञप्ति को विश्वविद्यालय की वेबसाइट पर अपलोड़ करने का कष्ट करें।
- निजी सचिव, कुलसचिव, कुलसचिव महोदय को सूचनार्थ।
- 9. मा० प्रति कुलपति, महोदय को सादर सूचनार्थ।
- 10. निजी सचिव, कुलपति, माननीय कुलपति महोदया को सादर सूचनार्थ।
- 11. गार्ड फाईल।

(प्रो0 अनिल कुमार नौटियाल) कोर्डिनेटर, प्रवेश परीक्षा

Dr. Ambedkar Centre of Excellence (DACE)

Hemvati Nandan Bahuguna Garhwal Central University

Srinagar Garhwal, Uttarakhand

Notice for the DACE Vacant Seats

All the interested aspirants are informed that the admission for the DACE program 2023-24 is underway and first merit list is notified on 30.11.2023 at the university website (www.hnbgu.ac.in).

After notification of first merit list, now the SAMRTH portal for admission to the remaining seats shall be opened from 01.12.2023 (Friday) to 03.12.2023 (Sunday). All the interested candidates may apply on the online admission portal.

Date of re-Opening of Admission Portal: 1st Dec. 2023

Date of Closing of Admission Portal: 3rd Dec. 2023

Online Admission Link: <u>https://hnbgudaceadmission.samarth.edu.in/</u>

Coordinator DACE

Dr. Ambedkar Centre of Excellence (DACE)

Hemvati Nandan Bahuguna Garhwal Central University Srinagar Garhwal

Merit List (Session 2023-24)

Based on merit and recommendation of selection committee, the following list of candidates were found suitable for admission in UPSC/SPSC coaching program for session 2023-24.

S.No.	Name of Candidate	Father's Name	Category
1	PRATIMA DAS	PRIYANATH DAS	SC
2	SAURABH KUMAR	UTTAM KUMAR	SC
3	NEETU VERMA	RAM JAGAT VERMA	SC
4	KM NEETU	RAJENDRA LAL ARYA	SC
5	SHAILJA SINGH	B L ARYA	SC
6	RAHUL KUMAR	SAJJAN LAL	SC
7	POOJA	DHEERAJ LAL	SC
8	SANDEEP	BISHU	SC
9	ANIL TAMTA	MAHESH CHANDRA TAMTA	SC
10	ANCHAL	VIRENDRA LAL	SC
11	VISHAL KUMAR	AWDHESH KUMAR	SC
12	AKANSHA	SUNIL KUMAR	SC
13	ABHISHEK KUMAR	SUNIL KUMAR	SC
14	RAVINDRA KISHOR	OM PRAKASH	SC

Select List of Scheduled Caste (SC) Candidates (Session 2023-24)

15	AKASH SINGHANIA	RESHAM PAL	SC
16	ARYAN VIKAS	PUSHKAR LAL	SC
17	NIKITA	BHOLA LAL	SC
18	KUMARI POONAM	RANVEER KUMAR	SC
19	DEEPAK SOURIYAL	NAND RAM	SC
20	POOJA SHAH	MANOHAR LAL SHAH	SC
21	SANJAY KUAMR	MOHAN LAL	SC
22	MUSKAN	VIKRAM	SC
23	KM USHA	PREM DAS	SC
24	RAJENDRA KUMAR	JASPAL LAL	SC
25	SAURABH	MUKESH KUMAR	SC
26	POOJA	VIJENDRA LAL	SC
27	HEMA	DINESH SINGH	SC
28	VISHAL KUMAR	PRADEEP KUMAR	SC
29	JITENDRA KUAMAR	JASPAL LAL	SC
30	SATENDRA SINGH	RAVINDRA SINGH	SC
31	RITIKA	VED PRAKASH	SC
32	SHAILESH RAJ	DINESH RAJ	SC
33	ROSHANI	VIRENDRA LAL	SC
34	RAMNEET	RAJVEER SINGH	SC
35	LALIT KUMAR	DAULAT RAM	SC
36	GANESH KUMAR	GORDHAN LAL	SC
37	DHARMENDRA KUMAR	UDAY LAL	SC
38	KM. USHA ARYA	BEERBAL LAL	SC

Select List of Other Backward Classes (OBC) Candidates

(Session 2023-24)

S.No.	Name of	Father's Name	Category	
	Candidate			
1	ADITYARAJ	ASHISH KUMAR	OBC	
	SHARMA	SHARMA		
2	KAREENA	JAGVEER SINGH	OBC	
	RAWAT	RAWAT		
3	RASHMI	RAKESH SINGH	OBC	
	CHAUHAN	CHAUHAN		
4	AMRENDRA	HARENDRA KUMAR	OBC	
	BEHERA	BEHERA		
5	MONA YADAV	HARIDAYNARAYAN	OBC	
		YADAV		
6	SIDDHARTHA	DINESH NAUTIYAL	OBC	
	NAUTIYAL			
7	OM PRABHA	VIJAY KUAMR	OBC	
		YADAV		
8	JITENDRA	PARAS YADAV	OBC	
	KUAMR			
9	DINESH	SHIV RAM	OBC	
	RAJPOOT			
10	TARUN	TULARAM	OBC	
	GANGWAR	GANGWAR		
11	ASHISH BYAHUT	TRILOKINATH	OBC	
		GUPTA		
12	SHIVANI YADAV	KRISHAN PAL	OBC	
		SINGH		
13	VIKASH	MOHAN LAL	OBC	
	CHAUDHARY			
14	GAURAV	PITAMBER DUTT	OBC	

	GOSWAMI	GOSWAMI	
15	JYOTI	MOHAN LAL	OBC
16	SHARAD YADAV	CHHOTE LAL YADAV	OBC
17	NIRBHAY	HARI NARAYAN	OBC
	KUMAR	PRAMANIK	
18	NEHA	PITAMBAR DUTT	OBC
		GOSWAMI	
19	ASHOK SINGH	BALBEER SINGH	OBC
20	DEEPAK	TEERATH SINGH	OBC
21	NISHANT GUPTA	YOGENDRA Y	OBC

Note:

1. The above candidates are required to report DACE office latest by 05/Dec/2023. The classes shall begin from 06/Dec/2023 onwards.

2. The formal admission to the DACE program shall be based on verification of all documents in original and fulfilling DACE program guidelines.

3. For further information candidates may contact on: 9891591788, 9805667711 or may contact through DACE email <u>dacehnbgu@gmail.com</u>

4. Information about the vacant seats is notified on university website.

Coordinator DACE

MoU with University of Haifa, Haifa, Israel

Potential of an intersectionality approach to climate change, medicinal/endemic plants, Himalayan/arid ecosystem and ethnography

Concept Note

Climate change is one of the most crucial environmental, social, and economic issues the world is facing today. Endemic and Medicinal plants (EMPs) of wild and agro origin provide valuable resources to indigenous communities throughout the Himalayan or/arid ecosystem for their food, traditional healthcare systems and cultural significance. Since ages, the indigenous/ traditional communities have been in close contact with the wild bio-resources and have developed sound knowledge about their surrounding environment which is deeply embedded in their sociocultural fabric. In the Himalayan/or arid ecosystems, different habitat types, varied microclimate conditions and a wide range of altitudes make this region an ideal place to grow and thrive EMPs. Endemic and Medicinal plants (EMPs) are also one of the critical components for livelihood and forms cultural identity for the people inhabited in this region. However, overexploitation and indiscriminate collection of EMPs species from their natural habitats have adversely affected their dominance and availability. Consequently, changing climatic conditions in the region threatened the status of many of the endemic and medicinal plants, continued provisional services of the ecosystem and traditional cultural integrity. The upward shift of plant species has already been reported from many parts of the Himalaya due to warming and loss of some species that formally were restricted to higher altitude ultimately, they may face the threat of extinction. Evidences indicated that many plants including EMPs have started blooming, fruiting and seed setting earlier the timing of their phenophases in response to the advancement of spring season due to changes in weather condition. Such kinds of events undermine the adaptive capacity and increased vulnerability to socio-ecological, traditional knowledge and cultural segments of the indigenous/traditional communities as they are poor and marginal and resides in remote and isolated area. Therefore, an intersectional approach is crucial to address the issues of threats to endemic and medicinal plants, degraded ecosystem and biodiversity and cultural integrity of the region through developing and advancing effective adaptation mechanism supported by evidence-based knowledge. The intersectional approach can open up new ways of understanding the socio-ecological, political, and cultural elements that make up a system vulnerable to change, and how the community perceives, experiences, and adapts to

change. The increasing level of concern about climate change and related forms of socialecological change affecting endemic and medicinal plant availability calls for an intersectional perspective for documenting and evaluating the impacts on indigenous community at individual, households and Institution's level and perceive, experience, and adapt to changing socioecological scenarios. Advancing such knowledge will help illuminate the various adaptation challenges faced by the indigenous/traditional community at individuals, households, and institutions within different adaptation measures i.e., location, social, economic, political and governance in which the indigenous society is trying to adapt. It is essential to translate scientific facts to the users' language, which is a major challenge. Scalability, diffusion, adoption and capacity building should all be considered in research and development efforts while following the intersectionality approach. "Sustainability and Culture in Himalayan Societies"

An International Workshop hosted by the University of Haifa, 13-15 March 2022

Goals

The aim of this present workshop is to promote and enhance academic collaboration and mutual projects between the University of Haifa and HNB Garhwal Srinagar (Uttarakhand) and the Indian Institute of Technology-Mandi (Himachal Pradesh), both of which are situated in the Indian Himalaya.

We focus on sustainability and culture in South Asian communities with an emphasis on Himalayan societies. Topics include resource management and reproduction, the use of new media in research, the politics of ethnic identities, and the uses of the Himalayan past.

1. Promoting sustainable environmental development and awareness among the local population of hill areas of Garhwal Himalayas, the reason being a large-scale migration from hill villages to plain areas, due to lack of natural resource conservation, unproductive agriculture, and natural disasters and calamities; 2. Conservation and promotion of tangible and intangible cultural heritage of disaster-affected hill areas; 4. Identification of gender disparity and vulnerability, promotion of equity; 5. Promoting ethnohistorical research and oral traditions in this area; 6. Preservation of the natural environment, traditional culinary practices and food consumption, and cultural heritage.

Participants:

University of Haifa, Israel: Prof Amos Megged, University of Haifa Prof Tali Katz-Gerro, University of Haifa Dr Arik Moran, University of Haifa

Hemvati Nandan Bahuguna Garhwal University (HNBGU), Srinagar Garhwal, Uttarakhand, India: Prof. Rajpal Singh Negi, HNB Garhwal Prof. RC Bhatt, HNB Garhwal Prof. R.K. Maikhuri, HNB Garhwal Dr. Prashant Kandari, HNB Garhwal (via Zoom

Indian Institute of Technology-Mandi, Himachal Pradesh, India: Dr Nilamber Chhettri, IIT-Mandi Dr Shyamasree Dasgupta, IIT-Mandi [1-2 more by zoom?] DAY 1. Sunday, 13 March, Senate Hall, University of Haifa:

Greetings

Resource management and reproduction (14:45-15:30):

14:45-15:30: Prof Talli Katz-Gerro, University of Haifa

The Cultural Politics of Household Sustainability

Research on the climate change and environmental sustainability has tended to pay significantly less attention to the cultural dimensions of adapting to climate change and environmental degradation. This means that little is known about how culturally-specific notions of sustainability, premised on reducing the impacts of Western overconsumption, are understood by immigrants to global North cities, by different religious groups, and by different lifestyle-communities. In this presentation I discuss the findings of two recent research projects. First, a mixed-methods research that explored the environmentally significant household practices of Somali immigrants living in Manchester, UK. Second, a project looking at household food practices of three social groups in Israel: religious Jews, secular Jews, and Muslim Arabs. I will discuss the way participants understand sustainability, how ideas around sustainability correspond to experiences of household resource use, how culture and religious norms shape household practices related to food behavior, and gendered and generational differences in participants' responses to policy messages about household sustainability. A main conclusion would be that cultural perspectives and practices can make important contributions to more inclusive sustainability governance.

15:30-16:15:

Prof. R.K. Maikhuri, HNB Garhwal

Socio-ecological approaches for bio-cultural and heritage resource conservation of Traditional/ethnic and indigenous communities of the Central Himalayan region–Uttarakhand

INTERMISSION (15 Minutes)

Environmentalism and Gender (16:30-18:00):

16:30-17:15: Dr Shyamasree Dasputa, IIT-Mandi TBC 17:15-18:00:

Dr. Prashant Kandari, HNB Garhwal (via Zoom): Symbiotic relationship between Mountain women and Natural resources: Sustainability, policy approach and emerging issues towards women empowerment.

DAY 2. Monday, 14 March, Zippori National Park/Senate Hall, University of Haifa.

Tour Zippori National Park (9:00-12:00):

Plck up from dormitories, tour of Zippori National Park

Lunch break (Zippori) [סנדוויצ'ים ארומה]

Session in the university [find topic, roundtable?] (14:30-16:00):

14:30-15:15

15:15-16:00

Add a speaker from IIT Mandi? TBC

DAY 3. Tuesday, 15 March, Jacobs Building, Room 305, University of Haifa

Ethnicity and Development in Himalayan Societies (9:00-12:00):

9:00-9:15: Gathering, Tea.

9:15-10:00

Dr Nilamber Chhettri, IIT-Mandi

Elusive Identities, Enduring Demands: Recognition struggle and scalar expression amongst the Hatti of Trans-Giri region in Himachal

10:00-10:45

Prof. Rajpal Singh Negi, HNB Garhwal Study of Multimedia Documentation of Folk Rituals and Processional Performances in Uttarakhand

10:45-11:30

Prof. RC Bhatt, HNB Garhwal "Territorial Jurisdiction and Interrelation among Gods and Goddess in Kinnaur, Himachal Pradesh, and Garhwal Central Himalaya, India"

LUNCH [on campus]

14:30-15:30: TOUR HECHT MUSEUM [תמי, להזמין סיור מודרך באנגלית]

[Retire to dorms]

20:30 – Workshop Dinner, Libira Restaurant (Downtown Haifa)

[pickup by Taxi from the dormitories at 20:00]

ABSTRACTS:

Dr Nilamber Chheettri, IIT-Mandi

Elusive Identities, Enduring Demands: Recognition struggle and scalar expression amongst the Hatti of Trans-Giri region in Himachal.

This paper is an empirical investigation into the claims made by the Hatti community of the Trans-Giri region in Sirmaur for recognition as a scheduled tribe. The paper traces the historical genealogy of this demand and discusses in detail Hatti's long quest to

secure the coveted ST status in the state. The paper examines the multifaceted domains of identity claims by elucidating their structure and content. The paper focuses on territorial demarcation that is redrawing of state boundaries and its impact on the constitution of ethnic boundaries in the region. Amidst the contested paradigm of recognition, this paper notes the scalar expression of such demands often embedded in the politics of place-making in the region. It will try to delineate their struggle and show how boundary drawing practices in South Asia have led to the contestation of identities in the Himalayan region.

Keywords: tribe, schedule tribe, ethnic groups, myths, memories, ritual, culture, history.

Prof. R.C. Bhatt, HNBU Garhwal

Territorial Jurisdiction and interrelation among Gods and Goddess in Kinnaur, Himachal Pradesh, And Garhwal Central Himalaya, India

Present study is a piece of research work conducted in Kinnaur region of Himachal Pradesh and Garhwal region of Uttarakhand to explore the role of divinity in defining the territory and also to understand the interrelationship between different gods and goddesses. It shows how a large geographical area was divided into many units which are marked and governed by divine characters. The major categories are principal deity and subordinate deity in which the largest deity is called Kunth who governs the entire Pargana and smallest deity is called Khimsu which is a household deity. The entire structure consists Paragana, Ghori, Gaon and Kim. Each subdivision is ruled by the deity with both civil and judicial rights. They ruled from their shrines with their own administrators and oracles as agents or mediums. The categorization of deities is not merely restricted in political hierarchy but it also categorised with the nature of their interest like- Adhikristh Devta, Isht devta, Kul Devta, Krishi Devta, pashu devta and more. It clearly indicates the separate area of interest of different deities which is full of specific rights and duties. In this inimitable idea of politico- religious territoriality some Gupt devta (hidden deity) also played their role which are not resided in any shrines. It has also come to light that most of the deities are movable in nature. Their procession called Jaat / yatra which moves in a prescribed way and a specified orbit within a defined geographical territory in the mountains as per the rituals after an interval of one and two years or sometimes over a long interval of twelve years. In this context it is also pertinent to mention that such practice of moving deities is also widely prevailing in Garhwal Central Himalayan region. And one of the well know Nanda RajJaat (Royal procession), which take place after an interval of 12 years is deep-rooted in the politicoreligious sphere of the Himalayan society as it was started in the early medieval period. Therefore, the present study also highlights the concept of the geo-political landscape of divine kingship of western and larger part of Central Himalayan region in a wider perspective which has been not delt before. These traditional practices will be highlighted in this presentation.

Key words: Divine kingship, geo-politics, territoriality, Village gods, Kinnaur

Prof. R.K. Maikhuri, HNB Garhwal

Socio-ecological approaches for bio-cultural and heritage resource conservation of Traditional/ethnic and indigenous communities of the Central Himalayan region–Uttarakhand

R.K.Maikhuri, Department of Environmental Sciences, HNBGU, Srinagar Garhwal, Uttarakhand, India (<u>rkmaikhuri89@gmail.com</u>).

The socio-ecological and natural characteristics of traditional/local communities are closely linked to this sensitive ecological setting. The extreme verticality of this region reverberates in major biophysical and socio-cultural peculiarities and globally significant for several reasons. With improvement in accessibility, bio-cultural and traditional systems are being increasingly impacted by external socio-economic political forces (conservative-developmental policies, economic globalization and democratization) coupled with the global environmental changes Conservation and management of natural resources under changed scenario is becoming a pressing challenge for sustainable development of the region. The traditional ecological knowledge and wisdom of the indigenous people have become a major focus of attention within the past decades. It is considered to have fundamental importance in management of local resources in the husbanding of the world' biodiversity and in providing locally valid models for sustainable living. It is now widely recognized that along with the conventional science and technology, the traditional knowledge products are of critical importance to overall development of the Himalayan region.

Globalization and homogenization have replaced local food cultures; high-yield crops and monoculture agriculture have taken the place of biodiversity; industrial and highinput farming methods have degraded ecosystems and harmed agro-ecosystem in diverse agro-ecological zones; and modern food industries have led to diet related chronic diseases and other forms of malnutrition. The stories show how traditional communities' food systems contain treasures of knowledge from long-evolved cultures and patterns of living in local ecosystems. However, these food systems which are intricately related to the complexities of social-cultural and economic circumstances are becoming increasingly more affected by the forces of globalization. With the passage of time, the knowledge and understanding on such a diversified food base has weakened considerably. This decline of knowledge base has wide range implication in view of resurgence of global interest on natural food and medicines. Considering this, it is high time to rejuvenate people's interest to harness the potential of these resources to counteract the impact of covid-19 and other diseases in future.

Further research to build on the present scenarios of traditional food systems could improve understanding of the forces driving negative environmental change that is decreasing availability of key food resources, and how to reverse these trends with local, regional and national policies. Understanding how to improve food choices, particularly among young generation youth, of both traditional and purchased foods with education and other incentives is greatly needed among the marginal and traditional communities of the Central Himalayan Region.

Socio-ecological approaches for bio-cultural and heritage resource conservation of Traditional/ethnic and indigenous communities of the Central Himalayan region–Uttarakhand



Prof. R.K. MAIKHURI

Dept. of Environmental Sciences HNB Garhwal University (A central University) Srinagar Garhwal ,Uttrakhand, India

Extent and biogeographic divisions of Indian Himalayan Region (IHR)







Bio-geographic representation of Indian Himalayan Region (IHR)

Bio-geographic Zones	Bio-geographic Provinces	Geographical area of India (%)	Major Biome Representation
Trans Himalaya	1A: Ladakh Mountains	3.3	Tundra
	1B: Tibetan Plateau	2.3	Alpine
	1C: Sikkim Trans Himalaya	<0.1	Alpine, Tundra
The Himalaya	2A: North West Himalaya	2.1	Alpine, Temperate, Sub-Tropical
	2B: West Himalaya	1.6	do
	2C: Central Himalaya	0.2	do
	2D: East Himalaya	2.5	do
North East India	9A: Brahmaputra Valley	2.0	Tropical Evergreen Forest, Very Moist Sal Forest, Tropical Grass Lands
	9B: Northeast Hills	3.2	Tropical Evergreen, Tropical Moist Deciduous, Sub-Tropical, Montane Temperate, Wetland



Diversity of wild relatives in the Himalayan sub-centers

Category	Distribution in Himalayan Sub-Centers				
	West Himalaya	East Himalaya	North-east Region		
Cereals and millets	29	07	16		
Legumes	09	05	06		
Fruits	37	32	51		
Vegetables	25	12	27		
Oilseeds	06	03	01		
Fibres	04	04	05		
Spices and condiments	10	09	13		
Miscellaneous	05	10	13		
Total spp. diversity	125	82	132		



Indian Himalayan Region(IHR)

- It covers a geographical area of 5.3 lakh km² which is 17% of the total geographical area of the country.
- The IHR inhabited by 49 million people which is about 4% of total population of the country, besides the region is home to more than 171 ethnic groups out of a total 573 schedule tribes reported in India.
- The IHR represent one-third of the total forest cover of India and nearly 45% of the very good forest cover in the country.
- The IHR is rich in biodiversity and represent diverse biomes and are about 1,740 species of medicinal and aromatic plants, 675 species of wild edibles and over 816 tree species reported.
- Temperatures across the IHR are expected to increase by 1–1.5°C (and in some higher altitudes, by up to 3.5°C) by the year 2050
- The IHR has the highest rate of outmigration, with mostly young men migrating for seeking employment and livelihoods





Managing system

Research & development (inter-& transdciplinary)
Education and information
Political decision/ governance
Implementation

Ecosystem

Driving forces (climate & environmental changes)

 Dynamics of natural process (risks and disaster)

 Natural resources (functions, diversity, values & services)

Impact of human activities

Natural system

Sustainable development

StrategiesIndicatorsmeasures

Local – regional system Driving forces (social, economic and political)

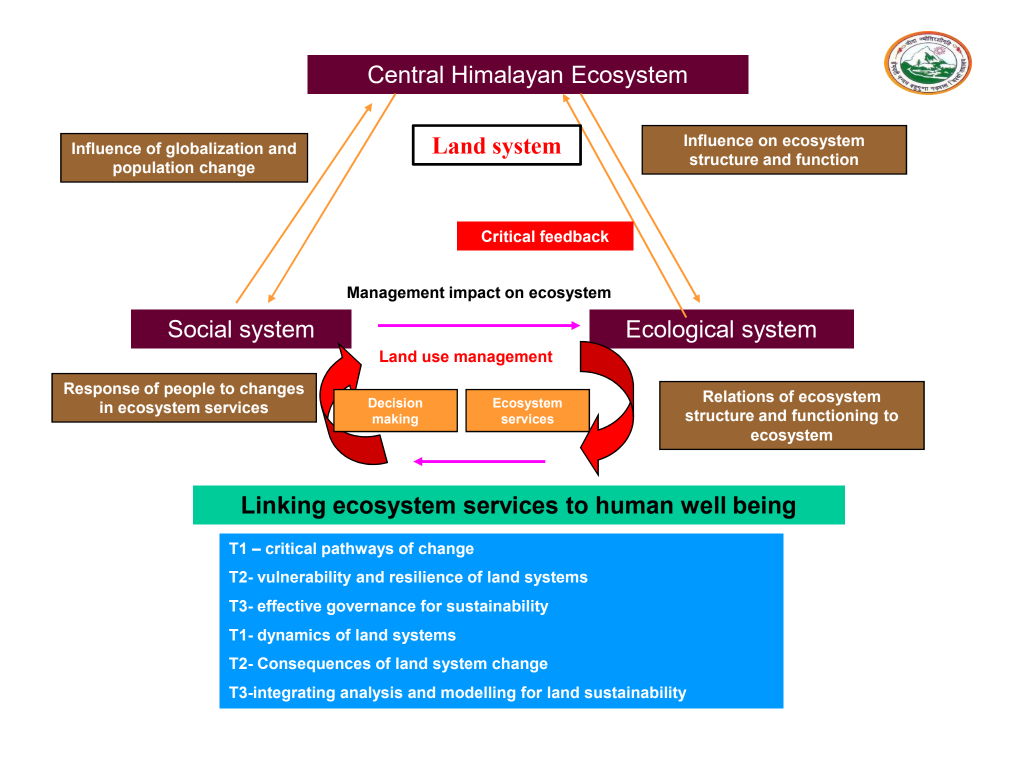
Cultural value systems

Landscape and land use change

Impact of natural process

Human system

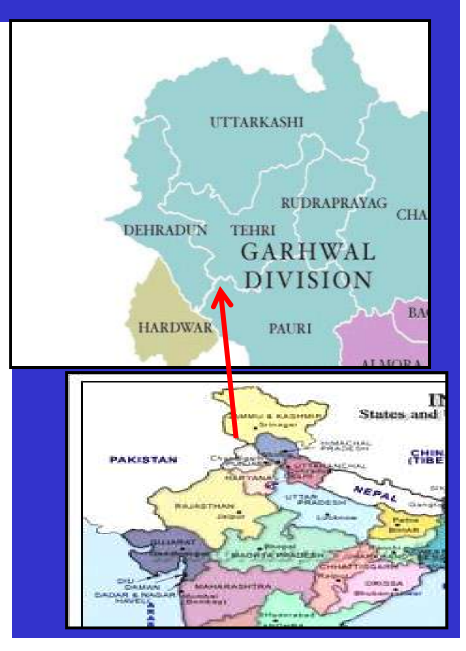
Element of an integrated approach to help understanding of the driving forces in natural and human systems from local to global level in a time of rapidly growing process of globalization and global change



The second secon

Central Himalayan Region (Uttarakhand)

- The Central Himalayan Region (CHR) is spread over 53,483 sq. km. and are susceptible to the impacts of natural perturbations such as natural disaster, climate change as other parts of IHR, HKH and elsewhere
- *The CHR is home of 10.1 million people and the communities living in the rural landscape are economically weak and marginalized.
- The region occupy 35,394 sq. km. forest area which constitutes 64.8% of its total geographical area.





Issues and challenges for achieving the Sustainable livelihood in the CHR, Uttarakhand

- Deforestation, land degradation, forest fire and declining carrying capacity of forest and rangelands.
- Biodiversity loss (forest and agro-biodiversity)/biological invasion.
- Hydrological imbalance (drying springs/ water resources, etc).
- Predominance of rain-fed agriculture, small and fragmented land holding-low agricultural yield.
- Human wildlife conflicts (crop raiding/livestock depredation by wildlife).
- Lack of small scale industries and poor micro-macro-economic condition of the region.
- Low skill/capacity among local people for local value addition in agro-based products for entrepreneurship development
- No employment opportunities, inadequate livelihood options that leads to out migration
- Low level of technological adoption and poor infrastructural facilities
- Natural disaster, extreme events, landslides, cloudburst/flash floods ,etc.
- Expansion of protected area network (PAs).
- Low access to technical education
- Social disintegration





Linking cultural diversity with biodiversity and livelihoods

- With whole range of traditional and tribal communities, the human dimension of bio-resource utilization and management is enormous.
- Bio-resources of agro and wild origin constitute an important source of livelihood for million of people across the IHR.
- Out of 17,000 recorded plant species, over 9,500 wild plants are recognised to be ethno-biological values (THCS), of these about 190 have marketability.
- Traditional agro-biodiversity Diverse culture have unique indigenous practices in the field of agriculture.





CHR – Rich in Bio-cultural/Ethnic Diversity

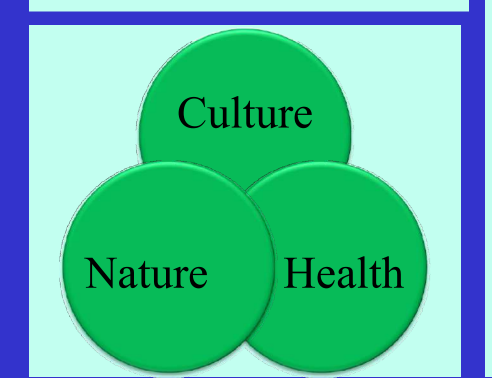
Ethnic/ Tribal and traditional Communities	Livelihood options		
Bhotiya	Agriculture		
(Marcha,Tolcha	• Settled agriculture		
, Jads)	(rain-fed and valley		
Van-Raji	land)	6 - C	
Boxa, Van	• Horticulture	it says	and a second
Gujjars,Jaunsa	Nomadic pastoralists		1.5.
ri, Barpatiyas,	Transhumant	A	
Garhwalis ,	Livestock	A STANDARD AND A STANDARD	
Kumaonis	• Cattle, sheep/goat, yak,		
	poultry, rabbit, etc.		A
	Handicrafts		
	Wild collection from		C'
	forests		
		VI WINE S	
			T. T

Culture, nature and health



•To understand the traditional knowledge in reality, the perspective culture-Nature-Health can be utilized

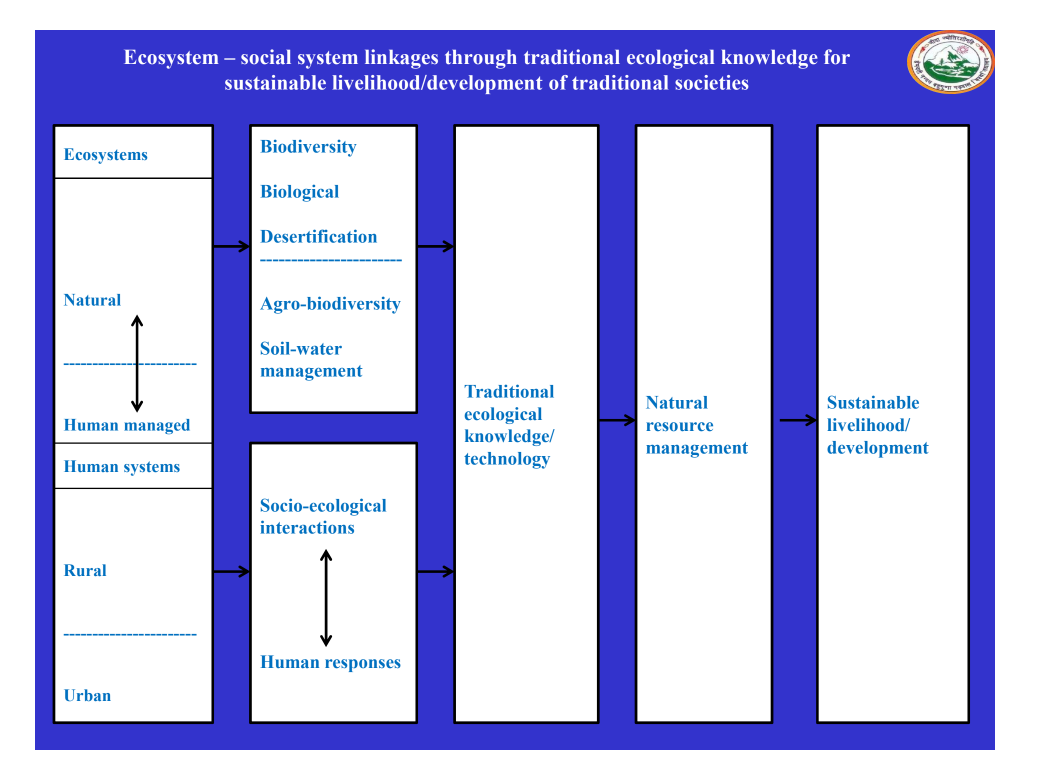
•However, to understand the interrelationship between the three concepts, one cannot fragment the analysis into independent categories.



There are four interstices among the three circles.

- **1. Relationship health-nature:** There is a close relation-ship between health and nature, eg., traditional health care systems by traditional communities.
- **2. Relationship nature-culture:** Ethnobiology interrelationship between nature and human culture.
- **3. Relationship culture-health:** Traditional medicine is understood as the medicine system used by indigenous or local communities to manage health and sickness.

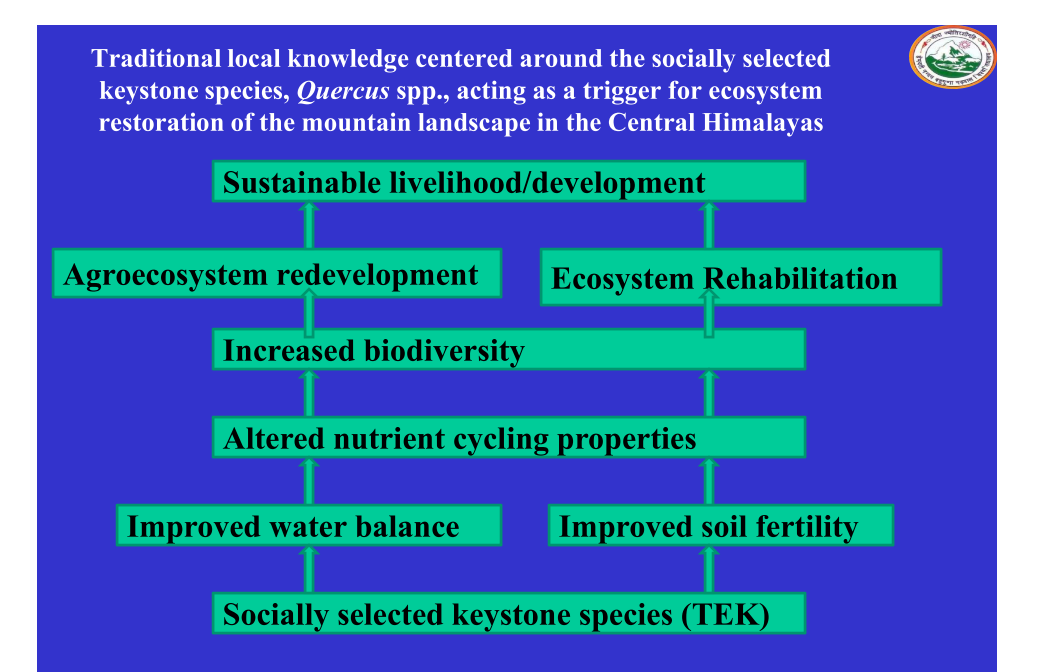
4. Relationship health-nature-culture: Western scientific thinking and its several disciplines still lack a science able to approach the integrality of the three relationships, but authentic shamanic systems can teach us about this integration of concepts.



Indigenous knowledge related to soil, space, water, soil fertility, crop and vegetation management systems developed by the farming communities of the Central Himalaya

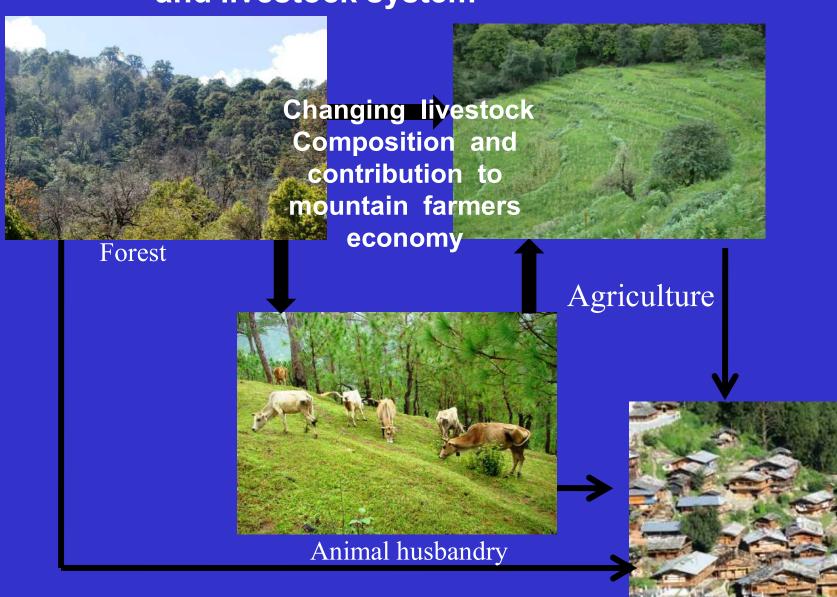


Environmental and others constraint	Use for	Indigenous practice
Limited space	Maximize use of environmental resources and land	Intercropping, agroforestry, multi-story cropping, altitudinal crop zonation and crop rotation
Steep slope	Control erosion and conserve water	Terracing, leveling, continuous crop and/or fallow cover, stone walls
Soil fertility maintenance	Replenish soil fertility and recycle organic matter	Farm yard manure, crop rotations, leaf litter gathering, composting, green manuring, mulching, in-situ manuring, ash and kitchen waste, mixed cropping with legumes, recycling weeds, burning biomass, fallowing etc.
Flooding or excess water	Integrate agriculture with water supply	Raised field agriculture, ditched fields, etc.
Excess water	Channel/direct available water	Control floodwater with canals, canal irrigation fed from streams, lakes and reservoirs.
Unreliable rainfall	Best use of the available moisture	Use of drought-tolerant traditional crop species and varieties, mulching, crops with short growing periods.
Temperature or radiation extremes	Ameliorate microclimate	Shade reduction or enhancement, plant spacing, thinning, shade-tolerant crops, increased plant densities, mulching, weeding, intercropping, agroforestry.
Pest incidence	Protect crops, minimize pest populations	Thick planting, crop watching, hedging or fencing, mixed cropping
Unavailability of the seed store/ cooperatives	Selection of healthy seed	The care of seeds always in the hands of women and based on field observations and IKS available with women.



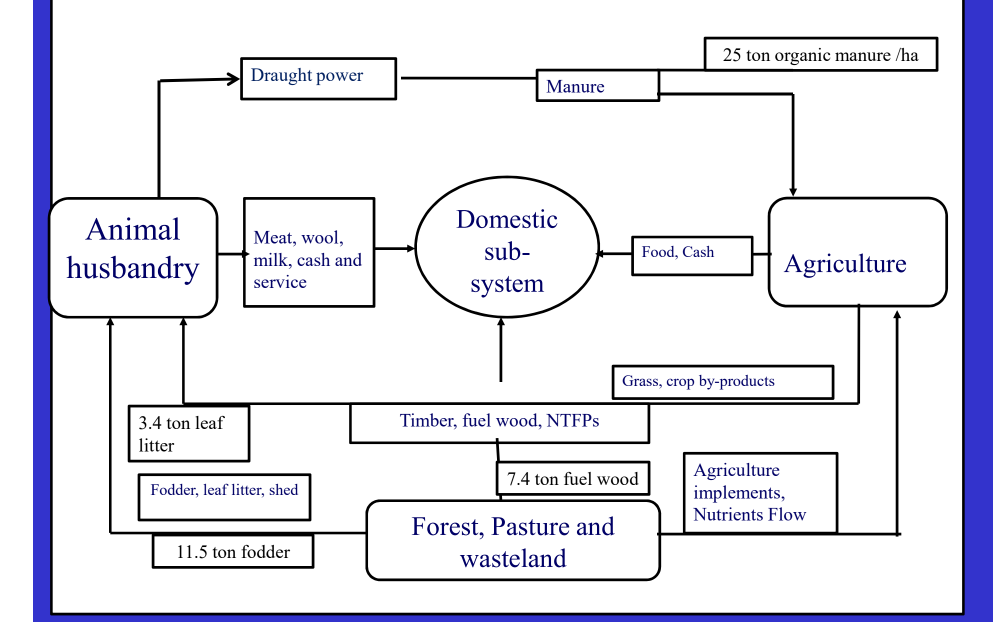
Inter-linkages between forest, agriculture and livestock system-





Domestic system

Inter-linkages between the various sub-systems in a village ecosystem



Central Himalayan Region: A storehouse of forest Biodiversity & Agro-biodiversity (300-4500 masl)



No. of vegetation types		8
No of plant categories	Trees	4248
	Ferns	241
	wild edibles	359
	Medicinal plants	850
	Crop wild relatives	132
	Crop plants	49
	Grains/Cereals-5	
	Millets- 5	
	Pseudo cereals- 6	
	Pulses -15	
	Oil seeds- 4	
	Tuber /bulbs-4	
	Others -10	

Traditional agro-diversity: key to sustainable agriculture





- Diversity and nutrient cycling
- Diversity and insect-pest management
- Diversity and plant disease and nematodes
- Diversity and weed control
- Diversity and nutrition
- Diversity and soil conservation
- Diversity and productivity, long term stability of the agroecosystem, food security, sustainability



Linkages between agro-biodiversity, provision of ecosystem services, and food security and resilience in the Central Himalayan Region

Agro-biodiversity	Ecosystem services	Contribution to food and nutrition security	Supplementary benefits
 Diversification of crops Integration of MPTs and fruit trees Revival of traditional crops Integration of livestock Introduction of honeybees of diverse species Conservation and management of palatable plant species in agricultural land 	 Provisioning services Diverse food items from various crops, animals livestock feed Medicinal plants and NTFPs Seeds/Genetic material Regulating services Pollination , Pest control Soil fertility maintenance Supporting services Soil protection Organic manure Cultural services Agro-tourism Cultural use of traditional crops as food and medicine 	 Improved dietary diversity and intake of micro- nutrients; Diverse crops and food security Alternative sources of income (purchasing power) Reduction of risks Enhanced agro- ecosystem resilience Improved production stability 	 Improved resilience of agricultural systems Higher diversity and flexibility in local production systems
		Increased income	

Major issues and changing scenario of Himalayan agro-ecosystems

- Himalayan agriculture transformation process.
- Subsistence farming, deterioration of the economy, environment, climate change and bio-resources.
- Illusions about quality of coarse and fine grains;
- Commercialization of agriculture: growth and economic prosperity.
- Unmindful introduction of HYVs and extinction scenario.
- Neglected mountain perspectives in agricultural policy and weak cross-sectoral linkages
- Research bias
- Unawareness about dimension of the threat.
- Lack of institutional arrangements, mechanisms and human capacity-building which would promote agro-biodiversity conservation
- Inadequate scientific and technical understanding and indigenous knowledge Himalayan agriculture
- Less attention on integrated components of hill agriculture



Area in ha/village under different traditional crops in *Kharif* and *Rabi* seasons during 1970-74 and 1990-94 in Central Himalaya (after Maikhuri *et al.* 1997,2001)

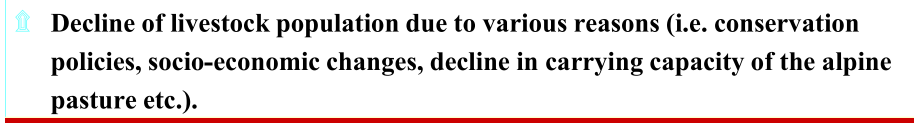
CROPS/ CROPPING SEASON	AREA IN ha/ VILLAGE		AREA IN ha REPLACED	AREA DECLINED IN %
KHARIF SEASON				
Panicum miliaceum	14.2	4.9	By high yielding rice varieties	65.5
Oryza sativa	14.2	14.2	Traditional rice varieties by HYV	-
Avenal sativa	15.8	3.4	By potato	78.5
Fagopyrum tataricum	8.6	1.5	By potata + rajama	82.5
Fagopyrum esculentum	4.1	0.3	By rajma	92.7
Perilla frutescense	1.3	-	By soyabean	100.0
Setaria italica	2.3	0.8	do	65.2
Oryza sativa	11.2	11.2	Traditional rice varieties by HYV	-
Eleusine coracana	9.6	6.1	By soyabean + amaranth	36.5

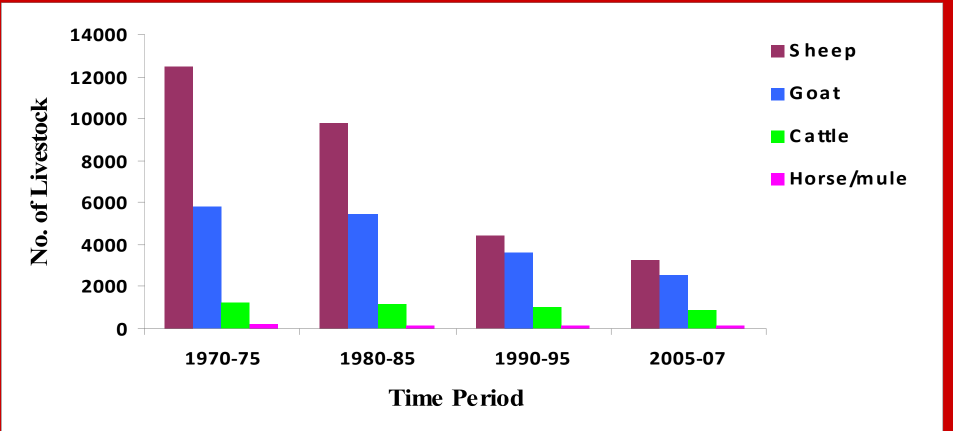
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Macrotyloma uniflorum	2.1	0.5	do	100.0
Echinochloa frumentacea	2.5	0.7	By pigeon pea	72.0
Vigna spp.	3.3	-	By pigeon pea + amaranth	100.0
RABI SEASON				
Triticum aestivam	14.2	14.2	Traditional wheat varieties by HYV	-
Hordeum himalayens	17.1	4.7	By potato, amaranth + rajama	-
Hordeum vulgare	7.0	1.1	By improved mustard varieties	-
Brassica compestris	2.0	2.0	No change	No change

B. Pastoralism & Transhumance Pastoralism





Changes in livestock population between the 1970-75 to 2005-07 period as reported by the people of Niti valley (10 villages).

Ecological, socio-economic drivers and policy issues responsible for agro-biodiversity loss in Central Himalaya, Uttarakhand



Uttarakhand			
Ecological drivers/indicators	Socio-economic drivers/indicators	Policy drivers/indicators	
• Decline in carrying capacity of forests and rangelands	• Small holding and land fragmentation	 Neglect of hill agriculture in policy and planning Research bias 	
• Increased abandoned land	• Out- and in-migration	Land use policiesSubsidies on food import and	
• Increased weed infestation/ invasive species	Change in food habits	credit policies • Strict Forest policies and	
• Climate change/variability	 Change in social values Increased female literacy 	wildlife conservation Act Subsidies on agricultural implements 	
• Soil erosion/run off	·	Fixed prices	
• Hydrological imbalances	• Dependence on wild collection of high value resources	 Govt. support services e.g., MENREGA and cheaper food 	
 Low crop yield /productivity 	• Decline traditional knowledge	to BPL families-PDS • Lack of expertise in agro-	
• Decline wild bio-resources affecting wildlife food chain systems	• Change in cropping pattern due to economic consideration	 technology transfer Lack of human resources in agri-business 	

Challenges to traditional food dietary diversity and nutrition security Challenge **Dimension of food and** Consequences nutrition security likely to be affected negatively Deterioration •Reduced food production & diversity •Food availability utilization of local food system Changing diets • Reduced dietary diversity •Food utilization Lingering •Reduced food intake & Dietary Diversity •Food accessibility & utilization poverty Abandonment •Low returns, land abandonment and loss •Food availability of production of cultivable land •Encroachment of agriculture land leading Rapid •Food availability to reduced agricultural production urbanization Depletion of •Loss to water resources & biomass manure •Food availability, availability, utilization, natural form forests •Reduced supply of wild edible, and accessibility resources reduced livestock production and income

Innovative (simple, cost effective and affordable) Technology Interventions used for livelihood enhancement and diversification in the CHR



Yield increasing

- Protected cultivation (Polyhouse, polypit, Poly trench)
- Bio-compost
- Vermicompost
- Vermiwash
- Cow pat pit
- Nadep compost

Income generating

- Vegetable cultivation
- Cash crop cultivation
- Integrated Fish –farming
- Horticulture
- Apiculture
- Floriculture
- Nursery development
- Mushroom –cultivation
- Medicinal Plant cultivation

Technology packages

Life Supporting Activities

- Water harvesting
- Management and improvement of waste land
- Fodder grasses
- Traditional art
- Sewing & knitting
- Multipurpose tree plantation

Other activities

- Bio briquetting
- Zero energy cool chamber
- Decorative items
- Bamboo propagation
- Drip irrigation
- Sprinkler
- PRA method
- Formulation of SHGs
- Bio- fencing

Types of traditional knowledge for adaptation in agroecosystem



Traditional knowledge about	How it helps adaptation in agro-ecosystem?
Resilient properties	Traditional farmers often live in marginal land where climate change impacts and selection pressures are greatest. This enables them to identify resilient crop species and varieties for adaptation
Farming practices	Traditional farming practices – conserve key resources for resilience and adaptation – such agro-biodiversity, water, soil and nutrients
Wild crops relatives	Local communities often draw on wild areas around farms for crop improvement and domestication, also provide food when crop fail
Plant breeding/seed selection	Traditional farmers particularly women conserving local landraces and selecting seeds for preferred and adaptive traits over generations
Climate forecasting	Traditional knowledge can help forecast local weather and provide accessible information to farmers at a local scale.

Climate change felt and response/adaptation using traditional knowledge and innovations in low elevation (300 – 3000 masl) in central Himalaya



- Low elevation (300 1000 masl)
- Nine hilly districts of Uttarakhand
- 54 villages, and 1620 HH
- Methods: People perception, workshop/PRA etc.
- Qualitative/quantitative studies of 6 years
- Higher temperature
- Increased exotic weeds
- Shift in season
- Drought
- New pests
- Unpredictable rain
- Low rainfall (peak season)

Adapting with T K & Innovations

Changes in climate

- Adopting traditional crops
- Tuber and rhizomatous crops under traditional agroforestry
- Monocropping of pulseszz

- Integration of *turmeric colocasia* and ginger under traditional agroforestry system
- Climate resilient traditional crops and pulses
- Irrigated land converted to rainfed due to water shortage
- Rainfed paddy replaced with millets (finger millet and barnyard millet)
- Delayed sowing of rainfed paddy (instead of 1st 3rd week of March)
- Horticulture (mango) in rainfed land
- Increased corn cultivation in few villages
- ***** Biocontrol (ash, cow urine and organic manure)

Evolution of the nutrition transition in the central Himalaya



Nutrition transition in full bloom: HYY displaced the indigenous varieties in the food systems and thus radically altered the dietary patterns and food habits of the population. This signaled the onset of the nutrition transition in the Himalayan region.

Is this trend reversible?

- The revitalization of traditional food systems
- Addressing important constraints to the production of traditional foods, TIME (Technology intervention in the mountain ecosystem), simple & affordable technology.



The transition from subsistence to commercial

production (Use attributes of millets, pseducereals and pulses)

Common and scientific name	Plant characteristics	Traditional use	Nutraceutial and commercial interest
Finger millet (Mandua) Elusine coracana	Herbaceous crop (seed- propagated)	Flour is used for making chapati & badi in Uttatakhand	High protein, Iron content, Gluten free
Amaranth (Cholai) <i>Amaranthus caudatus</i>	Herbaceous crop (seed- propagated)	The grain is roasted eaten, and flour is used as chapati	Rich source of Vitamins & Minerals
Buckwheat (Fafar) Fagopyrum tataricum	Herbaceous crop (seed- propagated)	The flour is used for chapati.	Good source of rutine used as medicine & highly nutritious.
Horse gram (Gahat) <i>Macrotyloma uniflorum</i>	Herbaceous crop (seed- propagated, legume crop)	Highly nutritious pulse consumed in winter and also cooked for a variety of dishes	The grain soup is used as a cure for kidney stones & rich in minerals.

Local value addition in agro-based products for entrepreneurship development





Maikhuri et al., 2001, The Environmentalist. Saxena et al., 2005, Journal of Mountain Science.

Ar	Area-specific approaches based on socio-agro-ecological potential and access to markets, information, and institutional services		
Agro- ecological	Access to markets, information, and institutional service	28	
potential and suitability	Good	Poor	
High	 Areas with high potential and good access to markets and services Promote large scale food production (where possible mixed cropping), horticulture, poultry farming, etc,. Enhance support for high value cash crops, MAPs and vegetables Integrate traditional food crops in cropping system in all agrozones Promote climate resilient crop varieties and adaptation measures Establish agro- processing and storage facilities Climate resilient technology and water management Encourage woman as entrepreneurs/managers in agriculture business 	 Areas with high potential but poor access to markets and services Improve marketing, storage and transport facilities, information systems, and extension services for fruit/ vegetable/livestock products Strengthen local food systems with a focus on traditional crops and support value chain development Promote high-value non-perishable agricultural products such as pulses, medicinal plants, and honey Promote livestock and livestock products and by products Improve credit, extension, and insurance facilities for crops and livestock 	
Low	 Areas with low potential but good access to markets and services Promote local products such as crafts (e.g. woodcarving, shawls, carpets) and services for markets Promote agro- technologies that enhance agricultural potential and utilize local niches Encourage agroforestry, NTFPs, and medicinal plants Develop local off-farm employment opportunities Encourage local breeds of livestock such as yak, goats, and sheep (mainly pastoralism) in high mountain ranges 	 Areas with low potential and poor access to markets and services Provide incentives for conservation and sustainable use of bio-resources for livelihood and income generation Encourage non-farm activities, e.g., tourist guides, homestay, local handicrafts Promote subsistence agriculture with zero-tillage, mixed cropping, and livestock production Promote rural, ecotourism and recreation Develop and harness environmental services 	

Wild bio-resources



➢ Medicinal and aromatic plants and traditional health care systems(THCS).

➢Wild edibles/Non timber forest product (NTFPs); Fruits, green leafy vegetables, etc.,

Traditional uses of medicinal plants for curing ailments in Alaknanda catchment of Uttarakhand (Phoondani and Maikhuri,2010).



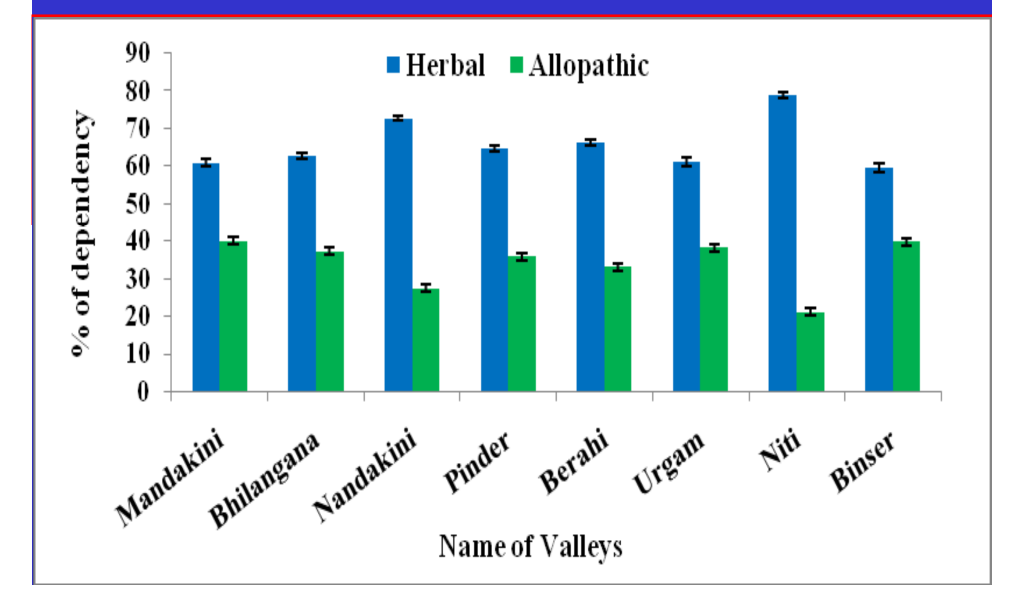
Name of plant species	Vernacular name	Ailments	Traditional uses
Picrorhiza kurrooa	Kutaki	Fever	50 gm.of dried roots milled along with 2 spoons sugar and taken with water.
Angelica glauca	Choru	Cold and cough	1 gm. root powder mixed with 1 cup tea as a cure for common cold.
Saussurea costus	Kuth	Tooth ache	100 ml decoction of the Saussurea costus tuber mixed with 4-5 drops of Prunus armenica oil and ½ spoon of salt.
Podophyllum hexandrum	Bankakri	Cancer	Root paste is applied
Arnebia benthami	Balchari	Baldness	5gm root milled and mix with 50ml mustard oil for applying on hair.
Aconitum hetrophyllum	Atis	Stomach ache	Root paste applied on the fore head to cure headache.
Dactylorhiza hatagirea	Hatajari	Wounds and cuts	Root paste is applied for treating cuts.
Swertia chirayita	Chirayata	Fever	Fresh leaves and stem milled juice and taken with water.

Collection period of some high value medicinal plants for curing various ailments in different communities/valleys of Alaknanda Catchment in Uttarakhand

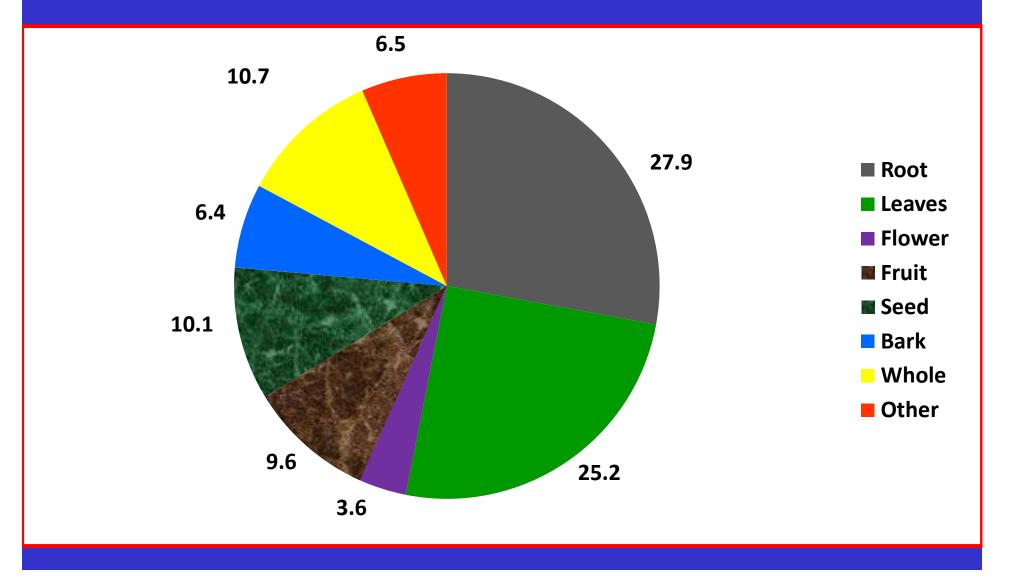


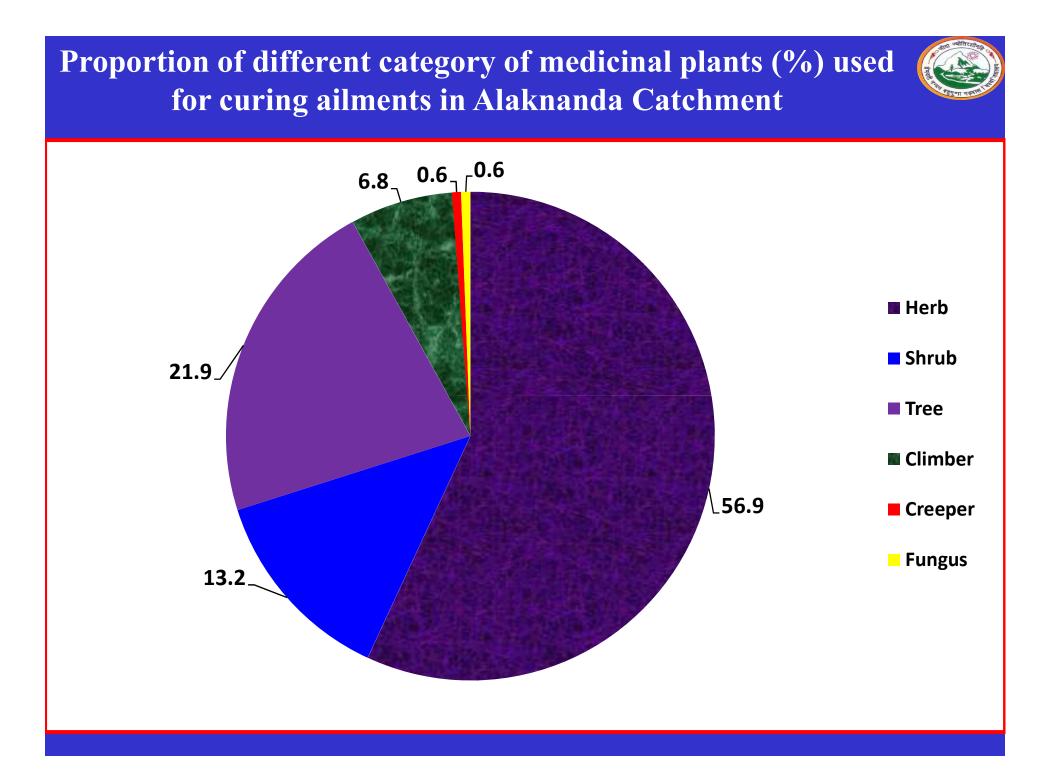
Name of plants	Part used	Medicinal uses	Collection period
Saussurea costus	Root	Toothache	Autumn season (September- October)
Delphinium denudatum	Root	Snakebite	Autumn season (September- October)
Saussurea ovallata	Flower	Leucorrhea, Mental disorder	Rainy season (August) When the flower is mature
Polygonatum verticillatum	Root	Anemia, Leucorrhea	Autumn season (September- October)
Podophyllum hexandrum	Root	Cancer	Autumn season (September- October)
Arnebia benthami	Root	Hair disease	When the above ground part is dry
Aconitum heterophyllum	Root	Fever, Stomachache	Autumn season (September- October)
Picrorhiza kurrooa	Root/Leaf	Typhoid fever, Jaundice	Autumn season (September- October)
Dactylorhiza hatagirea	Root	Cuts, Wounds	Autumn season (September- October)
Taxus baccata	Bark	Anti- cancer, Bone fracture	Winter season (November- December)

Total dependency of herbal and allopathic system of treatments for curing ailments in Alaknanda catchment of Uttarakhand



Medicinal plants and their parts utilized by the tribal and nontribal communities of Alaknanda catchment on traditional health care system.





Central Himalayan forests and alpine meadows safety nets mechanism as household livelihood coping strategies



Coping with adversities	Function	Description
Safety net	Insurance	Food and cash income in periods of unexpected food and income shortfalls
Support current consumption	Gap-filling	Regular and irregular food and income shortfall such as crop failures and seasonal shortages
Occupational activities	Source of employment and livelihood diversification	Unemployed youth/low income group people, off-cropping seasonal activities

Three critical functional categories – are major assets in responding food and nutritional security. As a source of livelihood security, these functions need to be included in planning for poverty reduction, nutritional security and climate adoption strategies, as well as forest and alpine management.



The main reasons for the lack of attention given to underutilize or wild gathered Species include

- A lack of information and reliable methods for measuring their contribution to farm households and the rural economy;
- The lack of guaranteed markets, except for a small number of products;
- The irregularity of supply of wild plant products;
- The lack of quality standards;
- Lack of standardization of the product;
- The lack of storage and processing technology for many of the products;
- The availability of substitutes;

Contribution of agro and wild bio resources in developing immune defenses to counter the impact of Covid-19 in central Himalayan rural landscapes



- Traditional mountain crops
- Medicinal plants of wild origin
- Wild fruits/edibles
- Spices/condiments of wild origin
- Wild greens used as a food/vegetables
- Wild and cultivated plants used as a Hot beverages (teas)

Traditional mountain crops used as major food by the local people during COVID-19 (as prescribed by local Vaidya's/herbalists) to increase immune defenses in Central Himalaya, Uttarakhand



Botanical	Local name	Uses and medicinal properties
name		
Amaranthus	Amaranth (Kedar-	The grain is considered very healthy and keeps body warm during
causdatus	chua)	winter season. Rich source of vitamins and minerals.
Fagopyrum	Buck wheat (Oggal)	The grains are rich source of vitamins and strengthen immune
esculentum		systems. High Medicinal use.
Fagopyrum	Buck wheat (Fafar)	It is a good source of rutine, healthy and improves immune defenses.
tataricum		
Macrotyloma	Horse gram (Gahat)	A highly nutritious pulse , the grain soup is used as a cure for kidney
uniflorum		stones.
Elusine	Finger Millet	The grain considered rich source of vitamins and healthy food.
coracana	(Manuduwa)	
Setaria italica	Foxtail Millet (Koni)	Substitute of rice, be good for patients suffering from typhoid fever,
Echinochloa	Domisional millat	pneumonia etc. The evolved grains when mixed with ourd given for patients suffering
	•	The cooked grains when mixed with curd given for patients suffering
frumentacea	(Jhangora)	from jaundice and also considered good for diabetic patient.
Parilla	Perilla (Bhangjeera)	Oil is edible and also used as a medicine and improve immune
frutescense		system,.
Vigna radiata	Green gram (Mung)	Soup of the grains are very nutritious, strengthens immune defenses.

Medicinal plants consumed /used by the local people to increase immune defenses during COVID-19 (as prescribed by local Vaidya's/herbalists) in Central Himalaya, Uttarakhand



Botanical Name of Medicinal Plants	Local Name	Uses
Origanum vulgare	Wild leaf Basil (Van tulsi)	Whole plant is used as an immunity booster by the local communities. Highly medicinal
Ocimum sanactum	Basil (Tulsi)	Whole plant parts are used to increase immunity and also used to cure colds and cough
Allium sativum	Garlic	Fresh bulb chewed which provide immunity to the body, and prevent pulmonary problems
Mentha arvensis	Mint (Podina)	Fresh leaves provide resistance against fever and very refreshing
Tinospora sinensis	Giloy	Whole plant used to cure fever, reduce body temperature and improve immune systems
Vibernum mullaha	Indian craneberry (Bhatmoliya)	Fruit juice used to provide strength, nutritious, rich in vitamin C and is very refreshing
Zingiber officinalis	Ginger (Adrak)	Rhizome used to cure cough and good immunity booster
Hippophae salicifolia	Sea-Buckthorn (Amesh)	Fruit juice is rich in vitamin C, provide strength to body and also used to treat colds and coughs
Phyllanthus emblica	Indian gooseberry (Amla)	Fruit is rich source of vitamin C and also used as blood purifier
Rhododendron arboreum	Rhodadendron (Burans)	Juice used to cure cardiac and respiratory disorder and very refreshing.



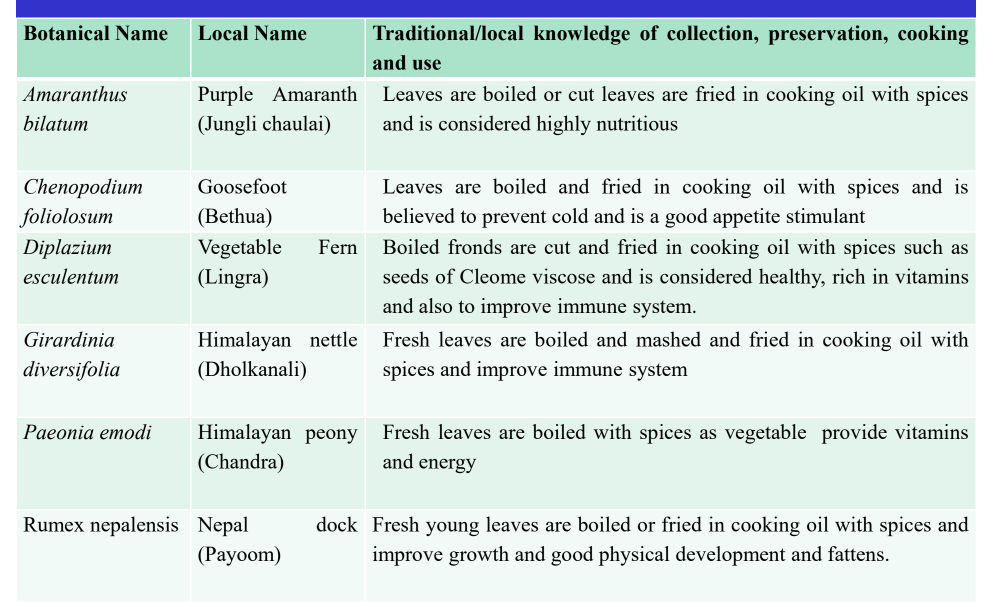
Wild fruits/edibles consumed /used by the local people to increase immune defenses during COVID-19 (as prescribed by local Vaidya's/herbalists) in Central Himalaya, Uttarakhand

Plant species	Local name	Part used/mode of application	Medicinal and other uses
Aegle marmelos	Stone apple (Bel)	Fruits used for juice and squash	The fruit is aromatic and is used in curing of peptic ulcer, constipation, scurvy and dysentery and is said to act as a tonic for the heart and brain.
Berberis asiatica	Indian barberry (Kingore)	Fruits used for juice and squash	It is good remedy for stomachache, diabetes, cold and cough fever.
Elaeagnus latifolia	Bastard Oleaster (Gewain)	Fruits used for juice, squash and sauce	Believed to be good in cough and bronchitis. It is capable of reducing the incidence of cancer.
Embilica officinalis	Indian gooseberry (Amla)	Fruits used for pickle and juice	The fruit juice is used to cure cough, anemia, piles and diabetic.
Spondias pinnata	Hog plum (Amara)	Fruits used for juice, squash and sauce	The fruit is good source of vitamin `C' and used to cure diabetes, heart ailment, urinary troubles etc.

Spices/condiments of wild origin consumed /used by the local people to increase immune defenses during COVID-19 (as prescribed by local Vaidya's/herbalists) in Central Himalaya, Uttarakhand		
Plant species & part used	Traditional and medicinal use	
<i>1. Allium humile (</i> Small alpine onion) Local name: Ladum	Traditionally fried paste is added to meat & culinary purposes, Medicinally it is useful to cure jaundice, cough and cold.	
 Alium rubellium (Purple flowered garlic) Local name: Doodhu 	Traditionally used for flavoring as condiments, and is reported to be good for the patients suffering from jaundice and also useful in cold and cough.	
 3. Angelica glauca (Angelica) Local name: Choru 4. Pleurospermum angelicoides Local name: Chhipi 	Traditionally the aromatic root is a flavoring agent and is commonly used as spice and condiment Leaves and stem of the plant are useful to cure dysentery and provide relieve from body pain caused due to extreme cold. Roots are used as spices and condiments. The decoction of the roots is used to cure typhoid fever, stomach pain, body pain, etc.	
5. <i>Carum Carvi</i> (Caraway) Local Name: Kala Jeera	The seeds are used as spice or condiments and also it is used to cure dyspepsia, cold and cough, an appetizer.	
 6. Cinnamomum tamala (Indian bay leaf) Local name: Tejpata, dalchini 	Leaves of C. tamala (tejpata) are widely used as a spice and condiments to improve the appetite and digestion.	

काषा ज्योतिरशीम्

Wild greens used as a food/vegetables by the local people to increase immune defenses during COVID-19 (as prescribed by local Vaidya's/herbalists) in Central Himalaya, Uttarakhand



Wild and cultivated plants and quantity of different ingredients (gm/liter) used in preparation of hot beverages (teas) by the traditional mountain communities to boost the immune system in treating corona disease



Name of plant and Ingredient used	Combination (gm/litre)
1. <i>Taxusbaccata</i> bark	1.00 gm
2. Common Salt	25gm
3. Purified butter (Ghee)	1-2 tea spoon (10-20 gm)
4. Pinna (dry mixture of walnut and apricot kernels, flour of fried wheat/barley/buckwheat is locally called pinna)	20-30 gm
5. <i>Bergenia ligulata</i> leaves	20gm
6. <i>Betula utilis</i> gum (resin)	10gm
7. Origanum vulgare leaves	30gm
8. Cinnamomum tamala (Tejpataa leaf+ bark)	Tejpataa leaf 5 gm and bark 20gm
9. Kadha (mixture of four different plant products such as ginger (<i>Zinger officinale</i>), black pepper (<i>Piper nigram</i>), cardamom (<i>Cardamom Elettaria</i>) and clove (<i>Syzygium aromaticum</i>) in different proportions)	Ginger (50 gm) Black pepper (5 gm) clove (2 gm) cardamom (2gm)

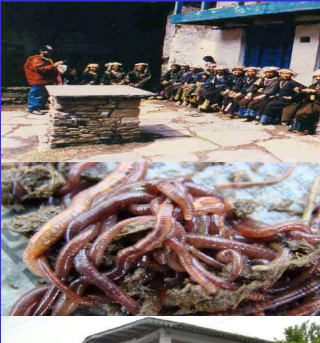
Success stories linked to Bio cultural Conservation & Sustainable Development



Garhwal Regional Centre

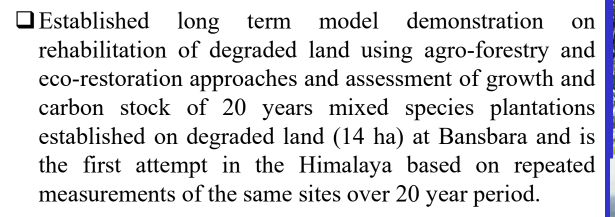
Unique model for livelihood enhancement and biodiversity conservation through participatory action research for resolving policy-people conflicts in Nanda Devi Biosphere Reserve (NDBR).

In-situ conservation of traditional mountain agrobiodiversity through village community participation in Urgam valley, Chamoli (a joint initiative of GBPIHED-GRC, NBPGR, New Delhi and village community).





Cont..



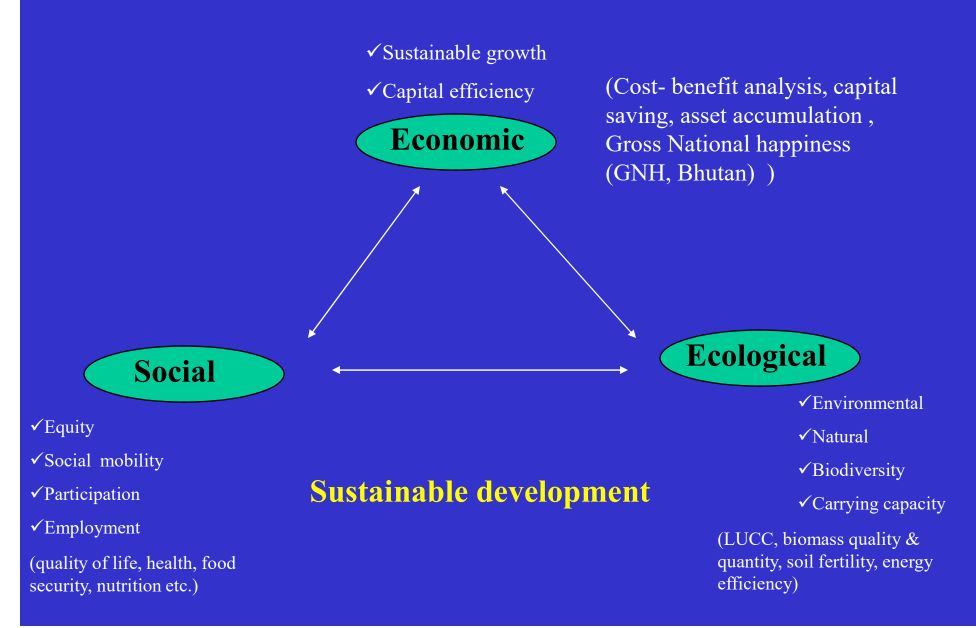
- □ Wild bio-resource utilization (38 potential wild spices) for livelihood enhancement, income generation and conservation.
- Promotion of medical plants sector through improvement of cultivation/ conservation practices and value addition of high value species.





Indicators of SD are varied:- therefore, monitoring & evaluation has to be done using a number of currencies.





Unsustainable development Insecure and Unsustainable development **Powerless** alienation and **Degradation** conflicts of Natural resources X **Environmental** Social **Economic** vulnerability **Vulnerability** vulnerability

Himalayan ecosystem and people inhabited there are subject to intense economic, social and physical/environmental vulnerabilities.

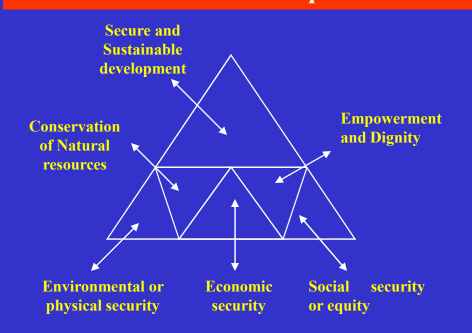
Don't have access to the pathways of economic growth

□Victims of natural calamities

- □Marginalized by most social and political resources
- Outsiders have control over their natural resources

□Poverty, hopelessness resulted into conflicts – which leads unsustainable development.

Sustainable development



Need for:

Decentralizing governance to local level

Developing policies and programmes which support sustainable use and environmentally sound management of natural resources

■Need for equal social rights and opportunities for participation in governance by all sections of the societies including women

□Involvement of diverse civil society and especially grassroots networks and associations to achieve more voice and influence in policy and public action choices.

Rural landscape Traditional agriculture (Food and Nutrition Security) in the CHR and the Sustainable Development Goals



- ➢ Goal 1- End poverty in all its forms everywhere
- Goal 2- End hunger, achieve food security and improved nutrition, and promote sustainable agriculture.
- ➢ Goal 3- Ensure healthy lives and promote well-being for all at all ages.
- ➢ Goal 5- Achieve gender equalities & empower all woman & girls.
- ➢ Goal 6- Ensure availability and sustainable management of water and sanitation for all.
- Goal 8- Promote sustained economic growth, full and productive employment & decent work for all.
- ➢ Goal 12- Ensure sustainable consumption and production patterns.
- ➢ Goal 13- Take urgent action to combat climate change and its impacts.
- Goal 15- Protect, restore & promote sustainable use of ecosystem, combat desertification and halt and reverse land degradation & halt biodiversity loss.

Priority action points for policy planning towards the sustainable development of hill agriculture of the central Himalayan region



- Develop decentralised approaches for the mobilisation and strengthening of formal and informal decision-making institutional mechanisms
- Redefi-ne research and development (R&D) priorities with a regional focus
- Develop strong linkages between R&D institutions, agricultural universities/NGOs and the private sector
- Improve integration of cross-sectoral linkages and interdependencies between different policies.
- Replicate success stories and identify lessons from failures
- Transfer appropriate hill-specific agro-technology to user groups
- Address human resource development issues in policies
- Properly implement extension and support services systems
- Ensure conservation of traditional agrobiodiversity and associated traditional knowledge
- Improve effectiveness of existing agricultural institutions, their arrangements and capabilities
- Promote organic cultivation, emphasising traditional hill crops and value addition

Priority interventions for conservation and management of bioresources of wild origin of the Central Himalaya



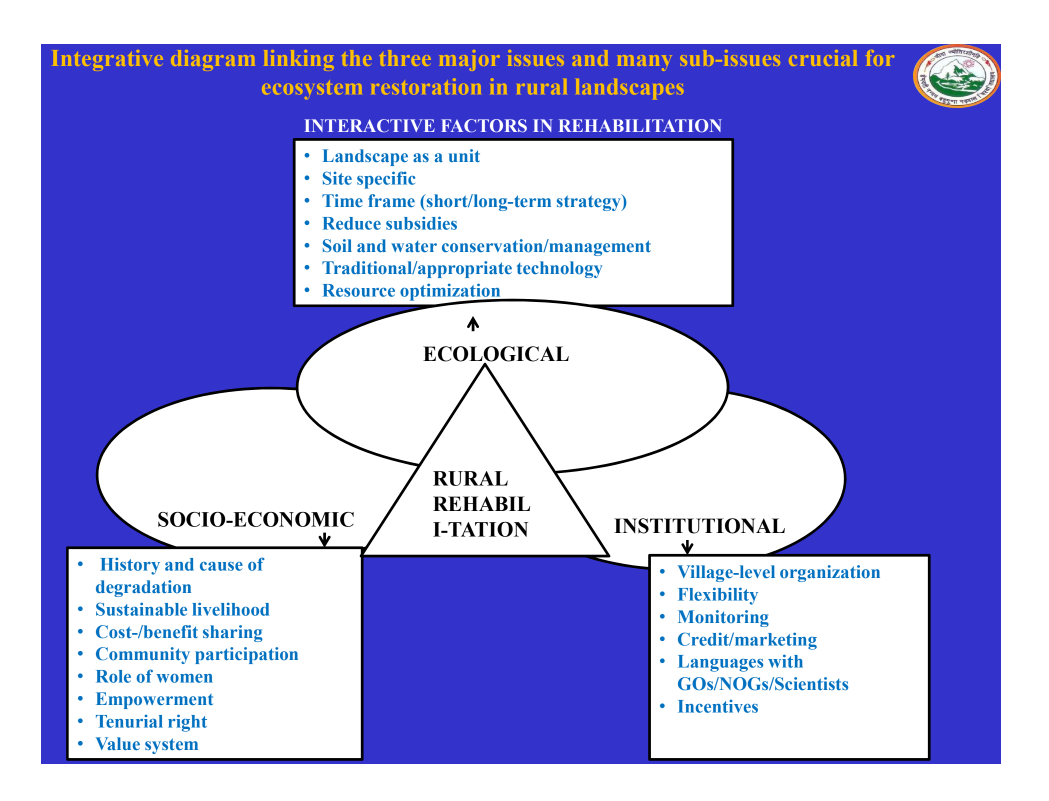
- Strengthen research and development into indigenous knowledge of medicinal and aromatic plants used in THCSs.
- Increase the quality and quantity of clinical trials and certification.
- Promote networking to facilitate communication and collaboration with traditional communities and researchers/experts.
- Establish appropriate and suitable frameworks and approaches for intellectual property rights, benefit sharing and to provide due recognition to the traditional societies for their creation of the indigenous knowledge system.
- Develop appropriate mechanisms of sustainable harvesting of medicinal and aromatic plants with the conservation and management of natural resources and promote cultivation of those plant species that are mostly used in the THCS.
- Develop a holistic policy to address conservation, cultivation, utilization, traditional knowledge, trade, intellectual property rights and research in Himalayan states.
- Create awareness of conservation and management of threatened medicinal and aromatic plants through training, workshops, publications, and school curricula.

Conclusion



- In this context, Mahatma Gandhi can be quoted, who aptly said.
- Earth provides enough to satisfy every man's need, but not every man's greed. If this mantra (statement/or principle or hymn) were adhered to, it would definitely make a significant contribution in augmenting the natural resource base of the Indian Himalayan Region.
- **The Isa Upanishad says.** The whole universe is pervade by God' Nature has spiritual significance. It has implications for the survival of humanity as well. Protection of nature is protection of self.

Thank You





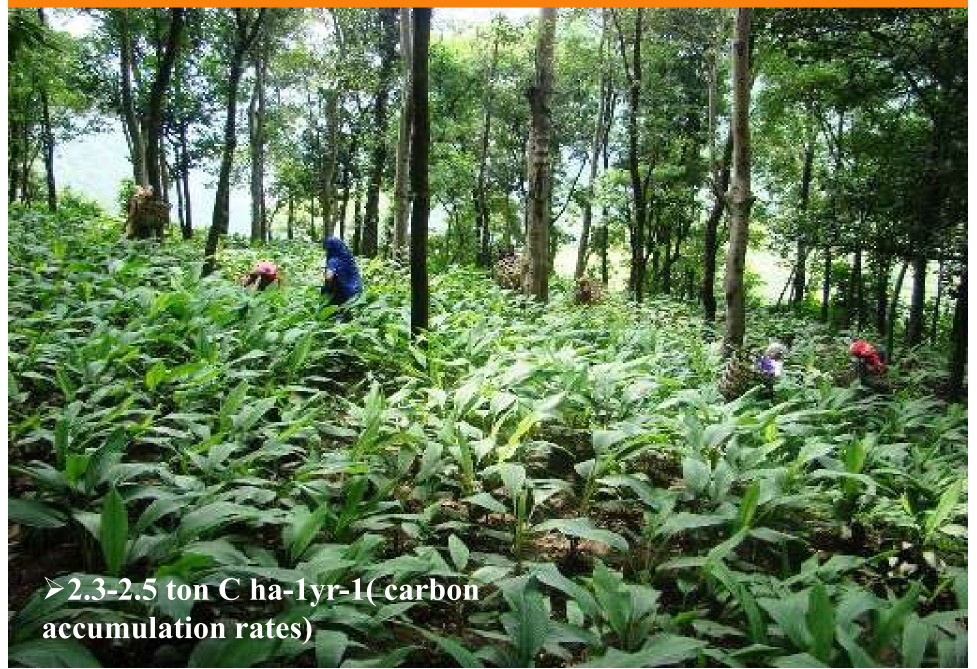
Priority R&D – Approaches for ecosystem restoration in the Himalayas

- Ecological restoration of degraded forest land
- Abandoned agriculture land restoration through Agroforestry
- Sylvi-pasture
- Agri-horticulture
- MAPs cultivation
- Environmental education (School Children primary and secondary level)

Banswara before Plantation



Integrated agro-forestry with MAPs cultivation



Land rehabilitation through Restoration ecology approach (6ha)

Help to mitigate climate change impact

The ecosystem restoration models demonstrated contributes in following major areas

- Reducing drudgery of women and reduce pressure on forests.
- Reduce crop damage by wildlife (turmeric cultivation).
- Improve carbon sequestration climate change mitigation.
- Improve economy and livelihoods

Need to tap the wisdom of traditional practices



- There is a need to tap the wisdom of traditional practices as well as the best of modern technology. Many traditional environmental practices are still relevant today.
- Dig wells
- Excavates water tanks
- Plants trees: whenever a body is cremated, plant a tree
- Don't cut green trees
- Create parks and flower garden (don't pick flowers at night)
- Don't disturb water at night (allow pollution to settle)
- Don't pollute (defecate on) river banks
- Adopt a simple, non-violent life style
- Reduce consumption and harm to the environment
- Recycle a reuse
- **The Isa Upanishad says.** The whole universe is pervade by God' Nature has spiritual significance. It has implications for the survival of humanity as well. Protection of nature is protection of self.

SS-5.Nanda Devi Biosphere Reserve: Resolving Policy Conflicts through Participatory Action Research and Model Demonstration



• Establishment of biosphere reserve resulted in curtailment of rights of local people, nor are they provided with adequate alternatives for meeting their resources.

Achievement and knowledge products

- Assessment of policy-People Conflicts
- The main reasons/causes of the conflict was that ban of expedition on Nanda Devi Peak, restriction on grazing in core zone, ban on NTFPs collection and crop and livestock depredation by wildlife.
- Scientific and participatory Action Research
- An integrated study was undertaken between 1992 to 2007 in 10 buffer zone villages considering people's perceptions, attitudes towards the reserve establishment and assessment of ecosystem function of buffer zone villages using ecological and economic currencies.
- The results of the altitudinal survey indicated that, majority of the respondents (75%) had negative attitude, about 15% were found neutral and 10% have positive attitude, despite the fact, that all respondents were receiving some benefits from eco-development programs implemented by NDBR authorities.
- > The impacts on people of establishment of biosphere reserve were studied. More than 95% of respondent expressed that agriculture and horticulture productivity is getting reduced, to a large extent, for reasons related to conservation policies.

Priority interventions for socio-economic development, conflict resolution and biodiversity conservation

Agriculture and Horticulture

- Based on the eco-energetic studies in agro-ecosystem suggested that mixed cropping of Solanum tuberosum + Phaseolus vulgaris + Amaranthus spp. Or Solanum + Phaseolus are ecologically and economically viable.
- Vegetable cultivation under protected condition.



- Value Addition in Wild Edibles and other Forest Products
- > Promoting cultivation of medicinal plants those have local market
- > Degraded land rehabilitation through linking appropriate cost-effective technologies
- Developed guidelines, strategies and action plan for ecotourism promotion and management: moving towards a community- centered approach
- Through our in-depth action research in the field of eco-tourism helped policy-planners to revisit the issues related to mountaineering/expedition in the core zone. Finally in 2003 the core zone of BR has been opened to regulated tourism with restrictions on tourist numbers (around 500 tourists per year).
- Conservation education and awareness
- In-depth study over last 25 years on various aspects changed the mindset of the people and now over 70% people have positive attitude towards biosphere reserve.
 - Developed strategies and action plan on "Promoting Eco-tourism in Nanda Devi Biosphere Reserve.
 - The wide scientific coverage on NDBR motivated UNSECO South Asia to document the outcome which was brought in the form of the film *"Invocations to the Mountain Goddess"*.
 - The action and participatory research work carried out in the reserve on various aspects is given due consideration by various line departments/agencies at district and state levels and most of our findings has been incorporated in the action plant of Forest and Tourism departments.

SS-3. Conservation and Sustainable Use of Medicinal Plants through Field based Model Demonstration, Cultivation and Outreach Programme



- > The medicinal and aromatic plants (MAPs) have huge economic potential
- Programme initiatives:
 - Sharing of indigenous knowledge of agronomic practices and use of MAPs.
 - Promoting farmers for nursery raising and large scale cultivation of potential species.

Achievements & Knowledge Products

- Documented traditional agronomic practices, wild collection and uses of 18 MAPs species.
- Promoting domestication/cultivation, sustainable use/value addition and conservation of high value low volume MAP species (*Picrorriiza kurooa, Saussurea costus, Vallenriana wallichii, Inula racemosa, Angelica glauca, Allium stracheyi, etc.*).
- Developed linkages of MAPs growers with value chain, marketing and buy back systems (Emami Pvt. Ltd., Kolkota).
- Skill development of local people through on-site training and field demonstration and organized 35 farmers training programme between 1996 to March 2015 in 9 hilly districts of Uttarakhand and trained about 889 participants.
- Established 5 medicinal plants model demonstration sites covering 4 ha. of village common land in four districts (i.e., Tehri, Uttarakashi, Chamoli and Rudraprayag).
- Raised about 5.0 lakhs seedlings of 8 MAPs species and about 3.25 lakhs seedlings distributed among the farmers and NGOs. As a result 94 farmers started cultivation of MAPs.
- The findings of this long term action research programme helped institutions involved in medicinal plant sector to develop enabling policy environment regarding cultivation/collection and marketing of MAPs for economic development.

SS-2. Bio-resource (wild origin) utilization and conservation for Livelihood Enhancement and enterprise development



- Forest bioresources constitute an important source of livelihood of millions of people across the Himalaya
 - Of the nearly 800 species of wild edibles reported from India, 344 species are found in central Himalaya.

Achievements and knowledge products

- > Potential bioresources and availability for enterprise development
 - More than 35 plant species having high economic value screened for value added product development (i.e., juice, squash, jelly, pickle, spice, condiments and medicines).
 - Information and database on distribution, traditional uses, phenophases, fruit yield, sustainable harvesting, conservation and cost-benefit analysis
 - Assessment of nutraceutical potential (food, vitamins, minerals, macro and micro nutrients, etc.) of 35 species.
 - Developed the package of practices for sustainable use of 35 wild bioresources for value addition and enterprise development.



- Capacity and skill development programme organized 18 trainings (each of 2 days) between 2008 2017 and trained about 662 people.
- Impact and replicability: About 525 households adopted value addition of wild bioresources as an off-farm activity and average additional income earned through this venture was estimated Rs. 11,700/HH/yr.
 - The approach and action research frame work for bioresources utilization has been included in the policy and action plan of the Uttarakhand government particularly livelihood extensions programme/activities.

Some barriers to the promotion and mainstreaming of agrobiodiversity for improved diets and nutrition



- •Disconnect between the biodiversity, agriculture and health sectors and other sectors (including education)
- Continued neglect by the international and national research and extension systems
- Biodiverse food-based approaches all too often fall outside the traditional scope of clinical nutrition and public health
- Lack of skills and institutional capacity necessary to promote multi sector approaches to fully exploit biodiversity, agriculture and health linkages
- Lack of data linking biodiversity to dietary diversity and improved nutrition outcomes
- Lack of evidence demonstrating or comparing the most (cost-) effective methods and approaches for delivering or mobilizing biodiversity for dietary and nutrition outcomes
- Poorly developed infrastructure and markets for the majority of biodiversity for food and nutrition
- Inadequate agricultural and food security policies and strategies that promote major cereal staples have often diminished the dietary role of more nutritious species such as millets, indigenous fruits and vegetables and roots and tubers
- Negative perceptions and attitudes to local, nutritionally-rich traditional biodiverse foods
- Non-tariff barriers and strict food safety assessment regulations such as the European Union's Novel Foods Regulation (NFR) which places a considerable burden of proof on those attempting to bring traditional biodiverse foods and their products to markets
- The 'artificial' cheap cost of exotic or imported foods which externalize their health and environmental costs.



Challenges to traditional food dietary diversity and nutrition security

Challenge	Consequences	Dimension of food and nutrition security likely to be affected negatively
Deterioration of local food system	•Reduced food production & diversity	•Food availability utilization
Changing diets	Reduced dietary diversity	•Food utilization
Climate change	•Risks to agriculture production, Diversity & Farm income	•Food availability, Stability, Utilization & Accessibility
Lingering poverty	•Reduced food intake & Dietary Diversity	•Food accessibility & utilization
Increased outmigration	•Labour shortages in agriculture leading to reduced production	•Food availability
Abandonment of cultivable land	•Low returns, land abandonment and loss of production	•Food availability
Rapid urbanization	•Encroachment of agriculture land leading to reduced agricultural production	•Food availability
Inadequate infrastructure and market centers	•Inadequate food distribution & post-harvest losses , Higher prices external food items	•Food accessibility & availability
Depletion of natural resources	 Loss to water resources & biomass manure form forests Reduced supply of wild edible, and reduced livestock production and income 	•Food availability, availability, utilization, accessibility
Constraints to internal food movement	•Reduced food supply to mountains, Higher prices of available food	•Food availability & accessibility
Inadequate access to improved drinking water, sanitation, and hygiene	•Higher prevalence of disease	•Health status



The major causes which are directly and indirectly responsible for this genetic erosion and creating imbalance in traditional agro-ecosystem are:

- 1. Infra-structural development, roads and consequent exposure;
- 2. Illusions about quality of coarse and fine grains;
- 3. Aspirations for off-farm employment;
- 4. Lack of specified programmes and incentives;
- 5. Abandonment of traditional agro-ecosystem by indigenous population;
- 6. Socio-economic and cultural change;
- 7. Migration of hill population to the plains in search of employment;
- 8. Deterioration of traditional agro-ecosystem caused by introduction of modern high-yielding crops and crop varieties;
- 9. Replacement of mixed cropping by mono-cropping;
- 10. Deterioration of natural habitat caused by man induced environmental changes;
- 11. Lack of scientific interest in these crops.

C. Cultivation and conservation of medicinal and aromatic plants (MAPs)



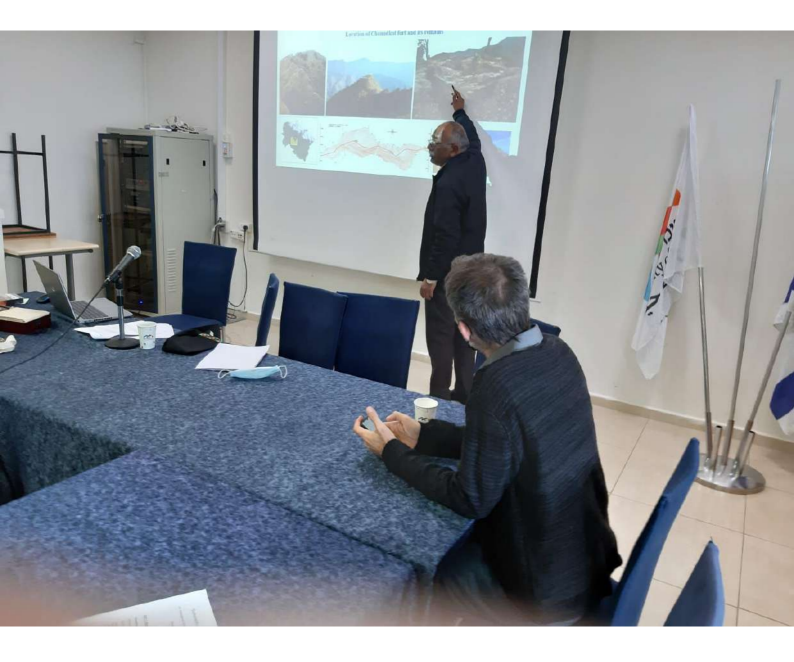


Large scale cultivation of *Picrorrhiza kurrooa*

Large scale cultivation of *Arnebia benthamii*









בית מגורים מפואר מהתקופה הרומית

Mansion from the Roman Period

בית המגורים המפואר נבנה לפי מתכונת אדריכלית, שהייתה רווחת בעולם הרומי במאה השלישית לסה״נ. במרכז הבית טרקלין, שהיה חדר אוכל ואירוח. יתר חדרי הבית, הבנויים רובם בשתי קומות, הקיפו את הטרקלין ואת חצר העמודים הסמוכה לו.

בחדרים רבים נחשפו רצפות פסיפס, מהן לבנות ומהן מעוטרות. המפוארת שבכולן היא רצפת הטרקלין, ובה תיאורים מחיי האל דיוניסוס ועיטורים נוספים.

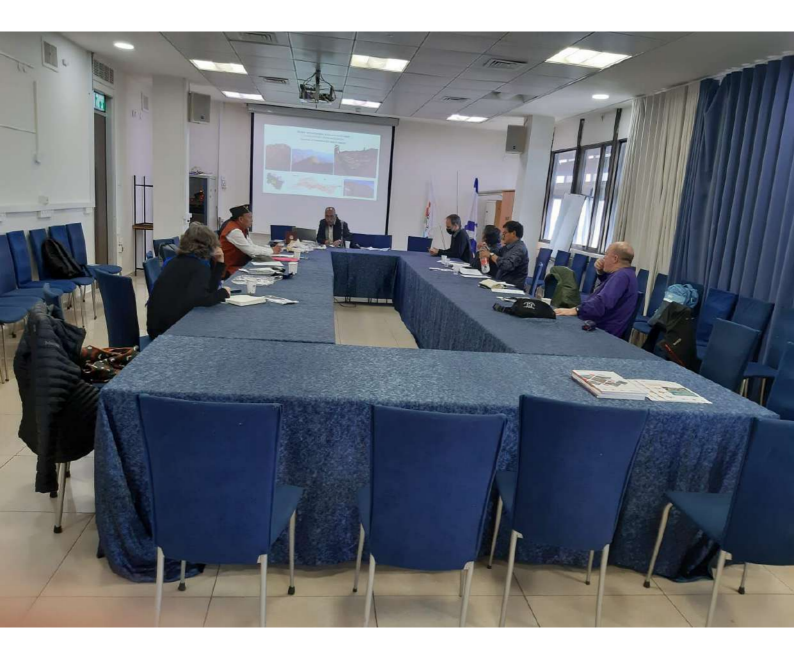
The mansion was built according to a plan that was popular in the Roman world in the third century CE. In the center of the house was a traclinium (living-dining room) and colonnaded courtyard. Surrounding them were other rooms.

Many of the rooms had mosaic floors, some white and others with colorful patterns. The most ornate of them all is the floor of the traclinium, containing scenes from the life of the god Dionysus.









Letter of Intent with

- 1. Central University of Kashmir
- 2. Tripura University
- 3. Tezpur University
- 4. Sikkim University, Gangtok
- 5. Rajiv Gandhi University, Itanagar
- 6. North Eastern Hill University
- 7. Nagaland University
- 8. Mizoram University
- 9. Manipur University
- 10. CSIR-Institute of Himalayan

Bioresource Technology, Palampur (HP)

- 11. Central University of Jammu
- 12. Central University of Himachal Pradesh
- 13. Assam University, Silchar



ONE HIMALAYA ONE POLICY INDIA'S G20 PRESIDENCY AND SUSTAINABLE **DEVELOPMENT IN THE HIMALAYA:**

OPPORTUNITIES, STRATEGIES AND POLICY RECOMMENDATIONS

SUBMITTED TO

NITI AAYOG, New Delhi, Government of India and PMO, Government of India, New Delhi

PREPARED BY

HNB Garhwal University, Srinagar, Garhwal, Uttarakhand (HNBGU) GB Pant National Institute of Himalayan Environment, Kosi Katarmal, Uttarakhand (NIHE) Research and Information System for Developing Countries (RIS) Indian Himalayan Central Universities' Consortium (IHCUC) Kalinga Institute of Indo Pacific Studies (KIIPS)



2023



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i			GBPNIHE, Kosi Katarmal, Almora	
		INDIA'S G20 PRESIDENCY		

INDIA'S G20 PRESIDENCY AND SUSTAINABLE DEVELOPMENT IN THE HIMALAYA: OPPORTUNITIES STRATEGIES AND POLICY RECOMMENDATIONS



connent prepared



(Thematic Study-I)



NITI Aayog **Enumeration and Valuation of the** Economic Impact of Female Labour in the Hills



A Study of Indian Himalayan Region (IHR) assigned by NITI AAYOG, New Delhi funded by UGC, New Delhi



Submitted by Indian Himalayan Central Universities Consortium (IHCUC) February, 2022



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(Thematic Study-II)



Agro-Ecology in Himalayan States with special emphasis on marketing





Submitted by Indian Himalayan Central Universities Consortium (IHCUC) February, 2022



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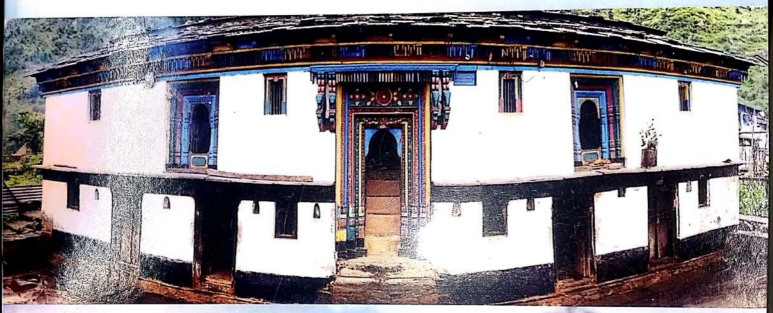
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(Thematic Study-III)



Development of Eco-friendly and Cost-effective Tourism in Hills



A Study of Indian Himalayan Region (IHR)

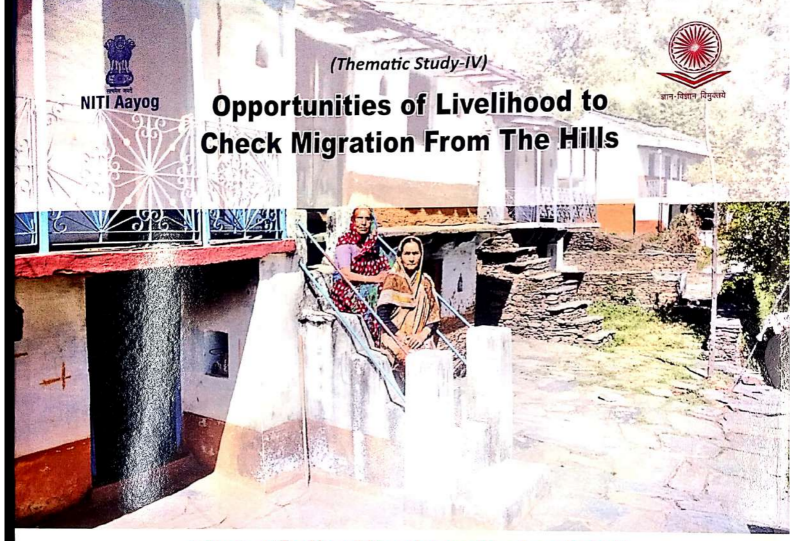
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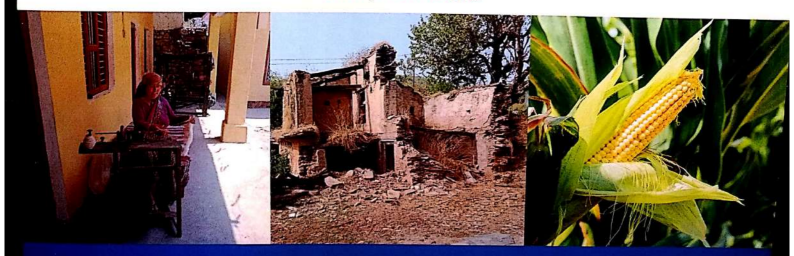


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A Study of Indian Himalayan Region (IHR) assigned by NITI AAYOG, New Delhi funded by UGC, New Delhi



Submitted by Indian Himalayan Central Universities Consortium (IHCUC) Feb, 2022



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(Thematic Study - V)

Water Conservation and Harvesting **Strategies**

> Study of Indian imalayan gion (IHR)

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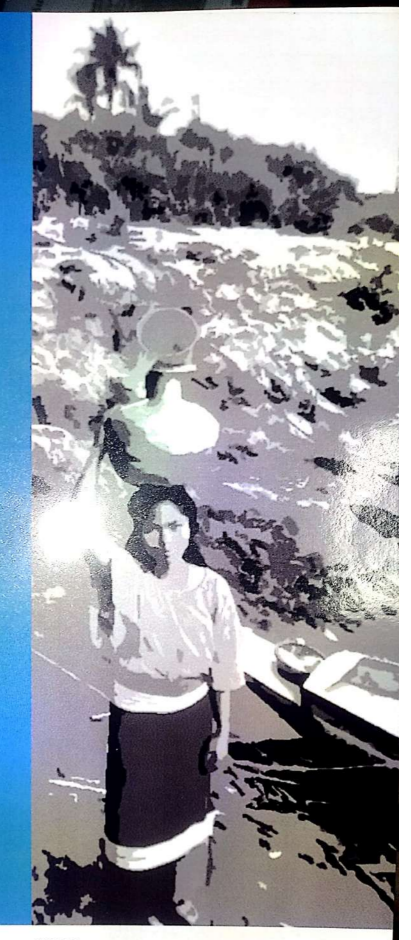


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February, 2022 Submitted by Indian Himalayan Central Universities Consortium (IHCUC)



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MoU with IIT Kanpur (2017, 2021)

MoU with IIT Kanpur signed in 2021

CHARACTERIZATION OF GROUNDWATER IN SEISMICALLY ACTIVE REGIONS OF UTTARAKHAND, INDIA: IMPLICATIONS FOR EARTHQUAKE INDUCED VARIATIONS

In this project we are collecting the water samples from 7 different sites of Himalayan region. We have selected two Groundwater Monitoring Stations in proximity of MCT, one is Agyasthamuni (Lat: 30.390725, Long: 79.026365), and another is Gopeshwar (Lat:30.409526, Long:79.319797). We have installed sophisticated monitoring instruments such as multiparameter water quality sensors with high frequency data transmission capabilities for analysis of pH, EC, ORP, Alkalinity, Water level, major anions (HCO₃, CO₃, SO₄, NO₃, Cl, F) and cations (Ca, Mg, Na, K, Fe, Al, Si) and stable isotopes ratios of Hydrogen (²H) and Oxygen (¹⁸O). We are collecting samples from 2 hot springs located in Gauri Kund (Rudraprayag) and Tapovan (Chamoli) for analysis of pH, EC, ORP, major anions (HCO₃, CO₃, SO₄, NO₃, Cl, F) and cations (Ca, Mg, Na, K, Fe, Al, Si) and stable isotopes ratios (Hydrogen (²H), Tritium and Oxygen ("O). Apart from it we are collecting 3 surface water samples near the monitoring stations, 2 samples of Mandakini River and 1 sample from Alakananda River for analysis of pH, EC, ORP, major anions (HCO₃, CO₃, SO₄, NO₃, Cl, F) and cations (Ca, Mg, Na, K, Fe, Al, Si) and stable isotopes ratios of Hydrogen $({}^{2}H)$ and Oxygen $({}^{18}O)$. All the samples are collected at a monthly interval and analysed in laboratory of National Institute of Hydrology, Roorkee. For the analysis advanced instruments such as Inductively Coupled Plasma Optical Emission Spectrometer, Ion chromatograph and Mass Spectrophotometer have been used. These instruments provide a full spectrum analysis of water chemistry in a cost and time effective manner.



