

# Divya Karade

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## Professional Summary

- AI researcher in drug discovery with over 6 years of experience in Cheminformatics, Computational Biology and Metabolomics
- National-award-winning innovator for “AIDrugApp: Artificial Intelligence-based Web-Application for Drug Discovery”
- Proficient knowledge in Drug Designing, Machine/Deep Learning, Data Analytics, Algorithms and small molecular datasets
- Excellent record of inter-disciplinary collaboration with researchers from biotechnology, chemistry & computational science fields
- Solid publication record with 4 first-author publications in peer-reviewed journals and many in the pipeline

## Technical Skills

- **Programming Languages:** Python, R, MySQL, HTML
- **Machine learning:** TensorFlow, Keras, scikit-learn, streamlit, NumPy, pandas, Matplotlib, seaborn, Building and deploying ML/DL algorithms, Feature engineering, Auto-ML, Auto-DL, Web-app development, cloud platform – Heroku, AWS
- **Data Science & Miscellaneous Technologies:** Biological and Chemical Data analysis, Data science pipeline (cleansing, wrangling, EDA, visualization, modeling, interpretation) Statistics, Hypothesis testing, Excel, GitHub
- **Cheminformatics:** Structure- & ligand-based drug design, data/ text mining, Network analysis, Chemical structure analysis, SAR/ QSAR, virtual screening, virtual libraries, ADME, scaffold analysis, MOE, ChemAxon, ChemDraw, Cytoscape, RDKit.
- **Bioinformatics:** Molecular docking and analysis using computational chemistry tools like Schrodinger, MOE, AutoDock, etc
- **Metabolomics:** Untargeted metabolomics, Mass spectrometric data analysis, Univariate and Multivariate statistical analysis
- **Genomics & Proteomics:** DNA/RNA/protein extraction, PCR, Phylogenetic statistical analysis, Electrophoresis tech.- PAGE
- **Patinformatics:** Prior art search, Freedom to operate analysis, Landscaping, Novelty check

## Innovation Project

### AIDrugApp: Artificial Intelligence-based Virtual Screening Web-App for Drug Discovery

*Sep 2020 - Present*

<https://sars-covid-app.herokuapp.com/>

(Python3 | TensorFlow | Keras | scikit-learn | Matplotlib | Streamlit | Heroku)

- Innovated and founded a user-friendly web application platform called ‘AIDrugApp’ for drug discovery
- Created and Implemented DL algorithms to AIDrugApp for bioactivity prediction of user-defined molecules against SARS-CoV-2
- Originated and integrated custom machine learning tools like Auto-Multi-ML, Auto-DL to AIDrugApp

## Education

### Ph.D. in Biological Science (Cheminformatics, Metabolomics and Computational Biology)

*Jan 2015 - Thesis submitted*

Academy of Scientific and Innovative Research (AcSIR)

*Pune, Maharashtra, India*

- Thesis: Cheminformatics based investigation of plant metabolites for their medicinal and crop protection values

### Master of Science (Plant Biotechnology) - Cumulative GPA: 8.14 /10.0

*Sep 2008 - Nov 2010*

A.N.G.R.A. University, Dept. of Biotechnology

*Hyderabad, India*

- Thesis: Characterization of genetic variability in cowpea using morphological, biochemical and molecular markers.

### Bachelor of Science (Agricultural Science) - Cumulative GPA: 7.68 /10.0

*Aug 2004 - Aug 2008*

Dr. P.D.K.V. University, College of Agriculture

*Nagpur, Maharashtra, India*

- Relevant Coursework: Statistics, Maths, Plant Biotechnology, Plant Biochemistry, Molecular Biology, Genetics, Microbiology

## Experience

### Senior Research Fellow

*Jan 2017 - Present*

CSIR National Chemical Laboratory, Chemical Engineering and Process Development Division

*Pune, India*

- Designed and executed laboratory biochemical experiments to optimize soybean tissue samples for LC/MS studies
- Conceptualized analyzing methods for mass spectrometric data (7185 mass features) using machine learning tools and techniques
- Discovered and reported the presence of 14 novel small molecules in soybean through tandem mass spectrometry analysis
- Developed virtual library of 815 novel drug-like molecules from soybean small molecules through cheminformatics analysis
- Investigated and designed 169 pesticide-like molecules from 162 plants using cheminformatics tools and QSAR approaches

### Junior Research Fellow

*Jan 2015 - Dec 2016*

CSIR National Chemical Laboratory, Chemical Engineering and Process Development Division

*Pune, India*

- Analyzed molecular data (n=3459) by text mining 91206 PubMed literature related to 104 Indian medicinal plants
- Formulated 4 novel computational protocols for designing novel molecules from small molecules of medicinal plants & food crops
- Created and Contributed virtual library of 4147 novel drug-like molecules for building database/ toolkit of Indian medicinal plants

## Patent Analyst (Life Science)

Mar 2013 - Dec 2014

CSIR Unit of Research & Development of Information Products, Life Science Division

