

# CURRICULUM VITAE

## Filip Miljković, MPharm, PhD

Gothenburg, Sweden

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### Work Experience

*July 2022 – Present*

#### Senior Scientist Computational Chemistry

Medicinal Chemistry, Early Cardiovascular, Renal and Metabolism (CVRM), AstraZeneca R&D, Gothenburg, Sweden

*January 2021 – Present*

#### Affiliate Scientist

Department of Life Science Informatics, b-it, LIMES Program Unit Chemical Biology and Medicinal Chemistry, Rheinische Friedrich-Wilhelms-Universität Bonn, Germany

*February 2020 – June 2022*

#### Senior Data Scientist

Imaging and Data Analytics, Clinical Pharmacology & Safety Sciences, AstraZeneca R&D, Gothenburg, Sweden

### Education

*June 2016 – December 2019*

#### PhD (Dr. rer. nat.) in Computational Life Sciences

Department of Life Science Informatics, b-it, LIMES Program Unit Chemical Biology and Medicinal Chemistry, Rheinische Friedrich-Wilhelms-Universität Bonn, Germany

Grade: 0.7 (*Magna cum laude*)

Thesis: Chemoinformatics-Driven Approaches for Kinase Drug Discovery

Supervisor: Prof. Jürgen Bajorath

*October 2009 – September 2014*

#### Master's degree in Pharmacy (MPharm, Integrated studies)

Department of Pharmacy, Faculty of Medicine, University of Niš, Serbia

Grade: 9.58/10.00

Thesis: An Overview of Dipeptidyl Peptidase-IV Inhibitors

Supervisor: Prof. Andrija Šmelcerović

### International Research Visits

*July 2013 – August 2013*

#### IPSF Student Exchange Program

Department of Analytical Chemistry, Medical University of Gdańsk, Poland

Description: Quality consistency evaluation of *Melissa officinalis* L. samples using HPLC fingerprints

Supervisor: Agnieszka Viapiana (née Arceusz), PhD

<b>Technical Skills</b>	Data extraction/curation (KNIME, RDKit, and OpenEye Chemistry Toolkit), SAR analysis, Network analysis (Cytoscape, Gephi, NetworkX Python package), 2D/3D ligand similarity (descriptor-/fingerprint-based, ROCS), Structure and ligand-based virtual screening, Molecular modeling suites (MOE, Schrödinger), Molecular docking (MOE Dock, Maestro AutoDock, PLANTS, DockTite for covalent docking), Protein-ligand interaction fingerprints, Machine learning (SVM, RF, KNN, XGB, DNN, GCNN; scikit-learn, TensorFlow)
<b>Medicinal Chemistry/Chemical Biology Skills</b>	Computational chemistry design in several ongoing drug projects, Chemical library enumeration, <i>In vitro</i> / <i>In vivo</i> assay knowledge and translation, Multi-parameter drug optimization, ADME/PK, Chemical toxicology/Safety sciences, Basic synthetic chemistry understanding
<b>Computer Skills</b>	Python (advanced knowledge), Bash (basic knowledge), R (basic knowledge), SQL (basic knowledge), ChemOffice, Biovia, KNIME/Pipeline Pilot, Office suite, Windows/Linux
<b>Languages</b>	Serbian (native) English (fluent): TOEFL (109/120) Swedish (good) German (good)
<b>Teaching Experience</b>	<p><b>Master's Program in Pharmacy at the Medical faculty, University of Niš, Serbia</b></p> <ul style="list-style-type: none"> <li>• "General Chemistry with Stoichiometry", October 2010 – January 2011.</li> <li>• "Organic Chemistry", March 2011 – June 2011.</li> </ul> <p><b>Master's Program in Life Science Informatics at the University of Bonn, Germany</b></p> <ul style="list-style-type: none"> <li>• "Introduction to Chemistry", September 2016 – October 2016.</li> <li>• "Structural Bioinformatics", October 2016 – January 2018.</li> <li>• "Molecular Modeling and Drug Design", April 2017 – February 2019.</li> <li>• "Programming Lab I" (Python), April 2019 – June 2019.</li> <li>• "Introduction to Machine Learning Tutorial", April 2019 – June 2019.</li> </ul> <p><b>Visiting Students Coming from Collaborative Projects between University of Bonn, Germany and Kyoto University, Japan or Waseda University, Japan</b></p> <ul style="list-style-type: none"> <li>• "Structural Bioinformatics and Molecular Modeling", October 2017 – December 2019.</li> </ul>
<b>Mentorship Experience</b>	<ul style="list-style-type: none"> <li>• Supervision of two Master theses at Rheinische Friedrich-Wilhelms-Universität Bonn, Germany</li> <li>• Individual development plan/Career development discussion (long-term) with two graduate scientists from Data Science &amp; AI Graduate Program, AstraZeneca R&amp;D, Sweden</li> </ul>