Groundwater monitoring station at Gopeshwar.

Groundwater monitoring station at Agasthyamuni.

Objectives:

(1)Monitoring groundwater level and groundwater quality in the active seismic belt of Uttarakhand in 2 sites.

(2) Analyse and characterize space-time variations in groundwater with respect to contemporary seismic events.

(3) Develop a process-based understanding of the observed anomalies and the mechanisms triggering them.

(4) Explore the potential to utilize groundwater anomalies for the development of precursor for Earthquakes.

Activity Report: MoU with IIT Kanpur, 2017

Himalayan Cloud observatory (HCO) BadshahiThaul, Tehri Garhwal

Joint Collaboration with Department of Physics, Srinagar and Tehri Campus, Hemvati Nandan Bahuguna Garhwal University Srinagar Garhwal & Department of Civil Engineering, Indian Institute of Technology Kanpur

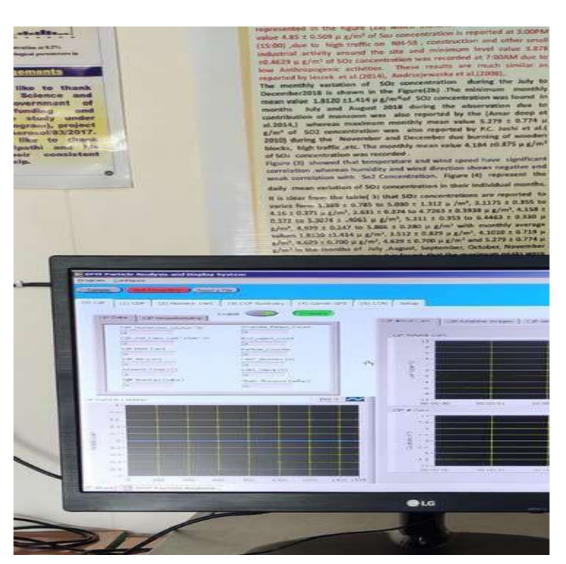
Funded by Government of India, Department of Science & Technology (DST)Strategic Programmes, Large Initiatives andCoordinated Action Enabler (SPLICE)CLIMATE CHANGE PROGRAMME

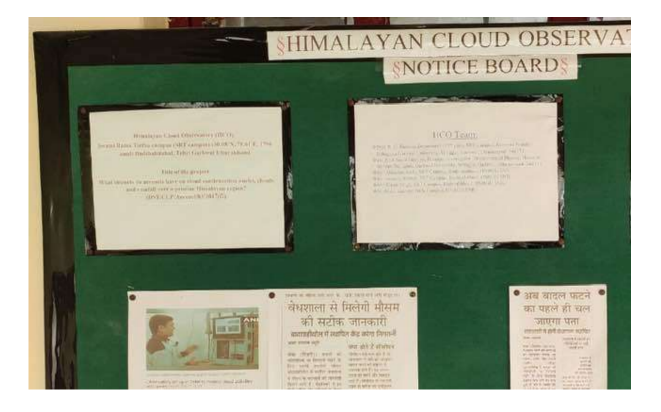
DST Funded project (SPLICE –Climate Change Program) research project entitled "What impacts do aerosols have on cloud condensation nuclei, clouds and rainfall over a pristine Himalayan region"

The Himalayan cloud observatory (HCO) is located on the slope of enchanted lesser Himalayan Mountain range surrounded by the dense forest of Alpine, Oak and Deodar trees. The site is situated in Department of physics HNB Garhwal University, SRT Campus Badshahithaul (30⁰34' N and 78⁰41'E) Tehri Garhwal District of Uttarakhand at altitude of 1725 m above mean sea level (ASL). The Himalayan cloud observatory (HCO) is situated on outskirts of new Tehri and Chamba cites (8 km and 3 km away from city center to the North East and Northwest respectively) with the population of 6.19Lakh and population Density of 170 km⁻² as per 2011 census report. The residential area is about 300 m apart from the HCO. The NH-34 passes with moderate traffic on other side of hill to the West direction. The site has no major industrial activities in Radius of 10 km, but Tehri Dam Hydro project is in the Range of 25 Km. The Himalayan cloud observatory is establishing to take ground-based observation for the study Aerosols, Cloud condensation nuclei (CCN) particles and meteorological Parameters over the large region of Uttarakhand to understand the complex mechanism of cloud bust and climate change in sensitive Himalayan region in all-weather condition.

Collaborating Team: Prof R C Ramola, Director, SRT Campus Badshahithaul Dr Alok Sagar Gautam, Srinagar Campus, HNB Garhwal University Prof S N Tripathi, IIT Kanpur Dr U C Dumka, Aries Nainital Dr Vijay Kanwade, University of Hyderabad Mr Anil Mandaria Mr Gaurav Mishara Mr Akash Rathi Mr Sanjeev Kumar Mr Karan Singh







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बादल बनने की प्रक्रिया पर ब



गढ़वाल केंद्रीरा विवि के भौतिक विज्ञान विभाग की हि साइजर यंत्र लगाते विवि भौतिक विज्ञान विभाग की रं खॅ.आलोक सागर गौतम व प्रो.आरसी रमोला • ज

भी काफी म	ादद मिलं	गेगी जि	ससे शोध	में
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जागरण संताददाता, श्रीनगर गढ़ताल : उत्तराखंड के पर्वतीय क्षेत्रों में मौसम में होने वाले बदलाव और बादल बनने की प्रक्रिया को समझने के लिए गढ़वाल केंद्रीय विश्वविद्यालय श्रीनगर के भौतिक विज्ञानियों ने शोध कार्य आरंभ किया है। विवि के चौरास परिसर में भौतिक विज्ञान विभाग की वातावरणीय भौतिकी प्रयोगशाला में यह शोध कार्य संचालित किया जा रहा है। वैज्ञानिक एक रिपोर्ट तैयार कर प्रदेश सरकार से शोध के आंकड़ों का विश्लेषण और अध्ययन के निष्कर्ष को साझा करेंगे।

विष्ठवविद्यालय के शोध



प्रक्रिया को मिलेगी। प आरसी रमो से शोध का और यह य साबित होग विज्ञान और मदद मिलेगं विज्ञान एवं सरकार द्व परियोजना जलवायु क मददगार स बताया कि विश्लेषण रचाप्तंट

• पहाड़ों के वातावरण, मौसम व जलवायु को समझने में मदद मिलेगी

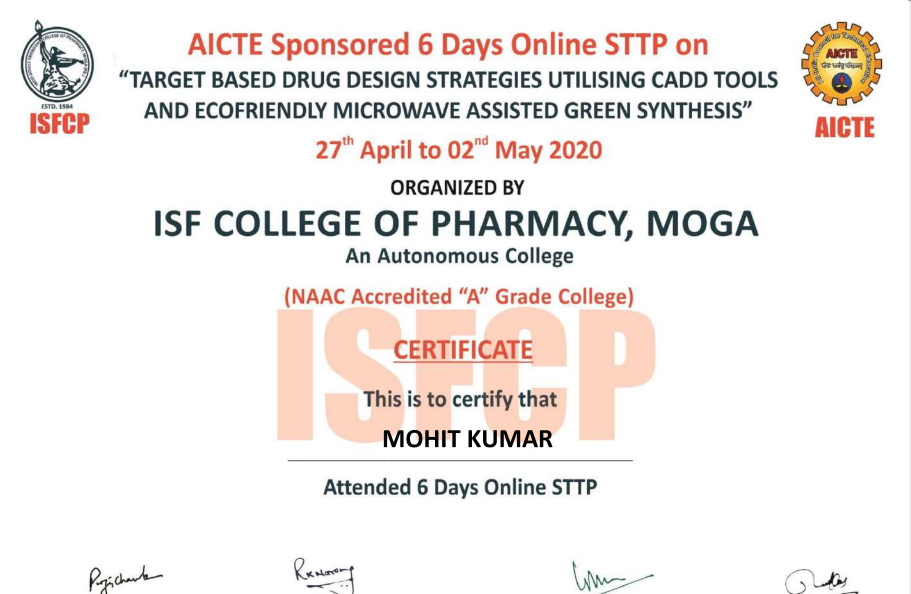
शाह टाइम्स संवाददाता श्रीनगर। हेमवती नंदन बहुगुणा गढ़वाल विवि श्रीनगर की ओर से एसआरटी कैंपस बादशाहीथौल में स्थापित हिमालयन क्लाउड आण्जरबैटरी में नैनो स्कैन स्कैनिंग मोबिलिटी पार्टिकल साइजर यंत्र का सफलतापूर्वक स्थापित किया गया। परियोजना विज्ञान एवं तकनीकी मंत्रालय भारत सरकार के निदेशक एंव भौतिक विभाग में सहायक पोफेसर टॉ आलोक सागर गौतम ने

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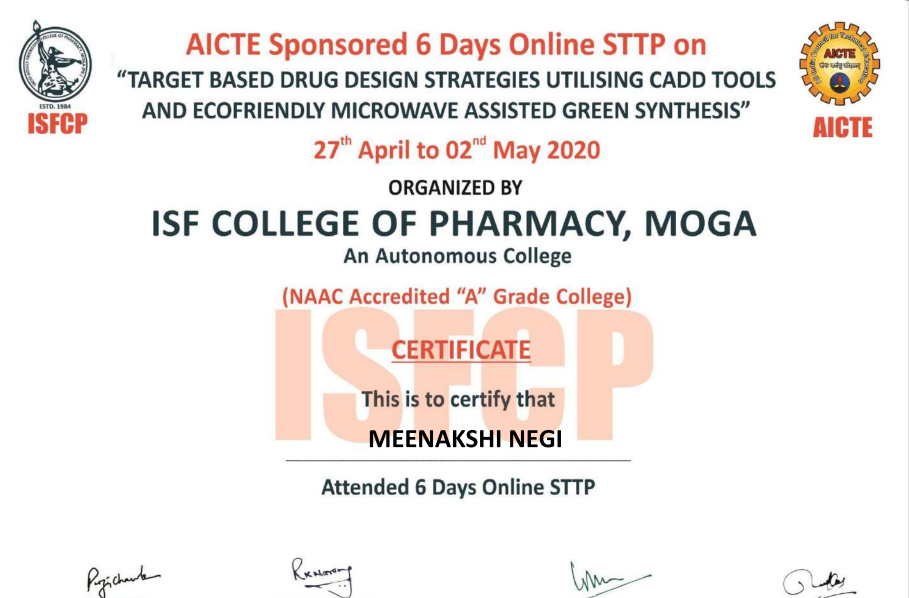
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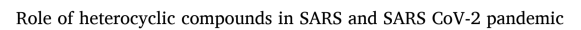
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ABSTRACT

Coronaviruses have led to severe emergencies in the world since the outbreak of SARS CoV in 2002, followed by MERS CoV in 2012. SARS CoV-2, the novel pandemic caused by coronaviruses that began in December 2019 in China has led to a total of 24,066,076 confirmed cases and a death toll of 823,572 as reported by World Health Organisation on 26 August 2020, spreading to 213 countries and territories. However, there are still no vaccines or medications available till date against SARS coronaviruses which is an urgent requirement to control the current pandemic like situations. Since many decades, heterocyclic scaffolds have been explored exhaustively for their anticancer, antimalarial, anti-inflammatory, antitubercular, antimicrobial, antidiabetic, antiviral and many more treatment capabilities. Therefore, through this review, we have tried to emphasize on the anticipated role of heterocyclic scaffolds in the design and discovery of the much-awaited anti-SARS CoV-2 therapy, by exploring the research articles depicting different heterocyclic moieties as targeting SARS, MERS and SARS CoV-2 coronaviruses. The heterocyclic motifs mentioned in the review can serve as crucial resources for the development of SARS coronaviruses treatment strategies.

1. Introduction

Heterocyclic scaffolds play a pivotal role in drug discovery and development as they constitute the key structural component of a majority of biologically active moieties. Their ability to interact with almost every cellular mechanism in living organism has been responsible for their versatile nature. Their interaction with different mechanistic pathways in viruses has continuously been exploited by researches for the designing of heterocycle-based antiviral agents. Several FDA approved drugs currently in the market comprise of different

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Review article



Abbreviations: 2019-nCoV-2, 2019-novel coronavirus; 3CLpro, 3chymotrypsin-like protease; 9-O-SIA, 9-O-acetyl-N-acetylneuraminic acid; ACE2, Angiotensin converting enzyme 2; COMFA, Comparative molecular field analysis; COMSIA, Comparative molecular similarity indices analysis; COVID 19, Corona virus disease 2019; CPE, Cytopathic effect inhibition assay; CRFK cells, Confluent crandel feline kidney cells; CVB3 3Cpro, Coxsackievirus B 3 cysteine protease; Dabcyl-Dabcyl-Lys-Thr-Ser-Ala-Val-Leu-Gln-Ser-Gly-Phe-Arg-Lys-Met-GluEdans, KTSAVLOSGFRKME-Edans. [4-(4-dimethylaminophenylazo) benzoic acid]-KTSAVLQSGFRKME-[5-[2'-(aminoethyl)amino]-naphthalenesulfonic acid]; DTT, 1,4-Dithio-D,L-threitol; E protein, Envelope protein; FDA, Food and Drug Administration; FIPV, Feline infectious peritonitis virus; FRET analysis, Fluorescence resonance energy transfer analysis; HBTU, Hexafluorophosphate benzotriazole tetramethyluronium; M protein, Membrane protein; MD simulation, Molecular dynamics simulation; MERS CoV, Middle east respiratory syndrome corona virus; MM-GBSA, Molecular Mechanics/Generalized Born Surface Area; MM-PBSA, Molecular mechanics Poisson-Boltzmann surface area; Mpro, Main protease; MTT, 3-(4,5-Dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide; N protein, Nucleocapsid protein; NFkB, Nuclear factor kappa-light-chain-enhancer of activated B cells; NIH (MLPCN) screening, National Institutes of Health (Molecular libraries probe production centers network) screening; nsp10, non-structural protein 10; nsp12, nonstructural protein 12, RNA dependent RNA polymerase; nsp13, non-structural protein 13, helicase; nsp14, non-structural protein 14, N-terminal exoribonuclease and Cterminal guanine-N7 methyl transferase; nsp15, non-structural protein 15, uridylate-specific endoribonuclease; nsp16, non-structural protein 16, 2'-O-methyl transferase; NTD, N-terminal domain; ORFs, open reading frames; PC 3Cpro, Picornavirus 3 cysteine protease; PDB, Protein data bank; PHEIC, Public Health Emergency of International Concern; PLpro, papain-like protease; PP1a, Polyprotein1a; PP1ab, Polyprotein1b; gRT-PCR, Quantitative reverse transcription polymerase chain reaction; QSAR, Quantitative structure-activity relationship; RASPD, Rapid Screening with Physicochemical Descriptors; RdRp, RNA dependent RNA polymerase; S protein, Spike protein; SARS CoV, Severe acute respiratory syndrome corona virus; WHO, World Health Organisation; Z-RLRGG-AMC, Z-Arg-Leu-Arg-Gly-Gly-7amido-4-methylcoumarin.

heterocyclic scaffolds [1].

Coronaviruses (CoV) are a family of viruses capable of causing mild to severe symptoms of respiratory distress. In the last two decades, the outbreak of two of the coronaviruses, Severe Acute Respiratory Syndrome (SARS) and Middle East Respiratory Syndrome (MERS), have emerged as epidemics with severe mortality. Both epidemics were of zoonotic origin, with SARS CoV transmission from civet cats to humans in 2002 in China and MERS CoV transmission from dromedary camels to humans in 2012 in Saudi Arabia [2]. There was emergence of cluster of pneumonia cases of unknown etiology in Wuhan city, Hubei province, China on 31 December 2019 and later declared by China that the outbreak is associated with a seafood market in Wuhan. China shared the genetic sequence of novel coronavirus responsible for the outbreak for diagnostic purposes on 12 January 2020 [3].

On 30 January 2020, World Health Organisation (WHO) declared this 2019-nCoV outbreak as a PHEIC (Public Health Emergency of International Concern) which was declared pandemic on 11 March 2020 [4]. On 11 February 2020, WHO named this novel coronavirus as COVID-19 (corona virus disease 2019) and later International Committee on Taxonomy of viruses renamed it as SARS CoV-2 [5]. There were 24,066,076 confirmed cases of COVID-19 and 823,572 deaths, globally as on 26 August 2020 [6].

Coronaviruses are single stranded positive sense RNA viruses. COVID-19 is caused by seventh of known coronaviruses which have infected humans, in the sequence: 229E, NL63, OC43, KKU1, MERS-CoV, SARS-CoV, and 2019-nCoV-2. The latter is a betacoronavirus of subgenus sarbecovirus, and is a severe acute respiratory syndrome coronavirus 2 with 96% genome similarity to other bat coronaviruses [7,8]. Among other coronaviruses, virus causing COVID-19 has an advantage of the presence of a unique polybasic cleavage site leading to its increased pathogenicity [9]. It has the largest RNA genome (30 kb) among all other RNA viruses with six to ten open reading frames (ORFs). It consists of some structural and some non-structural proteins. The structural proteins include: spike (S protein), envelope (E protein), membrane (M protein), nucleocapsid (N protein) while the nonstructural proteins include: main protease (\tilde{M}^{pro}), papain-like protease (PL^{pro}), non-structural protein 13 (nsp13, helicase), non-structural protein 12 (nsp12, RNA dependent RNA polymerase), N-terminal exoribonuclease and C-terminal guanine-N7 methyl transferase (nsp14), uridylate-specific endoribonuclease (nsp15), 2'-O-methyl transferase (nsp16) and nsp10 [10]. The N protein consists of viral genome while the other three proteins, S, E and M, make the viral envelope. The processing of two viral replicase polyproteins produced by ORF1a/b of COVID-19, PP1a and PP1ab, leads to the production of sixteen nonstructural proteins while the mRNA encodes for the formation of structural proteins [11]. The spike protein is responsible for attachment to the host cell membrane using the host cell's angiotensin converting enzyme-2 receptor thus initiating the infection process. Upon entering the host cells, the viral genome undergoes translation into viral polyproteins. The viral 3CL^{pro} and PL^{pro} then cleave these translated proteins into effector proteins. PL^{pro} has the capability to deubiquinate host's NFkB and interferon factor 3, resulting in suppression of host cell immunity. The binding of SARS-CoV-2 and ACE-2 has been found to be 10-20 times greater than that of SARS-CoV and ACE-2, favouring the higher transmissibility of SARS-CoV-2 [12].

Like other betacoronaviruses, SARS-CoV and MERS-CoV, SARS-CoV-2 also attacks the lower respiratory system of the patient, release the nucleocapsid in host cellular machinery and further undergoes replication in host cytoplasm leading to viral pneumonia. It can also lead to multiple organ damage affecting heart, kidney, gastrointestinal tract, liver and central nervous system of the patient [4,12].

The RNA genome sequence of SARS CoV-2 (GenBank ID: MN908947.3) has shown to exhibit 82% similarity with SARS CoV (GenBank ID: NC_004718.3) and also it is often seen that the key target enzymes in coronaviruses reveal some sequence similarities, like RdRp of SARS CoV-2 shows 96% identity with SARS CoVRdRp, 3CLpro of

SARS CoV-2 revealed 96.08% and 87.00% similarity with that of SARS CoV and MERS CoV, respectively, and though only 83% sequence similarity is seen between PLpro of SARS CoV-2 and SARS CoV but the active site of both the proteins do not show much variation, therefore the heterocyclic scaffolds showing effectiveness against these targets in SARS CoV and MERS CoV might also be repurposed and modified for healthcare emergencies like SARS CoV-2 and other such outbreaks of coronaviruses in future [13–15].

As the heterocyclic compounds have been rigorously involved in the ailments including viral infections, AIDS, cancer, there exists a profound scope of exploring these multiple nuclei to curb coronaviruses. Therefore, through this review we have tried to summarise some of the treatment options based on the heterocyclic nuclei researched and developed against SARS CoV, MERS-CoV and SARS-CoV-2 epidemics using in vitro, in vivo and in silico approaches, which may be of immense value at this hour of global emergency and in future.

2. Isatin and indole-based derivatives

As some isatin based compounds have shown potent activity against 3C protease of rhinoviruses and the cysteine proteases of both rhinovirus and SARS CoV possessed structural similarity at the active site, some isatin derivatives were designed and assayed for inhibitory activity against SARS CoV 3CLpro by fluorescence resonance energy transfer FRET analysis. The assay revealed the higher potency of 5-iodo or 7bromo isatin derivatives with benzothiophenemethyl side chain rather than with benzyl or alkyl side chains. The IC₅₀ value of the most potent inhibitor 1 came to be 0.95 µM. The compound exhibited efficient binding within the active site of the enzyme. The carbonyl and amine groups of isatin scaffold were involved in forming hydrogen bond with Gly-143, Ser-144 and Cys-145 as shown by the docking analysis. The authors reported that the bulky nature of side chain on isatin derivatives decreased the inhibitory activity due to steric hinderance of the side chain with His-164 and Met-165 of 3CLpro [16]. After a year, Nsubstituted-5-carboxamide derivatives of isatin scaffold were designed and compared with 5-iodo substituted analogues using in vitro and computational approaches. The colorimetric inhibition assay results reported **2** as the potent inhibitor with lowest IC₅₀value of 0.37 μ M. The hydrophobic naphthyl group at N-1 of isatin moiety was found to be well fitted in hydrophobic pocket, thus increasing the activity. Also the C-5 carboxamide substitution imparted 3 to 4 times higher potential compared to the 5-iodo derivatives while the C-5 ester and carboxylic analogues did not display any inhibitory activity which was explained to be due to their poor binding at P1 site of 3CLpro as a result of strong charge repulsion from Glu-166. In addition, the carboxamide group successfully formed two hydrogen bonds with His 163 and Phe 140. The synthesized derivatives possessed noncovalent reversible binding with 3CLpro of SARS CoV [17]. Other group of scientists synthesized 5-sulfonamide derivatives of isatin and docked within SARS CoV 3CLpro active pocket with an improvement in inhibitory potential. The results of FRET inhibition assay suggested a better activity of 5-substituted analogues compared to substitutions at other positions of isatin scaffold. The 5-(piperazin-1-ylsulfonyl)isatin analogues displayed the higher potential than the 5-halogen substituted derivatives. Replacing the 5piperazinyl moiety with 5-piperidinyl substitution further enhanced the antiviral potency with an IC_{50} of <5 μ M. The effect of substitutions at N1 of isatin moiety revealed the most promising activity of compound 3 $(IC_{50} = 1.04 \,\mu\text{M})$ having an *N*-benzyl group which was further confirmed by docking analysis using Glide 5.5 software. Compound 3 extended hydrogen bonds with residues Gly143 and Cys145 of 3CLpro (PDB ID: 1UK4), with a dock score of 8.70. The docking results unleashed the crucial role of N-1 and C=O at position 2 of isatin nucleus in hydrogen bond formation. The 5-sulfonyl and N1-benzyl substituents fitted well into the S2 and S1 hydrophobic pockets of 3CLpro, respectively, thus improving the inhibitory potential of the derivatives [18]. Reports are also available with tripeptidic inhibitors of SARS CoV 3CL protease. The

dipeptide-type analogues with position-3 N-arylglycyl moiety were developed resulting in enhanced activity due to the hydrogen bond formation between amine of glycyl and Glu166 residue of 3CL pro. The fluorometric inhibition assay showed that the presence of DL-pyroglutamyl or pyrrole-2-carbonyl in place of N-(3-methoxyphenyl) glycyl moiety as rigid position-3 analogue, decreased the inhibitory potential dramatically while the indole-2-carbonyl group as rigid position-3 moiety remarkably improved the potential. Compound 4 with substitutions on position-3 indole nucleus with a 4-methoxy substituent was reported as the most potent derivative (IC₅₀ value = 0.74 μ M) when compared to 5- or 6-methoxy or 4-hydroxyl/4-isopropoxyl/4-isobutyloxyl derivatives. Replacing the rigid position-3 indole scaffold with other heterocycles like benzothiazole or benzofuran greatly reduced the activity suggesting the crucial role of amine of indole nucleus in hydrogen bond formation. The docking analysis of compound 4 ($K_i =$ 0.0063 µM) with SARS CoVMpro (PDB ID: 1WOF) highlighted a favourable conformation of the rigid P3 4-methoxyindole moiety in the active site, compared to the flexible N-(3-methoxyphenyl)glycyl unit providing a 65-fold greater potential to the indole derivative [19]. Two series of pyrazole and pyrimidine fused indole derivatives were designed and analysed for their antiviral activity against SARS CoV 3CLpro, biosterically replacing isatin with indole. The FRET inhibition assay results revealed compound 5 as the most potent derivative with an IC₅₀ value of 0.12lM. It was concluded that 2,3-dihydroinden-1-one was a required for anti-SARS CoV activity whereas it's clubbing with pyrimidine nucleus provided greater enhancement in potency compared to that with pyrazole moiety and at the same time, isoxazole fusion lead to compounds with lesser potency [20]. The SARS CoV-2 helicase was homology modelled using 2019-nCoV/USA-WA1-F6/2020 (Gen Bank: QHU79203.1) helicase amino acid sequence against SARS CoV helicase (PDB: 6jyt.2.A) as template due to a 99.78% sequence similarity between the two and the modelled protein was analysed for antiviral properties of 23 clinically approved antiviral drug candidates using MOE software. The docking study revealed the highest binding affinity of compound 6 with a dock score of -9.84 kcal/mol involving a hydrogen bond formation with Gly79 of SARS CoV-2 helicase thus enabling the compound to be an efficient inhibitor of SARS CoV-2 helicase, thus interfering with the viral replication potential of helicase [21]. The disruption of the trimerization of SARS CoV-2 spike protein which is reported to be involved in host cell membrane fusion, can lead to antiviral activity. A portion of spike protein trimer S2 domain (947-1027 residues) was found to be structurally similar to influenza virus H3N2 haemagglutinin by comparative analysis, to which Arbidol7 binds for its anti-influenza action. The binding mode and mechanism of action of compound7, against SARS CoV-2 spike protein trimer (PDB: 6VSB), was analysed using HADDOCK2.2 and SWISS-DOCK molecular docking servers. The results of docking analysis revealed the potential interaction of compound 7 with the SARS CoV-2 spike protein S2 domain residues (K776, K780, K947, E1017, R1019, S1021, N1023, L1024, T1027) in the same way as with H3N2 HA protein (PDB: 5T6N), thus interfering with SARS CoV-2 trimerization and suggesting a potential role of compound 7 in the treatment of SARS CoV-2 infection [22]. The homology modelling of SARS CoV-2 RdRp structure using SWISS-MODEL server revealed its 97.05% sequence similarity with the template, SARS CoVRdRp (PDB: 6NUR). The modelled structure was then docked against 74 antiviral drugs using Autodock Vina and the binding interactions were analysed by PyMol and Chimera. Delavirdine 8, an anti-HIV drug, exhibited pronounced inhibitory activity against SARS CoV-2 RdRp forming hydrogen bonds with Ala576 and Asn 582, with a dock score of -8.5, thus emphasizing the imperative role of 8 in SARS CoV-2 treatment therapy by interfering with RdRp dependant viral replication [23]. Novel inhibitors of SARS CoV-2 Mpro (PDB: 2H2Z) were designed by analyzing the active site, which basically comprise of S1', S1, S2 and S4 subunits. In the newly synthesized compounds, aldehyde was taken as a warhead required to form covalent bond with the thiol of cysteine which keeps the inhibitor anchored at S1' site of the protein. The FRET

inhibition assay displayed a high potential of compound 9 with an IC_{50} value of 0.053 µM with no cytotoxicity. The electron density map examination of the crystal structure of SARS CoV-2 Mpro (PDB: 6LZE) in complex with compound 9 revealed a favourable conformation of compound **9** in the active site, involving a hydrogen bond of its indole moiety with Glu166 and hydrophobic interactions with Pro168 and Gln189. In vivo pharmacokinetic study in mice with 5 mg/kg i.p and 5 mg/kg i.v administration, demonstrated 87.8% bioavailability value while in vivo toxicity analysis in SD rats and beagle dogs indicated no toxicity and mortality at a dose of 40 mg/kg. Therefore, the results of analysis suggested a promising role of compound 9 for SARS CoV-2 clinical trial studies [24]. Six anti-influenza drugs namely arbidol, oseltamivir, baloxavir, zanamivir, peramivir and laninamivir were evaluated for their antiviral potential against SARS CoV-2. The cell counting kit-8 (CCK-8) assay demonstrated arbidol7 as a potent inhibitor with an EC₅₀ value of 4.11 μ M while a CC₅₀ value of 31.79 μ M. A sharp decline in the SARS CoV-2 induced cytopathic effect and viral NP expression by compound 7 was also observed using immunofluorescence staining assay. The qRT-PCR examination displayed an efficient inhibition of viral replication stages with more pronounced effect at entry (41%) compared to post-entry stage (61%) using viral multiplicity of infection (MOI) of 0.05. Arbidol 7 lowered the binding efficiency of virions, in virus infected Vero E6 cells (MOI of 0.05) upto 67%. The result of viral intracellular trafficking analysis by immunofluorescence microscopy revealed a decrease in release of SARS CoV-2 from endo lysosomes. Since no neuraminidase analogue was found in SARS CoV-2, therefore the neuraminidase inhibitors tested in this study failed to show any result. Also, since the cap snatching effect of endonuclease of viral polymerases is absent in coronaviruses, as a result of this, baloxavir which acts by inhibiting this particular mechanism of endonuclease, also failed in SARS CoV-2 infection [25]. Fig. 1 outlines various indole and isatin derivatives.

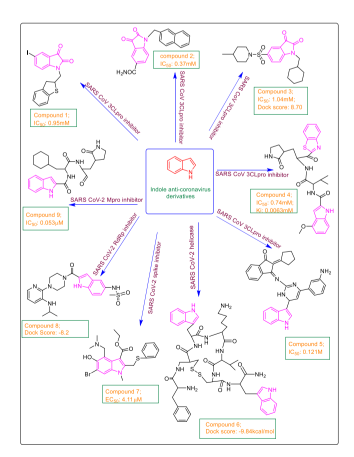


Fig. 1. Isatin and indole-based derivatives.

3. Quinoline and isoquinoline derivatives

In 2003, with the outbreak of SARS CoV epidemic, several drugs were repurposed against it. One such anti-malarial drug, chloroquine 10, was also evaluated for its antiviral potential against SARS CoV infection. The results of MTT assay in Vero E6 cells revealed a good potency with an IC₅₀ value of 8.8 μ M while a CC₅₀ value of 261.3 μ M, suggesting the use of chloroquine in SARS CoV like infections. The drug proved to be more effective in later stages of viral replication rather than effecting viral attachment or penetration [26]. Almost a year later, chloroquine 10 was further reported for its prophylactic and therapeutic use against SARS CoV activity by treating Vero E6 cells at a concentration of 0.1–10 μ M, both 20–24 h prior and 3–5 h after the viral infection. Immunofluorescence assay results indicated a 100% viral inhibition chloroquine pre-treated cells at 10 µM concentration while a 90-94% decrease in virus antigen-positive cells at 33-100 µM in post-treated cells with an ED_{50} of 4.4 $\mu M.$ The mechanism of action for antiviral action was depicted as the abrogation of terminal glycosylation of ACE-2 receptor thus interfering with SARS CoV spike-ACE-2 receptor binding necessary for viral entry in host cells by flow cytometry and immunoprecipitation analysis. As chloroquine is basic in nature, the inhibition of virus-endosome fusion due to increase in endosomal pH was presumed to be responsible for post-treatment anti-SARS CoV effects [27]. Keeping in view the antimalarial potential of ferroquine, even on chloroquine resistant strains and the antiviral efficacy of chloroquine was well established against SARS CoV, some ferroquine derivatives have been reported for their antimalarial and antiviral potential. The analysis results revealed a high anti-malarial potency of compound 11, a hydroxyferroquine analogue, with high lipophilicity and lesser toxicity than ferroquine. From the in vitro study results of ferroquine showing the good potency of its metabolites against chloroquine resistant Plasmodium falciparum strain, it was suggested that the de-alkylated metabolite of the metallocene 11 namely mono-N-desethyl-ferroquine, facilitated the activity of parent molecule. The compound 11 exhibited good anti-HIV and anti-SARS CoV activity with an EC_{50} value of 3.6 μ M in SARS CoV infected Vero cells, compared to the standard chloroquine showing an EC₅₀ value of 6.5 µM though it exhibited poor potential against other viruses like reovirus-1 and HSV-2, herpes simplex virus-1, sindbis virus, influenza A virus, vesicular stomatitis virus and vaccinia virus, thus revealing its high selectivity towards HIV and SARS CoV infections [28]. From the virtual screening of over six lakh compounds in a database, a quinolinone derivative was selected for its promising anti-SARS CoV 3CLpro activity, for which the in vitro SARS CoV inhibition analysis revealed an IC50 value of 0.44 µM/L. Therefore, taking it as lead, based on its structure and activity, 23 novel 4-quinolinone ester derivatives were synthesized and in vitro and in silico studies were conducted to evaluate their SARS CoV 3CLpro inhibition potential. Compound 12 with a methyl substitution at NH moiety of the quinolinone lead compound, demonstrated nearly 12-fold greater anti-SARS CoV 3CLpro activity using FRET analysis, with an IC₅₀ of 36.86 nM/L in comparison to lead. The docking analysis with SARS CoV 3CLpro (PDB: 3SN8) via Discovery Studio 3.0, indicated a more favourable binding of 12 in the active site of SARS CoV 3CLpro compared to the lead molecule which is proposed to be due to a hydrogen bond of amide oxygen with His 163 which remains conserved in both molecules and hydrogen bonds of ester carbonyl oxygen of 12 with Cys 145 and His 41 (the catalytic enzymes in SARS CoV 3CLpro active site necessary for proteolytic function). The structure activity relationship study of synthesized derivatives displayed a decrease in anti SARS CoV potential with the replacement of thiophene moiety of lead molecule with benzene, substituted benzene or alkyl carbon chains which is considered to be due to large size of these substituents which failed to fit within small S1' pocket of the protease [29]. With the outbreak of MERS CoV infection in 2012, some FDA approved drugs which have shown activity against other coronaviruses, were repurposed against MERS CoV infection also. One such drug namely chloroquine 10 from a database of 348 FDA approved compounds, was

also investigated for its in vitro potential against MERS CoV infected Vero and Huh7 cells using a colorimetric cell viability assay. The results unleashed the inhibition of MERS CoV mediated cytopathic effect by chloroquine with an EC₅₀ value of 3 µM while it exhibited an EC₅₀ value of 4.1 μ M in SARS CoV infected Vero E6 cells. To determine the pre- and post- treatment effects of chloroquine, the time-of-addition experiment using plaque assay was conducted by adding the drug both 1 h prior and 1 h post viral infection (multiplicity of infection, 1) which revealed an approximate decrease of 2-log in viral production in chloroquine pretreated Vero cells with no effect in post-treated cells, thus suggesting an early stage inhibition of MERS CoV infection via inhibition of clathrin mediated endocytosis [30]. The catalytic site of SARS CoV at Arg188/ Gln189, was reported to be degradation sensitive while its mutant strain with isoleucine residue at 188 in place of arginine renders the protease highly stable and catalytically more efficient. At the same time, a peptide-mimetic substrate-based inhibitor with an aldehyde group at Cterminal demonstrated a potent competitive inhibition of the mutant R188I SARS CoV 3CLpro strain.

Novel decahydroisoquinoline based non-peptide derivatives were designed taking the above peptide mimetic inhibitor as lead and evaluated for their anti-R188I SARS CoV 3CLpro activity. The compound 13, a (4aR,8aS) isomer of trans-decahydroquinoline diastereomers with Np-bromo-benzoyl substitution, displayed a high potential with an IC₅₀ value of 63 µM. The X-ray crystallographic study of compound 13 in complex with 3CLpro (PDB: 4TWW), revealed a compact fitting of compound 13 within the active site of SARS CoV 3CLpro comparable to the active lead molecule. The S2 pocket was largely covered with decahydroisoquinoline fused ring while the hydrogen bond between the N atom of imidazole ring of compound 13 and His163 of 3CLpro provided it a favourable conformation at the S1 site of protease. Also, the N-pbromobenzoyl moiety lies outward from 3CLpro where hydrophobic interactions might be possible [31]. From a dataset of FDA approved drugs, a set of 1528 compounds were screened and evaluated for their anti-SARS CoV-2 activity by combining cell viability assay and SARS CoV-2 ELISA. The Cell Titer Glo assay and cytopathic effect (CPE) inhibition assay (MOI: 0.01) in Vero E6 cells revealed promising effects of compound diiodohydroxyquinoline 14, (a quinoline derivative) used as a luminal amebicide, with a CC $_{50}$ value of ${>}100~\mu\text{M}$ and approximately 70% CPE inhibition, respectively. Compound 14 also showed an EC_{90} value of 4.50 µM in viral load assay using MOI of 0.01 in Vero E6 cells while an EC₅₀ value of 1.38 µM through plaque reduction assay. It was also observed that treatment with compound 14 greatly reduced the SARS CoV-2 N antigen expression at a non-toxic concentration of <10 μ M just like remdesivir which was taken as reference. The replication of SARS CoV-2 was also inhibited effectively at post entry. The in vivo studies revealed that compound 14 could be used as a potent luminal antiviral agent [32]. As it has been already reported that apart from using ACE-2 receptor, coronaviruses use sialic acid-containing glycoproteins and glycosides also as a source for entry into the host cell membrane and also that MERS CoV entry can be inhibited by depleting these sialic acids content. The efficiency of chloroquine 10 and hydroxychloroquine 15 against SARS CoV-2 was studied in silico (Hyperchem and Molegrow molecular viewer) using the mechanism of depletion of cell surface sialic acids containing gangliosides. Merging chloroquine with Neu5Ac depicted a good fit between the two, through interaction between negatively charged carboxylate of Neu5Ac and a cationic charge of chloroquine. The molecular modelling of 9-O-SIA (9-O-acetyl-N-acetylneuraminic acid), the preferable sialic acid for interaction with coronaviruses, also demonstrated a favourable fit among the two via interaction of nitrogen containing cationic ring of chloroquine with carboxylate of 9-O-SIA (interaction energy: -47 kJ/mol). Compound 15 showed a similar interaction with 9-O-SIA with enhanced binding owing to a hydrogen bond formation (interaction energy: -46 kJ/mol). The interaction affinities of the two drugs with ganglioside sialic acids, as sialic acid mostly forms a part of human respiratory tract gangliosides and glycoproteins was studied through molecular

modelling using ganglioside GM1. The results revealed good interaction of both the drugs within the two drug-binding sites in GM1, showing a high interaction energy of -108 kJ/mol and -120 kJ/mol, respectively. Finally to determine the ability of both the drugs to prevent the binding of SARS CoV-2 spike NTD with cell surface ganglioside, the NTD-GM1 complex was superposed with drug-GM1 complex, which indicated that both NTD and the drug binds at the same position in GM1 and with the same mechanism involving a hydrogen bond and a CH π interaction, thus unleashing the anti-SARS CoV-2 potential of the drugs [33]. Some active quinoline and isoquinoline derivatives are shown Fig. 2.

4. Pyridine derivatives

A library of 50,000 compounds was screened for their activity against SARS CoV 3CLpro using FRET analysis. The non-specific compounds were further screened by examining their activity against SARS CoV 3CLpro in the presence of BSA (bovine serum albumin) and out of the screened compounds, 69 compounds show specificity against SARS CoV 3CLpro. All of these screened compounds showed potential electrophilic centres like amides, nitriles which may be capable of forming covalent bond with the nucleophilic thiol of Cys 145 at the active site of 3CLpro. Therefore, they were further analysed in the presence of DTT (1,4-dithio-D,L-threitol) for their specificity against SARS CoV 3CLpro which depicted 5 inhibitors whose activities were least affected by DTT. Finally, the selected compounds were evaluated against other coronaviruses also namely hepatitis A virus 3Cpro, chymotrypsin, hepatitis C NS3pro and papain, to test their selectivity for SARS CoV 3CLpro. Compound 16 displayed the highest selectivity and potency against SARS CoV 3CLpro with an IC_{50} value of 0.5 μ M, thus emphasizing on the role of 5-chloropyridine derivatives as anti-SARS CoV agents [34]. The derivatives were evaluated for their activities against SARS CoV (Frankfurt-1 strain) and FIPV (feline infectious peritonitis virus) infections by MTT assay in Vero E6 cells and CRFK (confluent crandel feline kidney) cells infected with 100 CCID_{50} (50% cell culture infective dose) SARS CoV and FIPV, respectively. The highest potency was revealed by compound 17, with an EC50 of 17 mg/L against SARS CoV and with almost no cytotoxic effect in Vero cells (MIC > 100 mg/L). The compound showed post entry inhibition of viral infection in time of addition experiment in FIPV infected CRFK cells. The structure activity relationship study revealed that the sulphide and sulphoxide analogues displayed poor activity against both viruses. The reduced pyridine-N-

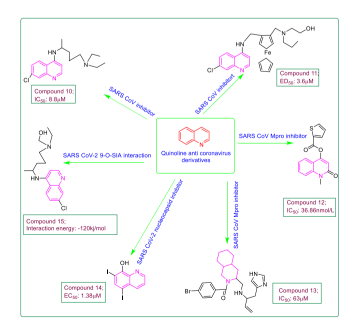


Fig. 2. Some of the promising Quinoline and Isoquinoline derivatives.

oxide analogues exhibited a complete loss of antiviral potential against both SARS CoV and FIPV, at their subtoxic concentrations, thus emphasizing on the imperative role of *N*-oxide in pyridine nucleus [35]. The 5-chloropyridinyl indolecarboxylate analogues were synthesized as anti-SARS CoV evaluated for their in vitro and in silico anti-SARS CoV 3CLpro inhibitory activity [36]. The FRET inhibition assay demonstrated compound 18, as the most active analogue having a 4-carboxylate indole substitution, with an IC_{50} value 0.03 μ M. The analysis results indicated a decrease in potential with the shift of carboxylate moiety on positions other than 4 in indole nucleus. The acylation of indole N also resulted in loss of anti-viral potency. In silico study was also conducted by docking of compound 18 within the binding pocket of SARS CoV 3CLpro (PDB: 2HOB) using GOLD 3.2 program which further confirmed its promising role as an anti-SARS CoV agent. The results of docking analysis revealed a good fit of compound 18 within the active site of protease due to the formation of three hydrogen bonds by carbonyl oxygen with Cys 145, Ser 144 and Gly 143 while the indole moiety fits into the hydrophobic S2 site involving the interaction of imidazole N with His 41 [37]. Two series of derivatives of 5 chloropyridine were synthesized using MAC-5576 16 (a potent anti-SARS CoV agent) as lead. In vitro SARS CoVMpro inhibition assay showed the derivatives of series 1 (derivatized benzene substituted furan fused to chloropyridyl ester) to be possessing higher inhibitory potential than those of series 2 (derivatized, six membered cyclic aromatic moiety clubbed with chloropyridyl ester) with compound 19 as the most promising inhibitor exhibiting an IC₅₀ value of 60 nM. The analysis results revealed a decrease in anti-SARS CoV potential with a change in the position of nitro group in compound 19 from para position to ortho or meta position of benzene ring. The molecular docking study using AutoDock depicted the binding of chloropyridine group within the S1 pocket of SARS CoV 3CLpro, with N of pyridine forming a hydrogen bond with N of His 163 while the furan moiety fits into the space between S1 and S2 subsites involving van der Waals interaction with Met 165. The *p*-substituted benzene moiety (Δ G: -9.34 kcal/mol) fitted well into the S2 pocket while the o- or m-substituted benzene rings in other derivatives were forced to move outside from S2 pocket due to steric hinderance with SARS CoVMpro residues, His 41 and Met 49 [38]. Fig. 3. shows various pyridine derivatives.

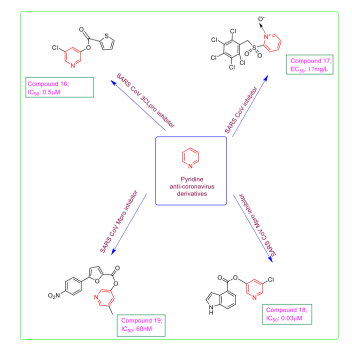


Fig. 3. Potential Pyridine derivatives.

5. Purine and pyrimidine derivatives

The Maybridge compound database, was virtually screened for a set of anti-SARSCoV 3CLpro (PDB: 1UK4) inhibitors within the binding site containing the catalytic dyad Cys 145 and His 41 which forms a part of subsites S1, S2 and S3, using DOCK 4.0.2 server. The top 93 compounds involving more than two hydrogen bonds with Mpro, were further analysed by SARS CoVMpro inhibition assay, among which 21 derivatives were found active exhibiting an IC₅₀ value of $<30 \,\mu$ M. Three of these 21 compounds shared a similar core structure, N-phenyl-2-(2pyrimidinylthio)acetamide, using which 25 more structural analogues were screened from Maybridge, SPECS_SC and ChemBridge databases and a total of 28 analogues were analysed for anti-SARS CoV inhibitory potential which displayed compound **20** as the most promising analogue with an IC_{50} value of 3 $\mu M.$ The results furnished from 3D-QSAR modelling using COMFA and COMSIA analysis completely matched with those from experimental analysis. The MM/GBSA analysis results also revealed the best fit of compound 20 within the binding pocket of SARS CoV 3CLpro with a binding free energy of -23.17 kcal/mol, due to the strong interactions formed by its thiazole and benzene moieties with protease residues namely, Gln 192, Leu 167, Glu 166 and Pro 168 [39]. 6-Chloropurine derivatives have been reported to exhibit potent antiviral activity against many viruses. Nucleoside analogues based on 6chloropurine moiety were designed and analysed for their activity against SARS CoV using plaque reduction assay in SARS CoV Frankfurt-1 strain infected Vero E6 cells. Compound 21 exhibited promising activity with an IC₅₀ value of 48.7 μ M and a decrease in virus yield to less than one-hundredth of the control, at 20 µM concentration. The structure activity relationship study showed the importance of chlorine atom at position-6 of purine nucleus and a loss of anti-SARS CoV activity with the 2-amino group substitution in 6-chloropurine moiety. The replacement of chlorine group with weaker leaving groups like --SMe or -OMe, led to a decrease in anti-SARS CoV potential which is attributed to be due to lack of formation of an irreversible covalent bond which 6chloropurine can form with the active site of SARS CoV owing to its electrophilic nature. The unprotected 5'-hydroxyl substitution in ribofuranosyl moiety is also imperative as it gets converted to the active triphosphate form leading to the antiviral activity. Also, the replacement of ribofuranosyl moiety with 2'-deoxy- or 3'-deoxyribonucleoside analogues led to a detrimental effect on the anti-SARS CoV potential of compound **21** [40]. From the compound library, Genesis plus collection, 960 compounds were screened against SARS CoVPLpro using in vitro and in silico approaches. The results of deubiquitination assay demonstrated that out of the screened compounds, only two derivatives, 6-Mercaptopurine 22, (IC₅₀ value: 21.60 µM) and 6-Thioguanine 23 (IC₅₀ value 5 µM) exhibited pronounced SARS CoVPLpro inhibition. The thiocarbonyl group of 22 and 23 was found to be crucial for SARS CoVPLpro inhibition using structure function relationship study by ISIS, as replacing it with hydroxyl or methylthio moiety led to loss of inhibition. The docking analysis confirmed the inhibition inhibition mechanism within the active sites of SARS CoVPLpro (PDB:2FE8) and 3CLpro (PDB: 1UK2) using DS modelling 1.7 program. The docking result suggested a good fit of both the compounds with a dock score for 22 (23.9) and 23 (24.4) within the active site of PLpro with the probability of formation of a hydrogen bond between sulphur atom of the compounds and Cys 1651 residue of SARS CoVPLpro resulting in reversible competitive inhibition. While the dock scores with SARS CoV 3CLpro were found to be 17.8 for 22 and 18.4 for 23, much lower than with SARS COV PLpro, thus emphasizing on the selectivity of both the compounds towards PLpro [41]. Another series of pyrimidine derivatives was synthesized and evaluated for their SARS CoV 3CLpro inhibitory potential. The FRET analysis using Dabcyl-KTSAVLQSGFRKME-Edans as fluoregenic substrate demonstrated compound 24 to be endowed with marked SARS CoV 3CLpro inhibitory potential exhibiting an IC₅₀ value of 6.1 µM, with no cytotoxicity as depicted by MTT assay. The molecular docking analysis of compound 24 with SARS CoV 3CLpro (PDB: 1UK4) using

Discovery studio modelling 1.2 SBD program unleashed a good fit of phenylnitro group within the S1 pocket with O atom of nitro group forming a hydrogen bond with Cys 145 and Gly 143 at the binding site of 3CLpro. The lack of nitro group led to a loss of inhibitory potential, emphasizing on the imperative role of nitro substituent. Moreover, electron withdrawing groups like chloro in phenyl ring favoured the inhibitory potential more than the electron releasing groups like methoxy or methyl [42]. The antineoplastic drug carmofur 25, a pyrimidine analogue was analysed for SARS CoV-2 Mpro inhibitory potential using the X-ray crystallographic study. The electron density map of the 25-SARS CoV-2 Mpro complex indicated a favourable conformation of the drug within the active site of protease with its fatty acid moiety forming a covalent bond with Cys 145 of the catalytic dyad leading to the release of 5-fluorouracil. The carbonyl O of 25 was observed to be involved in hydrogen bond formation with Cys 145 and Gly 143 while the fatty acid tail showed hydrophobic interaction with His 41, Met 165 and Met 49 at the S2 subsite of the protease. The in vitro inhibition assay in SARS CoV-2 infected Vero E6 cells revealed an EC₅₀ value of 24.30 µM while the cytotoxicity study indicated the selectivity of compound 25 with a CC_{50} value of 133.4 μ M [43]. A set of 6799 compounds from Pubchem and Asinex library were virtually screened and analysed by molecular docking against SARS CoV-2 nucleocapsid RNA binding domain (PDB: 6VYO). The high throughput virtual screening with docking was done using Schrodinger's molecular docking module at the active sites 1, 2 and 3 of the protein. From the screened ligands, compound zidovudine 26, a thymidine analogue, demonstrated the highest potential with a dock score of -9.75 while a binding free energy of -59.43 kcal/mol at site 3, as depicted by MM-GBSA approach. The molecular dynamic simulation study also revealed a stable interaction of 26-SARS CoV-2 N protein complex, thus suggesting the exploration of this potential nucleus as anti-SARS CoV-2 agent [44]. A series of diaryl pyrimidine derivatives were evaluated for their SARS CoV-2 inhibitory potential using in silico approaches. The molecular docking analysis of the derivatives with the SARS CoV-2 spike glycoprotein-human ACE-2 complex (PDB: 6VW1) was carried out using AutoDock Vina while the results were analysed by MGL tools and PyMol. The pyrimidine derivative 27 showed best binding affinity at the interface of the complex with a binding energy of -8.95 kcal/mol which was attributed to be due to formation of a hydrogen bond with ASP 350, Arg 393 and Asn 394 residues and hydrophobic interactions of its naphthyl and phenyl moieties with Phe 40, Trp 69, Leu 73, Phe 390 and Leu 391 amino acid residues. The molecular dynamic simulation study with GROMACS v5.1.4 biomolecular simulation package demonstrated a stable conformation of 27 within the spike glucoprotein-ACE2 complex throughout the simulation. Its high binding affinity of with the complex was further confirmed by binding free energy calculation using MM-PBSA, the results of which came out to be lowest (Δ G: -30.89 kcal/ mol), thus suggesting further evaluation of diaryl pyrimidines as effective anti-SARS CoV-2 agents [45]. Fig. 4 highlights some of the purine and pyrimidine derivatives active against coronaviruses.

6. Pyrazole derivatives

Pyrazolones have been exploited for activity against viruses. A set of 6800 compounds from Korea chemical bank database underwent high throughput screening to obtain the most potent compound against SARS CoV 3CLpro. After primary screening using Dabcyl-KTSAVLQSGFRKME-Edans as fluoregenic substrate, the compounds that showed > 50% inhibition of protease activity at 50 μ M further undergo secondary screening at 10 μ M. The inhibition assay revealed compound **28** as the most potent inhibitor of SARS CoV 3CLpro with an IC₅₀ value of 2.5 μ M. The docking analysis of compound **28** within the active site of SARS CoV 3CLpro (PDB: 1UK4) using Discovery studio modelling 1.2 SBD program, depicted a good fit of the molecule with its 4,5-dihydro-1H-pyrazole moiety occupying S1' and S2 sites and remaining part resting at S3 site of the protease [46]. Another series of pyrazolone analogues were

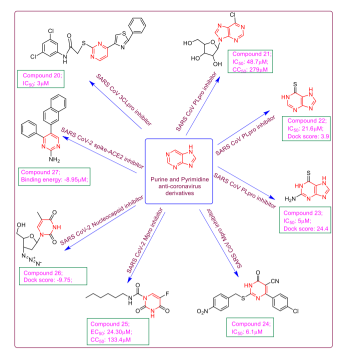


Fig. 4. Purine and Pyrimidine derivatives acting on coronaviruses.

reported for the antiviral potential against SARS CoV 3CLpro and CVB3 3Cpro. As a result of in vitro FRET analysis, compound 29, with a pbenzylidene aryl ring substitution at C4 of pyrazolone moiety, was observed to be endowed with marked activity against both 3CLpro and 3Cpro with IC_{50} values of 8.4 μ M and 9.6 μ M, respectively. The MTT cytotoxicity assay results revealed no cytotoxicity of the target compounds at 200 µM. To further confirm the inhibitory potential of the target compounds, molecular docking analysis via Discovery studio modelling 1.2 SBD program of Accelrys, of the derivatives within the binding site of SARS CoV 3CLpro (PDB: 1UK4) was performed which unleashed a favourable conformation of compound 29 with its N1 phenyl substituent fitted well in S1' pocket while the O of nitro group formed a hydrogen bond with Gly 143. The carbonyl oxygen of pyrazolone moiety was involves in H-bond formation with Glu 166. The phenyl ring at C3 of pyrazolone fitted well in hydrophobic S2 site while the p-carboxybenzylidene substituent at C4 of pyrazolone nucleus fitted easily in S3 pocket with carboxyl oxygen forming a hydrogen bond to Gln 192. The in vitro and in silico study results emphasized on the imperative role of carboxyl group in benzylidene ring as pyrazolone derivatives lacking this substituent lost their inhibitory potential. It was found that the presence of electron withdrawing substituents like nitro, cyano or fluoro at N1 phenyl ring led to an increase in inhibitory potential [47]. Some 5-pyrazolone derivatives were designed and evaluated for their inhibitory potential against SARS CoV and MERS CoV 3CLpro. The in vitro fluorometric analysis using Dabcyl-KTSAVLQSGFRKME-Edans as peptide substrate for 50 nM SARS CoV 3CLpro or 300 nM MERS CoV 3CLpro, unleashed the highest potential of compound **30** with IC₅₀ values of 5.8 μ M and 7.4 μ M, respectively. The structure activity relationship study proved that the replacement of bulky phenyl substituent from position 3 of pyrazolone moiety with smaller groups like methyl or CF₃ will led to loss of inhibitory potential. Also, the removal of carboxylate group from compound **30** resulted in derivatives with no inhibitory activity. The substitution with lipophilic group at p-position of phenyl ring at N1 of pyrazolone also led to a marked increase in activity. The in vitro assay results were further confirmed by docking analysis of the ligand within the active site of SARS CoV 3CLpro (PDB: 2ALV) using iGemdock v2.1 program. The docking results revealed a good fit of compound 30 within the active site

with carboxylate moiety resting in S1 site forming hydrogen bonds to Gly 143, Ser 144 and Cys 145 while furan moiety interacting with hydrophobic S1' site. The lipophilic tert-butyl group of compound 30 showed good interaction with hydrophobic S2 site, thus enhancing its inhibitory potential. Also, the carbonyl group of pyrazolone moiety is involved in hydrogen bond formation with His 41, thus destabilizing the catalytic dyad at the active site of SARS CoV 3CLpro [48]. The ZINC database was screened using RASPD web server with the aim of searching for potent SARS CoV-2 main protease inhibitors. The best two hit molecules selected as a result of RASPD score, were screened for their drug likeness using SwissADME and Molinspiration servers which revealed a good bioactive score and pharmacokinetics of both the ligands. Therefore, these two ligands were analysed for their binding affinities with SARS CoV-2 main protease (PDB: 6LU7) using ParDOCK server. The docking study results showed a good fit of both the ligands within the active site of protease with a higher binding affinity of compound 31 (Binding energy: -6.20 kcal/mol) comparable to ligand N3 (Binding energy: -6.43 kcal/mol), involving π -alkyl interactions with histidine residue of main protease [49]. Fig. 5 depicts pyrazole derivatives.

7. Thiazole derivatives

Thiazoles and their derivatives have been explored for activity against corona viruses. A series of trifluoromethyl ketone peptide derivatives were designed and evaluated for their activity against SARS CoV 3CLpro. The inhibition assay results revealed low inhibition potential of these derivatives with a minimum K_i value of 21 μ M. Regnier *et al.* designed another series of derivatives by replacing the trifluoromethyl ketone group with electron withdrawing thiazolyl or benzothiazolyl ketone moieties, with the objective of improving the covalent bond formation with Cys 145 of catalytic dyad at the active site of 3CLpro. The inhibition assay results proved the highest potency of compound **32**, a thiazolyl ketone peptide derivative, possessing a K_i value of 2.2 μ M. This inhibitory potential was further confirmed by computational docking study of **32** with SARS CoV 3CLpro (PDB: 1WOF) using MOE 2007.09 modelling package which demonstrated a good fit of ligand within the active site of protease with N of thiazole forming a

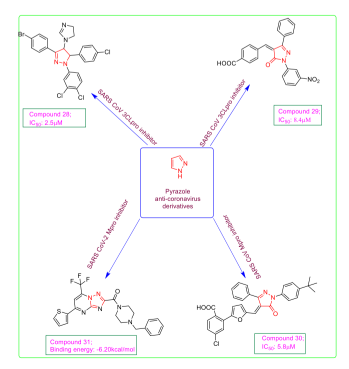


Fig. 5. Pyrazole derivatives acting as inhibitors of SARS CoV or SARS-CoV-2.

hydrogen bond to His 41 while the benzyloxycarbonyl and thiazolyl ketone groups involved in interaction with Cys 145 of the catalytic dyad [50]. The 5-arylidene-4-thiazolidinones were considered less selective towards biological targets because of the high reactivity of exocyclic double bond with the nucleophilic protein residues favouring the Michael addition reaction. Therefore, isosteric thiopyrano[2,3-d][1,3] thiazole derivatives were synthesized taking 5-substituted-4-thioxo-2thiazolidinone as precursors. Compound 33 displayed the highest anticancer potential with GI_{50} value of 0.309 μ M against MCF-7 breast cancer cell lines while it demonstrated a moderate potency against SARS CoV with IC_{50} value of 23 μM (visual assessment) and IC_{50} value of 14 µM (neutral red dye assessment) [51]. Konno et al. took compound 32 as lead and carried out the molecular modelling with 3CLpro (PDB: 1WOF) and found a vacant space in S1' pocket carrying the thiazolyl group of compound 32 and a protruding benzyloxycarbonyl (P4) moiety from the active site while cyclic amide (P1) and electron withdrawing thiazolyl (P1') moieties were observed to be crucial for inhibitory activity. The authors carried out optimization of P1' and P4 moieties to get more efficient anti-SARS CoV 3CLpro inhibitors. The fluorescence-based peptide cleavage assay results of the newly optimized derivatives using Dabcyl-KTSAVLOSGFRKME-Edans as fluorogenic substrate, indicated the pronounced inhibition effect of compound 34 (against SARS CoV 3CLpro possessing K_i value of 0.003 μ M. The authors reported that a benzothiazolyl moiety led to enhancement of activity owing to its good fit in the large S1' pocket. Also the presence of 4-N,N'-dimethylaminophenoxy acetyl moiety resulted in favourable hydrophobic interactions with Ala 191 at S4 pocket, with a unique folding conformation for improved anti-SARS CoV potential [52]. As 4-thiazolidinone and pyrazoline analogues exhibit potent antiviral activity, 2-pyrazoline-4-thiazolidinone hybrid derivatives were designed and evaluated in vitro for their anticancer and antiviral potential. Compound 35 emerged as the most potent anticancer and antiviral agent based on AACF (antimicrobial acquisition coordinating facility) programme, against Tacaribe TRVL 11,753 strain (EC_{50}: 0.71 $\mu\text{g/mL})$ but showed mild activity towards SARS CoV urbani strain (EC₅₀: 49 µg/mL) [53]. The same group (Havrylyuk et al., 2013) synthesized, another series of 5-pyrazoline conjugated 4-thiazolidinones analogues to get more active anticancer and antiviral agents. The evaluation of antiviral potency of compound 36 exhibited an EC_{50} value of 21.46 μ M and a CC₅₀ value of 34.67 μ M against SARS CoV urbani strain [54]. A set of 5-ylidene-4-thiazolidinone-3-carboxylic acid derivatives were synthesized and found to have low or moderate inhibitory potential of the target compounds where compound **37** displayed marked anti-SARS CoV activity with an EC₅₀ value of 27 µM and a selectivity index of >3.7 against SARS CoV urbani strain in Vero 76 cells [55]. A commercial database was screened to identify the novel inhibitors of SARS CoVMpro. Using the 3D structure of transmissible gastroenteritis Mpro (PDB: 1LVO) as template, the structure of SARS CoVMpro was simulated and its active site predicted owing to the sequence similarity between the two proteases. The molecular docking analysis result revealed the analogues bearing the core nucleus as compound 38 as inhibitors of SARS CoVMpro had the potential to inhibit Mpro infected Vero-E6 cells [56]. Some of such derivatives are depicted in Fig. 6.

8. Triazole derivatives

Triazoles have been reported in literature to possess remarkable antimicrobial activity. A series of benzotriazole esters was prepared by condensing (HBTU Hexafluorophosphate benzotriazole tetramethyluronium) with carboxylic acids and the target compounds were evaluated for their potential against SARS CoV 3CLpro using *in vitro* and *in silico* studies. The *in vitro* fluorometric assay revealed compound **39** to be highly active anti-viral agent (K_i value: 7.5 nM; CC₅₀: >100 μ M) with an irreversible inhibition of SARS CoV 3CLpro. The *in silico* molecular docking exposed a favourable conformation of compound **39** within the active site of SARS CoV3CLpro (PDB: 1uk4) with benzotriazole scaffold

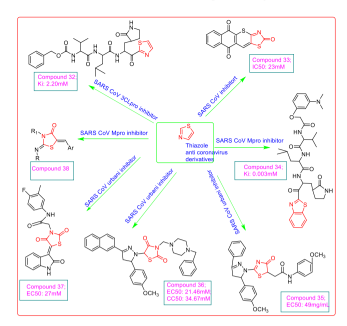


Fig. 6. Thiazole derivatives active against coronaviruses.

resting in pocket containing Gly 143, Ser 144 and Cys 145, facilitating a nucleophilic attack by the Cys 145 to the carbonyl group in benzotriazole ester. Also, the hydrogen bond formation between indole of 39 and -OH group of Thr 25 further stabilized the complex formation with SARS CoV 3CLpro. The benzotriazole ester derivatives exhibited a good inhibition of SARS CoV 3CLpro via acylation of Cys 145 in the catalytic dyad at active site [36]. Triazole based non-covalent inhibitors of SARS CoV 3CLpro were developed and analysed for their anti-SARS CoV 3CLpro inhibitory activity. Compound 40, a biaryl substituted triazole derivative possessed a ligand efficiency of >0.3 which on analysis demonstrated the highest potency with an IC_{50} value of 0.051 μM [57]. Some 1,5-disubstituted tetrazole-1,2,3-triazole conjugates were screened via docking analysis against SARS CoV-2 main protease (PDB:6LU7) possessing the favourable interactions similar to the cocrystalized ligand with the active site catalytic triad, Gly 143, Ser 144 and Cys 145 residues involving the hydrogen bonds and hydrophobic interactions. Ten 1,5-disubstituted tetrazole-1,2,3-triazole hybrids as SARS CoV-2 Mpro inhibitors were generated. Among the proposed hybrids, compound 41 having an isatin moiety exhibited the highest interaction energy (E: -255.79 kcal/mol) within the active site of 6LU7 involving hydrogen bond formation of 1,2,3 triazole moiety with Ser 144 and Cys 145 and that of tetrazole moiety with Ser 1 and Asn 142 and also electrostatic interactions with His 41, His 172, Glu 166 and His 163 while the higher number of rings in its structure enhance the hydrophobic interactions [58]. Fig. 7 depicts some triazole derivatives active against corona viruses.

9. Miscellaneous heterocycles

Keeping in view the anti-SARS CoV activity of aryl diketo acids, a series of bioisosteric dihydroxy chromones were designed and examined for their SARS COV inhibitory potential against ATPase and helicase using phosphate release assay method and FRET-based analysis, respectively. The *in vitro* assay results revealed a good inhibitory potential of compound **42**, a flavonol analogue with free catechol group, with IC₅₀ values of 25.4 μ M and 2.7 μ M against ATPase and helicase, respectively which proposed the involvement of two binding sites in Mpro i.e. one for hydrophobic interaction with arylmethyl moiety while other for hydrogen bond interaction with free catechol moiety [59]. Kim *et al.* developed a series of 2,6-bis-arylmethyloxy-5-hydroxychromones as anti-SARS CoV agents. The compounds displayed dual inhibitory

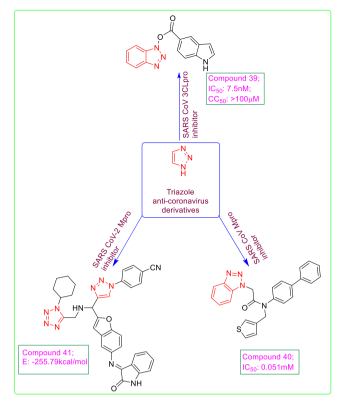


Fig. 7. Some active triazole anti-coronavirus derivatives.

activities against both nucleoside triphosphatases and helicases of SARS CoV. The compound 43 was found to be endowed with highest potential against both HCV (EC50: 4 µM) and SARS CoV (IC50: 4 µM for ATPase and 11 µM for helicase) with no cytotoxicity in HS27 (human normal fibroblast) cells ($CC_{50} > 50 \mu M$). According to structure activity relationship study, 3-iodo- or chloro-substituted benzyloxy moiety on 5-hydroxy chromone scaffold played a major role whereas derivatives with substitution at position 4 of benzyloxy ring displayed lower inhibition potential [60]. As nitric oxide (NO) derivatives have shown good antiviral effect against SARS CoV, hence phenyl furoxan derivatives, as NO donors, were evaluated by in silico approaches to test their potency against SARS CoV. The molecular docking analysis of some compounds with SARS CoVMpro (PDB: 6 W63) disclosed the best binding pose of compound 44 with binding affinity of -9.8 kcal/mol and dock score of -90.91. The docked conformation revealed hydrogen bond formation of 44 with Cys 145 and Ser 144 residues of the catalytic triad and also aromatic interactions involving His 41 and His 163. It was also concluded from docking results that spiroisoquinolino-piperidine substituted furoxan analogues exhibited better binding interactions compared to benzhydrylpiperazine substituted furoxan analogues. The molecular dynamic simulation demonstrated the stable complex formation between 44 and Mpro comparable to the co-crystalized ligand with ΔG value of -171.972 kj/mol [61]. A series of oxazine conjugate 9anilinoacridine derivatives was designed using molecular docking approach of Schrodinger suite 2019. All the docked ligands represented favourable conformation within the active site of SARS CoV-2 Mpro (PDB: 5R82) while compound 45 exhibited the highest binding affinity with a Glide score of -7.829 greater than the standard hydroxychloroquine (Glide score: -5.47) which could be attributed to enhanced lipophilicity and hydrogen bonding of the ligand. The --OH group of phenyl ring was found to be involved in H-bond formation with Asn 119 residue of Mpro. The determination of ADMET properties of all the designed ligands were within the acceptable values. The molecular dynamic simulation study depicted a stabilized complex formation of A38-5R82, with –OCH₃ group of **45** forming a hydrogen bond with Asn 119

while N of acridine hydrogen bonded with Arg 188 and Ser 144 and acridine moiety also showed π - π interaction with His 41 [62]. With the help of computational methods, the CAS COVID-19 antiviral compound database was screened for nearly 50,000 compounds to examine their potencies against SARS CoV-2 Mpro and RdRp. The compounds were virtually screened by docking analysis using the crystal structures of main protease (PDB: 6LU7) and RdRp (PDB: 6LM7) proteins, by SMINA software. The selected hits were further analysed for their pharmacokinetic and pharmacodynamic parameters using pkCSM model. The best possible compounds were subjected to molecular dynamic simulation for stabilization of the complexes with the proteins via GROMACS 2018 program and compounds 46 (Binding affinity: -9.064 kcal/mol) and 47 (Binding affinity 8.816 kcal/mol) showed best binding poses and good ADMET against Mpro and RdRp, respectively. The compound 46 was stabilized well in the active site of Mpro forming hydrogen bonds with Gly 143, Ser 144 and Cys 145 while the morpholine moiety and 1,3,5triazine group were involved in hydrophobic interactions with His 41 and Met 49 of catalytic dyad. The compound 47 also showed stable conformation within RdRp active site involving hydrogen bonds with Arg 553, Tvr 619 and Ser 682. There was formation of stable complexes 46 with 6LU7 and 47 with 6MU7 as per MD simulation studies [63]. Miscellaneous compounds are shown in Fig. 8.

10. Natural products

Natural products are being used as natural remedies for various ailments since time immemorial. Scientists have been actively involved in derivatization of these natural products to yield active compounds. A class of quinoid derivatives, tanshinones from Salvia miltiorrhiza, were evaluated for their antiviral potential against SARS CoV cystein proteases, 3CLpro and PLpro. The FRET based peptide cleavage assay results showed that compound 48, a dihydrotanshinone I, was found to be endowed with highest competitive inhibition potential against both SARS CoVMpro (IC50: 14.4 µM) and PLpro (IC50: 4.9 µM) and a good inhibition of deubiquitination (IC₅₀: 1.2μ M). The structure activity relationship study revealed the presence of a naphthalene ring and a dihydrofuran moiety in tanshinone 1 analogues to be crucial for anti-SARS CoV activity [64]. Twelve geranylated flavonoids were isolated from Paulownia tomentosa fruit extract with the view to examine their anti-SARS CoV potential against PLpro. The FRET inhibition assay results demonstrated compound 49 possessing a 3,4-dihydro-2H-pyran

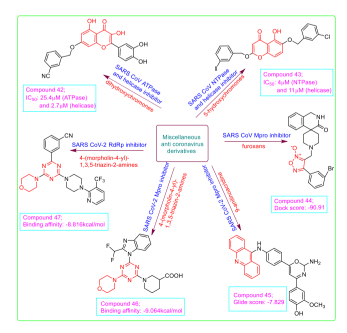


Fig. 8. Miscellaneous heterocyclic compounds with anti-coronavirus activity.

moiety, to be the most potent mixed type reversible inhibitor of SARS CoVPLpro exhibiting an IC₅₀ value of 6.1 μ M and K_i value of 3.5 μ M [65]. Another library of 64 flavonoids was examined using FRET based inhibition assay and induced fit docking analysis, to determine the inhibitory potential of the target compounds against SARS CoV 3CLpro. Herbacetin 50, a pentahydroxyflavone, showed good inhibitory activity with an IC₅₀ value of 33.17 µM. It showed good inhibition even in the presence of 0.01% Triton X-100, which is used to avoid the false bioassay results owing to the aggregating tendency of flavonoids. The induced-fit docking analysis of compound 50 within the active site of SARS CoV 3CLpro (PDB: 4WY3) revealed its good binding interaction with a glide score of -9.263. The 8 —OH group of **50** formed hydrogen bonds with Glu 166 (S1 pocket) and Gln 189 (S2 pocket) which imparted additional binding affinity to 50 with the 4WY3 compared to other analogues, kaempferol and morin which lack the hydroxyl group at position 8 [66]. Some plant alkaloids and terpenoids of African origin were analysed by molecular docking studies, for their antiviral potential against SARS CoV-2 main protease (PDB: 6LU7). Among the alkaloids, 10-hydroxyusambarensine (51) an indole alkaloid from Strychnos usambarensis (Binding affinity: -10.0 kcal/mol) and among the terpenoids, compound 6-oxoisoiguesterin (52), a bisnorterpene from Bisnor*terpenes* (Binding affinity: -9.1 kcal/mol) exhibited the highest binding potentials against SARS CoV-2 3CLpro even higher than the references taken, lopinavir (-8.3 kcal/mol) and ritonavir (-6.8 kcal/mol). There was hydrogen bond formation with Cys 145, Gln 166 and Gln 189 in 51 and in 52 with Thr 111 and Thr 292 leading to a favourable conformational fit within the active site of SARS CoV-2 Mpro. The ADMET study revealed good pharmacokinetics of both the ligands with a high gastrointestinal absorption index and no toxicity [67]. Fig. 9 highlights some natural products.

10.1. Authors' perspective

Based on the literature available, it was observed that some of the key enzymes of coronaviruses show high sequence similarity among themselves which can be exploited to target these coronaviruses by compounds with similar scaffolds and their bioisosteric analogues over a wide range. Therefore, keeping this in mind, this review gives an insight into the crucial role of heterocyclic moieties as anti-SARS CoV and anti-

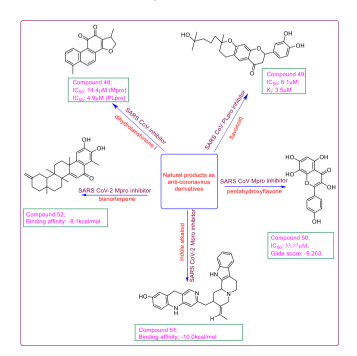


Fig. 9. Natural products as SARS CoV/CoV2 inhibitors.

SARS Cov-2 agents. Isatin nucleus has shown tremendous effects against SARS CoV Mpro involving hydrogen bond formation by carbonyl and amine groups of isatin scaffold within the active site of protease as demonstrated by docking analysis [16]. 5-carboxamide and 5-sulfonamide analogues of isatin also revealed better inhibitory potential compared to 5-iodo analogues [17,18]. As hybrids of different heterocycles have often been used to increase potency, similarly, the pyrimidine fused indole derivatives demonstrated great potential against SARS CoVMpro [20]. Drugs like vapreotide, arbidol and delavirdine have also shown good inhibition activities against SARS CoV-2 helicase, spike and RdRp proteins, respectively [21-23]. Quinoline based heterocycles are in great demand with the researches on chloroquine and hydroxychloroquine as anti-SARS CoV-2 agents [33]. The quinolinone derivatives have also shown promise against SARS CoV Mpro, thus emphasizing the need to think of more such derivatives as anti-SARS CoV-2 agents [29]. The 5-chloropyridines like MAC-5576 and pyridine N-oxides also exhibited good inhibition of SARS CoVMpro [34,35]. Further, the docking analysis revealed a good fit of 4-indolecarboxylate fused 5-chloropyrine derivatives within the binding pocket of SARS CoV Mpro [37]. Therefore, 5-chloropyridine moiety proved to be effective against SARS CoVMpro and can be further exploited against SARS CoV-2 infections. Purine analogues, 6-mercaptopurine and 6-thioguanine demonstrated reversible competitive inhibition of SARS CoV PLpro through the formation of strong hydrogen bonds within the active site of protease [41]. The *in silico* analysis of pyrimidine based drugs, carmofur, zidovudine and AP-NP unleashed a potent inhibition of SARS CoV-2 Mpro, SARS CoV-2 nucleocapsid RNA binding domain and SARS CoV-2 spike-ACE-2 complex, respectively, thus providing an insight into the need for greater research on purine and pyrimidine analogues as SARS CoV-2 inhibitors [43-45]. The pyrazole and 5-pyrazolone derivatives underwent a favourable conformation within the active site of SARS CoV 3CLpro through hydrogen bonds and hydrophobic interactions with S1, S1', S2 and S3 pockets of the protease, emphasizing on a crucial role of this moiety as SARS CoV inhibitory agent [46-48]. This review discloses a mild to moderate SARS CoV inhibitory potential of pyrazoline conjugated 4-thiazolidinones while 5-benzylidene-4-oxo-1,3-thiazolidine derivatives demonstrated good binding affinity with SARS CoV Mpro [havrylyuk; shen], thus asserting on a need for developing better 4-thiazolidinone derivatives as inhibitors of SARs coronaviruses. Some triazole derivatives also revealed potent inhibition of SARS CoV as indicated by docking analysis with SARS CoV Mpro active site [36]. Heterocyclic compounds of natural origin also play a crucial role in inhibiting the SARS CoV and SARS CoV-2 infections as depicted in this review [64-67].

11. Conclusion

With the emergence of SARS CoV-2 pandemic there is an urgent need for designing and developing safe, low-cost and potent anti-SARS CoV-2 agents and therapies with the aim to put an end to this global health crisis as early as possible. At present, a number of pharmaceutical industries and research centres throughout the world are working persistently to find a solution to the current pandemic situation. Some drugs are being repurposed based on their current therapeutic application while some are undergoing clinical trials to investigate their safety, efficacy and toxicity against SARs CoV-2 infections. Much focus has been given on pharmacotherapy, immunotherapy and plasma therapy as a means to combat SARS CoV-2 and other such life-threatening infections at present and in future. However, there is no vaccine or medication approved by FDA till date. Therefore, keeping in view the immense role of heterocyclic compounds as antiviral agents specially against coronaviruses, the better knowledge and understanding of the mechanism of heterocyclic scaffolds as anti-SARS CoV, anti-MERS CoV and anti-SARS CoV-2 agents may led to the discovery of an effective antiviral treatment thus minimizing the morbidity and mortality. As SARS CoV-2 show 82% sequence similarity with SARS CoV genome and also many target

enzymes in SARS CoV, MERS CoV and SARS CoV-2 also show similarity, therefore, many existing approved drugs and compounds revealing inhibitory potential against SARS CoV or MERS CoV can be exploited to analyse their potential against SARS CoV-2 pandemic. This review focuses on FDA approved or unapproved heterocyclic compounds involved in inhibiting SARS coronaviruses through *in vitro* or *in silico* approaches which may act as lead structures for the design and development of potent SARS CoV-2 inhibitors and other such pathogenic infections in future.

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References

- The importance of heterocyclic compounds in anti-cancer drug design. http s://www.ddw-online.com/therapeutics/p320375-the-importance-of-heterocyclic -compounds-in-anti-cancer-drug-design.html (accessed July 10, 2020).
- [2] H.A. Rothan, S.N. Byrareddy, The epidemiology and pathogenesis of coronavirus disease (COVID-19) Outbreak, J. Autoimmun. (2020) 1–4, https://doi.org/ 10.1016/j.jaut.2020.102433.
- [3] Novel coronavirus (2019-nCoV) situation report 1, 21 JANUARY 2020. https://www.who.int/docs/default-source/coronaviruse/situation-reports/20200121
 -sitrep-1-2019-ncov.pdf?sfvrsn=20a99c10_4 (accessed on July 10, 2020).
- [4] B. Shah, P. Modi, S.R. Sagar, In silico studies on therapeutic agents for COVID-19: drug repurposing approach, Life Sci. 252 (2020) 1–12, https://doi.org/10.1016/j. lfs.2020.117652.
- [5] A distinct name is needed for the new coronavirus. https://www.thelancet. com/pdfs/journals/lancet/PIIS0140-6736(20)30557-2.pdf (accessed on August 24, 2020).
- [6] Coronavirus disease (COVID-19) pandemic. World Health Organization (WHO). https://www.who.int/emergencies/diseases/novel-coronavirus-2019 (accessed on August 24, 2020).
- [7] Report of the WHO-China joint mission on coronavirus disease 2019 (COVID-19). World Health Organization (WHO). https://www.who.int/docs/default-sour ce/coronaviruse/who-china-joint-mission-on-covid-19-final-report.pdf (accessed on August 24, 2020).
- [8] 9 novel coronavirus (9-nCov) Update: uncoating the virus. https://asm.org/Articles/2020/January/9-Novel-Coronavirus-9-nCoV-Update-Uncoating (accessed on August 24, 2020).
- [9] K.G. Andersen, A. Rambaut, W.I. Lipkin, E.C. Holmes, R.F. Garry, Correspondence: the proximal origin of SARS-CoV-2, Nat. Med. 26 (2020) 450–452, https://doi.org/ 10.1038/s41591-020-0820-9.
- [10] R. Kong, G. Yang, R. Xue, M. Liu, F. Wang, J. Hu, X. Guo, S. Chang, COVID-19 docking server: an interactive server for docking small molecules, peptides and antibodies against potential targets of COVID-19, Bioinformatics (2020). https://ar xiv.org/ct?url=https%3A%2F%2Fdx.doi.org%2F10.1093%2Fbioinformatics%2F btaa645&v=22737329.
- [11] M. Prajapa, P. Sarma, N. Shekhar, P. Avti, S. Sinha, H. Kaur, S. Kumar, A. Bhattacharyya, H. Kumar, S. Bansal, B. Medhi, Drug targets for corona virus: a systematic review, Indian J. Pharmacol. 52 (2020) 56–65.
- [12] X. Liu, Z. Li, S. Liu, J. Sun, Z. Chen, M. Jiang, Q. Zhang, Y. Wei, X. Wang, Y. Huang, Y. Shi, Y. Xu, H. Xian, F. Bai, C. Ou, B. Xiong, A.M. Lew, J. Cui, R. Fang, H. Huang, J. Zhao, X. Hong, Y. Zhang, F. Zhou, H. Luo, Potential therapeutic effects of dipyridamole in the severely ill patients with COVID-19, Acta Pharm. Sin. B. 811 (2020) 1–11, https://doi.org/10.1016/j.apsb.2020.04.008.
- [13] R. Ghosh, A. Chakraborty, A. Biswas, S. Chowdhuri, Evaluation of green tea polyphenols as novel corona virus (SARS CoV-2) main protease (Mpro) inhibitors – an in silico docking and molecular dynamics simulation study, J. Biomol. Struct. Dyn. (2020) 1–13, https://doi.org/10.1080/07391102.2020.1779818.

- [14] W. Ko, J. Rolain, N. Lee, P. Chen, C. Huang, P. Lee, P. Hsueh, Arguments in favour of remdesivir for treating SARS-CoV-2 infections, Int. J. Antimicrob. Agents 55 (2020) 1–3, https://doi.org/10.1016/j.ijantimicag.2020.105933.
- [15] M.T. ul Qamar, S.M. Alqahtani, M.A. Alamri, L. Chen, Structural basis of SARS-CoV-2 3CLpro and anti-COVID-19 drug discovery from medicinal plants, J. Pharm. Anal. (2020) 1–7, https://doi.org/10.1016/j.jpha.2020.03.009.
- [16] L. Chen, Y. Wang, Y.W. Lin, S. Chou, S. Chen, L.T. Liu, Y. Wu, C. Kuo, T.S. Chen, S. Juang, Synthesis and analysis of isatin derivatives as effective SARS coronavirus 3CL protease inhibitors, Bioorg. Med. Chem. Lett. 15 (2005) 3058–3062, https:// doi.org/10.1016/j.bmcl.2005.04.027.
- [17] L. Zhou, Y. Liu, W. Zhang, P. Wei, C. Huang, J. Pei, Y. Yuan, L.J. Lai, Isatin compounds as noncovalent SARS coronavirus 3C-like protease inhibitors, J. Med. Chem. 49 (2006) 3440–3443, https://doi.org/10.1021/jm0602357.
- [18] W. Liu, H. Zhu, G. Niu, E. Shi, J. Chen, B. Sun, W. Chen, H. Zhou, C. Yang, Synthesis, modification and docking studies of 5-sulfonyl isatin derivatives as SARS-CoV 3C-like protease inhibitors, Bioorg. Med. Chem. 22 (2014) 292–302, https://doi.org/10.1016/j.bmc.2013.11.028.
- [19] P. Thanigaimalai, S. Konno, T. Yamamoto, Y. Koiwai, A. Taguchi, K. Takayama, F. Yakushiji, K. Akaji, S. Chen, A. Naser-Tavakolian, A. Schön, E. Freire, Y. Hayashi, Development of potent dipeptide-type SARS-CoV 3CL protease inhibitors with novel P3 scaffolds: design, synthesis, biological evaluation, and docking studies, Eur. J. Med. Chem. 68 (2013) 372–384, https://doi.org/10.1016/j. ejmech.2013.07.037.
- [20] S.F. Mohamed, A.A. Ibrahiem, A.E. Amr, M.M. Abdalla, SARS-CoV 3C-like protease inhibitors of some newly synthesized substituted pyrazoles and substituted pyrimidines based on 1-(3 Aminophenyl)-3-(1Hindol-3-yl)prop-2-en-1-one, Int. J. Pharmacol. 11 (2015) 749–756, https://doi.org/10.3923/ijp.2015.749.756.
- [21] J.F. Borgio, H.S. Alsuwat, W.M.A. Otaibi, A.M. Ibrahim, N.B. Almandil, L.I. A. Asoom, M. Salahuddin, B. Kamaraj, S. AbdulAzeez, State-of-the-art tools unveil potent drug targets amongst clinically approved drugs to inhibit helicase in SARS-CoV-2, Arch Med Sci. 16 (2020) 508–518, https://doi.org/10.5114/ aoms.2020.94567.
- [22] N. Vankadari, Arbidol: a potential antiviral drug for the treatment of SARS-CoV-2 by blocking trimerization of the spike glycoprotein, Int. J. Antimicrob. Agents 4 (2020) 1–3, https://doi.org/10.1016/j.ijantimicag.2020.105998.
- [23] M.A. Beg, F. Athar, Anti-HIV and Anti-HCV drugs are the putative inhibitors of RNA-dependent-RNA polymerase activity of NSP12 of the SARS CoV-2 (COVID-19), Pharm. Pharmacol. Int. J. 8 (2020) 163–172, https://doi.org/10.15406/ ppij.2020.08.00292.
- [24] W. Dai, B. Zhang, X. Jiang, H. Su, J. Li, Y. Zhao, X. Xie, Z. Jin, J. Peng, F. Liu, C. Li, Y. Li, F. Bai, H. Wang, X. Cheng, X. Cen, S. Hu, X. Yang, J. Wang, X. Liu, G. Xiao, H. Jiang, Z. Rao, L. Zhang, Y. Xu, H. Yang, H. Liu, Structure-based design of antiviral drug candidates targeting the SARS-CoV-2 main protease, Science 368 (2020) 1331–1335, https://doi.org/10.1126/science.abb4489.
- [25] X. Wang, R. Cao, H. Zhang, J. Liu, M. Xu, H. Hu, Y. Li, L. Zhao, W. Li, X. Sun, X. Yang, Z. Shi, F. Deng, Z. Hu, W. Zhong, M. Wang, The anti-influenza virus drug, arbidol is an efficient inhibitor of SARS-CoV-2 in vitro, Cell Discov. 6 (2020) 1–5, https://doi.org/10.1038/s41421-020-0169-8.
- [26] E. Keyaerts, L. Vijgen, P. Maes, J. Neyts, M.V. Ranst, In vitro inhibition of severe acute respiratory syndrome coronavirus by chloroquine, Biochem. Bioph. Res. Co. 323 (2004) 264–268, https://doi.org/10.1016/j.bbrc.2004.08.085.
- [27] M.J. Vincent, E. Bergeron, S. Benjannet, B.R. Erickson, P.E. Rollin, T.G. Ksiazek, N. G. Seidah, S.T. Nichol, Chloroquine is a potent inhibitor of SARS coronavirus infection and Spread, Virol. J. 2 (2005) 1–10, https://doi.org/10.1186/1743-422X-2-69.
- [28] C. Biot, W. Daher, N. Chavain, T. Fandeur, J. Khalife, D. Dive, E. de Clercq, Design and synthesis of hydroxyferroquine derivatives with antimalarial and antiviral activities, J. Med. Chem. 49 (2006) 2845–2849, https://doi.org/10.1021/ jm0601856.
- [29] Y. Sun, N. Zhang, J. Wang, Y. Guo, B. Sun, W. Liu, H. Zhou, C. Yang, Synthesis and biological evaluation of quinolinone compounds as SARS CoV 3CLpro inhibitors, Chin. J. Chem. 31 (2013) 1199–1206, https://doi.org/10.1002/cjoc.201300392.
- [30] A.H. de Wilde, D. Jochmans, C.C. Posthuma, J.C. Zevenhoven-Dobbe, S. van Nieuwkoop, T.M. Bestebroer, B.G. van den Hoogen, J. Neyts, E.J. Snijder, Screening of an FDA-approved compound library identifies four small-molecule inhibitors of middle east respiratory syndrome coronavirus replication in cell culture, Antimicrob. Agents Chemother. 58 (2014) 4875–4884, https://doi.org/ 10.1128/AAC.03011-14.
- [31] Y. Shimamoto, Y. Hattori, K. Kobayashi, K. Teruya, A. Sanjoh, A. Nakagawa, E. Yamashita, K. Akaji, Fused-ring structure of decahydroisoquinoline as a novel scaffold for SARS 3CL protease inhibitors, Bioorg. Med. Chem. 23 (2015) 876–890, https://doi.org/10.1016/j.bmc.2014.12.028.
- [32] S. Yuan, J.F.W. Chan, K.K.H. Chik, C.C.Y. Chan, J.O.L. Tsang, R. Liang, J. Cao, K. Tang, L. Chen, K. Wen, J. Cai, Z. Ye, G. Lu, H. Chu, D. Jin, K. Yuen, Discovery of the FDA-approved drugs bexarotene, cetilistat, diiodohydroxyquinoline, and abiraterone as potential COVID-19 treatments with a robust two-tier screening system, Pharmacol. Res. 159 (2020) 1–9, https://doi.org/10.1016/j. phrs.2020.104960.
- [33] J. Fantini, C. Di Scala, H. Chahinian, N. Yahi, Structural and molecular modelling studies reveal a new mechanism of action of chloroquine and hydroxychloroquine against SARS-CoV-2 infection, Int. J. Antimicrob. Agents 55 (2020) 1–8, https:// doi.org/10.1016/j.ijantimicag.2020.105960.
- [34] J.E. Blanchard, N.H. Elowe, C. Huitema, P.D. Fortin, J.D. Cechetto, L.D. Eltis, E. D. Brown, High-throughput screening identifies inhibitors of the SARS coronavirus main proteinase, Chem. Biol. 11 (2004) 1445–1453, https://doi.org/10.1016/j. chembiol.2004.08.011.

- [35] J. Balzarini, E. Keyaerts, L. Vijgen, F. Vandermeer, M. Stevens, E. de Clercq, H. Egberink, M.V. Ranst, Pyridine N-oxide derivatives are inhibitory to the human SARS and feline infectious peritonitis coronavirus in cell culture, J. Antimicrob. Chemother. 57 (2006) 472-481, https://doi.org/10.1093/jac/dki481.
- Chemother. 57 (2006) 472–481, https://doi.org/10.1093/jac/dki481.
 [36] C. Wu, K. King, C. Kuo, J. Fang, Y. Wu, M. Ho, C. Liao, J. Shie, P. Liang, C. Wong, Stable benzotriazole esters as mechanism-based inactivators of the severe acute respiratory syndrome 3CL protease, Chem. Biol. 13 (2006) 261–268, https://doi.org/10.1016/j.chembiol.2005.12.008.
- [37] A.K. Ghosh, G. Gong, V. Grum-Tokars, D.C. Mulhearn, S.C. Baker, M. Coughlin, B. S. Prabhakar, K. Sleeman, M.E. Johnson, A.D. Mesecar, Design, synthesis and antiviral efficacy of a series of potent chloropyridyl ester-derived SARS-CoV 3CLpro inhibitors, Bioorg. Med. Chem. Lett. 18 (2008) 5684–5688, https://doi.org/10.1016/j.bmcl.2008.08.082.
- [38] C. Niu, J. Yin, J. Zhang, J.C. Vederas, M.N.G. James, Molecular docking identifies the binding of 3-chloropyridine moieties specifically to the S1pocket of SARS-CoV Mpro, Bioorrg. Med. Chem. 16 (2008) 293–302, https://doi.org/10.1016/j. bmc.2007.09.034.
- [39] K. Tsai, S. Chen, P. Liang, I. Lu, N. Mahindroo, H. Hsieh, Y. Chao, L. Liu, D. Liu, W. Lien, T. Lin, S. Wu, Discovery of a novel family of SARS-CoV protease inhibitors by virtual screening and 3D-QSAR studies, J. Med. Chem. 49 (2006) 3485–3495, https://doi.org/10.1021/jm050852f.
- [40] M. Ikejiri, M. Saijo, S. Morikawa, S. Fukushi, T. Mizutani, I. Kurane, T. Maruyama, Synthesis and biological evaluation of nucleoside analogues having 6-chloropurine as anti-SARS-CoV agents, Bioorg. Med. Chem. Lett. 17 (2007) 2470–2473, https:// doi.org/10.1016/j.bmcl.2007.02.026.
- [41] C. Chou, C. Chien, Y. Han, M.T. Prebanda, H. Hsieh, B. Turk, G. Chang, X. Chen, Thiopurine analogues inhibit papain-like protease of severe acute respiratory syndrome coronavirus, Biochem. Pharmacol. 75 (2008) 1601–1609, https://doi. org/10.1016/j.bcp.2008.01.005.
- [42] R. Ramajayam, K. Tan, H. Liu, P. Liang, Synthesis, docking studies, and evaluation of pyrimidines as inhibitors of SARS-CoV 3CL protease, Bioorg. Med. Chem. Lett. 20 (2010) 3569–3572, https://doi.org/10.1016/j.bmcl.2010.04.118.
- [43] Z. Jin, Y. Zhao, Y. Sun, B. Zhang, H. Wang, Y. Wu, Y. Zhu, C. Zhu, T. Hu, X. Du, Y. Duan, J. Yu, X. Yang, X. Yang, K. Yang, X. Liu, L.W. Guddat, G. Xiao, L. Zhang, H. Yang, Z. Rao, Structural basis for the inhibition of SARS-CoV-2 main protease by antineoplastic drug carmofur, Nat Struct. Mol. Biol. 27 (2020) 529–532, https:// doi.org/10.1038/s41594-020-0440-6.
- [44] R. Yadav, M. Imran, P. Dhamija, K. Suchal, S. Handu, Virtual screening and dynamics of potential inhibitors targeting RNA binding domain of nucleocapsid phosphoprotein from SARS-CoV-2, J. Biomol. Struct. Dyn. (2020) 1–16, https:// doi.org/10.1080/07391102.2020.1778536.
- [45] J.S. Rane, P. Pandey, A. Chatterjee, R. Khan, A. Kumar, A. Prakash, S. Ray, Targeting virus-host interaction by novel pyrimidine derivative: an in silico approach towards discovery of potential drug against COVID-19, J. Biomol. Struct. Dyn. (2020) 1–11, https://doi.org/10.1080/07391102.2020.1794969.
- [46] C. Kuo, H. Liu, Y. Lo, C. Seong, K. Lee, Y. Jung, P. Liang, Individual and common inhibitors of coronavirus and picornavirus main proteases, FEBS Lett. 583 (2009) 549–555, https://doi.org/10.1016/j.febslet.2008.12.059.
- [47] R. Ramajayam, K. Tan, H. Liu, P. Liang, Synthesis and evaluation of pyrazolone compounds as SARS-coronavirus 3C-like protease inhibitors, Bioorg. Med. Chem. 18 (2010) 7849–7854, https://doi.org/10.1016/j.bmc.2010.09.050.
- [48] V. Kumar, K. Tan, Y. Wang, S. Lin, P. Liang, Identification, synthesis and evaluation of SARS-CoV and MERS-CoV 3C-like protease inhibitors, Bioorg. Med. Chem. 24 (2016) 3035–3042, https://doi.org/10.1016/j.bmc.2016.05.013.
- [49] D. Kumar, K. Kumari, V.K. Vishvakarma, A. Jayaraj, D. Kumar, V.K. Ramappa, R. Patel, V. Kumar, S.K. Dass, R. Chandra, P. Singh, Promising inhibitors of main protease of novel corona virus to prevent the spread of COVID-19 using docking and molecular dynamics simulation, J. Biomol. Struct. Dyn. (2020) 1–15, https:// doi.org/10.1080/07391102.2020.1779131.
- [50] T. Regnier, D. Sarma, K. Hidaka, U. Bacha, E. Freire, Y. Hayashi, Y. Kiso, New developments for the design, synthesis and biological evaluation of potent SARS-CoV 3CLpro inhibitors, Bioorg. Med. Chem. Lett. 19 (2009) 2722–2727, https:// doi.org/10.1016/j.bmcl.2009.03.118.
- [51] D. Atamanyuk, B. Zimenkovsky, V. Atamanyuk, R. Lesyk, 5-Ethoxymethylidene-4thioxo-2-thiazolidinone as versatile building block for novel biorelevant small molecules with thiopyrano[2,3-d] [1,3]thiazole core, Synth. Commun. 44 (2014) 237–244, https://doi.org/10.1080/00397911.2013.800552.

- [52] S. Konno, P. Thanigaimalai, T. Yamamoto, K. Nakada, R. Kakiuchi, K. Takayama, Y. Yamazaki, F. Yakushiji, K. Akaji, Y. Kiso, Y. Kawasaki, S. Chen, E. Freire, Y. Hayashi, Design and synthesis of new tripeptide-type SARS-CoV 3CL protease inhibitors containing an electrophilic arylketone moiety, Bioorg. Med. Chem. 21 (2013) 412–424, https://doi.org/10.1016/j.bmc.2012.11.017.
- [53] D. Havrylyuk, B. Zimenkovsky, O. Vasylenko, R. Lesyk, Synthesis and anticancer and antiviral activities of new 2-pyrazoline-substituted 4-thiazolidinones, J. Heterocycl. Chem. 50 (2013) E55–E62, https://doi.org/10.1002/jhet.1056.
- [54] D. Havrylyuk, B. Zimenkovsky, O. Vasylenko, C.W. Day, D.F. Smee, P. Grellier, R. Lesyk, Synthesis and biological activity evaluation of 5-pyrazoline substituted 4thiazolidinones, Eur. J. Med. Chem. 66 (2013) 228–237, https://doi.org/10.1016/ j.ejmech.2013.05.044.
- [55] D.V. Kaminskyy, Screening of the antiviral activity in the range of C5 and N3 substituted 4-thiazolidinone derivatives, J. Org. Pharm. Chem. 13 (2015) 64–69, https://doi.org/10.24959/ophcj.15.819.
- [56] J. Shen, H. Jiang, X. Shen, S. Li, W. Huang, C. Gui, J. Chen, T. Sun, F. Ye, D. Bai, H. Liu, X. Luo, K. Chen, Patent CN200410018418, 2005.
- [57] M. Turlington, A. Chun, S. Tomar, A. Eggler, V. Grum-Tokars, J. Jacobs, J. S. Daniels, E. Dawson, A. Saldanha, P. Chase, Y.M. Baez-Santos, C.W. Lindsley, P. Hodder, A.D. Mesecar, S.R. Stauffer, Discovery of N-(benzo[1,2,3]triazol-1-yl)-N-(benzyl)acetamido) phenyl) carboxamides as severe acute respiratory syndrome coronavirus (SARS-CoV) 3CLpro inhibitors: identification of ML300 and noncovalent nanomolar inhibitors with an induced-fit binding, Bioorg. Med. Chem. Lett. 23 (2013) 6172–6177, https://doi.org/10.1016/j.bmcl.2013.08.112.
- [58] C.J. Cortés-García, L. Chacón-García, J.E. Mejía-Benavides, E. Díaz-Cervantes, Tackling the SARS-CoV-2 main protease using hybrid derivatives of 1,5-disubstituted tetrazole-1,2,3-triazoles: an in silico assay, PeerJ Phys. Chem. 2 (e10) (2020) 1–16, https://doi.org/10.7717/peerj-pchem.10.
- [59] C. Lee, J.M. Lee, N. Lee, D. Kim, Y. Jeong, Y. Chong, Investigation of the pharmacophore space of Severe Acute Respiratory Syndrome coronavirus (SARS-CoV) NTPase/helicase by dihydroxychromone Derivatives, Bioorg. Med. Chem. Lett. 19 (2009) 4538–4541, https://doi.org/10.1016/j.bmcl.2009.07.009.
- [60] M.K. Kim, M. Y, H.R. Parka, K.B. Kim, C. Lee, S.Y. Cho, J. Kang, H. Yoon, D. Kim, H. Choo, Y. Jeong, Y. Chong, 2,6-Bis-arylmethyloxy-5-hydroxychromones with antiviral activity against both hepatitis C virus (HCV) and SARS-associated coronavirus (SCV), Eur. J. Med. Chem. 46 (2011) 5698–5704, https://doi.org/ 10.1016/j.ejmech.2011.09.005.
- [61] A.G. Al-Sehemi, M. Pannipara, R.S. Parulekar, O. Patil, P.B. Choudhari, M. S. Bhatia, P.K. Zubaidh, Y. Tamboli, Potential of NO donor furoxan as SARS-CoV-2 main protease (Mpro) inhibitors: in silico analysis, J. Biomol. Struct. Dyn. (2020) 1–15, https://doi.org/10.1080/07391102.2020.1790038.
- [62] K. Rajagopal, P. Varakumar, B. Aparna, G. Byran, S. Jupudi, Identification of some novel oxazine substituted 9-anilinoacridines as SARS-CoV-2 inhibitors for COVID-19 by molecular docking, free energy calculation and molecular dynamics studies, J. Biomol. Struct. Dyn. (2020) 1–12, https://doi.org/10.1080/ 07391102.2020.1798285.
- [63] A. Aouidate, A. Ghaleb, S. Chtita, M. Aarjane, A. Ousaa, H. Maghat, A. Sbai, M. Choukrad, M. Bouachrine, T. Lakhlifi, Identification of a novel dual-target scaffold for 3CLpro and RdRp proteins of SARS-CoV-2 using 3Dsimilarity search, molecular docking, molecular dynamics and ADMET evaluation, J. Biomol. Struct. Dyn. (2020) 1–14, https://doi.org/10.1080/07391102.2020.1779130.
- [64] J. Park, J.H. Kim, Y.M. Kim, H.J. Jeong, D.W. Kim, K.H. Park, H. Kwon, S. Park, W. S. Lee, Y.B. Ryu, Tanshinones as selective and slow-binding inhibitors for SARS-CoV cysteine proteases, Bioorg. Med. Chem. 20 (2012) 5928–5935, https://doi.org/10.1007/s12272-012-0108-9.
- [65] J. Keun Cho, M.J. Curtis-Long, K.H. Lee, D.W. Kim, H.W. Ryu, H.J. Yuk, Ki. H. Park, Geranylated flavonoids displaying SARS-CoV papain-like protease inhibition from the fruits of Paulownia tomentosa, Bioor. Med. Chem. 21 (2013) 3051–3057, https://doi.org/10.1016/j.bmc.2013.03.027.
- [66] S. Jo, S. Kim, D.H. Shin, M. Kim, Inhibition of SARS-CoV 3CL protease by flavonoids, J. Enzyme Inhib. Med. Chem. 35 (2020) 145–151, https://doi.org/ 10.1080/14756366.2019.1690480.
- [67] G.A. Gyebi, O.B. Ogunro, A.P. Adegunloye, O.M. Ogunyemi, S.O. Afolabi, Potential inhibitors of coronavirus 3-chymotrypsin-like protease (3CLpro): an in silico screening of alkaloids and terpenoids from African medicinal plants, J. Biomol. Struct. Dyn. (2020) 1–13, https://doi.org/10.1080/07391102.2020.1764868.

