BRIEF BIO-DATA OF DR. VISHAL ACHARYA

Senior Scientist, Functional Genomics & Complex System Lab, **Present Post:**

Biotechnology Division, CSIR-Institute of Himalayan Bioresource

Technology (IHBT), Palampur-176061 (H.P.), India

Email-id vishal@ihbt.res.in, acharya.vishalacharya@gmail.com

Awards i) Indian Science Congress Association (ISCA) Young Scientist

Award -2014 (New Biology)

ii) Young Scientist Award for Best Paper Presentation (2014) by

Society of Plant Biochemistry and Biotechnology, New Delhi iii) Bioclues Innovation, Research and Development (BIRD)

Award-2016 from International Society for Computational Biology and Asia-Pacific Bioinformatics Network under the age of 35.

iv) Awarded for 2nd Position – Incentive Awards for Publications-2016 for publications in Year 2015, from BITSnet, Department of Biotechnology, Govt. of India (including my 5 out of total 8 papers).

v) Elected as "Associate" of National Academy of Agriculture

Sciences (NAAS) under Social Sciences (Bioinformatics)-2021

i) Co-Chair of Bioclues organization (an affiliate of International Society of Computational Biology (ISCB)) for the year 2017-2018.

ii) Chairperson and Invited speaker for the "International Conference on Bioinformatics" held at Hans Raj Mahavidalaya

(HMV), Jalandhar, 2019.

iii) As a Co-co-ordinator for "Establishment of Bioinformatics infrastructure for biology through bioinformatics (BTBI) under BTISnet", Department of Biotechnology (DBT), Government of

India.

Fellowships Merits i). ICAR's JRF for MSc Programme-2003

ii).DBT-Fellow (M.Sc), Gov. of India, conducted by JNU-2003

iii). JRF (2005) for PhD granted by DBT, Gov. of India

iv). Travel fellowship by European Science Foundation (Availed)

Supervisory Experience i).Guided **Three Phd students** through Academy of

Scientific and Innovation Research (AcSIR) research program and

one DBT-Research Associate (RA)

ii) Presently, guiding 7 (Seven) PhD students through AcSIR-PhD

program

Computational Experience Perl, Python, Awk, Linux Server Administrator, R Package, GNU,

Matlab, Bioconductor, PHP

Professional Memberships 1.Life Member of the Indian Science Congress Association (ISCA)

Honours

2. Joint Secretary of Bioclues Organization affiliated with International Society for Computational Biology and Asia-Pacific Bioinformatics Network

Countries Visited: Germany; France; UAE (Dubai, Abu Dhabi)

Educational Qualifications

Examination	School/College	Year of	Percentage/
		Passing	Remarks
PhD (Bioinformatics)	Center for DNA Fingerprinting & Diagnostics (CDFD), Hyderabad	2005- 2011	PhD Degree Awarded
M.Sc. (Biotechnology)	Madurai Kamaraj University, Madurai (T.N)	2003-2005	75.4%

Scientific Contributions:

- * Developed **artificial intelligence based software** (Support vector Machine) named "*Hansa*" ("http://www.cdfd.org.in/HANSA/) for discrimination of pathogenic from nonpathogenic missense mutations with 10% more accurate than best known methods (**Human Mutation**, **2012**; **2013**)
- * "Hansa" has been highlighted in annual virtual issue of Human Mutation entitled "Evaluating Mutation Pathogenicity" for the year 2012.
- * The National Genetics Reference Laboratory (NGRL), Manchester, UK, on analyzing popular missense prediction tools advised usage of "Hansa" as one of the three tools, thus emphasizing the importance of this software that will be highly beneficial to the interested biological community.
- * His **Al-based logistic regression modeling** method suggested plant derived molecules from Himalayas as anti-oral cancer agents (**BMC medical genomics**, 2015, **highly accessed article**; **Molecular BioSystems**, 2015)
- * His stringent Hidden Markov model-based approach for prediction of disease-resistance in *Malus x domestica* (apple) genome (**Plos One**, 2014) was being successfully applied in many **other plant sequenced genomes** supported by ≥ 80 publications.
- *Discovered **novel domain** (animal like defense regulator NACHT domain) for the first time related to immune response in **early green plants** (**Molecular genetics & Genomics**, 2018; **PloS One**; 2016)
- *His conceived developed pipeline on molecular adaptational analysis (https://github.com/fgcsl/Molecular-adaptational-analysis-cold) can easily be applied to any sequenced genomes even by non-bioinformatics researchers (Genomics, 2020).
- *His another conceived **computational pipeline on prediction of transcription factors** (https://github.com/ShivalikaP/R_serpentina_transcriptome_annotation) led into the investigation and organization of secondary metabolites in *Rauvolfia serpentina* (**Plant Molecular biology reporter**, 2015) and abiotic stress (NAC) in potato (**DNA Research**, 2013 (8th most read article).
- *His conceived pipeline on first ever implemented deep learning approach (DLNet) to understand the plant immune genes network response architecture using genomics data https://gitlab.com/ravisaroch/dlnet-host-pathogen-interaction/ (iScience, 2022, IF =6.1)
- *His conceived pipeline https://gitlab.com/neeraj-24/A-HIOT on advanced Virtual Screening framework-automated hit identification and optimization tool (A-HIOT)-comprises chemical