## Publications

37. **Miljković, F.\***; Medina-Franco, J. L. Artificial Intelligence-Open Science Symbiosis in Chemoinformatics. *Artif. Intell. Life Sci.* **2023**, 5, 100096.
36. Chen, Y.; Seidel, T.; Jacob, R. A.; Hirte, S.; Mazzolari, A.; Pedretti, A.; Vistoli, G.; Langer, T.; **Miljković, F.**; Kirchmair, J. Active Learning Approach for Guiding Site-of-Metabolism Measurement and Annotation. *J. Chem. Inf. Model.* **2024**, 64, 348–358.
35. Bajorath, J.; Gardner, S.; Grisoni, F.; Horta Andrade, C.; Kirchmair, J.; Landon, M.; Medina-Franco, J.-L.; **Miljković, F.**; Montanari, F.; Rodríguez-Pérez, R. First-generation Themed Article Collections. *Artif. Intell. Life Sci.* **2023**, 4, 100088.
34. Xerxa, E.; **Miljković, F.**; Bajorath, J. Data-Driven Global Assessment of Protein Kinase Inhibitors with Emphasis on Covalent Compounds. *J. Med. Chem.* **2023**, 66, 7657-7665.
33. Bai, P.; **Miljković, F.**; John, B.; Lu, H. Interpretable Bilinear Attention Network with Domain Adaptation Improves Drug–Target Prediction. *Nat. Mach. Intell.* **2023**, 5, 126-136.
32. Gill, G.; Moullet, M.; Martinsson, A.; **Miljković, F.**; Williamson, B.; Arends, R. H.; Pilla Reddy, V. Evaluating the Performance of Machine-Learning Regression Models for Pharmacokinetic Drug–Drug Interactions. *CPT: Pharmacomet. Syst. Pharmacol.* **2023**, 1, 122-134.
31. Gill, G.; Moullet, M.; Martinsson, A.; **Miljković, F.**; Williamson, B.; Arends, R. H.; Pilla Reddy, V. Comparing the Applications of Machine Learning, PBPK, and Population Pharmacokinetic Models in Pharmacokinetic Drug–Drug Interaction Prediction. *CPT: Pharmacomet. Syst. Pharmacol.* **2022**, 11, 1560-1568.
30. Martínez Mora, A.; Mogemark, M.; Subramanian, V.; **Miljković, F.\*** Interpretation of Multi-Task Clearance Models from Molecular Images Supported by Experimental Design. *Artif. Intell. Life Sci.* **2022**, 2, 100048.
29. Trapotsi, M.-A.; Mouchet, E.; Williams, G.; Monteverde, T.; Juhani, K.; Turkki, R.; **Miljković, F.**; Martinsson, A.; Mervin, L.; Pryde, K. R.; Müllers, E.; Barrett, I.; Engkvist, O.; Bender, A.; Moreau, K. Cell Morphological Profiling Enables High-Throughput Screening for PROteolysis TArgeting Chimera (PROTAC) Phenotypic Signature. *ACS Chem. Biol.* **2022**, 17, 1733-1744.
28. Martínez Mora, A.; Subramanian, V.; **Miljković, F.\*** Multi-Task Convolutional Neural Networks for Predicting *In Vitro* Clearance Endpoints from Molecular Images. *J. Comput. Aided Mol. Des.* **2022**, 36, 443-457.
27. Obrezanova, O.; Martinsson, A.; Whitehead, T.; Mahmoud, S.; Bender, A.; **Miljković, F.**; Grabowski, P.; Irwin, B.; Oprisiu, I.; Conduit, G.; Segall, M.; Smith, G. F.; Williamson, B.; Winiwarter, S.; Greene, N. Prediction of *In Vivo* Pharmacokinetic Parameters and Time-Exposure Curves in Rats Using Machine Learning from the Chemical Structure. *Mol. Pharmaceutics* **2022**, 19, 1488-1504.
26. Rodríguez-Pérez, R.; **Miljković, F.**; Bajorath, J. Machine Learning in Chemoinformatics and Medicinal Chemistry. *Ann. Rev. Biomed. Data Sci.* **2022**, 5, 43-65.
25. Laufkötter, O.; Hu, H.; **Miljković, F.**; Bajorath, J. Structure- and Similarity-Based Survey of Allosteric Kinase Inhibitors, Activators, and Closely Related Compounds. *J. Med. Chem.* **2022**, 65, 922-934.
24. Yoshimori, A.; **Miljković, F.**; Bajorath, J. Approach for the Design of Covalent Protein Kinase Inhibitors via Focused Deep Generative Modeling. *Molecules* **2022**, 27, e570.