Review Article



Molnupiravir – A prospective silver bullet to mitigate severe acute respiratory syndrome corona virus-2

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ABSTRACT

The whole world is eagerly waiting for the unearthing of the best treatment strategy to put an end to the prevailing coronavirus disease-2019 pandemic. The pathogen responsible for this disease, that is, severe acute respiratory syndrome coronavirus-2 (SARS CoV-2) continues to be the most challenging issue that has kept the researchers and innovators all over the world in a dilemma of resolving it through finding the most efficacious, safe, and cost-effective therapy. A large number of drugs are under investigation as a part of drug repurposing approach for the treatment of SARS-CoV-2. One such drug, molnupiravir, is under Phase II/III clinical trials against SARS-CoV-2. Through this work, the authors will give an insight into the various aspects of molnupiravir as an antiviral agent including chemistry, pharmacokinetics, synthetic route, *in vitro*, *in vivo* studies, clinical trials, and probable mode of antiviral action of molnupiravir against SARS-CoV-2. The molecular docking approach has also been used to evaluate the binding interactions of the active form of molnupiravir, N-4-hydroxycytidine, with the RNA-dependent RNA polymerase of SARS-CoV-2 which emphasized on its good binding potential with the active site residues displaying a binding energy of -6.4 kcal per mol.

Keywords: Molnupiravir, severe acute respiratory syndrome coronavirus-2, clinical trials, docking studies

INTRODUCTION

Coronaviruses (CoV), a group of RNA viruses belonging to the family, Coronaviridae, are positive-sense single-strand RNA viruses having the potential to cause mild to fatal respiratory tract infections in mammals and birds.^[1] The outbreak of CoV began with severe acute respiratory syndrome coronavirus-2 (SARS CoV) in 2002 in China spreading from civet cats to humans which was later observed in the form of Middle-East respiratory syndrome coronavirus (MERS CoV) in the year 2012, in Saudi Arabia, transmitting from dromedary cats to humans. Most recently emerged are the cluster of cases having symptoms like pneumonia in the Wuhan city of China, in December 2019 which was later named by the World Health Organization as coronavirus disease 2019 (COVID-19) and declared as pandemic in March 2020.^[2]The CoVs comprise structural

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(spike, envelope, membrane, and nucleocapsid) and non-structural proteins (nsp) (main protease, RNA-dependent RNA polymerase [RdRp] [nsp 12], helicase, papain-like protease, N-terminal exoribonuclease, nsp 10, nsp 14, nsp 15, and nsp 16).^[3]The SARS-CoV-2 exhibits approximately 80% sequence similarity with other bat CoVs.^[4] It is more pathogenic than the other CoVs due to the presence of a unique polybasic cleavage site and has higher transmissibility due to the 10–20 times greater binding of SARS-CoV-2 with angiotensin-converting enzyme-2 (ACE-2) receptor compared to SARS-CoV with ACE-2.^[5,6]

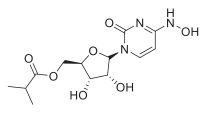
The SARS-CoV-2 releases its nucleocapsid into the patient's cells after attacking the host's lower respiratory tract, leading to viral replication responsible for the emanation of pneumonia-like symptoms.^[7] In critical cases, SARS-CoV-2 infection can lead to multiple organ damage adversely affecting the heart, liver, lungs, kidney, gastrointestinal system, and central nervous system. The membrane protein has a role in virus humoral response together with neutralizing developed antibodies while spike protein assists the viral entry into host cellular machinery using host ACE-2 receptor. Transmembrane protease, serine 2 (TMPRSS2),

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a transmembrane serine protease, cleaves the spike protein into S1 and S2 domain leading to its activation and viral entry into the host cell. The entry is followed by translocation which occurs through endocytic and non-endocytic pathways. Once the virus enters the host cell endosomes, the viral envelope protein supports the viral genome release into host cell as single-stranded positive RNA which further moves to transcription and translation processes using host cell machinery.

RdRp plays an important role in viral transcription and replication and has proved to be a crucial target in many antiviral therapies like remdesivir. RdRp shows high sequence similarity among the three CoVs while it is not expressed by the host cells, therefore, drugs targeting RdRp or RdRp inhibitors used against SARS-CoV and MERS-CoV can also be repurposed against SARS-CoV-2 with high potency and selectivity to viral entry.^[2,8,9] RdRp catalyzes viral RNA genome synthesis using *de novo* (primer dependent) or primer independent molecular mechanisms. The *de novo* RNA synthesis involves the phosphodiester bond formation between 3'-hydroxyl end of one nucleotide and the 5'-phosphate moiety from next nucleotide. While in primer dependent mechanism, an oligonucleotide or a protein primer acts as a template for the development of a new complimentary RNA, through base pairing.^[9]

Molnupiravir (MK-4482 and EIDD-2801) is an orally bioavailable 5'-isopropylester prodrug of β -D-N-hydroxycytidine (EIDD-1931, or N-hydroxycytidine, NHC), a ribonucleoside analog with potent anti-influenza activity. Molnupiravir was originally developed by Drug Innovation Ventures at Emory, a drug innovation company of Emory University, as an inhibitor of influenza viral replication. Later, the drug was acquired by Ridgeback Biotherapeutics which collaborated with Merck to develop it further.^[10,11] Afterward, the drug was tested for its activity against SARS-CoV and MERS-CoV also. In March 2020, molnupiravir was found to be active against SARS-CoV-2 and thereafter it underwent randomized, double-blind, placebocontrolled, first-in-human study, in the US and UK, to test its safety, tolerability, and pharmacokinetics in healthy volunteers. In October 2020, a phase II/III randomized, placebo-controlled, double-blind clinical study of the drug was started by Merck on hospitalized patients to test its efficacy, safety, and pharmacokinetics.^[12,13] Molnupiravir was found to be effective when given orally in SARS-CoV-2-infected ferrets blocking the viral transmission in them.^[14]



Drug Name: Molnupiravir (MK-4482 and EIDD-2801) IUPAC: ((2R,3S,4R,5R)-3,4-dihydroxy-5-(4-(hydroxyimino)-2-oxo-3,4-dihydropyrimidin-1(2H)yl)tetrahydrofuran-2-yl)methyl isobutyrate Formula:C₁₃H₁₉N₃O₇ Molecular mass: 329.31g/mol Category: Antiviral, a ribonucleoside analogue inhibitor of RNA virus

PATHOPHYSIOLOGY

SARS-CoV-2 is a β -coronavirus showing genomic similarity of nearly 80% with SARS-CoV-1 and 96.2% with bat coronavirus RaTG₁₃ suggesting bats as the main source of its transmission.^[15] It is a pleomorphic virus having a diameter of ~125 nm and the RNA genome of 30 kb(+) together with up to 10 open reading frames (ORFs). As per global initiative on sharing all influenza data database, three SARS-CoV-2 clades are identified as (a) G clade (spike S protein variant, D614G), (b) V clade (ORF3 variant, G251V), and (c) S clade (ORF8 variant, L84S). In India, A2a clade, S protein variant, D614G, was found to be the most prominent clade (48.6%). These clades show divergence in virulence thus affecting the effectiveness of repurposed drugs or of future vaccines and biologicals.^[8,16,17]

SARS-CoV-2 viral infection is mostly found to be transmitted from person to person through respiratory droplets and aerosols through coughing and sneezing. The nasopharyngeal swab and feces have shown presence of the virus, and therefore, fecal-oral route transmission can also be a possibility.^[18] It can also involve contact transmission that is by talking to the infected person or by inhaling the exhaled gas from infected person within a distance of about 6 feet and indirect transmission by coming in contact with contaminated droplets from mouth, nose, and eyes settled on different surfaces.^[16]

The SARS-CoV-2 infection begins with the entry of virus into host cells by the attachment of its spike protein with the host cell's ACE-2 cell surface receptor on alveolar epithelial type II (AT 2) cells in respiratory tract. The S protein comprises S1 (N-terminal domain) and S2 (C-terminal domain) domains. The S1 domain is responsible for receptor binding and is also termed as receptor binding domain. SD1 and SD2 are the two subdomains of S1 domain and allow the conformational changes in S2 domain on binding to the receptor.^[8]The life cycle of virus in host is believed to involve these major stages: Spike protein fusion with host ACE-2 receptor (attachment), spike protein cleavage by TMPRSS2 (activation), membrane fusion or endocytosis of virus by host cells (penetration), viral ssRNA entry into the host cell nucleus and synthesis of viral proteins by viral mRNA (biosynthesis), and maturation and release of new viral particles (maturation).^[8,19]

TMPRSS2, a transmembrane serine protease (type II), cleaves the spike protein into S1 and S2 domains, thus making it possible for S1 domain to interact with host ACE-2 for entry into the host cells. After entering the host cells, the ss(+) RNA undergoes replication using viral RdRp forming complimentary ss(-) RNA which further leads to the formation of new positive mRNA strands suitable for synthesis of new viral proteins in host cells. The translation of viral genome leads to the formation of viral polyproteins which are cleaved by viral main protease and papain like proteases into effector proteins. The binding of nucleocapsid protein to positive strand RNA forms a nucleoprotein complex while the spike, envelope, and membrane proteins move to the endoplasmic reticulum. Thus, the virion assembly gets completed in Golgi apparatus and is now ready for its release from infected cells through exocytosis, as depicted in Figure 1. The viral main protease and papain-like protease have the ability to deubiquitinase NFkB and interferon factor 3 in host

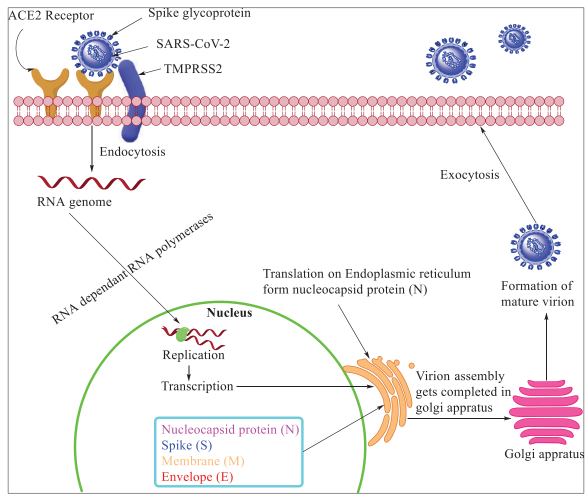


Figure 1: Pathophysiology of severe acute respivirus-2

cells, thus suppressing the host innate immunity.^[2,8,16,20,21] Both lung inflammation and immune deficiency are the two interconnected processes involved in SARS-CoV-2 pathogenesis. ACE-2 is a metalloproteinase which is expressed in organs such as lungs, CNS, cardiovascular system, kidneys, gastrointestinal tract, and adipose tissues. The whole viral life cycle in the host cells leads to the release of cytokines and inflammatory markers such as interleukins, interferons, TNF-α, macrophage inflammatory protein 1α , monocyte chemoattractant protein-1, and chemokines like CXCL10 which results into vasodilation and an increase in capillary permeability. The "cytokine storm" causes the recruitment of CD4+ helper T cells, neutrophils, and CD8 cytotoxic T cells which results in further lung inflammation and injury. The resultant apoptosis of the host cells leads to new viral particle release and infection of surrounding type II alveolar epithelial cells, ultimately resulting in acute respiratory distress syndrome.^[21,22]

MECHANISM OF ACTION

Molnupiravir is a prodrug of N-4-hydroxycytidine, therefore *in vivo*, it gets hydrolyzed to its active form which exists in two tautomeric forms: A cytidine mimic (pairs with guanosine) and a uridine mimic (pairs with adenosine) form. In the process of viral RNA replication

using RdRp, switching between these two mimic forms led to mismatches causing catastrophic mutations in the newly generated viral RNA transcripts thus rendering them non-functional, as shown in Figure 2.^[10,22]

METABOLISM

Molnupiravir gets hydrolyzed to β -D-N⁴-hydroxycytidine (NHC, EIDD-1931) in host cells which further gets phosphorylated to its active 5'-triphosphate form in tissues as depicted in Figure 3.^[10]

SYNTHETIC STRATEGY

Molnupiravir shows structural similarity with antiviral drug, remdesivir but both the drugs block RdRp in a different way and therefore can act complimentarily. Molnupiravir is an orally active drug and therefore can have an advantage of increased patient compliance in SARS-CoV-2 patients over remdesivir which is administered intravenously.

Molnupiravir was first developed by Emory University researchers through a five-step synthesis route using uridine as the precursor. In this patented route, initially, the protection of acetonide

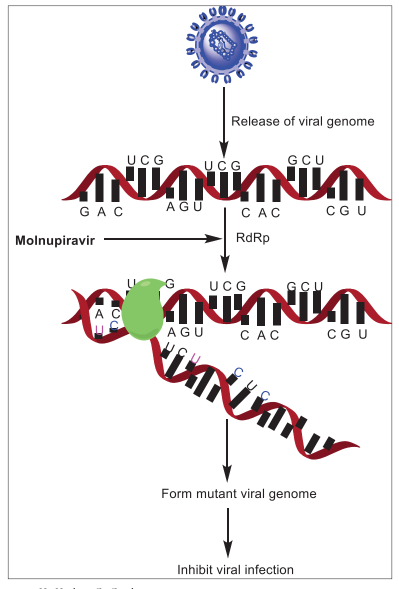


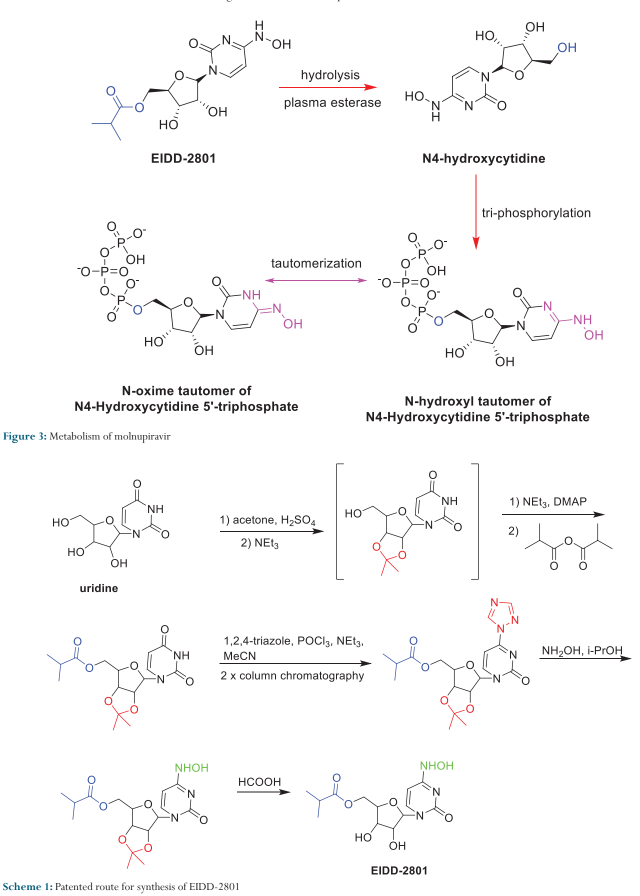
Figure 2: Mechanism of molnupiravir: U=Uridine, C=Cytidine

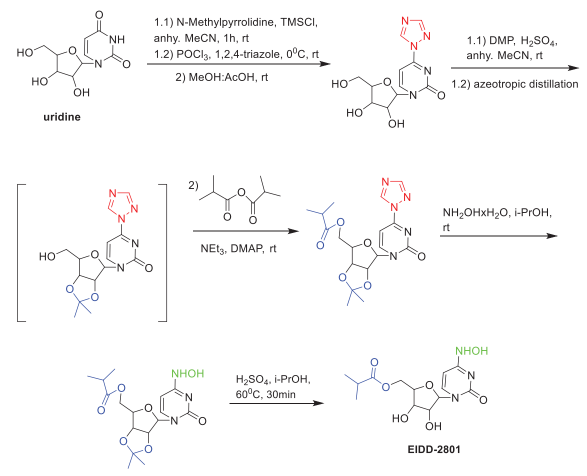
is done followed by selective esterification of the 5'-hydroxy group. Further, the activation of the molecule is performed by introduction of 1,2,4-triazole nucleus which gets replaced by hydroxylamine. This is followed by deprotection of the acetonide group yielding the final product, molnupiravir. This synthetic strategy has a disadvantage of very low yield at triazole coupling step (29%) [Scheme 1]. With an aim to improve the synthetic yield, a modification of the patented scheme was done. In this method, the triazole coupling was carried out in the first step using Et₂N as base, thus increasing the yield of this step from 29% to 88%. In the second step, the acetonide ester was obtained by stirring the triazole derivative with DMP and sulfuric acid in MeCN for 30 min. This was followed by conversion of acetonide ester to hydroxylamine by stirring it with hydroxylamine and iPrOH at room temperature. Finally, the deprotection of acetonide group was obtained by reaction with sulfuric acid and iPrOH as solvent at 60°C for 30 min resulting in the formation of title compound with 80% yield [Scheme 2].^[23]

IN VITRO AND *IN VIVO* STUDY OF MOLNUPIRAVIR

Urakova *et al.* evaluated the role of NHC, a nucleoside analogue, as an anti-VEEV (Venezuelan equine encephalitis virus) agent. In the plaque assay, NHC revealed a strong anti-viral action in VEEV treated Vero cells, when applied before, at the time of or 4 h post infection (p.i.) at $1-2\,\mu$ M concentration with an EC₅₀ value of 1 μ M. It was observed that NHC acts by inducing mutations in viral G RNA rendering them incapable of replication that is by causing viral lethal mutagenesis. The mutations acquired were mostly transition mutations like U-to-C or C-to-U transitions generated at the time of positive-strand RNA synthesis and A-to-G and G-to-A transitions occurring as a result of incorporation of NHC in negative-strand RNA.^[24]

Due to the low oral bioavailability of NHC in cynomolgus macaques, a 5'-isopropyl ester of NHC, EIDD-2801, was synthesized with the aim to improve its oral bioavailability. EIDD-2801 showed similar





Scheme 2: New route for synthesis of EIDD-2801

bioavailability in mice as that of NHC but an improved bioavailability in nonhuman-primates and ferrets. It was observed that EIDD-2801 was hydrolyzed to NHC *in vivo*. A decrease in group pandemic 1 and group 2 seasonal influenza A shed virus load, inflammation, fever, and airway epithelium histopathology was observed in influenza virus infected ferrets, on oral therapeutic administration of EIDD-2801. Further, whole genome deep sequencing analysis revealed lethal mutagenesis as the mechanism involved in NHC influenza virus inhibition together with high barrier to viral resistance. Antiviral activity analysis in human airway epithelia (HAE) model revealed a CC_{50} value of 137 μ M and a high therapeutic window of >1713 of NHC against different strains of influenza virus.^[10]

Toots *et al.* conducted experiments in influenza infected ferret models to determine the quantitative efficacy parameters of EIDD-2801 like its minimum effective dose, latest onset of effective treatment together with the minimum doses required for maximum effect. The analysis results demonstrated 7 mg/kg of EIDD-2801 given by oral route, as the lowest efficacious dose following a b.i.d dosing regimen. Furthermore, a 36 h time window was found to exist for the initiation of effective treatment p.i. in ferrets. The administration of 7 mg/kg dose of EIDD-2801 at intervals of 12 h was observed to be sufficient to achieve maximum therapeutic effect against pandemic influenza A virus.^[11]

of NHC, an active form of prodrug EIDD-2801, against coronavirus strains in mouse models. NHC showed potent anti-viral activity in MERS-CoV-infected Calu-3 2B4 human lung epithelial cell lines with an IC₅₀ value of 0.15 μ M and a CC₅₀ value of >10 μ M while an IC₅₀ value of 0.3 μ M and a CC₅₀ value of >10 μ M in SARS-CoV-2 (2019nCoV/USA-WA1/2020) infected Vero cells. Furthermore, the anti-SARS CoV-2 assay in Calu-3 cells suggested a dose-dependent decrease in viral titers and vial genomic RNA (IC₅₀ $0.09 \,\mu$ M). Further, antiviral assay study in HAE cells demonstrated no cytotoxicity up to a dose of 100 μ M of NHC while at the same time, a dose-dependent decrease in SARS-CoV-2 replication was also observed in SARS-CoV-2-infected HAE cells. Similarly, a reduction in viral titer and genomic (ORF1) and subgenomic (ORFN) RNA was revealed in MERS-CoV- and SARS-CoV-infected HAE cells with IC₅₀ values of 0.024 μ M and 0.14 μ M, respectively. The virus titer reduction assay in DBT cells unleashed an increased sensitivity to inhibition by NHC in coronavirus bearing resistance mutations to antiviral drug, remdesivir. NHC also displayed lethal mutagenicity and error catastrophe induction in RdRp, in MERS-CoV-infected HAE cells with A-to-G and U-to-C transitions in RNA. The in vivo study of EIDD-2801, an orally bioavailable form of NHC, in SARS-CoV- and MERS-CoV-infected C57BL/6 mice revealed a significant decrease in weight loss, lung titer, and hemorrhage at a dose of 500 mg/kg, both prophylactically and therapeutically.^[25]

Sheahan et al. demonstrated the anti-viral mechanism and efficacy

Due to the fact that ferrets transmit the SARS-CoV-2 virus effectively with minimum clinical disease manifestations resembling the SARS-CoV-2 asymptomatic or mildly symptomatic transmission in young human population, Cox *et al.* explored the efficacy of EIDD-2801 in SARS-CoV-2-infected ferret models. The *in vivo* study involved administration of EIDD-2801 as oral gavage, at doses of 5 or 15 mg/kg b.i.d. 12 h p.i. or 15 mg/kg 36 h p.i. in ferrets (*Mustela putorius furo*) inoculated with 1×10^5 plaque-forming units of SARS-CoV-2 2019-nCoV/USA-WA1/2020 clinical isolate. The assay results unleashed the high potential of EIDD-2801, significantly decreasing the SARS-CoV-2 viral load in upper respiratory tract together with suppressing the transmission of infection to untreated contact animals.^[14]

Abdelnabi *et al.* evaluated the anti-viral effect of EIDD-2801 in SARS-CoV-2-infected Syrian hamster model. The hamster models infected with 2×10^{6} TCID₅₀ SARS-CoV-2 (BetaCov/Belgium/GHB-03021/2020 [EPI ISL 109 407976 | 2020-02-03]), were treated with 75 or 200 mg/kg b.i.d. of EIDD-2801 for 4 consecutive days. The assay results revealed a significant reduction in virus titers and RNA loads in lungs when given at 200 mg/kg twice a day with improved lung histopathology while a mild antiviral effect was observed on starting treatment 1 or 2 days p.i.^[26]

MOLECULAR DOCKING

Enthused by the encouraging results from the literature, we performed the molecular docking using AutoDockVina software to predict the best fit possible biological conformation of molnupiravir in the active site of a SARS-CoV-2 protein. The structure of SARS-CoV-2 (Protein Data Bank [PDB] ID: 6M71) with resolution 2.6Å was downloaded from the PDB. The protein was prepared using AutoDock 4.2.6. The twodimensional (2D) structure of the active form of molnupiravir (NHC) was prepared using ChemDraw professional 16.0 software and converted to three-dimensional (3D) format by Chem3D Ultra 8.0. The energy of molnupiravir was minimized using Chem3D Ultra 8.0 and MOPAC then the pdbqt format of the active form of molnupiravir was prepared using PyMOL. The active form of molnupiravir was docked with active site of SARS-CoV-2 protein. A grid box was prepared with 30, 30, 30 centered (x, y, z) of (114.52, 114.11, 122.91) Å as reported.^[27] Discovery studio visualizer was employed to view the docking results. The active form of molnupiravir displayed good binding energy of -6.4 and hydrogen bond interactions with five amino acids such as Tyr619, Asp760, Asp761, Trp800, and Glu811. The 2D and 3D binding patterns of the active form of molnupiravir are depicted in Figures 4 and 5.

CLINICAL TRIALS

The success of any new molecule depends on the promise shown by it in the clinical trials. A number of such trials involving this drug have been undertaken.

Phase I clinical trial (clinical trial identifier: NCT04392219)

First-in-human randomized, double-blind, placebo-controlled study was conducted in 130 healthy volunteers to evaluate the safety,

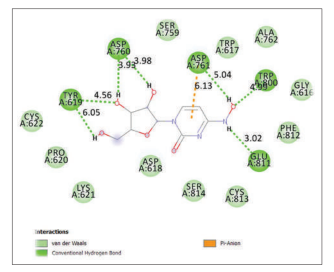


Figure 4:Two-dimensional interaction of target compound with severe acute respiratory syndrome coronavirus-2 protein (PDB ID: 6M71).

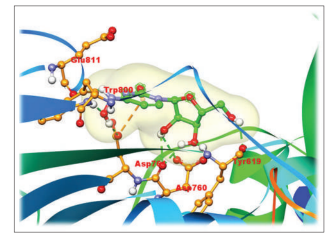


Figure 5: Three-dimensional interaction of target compound with severe acute respiratory syndrome coronavirus-2 protein (PDB ID: 6M71)

tolerability, and pharmacokinetics of orally administered EIDD-2801. For studying drug or placebo in single- and multiple-dose parts of the study, the randomization of eligible volunteers (age: 19-60 years; mean body mass index: 24.4-25.4 kg/m², mostly white, male) was done in a ratio of 3:1. For conducting the trial, each cohort comprising eight subjects was administered with 50-1600 mg single oral dose in single ascending dose part and 50-800 mg b.i.d. for 5.5 days in multiple ascending dose part. The study result displayed a high plasma concentration of EIDD-1931, the hydrolyzed form of prodrug EIDD-2801, possessing a median time of 1.00-1.75 h for maximum observed concentration. The Division of Microbiology and Infectious Diseases toxicity grading study demonstrated headache as the most frequent adverse event reported in 12.5% of molnupiravir administered subjects taking single ascending dose study and diarrhea in 7.1% of molnupiravir administered subjects taking multiple ascending dose study. In food-effect evaluation study also, only mild (grade 1) adverse event was observed. EIDD-2801 was found to be safe, well tolerated with no serious adverse effects. In case of single ascending doses, the administration of EIDD-2801 at doses between

Table 1: Clinical trials of molnupiravir for COVID-19						
NCT number	Sponsor	Brief title	Phase	Reference		
NCT04575584	Merck Sharp & Dohme Corp.	Efficacy and Safety of Molnupiravir (MK- 4482) in Hospitalized Adult Participants With COVID-19 (MK-4482-001)	II/III	[29]		
NCT04575597	Merck Sharp & Dohme Corp.	Efficacy and Safety of Molnupiravir (MK-4482) in Non-Hospitalized Adult Participants With COVID-19 (MK-4482-002)	II/III	[30]		
NCT04405570	Ridgeback Biotherapeutics, LP	A Safety, Tolerability and Efficacy of Molnupiravir (EIDD-2801) to Eliminate Infectious Virus Detection in Persons With COVID-19	IIa	[31]		
NCT04405739	Ridgeback Biotherapeutics, LP	The Safety of Molnupiravir (EIDD- 2801) and Its Effect on Viral Shedding of SARS- CaV 2 (END COVID)	IIa	[13]		

NC104405739	кидераск	The Safety of	lla	[13]
	Biotherapeutics, LP	Molnupiravir (EIDD-		
		2801) and Its Effect on		
		Viral Shedding of SARS-		
		CoV-2 (END-COVID)		
NCT04392219	Ridgeback	COVID-19 First	IIa	[12]
	Biotherapeutics, LP	in Human Study		
		to Evaluate Safety,		
		Tolerability, and		
		Pharmacokinetics of		
		EIDD-2801 in Healthy		
		Volunteers		

600 and 1600 mg gave mean C_{max} value of 13.2 ng/mL and median t_{max} value between 0.25 and 0.75 h. In multiple ascending dose study, a mean t_{1/2} of 7.08 h was observed following 800 mg b.i.d of EIDD-2801.Therefore, molnupiravir displayed good tolerability, when administered in healthy volunteers, with dose-proportional pharmacokinetics.After the success of Phase I clinical trial, few Phase II/III trials also began, as shown in Table 1.^[28]

CONCLUSION

COVID-19 pandemic has led to morbidities of millions of individuals till date. It is spreading worldwide with the emergence of different mutant strains. Several FDA approved drugs are under clinical trials against SARS-CoV-2 infection. Molnupiravir is one such drug which has been approved by FDA for influenza and is now under Phase II/ III trials for SARS-CoV-2. Molnupiravir exerts its antiviral action by causing catastrophic mutations during viral RNA replication using RdRp. The Phase I trial of molnupiravir has proved it to be safe in healthy human volunteers. The pharmacokinetic profile of molnupiravir is equally encouraging. This article has reviewed its chemistry, pharmacology, and the progress through various clinical trials. We have carried out the molecular docking analysis of this drug with RdRp protein of SARS-CoV-2 where the drug demonstrated a high binding affinity involving hydrogen bond interactions with the active site residues. Therefore, our work substantiates the literature claims on the high potential of molnupiravir as an anti-viral agent against SARS-CoV-2 and other such CoV.

REFERENCES

- 1. Fan Y, Zhao K, Shi Z, Zhou P. Bat Coronaviruses in China. Viruses 2019;11:1-14.
- Negi M, Chawla PA, Faruk A, Chawla V. Role of heterocyclic compounds in SARS and SARS CoV-2 pandemic. Bioorg Chem 2020;104:1-36.
- Kong R, Yang G, Xue R, Liu M, Wang F, Hu J, et al. COVID-19 docking server: An interactive server for docking small molecules, peptides and antibodies against potential targets of COVID-19. Bioinformatics 2020;36:5109-111.
- Report of the WHO-China Joint Mission on Coronavirus Disease 2019. Geneva: World Health Organization; 2020. Available from: https://www.who. int/docs/default-source/coronaviruse/who-china-joint-mission-on-covid-19-final-report.pdf. [Last accessed on 2020 Dec 16].
- Andersen KG, Rambaut A, Lipkin WI, Holmes EC, Garry RF. Correspondence: The proximal origin of SARS-CoV-2. Nat Med 2020;26:450-2.
- Liu X, Li Z, Liu S, Sun J, Chen Z, Jiang M, *et al.* Potential therapeutic effects of dipyridamole in the severely ill patients with COVID-19. Acta Pharm Sin B 2020;811:1-11.
- Shah B, Modi P, Sagar SR. *In silico* studies on therapeutic agents for COVID-19: Drug repurposing approach. Life Sci 2020;252:1-12.
- Thakur S, Sarkar BM, Ansari AJ, Khandelwal A, Arya A, Poduri R, et al. Exploring the magic bullets to identify Achilles' heel in SARS-CoV-2: Delving deeper into the sea of possible therapeutic options in COVID-19 disease: An update. Food Chem Toxicol 2021;147:1-21.
- Wang Y, Anirudhan V, Du R, Cui Q, Rong L. RNA-dependent RNA polymerase of SARS-CoV-2 as a therapeutic target. J Med Virol 2021;93:300-10.
- Toots M, Yoon J, Cox RM, Hart M, Sticher ZM, Makhsous N, et al. Characterization of orally efficacious influenza drug with high resistance barrier in ferrets and human airway epithelia. Sci Transl Med 2019;11:1-13.
- Toots M, Yoon J, Hart M, Natchus MG, Painter GR, Plemper RK. Quantitative efficacy paradigms of the influenza clinical drug candidate EIDD-2801 in the ferret model. Transl Res 2020;218:16-28.
- COVID-19 First in Human Study to Evaluate Safety, Tolerability, and Pharmacokinetics of EIDD-2801 in Healthy Volunteers; 2020. Available from: https://www.clinicaltrials.gov/ct2/show/nct04392219. [Last accessed on 2020 Dec 16].
- Efficacy and Safety of Molnupiravir (MK-4482) in Hospitalized Adult Participants with COVID-19 (MK-4482-001); 2020. Available from: https:// www.clinicaltrials.gov/ct2/show/nct04405739?term=molnupiravir and cond=covid19 and draw=2 and rank=4. [Last accessed on 2020 Dec 16].
- Cox RM, Wolf JD, Plemper RK. Therapeutically administered ribonucleoside analogue MK-4482/EIDD-2801 blocks SARS-CoV-2 transmission in ferrets. Nat Microbiol 2021;6:11-8.
- Cevik M, Kuppalli K, Kindrachuk J, Peiris M. Virology, transmission, and pathogenesis of SARS-CoV-2. BMJ 2020;371:1-6.
- Shang Z, Chan SY, Liu WJ, Li P, Huang W. Recent insights into emerging Coronavirus: SARS-CoV2. ACS Infect Dis 2020.
- Forstera P, Forsterd L, Renfrewb C, Forsterc M. Phylogenetic network analysis of SARS-CoV-2 genomes. Proc Natl Acad Sci U S A 2020;117:9241-3.
- Azer SA. COVID-19: Pathophysiology, diagnosis, complications and investigational therapeutics. New Microbes New Infect 2020;37:100738.
- Yuki K, Fujiogi M, Koutsogiannaki S. COVID-19 pathophysiology: A review. Clin Immunol 2020;215:108427.
- Vallamkondu J, John A, Wani WY, Ramadevi RS, Jella KK, Reddy PH, et al. SARS-CoV-2 pathophysiology and assessment of Coronaviruses in CNS diseases with a focus on therapeutic targets. Biochim Biophys Acta Mol Basis Dis 2020;1866:165889.
- Parasher A. COVID-19: Current understanding of its pathophysiology, clinical presentation and treatment. Postgrad Med J 2020;97:312-20.
- Hampton T. New flu antiviral candidate may thwart drug resistance. JAMA 2020;323:17.

- Steiner A, Znidar D, Otvos SB, Snead DR, Dallinger D, Kappe CO. A highyielding synthesis of EIDD-2801 from uridine. European J Org Chem 2020;22:6736-9.
- Urakova N, KuznetsovaV, Crossman DK, Sokratian A, Guthrie DB, Kolykhalov AA, et al. β-D-N4-Hydroxycytidine is a potent anti-alphavirus compound that induces a high level of mutations in the viral genome. J Virol 2018;92:e01965-17.
- Sheahan TP, Sims AC, Zhou S, Graham RL, Pruijssers AJ, Agostini ML, et al. An orally bioavailable broad-spectrum antiviral inhibits SARS-CoV-2 in human airway epithelial cell cultures and multiple coronaviruses in mice. Sci Transl Med 2020;12:eabb5883.
- Abdelnabi R, Foo CS, Kaptein SJ, Zhang X, Langendries L, Vangeel L, et al. Molnupiravir (EIDD-2801) Inhibits SARS-CoV2 Replication in Syrian Hamsters Model, bioRxiv; 2020.
- Ahmad J, Ikram S, Ahmad F, Rehman IU, Mushtaq M. SARS-CoV-2 RNA dependent RNA polymerase (RdRp)-a drug repurposing study. Heliyon 2020;6:e04502.

- Painter WP, Holman W, Bush JA, Almazedi F, Malik H, Eraut NC, et al. Human Safety, Tolerability, and Pharmacokinetics of a Novel Broad-Spectrum Oral Antiviral Compound, Molnupiravir, with Activity Against SARS-CoV-2, medRxiv; 2020.
- 29. Efficacy and Safety of Molnupiravir (MK-4482) in Hospitalized Adult Participants with COVID-19 (MK-4482-001); 2020. Available from: https:// www.clinicaltrials.gov/ct2/show/nct04575584?term=molnupiravir and cond=covid19 and draw=2 and rank=1. [Last accessed on 2020 Dec 16].
- 30. Efficacy and Safety of Molnupiravir (MK-4482) in Non-Hospitalized Adult Participants with COVID-19 (MK-4482-002); 2020. Available from: https:// www.clinicaltrials.gov/ct2/show/nct04575597?term=molnupiravir and cond=covid19 and draw=2 and rank=2. [Last accessed on 2020 Dec 16].
- 31. A Safety, Tolerability and Efficacy of Molnupiravir (EIDD-2801) to Eliminate Infectious Virus Detection in Persons; 2020. Available from: https:// www.clinicaltrials.gov/ct2/show/nct04405570?term=molnupiravir and cond=covid19 and draw=2 and rank=3. [Last accessed on 2020 Dec 16].



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The Role of 4-Thiazolidinone Scaffold in Targeting Variable Biomarkers and Pathways Involving Cancer

Meenakshi Negi¹, Pooja Chawla², Abdul Faruk¹, Viney Chawla³

Affiliations PMID: 34229596 DOI: 10.2174/1871520621666210706104227

Abstract

Background: Cancer can be considered as a genetic as well as a metabolic disorder. The current cancer treatment scenario looks like aggravating tumor cell metabolism, causing the disease to progress even with greater intensity. The cancer therapy is restricted to the limitations of poor patient compliance due to toxicities to normal tissues and multi-drug resistance development. There is an emerging need for cancer therapy to be more focused towards better understanding of genetic, epigenetic and transcriptional changes resulting in cancer progression and their relationship with treatment sensitivity.

Objective: The 4-thiazolidinone nucleus possesses marked anticancer potential towards different biotargets, thus targeting different cancer types like breast, prostate, lung, colorectal and colon cancers, renal cell adenocarcinomas and gliomas. Therefore, conjugating the 4-thiazolidinone scaffold with other promising moieties or directing the therapy towards targeted drug delivery systems like the use of nanocarrier systems, can provide the gateway for optimizing the anticancer efficiency and minimizing the adverse effects and drug resistance development, thus providing stimulus for personalized pharmacotherapy.

Methods: An exhaustive literature survey has been done to give an insight into the anticancer potential of the 4- thiazolidinone nucleus either alone or in conjugation with other active moieties, with the mechanisms involved in preventing proliferation and metastasis of cancer covering a vast range of publications of repute.

Conclusion: This review aims to summarise the work reported on anticancer activity of 4thiazolidinone derivatives covering various cancer biomarkers and pathways involved, citing the data from the year 2005 till now, which may be beneficial to the researchers for future development of more efficient 4-thiazolidinone derivatives.

Keywords: 4-Thiazolidinone; biotargets; cancer; cytotoxicity; genetic; transcriptional.

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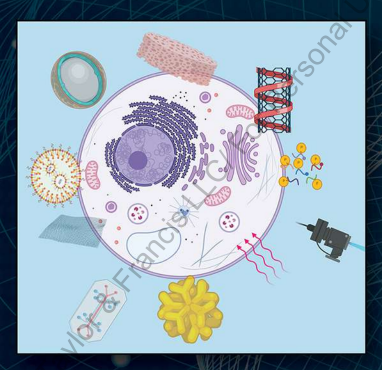
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NANOPHARMACEUTICALS IN REGENERATIVE MEDICINE



EDITED BY HARISHKUMAR MADHYASTHA DURGESH NANDINI CHAUHAN



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Nanopharmaceuticals in Regenerative Medicine

Edited by Harishkumar Madhyastha and Durgesh Nandini Chauhan



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Foreword



Nanopharmaceutics is a branch of nanobiotechnology with vast applications in diagnostics, regenerative medicine, and drug development in current science of medicine. Within a short span of two decades, the subject has expanded into a promising arena for clinical and translational medicine. The biomedical scientists show immense interest in nanomaterials due to their extraordinary surface to volume area, tunable optical emission, unique electrical, and magnetic behaviour, which particularly helps in drug discovery research. The hybridisation of nanotechnology and tissue regeneration will open a new path of innovation and will have potential application to treat incurable diseases. The book '*Nanopharmaceuticals in regenerative medicine*' is an informative compilation of nanomedicine, combining description of pharmaceutical formulations and their mechanisms of action. The book provides the comprehensive bundle of information and accurate scientific information on nanopharmaceutical use in regenerative medicine and would be

epochal to the scientific community, especially clinicians and pharmacists.

I applaud the editors, Dr. Harishkumar Madhyastha who has been my colleague for many years at University of Miyazaki and Smt. Durgesh Nandini Chauhan for the excellent compilation of chapters contributed by well-known scientists and academicians from different countries. All 18 chapters are different from each other in content, but share a single objective of nanopharmaceutical advancement. The most notable chapters include therapeutic applications, technological innovations, and tissue regeneration. The authors successfully navigate the chapter contents with updated literature. I believe '*Nanopharmaceuticals in regenerative medicine*' will remain a valuable resource for years to come.

Prof. Dr. Tsuyomu Ikenoue, MD. Ph.D.

President University of Miyazaki Miyazaki-Japan.

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Preface

Trajectory of scientific thoughts is propelling rapidly through good research communications. Research ideas will be broadcasted through good publications which are mainly dispersed by review manuscripts, book chapters, etc. A comprehensive scientific dissertation serves as a satellite stop reference book for budding academicians, scientists, professionals, and technologists. With extensive and annotated knowledge and information, the book is a gateway for knowledge dispersion and escalation, community curation, and finally betterment of society. With the advancement of scientific knowledge, a new paradigm of science, nanobiotechnology, is emerging in the area of biomedical science and regenerative medicine. In regenerative medicine arena, nanotechnologies have wide and high-impact benefits like drug development, diagnostics, and delivery system. This book provides an in-depth knowledge on applied nanobiomedical contents for university graduates, researchers, and technocrats with striking balance between fundamentals and applications for regenerative medicine. The book contains 18 chapters covering a wide range of topics related to chemistry, pharmacy, and material science. The chapters are broadly classified into three categories; potential insights into smart technologies, interpretations of different modes as delivery systems, and tissue engineering and generation aspects. Each chapter includes multidisciplinary approaches and recommendations to use the nanotechnologies for tissue regeneration with meaningful conclusions and attracts new ideas for future development. Chapters 1-5 emphasize the applications of nanoparticles in regenerative therapy. Chapters 6-12 focus on different technological approaches devoted to tissue recalcitrant engineering. Chapters 13-18 elucidate the updates on nanomaterials in the field of tissue regeneration, with special focus on osteoporosis, cancer, and cardiology with a pharmaceutical angle. Date: 16 April 2021

> Dr. Harishkumar Madhyastha Durgesh Nandini Chauhan

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Editors



Dr. Harishkumar Madhyastha Ph.D., FBRSI. Harishkumar Madhyastha, faculty at Department of Applied Physiology University of Miyazaki, Miyazaki, Japan. With two Ph.D. degrees, he ignited his research career as a scientist in *Spirulina* biotechnology at MCRC-Chennai. Later on, he pursued postdoctoral research at Miyazaki University that culminated in a faculty position in the Department of Applied Physiology at the University of Miyazaki from 2006. His current research interests include generation and delivery of nanosized metallic payloads for regenerative diseases application. His academic credentials are credited with more than 80 *Sci-E* indexed papers; *h*-value of 29, clarivate analytic cumulative impact factor of 204.5 and RG score of 33.76 with *six* international patents. His research has been presented in conferences more than 100 and has been frequently picked up by national and

international media. He is also actively involved in many international projects including ongoing Indo-Japan scientific and academic collaborations. He is Fellow of Biotechnology Research Society of India (FBRSI), Fellow of Royal Biological Society-London (FRBS-UK). He is an officially recognised Indo-Japan academic spokesperson of University of Miyazaki and engaged in outreach programs to further strengthen the cohesive relationship between Indian academicians and University of Miyazaki-Japan.



Mrs. Durgesh Nandini Chauhan, M.Pharma, has completed her B.Pharm degree in Pharmacy from the Rajiv Gandhi Proudyogiki Vishwavidyalaya, Bhopal, India and her M.Pharma in pharmaceutics from Uttar Pradesh Technical University, currently known as Dr. A.P.J. Abdul Kalam Technical University, Lucknow in 2006. She is presently working as Assistant Professor in Columbia Institute of Pharmacy, Raipur, Chhattisgarh, India. Mrs. Durgesh Nandini Chauhan has 14 years of academic (teaching) experience from Institutes of India in pharmaceutical sciences. She taught subjects as pharmaceutics, pharmacognosy, traditional concepts of medicinal plants, drug-delivery phytochemistry, cosmetic technology, pharmaceutical engineering, pharmaceutical packaging, quality assurance, dosage form designing and anatomy, and physiology.

She is member of Association of Pharmaceutical Teachers of India (APTI), SILAE: Società Italo-Latinoamericana di Etnomedicina (The

Scientific Network on Ethnomedicine, Italy), and so forth. She has written more than 10 publications in national and international journals, 16 book chapters, and has edited 7 books. She is also active as a reviewer for several international scientific journals and active participant in national and international conferences.

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Extracellular Matrix: The State of the Art in ial use only Regenerative Medicine

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Introduction

Regenerative medicine gained significant interest in the treatment of life-threatening diseases and disorders, especially in cardiovascular and neurodegenerative diseases (Mao and Mooney 2015). It is a multidisciplinary approach, which restores the normal physiological functions of the human body by replacement or repair of tissues and organs (Christ et al. 2013). Regenerative medicines are innovative therapies that involve various strategies of tissue engineering, stem cell biology, gene, and cellular therapeutics (Lorden et al. 2015). All regenerative medicine approaches depend upon cellular level events and their constituents, which are involved in various developmental or repair processes of human tissues, i.e. replacing damaged cells in the brain and pancreas (Mao and Mooney 2015). These transplanted cells perform all normal functions and functionally participate in the all tissue events (Chen and Liu 2016). Presently, regenerative medicine-based treatment is very expensive and not affordable by all (Mahalatchimy 2016).

Regenerative medicine is defined as a cellular therapeutic approach which "substitutes or repair human cells, various tissues or organ systems, to restore normal physiological function of human body" (Han et al. 2020; Sampogna et al. 2015).

There are a number of regulatory issues that influence the development of regenerative medicine and thus, in this scenario, need additional focus on legislation for regenerative medicine (Kleiderman et al. 2018). Recent research reports suggested that stem cell-based therapy has a promising role in the treatment of deadly human diseases, i.e. leukaemia, breast cancer, and others (Aly 2020). The ultimate objective of regenerative medicine is the isolation of specialised cell constituents and implanted into a patient where it replaces or repairs damage part of tissue or cells through self-repair remodelling (Mao and Mooney 2015). Therefore, it regulates the functioning of native tissues or cells. It offers transformative and effective outcomes for targeting life-threatening acute and chronic conditions and also an alternative for degenerative and genetic disorders (Mahla 2016).

According to the status of the Global Regenerative Medicine Market forecast, the international market of regenerative medicine is continuously growing and expected to reach USD 17.9 billion by 2025 (marketsandmarkets 2020). Food and Drug Administration (FDA, United States) implemented the 21st Century Cures Act in 2016 for the regulation of regenerative medicine therapies under a special section 3033, which describes the term and conditions for designation of drug under Regenerative Medicine Advanced Therapy (RMAT) (Barlas 2018). The Cures Act improves the ability of scientific, technical, and professional experts regarding clinical trial designs for regenerative medicine. It will accelerate the production of regenerative medicine products with safety of patients (FDA 2020).

The Cures Act defines the regenerative medicine as:

cell therapy, therapeutic tissue engineering product, human cell and tissue product, or any combination product using such therapies or products intended to treat, modify, reverse, or cure a serious or life threatening disease.

(FDA 2021b)

There are currently four main categories of stem cells that have the clone ability and differentiate into particular types of cells.

- i. **Embryonic stem cells:** Derived from the initial developmental phase of few days old embryos at the blastocyst stage. It has the potential to differentiate into various cells with a distinct biological response. Such cells are known as pluripotent (Romito and Cobellis 2016).
- ii. Foetal stem cells: Isolated from aborted human foetuses, especially foetal blood, foetal tissues, and also bone marrow. They have the ability to differentiate but not all cells. They are known as multipotent and have been utilised in the regeneration and repair of damaged tissues/organs (Biehl and Russell 2009).
- iii. Cord blood and placental stem cells: Obtained from umbilical cord blood and placentas. They possess the therapeutic potential and used in bone-marrow replacement therapies. They are not able to differentiate into all types of cells (Weiss and Troyer 2006).
- iv. Adult stem cells: They are the most abundant cells, which are used for various therapies/conditions. They are isolated from almost all human tissue and organs. They are known as "somatic stem cells" (Liras 2010).

There is no doubt that regenerative medicine products provide a better treatment option than conventional drugs. But still, there are certain limitations and challenges for researchers and pharmaceutical companies that need to be addressed for the improvement of these specialised products (Dodson and Levine 2015). The following are the few noticeable points that should be considered during the design and production of regenerative medicine (Herberts et al. 2011):

i. **Safety:** The derived product should be safe and effective without any tumour formation or production of unwanted cell types.

- ii. **Regulatory aspects and standardisation:** Must meet regulatory requirements which ensure product quality, safety, and efficacy as mention by standards (Rosemann et al. 2019).
- iii. Imaging and Monitoring: Need sophisticated techniques with the features of observing all the changes and variation during cell behaviour (Leahy et al. 2016) and also, monitoring the migration of cells after administration (Naumova et al. 2014).
- iv. **Manufacturing:** Manufacturing of viable (living) cells for regenerative medicine must follow through optimised process protocol to avoid cell variability (Martin et al. 2014).
- v. Multidisciplinary research involves in regenerative medicine requiring effective communication within all research communities for better outcomes (Shineha et al. 2017).
- vi. **Animal Models:** Appropriate animal models are needed for the comparison of animal embryos/ human genetic or cellular material information (Ribitsch et al. 2020).
- vii. Scale up/Technology Transfer: Large-scale production reduces the overall cost of the product. The scalable production processes provide safe and effective products (Pigeau et al. 2018).
- viii. Immunogenicity: In regenerative therapies, a major issue is the rejection of transplanted cell by the patient. This could be overcome by exploring the research for new generation of immunosuppressant drugs (Charron 2013).
- ix. Cell Viability: Cell viability and storage conditions (Yu et al. 2018).

A number of regenerative medicine which have already received FDA approval (FDA 2021a) and are commercially available are listed in Figure 9.1. This chapter explores the role of extracellular matrix (ECM) in regenerative medicine.

The Extracellular Matrix

Regenerative strategies mainly focus on stem cell-based or tissue engineering applications for remodelling and regeneration of defective cells, tissues, and organs. Stem cell differentiation is modulated by signals from the extracellular microenvironment including the extracellular matrix (ECM) (Chen and Liu 2016). Cellular migration and differentiation events are the main key factors that are considered for the design of regenerative medicine (Mata et al. 2017). The ECM is composed of several types of collagens, proteoglycans, glycoproteins, and glycosaminoglycans, which are assembled into a complex structure (Yue 2014). The composition of ECM varies from tissue to tissue and organ to organ (Kular et al. 2014). The distinctive functions of the ECM include cell adhesion, the physical barrier for different tissues. It also impacts many cellular functions, including mechanical stimulation from substrates, activation of intracellular signalling by cell adhesion molecules, and availability and action of soluble factor (Muncie and Weaver 2018).

The extracellular matrixes (ECM) define the tissue architecture and biochemical and biophysical features. The main organisational unit of the ECM called core matrisome, which includes different kinds of collagen (divided into several families), glycoproteins, and proteoglycans (Hynes and Naba 2012). Other than ECM, there are numerous non-ECM varying factors, which also participate in different cellular events, i.e. remodelling and cell behaviour. They mainly include proteases, growth factors, cytokines, and cross-linking enzymes (Vaday and Lider 2000). Collagen is the most abundant protein of mammalian ECM and accountable for the structural and functional integrity of the tissue (Frantz et al. 2010). Other structural molecules of ECM belong to the glycosaminoglycans class which includes hyaluronic acid (HA, non-sulphated glycosaminoglycan), chondroitin sulphate (CS, sulphated glycosaminoglycan), and heparin (natural glycosaminoglycan) Figure 9.2 (Pomin and Mulloy 2018). They play a vital role in elasticity, water retention, and resistance to compressive forces, while adhesion proteins play a significant role as molecular glue for a structural network of ECM complex. Examples of adhesion molecules are laminin, fibronectin, and tenascin-C (Walker et al. 2018).

Early in the 20th century, cell biologists worked in a two-dimensional (2D) framework, which includes separating and culturing cells from living tissue for replacing damaged or diseased tissue (O'Brien and Duffy 2015). With the advancement in the field of bioengineering and regenerative medicine, it is

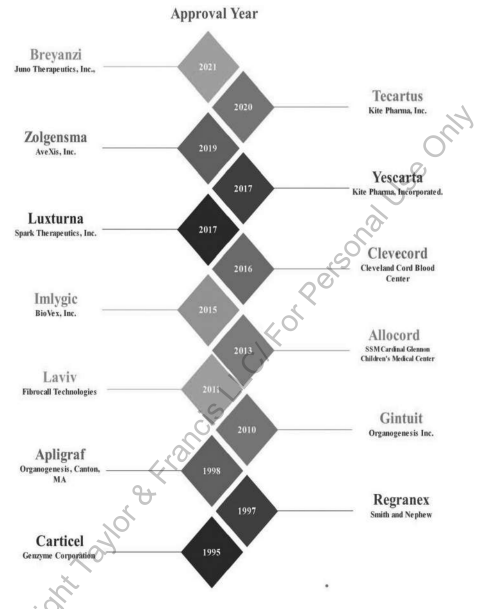


FIGURE 9.1 List of FDA Approved Products.

observed that multicellular organisms require a three-dimensional (3D) framework for structural integrity with specific microenvironments (Chen and Liu 2016). It is required to incorporate the knowledge of cell biology and cell transplantation with the discipline of material science for providing a 3D environment for growing cells and tissues. The evolution in the medical field has opened a new horizon for use of ECM in regenerative medicine. Various ECM analogues have been developed from synthetic scaffolds (Nikolova and Chavali 2019), hydrogels, and ceramic-based scaffolds (Hussey et al. 2018). These scaffolds are commonly made-up of synthetic and biodegradable polymers (Chaudhari et al. 2016). Commonly used polymers include polycaprolactone, polyethylene glycol, polyacrylic acid, hydroxyapatite or tricalcium phosphate, alginate, chitosan, and cellulose derived from plants (Hussey et al. 2018). Biomaterials used in regenerative medicine are broadly classified into two groups, i.e. naturally obtained and synthetic

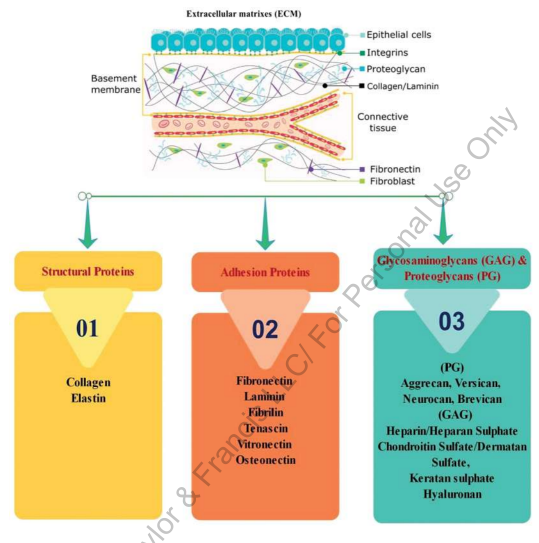


FIGURE 9.2 Composition of ECM (Kular et al. 2014).

materials (Fernandes et al. 2009). Natural materials are generally extracted or purified from ECM or its components such as collagen, laminin, and fibronectin (Frantz et al. 2010). Synthetic materials include polymers, metals or derived substrates. Both synthetic and natural materials have distinct pros and cons in regenerative medicine. Ideally, they are selected on the basis of condition and requirement of treatment. Biomaterials isolated from ECM show more unpredictability than synthetic polymers. In the case of synthetic polymers, immune response and their antigenicity is the major issue (Chen and Liu 2016). New trends and technologies in the bioengineering field reveal the functions of the ECM in regenerative medicine. This enriched the knowledge of ECM signalling in the functions of stem cells. These outcomes revealed the use of synthetic ECM scaffolds, which promote the endogenous stem cell repair and healing of damaged cells/tissues and mimic the native microenvironment (Chan and Leong 2008). A list of potential components of the extracellular matrix which are utilised in regenerative medicine (Traphagen and Yelick 2009) are summarised in Figures 9.3 and 9.4.

ECM-based biomaterials promote tissue remodelling in a precise and controllable manner. The decellularised ECM (DECM), which is water-insoluble matrix obtained after removal of cellular constituents

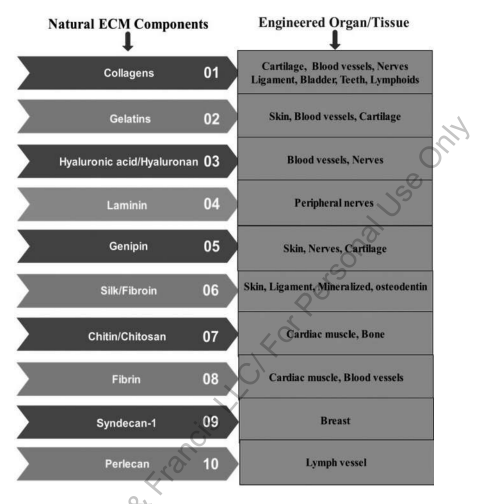


FIGURE 9.3 List of Engineered Organs and Tissues Based on ECM.

from ECM, also plays a significant role in the remodelling and repair process (Chakraborty et al. 2020). Due to biocompatibility and biodegradability, the DECM offers better results than other commonly used biomaterials (Liao et al. 2020). DECM-based tissue/organ, hydrogel, and microparticles have high demand in regenerative medicine (Parmaksiz et al. 2016).

Biomimetic materials can be fabricated using different techniques, i.e. soft lithography (Whitesides et al. 2001) (micro-contact printing), electrospinning (Braghirolli et al. 2014), and 3D printing (Atala and Forgaes 2019). Cellular constituents present within all tissues are required for tissue morphogenesis, differentiation, and the homeostasis process. Fundamentally, ECM can resolve various syndromes, physiological conditions, and defects in the body (Theocharis et al. 2019). In recent years, many studies indicate the role of native ECMs/DECM in regenerative medicine (Ramos and Moroni 2020). The main applications of ECMs include 3D tissue culturing (Edmondson et al. 2014), stimulate the wound healing process (Agren and Werthen 2007), activate stem cell differentiation (Gattazzo et al. 2014), and drug screening assays (Langhans 2018). It's also applied in cell repair pathways and functional recovery of kidney (Bulow and Boor 2019), adrenal glands (Ruiz-Babot et al. 2015), and reproductive organs (Yalcinkaya et al. 2014). ECMs have many applications due to their biocompatibility and *in vivo* replicate ability (Aamodt and Grainger 2016). This chapter summarises some research investigations based on EMCs in regenerative medicine.

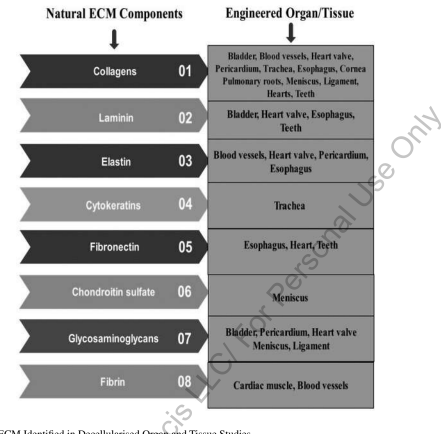


FIGURE 9.4 ECM Identified in Decellularised Organ and Tissue Studies.

Application of Extracellular Matrix

The chief proteins of the ECM are collagens and elastin. They are considered for biomedical applications because of their tensile strength and viscoelasticity to tissues. Other proteins include fibronectin, laminin, and nidogen, which act as connectors or linking proteins in the matrix network. Glycosaminoglycans (GAGs), proteoglycans, and growth factors are also promoting the *in vivo* construction of functional tissue (Mouw et al. 2014). Overall it is a challenging task due to limited knowledge and tissue to tissue variability. Ultimate goals of regenerative medicine can be achieved only if biomaterials maintain desired morphology, differentiation, proliferation, and metabolism of the cell.

Cardiac Extracellular Matrix

The heart is a vital and delicate organ of our body that requires sophisticated tissue architecture for normal functioning. It acts as a circulatory motor for our body that supplies the blood and fulfils the variable demands during the rest and exercise phase (Lee and Walsh 2016). This excitation-contraction, the cycle of the heart, developed a physical force at the cellular level of the myocardial structure. Overall, this process is regulated by the delicate organisation of the cardiac extracellular matrix. In each excitation and contraction cycle, a number of mechanical events are involved in myocardial elements (Stoppel et al. 2016).

Recent investigations suggest that ECM is found in all the segments of the heart. However, it is particularly present in mesenchyme structures and plays a role in valvuloseptal morphogenesis (Lockhart et al. 2011). Any impairment in the composition of ECM in mesenchyme structures often leads to congenital heart disease. Several reported animal studies describe the involvement of ECM in congenital heart diseases (Hacker 2018). Studies indicate the involvement of aggrecan, hyaluronan, versican, collagen type I-V, fibulin1, and fibronectin (Wight 2018). The common complications are vascular defects, blood vessel rupture, and cardiomyopathy. ECM involves in the regulation of cell differentiation and proliferation, which serve the cell survival (Ponticos and Smith 2014).

Extracellular Matrix in Brain

A major part of the brain is occupied by ECM, which contains collagens, fibronectin, vitronectin, laminin, and perlecan especially in amyloid deposits of the brain (Bonneh-Barkay and Wiley 2009). These ECM components play the main role in the development of nervous tissue and also regulate cell adhesion (Barros et al. 2011). Matrix proteins are almost absent in the adult brain (Ruoslahti 1996). Any change that occurs in the composition of ECM after neural injury may result in drastic consequences. Brain injury may induce changes in chondroitin sulphate proteoglycans, which influence myelin repair (Rhodes and Fawcett 2004). During the early stage of neural growth, the ECM provides structural support and stimulates signalling pathways of proliferation, especially by proteoglycans, laminins, and integrins. Proteoglycans provide structural support while laminins and integrins enhance neural progenitor proliferation (Bonnans et al. 2014). They also modify the shape of neural progenitors and neurons. In addition to this, ECM components affect the migration of newborn neurons during cortical growth. The role of the ECM in the brain is highly complicated (Lu et al. 2011). The same ECM component performs multiple roles during neural development and also influences the functioning of neighbouring cells (Rozario and DeSimone 2010). Recently reported evidence indicates the involvement of the ECM in several disease conditions, such as traumatic brain injury (George and Geller 2018), Alzheimer's disease (Lepelletier et al. 2017), age-associated cognitive deficits (Richard and Lu 2019), and schizophrenia (Lubbers et al. 2014). ECM-based regenerative approaches are widely used in the repair of peripheral soft tissue but not in the case of the brain due to the invasive route of administration. It requires a very specific narrow needle-guided administration approach for specific targeting. Current research efforts in regenerative medicine suggest that ECM-based biomaterials could serve as regenerative therapies in the brain (Hwang et al. 2020). A variety of underlying factors and mechanisms are still under observation and site-specific administration of ECM-based biomaterials is another issue in development of regenerative medicine (Chen and Liu 2016).

Pulmonary Extracellular Matrix

Pulmonary ECM is a structural complex system of protein molecules, which participate in various biochemical processes (Burgstaller et al. 2017). The remodelling mechanism is important for tissue homeostasis and any change in it may result in conditions like chronic obstructive pulmonary disease (COPD). Impaired expression of ECM proteins seen in COPD leads to the degradation and disruption of alveolar walls and stiffening of minor airways, which result in obstruction of airways (Ito et al. 2019). Alterations in ECM composition also influence the immune cell movement and their maintenance in the lung (Bonnans et al. 2014). Any abnormal functioning of ECM and response of inflammatory cell surface receptors may modify the collagen microstructure of the lung (Hussell et al. 2018). It is observed that there is a change in collagen organisation in COPD lung as compared to normal lung. The imbalance of enzymes like lysyloxidase and transglutaminase2 may involve structural changes of ECM during COPD (Burgess et al. 2016). ECM regulates normal interstitial fluid dynamics and strength and elasticity, tissue repair, and remodelling in the lungs. Versican and perlecan participate in the balancing of tissue fluid homeostasis (Pelosi et al. 2007). In the area of regenerative medicine, several studies reported lung scaffolds from small and large animals as an alternative to lung transplantation (Ohata and Ott 2020). These lung scaffolds were decellularised and reseed with lung perfusion culture in bioreactors. The resulting bioartificial lungs are probable to solve the problem of donor organ shortage and also reduced the immunogenicity (Panoskaltsis-Mortari 2015).

Extracellular Matrix in Inflammatory Bowel Disease

Inflammatory Bowel Disease (IBD) is a global health issue and the specific aetiology of IBD is unknown. Ulcerative colitis and Crohn's disease are the two main forms of IBD and they are characterised by an unusual immune response linked with defects functioning in the intestinal epithelial cell barrier (Zhang and Li 2014). Macroscopic tissue injury and clinical features of IBD are developed by changes in the ECM. Any change in ECM constituents may result in intestinal inflammation and progression of IBD (Petrey and de la Motte 2017). The conventional treatments merely target treatment of inflammation not repair/recovery of damaged tissue. Recently published work reports the use of hematopoietic or mesenchymal stem cells (HSCs or MSCs) for the management of IBD (Martinez-Montiel Mdel et al. 2014). It may help to establish an effective regenerative medicine for IBD patients. The development of decellularisation techniques in biomedical engineering greatly assisted the site-specific applications of ECM bio-scaffolds in the gastrointestinal tract (Almeida-Porada et al. 2013).

Conclusion

In conclusion, it is clear that the manipulation of ECM may serve as natural mimicking scaffolds in the arena of regenerative medicine. Regenerative medicine will change the traditional methods of management of various life-threatening diseases and conditions. Moreover, there is no doubt that all classes of stem cells (embryonic, adult, and induced pluripotent stem cells) have the potential to control the variety of diseases. ECM and ECM-like materials are biocompatible and having integration with the physiological microenvironment and mimic the ECM structure of the target tissues. ECM supports various biological functions and preserves the structures of entire organs. Ideally, they are preferred over synthetic polymers for biomedical engineering because of their immune tolerance. ECM regulation can play a significant role in several body conditions such as COPD, spinal cord injury, and neurodegenerative disorder. Furthermore, innovative interdisciplinary approaches and advancements in methodologies may lead to the improvement and discovery of new treatments for human disease.

REFERENCES

Aamodt J.M., Grainger D.W. 2016. Biomaterials 86: 68-82. doi:10.1016/j.biomaterials.2016.02.003.

- Agren M.S., Werthen M. 2007. The International Journal of Lower Extremity Wounds 6 (2): 82–97. doi:10.1177/1534734607301394.
- Almeida-Porada G., Soland M., Boura J., Porada C.D. 2013. Regenerative Medicine 8 (5): 631–644. doi:10.2217/rme.13.52.
- Aly R.M. 2020. Stem Cell Investigation 7: 8. doi:10.21037/sci-2020-001.
- Atala A., Forgacs G. 2019. Stem Cells Translational Medicine 8 (8): 744-745. doi:10.1002/sctm.19-0089.
- Barlas S. 2018, P & T: A Peer-Reviewed Journal for Formulary Management 43 (3): 149–179.
- Barros C.S., Franco S.J., Muller U. 2011. Cold Spring Harbor Perspectives in Biology 3 (1): a005108. doi:10.1101/cshperspect.a005108.
- Biehl J.K., Russell B. 2009. *The Journal of Cardiovascular Nursing 24* (2): 98–103; quiz 104–105. doi:10.1097/ JCN.0b013e318197a6a5.
- Bonnans C., Chou J., Werb Z. 2014. Nature Reviews. Molecular Cell Biology 15 (12): 786–801. doi:10.1038/ nrm3904.
- Bonneh-Barkay D., Wiley C.A. 2009. Brain Pathology 19 (4): 573-585. doi:10.1111/j.1750-3639.2008.00195.x.
- Braghirolli D.I., Steffens D., Pranke P. 2014. Drug Discovery Today 19 (6): 743–753. doi:10.1016/j. drudis.2014.03.024.
- Bulow R.D., Boor P. 2019. The Journal of Histochemistry and Cytochemistry: Official Journal of the Histochemistry Society 67 (9): 643–661. doi:10.1369/0022155419849388.
- Burgess J.K., Mauad T., Tjin G., Karlsson J.C. et al. 2016. *The Journal of Pathology 240* (4): 397–409. doi:10.1002/path.4808.

- Burgstaller G., Oehrle B., Gerckens M., White E.S. et al. 2017. *The European Respiratory Journal 50* (1). doi:10.1183/13993003.01805-2016.
- Chakraborty J., Roy S., Ghosh S. 2020. Biomaterials Science 8 (5): 1194–1215. doi:10.1039/C9BM01780A.

Chan B.P., Leong K.W. 2008. *European Spine Journal 17* Suppl 4: 467–479. doi:10.1007/s00586-008-0745-3. Charron D. 2013. *The Indian Journal of Medical Research 138* (5): 749–754.

- Chaudhari A.A., Vig K., Baganizi D.R., Sahu R. et al. 2016. International Journal of Molecular Sciences 17 (12): 1974.
- Chen F.M., Liu X. 2016. Progress in Polymer Science 53: 86–168. doi:10.1016/j.progpolymsci.2015.02.004.
- Christ G.J., Saul J.M., Furth M.E., Andersson K.E. 2013. *Pharmacological Reviews* 65 (3): 1091–1133. doi:10.1124/pr.112.007393.
- Dodson B.P., Levine A.D. 2015. BMC Biotechnology 15: 70. doi:10.1186/s12896-015-0190-4.
- Edmondson R., Broglie J.J., Adcock A.F., Yang L. 2014. Assay and Drug Development Technologies 12 (4): 207–218. doi:10.1089/adt.2014.573.
- FDA. 2020. 21st Century Cures Act. Retrieved 5/2/2021, 2021, from https://www.fda.gov/regulatory-information/selected-amendments-fdc-act/21st-century-cures-act
- FDA. 2021a. Approved Cellular and Gene Therapy Products. Retrieved 3 March 2021, 2021, from https://www. fda.gov/vaccines-blood-biologics/cellular-gene-therapy-products/approved-cellular-and-gene-therapyproducts
- FDA. 2021b. Regenerative Medicine Advanced Therapy Designation. Retrieved 20/2/2021, from https://www. fda.gov/vaccines-blood-biologics/cellular-gene-therapy-products/regenerative-medicine-advanced-therapy-designation
- Fernandes H., Moroni L., van Blitterswijk C., de Boer J. 2009. Journal of Materials Chemistry 19 (31): 5474–5484. doi:10.1039/B822177D.
- Frantz C., Stewart K.M., Weaver V.M. 2010. Journal of Cell Science 123 (Pt 24): 4195–4200. doi:10.1242/ jcs.023820.
- Gattazzo F., Urciuolo A., Bonaldo P. 2014. *Biochimica et biophysica acta 1840* (8): 2506–2519. doi:10.1016/j. bbagen.2014.01.010.
- George N., Geller H.M. 2018. Journal of Neuroscience Research 96 (4): 573–588. doi:10.1002/jnr.24151.
- Hacker T.A. 2018. Advances in Experimental Medicine and Biology 1098: 45–58. doi:10.1007/978-3-319-97421-7_3.
- Han F., Wang J., Ding L., Hu Y. et al. 2020. Frontiers in Bioengineering and Biotechnology 8: 83. doi:10.3389/ fbioe.2020.00083.
- Herberts C.A., Kwa M.S., Hermsen H.P. 2011. Journal of Translational Medicine 9: 29. doi:10.1186/1479-5876-9-29.
- Hussell T., Lui S., Jagger C., Morgan D. et al. 2018. European Respiratory Review: An Official Journal of the European Respiratory Society 27 (148). doi:10.1183/16000617.0032-2018.
- Hussey G.S., Dziki J.L., Badylak S.F. 2018. Nature Reviews Materials 3 (7): 159–173. doi:10.1038/s41578-018-0023-x.
- Hwang J., Sullivan M.O., Kiick K.L. 2020. Frontiers in Bioengineering and Biotechnology 8: 69. doi:10.3389/ fbioe.2020.00069.
- Hynes R.O., Naba A. 2012. Cold Spring Harbor Perspectives in Biology 4 (1): a004903. doi:10.1101/cshperspect.a004903.
- Ito J.T., Lourenco J.D., Righetti R.F., Tiberio I. et al. 2019. Cells 8 (4). doi:10.3390/cells8040342.
- Kleiderman E., Boily A., Hasilo C., Knoppers B.M. 2018. Stem Cell Research & Therapy 9 (1): 307. doi:10.1186/s13287-018-1055-2.
- Kular J.K., Basu S., Sharma R.I. 2014. Journal of Tissue Engineering 5: 2041731414557112. doi:10.1177/2041731414557112.
- Langhans S.A. 2018. Frontiers in Pharmacology 9: 6. doi:10.3389/fphar.2018.00006.
- Leahy M., Thompson K., Zafar H., Alexandrov S. et al. 2016. *Stem Cell Research & Therapy 7* (1): 57. doi:10.1186/s13287-016-0315-2.
- Lee R.T., Walsh K. 2016. Circulation 133 (25): 2618-2625. doi:10.1161/CIRCULATIONAHA.115.019214.
- Lepelletier F.X., Mann D.M., Robinson A.C., Pinteaux E. et al. 2017. *Neuropathology and Applied Neurobiology 43* (2): 167–182. doi:10.1111/nan.12295.
- Liao J., Xu B., Zhang R., Fan Y. et al. 2020. *Journal of Materials Chemistry B* 8 (44): 10023–10049. doi:10.1039/ D0TB01534B.

Liras A. 2010. Journal of Translational Medicine 8: 131. doi:10.1186/1479-5876-8-131.

- Lockhart M., Wirrig E., Phelps A., Wessels A. 2011. Birth Defects Research. Part A, Clinical and Molecular Teratology 91 (6): 535–550. doi:10.1002/bdra.20810.
- Lorden E.R., Levinson H.M., Leong K.W. 2015. Drug Delivery and Translational Research 5 (2): 168–186. doi:10.1007/s13346-013-0165-8.
- Lu P., Takai K., Weaver V.M., Werb Z. 2011. Cold Spring Harbor Perspectives in Biology 3 (12). doi:10.1101/ cshperspect.a005058.
- Lubbers B.R., Smit A.B., Spijker S., van den Oever M.C. 2014. Progress in Brain Research 214: 263–284. doi:10.1016/B978-0-444-63486-3.00012-8.
- Mahalatchimy A. 2016. Medical Law Review 24 (2): 234–258. doi:10.1093/medlaw/fww009.
- Mahla R.S. 2016. International Journal of Cell Biology 2016: 6940283. doi:10.1155/2016/6940283.
- Mao A.S., Mooney D.J. 2015. Proceedings of the National Academy of Sciences of the United States of America 112 (47): 14452–14459. doi:10.1073/pnas.1508520112.
- marketsandmarkets. 2020. Regenerative Medicine Market by Product. Retrieved 14 Feb, 2021, from https:// www.marketsandmarkets.com/Market-Reports/regenerative-medicine-market-65442579.html
- Martin I., Simmons P.J., Williams D.F. 2014. Science Translational Medicine 6 (232): 232fs216. doi:10.1126/ scitranslmed.3008558.
- Martinez-Montiel Mdel P., Gomez-Gomez G.J., Flores A.I. 2014. World Journal of Gastroenterology 20 (5): 1211–1227. doi:10.3748/wjg.v20.i5.1211.
- Mata A., Azevedo H.S., Botto L., Gavara N. et al. 2017. Current Stem Cell Reports 3 (2): 83–97. doi:10.1007/ s40778-017-0081-9.
- Mouw J.K., Ou G., Weaver V.M. 2014. Nature Reviews. Molecular Cell Biology 15 (12): 771–785. doi:10.1038/ nrm3902.
- Muncie J.M., Weaver V.M. 2018. Current Topics in Developmental Biology 130: 1–37. doi:10.1016/ bs.ctdb.2018.02.002.
- Naumova A.V., Modo M., Moore A., Murry C.E. et al. 2014. *Nature Biotechnology 32* (8): 804–818. doi:10.1038/nbt.2993.
- Nikolova M.P., Chavali M.S. 2019. Bioactive Materials 4: 271–292. doi:10.1016/j.bioactmat.2019.10.005.
- O'Brien F.J., Duffy G.P. 2015. Journal of Anatomy 227 (6): 705–706. doi:10.1111/joa.12401.
- Ohata K., Ott H.C. 2020. Surgery Today 50 (7): 633-643. doi:10.1007/s00595-020-02000-y.
- Panoskaltsis-Mortari, A. 2015. Current Transplantation Reports 2 (1): 90–97. doi:10.1007/s40472-014-0048-z.
- Parmaksiz M., Dogan A., Odabas S., Elcin A.E. et al. 2016. *Biomedical Materials 11* (2): 022003. doi:10.1088/1748-6041/11/2/022003.
- Pelosi P., Rocco P.R., Negrini D., Passi A. 2007. Anais da Academia Brasileira de Ciencias 79 (2): 285–297. doi:10.1590/s0001-37652007000200010.
- Petrey A.C., de la Motte C.A. 2017. Current Opinion in Gastroenterology 33 (4): 234–238. doi:10.1097/ MOG.00000000000368.
- Pigeau G.M., Csaszar E., Dulgar-Tulloch A. 2018. *Frontiers in Medicine 5*: 233. doi:10.3389/fmed.2018.00233. Pomin V.H., Mulloy B. 2018. *Pharmaceuticals 11* (1): 27.
- Ponticos M., Smith B.D. 2014. Journal of Biomedical Research 28 (1): 25–39. doi:10.7555/JBR.27.20130064.
- Ramos T., Moroni L. 2020. Tissue Engineering. Part C, Methods 26 (2): 91-106. doi:10.1089/ten. TEC.2019.0344.
- Rhodes K.E., Fawcett J.W. 2004. Journal of Anatomy 204 (1): 33-48. doi:10.1111/j.1469-7580.2004.00261.x.
- Ribitsch I., Baptista P.M., Lange-Consiglio A., Melotti L. et al. 2020. Frontiers in Bioengineering and Biotechnology 8: 972. doi:10.3389/fbioe.2020.00972.
- Richard A.D., Lu X.H. 2019. *Neural Regeneration Research 14* (4): 578–581. doi:10.4103/1673-5374.247459. Romito A., Cobellis G. 2016. *Stem Cells International 2016*: 9451492. doi:10.1155/2016/9451492.
- Rosemann A., Vasen F., Bortz G. 2019. Science as Culture 28 (2): 223–249. doi:10.1080/09505431.2018.155 6253.
- Rozario T., DeSimone D.W. 2010. Developmental Biology 341 (1): 126-140. doi:10.1016/j.ydbio.2009.10.026.
- Ruiz-Babot G., Hadjidemetriou I., King P.J., Guasti L. 2015. Frontiers in Endocrinology 6: 70. doi:10.3389/ fendo.2015.00070.
- Ruoslahti E. 1996. Glycobiology 6 (5): 489-492. doi:10.1093/glycob/6.5.489.
- Sampogna G., Guraya S.Y., Forgione A. 2015. Journal of Microscopy and Ultrastructure 3 (3): 101–107. doi:10.1016/j.jmau.2015.05.002.

- Shineha R., Inoue Y., Ikka T., Kishimoto A. et al. 2017. Regenerative Therapy 7: 89-97. doi:10.1016/j. reth.2017.11.001.
- Stoppel W.L., Kaplan D.L., Black L.D., 3rd 2016. Advanced Drug Delivery Reviews 96: 135-155. doi:10.1016/j. addr.2015.07.009.
- Theocharis A.D., Manou D., Karamanos N.K. 2019. The FEBS Journal 286 (15): 2830-2869. doi:10.1111/ febs.14818.
- Traphagen S., Yelick P.C. 2009. Regenerative Medicine 4 (5): 747-758. doi:10.2217/rme.09.38.
- Vaday G.G., Lider O. 2000. Journal of Leukocyte Biology 67 (2): 149-159. doi:10.1002/jlb.67.2.149.
- Walker C., Mojares E., Del Rio Hernandez A. 2018. International Journal of Molecular Sciences 19 (10). doi:10.3390/ijms19103028.
- Weiss M.L., Troyer D.L. 2006. Stem Cell Reviews 2 (2): 155-162. doi:10.1007/s12015-006-0022-v.
- Whitesides G.M., Ostuni E., Takayama S., Jiang X. et al. 2001. Annual Review of Biomedical Engineering 3: 335-373. doi:10.1146/annurev.bioeng.3.1.335.
- Wight T.N. 2018. Matrix Biology: Journal of the International Society for Matrix Biology 71-72: 396-420. doi:10.1016/j.matbio.2018.02.019.
- Yalcinkaya T.M., Sittadjody S., Opara E.C. 2014. Maturitas 77 (1): 12-19. doi:10.1016/j.maturitas.2013.10.007.
- Yu N.H., Chun S.Y., Ha Y.S., Kim H.T. et al. 2018. Tissue Engineering and Regenerative Medicine 15 (5): 639-647. doi:10.1007/s13770-018-0133-y.
- .0.1(_0(1):9, contrances Yue B. 2014. Journal of Glaucoma 23 (8 Suppl 1): S20-23. doi:10.1097/IJG.000000000000108.
- Zhang Y.Z., Li Y.Y. 2014. World Journal of Gastroenterology 20 (1): 91–99. doi:10.3748/wjg.v20.i1.91.

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Innovations in Plant Science for Better Health: From Soil to Fork

Herbs, Spices, and Medicinal Plants for Human Gastrointestinal Disorders

Health Benefits and Safety



Editors Megh R. Goyal | Preeti Birwal Durgesh Nandini Chauhan



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Ethnopharmacology and Therapeutic Potential of *Carica papaya*

GURPREET SINGH, POOJA CHAWLA, ABDUL FARUK, and VINEY CHAWLA

ABSTRACT

Papaya (*Carica papaya* Linn) has been widely used as traditional herbal remedy for the prevention and management of several conditions and diseases. During the past few decades, it has been used in the treatment of digestive problems, wounds, dengue, and jaundice, etc. Its major bioactive phytoconstituents are: papain, chymopapain, alkaloids, flavonoids, lycopene, carotenoids, anthraquinones glycoside, antioxidants, and vitamins. This chapter has highlighted various ethnopharmacological and traditional uses of different parts of *Carica papaya*.

1.1 INTRODUCTION

Carica papaya is a member of the family *Caricaceae* (a family of dicots plants with four genera).⁵⁶ *Papaya* is a delicious fruit in most tropical and semitropical countries and is cultivated mostly for its consumption as fresh fruit, and for use in drinks, jams, salads, and candies.² The papaya plant has been well-documented in the literature for a number of medicinal properties and has been used against diseases, such as gastroenteritis, urethritis, typhoid fever, wound infection, asthma, rheumatism, fever, diarrhea, boils, and hypertension, etc.^{7,11,37,59,60,67,71} All parts of papaya (seeds, roots, rinds, and fruits) have beneficial therapeutic and protective properties (Fig. 1.1). Different parts of the papaya plant have been used in

the food (nutraceuticals), skincare products, leather, and pharmaceutical production.⁴¹ Scientists have reported the activity of papaya for antifertility, anthelmintic, and anti-inflammatoryeffects.^{50,53,55,58,97} The latex of unripe fruit is widely used in pharmaceutical and cosmetics products.^{18,69,78}



FIGURE 1.1 Major parts of Carica papaya plant.

The Spanish chronicler Oviedo indicated the papaya on Panamanian and Colombian coasts in 1526. Due to the high viability of papaya seeds, the fruit was rapidly produced in the tropics.²³ During this century, papaya has been cultivated in tropical regions with fertile soils and heavy rainfall. Then, papaya seeds were introduced to Southeast Asia and India by Spanish and Portuguese mariners. Later, papaya seeds reached Hawaii between 1800 and 1820.⁷⁷ In the 20th century, papaya seeds were taken to Barbados, Jamaica, Mexico, and Florida.

1.2 GEOGRAPHICAL DISTRIBUTION

It is local to the tropics of the Americas, however, now it is generally developed all through the world, and is accessible consistently.³ It is cultivated in different parts of the world and the significant cultivators of papaya plants include India, tropical America, Europe, Australia, Hawaii, and South-East Asia.³⁰ Papaya is cultivated in all five continents due to its capability of growing in all soil types, but it requires good drainage.^{1,73}

The major contribution of its total production⁹⁸ comes from Asia, Central America, and other countries as shown in Figure 1.2, and major cultivars of papaya plant are listed in Table 1.1. Different vernacular names of *C. papaya* and the taxonomic hierarchy of *C. papaya* have been illustrated by many investigators.^{20,44,73}

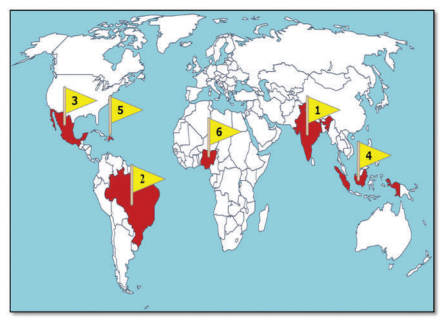


FIGURE 1.2 Major producers of papaya plant: (1) India, (2) Brazil, (3) Mexico, (4) Indonesia, (5) Dominican Republic, and (6) Nigeria.

1.3 MORPHOLOGY

Papaya is a small softwood and unbranched tropical fruit tree of 5–10 m in height with the spirally arranged leaves. The seven lobed leaves are large in diameter of about 50–70 cm, vary in sizes and shapes in different maturity stages. Fruits are commonly green while young and yellow-greenish or orange when ripe with the large ovoid smooth surface.^{1,10} The fruit has a hollow berry, which contains small black seeds that constitute about 15% of the total weight and the seeds are lined in five rows to the interior wall

of the fruit. Papaya tree starts to bear fruit within 1-2 years.⁷² It can be cultivated in either home gardens or outdoors.

Country	Variety
Australia	Improved Petersen, Guinea Gold
Barbados	Wakefield, Graeme
Cuba	Maradol
Dominican Republic	Cartagena
Florida	Cariflora, Betty, Homestead
Hawaii	Kapoho Solo, Waimanalo, Rainbow
India	Coorg Honey Dew, Coimbatore Varieties (CO1-CO8)
Indonesia	Semangka, Dampit
Malaysia	Eksotika, Sekaki
Mexico	Verde, Gialla, Cera, Chincona
Philippines	Cavite, Sinta
South Africa	Hortus Gold, Kaapmuiden
Taiwan	Tainung five
Thailand	Sai-nampueng, Khaek Dam
Trinidad	Santa Cruz Giant, Cedro
Venezuela	Paraguanera, Roja

TABLE 1.1Major Cultivars of Papaya in the World.

On the basis of reported literature, papaya plant is categorized into three primary sexes (Fig. 1.3), such as male (staminate) (\mathcal{J}), hermaphrodite (bisexual) (\mathcal{J}), and female (pistillate) (\mathcal{Q}). A typical male and female plants bear individual unisexual flowers, while hermaphrodite plant bears a combination of male unisexual and hermaphroditic flowers.^{33,54} The typical female flower is mostly large and conical in shape when it is mature with five petals spread from the base. The ovary is large in structure with a circular smooth surface, which produces spherical or ovoid-shaped flowers. Fruit progresses from globular to egg-shaped. In the case of hermaphrodite intermediate type, the flower is undefined and petals may be fused in their length or may be free from the base. Hermaphrodite elongated type of flower has fused petals from one-fourth to three-fourths of their total length with 10 anthers, out of which five are long and five are short. The long ovary contains five or more carpels and forms the fruit which is cylindrical to pear-shaped and is of great commercial value.

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The typical male flower has a long and thin corolla. It contains anthers in two series of five; one series is longer than the other. The male flowers have nonfunctional rudimentary pistil.^{33,61} Multiple species of papaya have been documented in the scientific literature, which belong to five genera, that is, *Jacaratia, Jarilla, Horovitzia, Carica*, and *Vasconcellea*.²⁴

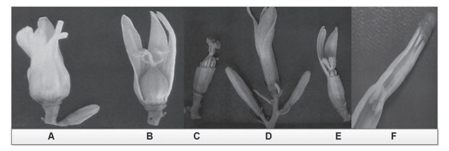


FIGURE 1.3 Six varieties of flowers of papaya plant: Typical female (A, B). Hermaphrodite intermediate (C). Hermaphrodite elongated (D). Hermaphrodite sterile (E). Typical male (F).

1.4 PHYTOCONSTITUENTS

Primary phytoconstituents reported from various parts of the *C. papaya* plant include papain (proteolytic enzyme), lycopene (tetraterpene), carotenoids, alkaloids, monoterpenoids, flavonoids, mineral (potassium, etc.), vitamins (A, C, and E; thiamine, niacin, and riboflavin), malic acid, and glycosides.^{1,34,69,74,81,96} Fresh fruit juice contains flavonoids, tannins, and anthocyanins with antioxidant ability as free radical scavengers.⁶⁸ Young leaves of papaya include carpaine, pseudocarpaine, dehydrocarpaine, choline, carposide, and vitamins (C and E). Phytochemical analysis of the different parts of the plant revealed the presence of various bioactive phytochemicals, which have pharmacological importance (Fig. 1.4).

Papaya fruit exhibits wide range of medicinal properties (i.e., antimicrobial, antiviral, anti-inflammatory, healing of wound and dressing aid, anticancer, neurodegenerative, diuretic, abortifacient agent, and contraceptive).⁴³ It is highly well-known for its nutritional values and it aids in digestion. Extract of the whole fruit contains immunity boosters (i.e., vitamin C, ferulic, caffeic acid, and *p*-coumaric) that protect human cells against oxidative stresses.¹³

Unripe fruit of papaya contains proteolytic enzyme papain (cysteine protease), which acts like pepsin in gastric juice. The papain is more active

in green fruit and shows extensive proteolytic activity toward proteins. The extract from the seeds of papaya shows antioxidant and anticancer activities due to the presence of various phenolic compounds, vanillic acid, and vitamin C.^{52,62,86}

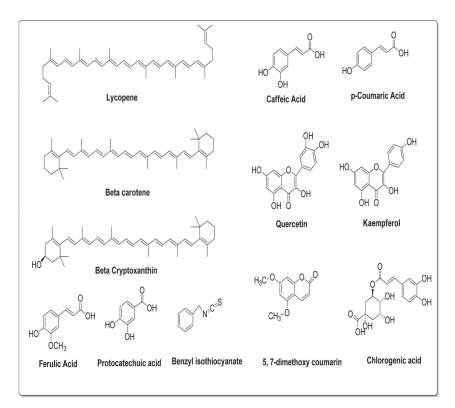


FIGURE 1.4 The structures of some phytoconstituents isolated from *C. papaya*.

Another source of papain is latex, which is harvested by incision on the surface of unripe fruit. After 4–5 days, latex is collected and further processed into dry powder for various uses in pharmaceutical and food industries.⁵¹ The process of isolation of papain from unripe fruit latex is shown in Figure 1.5.

The papaya fruit is suitable for human consumption due to its nutritional and digestive value, with a low caloric content, which provides a favorable cost-benefit to human health.⁶⁹ Furthermore, scientific studies report the nutritional content of 100 g of ripe and unripe papaya fruits as summarized

in Table 1.2. Results revealed that unripe papaya has the highest concentrations of different vitamins and minerals as compared with ripe fruits.^{22,79}

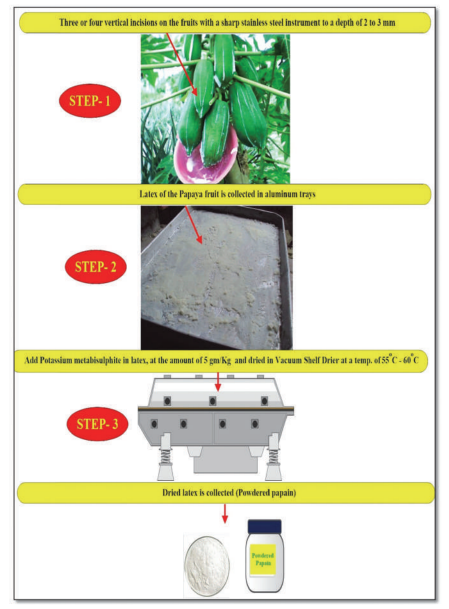


FIGURE 1.5 Isolation of papain powder from the latex from the papaya fruit.

Constituent	Ripe fruit (g)	Unripe fruit (g)	
Water	89.1	92.6	
Proteins	8.26	10.8	
Total lipid	0.93	1.35	
Ash	0.00459	6.76	
Carbohydrates	86.2	81.1	
Mineral Macronutrients:			
Sodium	0.1284	0.2838	
Potassium	1.238	2.743	
Magnesium	0.2294	0.6351	
Calcium	0.1468	0.4324	
Micronutrients:			
Iron	0.01284	0.00811	
Copper	0.00018	0.00014	
Zinc	0.00092	0	
Vitamins:			
Vitamin C	0.5688	0.0003919	
Thiamine	0.00028	0.00054	
Riboflavin	0.00028	0.026	
Niacin	0.0028	0.00405	
Carotene	7.807(µg)	0	

TABLE 1.2Nutritional Value of a Papaya Fruit.

1.5 PHARMACOLOGICAL ACTIVITIES AND THERAPEUTIC USAGES OF CARICA PAPAYA

Every part of papaya plant holds the therapeutic value from leaves to roots.^{56,69} The fruits, latex, and juice of papaya plant are the main source of many vitamins, which aid in dyspepsia, intestinal irritation, and habitual constipation.

The main constituent papain plays a vital role to improve the immune system.⁹ In traditional veterinary medicine, papaya seeds are used as de-wormers and is also used in tropical folk medicine. The fresh latex is used as a vermifuge.⁶ Papain is a proteolytic digestive enzyme that is used in several herbal formulations. Fresh juice of papaya prepared from peeled or unpeeled fruit is also sold as immunity booster drink because of its low cost, easy availability throughout the year and high nutritive value. In certain countries, the latex of the plant is used for tumors of uterus,

psoriasis, and ringworm. The root infusion is used against syphilis.⁸² Through several scientific studies, the traditional, pharmacological, and biological effects of *C. papaya* have been validated.^{10,44,74,78}

1.5.1 Anthelmintic Activity

A wide collection of papaya and their extracts have been used traditionally for the management of helminths (parasites). Papaya contains many biologically active compounds with varying properties in fruit, latex, leaves, and roots that aid in digestion. It has also been employed for treating intestinal worms.^{6,16} Papain, which is present in the latex of unripe green fruits of papaya, has been commercialized in various forms. Dried seeds of papaya have shown significant activity in the management of human intestinal parasites, which have increased the stool clearance rate of parasites without any side effects. It is represented as a novel class of antihelminthic due to the efficacy of papaya latex and cysteine proteinases against *Heligmosomoid espolygyrus* (nematode).⁸⁹ Shaziya et al. reported the antihelminthic action of papaya leaves on *A. Caninum* nematode infecting mice.⁸⁷

Papain is a protein enzyme with cysteine protease, chymopapain, and lysozyme, which can accelerate the reaction within body cells. During the digestion process, pancreas commonly produces enzymes in the human body, these enzymes break down the foods into micronutrients, which can be used by the body for energy and other functions.¹² Two main proteolytic enzymes (papain and chymopapain) in the latex of the papaya simply break down the proteins into amino acids through cleavage of the peptide bond. These proteins contained peptide bonds and can be easily broken down by enzymatic action into easily digestible micronutrients. It also helps to promote the digestion of wheat protein.⁴⁰

1.5.2 Antioxidant Activity

Antioxidant properties of aqueous extract of papaya leaves were evaluated in alcohol-induced acute gastric damage. The outcomes revealed that gastric ulcer index was significantly better in rats pretreated with the extract of papaya leaf as compared with the alcohol-treated rats. Further, leaf extract also offered reduced blood oxidative stress level in rats via the reduction of lipid peroxide levels in plasma and amplified red blood cell glutathione peroxidase activity.³⁹

Another study showed strong in vivo antioxidant actions of ethyl acetate fraction of unripe pulp of papaya on antioxidant enzymes (i.e., glutathione peroxidase (GPX), glutathione S-transferase (GST), glutathione reductase (GR), catalase, and glucose-6-phosphate dehydrogenase (G6PD)) in albino mouse. It has been suggested that it can be used for protection against gastric ulcer and oxidative stress.⁶⁴ Natural source of antioxidants may responsible for total antioxidant effect due to the presence of carotenoid, polyphenols, vitamin C, and vitamin E.⁵⁷ Several studies showed that the antioxidant property is related to the diminished DNA damage and decreased lipid peroxidation, which maintained the immune function.^{46,48}

1.5.3 Antiviral Activity

The published studies on dengue specified that the juice of papaya leaves could help to increase the platelets and white blood cells count in these patients.^{15,80} A study in 2012 has reported about in vitro studies of papaya leaf extracts on persons infected with dengue. Papaya leaf extract inhibited the heat- and hypotonicity-induced hemolysis of red blood cells and has membrane-stabilizing properties.⁷⁶ In a randomized controlled trial in dengue patients, there was an increment in platelets-related genes like arachidonate 12-lipoxygenase and platelet-activating factor receptor gene and that contributed to the prevention of platelet lysis. In folk medicine, papaya leaves have been used for the management of dengue fever with hemorrhagic symptoms.⁹¹

1.5.4 Antimicrobial Activity

Osato et al.⁶⁵ and Calzada et al.¹⁷ reported the ability of papaya seeds as antimicrobial agent against several Gram-positive and Gram-negative bacteria like *Trichomonas vaginalis* trophozoites, *Bacillus subtilis, Escherichia coli* and *Salmonella typhi*.^{17,65} The aqueous extract of papaya leaves and roots at different concentrations showed antimicrobial effects against pathogenic bacteria.⁸ The pulp and fruits of papaya also showed remarkable antibacterial effect against *B. subtilis, K. pneumonia, P. vulgaris, E. coli, P. aeruginosa, S. typhi, E. cloacae, and S. aureus*.¹¹

1.5.5 Antifungal Activity

The papaya leaves and seeds of ripe and unripe fruits were evaluated against phytopathogenic fungi (i.e., *R. stolonifer*, Fusarium spp. and *C. gloeosporioides*), which exhibited good antifungal activity. The antifungal activity was observed to increase in a concentration-dependent manner.¹⁹ The latex of papaya also inhibits the growth of *Candida albicans*. The latex shows antifungal activity due to partial degradation of the outermost layers of fungal cell wall, which lacks polysaccharides.²⁶ The synergistic effect of latex of papaya with fluconazole in *C. albicans* was also reported.²⁵

1.5.6 Anti-Inflammatory Activity

It has been well documented in the literature that the dried papaya leaves are used for the management of inflammation, arthritis, rheumatism, and as wound dressing material. The ethanolic extract of the leaves was examined in rats using a paw edema model with indomethacin-treated control group. The results showed that the extracts significantly reduced edema and amount of granuloma. Similar results were confirmed with other models, that is, cotton pellet granuloma model and formaldehyde-induced arthritis model.^{67,92,93}

Papaya leaves are a rich source of carpaine, nicotinic acid, which may be accountable for the anti-inflammatory effect. Ahmed et al.⁴ assessed the inflammation at acute, subchronic, and chronic phase using the cotton pellet granuloma model, formaldehyde-induced arthritis and carrageenaninduced paw edema models. They suggested that the anti-inflammatory activity of the ethyl alcohol extract of papaya was due to the inhibition of *prostaglandin*- mediated inflammation.⁴ Papaya leaf extract also exhibited anti-arthritic activity by the modulation of inflammatory mediators, such as, cytokines or chemokines, prostaglandins or leukotrienes.⁶⁷

1.5.7 Antifertility Effects

The antifertility activity of papaya fruit was evaluated in adult rat and pregnant rat model. The results revealed that the unripe fruit disturbed the estrous cycle and encouraged the abortion.²⁷ Seed extract showed antifertility activity due to gradual degeneration of Sertoli and Leydig cells,

which induced long-term azoospermia.⁹⁵ A recent report revealed that seeds possess reversible male contraceptive potential by directly rendered the spermatozoa process.⁹⁰ It is further reported that root extract exerts morphological changes in the endometrium of rat uterus⁸³ and the aqueous extract of seeds has shown miscarriage in female *Sprague Dawley* rats.

The crude extract of papaya bark showed antifertility activity in rats due to its effect on sperm motility; and while the aqueous/petroleum ether/ alcoholic extracts in rabbits inhibited ovulation cycle. Therefore, it can be utilized as an effective contraceptive in animals.⁴⁷ It was further reported that the unripe or half-ripe fruits contain a high concentration of the latex, which increased the uterine contraction. Normal consumption of ripe papaya is safe in pregnancy, but unripe papaya is unsafe.²

1.5.8 Anticancer Activity

Many studies scientifically validated the anticancer effects of papaya leaves. The aqueous extract of papaya leaves exhibits a dose-dependent significant activity against the cells of breast and lung adenocarcinoma, cervical, hepatocellular and pancreatic epithelial carcinoma, and mesothelioma. These results indicate that extracts may inhibit the growth of different types of cancer cell lines. However, the precise cellular mechanism of action remains unclear.^{29,66} Several studies have claimed that mechanisms in the inhibition of proliferation by papain include the production of cytokines by human peripheral blood mononuclear cells, interfering in cancer cell wall and cleavages of proteins into amino acid form.²¹

Leaves of *C. papaya* (which contain a high concentration of tocopherol, lycopene, flavonoid, and benzyl isothiocyanate) potentially contribute to anti-tumor activity.⁷⁹ Similarly, fermented product of papaya (FPP[®]) claimed the immunity booster and antioxidant activity. The role of free radicals in propagating cancer is fully documented. Thus, by acting as an antioxidant, it helps to control cancerous growth.⁴⁹

1.5.9 Antihypertensive Activity

Methanolic extract of papaya elicited the antihypertensive effects due to in vivo inhibition of angiotensin-converting enzyme and it improved the effect on the baroreflex. It was reported that angiotensin-converting enzyme inhibitory activity was similar to those of enalapril and reduced the cardiac hypertrophy.¹⁴

1.5.10 Antimalarial Activity

Daily consumption of papaya leaves is a common practice in tropical communities for preventing malaria caused by *Plasmodium* genus. In vitro antiplasmodial effect of the leaf extracts was reported to be due to carpaine, which is an alkaloid.^{42,94} Petroleum ether extract of the rind of papaya fruit also exhibits antimalarial activity.⁹⁹

1.5.11 Hematological Activity

The study revealed that phytochemicals in seed, leaf, and pulp produced significant effects on certain blood parameters in treated rats. A dose-dependent effect was observed, which could be attributed to the existence of folic acid, vitamin B_{12} , alkaloids, and glycosides. It can be used for the treatment of sickle-cell anemia.^{36,38}

1.5.12 Wound Healing Activity

Papaya latex contains papain, which can break down the necrotic tissue contributing to wound healing process. The study showed that the latex of *C. papaya* decreases the oxidative tissue damage thus ensuring the clot formation process during healing and the increase in di-hydroxyproline content.²⁸ It is also known to be effective in diabetic wound healing by preventing infection due to its antimicrobial activity.⁵

1.5.13 Hepatoprotective Activity

Ethanol and aqueous extracts of papaya fruit hold the hepatoprotective effect against carbon tetrachloride (CCl_4) -induced hepatotoxicity in rats. Results revealed significant hepatoprotection by reduction in biochemical parameters, such as, SGPT (serum glutamic pyruvic transaminase), SGOT (serum glutamic-oxaloacetic transaminase), ALP (alkaline phosphatase), and serum bilirubin, which are indicators of liver damage.⁷⁵

1.5.14 Topical Use

Various topical applications of papaya fruits have been used in developing countries, such as topical ulcer dressings and burn dressing. It is a cost-effective remedy for desloughing necrotic tissue and preventing burn wound infection.³¹ It also provides a granulating tissue, which is suitable for the application of skin graft. Now-a-days, papaya is commonly used in children's burns dressing. Papaya fruit is crushed and is daily applied on the infected burns as a layer.⁸⁸

1.6 SUMMARY

Scientists around the globe have focused on papaya plant for its high medicinal value with simple availability in nature. *C. papaya* has the potential of capturing the global market of herbal formulations for therapeutic potential in digestive disorders. However, this needs a clinical validation. The presence of secondary metabolites has been identified, which may help in the planning of such clinical studies, which are needed to understand and explore the exact pharmacological and molecular mechanisms action of *C. papaya* activity. It will also help to establish its toxicity profile along with drug interactions.

