# Divya Karade

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# **Professional Summary**

- AI researcher in drug discovery with 7+ years of experience in Biological Sciences (Cheminformatics, Comp. Biology & Biotech.)
- National-award-winning innovator for "AIDrugApp: Artificial Intelligence-based Web-Application for Drug Discovery"
- Skilled in the implementation of R or Python scripts for scientific data analysis and real-life drug discovery problems
- Proficient knowledge in Drug Designing, Machine/ Deep Learning, Data Analytics, Algorithms and small molecular datasets
- Excellent record of inter-disciplinary collaboration with researchers from biology, chemistry & computational science fields
- Solid publication record with first-author publications in peer-reviewed journals and many in the pipeline
- Well versed in popular data science python libraries such as TensorFlow, Keras, scikit-learn, NumPy, Pandas, RDKit, PubChemPy

## **Technical Skills**

- Programming Languages: Python, R/RStudio Bioconductor, HTML
- Machine learning: Building and deploying ML/ DL algorithms, Auto-ML, Auto-DL, Web-app development streamlit, Data analysis -TensorFlow, Keras, scikit-learn, NumPy, Pandas, Data visualization Matplotlib, seaborn, yellowbrick & ggplot2
- Data Science: Metabolomics and Chemical Data analysis, Data science pipeline (cleansing, wrangling, optimization, EDA, modeling, interpretation), Feature engineering, Statistics, Hypothesis testing, GitHub, Data curation, Cloud tech. Heroku, AWS
- Chemoinformatics: Structure & ligand-based drug design, Text mining, Network analysis, Chemical structure analysis, QSAR, virtual screening, virtual libraries, ADME, scaffold analysis, MOE, ChemAxon, ChemDraw, Cytoscape, RDKit, PubChemPy
- Bioinformatics: Molecular docking and analysis using computational chemistry tools like Schrodinger, MOE and AutoDock
- Metabolomics: Untargeted metabolomics, Mass spectrometric data analysis, Univariate and Multivariate statistical analysis

# **Innovation Project**

AIDrugApp: Artificial Intelligence-based Virtual Screening Web-App for Drug Discovery (v1.2.4)

Sep 2020 - Present

https://sars-covid-app.herokuapp.com/ (Python3| TensorFlow| Keras| sklearn | PubChemPy| RDKit | Streamlit | Heroku)

- Innovated and founded a user-friendly and novel web application platform called 'AIDrugApp' for drug discovery
- Bioactivity prediction: Built QSAR based DL algorithms for bioactivity prediction of user-defined molecules against SARS-CoV2
- DeepDocker: Originated and Implemented QSAR based DL algorithms for predicting binding affinity of user-defined molecules
- Custom ML Tools: ML tools integrated to AIDrugApp for user-defined model building, calculations and predictions on user data
  - 1. Auto-Multi-ML: Created for building and comparing user-defined multiple ML models and making predictions on users data
  - 2. Auto-DL: Developed to build user-defined deep learning models with neural networks and making predictions on users data
  - 3. DesCal: Developed for calculating user-defined 2-D descriptors and fingerprints using molecular SMILES of users data
  - 4. Mol\_Identifier: Developed to identify molecular information like chemical name, SMILES & similarity scores

# **Experience**

Senior Research Scientist (QSAR Modeller)

Syngenta Ltd, Jealott's Hill International Research Centre

May 2022 - Present Bracknell, England, UK

Senior Research Fellow Jan 2017 - Dec 2021

CSIR National Chemical Laboratory, Chemical Engineering and Process Development Division

Pune, India

- Bridging In-Silico and Experimental: Chemoinformatics Analysis of Mass Spectrometry-Based Metabolomics Study of Soybean
  - Discovered and reported the presence of 14 novel small molecules in soybean by mass spectrometry data analysis
  - Developed virtual library of 815 novel drug-like molecules from soybean small molecules through chemoinformatics analysis
  - Formulated 4 computational protocols for designing drug-like novel molecules from metabolites of medicinal & food
- <u>Chemoinformatics Investigation on Chemical Defense in Plants</u>
  - Investigated 280 allelochemicals from 162 plants for their pesticide properties by chemoinformatics and QSAR approaches
  - Designed virtual libraries of allelochemicals that were then quantitatively evaluated to screen 169 pesticide-like molecules