23. **Miljković, F.**; Rodríguez-Pérez, R.; Bajorath, J. Impact of Artificial Intelligence on Compound Discovery, Design, and Synthesis. *ACS Omega* **2021**, *6*, 33293-33299.
22. **Miljković, F.\***; Martinsson, A.; Obrezanova, O.; Williamson, B.; Johnson, M.; Sykes, A.; Bender, A.; Greene, N. Machine Learning Models for Human *In Vivo* Pharmacokinetic Parameters with In-House Validation. *Mol. Pharmaceutics* **2021**, *18*, 4520-4530.
21. Hu, H.; Laufkötter, O.; **Miljković, F.**; Bajorath, J. Data Set of Competitive and Allosteric Protein Kinase Inhibitors Confirmed by X-ray Crystallography. *Data in Brief* **2021**, *35*, e106816.
20. Hu, H.; Laufkötter, O.; **Miljković, F.**; Bajorath, J. Systematic Comparison of Competitive and Allosteric Kinase Inhibitors Reveals Common Structural Characteristics. *Eur. J. Med. Chem.* **2021**, *214*, e113206.
19. **Miljković, F.**; Chaudhari, R. Members of our Early Career Panel Highlight Key Research Articles on the Theme of Computer-Aided Drug Design. *Future Drug Discov.* **2020**, *2*, FDD52.
18. Rodríguez-Pérez, R.; **Miljković, F.**; Bajorath, J. Assessing the Information Content of Structural and Protein–Ligand Interaction Representations for the Classification of Kinase Inhibitor Binding Modes via Machine Learning and Active Learning. *J. Cheminform.* **2020**, *12*, e36.
17. **Miljković, F.**; Xiong, R.; Sivakumar, D.; Brown, C. A. Members of our Early Career Panel Highlight Key Research Articles on the Theme of Drug Repurposing. *Future Drug Discov.* **2020**, *2*, FDD39.
16. **Miljković, F.**; Rodríguez-Pérez, R.; Bajorath, J. Machine Learning Models for Accurate Prediction of Kinase Inhibitors with Different Binding Modes. *J. Med. Chem.* **2020**, *63*, 8738-8748.
15. **Miljković, F.**; Bajorath, J. Data Structures for Computational Compound Promiscuity Analysis and Exemplary Applications to Inhibitors of the Human Kinome. *J. Comput. Aided Mol. Des.* **2020**, *34*, 1-10.
14. González-Medina, M.; **Miljković, F.**; Haase, G. S.; Drueckes, P.; Trappe, J.; Laufer, S.; Bajorath, J. Promiscuity Analysis of a Kinase Panel Screen with Designated p38 alpha Inhibitors. *Eur. J. Med. Chem.* **2020**, *187*, 112004.
13. Feldmann, C.; **Miljković, F.**; Yonchev, D.; Bajorath, J. Identifying Promiscuous Compounds with Activity Against Different Target Classes. *Molecules* **2019**, *24*, e4185.
12. **Miljković, F.**; Bajorath, J. Data Structures for Compound Promiscuity Analysis: Cliffs, Pathways, and Hubs Formed by Inhibitors of the Human Kinome. *Future Sci. OA* **2019**, *5*, FSO404.
11. **Miljković, F.**; Vogt, M.; Bajorath, J. Systematic Computational Identification of Promiscuity Cliff Pathways Formed by Inhibitors of the Human Kinome. *J. Comput. Aided Mol. Des.* **2019**, *33*, 559-572.
10. Blaschke, T.; **Miljković, F.**; Bajorath, J. Prediction of Different Classes of Promiscuous and Nonpromiscuous Compounds Using Machine Learning and Nearest Neighbor Analysis. *ACS Omega* **2019**, *4*, 6883-6890.
9. **Miljković, F.**; Bajorath, J. Computational Analysis of Kinase Inhibitors Identifies Promiscuity Cliffs across the Human Kinome. *ACS Omega* **2018**, *3*, 17295–17308.
8. **Miljković, F.**; Bajorath, J. Data-Driven Exploration of Selectivity and Off-Target Activities of Designated Chemical Probes. *Mol.* **2018**, *23*, e2434.
7. **Miljković, F.**; Bajorath, J. Evaluation of Kinase Inhibitor Selectivity Using Cell-Based Profiling Data. *Mol. Inform.* **2018**, *37*, e1800024.