KEYWORDS

- abortifacient
- anthelmintic activity
- antifertility
- Carica papaya
- caricaceae
- chymopapain
- dengue
- digestion enhancer
- nutraceutical
- papain powder

REFERENCES

- Abdulazeez, M. A.; Sani, I. Use of Fermented Papaya (*Carica papaya*) Seeds as a Food Condiment, and Effects on Pre- and Post-implantation Embryo Development. In *Nuts and Seeds in Health and Disease Prevention*; Preedy, V. R., Watson, R. R., Patel, V. B., Eds.; Academic Press: San Diego, 2011; pp 855–863.
- Adebiyi, A.; Adaikan, P. G.; Prasad, R. N. Papaya (*Carica papaya*) Consumption Is Unsafe in Pregnancy: Fact or Fable? Scientific Evaluation of a Common Belief in Some Parts of Asia using a Rat Model. *Br. J. Nutr.* 2002, *88* (2), 199–203.
- Adeneye, A. A. The 6-Subchronic and Chronic Toxicities of African Medicinal Plants. In *Toxicological Survey of African Medicinal Plants*; Kuete, V., Ed.; Elsevier, 2014; pp 99–133.
- 4. Ahmed, M.; Ramabhimalah, S. Anti-Inflammatory Activity of Aqueous Extract of *Carica papaya* Seeds in Albino Rats. *Biomed. Pharmacol. J.* **2012**, *5*, 173–177.
- Ajani, R.; Ogunbiyi, K. *Carica papaya* Latex Accelerates Wound Healing in Diabetic Wistar Rats. *Eur. J. Med. Plants* 2015, *9*, 1–12.
- Ameen, S.; Azeez, O. M.; Baba, Y.; Raji, L. Anthelminitic Potency of *Carica papaya* Seeds Against Gastro-intestinal Helminths in Red Sokoto goat. *Ceylon J. Sci.* 2018, 47, 137–143.
- Amzad-Hossain, M.; Hitam, S.; Hadidja I. A. Pharmacological and Toxicological Activities of the Extracts of papaya Leaves used Traditionally for the Treatment of Diarrhea. J. King Saud Univ. – Sci. 2020, 32 (1), 962–969.
- Anibijuwon, I.; Augustine, U. Antimicrobial Activity of *Carica papaya* on Some Pathogenic Organisms of Clinical Origin from South-Western Nigeria. *Ethnobot. Leaflets* 2009, 13, 850–864.
- Anjum, V.; Arora, P.; Ansari, S. H.; Najmi, A. K.; Ahmad, S. Antithrombocytopenic and Immunomodulatory Potential of Metabolically Characterized Aqueous Extract of *Carica papaya* Leaves. *Pharm. Biol.* 2017, 55 (1), 2043–2056.
- Aravind, G.; Bhowmik, D.; S, D.; Harish, G. Traditional and Medicinal uses of Carica papaya. J. Med. Plants Stud. 2013, 1, 7–15.
- 11. Asghar, N.; Naqvi, S. A. R.; Hussain, Z. Compositional Difference in Antioxidant and Antibacterial Activity of all Parts of the *Carica papaya* using Different Solvents. *Chem. Central J.* **2016**, *10* (1), 5–9.
- Azarkan, M.; Moussaoui, A.; Wuytswinkel, D.; Dehon, G.; Looze, Y. Fractionation and Purification of the Enzymes Stored in the Latex of *Carica papaya*. J. Chromatogr. B Anal. Technol. Biomed. Life Sci. 2003, 790, 229–238.
- Boshra, V.; Tajul, A. Y. Papaya-an Innovative Raw Material for Food and Pharmaceutical Processing Industry. *Health Environ. J.* 2013, 4, 68–75.
- Brasil, G. A.; Ronchi, S. N.; do Nascimento, A. M. Antihypertensive Effect of *Carica papaya* via a Reduction in ACE Activity and Improved Baroreflex. *Planta Medica* 2014, 80 (17), 1580–1587.
- 15. Bsr, D.; Kj, G.; Lakshmiprasad, J. Effect of Papaya Leaf Juice on Platelet and WBC Count in Dengue Fever: A Case Report. *J. Ayurveda Holistic Med.* **2013**, *1*, 44–47.
- Buttle, D. J.; Behnke, J. M.; Bartley, Y. Oral Dosing with Papaya Latex is an Effective Anthelmintic Treatment for Sheep Infected with *Haemonchus contortus*. *Parasites Vectors* 2011, 4 (1), 36–40.

- Calzada, F.; Yepez-Mulia, L.; Tapia-Contreras, A. Effect of Mexican Medicinal Plant Used to Treat Trichomoniasis on Trichomon as vaginal istrophozoites. J. Ethnopharmacol. 2007, 113 (2), 248–251.
- Chandrasekaran, R.; Seetharaman, P. *Carica papaya* (Papaya) Latex: A New Paradigm to Combat Against Dengue and Filariasis Vectors Aedesaegypti and Culexquinque fasciatus (Diptera: Culicidae). *3 - Biotech* 2018, *8* (2), 83–91.
- Chavez-Quintal, P.; Gonzalez-Flores, T.; Rodriguez-Buenfil, I.; Gallegos-Tintore, S. Antifungal Activity in Ethanolic Extracts of *Carica papaya* L. cv. Maradol Leaves and Seeds. *Ind. J. Microbiol.* 2011, *51* (1), 54–60.
- Cotruţ, R.; Butcaru, A.; Mihai, C.; Stănică, F. Carica papaya L. Cultivated in Greenhouse Conditions. J. Horticulture Forestry Biotechnol. 2017, 21 (3), 130–136.
- Desser, L.; Rehberger, A.; Paukovits, W. Proteolytic Enzymes and Amylase Induce Cytokine Production in Human Peripheral Blood Mononuclear Cells in vitro. *Cancer Biother.* 1994, 9 (3), 253–263.
- 22. Fauziya, S.; Krishnamurthy, R. Papaya (*Carica papaya*): Source Material for Anticancer. *CIB Tech J. Pharma. Sci.* **2013**, *2*, 25–34.
- Fuentes, G.; Santamaría, J. M. Papaya (*Carica papaya* L.): Origin, Domestication, and Production. In *Genetics and Genomics of Papaya*. Ming, R., Moore, P. H., Eds.; Springer: New York, NY, 2014; pp 3–15.
- Geetika, S.; Ruqia, M.; Harpreet, K.; Neha, D.; Shruti, K.; Singh, S. P. Genetic Engineering in Papaya. Chapter 7; In *Genetic Engineering of Horticultural Crops*; Rout, G. R., Peter, K. V., Eds.; New York: Academic Press, 2018; pp 137–154.
- Giordani, R.; Gachon, C.; Moulin-Traffort, J.; Regli, P. Synergistic Effect of *Carica papaya* Latex Sap and Fluconazole on *Candida albicans* Growth. *Mycoses* 1997, 40 (11–12), 429–437.
- Giordani, R.; Siepaio, M.; Moulin-Traffort, J.; Regli, P. Antifungal Action of *Carica papaya* Latex: Isolation of Fungal Cell Wall Hydrolyzing Enzymes. *Mycoses* 1991, 34 (11–12), 469–477.
- Gopalakrishnan, M.; Rajasekharasetty, M. R. Effect of papaya (*Carica papaya* Linn) on Pregnancy and Estrous Cycle in Albino Rats of Wistar strain. *Ind. J. Physiol. Pharmacol.* 1978, 22 (1), 66–70.
- Gurung, S.; Škalko-Basnet, N. Wound Healing Properties of *Carica papaya* latex: In vivo Evaluation in Mice Burn Model. J. Ethnopharmacol. 2009, 121 (2), 338–341.
- Hadadi, S.; Li, H.; Rafie, R.; Kaseloo, P.; Witiak, S.; Siddiqui, R. Anti-oxidation Properties of Leaves, Skin, Pulp, and Seeds Extracts from Green papaya and Their Anti-cancer Activities in Breast Cancer Cells. J. Cancer Metastasis Treat. 2018, 4, 25–29.
- Hewajulige, I.; Wijeratnam, S.; Hutchinson, M. Hexanal Compositions for Enhancing Shelf-life and Quality in Papaya Chapter 9, In *Postharvest Biology and Nanotechnology*; Paliyath, G., Subramanian, J., Lim, L.T., Subramanian, K., Handa, A.K., Mattoo, A.K., Eds.; Wiley: USA, 2018; pp 199–214.
- 31. Hewitt, H.; Whittle, S.; Lopez, S.; Bailey, E.; Weaver, S. Topical Use of papaya in Chronic Skin Ulcer Therapy in Jamaica. *West Ind. Med. J.* **2000**, *49* (1), 32–3.
- Hounzangbe-Adote, M. S.; Paolini, V.; Fouraste, I.; Moutairou, K.; Hoste, H. In vitro Effects of Four Tropical Plants on Three Life-cycle Stages of the Parasitic Nematode, Haemonchus contortus. *Res. Veterin. Sci.* 2005, 78 (2), 155–160.

- https://www.itfnet.org/v1/2016/05/papaya-name-taxonomy-botany/; Accessed on March 02, 2020.
- Hussain, S. Z.; Razvi, N.; Ali, S. I.; Hasan, S. M. F. Development of Quality Standard and Phytochemical Analysis of *Carica papaya* Linn leaves. *Pakistan J. Pharma. Sci.* 2018, *31* (5), 2169–2177.
- Ianiro, G.; Pecere, S.; Giorgio, V.; Gasbarrini, A.; Cammarota, G. Digestive Enzyme Supplementation in Gastrointestinal Diseases. *Curr. Drug Metabol.* 2016, 17 (2), 187–193.
- Ikpeme, E. V.; Ekaluo, U. B.; Kooffreh, M. E.; Udensi, O. Phytochemistry and Hematological Potential of Ethanol Seed Leaf and Pulp Extracts of *Carica papaya* (Linn.). *Pak. J. Biol. Sci.* 2011, *14* (6), 408–411.
- Ikram, E. H. K.; Stanley, R.; Netzel, M.; Fanning, K. Phytochemicals of Papaya and Its Traditional Health and Culinary Uses: Review. J. Food Compos. Anal. 2015, 41, 201–211.
- Imaga, N.; Gbenle, G.; Okochi, V.; Akanbi, S. Antisickling Property of Carica papaya Leaf Extract. Afr. J. Biochem. Res. 2009, 3 (4), 102–106.
- Indran, M.; Mahmood, A. A.; Kuppusamy, U. R. Protective Effect of *Carica papaya* L Leaf Extract against Alcohol-induced Acute Gastric Damage and Blood Oxidative Stress in Rats. *West Ind. Med. J.* 2008, *57* (4), 323–326.
- 40. Islam, R. Isolation, Purification and Modification of Papain Enzyme to Ascertain Industrially Valuable Nature. *Int. J. Bio-Technol. Res.* **2013**, *3*, 11–22.
- Jagtiani, J.; Chan, H. T.; Sakai, W. S. The 4-Papaya. In *Tropical Fruit Processing*; Jagtiani, J., Chan, H. T., Sakai, W. S., Eds.; Academic Press: New York, 1988; pp 105–147.
- Julianti, T.; Oufir, M.; Hamburger, M. Quantification of the Antiplasmodial Alkaloid Carpaine in papaya (*Carica papaya*) Leaves. *Planta Medica* 2014, 80 (13), 1138–1142.
- Kadiri, O.; Olawoye, B.; Samson, O.; Adalumo, O. Nutraceutical and Antioxidant Properties of the Seeds, Leaves and Fruits of *Carica papaya*: Potential Relevance to Humans Diet, the Food Industry and the Pharmaceutical Industry–A Review. *Turkish J. Agric.*—Food Sci. Technol. 2016, 4, 1039–1052.
- Kaliyaperumal, K.; Kim, H. M.; Jegajeevanram, K.; Xavier, J.; Vijayalakshmi, J. Papaya: Gifted Nutraceutical Plant—Critical Review of Recent Human Health Research. *Int. J. Genuine Trad. Med.* 2014, *4*, 1–17.
- Kermanshai, R.; McCarry, B.; Rosenfeld, J.; Summers, P.; Weretilnyk, E.; Sorger, G. Benzyl Isothiocyanate is the Chief or Sole Anthelmintic in papaya Seed Extracts. *Phytochemistry* 2001, *57*, 427–435.
- Kurutas, E. B. The Importance of Antioxidants Which Play the Role in Cellular Response Against Oxidative/ Nitrosative Stress: Current State. *Nutr. J.* 2016, *15* (1), 71–77.
- Kusemiju, O.; Noronha, C.; Okanlawon, A. The Effect of Crude Extract of the Bark of *Carica papaya* on the Seminiferous Tubules of Male Sprague-Dawley Rats. *Nigerian Postgraduate Med. J.* 2002, 9 (4), 205–209.
- Lobo, V.; Patil, A.; Phatak, A.; Chandra, N. Free Radicals, Antioxidants and Functional Foods: Impact on Human Health. *Pharmacognosy Rev.* 2010, *4* (8), 118–126.
- Logozzi, M.; Di Raimo, R.; Mizzoni, D. Beneficial Effects of Fermented Papaya Preparation (FPP®) Supplementation on Redox Balance and Aging in a Mouse Model. *Antioxidants* 2020, 9 (2), 144–149.

- Lohiya, N. K.; Goyal, R. B.; Jayaprakash, D.; Ansari, A. S.; Sharma, S. Antifertility Effects of Aqueous Extract of Carica papaya Seeds in Male Rats. *Planta Medica* 1994, 60 (5), 400–404.
- Macalood, J.; Vicente, H.; Boniao, R.; Gorospe, J.; Roa, E. Chemical Analysis of Carica papaya L. Crude Latex. Am. J. Plant Sci. 2013, 4, 1941–1948.
- Malek, K.; Norazan, M.; Ramaness, P. Cysteine Proteases from *Carica papaya*: An Important Enzyme Group of Many Industrial Applications. *IOSR J. Pharm. Biol. Sci.* 2016, 11, 11–16.
- Manpreet, K.; Naveen C. T.; Seema, S.; Arvind, K.; Elena, E. S. Ethnomedicinal Uses, Phytochemistry and Pharmacology of *Carica papaya* Plant: Review. *Mini-Rev. Org. Chem.* 2019, *16* (5), 463–480.
- Ming, R.; Yu, Q.; Moore, P. H. Sex Determination in Papaya. Semin. Cell Devel. Biol. 2007, 18 (3), 401–408.
- 55. Mohansrinivasan, V.; Janani, S. V. Exploring the Bioactive Potential of *Carica papaya*. *Nat. Prod. J.* **2017**, *7* (4), 291–297.
- Moy, J. H. Papayas. In *Encyclopedia of Food Sciences and Nutrition*; Caballero, B., Ed., 2nd ed.; Academic Press: Oxford, 2003; pp 4345–4351.
- Mutalib, M.; Amira, B.; Asmah, R.; Othman, F. Antioxidant Analysis of Different Parts of Carica Papaya. *Int. Food Res. J.* 2013, 20, 1043–1048.
- Nafiu, A. B.; Alli-Oluwafuyi, A. Papaya (*Carica papaya* L., Pawpaw). Chapter 3.32; In *Nonvitamin and Nonmineral Nutritional Supplements*; Nabavi, S. M., Silva, A. S., Eds.; Academic Press: New York, 2019; pp 335–359.
- O'Hare, T. J.; Williams, D. J. Papaya as a Medicinal Plant. In *Genetics and Genomics of Papaya*; Ming, R., Moore, P. H., Eds; Springer: New York, NY, 2014; pp 391–407.
- Odoh, U. E.; Uzor, P. F.; Eze, C. L.; Akunne, T. C. Medicinal Plants used by the People of Nsukka Local Government Area, South-eastern Nigeria for the Treatment of Malaria: An Ethnobotanical Survey. *J. Ethnopharmacol.* 2018, *218*, 1–15.
- Odu, E. A.; Adedeji, O.; Adebowale, K. Occurrence of Hermaphroditic Plants of *Carica papaya* L. in Southwestern Nigeria. J. Plant Sci. 2010, 5, 335–344.
- 62. Ojimelukwe, P.; Eji, C. Chemical Composition of Leaves, Fruit Pulp and Seeds in *Carica papaya* (L) Morphotypes. *Int. J. Med. Arom. Plants* **2012**, *2*, 200–206.
- Okeniyi, J. A.; Ogunlesi, T. A.; Oyelami, O. A.; Adeyemi, L. A. Effectiveness of Dried Carica papaya Seeds Against Human Intestinal Parasitosis: A Pilot Study. J. Med. Food 2007, 10 (1), 194–196.
- Oloyede, O.; Franco, J.; Roos, D. Antioxidant Properties of Ethyl Acetate Fraction of Unripe Pulp of *Carica papaya* in Mice. *J. Microbiol., Biotechnol. Food Sci.* 2011, 1, 409–425.
- Osato, J. A.; Santiago, L. A.; Remo, G. M.; Cuadra, M. S.; Mori, A. Antimicrobial and Antioxidant Activities of Unripe Papaya. *Life Sci.* 1993, 53 (17), 1383–1389.
- Otsuki, N.; Dang, N. H.; Kumagai, E. Aqueous Extract of *Carica papaya* Leaves Exhibits Anti-tumor Activity and Immunomodulatory Effects. *J. Ethnopharmacol.* 2010, *127* (3), 760–767.
- Owoyele, B. V.; Adebukola, O. M. Anti-inflammatory Activities of Ethanolic Extract of *Carica papaya* Leaves. *Inflammo Pharmacol.* 2008, *16* (4), 168–173.

- Panzarini, E.; Dwikat, M.; Mariano, S.; Vergallo, C.; Dini, L. Administration Dependent Antioxidant Effect of *Carica papaya* Seeds Water Extract. *Evidence-Based Comple. Altern. Med.: eCAM* 2014, 2014, article ID: 281508.
- 69. Parle, M.; Gurditta. Basketful Benefits of Papaya. Int. Res. J. Pharm. 2011, 2, 6-12.
- Pavan, R.; Jain, S.; Shraddha; Kumar, A. Properties and Therapeutic Application of Bromelain: A Review. *Biotechnol. Res. Int.* 2012, 2012, article ID: 976203.
- Pendzhiev, A. M. Proteolytic Enzymes of Papaya: Medicinal Applications. *Pharm. Chem. J.* 2002, *36* (6), 315–317.
- 72. Plantvillage; https://plantvillage.psu.edu/topics/papaya-pawpaw/infos; Accessed on March 12, 2020.
- Robert, E. P.; Odilo, D. Papaya. Chapter 11; In *Tropical Fruits.*, *Volume 1*; 2nd ed.; CABI: Wallingford, 2011; pp 389–400.
- Rajasekhar, P. Nutritional and Medicinal Value of Carica papaya. World J. Pharm. Pharm. Sci. 2017, 6 (8), 2559–2578.
- Rajkapoor, B.; Jayakar, B.; Kavimani, S.; Murugesh, N. Effect of Dried Fruits of Carica papaya Linn on Hepatotoxicity. *Biol. Pharm. Bull.* 2002, 25 (12), 1645–1646.
- Ranasinghe, P.; Ranasinghe, P.; Abeysekera, W. P. In vitro Erythrocyte Membrane Stabilization Properties of *Carica papaya* Leaf Extracts. *Pharm. Res.* 2012, 4 (4), 196–202.
- 77. Richard, M., The Papaya in Hawaii. HortScience Horts 2012, 47 (10), 1399-1404.
- 78. Saeed, F.; Arshad, M.; Pasha, I.; Naz, R. Nutritional and Phyto-Therapeutic Potential of Papaya: Overview. *Int. J. Food Prop.* **2014**, *17* (7), 1637–1653.
- 79. Santana, L. F.; Inada, A. C.; Espirito-Santo, B. Nutraceutical Potential of *Carica papaya* in Metabolic Syndrome. *Nutrients* **2019**, *11* (7), 1608–1613.
- Sarala, N.; Paknikar, S. Papaya Extract to Treat Dengue: Novel Therapeutic Option? Ann. Med. Health Sci. Res. 2014, 4 (3), 320–324.
- Saran, P. L.; Choudhary, R. Drug Bioavailability and Traditional Medicaments of Commercially Available Papaya-A Review. *Afr. J. Agric. Res.* 2013, *8*, 3216–3223.
- Saran, P. L.; Choudhary, R.; Solanki, I.; Devi, G. Traditional Medicaments Through Papaya in Northeastern Plains Zone of India. *Ind. J. Trad. Knowledge* 2015, 14, 537–543.
- Sarma, H. N.; Mahanta, H. C. Modulation of Morphological Changes of Endometrial Surface Epithelium by the Administration of Composite Root Extract in Albino Rat. *Contraception* 2000, 62 (1), 51–54.
- Satrija, F.; Nansen, P.; Bjorn, H. Effect of Papaya Latex against Ascarissuum in Naturally Infected Pigs. J. Helminthol. 1994, 68 (4), 343–346.
- 85. Satrija, F.; Nansen, P.; Murtini, S.; He, S. Anthelmintic Activity of Papaya Latex Infections in Mice. J. Ethnopharmacol. **1995**, 48 (3), 161–164.
- Senthilvel, P.; Pandian, L.; Kumar, K. Flavonoid from *Carica papaya* Inhibits NS2B-NS3 Protease and Prevents Dengue-2 viral assembly. *Bioinformation* 2013, *9*, 889–895.
- Shaziya, B.; Goyal, P. K. Anthelmintic Effect of Natural Plant (*Carica papaya*) Extract against the Gastrointestinal Nematode, Ancylostoma caninum in Mice. *Int. Res. J. Biol. Sci.* 2012, 1 (1), 2–6.
- Starley, I. F.; Mohammed, P.; Schneider, G.; Bickler, S. W. The Treatment of Pediatric Burns using Topical Papaya. *Burns: J. Int. Soc. Burn Inj.* 1999, 25 (7), 636–639.

- Stepek, G.; Buttle, D. J.; Duce, I. R.; Lowe, A. Assessment of the Anthelmintic Effect of Natural Plant Cysteine Proteinases Against the Gastrointestinal Nematode, Heligmosomoid espolygyrus, in vitro. *Parasitology* 2005, *130* (2), 203–211.
- 90. Stokes, T. Papaya Male Contraceptive. Trends Plant Sci. 2001, 6 (4), 143-148.
- Subenthiran, S.; Choon, T. C.; Cheong, K. C. *Carica papaya* Leaves Juice Significantly Accelerates the Rate of Increase in Platelet Count among Patients with Dengue Fever and Dengue Hemorrhagic Fever. *Evidence-Based Complem. Altern. Med.: Ecam* 2013, article ID: 616737.
- Sultana, A.; Afroz, R.; Yasmeen, O.; Aktar, M.; Yusuf, M. Anti-Inflammatory Effect of Ethanolic Extract of *Carica papaya* Leaves and Indomethacin in Cotton Pellet Induced Granuloma in Animal Model. *J. Curr. Adv. Med. Res.* 2019, *6*, 2–5.
- Sultana, A.; Khan, A.; Afroz, R.; Yasmeen, O.; Aktar, M.; Yusuf, M. A. Comparison of Anti-Inflammatory Effect of Ethanolic Extract of *Carica papaya* Leaves and Indomethacin in Carrageenan Induced Rat Paw Edema Animal Model. *J. Sci. Foundation* 2019, *16*, 49–53.
- Teng, W. C.; Chan, W.; Suwanarusk, R. In vitro Antimalarial Evaluations and Cytotoxicity Investigations of *Carica papaya* Leaves and Carpaine. *Nat. Product Commun.* 2019, 14 (1), 33–36.
- Udoh, P.; Essien, I.; Udoh, F., Effects of *Carica papaya* Seeds Extract on the Morphology of Pituitary-gonadal Axis of Male Wistar Rats. *Phytotherapy Res.* 2005, 19 (12), 1065–1068.
- Verma, S.; Varma, R.; Singh, S., Medicinal and Pharmacological Parts of *Carica papaya*: A Review. *Ind. J. Drugs* 2017, 5, 88–93.
- 97. Vij, T.; Prashar, Y. Review on Medicinal Properties of *Carica papaya* Linn. *Asian Pacific J. Trop. Dis.* **2015**, *5* (1), 1–6.
- Wikipedia Papaya; https://en.wikipedia.org/wiki/List_of_countries_by_papaya_ production; Accessed on March 16, 2020.
- 99. Zeleke, G.; Kebebe, D.; Mulisa, E.; Gashe, F. In Vivo Antimalarial Activity of the Solvent Fractions of Fruit Rind and Root of *Carica papaya* Linn against *Plasmodium berghei* in Mice. *J. Parasitol. Res.* **2017**, *2017*, 1–9.

REVIEW ARTICLE

Computational Design of Molecularly Imprinted Polymers in Drug Delivery Systems: A Comprehensive Review

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> Abstract: *Background:* Nowadays, biomedical research has been focusing on the design and development of new drug delivery systems that provide efficient drug targeting. The molecularly imprinted polymers (MIPs) have attracted wide interest and play an indispensable role as a drug carrier. Drug delivery systems based on MIPs have been frequently cited in the literature. They are cross-linked polymers that contain binding sites according to the complementary structure of the template molecules. They possess distinctive features of structure predictability and site recognition specificity. Versatile applications of MIPs include purification, biosensing, bioseparation, artificial antibodies, and drug delivery. An ideal MIPs should include features such as biocompatibility, biodegradability, and stability.

ARTICLE HISTORY

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DOI: 10.2174/1567201819666220427134549 **Objective:** In this article, we elaborate on the historic growth, synthesis, and preparation of different MIPs and present an updated summary of recent advances in the development of new drug delivery systems which are based on this technique. Their potential to deliver drugs in a controlled and targeted manner will also be discussed.

Conclusion: MIPs possess unique advantages, such as lower toxicity, fewer side effects, and good therapeutic potential. They offer administration of drugs by different routes, *i.e.*, oral, ocular or transdermal. Despite several advantages, biomedical companies are hesitant to invest in MIPs based drug delivery systems due to the limited availability of chemical compounds.

Keywords: Molecular imprinted polymer, Methacrylic acid, density functional theory, polymerization process.

1. INTRODUCTION

Molecularly Imprinted Polymers (MIPs) are cross-linked polymers having specific binding sites/cavities for the target molecule [1]. They have been widely used for various pharmaceutical applications due to their specific molecular recognition and stable physicochemical properties [2-5]. They are man-made synthetic polymers that possess the capacity to distinguish and bind specific substrates with high accuracy. MIPs are known for their robustness and resembling antibodies capabilities. They are designed with the involvement of various interdisciplinary techniques e.g., polymer, organic, analytical, physical, and biochemistry [6-8]. Nowadays, MIPs play a vital role in the design of sustained and controlled drug delivery systems [9, 10]. They are stable and biocompatible in nature but need to be further studied because they are mostly synthesized using organic solvents. Current researches focus on the utilization of computer-based high-throughput screening techniques for transformation from lab to clinical applications [11, 12]. MIPs have significant prospects in the targeted diagnosis and treatment of tumors [13]. The use of molecular imprinting technology with computer-based design has potential in drug delivery systems [10]. There have been extensive papers published on the topic of molecularly imprinted polymers in the various journals which have been surveyed through SciFinder® and Scopus. From the explored search term, "Molecularly Imprinted Polymer", the total number of papers was 10143 in Scopus, and for SciFinder[®] the number was 21775. Similar research is accomplished for "Molecularly Imprinted Polymer in Drug Delivery". From the explored search, it was noted that there is a sharp growth in the number of published papers (Fig. 1). This emphasized the importance of molecularly imprinted polymers in drug delivery systems and the growing interest of the scientific community in this field. The renowned scientists Wulff, Mosbach, and Klaus Mosbach reported the landmark research on molecularly imprinted polymers which can be synthesized through the copolymerization process [14-16].

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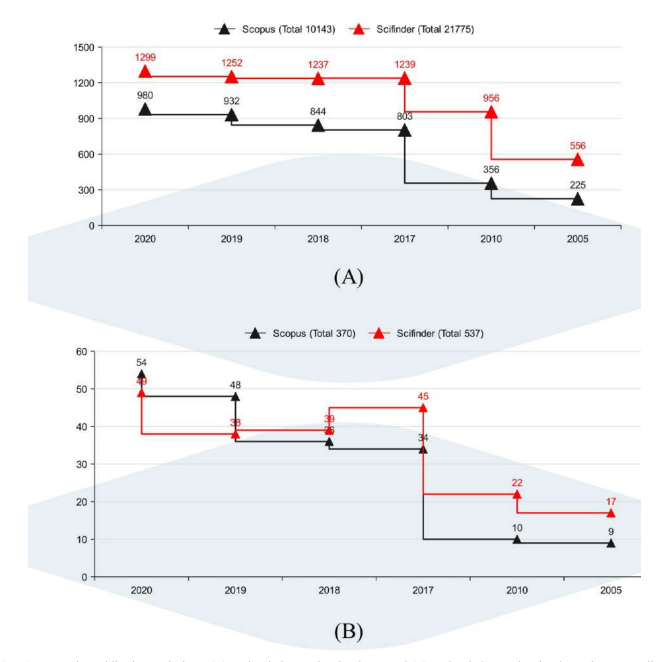


Fig. (1). Year wise publication analysis on (A) Molecularly Imprinted Polymer and (B) Molecularly Imprinted Polymer in Drug Delivery (Scopus & SciFinder[®] database). (*A higher resolution / colour version of this figure is available in the electronic copy of the article*).

MIPs are cross-linked polymers prepared by the processing of cross-linkers and the functional monomers in the presence of the template molecules. Their ability to respond to a variety of external stimuli (such as pH, temperature, photonic irradiation, electronic, and magnetic field interference) has earned them significant attention in drug delivery [17]. Ideal MIPs require mainly molecular recognition ability and exhibit specific site binding capability for targeting and controlled release of drugs [18]. Molecular recognition features of MIPs mainly depend upon the shape, size, and interactions of the template and imprinted cavities [19]. The fabrication of MIPs involves three main components which are responsible for their specific functionality *i.e.*, template (target molecule), functional monomer, and excess of crosslinker [20]. The various potential applications of MIPs in the field of biomedicine [4, 7, 21] are illustrated in Fig. (2).

1.1. Preparation of MIPs

MIPs are usually prepared by polymerization process through covalent or non-covalent interactions and in some cases a combination of both [20, 22]. MIPs utilize the template- mediated polymerization process. The main production components of MIPs include backbone, and functional monomer, a template molecule and a cross-linker [3]. The fabrication mechanism of MIPs involves the preparation of a solution that contains a backbone monomer, functional mon-

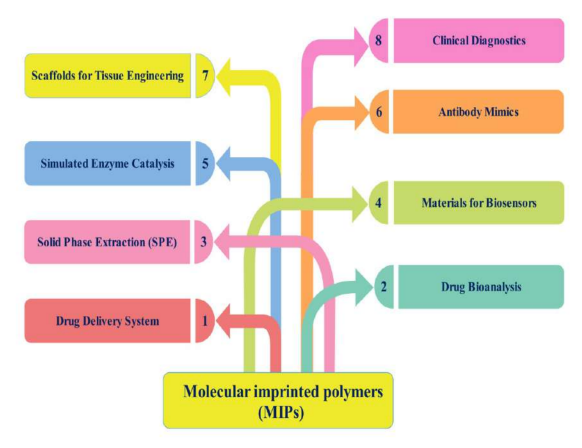


Fig. (2). Applications of Molecularly imprinted polymers (MIPs). (A higher resolution / colour version of this figure is available in the electronic copy of the article).

omers, and cross-linker in the presence of the template molecule (target molecule) [23]. The overall process is divided into three essential steps which are shown in Fig. (3).

Step 1 involves the assembly of the pre-polymerization network, followed by polymerization in step 2 and the removal of the template in step 3, thus liberating the binding site [24]. During this process, the template molecule is allowed to interact with functional groups of monomer solutions in the presence of a cross-linker so as to form a stable self-assembled complex [25]. Specific recognition of template molecules is influenced by the type of interaction established between them whether covalent or non-covalent [26]. As such covalent interactions are selective and stronger than non-covalent interactions. After the polymerization, the template molecule is cleaved from the resultant molecular imprinting polymers.

1.2. Building Blocks for MIPs

The main building blocks for the assembly of MIPs are backbone monomers, functional monomers, and crosslinkers. They all play a specific and critical role in the performance efficiency of smart polymers.

1.2.1. Backbone Monomers

Backbone monomers are the moieties that bind to the template molecule and may offer non-specific binding opportunities. They play a role in the swelling of the system and the nature of the monomers used influences the intensity of the swelling effect [20]. The backbone monomers also control the manner in which the template molecule is released from polymer assembly. They facilitate the multiple point interaction with the template molecule and enhance template binding capacity [27].

1.2.2. Functional Monomers

Functional monomers are another important part of the imprinted polymer process which provides complementary interactions with the template molecules. These monomers are used to improve the functionality features and properties of the final imprinting polymers [3].

The functional monomer mainly involves an arrangement of a cross-linked polymer matrix around the template molecule in the presence of a cross-linker [27]. Overall stability and rigid polymer matrix structure depend upon the crosslinker to the functional monomer molar ratio (C/M). The specificity, selectivity, and efficiency of imprinted polymers also depend upon the C/M ratio. Thus, a low C/M ratio in the range of 1 to 3 results in the formation of closely located binding sites while a large C/M ratio in the range of 7 to 10 results in a decrease in the number of formed binding sites especially in the case of non-covalent interactions. The optimal ratio of functional monomer to template molecule reported was 4:1 [28]. This rigid polymer matrix structure facilitates the subsequent step of removal of the template molecule. The selectivity and binding capacity of imprinted polymer is also influenced by the type and amount of the cross-

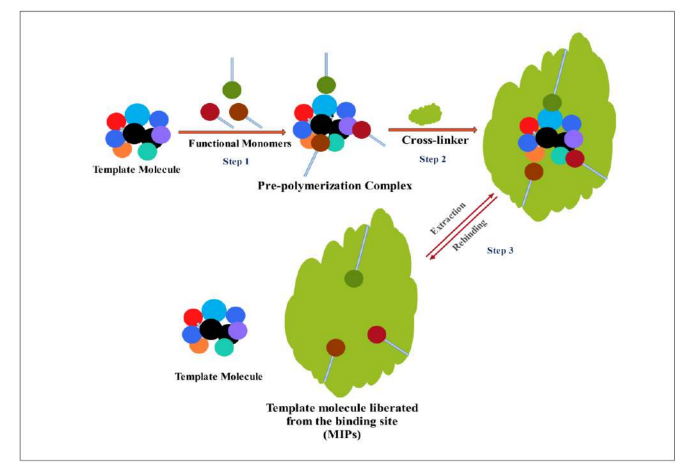


Fig. (3). Schematic interpretation of the formation of MIPs. (A higher resolution / colour version of this figure is available in the electronic copy of the article).

linker used in the polymerization process. Thus, the selection of functional monomers is important for the specificity, selectivity, and efficiency of imprinted polymers [3].

1.2.3. Cross-linkers

Cross-linker controls the morphology and serves to stabilize the imprinted binding sites in imprinted polymers. They provide mechanical stability and retain their molecular recognition capability [29]. They are also accountable for fixing the special orientation of the functional monomer relative to the template molecule and providing rigidity to the polymer structure. Optimizing the amount and type of crosslinker strongly influences the final size, yield, and type of final product *i.e.*, gel-type, macroporous, or a microgel powder [26]. The mechanical stiffness or flexibility of the polymer network and site confirmation is also controlled by the cross-linker [30]. The quantity of the cross-linking agent is a key factor and it should be high enough to maintain overall stability even after template removal [31]. The hydrophilicity and hydrophobicity of a cross-linker with respect to reaction medium influences physicochemical stability and sitespecificity of the imprinted polymer [27].

1.2.4. Templates

The template molecule also plays a vital role throughout the molecular imprinting processes. A template molecule should have the following features [20, 23].

- i. It should contain polymerizable groups.
- ii. It must be resistant to moderately elevated temperatures or UV radiation exposure.
- iii. It should be chemically inert during polymerization.
- iv. It must attain a definite orientation during molecular imprinting processes.
- v. It should leave cavities without any change in size, shape, and molecular interactions of MIPs.

1.2.5. Photo-initiators

A photo-initiator initiates the polymerization process upon irradiation after the absorption of light. During this process, the photo-initiator absorbs the photons and forms reactive species, which convert from a single state to a triplet state and thus initiate a chemical reaction [32]. Photoinitiators should exhibit several important features [23, 33]:

- 1. High absorption at the exposure wavelength.
- 2. High reactivity towards the monomer.
- 3. Adequate solubility in the system.
- 4. Thermal stability.
- 5. It should be non-toxic, biocompatible, and easy to handle.
- 6. Cheap and low production cost.

Table 1. List of commonly used building blocks for assembly of MIPs.

Functional monomers (FM)	Acrolein, Acrylamide, Acrylic acid, Acrylonitrile, Allylamine p-Divinylbenzene, N, N-Diethylamino ethyl methacrylate Ethylene glycol dimethacrylate, Itaconic acid, Methacrylic acid
	N,N'-Methylene bisacrylamide, Urocanic acid, Vinyl benzene1-Vinylimidazole, 2-Vinylpyridine, 4-Vinylpyridine
Cross-Linkers (CL)	Ethylene glycol dimethacrylate (EGDMA), Trimethylolpropane trimethacrylate (TRIM), <i>p</i> -divinylbenzene (DVB), and <i>N</i> , <i>N</i> ⁻ methylene bisacrylamide (MBAA)
Photo-initiators	2,2' -Azobis(2-methylpropixonitrile) (xAIBN), Benzoin, 2-Isopropylthioxanthone, Benzoyl peroxide, Ethyl-2-chloro- propionate, Ammonium persulphite, Azo-bi- isobutyro nitrile

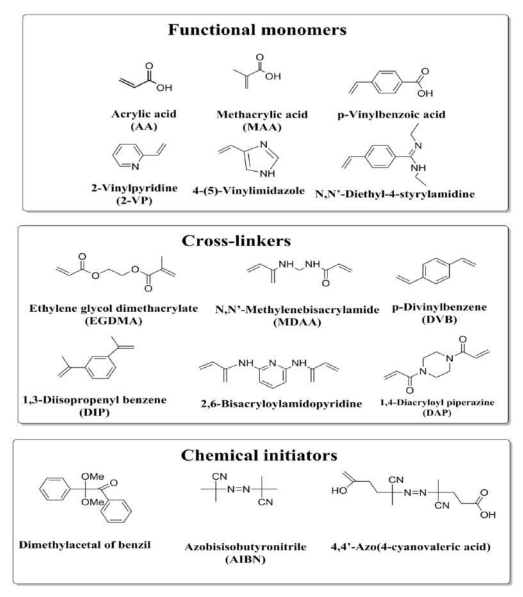


Fig. (4). The chemical structures of some functional monomers, cross-linkers, and chemical initiators used in MIPs synthesis.

Commonly used functional monomers (FM), crosslinkers (CL), and photo-initiators [3, 23] are listed in Table 1 and Fig. (4).

1.3. Computer-assisted Design of MIPs

The design of molecular imprinting polymers through computational tools is very effective. The main advantage of the computer-assisted rational design of MIPs is cost effectiveness especially in the screening of functional monomers [34]. Computer-assisted design of MIPs plays a significant role in the development of the biomedical field. At present, experimental trials are time-consuming and cause the wastage of chemicals [35]. It is possible to calculate energy and carry out a preliminary screening of the structural configuration of polymer before the start of the actual experiment through the use of computer simulations. This strategy saves a lot of time and money [36]. The main focus lies in the comparison of the binding energy of complexes formed between a template molecule and functional monomers. The success of MIPs generally depends on the selection of an optimal functional monomer and its configuration with the template molecule [23]. The selection of optimal functional monomers is a time-consuming process. Many computational approaches and a virtual library of functional monomers are reported in the literature as guiding documents for optimizing the imprinting conditions [37]. Various molecular models are reported for the design of MIPs such as density functional theory (DFT) [38, 39], Molecular Operating Environment (MOE) [35], and the Hartree-Fock (HF) method [40, 41]. Computational tools facilitate the understanding of intermolecular interactions in molecular imprinting. In order to understand the configuration of template-monomer complexes in MIPs at the molecular level, a model is a setup in computational tools. Before the start of computational tools, a virtual library of a few functional monomers is developed for the conformational optimization process with template molecules [42]. Then, the most stable template-monomer complex is searched on the basis of the interaction energy (E) of a specific feature as per the selected model. A higher value of interaction energy indicates the higher affinity and selectivity between template-monomer complexes [43]. Table 2 records some of the success stories involving the computer design of MIPs.

1.4. Characterization of the MIPs

Different techniques have been used to characterize the physicochemical properties of molecular imprinting polymers [53]. In some cases, more than one techniques are used to evaluate these properties. They are selected on the basis of synthesis and further processes involved in the polymerization step. The techniques are summarized in Fig. (5) [54]. The interdisciplinary nature of MIPs greatly affects the se-

lection of suitable methods for commercial applications and the interpretation of the data for regulatory requirements [55].

1.4.1. Thermogravimetric Analysis (TGA)

TGA is used to characterize the decomposition and thermal stability of MIPs. Physical and chemical changes of MIPs due to heating are examined in terms of the percentage of weight loss as a function of temperature. The TGA thermogram represents the difference between the decomposition stage of monomer and MIP at high-temperature conditions. Under dry nitrogen environment conditions, the samples are heated to a particular temperature at a rate of 10°C/min for examination. The weight percentage of each ensuing mass change of MIPs increases as the temperature rises [56, 57].

1.4.2. Adsorption Isotherm

The adsorption isotherms studies play an important role in the theoretical evaluation and interpretation of thermodynamic parameters of MIPs. A number of isotherms have been successfully employed to calculate the binding properties of MIPs. The adsorption isotherm equations give details regarding the binding equilibrium between target molecules and MIPs. With the help of these equations, the binding affinity and homogeneity of the binding-site distribution can be measured [58-59]. Several studies have been reported in the literature that use different models *i.e.*, simple Langmuir, SIPs models, Bi-Langmuir models, Jovanovic model, Bi-Jovanovic model, and Freundlich-Jovanovic model [60].

1.4.3. Scanning Electron Microscopy (SEM)

Scanning Electron Microscopy is used to observe the surface microscopic characteristics of the imprinted polymers (shape and size). It provides a direct image of the topographical nature of the surface from all the emitted secondary electrons. For surface morphology, samples are mounted on the sample holder and a thin layer of gold is applied to the sample surface for imaging purposes. Its high resolution

S. No.	Template	Computational Approach/Theory Software		Year	Refs.
1	Furosemide	Density functional theory (DFT) Gaussian 3		2010	[44]
2	Chlorogenic Acid	Hartree-Fock (HF) method	Hartree-Fock (HF) method Gaussian 3		[45]
3	Allopurinol	Hartree-Fock (HF) method Gaussian 9		2012	46]
4	Methadone	Density functional theory (DFT)	Gaussian 3	2012	[47]
5	Capsaicin	Simplified molecular-input files entry string ChemDB tool		2014	[48]
6	Melamine	Density functional theory (DFT) Gaussian 9		2015	[49]
7	Andrographolide	Restricted Hartree-Fock (RHF) semi- empirical method HyperChem 8.0.10		2017	[50]
8	Levetiracetam	acetam MMFF94x force field, Molecular Operating Environment (MOE) Gaussian 9		2018	[35]
9	Clenbuterol	nbuterol Density functional theory (DFT) Gaussian 9		2018	[34]
10	Bilobalide	balide Molecular orbital (MO) calculations Gaussian 16		2020	[51]
11	Bisphenol A	Hartree-Fock restricted (RHF)	Gaussian 9	2020	[52]

Table 2.Computer-assisted design of MIPs.

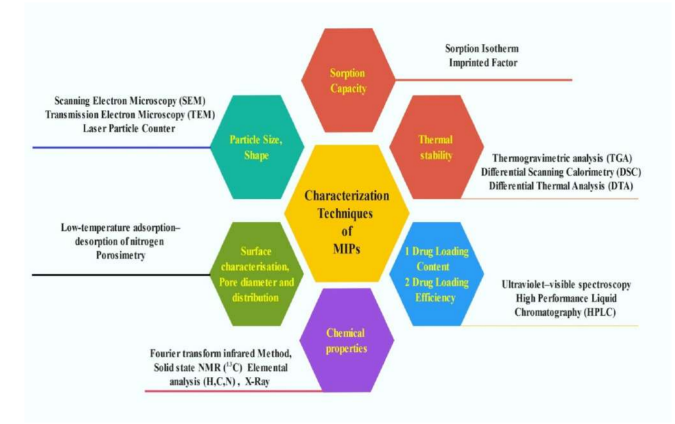


Fig. (5). Techniques used for the characterization of the MIPs. (A higher resolution / colour version of this figure is available in the electronic copy of the article).

makes it one of the best suitable methods for surface morphology studies [61-62].

1.4.4. Fourier Transform Infrared Method

FTIR spectroscopy is a suitable method to determine the functional groups and types of bonds present in MIP [63]. The FTIR spectra (KBr pellet) of the prepared polymers MIP, monomer, the cross-linker, and the template coating are recorded in the range of 4000-400cm⁻¹. It also provides information about the formation of a new bond in the MIP and rationalizes the mechanisms of recognition during the imprinting process [64, 65].

1.4.5. UV Spectroscopy

UV spectroscopy is providing information regarding the saturation of template molecules with functional monomer building blocks. It mainly provides information about the binding capacity of functional monomers with the template molecule. As per the literature reviewed, several studies have been reported for the determination of the binding capacity of MIPs by the UV Spectroscopy method [23, 66].

1.4.6. Nuclear Magnetic Resonance (NMR) spectroscopy

The NMR spectroscopy is a useful tool to investigate the interaction between functional monomer and template in the pre-polymerization process. The chemical shift studies and nuclear overhauled effect (NOE) allow the calculation of dissociation constants and the types of interactions occurring in the pre-polymerization mixture. It also recognizes the specific sites in interacting structures that engage in the formation of MIPs [67-69].

2. MIPS-MEDIATED DRUG DELIVERY SYSTEM

MIPs have a potential role in site-specific drug delivery systems due to their good molecular recognition performance [70]. They are perfectly complementary to the target biomolecule and are also known as "artificial antibodies" [71]. Various key features which are worth consideration for MIPs development include the rigidity of the polymer structure, high flexibility, Good accessibility, mechanical, thermal, and chemical stability [9,72].

MIPs are showing an increase in popularity, owing to their recognition characteristics, as can be observed from the increased number of investigated reports related to the application of MIPs in drug delivery systems [3, 10]. We reviewed all available databases and comprehensively summarize the utility of MIPs as innovative pharmaceutical polymers in Table **3**.

Moreover, stimuli-responsive molecularly imprinted polymers such as temperature-responsive, pH-responsive, photo-responsive, and magnetism responsive has drawn the greatest attention. Stimuli-responsive MIPs are comprehensively summarized in Table **4**. Whereas, methacrylic acid (MAA) is the most commonly used functional monomer, prevalent cross-linkers are ethylene glycol dimethacrylate (EGDMA) and N,N O-methylene bisacrylamide (MBA). It is

Table 3.	List of MIPs-based drug carrier systems.
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S. No.	Drug/Molecule	Carrier System	Main Components FM/ CL/ Initiator	Year	Refs.
1.	Atropine	Microspheres	MAA,TRIM/ EPI, Genipin		[73]
2.	Sunitinib	Theranostic systems	MAA/ EGDMA/AIBN Fluorescent marker: Rhodamine 6G	2020	[13]
3.	Diclofenac	Nanospheres	MAA/ EGDMA/ AIBN	2018	[74]
4.	Fenbufen	Carbon Nanotubes	4-vinylpyridine (4-VP)/ EGDMA	2018	[75]
5.	Mitomycin C	Cryogel Membranes	MAH/ HEMA, MBAAm	2018	[76]
6.	Donepezil	Microparticles	MAA, GMA, HEMA/ EGDMA	2016	[77]
7.	Trinitroglycerin	Nanoparticles	MAA/ TRIM	2016	[78]
8.	L-DOPA	Nanosponge	β -cyclodextrin/ 1,1'-Carbonyldiimidazole	2016	[79]
9.	Olanzapine	Nanoparticles	MAA/ EGDMA/ AIBN	2016	[80]
10.	Azithromycin	Nanoparticles	FMAA/ EDMA/AIBN	2015	[81]
11.	Sitagliptin and Metformin	Nanoparticles	MAA,MMA/ EGDMA	2015	[82]
12.	Nicotine	MIPs	MAA/ EGDMA	2014	[83]
13.	Propranolol HCL	Polymer complex	MAA/ EGDMA	2013	[84]
14.	Tetracycline	Microporous MIPs	MAA/ EGDMA/ AIBN	2013	[85]
15.	Dipyridamole	Microspheres	MAA/ EGDMA/AIBN	2011	[86]
16.	Bromhexine	MIPs	MAA/ EGDMA/AIBN	2011	[87]
17.	Citalopram	MIPs	MAA/ EGDMA/AIBN	2011	[88]
18.	Flufenamic Acid	MIPs	MAA, NIPAAm/EGDMA/ AIBN	2011	[89]
19.	Glycyrrhizic Acid	MIPs	MAA, DMAEMA, HEMA/EGDMA/AIBN	2010	[90]
20.	Tramadol	MIPs	MAA/EGDMA/ AIBN	2010	[91]
21.	Nicotinamide	Microspheres	MAA/EGDMA/AIBN	2010	[92]
22.	5-Fluorouracil	Hydrogel nanospheres	MAA/ EGDMA/AIBN	2009	[93]
23.	α-Tocopherol	MIPs	MAA/TRIM/AIBN	2008	[94]
24.	Levonorgestrel	MIPs	MAA/TRIM/AIBN	2008	[95]
25.	Propranolol	Granules and beads for matrix tablets	MAA, NAA/ EGDMA/AIBN	2000	[96]

S. No	Drug/ Molecule	Carrier System	Stimuli-responsive	Main Components FM/ CL/ Initiator	Year	Refs.
1.	Curcumin	Nanocomposite	pH-responsive	pH-responsive AA, β-CD/TEOS,HEMA/AIBN		[97]
2.	5-Fluorouracil	Nanoparticles	Reduction-responsive	iPOx/ DTDPA	2020	[98]
3.	Doxorubicin	Graphene Quantum Dots	pH-responsive	HEMA/MBA	2020	[99]
4.	Acyclovir Valacyclovir	Hydrogel Contact Lenses pH-responsive		2020	[100]	
5.	Paclitaxel	Microparticles	pH-responsive	MAA, HEMA/ EGDMA, TRIM	2019	[101]
6.	Capecitabine	Capecitabine Floating MIPs pH-responsive MAA/MPDE,MPDB, CPCE, CPCP/AIBN		2019	[102]	
7.	Acyclovir	Microspheres	Photo- responsive	ADDDM/TEA, TMA	2018	[103]
8.	5-Fluorouracil	Microspheres	Thermo- Magnetic bi- responsive	NIPAM/MBA	2017	[104]
9.	Quercetin	Nanogel	Magnetic responsive	VI/TG	2016	[105]
10.	Letrozole	Nanoparticles	Magnetic responsive	MAA/TRIM/AIBN	2016	[106]
11.	5-Fluorouracil	Carbon Nanospheres	Thermo- Magnetic bi- responsive	NIPAM/ MBA/APS	2016	[107]
12.	5-Fluorouracil	Microspheres	Thermo- Magnetic bi- responsive	NIPAM/ MBAA/ APS	2015	[108]
13.	Salicylic Acid	Sol-Gel Polymers	pH-responsive	1-(4-vinylphenyl)-3-(3,5- bis(trifluoromethyl)phenyl)urea, APTES TMPS/ TEOS	2014	[109]
14.	Mitomycin C	Nanoparticles	Thermo responsive	MAH/ HEMA, EGDMA	2014	[110]
15.	Diclofenac	Carbon nanotubes (CNTs)	Electro responsive	MAA/ EGDMA/AIBN	2013	[111]
16.	Aspirin	Magnetic nanoparticles	Magnetic responsive	MAA/TRIM	2009	[24]

 Table 4.
 List of Stimuli-responsive MIPs-based drug carrier systems.

generally observed that pH-responsive MIPs offer great potential than other stimuli-responsive ones.

CONCLUSIONS AND FUTURE PERSPECTIVE

In recent years, novel MIPs have received substantial attention and investigation due to their excellent recognition properties. Stimuli-responsive molecularly imprinted polymers have drawn the greatest attention and the mechanism of response can be understood in both theoretical and practical terms. Various stimuli-responsive molecularly imprinted polymers include thermo-responsive, pH-responsive, photo responsive MIPs, biomolecule responsive and ion responsive MIPs. They possess unique advantages, such as lower toxici-

ty, fewer side effects, and good therapeutic potential. They offer administration of drugs by different routes, *i.e.*, oral, ocular, or transdermal. Despite several advantages, biomedical companies are hesitant to invest in MIPs-based drug delivery systems due to the limited availability of chemical compounds. Consequently, MIPs have not yet reached clinical trial phases, although this technology has a vast prospective for creating novel dosage forms and devices that may be useful for the treatment and diagnosis of various diseases. Future studies and the development of more clinical trials will lead to the use of MIPs as dual integration tools for therapy and diagnostic (theranostic) purposes for patients. More strategies are needed to realize the targeted efficacy of MIPs.

LIST OF ABBREVIATIONS

AA	=	Acrylic acid
ADDDM	=	<i>N</i> -(4-((4-amino-2,6- dimethoxyphenyl)diazenyl)-3,5- dimethoxyphenyl)methacrylamide
ADDDMN	=	(4-((4-amino-2,6- dimethoxy- phenyl)diazenyl)-3,5- dimethoxyphenyl)methacrylamide
AIBN	=	2,2'-Azoisobutyronitrile
APS	=	Ammonium persulfate
APTES	=	3-Aminopropyl)triethoxysilane
CPCE	=	4-Myanophenyl cyclohexyl ethylene
CPCP	=	4-Cyanophenyl cyclohexyl propylene
DMAEMA	=	2-(dimethylamino)ethyl methacrylate
DTDPA	=	3,3'-dithiodipropionic acid
EGDMA	=	Ethylene glycol dimethacrylate
EPI	=	Epichlorohydrin
GMA	=	Glycidyl methacrylate
HEMA	=	2-hydroxyethylmethacrylate
HEMA	=	Hydroxyethyl methacrylate
iPOx	=	2-Isopropenyl-2-oxazoline
MAA	=	Methacrylic acid
MAH	=	N-Methacryloyl-L-histidine methyl ester
MBA	=	N,N'-Methylenebisacrylamide ethylene
MBAA	=	N,N'-Methylenebisacrylamide
MBAAm	=	methylene bisacrylamide
MPDB	=	4-Methylphenyl dicyclohexyl butylene
MPDE	=	4-methylphenyl dicyclohexyl ethylene,
NAA	=	N-acryloyl-alanine
NIPAAm	=	N-Isopropyl acrylamide
NIPAM	=	N-Isopropylacrylamide
TEA	=	Triethanolamine,
TEOS	=	Tetraethyl orthosilicate
TG	=	Tragacanth Gum
TMA	=	Trimethacrylate
TMPS	=	Trimethoxyphenylsilane
TRIM	=	Trimethylolpropane trimethacrylate
VI	=	N-Vinyl imidazole
β-CD	=	Acrylated β-cyclodextrin
	ITC	

HIGHLIGHTS

Concept of Molecularly Imprinted Polymer

Role in drug delivery systems

Focus on Computer-Aided Design

Methacrylic acid as main functional monomers

Recent application of Molecularly Imprinted Polymer in drug delivery system

CONSENT FOR PUBLICATION

Not applicable.

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CONFLICT OF INTEREST

The authors declare no conflict of interest, financial or otherwise.

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REFERENCES

- Orowitz, T.E.; Ana Sombo, P.P.A.A.; Rahayu, D.; Hasanah, A.N. Microsphere polymers in molecular imprinting: Current and future perspectives. *Molecules*, 2020, 25(14), E3256. http://dx.doi.org/10.3390/molecules25143256 PMID: 32708849
- [2] Guan, G.; Liu, B.; Wang, Z.; Zhang, Z. Imprinting of molecular recognition sites on nanostructures and its applications in chemosensors. *Sensors (Basel)*, 2008, 8(12), 8291-8320. http://dx.doi.org/10.3390/s8128291 PMID: 27873989
- [3] Vasapollo, G.; Sole, R.D.; Mergola, L.; Lazzoi, M.R.; Scardino, A.; Scorrano, S.; Mele, G. Molecularly imprinted polymers: Present and future prospective. *Int. J. Mol. Sci.*, 2011, *12*(9), 5908-5945. http://dx.doi.org/10.3390/ijms12095908 PMID: 22016636
- [4] Rutkowska, M.; Płotka-Wasylka, J.; Morrison, C.; Wieczorek, P.P.; Namieśnik, J.; Marć, M. Application of molecularly imprinted polymers in analytical chiral separations and analysis. *Trends Analyt. Chem.*, 2018, 102, 10291-102. http://dx.doi.org/10.1016/j.trac.2018.01.011
- [5] Vaneckova, T.; Bezdekova, J.; Han, G.; Adam, V.; Vaculovicova, M. Application of molecularly imprinted polymers as artificial receptors for imaging. *Acta Biomater.*, **2020**, *101*, 444-458. http://dx.doi.org/10.1016/j.actbio.2019.11.007 PMID: 31706042
- [6] Mayes, A.G.; Whitcombe, M.J. Synthetic strategies for the generation of molecularly imprinted organic polymers. *Adv. Drug Deliv. Rev.*, 2005, 57(12), 1742-1778. http://dx.doi.org/10.1016/j.addr.2005.07.011 PMID: 16225958
- [7] Takeuchi, T.; Sunayama, H. Molecularly Imprinted Polymers. In: Kobayashi, S.; Müllen, K., Eds.; Encyclopedia of Polymeric Nanomaterials; Springer Berlin Heidelberg: Berlin, Heidelberg, 2021, pp. 1-5.
- [8] Bedwell, T.S.; Whitcombe, M.J. Analytical applications of MIPs in diagnostic assays: Future perspectives. *Anal. Bioanal. Chem.*, 2016, 408(7), 1735-1751.
 - http://dx.doi.org/10.1007/s00216-015-9137-9 PMID: 26590560
- Zaidi, S.A. Molecular imprinted polymers as drug delivery vehicles. *Drug Deliv.*, 2016, 23(7), 2262-2271. http://dx.doi.org/10.3109/10717544.2014.970297 PMID: 25317753
- [10] Bodoki, A.E.; Iacob, B.C.; Bodoki, E. Perspectives of molecularly imprinted polymer-based drug delivery systems in cancer therapy. *Polymers (Basel)*, 2019, 11(12), E2085.
- http://dx.doi.org/10.3390/polym11122085 PMID: 31847103 [11] Khodadadian, M.; Ahmadi, F. Computer-assisted design and synthe
 - sis of molecularly imprinted polymers for selective extraction of acetazolamide from human plasma prior to its voltammetric determination. *Talanta*, **2010**, *81*(4-5), 1446-1453. http://dx.doi.org/10.1016/j.talanta.2010.02.049 PMID: 20441921
- [12] Saylan, Y.; Akgönüllü, S.; Yavuz, H.; Ünal, S.; Denizli, A. Molecularly imprinted polymer based sensors for medical applications. *Sensors (Basel)*, 2019, 19(6), E1279. http://dx.doi.org/10.3390/s19061279 PMID: 30871280
- [13] Parisi, O.I.; Ruffo, M.; Malivindi, R.; Vattimo, A.F.; Pezzi, V.; Puoci, F. Molecularly Imprinted Polymers (MIPs) as theranostic systems for sunitinib controlled release and self-monitoring in cancer therapy. *Pharmaceutics*, **2020**, *12*(1), E41.

http://dx.doi.org/10.3390/pharmaceutics12010041 PMID: 31947815

- [14] Wulff, G.; Sarhan, A.; Zabrocki, K. Enzyme-analogue built polymers and their use for the resolution of racemates. Tetrahedron Lett., 1973, 14(44), 4329-4332.
 - http://dx.doi.org/10.1016/S0040-4039(01)87213-0
- [15] Wulff, G. The role of binding-site interactions in the molecular imprinting of polymers. Trends Biotechnol., 1993, 11(3), 85-87. http://dx.doi.org/10.1016/0167-7799(93)90056-F PMID: 7763512
- [16] Mosbach, K. The promise of molecular imprinting. Sci. Am., 2006, 295(4), 86-91. http://dx.doi.org/10.1038/scientificamerican1006-86 PMID: 16989485
- [17] Xu, S.; Lu, H.; Zheng, X.; Chen, L. Stimuli-responsive molecularly imprinted polymers: Versatile functional materials. J. Mater. Chem. C Mater. Opt. Electron. Devices, 2013, 1(29), 4406-4422. http://dx.doi.org/10.1039/c3tc30496e
- Puoci, F.; Cirillo, G.; Curcio, M.; Parisi, O.I.; Iemma, F.; Picci, N. [18] Molecularly imprinted polymers in drug delivery: State of art and future perspectives. Expert Opin. Drug Deliv., 2011, 8(10), 1379-1393.
- http://dx.doi.org/10.1517/17425247.2011.609166 PMID: 21933031 [19] Chen, L.; Xu, S.; Li, J. Recent advances in molecular imprinting
- technology: Current status, challenges and highlighted applications. Chem. Soc. Rev., 2011, 40(5), 2922-2942. http://dx.doi.org/10.1039/c0cs00084a PMID: 21359355
- [20] Sellergren, B.; Hall, A.J. Molecularly imprinted polymers. In: Gate, P.A.; Steed, J.W.; Eds.; Supramolecular Chemistry: From Molecules to Nanomaterials; Wiley: Chichester, West Sussex, 2012. http://dx.doi.org/10.1002/9780470661345.smc137
- El-Schich, Z.; Zhang, Y.; Feith, M.; Beyer, S.; Sternbæk, L.; [21] Ohlsson, L.; Stollenwerk, M.; Wingren, A.G. Molecularly imprinted polymers in biological applications. Biotechniques, 2020, 69(6), 406-419.

http://dx.doi.org/10.2144/btn-2020-0091 PMID: 33000637

- [22] Fresco-Cala, B.; Batista, A.D.; Cárdenas, S. Molecularly imprinted polymer micro- and nano-particles. A review. Molecules, 2020, 25(20), E4740. http://dx.doi.org/10.3390/molecules25204740 PMID: 33076552
- Chen, L.; Wang, X.; Lu, W.; Wu, X.; Li, J. Molecular imprinting: [23] Perspectives and applications. Chem. Soc. Rev., 2016, 45(8), 2137-2211.
 - http://dx.doi.org/10.1039/C6CS00061D PMID: 26936282
- Kan, X.; Geng, Z.; Zhao, Y.; Wang, Z.; Zhu, J.J. Magnetic molecu-[24] larly imprinted polymer for aspirin recognition and controlled release. Nanotechnology, 2009, 20(16), 165601. http://dx.doi.org/10.1088/0957-4484/20/16/165601 PMID: 19420571
- Wulff, G.; Vesper, W.; Grobe-Einsler, R.; Sarhan, A. Enzyme-[25] analogue built polymers, 4. On the synthesis of polymers containing chiral cavities and their use for the resolution of racemates. Makromol. Chem., 1977, 178(10), 2799-2816. http://dx.doi.org/10.1002/macp.1977.021781004
- Algieri, C.; Drioli, E.; Guzzo, L.; Donato, L. Bio-mimetic sensors [26] based on molecularly imprinted membranes. Sensors (Basel), 2014, 14(8), 13863-13912. http://dx.doi.org/10.3390/s140813863 PMID: 25196110
- [27] Yan, H.; Row, K.H. Characteristic and synthetic approach of molecularly imprinted polymer. Int. J. Mol. Sci., 2006, 7(5), 155-178. http://dx.doi.org/10.3390/i7050155
- Kotrotsiou, O.; Kiparissides, C. Chapter 7 Water treatment by [28] molecularly imprinted materials. In: Thomas, S.; Pasquini, D.; Leu, S-Y.; Gopakumar, D.A., Eds.; Nanoscale Materials in Water Purification; Elsevier: Amsterdam, 2019, pp. 179-230. http://dx.doi.org/10.1016/B978-0-12-813926-4.00012-4
- Kupai, J.; Razali, M.; Buyuktiryaki, S.; Kecili, R.; Szekely, G. [29] Long-term stability and reusability of molecularly imprinted polymers. Polym. Chem., 2017, 8(4), 666-673. http://dx.doi.org/10.1039/C6PY01853J PMID: 28496524
- [30] McCluskey, A.; Holdsworth, C.I.; Bowyer, M.C. Molecularly imprinted polymers (MIPs): Sensing, an explosive new opportunity? Org. Biomol. Chem., 2007, 5(20), 3233-3244. http://dx.doi.org/10.1039/b708660a PMID: 17912377
- [31] Yoshimatsu, K.; Reimhult, K.; Krozer, A.; Mosbach, K.; Sode, K.; Ye, L. Uniform molecularly imprinted microspheres and nanoparti-

cles prepared by precipitation polymerization: The control of particle size suitable for different analytical applications. Anal. Chim. Acta, 2007, 584(1), 112-121.

- http://dx.doi.org/10.1016/j.aca.2006.11.004 PMID: 17386593
- [32] Ravve, A. Photosensitizers and Photoinitiators. In: Ravve, A.; Ed; Light-Associated Reactions of Synthetic Polymers; Springer New York: New York, NY, 2006, pp. 23-122. http://dx.doi.org/10.1007/0-387-36414-5 2
- [33] Schwalm, R. Photoinitiators and Photopolymerization. In: Buschow, K.H.J.; Cahn, R.W.; Flemings, M.C.; Ilschner, B.; Kramer, E.J.; Mahajan, S.; Veyssière, P., Eds.; *Encyclopedia of Materials: Science and Technology*; Elsevier: Oxford, **2001**, pp. 6946-6951. http://dx.doi.org/10.1016/B0-08-043152-6/01230-4
- [34] Zhang, B.; Fan, X.; Zhao, D. Computer-aided design of molecularly imprinted polymers for simultaneous detection of clenbuterol and its metabolites. Polymers (Basel), 2018, 11(1), E17. http://dx.doi.org/10.3390/polym11010017 PMID: 30960001
- [35] Attallah, O.A.; Al-Ghobashy, M.A.; Ayoub, A.T.; Tuszynski, J.A.; Nebsen, M. Computer-aided design of magnetic molecularly imprinted polymer nanoparticles for solid-phase extraction and determination of levetiracetam in human plasma. RSC Advances, 2018, 8(26), 14280-14292. http://dx.doi.org/10.1039/C8RA02379D
- Sliwoski, G.; Kothiwale, S.; Meiler, J.; Lowe, E.W., Jr. Computa-[36] tional methods in drug discovery. Pharmacol. Rev., 2013, 66(1), 334-395

http://dx.doi.org/10.1124/pr.112.007336 PMID: 24381236

- Khan, M.S.; Wate, P.S.; Krupadam, R.J. Combinatorial screening of [37] polymer precursors for preparation of $benzo[\alpha]$ pyrene imprinted polymer: An ab initio computational approach. J. Mol. Model., 2012, 18(5), 1969-1981.
- http://dx.doi.org/10.1007/s00894-011-1218-x PMID: 21877152
- [38] Wungu, T.D.K.; Marsha, S.E.; Widayani; Suprijadi. Density Functional Theory (DFT) study of Molecularly Imprinted Polymer (MIP) Methacrylic Acid (MAA) with D-glucose. IOP Conf. Series Mater. Sci. Eng., 2017, 214, 012004. http://dx.doi.org/10.1088/1757-899X/214/1/012004
- Ren, X.; Yang, L.; Li, Y.; Cheshari, E.C.; Li, X. The integration of [39] molecular imprinting and surface-enhanced Raman scattering for highly sensitive detection of lysozyme biomarker aided by density functional theory. Spectrochim. Acta A Mol. Biomol. Spectrosc., 2020, 228, 117764.

http://dx.doi.org/10.1016/j.saa.2019.117764 PMID: 31727516

- [40] Abdel Ghani, N.T.; Mohamed El Nashar, R.; Abdel-Haleem, F.M.; Madbouly, A. Computational design, synthesis and application of a new selective molecularly imprinted polymer for electrochemical detection. Electroanalysis, 2016, 28(7), 1530-1538. http://dx.doi.org/10.1002/elan.201501130
- [41] Sanadgol, N.; Wackerlig, J. Developments of smart drug-delivery systems based on magnetic molecularly imprinted polymers for targeted cancer therapy: A short review. Pharmaceutics, 2020, 12(9), E831

http://dx.doi.org/10.3390/pharmaceutics12090831 PMID: 32878127

- [42] Huang, Y.; Zhu, Q. Computational modeling and theoretical calculations on the interactions between spermidine and functional monomer (Methacrylic Acid) in a molecularly imprinted polymer. J. Chem., 2015, 2015216983, 1-9. http://dx.doi.org/10.1155/2015/216983
 - Nicholls, I.A.; Adbo, K.; Andersson, H.S.; Andersson, P.O.; Ankar-
- [43] loo, J.; Hedin-Dahlström, J.; Jokela, P.; Karlsson, J.G.; Olofsson, L.; Rosengren, J.; Shoravi, S.; Svenson, J.; Wikman, S. Can we rationally design molecularly imprinted polymers? Anal. Chim. Acta, 2001, 435(1), 9-18.

http://dx.doi.org/10.1016/S0003-2670(01)00932-1

- [44] Gholivand, M.B.; Khodadadian, M.; Ahmadi, F. Computer aidedmolecular design and synthesis of a high selective molecularly imprinted polymer for solid-phase extraction of furosemide from human plasma. Anal. Chim. Acta, 2010, 658(2), 225-232 http://dx.doi.org/10.1016/j.aca.2009.11.019 PMID: 20103099
- [45] Li, X.F.; Zhong, S.A.; Chen, L.; Whittaker, A. Computer simulation and preparation of molecularly imprinted polymer membranes with chlorogenic acid as template. Polym. Int., 2011, 60(4), 592-598. http://dx.doi.org/10.1002/pi.2985

12 Current Drug Delivery, XXXX, Vol. XX, No. XX

[46] Tabandeh, M.; Ghassamipour, S.; Aqababa, H.; Tabatabaei, M.; Hasheminejad, M. Computational design and synthesis of molecular imprinted polymers for selective extraction of allopurinol from human plasma. J. Chromatogr. B Analyt. Technol. Biomed. Life Sci., 2012, 898, 24-31.

http://dx.doi.org/10.1016/j.jchromb.2012.04.009 PMID: 22565062

- [47] Ahmadi, F.; Rezaei, H.; Tahvilian, R. Computational-aided design of molecularly imprinted polymer for selective extraction of methadone from plasma and saliva and determination by gas chromatography. J. Chromatogr. A, 2012, 1270, 9-19. http://dx.doi.org/10.1016/j.chroma.2012.10.038 PMID: 23159198
- [48] Tahir, I.; Wijaya, K.; Islam, A.K.M.; Ahmad, M. Computer aided design of molecular imprinted polymer for selective recognition of capsaicin. Indones. J. Chem., 2014, 1485(1), 85-93. http://dx.doi.org/10.22146/ijc.21272
- [49] Wang, Y.; Liu, J-B.; Tang, S-S.; Jin, R-F. Preparation of melamine molecularly imprinted polymer by computer-aided design. J. Sep. Sci., 2015, 38(15), 2647-2654. http://dx.doi.org/10.1002/jssc.201500375 PMID: 25964122
- [50] Krishnan, H.; Islam, K.M.S.; Hamzah, Z.; Ahmad, M.N. Rational computational design for the development of andrographolide molecularly imprinted polymer. AIP Conf. Proc., 2017, 1891(1), 020083

http://dx.doi.org/10.1063/1.5005416

- Huang, X.; Zhang, W.; Wu, Z.; Li, H.; Yang, C.; Ma, W.; Hui, A.; [51] Zeng, Q.; Xiong, B.; Xian, Z. Computer simulation aided preparation of molecularly imprinted polymers for separation of bilobalide. J. Mol. Model., 2020, 26(8), 198. http://dx.doi.org/10.1007/s00894-020-04460-y PMID: 32632503
- Zhang, Y.; Huang, W.; Yin, X.; Sarpong, K.A.; Zhang, L.; Li, Y.; [52] Zhao, S.; Zhou, H.; Yang, W.; Xu, W. Computer-aided design and synthesis of molecular imprinting polymers based on doubly oriented functional multiwalled carbon nanotubes for electrochemically sensing bisphenol A. React. Funct. Polym., 2020, 157, 104767. http://dx.doi.org/10.1016/j.reactfunctpolym.2020.104767
- [53] Boysen, R.I. Advances in the development of molecularly imprinted polymers for the separation and analysis of proteins with liquid chromatography. J. Sep. Sci., 2019, 42(1), 51-71. http://dx.doi.org/10.1002/jssc.201800945 PMID: 30411488
- [54] Joke Chow, A.L.; Bhawani, S.A. Synthesis and characterization of molecular imprinting polymer microspheres of cinnamic acid: Extraction of cinnamic acid from spiked blood plasma. Int. J. Polym. Sci., 2016, 2016, 2418915. http://dx.doi.org/10.1155/2016/2418915
- [55] Shadabfar, M.; Abdouss, M.; Khonakdar, H.A. Synthesis, characterization, and evaluation of a magnetic molecular imprinted polymer for 5-fluorouracil as an intelligent drug delivery system for breast cancer treatment. J. Mater. Sci., 2020, 55(26), 12287-12304. http://dx.doi.org/10.1007/s10853-020-04887-x
- Chen, Q.; Liu, X.; Yang, H.; Zhang, S.; Song, H.; Zhu, X. Prepara-[56] tion and evaluation of magnetic graphene oxide molecularly imprinted polymers (MIPs-GO-Fe₃ O_4 @SiO₂) for the analysis and separation of tripterine. React. Funct. Polym., 2021, 169, 105055. http://dx.doi.org/10.1016/j.reactfunctpolym.2021.105055
- Kushwaha, A.; Singh, S.; Gupta, N.; Singh, A. K.; Singh, M. Syn-[57] thesis and characterization of antipyrine-imprinted polymers and their application for sustained release. Polym. Bull, 2018, 75(11), 5235-5252 http://dx.doi.org/10.1007/s00289-018-2326-x
- Umpleby, R.J., II; Baxter, S.C.; Chen, Y.; Shah, R.N.; Shimizu, [58] K.D. Characterization of molecularly imprinted polymers with the Langmuir-Freundlich isotherm. Anal. Chem., 2001, 73(19), 4584-4591
 - http://dx.doi.org/10.1021/ac0105686 PMID: 11605834
- [59] Nishitani, S.; Sakata, T. Potentiometric adsorption isotherm analysis of a molecularly imprinted polymer interface for small-biomolecule recognition. ACS Omega, 2018, 3(5), 5382-5389. http://dx.doi.org/10.1021/acsomega.8b00627 PMID: 30023917
- Baggiani, C.; Giraudi, G.; Giovannoli, C.; Tozzi, C.; Anfossi, L. [60] Adsorption isotherms of a molecular imprinted polymer prepared in the presence of a polymerisable template: Indirect evidence of the formation of template clusters in the binding site. Anal. Chim. Acta, 2004, 504(1), 43-52.

http://dx.doi.org/10.1016/S0003-2670(03)00671-8

- [61] González, G.P.; Hernando, P.F.; Alegría, J.S.D. A morphological study of molecularly imprinted polymers using the scanning electron microscope. Anal. Chim. Acta, 2006, 557(1), 179-183. http://dx.doi.org/10.1016/j.aca.2005.10.034 PMID: 17386663
- [62] Chen, X.; Ye, N. A graphene oxide surface-molecularly imprinted polymer as a dispersive solid-phase extraction adsorbent for the determination of cefadroxil in water samples. RSC Adv., 2017, 7(54), 34077-34085.

http://dx.doi.org/10.1039/C7RA02985C

- [63] Roland, R.M.; Bhawani, S.A.; Wahi, R.; Ibrahim, M.N.M. Synthesis, characterization, and application of molecular imprinting polymer for extraction of melamine from spiked milk, water, and blood serum. J. Liq. Chromatogr. Relat. Technol., 2020, 43(3-4), 94-105. http://dx.doi.org/10.1080/10826076.2019.1672077
- [64] Hasanah, A.N.; Safitri, N.; Zulfa, A.; Neli, N.; Rahayu, D. Factors affecting preparation of molecularly imprinted polymer and methods on finding template-monomer interaction as the key of selective properties of the materials. Molecules, 2021, 26(18), 5612. http://dx.doi.org/10.3390/molecules26185612 PMID: 34577083
- Awokoya, K.N.; Okoya, A.A.; Elujulo, O. Preparation, characteriza-[65] tion and evaluation of a styrene-based molecularly imprinted polymer for capturing pyridine and pyrrole from crude oil. Sci. Afr., 2021, 13, e00947. http://dx.doi.org/10.1016/j.sciaf.2021.e00947
- Scorrano, S.; Mergola, L.; Del Sole, R.; Vasapollo, G. Synthesis of [66] molecularly imprinted polymers for amino acid derivates by using different functional monomers. Int. J. Mol. Sci., 2011, 12(3), 1735-1743.

http://dx.doi.org/10.3390/ijms12031735 PMID: 21673919

Svenson, J.; Zheng, N.; Föhrman, U.; Nicholls, I.A. The role of [67] functional monomer-template complexation on the performance of atrazine molecularly imprinted polymers. Anal. Lett., 2005, 38(1), 57-69

http://dx.doi.org/10.1081/AL-200043443

- [68] Madikizela, L. M.; Zunngu, S. S.; Mlunguza, N. Y.; Tavengwa, N. T.; Mdluli, P. S.; Chimuka, L. Application of molecularly imprinted polymer designed for the selective extraction of ketoprofen from wastewater. Water SA, 2018, 44(3), 406-418. http://dx.doi.org/10.4314/wsa.v44i3.08
- [69] Mahony, J.O.: Nolan, K.: Smyth, M.R.: Mizaikoff, B. Molecularly imprinted polymers-potential and challenges in analytical chemistry. Anal. Chim. Acta, 2005, 534(1), 31-39. http://dx.doi.org/10.1016/j.aca.2004.07.043
- [70] Chen, W.; Tian, X.; He, W.; Li, J.; Feng, Y.; Pan, G. Emerging functional materials based on chemically designed molecular recognition. BMC Materials, 2020, 2(1), 1. http://dx.doi.org/10.1186/s42833-019-0007-1
- Ansell, R.J.; Ramström, O.; Mosbach, K. Towards artificial antibod-[71] ies prepared by molecular imprinting. Clin. Chem., 1996, 42(9), 1506-1512.

http://dx.doi.org/10.1093/clinchem/42.9.1506 PMID: 8787721

- Zaidi, S.A. Molecular imprinting: A useful approach for drug deliv-[72] ery. Mater. Sci. Energy Technol., 2020, 3, 372-377. http://dx.doi.org/10.1016/j.mset.2019.10.012
- He, Y.; Zeng, S.; Abd El-Aty, A.M.; Hacımüftüoğlu, A.; Kalekristos [73] Yohannes, W.; Khan, M.; She, Y. Development of water-compatible molecularly imprinted polymers based on functionalized β cyclodextrin for controlled release of atropine. Polymers (Basel), 2020, 12(1), E130.

http://dx.doi.org/10.3390/polym12010130 PMID: 31935897

- [74] Zheng, L.; Wang, H.; Cheng, X. Molecularly imprinted polymer nanocarriers for recognition and sustained release of diclofenac. Polym. Adv. Technol., 2018, 29(5), 1360-1371. http://dx.doi.org/10.1002/pat.4247
- [75] Liu, X.L.; Yao, H.F.; Chai, M.H.; He, W.; Huang, Y.P.; Liu, Z.S. Green synthesis of carbon nanotubes-reinforced molecularly imprinted polymer composites for drug delivery of fenbufen. AAPS PharmSciTech, 2018, 19(8), 3895-3906. http://dx.doi.org/10.1208/s12249-018-1192-z PMID: 30324359
- [76] Bakhshpour, M.; Yavuz, H.; Denizli, A. Controlled release of mitomycin C from PHEMAH-Cu(II) cryogel membranes. Artif. Cells Nanomed. Biotechnol., 2018, 46(sup1), 946-954. http://dx.doi.org/10.1080/21691401.2018.1439840

- [77] Ruela, A.L.; de Figueiredo, E.C.; de Araújo, M.B.; Carvalho, F.C.; Pereira, G.R. Molecularly imprinted microparticles in lipid-based formulations for sustained release of donepezil. *Eur. J. Pharm. Sci.*, 2016, 93, 114-122. http://dx.doi.org/10.1016/j.ejps.2016.08.019 PMID: 27519666
- [78] Mohebali, A.; Abdouss, M.; Mazinani, S.; Zahedi, P. Synthesis and characterization of poly(methacrylic acid)-based molecularly imprinted polymer nanoparticles for controlled release of trinitroglycerin. *Polym. Adv. Technol.*, **2016**, *27*(9), 1164-1171. http://dx.doi.org/10.1002/pat.3778
- [79] Trotta, F.; Caldera, F.; Cavalli, R.; Soster, M.; Riedo, C.; Biasizzo, M.; Uccello Barretta, G.; Balzano, F.; Brunella, V. Molecularly imprinted cyclodextrin nanosponges for the controlled delivery of L-DOPA: Perspectives for the treatment of Parkinson's disease. *Expert Opin. Drug Deliv.*, **2016**, *13*(12), 1671-1680.
- http://dx.doi.org/10.1080/17425247.2017.1248398 PMID: 27737572
 [80] Jafary Omid, N.; Morovati, H.; Amini, M.; Dehpour, A.R.; Partoazar, A.; Rafiee-Tehrani, M.; Dorkoosh, F. Development of molecularly imprinted olanzapine nano-particles: *In vitro* characterization and *in vivo* evaluation. *AAPS PharmSciTech*, **2016**, *17*(6), 1457-1467.

http://dx.doi.org/10.1208/s12249-016-0480-8 PMID: 26831447

- [81] Sheybani, S.; Hosseinifar, T.; Abdouss, M.; Mazinani, S. Mesoporous molecularly imprinted polymer nanoparticles as a sustained release system of azithromycin. *RSC* <u>4dv.</u>, 2015, 5(120), 98880-98891. http://dx.doi.org/10.1039/C5RA11970G
- [82] Haq, I.; Mujahid, A.; Afzal, A.; Iqbal, N.; Bajwa, S.Z.; Hussain, T.; Shehzad, K.; Ashraf, H. Developing imprinted polymer nanoparticles for the selective separation of antidiabetic drugs. *J. Sep. Sci.*, **2015**, *38*(19), 3469-3476. http://dx.doi.org/10.1002/jssc.201500506 PMID: 26179897
- [83] Ruela, A.L.M.; Figueiredo, E.C.; Pereira, G.R. Molecularly imprinted polymers as nicotine transdermal delivery systems. *Chem. Eng. J.*, 2014, 248, 2481-2488.

http://dx.doi.org/10.1016/j.cej.2013.12.106

- [84] Barde, L.N.; Ghule, M.M.; Roy, A.A.; Mathur, V.B.; Shivhare, U.D. Development of molecularly imprinted polymer as sustain release drug carrier for propranolol HCL. *Drug Dev. Ind. Pharm.*, 2013, 39(8), 1247-1253.
- http://dx.doi.org/10.3109/03639045.2012.710236 PMID: 22871098
 [85] Mirzaei, M.; Najafabadi, S.A.H.; Abdouss, M.; Azodi-Deilami, S.; Asadi, E.; Hosseini, M.R.M.; Piramoon, M. Preparation and utilization of microporous molecularly imprinted polymer for sustained release of tetracycline. J. Appl. Polym. Sci., 2013, 128(3), 1557-1562. http://dx.doi.org/10.1002/app.3831
- [86] Javanbakht, M.; Mohammadi, S.; Esfandyari-Manesh, M.; Abdouss, M. Molecularly imprinted polymer microspheres with nanopore cavities prepared by precipitation polymerization as new carriers for the sustained release of dipyridamole. J. Appl. Polym. Sci., 2011, 119(3), 1586-1593. http://dx.doi.org/10.1002/app.32798
- [87] Azodi-Deilami, S.; Abdouss, M.; Javanbakht, M. The syntheses and characterization of molecularly imprinted polymers for the controlled release of bromhexine. *Appl. Biochem. Biotechnol.*, 2011, 164(2), 133-147.

http://dx.doi.org/10.1007/s12010-010-9121-y PMID: 21076945

- [88] Abdouss, M.; Asadi, E.; Azodi-Deilami, S.; Beik-mohammadi, N.; Aslanzadeh, S.A. Development and characterization of molecularly imprinted polymers for controlled release of citalopram. *J. Mater. Sci. Mater. Med.*, 2011, 22(10), 2273-2281. http://dx.doi.org/10.1007/s10856-011-4395-3 PMID: 21833610
- [89] da Silva, M.S.; Nobrega, F.L.; Aguiar-Ricardo, A.; Cabrita, E.J.; Casimiro, T. Development of molecularly imprinted co-polymeric devices for controlled delivery of flufenamic acid using supercritical fluid technology. J. Supercrit. Fluids, 2011, 58(1), 150-157. http://dx.doi.org/10.1016/j.supflu.2011.05.010
- [90] Cirillo, G.; Parisi, O.I.; Curcio, M.; Puoci, F.; Iemma, F.; Spizzirri, U.G.; Picci, N. Molecularly imprinted polymers as drug delivery systems for the sustained release of glycyrrhizic acid. *J. Pharm. Pharmacol.*, 2010, 62(5), 577-582. http://dx.doi.org/10.1211/jpp.62.05.0003 PMID: 20609058

- [91] Azodi-Deilamia, S.; Abdoussa, M.; Rezvaneh Seyedib, S. Synthesis and characterization of molecularly imprinted polymer for controlled release of tramadol. *Open Chem.*, **2010**, 8(3), 687-695. http://dx.doi.org/10.2478/s11532-010-0035-x
- [92] Del Sole, R.; Lazzoi, M.R.; Vasapollo, G. Synthesis of nicotinamide-based molecularly imprinted microspheres and *in vitro* controlled release studies. *Drug Deliv.*, 2010, 17(3), 130-137. http://dx.doi.org/10.3109/10717541003587418 PMID: 20163194
- [93] Cirillo, G.; Iemma, F.; Puoci, F.; Parisi, O.I.; Curcio, M.; Spizzirri, U.G.; Picci, N. Imprinted hydrophilic nanospheres as drug delivery systems for 5-fluorouracil sustained release. *J. Drug Target.*, 2009, 17(1), 72-77.

http://dx.doi.org/10.1080/10611860802455813 PMID: 19016107

- [94] Puoci, F.; Cirillo, G.; Curcio, M.; Iemma, F.; Parisi, O.I.; Castiglione, M.; Picci, N. Molecularly imprinted polymers for alphatocopherol delivery. *Drug Deliv.*, **2008**, *15*(4), 253-258. http://dx.doi.org/10.1080/10717540802006724 PMID: 18446571
- [95] Khorrami, A.R.; Mehrseresht, S. Synthesis and evaluation of a selective molecularly imprinted polymer for the contraceptive drug levonorgestrel. J. Chromatogr. B Analyt. Technol. Biomed. Life Sci., 2008, 867(2), 264-269.
- http://dx.doi.org/10.1016/j.jchromb.2008.04.017 PMID: 18456579 [96] Suedee, R.; Srichana, T.; Martin, G.P. Evaluation of matrices con-
- taining molecularly imprinted polymers in the enantioselectivecontrolled delivery of beta-blockers. *J. Control. Release*, **2000**, *66*(2-3), 135-147. http://dx.doi.org/10.1016/S0168-3659(99)00261-8 PMID: 10742575
- [97] Sedghi, R.; Ashrafzadeh, S.; Heidari, B. pH-sensitive molecularly imprinted polymer based on graphene oxide for stimuli actuated controlled release of curcumin. J. Alloys Compd., 2021, 857157603, 157603.

http://dx.doi.org/10.1016/j.jallcom.2020.157603

- [98] Cegłowski, M.; Jerca, V.V.; Jerca, F.A.; Hoogenboom, R. Reduction-responsive molecularly imprinted poly(2-isopropenyl-2oxazoline) for controlled release of anticancer agents. *Pharmaceutics*, **2020**, *12*(6), E506. http://dx.doi.org/10.3390/pharmaceutics12060506 PMID: 32498326
- [99] Javanbakht, S.; Saboury, A.; Shaabani, A.; Mohammadi, R.; Ghorbani, M. Doxorubicin imprinted photoluminescent polymer as a phresponsive paperarier. *ACS Appl. Bio. Mater.* 2020, 3(7), 4168-

responsive nanocarrier. ACS Appl. Bio Mater., **2020**, 3(7), 4168-4178. http://dx.doi.org/10.1021/acsabm.0c00254 PMID: 35025419

[100] Varela-Garcia, A.; Gomez-Amoza, J.L.; Concheiro, A.; Alvarez-Lorenzo, C. Imprinted contact lenses for ocular administration of an-

tiviral drugs. *Polymers (Basel)*, **2020**, *12*(9), E2026. http://dx.doi.org/10.3390/polym12092026 PMID: 32899893

[101] Cegłowski, M.; Kurczewska, J.; Ruszkowski, P.; Schroeder, G. Application of paclitaxel-imprinted microparticles obtained using two different cross-linkers for prolonged drug delivery. *Eur. Polym.* J., 2019, 118, 118328-118336.

http://dx.doi.org/10.1016/j.eurpolymj.2019.06.010 [102] Mo, C.E.; Chai, M.H.; Zhang, L.P.; Ran, R.X.; Huang, Y.P.; Liu,

- [102] Mo, C.E., Chai, M.H., Zhang, L.F., Ran, K.A., Huang, T.F., Liu, Z.S. Floating molecularly imprinted polymers based on liquid crystalline and polyhedral oligomeric silsesquioxanes for capecitabine sustained release. *Int. J. Pharm.*, **2019**, *557*, 293-303. http://dx.doi.org/10.1016/j.ijpharm.2018.12.070 PMID: 30599225
- [103] Liu, L.; Li, N.; Chen, M.; Yang, H.; Tang, Q.; Gong, C. Visiblelight-responsive surface molecularly imprinted polymer for acyclovir through chicken skin tissue. ACS Appl. Bio Mater., 2018, 1(3), 845-852.

http://dx.doi.org/10.1021/acsabm.8b00275 PMID: 34996176

- [104] Zhang, L.; Chen, L.; Zhang, H.; Yang, Y.; Liu, X. Recognition of 5fluorouracil by thermosensitive magnetic surface molecularly imprinted microspheres designed using a computational approach. J. Appl. Polym. Sci., 2017, 134(43), 45468. http://dx.doi.org/10.1002/app.45468
- [105] Hemmati, K.; Masoumi, A.; Ghaemy, M. Tragacanth gum-based nanogel as a superparamagnetic molecularly imprinted polymer for quercetin recognition and controlled release. *Carbohydr. Polym.*, **2016**, *136*, 630-640.

http://dx.doi.org/10.1016/j.carbpol.2015.09.006 PMID: 26572395

[106] Kazemi, S.; Sarabi, A.A.; Abdouss, M. Synthesis and characterization of magnetic molecularly imprinted polymer nanoparticles for controlled release of letrozole. *Korean J. Chem. Eng.*, **2016**, *33*(11), 3289-3297.

http://dx.doi.org/10.1007/s11814-016-0171-x

- [107] Karimi, A.R.; Khodadadi, A.; Hadizadeh, M. A nanoporous photosensitizing hydrogel based on chitosan cross-linked by zinc phthalocyanine: An injectable and pH-stimuli responsive system for effective cancer therapy. *RSC Adv.*, 2016, 6(94), 91445-91452. http://dx.doi.org/10.1039/C6RA17064A
- [108] Li, L.; Chen, L.; Liu, W.; Yang, Y.; Liu, X.; Chen, Y. Preparation and characterization of 5-fluorouracil surface-imprinted thermosensitive magnetic microspheres. *Monatsh. Chem.*, **2015**, *146*(3), 441-447.

http://dx.doi.org/10.1007/s00706-014-1335-1

[109] Li, B.; Xu, J.; Hall, A.J.; Haupt, K.; Tse Sum Bui, B. Watercompatible silica sol-gel molecularly imprinted polymer as a potential delivery system for the controlled release of salicylic acid. *J. Mol. Recognit.*, **2014**, *27*(9), 559-565. http://dx.doi.org/10.1002/jmr.2383 PMID: 25042710

[110] Türkmen, D.; Bereli, N.; Çorman, M.E.; Shaikh, H.; Akgöl, S.; Denizli, A. Molecular imprinted magnetic nanoparticles for controlled

delivery of mitomycin C. Artif. Cells Nanomed. Biotechnol., 2014, 42(5), 316-322.

http://dx.doi.org/10.3109/21691401.2013.823094 PMID: 23937455

[111] Puoci, F.; Hampel, S.; Parisi, O.i.; Hassan, A.; Cirillo, G.; Picci, N. Imprinted microspheres doped with carbon nanotubes as novel electroresponsive drug-delivery systems. J. Appl. Polym. Sci., 2013, 130(2), 829-834. http://dx.doi.org/10.1002/app.39212















ਫਾਰਮੇਸੀ ਕਾਲਜ ਨੇ ਗੜ੍ਹਵਾਲ ਯੂਨੀਵਰਸਿਟੀ ਨਾਲ ਕੀਤਾ ਸਮਝੌਤਾ

 6 भाषाच लोगी राषिये toron worth man mite want for how we want



संभवभेत पृष्टीत ततवा, वाहितीवत्व वर्ग सी वी कुधवा, बागीम शिंमीधम वर्ग भाव, वे, तार्वल लखींचे धंजव सी बाग्वे डिफाईमें वेवे। वार्वलाग

त्रिम से राज्य से विमसस पुर्गाबर, जबरतीब मिंधत स' मेरेगा । वेवजते अवगसेस प्रेतवाभ से जीवत । प्रिप्त जीवलाग्वर प्रियं प्रतिविज्ञानी बिस्टिमव्योभ अन्त्र मठाव ठूं तथी वयाभामिट्रिटीयस सिभारिज राज

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आई.एस.एफ. कॉलेज ऑफ फार्मेसी ने गढ़वाल विश्वविद्यालय से किया समझौता

मौए, 5 मार्थ (अकलियांवाला): प्राक्षतां में स्थित संस्था अई एक.एफ इउंतेन अगेर फार्मेसो ने डिक्टॉर्ट ऑफ फा मॉस्च् टिकल साइंस एन.एन.चे. पद्माल चूनिवस्टिंगे (सेंट्रा पूर्विचसिटो) ओराजर के साथ सच्चतील एन पर हात्वास किए हैं।

संस्थ के उसकेरर का भी है। मुझ्ल के उसकेरर का भी है। किस्टोर अभ भवते प्रतुभिक्त साहत 1 काई पर, पर, कोनेन और क्यमित्र के के से का के कार्य पर साहती प्रका के के से के कार्य करने पर साहती प्रका के के से के कार्य करने पर साहती प्रका को 1 इस स्वादी के दाज मि बीच कार एवं बई एस एस, कोनेन के जनका सार्य मुरियमों का कार उनके पर संबंधी 1 स्वारं का का ती प्रवां प्रतिक किस्टो 1 स्वारं का साह ने सिप्ट के कार्य का स्वारंग के साहत कार्य पर देवर की में दाक के साह कार कार्य साहजा के साहत



FRIDAY 6 HIT 2020

तम्ब्रीता पत्र की कॉर्पी डिखाने आई.एस.एक. कॉर्सेज ऑफ कार्परी के पेशामेंन प्रथिष गर्ग, रामरेक्टर या. की डी.नुप्त, बहुल डिसीफल या. आर.के. नार्रम (अव्याण्याय

हिमिटेंट के साथ समझीत पत्र भा हस्वावर सिंग, यह फार्मायुटिकस कबर्ग्न कार्ज के ट्रीन, एसप्रीट पर बॉस्टेंटिक दुग डिसीको के सिंग जानरकस साथाी उपस्त्रथ करवाणी वॉक रू प्रोठसरों को लेवि में माझीत स्व भा का से के सोवेंस्टा में ने देनींबेंग डाइसेस्टर वित्त प्रोडास्टान ने इस मासहीट पर युको प्राठ करते करत कि संधान की पित्र पर जीत प्राठ करते करत को गॉडॉविश्वये में और तेवे आगरी व जावें को रोजगार व स्व-रोजगार छाथ करने के लिए स्वर्थना पिलोगा इस मौके पर संस्था के पोलमैन इसी पार्ग, सार्थ्य देवी, बनेक गर्ग, उस, मुक्कत गर्ग, डायेफ्ल ड, आर.के, नगेर पर मुझ, केक्टरी ठाउक ने समझौत पज म, हनराक्ष के लिए, खो को पर म, हनराक्ष के लिए,



आइएसएफ ने गढवाल विश्वविद्यालय से किया समझौता

संसु स्वान् । संसु स्वान् । संस्वार्थन से अंदर वे जिन्दरिय सांक सामान्द्रियन साहित्य राजवंद गुजम देवरायन कि ता संस्वार कर सामान्द्र यो सीचे कुल ने साहत कि तो स्वानं साहित्व ने अनुस्वरण सीचे सीच साहित्व ने अनुस्वरण सीचेल की साहित्व ने अनुस्वरण सीचेल की साहित्व ने साह स्वार्थन कर सामाने अस्य मा

स्म अवस्ती के तथा दिवाधिकाला पूर्व आवस्त्रप्र भौतित में उपताल दिवर्थ सुविधायों का स्वय जयक का स्वर्थन सुविधायों का स्वय जयक का



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समागम का उदघाटन डायरैक्टर , गुप्ता, उप प्रिंसीपल ढा ,नारंग आदि ने किया आई.एस.एफ. कालेज में राष्ट्रीय कांफ्रैंस करवाई

मोना, 27 फरसरी (मलादी) । राज्य को प्रमुख सिक्षण संस्थ and the other and the second base ü sichman und mendal ü धोवापटी आफ प्रजाशिहरूल व हिन्दर्थ के बाल इंग्रेटी-व्यूलसन क्वारिनरी could a strong the franks realist नेमाल कांग्रेस का आयोजन arefebfern rus it arithm fam गकः हम बाउँम का शावरंत माल हे चेवारेन प्रतेष पर जा आवान फास्ता गठावाल वुविधनिती सीवगर राष्ट्राल जनेला बाईटिस्ट आई.पी.अट. जीन बेज्यान पैनेजिन वापरीक्टर करनेर करेरनों प्रायवेश विविदेश एम से आग के आयल एव रतेना के उपरीक्तर या जो हो। एस en fraften in ann it ann प्रमाणी अगर भी इटेन अभवध मा



ता. राहल हनेजा को सम्मानित करते वेकामेन प्रतीण गर्न जावरेकटर जा जी जी. THE & SHE | VANH

विद्यूच केल ने संपत्र के पर लोग प्रस्तनित आये दिन्दा पर सीचे प ता. विद्यानं बेहन ने आग गए सभी वेतमानों का ल्वागत किया। प्राप्तिन को

विययत जानवारी दी। इस सीचे unfrer et mein nite 2 ber ab संबोधित करने सप्टरेक्टर जा भी थीं जयपंत्रित प्राइतिर पर उसी थे आग

को उपयोगित जो अक्षम करावय प्रयोगे कहा कि देख एवं चिटेल में and the same will rate forther way ref है। इस केरे क जा, आरत्म प्रमान und fit ann un warm immit mu iture are fic-flash favora a frand at \$70 minut Parents तथा इंटरली तक पोलने पर ओर दिया त्य कीरे पर से जीपन जोते है आप in after the ferrers for का होते पर प्रनाहर किंग it and an abortion fires प्रवर्तन एवं देखें जरीता में जगुनी दर से किया इस जीवे वे सीवत या जोन्द् ज्यान ने एक पी जार पंजाब जांच को कांग्रेंग के आयोजन पर कहा लें। या कीचे का कैतिवाली प्रताह एक fielde aftere its

आई एस.एफ. कालेज ने गढवाल विश्वविद्यालय व कोनोरकोरमो से किया समझौता कोनोरकोस्मो कंपनी छात्रों को देगी द्रेगिंग व करवाएगी प्लेसमैंट

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जेन्द्रतीय वर्ग अस्टीकेटर जा जी ती राज तर्ह दिसीयल जा ज प्रमाई पर की कार्य दिवाने वा

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ਇਕ ਰੋਜ਼ਾ ਰਾਸ਼ਟਰੀ ਕਾਨਫਰੰਸ ਦਾ ਆਯੋਜਨ

ਵਰਬਰੀ (ਹੈ।) ਰਾਊਕੇ।-ਸੂਸੇ ਦੀ ਪ੍ਰਮੁੱਖ ਵਿਦਿਅਕ ਸੰਸਥਾ ਆਈ ਐੱਸ ਐੱਫ. ਕਾਲਜ ਆਫ ਵਾਧਮੰਸੀ ਵਿਧ ਹਿਟਸੈਕਚੁਅਲ ਪ੍ਰਾਪਰਟੀ ਰਾਈਟਸ 'ਤੇ ਸੋਸ਼ਾਇਟੀ ਆਬ ਵਾਧਮਾਸਿਊਟੀਕਲ



ਰਿਸਟਦ ਦੇ ਨਾਲ ਇੱਕ duxu areada מד שולמא ਆਡੀਟੋਗੋਅਮ ਹਾਲ ਵਿਚ ਕੀਤਾ ਗਿਆ। ਇਸ ਕਾਨਵਰੇਸ ਦੀ ਤੁਰੂਆਤ ਸੋਸਥਾ ਜ ਅਰਮੈਨ ਪ੍ਰਧੀਨ ਗੁਰੂਗ, ਡਾ. ਅਸਦ रपार सहकार प्रयोगती होत हा. हाएक जर्तेसा माहिटिमट आर्थ ਹੈ ਆਵਾ ਦਰਿਨ ਸ਼ੇਰਵਪਾਲ ਮੈਨੇਰਿੰਦ ਾਇਰੈਸਟਰ ਕਨੋਰ ਜੋਸਮੇ ਪਾਈਵੇਟ ਜਿਸਟਿਡ, ਐੱਸ. ਪੀ. ਆਰ. ਦੇ ਪਧਾਨ मधा से वाहितेवतन जा सी ते. वापजा, बाहीम प्रिंमीपत ज से राजेन और धे unter at footantiers à ut जीनी साम के कीनी। हैन ਜ਼ਾਰਜ਼ ਨੇ ਆਏ ਹੋਵੇ। ur üxmu aları वाठवर्तम के मंब्रेयक जनवि वाफ़िकेवटर बा, सी, जी, संपत्र र ਐੱਡ ਸੀ ਅਤਰ ਕੋਲੋਂ ਜੋਲ ਕ multimite sur termitum

प्राप्तके से प्रकार कर केवा हे केवरि पहरे तका। अन

ਦਿੰਗੀ। ਇਸ र जा, जायस अर्देश से प्रेटेट ਮਹੱਤਣਾ, ਵਾਈਇੰਗ ਅਤੇ ਆਈ ਦੀ ਮਹੱਤਰਾ ਸਾਰੇ ਕਰਵਾਇਆ। ਉਨਾਂ ਕਿਹਾ ਕਿ ਦੇਸ਼ ਅਤੇ ਵਿਧੇਸ਼ ਵਿਚ ਆਈ ਪੀ ਆਰ ਦੀ ਮੰਗ ਇਕੇ ਕਰ ਕੀਏ ਕਰੀ ਹੈ। ਇਸ ਕੋਰਾਨ ਕਾ करने मैमलिलां रेवल हरेत वरिषती דעראי אים הערוד אות א נוצר ਤੇਸ਼ਨਾਲਗੀ ਅਤੇ ਇੰਡਸਟ ਤੇ ਦੇਸ਼ ਇੱਕਾ। ਹੈ ਜੋਸ अपने तेने माते अतिमालं स לום למים וביעים לוסי "ה להכמים बीजगः स्टेन araand विप्रवधानीय भागे होती भगीवा रे ਸੰਗਾਣੀ। ਇਸ ਮੌਕੇ ਐੱਸ. ਪੀ ਸੈਕਟਰੀ ਕਾ ਉਪਿੰਦਰ ਪੀਕਾਲ ਸਟੇਟ ਭਾਂਦ ਨੇ ਕਾਨਫਰੇਸ गरंबर समी लगानी ਹੋਰਾਨ ਵੇਇਲਟੀ ਸਟਾਫ ਅਹ विकिश्वाची तरस्य प्रहा



आइएसएफ कॉलेग औरू प्रार्थनी में पंचरमैन ब्रॉन को, जी, अब्दुत फाल खा। वी, सहुत तनेजा कड़िस के बाद ही, जीडी नुप्रा को सम्मानिश करते सुर 4 जिल्ली

आइएसफ कॉलेज ऑफ फार्मेसी में करवाई कांफ्रेंस

सवाद रूखोगे, मंत्रा : आवश्मसण् अधिक अधिक प्रमर्थेस में इंटलेक्युवल प्रॉपटी राहट्स पर सांसायरी और प्रमासंबुटिकल पर्व सित्ता के साथ इंस्टोट्यूरानल क्यांलिटी इंटकेंस इस एक टिक्सीय संपूर्णि के साथ कॉर्डेस क्र अवोज्ज ऑडीटोरियन पाल में किया क्या।

कांग्रेस का राप्यरंग संस्था के प्रेवस्मेंन प्रवीण गर्म, डी. अल्युल प्रास्थवा वर्युवाल प्रत्नेत खडीव्स आपिआर, तीन- विश्वापन मैनीजन प्रायंग्रेस्ट कुनारं कांस्था स्वर्धव्य-सिमिटिइ, एसप्रेअवर के अव्यक्ष एवं संस्था के प्रायंग्रेस्ट की जीवे गुरा, वायस विस्थित की जीवे राज्य की नोंग, प्रायंग्रेस्ट की टी. आर्ग्ते नोंग, मेठन ने संमुका तौर पर ज्योति प्रत्यतित करके किया। इस दीवन उंद्य तीर्द्र गतविधियाँ व संस्था के बारे स्टी गतिबिधियाँ व संस्था के बारे से पिरस्तुत जानकार्य दी इस मेंके पर साईदेस्ट तो उष्ट्रल संग्रे में प्रेंटर को उपयोगित, काइलिंग पर्य अइतिप्रस को उपयोगित बारे अटकार करनका।

उन्होंने कहा कि देव एवं विदेश में आइवेआर को मंगनितंतर कड़ एसे हैं। कु अब्दुल कहर के आइयेआर पर प्रकात उलते हुए नेवल दुब डिलीयर्स बिसटम च सिर्फ वर्ष को पेटेंट, दुसपर टेबनालजों तब इंट्रदी तक धेजने पर जोर दिया। इस मौबे पर कैकटरों स्टाफ एवं विद्यार्थी उनस्थित बे।

ਇਕ ਰੋਜ਼ਾ ਕੌਮੀ ਕਾਨਫਰੰਸ ਕਰਵਾਈ

ਵਕੀਲ ਮਹਿਰੋਂ, ਮੋਗਾ

ਸ਼ਬੇ ਦੀ ਪ੍ਰਮੁੱਖ ਵਿਦਿਅਕ ਸੰਸਥਾ ਆਈਐੱਸਐੱਟ ਕਾਸ਼ਜ ਆਰ ਰਾਨਮੀ ਵਿੱਚ ਇੰਟਲੈਕਚਅਲ ਪਾਪਟੀ ਰਾਈਟਸ ਸਸਾਇਟੀ ਆਫ ਫ਼ਚਮਾਸਿਊਟਿਡਲ ਤੇ ਕਿਸਰਚ ਦੇ ਨਾਲ ਇੰਸਟੀਚਿਸ਼ਨਲ ਕਆਸਿਟੀ ਇੰਸੋਰੈਂਸ ਵੋਲੋਂ ਇੱਕ ਰੋਵਾ ਰਾਸਟਰੀ ਨੈਸ਼ਨਲ ਕਾਨਰਹੇਸ਼ ਆਈਟਰੀਅਮ ਹਾਲ 'ਚ ਕਰਵਾਈ ਗਈ। ਕਾਨਫਰੇਸ ਦੀ ਸ਼ਰਆਤ ਸੰਸਥਾ ਦੇ ਚੇਅਰਮੈਨ ਪਵੀਨ ਗਰਗ, ਡਾ. ਅਬਦਲ ਵਾਰਖ ਗੜਵਾਲ ਯਨੀਵਰਸਿਟੀ ਸ਼ੀਨਗਰ, ਡਾ. ਰਾਹਲ ਤਨੇਜਾ ਸਾਇੰਟਿਸਟ ਆਈਪੀਆਰ ਜੀਤਨ ਸ਼ੇਤਰਪਾਲ ਮੈਨੇਜਿੰਗ ਡਾਇਰੈਕਟਰ ਕਨੋਰ ਕੋਸਮੇ ਪਾਈਵੇਟ ਲਿਮਿਟਡ, ਐੱਸਪੀਆਰ ਦੇ ਪਹਾਨ ਕੇ ਲੰਜਬਾ ਦੇ ਡਾਇਲੈਕਟਰ ਡਾ ਜੀਡੀ ਗੁਪਤਾ, ਵਾਈਸ ਪਿੰਸੀਪਲ ਡਾ. ਆਰਕੇ ਨਾਰੰਗ, ਐੱਸਪੀਆਰ ਦੇ ਲਬਾ ਪ੍ਰਧਾਨ ਡਾ. ਸਿਰਾਰਥ ਮੰਗਨ ਨੇ ਸਾਂਬੇ ਤੌਰ 'ਤੇ ਦੋਤੀ ਜਗਾ

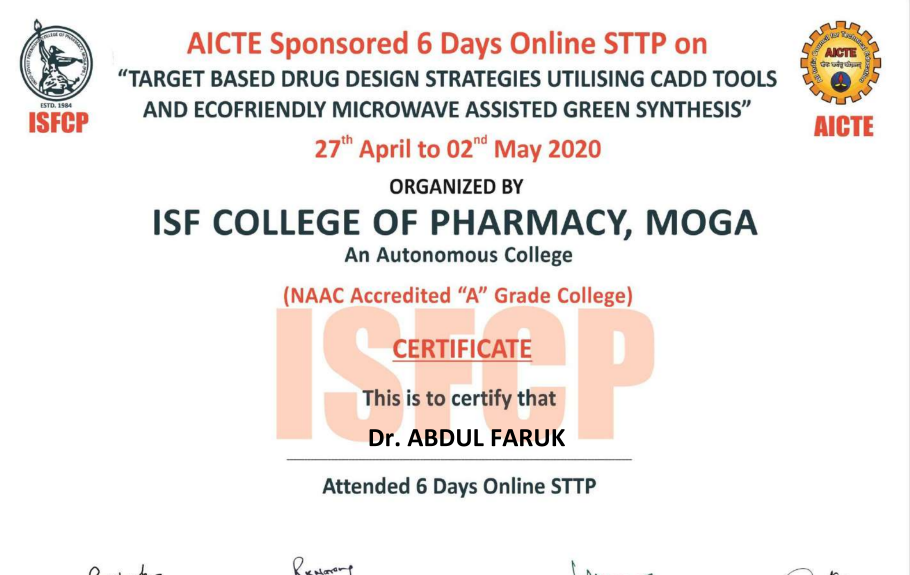
ਇਸ ਸੌਕੇ ਡਾ. ਸਿਧਾਰਥ ਮੇਹਨ ਨੇ ਆਏ ਹੋਏ ਸਾਰੇ ਮਹਿਮਾਨਾਂ ਦਾ ਧੋਨਵਾਦ ਕੀਤਾ। ਕਾਨਫਰੰਸ ਨੇ ਸੰਬੋਧਨ ਕਰਦੇ



ਸੰਸਥਾ ਦੇ ਚੇਅਰਮੇਨ ਪ੍ਰਵੀਨ ਗਰਗ,ਵਾਈਸ ਪ੍ਰਿੰਸੀਪਲ ਡਾ. ਆਰ.ਕੇ ਨਾਰੰਗ ਡਾ ਅਬਦੁਲ ਵਾਰੁਪ ਗੜ੍ਹਵਾਲ ਯੂਨੀਵਰਸਿਟੀ ਸ਼੍ਰੀਨਗਰ ਨੂੰ ਸਨਮਾਨਤ ਕਰਦੇ ਹੋਏ।

ਰੇਏ ਡਾਇਰੈਕਟਰ ਡਾ ਜੀਡੀ ਗੁਪਤਾ ਨੇ ਐਸਪੀਆਰ ਏਲੋਂ ਚੋਲ ਹੀ ਗਤੀਵਿਧੀਆਂ ਤੇ ਸੰਸਾਬ ਬਾਰੇ ਸਾਣਕਾਰੀ ਇਤੀ। ਇਸ ਮੈਂਕੇ ਸਾਇਟਿਸਟ ਡਾ ਰਾਪੂਲ ਤਠੇਜਾ ਨੇ ਪੈਟੇਟ ਦੀ ਉਪਸ਼ੀਨਿਤਾ, ਭਾਸ਼ੀਲਿੰਗ ਤੇ ਆਈਪੇਅਰ ਦੀ ਉਪਸ਼ੀਨਿਤਾ ਬਾਰੇ ਜਟੂ ਕਰਵਾਇਆ। ਉਨ੍ਹਾਂ ਕਿਹਾ ਕਿ ਦੇਸ਼ ਤੋਂ ਵਿਦੇਸ਼ 19 ਆਈਪੈਆਰ ਦੀ ਮੰਗ ਨਿਰੰਤਰ ਚੋਧ ਜ਼ਰੀ ਹੈ।

ਇਸ ਮੌਕੇ ਡਾ. ਅਬਦੁਲ ਫਾਰੁਖ ਨੇ ਆਈਪੀਆਰ ਬਾਰੇ ਦੋਸਦੇ ਹੋਏ ਨੋਬਲ ਡੇਗਰ ਜ਼ਿਲੀਵਜੀ ਸਿਸਟਮ 'ਤੇ ਸਿਸਦਰ ਵਰਕ ਨੂੰ ਪੈਂਟੋਟ, ਹਰਾਮਕਰ ਟੈਕਨਾਲੋਜੀ ਅਤੇ ਤੈਡਿੰਡਸਟਰੀ ਤੋਂ ਕੇ ਜੱਸਦ' ਤੇ ਰੋਲ ਦਿਤਾ। ਇਸ ਸਿੰਧ ਦ੍ਰੇ ਸਿੰਗ ਕਮੇਨੇ ਨੇ ਆਏ ਹੋਏ ਸਾਂਦੇ ਮਹਿਲਾਰਾ ਕੀਤਾ। ਕਾਨਕੰਸ਼ਨਾਂ 'ਚ ਸਨੇਲਾ ਦੇ ਟੇ ਪੈਕਿਲਾਰ ਕੀਤਾ। ਕਾਨਕੰਸ਼ਨਾਂ 'ਚ ਸਨੇਲਾ ਦੀ ਕਾਰਵਾਈ ਵਿਸ਼ ਸਿੰਘ ਸਿਆਇਆ ਟੈਕਨਰਨੀ ਡਾ. ਉਪਿੰਦਰ ਨਗਾਇਜ਼ ਨੇ ਐੱਸਪੀਆਰ ਪੰਜਾਬ ਸਟੇਟ ਪ੍ਰਾਚ ਨੂੰ ਕਾਨਕੰਸ਼ ਦੇ ਅਨਜਲ ਸ਼ਲੇ ਵਧਾਈ ਇਨੀ।



Dr. Pooja Chawla Professor ISF College of Pharmacy, Moga

Organizing Secretary

Dr. R. K. Narang Vice- Principal ISF College of Pharmacy, Moga

Convener

Dr. G. D. Gupta Director-cum-Principal ISF College of Pharmacy, Moga



Parveen Garg Chairman ISF College of Pharmacy, Moga

MoU with Jivanti Welfare and Charitable Trust

<u>Progress Report (Financial Year, 2022 – 2023) with Major</u> <u>Achievements and Observations</u>

Development of large scale seedlings and promotion of cultivation of selected high altitude medicinal plants (*Nardostachys grandiflora*, *Aconitum balfourii*, *Aconitum heterophyllum*, *Picrorhiza kurrooa*, *Saussurea costus* and *Valeriana wallichii*) in farmer's field in high altitude region of Uttarakhand" for easy availability of raw materials as well as conservation of species in natural habitats

Memorandum of Understanding (MOU) between HAPPRC & JWCT (15/9/2020)



Project Sponsored by

Jivanti Welfare and Charitable Trust (JWCT), New Delhi M/S Dabur Research & Development Centre (DRDC), Dabur India Limited (DIL), Sahibabad, Ghaziabad (U.P.), India

At

High Altitude Plant Physiology Research Centre (HAPPRC) Hemvati Nandan Bahuguna Garhwal University, Srinagar (Garhwal), Uttarakhand Ph. 01346-252172, 253760; Fax: 01346- 252070 Principal Investigator: Dr. Vijay Kant Purohit

Project Details

1. Project Proposal: Development of large scale seedlings and promotion of cultivation of selected high altitude medicinal plants (*Nardostachys grandiflora*, *Aconitum balfourii*, *Aconitum heterophyllum*, *Picrorhiza kurrooa*, *Saussurea costus* and *Valeriana wallichii*) in farmer's field in high altitude region of Uttarakhand" for easy availability of raw materials as well as conservation of species in natural habitats.