space-driven stacked ensemble for identification and protein space-driven deep learning architectures for optimization of an array of specific hits for fixed protein receptors (Journal of Cheminformatics, 2022, IF =8.48)

*His conceived protocol for network and expression integration to identify potential defense gene in host-pathogen interactions https://gitlab.com/ravisaroch/star-protocols (Star Protocols, 2023)

*His conceived pipeline on clustering of protein molecule into active/inactive targets (https://gitlab.com/neeraj-24/clustering-of-uncharacterised-data) by applying unsupervised machine learning using atom pair, fingerprints and similarity indices as features (Computers in Biology & Medicine, 2023, IF=6.698)

Technology/Software Developed

SI. No	Details of Product/Technology/Software * (Also write in one bullet point its impact in terms of adoption, commercialization and revenue generated)	Main role as a Developer, Co-developer or Associate
1.	Software: His conceived pipeline https://gitlab.com/neeraj-24/A-HIOT on advanced Virtual Screening framework-automated hit identification and optimization tool (A-HIOT)-comprises chemical spacedriven stacked ensemble for identification and protein space-driven deep learning architectures for optimization of an array of specific hits for fixed protein receptors (Journal of Cheminformatics, 2021)	Developer
2.	Software: His conceived pipeline on plant-pathogen interactions can be used to investigate network responses in various stress/plant situations https://gitlab.com/ravisaroch/dlnet-host-pathogen-interaction/ (iScience, 2022)	Developer
3.	Software: His conceived protocol for network and expression integration to identify potential defense gene in host-pathogen interactions https://gitlab.com/ravisaroch/star-protocols (Star Protocols, 2023)	Developer
4.	Software: His conceived pipeline on clustering of protein targets into active/inactive (https://gitlab.com/neeraj-24/clustering-of-uncharacterised-data) by applying unsupervised machine learning using atom pair, fingerprints and similarity indices as features (Computers in Biology & Medicine, 2023)	Developer
5.	Software: His conceived computational pipeline on prediction of transcription factors (https://github.com/ShivalikaP/R serpentina transcriptom	Developer

6.	 e_annotation) led into the investigation and organization of secondary metabolites in <i>Rauvolfia serpentina</i> (Plant Molecular biology reporter, 2015) and abiotic stress (NAC) in potato (DNA Research, 2013 (8th most read article) Software: His conceived developed pipeline on molecular 	Developer
	adaptational analysis (https://github.com/fgcsl/Molecular-adaptational-analysis-cold) can easily be applied to any sequenced genomes even by non-bioinformatics researchers (Genomics , 2020).	Develope.
7.	Software & Technology: AI based-software "Hansa" http://www.cdfd.org.in/HANSA/ (Human Mutation,2012; 2013) wherein novel biological-based features implemented (Gribskov's score) has been applied for future prediction of disease-resistance genes	Developer
8.	<u>Software & Products:</u> Developed novel computational pipeline for screening available Himalayan plant-derived molecules by means of the advanced Artificial Intelligence AI-based logistic regression modeling method suggested plant derived molecules from Himalayas (Molecular BioSystems, 2015; BMC medical genomics , 2015, highly accessed article). for pharmaceutical discovery as an antioral cancer agents	Developer
9.	<u>Products:</u> By means of the integrated Artificial intelligence- and system biology- approaches, novel animal-like defense apoptosis/cell death regulator gene "NACHT NTPase" discovered for the first time in the early green plant lineages (green algae and lycophyte) can be used for the enhancement of disease-resistance genes in many crop plants by transgenic approaches (Molecular Genetics and Genomics, 2017; PloS One, 2016)	Developer
10.	Software: HNHdb database (http://bicmku.in:8081/hnh.) associated with a range of DNA binding proteins domain HNHc (SMART id: SM00507) which performs a variety of binding and cutting functions especially used in CRISPR/Cas9 system (Bioinformation, 2009)	Developer