6. **Miljković, F.**; Bajorath, J. Reconciling Selectivity Trends from a Comprehensive Kinase Inhibitor Profiling Campaign with Known Activity Data. *ACS Omega* **2018**, 3, 3113–3119.
5. **Miljković, F.**; Bajorath, J. Exploring Selectivity of Multikinase Inhibitors across the Human Kinome. *ACS Omega* **2018**, 3, 1147–1153.
4. **Miljković, F.**; Kunimoto, R.; Bajorath, J. Identifying Relationships between Unrelated Pharmaceutical Target Proteins on the Basis of Shared Active Compounds. *Future Sci. OA* **2017**, 3, FSO212.
3. Smelcerovic, A.; **Miljkovic, F.**; Kolarevic, A.; Lazarevic, J.; Djordjevic, A.; Kocic, G.; Anderluh, M. An Overview of Recent Dipeptidyl Peptidase-IV Inhibitors: Linking Their Structure and Physico-Chemical Properties with SAR, Pharmacokinetics and Toxicity. *Curr. Top. Med. Chem.* **2015**, 15, 2342-2372.
2. Toropov, A. A.; Veselinović, J. B.; Veselinović, A. M.; **Miljković, F. N.**; Toropova, A. P. QSAR Models for 1,2,4-Benzotriazines as Src Inhibitors Based on Monte Carlo Method. *Med. Chem. Res.* **2015**, 24, 283-290.
1. Toropova, A. P.; Toropov, A. A.; Veselinović, J. B.; **Miljković, F. N.**; Veselinović, A. M. QSAR Models for HEPT Derivates as NNRTI Inhibitors Based on Monte Carlo Method. *Eur. J. Med. Chem.* **2014**, 77, 298-305.

#### Selected Abstracts and Conference Publications

2. "Proceedings of IEEE International Conference on Bioinformatics and Biomedicine (BIBM 2021). IEEE International Conference on Bioinformatics and Biomedicine (BIBM 2021)", December 9-12, 2021, Institute of Electrical and Electronics Engineers (IEEE), Virtual. Conference Paper: Bai, P.; **Miljković, F.**; Ge, Y.; Greene, N.; John B.; Lu. H. "Hierarchical Clustering Split for Low-Bias Evaluation of Drug-Target Interaction Prediction".
1. "Chemoinformatics Strasbourg Summer School", June 25-29 June 2018, University of Strasbourg, Strasbourg, France. Poster: **Miljković, F.**; Bajorath, J. "Exploring Selectivity of Multi-kinase Inhibitors across the Human Kinome", awarded as the best poster by public choice.

#### Selected Oral Presentations

3. ACS Fall 2023, Division of Chemical Information, Chemical Informatics (R)evolution: Towards Democratization and Open Science, August 13-17, 2023, American Chemical Society (ACS), In Person, Oral Presentation: **Miljković, F.** "Machine Learning Models for Predicting Human *In Vivo* PK Parameters Using Chemical Structure and Dose".
2. Industry Symposium on "AI in the Life Sciences", June 26-27, 2023, Bonn-Aachen International Center for Information Technology (B-IT), University of Bonn, Bonn, Germany, In Person, Oral Presentation: **Miljković, F.** "Machine Learning Models for Predicting Human *In Vivo* PK Parameters Using Chemical Structure and Dose".
1. "19<sup>th</sup> International Workshop on (Q)SAR in Environmental and Health Sciences – QSAR 2021 From QSAR to New Approach Methodologies (NAMs)", June 7-9, 2021, American Society for Cellular and Computational Toxicology (ASCCT), Virtual. Oral Presentation: **Miljković, F.**; Martinsson, A.; Obrezanova, O.; Williamson, B.; Johnson, M.; Oprisiu, I.; Bender, A.; Greene N. "Machine Learning Models for Predicting Human *In Vivo* PK Parameters Using Chemical Structure and Dose".