2. Financial Assistant Providing Agency: Jivanti Welfare and Charitable Trust (JWCT), New Delhi through M/S Dabur Research & Development Centre (DRDC), Dabur India Limited (DIL), Sahibabad, Ghaziabad (U.P.), India

3. Project Implementing Agency: High Altitude Plant Physiology Research Centre (HAPPRC), Hemvati Nandan Bahuguna Garhwal University, Srinagar (Garhwal)-246 174

4. Project Executing Authority: Director, HAPPRC

5. Project Principal Investigator: Dr. Vijay Kant Purohit, Sr. Scientific Officer, HAPPRC

6. Project Period: 3 years

7. Project Cost: Rupees Thirty one lakh three thousand only (Minimum Rupees 31.03 Lakh)

8. Grant Received: Rs. 24,88,300 (Rupees twenty four lakh eighty eight thousnad three hundred only)

9. Financial year: March to March

10. Major objective of the project: Development of large scale seedlings and promotion of cultivation of selected medicinal plants in farmers' fields for easy availability of raw materials as well as conservation of species in natural habitats.

11. Details of objectives/technicalities of the project: Mass scale planation of quality planting materials of high value threatened medicinal species is one of the viable options for conservation and sustainable utilization. It will also provide great livelihood opportunities to the poor and marginal farmers in rural areas. The present project has been proposed in these lines with the following technicalities.

- Mass scale production and multiplication arrangement of quality planting material through nursery development and utilization of other regional resource of public and private sector.
- > Distribution of plantlets for extensive plantations of the species in suitable areas.
- Training and skill development of the farmers for plantation, maintenance/ after care harvesting and post- harvest management of the selected species.
- > To develop plantations and harvesting protocols of the species for future replications.
- To ensure people's participation in long term maintenance of the plantation for their optimum production.
- Income augmentation through livelihood support generation for small and marginal farmers (especially women).

- > Conduct Research and Domestication of the selected species for promoting cultivation.
- Rhizospheric studies of Vatsanabh, Jatamansi etc. will be carried out in joint collaboration between Dabur & HAPPRC.

12. Project Deliverables:

- 1. The plantation in the selected areas of Uttarakhand with an aim to natural resource augmentation and supplement the livelihood of local community through sustainable harvesting.
- 2. Methods to develop sustainable plant part collection in an eco-friendly manner may be by engaging with other partners.
- 3. Both the signing parties to abide by the same and work together to ensure that, the targets of the project are met effectively.
- 4. Submission of report has to be ensured by HAPPRC on quarterly basis as per the prescribed format under CSR norms.
- 5. Social benefit aspects for the outcome of project.
- 6. Sharing of scientific data outcome and publication from this project.

13. Work done so far upto 31 March 2023: To develop the mass scale Quality Planting Material (QPM), i.e. seedlings and further promotion of cultivation in farmers and community owned land preferable in high altitude region of Chamoli, Rudraprayag, Pauri, Tehri, Bageshwar, Nainital, and Pithoragarh districts of Uttarakhand, six highly important medicinal plants have been selected (**Table 1**). Besides the development of mass scale seedlings and promotion of cultivation, the rhizospheric studies of the selected medicinal and aromatic plants have been also proposed. In continuation of the proposed work, the total 6ha (300 nali) of land has been attempted/covered under cultivation. Approximate 2,78,625 seedlings were developed and approximate 1391.62 gm of seeds were collected. During the report period 15 villages, 4 development block and three districts with 577 farmers/villages has been covered through organizing the 15 farmers workshop/training/ plant distribution programme so far. The details of the work performed with photographs are depicted as follows.

Sr.No.	Plant species	Local Name	Geographical Distribution (m asl)	Survival Status	Cultivation Status in Uttarakhand	Specimen Photo of the selected species
	Nardostachys grandiflora	Jatamansi/ Masi	3200-5000	EN	Very poor	
	Aconitum balfourii	Vatsanabha, Mitha Vish	2800-4200	EN	Poor	

 Table 1. List of proposed medicinal and aromatic plants under project.

3.	Aconitum heterophyllum	Atis, Atvika	3000-4500	EN	Moderate	
4.	Picrorhiza kurrooa	Kutki, Kedar kadwi	3000-4500	R-EN	Good	
5.	Saussurea costus	Kuth	2600-4000	R-EN	Good	
6.	Valeriana wallichii	Tagar, Sugandhbala	1500-2500		Very poor	

14. Photographs of the seedlings developed and kept for further growth in shade house at Baniyakund (2460m asl), cultivation in farmers field, monitoring of the works with some other activities performed during the year (April – June, 2022) under J.W.C.T. project.



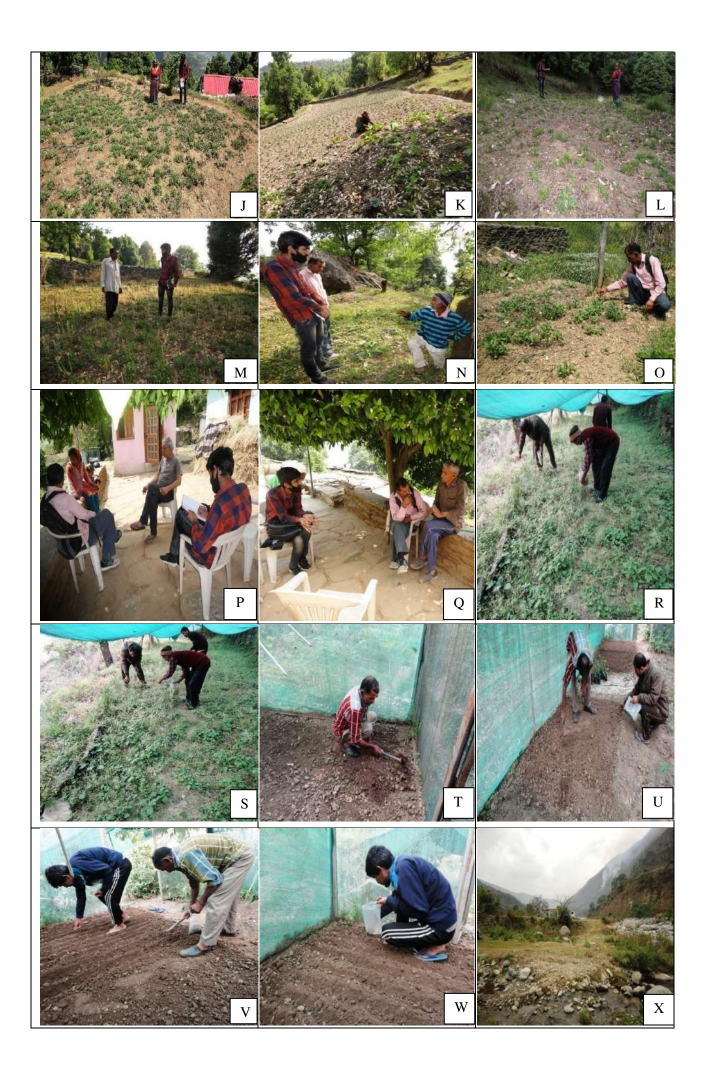






Figure 1. A-D Field visit of village of Partha, E-I Field visit of village of Rushiyana, J-K Field visit of village of Kasbinagar, L-N Field visit of village of Sunaumallah, O-P Seed collection of *Valeriana jatamansi* at our Nursery Kulsari (Chamoli), Q-T Seed sowing of *Saussurea costus*, *Nardostacys grandiflora*, at Kulsari nursery (Chamoli), U-X transport of project related materials from Srinagar to Kulsari nursery (Chamoli), Y-Z,a Field visit of village of Tyuri, b-g,m Field visit of village of Shyalmi, h-I Seed sowing of *Nardostachys grandiflora* and *Aconitum heterophyllum* in Poly house at Baniyakund, k Seedling of *Saussurea costus*, n-s land preparationand construction of Polytunalsat Baniyakund for seed sowing work and seeds of *Aconitum balforii*, *Nardostachys grandiflora* and *Saussurea costus* sowed inside the constructed polytunnels.

Sr. No.	Name of the farmer	Name of the village	Contact number of the farmer	Name of the species cultivating by farmer	Land occupied under cultivatio n (Nali)	Status of the cultivatio n	Additional crop cultivating by farmers
1.	Shri Dalveer Singh	Partha	8057691089	Saussurea costus, Valeriana jatamansi,	0.5	Initiated	Potato, Barley, wheat, Rice, Amaranthus
2.	Smt. Gita Devi	,,	9999620298	Picrorhiza kurrooa	1.0	Well performan ce	Potato, Barley, wheat, Rice, Amaranthus
3.	ShriRaghuv eer Singh	,,		Picrorhiza kurrooa	0.5	Initiated	Potato, Barley, wheat, Rice, Amaranthus
4.	Shri Govind Singh	"		Picrorhiza kurrooa	0.5	Initiated	Potato, Barley,

							wheat, Rice, Amaranthus
5.	Shri Pushkar Singh	,,	8755047543	Picrorhiza kurrooa	0.5	Initiated	Potato, Barley, wheat, Rice
6.	Shri Paan Singh	"			0.5	Initiated	Potato, Barley, wheat, Rice
7.	Shri Gajae Singh	Tran Partha	8979705536	Picrorhiza kurrooa, Rheum emodi	0.5	Initiated	Potato, Barley, wheat
8.	Shri Balwant Singh Negi	"	8192862863	Valerianajatam ansi	0.5	Initiated	Initiated
9.	Smt. Bhawani Devi	"	9756240196	Picrorhiza kurrooa	0.5	Initiated	Potato, Barley, wheat, Rice
10.	ShriRatan Singh	"	-	Picrorhiza kurrooa	0.5	Initiated	Potato, Barley, wheat, Rice
11.	Shri Anand Singh	"	-	Saussurea costus, Valeriana jatamansi, Picrorhiza kurrooa	1.0	Well performan ce	Potato, Barley, wheat, Rice
12.	Shri Madho Singh	"	-	Picrorhiza kurrooa	0.5	Initiated	Potato, Barley, wheat, Rice
13.	Shri Bhuwan Singh Pimoli	"	-	Saussurea costus, Valeriana jatamansi, Picrorhiza kurrooa	0.5	Initiated	Potato, Barley, wheat, Rice
14.	Shri Kuwar Singh (Pardhan)	"	-	Saussurea costus, Picrorhiza kurrooa	1.0	Initiated	Potato, Barley, wheat, Rice
15.	Shri PitamberDu tt	>>	9871084382	Picrorhiza kurrooa	1.0	Initiated	Potato, Barley, wheat, Rice
16.	Shri RamChande r	"	7668581366	Picrorhiza kurrooa	1.5	Well performan ce	Potato, Barley, wheat, Rice
15	<u>(1) : 51</u>	Rusiyana	7 0/04/00/1			XX / 11	
17.	Shri Bharat Singh Rana	"	7060449964	Picrorhiza kurrooa, Aconitum hetrophyllum, Valeriana jatamansi	2.0	Well performan ce	Potato, Barley, wheat, Rice
18.	Shri Sudershan Singh Rana	"	7456966732	Picrorhiza kurrooa, Saussurea costus, Rheum emodi, Valeriana jatamansi	1.0	Well performan ce	Potato, Barley, wheat, Rice
19.	Shri Mohan Singh	"	9634123066	Picrorhiza kurrooa	1.5	Well performan	Potato, Barley,

	Rauthan					ce	wheat, Rice
21.	Shri Mohan	,,	7500677806	Picrorhiza	1.0	Well	Potato,
	Singh		21	kurrooa		performan	Barley,
	Rawat					ce	wheat, Rice
23.	Shri Dinesh	,,	-	Picrorhiza	2.0	Well	Potato,
	Singh			kurrooa		performan	Barley,
	Rawat					ce	wheat, Rice
24.	Shri	,,	-	Picrorhiza	1.5	Well	Potato,
	Pushkar			kurrooa		performan	Barley,
	Singh Rana					ce	wheat, Rice
25.	Shri Puna	"	-	Picrorhiza	0.5	Initiated	
	Singh			kurrooa			
	Rawat						
26.	Shri	"	9878149527	Saussurea	-	Interested	
	Mahendar			costus			
	Singh						
27	G1 .	Kasbinagar	0065040100	D: /:	2.0	XX 7 11	D 1
27.	Shri	Kasbinagar	8865042109	Picrorhiza	2.0	Well	Barley,
	Balwant			kurrooa		performan	wheat, Rice
20	Ram		9690499157			ce	W-1+
28.	Shri Gajpal Ram	"	9090499137	-	-	Interested	Walnut, Rice,
	Kalli						Wheat,
29.	Shri	Sunaumall	9557461512	-		Interested	Potato,
27.	DevSinghB	a	JJJ7401312	_	_	interested	wheat, Rice
	handari	a					wheat, Rice
30.	Gudi Devi	,,		-	_	Interested	,,
31.	Kundan	,,	8979011384	-	_	Interested	Potato,
	Singh	"					Coriander,
	U						Wheat, Rice
32.	Shri Kalm	Tyuri	7455807117	-	-	Interested	Wheat,
	Singh	-					Rice,Barley,
	Semwal						Millet
33.	Smt. Sharita	,,	7895949115	-	-	Interested	Wheat,
	Devi						Rice,Barley,
							Millet
34.	Dr. D.S.	"	9412404077	Picrorhiza	50.0	Well	Wheat,
	Rawat			kurrooa		performan	Rice,Barley,
25	G1 .	T T 1	7500650201	D: /:	0.5	ce	Millet
35.	Shri	Ushara	7500659381	Picrorhiza	0.5	Well	Wheat,
	Yogendar	(Shayamli)		kurrooa		performan	Rice,Barley,
26	Singh Shri Parmod					ce	Millet
36.	Shri Parmod Singh	"	-	-	-	Interested	Wheat, Rice,Barley,
	Bejwal						Millet
37.	ShriPardeep		_	-		Interested	Wheat,
57.	Singh	"			-	meresteu	Rice,Barley,
	Bejwal						Millet
38.	Shri	,,	_	_	_	Interested	Wheat,
	Rajender	,,					Rice,Barley,
	Singh						Millet
39.	Shri Veer	,,		Picrorhiza	2.0	Well	Wheat,
	Singh			kurrooa		performan	Rice,Barley,
	Bejwal					ce	Millet
40.	Shri Perbal	,,		Picrorhiza	2.0	Well	Wheat,
	Singh			kurrooa		performan	Rice,Barley,
	Bejwal					ce	Millet
41.	Shri Satveer	"	9389307557	Picrorhiza	1.0	Well	Wheat,
	Singh			kurrooa		performan	Rice,Barley,
10	Bejwal		0.0007001	D . 7.	1.0	ce	Millet
42.	Shri Kalm	"	9690079914	Picrorhiza	1.0	Well	Wheat,

	Singh Bejwal			kurrooa		performan ce	Rice,Barley, Millet
43.	Shri Pardeep Singh Bejwal	,,		Picrorhiza kurrooa	0.5	Well performan ce	Wheat, Rice,Barley, Millet
44.	Shri Satyender Singh Bejwal	,,		Picrorhiza kurrooa	0.5	Well performan ce	Wheat, Rice,Barley, Millet
45.	Shri noop Singh Negi	"		Picrorhiza kurrooa	2.0	Well performan ce	Wheat, Rice,Barley, Millet
45.	Shri Ramesh Singh Bejwal	,,		Picrorhiza kurrooa	1.5	Well performan ce	Wheat, Rice,Barley, Millet
47.	Shri Kuwar Singh (Pardhan)	"		Picrorhiza kurrooa	3.0	Well performan ce	Wheat, Rice,Barley, Millet
48.	Shri Darshan Singh Bejwal	,,		Picrorhiza kurrooa	-	Interested	Wheat, Rice,Barley, Millet
49.	ShriGajae Singh			Picrorhiza kurrooa	0.5	Well performan ce	Wheat, Rice,Barley, Millet
50.	ShriVirenda r Singh Negi	"		Picrorhiza kurrooa	-	Interested	Wheat, Rice,Barley, Millet
51.	Shri Jai Singh Bejwal			Picrorhiza kurrooa	-	Interested	Wheat, Rice,Barley, Millet
52.	ShriSumant Singh		8958342983	Picrorhiza kurrooa	0.5	Initiated	Wheat, Rice,Barley, Millet
53.	Shri Mohan Singh (Teacher)		-	-	-	Interested	Wheat, Rice,Barley, Millet
54.	ShriDilip Singh		-	-	-	Interested	Wheat, Rice,Barley, Millet

15. Information about sale of raw as well as planting material of selected species recorded during field visit.

Farmers get income for their livelihood:

1. Shri Balwant Ram (Kasbinagar) sold 18 kg dry roots of kutki worth of rupees 27000 and also sold 20000 cuttings of kutki worth of rupees 20000.

2. Shri Gajpal Ram (Kasbinagar) sold 10 kg seed of kuth@ 500/kg and earn rupees 5000.

3. Shri Sudarshan Singh Rana (Ruisiyaan) sold 20 kg dry roots of kutki worth of rupees 29000.

4. ShriYogendrasingh (Shayalmi, ushara) sold 20 kg seed of kuth@ 200/ kg worth of rupees 2000 and also sold 50 kg dry root of kutki@ Rs. 1200/ kg and earn rupees 60000 through HRDI, Gopeshwar, mandal.

5. Anoop Singh Negi (Shayalmi, ushara) sold 15 kg of kutki@ Rs. 1200/ kg worth of rupees 18000 throughHRDI, GopeshwarMandal.

6. Tungnath group of farmers (Shayalmi, ushara) sold 8 lakh kutki cutting@ Rs. 1.0/cutting worth of rupees 800000 through HRDI, Gopeshwar, Mandal.

7. Dr. D.S. Rawat (Tyuri, Guptkashi) sold 2.0quntals kutki @ Rs. 1450/ kg worth rupees 290000 through HRDI, Gopeshwarmandal.

16. Details of seedlings distributed to faermers for promotion of cultivation of selected species 2022-2023.



Figure 2. Uprooting and packing of quality planting material of *P. kurooa*, (*A-D*), *N. grandiflora* (F-H), *A. heterophyllum* (I-K) and *V.wallichii* from field stations Baniyakund, Pothivasa and Kulsari of HAPPRC for farmers distribution under JWCT project.

Sr. NO.	Name of the species	Number of seedlings distributed
1.	P. kurrooa	1,91,550
2.	N. grandiflora	19.930
3.	A. heterophyllum	14,045
4.	V. wallichii	48,100
5.	S. costus	5000
6.	Total Number of seedlings distributed	2,78,625

Table 3. Summary of seedlings distributed to farmers in the month of July-September 2022.

17. Details of farmers workshop/training/ plant distribution programme organized during the 2022-2023.

Sr.No.	Activity	No.
1.	On-farm workshop cum training/plant distribution programme	15
2.	Toatl Number of district covered under cultivation	03
3.	Total number of institutional members (HAPPRC) attended	68
	programme	
4.	Total number of Males farmers (GEN) attented programme	280
5.	Total number of Females farmers (GEN) attended programme	235
6.	Total number of Males farmers (SC) attended programme	41
7.	Total number of Females farmers (SC) attended programme	21
8.	Total number of farmers/participants attended meetings	645 (577 farmers)

Table 4. Summary of Farmer's on-farm workshop cum training/plant distribution programme.

Note- GEN (General), SC (Scheduled Cast)

Table 5. Summary of seedlings distributed to farmers in the month of July-September 2022.

Sr. NO.	Name of the species	Area covered under cultivation
1.	P. kurrooa	4.75 acre (96 Nali)
2.	N. grandiflora	0.50 acre (10 Nali)
3.	A. heterophyllum	0.35 acre (7.02 Nali)
4.	V. wallichii	1.19 acre (24.05Nali)
5.	S. costus	0.12 acre (2.5 Nali)
6.	Total area occupied under cultivation/	6.91 acre (139.57 Nali)
	plantation	

Table 6. List and Summary of villagers/farmers and other participants attended one day on farm workshop/Training/plants distribution programme organized during July to August 2022.

Sr.No.	Name of	Participants				Other	Institutional	Total
	the village	Male (Gen)	Female (Gen)	Male (SC)	Female (SC)	particip ants	members (HAPPRC)	partici pants
1.	Teela	37	04	08	10	-	05	64
2.	Pala-Kurali	24	36	0	0	06	05	71
3.	Gainthnda	23	14	10	5	01	05	58
4.	Kaviltha	9	19	0	0	03	05	36
5.	Jaal Malla	11	18	0	0	02	05	36
6.	Jaal Talla	09	28	0	0	02	05	44
7.	Chaumasi	11	20	0	0	02	05	38
8.	Kulpudi	11	05	01	0	0	04	21
9.	Rushyan	10	0	0	0	0	03	13

10.	Ratgaov	13	0	02	0	0	03	18
11.	Taal	12	8	05	0	0	04	29
12.	Syanri Bang	27	24	0	0	05	04	60
13.	Pagna	25	17	01	04	-	04	51
14.	Sitel	31	25	03	-	11	07	77
15.	Syanri Bhair	27	17	11	02	01	04	62
Total	participants	280	235	41	21	33	68	678

Table 7. Seedlings/plants provided /distributed to villagers/farmers during on farm levelworkshop /training/plant distribution programme organized during July to August 2022.

Sr.No.	Name		Name of the plants distributed				
	of the	<i>P</i> .	<i>A</i> .	<i>N</i> .	<i>V</i> .	S. costus	of plants
	villages	kurrooa	heterophyllu	grandiflora	wallichii		distributed
			т				
1.	Teela	14,750	2950	-	5900	-	
2.	Pala	30,000	3000	6000	6000	-	23,600
	Kurali						
3.	Gaithana	10,400	1000	2000	5,200	-	45,000
4.	Kaviltha	2,800	-	-	5,600	-	18,600
5.	Jaal	5,800	-	-	5,800	-	8,400
	Malla						
6.	Jaal Talla	7,400	-	-	7,400	-	11,600
7.	Chaumasi	6,200	775	1,550	-	-	14,800
8.	Kulpudi	5,100	-	-	1700	-	8,525
9.	Rushyan	2,000	1000		2500	-	6,800
10.	Ratgaov	22,500	1000	-	3,000		5,500
11.	Taal	10,000	-	-	5,000	5,000	26,500
12.	Syanri	10,200	2,550	1,530			20,000
	Bangali						
13.	Pagna	23,500	-	-	-	-	14,280
14.	Sitel	29,500	1,770	8,850	-	-	23,500
15.	Syanri	11,400		-			40,120
	Bhainti						
Total		1,91,550	14,045	19,930	48,100	5000	2,78,625

18. Details of farmers workshop cum training/plant distribution programme organsied during the year 2022-2023

The numbers of farmers workshop cum training/plant distribution programme were organized under JWCT project in different area of Uttarkhand for acheiveing the cultivation target of selecdted species. During the entire period of workshop cum training programme the project staff and representative of institute briefs about the aim of the project, role of HAPPRC, work plan of participants followed by diffusion of technical information about cultivation and distribution of plants of selected species to farmers.

Table 8. List of villagers/farmers participated in one day On-farm farmer's workshop cum training programme organized at Village Pala Kurali, Block, Jakholi, Rudraprayag on 12/July/2022 under J.W.C.T. Project.

Sr.No.	Villagers Name	Father/Husband Name	Address
1.	Shri Suryapal Singh Rana	Shri Asad Singh	Vill-Pala Kurali
			Block & Distt Jakholi, Rudraprayag
			Contact No 7465964681
			Aadhar No 304465923973

2.	Shri Ravindra Singh Rana	Shri Jabbar Singh Rana	Vill-Pala Kurali
2.	Shiri Kuvinaru Singii Kunu	Shiri Jubbar Shirgh Kuna	Block & Distt Jakholi, Rudraprayag
			Contact No 9389703778
			Aadhar No 787809975948
3.	Shri Narottam Singh Rana	Shri Ameer Singh Rana	Vill-Pala Kurali
5.	Shiri Nafottalli Shigli Kalla	Shiri Ameer Shigh Kana	Block & Distt Jakholi, Rudraprayag
			Contact No 7818039564
			Aadhar No 893805772806
4.	Shri Rakesh Singh Rana	Shri Umrav Singh Rana	Vill-Pala Kurali
			Block & DisttJakholi, Rudraprayag
			Contact No 7248625456
5.	Shri Bhagwan Singh Rana	Shri Gabbar Singh Rana	Vill-Pala Kurali
			Block & DisttJakholi, Rudraprayag
			Contact No 7534063904
			Aadhar No 533744433892
6.	Smt. Beena Devi	Shri Dinesh Rana	Vill-Pala Kurali
			Block & Distt Jakholi, Rudraprayag
			Contact No 7055968537
			Aadhar No 910561568370
7.	Smt. Pushpa Devi	Shri Surendra Rana	Vill-Pala Kurali
7.			
			Block &DisttJakholi, Rudraprayag Contact No 7505824758
			Aadhar No 805532445130
8.	Smt. Sunita Devi	Shri Trilok Singh	Vill-Pala Kurali
			Block & DisttJakholi, Rudraprayag
			Contact No 6398821034
			Aadhar No 626478618985
9.	Shri Vinod Singh	Shri Vachan Singh	Vill-Pala Kurali
	6	6	Block & DisttJakholi, Rudraprayag
			Contact No 8192959490
			Aadhar No 458712587278
10.	Smt. Chaita Devi	Shri Meharvan Singh	Vill-Pala Kurali
10.	Shit. Chata Devi	Shiri Wenar van Shigh	Block & DisttJakholi, Rudraprayag
			Aadhar No 727263497923
11.	Smt. Shashi Devi	Classi Maathaa aa laa Cimah	
11.	Smt. Snasni Devi	Shri Madhusudan Singh	Vill-Pala Kurali
			Block & DisttJakholi, Rudraprayag
			Contact No 7505123979
			Aadhar No 604241638671
12.	Smt. Rinki Devi	Shri Rajesh Singh	Vill-Pala Kurali
			Block & DisttJakholi, Rudraprayag
			Contact No 8941886509
			Aadhar No 625309287735
13.	Smt. Vijaya Devi	Shri Satpal Singh	Vill-Pala Kurali
			Block & DisttJakholi, Rudraprayag
			Contact No 9012268140
			Aadhar No 750765191651
14.	Smt. Jasdei Devi	Shri Jai Singh	Vill-Pala Kurali
14.			
			Block & DisttJakholi, Rudraprayag
			Contact No 6396742801
			Aadhar No 796011362357
15.	Smt. Rajni Devi	Shri Kuldeep Singh	Vill-Pala Kurali
			Block & DisttJakholi, Rudraprayag
			Contact No 8476944137
			Aadhar No 676776074299
16.	Smt. Guddi Devi	Shri Balveer Singh	Vill-Pala Kurali
			Block & DisttJakholi, Rudraprayag
			Contact No 7409640836
			Aadhar No 610278932538
17.	Shri Govind Singh	Shri Bhagwan Sirah	Vill-Pala Kurali
1/.	DILL OOVIIIQ DILIZII	Shri Bhagwan Singh	v III=F ala INUI all

			Block &DisttJakholi, Rudraprayag Contact No 9625075469 Aadhar No 729979632867
18.	Shri Manveerendra Singh	Shri Bachan Singh	Vill-Pala Kurali Block &DisttJakholi, Rudraprayag Contact No 8958135079 Aadhar No 554154845552
19.	Smt. Veena	Shri Raghuveer Singh	Vill-Pala Kurali Block &DisttJakholi, Rudraprayag Contact No 8865864816 Aadhar No 549153943622
20.	Smt. Vineeta Devi	Shri Narendra Singh	Vill-Pala Kurali Block &DisttJakholi, Rudraprayag Contact No 7417018521 Aadhar No 803352129529
21.	Smt. Anita Devi	Shri Bhupendra Singh	Vill-Pala Kurali Block &DisttJakholi, Rudraprayag Contact No 8449135591 Aadhar No 760917897450
22.	Smt. Raji Devi	Shri Dhan Singh	Vill-Pala Kurali Block &DisttJakholi, Rudraprayag Contact No 8445671828 Aadhar No 260508792242
23.	Smt. Roopa Devi	Shri Rukam Singh	Vill-Pala Kurali Block &DisttJakholi, Rudraprayag Contact No 6397728517 Aadhar No 676265296251
24.	Smt. Pushpa Devi	Shri Guddu Singh	Vill-Pala Kurali Block &DisttJakholi, Rudraprayag Contact No 7055631279 Aadhar No 783159581632
25.	Smt. Sumitra Devi	Shri Uday Singh	Vill-Pala Kurali Block &DisttJakholi, Rudraprayag Contact No 7247821212 Aadhar No 380423562797
26.	Smt. Sarita Devi	Shri Chain Singh	Vill-Pala Kurali Block &DisttJakholi, Rudraprayag Contact No 7088198816 Aadhar No 935920306561
27.	Smt. Darshni Devi	Shri Bachan Singh	Vill-Pala Kurali Block &DisttJakholi, Rudraprayag Contact No 7533937834 Aadhar No 457527648630
28.	Smt. Anuradha Devi	Shri Jairaj Singh	Vill-Pala Kurali Block &DisttJakholi, Rudraprayag Contact No 8006531823 Aadhar No 628513216638
29.	Smt. Suneeta Devi	Shri Nathi Singh	Vill-Pala Kurali Block &DisttJakholi, Rudraprayag Contact No 7900582560 Aadhar No 879315254402
30.	Smt. Lakshmi	Shri Balwant Singh	Vill-Pala Kurali Block &DisttJakholi, Rudraprayag Contact No 7505177982 Aadhar No 228204243104
31.	Shri Padam Singh	Shri Karan Singh	Vill-Pala Kurali Block &DisttJakholi, Rudraprayag Contact No 9411398556 Aadhar No 327512788635

32.	Shri Ravindra Singh	Shri Pratap Singh	Vill-Pala Kurali
52.	Shiri Kavindia Shigh	Shiri Fidup Shigh	Block & DisttJakholi, Rudraprayag
			Contact No 8938928927
			Aadhar No 576834695884
33.	Shri Baishak Singh	Shri Musha Singh	Vill-Pala Kurali
55.	Shiri Duishak Shigh	Shiri Mushu Shigh	Block & DisttJakholi, Rudraprayag
			Aadhar No 301941080183
34.	Shri Suresh Singh	Shri Chhota Singh	Vill-Pala Kurali
51.	Shiri Sureshi Shirgh	Shiri Chinota Shigh	Block & DisttJakholi, Rudraprayag
			Contact No 9355495406
			Aadhar No 653669977356
35.	Shri Sate Singh	Shri Tirpal Singh	Vill-Pala Kurali
55.	Shiri Sute Shigh	Sini inpu Singn	Block & DisttJakholi, Rudraprayag
			Contact No 9967377934
			Aadhar No 726808914400
36.	Shri Jaspal Singh	Shri Gabar Singh	Vill-Pala Kurali
	B	2 2B	Block & DisttJakholi, Rudraprayag
			Contact No 8449627193
			Aadhar No 776062928026
37.	Shri Leelanand Thapliyal	Shri Narayan Dutt	Vill-Pala Kurali
	1 5	5	Block & DisttJakholi, Rudraprayag
			Contact No 9720756117
			Aadhar No 655872115578
38.	Smt. Roshani Devi	Shri Pushkar Singh	Vill-Pala Kurali
		C	Block & DisttJakholi, Rudraprayag
			Contact No 9548441648
			Aadhar No 595553937137
39.	Smt. Maya Devi	Shri Narendra Singh	Vill-Pala Kurali
			Block & DisttJakholi, Rudraprayag
			Contact No 7251926729
			Aadhar No 560928341678
40.	Shri Rakesh Singh	Shri Chhota Singh	Vill-Pala Kurali
			Block & DisttJakholi, Rudraprayag
			Contact No 9756823301
			Aadhar No 298036357588
41.	Shri Mukesh Singh	Shri Chhota Singh	Vill-Pala Kurali
			Block & DisttJakholi, Rudraprayag
			Contact No 8958560566
			Aadhar No 752890057445
42.	Shri Digraj Singh Rana	Shri Dinesh Singh	Vill-Pala Kurali
			Block & DisttJakholi, Rudraprayag
			Contact No 8006239765
40			Aadhar No 816754735614
43.	Shri Virendra Singh	Shri Amar Singh	Vill-Pala Kurali
			Block & DisttJakholi, Rudraprayag
			Contact No 8126634395
A_A	Smt Kurner Der	Chui Dalah Chu-h	Aadhar No 778080416049
44.	Smt. Kuwari Devi	Shri Daleb Singh	Vill-Pala Kurali Block & Diett, Jakholi, Budraprayag
			Block &DisttJakholi, Rudraprayag Contact No 97560069779
			Aadhar No 527869878628
45.	Smt. Anari Devi	Shri Chait Sinch	Vill-Pala Kurali
43.	Sint. Anari Devi	Shri Chait Singh	
			Block &DisttJakholi, Rudraprayag Contact No 7055192088
			Aadhar No 969845889216
46.	Smt. Deepa Devi	Shri Pradeep Singh	Vill-Pala Kurali
40.		Sint radeep Singh	Block & DisttJakholi, Rudraprayag
			Contact No 7536069550
			Aadhar No 962460998378
L	I		1100- 702T00770570

47.	Smt. Lakshmi Devi	Shri Ajay Shankar Singh	Vill-Pala Kurali
			Block & DisttJakholi, Rudraprayag
			Contact No 9627625954
			Aadhar No 248186208697
48.	Smt. Kavita Devi	Shri Makan Singh	Vill-Pala Kurali
		6	Block & DisttJakholi, Rudraprayag
			Contact No 7409438199
			Aadhar No 559933753205
49.	Smt. Usha Devi	Shri Jatan Singh	Vill-Pala Kurali
12.		Shiri Suturi Shigh	Block & DisttJakholi, Rudraprayag
			Contact No 8475865365
			Aadhar No 661871755461
50.	Shri Chain Singh	Shri Avtar Singh	Vill-Pala Kurali
50.	Shiri Chum Shigh	Shiri Avtar Shigh	Block & DisttJakholi, Rudraprayag
			Contact No 7088198816
			Aadhar No 341313510998
51.	Shri Kamana Bhandari	Shri D.S.Bhandari	Vill-Pala Kurali
51.	Shifi Kamana Dhandan	Shiri D.S.Bhandari	Block & DisttJakholi, Rudraprayag
			Contact No 8475980471
50	Shri Sunil Kumar Maithani	Shri D.N.Maithani	Vill-Pala Kurali
52.	Shri Sunil Kumar Maithani	Shri D.N.Maithani	
50	Smt. Bharti Devi	Chui Douis et De 1	Block &DisttJakholi, Rudraprayag
53.	Smt. Bharti Devi	Shri Ranjeet Rawla	Vill-Pala Kurali
			Block & DisttJakholi, Rudraprayag
			Contact No 8954305464
54.	Shri Ashwal Gaur	Shri H.M.Gaur	Vill-Pala Kurali
			Block & DisttJakholi, Rudraprayag
			Contact No 7467843141
55.	Shri Subhash Singh Rana	Shri Pratap Singh	Vill-Pala Kurali
			Block & DisttJakholi, Rudraprayag
			Contact No 7500301843
			Aadhar No 491433819868
56.	Shri Hayat Singh	Shri Chatar Singh	Vill-Pala Kurali
			Block & DisttJakholi, Rudraprayag
			Contact No 8954392188
			Aadhar No 554510156857
57.	Smt. Sona Devi	Shri Roshan Singh	Vill-Pala Kurali
			Block & DisttJakholi, Rudraprayag
			Contact No 7466042645
			Aadhar No 416194559931
58.	Shri Raghuveer Singh	Shri Jeet Singh	Vill-Pala Kurali
			Block & DisttJakholi, Rudraprayag
			Contact No 9675347529
			Aadhar No 829643787949
59.	Shri Madan Singh	Shri Shyam Singh	Vill-Pala Kurali
		_	Block & DisttJakholi, Rudraprayag
			Contact No 9645347529
			Aadhar No 431222770664
60.	Smt. Sulochana Devi	Shri Sukhchain Singh	Vill-Pala Kurali
			Block & Distt Jakholi, Rudraprayag
			Contact No 7617576523
			Aadhar No 283472947749
61.	Smt. Saukari Devi	Shri Mangal Singh	Vill-Pala Kurali
			Block & DisttJakholi, Rudraprayag
			Contact No 9818623846
			Aadhar No 443187721665
62.	Smt. Mamta Devi	Shri Chaman Singh	Vill-Pala Kurali
02.			Block & DisttJakholi, Rudraprayag
			Contact No 7983655985
			Aadhar No 961530882478
			maunai 110,- 701550002470

63.	Smt. Thuma Devi	Shri Puran Singh	Vill-Pala Kurali
		_	Block & Distt Jakholi, Rudraprayag
			Aadhar No 623062754883
64.	Smt. Darshni Devi	Shri Veer Singh	Vill-Pala Kurali
			Block & Distt Jakholi, Rudraprayag
			Aadhar No 871325568537
65.	Shri Beerbal Singh Rana	-	Vill-Pala Kurali
			Block & Distt Jakholi, Rudraprayag
66.	Dr. Vijay Kant Purohit	Shri A. P. Purohit	HAPPRC, Contact No 9456531715
67.	Shri Kailash Kandpal	Shri B. P. Kandpal	HAPPRC, Contact No 7302808941
68.	Shri Jaidev Chauhan	Shri B. S. Chauhan	HAPPRC, Contact No 8126211560
69.	Shri Vipin Rawat	Shri P. S. Rawat	HAPPRC, Contact No 9458113893
70.	Shri Kamal Pundir	Shri G. S. Pundir	HAPPRC, Contact No 9540468782



Figure 3. Photographs of On-farm farmers workshop cum training/plant distribution programme organized at Pala Kurali village, district Rudraprayag on 12/07/2022. Delivered technical knowledge by Dr. V.K. Purohit (SSO) to farmers/participants (A-G), distributed plant material to farmers (H-J), farmers/participants group photograph (K-L).

Table 9. List of Villagers/Farmers participated in one day On-farm farmers workshop cumtraining/plant distribution programme for promotion of cultivation of MAP's at Village GainthanaBlock, Jakholi, Rudraprayag on 12/7/2022 under J.W.C.T. Project.Total participants: 54 (male- 35, female-19)

Sr.No.	Villagers Name	Father/Husband Name	Address
1.	Shri Sarveer Singh	Late Shri Buddhi Singh	Vill-Gainthana
	Mengwal		Block & Distt Jakholi, Rudraprayag
			Contact No 9758634254
2.	Shri Anusuya Prasad	Shri Balveer Chomwal	Vill-Gainthana
			Block & Distt Jakholi, Rudraprayag
			Contact No 7251079688
3.	Shri Harendra Singh	Shri Govind Singh	Vill-Gainthana
			Block & Distt Jakholi, Rudraprayag
			Contact No 9758656459
4.	Shri Mohan Lal Shah	Shri Dillu Shah	Vill-Gainthana
			Block & Distt Jakholi, Rudraprayag
			Contact No 8958395424
5.	Shri Shivraj Singh	Shri Katag Singh	Vill-Gainthana
			Block & Distt Jakholi, Rudraprayag
			Contact No 9759412169
6.	Shri Virendra Singh	Shri Mahendra Singh	Vill-Gainthana
			Block & Distt Jakholi, Rudraprayag
			Contact No 9702560573
			Aadhar No 308359136715
7.	Shri Sajjan Singh	Late Shri Dal Bahadur Sing	Vill-Gainthana
			Block & Distt Jakholi, Rudraprayag
			Contact No 8393931593
			Aadhar No 566212677130
8.	Shri Prakash Chand	Shri Prem Shah	Vill-Gainthana
			Block & Distt Jakholi, Rudraprayag
			Contact No 7409982828
9.	Shri Pritam Singh	Late Shri Umrav Singh	Vill-Gainthana
			Block & Distt Jakholi, Rudraprayag
			Contact No 9675730456
10.	Shri Gambhir Singh	Late Shri Jagat Singh	Vill-Gainthana
			Block & Distt Jakholi, Rudraprayag
			Contact No 9619581358
11.	Shri Mahesh Lal	Shri Shobha Lal	Vill-Gainthana
			Block & Distt Jakholi, Rudraprayag
			Contact No 7830956067
12.	Shri Chiranji Lal	Shri Shibbu Lal	Vill-Gainthana
			Block & Distt Jakholi, Rudraprayag
			Contact No 8006027450
13.	Shri Ranvir Singh	Late Shri Keshar Singh	Vill-Gainthana
			Block & Distt Jakholi, Rudraprayag
			Contact No 8650702284
14.	Shri Lakhan Singh	Late Shri Bhauh Singh	Vill-Gainthana
	-	-	Block & Distt Jakholi, Rudraprayag
			Contact No 9675346365
15.	Shri Ram Lal	Shri Udai Lal	Vill-Gainthana

			Block & Distt Jakholi, Rudraprayag
			Contact No 8393857913
16.	Shri Guddu Singh	Shri Chand Singh	Vill-Gainthana
10.	Shiri Guddu Shigii	Shiri Chand Shigh	Block &DisttJakholi, Rudraprayag
			Contact No 7902146678
17.	Shri Ranjeet Singh	Late Shri Hayat Singh	Vill-Gainthana
17.	Shiri Kanjeet Singh	Late Shiri Hayat Shigh	
			Block & DisttJakholi, Rudraprayag
10			Contact No 9568449256
18.	Shri Kapoor Lal	Late Shri Keshru Lal	Vill-Gainthana
			Block & DisttJakholi, Rudraprayag
			Contact No 9719149034
19.	Smt. Mamta Devi	Shri Mohan Singh	Vill-Gainthana
			Block & DisttJakholi, Rudraprayag
			Contact No 7249942581
20.	Smt. Seema Devi	Shri Vikram Singh	Vill-Gainthana
			Block & DisttJakholi, Rudraprayag
			Contact No 7830128895
21.	Smt. Pinki Devi	Shri Virendra Singh	Vill-Gainthana
			Block & DisttJakholi, Rudraprayag
			Contact No 8954324932
22.	Smt. Shashi Devi	Shri Pradeep	Vill-Gainthana
		1	Block & DisttJakholi, Rudraprayag
			Contact No 7409344002
23.	Smt. Rukma	Shri Sabbal	Vill-Gainthana
25.	Sint. Rukina	Shiri Subbul	Block & DisttJakholi, Rudraprayag
24.	Smt. Chhoti Devi	Shri Indra Singh	Vill-Gainthana
27.		Shiri indra Singh	Block & DisttJakholi, Rudraprayag
			Contact No 8192065392
25.	Smt. Neema Devi	Shri Dailanun Sinah Manar	
23.	Sint. Neema Devi	Shri Rajkapur Singh Mengy	
			Block & DisttJakholi, Rudraprayag
26			Contact No 9627484106
26.	Smt. Rajni Devi	Shri Meharban Singh	Vill-Gainthana
~ ~ ~			Block &DisttJakholi, Rudraprayag
27.	Smt. Sharmila Devi	Shri Vijaypal Singh Mengw	
		Singh	Block & DisttJakholi, Rudraprayag
			Contact No 9536661865
28.	Smt. Rajni Devi	Shri Surjeet Singh Mengwa	
			Block & DisttJakholi, Rudraprayag
			Contact No 8057052123
29.	Smt. Sharmila Devi	Shri Surjeet Singh Mengwa	Vill-Gainthana
			Block & DisttJakholi, Rudraprayag
			Contact No 9627484106
30.	Smt. Rajni Devi	Shri Surjeet Singh Mengwa	Vill-Gainthana
	-		Block & DisttJakholi, Rudraprayag
			Contact No 8057052123
31.	Shri Prakash Singh	Shri Umed Singh	Vill-Gainthana
			Block & DisttJakholi, Rudraprayag
			Contact No 7251804850
32.	Shri Avtar Singh	Shri Mohan Singh	Vill-Gainthana
54.		Shiri Wonan Shigh	Block & DisttJakholi, Rudraprayag
			Contact No 7251925681

	~ ~~		Aadhar No 306624176354
33.	Smt. Kastura	Late Shri Prem Singh	Vill-Gainthana
			Block & DisttJakholi, Rudraprayag
34.	Shri Man Singh	Shri Nakul Singh	Vill-Gainthana
	Nepali		Block & DisttJakholi, Rudraprayag
			Contact No 7088496507
35.	Shri Vikram Singh	Shri Bal Singh	Vill-Gainthana
			Block & DisttJakholi, Rudraprayag
36.	Shri Kripal Singh	Late Shri Umrav Singh	Vill-Gainthana
		_	Block & DisttJakholi, Rudraprayag
			Contact No 8006626928
37.	Shri Pratap Singh	Late Shri Hukam Singh	Vill-Gainthana
			Block & DisttJakholi, Rudraprayag
38.	Shri Rajkumar	Late Shri Umrav Singh	Vill-Gainthana
20.	Sini Rujitaniai	Luc Shir China Shigh	Block & DisttJakholi, Rudraprayag
39.	Shri Rajendra Shah	Shri Prem Shah	Vill-Gainthana
57.	Shiri Kajendra Shan	Shiri i tem Shan	Block &DisttJakholi, Rudraprayag
40.	Shri Jeewan Lal	Shri Paatu Lal	Vill-Gainthana
4 0.	Shiri Jeewali Lal	SIIII I aalu Lai	Block & DisttJakholi, Rudraprayag
			Contact No 8194049387
41	Chai Kabutan Cinah	Chai Chin Cin ab	
41.	Shri Kabutar Singh	Shri Shiv Singh	Vill-Gainthana
			Block & DisttJakholi, Rudraprayag
			Contact No 9548826928
42.	Smt. Phooldei	Late Shri Amar Singh	Vill-Gainthana
			Block & DisttJakholi, Rudraprayag
43.	Smt. Mamta Devi	Shri Om Prakash	Vill-Gainthana
			Block & DisttJakholi, Rudraprayag
44.	Smt. Pooja	Shri Rajendra Chomwal	Vill-Gainthana
			Block & DisttJakholi, Rudraprayag
			Contact No 8650903653
45.	Smt. Rukmani Devi	Shri Jagori Chomwal	Vill-Gainthana
			Block & DisttJakholi, Rudraprayag
46.	Smt. Prabha Devi	Shri Manoj Kumar	Vill-Gainthana
			Block & DisttJakholi, Rudraprayag
			Contact No 7617643841
47.	Smt. Vineeta Devi	Shri Kmalesh Shah	Vill-Gainthana
			Block & DisttJakholi, Rudraprayag
			Contact No 9536747874
48.	Kumari Sweta	Shri Manohar Chomwal	Vill-Gainthana
			Block & DisttJakholi, Rudraprayag
			Contact No 7830163448
49.	Shri Kapoor Shah	Shri Bhupati Shah	Vill-Gainthana
	T	T	Block & DisttJakholi, Rudraprayag
			Contact No 9761681259
50.	Smt. Gaura Devi	Shri Shurveer Singh	Vill-Gainthana
50.		Shiri Shur voor Shigh	Block & DisttJakholi, Rudraprayag
			Contact No 7251811323
51.	Shri Dhan Lal	Shri Bachhu Lal	Vill-Gainthana
51.	SIIII DIIAII LAI	SIIII Daciiliu Lai	
	1		Block & DisttJakholi, Rudraprayag
			Contact No 7618465372

			Block & DisttJakholi, Rudraprayag
53.	Smt. Manju Devi	Shri Kutma	Vill-Gainthana
			Block & DisttJakholi, Rudraprayag
			Contact No 7830180194
54.	Smt. Jas Devi	Shri Guddu Lal	Vill-Gainthana
			Block & DisttJakholi, Rudraprayag
			Contact No 7830180194
55.	Dr.V.K. Purohit	Late Shri A.P. Purohit	HAPPRC, Contact No 9456531715
56.	Shri Kailash Kandpal	Shri B. P. Kandpal	HAPPRC, Contact No 7302808941
57.	Shri Jaidev Chauhan	Shri B. S. Chauhan	HAPPRC, Contact No 8126211560
58.	Shri Vipin Rawat	Shri P. S. Rawat	HAPPRC, Contact No 9458113893
59.	Shri Kamal Pundir	Shri G. S. Pundir	HAPPRC, Contact No 9540468782



Figure 4. Photographs of On-farm farmers workshop cum training/plant distribution programme organsied at Village Gainthana Block, Jakholi, Rudraprayag on 12/7/2022 under J.W.C.T. Project. Delivered technical knowledge by Dr. V.K. Purohit (SSO) to farmers/participants (A-B), plant material distributed to farmers (C-E), group photograph of farmers/participants (F).

Table 10. List of Villagers/Farmers participated in one day On-farm farmers workshop cum training/plant distribution programme for promotion of cultivation of MAP's at Village Kaviltha Block, Ukhimath, Rudraprayag on 19/July/2022 under J.W.C.T. Project.

S.N.	Villagers Name	Father/Husband Name	Address
1.	Shri Devendra Singh Negi	Shri Makar Singh Negi	Vill-Kaviltha Block &DisttUkhimath, Rudraprayag Contact No 9045183495
2.	Shri Kuldeep Singh	Shri Kunwar Singh	Vill-Kaviltha Block & DisttUkhimath, Rudraprayag Contact No9675852923
3.	Shri Arvind Singh	Shri Abbal Singh	Vill-Kaviltha Block &DisttUkhimath, Rudraprayag Contact No9720592827 Aadhar No663105982011
4.	Shri Digvijay Rawat	Shri Narendra Singh Rawat	Vill-Kaviltha

Total participants: 35 (male- 17, female-19)

			Block & DisttUkhimath, Rudraprayag
			Contact No8194095634
5.	Shri Balwant Singh	Shri Bagh Singh	Vill-Kaviltha
	Rawat	0 0	Block & DisttUkhimath, Rudraprayag
			Contact No9761868101
			Aadhar No234015000424
6.	Shri Mahendra Singh	Shri Kalam Singh Rawat	Vill-Kaviltha
	Rawat	~8	Block & DisttUkhimath, Rudraprayag
			Contact No8941912851
			Aadhar No498885695501
7.	Shri Pradeep Rawat	Shri Dev Singh Rawat	Vill-Kaviltha
7.	Shiri Fradcep Kawai	Shiri Dev Shigh Rawat	Block & DisttUkhimath, Rudraprayag
			Contact No7895728859
8.	Smt. Bhama Devi	Shri Rajesh Gaur	Vill-Kaviltha
0.	Sint. Bhana Devi	Shiri Kajesh Gaul	
			Block & DisttUkhimath, Rudraprayag
-			Contact No8393027651
9.	Smt. Sangeeta Devi	Shri Chandramohan Bhatt	Vill-Kaviltha
			Block & DisttUkhimath, Rudraprayag
			Contact No 7037391051
10.	Smt. Puja Devi	Shri Dinesh Chandra Gaur	Vill-Kaviltha
			Block & DisttUkhimath, Rudraprayag
			Contact No 9720026627
11.	Smt. Sulochana Devi	Shri Jaykrishna Chamola	Vill-Kaviltha
			Block & DisttUkhimath, Rudraprayag
			Contact No8979039669
12.	Smt. Beena Devi	Lt. Shri Purosottam Bhatt	Vill-Kaviltha
12.	Sint. Deena Devi		Block & DisttUkhimath, Rudraprayag
13.	Smt. Sudha	Shri Kunwar Singh Rawat	Vill-Kaviltha
15.	Sint. Sudna	Shiri Kuliwai Shigii Kawat	Block & DisttUkhimath, Rudraprayag
			Contact No 6978510432
1.4	Crut Inst' Dani	Shai De en els Cassa	
14.	Smt. Jyoti Devi	Shri Deepak Gaur	Vill-Kaviltha
			Block & DisttUkhimath, Rudraprayag
			Contact No 9897813793
15.	Smt. Asha Devi	Shri Anil Singh Rana	Vill-Kaviltha
			Block & DisttUkhimath, Rudraprayag
			Contact No8476833017
16.	Smt. Priyanka Gaur	Shri Subhash Gaur	Vill-Kaviltha
			Block & DisttUkhimath, Rudraprayag
			Contact No 8979399748
17.	Smt. Shanta Devi	Shri Bhagat Singh	Vill-Kaviltha
			Block & DisttUkhimath, Rudraprayag
			Contact No 9758268906
18.	Smt. Shashi Devi	Shri Sarvesh Singh Rawat	Vill-Kaviltha
10.		Sint Sur (Shi Shigh Ruwat	Block & DisttUkhimath, Rudraprayag
			Contact No 8588929568
19.	Smt. Sarita Devi	Shri Kalam Singh	Vill-Kaviltha
19.	Sint. Salita Devi	Shiri Kalalli Shigli	
			Block & DisttUkhimath, Rudraprayag
20	$\Omega_{1} = \frac{1}{2} N_{1} = 1$		Contact No 9639343056
20.	Shri Yogendra Singh	Lt. Shri Balak Singh Rawat	Vill-Kaviltha
			Block & DisttUkhimath, Rudraprayag
			Contact No 7466875457
21.	Smt. Chaita Devi	Late Shri Shambhu Prasad	Vill-Kaviltha
			Block & DisttUkhimath, Rudraprayag
			Contact No 7895670295
22.	Smt. Sushila Devi	Shri Pradeep Chauhan	Vill-Kaviltha
-		L	Block & DisttUkhimath, Rudraprayag
			Contact No 7451983629
23.	Shri Sudanand	Lt. Shri Purosotatam	Vill-Kaviltha
23.	Sini Suuananu		Block & DisttUkhimath, Rudraprayag
24	Chai Abbal Charl	I t Chai Charana Charan	Contact No 8171467246
24.	Shri Abbal Singh	Lt. Shri Shyam Singh	Vill-Kaviltha
			Block & DisttUkhimath, Rudraprayag

			Contact No 8958239777
25.	Shri Sanjay Singh	Lt. Uday Singh	Vill-Kaviltha
			Block &DisttUkhimath, Rudraprayag Contact No 9627004558
26.	Smt. Roshani Devi	Shri Diwakar Gairola	Vill-Kaviltha
20.	Sint. Koshani Devi	Sin Diwakar Ganola	Block & DisttUkhimath, Rudraprayag
			Contact No 9759723629
27.	Smt. Mukhari Devi	Shri Prabal Singh	Vill-Kaviltha
		C	Block & DisttUkhimath, Rudraprayag
			Contact No 9634346751
28.	Shri Dinesh Rawat	Shri Dev Singh Rawat	Vill-Kaviltha
			Block & DisttUkhimath, Rudraprayag
			Contact No 7895037902
29.	Smt. Vijaya Devi	Lt. Shri Govind Singh	Vill-Kaviltha
		Rawat	Block & DisttUkhimath, Rudraprayag
20			Contact No 7037402781
30.	Smt. Anita Devi	Shri Abbal Singh	Vill-Kaviltha
			Block & DisttUkhimath, Rudraprayag
31.	Smt. Sarla Devi	Shui Kummun Singh Domot	Contact No9719568284 Vill-Kaviltha
51.	Smt. Saria Devi	Shri Kunwar Singh Rawat	Block & DisttUkhimath, Rudraprayag
			Contact No 8057139089
32.	Shri Jaidev Chauhan	Shri B.S.Chauhan	HAPPRC, Contact No8126211560
33.	Shri Kailash Kandpal	Shri Bhagwati Prasad	HAPPRC, Contact No 7302808941
34.	Shri Vipin Rawat	Shri P.S.Rawat	HAPPRC, Contact No 9458113893
35.	Shri Ajay Hemdan	Shri Ajay Hemdan	HAPPRC, Contact No 8272820207
36.	Shri Mukesh	Shri U.S. Karasi	HAPPRC, Contact No 9675418245



Figure 5. Photographs of the one day On-farm farmers workshop cum training/plant distribution programme for promotion of cultivation of MAP's at Village Kaviltha Block, Ukhimath, Rudraprayag on 19/July/2022 under J.W.C.T. Project. Registration of participants (A), diffusion pf technical knowledge to farmers (B-C), chief guests address to the farmers (D), distribution of *P. Kurrooa* seedlings plants to farmers (E) and group photo of participants (F).

Table 11. List of villagers/farmers participated in one day On-farm farmers workshop cum training/plant distribution programme for promotion of cultivation of MAP's at Village Jaal Malla Block, Ukhimath, Rudraprayag on 19/7/2022 under J.W.C.T. Project.

Sr.No.	Villagers Name	Father/Husband Name	Address
1.	Shri Chandra	Lt. Shri Balak Singh Rawa	Vill-Jaal Malla
	Singh Rawat		Block & DisttUkhimath, Rudraprayag
			Contact No 7217433742
			Aadhar No 304548552222
2.	Shri Trilok Singh	Shri Baiker Singh	Vill-Jaal Malla
	C	C	Block & DisttUkhimath, Rudraprayag
			Contact No 8006346583
			Aadhar No690891288502
3.	Shri Madan Singh	Lt. Shri Gokal Singh	Vill-Jaal Malla
0.	Rawat		Block & DisttUkhimath, Rudraprayag
			Contact No7830966432
4.	Shri Mangal Singh	Shri Ghanshyam Singh	Vill-Jaal Malla
	Shiri Mungui Shigh	Shiri Shunshyuni Shigh	Block & DisttUkhimath, Rudraprayag
			Contact No7302161380
5.	Shri Madhavar	Lt. Shri Dhoom Singh	Vill-Jaal Malla
5.	Singh	Lt. Shiri Dhoom Shigh	Block & DisttUkhimath, Rudraprayag
	Singn		
6.	Shri Dinach Sinat	Shri Changyam Singh	Contact No8650805288 Vill-Jaal Malla
0.	Shri Dinesh Singh	Shri Ghansyam Singh	
			Block & DisttUkhimath, Rudraprayag
			Contact No7060165953
7			Aadhar No369431198374
7.	Shri Vipin Singh	Shri Lakhan Singh Rawat	Vill-Jaal Malla
	Rawat		Block & DisttUkhimath, Rudraprayag
			Contact No7043449663
			Aadhar No573410422324
8.	Shri Mohan Singh	Lt. Shri Bhopal Singh	Vill-Jaal Malla
	Rawat		Block & DisttUkhimath, Rudraprayag
			Contact No7895399593
9.	Shri Daulat Singh	Lt. Shri Gokal Singh	Vill-Jaal Malla
	Rawat	Rawat	Block & DisttUkhimath, Rudraprayag
10.	Smt. Mangsiri	Lt. Shri Thepad Singh	Vill-Jaal Malla
	Devi	Rawat	Block & DisttUkhimath, Rudraprayag
			Contact No 7895026624
11.	Smt. Anita Rawat	Shri Rajendra Singh Rawat	Vill-Jaal Malla
			Block & DisttUkhimath, Rudraprayag
			Contact No8755725353
12.	Smt. Gaytri Devi	Lt. ShriJaman Singh Rawat	Vill-Jaal Malla
	~		Block & DisttUkhimath, Rudraprayag
13.	Km. Bhavna	Shri Shivraj Singh Panwar	Vill-Jaal Malla
	Panwar		Block & DisttUkhimath, Rudraprayag
			Contact No 7310888055
14.	Smt. Anusuya	Shri Vikram Singh	Vill-Jaal Malla
17.	Devi		Block & DisttUkhimath, Rudraprayag
			Contact No 8449986190
15.	Smt. Laxmi Devi	Shri Trilak Singh Dowot	Vill-Jaal Malla
13.	Sint. Laxini Devi	Shri Trilok Singh Rawat	
			Block & DisttUkhimath, Rudraprayag
			Contact No8006346583
17			Aadhar No 854811279271
16.	Smt. Asha Devi	Shri Dinesh Singh	Vill-Jaal Malla
			Block & DisttUkhimath, Rudraprayag
			Contact No 7302161373
17.	Smt. Pushpa Devi	Shri Pramod Singh	Vill-Jaal Malla
	_	-	Block & DisttUkhimath, Rudraprayag
			Contact No 8791837310
18.	Smt. Anita Devi	Shri Jaspal Singh	Vill-Jaal Malla
		i C	Block & DisttUkhimath, Rudraprayag

19.	Smt. Deepa Devi	Shri Arvind Singh Panwar	Vill-Jaal Malla
	1	8	Block & DisttUkhimath, Rudraprayag
			Contact No 8755587122
20.	Smt. Sunita Devi	Shri Sate Singh	Vill-Jaal Malla
		ç	Block & DisttUkhimath, Rudraprayag
			Contact No 7302163977
21.	Smt. Prema Devi	ShriBhagat Singh Panwar	Vill-Jaal Malla
			Block & DisttUkhimath, Rudraprayag
			Contact No9557443334
22.	Shri Virendra	Lt. Shri Jeth Singh	Vill-Jaal Malla
	Singh Panwar		Block & DisttUkhimath, Rudraprayag
			Contact No8006491086
23.	Smt. Hema Devi	Shri Shivraj Singh Panwar	Vill-Jaal Malla
			Block & DisttUkhimath, Rudraprayag
			Contact No7310888055
24.	Shri Shivraj Singh	Lt. Shri Jagat Singh	Vill-Jaal Malla
		Panwar	Block & DisttUkhimath, Rudraprayag
			Contact No 7310888055
25.	Smt. Kushma Devi	Shri Bheem Singh Panwar	Vill-Jaal Malla
			Block & DisttUkhimath, Rudraprayag
26.	Shri Surendra	Shri Daulat Singh Rawat	Vill-Jaal Malla
	Singh	<u> </u>	Block & DisttUkhimath, Rudraprayag
27.	Smt. Vinita Devi	Shri Umed Singh	Vill-Jaal Malla
20			Block & DisttUkhimath, Rudraprayag
28.	Smt. Vimla Devi	Shri Dev Singh	Vill-Jaal Malla
20			Block & DisttUkhimath, Rudraprayag
29.	Smt. Ranju Devi	Lt. Shri Dhan Singh Rawat	
20		01 : 0 : 0: 1	Block & DisttUkhimath, Rudraprayag
30.	Smt. Kunwari	Shri Suraj Singh	Vill-Jaal Malla
21	Devi	<u> </u>	Block & DisttUkhimath, Rudraprayag
31.	Shri Puran Singh	Shri Ram Singh	Vill-Jaal Malla
	Rana		Block & DisttUkhimath, Rudraprayag
32.	Shri Jaidev	Shri B.S.Chauhan	Contact No 7895026803 HAPPRC, Contact No8126211560
32.	Chauhan	SIIII B.S.Chaunan	ПАГГКС, Contact No8120211500
33.	Shri Kailash	Shri Dhagwati Dragg d	HADDDC Contract No. 7202808041
35.		Shri Bhagwati Prasad	HAPPRC, Contact No 7302808941
34.	Kandpal Shri Vipin Rawat	Shri P.S.Rawat	HAPPRC, Contact No 9458113893
35.	Shri Ajay Hemdan	Shri Ashok Kumar	HAPPRC, Contact No 8272820207
55.	Sini Ajay menidan	Hemdan	11A1 I NU, UUITAU 110 02/2020/
36.	Shri Mukesh	Shri U.S. Karasi	HAPPRC



Figure 6. Photographs of the one day On-farm farmers workshop cum training/plant distribution programme for promotion of cultivation of MAP's at Jaal Malla, Block, Ukhimath, Rudraprayag on 19/7/2022 under J.W.C.T. Project. Registration of participants (A), diffusion of technical knowledge to farmers (B), distribution of *P. Kurrooa* plants to farmers(C-E), and group photo of participants (F).

Table 13. List of Villagers/Farmers participated in one day On-farm farmers workshop cum
training/plant distribution programme for promotion of cultivation of MAP's at Village Jaal Talla
Block, Ukhimath, Rudraprayag on 19/7/2022 under J.W.C.T. Project.Total participants:41 (male- 13, female-28)

Sr.No.	Villagers Name	Father/Husband Name	Address
1.	Shri Pradeep Rana	Shri Chandra Singh	Vill-Jaal Talla
1.	~r	2	Block & DisttUkhimath, Rudraprayag
			Contact No 8755932742
			Aadhar No 633462305263
2.	Shri Hukum Singh	Lt. Shri Narayan Singh	Vill-Jaal Talla
۷.	Shiri Hukuni Shigh	Lt. Shiri Narayan Shigh	
2			Block & DisttUkhimath, Rudraprayag
3.	Smt. Kusum Devi	Shri Pradeep Singh Rana	Vill-Jaal Talla
			Block & DisttUkhimath, Rudraprayag
			Contact No7895089752
4.	Smt. Anita Devi	Shri Pramod Singh Rawat	Vill-Jaal Talla
			Block & DisttUkhimath, Rudraprayag
			Contact No7302236522
5.	Smt. Ritu Devi	Shri Sandeep Singh	Vill-Jaal Talla
		1 8	Block & DisttUkhimath, Rudraprayag
			Contact No9997821325
6.	Smt. Sateshwari Devi	Shri Ranjit Singh	Vill-Jaal Talla
0.	Sint. Satesiiwali Devi	Shiri Kanjit Singh	
			Block & DisttUkhimath, Rudraprayag
-			Contact No8791803662
7.	Smt. Indra Devi	Shri Suraj Singh	Vill-Jaal Talla
			Block & DisttUkhimath, Rudraprayag
			Contact No8126387274
8.	Smt. Deepa Devi	Shri Tejpal Singh	Vill-Jaal Talla
	-		Block & DisttUkhimath, Rudraprayag
			Contact No8979485245
9.	Smt. Vijaya Devi	Shri Prakash Singh	Vill-Jaal Talla
	Since + ijuju 20+1	Shiri i ranash Shigh	Block & DisttUkhimath, Rudraprayag
			Contact No 8273527695
10.	Smt. Surji Devi	Shri Virendra Singh	Vill-Jaal Talla
10.	Sint. Surji Devi	Shiri vitendra Singh	
			Block & DisttUkhimath, Rudraprayag
			Contact No 8532880434
11.	Smt. Pushpa Devi	Shri Bhagat Singh	Vill-Jaal Talla
			Block & DisttUkhimath, Rudraprayag
			Contact No8755873318
12.	Smt. Mitla Devi	Shri Rakesh	Vill-Jaal Talla
			Block & DisttUkhimath, Rudraprayag
			Contact No 9119041988
13.	Smt. Vimla Devi	Shri Virendra	Vill-Jaal Talla
15.		Sini (nondru	Block & DisttUkhimath, Rudraprayag
1 4	Quet Dama'l D		Contact No 8923347857
14.	Smt. Permila Devi	Shri Suresh Singh	Vill-Jaal Talla
			Block & DisttUkhimath, Rudraprayag
			Contact No 8755644676
15.	Smt. Vinita Devi	Shri Prakash Singh	Vill-Jaal Talla
		~	Block & DisttUkhimath, Rudraprayag
			Contact No8954612563
16.	Km. Monika	Shri Prem Singh	Vill-Jaal Talla
		ongn	Block & DisttUkhimath, Rudraprayag
17	Cast Vanla D	Chai Damar Charle	Contact No 9997373876
17.	Smt. Kamla Devi	Shri Puran Singh	Vill-Jaal Talla
			Block & DisttUkhimath, Rudraprayag
			Contact No 8532801988
18.	Smt. Hema Devi	Shri Kunwar Singh	Vill-Jaal Talla
		_	Block & DisttUkhimath, Rudraprayag
			Contact No9634783051
	Smt. Maheshi Devi	Shri Prabal Singh	Vill-Jaal Talla

			Block & DisttUkhimath, Rudraprayag
20	Smt. Uma Devi	Shri Suroi Sinah	Contact No 7060502073 Vill-Jaal Talla
20.	Smt. Uma Devi	Shri Suraj Singh	
			Block & DisttUkhimath, Rudraprayag
21	Smt. Guddi Devi	ShriDinaah Sinah	Contact No 8923346387
21.	Smt. Guddi Devi	ShriDinesh Singh	Vill-Jaal Talla
			Block & DisttUkhimath, Rudraprayag
22	Cust Use also Deed		Contact No8755615887
22.	Smt. Umesha Devi	Shri Keshar Singh	Vill-Jaal Talla
			Block & DisttUkhimath, Rudraprayag
22	Cast Dais Dani		Contact No9759485205
23.	Smt. Puja Devi	Shri Sunil	Vill-Jaal Talla
			Block & DisttUkhimath, Rudraprayag
24	Cast Issails Dara'	ChaiDealat Circel	Contact No8393027440
24.	Smt. Jasoda Devi	ShriDaulat Singh	Vill-Jaal Talla
			Block & DisttUkhimath, Rudraprayag
25	Curt Americ Duri	L t. Chui Malagui Cingh	Contact No 8650761282
25.	Smt. Amra Devi	Lt. Shri Makani Singh	Vill-Jaal Talla
			Block & DisttUkhimath, Rudraprayag
26	Smt Dears Dear	Shui Duong Sin ah	Contact No 8393895891
26.	Smt. Beena Devi	Shri Prem Singh	Vill-Jaal Talla Plaak & Diett Ukbimath Budraproved
			Block & DisttUkhimath, Rudraprayag
77	Smt Mulikan' Deed	Shri Dojor 470	Contact No 8954022101
27.	Smt. Mukhari Devi	Shri Rajendra	Vill-Jaal Talla Plack & Diett, Ulthimath, Dudramawaa
			Block & DisttUkhimath, Rudraprayag
20	V O'	<u> </u>	Contact No 8393895891
28.	Km. Siya	Shri Shiv Singh Siya	Vill-Jaal Talla
			Block & DisttUkhimath, Rudraprayag
20	0		Contact No 9897867708
29.	Smt. Amra	Shri Mangal Singh	Vill-Jaal Talla
			Block & DisttUkhimath, Rudraprayag
20			Contact No7830265350
30.	Smt. Vimla Devi	Shri Ramchandra Singh	Vill-Jaal Talla
			Block & DisttUkhimath, Rudraprayag
2.1			Contact No 8393055336
31.	Shri Amit Negi	Shri O.P.Negi	Vill-Jaal Talla
			Block & DisttUkhimath, Rudraprayag
22			Contact No 8057890309
32.	Shri Anoop	Shri Abbal Singh	Vill-Jaal Talla
			Block & DisttUkhimath, Rudraprayag
22			Contact No7078508293
33.	Shri Sandeep Singh	Lt. Shri Jaspal Singh	Vill-Jaal Talla
			Block & DisttUkhimath, Rudraprayag
<u>.</u>			Contact No 7060549815
34.	Shri Pramod Singh	Lt. Shri Shivraj Singh	Vill-Jaal Talla
			Block & DisttUkhimath, Rudraprayag
			Contact No 9068831984
35.	Shri Dharmendra	Shri Hayat Singh	Vill-Jaal Talla
	Singh		Block & DisttUkhimath, Rudraprayag
			Contact No 9634630265
36.	Shri Surat Singh	Shri Jauna Singh	Vill-Jaal Talla
			Block & DisttUkhimath, Rudraprayag
			Contact No 8923346387
37.	Shri Trilok Singh	Shri Gabbar Singh	Vill-Jaal Talla
	_	_	Block & DisttUkhimath, Rudraprayag
			Contact No8954950231
38.	Shri Jaidev Chauhan	Shri B.S.Chauhan	HAPPRC, Contact No 8126211560
39.	Shri Kailash Kandpal	Shri Bhagwati Prasad	HAPPRC, Contact No 7302808941
40.	Shri Vipin Rawat	Shri P.S.Rawat	HAPPRC, Contact No 9458113893
τυ.	Sini vipin Kawat		
41.	Shri Ajay Hemdan	Shri Ashok Kumar Hemdan	HAPPRC, Contact No8272820207



Figure 7. Photographs of the one day On-farm farmers workshop cum training/plant distribution programme for promotion of cultivation of MAP's at Jaal Talla, Block, Ukhimath, Rudraprayag on 19/7/2022 under J.W.C.T. Project. Delivered technical knowledge by Mr. Jaidev Chauhan to farmers/participants (A-B), distributed plant material to farmers (C-E), group photograph farmers/participants (F).

Table 14. List of Villagers/Farmers participated in one day On-farm farmers workshop cum					
training/plant distribution programme for promotion of cultivation of MAP's at Village Chaumasi					
Block, Ukhimath, Rudraprayag on 19/7/2022 under J.W.C.T. Project.					

Sr.No.	Villagers Name	Father/Husband Name	Address
1.	Shri Praveen	Shri Vikram Singh	Vill-Chaumasi
		_	Block & DisttUkhimath, Rudraprayag
			Contact No 7830543661
2.	Shri Rahul	Shri Bhagat Singh	Vill-Chaumasi
			Block & DisttUkhimath, Rudraprayag
			Contact No 7302175567
3.	Shri Vipin Singh	Shri Ranjeet Singh	Vill-Chaumasi
			Block & DisttUkhimath, Rudraprayag
			Contact No9997330540
4.	Shri Naveen	Shri Mahipal Singh	Vill-Chaumasi
			Block & DisttUkhimath, Rudraprayag
			Contact No8755153669
5.	Shri Mulayam Singh	Shri Suraj Singh	Vill-Chaumasi
			Block & DisttUkhimath, Rudraprayag
			Contact No8859125618
6.	Shri Badri	Shri Rajendra Singh	Vill-Chaumasi
			Block & DisttUkhimath, Rudraprayag
			Contact No7302228369
7.	Shri Mohan Singh	Lt. Shri Chandra Singh	Vill-Chaumasi
	Tindori		Block & DisttUkhimath, Rudraprayag
			Contact No9410145575
8.	Shri Jay Singh	Shri G.S.Rawat	Vill-Chaumasi
	Rawat		Block & DisttUkhimath, Rudraprayag
			Contact No9412949214
9.	Smt. Kamla Devi	Shri Gabbar Singh	Vill-Chaumasi
			Block & DisttUkhimath, Rudraprayag
			Contact No 7830261039
10.	Smt. Chhoti Devi	Lt. Shri Jeet Singh	Vill-Chaumasi

Total participants: 38 (male-18, female-20)

			Block & DisttUkhimath, Rudraprayag
			Aadhar No
11.	Smt. Katgi Devi	Shri Jeet Singh	Vill-Chaumasi Block &DisttUkhimath, Rudraprayag
12.	Smt. Prabha Devi	Shri Ravindra Singh	Vill-Chaumasi
12.	Sint. I fabila Devi	Shiri Kavindra Shigh	Block & DisttUkhimath, Rudraprayag
			Contact No 9557065755
13.	Smt. Uma Devi	Shri Akhilesh Singh	Vill-Chaumasi
		C C	Block & DisttUkhimath, Rudraprayag
			Contact No 8791104542
14.	Smt. Anita Devi	Lt. Shri Jaspal Singh	Vill-Chaumasi
			Block & DisttUkhimath, Rudraprayag
15.	Smt. Kamla Devi	Shri Vijay Singh	Vill-Chaumasi
			Block & DisttUkhimath, Rudraprayag
			Contact No8979052373
16.	Smt. Mamta Devi	Shri Pramod Singh	Vill-Chaumasi
			Block & DisttUkhimath, Rudraprayag
			Contact No 8755163645
17.	Smt. Anita Devi	Shri Rai Singh	Vill-Chaumasi
			Block & DisttUkhimath, Rudraprayag
			Contact No 9458945941
18.	Smt. Madhu Devi	Shri Sarvesh Singh	Vill-Chaumasi
			Block & DisttUkhimath, Rudraprayag
			Contact No8755847580
19.	Smt. Roshani Devi	Shri Yogember Singh	Vill-Chaumasi
			Block & DisttUkhimath, Rudraprayag
20			Contact No 8445974761
20.	Smt. Shanta Devi	Shri Katig Singh	Vill-Chaumasi
			Block & DisttUkhimath, Rudraprayag
21.	Smt. Soni Devi	ShriRajendra Singh	Vill-Chaumasi
	Smt. Gaini Devi		Block &DisttUkhimath, Rudraprayag
22.	Smt. Gaini Devi	Shri Gulab Singh	Vill-Chaumasi
			Block & DisttUkhimath, Rudraprayag
23.	Smt. Narvada Devi	Shri Rajpal Singh	Contact No8979835767 Vill-Chaumasi
23.	Sint. Ival valia Devi	Shiri Kajpai Shigh	Block & DisttUkhimath, Rudraprayag
			Contact No9068512373
24.	Shri Ajay Singh	Shri Virendra Singh	Vill-Chaumasi
27.	Shiri Ajay Shigh	Shiri virendra Shigh	Block & DisttUkhimath, Rudraprayag
			Contact No 8755847845
25.	Shri Anil	Shri Prabal Singh	Vill-Chaumasi
23.	Shiri 7 tini	Shiri Frabar Shirgh	Block & DisttUkhimath, Rudraprayag
			Contact No 8979823801
26.	Smt. Babita Devi	Shri Mulayam Singh	Vill-Chaumasi
		singh	Block & DisttUkhimath, Rudraprayag
			Contact No 8859125618
27.	Smt. Vijaya Devi	Lt. Shri Dayal Singh	Vill-Chaumasi
			Block & DisttUkhimath, Rudraprayag
28.	Smt. Shakha Devi	Shri Prabal Singh	Vill-Chaumasi
		gine	Block & DisttUkhimath, Rudraprayag
29.	Smt. Vimala Devi	Shri Mahesh Singh	Vill-Chaumasi
			Block & DisttUkhimath, Rudraprayag
30.	Smt. Laxmi Devi	Shri Devendra Singh	Vill-Chaumasi
			Block & DisttUkhimath, Rudraprayag
31.	Shri Gajpal Singh	Shri Dayal	Vill-Chaumasi
	Sint Suppar Singh		Block &DisttUkhimath, Rudraprayag
			Contact No 9084412374
32.	Shri Jaidev Chauhan	Shri B.S.Chauhan	HAPPRC
			Contact No8126211560
33.	Shri Kailash	Shri Bhagwati Prasad	HAPPRC
	Kandpal	0	Contact No 7302808941

34.	Shri Vipin Rawat	Shri P.S.Rawat	HAPPRC
	_		Contact No 9458113893
35.	Shri Ajay Hemdan	Shri Ashok Kumar Hemdan	HAPPRC
			Contact No 8272820207
36.	Shri Mukesh	Shri U.S. Karasi	HAPPRC
			Contact No



Figure 8. Photographs of the one day On-farm farmers workshop cum training/plant distribution programme for promotion of cultivation of MAP's organized at Chaumasi, Ukhimath, Rudraprayag on 19/7/2022 under J.W.C.T. Project. Distribution of plant material to farmers (A-B), group photograph of farmers/participants (C-D).

Table 15. List of Villagers/Farmers participated in one day On-farm farmers workshop cum					
training/plant distribution programme for promotion of cultivation of MAP's organized at at Village					
Kulpudi, Block, Tharali, Chamoli on 16/7/2022 under J.W.C.T. Project.					
Total participants: 21 (mala 16 famala 05)					

Sr.No.	Villagers Name	Father/Husband Name	Address
1.	Shri Jaiveer Singh	Shri Bhawan Singh	Vill- Kulpudi
		_	Block & DisttTharali, Chamoli
			Contact No 9119047850
			Aadhar No 720575991078
2.	Shri Gaur Singh	Shri Khilaph Singh	Vill- Kulpudi
			Block & DisttTharali, Chamoli
			Contact No 9568888047
			Aadhar No 765905017402
3.	Smt. Laxmi Devi	Shri Kalyan Singh	Vill- Kulpudi
			Block & DisttTharali, Chamoli
4.	Smt. Deepa Devi	Shri Dilwar Singh Negi	Vill- Kulpudi
			Block & DisttTharali, Chamoli
			Contact No7454879012
5.	Smt. Rekha Devi	Shri Amar Singh Negi	Vill- Kulpudi
			Block & DisttTharali, Chamoli
			Contact No9068514501
6.	Smt. Aruni Devi	Lt. Shri Ram Singh	Vill- Kulpudi
			Block & DisttTharali, Chamoli
7.	Smt. Laxmi Devi	Shri Gaur Singh Negi	Vill- Kulpudi
			Block & DisttTharali, Chamoli
			Contact No8755810416
8.	Shri Rajendra Singh	Shri Avtar Singh	Vill- Kulpudi

Total participants: 21 (male- 16, female-05)

			Block & DisttTharali, Chamoli Contact No7248475239
9.	Chui Mahinal Cinah	Shui Doulat Sin ah	
9.	Shri Mahipal Singh	Shri Daulat Singh	Vill- Kulpudi
			Block &DisttTharali, Chamoli
10		<u>01 : 11: (0: 1</u>	Contact No 9634453637
10.	Shri Kundan Singh	Shri Himmat Singh	Vill- Kulpudi
11			Block &DisttTharali, Chamoli
11.	Shri Paar Singh	Shri Baag Singh	Vill- Kulpudi
			Block &DisttTharali, Chamoli
10			Contact No8979748967
12.	Shri Pushkar Singh	Shri Paan Singh	Vill- Kulpudi
			Block &DisttTharali, Chamoli
			Contact No 8865040384
1.2			Aadhar No 260181124259
13.	Shri Parvendra Singh	Lt. Shri Digpal Singh Rawat	
	Rawat		Block & DisttTharali, Chamoli
			Contact No 7895789059
14.	Shri Laxman Singh	Shri Avtar Singh	Vill- Kulpudi
			Block & DisttTharali, Chamoli
			Contact No 9755076466
			Aadhar No 473694514402
15.	Shri Subhash Chandra	Shri Avtar Ram	Vill- Kulpudi
			Block & DisttTharali, Chamoli
			Aadhar No 467790326187
16.	Dr.Vijay Kant Purohit	Shri A.P.Purohit	HAPPRC
			Contact No 94565317115
17.	Shri Mahaveer Singh	Shri Raghuveer Singh	HAPPRC
	Rawat		
18.	Shri Ajay Hemdan	Shri A.K.Hemdan	HAPPRC
			Contact No 8272820207
19.	Shri Bhawani Dutt	Lt. Shri Bhairav Dutt Kotha	HAPPRC
	Kothari		Contact No 8979525629
20.	Shri Kamal Singh	Shri G. S. Pundir	HAPPRC
	Pundir		Contact No 9540468782
21.	Shri Arun Singh	Shri Kalam Singh Gusain	HAPPRC
	Gusain	č	Contact No9548194033

Table 16. List of villagers/farmers participated in one day On-farm farmers workshop cum training/plant distribution programme for promotion of cultivation of MAP's Cultivation organised at Village Ratgaov, Taalger Block, Tharali, Chamoli on 22/7/2022 under J.W.C.T. Project.

Total participants: 18 (male- 18, female-0)

Sr.No.	Villagers Name	Father/Husband Name	Address
1.	Shri Gajendra Prasasd	Shri Khilaph Ram	Vill-Ratgaov Taalger
			Block & Distt Tharali, Chamoli
			Contact No 8859673676
2.	Shri Bhawani Dutt	Shri Bhairav Dutt Kothari	Vill- Ratgaon Taalger
	Kothari		Block & DisttTharali, Chamoli
3.	Shri Arun Singh	Shri Kamal Singh Gusain	Vill- Ratgaon Taalger
	Gusain		Block & DisttTharali, Chamoli
			Contact No 9548194033
4.	Shri Prithvi Singh	Shri Kedar Singh Pharshwar	Vill- Ratgaon Taalger
			Block & DisttTharali, Chamoli
			Contact No8449855389
			Aadhar No668123293888
5.	Shri Balwant Ram	Shri Kalam Ram	Vill- Ratgaon Taalger
			Block & DisttTharali, Chamoli
6.	Shri Rahul Singh	Shri Devendra Singh	Vill- Ratgaon Taalger
			Block & DisttTharali, Chamoli
			Contact No 7895322396

7.	Shri Avtar Singh Rawat	Shri Kanak Singh	Vill- Ratgaon Taalger Block & DisttTharali, Chamoli
8.	Shri Manohar Singh	Shri Mohan Singh	Vill- Ratgaon Taalger Block &DisttTharali, Chamoli Contact No8630154715 Aadhar No443445748901
9.	Shri Chandramohan Mishra	Shri Dayakrishna Mishra	Vill- Ratgaon Taalger Block &DisttTharali, Chamoli Contact No 7335051277 Aadhar No518282191269
10.	Shri Mahaveer Singh	Lt. Shri Chandra Singh	Vill- Ratgaon Taalger Block &DisttTharali, Chamoli Contact No 9639402314 Aadhar No 872769453018
11.	Shri Narendra Singh	Lt. Shri Darban Singh	Vill- Ratgaon Taalger Block & DisttTharali, Chamoli Contact No8938907106
12.	Shri Pushkar Singh	Lt. Shri Narayan Singh	Vill- Ratgaon Taalger Block &DisttTharali, Chamoli Contact No 9084911891
13.	Shri Madan Singh	Shri Gopal Singh	Vill- Ratgaon Taalger Block & DisttTharali, Chamoli Contact No 9927609759
14.	Shri Jaiveer Singh	Shri Balwant Singh	Vill- Ratgaon Taalger Block &DisttTharali, Chamoli Contact No 9520245030
15.	Shri Mahaveer Singh Rawat	Lt. Shri Kishan Singh	Vill- Ratgaon Taalger Block &DisttTharali, Chamoli Contact No8979815419 Aadhar No 467790326187
16.	Dr.Vijay Kant Purohit	Shri A.P.Purohit	HAPPRC SRINAGAR GARHWAL Contact No 9456531715
17.	Shri Kamal Pundir	Shri G.S.Pundir	HAPPRC SRINAGAR GARHWAL Contact No9540468782
18.	Shri Ajay Hemdan	Shri A.K.Hemdan	HAPPRC SRINAGAR GARHWAL Contact No8272820207

Table 17. List of villagers/farmers participated in one day On-farm farmers workshop cum training/plant distribution programme for promotion of cultivation of MAP's organsied at Village Rushyan Block, Tharali, Chamoli on 22/7/2022 under J.W.C.T. Project.

Sr.No.	Villagers Name	Father/Husband Name	Address
1.	Shri Bharat Singh	Shri Narayan Singh	Vill-Rusiyan
			Block & Distt Tharali, Chamoli
			Contact No 7060449964
2.	Shri Kuldeep Singh	Shri Raghuveer Singh Bisht	Vill- Rusiyan
			Block & Distt Tharali, Chamoli
			Contact No 9718591046
3.	Shri Mohan Singh	Shri Balwant Singh	Vill-Rusiyan
	Rawat		Block & Distt Tharali, Chamoli
			Contact No 7500677806
4.	Shri Digamber Singh	Shri Ishwari Dutt	Vill-Rusiyan
			Block & Distt Tharali, Chamoli
			Contact No7060081661
			Aadhar No 777651358872
5.	Shri Manoj Singh	Shri Mahipal Singh	Vill-Rusiyan
			Block & Distt Tharali, Chamoli
6.	Shri Sudarshan	Shri Kedar Singh	Vill-Rusiyan
	Singh		Block & Distt Tharali, Chamoli
			Contact No 7456966732

Total participants: 13 (male- 13, female-0)

7.	Shri Duli Ram	Shri Trimanu Ram	Vill-Rusiyan Block &DisttTharali, Chamoli
8.	Shri Mohan Singh Rawat	Shri Balwant Singh	Vill-Rusiyan Block &DisttTharali, Chamoli Contact No 7500677806
9.	Shri Bharat Singh	Shri Narayan Singh	Vill-Rusiyan Block &DisttTharali, Chamoli Contact No7080449964
10.	Shri Manoj Rana	Shri Mahipal Singh	Vill-Rusiyan Block &DisttTharali, Chamoli Contact No 9953271800
11.	Dr.Vijay Kant Purohit	Shri A.P.Purohit	HAPPRC Contact No 94565317115
12.	Shri Ajay Hemdan	Shri A.K.Hemdan	HAPPRC Contact No 8272820207
13.	Shri Kamal Singh Pundir	Shri G. S. Pundir	HAPPRC Contact No 9540468782

Table 18. List of villagers/farmers participated in one day On-farm farmers workshop cum training/plant distribution programme for promotion of cultivation of MAP's organized at Village Taal, block, Tharali, Chamoli on 23/7/2022 under J.W.C.T. Project. Total participants: 28 (male- 20, female-08)

S.N.	Villagers Name	Father/Husband Name	Address
1.	Smt. Munni Devi	Shri Balwant Singh Gariya	Vill- Taal Block & Distt Tharali, Chamoli Contact No 9568662231
			Aadhar No 878076804085
2.	Smt. Babli Devi	Shri Anil Singh Gariya	Vill- Taal Block &DisttTharali, Chamoli Contact No 9971353648
3.	Smt. Renu Devi	Shri Devendra Singh Gariya	
4.	Smt. Rajeshwari Devi	Shri Mahaveer Singh	Vill- Taal Block &DisttTharali, Chamoli Contact No9520235664
5.	Smt. Kalpeshwari Devi	Shri Darshan Singh Gariya	Vill- Taal Block &DisttTharali, Chamoli Contact No8445613358
6.	Shri Prem Singh Gariya	Shri Balwant Singh Gariya	Vill- Taal Block &DisttTharali, Chamoli Contact No 9012292138
7.	Shri Khushal Singh Gariya	Shri Balwant Singh Gariya	Vill- Taal Block &DisttTharali, Chamoli Contact No8193056373
8.	Shri Anil Singh	Shri Bhupal Singh Gariya	Vill- Taal Block &DisttTharali, Chamoli Contact No9568340283 Aadhar No 918967545019
9.	Shri Indra Singh Gariya	Lt. Shri Kedar Singh	Vill- Taal Block &DisttTharali, Chamoli Contact No 9568639595 Aadhar No 811537110324
10.	Shri Devu Lal	Lt. Shri Chotanu Lal	Vill- Taal Block &DisttTharali, Chamoli Contact No 7055564762 Aadhar No 619440175381
11.	Shri Raghuveer Singh	Shri Darvaan Singh	Vill- Taal Block &DisttTharali, Chamoli Contact No 9997339615

12.	Shri Ranjeet Lal	Shri Hayat Lal	Vill- Taal Block &DisttTharali, Chamoli Contact No 9997348799 Aadhar No 404955092168
13.	Shri Ratan Singh	Lt. Shri Narayan Singh	Vill- Taal Block &DisttTharali, Chamoli Contact No 7078425063 Aadhar No 823859962940
14.	Shri Ganesh Lal	Shri Shyam Lal	Vill- Taal Block &DisttTharali, Chamoli Contact No 8958720581 Aadhar No 551239041503
15.	Shri Mohan Singh	Shri Narayan Singh	Vill- Taal Block &DisttTharali, Chamoli Aadhar No 652902849566
16.	Shri Govind Lal	Shri Shankar Lal	Vill- Taal Block &DisttTharali, Chamoli Contact No 8445613230 Aadhar No 596771896121
17.	Smt. Beena Pandey	Shri Katika Prasad Pandey	Vill- Taal Block &DisttTharali, Chamoli Contact No 7037693744
18.	Shri M.L.Dhuniyal	Shri J.L.Dhuniyal	Vill- Taal Block &DisttTharali, Chamoli Contact No 97609633397
19.	Smt. Urmila Rawat	Shri Shiv Singh	Vill- Taal Block &DisttTharali, Chamoli Contact No 8057220898
20.	Smt. Baleshwari Devi	Shri Himat Singh Gariya	Vill- Taal Block &DisttTharali, Chamoli Contact No 7500231750
21.	Shri Balwant Singh	Shri Netra Singh	Vill- Taal Block &DisttTharali, Chamoli Contact No 8755618748 Aadhar No 813348420521
22.	Shri Maheshanand Pandey	Lt. Shri Ratnamani Pandey	Vill- Taal Block &DisttTharali, Chamoli Contact No 9760289538 Aadhar No 954767722001
23.	Shri Pradhuman Singh Gariya	Shri Gopal Singh	Vill- Taal Block &DisttTharali, Chamoli Contact No 9897652396 Aadhar No 980751802318
24.	Shri Anil Agri	Shri Kayar Agri	Vill- Taal Block &DisttTharali, Chamoli Contact No 7457818032 Aadhar No 418282538036
25.	Shri R.P.Joshi	Lt. Shri Dharmdutt Joshi	Vill- Taal Block &DisttTharali, Chamoli Contact No 7055859304 Aadhar No 839067955697
26.	Dr.Vijay Kant Purohit	Shri A.P.Purohit	HAPPRC Contact No 94565317115
27.	Shri Ajay Hemdan	Shri A.K.Hemdan	HAPPRC Contact No 8272820207
28.	Shri Kamal Singh Pundir	Shri G. S. Pundir	HAPPRC Contact No 9540468782



Figure 9. Photographs of the one day On-farm farmers workshop cum training/plant distribution programme for promotion of cultivation of MAP's organized at village Taal, Tharali, Chamoli on 23/07/2022. Delivered technical knowledge by Dr. V.K. Purohit (SSO) to farmers/participants (A-C), distributed plant material to farmers (D-E), group photograph of farmers/participants (F).

Table 19. List of villagers/farmers participated in one day On-farm farmers workshop cum				
training/plant distribution programme for promotion of cultivation of MAP's organized at Village				
Syanri Bangali, Block, Ghat, Chamoli on 14/7/2022 under J.W.C.T. Project.				