Selected Publications

- Neeraj Kumar & Vishal Acharya*, Machine intelligence-guided selection of optimized inhibitor for human immunodeficiency virus (HIV) from natural products. Computers in Biology and Medicine Volume 153, February 2023, 106525 (IF=6.698)
 Software link: https://gitlab.com/neeraj-24/clustering-of-uncharacterised-data
- 2. Ravi Kumar, Vishal Acharya*, Deep learning based protocol to construct an immune-related gene network of host-pathogen interactions in plants. STAR Protocols, Cell Press 4,1, 2023.

Software link: https://gitlab.com/ravisaroch/star-protocols

- 3. Neeraj Kumar & Vishal Acharya* Machine intelligence-driven framework for optimized hit selection in virtual screening. Journal of Cheminformatics 2022 Jul 22;14(1):48. doi: 10.1186/s13321-022-00630-7. PMID: 35869511; PMCID: PMC9306080 (IF =8.48) Software link: https://gitlab.com/neeraj-24/A-HIOT
- 4. Ravi Kumar, Abhishek Khatri & **Vishal Acharya*** Deep learning uncovers distinct behavior of rice network to pathogens response **iScience**, Volume 25, Issue 7, 104546 (**IF** =**6.1**)

Software link: https://gitlab.com/ravisaroch/dlnet-host-pathogen-interaction/

Database link: https://fgcsl.ihbt.res.in/RicePathDLNet

- 5. Meetal Sharma, Prince Anand, Yogender Padwad, Vivek Dogra, Vishal Acharya* DNA damage response proteins synergistically affect the cancer prognosis and resistance Free Radical Biology & Medicine 2021 Nov 27;S0891-5849 (21) 00831-5. doi:10.1016/j.freeradbiomed.2021.11.033 (IF=8.1)
- **6.** Srijana Mukhia#, Abhishek Khatri#, **Vishal Acharya***, Rakshak Kumar* Comparative genomics and molecular adaptational analysis of Arthrobacter from Sikkim Himalaya provided insights into its survivability under multiple high-altitude stress **Genomics** (2020) https://doi.org/10.1016/j.ygeno.2020.12.001 (**IF=4.31**)
 Software Link: https://github.com/fgcsl/Molecular-adaptational-analysis-cold
- 7. Vinay Randhawa, Anil Kumar Singh & Vishal Acharya* A systematic approach to prioritize drug targets using machine learning, a molecular descriptor-based classification model, and high-throughput screening of plant derived molecules: a case study in oral cancer Molecular Omics (RSC), 2015, 3362-3377 DOI: 10.1039/C5MB00468C (IF= 4.212)

Details of Publications is provided at the link: https://fgcsl.ihbt.res.in/fgcsl_lab/publication.php

Books Authored:

1. VijayKumar Muley & Vishal Acharya (2013) Genome-Wide Prediction and Analysis of Protein-Protein Functional Linkages in Bacteria, Springer Briefs in Systems Biology Volume 2,1-2, https://doi.org/10.1007/978-1-4614-4705-4 1 Online ISBN 978-1-4614-4705-4

Book Chapters:

 Neeraj Kumar & Vishal Acharya* Metabolomics Resources: An Introduction of Databases and Their Future Prospective (2018) Recent Trends and Techniques in Plant Metabolic Engineering Springer DOI (Editors: Sudesh Kumar Yadav, Vinay Kumar, Sudhir P Singh) DOI: https://doi.org/10.1007/978-981-13-2251-8

^{*}Corresponding author