Pune, India

- Examined current technology alerts and patent updates in the field of life sciences for 5 projects
- Interpreted comparative and competitive patent portfolio after novelty, patentability and prior art search
- Mapped data for constructing a technological patent landscape

Reviewer for the “Journal of Experimental and Theoretical Artificial Intelligence” (Taylor & Francis)

Feb 2021- Present

## Honors & Awards

### National & International

- National-level AWS DeepRacer (**Selected in top 100**) at Woman's league for community car races driven by reinforcement learning codes in python through AWS cloud-based 3D racing simulator organized by AWS Educate Apr 2021
- National Award (Consolation - **Selected in top 10 out of 377 shortlisted applicants**) for “AIDrugApp” at the open-source “Drug Discovery Hackathon-2020” against Covid-19 in phase-1 under moonshot category organized by Govt. of India. Mar 2021
- National Award (Certificate of Merit - **Selected in top 4 out of 31 finalists**) for “AIDrugApp” in student engineering model competition under “Digital India” theme held at “Indian International Science Festival” organized by Govt. of India Dec 2020

### Academic

- Awarded Senior Research Fellowship for continuing Ph.D., CSIR-UGC (National Eligibility Test) NET exam, India 2017
- All India rank **90** for Junior Research Fellowship for Ph.D. admission in Life-Science, CSIR-UGC NET exam, India 2013
- Awarded for Graduate Aptitude Test in Engineering (Biotechnology) for Ph.D. admission, MHRD, India 2011, 2012
- All India rank **64** in National Eligibility for Lectureship in Life-Science, CSIR-UGC NET, India 2011
- Awarded National Eligibility for Lectureship in Plant biotechnology by ASRB NET, India 2011
- All India rank **31** in Plant Biotechnology common entrance exam for P.G. admission, JNU, India 2008
- All India rank **14** in Plant Biotechnology entrance exam for P.G. admission, ICAR, India 2008

## Publications

- **Divya Karade**, D. V., N. Kadoo, R. Vyas, P. K. Ingle and M. Karthikeyan. Design of Novel Drug-like Molecules using Informatics Rich Secondary Metabolites Analysis of Indian Medicinal and Aromatic Plants. *Combinatorial Chemistry & High Throughput Screening* **2020**, 23 (10), pp. 1113-1131. (Impact factor: 1.2, [click here](#))
- **Divya Karade**, Vikas Karade. AIDrugApp: Artificial Intelligence-based Web-App for Virtual Screening of Inhibitors against SARS-COV-2. *ChemRxiv. Preprint* **2020**. (Web-App: [click here](#))
- **Divya Karade**. AutoDL: Automated Deep Learning (Machine learning module of AIDrugApp - Artificial Intelligence Based Virtual Screening Web-App for Drug Discovery) (Version 1.0.0). *Zenodo*. **2021**. (Software: [click here](#))
- **Divya Karade**, S. Sivaramakrishnan, K. Venkateswaran, R.S. Reddy. Characterization of genetic variability among cowpea (*Vigna Unguiculata* L. Walp) germplasm using morphological, biochemical and molecular markers. *Jour. Of Med. Sci. & Tech.* **2012**, 1 (2), pp. 43-61. ([click here](#))
- **Divya Karade**, Vikas Karade. AIDrugApp: Artificial Intelligence-based Web-App for Virtual Screening of Inhibitors against SARS-COV-2. *Journal of Experimental and Theoretical Artificial Intelligence*. **2021**. (Impact factor: 2.11, Accepted)
- **Divya Karade**, S. Mundhe, N. Kadoo, M. Ratnaparkhe, R. Vyas and M. Karthikeyan. Bridging In-Silico and Experimental: Chemoinformatics Analysis for Mass Spectrometry-Based Metabolomics Study of Soybean. *Metabolomics*. **2021**. (Impact factor: 3.167, Under review)

## Research Presentations

### Posters

- **Divya Karade**, N. Kadoo, M. Karthikeyan (**2019**) ‘Antidiabetic Drug Designing Based on Organic Metabolites from Fenugreek: A Chemo- and Bioinformatics Approach’ presented at an international symposium on “Accelerating Biology: Towards Thinking Machines” conducted by C-DAC, Pune
- **Divya Karade**, N. Kadoo, M. Karthikeyan (**2017**) ‘Chemoinformatics Investigation of Organic Metabolites from Soybean Reveals Common Drug Scaffolds’ presented at CSIR-National Chemical Laboratory, Pune on the “National Science Day”
- **Divya Karade**, N. Kadoo, M. Karthikeyan (**2017**) ‘Drug Design based on Metabolomics of Indian Medicinal and Aromatic Plants: A Chemoinformatics Approach’ presented at CSIR-National Chemical Laboratory, Pune on the “National Science Day”

### Workshop

**Instructor** (10 people) and participated in “Chemoinformatics-2018” for skill development program by CSIR-NCL, Pune