Sr.No.	Villagers Name	Father/Husband Name	Address
1.	Shri Bharat Singh	Shri Pushkar Singh	Vill-Syanri Bangali
	-	-	Block-Ghat
			DisttChamoli
			Contact No 7351937845
			Aadhar No 210836256127
2.	Shri Bharat Singh	Shri Ganga Singh	Vill-Syanri Bangali
			Block-Ghat
			DisttChamoli
			Contact No 9837436543
			Aadhar No925456340322
3.	Shri Dinesh Singh	Shri Rajendra Singh	Vill-Syanri Bangali
			Block-Ghat
			DisttChamoli
			Contact No 9837550384
			Aadhar No285678568481
4.	Shri Narendra Singh	Shri Gopal Singh	Vill-Syanri Bangali
			Block-Ghat
			DisttChamoli
			Contact No8449801975
			Aadhar No978851500093
5.	Shri Vikram Singh	Shri Puran Singh	Vill-Syanri Bangali
			Block-Ghat,
			DisttChamoli
			Contact No 7055947688
			Aadhar No 222061825187
6.	Shri Puran Singh	Shri Umed Singh	Vill-Syanri Bangali
	-	-	Block-Ghat,
			DisttChamoli
			Contact No 7351246174
			Aadhar No 626184504520

Total participants: 60 (male- 35, female-25)

7.	Shri Surendra Singh	Shri Karan Singh	Vill-Syanri Bangali
/.			Block-Ghat,
			DisttChamoli
			Contact No 7535820929
			Aadhar No370799476679
8.	Smt. Hema Devi	Shri Digpal Singh	Vill-Syanri Bangali
			Block-Ghat,
			DisttChamoli
			Contact No 9917992566
9.	Smt. Shanta Devi	Lt. Shri Manbar Singh	Vill-Syanri Bangali
			Block-Ghat,
			DisttChamoli
10			Contact No 9568054901
10.	Smt. Pushpa Devi	Shri Vijendra Singh	Vill-Syanri Bangali
			Block-Ghat, DisttChamoli
			Contact No 8958253295
11.	Smt. Kusum Negi	Shri K.S.Negi	Vill-Syanri Bangali
11.	Sint. Rusuin Negi	Shiri K.S.Negi	Block-Ghat,
			DisttChamoli
12.	Shri Darshan Singh	Shri Raghuveer Singh	Vill-Syanri Bangali
			Block-Ghat,
			DisttChamoli
			Contact No 9756258483
13.	Shri Mahipal Singh	Shri Bhajan Singh	Vill-Syanri Bangali
	1 0	5 6	Block-Ghat,
			DisttChamoli
			Contact No 9917696080
14.	Smt. Sarita Devi	Shri Bharat Singh	Vill-Syanri Bangali
			Block-Ghat,
			DisttChamoli
			Contact No8650705542
15.	Smt. Geeta Devi	Shri Ganga Singh	Vill-Syanri Bangali
			Block-Ghat,
			DisttChamoli
16	Smt. Chuchri Devi	Lata Shri Mahan Singh	Contact No 8958425426
16.	Sint. Chuchin Devi	Late Shri Mohan Singh	Vill-Syanri Bangali Block-Ghat,
			DisttChamoli
			Contact No 9639012295
17.	Smt. Parvati Devi	ShriVirendra Singh	Vill-Syanri Bangali
			Block-Ghat,
			DisttChamoli
			Contact No 9917991611
			Aadhar No 255269479523
18.	Smt. Basanti Devi	Shri Ranjeet Singh	Vill-Syanri Bangali
			Block-Ghat,
			DisttChamoli
			Contact No 7351263875
			Aadhar No 952413843915
19.	Smt. Geeta Devi	Shri Sujan Singh Bisht	Vill-Syanri Bangali
			Block-Ghat,
			DisttChamoli
			Contact No. -8954512094
20	Smt Aniali Dari	Shri Anond Sirch	Aadhar No433274646110
20.	Smt. Anjali Devi	Shri Anand Singh	Vill-Syanri Bangali Block-Ghat,
			Block-Gnat, DisttChamoli
			Contact No 7533967634
21.	Smt. Manisha devi	Shri Bhopal Singh	Vill-Syanri Bangali
<i>L</i> 1.			Block-Ghat,
			DisttChamoli
l			

			Contact No 7536860181
			Aadhar No347714931297
22.	Smt. Godambari Devi	Shri Surendra Singh	Vill-Syanri Bangali
			Block-Ghat,
			DisttChamoli
			Contact No 8449936523
23.	Smt. Prema Devi	Shri Madan Singh	Aadhar No 305362407525Vill-Syanri Bangali
23.	Sint. I Tenna Devi	Shiri Wadan Shigh	Block-Ghat,
			DisttChamoli
			Contact No 9756886373
24.	Smt. Bhaguli Devi	Shri Mahendra Singh	Vill-Syanri Bangali
			Block-Ghat,
			DisttChamoli Contact No 8981514121
25.	Shri Trilok Singh	Shri Chandra Singh	Vill-Syanri Bangali
23.	Shiri Tinok Shigh	Shiri Chandra Shigh	Block-Ghat,
			DisttChamoli
			Contact No8057543042
26.	Shri Deepak Negi	Shri Jay Singh	Vill-Syanri Bangali
			Block-Ghat,
			DisttChamoli
27.	Smt. Kamla Devi	Shri Vilok Singh Bisht	Contact No 8449971048 Vill-Syanri Bangali
27.	Sint. Kanna Devi	Shiri Vilok Shigh Bisht	Block-Ghat,
			DisttChamoli
			Contact No 9927572633
28.	Smt. Leela Devi	Shri Vilok Singh Negi	Vill-Syanri Bangali
			Block-Ghat,
			DisttChamoli
29.	Smt. Basanti Devi	Shri Dhunal Singh	Contact No8958018871
29.	Sint. Basanti Devi	Shri Bhupal Singh	Vill-Syanri Bangali Block-Ghat,
			DisttChamoli
			Contact No 7351959762
30.	Shri Chandramohan Bisht	Shri Vikram Singh	Vill-Syanri Bangali
			Block-Ghat,
			DisttChamoli
31.	Shri Shishupal Singh	Shri Sulabh Singh	Contact No 9756320022 Vill-Syanri Bangali
51.	Sill'i Shishupat Shigh	Shiri Sulaoli Shigh	Block-Ghat,
			DisttChamoli
32.	Shri Bhopal Singh	Shri Dilbar Singh	Vill-Syanri Bangali
			Block-Ghat,
			DisttChamoli
22	Shui Magandan Singh	Chui Anond Cinch	Contact No 8057887339
33.	Shri Narendra Singh	Shri Anand Singh	Vill-Syanri Bangali Block-Ghat,
			DisttChamoli
			Contact No 7465920280
34.	Shri Khushal Negi	Shri Bhupal Negi	Vill-Syanri Bangali
			Block-Ghat,
			DisttChamoli
			Contact No 7351279292
35.	Shri Deewon Sinch	Shri Kishan Singh Bisht	Aadhar No 368096331489
55.	Shri Deewan Singh	SIIIT AISHAII SIIIGH BISHI	Vill-Syanri Bangali Block-Ghat,
			DisttChamoli
			Contact No 8057063819
			Aadhar No 655234571967
36.	Smt. Parvati Bisht	Shri Virendra Singh	Vill-Syanri Bangali
			Block-Ghat,

			DisttChamoli
			Contact No 9917991611
37.	Shri Manoj Bisht	Shri Uday Singh	Vill-Syanri Bangali
			Block-Ghat,
			DisttChamoli
			Contact No 8449358686
38.	Shri Mahipal Negi	Shri Khilaf Singh	Vill-Syanri Bangali
			Block-Ghat,
			DisttChamoli Contact No7088681501
			Aadhar No 833503750036
39.	Shri Avtar Singh	Shri Hoyan Singh	Vill-Syanri Bangali
57.	Shiri Avtar Shigh	Shiri Hoyan Shigh	Block-Ghat,
			DisttChamoli
			Contact No. - 9690028938
			Aadhar No 503346143780
40.	Shri Trilok Singh	Shri Kanchan Singh	Vill-Syanri Bangali
			Block-Ghat,
			DisttChamoli
			Contact No 7533859264
41.	Shri Trilok Singh	Shri Gaur Singh	Vill-Syanri Bangali
			Block-Ghat,
			DisttChamoli
42.	Smt. Vimla Devi	Shri Virondro Naci	Contact No 8449607826
42.	Smt. Vimia Devi	Shri Virendra Negi	Vill-Syanri Bangali Block-Ghat,
			DisttChamoli
			Contact No 7252805607
43.	Shri Himmat Singh	Shri Rupchandra Singh	Vill-Syanri Bangali
		Sin rependent Singh	Block-Ghat,
			DisttChamoli
			Contact No 9690057249
44.	Shri Mohan Singh	Shri Dev Singh	Vill-Syanri Bangali
			Block-Ghat,
			DisttChamoli
45.	Smt. Shakuntala Devi	Shri Anand Singh	Vill-Syanri Bangali
			Block-Ghat,
			DisttChamoli Contact No 8755369106
46.	Smt. Deepa Devi	Shri Virendra Singh	Vill-Syanri Bangali
40.	Shit. Deepa Devi	Shiri vitendra Shigh	Block-Ghat,
			DisttChamoli
			Contact No 8192043924
47.	Smt. Sunita Devi	Shri Rajendra Singh	Vill-Syanri Bangali
		5 0	Block-Ghat,
			DisttChamoli
			Contact No 9527287630
48.	Smt. Surati Devi	Shri Prem Singh	Vill-Syanri Bangali
			Block-Ghat,
40			DisttChamoli
49.	Shri Lakshman Singh	Shri Kheem Singh	Vill-Syanri Bangali
			Block-Ghat,
			DisttChamoli Contract No. 8036001007
50.	Shri Gaur Singh	Shri Beer Singh	Contact No8936991997 Vill-Syanri Bangali
50.		Sini Deel Singli	Block-Ghat,
			DisttChamoli
			Contact No. -7467021996
51.	Smt. Deepa Negi	Shri Vijay Singh	Vill-Syanri Bangali
			Block-Ghat,
1			DisttChamoli

52.	Shri Sanjay Negi	Shri Gabbar Singh	Vill-Syanri Bangali
	July 1.6		Block-Ghat,
			DisttChamoli
			Contact No 9417663033
53.	Shri Mahipal Negi	Shri Khilap Singh	Vill-Syanri Bangali
			Block-Ghat,
			DisttChamoli
			Contact No 7088681501
54.	Shri Rajendra Singh	Shri Prem Singh	Vill-Syanri Bangali
			Block-Ghat,
			DisttChamoli
			Contact No 9690028937
55.	Smt. Basanti Devi	Shri Dilwar Singh	Vill-Syanri Bangali
			Block-Ghat,
			DisttChamoli
56.	Shri Trilok Singh	Shri Balwant Singh	Vill-Syanri Bangali
			Block-Ghat,
			DisttChamoli
57.	Shri Pradeep Dobhal	Shri Rudramani Dobhal	HAPPRC
			Contact No 8650843550
			Aadhar No 420085133327
58.	Shri Rajeev Ranjan Kumar	Shri Dharmnath Singh	HAPPRC
	5 5	6	Contact No 9412974451
			Aadhar No415953645662
59.	Shri Ajay Hemdan	Shri Ashok Kumar Hemdan	HAPPRC
			Contact No8272820207
60.	Shri Kuldeep Rawat	Shri Mahaveer Singh Rawat	HAPPRC
			Contact No9760944027
			Aadhar No944171489054



Figure 10. Photographs of the one day On-farm farmers workshop cum training/plant distribution programme organized for promotion of cultivation of MAP's at village Syanri Bangali, Block, Ghat, Chamoli on 14/7/2022. Registration of participants (A), Difussion of technical knowledge to farmers (B), chief guests address to the farmers (C-E), distribution of plants of *P. kurrooa* (G-H), *N. jatamansi* (I) and *A. heterophyllum* (J) to farmers amd group photograph of participants (K-L).

Table 20. List of villagers/farmers participated in one day On-farm farmers workshop cum training/plant distribution programme organized for promotion of cultivation of MAP's at Village Pagna Block, Nandanagar, Chamoli on 27/7/2022 under J.W.C.T. Project. Total participants: 51(male-30, female-21)

Sr.Ño	Villagers Name	Father/Husband Name	Address
1.	Smt. Saraswati Devi	Shri Madho Ram	Vill- Kanol Block & DisttNandanagar, Chamol Contact No 9927115094 Aadhar No 798908278461
2.	Smt. Bharti Pharswan	Shri Tribhuwan Pharswan	Vill- Pagna Block &DisttNandanagar, Chamoli Contact No 7500673346
3.	Smt. Nandita Rawat	Shri Karendra Rawat	Vill- Malkot Block &DisttNandanagar, Chamoli Contact No 80778421769
4.	Shri Yashpal Agri	Shri Mahaveer Singh	Vill- Guladi Block &DisttNandanagar, Chamoli Contact No9012045708
5.	Shri Vijay Mendoli	Shri Sundarmani	Vill- Sainti Nandanagar Block &DisttNandanagar, Chamoli Contact No9568035833
6.	Shri Rakesh Kumar	Shri Shriram	Vill- Bhesaj Bhawan Block & DisttKranprayag, Chamoli Contact No 9927571214 Aadhar No669051011254
7.	Shri Umrav Negi	Shri Balwant Singh	Vill- Sitel Block & DisttNandanagar, Chamol: Contact No7500216430
8.	Shri Jitendra Rawat	Shri Nandan Singh Rawat	Vill- Sitel Block & DisttNandanagar, Chamoli Contact No7060319014 Aadhar No 90883718063
9.	Smt. Deepa Negi	Shri Vijay Singh	Vill- Sitel Block &DisttNandanagar, Chamoli Contact No 8439157977
10.	Shri Sandeep Singh	Shri D.S.Sajwan	Vill- Sitel Block &DisttNandanagar, Chamoli Contact No 8279808581
11.	Smt. Kamla Devi	Shri Pushkar Singh	Vill- Sitel Block &DisttNandanagar, Chamoli Contact No 9520218041
12.	Smt. Rameshwari Devi	Shri Abbal Singh	Vill- Sitel Block & DisttNandanagar, Chamoli Contact No 9520218062 Aadhar No 616701084383
13.	Smt. Kanti Devi	Shri Sujan Singh	Vill- Sitel Block &DisttNandanagar, Chamoli Contact No 7533895526
14.	Smt. Laxmi Devi	Shri Abbal Singh	Vill- Sitel Block & DisttNandanagar, Chamoli Contact No 7351439477
15.	Smt. Savitri Devi	Shri Khilap Singh	Vill- Sitel Block &DisttNandanagar, Chamoli Contact No8791778908 Aadhar No 263687921886
16.	Smt. Geeta Devi	Shri Indra Singh	Vill- Sitel Block & DisttNandanagar, Chamoli Contact No 7453967537
17.	Smt. Anshi Devi	Lt. Shri Harsh Singh	Vill- Sitel Block &DisttNandanagar, Chamoli

			Contact No 8923032403
18.	Smt. Budli Devi	Shri Kedar Singh	Vill- Gairi Block &DisttNandanagar, Chamoli
19.	Smt. Urmila Devi	Shri Laxman Singh	Vill- Sitel Block &DisttNandanagar, Chamoli Contact No 8449854799 Aadhar No 497032590843
20.	Smt. Sunita Devi	Shri Yogendra Singh	Vill- Sitel Block &DisttNandanagar, Chamoli Contact No 9084325218 Aadhar No 818330837236
21.	Smt. Heera Devi	Shri Narendra Singh	Vill- Sitel Block &DisttNandanagar, Chamoli Contact No 7037554612 Aadhar No 460088923545
22.	Smt. Jamuna Devi	Shri Jitendra Singh	Vill- Sitel Block &DisttNandanagar, Chamoli Contact No 7500323660
23.	Km. Mathura	Shri Dev Singh	Vill- Sitel Block &DisttNandanagar, Chamoli Contact No 8477082468
24.	Smt. Gaytri Devi	Shri Rai Singh	Vill- Sitel Block & DisttNandanagar, Chamoli Contact No 7500363213
25.	Smt. Deepa Devi	Shri Deepak Singh	Vill- Sitel Block &DisttNandanagar, Chamoli Contact No 8057219226
26.	Smt. Parvati Devi	Shri Surendra Singh	Vill- Sitel Block &DisttNandanagar, Chamoli Contact No 7088756474 Aadhar No 489277086965
27.	Smt. Ganeshi Devi	Shri Sanjay Singh	Vill- Sitel Block &DisttNandanagar, Chamoli Contact No 8979307219
28.	Smt. Janki Devi	Shri Chandra Singh	Vill- Sitel Block &DisttNandanagar, Chamoli Contact No 7819012477 Aadhar No 417905779520
29.	Smt. Sulochana Devi	Shri Narayan Singh	Vill- Sitel Block &DisttNandanagar, Chamoli Contact No 9520218013 Aadhar No 945725865692
30.	Shri Pradeep Singh	Shri Narendra Singh	Vill- Prandmati Block & DisttNandanagar, Chamoli Contact No 8193945422 Aadhar No 929693693343
31.	Shri Sabar Singh	Shri Alam Singh	Vill- Pairi Block &DisttNandanagar, Chamoli Contact No 7253939628 Aadhar No 364091627838
32.	Shri Surendra Singh	Shri Kedar Singh	Vill- Pairi Block & DisttNandanagar, Chamoli Contact No 7983715406 Aadhar No 957847336456
33.	Smt. Kalli Devi	Shri Kanchan Singh	Vill- Sitel Block &DisttNandanagar, Chamoli Contact No 7500870532
34.	Smt. Nandi Devi	Shri Bakhtawar Singh	Vill- Sitel Block &DisttNandanagar, Chamoli Contact No 8937808436
35.	Shri Kanchan Singh	Shri Meharban Singh	Vill- Sitel Block &DisttNandanagar, Chamoli

			Contact No 8868056409
36.	Shri Sachin Singh	Shri Pushkar Singh	Vill- Sitel Block &DisttNandanagar, Chamoli Contact No 7453967305 Aadhar No 713407125416
37.	Shri Gaur Singh	Shri Umrav Singh	Vill- Sitel Block &DisttNandanagar, Chamoli Contact No 7500750841 Aadhar No 472258959007
38.	Smt. Dhamti Devi	Shri Diwan Singh	Vill- Sitel Block & DisttNandanagar, Chamoli Contact No 9837150806
39.	Smt. Nandi Devi	Shri Pushkar Singh	Vill- Sitel Block &DisttNandanagar, Chamoli Contact No 8474944989 Aadhar No 215251248943
40.	Shri Pratap Singh	Shri Kotwal Singh	Vill- Guladi Block &DisttNandanagar, Chamoli Contact No 7417572885 Aadhar No 372552996423
41.	Shri Suraj Singh	Shri Amar Singh	Vill- Waduk Block &DisttNandanagar, Chamoli Contact No 9837805538 Aadhar No 886362898160
42.	Shri Rajendra Singh	Shri Laxman Singh	Vill- Waduk Block &DisttNandanagar, Chamoli Contact No 9568035927 Aadhar No 645215975101
43.	Shri Virendra Kumar	Shri Dulpi Ram	Vill- Prandmati Block &DisttNandanagar, Chamoli Contact No 8958169069 Aadhar No 349166612193
44.	Shri Jagdish Prasad	Shri Vidya Dutt	Vill- Guladi Block &DisttNandanagar, Chamoli Contact No 8958155523 Aadhar No 910665786672
45.	Shri Anand Singh	Shri Chuyya Singh	Vill- Gairi Block &DisttNandanagar, Chamoli Aadhar No 724853295119
46.	Shri Balwant Singh	Shri Meharwan Singh	Vill- Guladi Block &DisttNandanagar, Chamoli Contact No 7351264118 Aadhar No 552315404069
47.	Shri Kamal Singh	Shri Deewan Singh	Vill- Gairi Block &DisttNandanagar, Chamoli Contact No 9837805442 Aadhar No 953006035660
48.	Shri Kunwar Singh	Shri Balwant Singh	Vill- Kanol Block &DisttNandanagar, Chamoli Contact No 7500841277 Aadhar No 944316304493
49.	Shri Veer Singh	Shri Khilap Singh	Vill- Kanol Block &DisttNandanagar, Chamoli Contact No 8192858022 Aadhar No 632872334704
50.	Shri Alam Singh	Shri Hukum Singh	Vill- Pairi Block &DisttNandanagar, Chamoli Contact No 9639741400 Aadhar No 840529247896
51.	Shri Ranjit Singh	Shri Dulap Singh	Vill- Guladi Block &DisttNandanagar, Chamoli Contact No 7453927478

			Aadhar No 653174801062
52.	Shri Alam Ram	Shri Ashad Ram	Vill- Prandmati Block & DisttNandanagar, Chamoli Contact No 8958539960 Aadhar No 510783640548
53.	Shri Pratap Singh	Shri Deewan Singh	Vill- Guladi Block &DisttNandanagar, Chamoli Contact No 9012769602
54.	Smt. Pushpa Devi	Shri Balwant Singh	Vill- Sitel Block & DisttNandanagar, Chamoli Contact No 7037257267 Aadhar No 359202351701
55.	Smt. Saruja Devi	Shri Kunwar Singh	Vill- Sitel Block &DisttNandanagar, Chamoli Contact No 8791135542
56.	Shri Seri Ram	Shri Banwa Ram	Vill- Sitel Block & DisttNandanagar, Chamoli Contact No 9756266375
57.	Shri Kanchan Negi	Shri Ganga Singh	Vill- Kanol Block & DisttNandanagar, Chamoli Contact No 9568186162 Aadhar No 703164638014
58.	Shri Nanda Gaur	Shri Bhawani Dutt Gaur	Vill- Sainti Block & DisttNandanagar, Chamoli Contact No 8191978328 Aadhar No 232325848714
59.	Shri Kalpeshwar Prasad Sati	Shri Peetambar Prasad Sati	Vill- Sainti Block &DisttNandanagar, Chamoli Contact No 8192012865
60.	Shri Daulat Singh	Shri Sabbal Singh Bisht	Vill- Aala Block & DisttNandanagar, Chamoli Contact No 89419896307 Aadhar No 627960890461
61.	Shri Khilap Singh	Shri Balwant Singh	Vill- Kanol Block & DisttNandanagar, Chamoli Contact No 9690528062 Aadhar No 267949322010
62.	Shri Sanjay Singh	Shri Alam Singh	Vill- Sitel Block & DisttNandanagar, Chamoli Contact No 7078891174 Aadhar No 635586366937
63.	Shri Balwant Singh	Shri Dhan Singh	Vill- Gairi Block &DisttNandanagar, Chamoli Contact No 8958662515 Aadhar No 497494469674
64.	Shri Alam Singh	Shri Hari Singh	Vill- Prandmati Block & DisttNandanagar, Chamoli Contact No 7351624159 Aadhar No 391544560153
65.	Shri Abbal Negi	Shri Alama Singh	Vill- Morav Malla Block &DisttNandanagar, Chamoli Contact No 9690404341 Aadhar No 449242522329
66.	Shri Mangal Singh	Shri Hayat Singh	Vill- Sitel Block &DisttNandanagar, Chamoli Aadhar No 723551807678
67.	Shri Vikram Singh	Shri Sarup Singh	Vill- Waduk Block & DisttNandanagar, Chamoli Contact No 9639453953
68.	Shri Umed Singh Panwar	Shri Gulab Singh	Vill- Gairi Block &DisttNandanagar, Chamoli Contact No 9997637660

			Aadhar No 263043645429	
69.	Shri Gabbar Singh	Shri Ratan Singh	Vill- Pairi Block & DisttNandanagar, Chamoli Contact No 8958661718	
70.	Shri Rajendra Singh	Shri Daulat Singh	Vill- Sitel Block &DisttNandanagar, Chamoli Contact No 9756256210 Aadhar No 488785576830	
71.	Dr. V. K. Purohit	Shri A.P. Purohit	HAPPRC Contact No 9456531715	
72.	Shri Pradeep Dobhal	Shri R. M. Dobhal	HAPPRC Contact No 8650843550	
73.	Shri Jaidev Chauhan	Shri B. S. Chauhan	HAPPRC Contact No 8126211560	
74.	Shri Kailash Kandpal	Shri B. P. Kandpal	HAPPRC Contact No 7302808941	
75.	Shri Ajay Hemdan	Shri Ashok Kumar Hemda	HAPPRC Contact No 8272820207	
76.	Shri Kamal Pundir	Shri G. S. Pundir	HAPPRC Contact No 9540468782	
77.	Shri Kuldeep Rawat	Shri Mahaveer Singh	HAPPRC Contact No 9760944027	



Figure 11. Photographs of the one day On-farm farmers workshop cum training/plant distribution programme organized for promotion of cultivation of MAP's at Sitel Block, Nandanagar (Ghat), Chamoli on 28/7/2022. Registration of participants (A), Difussion of technical knowledge to farmers by Dr. V.K. Purohit Senior Scientific Officer (B-C), address of chief guests to the farmers (D-F), distribution of plants of *P. kurrooa* to farmers (G-J) and group photograph of participants (K-L).



Figure 12. Photographs of the one day On-farm farmers workshop cum training/plant distribution programme organized for promotion of cultivation of MAP's at village Pagna Block, Nandanagar, Chamoli on 27/7/2022. Registration of participants (A), Difussion of technical knowledge to farmers (B), chief guests address to the farmers (C-E), distribution of plants of *P. kurrooa* to farmers (D-E), and group photograph of participants (F).

Table 21. List of villagers/farmers participated in one day On-farm farmers workshop cum training/plant distribution programme organized at village Syanri Bhainti, Nandanagar, Chamoli on 04/08/2022 under J.W.C.T. Project.

Sr.No.	Villagers Name	Father/Husband Name	Address	
1.	Smt. Manisha Kaithait	Shri Surendra Singh	Vill- Syanri Bhainti	
			Block-Nandanagar, DisttChamoli	
			Contact No 8194024496	
			Aadhar No 644156741861	
2.	Shri Ram Bisht	Shri Digambar Singh	Vill- Syanri Bhainti	
			Block-Nandanagar, DisttChamoli	
			Contact No 7351500201	
			Aadhar No 455294685807	
3.	Shri H.N. Mainduli	Shri M.R. Mainduli	Vill- Syanri Bhainti	
			Block-Nandanagar, DisttChamoli	
			Contact No 7500300112	
4.	Smt. Pushpa Devi	Shri Vikram Singh	Vill- Syanri Bhainti	
			Block-Nandanagar, DisttChamoli	
5.	Smt. Anjani Devi	Shri Jaspal Singh	Vill- Syanri Bhainti	
			Block-Nandanagar, DisttChamoli	
			Contact No 8449438950	
6.	Smt. Anita Devi	Shri Gopichand	Vill- Syanri Bhainti	
			Block-Nandanagar, DisttChamoli	
			Contact No 8958187479	

Total participants: 62 (male- 43, female-19)

			Aadhar No981996582558
7.	Smt. Anusuya Devi	Shri Digamber Singh	Vill- Syanri Bhainti
			Block-Nandanagar, DisttChamoli
			Contact No 7500057436
			Aadhar No857094032313
8.	Smt. Radha Devi	Shri Surendar Rawat	Vill- Syanri Bhainti
			Block-Nandanagar, DisttChamoli
9.	Smt. Dhanuli Devi	Shri Mahendar Singh	Vill- Syanri Bhainti
		C	Block-Nandanagar, DisttChamoli
			Contact No 9758654605
10.	Smt. Radha Devi	Shri Khilaf Singh	Vill- Syanri Bhainti
		e	Block-Nandanagar, DisttChamoli
			Contact No 7248585265
11.	Shri Surendra Singh	Shri Dayal Singh	Vill- Syanri Bhainti
			Block-Nandanagar, DisttChamoli
			Contact No 9639321778
12.	Shri Amar Singh	Shri Khem Singh	Vill- Syanri Bhainti
	String to the string to		Block-Nandanagar, DisttChamoli
13.	Shri Harendra Singh	Shri Diwan Singh	Vill- Syanri Bhainti
	singht singht		Block-Nandanagar, DisttChamoli
			Contact No 9837757467
14.	Shri Narendra Singh	Shri Mahipal Singh	Vill- Syanri Bhainti
	Shiri (alenara Shigh	Sur tranpa Suga	Block-Nandanagar, DisttChamoli
			Contact No 9639634242
15.	Shri Pushkar Singh	Shri Raghubir Singh	Vill- Syanri Bhainti
10.	Shiri i ushkur Shigh		Block-Nandanagar, DisttChamoli
16.	Shri Pradeep Kumar	Shri Raghubir Ram	Vill- Syanri Bhainti
100	Shiri Fudeep Human		Block-Nandanagar,
			DisttChamoli
			Contact No 7055905956
17.	Shri Himmat Singh	Shri Gyan singh	Vill- Syanri Bhainti
1,1	Sini ininin Singh		Block-Nandanagar, DisttChamoli
			Contact No 9927941430
18.	Shri Vikram Ram	Shri Hukumi Ram	Vill- Syanri Bhainti
100			Block-Nandanagar, DisttChamoli
			Contact No 6399226328
19.	Shri Vikram Singh	Shri Mahipal Singh	Vill- Syanri Bhainti
_~ •		F	Block-Nandanagar, DisttChamoli
			Contact No 9670354242
			Aadhar No608070662137
20.	Shri Dilbar Singh	Shri Balak Singh	Vill- Syanri Bhainti
	<u>-</u>		Block-Nandanagar, DisttChamoli
			Contact No 7500639926
21.	Shri Manoj Singh	Shri Mahendar Singh	Vill- Syanri Bhainti
	series series series		Block-Nandanagar, DisttChamoli
			Contact No 7618619006
			Aadhar No 207739449507
22.	Shri Vikram Singh	Shri Dalbeer Singh	Vill- Syanri Bhainti
		Shiri Duibber Shigh	Block-Nandanagar,
			DisttChamoli
			Contact No 9690267796
			Contact 100 9090207790

23.	Shri Akshay Kumar	Shri Raj Kumar	Vill- Syanri Bhainti
	Shiri monuji Kumu	Sini Kuji Kunu	Block-Nandanagar, DisttChamoli
			Contact No7618354621
			Aadhar No 386441198214
24.	Shri Sanju Ram	Shri Bhajni Ram	Vill- Syanri Bhainti
24.	Siiri Saiiju Kaiii	Siiri Dilajiii Kalii	Block-Nandanagar, DisttChamoli
			Contact No 9105625742
			Aadhar No 275358745229
25.	Shri Anond Singh	Shri Harak Singh	Vill- Syanri Bhainti
23.	Shri Anand Singh		Block-Nandanagar, DisttChamoli
			Contact No 7248646629
26.	Shri Kamal Singh	Shri Gopal Singh	Vill- Syanri Bhainti
20.	Sini Kamai Singh		-
			Block-Nandanagar, DisttChamoli Contact No 8193915761
27	Chri Vinin Cinch	Shri Dianal Sirah	Aadhar No 851002575508
27.	Shri Vipin Singh	Shri Digpal Singh	Vill- Syanri Bhainti Black Nondonagor Distt Chamali
			Block-Nandanagar, DisttChamoli
			Contact No 7351959466
20	Chai Darra da C' 1		Aadhar No313627468887
28.	Shri Devendra Singh	Shri Kuwar Singh	Vill- Syanri Bhainti Black Nandanagar, Digtt, Chamali
			Block-Nandanagar, DisttChamoli
			Contact No 8958583396
			Aadhar No 739208719361
29.	Shri Kamleshwar	Shri Indra Ram	Vill- Syanri Bhainti
			Block-Nandanagar, DisttChamoli
			Aadhar No 615953914223
30.	Shri Padmi Ram	Shri Jagdish Ram	Vill- Syanri Bhainti
			Block-Nandanagar, DisttChamoli
			Contact No. -7534084861
			Aadhar No 931215619444
31.	Shri Mukesh Ram	Shri Chaitu Ram	Vill- Syanri Bhainti
			Block-Nandanagar, DisttChamoli
			Contact No 8476924561
			Aadhar No707316614730
32.	Shri Kalyan Singh	Shri Prem Singh	Vill- Syanri Bhainti
			Block-Nandanagar, DisttChamoli
			Contact No 7618529037
			Aadhar No 846725012176
33.	Shri Vijay Ram	Shri Gandhi Ram	Vill- Syanri Bhainti
			Block-Nandanagar, DisttChamoli
			Contact No 9927176755
			Aadhar No 658896307409
34.	Shri Ratan Singh	Shri Tan Singh	Vill- Syanri Bhainti
			Block-Nandanagar, DisttChamoli
35.	Shri Karan Singh	Shri Khyati Singh	Vill- Syanri Bhainti
			Block-Nandanagar, DisttChamoli
			Contact No 7535977745
36.	Smt. Hema Devi	Shri Vinod Singh Bisht	Vill- Syanri Bhainti
			Block-Nandanagar, DisttChamoli
			Contact No 9690020172
37.	Smt. Pooja Devi	Shri Deepak Singh	Vill- Syanri Bhainti
			- ,

			Block-Nandanagar, DisttChamoli	
38.	Shri Raghuveer Singh	Shri Fathe Singh	Vill- Syanri Bhainti	
			Block-Nandanagar, DisttChamoli	
			Contact No 8958675358	
			Aadhar No 622765820727	
39.	Smt. Kalawati Devi	Shri Sishupal Singh	Vill- Syanri Bhainti	
			Block-Nandanagar, DisttChamoli	
40.	Smt. Ghulli Devi	Shri Khilaf Singh	Vill- Syanri Bhainti	
			Block-Nandanagar, DisttChamoli	
41.	Smt. Dhanuli Devi	Shri Matendar Singh	Vill- Syanri Bhainti	
			Block-Nandanagar, DisttChamoli	
42.	Smt. Naumi Devi	Shri Mahipal Singh	Vill- Syanri Bhainti	
-		1 0	Block-Nandanagar, DisttChamoli	
43.	Shri Mahipal Ram	Shri Kutti Ram	Vill- Syanri Bhainti	
			Block-Nandanagar, DisttChamoli	
			Contact No 7500956263	
44.	Shri Vinod Kumar	Shri Hukumi Ram	Vill- Syanri Bhainti	
			Block-Nandanagar, DisttChamoli	
			Contact No 7088327762	
45.	Shri Ratan Singh	Shri Prem Singh	Vill- Syanri Bhainti	
			Block-Nandanagar, DisttChamoli	
			Contact No 9690122621	
46.	Shri Sunil Singh	Shri Kamal Singh	Vill- Syanri Bhainti	
40.	Shiri Suhiri Shirgh	Shiri Kumur Shigh	Block-Nandanagar, DisttChamoli	
			Contact No 8449440898	
47.	Shri Gopal Ram	Shri Kamal Ram	Vill- Syanri Bhainti	
-7.	Shiri Gopai Kani		Block-Nandanagar, DisttChamoli	
			Contact No 8941067172	
			Aadhar No402049466343	
48.	Shri Gabbar Singh	Shri Dholya Singh	Vill- Syanri Bhainti	
-0.	Shiri Gabbai Shigh	Silli Dilorya Siligi	Block-Nandanagar, DisttChamoli	
			Aadhar No 347952718954	
49.	Smt. Basanti Devi	Shri Shankar Singh	Vill- Syanri Bhainti	
42.	Sint. Dasanti Devi	Silli Shankar Siligi	Block-Nandanagar, DisttChamoli	
50.	Smt. Gomti Devi	Shri Girdhari Ram	Vill- Syanri Bhainti	
50.			Block-Nandanagar, DisttChamoli	
			Aadhar No 919002102139	
51.	Smt. Savitri Devi	Shri Jagdish Ram	Vill- Syanri Bhainti	
51.			Block-Nandanagar, DisttChamoli	
			Aadhar No 218934905873	
52.	Shri Balwant Singh	Shri Thuni Singh	Vill- Syanri Bhainti	
32.	Silli Daiwalit Siligli		Block-Nandanagar, DisttChamoli	
			Contact No 7055527591	
53	Chui Donicat Circ-1	Chui Derman Circe-1	Aadhar No 355187853635	
53.	Shri Ranjeet Singh	Shri Puran Singh	Vill- Syanri Bhainti Black Nandens con Diette Chamali	
			Block-Nandanagar, DisttChamoli	
			Contact No 7351928523	
54.	Smt. Vimla Devi	Shri Sobhan Singh	Vill- Syanri Bhainti	
			Block-Nandanagar, DisttChamoli	
			Aadhar No 540329684015	
55.	Shri Pushkar Singh	Shri Umed Singh	Vill- Syanri Bhainti	

			Block-Nandanagar, DisttChamoli	
			Contact No 7500015449	
56.	Smt. Bhaduli Devi	Shri Mohan Singh	Vill- Syanri Bhainti	
			Block-Nandanagar, DisttChamoli	
			Contact No 8057542318	
			Aadhar No 351857574202	
57.	Shri Dheeraj Pal	Shri Shri Feti Ram	Vill- Syanri Bhainti	
			Block-Nandanagar, DisttChamoli	
			Contact No 8191824510	
			Aadhar No 450390409346	
58.	Shri Bhajan Singh	Shri Shri Thepad Singh	Vill- Syanri Bhainti	
			Block-Nandanagar, DisttChamoli	
			Contact No 8938953748	
			Aadhar No 539319209892	
59.	Shri Pradeep Dobhal	Shri R. M. Dobhal	HAPPRC, Contact No 8650843550	
60.	Shri Kailash Kandpal	Shri B. P. Kandpal	HAPPRC, Contact No 7302808941	
61.	Shri Ajay Hemdan	Shri A. K. Hemdan	HAPPRC, Contact No 8272820207	
62.	Shri Kuldeep Rawat	Shri Mahaveer Singh	HAPPRC, Contact No 9760944027	



Figure 13. Photographs of the one day On-farm farmers workshop cum training/plant distribution programme organized for promotion of cultivation of MAP's at Syanri Bhainti, Nandanagar, Chamoli on 04/08/2022. Registration of participants (A), Difussion of technical knowledge to farmers (B), address of chief guests to the farmers (C-D), distribution of plants of *N. grandiflora* and *P. kurrooa* to farmers (E), group photograph of participants (F).

Table 22. List of villagers/farmers participated in one day On-farm farmers workshop cum training/plant distribution programme organized for promotion of cultivation of MAP's at Village Teela, Block Thalisain, District Pauri Garhwal on 5/7/2022.

Sr.No.	Villagers Name	Father/Husband Name	Address
1.	Shri Belam Singh	Shri Jaswant Singh	Vill-Teela Block- Thalisain, DisttPauri Garhwal Contact No 9690923649 Aadhar No 593324158176
2.	Shri Satye Singh	Shri Chetriya Singh	Vill-Teela

Total participants: 64 (male-50, female-14)

			Block-Thalisain, DisttPauri Garhwal	
			Contact No 8879482627	
3.	Shri Umed Singh	Shri Ghwan Singh	Vill-Teela Block-Thalisain, Distt Pauri Garhwal	
4.	Shri Kartik Singh	Shri Jomal Singh	Vill-Teela Block- Thalisain, DisttPauri Garhwal	
5.	Shri Guman Singh	Shri Chetrya Singh	Vill-Teela Block- Thalisain, DisttPauri Garhwal	
6.	Shri Gagan Singh	Shri Kartik Singh	Vill-Teela Block- Thalisain, DisttPauri Garhwal	
7.	Shri Hansa Singh	Shri Gudal Singh	Vill-Teela Block- Thalisain, DisttPauri Garhwal	
8.	Shri Uday Singh	Shri Matbar Singh	Vill-Teela Block- Thalisain, DisttPauri Garhwal	
9.	Shri Govind Singh	Shri Morkhala Singh	Vill-Teela Block- Thalisain, DisttPauri Garhwal Contact No 8475860031	
10.	Shri Bharat Singh	Shri Uday Singh	Vill-Teela Block- Thalisain, DisttPauri Garhwal Contact No 9654729406 Aadhar No 694119868318	
11.	Smt. Ganeshi Devi	Shri Vijay Singh	Vill-Teela Block- Thalisain, DisttPauri Garhwal	
12.	Shri Madan Singh	Shri Darshan Singh	Vill-Teela Block- Thalisain, DisttPauri Garhwal Contact No 8302788058 Aadhar No 210751927693	
13.	Shri Chhora Singh	Shri Gabar Singh	Vill-Teela Block- Thalisain, DisttPauri Garhwal	
14.	Shri Virendra Singh	Shri Darshan Singh	Vill-Teela Block- Thalisain, DisttPauri Garhwal Contact No 9917568377	
15.	Shri Ram Lal	Shri Kala Lal	Vill-Teela Block- Thalisain, DisttPauri Garhwal Aadhar No 657259955374	
16.	Shri Narayan Singh	Shri Nathi Singh	Vill-Teela Block- Thalisain, DisttPauri Garhwal Contact No 9690732048	
17.	Shri Sangram Singh	Shri Chhota Singh	Vill-Teela Block- Thalisain, DisttPauri Garhwal Contact No 7500566744	
18.	Shri Bhupal Lal	Shri Kala Lal	Vill-Teela Block- Thalisain, DisttPauri Garhwal Aadhar No 382715992946	
19.	Shri Barjan Lal	Shri Jangali Lal	Vill-Teela Block- Thalisain, DisttPauri Garhwal Contact No 6395677901 Aadhar No 414115825286	
20.	Smt. Usha Devi	Shri Kala Lal	Vill-Teela Block- Thalisain, DisttPauri Garhwal Contact No 7298105622 Aadhar No 805040324898	
21.	Shri Prem Lal	ShriFagunu Lal	Vill-Teela Block- Thalisain, DisttPauri Garhwal Aadhar No 255269479523	
22.	Shri Kunjpal Lal	Shri Idhala Lal	Vill-Teela Block- Thalisain, DisttPauri Garhwal Aadhar No 329852275768	
23.	Shri Khushal Singh	Shri Darshan Singh	Vill-Teela Block- Thalisain, DisttPauri Garhwal	

24.	Shri Madan Singh	Shri Akal Singh	Vill-Teela
			Block- Thalisain, DisttPauri Garhwal Contact No 9627431890
25.	Shri Matbar Singh	Shri Gabar Singh	Vill-Teela
			Block- Thalisain, DisttPauri Garhwal Contact No 8791891712
26.	Shri Chakraveer Singh	Shri Kutal Singh	Vill-Teela
			Block- Thalisain, DisttPauri Garhwal Contact No 7055152628
			Aadhar No 916334742489
27.	Shri Digambar Singh	Shri Thep Singh	Vill-Teela Block- Thalisain, DisttPauri Garhwal
			Contact No 9917568095
28.	Shri Alam Singh	Shri Darman Singh	Vill-Teela
			Block- Thalisain, DisttPauri Garhwal Contact No 8392848231
29.	Shri Raje Singh	Shri Chandra Singh	Vill-Teela Block- Thalisain, DisttPauri Garhwal
30.	Shri Kunwar Singh	Shri Jagat Singh	Vill-Teela
21	Chui Honor des Circili	Chui Caia Cira I	Block- Thalisain, DisttPauri Garhwal Vill-Teela
31.	Shri Harendra Singh	Shri Gaje Singh	Block- Thalisain, DisttPauri Garhwal
			Contact No 9193329875
32.	Shri Gaur Singh	Shri Pancham Singh	Vill-Teela Block- Thalisain, DisttPauri Garhwal
33.	Shri Suraj Singh	Shri Jeewan Singh	Vill-Teela
			Block- Thalisain, Distt Pauri Garhwal Contact No 7536867047
34.	Shri Meharban Singh	Shri Prem Singh	Vill-Teela
			Block- Thalisain, DisttPauri Garhwal Contact No 9720510171
			Aadhar No 958970279282
35.	Shri Virendra Negi	Lt. Shri Gaina Singh	Vill-Teela
			Block- Thalisain, DisttPauri Garhwal Contact No 8447107954
			Aadhar No 394908301287
36.	Shri Mahaveer Singh	Shri Kunwar Singh	Vill-Teela Block- Thalisain, DisttPauri Garhwal
			Contact No 8279947131
			Aadhar No 603693354694
37.	Shri Surendra Lal	Shri Shishpal Lal	Vill-Teela Block- Thalisain,
			DisttPauri Garhwal
20	Shri Suresh Lal	Shri Mahiya Lal	Aadhar No 677220545252 Vill-Teela
38.		Shri Mohiya Lal	Block- Thalisain, DisttPauri Garhwal
			Contact No 9557976041
39.	Shri Vikram Singh	Lt. Shri Shyam Singh	Aadhar No 386528503518 Vill-Teela
57.			Block- Thalisain,
			DisttPauri Garhwal Contact No 9389183695
			Aadhar No 593397476580
40.	Shri Sobat Singh	Lt. Shri Saupa Singh	Vill-Teela
			Block- Thalisain, DisttPauri Garhwal Contact No 9557211771
41.	Smt. Srimati Devi	Shri Darban Lal	Vill-Teela
			Block- Thalisain, DisttPauri Garhwal
42.	Smt. Savitri Devi	Shri Dhol Lal	Aadhar No 545010081427 Vill-Teela
			Block- Thalisain, DisttPauri Garhwal

			Aadhar No 901396618325
43.	Smt. Sureshi Devi	Shri Radhe Shyam	Vill-Teela Block- Thalisain, DisttPauri Garhwal Contact No 8126302115 Aadhar No 740601188704
44.	Smt. Bhama Devi	Shri Mahesh Lal	Vill-Teela Block- Thalisain, DisttPauri Garhwal Contact No 9037342983 Aadhar No 503341004878
45.	Smt. Usha Devi	Shri Narendra Tamta	Vill-Teela Block- Thalisain, DisttPauri Garhwal Contact No 9557968795 Aadhar No 714768632245
46.	Smt. Choma Devi	Shri Harish Lal	Vill-Teela Block- Thalisain, DisttPauri Garhwal Aadhar No 973260071236
47.	Smt. Guddi Devi	Shri Mani Lal	Vill-Teela Block- Thalisain, DisttPauri Garhwal Contact No 9761667599 Aadhar No 201892968325
48.	Shri Pravesh Lal	Shri Chandri Lal	Vill-Teela Block- Thalisain, DisttPauri Garhwal Aadhar No529599159654
49.	Shri Hari Prasad Pant	Shri Tulsiram Pant	Vill-Teela Block- Thalisain, DisttPauri Garhwal Contact No 9990035816 Aadhar No976217962575
50.	Smt. Meena Devi	Shri Jagdish Aagre	Vill-Teela Block- Thalisain, DisttPauri Garhwal Aadhar No528700365765
51.	Shri Vinod Lal	Shri Vinta Lal	Vill-Teela Block- Thalisain, DisttPauri Garhwal Aadhar No258666362292
52.	Shri Dinesh Singh	Shri Rajendra Singh	Vill-Teela Block- Thalisain, DisttPauri Garhwal Contact No9068532267 Aadhar No469159184518
53.	Smt. Kusum Devi	Shri Balwant Singh	Vill-Teela Block- Thalisain, DisttPauri Garhwal Contact No9917029919
54.	Smt. Dupa Devi	Shri Chand Lal	Vill-Teela Block- Thalisain, DisttPauri Garhwal Contact No8859933882 Aadhar No534058368553
55.	Shri Kalam Singh	Lt. Shri Dham Singh	Vill-Teela Block- Thalisain, DisttPauri Garhwal Contact No939412375
56.	Smt. Sarita Devi	Shri Dinesh Singh	Vill-Teela Block- Thalisain, DisttPauri Garhwal Contact No7900712264
57.	Smt. Sarojani Devi	Shri Kundan Singh	Vill-Teela Block- Thalisain, DisttPauri Garhwal Aadhar No651833412252
58.	Shri Padam Singh	Shri Bhawan Singh	Vill-Teela Block- Thalisain, DisttPauri Garhwal Contact No7088574637
59.	Shri Darshan Singh	Shri Jagat Singh	Vill-Teela Block- Thalisain, DisttPauri Garhwal
60.	Dr. Vijay Kant Purohit	Shri A. P. Purohit	HAPPRC (Senior Scientific Officer) Contact No 9456531715
61.	Shri Pradeep Dobhal	Shri R.M. Dobhal	HAPPRC, Contact No 8650843550



Figure 14. Photographs of the one day On-farm farmers workshop cum training/plant distribution programme organized for promotion of cultivation of MAP's at Teela village, district Pauri Garhwal on 05/07/2022. Diffusion of technical knowledge by Dr. V.K. Purohit (SSO) to farmers/participants (A-C), distribution of planta of *P. kurooa* to farmers (D-E) and group photograph of farmers/participants (F).

19. Monitoring of the project progress by JWCT nominee/representative: To assess the on spot progress of the JWCT project running by HAPPRC, the JWCT nominee Dr C. S. Rana (Senior Scientist – Bioresource), Mr. Aman deep (Scientist-Bioresource), Smt. Sakshi Sharma (Scientist-Bioresource), Surendra Singh Bhagat (Scientist- Bioresource), Mr. Yasveer Singh Negi (Scientist-Bioresource) and Mr. Amit Bhatt (Field Supervisor - Bioresource) visited the farmers field (Tyuni, Setail, Gairi and Rusiayan) in Rudraprayag and Chamoli District and in field stations of Pothivasa, Nature Interpretation Centre Baniyakund and Alpine Research Centre at Tungnath and Model nursery, Kulsari of HAPPRC from 23 July 2022 to 30 July 2022. The entire expert team physically monitors the seedling development activity to cultivation in farmers field under the project and critically examines the progress on other aspects of the project with face to face interaction with farmers, project staff and director of the Centre (HAPPRC). They also suggest the solutions of the difficulties, particularly low germination and high mortality of seedlings in some species (Figure).

Dr. CS Rana, ream leader of the monitoring team suggested that the, HAPPRC need to prepare a separate block under JWCT project for seed production of *N. grandiflora* at Tungnath. To consider the suggestion provided by the mnonitoring team, a work on development of separate block for seed production was estabilihed at Tungnath in the month of September-October 2023.



Figure 15. Monitoring of field stations (A), farmers fields at Rudraprayag and Chamoli district (B-F) and interaction with Prof. M.C. Nautiyal, Director, HAPPRC and Dr. Vijay Kant Purohit, PI, JWCT (G-J) project about the future work plan and fund disbursement.



Figure 16. Preparation of nursery bed's for estabilihement of separate block for seed development of Jatamansi (*Nardostachys grandiflora*) seedling at Alpine Research Centre Tungnath, recommendations by Dr. C. S. Rana (Principal Scientist-Bioresource) and their teams for Dabur Research & Development Centre.



Figure 17. Plantation of Jatamansi (*Nardostachys grandiflora*) seedling in separate block developed for seed production of jatamansi at Alpine Research Centre Tungnath (3400 m asl) under JWCT project.

Table 23. Seed collected (gm) from different field sites during the current report period (October-November, 2022) for seedling dvelopment.

Sr.No.	Name of species	Seed collection sites	Approx. seeds collected (gm)
1.	Picrorhiza kurrooa	Tungnath	200gm
2.	Nardostachys grandiflora	Tungnath	70 gm
3.	Aconitum balfourii	Tungnath(49.66gm), Baniyakund (17.8gm) and Kilpur (19.57gm)	87.03gm
4.	Aconitum heterophyllum	Tungnath (8gm), Baniyakund (22gm) and Kilpur (4.59gm)	34.59gm
5.	Sassurea costus	Pothivasa	10000 gm
		Total weight of collected seeds	1391.62 gm



Figure 18. Drying of collected seeds of different species at HAPPRC, Srinagar (Garhwal, 550m asl).





Figure 19. Packed and labeled seeds of selected species for storage and further use. A. balfourii (A-C), A. heterophyllum (D-F), P. kurrooa (G), N. grandiflora (H), and S. costus (I)

20. Seedling development and Transfer to Field for Further growth and distribution to farmers: To fulfill the demand of seedlings under project, the collected seeds were sowed inside the green house condition and approximate 80000 seedlings of selected species (2500-*A. balfouri*, 8000-*A. heterophyllum*, 30000-*P. kurrooa*, 195000- *N. grandiflora*, and 20000-*S. costus*) has been developed and transfered to field station Pothibasa (2200m asl) for further growth.





Figure 20. Photographs of seedlings development of selected species and transfered of seedlings to field station Pothibasa (2200 m asl) for further growth.

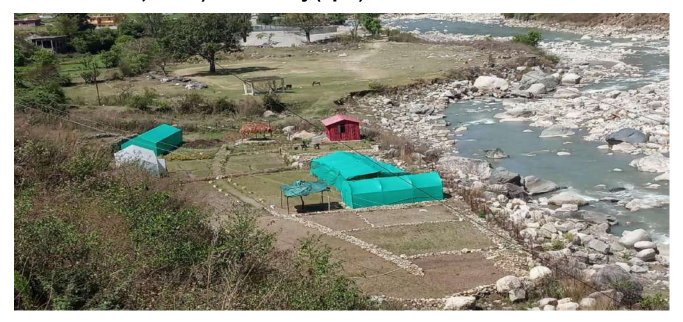
21. Establishment of satellite nursery: Keeping in view the development and maintenance of seedlings of selected species in nearby areas of farmers village/ land, establishment work of one satellite nursery in 0.5ha land has been done at Lumkundi, Kulsari under project. The land preparation and construction of polyhouse, shadehouse work and bed preparation work is completed (**Figure**). This year onwards the seedlings development and maintenance work of the selected species in JWCT project and other medicinally important species will be start.



Figure 21. View of satellite Nursery Established Under JWCT, Dabur India Ltd. Project at Lumkundi, Kulsari (1200 m asl), Chamoli.

22. Work has been done under JWCT Project (2020-2023)

- Collection of seeds (4652.8 gm) of selected species from different high altitude region/field nursery of HPPRC
- Development and maintenance of 6,20,000 seedlings in nursery condition and their transfer in farmers fields.
- > Organisation of 15 farmers meeting/workshop/training/plant distribution programme.
- Sensitization and cultivation of selected species in three districts, 18 villages and four developmental blocks of Uttarakhand with 720 villagers/farmers.
- Promotion of cultivation of selected species in 6.16 ha (312.07 nali) of farmers land.
- > Estabilishment of one satellite nursery in 0.2ha of land for future use.
- Estabilishment of one separate block for seed production of Jatamansi in
 0.1 ha (5.0 nali) at Tungnath
- Regular monitoring of project work, data collection, report writing and submission to funding agency (JWCT) – Quaterly (June, September, December, March) and Annualy (April).



Close view of Satellite Nursery at Lumkundi (1200 m asl), Kusari, Chamoli



Close view of Separate Block for Seed Production of Jatamansi at Tungnath (3400 m asl)

23. Justification/Remarks: The JIVANTI WELFARE AND CHARITABLE TRUST (JWCT) is working as a part of Dabur India Ltd. with focus on fulfilment of welfare and charitable obligations towards society at large through advancement of equal opportunities for education, providing food, healthcare, medical care, ensuring environmental sustainability, enhanced vocational skills and advancement of any other objects of general public welfare. Simultaneously, Dabur India Ltd. is a leading organization in the field of herbal medicine and therefore the firm adopted conservation and sustainable development of biological diversity as part of Corporate Social responsibility and has designed an integrated programme through a community centric Project based approach. Under this approach JWCT, Dabur India Ltd. and High Altitude Plant Physiology Research Centre (HAPPRC) is collaboratively working for promotion of cultivation and conservation of high value medicinal herbs in different parts of Uttarakhand. The role of HAPPRC is clearly defined as production of planting materials, their distribution to NGOs/farmers and promotion of cultivation in farmers field so that the produce raw materials will be easily available to Dabur India Ltd. and other concern firms of herbal medicine and livelihood opportunities for local farmers/villagers.





24. Media released/Publication: The work done under JWCT project has been highlighted through writing and publishing of scientific research papers and news publishing through print as well as electronic media for public domain.

25. Final observations from farmers field

To take up the cultivation of medicinal and aromatic plants as option/source of additional income to local inhabitants, numbers of villagers/farmers are interested to do the cultivation of selected species, but there are some concern about irrigation facility during the summer months particularly April to June, planting materials of the important species and intime marketing of the raw produce. To address all these concerns of the farmers, the cultivation of medicinal and aromatic plants can boost the local economy with conservation of RET species in natural habitats.

Dr. V.K. Purohit PI, JWCT, Dabur India Ltd. Project HAPPRC, HNBGO, Srmagan (Garhwal) Scientific Officer Althude Plant Physikogy Research Centre H.M.B. Garhwel University unneer (Garhwel) Uterathand - 246174

(Dr. Vijay Kant Purohit) Sr. Scientific Officer & Principal Investigator JWCT, Dabur India Ltd. Project

(Director/HOD)

MoU with Rishikesh Yogpeeth

एक्सटेंशन सेंटर (विस्तार केन्द) ऋषिकेश योग पीठ हेमवती नंदन बहुगुणा गढवाल विश्वविद्यालय, श्रीनगर गढवाल (केन्द्रीय विश्वविद्यालय)

रिपॉट

ऋषिकेश योग पीछ व हे०न०व०ग०वि०वि० श्रीनगर,गढवाल के मध्य दिनांक - 16 मार्च, 2021 में हुए एम०ओ०यू० के आधार पर तथा 28 मई 2021 विभागीय (बीठओठएस०), 30 जून 2021 विश्वविद्यालय विद्या परिषद व 23 अगस्त 2021 विश्वविद्यालय कार्य परिषद की बैठक में मिली स्वीकृति के पश्चात ऋषिकेश योग पीठ को विश्वविद्यालय का एक्सटेंशन सेंटर (विस्तार केन्द्र), स्वीकृत किया गया है तथा योग में प्रमाण पत्र/डिप्लोमा/योग प्रशिक्षण कार्यकम का संचालन प्राकृतिक चिकित्सा एवं योग विभाग, चौरास परिसर की देख-रेख में करने की स्वीकृति ऋषिकेश योग पीठ को दी गई है। इस कार्य को सुचारू रूप से संचालन करने हेतु डॉ० विनोद प्रसाद नौटियाल, योग प्रशिक्षक, योग विभाग को विभागीय कार्यों के साध-साथ ऋषिकेश योग पीठ का अतिरिक्त कार्यभार देकर संपर्क अधिकारी, नियुक्त किया गया है। योग प्रशिक्षण कार्यकम (15, 28 व 30 दिवस) में विदेशी नागरिक का शुल्क प्रति रू० 10,000 तथा भारतीय नागरिक का शुल्क प्रति रू० 3000 निर्धारित है। इनसे प्राप्त 40 प्रतिशत शुल्क वित्तअधिकारी,हे० न० ब० ग० वि० वि०, श्रीनगर के नाम योग फन्ड 585, 526002011015013 यूनियन बैंक, विश्वविद्यालय, चौरास शाखा में जमा किया जाता है। वर्तमान समय तक उपरोक्त कार्यक्रम से प्राप्त कुल आय रू० 3,45,600.00 (रू० तीन लाख पैतालिस हजार छ' सौ मात्र) उपरोक्त एकाउन्ट मे जमा है। 60 प्रतिशत शुल्क एक्सटेंशन सेंटर छात्र/छात्राओं से अपने द्वारा देय व्यवस्था जैसे रहने-खाने, शिक्षण, भवन इत्यादि पर व्यय के लिए लेता है। विभाग द्वारा योग प्रशिक्षण कार्यकम में प्रतिमाग करने वाले छात्र/छात्राओं को योग प्रशिक्षण प्रमाण-पत्र प्रदान किया जाता है। भविष्य में प्रमाणपत्र/ डिप्लोमा पाठयकम एक्सटेशन सेंटर में शीघ्र प्रारम्भ किए जाने है। विदेशी छात्र/छात्राओं से इस कार्यकम के पंजीकरण शुल्क के रूप में वर्तमान समय तक प्रति 20 डॉलर लगभग रू0 1,30,000.00 विश्वविद्यालय के Foreign Students कार्यालय में जमा है। योग प्रशिक्षण कार्यकम एक माह के छात्र/छात्राओं का वर्षवार विवरण -

BATCH	DATE	STUDENT		TOTAL	FEES
		FOREIGN	INDIAN	STUDENT	1000
lst	3 January 2022 to 28 January 2022	02	02	04	Rs 10,400.00
2 nd	7 March to 3 April 2022	05	02	07	D. 33 100 00
3rd	10 April to 7 May 2022	07	02		Rs.22,400.00
			02	09	Rs 30,400.00
4 th	09 May to 04 June 2022	04	05	09	Re 33 000 00
-th				05	Rs.22,000.00
5 th	11 July,2022 to 06 August 2022	02	03	05	Rs. 11,600.00
6 th	14 August to 10 September 2022	05	01	06	Rs.21,200.00
7%	122				13:24,200,00
8 th	12 September to 15 October 2022	02	NIL	02	Rs. 8,000.00
9 th	17 October to 12 November 2022	04	NIL	04	Rs. 16,000.00
	14 November to 17 December, 2022	03	NIL		Rs 12,000.00
10 th	10 December 14				10 12,000.00
10	19 December to 14 January 2023	04	NIL	04	Rs. 16,000.00
11 th	26 Feb to 25 March 2023				
2 th	3 April to 29 April 2023	NIL	02	02	Rs. 2,400.00
3 th	12 June to 8 July 2023	09	01	10	Rs. 37,200.00
-	12 Julie to 8 July 2023	NIL	01	01	Rs. 1,200.00
4 th	14th 19 100 to 02 August 2022000 0				1.00.00
2-1-1-1	14th 19 July to 02 August, 2023(15 Days Rs 1200)	01	NIL	01	Rs. 1,200.00
5 th	17.00.00.00.0			la contra de la co	
2011	17 July to 12 August 2023	02	01	03	Pr 0 200 00
6 th	21.4			0.5	Rs. 9,200.00
th	21 August to 16 September 2023	NIL	03	03	Pr 2 C00 00
3 th	26 September to 16 October 2023	07	NIL	07	Rs. 3,600.00
p ^{ch}	04 October to 24 October 2023	07	03	10	Rs. 28,000.00
th	27 October to 17 November 2023	05	NIL	05	Rs. 31,600.00
	19 November to 09 December 2023	09	01		Rs. 20,000.00
	12 December to 02 January 2024	01	NIL	10	Rs. 37,200.00
_		79	27	01	Rs. 4,000.00
	and the second		21	TOTAL	Rs. 3,45,600

उपरोक्त योग प्रशिक्षण कार्यकम के छात्र / छात्राओ हेतु वर्कशॉप का आयोजन संख्या - 02

- 1. यौगिक चिकित्सा पर दिनांक 24 एवं 25 मार्च 2022
- 2. यौगिक चिकित्सा पर दिनांक 15,16 एवं 17 मार्च 2023

वर्चुवल तीन दिवसीय अन्तराष्ट्रीय बेबिनार का आयोजन संख्या – 01

दिनांक 23 से 25 जून 2021 विषय – "हॉलेस्टिक हेल्थ थ्रू योगा ड्यूरिंग कोविड 19"

प्रत्येक माह विभाग की फैक्लटी के द्वारा ऑफ व वर्चुवल स्तर पर अतिथि शिक्षक के रूप में योग पर व्याख्यान व प्रशिक्षण उपरोक्त कार्यक्रम के छात्र/छात्राओ को दिया जाता है तथा एक्सटेंशन सेंटर व योग विभाग के छात्र/छात्राओ के मध्य योग विद्या पर व शोध कार्यों पर विचार विमर्श समयानुसार चलता रहता है।









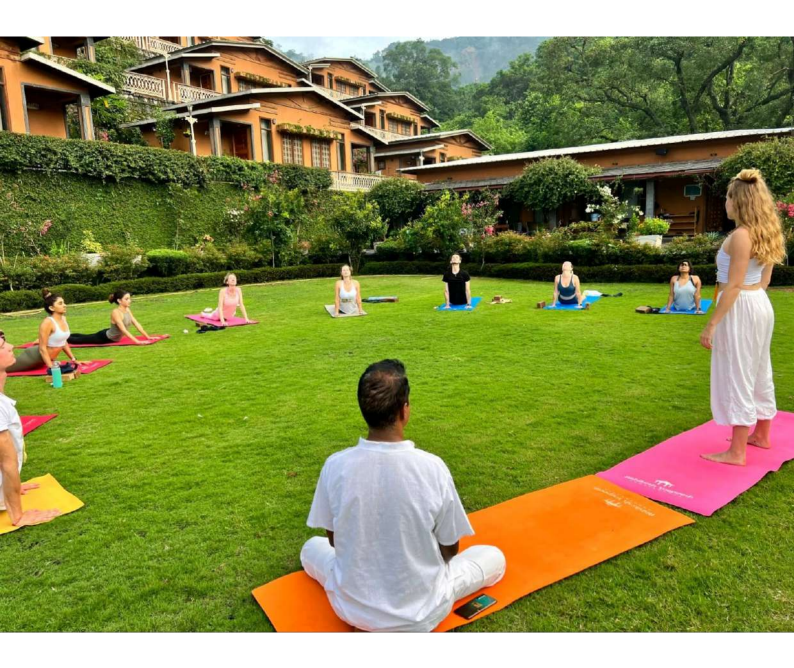


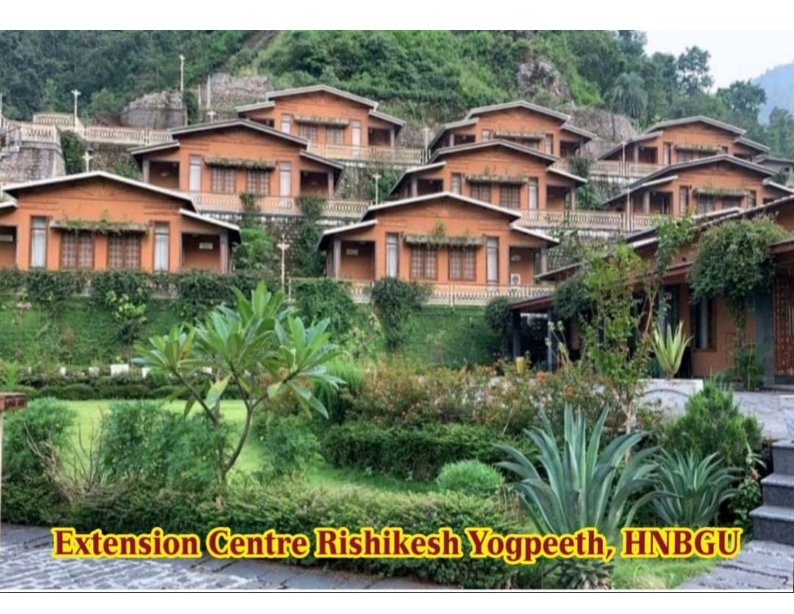














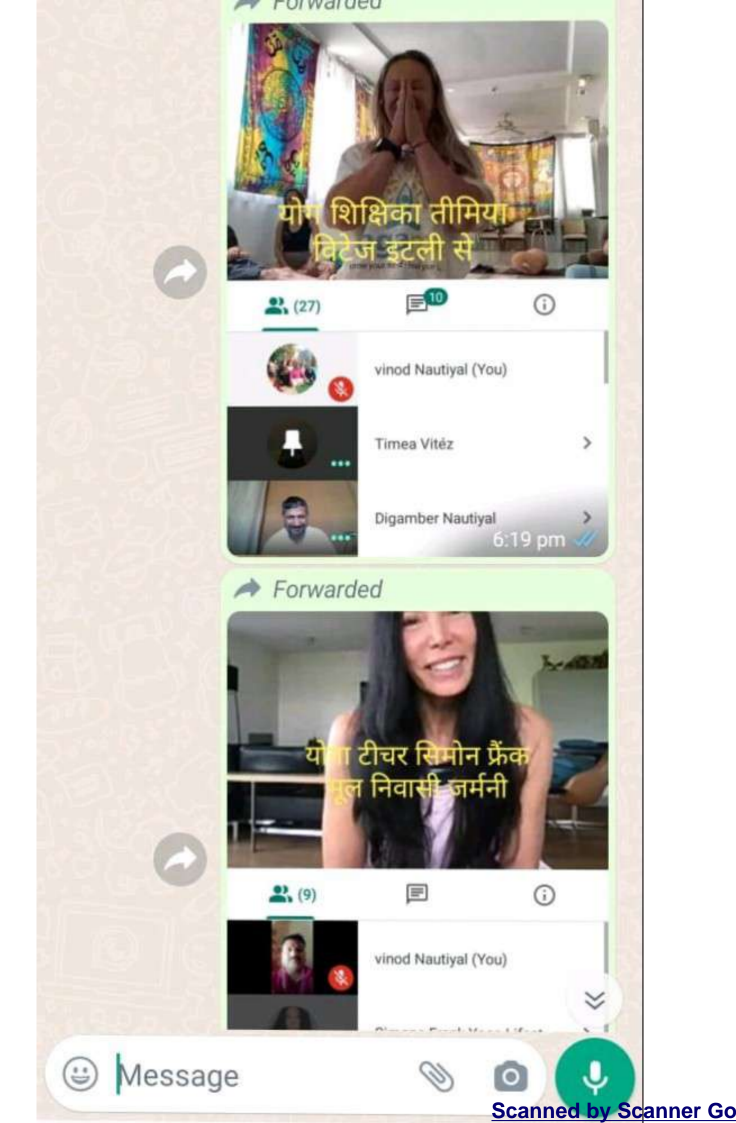
International Webinar on Yoga Organized by Naturopathy and Yoga Dept HNBGU With Rishikesh Yoga Peeth, 23 June 2021 6:19 pm 🗸



Message

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कोरोना काल में योग के महत्व पर किया मंथन

जनगण संघादराता, भीनगर गढवातः णहणाल केंद्रीय विस्वविद्यालय को कुलपति हो, अञ्चपूर्ण जीटियाल,ने बजा कि कोरोता महामारी से बचाव को लेकर शरीर को रोग प्रतिरोधक क्षमता बहाने के सिए खेग और प्राणमां अहत प्रधावी होने के साथ ही साथकारी भी हैं। ग्रे. मीटियाल ने चुधावल को गढावाल केंग्रीय विश्वविद्यालन अप्रेनगर के योग विष्यान की और से योग योठ ऋषिकेश स्थित विश्वविद्यालय धुम्सटेशन योग सेंटर के संहयोग से तीन दिवसीय आंतरराष्ट्रीय योग वेबिनार का उद्धाटन करते हुए यह ৰাণ করী।

रामग्र स्थास्थ्य को लेकर 'हॉलेंस्टिक हेल्स यू योगा इयुरिंग करेथिड 19' विषय धर आयोजित अंतरराष्ट्रीय वेबिन्स में पहले दिन देश-बिदेश के 100 से ज्याच प्रतिधागी जुहै। वर्षुआल माध्यम झे



गटवान सिरि के रोग विधान की और में आवेतित अंसरसदीय येंग वेंब्रेनन में आवोजकों का अध्यस असने के साथ प्रतिमान करती इटनी के जिन्सीलिक शहर की चेन लिकिका टिनिक सिर्डड + अवस्था

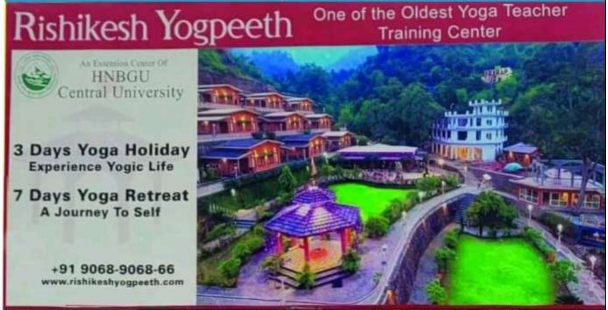
प्रतियन 25 जून तक दोखर एक बजे को पांच करते हुए कुलपति जो. नीटियाल से अपग्रम हीन बजे तक वैवितार का ने कहा कि भारत ने पूरे विसय को यौग संग्रालन किया जाएया। योग को विभिन्न जैसा अनुषम् उपरार देकर संपूर्व मानव विधाओं से शरीर की मिलने वाले लाघों

य के लिए देखि 10000 0000

भी किया है। अंतरराष्ट्रीय वेश्विन्तर के प्रथम रिसोर्स पर्सन और हिमाचल विवि बोग विभाग के पूर्व अध्यक्ष हो, जोती शर्मा ने बोगिक जीवन जीते हुए कोरोना काल में स्वस्थ रहने को लेकर व्याख्यान दिया। दूसरे रिसोर्स पर्सन और वैसिणिक विनि राजस्थान के प्रे. नरेंद्र कुमार ने 122 योग की विभिन्म कलाओं का आकर्षक प्रदर्शन किया। प्रो. नरेंद्र ने शरीर की रोग -प्रतिरोधक क्षमता बढाने में योग के महत्व -के बारे में बताया। गढवाल केंद्रीय विवि के योग विभाग की अध्यक्ष हा. अनुजा ीवन ने वेबिनार के उद्देश्यों पर विस्तार से 111 प्रकास प्राला। आयोजक सचित्र और 57 गडवाल केंद्रीय विवि योग विष्णग के 122 मरम प्रशिक्षक हा, विमोर मीटियाल मे 121

ACHOIRS

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विश्वविद्यालय के विस्तार केंद, ऋषिकेश योग पीठ में आयोजित योग प्रशिक्षण व कार्यशाला



स्पष्टएक्सप्रेस।

उप्राधिकेश (20 जून 2023): विश्वविद्यालय कुलपति मसेदया प्रोफेसर अन्नपूर्णा नौटियाल के मार्गदर्शन में विस्तार केंद्र में योग पर कार्यशाला व प्रशिक्षण संचालित किए जा रहेहैं।

संचालन के लिए प्राकृतिक चिकित्सा एवं योग विभाग के डॉ विनोद नौटियाल को संपर्क अधिकारी, विस्तार केंद्र हेतु अतिरिक्त कार्यभार देकर नियुक्त किया गया। 16 मार्च 2021 से वर्तमान समय जून 2023 तक योग विभाग द्वारा आयोजित एक माह का योग प्रशिक्षण कार्यक्रम

में विदेशी नागरिकों ने बढ चढकर प्रतिभाग किया। जून 2023 तक 49 विभिन्न देशों के नागरिकों और 21 भारतीय नागरिको ने एक माह के योग प्रशिक्षण कार्यकम में प्रतिभाग किया, जिससे लगभग २ लाख रुपए शल्क के रूप में प्राप्त हुए और विश्वविद्यालय के विदेशी नागरिकों के रजिस्टेशन शल्क में लगभग 75 हजार रुपए के रूप में प्राप्त हुए। विश्वविद्यालय के प्राकृतिक चिकित्सा एवं योग विभाग और विस्तार केंद्र के समन्वय से दिनांक 23 से 25 जून २०२१ तक तीन दिवसीय,

द्वितीय अंतरराष्ट्रीय स्तर पर समग्र स्वास्थ्य हेतु योग पर वेबीनार का भी आयोजन किया गया तथा विभिन्न प्रकार की योग चिकित्सा पर कार्यशाला और योग प्रशिक्षण कार्यक्रम प्रत्येक माह डॉ विनोव नौटियाल, डॉ रजनी नौटियाल के माध्यम से एवं विभागाध्यक्ष डॉ अनुजा रावत की देखरेख में आयोजित किए जा रहेहैं।

विश्व के विभिन्न देशों से आए हुए नागरिकों ने इन कार्यशाला व प्रशिक्षण में भाग लेकर विश्वविद्यालय के प्रति अपना आभार और धन्यवाद ज्ञापित

^{किटा।}Scanned by Scanner Go

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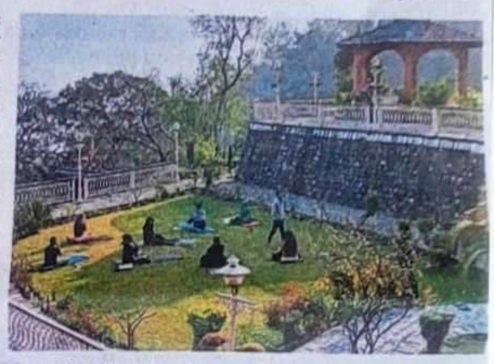
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वावक पदम सिंह = सामारः स्वर्य

योग के प्रति विदेशी छात्रों का रुझान

जागरण संवाददाता. श्रीनगर गढ़तालः गढ़वाल केंद्रीय विश्वविद्यालय श्रीनगर के योग विभाग के विषय विशेषज्ञों से योग प्राणायाम सीखने को लेकर विदेशी छान्न-छान्नाओं की संख्या में अब हर वर्ष इजाफा होता जा रहा है। विदेशी छान्नों के लिए गढ़वाल केंद्रीय विवि से संचालित योग सर्टिफिकेट कार्यक्रम में पिछले वर्ष जहां पांच विदेशी छान्नों ने प्रवेश लिया वहीं अब आगामी अप्रैल महीने से शुरू होने वाले कोर्स के लिए अब तक 22 विदेशी छान्न-छान्नाएं संपर्क कर चुके हैं।

विदेशों के छात्र-छात्राओं को योग प्राणायाम पाठ्यक्रम के प्रति आकर्षित करने के लिए गढ़वाल केंद्रीय विश्वविद्यालय की कुलपति प्रो. अन्नपूर्णा नौटियाल की पहल पर पिछले वर्ष विश्वविद्यालय ने यह कार्ययोजना शुरू की। जिसमें योगपीठ ऋषिकेश के साथ एमओयू कर योग प्राणायाम को लेकर विस्तार केंद्र संचालित किया जा रहा है। कुलपति ने इस केंद्र के लिए गढ़वाल विवि योग विभाग के वरिष्ठ डा. विनोद



गढ़वाल केंद्रीय विवि के योग विभाग के तत्वावचान में योगपीठ ऋषिकेश में संचालित योग केंद्र में डा . विनोद नीटियाल से योग प्रशिक्षण लेते विदेशी छात्र = जागरण

नौटियाल को प्रभारी की जिम्मेदारी दी है। डा. विनोद नौटियाल ने कहा कि योग में अल्पकालिक पाठ्यक्रम संचालन के साथ ही अब विस्तार केंद्र में योग डिप्लोमा पाठ्यक्रम भी शुरू करने पर गंभीरता से विचार किया जा रहा है।

जिससे अधिक से अधिक विदेशी

छात्र-छात्राएं योग में डिप्लोमा हासिल कर सकें। डा. नौटियाल ने कहा कि अप्रैल माह से शुरू हो रहे नए रुज में ऋषिकेश योगपीठ में प्रशिक्षित हो रहे विदेशी छात्रों को गढ़वाल केंद्रीय विवि योग विभाग का भी भ्रमण कराने के साथ ही उन्हें यहां भी प्रशिक्षण दिया जाएगा।



भीनगर । गढवात थिव के जिन्तर कव आपकार में विज्ञान है। छात्रों हेंचु योग शिविर का आवोजन किया गया। जिसमें योजनाव हो, दिनोद नोटियाल द्वारा तीन दिवसीय योग विकित्स शिविर में दिटेशी खाई को योग के नोटियाल द्वारा तीन दिवसीय योग विकित्स शिविर में दिदेशी खाई को योग के गुर सिरवारि गये। जिसमें विदेशी छात्र खात्रओं की विभिन्न रोगों पर किस प्रवार से योग कियाओ द्वारा सिरवात याई जा सकती है यह विस्तार से बताया।

MoU with Texas Tech University, USA

	DR. DEVENDRA NEGI'S ITINERARY								
Sunday, Sept 1	Monday, Sept 2	Tuesday, Sept 3rd	Wednesday, Sept 4th	Thursday, Sept 5th	Friday, Sept 6th				
	Texas Tech Holiday	8:30am Shelley will pick up Dr. Negi	8:15am Dr. Duncan will pick up Dr. Negi	8:15am Shelley will pick up Dr. Negi	8:45am Dr. Duncan will pick up Dr. Negi				
	Labor Day	from Woodrow and deliver to Holden Hall	from Woodrow and deliver to Chem. Eng.	from Woodrow and deliver to Admin Bldg	from Woodrow				
			Chemical Engineering	8:30-8:55am Dr. Galyean, Admin, Room 107					
Dr. Negi's flight arrives	Rest Day		Mahdi Malmali, MERC 212	Shelley will deliver Dr. Negi to Chemistry	-				
@ 7:50pm on	Breakfast is served at	9:00-9:55am			7				
American Airlines	Woodrow B&B	Dean Lindquist & Dr. Akchurin	9:00-10:00am						
\$5995		Holden Hall 202	Dr. Siva Vanapalli, IMSE 202FA	Chemistry					
	(Tentative)	Deaven will escort Dr. Negi to Human	Dr. Duncan will pick up Dr. Negi from	9:00 – 11:00am	9:00 – 11:00am				
Check-In at	Dinner with Dr. Cuikun Lin	Sciences, Room 407	Chem Eng.	- Chemistry 102 conference room	Ranching Heritage Center tour with				
Noodrow House	(Scientist in Dr. Duncan's	10:00-10:30am Dr. Naima Moustaid-Moussa	10:00-10:30am OPEN	(4-6 faculty members to meet	Dr. Duncan				
Bed & Breakfast (806) 793-3330	lab) - Time and Place TBD	Human Sciences, Room 407		with Dr. Negi)					
000) 793-5550		TBD will escort Dr. Negi to ESB 153	10:03-11:00am OPEN						
	Cuikun.lin@ttu.edu, or			Dr. Duncan will pick up Dr. Negi for					
	605-659-4195	10:45-11:45am	10:30-11:00am OPEN	transport to Animal Sciences					
		Dr. Duncan – Tour of CEES		_					
		12:00-1:30pm LUNCH @ Tech Club							
		Drs. Negi		11:15–11:45am					
		Duncan	11:30-1:00pm <mark>LUNCH_TBD</mark>	Dr. Sarturi					
		Snow Lee		Animal & Food Sciences, Room 207 (West side of the United					
		Dr. Duncan will deliver Dr. Negi to HSC		Supermarket Arena (right in front of	11:00am – 12:30pm LUNCH TBD				
				the construction site).					
		2:00 – 4:00pm	1:30-2:15pm	Dr. Sarturi will bring Dr. Negi to ESB					
		Conference Room 2B158 (SVPR office)	Dr. Heppert						
		Drs. Leslie Shen	2:15pm	12-1:30pm LUNCH TBD					
		Min Kang	Dr. Duncan will deliver Dr. Negi to		- 1:00pm				
		Sam Prien	BayerBldg		 TBD will Transport Dr. Negi to Airport 				
			Plant & Soil Science (Room 122F)	1:30-2:15pm OPEN					
			2:30-2:55pm Dr. Jyotsna Sharma	0.00.0.00					
		4:00pm Shelley will pick up Dr. Negi	3:00-3:25pm Dr. Luis Estrella-Herrera	2:30-3:00pm	Dr. Negi departs LBB at 2:29pm on				
			3:30-3:55pm Dr. Haydee Laza	Coffee/Refreshments, ESB 120	American Airlines #5967				
		DINNER at Café J's	4:00pm	3:00-4:30pm	-				
		Dr. Negi, Drs. Duncan and Sobel	Pick up Dr. Negi from Bayer Bldg	Dr. Negi's Colloquium, ESB 120					
			DINNER at Duncan/Sobel Residence w/	DINNER at Triple J's					
			Medical Community (Hendershots, and	Drs. Negi, Duncan, Sobel, Naima					
			Walters)	Moustaid-Moussa, Prien					



When I despair, I remember that all through history the way of truth and love have always won. There have been tyrants and murderers, and for a time, they can seem invincible, but in the end, they always fall. Think of it--always. Mahatma Gandhi



Web Panel Discussion On **Understanding Gandhian way of life: A timeless** lesson for everyone to emulate

Organized by

H.N.B. Garhwal University

In collaboration with

Texas Tech University, Texas, USA

Date: 18th August, 2020 Timings: 7:00 PM (IST)



Patron **Prof. Annpurna Nautival** Vice Chancellor **HNB** Garhwal University



Guest of Honour Prof. Robert V. Duncan Texas Tech University, Lubbock Texas, USA



Chief Guest Prof. Mehraj Uddin Mir Vice Chancellor Central University of Kashmir



Special Guest Prof. Michael San Francisco. Texas Tech University, Lubbock Texas, USA

Celebrating 150 Birth anniversary of Mahatma Gandhi

Abstract of the talk

Special Talk-1 A Man of His Time or a Man for Our Time? Mohandas Gandhi in Africa **Prof. Paul Bjerk**

Mohandas Gandhi came of age in Natal in South Africa, at the turn of the 20th century. He came of age as both a global citizen, or more specifically as a subject of the British Empire, but also, quite literally as the representative of Indians now defined in relation to that empire. His life in Natal grew in parallel to his contemporary global citizen, the Zulu educator John Langalibalele Dube. Considering Gandhi's and Dube's lives in their Natal context, we find illumination for the birth of that dubious signifier: the "modern." Modernity is a 20th century cultural complex, bearing the birthmark of European Enlightenment, but only conceivable as something fundamentally global, and contentious.

Special talk-2 Gandhi's Experiments with Failure **Prof. Costica Bradatan**

Mahatma Gandhi was one of the most successful men of his time. He came to be venerated as a political genius, secular saint, India's Messiah, among other things. At the same time, however, Gandhi had a life-long, intimate relationship with failure. Indeed, in a certain sense, his failures are more revealing and more fascinating than his successes. In my contribution I will examine some moments of Gandhi's complex relationship with failure.

Programme Schedule

Welcome by Prof. D.S. Negi About the program by VC, HNBGU Inaugural address by Chief Guest Address by Guest of Honour Address by Special Guest Special talk-1 Prof. Paul Bjerk Special talk-2 Prof. Costica Bradatan Each special talk will be followed with a 5-7 minutes Q & A Session

Please Note

- Registration is essential for attending the lecture (kindly use the following link for registration)
- Link for Registration: https://forms.gle/T2B8ST88ibn8SV877
- Meeting ID and Password will be provided to the registered participants by 17th August 2020 through E-Mail
- Last date of Registration : 17th August, 2020 till 5:00 PM





LIVE https://www.facebook.com/hnbgu.uttarakhand.3?epa=SEARCH BOX

About the Guest Speakers

Prof. Paul Bierk

Paul Bierk is Professor of History at College of Arts and Sciences, Texas Tech University, Lubbock, Texas. He teaches African History, with a particular emphasis on the continuities across the ruptures of the twentieth century. Dr. Bjerk received a Fulbright Fellowship for dissertation research in Tanzania, and recently received a Fulbright Faculty Fellowship to teach at the University of Iringa, and do research on a second project that will look at the socialist economy of the 1960s and 1970s in Tanzania.

Professor Costica Bradatan

Professor Costica Bradatan is Professor of Humanities in the Honors College at Texas Tech University and an Honorary Research Professor of Philosophy at the University of Queensland, Australia. He has also held faculty appointments at Cornell University, University of Notre Dame, University of Wisconsin-Madison, Miami University, and Arizona State University, as well as at universities in Europe, South America, and Asia.



Head, Department of Chemistry, HNBGU

Organizing Committee

- Dr. Prashant Kandari (Coordinator)
- Dr. Nagendra Rawat (Member)
- Dr. Rakesh Negi (Member)
- Dr. Shweta Verma (Member)
- Dr. Naresh Kumar (Member)

https://us02web.zoom.us/j/85408128617?pwd=dm1NTWVhdW RPeXVpU3k1Z2hQWGR5UT09





Report of the activity conducted by HNBGU and Texas Tech University, USA

A webinar for web panel discussion was conducted on the occasion of 150th Birth anniversary year of Mahatma Gandhi on 18th August 2020. The topic of the discussion was "Understanding Gandhian way of life: A timeless lesson for everyone to emulate." **This webinar was jointly organized by H.N.B. Garhwal University and Texas Tech University, Texas, USA as a part of MoU between both institutions.** The Convener & Moderator of this this discussion, Prof. D.S. Negi welcomed all of the distinguished guests, speakers, and participants. Hon'ble Vice Chancellor, HNBGU, Prof. Annpurna Nautiyal briefly talked about the program and welcomed all of the esteemed delegates from India and abroad. The program started with the inaugural address by Chief Guest Prof. Mehraj Uddin Mir; Hon'ble Vice Chancellor, Central University of Kashmir followed by the address by Guest of Honor, Prof. Robert V. Duncan, Texas Tech University (TTU), Lubbock Texas, USA, and address by Special Guest Prof. Michael San Francisco, TTU, USA.

The first special talk was delivered by Prof. Paul Bjerk, a Professor of History at College of Arts and Sciences, Texas Tech University, Lubbock, Texas. His title of the talk was "A Man of His Time or a Man for Our Time? Mohandas Gandhi in Africa." He talked about that how Gandhi came of age as both a global citizen, or more specifically as a subject of the British Empire, but also, quite literally as the representative of Indians now defined in relation to that empire. Gandhi life in natal grew in parallel to his contemporary global citizen.

The second special talk was delivered by Prof. Costica Bradatan, a Professor of Humanities in the Honors College at Texas Tech University and an Honorary Research Professor of Philosophy at the University of Queensland, Australia. His topic of the discussion was "Gandhi's Experiments with Failure." Prof. Bradatan talked about the success of Gandhi and how he came to be venerated as a political genius, secular saint, India's Messiah, among other things. At the same time, however, Gandhi had a lifelong, intimate relationship with failure. Indeed, in a certain sense, his failures are more revealing and more fascinating than his successes. In his contribution, he examined some moments of Gandhi's complex relationship with failure.

This wonderful web panel discussion on Gandhi was witnessed by around 90 participants, which includes faculties, students, and staff members of various institutions. The recorded videos of the webinar can be found in the following Facebook and YouTube links: <u>https://www.facebook.com/hnbgu.uttarakhand.3?epa=SEARCH_BOX</u> <u>https://www.youtube.com/watch?v=acmca9Tins0&ab_channel=HNBGarhwalUniversity%28offi</u> <u>cialchannel%29</u>

Photos of the Event









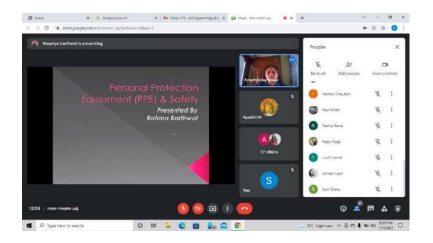


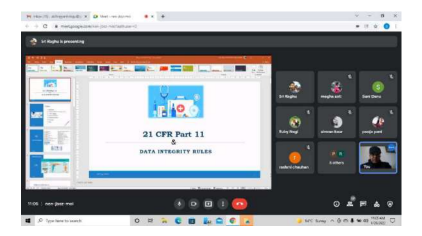


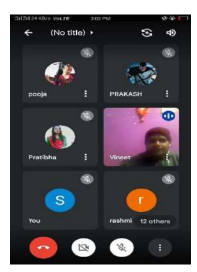
MoA with Uttarakhand State Council for Science & Technology

Student Skill Training Program on 'Quality Control Chemist-Microbiology'

A 3-months Student Skill Training Program on 'Quality Control Chemist-Microbiology' was organized from 21 December 2021 in the Department of Biotechnology, HNB Garhwal University as per the MoA (Memorandum of Agreement) signed between UCOST and HNBGU. The training was conducted under the Skill Vigyan Program sanctioned to the University with Prof. G.K. Joshi as Program Coordinator, by Department of Biotechnology (DBT), Govt. of India and Uttarakhand Council of Science and Technology (UCOST). A total number of 18 participants with graduate degree in biological or related sciences underwent this training. Each trainee received a studentship of Rs. 5000 pm during the training. During the training, various internal and external subject experts as well as industry professionals from India as well as abroad have interacted with trainees and shared their knowledge and skills. In addition, prolonged hands on/practical training has been provided to the trainees in the domain area. On Job Training was also given to the trainees under the actual work environment in industries to gain required expertise and skills. The certification of this course was done by Life Sciences Sector Skill Development Council(LSSSDC). After having completed the training, many of the trainees got placement in industries and academic sectors.





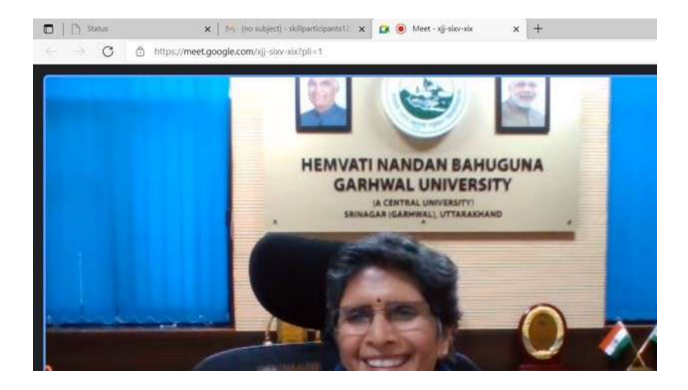














Skill Vigyan Program Batch-1 Quality Control Chemist - Microbiology at Department of Biotechnology HNB Garhwerd University, Singer Garhad Prakash =Staff= _____Attendance = Register=
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Signed MoUs:



हिमाचल प्रदेश केन्द्रीय विश्वविद्यालय CENTRAL UNIVERSITY OF HIMACHAL PRADESH

दूरभाष	1	+91-1892-229330
फैक्स	3	+91-1892-229331
मोबाइल	6	+91-94181-77778
ईमेल	3	kuldeepagnihotri@gmail.com vc.cuhimachal@gmail.com
वेबसाइट	3	www.cuhimachal.ac.in

डॉ. कुलदीप चंद अग्निहोत्री कुलपति

फाइल सं. : 1-8/हि.प्र.के.वि./शैक्षणिक/ 2010/समझौता ज्ञापन/ 3164

दिनांक : 03.06.2020

सेवा में,

प्रो.अन्नपूर्णा नौटियाल कुलपति हेमवती नंदन बहुगुणा गढ़वाल विश्वविद्यालय श्रीनगर गढ़वाल उत्तराखंड

महोदया,

हिमाचल प्रदेश केंद्रीय विश्वविद्यालय की ओर से हार्दिक शुभकामनाएं !

हिमाचल प्रदेश केंद्रीय विश्वविद्यालय प्रस्तावित इंडियन हिमालयन सेंट्रल यूनिवर्सिटी कंसोर्टियम (IHCUC), जिसका एक सदस्य हिमाचल प्रदेश केंद्रीय विश्वविद्यालय भी है, की ओर से कंसोर्टियम के संस्थापक सदस्यों के परामर्श से विश्वविद्यालय अनुदान आयोग तथा अन्य एजेंसियों में प्रस्तावों को जमा कराने तथा इस कंसोर्टियम को बनाने हेतु नीति आयोग तथा/अथवा अन्य एजेंसियों के समक्ष प्रतिनिधित्व करने के लिए कुलपति, हेमवती नंदन बहुगुणा गढ़वाल विश्वविद्यालय को प्राधिकृत करते हुए संलग्न आशय पत्र पर अपनी सहमति प्रदान करता है।

प्रो. (डॉ.) कुलदीप चंद अग्निहोत्री कुलपति, हिमाचल प्रदेश केंद्रीय विश्वविद्यालय

Ziana: Zalyiz

संलग्नक

(फाइल सं. : 1-8/हि.प्र.के.वि./शैक्षणिक/ 2010/समझौता ज्ञापन/ दिनांक : 03.06.2020)

आशय पत्र (Letter of Intent)

भारत के हिमालयी क्षेत्र में स्थित निम्नलिखित 13 केंद्रीय विश्वविद्यालय और सीएसआइआर-आईएचबीटी, पालमपुर, हिमाचल प्रदेश द्वारा आपसी सहयोग से "इंडियन हिमालयन सेंट्रल यूनिवर्सिटीज कंसोर्टियम" (IHCUC) नाम से, जिसे आगे कंसोर्टियम के रूप में संदर्भित किया गया है, एक कंसोर्टियम गठित करने का आशय है, जिसकी रूपरेखा और उद्देश्य नीचे प्रस्तुत हैं । निम्नोक्त विश्वविद्यालय और सीएसआईआर-आईएचबीटी पालमपुर दिए गए उद्देश्यों की पूर्ति के लिए एक संयुक्त समझौता ज्ञापन करने का भी आशय रखते हैं।

- 1. असम विश्वविद्यालय, सिलचर
- 2. हिमाचल प्रदेश केंद्रीय विश्वविद्यालय धर्मशाला
- 3. जम्मू केंद्रीय विश्वविद्यालय
- 4. कश्मीर केंद्रीय विश्वविद्यालय
- 5. सीएसआइआर –आईएचबीटी, पालमपुर हिमाचल प्रदेश
- 6. हेमवती नंदन बहुगुणा गढ़वाल विश्वविद्यालय
- 7. मणिपुर विश्वविद्यालय
- 8. मिजोरम विश्वविद्यालय
- 9. नागालैंड विद्यालय
- 10. नॉर्थ ईस्टर्न हिल यूनिवर्सिटी
- 11. राजीव गांधी विश्वविद्यालय, ईटानगर
- 12. सिक्किम विश्वविद्यालय, गंगटोक
- 13. तेजपुर विश्वविद्यालय
- 14. त्रिपुरा विश्वविद्यालय

रूपरेखाः

उपर्युक्त केंद्रीय विश्वविद्यालय और सीएसआइआर-आईएचबीटी, पालमपुर कंसोर्टियम के संस्थापक सदस्य होंगे ।

उद्देश्य:

- कंसोर्टियम के सदस्यों के मध्य आपसी विचारों का आदान-प्रदान बढ़ाना, विशेषज्ञता साझा करना और वृत्तिक उत्कृष्टता को बढ़ावा देना
- कंसोर्टियम अथवा अन्य एजेंसियों द्वारा वित्तपोषण के लिए संयुक्त प्रस्ताव तैयार एवं जमा कराना
- भारतीय हिमालयी क्षेत्र के सामाजिक-आर्थिक तथा पर्यावरणीय मुद्दों में शोध को बढ़ावा देना
- भारतीय हिमालयी क्षेत्र से जुड़े मुद्दों पर, जहां भी आवश्यक हो, अन्य अतिरिक्त साझेदारों सहित संयुक्त रूप से क्षेत्रीय/ राष्ट्रीय कार्यशाला/ सेमिनार/ सम्मेलनों का आयोजन

हिमाचल प्रदेश केंद्रीय विश्वविद्यालय उपर्युक्त 14 संस्थापक सदस्यों द्वारा संयुक्त रूप से दिए गए इस आशय पत्र के लिए अपनी सहमति प्रदान करता है ।

प्रो. (डॉ.) कुलदीप चंद अग्निहोत्री कुलपति, हिमाचल प्रदेश केंद्रीय विश्वविद्यालय

स्थान: धर्मशाला दिनांक: 3 जून 2020



सी.एस.आई.आर-हिमालय जैवसंपदा प्रौद्योगिकी संस्थान (वैज्ञानिक तथा औद्योगिक अनुसंधान परिषद्) पो॰ बॉ॰ न॰ 6 पालमपुर - 176 061 (हि.प्र.) भारत CSIR-INSTITUTE OF HIMALAYAN BIORESOURCE TECHNOLOGY (Council of Scientific & Industrial Research) Post Box No. 6 Palampur (H.P.) 176 061 INDIA



डाँ. सजय कुमार, एक.एन.ए.एस.सी.,एक.एन.ए.ए.एस.सी.,एफ.सी.आई.एस.आई.

निदेशक

Dr. Sanjay Kumar, FNASC, FNAASC, FCISI

Director

Letter of Intent

Listed below 13 Central Universities and CSIR-IHBT, Palampur (HP) in Himalayan Region of India intent to collaborate to form a consortium designated as "Indian Himalayan Central Universities Consortium (IHCUC) herein referred as "Consortium", with framework and objectives as given under. These Universities and CSIR-IHBT, Palampur also intend to participate in a joint Memorandum of Understanding (MoU) in fulfilling the stated objectives.

- 1. Assam University, Silchar
- 2. Central University of Himachal Pradesh
- 3. Central University of Jammu
- 4. Central University of Kashmir
- 5. CSIR-Institute of Himalayan Bioresource Technology, Palampur (HP)
- 6. HNB Garhwal University
- 7. Manipur University
- 8. Mizoram University
- 9. Nagaland University
- 10. North Eastern Hill University
- 11. Rajiv Gandhi University, Itanagar
- 12. Sikkim University, Gangtok
- 13. Tezpur University
- 14. Tripura University

Framework:

 The above stated Central Universities and CSIR-IHBT, Palampur will be the founding members of the 'Consortium'.

Objectives:

- Promote exchange of ideas, share expertise and professional excellence among members of Consortium
- To develop and submit joint proposals for funding by the Consortium or other agencies
- To promote research on socio-economic and environmental issues in Indian Himalayas
- Jointly organize, also with other additional partners, wherever required, regional/national workshops/seminars/ conferences on issues of Indian Himalayan Region.

CSIR-IHBT, Palampur gives its consent to the joint intent of 14 founding members as given above.

Place: Palampur Date: 09 June 2020

(Sanjay Kumar)

प्रो॰ अशोक ऐमा कुलपति Prof. Ashok Aima Vice Chancellor



जम्मू केंद्रीय विश्वविद्यालय Central University of Jammu टेलिफैक्स/Telefax: 01923-249634(0) ई-मेल /e-mail: vicechancellor.cuj@gmail.com No. ८७५/ ४८/ ९५(२०१९/१८/१२५ Dated: 02-06-2020

Central University Jammu gives its consent to the Vice-Chancellor HNB Garhwal University to represent the proposed Indian Himalayan Central University Consortium (IHCUC) to Niti Aayog and/or other agencies for submitting proposals to the UGC or other agencies and formalsing the said Consortium in consultation with the founding members.

(Prof. Ashok Aima) 2/6/2020

Prof. Annpurna Nautiyal Vice-Chancellor, HNB Garhwal University, Srinagar Garhwal, Uttarakhand प्रो॰ अशोक ऐमा कुलपति Prof. Ashok Aima Vice Chancellor



जम्मू केंद्रीय विश्वविद्यालय Central University of Jammu टेलिफैक्स/Telefax: 01923-249634(0) ई-मेल /e-mail: vicechancellor.cuj@gmail.com

Letter of Intent

Listed below 13 Central Universities and CSIR-IHBT, Palampur (HP)in Himalayan Region of India intent to collaborate to form a consortium designated as "Indian Himalayan Central Universities Consortium (IHCUC) herein referred as "Consortium", with framework and objectives as given under. These Universities and CSIR IHBT, Palampur also intend to participate in a joint Memorandum of Understanding (MoU) in fulfilling the stated objectives.

- 1. Assam University, Silchar
- 2. Central University of Himachal Pradesh
- 3. Central University of Jammu
- 4. Central University of Kashmir
- 5. CSIR-IHBT Palampur (HP)
- 6. HNB Gharwal University
- 7. Manipur University
- 8. Mizoram University
- 9. Nagaland University
- 10.North Eastern Hill University
- 11. Rajiv Gandhi University, Itanagar
- 12. Sikkim University, Gangtok
- 13. Tezpur University
- 14. Tripura University

Framework:

• The above stated Central Universities and CSIR, HBT, Palampur will be the founding members of the 'Consortium'.

Objectives:

- Promote exchange of ideas, share expertise and professional excellence among members of Consortium
- To develop and submit joint proposals for funding by the Consortium or other agencies
- To promote research on socio-economic and environmental issues in Indian Himalayas

Ashele aima

बागला (राया-सूचानी), जिला सांबा-181143, जम्मू (जम्मू एवं कश्मीर) Bagla (Rahya- Suchani), District Samba-181143, Jammu (Jammu & Kashmir)

-1-





जम्मू केंद्रीय विश्वविद्यालय Central University of Jammu टेलिफैक्स/Telefax: 01923-249634(0) ई-मेल /e-mail: vicechancellor.cuj@gmail.com

• Jointly organize, also with other additional partners, wherever required, regional/national workshops/seminars/ conferences on issues of Indian Himalayan Region.

Central University Jammu gives its consent to the joint intent of 14 founding members as given above.

Place: Janune Date: 2/6/2020

(Prof. Ashok Aima) Vice-Chancellor



कश्मीर केन्द्रीय विश्वविद्यालय Central University of Kashmir

केन्द्रीय विश्वविद्यालय (संज्ञोधन) अधिनियम, २००९ के अंतर्गत स्यापित Established Under Central Universities (Amendment) Act, 2009



Prof. Mehraj Uddin Mir Vice-Chancellor NO: cuk ma/ve-26/2015/523 Dt: 18.06.2020

Letter of Consent

Central University of Kashmir gives its consent to the Vice Chancellor HNB Garhwal University to represent the proposed Indian Himalayan Consortium (IHUCC) to Nitiayog and/or other agencies for submitting proposals to UGC or other agencies & formalising the said consortium in consultation with the funding members

ProfWC केन्द्रीय दिन्द्रीव आपन्धी संस्थानार CHTRAL UNIVERSITY OF KASHI





कश्मीर केन्द्रीय विश्वविद्यालय

Central University of Kashmir

केन्द्रीय विश्वविद्यालय (संशोधन) अधिनियम, २००९ के अंतर्गत स्थापित Established Under Central Universities (Amendment) Act, 2009



Prof. Mehraj Uddin Mir Vice-Chancellor Letter of Intent

No: eulins/ve-26/2015/50 Df: 18.06.2020

Listed below 13 Central Universities and CSIR-IHBT, Palmpur (HP) in Himalayan Range of India intent to collaborate to form a consortium designated as "INDIAN HIMALAYAN CENTRAL UNIVERSITIES CONSORTIUM (IHUCC) herein referred as consortium, with frameworks and objectives as given under these universities and CSIR (IHBT), Palmpur also intent to participate in a join memorandum of understanding (MoU) in fulfilling the stated objectives.

- 1. Assam University, Silchar
- 2. Central university of Himachal Pradesh
- 3. Central university of Jammu
- 4. Central University of Kashmir
- 5. CSIR-IHBT Palmpur (HP)
- 6. HNB Garhwal University UT
- 7. Manipur University
- 8. Mizoram University
- 9. Nagaland University
- 10. North Eastern Hill University (NEHU), Meghalaya.
- 11. Rajiv Gandhi University Itanagar
- 12. Central Sikkim University, Gangtok
- 13. Tezpur University
- 14. Tripura University

Framework

The above stated universities and CSIR, HBT, Palmpur will be the founding members of the 'Consortium'.

Objectives

- Promote exchange of ideas share expertise & professional excellence among members of consortium,
- To develop & submit joint proposals for funding by the Consortium or other agencies.
- To promote research on socio-economic & environment issues in Indian Himalayas.
- Jointly organise, also with other additional partners wherever required, Regional/National Workshops/Seminars conferences issues of Indian Himalayan Region.

Central University of Kashmir, gives its consent to the joint intent of 14 founding members as given above.

Place In Ganderbal

Date 18.06.2020

ProMUEHAHDELLYOR संस्ट्रीयीविष्याधिकारय कशजीर CENTRAL UNIVERSITY OF KASHE



प्रशासनिक परिसर : नवगाम, बाय—पास निकट पुहरू क्रॉसिंग, श्रीनगर — 190015 (जे एण्ड के) Admin. Campus : Nowgam, By Pass Near Puhroo Crossing, Srinagar - 190015 (J&K) Phone : 0194-2315271, email : vc@cukashmir.ac.in ; website : www.cukashmir.ac.in

Scanned with CamScanner

6 माइल, सामदुर, तादोंग -737102 गंगटोक, सिक्किम, भारत फोन-03592-251067, 251073 वेबसाइट - <u>www.cus.ac.in</u>



6th Mile, Samdur, Tadong -737102 Gangtok, Sikkim, India Ph. 03592-251067, 251073 Website: www.cus.ac.in

Date: 2/06/2020

(भारत के संसद के अधिनियम द्वारा वर्ष 2007 में स्थापित और नैक (एनएएसी) द्वारा वर्ष 2015 में प्रत्यायित केंद्रीय विश्वविद्यालय) (A central university established by an Act of Parliament of India in 2007 and accredited by NAAC in 2015)

The Vice-Chancellor

Sa/2019-26/126

Τo,

Prof. Annapurn Nautiyal Vice-Chancellor HNB Garhwal University Srinagar Garhwal Uttarakhand

Dear Madam,

Greetings from Sikkim University!

Sikkim University, Gangtok gives its consent to the Vice-Chancellor HNB Garhwal University to represent the proposed Indian Himalayan Central University Consortium (IHCUC) to Niti Aayog and/or other agencies for submitting proposals to the UGC or other agencies and formalsing the said Consortium in consultation with the founding members.

Yours sincerely,

(Prof. Avinash Khare) Vice-Chancellor

कुलपति Vice-Chancellor रिकिकम विश्वविद्धालय Sikkim University

Letter of Intent

Listed below 13 Central Universities and CSIR-IHBT, Palampur (HP) in Himalayan Region of India intent to collaborate to form a consortium designated as "Indian Himalayan Central Universities Consortium (IHCUC) herein referred as "Consortium", with framework and objectives as given under. These Universities and CSIR IHBT, Palampur also intend to participate in a joint Memorandum of Understanding (MoU) in fulfilling the stated objectives.

- 1. Assam University, Silchar
- 2. Central University of Himachal Pradesh
- 3. Central University of Jammu
- 4. Central University of Kashmir
- 5. CSIR-IHBT Palampur (HP)
- 6. HNB Gharwal University
- 7. Manipur University
- 8. Mizoram University
- 9. Nagaland University
- 10.North Eastern Hill University
- 11. Rajiv Gandhi University, Itanagar
- 12. Sikkim University, Gangtok
- 13. Tezpur University

14. Tripura University

Framework:

• The above stated Central Universities and CSIR, HBT, Palampur will be the founding members of the 'Cosortium'.

Objectives:

- Promote exchange of ideas, share expertise and professional excellence among members of Consortium
- To develop and submit joint proposals for funding by the Consortium or other agencies
- To promote research on socio-economic and environmental issues in Indian Himalayas
- Jointly organize, also with other additional partners, wherever required, regional/national workshops/seminars/ conferences on issues of Indian Himalayan Region.