Junior Research Fellow Jan 2015 - Dec 2016

CSIR National Chemical Laboratory, Chemical Engineering and Process Development Division

Pune, India

- Design of Novel Drug-like Molecules using Informatics Rich Secondary Metabolites Analysis of Indian Medicinal Plants
  - Analyzed molecular data (n=3459) by text mining 91206 PubMed literature related to 104 Indian medicinal plants
  - Created and Contributed a virtual library of 4147 novel drug-like molecules for building a database 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 and patent updates for constructing a technological patent landscape for 5 life sciences projects
- Interpreted comparative and competitive patent portfolio after novelty, patentability and prior art search

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## **Education**

Thesis: Chemoinformatics 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

## **Honors & Awards**

#### **Invited Talks**

- Invited speaker for an expert talk on "Revolutionising Drug Discovery with AI" as a part of Capacity Building Programmes for NextGen Entrepreneurs organised by Marwadi University Centre for Innovation, Incubation and Research (MUIIR)
- Invited speaker for a national webinar on "Development of web application for drug discovery" organised by Dept. of Biotechnology of Vignan's Foundation of Science, Technology & Research.

  May 2022

#### **National & International**

- National Award (<u>Consolation</u> 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 (<u>Certificate of Merit</u> **Selected in top 4 out of 31 finalists**) for "AIDrugApp" in student engineering model competition under "Digital India" theme held at "India International Science Festival" organized by Govt. of India 

  \*Dec 2020\*\*

#### **Academic**

•	Awarded Senior Research Fellowship for Ph.D. Life-Science, CSIR-UGC (National Eligibility Test) NET exam, Indi	a 2017
•	All India rank 90 for Junior Research Fellowship for Ph.D. admission in Life-Science, CSIR-UGC NET exam, India	2013
•	Awarded in Graduate Aptitude Test in Engineering (Biotechnology) for Ph.D. admission, MHRD, India	2011 & 2012
•	All India rank 64 with National Eligibility for Lectureship in Life-Science, CSIR-UGC NET, India	2011
•	Awarded National Eligibility for Lectureship in Plant Biotechnology, 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**

A list of publications can be viewed on my Google Scholar page. The following are a few recent publications:

- **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* **2022**. (Impact factor: 2.11, <u>click here</u>)
- **Divya Karade**. Custom ML Module of AIDrugApp for Molecular Identification, Descriptor Calculation, and Building ML/DL QSAR Models. *ChemRxiv. Cambridge: Cambridge Open Engage;* **2021**. (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: <a href="click here">click here</a>)
- **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, <u>click here</u>)
- Damini Kolhe, **Divya Karade**. DeepDocker: AIDrugApp module for predicting the binding affinity of molecules against SARS-CoV-2. **2022**. (In preparation)

#### **Research Presentations**

- **Oral presentation** of 'AIDrugApp: Artificial Intelligence Based Virtual Screening Web-App for Drug Discovery' at "Drug Discovery Hackathon-2020" organized by GOI and awarded consolation cash prize (**2021**)
- **Oral presentation** of 'AIDrugApp: Artificial Intelligence Based Virtual Screening Web-App for Drug Discovery' at "Indian International Science Festival-2020" organized by GOI and awarded Certificate of Merit with cash prize (**2020**)
- Poster presentation on 'Antidiabetic Drug Designing Based on Organic Metabolites from Fenugreek: A Chemo-and Bioinformatics Approach' at "International Symposium on Accelerated Biology" conducted by C-DAC, Pune (2019)
- Poster presentation on 'Chemoinformatics Investigation of Organic Metabolites from Soybean Reveals Common Drug Scaffolds presented at CSIR-National Chemical Laboratory, Pune on the "National Science Day" (2017)
- **Poster presentation** on '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" (2017)

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