Sikkim University, Gangtok gives its consent to the joint intent of 14 founding members as given above.

(Prof. Avinash Khare)

Vice-Chancellor

कुलपति Vice-Chancellor रिरकि**कन विश्वविद्धालय** Sikkim University

Place: Gaugtok Date: 2/06/2020

प्रोफेसर दिलीप चन्द्र नाथ



असम विश्वविद्यालय (एक केंन्द्रीय विश्वविद्यालय) सिलचर 788011 असम, भारत

ASSAM UNIVERSITY (A Central University) Silchar 788011, Assam, India

F.: 7-45/VCS/2020/ Date : 2.6.2020

Letter of Intent

Listed below 13 Central Universities and CSIR-IHBT, Palampur (HP) in Himalayan Region of India intent to collaborate to form a consortium designated as "Indian Himalayan Central Universities Consortium (IHCUC) herein referred as "Consortium", with framework and objectives as given under. These Universities and CSIR IHBT, Palampur also intend to participate in a joint Memorandum of Understanding (MoU) in fulfilling the stated objectives.

- 1. Assam University, Silchar
- 2. Central University of Himachal Pradesh
- Central University of Jammu
- 4. Central University of Kashmir
- 5. CSIR-IHBT Palampur (HP)
- HNB Gharwal University
- 7. Manipur University
- 8. Mizoram University
- 9. Nagaland University
- 10. North Eastern Hill University
- 11. Rajiv Gandhi University, Itanagar
- 12. Sikkim University, Gangtok
- 13. Tezpur University
- 14. Tripura University

Framework:

The above stated Central Universities and CSIR, HBT, Palampur will be the founding • members of the 'Consortium'.

Objectives:

- Promote exchange of ideas, share expertise and professional excellence among members of Consortium
- To develop and submit joint proposals for funding by the Consortium or other agencies
- To promote research on socio-economic and environmental issues in Indian Himalayas
- Jointly organize, also with other additional partners, wherever required, regional/national ۰ workshops/seminars/ conferences on issues of Indian Himalayan Region.

Assam University, Silchar gives its consent to the joint intent of 14 founding members as given above.

Place: Silchar Date: 4/6/2020

(Prof. Dilip Ch Nath) Vice-Change

Prof. Dilip Chandra Nath



Tezpur University

(A Central University) Tezpur - 784028 Assam, India

- Visitor's Best University Award 2016
- NIRF India Rankings 2016 : 05
- NIRF 2019, Univ : 29, Overall : 48

Fax : 03712-267006 (0) *E-mail* : vc@tezu.ernet.in

Phone: 03712-267003 (0)

Accredited with 'A' Grade by NAAC
THE Asia Rankings 2018 : 100

• THE World Best Small Univ. 2018 - 20th

- QS Ranking 2020 : IND 35, BRICS 146
- QS Ranking 2020 : IND 35, BRICS 146

16 June 2020

Professor V. K. Jain, D.Phil (UK) Vice-Chancellor

TU/VC/1(I)/2020/128

To,

Prof. Annpurna Nautiyal Vice-Chancellor HNB Garhwal University Srinagar, Garhwal Uttarakhand, India, 246174

Dear Madam,

Greetings from Tezpur University!

Tezpur University, Tezpur gives its consent to the Vice Chancellor HNB Garhwal University to represent the proposed Indian Himalayan Central University Consortium (IHCUC) to Niti Aayog and /or other agencies for submitting proposals to the UGC or other agencies and formalizing the said Consortium in Consultation with the founding members.

Yours sincerely,

(V. K. Jain) Vice- Chancellor



(A Central University) Tezpur - 784028 Assam. India

Professor V. K. Jain, D.Phil (UK) Vice-Chancellor

- **Tezpur University** Visitor's Best University Award 2016
 - NIRF India Rankings 2016 : 05
 - NIRF 2019, Univ : 29, Overall : 48

Phone: 03712-267003 (0) Fax : 03712-267006 (0) E-mail: vc@tezu.ernet.in

- Accredited with 'A' Grade by NAAC • THE Asia Rankings 2018 : 100
- THE World Best Small Univ. 2018 20th
- QS Ranking 2020 : IND 35, BRICS 146

Letter of Intent

Listed below 13 Central Universities and CSIR- IHBT, Palampur (HP) in Himalayan Region of India intent to collaborate to form a consortium designated as "Indian Himalayan Central Universities Consortium (IHCUC) here in referred as "Consortium", with framework and objectives as given under. These Universities and CSIR IHBT, Palampur also intend to participate in a Joint Memorandum of Understanding (MoU) in fulfilling the stated objectives.

- 1) Assam University, Silchar
- 2) **Central University of Himachal Pradesh**
- 3) Central University of Jammu
- 4) Central University of Kashmir
- 5) CSIR- IHBT Palampur (HP)
- 6) HNB Gharwal University
- 7) Manipur University
- 8) **Mizoram University**
- 9) Nagaland University
- 10) North eastern Hill University
- 11) Rajiv Gandhi University, Itanagar
- Sikkim University, Gangtok 12)
- **Tezpur University** 13)
- 14) **Tripura University**

Frame work:

The above stated Central Universities and CSIR, HBT, Palampur will be the founding members of the 'Consortium'.

Objectives:

- Promote exchange of ideas, share expertise and professional excellence among members of consortium
- To develop and submit joint proposals of funding by the Consortium or other agencies.
- To promote research on socio- economic and environmental issues in Indian Himalayas
- Jointly organize, also with other additional partners, wherever required, regional/national workshops/ seminars/ conferences on issues of Indian Himalavan Region.

Tezpur University, Tezpur gives its consent to the Joint intent of 14 founding members as given above.

(V. K. Jain)

Place : Tezpur University Date : 16 June 2020

Vice Chancellor



पूवोत्तर पर्वतीय विश्वविद्यालय ५० प॰ विवि॰ परिसर, शिलांग-७९३०२२ (मेघालय) Phone : Grams : NEHU

North-Eastern Hill University

NEHU Campus, Shillong - 793 022 (Meghalaya)

Prof. S.K. Srívastava Více-Chancellor Telephone Nos 0364-2550101/2721003/4(0) Fax No.0364-2550076/2551634 e-maíl: <u>vonehu@nehu.ac.ín</u>

> No.F.28-6/Acad/AC/2019/66 June 18, 2020

Letter of Consent

The North-Eastern Hill University, Shillong gives its consent to the Vice-Chancellor, HNB Garhwal University, Srinagar (Garhwal) to represent the proposed Indian Himalayan Central Universities' Consortium (IHCUC) to NITI Aayog and / any other agency to submit proposal to the UGC / any other agency, and formalising the same on behalf of the Consortium of the Founding Members.

(Sri Krishna Srivastava)

कुलपति/Vice-Chancellor पूर्वोत्तर पर्वतीय विश्वविद्यालय North Eastern Hill University शिलांग/Shillong

राजीव गाँधी विश्वविद्यालय

केंद्रीय विश्वविद्यालय

रोनोहिल्स,दोइमुख- ७९१ ११२ अरुणाचल प्रदेश, भारत

Rajiv Gandhi University

Central University Rono Hills, Doimukh – 791 112 Arunachal Pradesh, India

No.RGU/VCS/CHU/2020 June 2, 2020

То

प्रो. साकेत कुशवाहा

Vice Chancellor

Prof. Saket Kushwaha

कुलपति

Prof. Annpurna Nautiyal Vice-Chancellor HNB Garhwal University Srinagar Garhwal Uttarakhand

Dear Madam,

Greetings from Rajiv Gandhi University!

Rajiv Gandhi University, Arunachal Pradesh gives its consent to the Vice-Chancellor HNB Garhwal University to represent the proposed Indian Himalayan Central University Consortium (IHCUC) to Niti Aayog and/or other agencies for submitting proposals to the UGC or other agencies and formalsing the said Consortium in consultation with the founding members.

Yours sincerely,

Saket Kushwaha



राजीव गाँधी विश्वविदयालय

केंद्रीय विश्वविदयालय

रोनोहिल्स,दोइमुख- ७९१ ११२ अरुणाचल प्रदेश, भारत

Rajiv Gandhi University

Central University Rono Hills, Doimukh – 791 112 Arunachal Pradesh, India

No.RGU/VCS/CHU/2020 June 2, 2020

Letter of Intent

Listed below 13 Central Universities and CSIR-IHBT, Palampur (HP)in Himalayan Region of India intent to collaborate to form a consortium designated as "Indian Himalayan Central Universities Consortium (IHCUC) herein referred as "Consortium", with framework and objectives as given under. These Universities and CSIR IHBT, Palampur also intend to participate in a joint Memorandum of Understanding (MoU) in fulfilling the stated objectives.

- 1. Assam University, Silchar
- 2. Central University of Himachal Pradesh
- 3. Central University of Jammu
- 4. Central University of Kashmir
- 5. CSIR-IHBT Palampur (HP)
- 6. HNB Gharwal University
- 7. Manipur University
- 8. Mizoram University
- 9. Nagaland University
- 10.North Eastern Hill University
- 11. Rajiv Gandhi University, Itanagar
- 12. Sikkim University, Gangtok
- 13. Tezpur University
- 14. Tripura University

Framework:

प्रो. साकेत कुशवाहा

Vice Chancellor

Prof. Saket Kushwaha

कलपति

• The above stated Central Universities and CSIR, HBT, Palampur will be the founding members of the 'Consortium'.

Objectives:

- Promote exchange of ideas, share expertise and professional excellence among members of Consortium
- To develop and submit joint proposals for funding by the Consortium or other agencies
- To promote research on socio-economic and environmental issues in Indian Himalayas
- Jointly organize, also with other additional partners, wherever required, regional/national workshops/seminars/ conferences on issues of Indian Himalayan Region.

Rajiv Gandhi University, Rono Hills, Arunachal Pradesh gives its consent to the joint intent of 14 founding members as given above.

Place: Rono Hills Date :02.06.2020 Saket Kushwaha Vice-Chancellor







Speed Post

(संसद द्वारा पारित अधिनियम 1989, क्रमांक 35 के अंतर्गत स्थापित केन्द्रीय विश्वविद्यालय) (A Central University Established by the Act of Parliament No.35 of 1989)

मुख्यालय : लुमामी, जिला : जुन्हेबोटो (नागालैण्ड), पिन कोड - 798 627

Headquarters: Lumami, District: Zunheboto (Nagaland), Pin Code - 798 627

No. NUL/RDC-26/IHCUC/2020 - 365

Dated: 03.06.2020

То

Prof. Annpurna Nautiyal Vice-Chancellor HNB Garhwal University Srinagar Garhwal Uttarakhand - 246174

Sub: Indian Himalayan Central University Consortium (IHCUC) - reg.

Madam,

Nagaland University, Lumami gives its consent to the Vice-Chancellor HNB Garhwal University to represent the proposed Indian Himalayan Central University Consortium (IHCUC) to Niti Aayog and/or other agencies for submitting proposals to the UGC or other agencies and formalising the said Consortium in consultation with the founding members.

Yours faithfully,

a

(PARDESHI LAL) Vice-Chancellor



विश्वविद्यालय UNIVERSITY

(संसद द्वारा पारित अधिनियम 1989, क्रमांक 35 के अंतर्गत स्थापित केन्द्रीय विश्वविद्यालय) (A Central University Established by the Act of Parliament No.35 of 1989)

मुख्यालय : लुमामी, जिला : जुन्हेबोटो (नागालैण्ड), पिन कोड - 798 627

Headquarters: Lumami, District: Zunheboto (Nagaland), Pin Code - 798 627

No. NUL/RDC-26/IHCUC/2020 - 366

Dated: 03.06.2020

Letter of Intent

Listed below 13 Central Universities and CSIR-IHBT, Palampur (HP) in Himalayan Region of India intent to collaborate to form a consortium designated as "Indian Himalayan Central Universities Consortium (IHCUC) herein referred as "Consortium", with framework and objectives as given under. These Universities and CSIR IHBT, Palampur also intend to participate in a joint Memorandum of Understanding (MoU) in fulfilling the stated objectives.

- 1. Assam University, Silchar
- 2. Central University of Himachal Pradesh
- 3. Central University of Jammu
- 4. Central University of Kashmir
- 5. CSIR-IHBT Palampur (HP)
- 6. HNB Gharwal University
- 7. Manipur University
- 8. Mizoram University
- 9. Nagaland University
- 10. North Eastern Hill University
- 11. Rajiv Gandhi University, Itanagar
- 12. Sikkim University, Gangtok
- 13. Tezpur University
- 14. Tripura University

Framework:

 The above stated Central Universities and CSIR, HBT, Palampur will be the founding members of the 'Consortium'.

Objectives:

- Promote exchange of ideas, share expertise and professional excellence among members of Consortium
- To develop and submit joint proposals for funding by the Consortium or other agencies
- To promote research on socio-economic and environmental issues in Indian Himalayas
- Jointly organize, also with other additional partners, wherever required, regional/national workshops/seminars/ conferences on issues of Indian Himalayan Region.

Nagaland University, Lumami gives its consent to the joint intent of 14 founding members as given above.

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(PARDESHI LAL) Vice-Chancellor



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No MU/VC/Admtr/2018/1

Dated 19 May 2020

Letter of Intent

Listed below 13 Central Universities and CSIR-IHBT, Palampur Range of India intent to collaborate to form a consortium designated as "Indian Himalayan Central Universities Consortium (IHUCC), herein referred as consortium, with framework and objectives as given under these universities and CSIR (IHBT), Palampur also intent to participate in a join Memorandum of Understanding (MoU) in fulfilling the stated objectives.

- 1. Assam University, Silchar
- 2. Central University of Himachal Pradesh
- 3. Central University of Jammu
- 4. Central university of Kashmir
- 5. CSIR-IHBT Palampur (HP)
- 6. HNB Garhwal University
- 7. Manipur University
- 8. Manipur University
- 9. Nagaland University
- 10. North Eastern Hill University(NEHU), Meghalaya
- 11. Rajiv Gandhi University, Itanagar
- 12. Central Sikkim University, Gangtok
- 13. Tezpur University
- 14. Tripura University

Framework

The above stated universities and CSIR-IHBT Palampur will be the founding members of the 'Consortium'.

Objectives

- Promote exchange of ideas share expertise & professional excellence among members of consortium
- To develop & submit joint proposal for funding by the consortium or other agencies.
- To promote research on socio-economic & environment issues in Indian Himalayas
- Jointly organise, also with other additional partners wherever required, Regional/National Workshops/Seminars Conferences issues of Indian Himalayan Region.

Manipur University gives its consent to joint Intent of 14 founding members as stated above.

Place Imphal Date 19-06-2020

(Jarnail Singh) Vice Chancellor

र्मा केंद्र भूम केंद्र भूमी जम ॥ २००९२४ - ह्वेमद भूमी प्रजा

Phone - Office (0385) EPABX : 2435276/2435055 Fax Modem : (0385) 2435145 : Telegrams; Manvarsity



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No MU/VC/Admtr/2018/1

Dated 19 May 2020

Letter of Consent

Manipur University gives its consent to the Vice Chancellor HNB Grahwal University to represent the proposed Indian Himalayan Consortium (IHUCC) to Nitiayog and /or other agencies for submitting proposal to UGC or other agencies & formalising the said consortium in consultation with the funding members.

(Jarnail Singh) 19-06-2020 Vice Chancellor



मिजोरम विश्वविद्यालय MIZORAM UNIVERSITY AIZAWL - 796004

Post Box No.190 Gram : MZU Phone:0389-2330654 Fax : 0389-2330834 Email:registrar@mzu.edu.in Website : www.mzu.edu.in

A CENTRAL UNIVERSITY CREATED BY AN ACT OF PARLIAMENT OF INDIA भारतीय संसद दूारा पारित एफ्ट के अर्न्तगत रथापित एक केन्दीय विश्वविद्यालय

To,

Prof. Annpurna Nautiyal Vice-Chancellor HNB Garhwal University Srinagar Garhwal Uttarakhand

Dear Madam,

Greetings from Mizoram University!

Mizoram University, Aizawl gives its consent to the Vice-Chancellor HNB Garhwal University to represent the proposed Indian Himalayan Central University Consortium (IHCUC) to Niti Aayog and/or other agencies for submitting proposals to the UGC or other agencies and formalsing the said Consortium in consultation with the founding members.

Yours sincerely,

15506

(Prof. K. R. S. Sambasiva Rao) Vice-Chancellor

> कुलपति Vice - Chancellor मिज़ोरम विश्वविद्यालय Mizoram University तन्हिल, आइज़ोल Tanhril, Aizawl.

Letter of Intent

Listed below 13 Central Universities and CSIR-IHBT, Palampur (HP) in Himalayan Region of India intent to collaborate to form a consortium designated as "Indian Himalayan Central Universities Consortium (IHCUC) herein referred as "Consortium", with framework and objectives as given under. These Universities and CSIR IHBT, Palampur also intend to participate in a joint Memorandum of Understanding (MoU) in fulfilling the stated objectives.

- 1. Assam University, Silchar
- 2. Central University of Himachal Pradesh
- 3. Central University of Jammu
- 4. Central University of Kashmir
- 5. CSIR-IHBT Palampur (HP)
- 6. HNB Gharwal University
- 7. Manipur University
- 8. Mizoram University
- 9. Nagaland University
- 10.North Eastern Hill University
- 11. Rajiv Gandhi University, Itanagar
- 12. Sikkim University, Gangtok
- 13. Tezpur University

14. Tripura University

Framework:

• The above stated Central Universities and CSIR, HBT, Palampur will be the founding members of the 'Consortium'.

Objectives:

- Promote exchange of ideas, share expertise and professional excellence among members of Consortium
- To develop and submit joint proposals for funding by the Consortium or other agencies
- To promote research on socio-economic and environmental issues in Indian Himalayas
- Jointly organize, also with other additional partners, wherever required, regional/national workshops/seminars/ conferences on issues of Indian Himalayan Region.

Mizoram University, Aizawl gives its consent to the joint intent of 14 founding members as given above.

Place: Aizand Date: 3/6/2020

45DC

(Prof. K. R. S. Sambasiva Rao) Vice-Chancellor

कुलपति Vice - Chancellor मिज़ोरम विश्वविद्यालय Mizoram University तन्हिल, आइज़ोल Tanhril, Aizawi



Suryamaninagar, Agartala, Tripura, INDIA पिन Pin - 799022

237 9024 फैक्स Fax : (0381) 237 4802 237 4803 237 5355 237 4804 e-mail : tuoffice@tripurauniv.in website : www.tripurauniv.in

No. F.TU/VCS/MISC/01/2018

Date: 19th June, 2020

Letter of Consent

Tripura University (A Central University) gives its consent to the Vice-Chancellor HNB Garhwal University to represent the proposed Indian Himalayan Consortium (IHUCC) to Nitiayog and/or other agencies for submitting proposal to UGC or other agencies and formalizing the said consortium with the funding members.

106/25

(Prof. M. K. Singh) कुलपति/Vice-Chancellor(i/c) त्रिपुरा विश्वविद्यालय/Tripura University

त्रिपुरा विश्वविद्यालय TRIPURA UNIVERSITY

(केन्द्रीय विश्वविद्यालय) (A CENTRAL UNIVERSITY)

सूर्यमणिनगर, अगरतला, त्रिपुरा, भारत Suryamaninagar, Agartala, Tripura, INDIA पिन Pin - 799022



फोन Phone: (0381) 237 4801 237 9002 237 9003 237 9004 237 9024 फैक्स Fax : (0381) 237 4802 237 4803 237 5355 237 4804 e-mail : tuoffice@tripurauniv.in website : www.tripurauniv.in

No. F.TU/VCS/MISC/01/2018

Date: 19th June, 2020

Letter of Intent

Listed below 13 Central Universities and CSIR-IHBT, Palmpur (HP) in Himalayan Range of India intent to collaborate to form a consortium designated as "INDIAN HIMALAYAN CENTRAL UNIVERSITIES CONSORTIUM (IHUCC) herein referred as consortium, with frameworks and objectives as given under these universities and CSIR (IHBT), Palmpur also intent to participate in a join memorandum of understanding (MoU) in fulfilling the stated objectives.

- 1. Assam University, Silchar
- 2. Central university of Himachal Pradesh
- 3. Central university of Jammu
- 4, Central University of Kashmir
- 5. CSIR-IHBT Palmpur (HP)
- 6. HNB Garhwal University UT
- 7. Manipur University
- 8. Mizoram University
- 9. Nagaland University
- 10. North Eastern Hill University (NEHU), Meghalaya.
- 11. Rajiv Gandhi University Itanagar
- 12. Central Sikkim University, Gangtok
- 13. Tezpur University
- 14. Tripura University

Framework

The above stated universities and CSIR, HBT, Palmpur will be the founding members of the 'Consortium'.

Objectives

- Promote exchange of ideas share expertise & professional excellence among members of consortium,
- To develop & submit joint proposals for funding by the Consortium or other agencies.
- To promote research on socio-economic & environment issues in Indian Himalayas.
- Jointly organise, also with other additional partners wherever required, Regional/National Workshops/Seminars conferences issues of Indian Himalayan Region.

Tripura University, gives its consent to the joint intent of 14 founding members as given above.

106/200 (Prof. M. K. Singh)

कुलपति/Vice-Chancellor (i/c) निग्रम निश्चनितालम् /Trinura University.





CONTACT INFORMATION

TEXAS TECH UNIVERSITY

College of Agricultural Sciences and Natural Resources

Name: Dr. Glen Ritchie

Email: Glen.ritchie@ttu.edu

Chair, Department of Plant and Soil Sciences

College of Human Sciences

Name: Dr. Nikhil V. Dhurandhar

Email: Nikhil.dhurandhar@ttu.edu

Chair, Department of Nutritional Sciences

Office of International Affairs

Name: Michael Q Johnson

Email: Michael johnson@ttu.edu

Senior International Partnerships Administrator

24-hour emergency line

806-742-3931

TTU and HNB-G Letter of Intent Page 3 of 3

HNB GARHWAL UNIVERSITY

School of Sciences

Name: Prof. D.S. Negi

Email: Devendra negi@vahoo.com

Head, Department of Chemistry

01346-252143





LETTER of INTENT

Between

TEXAS TECH UNIVERSITY, USA

and

HNB GARHWAL UNIVERSITY, INDIA

Representatives of Texas Tech University (hereafter referred to as TTU), 601 Indiana Ave, 79409 Lubbock, Texas, U.S.A. (phone 001.806.742.3667) and HNB Garhwal University (hereafter referred to as HNB-G), Srinagar, Uttarakhand 246174, India (phone +01346.252143), have discussed the desirability and feasibility of an interactive relationship designed to strengthen the bonds between the institutions. This Letter of Intent (LoI) is intended to help the parties facilitate future agreements that may formally establish an international exchange of faculty and students, research data, and educational and development programs.

OBJECTIVES:

- Desired relationship. The parties would like to further discuss an alliance between the institutions with regard to academic programs and research activities, particularly with the following departments:
- Purpose. The parties wish to broaden the experience of their respective faculty and students, providing each group with opportunities for increased cultural understanding.
- Future binding agreement. The parties wish to work toward a formal agreement that contains specific and binding provisions.
- 4. No exchange of funds. Neither party will incur, as a result of this Lol, any financial obligations resulting from the actions of the other party without an agreement in writing stating specific financial obligations.
- Term. This Lol will be effective for 2 years from the last signature date and either party may terminate at any time by giving the other party written notice.
- Letter Not Binding. This letter is intended solely as a basis for further discussion and is not intended to be and does not constitute a legally binding agreement.

TTU and HNB-G Letter of Intent Page 1 of 3 For Texas Tech University

For HNB Garhwal University

Sukant Misra Ph.D., Vice Provost for International Affairs

, 1, 2019

Month/Day/Year

Quan do

Dr. Annpurna Nautiyal, Vice Chancellor

1 13 1 2020 Month/Day/Year

Prof. Dr. D.S. Negi, Head. Department of Chemistry

103,2020 01

Month/Day/Year

TTU and HNB-G Letter of Intent Page 2 of 3



सी एस आई आर – राष्ट्रीय वनस्पति अनुसंधान सस्थान SPEED I

(वैज्ञानिक तथा औद्योगिक अनुसंधान परिषद) राणा प्रताप मार्ग, पोस्ट बाक्स सं0 436, लखनऊ-226 001 (भारत)

CSIR - NATIONAL BOTANICAL RESEARCH INSTITUTE

(Council of Scientific & Industrial Research) Rana Pratap Marg, P.B. No. 436, Lucknow-226 001 (India)

HO/NO.

Vivek Srivastava Principal Scientist Technology Transfer & Business Development email: vivek@nbri.res.in

26

दिनांक / Date

<u>T.T.B.D./C-02/2018/</u>33/ April 24, 2018

To,

The Registrar HNB Garhwal University (A Central University) Srinagar, Garhwal – 246174(Uttarakhand)

Subject : Signed Copy of Memorandum of Agreement - reg.

Reference : Your letter dated 29.03.2018 along with signed copies of MoU at your end.

Sir,

This has reference to the above mentioned subject. In this connection, please find enclosed herewith duly signed copy of MoU from our side for your record.

With regards

Yours Sincerely,

(Vivek Srivastava)

Encl. : As above



948X Phones 2208531, 32/33/2297800-2297999

Phone: 0522 - 120/206, 984

Gram - BAGH, Lucknow Fax (0622) 2205836 (2206839 Waneka (www.com



INDIA NON JUDICIAL Government of Uttarakhand

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- Article 5 Agreement or Memorandum of an agreement
- : H N B G U SRINAGAR PAURI GARHWAL
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- REGISTRAR
- REGISTRAR
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- (One Hundred only)

STAND

MEMORANDUM OF UNDERSTANDING

CL. 1. THE MoU

CL.1.1 This MoU made and entered into on.2.3. Day of April. 2019 between CSIR-National Botanical Research Institute (Council of Scientific & Industrial Research, a Society registered under the Societies Registration Act (XXI of 1860), having its registered office at Anusandhan Bhavan, 2, Rafi Marg, New Delhi-110001) having its office at Rana Pratap Marg, Lucknow-226001 (hereinafter called

Statutory Alert:

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CSIR-NBRI which expression shall where the context so admits include its successors and permitted assigns) of the one part,

And

CL.1.2 H. N. B. GARHWAL UNIVERSITY, Srinagar, a Central University having its office at Srinagar-246174 Dist. Garhwal (Uttarakhand) (hereinafter called the HNB Garhwal University which expression shall where the context so admits include its successors and permitted assigns) of the other part.

CL. 2. PREAMBLE

CSIR-NBRI, Lucknow and HNB Garhwal University, Srinagar have realized the need for a close linkage between the National Laboratories and the Universities in the terms of research and academic activities. In compliance to this idea, both have decided to sign a Memorandum of Understanding (MoU) as per the following terms and conditions:

CL. 3. GENERAL

CL.3.1 CSIR-National Botanical Research Institute (CSIR-NBRI) scientists may participate in the teaching program in the area of Botany, Plant and Animal Biotechnology, Zoology, Microbiology, Chemistry, Biochemistry, Pharmaceutical Sciences, Pharmaceutical Chemistry, Environmental Sciences, High Altitude Plant Physiology, Forestry, Horticulture, etc. at the Departments of Botany and Microbiology, Zoology and Biotechnology, Chemistry, Biochemistry, Pharmaceutical Sciences, Pharmaceutical Chemistry, Environmental Sciences, High Altitude Plant Physiology Research Centre, Forestry, Horticulture, etc. of HNB Garhwal University on request and subject to mutual convenience.

CL.3.2 CSIR-NBRI scientists and faculty members of Departments of Botany and Microbiology, Zoology and Biotechnology, Chemistry, Biochemistry, Pharmaceutical Sciences, Pharmaceutical Chemistry, Environmental Sciences, High Altitude Plant Physiology Research Centre, Forestry, Horticulture, etc. of HNB Garhwal University, will work towards joint research projects thereby mutually using leading research facilities developed/to be developed at CSIR-NBRI and HNB Garhwal University.

CL.3.3 HNB Garhwal University will recognize and accept CSIR-NBRI scientists as Supervisor / Co-Supervisor (as applicable) for Ph. D degree of HNB Garhwal University. One faculty member of the concerned department shall be Supervisor /Co-Supervisor (as applicable) with the student of CSIR-NBRI registered with HNB Garhwal University.

The Board of Studies, HNB Garhwal University shall decide Supervisor /Co-Supervisor for the student as applicable.

CL.3.4 The faculty and students of the above mentioned departments (as mentioned at 3.2) of HNB Garhwal University, who work in collaboration with the scientists of CSIR-NBRI shall be allowed to use laboratories and library of CSIR-National Botanical Research Institute (CSIR-NBRI) subject to the prior permission of the Director. Scientists and Ph. D students of CSIR-NBRI, who work in collaboration with the faculty of HNB Garhwal University shall likewise to be given the laboratory and library facilities of HNB Garhwal University with prior permission of competent authority.

CL. 4. ACADEMIC AND DEVELOPMENT PROGRAMMES

CL.4.1 For the mutual benefit of the two institutions i.e. CSIR-National Botanical Research Institute (CSIR-NBRI) and HNB Garhwal University, there will be exchange of CSIR-NBRI scientists and faculty members of the mentioned departments (as mentioned at point 3.2) of HNB Garhwal University.

CL.4.2 CSIR-NBRI and HNB Garhwal University scientists/faculty members will help each other in preparing common research programmes. CL.4.3 While registering CSIR-NBRI students for Ph. D degree in HNB Garhwal University, the ordinance related to Ph. D degree of HNB Garhwal University will be applicable. The students/researchers of CSIR-NBRI shall be required to qualify the entrance examination/ GATE/NET-JRF followed by interview for registration to Ph. D as per the norms of HNB Garhwal University, Srinagar, if any.

CL.4.4 CSIR-NBRI and HNB Garhwal University will work closely to generate appropriately trained manpower in the related areas through organization of formal academic curricula, symposia, short term training courses, etc.

CL.4.5 CSIR-NBRI as a national laboratory provides necessary regorous training of research methodology, literature survey, computation of data at par with the course work (one semester training course) required as per UGC norms/ordinance of HNB Garhwal University, Srinagar. Hence, research students of CSIR-NBRI will be completing this course at CSIR-NBRI but will have to give the Pre Ph. D examination at HNB Garhwal University.

CL.4.6 Honorarium, TA/DA (if applicable) will be provided to the faculty members/Scientists/technical staff of HNB Garhwal University and CSIR-NBRI as per rules by requesting organizations.

CL.5. DURATION OF MoU

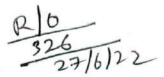
CL.5.1 This MoU shall be valid for a period of 60 months from the date of signing. The duration may be extended with the consent of both the parties under mutually agreed terms and conditions.

CL.5.2 It is further agreed that the Vice-Chancellor, HNB Garhwal Universith and Director, CSIR-National Botanical Research Insitute (CSIR-NBRI), through periodic meetings will monitor the process of this MoU. Any change/modification/termination as introdced/suggested by the Review Committee (comprising of Vice-Chancellor, HNB Garhwal University and Director, CSIR-National Botanical Research Institute (CSIR-NBRI) will be binding on both the organizations.

SEAL OF PARTIES

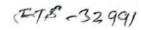
In WITNESS WHEREOF the parties hereto have signed this agreement of understanding the day, month and year as mentioned hereinbefore.

For and on behalf of CSIR-NBRI	For and on behalf of HNB Garhwal University
Signature:	Signature:
Name: A.K. GAUNIYAL	Name:A. K. THA
Designation: Head, P.M.E.	Designation: Registran
DR. ANIL KUMAR GAUNIYAL Seal:— Head, Planning, Monitoring & Evaluation CSIR-National Botanical Research Institu Lucknow-226001, Uttar Pradesh	H.N B. Garhwel University Div@ell:(A Central University)
Witnesses: (Name &Address)	Witnesses: (Name &Address)
1 Kg	1 Winter
1. Viveh Smandaula Scientin, TTISO, CSIE-NORI.	1. Int IK Thorthi Dept of Betany & Microbiology, HNB Genhurd Univ Smage
2. Sweet Sharing To, CSIR-NBRI, Luckmin	L
	Dr. A.K. Motionly
Date:	Date: Joint Registion (Academic) JANB Garhard (Mir.
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- Article 4 Affidavit
- HNBGU SRINAGAR GARHWAL
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- Registrar Dr Ajay Kumar Khunduri SO Govind Prasad
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......Please write or type below this line------MOU OF DR. AMBEDKAR CENTRE OF EXCELLENCE SCHEME (DACE)

MEMORANDUM OF UNDERSTANDING (MoU) FOR FUNCTIONING OF DR. AMBEDKAR CENTRE OF EXCELLENCE IN CENTRAL UNIVERSITY OF HEMVATI NANDAN BAHUGUNA GARHWAL UNIVERSITY, SRINAGAR GARHWAL, UTTARAKHAND

AND DR. AMBEDKAR FOUNDATION, MINISTRY OF SOCIAL JUSTICE AND EMPOWERMENT, GOVERNMENT OF INDIA, NEW DELHI

- The authenticity of this Stamp certificate should be venified at 'www.sholeslamp.com' or using e-Stamp Mobile App of Stock Holding Any discrepancy in the details on this Certificate and as available on the website / Mobile App remarks it invalid.
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 In case of the direction of the certificate
- 3. In case of any discrepancy please inform the Competent Authority

Dr. Bhimrao Ramji Ambedkar (1891-1956), fondly known as Babasaheb, is one 2. of the most illustrious sons of India and a great National Leader. He is considered as the champion for the Dalit cause, an erudite scholar, extraordinary statesman and a visionary who contributed greatly to the building of the modern Nation. Dr. Ambedkar left an indelible impression in the history of India as a messiah who unfettered the oppressed masses and secured human rights for millions of weaker and oppressed classes that were path-breaking in its essence and strived towards the monumental endeavors of freedom. He was the Chief Architect of the Constitution of India, wherein Babasaheb left emancipatory provisions for the justice and empowerment of the oppressed classes. He is the symbol of the struggle for justice and empowerment of the weaker and downtrodden population in India and laid the foundation stones of building a just society. Babasaheb's groundbreaking ideas led to the formation of the Reserve Bank of India during British rule. As a Labour leader, he promoted the revolutionary idea of "fair condition of life of labour" as opposed to "fair condition of work" which provided the outline of the future labour laws in India. Babasaheb was also a champion of the cause of gender parity as he initiated reforms for lessening of working hours to 48 hours per week, removed the ban of engaging the women in various forms of

employment and coded the principle of "equal pay for equal work" irrespective of gender. His idea of the Hindu Code Bill of emancipatory nature. Babasaheb also Ieft a lasting impression as a social reformer through his role in the movements like Mahad Satyagraha, the Anti-Khoti movement and the Dalit Buddhist movement.

3. The Governing Body of Dr. Ambedkar Foundation in its meeting held on 17.11.2021 had decided to implement a New Scheme called "Dr. Ambedkar Centre of Excellence"

4. In pursuance of the decisions of the Governing Body, the scheme is aimed to establish Dr. Ambedkar Centre for Excellence in at least 3\ Central Universities one from every State/UT of India. Through the Scheme, the Centre shall empower Scheduled Caste students in competitive exams by providing the best and free coaching facilities

AND WHEREAS, this scheme provides specialized coaching ONLY to the Scheduled Caste students for the Civil Services examination conducted by the UPSC.

5. AND WHEREAS the Scheme of Dr. Ambedkar Centre of Excellence includes the following criteria:

Implementing Universities and Other Conditions:

This scheme will run through at least \exists_{i} Central Universities one from every State/UT of India and provide financial assistance as per the following:

- i. DACE shall be established after the acceptance of Vice-Chancellor of the Central University.
- ii. In order to conduct coaching classes for the subject applied for, the Universities must possess necessary infrastructure such as a separate classrooms, library, Hi-Speed wifi connectivity, and other requisite equipment, etc.

Eligibility Criteria for Students and Selection Procedure for the Free Coaching

- i. In order to be considered for admission into the center for desired subject, candidates must have achieved the required minimum percentage of marks in the qualifying examination.
- ii. The Scheme is open to only Scheduled Caste students. University shall obtain/verily requisite caste certificate from the concerned candidate/ student/ issued by the competent authorities in his/her respective States/UTs or as the case may be.

Page 3 of 8

- iii. The coaching benefits under the scheme may only be utilized by a student ONCE regardless of the number of chances that he/she may be entitled to in a particular Competitive Examination. The student must submit to the University an affidavit stating that he/she has not received any monetary benefit from any of the other schemes by Government of India, State/UTs or any funding agency.
- iv. The candidates enrolled under this scheme shall have to attend all the classes. In case of remaining absent for more than 15 days without any valid reason, or leaving the coaching midway without prior approval of the competent authority, the total expenditure incurred the on candidate will be recovered from the University/student/candidate concerned. University shall be responsible for the recovery.
- v. A total of hundred seats (100) sanctioned for coaching per Centre. 33% of the total sanctioned seats for coaching may preferably be given to the eligible female candidates of SC category. In case the sufficient number of eligible female candidates do not apply, University may allocate the vacant seats to the male/transgender candidates (SC category only).
- vi. The concerned University shall widely publicize the scheme through open advertisement and make a public notice for a Common Entrance Test (CET). The student shall be selected through an entrance test conducted by the University as per merit.
- vii. The University will be required to maintain Attendance Registers for all candidates enrolled for the coaching classes as well as subject-wise results of practice tests, or any other evaluation.

Faculty/ Resources

i. The Competent Authority of the Central University shall appoint a "Program Coordinator" for the Doctor Ambedkar Centre for Excellence, for a minimum term of 2 years. He/she must be a permanent faculty of the same University who may be entrusted with this additional assignment. He/she may also be paid a token honorarium/Special pay @ of Rs. 20,000/- per month from the total budget of the scheme. His/her role of responsibilities/ duties etc. shall be defined by the concerned University, however, his/her Annual Performance Appraisal Report (APAR) shall be submitted to the DAF, for monitoring and records.

- ii. The University will be required to appoint at least Three (03) contractual/term appointment/ teachers, who must be eminent and professional scholars in the concerned/allied/relevant discipline and must have published work of high quality and actively engaged in research and training.
- iii. It should be necessary for a teacher to remain available at least 06 hours daily in the University. The minimum direct teaching- learning process must involve 20 to 25 hours per week.
- iv. Contractual appointee teachers may be entitled for 12 leaves in a calendar year excluding the public holidays. All other leaves shall be counted as leave without pay, subject to the maximum period of one month. The other relevant 'terms and conditions' may be fixed by the University and a copy of the same shall be shared with DAF.
- v. For the above mentioned post, he/she will be paid a consolidated fixed pay salary of Rs. 1,15,000/ per month (Taxes as applicable)

Nature of Financial Assistance

Dr Ambedkar Foundation shall fund the entire expenditure to be incurred on the establishment of Dr. Ambedkar Centre for Excellence as per the terms and conditions of the scheme.

- i. The grant will be provided per student basis to the University.
- ii. The expenditure of the scheme which includes all expenses may be released to Dr. Ambedkar Centre for Excellence in **Two installments** every year. The first installment @ 75% will be released immediately to the University as soon as it is notified/empanelled by the DAF and the students are selected and finalized by the University.
- iii. The remaining second installment of 25% to the Dr Ambedkar Centre for Excellence will be released after the submission of Utilization Certificate, Audited statement of accounts, receipts if any, and other payments details certified by a Chartered Accountant on the panel of CAG along with the half yearly report outcome of the programmes initiated by Dr Ambedkar Centre for Excellence.
- iv. The Central University must inform DAF about the date wise commencement of the coaching programs including nominal role of Students batch wise and any other details.

v. All the funds shall be released as per the norms prescribed in the GFR 2017 and amendments from time to time. The Central University will make expenditure only through Public Financial Management System (PFMS) DBT mode.

SI No.	Type of Coaching	Coaching Fees	Duration of Coaching/ Course	Minimum Annual Success Rates
a)	The Civil Services examination Conducted by UPSC	Rs. 75,000/- per annum / per student	01 year	10%

Subject wise earmarking of Universities including allocation of Students shall be done post declaration of **CET** results under a separate notification by DAF/University.

The Universities shall appoint subject wise faculty to meet the requirements of students selected in that University.

Financial assistance for each DACE per year

Number of Universities	31	
Number of Students (per University)	100	
Number of Teachers per University	03 + 01 (Programme Coordinator)	
Grant per student provided	INR 75,000/-	
Total	INR 75,00,000/- (per University) all inclusive	

Monitoring

At the end of each academic year, the Coordinator of the programme will submit, through the Registrar, an appraisal report, indicating the performance of each candidate. The appraisal report should also indicate:

- The duration for which the coaching was organized, classes /period, and the number of the candidates who actually participated in the programme.
- ii. The number of candidates who actually appeared in the examination.
- iii. The number of successful candidates in each paper along with the problems faced by the University in the implementation of the scheme.
- iv. The subjects in which they were tutored, names of Teachers and their subject discipline.
- v. Overall comments of the Coordinator

Additional Notes:-

- i. The University shall be monitored by the DAF and Do SJ&E. The sponsoring agency may conduct physical inspection, at any point of time and even without prior information, or it may be done through Independent agencies.
- ii. The University shall provide all details on its Website for Dr. Ambedkar Centre for Excellence and furnish website address to the DAF. The University shall also upload year-wise and course-wise photographs of coaching programnme enrolled students on their website for ready reference.
- iii. This University shall open a separate "Current Bank Account" for this scheme. All expenditures related to this program should be made from this account only. The account should be audited as per Government norms in each financial year, and confirm the same to Dr. Ambedkar Foundation by the month of May of next financial year.

6. WHEREAS, the FOUNDATION and the Central University of

for implementation of the above objectives, have decided to set up the Dr. Ambedkar Centre of Excellence and has agreed to act jointly with the FOUNDATION by extending necessary cooperation for proper implementation of the above Scheme.

7. NOW, BY THIS MEMORANDUM OF UNDERSTANDING the FOUNDATION and the CENTRAL. UNIVERSITY hereby agree as follows:

- i. That, the University and the Foundation shall cooperate with each other and jointly extend all cooperation for proper implementation on the objectives as aforesaid.
- ii. That, the University shall comply with the terms of the MoU of the Foundation for better functioning of the Scheme.
- iii. That, the University shall extend all infrastructural facilities as proposed in the MoU of the Foundation.
- iv. That, both the parties have agreed to comply with all financial and other liabilities as stipulated in the MoU of the Foundation for effective functioning of the Scheme.
- v. That, the compliances of the Scheme of the Foundation shall be treated as part of the MoU, which shall not be disputed by the University in any manner.

- vi. That, the performance shall be reviewed each year by the DAF/D/o SJE or through any other Agency.
- vii. That, this MoU shall come into force/ operational with effect from 01.04.2022.
- viii. In witnesses whereof, the parties hereto have put their respective seal and signature on the day, month, year written hereinabove.

Vice Chancellor/Nominee of the Vice Chancellor Poof . RC Bhatt chancellor HNBGI 1 Dean/ DR Dr. Sanjay Kumar Ahyin DR (Jegal) HNBGU 2

Member/Secretary, DA 1 ...

Representative, DAF

In witnesses whereof

 \mathcal{N} 2

22104/K

Registrar/Deputy Registrar University/Institute Dr. Ajay Kurner Hchandern Registrar, HHBGU

Director Dr. Ambedkar Foundation विकास जिवेदी/VIKAS TRIVEDI निदेशक Director औं अम्बेटकर अंतर जीत केन्द्र Dr. Ambedkar Infernational Cantra सामाजिक प्याद और अध्यक्षा के स्वारम्ब Ministry of Social Justice & Encountrant मारस सरकार, नई हि उन्हों Govt of India, New Delthi



हेमवती नन्दन बहुगुणा गढ़वाल किंवविद्यालय Hemvati Nandan Bahuguna Garhwal University श्रीनगर गढ़वाल (उत्तराखण्ड)–246174 Srinagar Garhwal (Uttarakhand) - 246174 (केन्द्रीय किंवविद्यालय) (A Central University)

31-08-2021

The Chairman, RCC Cell, HNBGU

Sub: Regarding Memorandum of Agreement (MOA) related to Skill Vigyan Program

Dear Sir,

This is to inform you that a Memorandum of Agreement (MOA) has been signed between HNBGU and UCOST for running a Skill Vigyan Program at HNBGU with financial support from DBT, Govt. of India and UCOST. A copy of the MOA is attached herewith for your kind perusal and information.

Thanking you,

Yours sincerely. Dr. GK Joshi

Nodal Coordinator Skill Vigyan Program, Assistant Professor. Department of Biotechnology HNBGU



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SIGN. . DEEPAK RAJPUT STAMP VENDOR T.G.M.O.U. BUS STAND SRINAGAR GARHWAL 9410536904



Statutory Aler

---Please write or type below this line-----MEMORANDUM OF AGREEMENT

This MEMORANDUM OF AGREEMENT is made on this 9th March day of Two thousand and Twenty One BY AND BETWEEN Uttarakhand State Council for Science and Technology (UCOST), Dehradun, Govt. of Uttarakhand having its registered office at "Vigyan Dham, Post-Jhajra (Suddhowala), Dehradun, Uttarakhand-248007" acting through Director General, Uttarakhand State Council for Science and Technology (UCOST), Dehradun, hereinafter referred to as the 'UCOST' (which expression unless excluded by or repugnant to the subject shall mean and include its successor-in-office and assignees) of the FIRST PARTY; H.N.B. Garhwal (Central) University, having its registered office at Srinagar-Garhwal–246174, Distt.- Pauri Garhwal (Uttarakhand), hereinafter referred to as HNBGU (which expression shall where the context so admits include its successors and permitted assignees) of the SECOND PARTY;

WHEREAS UCOST being desirous of capacity building in the area of Biotechnology teaching decided to support a Skill Vigyan Program submitted by HNBGU.

This Memorandum of Agreement (MoA) defines the role and responsibilities of the participating agencies, monitoring and other matters related to the Skill Vigyan Program.

NOW THE PARTIES HERETO AGREE AS FOLLOWS:-

1.0 ROLE OF UCOST, DEHRADUN

- 1.1 To provide funds to the extent of Rs. 19.80 lakh (Rs Nineteen lakh eighty thousand only) over a period of three years from the date of Administrative sanction of the project, (31.5.2019), for conducting Skill Vigyan Programme. Details of the grant, number of seats per year and annual recurring provision are given at Annexure-I (Administrative Sanction Order).
- 1.2 To provide additional funds amounting to Rs. 2.00 Lakh (Rs Two Lakh only) per year to meet any shortfall in miscellaneous expenses during implementation of project, viz., honorarium, travel, field visits, contingency, etc. The money would be released in ratio of 60: 40.

2.0. ROLE OF HNBGU

- 2.1. To provide existing facilities as mentioned in the project document.
- 2.2. To accomplish and fulfill the terms and conditions listed at Annexure-II.
- 2.3. NSQF aligned courses framed by Life Sciences Sector Skill Development Council (LSSSDC) and training duration specified therein would be strictly implemented by the institute.
- 2.4. Pre-placement and post placement initiatives would be submitted to UCOST mentioning the employment status of trained students (self-employment/ opting higher education/ job placement).
- 15 days before the completion of training program, UCOST would be informed regarding assessment of students.
- 2.6. For certification of trainers, the nodal coordinator and at least one team member would undergo Training-of-Trainer (TOT) program to be organized by UCOST.
- 2.7. Timely on-boarding of institute and registration of dedicated trainers on Skill India portal (https://skillindia.nsdcindia.org/) for affiliation and Train the Trainers program.
- 2.8. Institute should ensure timely purchase and distribution of LSSSDC Job Roles Participating handbooks to each trainee within a timeline.
- 2.9. To submit an annual audited statement of expenditure incurred under this Skill Vigyan program.
- 2.10. To ensure effective utilization of the grant given by UCOST for the purpose for which it was granted and to ensure timely completion of course work/ examination and placement.

2.11. HNBGU will take up all the liabilities of the Skill Vigyan program including Manpower (if any), upto the sanction period specified in the sanction order.

3.0 DURATION OF PROJECT

3.1 Duration of project shall be three years from the date the Program has been sanctioned by UCOST.

4.0 RIGHTS OF OWNERSHIP/TECHNOLOGY TRANSFER AND UTILIZATION

- 4.1 All the assets including the equipment and produce acquired will be the property of UCOST and shall not be utilized for purposes other than teaching/capacity building at **HNBGU**. The rights of **HNBGU** under this MoA shall not be transferred to any other party without prior approval in writing of UCOST.
- 4.2 It shall be the responsibility of HNBGU to ensure that support of UCOST is suitably acknowledged in all the publications (papers, reports, newspapers, etc.) arising out of the program.

5. MONITORING

- 5.1 The progress of implementation of the Skill Vigyan Program and proper utilization of grant shall be reviewed by UCOST and by the Advisory Committee set up by HNBGU on yearly basis.
- 5.2 The periodic progress of physical achievements and the utilization of funds, statement of expenditure shall be evaluated by the Advisory Committee and UCOST.
- 5.3 The Comptroller and Auditor General of India, at his discretion shall have the right of access to the books and accounts of HNBGU for the grants received from UCOST for this program.
- 5.4 UCOST may terminate the grant at any stage if it is convinced that the grant has not been properly utilized or appropriate progress has not been made. In the event UCOST terminates the grant, **HNBGU** shall hand over all documents including technical details and equipment purchased related to the programme.

6.0 DURATION OF MEMORANDUM OF AGREEMENT

This MoA will remain in force for the duration of the program.

7.0 ARBITRATION

In the event of any question, dispute or difference whatsoever arising between the parties to this Agreement out of or relating to the construction, meaning, scope, operation or effect of this Agreement or the validity of the breach thereof shall be referred to an Arbitrator to be appointed by mutual consent of both the parties herein. If the parties cannot agree on the appointment of the Arbitrator within a period of one month from the notification by one party to the other of existence of such dispute, then the Arbitrator shall be nominated by the Secretary, Law Department, Government of Uttarakhand. The provisions of the Arbitration and Conciliation Act, 1996 will be applicable and the award made there under shall be final and binding upon the parties hereto, subject to legal remedies available under the law. Such differences shall be deemed to be a submission to arbitration under the Indian Arbitration and Conciliation Act, 1996, or of any modifications or reenactments thereof.

GOVERNING LAW

This Contract shall be governed by the Law of India for the time being in force.

IN WITNESS WHEREOF the parties hereto have signed, sealed and delivered this Agreement on the day, month and year first above written in presence of:

Signed and Stamped by - Clockuby Witnesses: 1. 2. (Designation) महानिदेशक/Director General वि. प्रौ. परि./U-COST For and on behalf of UCOST, Dehradun Witnesses: Signed and Stamped by --103/21 1. Dr. G. K. Joli - Registrar Department of Ario tehnley in MARAU 2. Do Sudher Kuman Sudur-Dependent of Bestechnology H.N.B.Garhwal University (Designation) (A Central University) Srinagar (Sathwal) **HNBGU, Srinagar**



सी.एस.आई.आर-हिमालय जैवसंपदा प्रौद्योगिकी संस्थान पो॰ बॉ॰ न॰ 6 पालमपुर - 176 061 (हि.प्र.) भारत CSIR-INSTITUTE OF HIMALAYAN BIORESOURCE TECHNOLOGY Post Box No. 6 Palampur (H.P.) 176 061 INDIA

BY SPEED POST

No. CSIR-IHBT/BDMU/2021

15th November, 2021 /4 45

Subject:- Memorandum of Understanding.

Dear Sir,

Please find attached herewith a signed copy of Memorandum of Understanding made on 12th November, 2021 between CSIR-IHBT, Palampur and Hemvati Nandan Bahuguna Garhwal University (Central University), Srinagar in original. It is requested to kindly sign the MoU and send back your signed copy to this Institute for our record.

Kindly acknowledge the receipt.

Yours sincerely,

(Dr. Sukhjinder Singh) Coordinator Business Development & Marketing Unit

Encl: As above

Prof. M. C. Nautiyal Convener, RCC Cell Hemvati Nandan Bahuguna Garhwal University (Central University) Srinagar-246 174 Distt. Garhwal (Uttrakhand), India



MEMORANDUM OF UNDERSTANDING



C.S.I.R.-IHBT

BETWEEN

HEMVATI NANDAN BAHUGUNA GARHWAL UNIVERSITY (CENTRAL UNIVERSITY), SRINAGAR

AND

CSIR-INSTITUTE OF HIMALAYAN BIORESOURCE TECHNOLOGY (CSIR-IHBT), PALAMPUR

FOR

COLLABORATION ON RESEARCH & DEVELOPMENT & ACADEMICS INCLUDING FACULTY AND STUDENT EXCHANGE

In furtherance of their mutual interest in the fields of education and research and as a contribution towards increasing national cooperation, Hemvati Nandan Bahuguna Garhwal University (Central University), having its registered address at Srinagar- 246174, Dist. Garhwal (Uttrakhand), India (here in after referred to as H.N.B. Garhwal University).

And

CSIR-Institute of Himalayan Bioresource Technology having its registered address at Palampur-176061, District Kangra, Himachal Pradesh, India, Estt. By Council of Scientific and Industrial Research (here in after referred to as CSIR-IHBT) have entered into this Memorandum of Understanding (MoU) on 12th November, 2021 with broad terms and conditions as follows:

ARTICLE - I

The MoU involves academic and R&D collaboration between H.N.B. Garhwal University and CSIR-IHBT (also referred to as institutions) in areas of relevance to each other.

Accordingly, the institutions as per their extant Guidelines shall seek to promote or facilitate

H.N.B.G.U.

- 1. Exchange of staff and students (faculty, research scholars, and research project employees) regarding academics and research in the areas of mutual interest for institutions.
- 2. Exchange of students for pursuing course study and research programmes for mutual benefit of institutions.
- 3. Collaboration in research & development programs and consultancy activities.
- 4. Exchange of academic and research material and publications/IPs (exists in Public domain).
- 5. Cooperation in project proposals and research activities of mutual interest.
- 6. Collaboration in research & development in the areas of Food and Neutraceuticals, models for animal studies, ecology, medicinal aromatic plants, floriculture, tea sciences and other mutually decided areas. This also includes collaboration in setting-up and upkeep of the relevant infrastructure among the institutions.
- 7. Joint collaboration with Government organizations/ private entities/ PSUs in the mutually decided areas.
- 8. Jointly organization of scientific events like seminars/conferences/workshops etc.
- 9. Exchange of students for summer/winter internships
- 10.Intellectual Properties (IPs) and benefits arising out of commercialization of technologies generated out of co-operation under this MoU shall be jointly shared by H.N.B. Garhwal University and CSIR- IHBT on mutual consent in each case.
- 11.H.N.B. Garhwal University and CSIR-IHBT joint outreach programme for Science education and related technological advances for social benefit.

ARTICLE - II

Implementation of cooperation based on this MoU shall be dealt with between the relevant faculties and divisions/departments of the institutions. Wherever necessary and subject to availability of resources, a specific plan shall be

H.N.B.G.U.

worked-out for each activity mentioned above in Article – I setting forth detailed arrangements for collaboration. Such plans shall be subject to approval of the appropriate authorities of each institution. To facilitate development of such plans, each institution shall nominate a member of its staff to coordinate activities arising under this MoU.

ARTICLE – III

The institutions agree and undertake to keep confidential, at all times, any information and/or data that may be exchanged, acquired and/or shared in connection with the area of cooperation as mentioned above in ARTICLE - I (3) unless otherwise the same information already exists in the public domain.

ARTICLE - IV

Ownership of any research findings of joint research work specified shall be vested with the institutions specified in this MoU. Notwithstanding anything stated in the MoU, the MoU is free from any financial commitment of one party towards the other and also towards any other party.

ARTICLE - V

The MoU shall remain in force for a period of 05 (five) years commencing from the date of signing and may be reviewed/renewed by mutual consent by serving 3 (Three) months written notice to the other institution. The MoU may be amended with mutual consent of both the institutions in writing.

ARTICLE - VI

H.N.B., Garhwal University and CSIR-IHBT reserve the right to terminate this MoU by either institution giving 3 (Three) months written notice to the other. Where such termination occurs, the provisions of this MoU shall continue to apply to ongoing activities until their completion.

ARTICLE - VII

Participating staff and students involved in any activities under this MoU must adhere to laws of the country and rules & regulations of the host institutions.

Siloza 8 S.I.R.-IHB

H.N.B.G.U.

ARTICLE - VIII

H.N.B., Garhwal University and CSIR-IHBT welcome establishment of this MoU for cooperation and jointly agree to the provisions as set-out above. There will be two copies of this MoU equally valid, one for each institution, effective from the date of its signing.

Signed for and on behalf of H.N.B., Garhwal University

Jan Jan 12.11. 2021

(Prof. Annpurna Nautiyal) Vice-Chancellor H.N.B., Garhwal University

Witness

(**Dr. Ajay Kumar Khanduri**) Registrar H.N.B., Garhwal University

Thij-

(Prof. J. S. Chauhan) Dean, Agriculture & Allied Sciences H.N.B.Garhwal University

Muantinga

(Prof. M. C. Nantiyal) Convener, RCC Cell H.N.B., Garhwal University

Signed for and on behalf of CSIR-IHBT, Palampur

anijan 1 12: 11: 20 m

(Dr. Sanjay Kumar) Director CSIR-IHBT, Palampur

Witness

Antham

(**Dr. Ram Kumar Sharma**) Coordinator, Academy of Scientific & Innovation Research (AcSIR) CSIR-IHBT, Palampur

3320 20 12/11/202

(**Dr. Sukhjinder Singh**) Coordinator, BDMU, CSIR-IHBT, Palampur

Administ CSIR-IHBT, Palampur

H.N.B.G.U.

C.S.I.R.-IHBT







MEMORANDUM OF UNDERSTANDING

BETWEEN



Ahmedabad Textile Industry's Research Association (ATIRA)

Dr. Vikram Sarabhai Road, PO-Ambawadi Vistar Ahmedabad – 380015 Gujarat, India

AND



HEMVATI NANDAN BAHUGUNA GARHWAL UNIVERSITY

(A CENTRAL UNIVERSITY) Srinagar (Garhwal), Uttarakhand





This Memorandum of Understanding (hereinafter referred to as the "MoU") is entered into this <u>10</u> day of <u>Feb</u> 2022.

BETWEEN

AHMEDABAD TEXTILE INDUSTRY'S RESEARCH ASSOCIATION, AHMEDABAD

AND

HEMVATI NANDAN BAHUGUNA GARHWAL UNIVERSITY, SRINAGAR

Preamble:

Hemvati Nandan Bahuguna Garhwal University (HNBGU)

Hemvati Nandan Bahuguna Garhwal University was established as a State University vide U.P. State Government notification no. (10)/(865)/15/(75)(85)/64 dated 23 November 1973. The University has subsequently been upgraded to Central University by an Act of Parliament i.e. the Central Universities Act 2009. Since its inception, the University has shown commitment to regional and community development inherent in its teaching courses, research agenda, and other outreach and extension initiatives. The University, nestled in the lap of Himalayan ranges in the Garhwal region of Uttarakhand, is a residential cum affiliating institution of higher learning. The University has following three campuses distantly located from each other where the undergraduate, post-graduate, and research programmes are being offered in different disciplines.

1. Birla Campus, Srinagar Garhwal with its extension at Chauras Campus,

2. B. Gopal Reddy (BGR) Campus, Pauri and

3. Swami Ram Teerth (SRT) Campus, Badshahithaul, Tehri.

The university having its main campus at Srinagar – 246174, Dist.Garhwal (Uttarakhand), India – 246174 (herein after referred to as "HNBGU", which expressions shall mean and include, unless repugnant to the context or meaning thereof its successors and permitted assigns) on the first party;





Ahmedabad Textile Industry's Research Association (ATIRA), Ahmedabad

ATIRA is an autonomous, non-profit Research Institute, linked to Ministry of Textiles Government of India, for textile and allied research. It is the largest of its kind in India as Co-Operative Research Organization. ATIRA was declared as "Centre of Excellence in composite" by Ministry of Textiles, Government of India in March 2011. This is also supported by Govt. of Gujarat. ATIRA has so far established state-of-art facilities for Testing of Mechanical properties like strength, elongation, fatigue etc. related to composite materials and fibers like Glass, Carbon, and Aramid etc. and heat & flame properties of composite materials. ATIRA has unique facility to perform NDT by ultra-sonic, radiography, X-ray method. ATIRA is in the process of setting up testing facilities for evaluating electrical properties of composite materials. The processing facility covers the Pultrusion laboratory and vacuum infusion laboratory. It is having its registered office at Ambawadi Vistar, P.O., Ahmedabad 380 015 and (hereinafter referred to as "ATIRA" which expression shall include its successor-in-interest and assigns) on the second party.

Either or both may also be hereinafter referred to, individually as the "Party," and collectively as the "Parties."

2. Scope of Work

The primary objectives of this MoU are:

- 2.1. HNBGU and ATIRA to recognize each other as Centres of Excellence for Academic and Research collaborations and to share their facilities and resources for research and other mutual benefits.
- 2.2. To undertake joint research activities with National and International Funding support (for ex. GOI S&T agencies like DST, CSIR, SERC, etc.)
- 2.3. The scope covers mutual co-operation between ATIRA & HNBGU to collaborate with each other as strategic partners for undertaking collaborative R&D activities in composite materials and related thrust areas of technology for the benefit of the student fraternity and Industry in particular and the composite Sector at large.





- 2.4. Both the parties agree that they shall harness their complimentary resources (the resource persons/faculty, the Graduate/PG/Doctoral student fraternity of HNBGU and the Scientists of ATIRA would collaborate and leverage the facilities available at ATIRA and HNBGU) and expertise to work together to pursue co-operative research activities for the benefit of the composite and similar Sectors at large.
- a. Defining new areas of collaboration that have not been foreseen, but can be beneficial to the Parties.

The areas of cooperation may be revised by mutual consent. However, specific programmes may require separate agreements detailed out and documented as annexures to this MoU.

3. Responsibilities of ATIRA & HNBGU

- a) To identify specific contextual research themes/projects in consultation with stake holders including industry, as appropriate.
- b) Form Research teams consisting of ATIRA scientists and HNBGU faculty, on a project-to-project basis.
- c) Identify and establish a cost model for sourcing necessary materials, facilities/ equipment, consumables, etc. The facilities/resource persons can be sourced either from sponsor, ATIRA or HNBGU, or others on a project-to-project basis.
- d) To formulate, design and offer co-branded, Industry focused programmes under Academy of Excellence for Advanced composite Technologies under ATIRA and HNBGU.

4. Outcomes / Deliverables:

- a) Knowledge Up gradation and Research Capability Enhancement of the HNBGU and ATIRA people.
- b) Development of capabilities and institutionalized platforms to work on the futuristic requirements of the composite and similar Sectors.





5. Financial Terms

The parties agreed to co-operate and otherwise act in good faith with the view to successful implementation of this MOU. There is no financial binding on either Party.

This MoU is intended to be a mutual undertaking of the parties hereto as at the date thereof.

6. Confidentiality:

a. Both HNBGU and ATIRA acknowledge that certain Confidential information may be disclosed by one party to the other (the Party that owns and/or discloses the Confidential Information is hereinafter referred to as the "disclosing party" and the Party receiving or accessing such Confidential Information is referred to as "receiving party") during the tenure of this MoU or performance of the respective obligations under the resultant definitive agreements hereunder.

Confidential Information means all information identified as "Confidential", including but not limited to information concerning the trade secrets, intellectual property rights, know-how, formulae, processes, inventions, data, network configurations, system architecture, designs, flow charts, drawings, proprietary information, data or materials related to business, services, products, customers, employees, finances or operational information of either party, and any other confidential or proprietary information the disclosure of which might harm or destroy a competitive advantage of the disclosing party. The mode of communication of information:

- i. Written
- ii. Oral
- iii. Electronic or any other form

iv. Both Technical & Non-Technical information

The receiving party shall not, directly or indirectly, disclose to any third party other than its employees, affiliated companies, and authorized agents any information concerning the disclosing party's business methods, products, customers or



finances, or any other Confidential Information which is disclosed to it by the disclosing party, without the prior written permission of the disclosing party, unless such disclosure is specifically required in the course of the performance by the receiving party of its obligations hereunder or under the resultant definitive agreements. The obligations of receiving party under this Section shall not extend to any information which: (i) is or becomes a matter of public knowledge, not as a result of any action of the receiving party; (ii) is lawfully in the possession of the receiving party prior to a disclosure hereunder; (iii) is received from a third party who lawfully acquired such information without restriction, and without a breach hereof, by the receiving party; (iv) is disclosed by the receiving party with the disclosing party's prior written approval (v) the information is independently developed by the receiving party, without use of the disclosing party's Confidential Information;(vi) is disclosed by the receiving party under operation of law or regulation or legal process;

- b. The Parties acknowledge that this MoU contains confidential information that shall be considered proprietary by both Parties, and agree to limit distribution of or disclosure about the Confidential Information hereunder and/or this MoU to those persons who have the access to confidential information within their respective organizations with a legitimate need to know the contents of this MoU. Neither party shall publicize or make any public announcement concerning the terms or nature of the relationship or this MoU without the prior written consent of the other party.
- c. HNBGU and ATIRA both acknowledge that any breach by them of their respective obligations under this Section may cause irreparable harm to the other party for which its remedies at law may be inadequate and that in the event of any such breach either party shall be entitled to seek equitable relief (including without limitation injunctive relief and specific performance) in addition to other remedies provided hereunder or available at law.

Upon termination or dissolution of this MoU, or upon earlier demand thereof, each party shall at the other party's option, either destroy under written certification of such destruction or return to the other party all properties



containing the other party's confidential information and copies thereof in its possession. The deletion of information from the following sources:

- i. Computers
- ii. Servers
- III. Storage devices, to the extent of failure to destroy the data, corresponding obligations are preserved.
- d. The Confidentiality Obligations under this Section shall survive during the term of this MoU and 3 years thereafter.

7. Guarantee, Liability

The parties shall properly perform, to the best of their knowledge and taking into account the current state of the art, all work assumed by them under this Agreement. Neither of the Parties gives any warranties concerning the accuracy and completeness of information disclosed and of objects transferred or concerning the non-existence of rights of third parties.

The Parties shall mutually waive any claims for themselves and their staff members in respect of any damage resulting from the performance of this Agreement excepting in case of intention or gross negligence. The respective liability for their interactions with the third parties shall not be affected by this waiver.

8. Intellectual Property Rights

- 8.1 For purposes of this MoU, the term "Intellectual Property" shall mean registered and unregistered inventions, copyrighted works, design, trade secrets, know-how, technical information and any other proprietary information of either party.
- 8.2 Each Party shall retain title to any Intellectual Property if developed authored, conceived or reduced to practice independently and solely by that Party during the performance of this MoU without the other Party's Intellectual Property. In such event, no license, express or implied, shall inure to the benefit of the other participating Party to prepare copies and derivative works of such copyrighted works or to make, use, sell and export/import products or processes incorporating such Intellectual Property, except as expressly provided herein.





9. Force Majeure

Neither Party shall be responsible for non-fulfillment of its obligations under this MoU due to the exigency of one or more of the force majeure events such as but not limited to acts of God, War, Flood, Earthquakes, Epidemics, Riots, Civil Commotions, Strikes and Lockouts, provided on the occurrence and cessation of any such event, either Party shall give a notice in writing to the other within 48 hours of such occurrence or cessation, if the force majeure conditions continue beyond six months, the parties shall jointly decide about the future course of action.

10. Governing Law and Dispute Resolution

This MoU shall be governed, interpreted and construed in accordance with laws of India. In the event of any dispute arising between the Parties, the dispute shall be resolved between Heads of the institutions.

11. Termination

In case it is found by the two parties that the desired objectives cannot be achieved due to any reason(s) whatsoever this MoU will be treated as terminated by a mutually agreed procedure without any liability on either of the party.

12. Non- exclusive nature of this MoU.

This MoU shall be non-exclusive in its nature for all purpose.

13. Effective date & Duration of the MoU

This MoU comes in to effect from the date of its signing and will remain in force for three years, unless it is concluded or terminated on mutual consent by both the parties. However, the terms related to confidentiality/ secrecy will be respected by both the parties for a period of three (03) years even after such termination as provided in this MoU.



14. Notices.

All notices, request, demands, consents, waivers or other communications required to be given by either party to the other pursuant to this Agreement shall be in English, in writing and shall be deemed to have been given when hand delivered by messenger or a courier or sent by registered post or speed post or facsimile (to be subsequently confirmed by a registered letter or by hand delivery) to the other Party at the following address:

In witness whereof the Parties here to have executed this MoU through their authorized representatives.

Signed For and on behalf of Hemvati Nandan Bahuguna Garhwal University Signed For and on behalf of Ahmedabad Textile Industry's Research Association

Signature:

Witness:

Dr. Ajay Kumar Khandun Registrar Name : H.N.B. Garhwal University

(A Contral University)

Srinagar (Garmwal)

ATIRA

Name: Dr. TANMOY GANGOPADHTAT Dy Director ATIRA

Signature: Jan ~

and and Technology N.B. Garnwart Persity Signature M A Central University) Grinogar (Garhwall Name & Address: School Signature Name & Add ress

Name & Address: Dr. Ankush sharma Scientific officer, ATIRA

Signature:

Signature :

Name & Address: ASHCK KUMAR BHUYAN

Manager





This MEMORANDUM OF UNDERSTANDING DATED April 1st 2019

Between:

UNIVERSITY OF APPLIED FOREST SCIENCES ROTTENBURG, GERMANY

- and -

HNB-GARHWAL UNIVERSITY, SRINAGAR (GARHWAL), INDIA

WHEREAS:

A. The Head of International Office of Applied Forest Sciences Rottenburg and the Vice Chancellor of the HNB-GARHWAL UNIVERSITY, SRINAGAR (GARHWAL), for the purpose of furthering cooperation through educational and academic exchanges, hereby affirm their intent to promote such cooperation activities as will be of mutual benefit for their respective institutions.

Cooperation activities are considered here to include but not be limited to:

- Development of mutually beneficial academic programs and courses;
- Coordination of academic staff mobility for purposes of teaching, research and training;
- Coordination of student mobility programs for study and research;

Memorandum of Understanding

- A one month training programme with provision of boarding and lodging shall be provided;
- Coordination of academic activities such as joint research, publication and symposia;
- Exchange of documentation and research materials in fields of mutual interest provided that, to the best knowledge of the respective institutions, there is no prohibition at law or otherwise against the exchange;
- Other activities considered by the parties to be of benefit to each party's education and research programs; and
- 8. Supporting arrangement for practical internships for students of both Universities.
- B. Details of the implementation of any particular cooperation activity resulting from this Memorandum of Understanding shall be negotiated between the two institutions as and such specific case may arise, and will be outlined in a Supplementary Agreement between the institutions to be signed by designated offices. Supplementary Agreements are subject always to availability of sufficient funds at the respective institutions.
- C. The institutions recognize that this cooperative relationship may result in the development of various types of intellectual property and technology transfer. The institutions are committed to working in good faith to develop fair principles for dealing with intellectual property and technology transfer, including ownership, use, publication, and confidentiality. These principles will be developed in accordance with the parties' respective policies and collective agreements and will be incorporated into the Supplementary Agreements.
- D. Both parties shall designate a liaison office for this Memorandum of Understanding and for any Supplementary Agreements. For the University of Applied Forest Sciences Rottenburg the office shall be the Akademisches Auslandsamt (International Office). For the University of H.N.B Garhwal University, the liaison office shall be the Department of Forestry, Natural Resources.
- E. Both universities shall not charge enrollment fees for visiting students except administration fees. Travel expenses are borne by visiting students and staff. Both universities support students in application for scholarships.

Memorandum of Understanding

Page 2 of 3

- F. This Memorandum of Understanding reflects the commitment of the institutions to academic educational and research cooperation as of the date first written above.
- G. This Memorandum of Understanding may be amended by mutual written agreement.
- H. This Memorandum of Understanding shall continue for next 5 years and may be terminated at any time by either party, provided that notice of termination is provided by the notifying party to the other party at least ninety (90) days in advance of the date on which the termination is intended to become effective. Any termination of this Memorandum of Understanding shall not have effect on any arrangement in place at the time that the notice is provided, where the arrangement arises from any Supplementary Agreements resulting from this Memorandum of Understanding. Supplementary Agreements may only be terminated in accordance with the terms contained therein.

IN WITNESS WHEREOF the parties hereto have executed the Memorandum of Understanding as of the date first written above.

Witness Dr. A. K. Negil Head, Deptt of Foreolige NR. HNB-Gashval Driversty Sringer-Gashval HNB-GARHWAL UNIVERSITY, SRINAGAR (GARHWAL)

Per:

Prof. Annopurna Nautiya

UNIVERSITY OF APPLIED FORES

tund

Witness

Mrs. Tanja Münch International Office

FUR FORSTAIR SCHAFT

Per:

Itofan Prege

Prof. Stefan Ruge Head of International Office

Memorandum of Understanding

Garhw

hwal

कार्यालय आदेश

डा० आर०के० मैखुरी, साइंटिस्ट–जी, प्रभारी–क्षेत्रीय केन्द्र, जी०बी०पी०आई०एच०ई०एस० डी० श्रीनगर गढ़वाल को हाइफा विश्वविद्यालय हाइफा, इजराइल व हे०न०ब० गढ़वाल विश्वविद्यालय के मध्य हुए एम०ओ०यू० की प्रति एवं इस हेतु गठित समिति का कार्यालय आदेश संo HNBGU/RO/2019/529 व HNBGU/RO/2019/530 दिनांक 26.11.2019 इस आशय के साथ संलग्न कर प्रेषित किया जा रहा है कि उक्त समिति में आपको माननीय कुलपति महोदया के आदेशानुसार सह–सदस्य नियुक्त किया जाता है।

> आज्ञा से, कुलपति

हेमवती नन्दन बहुगुणा गढ़वाल विश्वविद्यालय, श्रीनगर गढ़वाल

संदर्भ सं0 : शैक्षणिक / 2020 / 🔿 /

दिनांक :01 जनवरी, 2020

प्रतिलिपि निम्नलिखित के सूचनार्थ एवं आवश्यक कार्यवाही हेतु प्रेषित :--

- 1. समिति के समस्त सदस्य महानुभावों को।
- 2. निदेशक-आई०क्यू०ए०सी० / एफ०डी०सी०।
- 3. संयुक्त कुलसचिव/समस्त उप कुलसचिव/सहायक कुलसचिव।
- सिस्टम मैनेजर को इस आशय के साथ प्रेषित कि उक्त अधिसूचना को विश्वविद्यालय की वेबसाइट पर अपलोड करना सुनिश्चित करें।
- 5. निजी सचिव-कुलपति, माननीय कुलपति महोदया के सादर सूचनार्थ।
- 6. निजी सचिव-कुलसचिव/वित्त अधिकारी।

1/1/202-कुलसचि



FRAMEWORK FOR ACADEMIC COLLABORATION

Between

HNB Garhwal (Central) University Srinagar, Garhwal, Uttarakhand, India

And University of Haifa, Haifa, Israel,

The HNB Garhwal (Central) University Srinagar, Garhwal, Uttarakhand, India represented by Prof. Annpurna Nautiyal), Vice-Chancellor, who is fully authorized to execute this agreement in her position(s) as Vice-Chancellor,

and

The University of Haifa, Israel, represented by Prof. Ron Robin, and Prof. Gustavo Mesch, who are fully authorized to execute this Agreement in their respective positions as President, and Rector.

(hereinafter referred to collectively as the "Parties" or individually as the "Party"),

Whereas

The Parties have decided to enter into this Framework of Academic Collaboration (the "Agreement") in order to promote, facilitate and consolidate international cooperation in education and research, based on the principles of reciprocity and mutual benefit.

Decide as follows:

Article 1. Areas of Cooperation

The areas of cooperation shall include, subject to mutual consent, any activity or program at either institution as considered feasible and desirable on either side in order to foster and develop the cooperative academic relationship between the two parties.

Article 2. Modes of Cooperation

The implementation of research and study programs determined in conformity with Article 1 may be carried out through the joint realization of research projects and teaching programs at the location of both Parties, according to the modalities specified in Article 4 of this Agreement.

More specifically, these are the modes of cooperation that may be used in a separate appendix to which the parties agree to, included the following:

- Exchange of scholars
- Joint research projects
- Joint teaching and/or supervision of students
- Joint participation in workshops and/or conference days
- Student mobility and prospective exchange of students (as may be mutually agreed upon)
- Participation of students in Study Abroad programs hosted by each institution.

Article 3. Liaison

Where needed, the Parties will appoint a faculty member as coordinator of the activities entailed in this general framework agreement who will also be responsible for facilitating and maintaining communication with the other Party for that purpose.

Article 4. Specific Cooperation Agreements

Nothing in this Framework Agreement shall be construed as creating any legal or financial relationship or commitment between the two Parties.

The terms of collaboration and the necessary funding for each program and activity shall be mutually negotiated, discussed and agreed upon in writing by both Parties in separate specific cooperation agreements prior to the initiation of any particular program or activity. Each institution will designate where needed, a lead coordinator for the activities or programs of potential cooperation that may form the basis for specific cooperation agreements to be executed between the parties.

Besides implementation modalities specifying academic, organizational, technical and financial aspects, specific cooperation agreements shall contain terms relating to intellectual property rights and procedures for publication etc.

Article 5. Principles

All participants will be treated in the same non-discriminatory manner in carrying out the provisions of the agreement, subject to the provisions of the policies and requirements of each of the institutions.

Article 6. Terms of Cooperation

6.1 Duration

This Framework Agreement shall commence on the later of the two dates of signature of the parties, this being the date when both parties have duly accepted the conditions laid down in this Agreement. The Agreement shall be in force for five (5) years and may be renewed or extended by mutual written consent of the authorized officials of the parties to it.

6.2 Amendments

Any amendment and/or modification of the Framework Agreement will require written approval of both Parties and shall be appended hereto.

6.3 Termination for convenience

Either Party reserves the right to terminate this Agreement at any time, for any or no reason, subject to ninety (90) days prior written notice to the other party hereto. Any program or activity that has already commenced pursuant to specific cooperation agreements executed between the parties hereto, if executed, shall survive termination of this Agreement and be completed to the best of both parties' abilities pursuant and subject to the terms of the relevant specific cooperation agreement.

Date: 24.11.2019

Signed: Orguntyn

HNB Garhwal (Central) University Srinagar, Garhwal, Uttarakhand, India

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Vice-Chancellor, Prof. Annpurna Nautiyal



Date: 24.11 2019.

Signed:

University of Haifa

President Prof. Ron Robin

Recto Prof. Gustavo Mesch

OFFICE ORDER

With approval of Hon'ble Vice Chancellor, a task force committee is hereby constituted with the following members for establishment and meetings of a Consortium of the Central Universities of the Indian Himalayan States.

Constitution of committee is as follows:

- 1. Prof. Y.P. Sundriyal, Deptt. of Geology Convener
- 2. Prof. A.R. Nautiyal, HAPPRC
- 3. Prof. R.P.S. Negi, Deptt. of History
- 4. Dr. Prashant Kandari, Deptt. of Economics

Dr. (A.K. Jha) Registrar

H.N.B. GARHWAL UNIVERSITY, SRINAGAR (GARHWAL) UTTARAKHAND (A Central University) Ref. No.: HNBGU/RO/2019/ SR9 Dated : 26/ 11 / 2019

Copy for information and necessary action to:-

- 1. All above concerned.
- 2. PS to VC for kind information of Hon'ble Vice-Chancellor.
- 3. Guard File.

Dr. (A.K. Jha) Registrar

OFFICE ORDER

With approval of Hon'ble Vice Chancellor, a task force committee is hereby constituted with the following members to work on joint projects and MoU with University of Haifa, Israel.

Constitution of committee is as follows:

- 1. Prof. M.C. Nautiyal, HAPPRC
- 2. Prof. Y.P. Sundriyal, Deptt. of Geology
- 3. Prof. R.C. Bhatt, Deptt. of History
- 4. Prof. R.P.S. Negi, Deptt. of History
- 5. Dr. B.S. Bhandari, Deptt. of Botany
- 6. Dr. R.S. Fartyal, Deptt. of Zoology
- 7. Dr. Prashant Kandari, Deptt. of Economics
- 8. Dr. Alok Sagar Gautam, Deptt. of Physics

Dr. (A.K. Jha) Registrar

H.N.B. GARHWAL UNIVERSITY, SRINAGAR (GARHWAL) UTTARAKHAND (A Central University)

Ref. No.: HNBGU/RO/2019/530

Dated : 26/ 11 /2019

Copy for information and necessary action to:-

- 1. All above concerned.
- 2. PS to VC for kind information of Hon'ble Vice-Chancellor.
- 3. Guard File.

Dr. (A.K. Jha) Registrar

BETWEEN

Department of Pharmaceutical Sciences, HNB Garhwal University (A Central University) Srinagar-Garhwal (UK) herein after called as HNBGU represented by its Principal herein named as party one

AND

ISF College of Pharmacy (An Autonomous College), Moga (Punjab) hereafter called ISFCP which is represented by its Director herein named as <u>party Two</u>

1. Preamble

Department of Pharmaceutical Sciences, HNB Garhwal University (A Central University) Srinagar-Garhwal is charged with responsibility of training technical and scientific manpower in front-line areas of pharmacy and is also contributing to the rapidly growing scientific and technological knowledge and professional excellence in pharmacy.

WHEREAS, ISFCP is premier Pharmacy Institute and engaged in providing platform to emerging scientists by day to day research, innovations, design, development and consultancy in the field of Pharmacy and related areas. UGC granted Autonomous College Status to Institute. NIRF-2019 ranked 23 in Pharmacy category in India.

WHEREAS, both HNBGU and ISFCP, now

- Recognizing the importance of research and development in the areas TO BE INTRODUCED HERE, as well as
- Imparting practical training to the Pharmacy students.
- Appreciating the need for creation of large reservoir of highly qualified manpower in all fields related to Pharmacy
- Desiring to club their efforts by pooling their expertise and resources,
- Thrend to form a nucleus for promoting excellent quality manpower in the fields of pharmacy.

NOW, THEREFORE, in consideration of the mutual promises made herein and of good and valuable consideration, the receipt and sufficiency of which both HNBGU and ISFCP hereby acknowledge, HNBGU and ISFCP hereby agree to sign a memorandum of understanding (MOU).

2. Objectives:

- 1. ISFCP will facilitate to provide Industry defined projects/Trouble shooting projects to research students /faculty of HNBGU
- 2. ISFCP will facilitate onsite Training and Industrial visit to Pharmacy students of HNBGU
- 3. ISFCP will facilitate for project work of research students
- 4. HNBGU and ISFCP will jointly organize Technical Seminars/ Workshops.
- 5. HNBGU will invite ISFCP experts for examination of M.Pharm. course.
- 6. HNBGU will schedule class room mentoring session by ISFCP members.
- 7. HNBGU will invite members of ISFCP as a part of Syllabus Committee.

Sent

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The parties hitherto agree as follows:

ARTICLE-I: SCOPE OF THE MOU

This MOU details the modalities and general conditions regarding collaboration between HNBGU and ISFCP for enhancing, within the country, the availability of highly qualified manpower in the areas of TO BE INTRODUCED HERE without any prejudice to prevailing rules and regulations in HNBGU and ISFCP without any disregard to any mechanism evolved and approved by the competent authorities under Govt. of India in so far as such mechanism applies to HNBGU and ISFCP. The areas of cooperation can be extended through mutual consent.

ARTICLE-II: SCOPE AND TERMS OF INTERACTIONS

Both HNBGU and ISFCP shall encourage interactions between the Pharmacy Students, Research fellows, faculty members and students of both the organizations through the following arrangements:

- a) Exchange of personnel through deputation as per the rules of the respective institute, for limited periods as mutually agreed upon;
- b) Organization of joint conferences and seminars;
- c) Practical training of HNBGU students at ISFCP;
- d) Joint guidance of student projects/thesis in TO BE INTRODUCED HERE and other areas of national interest at HNBGU by ISFCP on mutually agreeable terms.
- e) ISFCP would accommodate B. Pharm. & M. Pharm. students who have completed the 5th & 2nd semester respectively, of their programme in such a number that ISFCP deems convenient to it for the purpose of imparting practical training.
- f) ISFCP may depute its personnel as visiting faculty at HNBGU to teach any of the regular Course or specialized topics.
- g) HNBGU faculty, as well as research scholars, may also be allowed to enroll for their Ph.D. /M. Pharm. at IFSCP, subject to availability of seats, research facilities and subject to their fulfilling eligibility criteria and all other academic regulations of IFSCP. Further, IFSCP may request to design and teach a Course or Courses which it deems fit to enhance quality and performance of its employees. Such Courses may be run at any mutually convenient premises.
- h) ISFCP may avail library, Internet, computational facilities at HNBGU & reverse.
- i) The students will carry out part of their Ph.D. research work project at HNBGU and ISFCP depending on the nature of the work as per rules of the respective institute depending on facilities and requirements.
- j) Both HNBGU and IFSCP will be free to independently carry out follow-up research on the thesis work conducted under this scheme.
- k) If the outcome of a project related to product development, process, technology and design etc. which involves matter of secrecy and concern with security of the State and the Country, the same will not be allowed for publication/printing in any form.. If the outcome of a project results into an intellectual property, for which rights can be secured, it will be decided on case to case basis. Similarly, sharing of expenditure in securing such rights and income accrued through royalty etc by the parties under the law will be decided on case to case basis after mutual consultation.
- Research supervisors from both the Institutes will be the corresponding authors in any publication resulting from the collaborative work. All the efforts put by the student/s as a part of this MOU will be accounted for by way of reporting the work in thesis and/or paper publication except the part for which IPR needs be claimed.
- m) Neither of the supervisors will publish the work carried out under this MOU without knowledge of the other.

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payment conditions, etc. would be spelt out clearly before starting the activity.

ARTICLE-III: SHARING OF FACILITIES

- a) HNBGU and ISFCP shall make provisions to share their respective important R&D facilities in order to promote academic and research interaction in the areas of cooperation.
- b) HNBGU and ISFCP shall permit the sharing software and other materials and components developed in-house in the areas of cooperation, if permissible within the rules governing the two institutions. However, responsibility for safety of software and other materials during the exchange will rest on respective Head of academic department/section.
- c) HNBGU and ISFCP shall provide access to the library facilities to scientists, members of faculty and students as per the prevailing rules and norms in the respective institutes.

ARTICLE-IV: CO-ORDINATION OF THE PROGRAMME INCLUDING FINANCIAL ARRANGEMENTS

- a) The collaborative programme between HNBGU and ISFCP shall be coordinated by a coordination committee appointed by Directors of both the Institutes.
- b) Financial arrangements for each specific collaboration will be decided on a case to-case basis and brought on record in each case after due approval from heads of both the Institutions.

ARTICLE-V: EFFECTIVE DATE AND DURATION OF MOU

- a) This MOU shall be effective from the date of its approval by competent authorities at both ends.
- b) The duration of the MOU shall be for a period of 5 years from the effective date.
- c) During its tenancy, the MOU may be extended or terminated by a prior notice of not less than six months by either party. However, termination of the MOU will not in any manner affect the interests of the students/faculty/scientists who have been admitted to pursue a programme under the MOU.
- Any clause or article of the MOU may be modified or amended by mutual agreement of ISFCP and HNBGU.

ARTICLE-VI: IPR

Rights regarding publications, patents, royalty, ownership of software/design/product developed etc. under the scope of this MOU, shall be decided by the two parties by mutual consent.

ARTICLE-VII: CONFIDENTIALITY

During the tenure of the MOU both HNBGU and ISFCP will maintain strict confidentiality and prevent disclosure of all the information and data exchanged under the scope of this MOU for any purpose other than in accordance with this MOU.

Both HNBGU and IFSCP shall bind their respective personnel who come into possession or knowledge of any confidential information not to disclose the same to third parties without written approval of the disclosing party or use such confidential information for any use other than intended under this agreement or projects.

Further both HNBGU and ISFCP shall put in place adequate and reasonable measures to keep and store confidential information secure so as to prevent any unauthorized use.

Any amendment and/or addenda to the AGREEMENT shall be in writing and signed by the PARTIES hereto and shall only after such execution be deemed to form part of the AGREEMENT and have the effect of modifying the AGREEMENT to the extent required by such amendment or addenda.

ARTICLE-IX: RESOLUTION OF DISPUTES

- a) This agreement shall take effect and be construed in accordance with the Laws of India.
- b) The dispute or difference whatsoever arises between PARTIES in relation to or in connection with this AGREEMENT both the parties shall first try to resolve the dispute/difference amicably between themselves, failing which the matter shall be referred to and settled through arbitration. The arbitration proceedings shall be held in accordance with the provision of Indian Arbitration and Reconciliation ACT, 1996.

ARTICLE-X: MISCELLANEOUS

- The headings and sub-headings are inserted for convenience only and shall not affect the a) construction of this Agreement.
- Both HNBGU and ISFCP shall not, during the term of this Agreement directly or indirectly, b) solicit or offer employment or engagement to any of the personnel of other party without the prior consent in writing of that other party.
- No failure to exercise and no delay in exercising, on the part of a Party, and right, remedy, 0) power or privilege hereunder shall operate as a waiver thereof, nor shall any single or partial exercise of any right, remedy, power or privilege hereunder preclude any other or further exercise thereof or the exercise of any other right, remedy, power or privilege. The rights, remedies, power and privileges herein provided are cumulative and not exclusive of any right, remedies, powers and privileges provided by law.
- d) After this Agreement has been signed, all preceding understandings/negotiations and correspondence pertaining to it shall become null and void.

IN WITNESS WHEREOF PARTIES HERE TO HAVE ENTERED INTO THIS AGREEMENT EFFECTIVE AS ON THE DATE AND YEAR FIRST WRITTENABOVE.

Port: Abdul Faruk Signature

Mead Withmenkindidatenarmacautical sciences H.N.B. Garhwal University Srinagar (Garnwal), Utterakhand

Signature

Wilness: Pallei S. CINDU, ASR

Signature 22/2/2020 With seal and date . D. Gupta)

Director/Principal ISF College of Pharmacy, Mpga (ASignaturenous College)

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Sponsored Project

Memorandum of Understanding (MoU) Between Indian Institute of Technology Kanpur And

Hemvati Nandan Bahuguna Garhwal University, Srinagar Garhwal

This MoU is effective as of 15th Dce, 2021 (date) (Effective Date) by and between

Indian Institute of Technology Kanpur, an Institute established under a special act of Parliament of Republic of India, incorporated under the Institutes of Technology Act, 1961, having its office at Kanpur 208016, India hereinafter referred to as IITK, of the FIRST PART.

And

Hemvati Nandan Bahuguna Garhwal University (A Central University), an Institute under UGC Act 1956 and having its registered office at Srinagar, Uttarakhand, India, hereinafter referred to as HNBGU, of the SECOND PART.

The aforesaid institutions are hereinafter referred to individually as Party and collectively as the Parties.

Whereas Prof. T. H. Syed, Department of Earth Sciences, (hereinafter referred to as IITK **Principal Investigator (IITK PI)**) will initiate <u>the Ministry of Earth Science (MoES)</u> sponsored project titled "Characterization of Groundwater in seismically active regions of Uttarakhand, India: implications for earthquake induced variations (Project No. MOES-ES-2021185) (hereinafter referred to as Project). He and his research team at IITK will receive/disclose Confidential Information on behalf of IITK and will be solely responsible for non-disclosure of the Confidential Information received from HNBGU.

Whereas **Prof. H. C. Nainwal**, Department of Geology, School of Earth Science, HNB Garhwal University (hereinafter referred to as HNBGU **Principal Collaborator (PC)**) will facilitate all intellectual interaction and administer personnel/staff hired for the Project.

Whereas the Parties desire to record the broad terms and conditions that are jointly accepted and agreed to in this MoU as contained hereunder.

1. Definition:

(a) HNBGU know-how shall mean and include all know-how of methods, material, software,



Page 1 of 5

designs, patterns, formats, proprietary technical literature, and information developed, owned and provided by HNBGU, which are required for the Project.

(b) IITK know-how shall mean and include all know-how of methods, material, software, designs, patterns, formats, proprietary technical literature, and information developed, published or otherwise owned and provided by IITK, which are required for the Project.

(c) HNBGU Team shall mean Principal Collaborator (PC) and the personnel or staff deputed by HNBGU for the Project.

(d) Principal Investigator Research Team shall comprise the Principal Investigator (IITK PI), Co-Investigators and research personnel participating in the Project(s) from IITK.

2. Items/areas of collaboration/deliverables:

Technical specifications of the Project are given in Annexure A to this MoU.

3. Activities and roles of the Parties:

(a) IITK shall recruit the project assistant with the help of HNBGU. The recruitment formalities will be completed by IITK. It will have the financial responsibility for the payment of Rs. 20,000 + 16% HRA (per month) to the project assistant (personnel/staff) hired for the Project. Additionally, an amount up to Rs. 5,000/- per month will be provided to project assistant by IITK PI towards the TA/DA expenses to collect the samples form Agastyamuni, Gopeshwar and Saldhar (hot spring) from the project fund.

(b) HNBGU shall facilitate and guide the Project Assistant or staff hired by IITK for monthly collection of groundwater, surface water and hot spring samples from locations specified by the IITK Pl.

(c) HNBGU shall facilitate and guide the Project Assistant or staff hired by IITK to deliver the collected samples to the laboratory in National Institute of Hydrology (NIH), Roorkee for chemical analysis.

(d) The Project Assistant or staff hired by IITK will be stationed at HNBGU Srinagar and will work/collect the samples under the supervision of the PC in HNBGU.

(e) HBNGU will help IITK PI to make separate provisions with Govt. PG College Agastyamuni and Institute of Technology, Gopeshwar for the safety and security of the instrument and equipment installed at Agastyamuni and Gopeshwar, respectively.

(f) IITK shall be responsible for the maintenance and upkeep of the equipment installed or established for the Project.

(g) IITK shall have the ownership rights of all the monitoring stations established, and the equipment in it, utilizing Project funds.

4. Intellectual Property Rights:



Page 2 of 5

Ownership of and any intellectual property (including but not limited to data generated, confidential information, know-how, patents, copyrights, design rights, rights relating to computer software, and any other intellectual property rights) developed during the course of this Project shall be vested in IITK.

The authorship of scientific publication(s) resulting from this project will be shared with HBNGU PC and other faculties involved in the project.

Any utilization or publication regarding such intellectual property shall be possible only with the prior approval from IITK PI.

5. Effective date, duration, termination of the MoU:

The MoU shall be effective from the Effective Date upon signatures and shall remain in force for 3 years or the till the project continues whichever is less. The Parties shall extend the MoU by such a period equal to the period in case the Project gets extended by written agreement signed by both the Parties.

The project work may be terminated by either party by giving the other party a written notice of 60 days. However, both parties will ensure that the provisions of this MoU shall continue to apply to all activities in progress until their completion. Clauses relating to Intellectual Property Rights, governing laws shall survive the termination or expiration of this MoU.

6. Conflict Resolution:

This MoU is subject to Indian law. The parties will try to settle all disputes concerning this MoU in an amicable way. In case of any dispute, the same shall be referred to the Vice Chancellor of HNBGU and the Director, IITK or his nominee for arbitration. Reference made shall be deemed to have been made under the provisions of the Arbitration and Conciliation Act, 1996 or any statutory modification/re-enactment thereof and rules made there under. The award of the arbitrator shall be binding on both the parties. In case, however, the arbitrators are unable to come to a conclusion, then they will appoint an umpire whose decision shall be final and binding on both the parties.

7. Force Majeure:

Each party shall be excused from performance of the MoU only to the extent that the performance is prevented by conditions beyond reasonable control of the affected party. The party claiming excuse for the delayed performance will promptly notify the other party and will resume its performance as soon as performance is possible.



Page 3 of 5

Seal of the parties

In witness thereof, the parties hereto have signed this MoU on the date, month and year mentioned hereinbefore.

For and on behalf of HNBGU

Signature Name: Dr. A. K. Khanduri Designation: Registrar (HNBGU) Date: 14/102/1 Registrar ** (A Central University) Designation: Registrar (HNBGU) Date: 14/102/2021 For and on behalf of IITK

अधिष्ठाता DEAN अनुसंधान एवं विकास Research & Development आई० आई० टी० कानपुर I, I. T. KANPUR

Signature Name: Prof. A. R. Harish Designation: Dean, R & D, IIT Kanpur Date:

In the presence of Witness (from HNBGU)

Signature Name: Dr. H. C. Nainwal Designation: Professor Date: 24 In the presence of Witness (from IITK)

Syed Tajdan Hassan

Signature Name: Dr. Tajdarul Hassan Syed Designation: Associate Professor (PI) Date:

Enclosed:

Annexure A: Technical specifications of the project Annexure B: Project sanction letter from Ministry of Earth Science (MoES)



Page 4 of 5

Annexure A

Technical specifications of the Project

- 1. Objectives: To ensure successful completion of the project <u>titled "Characterization of</u> <u>Groundwater in seismically active regions of Uttarakhand, India: implications for</u> <u>earthquake induced variations</u>
- 2. Proposed activity with timeline:

SI. No.	Proposed activity	Timeline
1	Obligations/commitments on the part of the HNBGU:	January, 2022 – December, 2024
	To provide expertise for field sampling, exchange of ideas, hiring of necessary personnel (project assistant) to ensure monthly/bi-monthly collection of groundwater, surface water and hot spring samples, and delivering the samples to Laboratory of Dr. Sudhir Kumar in National Institute of Hydrology, Roorkee for chemical analysis	
2.	Obligations/commitments on the part of IITK: To provide monthly financial support for the salary of the project assistant and maintenance of equipment, and intellectual support to enable future collaboration.	January, 2022 – December, 2024



B Anneuve

MoES/P.O.(Seismo)/1(321)/2017 Government of India Ministry of Earth Sciences Seismology Division

Prithvi Bhawan, Lodhi Road, New Delhi – 110003. Dated: 30/04/2020

ADMINISTRATIVE ORDER

Sub.: Financial assistance for the research project entitled, "Characterization of Groundwater in seismically active regions of Uttarkhand, India: Implications for earthquake induced variations."

P1: Dr. Tajdarul Hassan Syed, Department of applied Geology, IIT (ISM), Dhandbad. Co-P1: Prof. Dr. Sudhir Kumar, National Institute of Hydrology (NIH), Roorkee. Co-P1: Shri Tathagata Chakraborty, SAC- ISRO, Ahemdabad.

Approval of the Competent authority is hereby conveyed under Rule 20 of the Delegation of Financial Power Rules, 1978, for the above-mentioned project at a total cost of Rs.96,41,146/- (Rupees ninety six lakh forty one thousand and one hundred forty six only) for a period of three years i.e.2020-2023. The items of expenditure for which the total allocation of Rs. 96,41,146/- has been approved are given below:

S.No.	ltem	l year	II year	III year	Total
А	Recurring				Amount in Rs
1,	Manpower JRF @31,000/- pm with HRA @1 6% SRF @35,000/- pm with HRA @1 6%	4,31,520/-	4,31,520/-	4,87,200/-	13,50,240/-
	Project Assistant@20,000/- pm wi th HRA @16%	2,78,400/-	2,78,400/-	2,78,400/-	8,35,200/-
2.	Consumables	20,000/-	20,000/-	20,000/-	60,000/-
3.	Travel	1,50,000/-	1,70,000/-	1,70,000/-	4,90,000/-
4.	Contingency	50,000/-	50,000/-	50,000/-	1,50,000/-
5.	Other Costs (a)Analytical charges	5,00,000/-	5,00,000/-	5,00,000/-	15,00,000/-
	(b) Drilling of two boreholes upto 100 m depth and Construction of t wo Piezometers	19,61,800/-			19,61,800/-
	Total (A)	33,91,720/-	14,49,920/-	15,05,600/-	63,47,240/-
B.	Non Recurring				
1.	Groundwater monitoring sonde with telemetry and two years AMC	26,01,406/-	-	-	26,01,406/-
2.	Portable water quality analysis kit	1,65,500/-		-	1,65,500/-
3.	Computers and peripherals	2,50,000/-			2,50,000/-
4.	UPS	1,27,000/-			1,27,000/-
	Total (B)	31,43,906/-	Nil	Nil	31,43,906/-
C.	Overhead	50,000/-	50,000/-	50,000/-	1,50,000/-
	Grand Total(A+B+C)	65,85,626/-	14,99,920/-	15,55,600/-	96,41,146/-

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2. Objectives

- Monitor groundwater level and groundwater quality in the active seismic belts of Uttarakahand at 2 sites.
- Analyze and characterize space-time variations in groundwater with respect to contemporary seismic events.
- III. Develop a process-based understanding of the observed anomalies and the mechanisms triggering them.
- Explore the potential to utilize groundwater anomalies for the development of precursor for earthquakes.

")

3. The expenditure will be booked under the following head:

3425	- Other Scientific Research (Major Head)
60.200	- Assistance to Other Scientific Bodies (Minor Head)
52	- Research Education and Training Outreach (REACHOUT
52.00.31	- Grants-in-aid-General for the year 2020-21

Other Terms & conditions:

- Utilization certificate and statement of expenditure for the expenditure incurred shall be submitted periodically in the prescribed format.
- II. The grantee organization will maintain separate audited account for the project and the entire amount of grant will be kept in an interest bearing bank account. The interest earned/accrued should be reported to MoES (financial year wise) while submitting the Statement of Expenditure/ Utilization Certificate.
- III. MoES reserves sole right on the assets created out of grants. Assets acquired wholly or substantially out of government grants (except those declared as obsolete and unserviceable or condemned in accordance with the procedure laid down in GFR 2017), shall not be disposed of without obtaining the prior approval of MoES.
- Allocation of budget is tentative and subject to changes after review of progress, both physical and financial as recommended by the Committee of Experts.
- V. As per Rule 230(8) of GFR 2017, all interest or other earning against Grant-in-aid released to the grantee Institution should be mandatorily remitted to the Consolidated Fund of India immediately after finalization of the account. It should not be allowed to be adjusted against future release.
- VI. A list of equipments procured and other assets created for this project from the funds released shall be provided to MoES in the prescribed format as per GFR. The ownership in the physical and intellectual assets created or acquired out of MoES funds shall vest in MoES.
- VII. All equipments / assets procured out of MoES grant shall be maintained by the grantee institute and will be available for use by other researchers.
- VIII. All equipments / assets procured from MoES fund shall be maintained in the stock register of the grantee Institute and should not treat such assets as their own assets in their Book of Accounts but should disclose their holding and using such assets in the Notes to Accounts specifically. No asset/equipment shall be diverted and /or disposed of without prior approval of the competent authority in MoES.
- Codal Provisions as conveyed in GFR, manual on Policies and Procedures for purchase of Goods and Gol instructions issued from time to time for services and requisition of research personnel shall be ensured.
- IX. This sanction is subject to the condition that the grantee organization will furnish to the Ministry of Earth Sciences, financial year wise Utilization Certificate (UC) in the proforma prescribed as per GFR 2017, audited statement of expenditure (SE) along with up to date progress report at the end of each financial year duly reflecting the interest earned / accrued on the grants received under the project and DBT details of the manpower as per the enclosed proforma.

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- X. Expenditure, Advance and Transfer (EAT) module of PFMS should be activated for the grants released under the project and expenditure under the project shall be monitored through EAT module.
 XI. While submitting Utility of the submitting Utility of t
- X1. While submitting Utilisation Certificate/Statement of Expenditure, the organisation must ensure submission of supporting documentary evidences with regard to purchase of equipment/capital assets as per the provisions of GFR 2017. Subsequent release of grants under the project shall be considered only on receipt of the said documents.
- XII. The Position of project staff is co-terminus with the duration of the project & MoES would have no liability towards such manpower costs beyond the duration of the project.
- XIII. Overhead expenses are meant for the host Institute towards the cost for providing infrastructural facilities and benefits to the staff employed in the project, etc.
- XIV. While implementing the programme all relevant procedures will be followed and the Ministry shall be apprised of the progress of the project from time to time.
- XV. The account of the grantee organisation shall be open to inspection by the sanctioning authority and audit (both by C&AG of India and Internal Audit by the Principal Accounts Office of the MoES), whenever the organisation is called upon to do so, as laid down under Rule 236(1) of General Financial Rules 2017.

5. This issues with the concurrence of IFD Vide their Concurrence Dy. No. 30(R)/IFD/6660/2020-21/REACHOUT dated 28/04/2020 and approval of the Secretary vide Dy. No.6660/Secy./2020-21 dated 29/04/2020.

(R. Bagavathi) Under Secretary to the Govt. of India

To,

Pay and Accounts Officer, MoES, New Delhi - 110 003

Copy forwarded for information and necessary action to:

- The Principal Director of Audit, Scientific Department, IIIrd Floor, AGCR Building, IP Estate, New Delhi – 110 002.
- 2. The Director, Indian Institute of Technology (Indian School of Mines) Dhanbad, Dhanbad-826004, Jharkhand.
- 3. Dr. Tajdarul Hassan Syed, Department of Applied Geology, Indian Institute of Technology (Indian School of Mines) Dhanbad, Dhanbad-826004, Jharkhand.
- 4. Prof. Dr. Sudhir Kumar, Scientist-G, Head, Hydrological Investigations Division, National Institute of Hydrology (NIH), Roorkee-247667, Uttarkhand.
- 5. Shri Tathagata Chakraborty, Scientist/Engineer 'SC', Space Applications Centre (SAC) ISRO, Ahemdabad-380015, Gujarat.
- 6. Cash section, MoES (with two spare copy of the sanction for making necessary payment to the grantee.)
- 7. Sanction Folder/ File copy.
- 8. Head (REACHOUT), MoES, New Delhi.

(R. Bagavathi) 30/4/2020

Under Secretary to the Govt. of India

MoES/P.O.(Seismo)/1(321)/2017 Government of India Ministry of Earth Sciences Seismology Division

Prithvi Bhawan, Lodhi Road, New Delhi – 110003. Dated: 30/04/2020

SANCTION ORDER

Sub.: Financial assistance for the research project entitled, "Characterization of Groundwater in seismically active regions of Uttarkhand, India: Implications for earthquake induced variations."

Pl: Dr. Tajdarul Hassan Syed, Department of applied Geology, IIT (ISM), Dhandbad. Co-PI: Prof. Dr. Sudhir Kumar, National Institute of Hydrology (NIH), Roorkee. Co-PI: Shri Tathagata Chakraborty, SAC- ISRO, Ahemdabad

In pursuance of A.O. of even no. dated 30/04/2020, Sanction of Competent authority is hereby conveyed under Rule 20 of the Delegation of Financial Power Rules, 1978, for the release of an amount of Rs.65,85,626/- (Rupees sixty five lakh eighty five thousand and six hundred twenty six only) to the Director, Indian Institute of Technology (Indian School of Mines) Dhanbad, Dhanbad as the first installment in the CFY 2020-21 for implementation of the project as follows:

S. No	Head	Amount in Rupees
1.	Salary	7,09,920/-
2.	Consumables	20,000/-
3.	Travel	1,50,000/-
4.	Contingency	50,000/-
5.	Other Costs	24,61,800/-
6.	Non-Recurring	31,43,906/-
7.	Overhead	50,000/-
8.	Total	65,85,626/-

2. This sanction is subject to the condition that the grantee organisation will furnish to the Ministry of Earth Sciences, Financial year wise Utilization Certificate (UC) in the proforma prescribed as per GFR 2017 and audited statement of expenditure (SE) along with up to date progress report at the end of each financial year duly reflecting the interest earned/ accrued on the grants received under the project. This is also subject to the condition of submission of the final statement of expenditure, utilization certificate.

3. The grantee organisation will have to enter & upload the Utilization Certificate in the PFMS portal besides sending it in physical form to this Division. The subsequent/final installment will be released only after confirmation of the acceptance of the UC by the Division and entry of previous Utilization Certificate in the PFMS.

4. If the grant has been released under capital head through separate sanction order under the same project for purchase of equipments(s), separate SE/UC has to be furnished for the released Capital head grant.

5. Overhead expenses are meant for the host institute towards the cost for providing infrastructural facilities and benefits to the staff employed in the project, etc.

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6.

The Grant-in-aid being released is subject to the condition that a.

A transparent procurement procedure in line with the Provisions of General Financial Rules 2017 will be followed by the Institute/Organisation under the appropriate rules of the grantee organisation while procuring capital assets sanctioned for the above mentioned project and a certificate to this effect will be submitted by the Grantee organisation immediately on receipt of

b. While submitting Utilization Certificate/Statement of Expenditure, the organisation has to ensure submission of supporting documentary evidences with regard to purchase of equipment/capital assets as per the provisions of GFR 2017. Subsequent release of grants under the project shall be considered only on receipt of the said documents.

7. It is desirable to have MoES nominee in the selection process for recruitment of JRF/SRF/RA/Scientists in the project.

8. The position of project staff is co-terminus with the duration of the project & MoES would have no liability towards such manpower costs beyond the duration of the project.

9. The grantee organisation will maintain separate audited account for the project and the entire amount of grant will be kept in an interest bearing bank account. The interest earned/accrued should be reported to MoES (financial year wise) while submitting the Statement of Expenditure/Utilization Certificate. As per rule 230(8) of GFR 2017, all interests or other earnings against Grants-in-aid released to the grantee Institution should be mandatorily remitted to the Consolidated Fund of India immediately after finalization of the accounts. It should not be allowed to be adjusted against future releases.

MoES reserves sole right on the assets created out of grants. Assets acquired wholly or 10. substantially out of government grants (except those declared as obsolete and unserviceable or condemned in accordance with the procedure laid down in GFR 2017), shall not be disposed of without obtaining the prior approval of MoES.

Expenditure, Advance and Transfer (EAT) module of PFMS should be activated for the 11. grants released under the project and expenditure under the project shall be monitored through EAT module.

The account of the grantee organisation shall be open to inspection by the sanctioning 12. authority and audit (both by C&AG of India and Internal Audit by the Principal Accounts Office of the MoES), whenever the organisation is called upon to do so, as laid down under Rule 236(1) of General Financial Rules 2017.

All the future correspondence regarding the project may be addressed to Head 13. (REACHOUT), MoES.

Data acquired under the project needs to be sent to Ministry regularly and it should not be 14. shared with any private agency/foreigner, without prior approval of MoES.

The expenditure involved is debit able to Demand No. 23 Ministry of Earth Sciences 15.

3425	- Other Scientific Research (Major Head)
60.200	- Assistance to Other Scientific Bodies (Minor Head)
52	- Research Education and Training Outreach (REACHOUT)
52.00.31	- Grants-in-aid-General for the year 2020-21

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16. The amount of Rs.65,85,626/- (Rupees sixty five lakh eighty five thousand and six hundred twenty six only) sanctioned in para 1, will be drawn by DDO, MoES and will be disbursed to the Director, Indian Institute of Technology (Indian School of Mines) Dhanbad, Dhanbad through RTGS as per following details:

Name of the Bank	:	Canara Bank
Type of Account	:	Savings Account
IFSC Code	:	CNRB0000986
Account No.	:	0986101009746
MICR No.	:	826015003
Branch Name	;	Canara Bank, Serraidhela Branch, Dhanbad-828127

17. As per Rule 234 of GFR 2017, this sanction has been entered at S. No. 04 Page No. 66 in the register of grants maintained in the Division of the scheme (**REACHOUT**).

18. This issues with the concurrence of IFD Vide their Concurrence Dy. No. 30(R)/IFD/6660/2020-21/REACHOUT dated 28/04/2020 and approval of the Secretary vide Dy. No.6660/Secy./2020-21 dated 29/04/2020.

Barrow Solywa

(R. Bagavathi) Under Secretary to the Govt. of India

To,

Pay and Accounts Officer, MoES, New Delhi - 110 003

Copy forwarded for information and necessary action to:

- The Principal Director of Audit, Scientific Department, Illrd Floor, AGCR Building, IP Estate, New Delhi – 110 002
- The Director, Indian Institute of Technology (Indian School of Mines) Dhanbad, Dhanbad-826004, Jharkhand.
- Dr. Tajdarul Hassan Syed, Department of Applied Geology, Indian Institute of Technology (Indian School of Mines) Dhanbad, Dhanbad-826004, Jharkhand.
- Prof. Dr. Sudhir Kumar, Scientist-G, Head, Hydrological Investigations Division, National Institute of Hydrology (NIH), Roorkee-247667, Uttarkhand.
- Shri Tathagata Chakraborty, Scientist/Engineer 'SC', Space Applications Centre (SAC) -ISRO, Ahemdabad-380015, Gujarat.
- 6. Cash section, MoES (with two spare copy of the sanction for making necessary payment to the grantee.)
- 7. Sanction Folder/ File copy.
- 8. Head (REACHOUT), MoES, New Delhi

Under Secretary to the Govt. of India





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- JIVANTI WELFARE AND CHARITABLE TRUST
- Article Others
- Not Applicable
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- : JIVANTI WELFARE AND CHARITABLE TRUST
- Not Applicable
- JIVANTI WELFARE AND CHARITABLE TRUST
- : 100
 - (One Hundred only)

Director



MEMORANDUM OF UNDERSTANDING

Please write or type below this line __

This Memorandum of Understanding is made and executed on this 15th September of Two Thousand Twenty at New Delhi BETWEEN JIVANTI WELFARE AND CHARITABLE TRUST (JWCT), a Trust under India Trust Act and having its registered office at 8/3 Asaf Ali Road, New Delhi-110002- hereinafter referred to as the "JWCT" in this MoU which expression shall unless repugnant to the context include its successors, nominees and assignees, the PARTY OF THE FIRST PART, For Jivanti Welfare And Charitable Trust

Statutory Alert:

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- The authenticity of this Stamp Certificate should be verified at "www.shcilestamp.com". Any discrepancy in the details on this Certificate and as available on the website renders it invalid.
 The onus of checking the legitimacy is on the users of the certificate.
- 3. In case of any discrepancy please inform the Competent Authority.

AND

HIGH ALTITUDE PLANT PHYSIOLOGY RESEARCH CENTRE, Post Box 14, Srinagar-246174, Uttarakhand, a research centre established by Hemvati Nandan Bahuguna Garhwal University (A Central University), hereinafter the research centre referred to as the "HAPPRC" the PARTY OF THE SECOND PART in this MoU, represented by its Director, which expression shall unless excluded by or repugnant to the context be deemed to mean and include its executors, administrators, representatives, successors and assigns etc.

WHEREAS the JWCT is focused mainly on fulfilment of welfare and charitable obligations towards society at large through advancement of equal opportunities for education, providing food, healthcare, medical care, ensuring environmental sustainability, enhanced vocational skills and advancement of any other objects of general public welfare.

WHEREAS JWCT has adopted conservation and sustainable development of biological diversity as part of **Dabur India Limited** Corporate Social Responsibility and has designed an integrated programme through a community centric Project based approach and for this purpose approached the HAPPRC for a collaborative cultivation and conservation programme in Uttarakhand.

WHEREAS, both JWCT and HAPPRC have mutually agreed for a collaborative project in Uttarakhand state for consecutive three financial years i.e. 2020-21, 2021-22 & 2022-23.

AND WHEREAS the parties herein have arrived at the modalities and agreed to document operative modalities, mutual responsibilities and obligations for these activities through this Memorandum of Understanding.

NOW THIS MEMORANDUM OF UNDERSTANDING WITNESSETH AS UNDER: -

- 1. That the signing parties to work together with a common goal of ensuring sustainable resource augmentation of few selected medicinal plant species through mass scale propagation, plantation and conservation of species and thereby ensure improvement in livelihood opportunities for regional communities and environmental sustainability.
- The species were identified for inclusion under the project jointly by JWCT and HAPPRC. Both the signing parties agree to promote all conservation approach and resource augmentation for the selected species in the private lands owned by communities and other mutually agreed locations within the selected region of Uttarakhand state.
- 3. In the context of Medicinal plants, the signing parties have identified the following species and same to be included on priority under this Memorandum of Understanding:
 - 3.1. Species for standardization of mass scale development of Quality Planting Material (QPM) and further cultivation programme in farmers and community owned land. Further an important aspect WEIKHIZOSPHERIC Studies of the below listed plants:

Director

- 3.1.1. Jatamansi (Nardostachys grandiflora)
- 3.1.2. Vatsanabha/ Mitha vish (Aconitum balfourii)
- 3.1.3. Atis (Aconitum heterophyllum)
- 3.1.4. Kutki (Picrorhiza kurrooa)
- 3.1.5. Kuth (Saussurea costus)
- 3.1.6. Sugandhbala (Valeriana wallichii)
- 3.2. The above species were identified for inclusion under the project jointly by JWCT with consultation with Dabur India Limited and HAPPRC. Both the signing parties agree to promote the conservation and resource augmentation for the above species in the lands owned by tribal community/farmers preferable in the high altitude region as-Bageshwar, Nainital, Tehri, Rudraprayag, Pauri, Pithoragarh and Chamoli districts of Uttarakhand (As per suitable climatic conditions for plantation of these selected species).

4. Technical Programme:

1

Mass scale planation of quality planting materials of high value threatened medicinal species is one of the viable options for conservation and sustainable utilization. It will also provide great livelihood opportunities to the poor and marginal farmers in rural areas. The present project has been proposed in these lines with the following technicalities

- 4.1. Mass scale production and multiplication arrangement of quality planting material through nursery development and utilization of other regional resource of public and private sector.
- 4.2. Distribution of plantlets for extensive plantations of the species in suitable areas.
- 4.3. Training and skill development of the farmers for plantation, maintenance/ after care harvesting and post- harvest management of the selected species.
- 4.4. To develop plantations and harvesting protocols of the species for future replications.
- 4.5. To ensure people's participation in long term maintenance of the plantation for their optimum production.
- 4.6. Income augmentation through livelihood support generation for small and marginal farmers (especially women).
- 4.7. Conduct Research and Domestication of the selected species for promoting cultivation.
- 4.8. Rhizospheric studies of Vatsanabh, Jatamansi etc. will be carried out in joint collaboration between Dabur & HAPPRC

5. Responsibilities of JWCT:

- 5.1. All the responsibilities assigned to **JWCT** under this MoU will be fulfilled through assistance from M/s Dabur India Limited.
- 5.2. Technical support to **HAPPRC** in project for effective deployment of the project through our regional resource and from HO on need basis.
- 5.3. Selection of elite germplasm basis the scientific assessments & technical inputs for mother stocks availability and collection assistance as well. For Jivanti Weifare And Charitable Trust

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5.4. Financial grants as shown under clause-7.

6. Responsibilities of HAPPRC:

- 6.1. Awareness creation among local communities about high conservation value of the MAP species.
- 6.2. To initiate people's participation-based activities on niche area plantations, sustainable harvesting and management practices on the species.
- 6.3. Identification of planting location in the project areas and community mobilization. For this purpose, **HAPPRC** is free to assign the project to existing village institutions like Self Help Groups (SHGs) and Village Organizations (VOs). They will be working as "Supervisors" and report plantation status on a regular basis.
- 6.4. Establishment of nursery for mass scale development of Quality Planting Material (QPM) as per targets.
- 6.5. Arrangement of QPM for the selected species through own nursery production/arrangement, distribution and monitoring. For the purpose the society can explore the other facilities and outsource the QPM from authentic source. **HAPPRC** will ensure the number of sapling as per the plantation target mentioned in Annexure 1.
- 6.6. Extensive plantation of the species on suitable sites with proper long-term monitoring and maintenance.
- 6.7. Best planting season should be ensured by **HAPPRC** to overcome the mortality. Any number difference in the planting target will meet out in next consecutive year.
- 6.8. Training and capacity building of the selected farmers for plantations and sustainable harvesting.
- 6.9. Development and publication of plantation and harvesting protocols on the species for future replications.
- 6.10. Documentation of baseline data with reference to demographic profile, income levels and goal setting in terms of livelihood promotion.
- 6.11. Periodical project monitoring in terms of progression and to redress the constraints in programme implementation.
- 6.12. Networking with Government and Village level institutions for smooth implementation of programme
- 6.13. HAPPRC is free to arrange the targeted saplings from Forest department nursery/other reliable agencies in case of non-production/less production at their own nursery.

7. Financial Aspects:

- 7.1. The financial needs of the project have been jointly worked out by the signing parties. Accordingly, the total project cost is pegged at Rs. 31,03,100/- (Thirty one lakhs three thousand and one hundred only) to be spent over a period of three financial years, detailed in Annexure 1.
- 7.2. The fund will be released as annual grant of Rs. 8,52,500/-, 10,23,000/- and 12,27,600/- respectively for 3 consecutive financial year of the tenure (2020-21, 2021-22 and 2022-23). The total annually grant will be released in two equal For Jivanti Welfare And Charitable Trust

Director

instalments. First 50% will be released within three weeks of signing the MoU and rest 50% grants will be released after due physical verification and receiving of UC of last given grant/s as on authentic format along with technical report till time. Financial obligation will be same for the year 2 and 3.

- 7.3. The cost assessments shown for Year-2 and Year-3 are tentative & calculated on the assumption that overall scope & deliverables of the project would be maintained as such. The budgetary components shall be revisited in the event of any revision in the project component.
- 7.4. HAPPRC will ensure the submission of receipt of the grant and utilization certificate after the completion of targeted work with complete report on quarterly basis.
- 7.5. The HAPPRC is also supposed to adhere with the timelines proposed under the MOU detailed in Annexure 2 and Annexure 3.

8. The Project Deliverables:

- 8.1. The plantation in the selected areas of Uttarakhand with an aim to natural resource augmentation and supplement the livelihood of local community through sustainable harvesting.
- 8.2. Methods to develop sustainable plant part collection in an eco-friendly manner may be by engaging with other partners.
- 8.3. Both the signing parties to abide by the same and work together to ensure that, the targets of the project are met effectively.
- 8.4. Submission of report has to be ensured by HAPPRC on quarterly basis as per the prescribed format under CSR norms.
- 8.5. Social benefit aspects for the outcome of project.
- 8.6. Sharing of scientific data outcome and publication from this project.

9. Amendments & Termination Clauses:

- 9.1. The MoU shall stand valid up to 31st March 2023 only.
- 9.2. The MoU can be amended by mutual consent among the signing parties at any point of time during its tenure.
- 9.3. Either of the signing parties may express its intent to terminate the agreement with a prior notice of at least, 2 months. During such notice period, both parties discuss and agree upon modalities for terminating the same. HAPPRC agrees to refund the amount released by JWCT after deducting the costs of the project on pro-rata basis and also submit the Utilization Certificate against the grant spent already.
- 10. Force Majeure Clause: Both parties agree to terminate the project amicably and without touching any contentious issues in the events of major adverse situations which may not allow the sustenance of the project like but not limited to economic, political and natural factors which go beyond the control of both the parties.

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- 11. Publications & Publicity:
 - 11.1. Both the signing parties agree that, researcher involved in this study can publish research findings in public domain. Investigators/Institute will properly acknowledge the financial support received from the JWCT in the publications emanating from the research findings of the project.
 - 11.2. Patentable information will be pursued with mutual consultation between both parties.
 - 11.3. Similarly, both parties agree to desist from availing any publicity mileage upon the organizational reputation of the other party- by methods like but not limited to, usage of emblems/logos etc. without express consent.
- 12. That, both the parties agree to settle any differences of opinion through a process of mutual discussions and consultations.
- 13. That, in the event of any dispute and/or differences between the parties hereto, the same shall be referred to arbitration under the provisions of the Arbitration and Conciliation Act, 1996 and any amendment or statutory modifications made thereto. The courts at New Delhi alone shall have the jurisdiction for this purpose.

IN WITNESS WHEREOF the parties hereto have signed this Memorandum of Understanding on this day, month and year first above written.

Witness-1(i): Witness-1(ii): DR. Pankal 1 FlatNo, 417 Mahagun Mostro, Seep CC NU 6/a, U.P. India 201304 Witness-

For and On Behalf of Jivanti Welfare and Charitable Trust (JWCT)

A. SUD HARAM Name:

For and On behalf of High Altitude Plant Physiology Research Centre: (HAPPRC)

Prof. A. R. Nautiyal Central University Director/HOD, HAPPRCar (Garhwal) U.K. PIN - 246174

Prof. M.C. Nautiyal

Incharge, Research Cell, HNBGU

Prof. Annapurna Nautiyal Vice-Chancellor, HNBGU Vice Chancellor

H.N.B. Garhwei University (A Central University) Srinegar Gashwei-245174 Utbrokhand (Indiz)

Annexure-1: Financial Implications

Items	Year I	Year II	Year III	Total
A) Non-Recurring expenditure	X.		-	
B) Recurring expenditure				
 Planting materials and agriculture tools 	70,000	84,000	1,00,800	2,54,800
2. Salaries/Wages	6,00,000	7,20,000	8,64,000	21,84,000
3. Travel (POL and TA/DA) for visit to fields stations and farmers' fields	75,000	90,000	1,08,000	2,73,000
 Contingency (stationary, typing, printing, etc.,), consumables and miscellaneous items 	30,000	36,000	43,200	1,09,200
Total Rs. (A+B)	7,75,000	9,30,000	11,16,000	28,21,000
Institutional overhead charges (10%)	77,500	93,000	1,11,600	2,82,100
Total (Rs.)	8.52 lac	10.23 Lac	12.27 Lac	31.03 Lac

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Sr. No	Species	1 st	2 nd	3 rd	Scope in subseque nt years	Remarks
1.	Jatamasi (Nardostac hys grandiflora)	Seed collecti on and seed sowing (250 gm)	Seed collection and seed sowing. Sustaining seedlings and Transplantatio n (150000)	Maintenance of Saplings in the farmer's field and observation and trials for optimization of production (150000 saplings). Transplantation of seedlings against mortality.	quintal from 1.0 ha of land on the basis of 25 Kg productio	The grant will be released in two equal installme nts. The first grant will be
2.	Kuth (Saussurea costus)	Seed collecti on and seed sowing (1.0 Kg)	Seed collection and seed sowing. Sustaining seedlings (33000) and Transplantatio n (28000)	Maintenance of Saplings in the farmer's field and observation and trials for optimization of production (28000 saplings, 35 quintal from 1.0ha of land). Transplantation of seedlings against mortality.	35.0 quintal from 1.0 ha of land on the basis of 70 Kg productio n from one nali of land.	released after signing of MoU and second installme nt will be released after physical verificati on/evalu ation i.e. based on UC proportio nate to farmers training/ mobilizat ion, infrastruc ture/ logistic
3.	Kutki (Picrorhiza kurrooa)	Seed collecti on and seed sowing (200 gm)	Seed collection and seed sowing. Sustaining seedlings (13,80,000) and Transplantatio n (1,10,0000)	Maintenance of Saplings in the farmer's field and observation and trials for optimization of production (1, 10,000 saplings, 10 quintal production from 1.0 ha of land). Transplantation of seedlings against mortality.	10.0 quintal from 1.0 ha of land on the basis of 20 kg productio n from one nali of land.	
4.	Atis (Aconitum heterophyll um)	Seed collecti on and seed	Seed collection and seed sowing. Sustaining no on	Maintenance of Saplings in the farmer's field and observation and trials	2.5 quintal from 1.0 ha of land on the	progress and no. of plantatio

Annexure-2: Physical Target and Progress Verification

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		sowing (1.0 Kg)	seedlings (2,50,000) and Transplantatio n (1,60,000)	for optimization of production (1,60,000 saplings and 2.5 quintal from 1.0 ha of land). Transplantation of seedlings against mortality.	Kg productio n from one nali of	n and producti on achieved,
5.	Vatsnabh (Aconitum balfourii)	Seed collecti on and seed sowing (100 gram)	Seed collection and seed sowing. Sustaining seedlings and Transplantatio n (2500)	Maintenance of Saplings in the farmer's field and observation and trials for optimization of production (2500 saplings, 9 Kg from one nali of land). Transplantation of seedlings against mortality.	productio n from one nali of	
6.	#Tagar/Sug andhbala (Valeriana wallichii)	Seed collecti on and seed sowing (1.0 Kg)	Seed collection and seed sowing. Sustaining seedlings and Transplantatio n (1,70,000)	Maintenance of Saplings in the farmer's field and observation and trials for optimization of production (1,70,000 saplings, 22 quintal production from 1.0ha of land). Transplantation of seedlings against mortality	22.0 quintal from 1.ha of land on the basis of 22 Kg productio n from one nali of land.	

*The expected work should be completed within the stipulated/decided time. However, under certain special circumstances, in case the work is not completed, it may be carried forward to next year. #Seeds will be collect in the month of April-May in IInd year.

Points need to be noted:-

1. Cultivation target: Initially 1.0 ha of land for each species.

2. Large scale seedlings development: Depends on availability of seeds, germination and seedlings survivability.

3. Production: Expected production depends on growth and care of saplings/plants in farmers'

fields.

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4. Harvesting: Possibly after 3rd year but same may be harvested in fourth to fifth year also as per climatic condition of the area. The first right of refuse will of JWCT on procurement of the RM of the selected species.

5. Raising and maintenance of seedlings for transplantation: Seedlings of the proposed species will be raised inside the glass house facility of the institute in winter months and then transfer to the Field Stations, Pothibasa (2200m asl), Baniyakund (2460m asl), Chopta and Lumkundi (1250m asl) in the month of March-April for further growth and transplantation to farmers field as and when required.

6. Exchange of quality germplasm (Seeds/Roots/Rhizomes): Funding agency will provide the improved germplasm, if any, of proposed species to implementing agency for multiplication and transplantation into farmers' field.

6. Assurance for completing the proposed work: All the work related to project will be done honestly and on propriety basis for achieving the target within a stipulated time period.

7. Assurance for timely purchasing of farmers produce: On behalf of funding agency, project implementing agency will assure to farmers for timely purchasing of their produce.

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	Annual Mark Deservator	Activity Months		
S. No.	Annual Work Parameter	April-September	October-March	
1.	Seed collection and seed sowing			
	Jatamasi (Nardostachys grandiflora)		October - November	
	Kuth (Saussurea costus)	August-September to Mid-October		
	Kutki (Picrorhiza kurrooa)	September		
	Atis (Aconitum heterophyllum)		October- November	
	Vatsnabh (Aconitum balfourii)		October-Novembe	
	Tagar/Sugandhbala (Valeriana wallichii)	April-May		
2.	Sustainability of the seedlings (QPM) numbers in the nursery	To be done	To be done	
	Transplantation of the seedlings in the farmers' fields			
	Jatamansi (Nardostachys grandiflora)	April-August		
3.	Kuth (Saussurea costus)	April-August		
	Kutki (Picrorhiza kurrooa)	April-August		
	Atis (Aconitum heterophyllum)	April-August		
	Vatsnabh (Aconitum balfourii)	April-August		
	Tagar/Sugandhbala (Valeriana wallichii)	July-August		
4.	Sustainability of the saplings and replacements of mortality if any (species wise)	To be done	To be done	
5.	Sustainability of the saplings and production	To be done	To be done	

Annexure-3: Proposed work with Timeline

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(Dr. Vijay Kant Purohit) Sr. Scientific Officer & Project Investigator







Memorandum of Understanding (MoU) Between

High Altitude Plant Physiology Research Centre (HAPPRC) {Post Box No. 14, HNB Garhwal University, Srinagar 246174 Uttarakhand}

and

Cultivator Natural Products Private Limited (CNPPL)

{Plot No. 24 to 31 & 25 to 30, Khasra No. 135/1, Sonamukhi Nagar, Sangaria Fanta, Jodhpur 342013 Rajasthan}

As part of a constant endeavor to promote mutual understanding, promotion of scientific/ industrial collaboration and commercial cultivation of medicinal herbs, an agreement is hereby entered into on cooperation between High Altitude Plant Physiology Research Centre (HAPPRC), Hemvati Nandan Bahuguna Garhwal University, Srinagar (Garhwal), Uttarakhand, India represented by its Director Prof. A. R. Nautiyal and Cultivator Natural Products Private Limited (CNPPL), Jodhpur, Rajasthan represented by its Director Mr. Tarun Prajapati / Executive-CSR, Mr. Bhoja Ram.

This MoU is entered on the day 0.3/12/2.0.2.0 (DD/MM/YYYY) between High Altitude Plant Physiology Research Centre (HAPPRC) and Cultivator Natural Products Private Limited (CNPPL) empaneled for promotion of commercial cultivation and facilitating buyback farming arrangements of medicinal plants and exchange of industrial exposure to students of M.Sc., Medicinal and Aromatic Plants.

PREAMBLE

Whereas the High Altitude Plant Physiology Research Centre (HAPPRC) has decided to engage Cultivator Natural Products Private Limited (CNPPL) for the promotion of commercial cultivation and facilitating buyback farming arrangements of medicinal plants in the state of Uttarakhand under Organic and Fair For Life programme: certification standard for Fair Trade and responsible supply-chains (Name of programme) covering approximately 1000 farmers. Simultaneously, CNPPL will facilitate the students of M.Sc. Medicinal and Aromatic Plants about industrial exposure of the students.

High Altitude Plant Physiology Research Centre (HAPPRC) and Cultivator Natural Products Private Limited (CNPPL) will ensure that following key activities involved under commercial cultivation are duly followed: -

- 1. The selection of districts, blocks and villages will be done by the CNPPL in consultation with the HAPPRC.
- 2. The staffing and expenditure on the specified activities should be strictly in conformity with the process guideline of CNPPL.
- 3. A detailed survey of selected villages will be conducted regarding the socio-economic conditions, prospects and problems in agriculture production.
- Selection of farmers will be in such a way that Organic Grower Group (OGGs) will be organized on the basis
 of homogeneity of crop production. Each OGG so formed will consist of 25-500 farmers with common interest.
- CNPPL will work to encourage as a procurement agency under the Minimum Support Price (MSP) procurement.
- 6. The expenditure on Organic and other types of certificates will be borne by CNPPL.
- A labour type manpower (01 No.) should be deployed by the CNPPL for the said task (raising and maintenance of planting materials), which should be supervised and guided by technically qualified and experienced persons of HAPPRC.
- Pre decided payment to labour will be released by CNPPL after approved by HAPPRC. For Cultivator Natural Products Py. Ltd.

Director

- 9. HAPPRC will work with CNPPL to direct support (likes e.g. Agrotechniques of medicinal plants; Suggestions for project location of various medicinal plant species; arranging planting materials free of cost; technical supports; training to Farmers/Farmer Groups; etc.) for working needs of OGGs. HAPPRC will also work with all relevant stakeholders to achieve 100% empowerment and capacity building for members of OGGs.
- 10. HAPPRC to act as a single window for technical support, training needs, research and knowledge management to CNPPL. HAPPRC will provide all- round support to CNPPL engaged in promotion and development of commercial cultivation of medicinal plants. In particular, HAPPRC will create sustainable linkages between CNPPL and Farmers/Farmer Groups.
- 11. HAPPRC will provide seed, saplings and other planting material free of cost to producers for extending production.
- 12. CNPPL will be facilitate the M.Sc. Students of the HAPPRC about industrial exposure.

The MoU will come into effect from the date of most recent signature and is entered into for a period of 3 years. Thereafter it shall be automatically extended for an additional time period, if required. The period of validity of this agreement may be terminated with in this period by either party by giving at least six months' notice in writing to other party.

Notification of any change in Director's role may be made by letter without amending this MoU.

Cultivator Natural Products Private Limited (CNPPL), Jodhpur, Rajasthan and High Altitude Plant Physiology Research Centre (HAPPRC), HNB Garhwal University, Srinagar (Garhwal), Uttarakhand shall execute this MoU to the best of their knowledge and ability.

Place: Sninagar Garhwal

Date of Signing: 03/12/ 2020

On behalf of Cultivator Natural Products Private Limited (CNPPL), Jodhpur, Rajasthan

Director

Mr. Tarun Prajapati Director

On behalf of High Altitude Plant Physiology Research Centre (HAPPRC), HNB Garhwal University, Srinagar (Garhwal), Uttarakhand

Prof. A. R. Nautiyal Director/HODDirector

Prof. N. C. Nautiyal In-charge Research Cell

3-12.2020

Prof. Annapurna Nautiyal Vice-Chancellor Vice Chancellor HAUD Chancellor HAUD Chancellor HAUD Chancellor

MEMORANDUM OF UNDERSTANDING BETWEEN DEPARTMENT OF PHYSICS HEMVATI NANDAN BAHUGUNA GARHWAL UNIVERSITY AND THE GRADUATE SCHOOL OF HUMAN HEALTH SCIENCES TOKYO METROPOLITAN UNIVERSITY

The Department of Physics, Hemvati Nandan Bahuguna Garhwal University in India and the Graduate School of Human Health Sciences, Tokyo Metropolitan University in Japan entered into this Memorandum of Understanding (hereinafter referred to as "MoU") with the objective to promote academic cooperation and exchange, based upon the principles of equality and reciprocity, between the two institutions.

1. Scope of Cooperation and Exchange

A. Exchange of Academic Staff

Each institution may receive members of the academic staff of the other institution to give lectures, to engage in joint or individual research, or for other educational purposes.

B. Exchange of Students

Each institution agrees to accept students of the other institution for a short term study program and research visits within a period up to ninety (90) days (hereinafter referred to as "Short Term Program"). Any student exchange activities involving credit-bearing courses for one semester or one academic year shall not be included in the Short Term Program. The details of each Short Term Program shall be set forth in a separate written individual agreement between the two institutions.

C. Cooperation in Research

Both institutions shall strive to cooperate in research by exploring means of promoting symposia, research conferences, the exchange of academic information, the publication of research, and the collection and sharing of research materials and data.

D. Other Exchange

The institutions shall strive to cooperate in any other educational and academic exchanges to which they may both agree.

2. Financial Matters

This MoU imposes no financial obligation on either institution.

3. Data Protection

Both institutions acknowledge and agree that they need to collect, process, use, disclose and manage personal data relating to academic staff and students in accordance with the policies, rules and regulations of the host Institution, which that has agreed to receive students and staffs from the other institution for a period of exchange, and all applicable laws of the country where it is located solely for the purposes of the administration of academic cooperation and exchange under this MoU. Both institutions shall ensure that such personal data are held securely and confidentially at all times.

4. Intellectual Property Rights

This MoU shall not grant to either institution any license or right to the intellectual property of the other institution. Intellectual property rights associated with activities under this MoU should be determined based upon the existing policies of both institutions and subject to applicable laws. In the case of a conflict, intellectual property rights should be negotiated on a case by case basis.

5. Term, Termination, and Amendment

This MoU shall commence on 04/01/2021 and remain in force for five (5) years. Thereafter this MoU shall be renewed for a specific period by mutual agreement in writing between both institutions.

Either institution may terminate this MoU during the term of the MoU by giving written notice to the other institution at least twelve (12) months prior to the date such institution wishes to terminate. However, exchange students and staffs who have already been admitted prior to the proposed termination date shall be allowed to complete his/her study or research within twelve (12) months following the proposed termination date.

No amendments and modifications of any provision of this MoU shall be effective unless made in writing and duly signed by both institutions.

6. Dispute Resolution

Any dispute or claim arising out of or in relation to this MoU shall be resolved in good faith between the two institutions. If the dispute cannot be resolved informally, it shall be referred to a senior post-holder within each institution.

Any dispute or claim not resolved to the satisfaction to both institutions further to the dispute resolution efforts above shall be submitted to the Tokyo District Court, Japan.

7. Governing Law

Both institutions agree this MoU is governed by and construed in accordance with the laws of Japan without regard to the principles of the conflict of laws thereof. unless otherwise agreed and confirmed in writing by both institutions.

This MoU is drawn up in duplicate in English. Each institution shall retain each one (1) copy.

IN WITNESS THEREOF, both institutions have caused this MoU to be executed by their duly authorized representatives.

Signature

NAUTIYAL M C RCC Coordinator Hemvati Nandan Bahuguna Garhwal University

(DATE) MM /DD /YYYY

Signature

WATANABE Masaru Dean Tokyo Metropolitan University

(DATE) MM /DD /YYYY 02 122 12





INDIA NON JUDICIAL Government of Uttarakhand

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- Registrar HNBGU Srinagar Garhwal
- Article 4 Affidavit
- NA

0 (Zero)

- Registrar HNBGU Srinagar Garhwal
- NA
- Registrar HNBGU Srinagar Garhwal
- (One Hundred only)



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MEMORANDUM OF UNDERSTANDING



FOR INSTITUTIONAL COLLABORATION BETWEEN HEMVATI NANDAN BAHUGUNA GARHWAL UNIVERSITY (A CENTRAL UNIVERSITY) SRINAGAR (GARHWAL) UTTARAKHAND And

NATIONAL INSTITUTE OF TECHNOLOGY, UTTARAKAHD



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MEMORANDUM OF UNDERSTANDING

FOR INSTITUTIONAL COLLABORATION

BETWEEN

HEMVATI NANDAN BAHUGUNA GARHWAL UNIVERSITY (A CENTRAL UNIVERSITY) SRINAGAR (GARHWAL), UTTARAKHAND

AND

NATIONAL INSTITUTE OF TECHNOLOGY, UTTARAKHAND (NITUK)

Preamble:

This Memorandum of Understanding (MoU) is drawn up and agreed upon to establish the initial framework for cooperation between National Institute of Technology Uttarakhand, Temporary Campus, Government Polytechnic, Srinagar (Garhwal)-246174 and Hemvati Nandan Bahuguna Garhwal University, Srinagar (Garhwal) Uttarakhand.

Hemvati Nandan Bahuguna Garhwal University (HNBGU)

Hemvati Nandan Bahuguna Garhwal University was established as a State University vide U.P. State Government notification no. (10)/(865)/15/(75)(85)/64 dated 23 November 1973. The University has subsequently been upgraded to Central University by an Act of Parliament i.e. the Central Universities Act 2009. Since its inception, the University has shown commitment towards regional and community development which is inherent in its teaching courses, research agenda and other outreach and extension initiatives. The University, nestled in the lap of Himalayan ranges in the Garhwal region of Uttarakhand, is a residential cum affiliating institution of higher learning. The University has following three campuses distantly located from each other where the undergraduate, post-graduate and research programmes are being offered in different disciplines.

- 1. Birla Campus, Srinagar Garhwal with its extension at Chauras Campus,
- 2. B. Gopal Reddy (BGR) Campus, Pauri and
- 3. Swami Ram Teerth (SRT) Campus, Badshahithaul, Tehri.

National Institute of Technology, Uttarakhand (NITUK)

National Institute of Technology, Uttarakhand (NITUK), an Institute of National Importance is established in 2009 under the Act of Parliament by Ministry of Human

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Resource Development and is one among the 31 National Institutes of Technologies in the country. The Institute offers full time B. Tech. and M. Tech. Programmes in 05 different Engineering disciplines as well as Ph.D. programme in all Engineering, Sciences and Humanities disciplines.

Hemvati Nandan Bahuguna Garhwal University (herein after referred to as "HNBGU", which expressions shall mean and include, unless repugnant to the context or meaning thereof its successors and permitted assigns) has authorized Prof. Annapurna Nautiyal, Vice Chancellor, to enter into this MoU as the FIRST PARTY;

National Institute of Technology, Uttarakhand (herein after referred to as "NITUK", which expressions shall mean and include, unless repugnant to the context or meaning thereof its successors and permitted assigns) has authorized **Prof. Shyam Lal Soni, Director, NITUK** to enter into this **MoU** as the **SECOND PARTY**;

"HNBGU" and "NITUK" are hereinafter, wherever the context so admits, collectively referred to as the "Parties" and individually as a "Party".

AND WHEREAS the purpose of MoU is to establish an understanding of mutual cooperation between NITUK and HNBGU, providing a common platform for deriving mutual advantages in their pursuit of higher learning in general and benefiting their respective students and faculties, by way of exposure to each other's programmes.

1. SCOPE OF MoU:

The following areas of collaboration have been identified under this MoU.

- (i) <u>Faculty Exchange Programme</u>: The two parties will explore opportunities for interaction among members of faculty The total duration of visits from each side is expected to be approximately equal. Each such visit shall require approval of the respective Institutions/University.
- (ii) <u>Student Exchange Programme</u>: The two parties agree to participate in student exchange programme at the respective Institutions/University. Visits under such programmes may be for a short duration, such as summer/winter terms or short courses, A detailed proposal document on this has to be approved by the respective competent of the two Institutions before its implementation.
- (iii) Joint Research Projects: The two parties will explore opportunities of undertaking joint research projects including co-author publications and may seek research funding from external funding agencies. Each such research proposal shall require approval of the respective Institutions/University.

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- (iv) Joint Research Guidance for Ph.D/PG/UG Students: The faculty members of HNBGU and NITUK will explore opportunities for the joint research guidance to the Ph.D./PG/UG students. A detailed document related to such research guidance will be required by the competent authorities of both the organizations before implementation.
 - (v) Joint Academic Activities and Events: HNBGU and NITUK may formulate joint academic activities such as short course, faculty development programme, seminars, workshops, conferences, extracurricular activities etc. based on mutual interests and available expertise in respective Institute/University. They may also share and carry out joint research in technology for distance and computer-based learning.

(vi) <u>Miscellaneous</u>:

- a. Both the organizations shall endeavour to, as much as possible make available such facilities that will enable the faculty to obtain experience and training in respective departments and Institutes, including the use of its laboratories, libraries and e-resources etc.
- b. Both the organizations shall share the infrastructure facilities like hostels, guest house, playgrounds, complexes, seminar halls, etc. available with either Institute/University with permission of the competent authority.
- c. Defining new areas of collaboration that have not been foreseen, but can be beneficial to the Parties.

The areas of cooperation may be revised by mutual consent. However, specific programmes may require separate agreements detailed out and documented as annexures to this MoU.

2. NON-DISCRIMINATION

NITUK and HNBGU agree not to discriminate against any person because of age, sex, national origin, race, ancestry, color, religious creed disability or handicap, and sexual orientation. Neither Institution shall impose criteria for the exchange of faculty, staff and students that would violate the principles of non-discrimination.

3. CODE OF CONDUCT

Visiting student and faculty will abide by the codes of conduct of the host Institution/University.

4. INTELLECTUAL PROPERTY

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Each Institution/University will adhere to the intellectual property laws of its institute/nation. Intellectual property developed during the visit of an exchange student/researcher/faculty/staff will be governed by the rules of the host Institute unless otherwise specified.

Any Background Intellectual Property (BGIP) shall remain the sole and exclusive property of the Party to whom it belonged prior to the commencement of this MoU. If one Party receives any BGIP from the partner under a clearly defined non-disclosure agreement, necessary and reasonable care will be taken to protect the intellectual property received.

The two Institutions shall jointly own results and Intellectual Property generated thereof for clearly defined collaborative projects and exchange programmes. This joint ownership of Jointly Developed Intellectual Property (JDIP) also entitles each party to explore commercialization. However, transfer or sale of JDIP and associated sharing of revenue shall be governed by a separate agreement.

5. COORDINATION:

HNBGU and NITUK hereby nominate Chairman, RCC and Dean (R&C), NITUK as coordinators respectively for operating the various provisions of this MoU on behalf of their respective organizations. All formal communications will be exchanged only through these nominated representatives. The participants and Coordinators may change from time to time as decided by appropriate authority of the respective Institutions/University.

6. LEGAL STATUS

This document is a statement of intent to foster genuine and mutually beneficial cooperation and is not legally binding on both the parties. In the event of any dispute or difference arising in the implementation of the MoU, such disputes shall be resolved amicably by mutual discussions by the Head of both the Institutions. All such decisions shall be taken into account the status of students working/projects under this arrangement and the interest of such students/projects shall be guarded as much as possible.

7. DURATION, AMENDMENT AND TERMINATION:

a. The MoU remains in place for a period of five years (05) from the date of signing of the agreement and may be renewed for a further period of five years or such time period by mutual consent of the Parties, expressed in writing.

H.N.B.Gachwal University (A Central University) Srinegar (Gamwal)





- **b.** The terms of MoU may be modified/ amended at any time subject to mutual written agreement. Such modification/change shall be effective from the date on which both the parties execute them in writing.
- c. The MoU may be terminated by either party by giving three months advance notice in writing to the other Party.
- d. The amendment, termination and expiration of this MoU will not affect the terms of activities ongoing at the time of notification of amendment, termination and expiration, unless otherwise agreed upon between the Parties. In such case, both the parties shall ensure that the interest of students/projects under this MoU are safe guarded to the extent possible.

However, specific commitments made prior to such intimation shall be honored by both the partners including ensuring that any student at that time participating in the Programme is able to complete the term of the assignment and be assessed for it.

Now based on the aforementioned promise(s) the parties put their signatures on this MoU on March 16, 2021.

Signature: Date: 16 Marc

H.N.B.Garnwal University Dr. Ajay Kumar Khanduri,(A Conitai University) Registrar, Srinagar (Contrait HNBGU, Srinagar, Uttarakhand. Email ID: registrar@hnbgu.ac.in

Witness 1

Dr. Manoj Kumar Gupta Assistant Registration Processor Department of Mechanical Engineering HNBGU, Srinagar, Uttarakhand. Signature:

Dr. P. M. Kala Registrar, National Institute of Technology, Uttarakhand Email ID: <u>registrar@nituk.ac.in</u>

Witness 1 A Tore 16/03/2021 Dr. Dharmendra Tripathi Dean Research & Consultancy National Institute of Technology, Uttarakhand





Memorandum of Understanding (MoU)

between

the Swiss Federal Research Institute WSL, Switzerland

and

the Department of Geology, Hemvati Nandan Bahuguna Garhwal University, Srinagar Garhwal, 246174 India

The Swiss Federal Research Institute WSL, Zürcherstrasse 111, 8903 Birmensdorf, Switzerland (hereafter WSL) and the Department of Geology, Hemvati Nandan Bahuguna Garhwal University, Srinagar Garhwal, 246174 Uttarakhand, India (hereafter HNBGU) desire to engage in cooperative educational and research activities, for the mutual benefit of both institutions. The Parties have agreed upon the following:

- To encourage visits by faculty for the purpose of engaging in research and educational activities
- To support the exchange of Master and Doctoral students
- To foster the exchange of academic publications and scholarly information
- To develop joint research activities and to promote other academic activities, which enhance the above mentioned goals.

Terms of Cooperation

- The terms of cooperation for each activity implemented under this Memorandum shall be mutually discussed and agreed upon in writing by both parties prior to the initiation of that activity. Any such agreement entered into, as outlined above, will form an Appendix to this Memorandum of Understanding.
- This Memorandum does not result in any financial obligations. Each institution will be responsible for seeking funds to support its involvement in the cooperative activities contemplated under this Memorandum of Understanding, and all such activities will be dependent upon the budgetary appropriations of the parties.
 All activities shall be
- 3. All activities shall be in accordance with the regulations and policies of the WSL and HNBGU.

Development, Co-ordination and Execution of the Exchange

Responsible for the development, co-ordination and execution the specific activities for the purposes of this Memorandum of Understanding shall be:

Swiss Federal Research Institute WSL High Mountain Glaciers and Hydrology Group (HIMAL) Current Group Leader: Francesca Pellicciotti

Zuercherstrasse 111

CH-8903 Birmensdorf

E-Mail: francesca.pellicciotti@wsl.ch

Hemvati Nandan Bahuguna Garhwal University Department of Geology

Current Group Leader: Prof. H. C. Nainwal Srinagar Garhwal, Uttarakhand 246174, India nainwa61@gmail.com

Intellectual Property Rights

Information on research results and scientific materials (reports, articles, books) will be exchanged freely keeping in mind the mutually agreed provision of Intellectual Property Rights. All intellectual property solely conceived and/or developed by WSL during the course of this Agreement shall be owned by WSL. All intellectual property solely conceived and/or developed by HNBGU during the course of this Agreement shall be owned by WSL. All intellectual property solely conceived and/or developed by HNBGU during the course of this Agreement shall be owned by HNBGU. Intellectual property jointly conceived and/or developed by WSL and HNBGU will be jointly owned by WSL and HNBGU. Each party may use such property for research and scholarly purposes. WSL and HNBGU agree to collaborate towards the protection, if appropriate, and application of such intellectual property for commercial or other purposes on mutually acceptable terms to be negotiated in good faith between the parties.

Renewal, Termination, and Amendment

- This Memorandum becomes effective from the day the representatives of both universities affix their signatures and continue for an initial period of three (3) years, subject to review from time to time. At the end of three years, the agreement will automatically be extended for another three-year period unless otherwise determined.
- 2. This Memorandum may be revised through the mutual agreement of both parties and may be terminated by either party upon giving six months written notice signed by the presiding office of the notifying party. All modifications to this agreement must be in writing and signed by both parties.

On behalf of the

Swiss Federal Research Institute, WSL

Prof. Dr. Konrad Steffen

Director

On behalf of the

Hemvati Nandan Bahuguna Garhwal University

Prof. N.S. Panwar

Registrar

Date

Prof. Y. P. Sundriyal Head of Department of Geology

Dr. Francesca Pellicciotti

Date 22nd April 2020

Group Leader

2 April 2020

23 July 2020

MEMORANDUM OF UNDERSTANDING

between



Indian Council of Forestry Research and Education, Dehradun and



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Hemvati Nandan Bahuguna Garhwal University, Srinagar

Indian Council of Forestry Research and Education (hereinafter referred to as "ICFRE", which expression, shall unless repugnant to the context thereof, shall mean and include its successors and assignees) through the **Director General**, ICFRE, **Dehradun** of the First Part.

and

Hemvati Nandan Bahuguna Garhwal University (hereinafter referred to as "HNBGU", which expression, unless repugnant to the context thereof, shall mean and include its successors and assignees) through the Vice Chancellor, HNBGU, Srinagar of the Second Part.

ICFRE and HNBGU are collectively referred to as "Parties" and individually as "Party".

Background

This Memorandum of Understanding (MoU) sets for the terms and understanding between Indian Council of Forestry Research and Education and Hemvati Nandan Bahuguna Garhwal University (a Central University)to collaborate in applied research programs, capacity building and knowledge sharing on forests, biodiversity conservation, environment, ecosystems, climate change vulnerability, climate change mitigation and adaptation and livelihood of forest dependent communities and other subjects mutually agreed between the two parties.

Preamble

Whereas

Indian Council of Forestry Research and Education is an autonomous body of Ministry of Environment, Forest and Climate Change, Govt. of India and is an apex body in the national forestry research system. It has been undertaking the holistic development of forestry research through need based planning, promoting, conducting and coordinating research, education and extension covering various aspects of forestry. The council deals with the solutions based forestry research in tune with the emerging issues in the sector, including global concerns such as climate change, conservation of biological diversity, combating desertification and sustainable management and development of resources. Topical research by the council enhances public confidence in the ability of forest managers and researchers to successfully handle challenges related to natural resource management. ICFRE has nine Research Institutes and five Research Centres located in different bio-geographical regions of the country for catering to the forestry research needs of the nation. ICFRE has the following Institutes and Centres:

- Forest Research Institute (FRI), Dehradun 1.
- Arid Forest Research Institute (AFRI), Jodhpur 2.
- Tropical Forest Research Institute (TFRI), Jabalpur 3.
- Institute of Forest Genetics and Tree Breeding (IFGTB), Coimbatore 4. 5.
- Himalayan Forest Research Institute (HFRI), Shimla
- Rain Forest Research Institute (RFRI), Jorhat 6.
- Institute of Forest Productivity (IFP), Ranchi 7.
- Institute of Wood Science & Technology (IWST), Bengaluru 8. 9.
- Institute of Forest Biodiversity (IFB), Hyderabad
- 10. Forest Research Centre for Skill Development, Chhindwara
- 11. Forest Research Centre for Eco-Rehabilitation, Prayagraj
- 12. Forest Research Centre for Bamboo & Rattan, Aizawl
- 13. Forest Research Center for Livelihood Extension, Agartala
- 14. Forest Research Centre for Coastal Ecosystem, Visakhapatnam

Objectives of ICFRE:

- To undertake, aid, promote and coordinate forestry research, education and 1. extension leading to scientific and sustainable management of forest resources in the country.
- To align forestry research programs in the council with national priorities 2. including achievement of Sustainable Development Goals and combating climate change.
- To provide scientific advice and policy support to the central and state 3. governments aiding informed decision making in forestry matters of national importance and international commitments.
- To act as a repository of scientific knowledge related to forestry, environment 4. and climate change, and disseminate such knowledge to various stakeholders.
- To provide technical assistance and support to states, forest- based industries, 5. tree growers, farmers and others for forest protection, afforestation, agroforestry and allied activities.
- To develop appropriate forest based technologies, processes and products for 6. sustainable resource use, livelihoods and economic growth.
- To provide livelihood support to forest dependent communities through transfer 7. of scientific knowledge and appropriate forest-based technologies.
- To develop technically qualified human resource for forestry sector. 8.
- To promote forestry education in the country and facilitate universities in 9. improving quality through technical and financial support including development of uniform curricula.
- To provide consultancy and capacity building services in environment and forest 10. sector.

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- 11. To develop and maintain National Forest Library and Information Centre for forestry and allied sciences.
 - 12. To develop environment and forest extension programmes and promote the same through mass media and audio-visual aids.
 - 13. To support and advice Government on technical aspects of international conventions and treaties.
 - 14. To conduct other activities incidental and conducive to attainment of abovementioned objectives, which the council may consider necessary.

Also the council through Forest Research Institute (Deemed) University is running Post Graduate programmes in Forestry, Wood Science & Technology, Environment Management, Cellulose & Paper Technology and PhD. courses.

Whereas

The **Hemvati Nandan Bahuguna Garhwal University** (formerly known as **Garhwal University**) is an academic educational university established in Srinagar (Garhwal), Uttarakhand in North India that came into existence in 1973 (vide U.P. State Government notification no. (10)/(865)/15/(75)(85)/64, dated 23 November 1973) after several years of community movement with an aspiration to start an Institution of higher education in the region. It was renamed as Hemvati Nandan Bahuguna Garhwal University in 1989 in commemoration of the memory of a leading statesman Shri Hemvati Nandan Bahuguna. The University was upgraded as a Central University on 15th January 2009 by an Act of Parliament i.e. the Central Universities Act 2009. At present it is only Central University in the state of Uttarakhand and considered amongst ten largest universities in India.

Hemvati Nandan Bahuguna Garhwal University functions under University Grants Commission (UGC). The University is made up of four independent campuses Birla Campus Srinagar (BCC Srinagar- HQs) in Srinagar, Chauras Campus (Tehri Garhwal), B. Gopal Reddy Campus Pauri (BGR Pauri), and Swami Ram Teerth Campus Badshahithaul Tehri (SRT Tehri). All campuses functions as independent institutions within the university. All campuses are equipped with proper infrastructure such as buildings, well-furnished class rooms, lecture halls, laboratories, libraries, seminar/meeting halls, auditorium, grounds, hostels (boys and girls) and other related facilities such as transport, medical, wireless fidelity etc. A total of 117 government, semi-autonomous and private constituent colleges are also affiliated to the HNB Garhwal University. The university conducts exams and other activities simultaneously across all its campuses and constituent colleges.

The **Vision** of HNB Garhwal University is "to achieve excellence by empowering all stakeholders through promotion of innovations in the field of higher education by imparting training and education, and encouraging research for the development of the country with specific attention to the mountain region". The main **Mission** of the University is "to stimulate the academic environment for promotion of holistic learning and research and contribute to the nation's growth. To inculcate values and

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impart skills for shaping able and responsible individuals committed towards the intellectual, academic and cultural development of society".

The synergy derived from its vision & mission, and being a Central University, HNB-GU has a Pan-India appeal, therefore draws students from all over the country. The broad objectives of the HNB Garhwal University are as follows:

Objectives

- 1. To disseminate and advance knowledge by providing instructional and research facilities in the areas of humanities, social sciences, science and technology and other branches of learning.
- 2. To make special provisions for integrated courses in its educational programmes.
- 3. To take appropriate measures for promoting innovations in teaching-learning process and inter-disciplinary studies and research; to educate and train manpower for the development of the country.
- 4. To establish linkages with industries for the promotion of science and technology.
- 5. To pay special attention to the improvement of the social and economic conditions and welfare of the people, their intellectual, academic and cultural development.

The teaching and educational activities are undertaken through fifty-one academic departments distributed in eleven different schools (viz. Agriculture and Allied Sciences; Sciences; Life Sciences; Earth Sciences; Engineering and Technology; Humanities and Social Sciences; Management; Arts, Communication and Languages; Education; Commerce; and Law). Altogether there are over 350+ faculties involved in teaching, education, research, and diverse other co-curricular activities. It houses over 12,000 students annually. The profiles of subjects & courses offered in three different campuses is as follows:

	School	Departments	University Campus & course offered		
			BCC Srinagar	BGR Pauri	SRT Tehri
1.	School of Agriculture	Forestry & Natural Resources	*		
	and Allied Science	High Altitude Plant Physiology Research Center	*		
		Horticulture	*		
		Rural Technology	*		
		Seed Science & Technology	*		
2.	School of Sciences	Chemistry	*	*	*
		Home Science	*	*	*
		Mathematics	*	*	*
		Physics	*	*	*
		Pharmaceutical Science	*		
		Pharmaceutical Chemistry	*		
		Statistics	*	*	*
3.	School of Life Sciences	Botany & Microbiology	*	*	*
		Bio-Chemistry	*		
		Environment Sciences	*		
		Zoology	*	*	*
u		Himalayan aquatic biodiversity	*		
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<i>.</i>		Biotechnology	*		
4.	School of Earth Science	Geology	*		
		Geography	*	*	*
		Remote Sensing and GIS	*	*	*
		Defense & Strategic Studies	*		
5.	School of Management	Centre for Mountain Tourism &	*		*
	0	Hospitality Studies	*		
		Business Management			
6.	School of Humanities	Economics	*		
	and Social Sciences	Political Science	*	*	*
	of the other other of the other other of the other other of the other other other other other of the other ot		*	*	*
		Anthropology	*		*
		Sociology and social work	*	*	*
		Psychology	*		
		History, culture, and archeology	*	*	
7.	School of Engineering	Philosophy	*		
/.	School of Engineering	Computer Science and	*		
	and Technology	Engineering			
		Electronics and Communication	*		
		Engineering			
		Instrumentation Engineering	*		
		Mechanical Engineering	*		
		Information Technology	*		
8.	School of Law	Law		*	-
9.	School of Commerce	Commerce	*	*	*
		Secretarial Practices	*	100	4
10.	School of Education	Adult Education & Extension	*		Ŧ
		Education	*	4	
		Naturopathy & Yoga	*	Ŧ	*
		Physical Education	*		
11.	School of Arts,	Drawing & Painting	*		
	Communication and	Center for Journalism & Mass	*		*
	Languages	communication	Ŧ		
		English	12		
		0	*	*	*
		Centre for Folk and performing art	*		
		Hindi	*	*	
		Library & Information Science	*		
		Music	*		
		Sanskrit	*	*	*

University intake of students either on merit based or through competitive exams conducted by the university (such as Ph.D. entrance) and other agencies (i.e. Joint Entrance Examination Main (JEE Main) for School of Engineering & Technology, CAT, MAT etc.). The courses offered in university campuses comprise undergraduate, post graduate, doctoral and diploma courses. The **Under Graduate Programme** (3 years degree and 4 years honors courses) comprise Bachelor of Science (B.Sc. and B.Sc. Hon.); Bachelor of Technology (B. Tech.), Bachelor of Pharmaceutical Sciences (B. Pharma), Bachelor of Arts (B.A. and B.A. Hon.), Mass Communication (Integrated course); Bachelor of Commerce (B.Com.), Bachelor of Library and Information Science (B.Lib.), and Bachelor of Business Administration (B.B.A.).

The **Post Graduate Programme** (2 years) cover Master of Science (M.Sc.), Master of Arts (M.A.), Master of Business Administration, Tourism, Computer Application, Master of Commerce (M.Com.), M. Pharma, Master of Education (M.Ed.), and Master of Extension Education (MEE). The University also offers Under Graduate Diploma in Advertising and Sales Promotion and Sales Management, and Post Graduate Diploma in Company Administration, Business Administration, Tourism and Hoteliering, Yoga, Rural Technology, Bio techniques of Medicinal and Aromatic Plants, Human Rights, Environmental Economics, Journalism and Mass Communication, Folk Music, Folk Dance. The University also takes **Research Programme** M.Phil. (in English and Environmental Plant Biology) and Doctor of Philosophy (Ph.D.) in all disciplines.

Over the years the University has shown commitment towards regional and community development in its teaching/learning/educational courses, research programs, and other outreach and extension initiatives thus contributing immensely to the development of the society and the nation. H.N.B Garhwal University is 'A' Graded with CGPA of 3.11 by National Assessment and Accreditation Council (NAAC). It is trying hard to achieve excellence in academics and strive for all round development of the students. The focus of the university is to achieve i) excellence in higher education and research relevant to both nation, as a whole, and the hill states, including Uttarakhand, in particular; ii) expansion and inclusion at UG and PG levels of teaching for better outreach in rural areas, especially to the women in the hill districts of Uttarakhand; and iii) promotion & development of extension as third dimension of higher education.

The University is eager to play an important role as leaders in teaching, education, research and technology, training and capacity building, and strengthening institutions of cultural & societal values. For this purpose it is willing to extend its network with institutions of higher learning and research having common profession or academic interest by signing memorandum of understanding (MoU).

And Whereas

Both the parties have expressed their willingness to consider implementing joint programmes to support forestry research, education and extension through their respective institutions, namely Indian Council of Forestry Research and Education and Hemvati Nandan Bahuguna Garhwal University.

Article I: Objective

1.1 Providing a general framework for long term collaboration and mutual understanding in areas of forestry research, education and extension between Indian Council of Forestry Research and Education and Hemvati Nandan Bahuguna Garhwal University.

Article II: Scope of Collaboration

Subject to the terms this MoUs signed by authorized representatives of each party, the parties hereby agree to cooperate as follows:

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2.1 To offer collaboration as felt desirable and feasible on either side and both sides would contribute to the fostering and development of the cooperative relationship between respective organization on identified areas of research, education and

Article III: Areas of Collaboration

3.1 To develop programs as mutually agreed by both the parties in the following areas:

- Knowledge sharing
- Capacity building of stakeholders in the forestry and natural-resourcesii. related sectors
- Workshops, Conferences, Meetings to draw lessons on practices, approaches iii. and interventions
- Facilitation of research and education projects and programs iv. V.
- Joint efforts for synthesizing knowledge in prioritized areas vi.
- Joint seminars/webinars and panels in events, conferences and meetings etc. vii.
- Seeking funding and in-kind resources for these cooperation activities viii.
 - Exchange of scientists, Research Scholars and PG students

Article IV: Scope of MoU

4.1 Indian Council of Forestry Research and Education and Hemvati Nandan Bahuguna Garhwal University through this general Memorandum of Understanding have agreed to discuss opportunities to share resources and jointly undertake consultation, advisory and R&D programmes for their mutual benefit. Programmes and projects will be decided on merits and mutual understanding with respect to operative period of MoU for financial obligations and extent of cooperation. Periodic meetings of joint working groups shall be held as per mutual discussion to review and decide next course of action.

4.2 Both parties agree to provide logistic support to each other. Both parties may submit collaborative programmes for assignments on complementary basis. However, each case will be decided on its own merit, expertise available with the institutes and with mutual consent.

4.3 Structural composition of team members from respective parties will be decided mutually and competent professionals from both the organization will be offered to perform the job with professional prowess. Each programme and case will be decided on merit and mutual understanding and separate agreements will be signed for each assignment after negotiations on case to case basis agreed through mutual consent.

4.4 Both the parties shall maintain their sovereignty and share no liability unless agreed through mutual consent, by way of separate agreement drawn for the purpose.

4.5 In the event that the parties wish to carry out a programme or activity pursuant to this MoU (including those mentioned above), the periodicity, extent of bearing liabilities and terms of such cooperation shall be mutually discussed and agreed upon in writing by both parties through separate agreements prior to the initiation of the particular programme or activity. Any such agreement will detail the terms and nature of the

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activity and/or project, the responsibilities of each party, the necessary budget, and the financial responsibilities, if any, of the parties.

4.6.In the event of any dispute/s that may arise in connection with this MoU, the same shall be mutually settled by the Director General, ICFRE and Vice Chancellor, HNBGU.

4.7. This MoU shall also construe both the institutions to help, compliment and collaborate in areas of research and extension in the areas of common interest at the level of their regional institute/offices located all over India.

Article V: Protection of Intellectual Property

It is recognized by both the parties that the allocation of intellectual property rights will occur on the basis of a research and technology management plan developed for the research programmes executed by both the organizations. 5.2.

Research outcome will be jointly patented and outcome of the patent will be jointly shared in mutually agreed proportions. 5.3.

It is agreed by the parties to undertake not to disclose, divulge, part with, copy in any form for the period as specified in the MoU/agreement any scientific research document, planning, execution and appropriate financial, legal documents involved in the MoU. All personnel involved in collaborative projects from Indian Council of Forestry Research and Education and Hemvati Nandan Bahuguna Garhwal University are subject to this non-disclosure clause for the period of this MoU.

5.4. The outcome of the research in the form of publications should be shared by both the organizations and should have the authorship from both the organization with proportional credit to contributors.

Article VI: Limitation of Personnel Activities

The Parties shall ensure that any personnel engaged in the activities and/or 6.1. programs under this MoU shall not interfere with the internal political independence, sovereignty, and territorial integrity of the latter, any commercial ventures, and avoid any activities outside the aims of this MoU and any separate written arrangement.

Article VII: Confidential Information

Each Party's rights, title, interest in its confidential information remains 7.1. unaffected by the existence of this MoU.

Neither Party is obliged to disclose any confidential information to the other 7.2. Party pursuant to this MoU and each Party undertakes to observe confidentiality and secrecy of any information and other data received from the other Party, and any outputs jointly obtained, except as mutually consented to in writing by the other Party, during the period of the implementation of activities under this MoU or any arrangements made pursuant to this MoU.

7.3. Any disclosure or exchange of confidential information will be covered in separate subsidiary arrangements.

ICFRE with HNBGU

5.1.

Article VIII: Amendment

This MoU can be reviewed or amended at any time by mutual written consent by the Parties. Such revisions or amendments shall enter into force on such date as determined by the Parties and shall form an integral part of this MoU.

Article IX: Dispute Settlement

Any dispute, controversy or difference as to the interpretation of this MoU will be settled amicably by mutual consent between the Parties.

Article X: Validity of MoU

10.1 This MoU shall enter into force on the date of the completion of the signing by the Parties. It shall remain in force for the duration of 05 (Five) years from the date of signing of this MoU and may be extended by mutual written agreement of the Parties.
10.2 Either Party may terminate this MoU at any time by notifying the other Party of the intention to terminate this MoU in writing.

10.3 Unless agreed otherwise by the Parties, the termination of this MoU shall not affect the validity and duration of any ongoing arrangement, activity and contract made under the MoU until the completion of such arrangement, activity and contract.

IN WITNESS WHEREOF, the undersigned have signed this MoU.

For ICFRE

Director General Indian Council of Forestry Research and Education (ICFRE), Dehradun, Uttarakhand

Director General Indian Council of Forestry Research and Education P.O.New Forest, Dehradun-248 006 (Uttarakhaad) Witness:

1. Shri Anurag Bhardwaj Director (IC), ICFRE Director (IC) भा॰वा॰अःशि॰प॰, देहरादून-248006 2. E. Vikram

ADG (External Project), ICFRE सहायक महानिदेशक Assistant Director General बाह्य परियोजना External Project भा.वा.अ.शि.प. For HNBGU

Prof. Annpurna Nautiyal Vice Chancellor Hemvati Nandan Bahuguna Garhwal University (HNBGU), Srinagar

Vice Chancellor H.N.B. Garhwal University (A Central University) Srinagar Garhwal-246174 Witnes Uttarakhand (India)

1. Prof. M.C. Nautiyal HAPPRC, HNB-GU

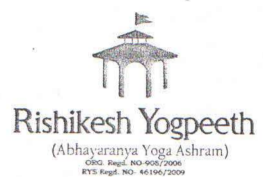
2. Prof. R.K. Maikhuri Head, Dept. of Environment HNB-GU, Srinagar (Garhwal)

Page 9 of 9

MEMORANDUM OF UNDERSTANDING

R·C·C

BETWEEN



Registered Office:

Lane No- 20 Amitgram, Gumaniwala, Rishikesh, Distt. Dehradun (Uttrakhand) India

AND



Herrivati Nandan Bahuguna Garhwal University (Central University recognized by the University Grants Commission)

H.N.B.Garhwal University, Srinagar - 246174, Dist.Garhwal (Uttarakhand), India

MEMORANDUM OF UNDERSTANDING

BETWEEN

RISHIKESH YOGPEETH

AND

HEMVATI NANDAN BAHUGUNA GARHWAL UNIVERSITY

Rishikesh Yogpeeth (hereinafter referred to as "RYP", which expression, shall unless repugnant to the context thereof, shall mean and include its successors and assignees) through the Secretary, RYP, Rishikesh of the First Part.

and

Hernvati Nandan Bahuguna Garhwal University (hereinafter referred to as "HNBGU", which expression, unless repugnant to the context thereof, shall mean and include its successors and assignees) through the Vice Chancellor, HNBGU, Srinagar of the Second Part.

RYP and HNBGU are collectively referred to as "Parties" and individually as "Party".

BACKGROUND

This Memorandum of Understanding (MoU) sets for the terms and understanding between Rishikesh Yogpeeth and Hemvati Nandan Bahuguna Garhwal University (a Central University) to collaborate in the field of Yogic Sciences and related themes and topics at mutually agreed to terms and conditions between the two parties.

PREAMBLE

WHEREAS

Rishikesh Yogpeeth (Street No. 20, Amit Gram, PO Gumaniwala, Rishikesh, Dehradun) was established in 2005 as a non-profit organization registered under the Indian Societies Act 1860 (No. 404/2016-2017, file no. 23537D, dt. 7/4/2017) with an aim to spread the awareness of holistic living with the help of Vedic wisdom in the form of Yoga, Meditation and Ayurvedic practices.

The main objective of Rishikesh Yogpeeth is to spread awareness of holistic living with the help of Vedic wisdom in the form of Yoga, meditation and Ayurvedic practices. It conducts 200, 300, and 500 hrs Hatha Yoga Teacher Training Program, yoga courses for beginners, and yoga retreats in Rishikesh, which is popularly known as the Yoga capital of the world.

The Core team of RYP comprises yogis, teachers, and yoga practitioners.

Rishikesh Yogpeeth comprises all basic facilities and infrastructure with well-equipped lecture halls, meditation ground/galleries, hostels, and all other logistics (including boarding and lodging facilities). It has a focused campus at Abhayaranya Yog Ashram, Neelkanth Temple Road, Patna, Rishikesh, Uttarakhand-249302.

Rishikesh Yogpeeth has been pursuing following programs/ courses:

- Yoga Teachers Training- The course work comprises principles, practices and concepts of Yoga, Asana, Pranayama, Meditation, Yogic cleansing techniques, philosophy of Yoga, modern anatomy
- Yoga retreats- Various yoga retreats focusing on current lifestyles challenges, full body detox, weight loss, etc.

As a yoga school, Rishikesh Yogpeeth is also registered with Yoga Alliance USA, a most recognized Yoga organization worldwide that sets up minimum criteria for Yoga school or a teacher to have international credentials. Rishikesh Yogpeeth has so far trained over 10,000 personnel in Yoga across 120 countries.

WHEREAS

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The Hemvati Nandan Bahuguna Garhwal University (formerly known as Garhwal University) is an academic educational university established in Srinagar (Garhwal), Uttarakhand in North India that came into existence in 1973. The University was upgraded as a Central University in 2009. The University is made up of four independent campuses Birla Campus Srinagar (BCC Srinagar- HQs) in Srinagar, Chauras Campus (Tehri Garhwal), B. Gopal Reddy Campus Pauri (BGR Pauri), and Swami Ram Teerth Campus Badshahithaul Tehri (SRT Tehri). All campuses are equipped with proper infrastructure such as buildings, well-furnished classrooms, lecture halls, laboratories, libraries, seminar/meeting halls, auditorium, grounds, hostels (boys and girls) and other related facilities such as transport, medical, wireless fidelity, etc.

The teaching and educational activities are undertaken through fifty-one academic departments distributed in eleven different schools (viz. Agriculture and Allied Sciences; Sciences; Life Sciences; Earth Sciences; Engineering and Technology; Humanities and Social Sciences; Management; Arts, Communication and Languages; Education; Commerce; and Law).

Under the School of Education, the University has a Department of Naturopathy & Yoga that undertakes postgraduate (2-year course), undergraduate (3-year course), PG Diploma (1-year course), certificate (6-month), and short-term (30 and 45 days) courses.

With the enactment of New Education Policy 2020 in the country that aims, among others, to increase quality of education and capacity building of students for employability, the University is eager to play an important role in extending quality education, research, and training programs.

The 10th Academic Council has advocated for the opening of University's Extension Centre at Rishikesh under self-finance mode for imparting a 'Six-month Certificate course in Yogic Science' and 'Six-month Certificate course in Mental Health and Spirituality'.

University plans to take up such courses in PPP (public-private-partnership) mode so as to avoid logistic and infrastructure costs. For this purpose it is willing to extend its network with institutions and agencies having similar vocational, theoretical & practical interests by signing memorandum of understanding (MoU).

AND WHEREAS

AFFIRMING that in recent time yoga teaching and practices have become symbolic to good health and mental well-being, wider dissemination of such education and information would be immensely beneficial to the society;

EMPHASIZING that there is an exigent need to strengthen the institutional delivery mechanism to train and build capacity among people on understanding of principles and practices of Yoga; and

PERCEIVING the need for close-linking of Institutions of similar learning, education, and capacity building to expand the knowledge base and outreach to develop human potential in PPP (public-private-partnership) mode;

both the parties have expressed their willingness to consider implementing joint programmes to support Yoga science, training and capacity building through their respective institutions, namely Rishikesh Yogpeeth and Hemvati Nandan Bahuguna Garhwal University

have reached the following understanding in compliance to the above-said prerequisites:

Article I: OBJECTIVES

- 1. To conduct yoga courses (short-term, certificate, degree, and PG level) at Rishikesh Yogpeeth so as to enhance the opportunity of more effective teaching in Yoga Science/subject.
- 2. To collaborate between themselves for strengthening knowledge and learning in Yoga science through joint R&D projects, national/international seminars, workshops and training programs
- 3. To conduct surveys and awareness on the effect of yoga on health, wellness, and performance of individuals

Article II: SCOPE OF COOPERATION

Subject to the terms this MoU signed by authorized representatives of each party, the parties hereby agree to cooperate as follows:

- 1. Both the parties would contribute to the fostering and development of the cooperative relationship between respective organization through planning, training, promotion and coordination of Yoga education, therapy, and research.
- 2. The Parties will collaborate to take up approved Yoga courses (short-term, certificate, UG, PG) at RYP campus in Rishikesh as per the broad parameters outlined in the MoU. Details will be worked out through mutual consultations between the parties.
- 3. After the signing of MoU, the RYP will undertake course work and activities in collaboration with HNBGU in accordance with this MoU at their centre in Rishikesh.
- 4. Both parties will take up work related to teaching, learning, capacity building, and certificate course as per approved norms and regulations of HNBGU

- 5. Both parties will organize joint workshops, conferences, seminars, events, training on Yoga and related subjects on mutually agreed terms and conditions
- 6. Both parties will facilitation joint research & education, knowledge synthesis, and formulation of policy documents
- 7. Both parties may exchange faculties, teachers, Research Scholars, and students to accomplish the above said

Article III: OBLIGATIONS OF THE PARTIES

- Both the parties agree to conduct short-term (defined as 30 and 45 days) and certificate and diploma courses (defined as 6 months to 1 year) at RYP centre* in Rishikesh in self-finance mode. The syllabus will be implemented with mutual discussion between RYP and HNBGU. The number of students per course and number of courses per year will be decided after mutual consultation.
- * RYP centre operating from two locations stated as Abhayaranya Yoga Ashram (Patna Village, Rishikesh) and Sri Vithal Ashram (Rishikesh)
- 2. All necessary logistics and infrastructure for conducting Yoga courses shall be provided by RYP with no cost to HNBGU.
- 3. The fee (tuition/examination fee) for the course will be as per the existing norms of the HNBGU and will be distributed 60% (RYP) and 40% (HNBGU).
- 4. Both the parties also agree to exchange faculties/experts for conducting courses/programs between RYP and HNBGU through mutually agreed modalities.
- 5. HNBGU may depute teachers/faculty/officials to monitor/conduct courses and training programs. RYP will allow it to do so and provide local hospitality, accommodation, and other facilities to the officials deputed by HNBGU to monitor the Yoga Course.
- 6. For any additional activity both the parties will finalize modalities through mutual consultations and discussion.

Article IV: AMENDMENT TO MOU

Either party, if a desire to modify or amend any or all part of this MoU, may do so on a written request for the revision. The mutually agreed revision, modification, or amendment shall be reflected in writing and form a part of this MoU, which shall enter into force on such date as may be determined by the parties.

Article V: SETTLEMENT OF DISPUTES

This MoU is not intended to create any legal obligations between the parties. In case of any difference or dispute between the parties concerning the interpretation and/or implementation and/or application of any of the provisions of this MoU shall be settled amicably through mutual consultations and negotiations between the parties.

Article VI: ENTRY INTO FORCE AND DURATION

- 1. This MoU shall come into effect on the date of its signature and shall be valid for a period of five years. Thereafter, both Parties shall review the status of this cooperation and may extend it on such terms as mutually agreed.
- 2. Notwithstanding Article VI(1), either Party may terminate this MoU by notifying the other Party in writing at least three months in advance notice of termination. Although such termination must meet with the last day of the course/s in progress.
- 3. Unless otherwise agreed by the Parties, the termination of this MoU shall not affect the implementation of on-going activities and/or programmes, which have been agreed upon before the date of termination of the MoU.

Article VII: OTHER PROVISIONS

- 1. The Parties agree to comply with existing rules and regulations of HNBGU applicable to the respective clause in the implementation of this MoU.
- 2. This MoU is intended to create a partnership, joint venture or employability for the students/beneficiaries.

3. Neither party shall have any right or authority to bind, speak for or contract on behalf of another party.

IN WITNESS WHERE OF

Signed on

day Musth 2021 at Srinagar in two originals in English.

For RYP

Secretary

12 de la

Rishikesh Yogpeeth, Street No. 20, Amit Gram, PO Gumaniwala, Rishikesh, Dehradun, Uttarakhand For HNBGU

Vice Chancellor / Pagistoan

Hemvati Nandan Bahuguna Garhwal University (HNBGU), Srinagar

1. RAJESH KANDARI ZTGIZ, FIE.

2. SANJEEV NAME

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2. Int S.S. Rowal Sorvi of concention



हेमवती नन्दन बहुगुणा गढ़वाल विश्वविद्यालय Hemwati Nandan Bahuguna Garhwal University श्रीनगर गढ़वाल (उत्तराखण्ड) 246174 Srinagar Garhwal (Uttarakhand) 246174 (केन्द्रीय विश्वविद्यालय) (A Central University)

पत्रांकः हे०न०ब०ग०वि०वि०/2021/जन्तु विज्ञान 2 39

दिनांक: 27/08/2021

सेवा में,

सहायक कुलसचिव आर0सी0सी0 प्रकोष्ठ हे0न0ब0ग0वि0वि0

विषयः – विभाग के एम.ओ.यू एवं प्रस्तावित एम.ओ.यू की छाया प्रतियों के सम्बन्ध।

महोदय,

आपके पत्रांक HNBGU/RCC/2021/774 dated 26.08.2021 के सन्दर्भ में हे0न0ब0ग0वि0वि0 (केन्द्रीय विश्वविद्यालय) के जन्तु विज्ञान विभाग के साथ कोपेनहेगन विश्वविद्यालय (डेनमार्क) और राष्ट्रीय मत्स्य अनुवांशिक संसाधन ब्यूरो (लखनऊ) के मध्य अनुबन्ध (एम.ओ.यू) की छाया प्रतिलिपि एवं डी0सी0एफ0आर0 (आईसीएआर) भीमताल का एम.ओ.यू आगामी विद्या परिषद में प्रस्तावित हेतु संलग्न कर प्रेषित किये जा रहे हैं।

भवदीय

विर्मागाध्यक्ष प्रो0 प्रकाश नौटियाल जन्तु विज्ञान विभाग

संलग्नः एम.ओ.यू की छाया प्रतियां

MEMORANDUM OF UNDERSTANDING ON ACADEMIC COOPERATION, STAFF AND RESEARCH EXCHANGE ETC.

Sh mou ()

BETWEEN

HEMVATI NANDAN BAHUGUNA GARHWAL UNIVERSITY, INDIA (A CENTRAL UNIVERSITY) AND

UNIVERSITY OF COPENHAGEN, DENMARK

In furtherance of their mutual interests in the fields of education and research and as a contribution to increased international cooperation, the Memorandum of Understanding is executed on 11.10.2019 between HEMVATI NANDAN BAHUGUNA GARHWAL UNIVERSITY (A CENTRAL UNIVERSITY), India, acting through its School of Life Sciences, Department of Zoology & Biotechnology AND the UNIVERSITY OF COPENHAGEN, DENMARK, acting through its Faculty of Science, Department of Biology. This MoU provides a framework for academic cooperation in general between the signatories, for different types of cooperation within the areas of Freshwater ecosystems function and services. Both institutions approve the following Memorandum of Understanding which shall be for a period of 5 years with effect from October 11th, 2019.

- 1. Both institutions will cooperate in the exchange of information relating to their activities in research in fields of mutual interest mentioned above.
- 2. Both institutions will promote appropriate joint research projects and joint courses of study.
 - The two institutions have already agreed to collaborate within the activities outlined in the appendix attached to this Memorandum of Understanding. The two institutions undertake to participate in good faith in the completion of the described activities; however, neither party can guarantee any specific results as an outcome of the planned activities.
 - When relevant the two institutions will collaborate with other institutions or individuals as partners in the activities within the fields of mutual interest mentioned above. When necessary separate MOUs or collaboration agreements with third parties will be entered.
- 3. In the event of a joint research project both the institutions agree to enter into a cooperation agreement covering the terms and conditions for the specific research project.
- 4. Both institutions will endeavour to develop and implement a programme of research personnel exchange in accordance with the principles described below. Each research personnel exchanges shall be subject to a specific agreement. Joint publications will be made to disseminate the outcomes of research programme.
- 5. Faculty members and other research personnel, including Ph. D. students, may be exchanged for mutually agreed periods. The host institution will waive the relevant fees for the period of research stay, while the home institution continues to undertake the appropriate salaries of the participant.



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Memorandum of Understanding between

National Bureau of Fish Genetic Resources And Hemvati Nandan Bahuguna Garhwal University, (A Central University) Srinagar Garhwal

This Memorandum of Understanding (hereinafter MoU) is made on this 5th day of the month, of December in the year 2019 by and between the National Bureau of Fish Genetic Resources having its head office at Lucknow (hereinafter called "NBFGR"), a constituent laboratory of the Indian Council of Agricultural Research, Krishi Bhavan, New Delhi 110 001 on the ONE PART and the Hemvati Nandan Bahuguna Garhwal University (A Central University) having its headquarters at Srinagar (Pauri Garhwal) (hereinafter called HNBGU) on the OTHER PART (who for the purpose of this MoU are hereinafter collectively referred to as the Parties.

The Parties, having discussed fields of common research interests and allied activities between the two institutions, have decided to enter into long-term collaboration in the areas of research, teaching and training.

WHEREAS "NBFGR" is involved in the studies on fish biology, genetics, biotechnology, Conservation and disease diagnosis of freshwater and marine fish species.

AND WHEREAS HNBGU at its Department of Zoology and Biotechnology is involved in many disciplines of Zoology including in Fish Biology, Fisheries Science and Ecology and Fish biotechnology of freshwater fishes.

AND WHEREAS the Parties have agreed to collaborate in undertaking projects related to hitherto unattained aspects of fisheries science and related biotechnology.

AND WHEREAS is has been considered expedient to agree in writing to participate jointly in the projects requiring expertise and logistics from both the Parties.

Seen. MOU National Bureau of Fish Genetic Resources, Lucknow and Department of Zoology and Biotechnology, HNB Garhwal University (A Central University) Srinagar Garhwal मत्त्य आनुवारीक संसाथ A Burecu of Fish Generic Res. (1)न रिग रोड आ. दिलकुशा

-Article 1. Scope

- 1.1 Research instrumentation facility and library facilities available in NBFGR and HNBGU will be made available to the faculty and research scholars. However, the costs of specific consumables will be borne by the respective organizations.
- 1.2 Detailing of research effort and collaboration in common research programmes and /or projects.
- 1.3 There shall be an exchange of faculty members and scientists for academic and research tenure purposes. Accommodation in the guest house shall be arranged whenever possible. Joint visitor programmes for distinguished scholars shall also be arranged.
- 1.4 Instrumentation facility developed in NBFGR may be made available to HNBGU at mutually acceptable rates.
- 1.5 Conduct of seminars and workshops.
- 1.6 Organization and teaching inputs in training programmes.

Article 2. Execution and Management

- 2.1 To manage the execution of this MoU, the Parties agree to constitute and Advisory Committee as follows:
- 2.1.1 The Vice-Chancellor, HNB Garhwal University (A Central University) Srinagar Garhwal, as Chairman.
- 2.1.2 The Director, National Bureau of Fish Genetic Resources as Co-Chairman.
- 2.1.3 The Head of concerned Department, Garhwal University, as Convener.
- 2.1.4 NBFGR Scientist nominated by the Director, NBFGR, as Member,
- 2.1.5 One senior faculty member of the concerned Department of the HNB Garhwal University in cognate area of research to be nominated by the Vice-Chancellor.
- 2.1.6 One Senior Scientist of the concerned department in cognate area of research to be nominated by the Director, NBFGR.

Seen. MOU National Bureau of Fish Genetic Resources, Lucknow and Department of Zoology and Biotechnology, HNB Garhwal University (A Central University) Srinagar Garhwal



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Article 3. Exchange of Information

- 3.1 The term "information" includes scientific or technical data, results and/or methods of investigation, and other information intended to be provided, exchanged or arising under project descriptions entered into pursuant to this MoU.
- 3.2 The Parties support the widest possible dissemination of information. Each party in joint projects shall be given the right to use, disclose, publish or disseminate such information for any and all purposes.

Article 4. General Provisions

- 4.1 Enrolment and Award of D.Phil. degree for candidates working at NBFGR:
 - (i) Ordinances and regulations of the University for the admission and award of Ph. D.degree of the HNB Garhwal University will be duly observed. The criterion for allocation of Supervisor and Co-Supervisorwill primarily be governed these ordinances.
- 4.2 In the case of training or teaching programmers and research projects, the host institution shall provide all necessary infrastructure and facilities, including audio-visual equipments and guest accommodation etc., as per the rules.
- 4.3 The institution organizing an item of work shall have the nodal responsibility for mobilizing resources and the conduct of such works. Funds earmarked for the collaborative activity will be used for the purpose for which they are meant, and will be disbursed in accordance with the financial rules and procedures of the respective institutions. In the HNB Garhwal University, the funds for collaborative work will be managed by Head of the Department and the Registrar, through a separate bank account to be operated jointly by them. The budget for the activity shall be approved by the Advisory Committee and the final statement of accounts shall also be submitted to the Advisory Committee.

Seen. MOU National Bureau of Fish Genetic Resources, Lucknow and Department of Zoology and Biotechnology, HNB Garhwal University (A Central University) Srinagar Garhwal



(3)

- 4.4 Any research publications and commercially exploitable innovations arising out of this joint effort will be jointly published or patented as appropriate sharing.
- 4.5 All questions related to the MoU arising during its term will be settled by the Parties by mutual agreement. Disagreements at the operating level shall be forwarded to respective higher officials for appropriate resolution failing which an arbitrator of mutual acceptance may be identified.
- 4.6 All questions not foreseen related to this MoU will be handled by the Parties by mutual agreement.
- 4.7 Nothing in this MoU is intended to affect other cooperation or collaborations between the Parties.

Article 5. Entry into Effect and Termination

- 5.1 This MoU shall become effective on the date it is signed by the Parties and shall be valid for five years. Its term shall be extended further by mutual agreement after the expiry of the initial five-year's term. This MoU may be amended by mutual written agreement and may be terminated at any time by either Party upon six months written notice to the other Party.
- 5.2 All joint activities not completed at the expiration or termination of the MoU may be continued until their completion under the terms this MoU.

Article 6. Amendments:

- 6.1 This MoU represents the entire understanding between the Parties and supersedes any and all understandings either oral or written hitherto with respect to the subject matter of the Agreement.
- 6.2 No amendment or modification of the MoU shall be valid unless the same is made in writing by both the <u>Parties</u> or their authorized representatives and specifically stating the same to be amendment of the MoU. The modifications/changes shall become part of the MoU and shall be effective from the date on which they are made/executed, unless otherwise agreed to.

This MoU has been executed in two originals, one of which has been retained by NBFGR and the other by Department of Zoology and Biotechnology of Garhwal

University.

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MOU National Bureau of Fish Genetic Resources, Lucknow and Department of Zoology and Biotechnology, HNB Garhwal University (A Central University) Srinagar Garhwal

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WITNESS THEREOF, the parties have executed this MoU and represent that they approve, accept and agree to terms and contained herein.

National Bureau of Fish Genetic Resources Canal Ring Road, PO Dilkusha, Lucknow/ Indian Council of Agricultural Research

Dr. Kuldeep K. Lal Director, ICAR, NBFGR, Lucknow

Date

5.12 J.9. skiloop K. Lai Director

ICAR-National Eureau of Fish Ganetic Resources

Conel Ring Road, Telibagh, P. O. Dilkusha Lucknow-226 CO2, U. P., India

Seal

Dr. Gaurav Rathore Principal Scientist & Head FHM & Exotics Division Nominated by the Director

Date 5-12-19.

Seal

मल्य आनुवांशिक संसाध Butecu of Fish Genetic Reson कैनाल रिंग रोड पो. आ. दिलकुशा METTS / LUCKNOW-226

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MOU National Bureau of Fish Genetic Resources, Lucknow and

Department of Zoology and Biotechnology, HNB Garhwal University (A Central University) Srinagar Garhwal



HNB Garhwal University

Prof. Annpurna Nautiyal Vice Chancellor

Date 5.12.2019

Vice Chancellor H, N.B. Gardievol University Seal (A Control University Srinaght Cariwal-246174

Prof. Prakash Nautiyal I/C Head Dept. of Zoology & Biotechnology HNB Garhwal University Srinagar Garhwal

Date 5th December 2019.

Professor & Flead Seal Deptt. of Zoology & Blotechne > **ENB** Garhwal University Srinagai-Garbwal (Uttarakhand)



भा.कृ.अनु.प.–राष्ट्रीय कृषिउपयोगी सूक्ष्मजीवब्यूरो (भारतीय कृषिअनुसंघान परिषद्) ICAR-NATIONAL BUREAU OF AGRICULTURALLY IMPORTANT MICROORGANISMS

CAR-NATIONAL BUREAU OF AGRICULTURALLY IMPORTANT MICROORGANISMS (Indian Council of Agricultural Research) कुशुमौर, मऊउत्तर प्रदेश-275103 Kushmaur, Mau Uttar Pradesh- 275103 Tel (दुरमाष): (0547) 2530080, FAX (फेक्स): (0547) 2530381, E-Mail (ई-मेल): <u>director nbaim@icar.gov.in</u>,(Web): <u>www.nbaim.org.in</u>



Date 14.01.2020

File No. 10-1/2013/Dir/NBAIM/2043

To,

Prof. Annpurna Nautiyal, Vice-Chancellor, H.N.B. Garhwal University (A Central University) Srinagar Garhwal-246174 Uttarakhand (India).

Sub: Submission of signed copy of MoU reg.

Dear Madam,

Please find enclosed herewith the signed copy of MoU for your Kind information and record.

With regards

Yours Faithfully

14 1. 2020 In-Charge PME

In-Charge PME ICAR-NBAIM

CC: Dr. Rahul Kunwar Singh, Assistant Professor of Microbiology, Department of Botany & Microbiology, H.N.B. Garhwal University.

22/01/2020

Memorandum of Understanding

between

ICAR-National Bureau of Agriculturally Important Microorganisms (NBAIM) and

Hemvati Nandan Bahuguna Garhwal University (A Central University) for facilitating Training and Research

This Memorandum of Understanding (hereinafter referred to as MoU) is made on this

The parties, having discussed fields of common research interests and allied activities between the two institutions, have decided to enter into long-term collaboration for promotion of training and quality research in cutting edge areas in accordance with the provisions contained in the Guidelines issued *vide* Letter No. 2-8/2012-HRD dated 11th December, 2012 or as revised from time to time.

WHEREAS the "First Party" is involved in the studies on Microbiology, Microbial Biotechnology, Extremophile Biology, Bioinformatics (specific mandated domain within the approved disciplines/divisions), AND WHEREAS the "Second Party", established by Govt. of India *vide* Act No. 25 of 2009 (Central University Act 2009) dated March 20, 2009 and recognized by University Grants Commission at its Department of Botany and Microbiology is involved in the studies on Microbiology, Microbiology, Extremophile Biology and Cyanobacterial biotechnology

AND THEREAS it has been considered expedient to agree in writing to participate jointly in the participate requiring expertise and logistics from both the parties.

Article 1. Scope

- 1.1. The Second party will recognize Scientists of the NBAIM as recommended by its Director in accordance with the provisions laid down in respective ordinances of the Second party for co-guiding students working for the M.Sc. and Ph.D. degree.
- 1.2. Operational details of research effort and collaboration will be made in common research programmes and/or joint projects restricted to specific mandated domain within the approved disciplines/divisions. The objective(s) for research work for a student coming from a Second party outside National Agriculture Research System (NARS) should be exclusively different as far as possible.
- 1.3. Research instrumentation facility and library facilities available with the First party and the Second party will be made available to the faculty and research scholars of both the parties. However, the costs of specific consumables will be borne by the respective organizations.
- 1.4. There shall be an exchange of students for academic, research and training purposes. Accommodation in the Hostel shall be arranged, wherever possible, as per extant rates. The duration of exchange visits will be determined by mutual consent between both the parties.
- 1.5. Both the parties will work closely to develop and execute joint research proposals.
- 1.6. Both the parties will work closely to generate appropriately trained manpower in the identified common areas through organization of symposia, workshop, short term training etc.

Article 2. Management

- 2.1. On the recommendation of respective Head of the Organization, an Advisory committee will be constituted for the effective execution of this MoU.
- 2.2. The Advisory Committee will meet at least once in a year alternatively in the institutions of the First party and the Second party to review the activities. This meeting shall include presentation on the academic and research activities, which should be open to the students, faculty and scientists.

Article 3. Exchange of Information

3.1. The term "information" includes scientific or technical data, results and/or methods of investigation, and other information intended to be provided, exchanged, or arising under the joint project descriptions entered into pursuant to this MoU.

3.2. The parties support the widest possible dissemination of information. Each party in joint joint projects shall be given the right to use, disclose, publish or disseminate such information for any and all purposes.

Article 4. General Provisions

- **4.1.** It is understood that the First party and the Second party subscribe to the principle of equal opportunity and do not discriminate on the basis of race, sex, age, caste or religion. Both the Institutions shall abide by these principles in the administration of this agreement and neither party shall impose criteria for exchange of scholars or students, which violate principles of non-discrimination.
- **4.2.** Both parties understand that all financial agreements will have to be negotiated separately and will depend on the availability of funds.
- **4.3.** Both parties acknowledge that exchange of students from one party to the other shall be subject to the availability of funds and shall comply with the regulations and policies of the First party and the Second party.
- 4.4. Any research publications arising will be jointly published by mutual consent of student and advisors at both the parties and/or in accordance with the provisions laid out in Item 3.2.1C of the Guidelines for the students to conduct research for their degree programme as trainees at ICAR institutions as notified *vide* Letter No. 2-8/2012-HRD dated 11th December, 2012 or revised guidelines, if any, as may be issued from time to time.
- **4.5.** A copy of the thesis/dissertation will be submitted to the First Party after the award of the degree by the Second party.
- **4.6.** All questions related to this MoU arising during its term will be settled by the parties by mutual agreement. Disagreements at the operating level shall be forwarded to respective higher officials for appropriate resolution failing which an arbitrator of mutual acceptance may be identified for the settlement of dispute, if any.
- 4.7. All questions not foreseen related to this MoU will be handled by the parties by mutual agreement.
- **4.8.** Nothing in this MoU is intended to affect other cooperation or collaborations between the parties.

Article 5. Intellectual Property Rights .

5.1. Both the parties will be expected to protect the Intellectual Property Rights of each other, generated or likely to be generated during the student's research work. The party, where the major part of the research work was carried out, will be the first applicant and the Second party shall be the joint applicant for IPRs and the students and involved scientific staff shall be included as the inventor/breeder/author. The sharing of benefits from the generated IPR shall be mutually decided in each case.

Article 6. Admission and Fees

- 6.1. All those who wish to register for Master/Doctoral programme under this MoU must apply for admission at the Second party. The allocation of Research Advisors would be finalized before the registration and will be governed by the provisions laid out in respective ordinances of Second party or revised Guidelines, if any, as may be issued from time to time.
- 6.2. Admission of the students and the award of degrees for different programmes will be the responsibility of the Second party as per the rules and regulations.
- 6.3. Allotment of the students at the First party will be done by the approval of Director of the First party and Vice-Chancellor/Head of the Institution of the Second party.
- 6.4. The Second party would have the right to screen the student's eligibility for admission based on their academic period.
- 6.5. The concerned Board of studies of the Second party in consultation with the PME Cell of the First party shall decide the location and sharing quantum of research work. However, minimum residential period may be relaxed up to maximum of 50% for only those doctoral students having their Co-advisors at First party.
- 6.6. The number of student(s) at any particular time will be subjected to the availability of research facilities and scientists' time to guide thesis research at the First party institution.
- 6.7. Any student(s) admitted to the First/Second party for training/ research, if found violating the rules and regulations laid down by the respective party or indulge in such activities that amount to tarnishing the image of the Institute, or cause damage to the property, the registration of such student(s) would be summarily terminated. Any party will not complete the formalities of issuing the certificates to such 'students until they compensate the losses to the other party.

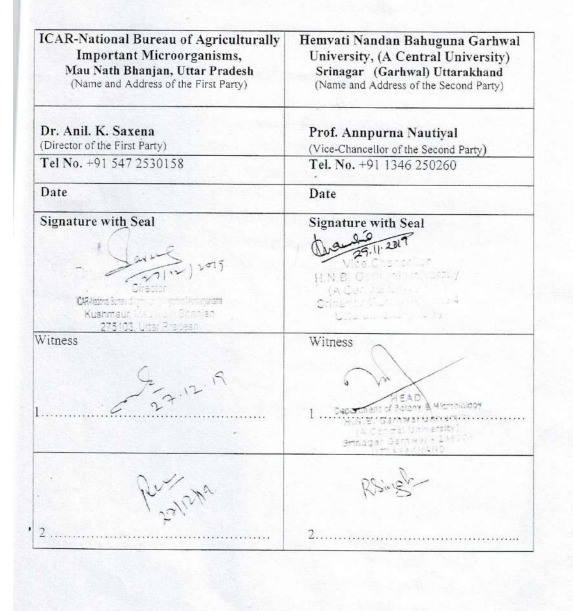
6.8. Fees will be charged from the students by the First and Second party as per the guidelines effective at the respective parties.

Article 7. Entry into effect, modification and termination

- 7.1. This MoU shall become effective on the date it is signed by the parties and shall be valid for three years. Both parties shall review the status of the MoU at the end of each three year period to determine any modification, whenever necessary. The period of validity of this MoU may be extended by mutual consent up to six years. This MoU may be amended by mutual written agreement and may be terminated at any time by either party upon written notification signed by the competent authority of the party initiating termination. Such notification must be given to the other party at least six months in advance from the effective date of termination.
- 7.2. All joint activities not completed at the expiration or termination of the MoU may be continued until their completion under the terms of this MoU.
- 7.3. No amendment or modification of the MoU shall be valid unless the same is made in writing by both the parties or their authorized representatives and specifically stating the same to be amendment of the MoU. The modifications/changes shall become part of the MoU and shall be effective from the date on which they are made/ executed, unless otherwise agreed to.

This MoU has been executed in two originals, one of which has been retained by the First party and the other by the Second party).

IN WITNESS WHEREOF, the parties have executed this MoU and represent that they approve, accept and agree to terms contained herein.



SUDASIEN-INSTITUT



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Südasien-Institut | Im Neuenheimer Feld 330 | D-69120 Heidelberg

Prof. Dinesh Saklani Department of History HNB Garhwal University Shrinagar, Uttarakhand Südasien-Institut (SAI) Prof. Dr. William Sax Geschäftsführender Direktor und Leiter der Abteilung Ethnologie T: +49 (0) 6221-54 89 31 F: +49 (0) 6221-54 49 98 M: william.sax@urz.uni-heidelberg.de

15 November, 2018 Heidelberg

Dear Professor Saklani,

this is to confirm that we will be working together on the German Research Foundation - supported project "Tantrik Text Practices: a comparative ethnography of "tantra" in South and Southeast Asia," which is led by me and Prof. Annette Hornbacher, also of Heidelberg University.

Our work will begin this coming February, when I plan to spend a few weeks with Pt. Devi Ram Paboch in Dehra Dun. The duration of the project is three years. We plan to invite you to Heidelberg during the final months of the project, to work on joint publications.

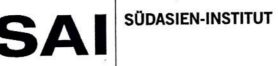
With best wishes,

1

William S. Sax Professor and Head Department of Anthropology, and Executive Director South Asia Institute Heidelberg University

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William S. Sax Professor and Head Department of Anthropology, and Executive Director South Asia Institute Heidelberg University

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MEMORANDUM OF UNDERSTANDING (MoU)

Between the



Hemvati Nandan Bahuguna Garhwal University (A Central University) Srinagar Garhwal Uttarakhand - 246 174

and



Indian Institute of Technology Kanpur Kalyanpur, Kanpur -208 016, India

1-5

Hemvati Nandan Bahuguna Garhwal University (A Central University) Srinagar Garhwal Uttarakhand (hereinafter referred to as "HNBGU") and Indian Institute of Technology, Kanpur, India (hereinafter referred to as "IIT Kanpur") have agreed on the following:

HNBGU is, nestled in the lap of Himalayan ranges in the Garhwal region of Uttarakhand is a residential cum affiliating institution of higher learning. The university has state-of-the-art research facilities, some of the best teachers and close academic relations with premier institutions. Its headquarters are at Srinagar some 108 km from Rishikesh along the Delhi-Niti National Highway; Birla Campus at Srinagar (bisected into the Srinagar Campus and the Chauras Campus by the river Alaknanda); BGR Campus set on enticing mountain slopes at Pauri (30 km from Srinagar and some 100 km from Kotdwar, the nearest railway station); and SRT Campus at Badshahi Thaul (Tehri Garhwal) amidst dense pine forest (85 km from Srinagar and 65 km from Rishikesh on the route to Gangotri).

One of the HNBGU Campus which is proposed for the observations, SRT Campus, is situated at the height of 5600 feet from the sea level and located at 30.08°N 78.61°E., the University Campus is surrounded by the lush green trees of cedar, pines, oaks and rhododendrons. This institution is the premier institute of higher learning in the catchments area of Tehri Dam, the largest rock fill dam of Asia. It is located at a distance of 65 kilometres from Rishikesh on Rishikesh-Gangotri National Highway near the hilly town Chamba from where the Campus can be approached at a distance of 3 kilometres drive.

The Indian Institute of Technology-Kanpur (IIT-K), is a leading academic institute having expertise in the area of scientific and technological education and research and the application of scientific knowledge and technology to human advancement and social development, established under a special act of Parliament of Republic of India, incorporated under the Institute of Technology Act, 1961, India.



AGREEMENT FOR SCIENTIFIC AND CULTURAL COLLABORATION

(Objective)

The HNBGU and IIT Kanpur will endeavour to enhance research activities by supporting the other Party's research efforts under the principle of respecting mutual autonomy. In particular, both Parties will contribute to the monitoring of Cloud Condensation Nuclei (CCN) and data streaming from cloud droplet spectra and Understanding the physico-chemical properties responsible for CCN activation and their relation to cloud microphysical properties over high altitude station SRT Campus HNB Garhwal University Tehri Garhwal Uttarakhand.

(Administrative Aspects and Infrastructure).

The HNBGU will provide all possible infrastructural facilities (in kind) at the existing at SRT Campus, Badshahithaul to keep instruments as well as necessary furniture to keep scientific and power supply instruments and manage through the existing staff at campus, which may be decided mutually.

Department of Physics, Birla Campus and SRT Campus HNBGU will act as collaborating Department to look after this project locally and Prof. R C Ramola / Alok sagar Gautam, from the HNBGU will act as local coordinator responsible for scientific as well as administrative work from HNBGU. From IIT Kapur side, Prof S N Tripathi will coordinate the mutual programme.

(Areas for and contents of collaboration)

The specific collaboration of HNBGU with IIT Kanpur involves activities:

- Long-Term observations of the aerosol composition and source apportionment for 3 Years. This involves the deployment of Single Particle Soot Spectrometer, Time of flight High Resolution Aerosol Mass Spectrometer, Cloud Condensation Nuclei Counter, Cloud Combination Probe and Hygroscopic Tandem Differential Mobility Analyzer at SRT Campus Badshahi Thaul, Tehri Garhwal by Indian Institute of Technology, Kanpur.

 Capacity building of HNBGU in cloud and aerosol research with help of IIT-K



3-5

(Exchange of research staff)

HNBGU confirms that only qualified personnel will operate the instrument and that it will be handled with care. In addition, for quality control and external checks, the computer of the instrument has to be attached to the internet at the station.

The exchange of data and knowhow will help to start a more widespread research activity using such powerful instruments in Himalayan region and the experience gained will be invaluable to both parties. Data sharing with any other institution outside of this agreement should be done on mutual consultation and agreement.

A visit of PhD students/Postdocs from IIT Kanpur at the HNBGU for in-depth exchange is foreseen. HNBGU will support the local hospitality for such visits.

In the event of any dispute or difference of opinion between the Parties arising out of or in connection with this Memorandum of Understanding, each of the Parties shall use its best efforts to settle such dispute or difference of opinion amicably by negotiation between the Parties.

(Term of the agreement)

This Memorandum is executed in duplicate in English, one original for each Party. The Memorandum will come into effect from the date of signing and will be valid for the period of three (3) years and may be extendable on mutual agreement. The agreement will come into act with the signatures of the appointed representatives of the two Parties & will have duration of five years. HNBGU/IIT Kanpur reserves the right to terminate the agreement. Present agreement may be renewed for additional terms, if mutually desired.

Singed this day:

Registrar, HNB Garhwal University Srinagar Uttarakhand India

H.N.B. Garhwal University (A Central University) Srinagar (Garhwal) 246174 UK

Professor S. Ganesh Dean, Research & Development, Indian Institute of Technology Kanpur, India 222207

> अधिष्ठाता DEAN अनुराधान एवं विकास Research & Development आई० टो० कानपुर I.A. T. KANPUR

4-5

Project Coordinators/collaborators Name (HNBGU) Signature Name (IIT Signature Kanpur) 1. Prof R C Ramola, Prof S N Tripathi Department of Civil Department of Physics, SRT Campus Engineering, IIT Kanpur Tehri Garhwal HNB Garhwal University S. N. TR:PATHI Professor Department of Civil Engineering IIT Kanpur, 208 016 UP INDIA Srinagar Uttarakhand) 2. Dr Alok Sagar Gautam, (Assistant Professor, Department Stimio Carpinov Birla of Physics, HNB Campus Garhwal University Srinagar) H.N.B. Garhwal University (A Central University) (A Central University) Inagar (Garbwal) 246174 Uk 5-5