

Derek van Tilborg www.derekvantilborg.com

Conference

contributions

PhD-trained scientist specialized in molecular machine learning for drug discovery. Experienced in deep learning, chemoinformatics, and interdisciplinary collaboration with wet-lab teams.

Education

2021-2025 (expected)

PhD, Biomedical Engineering

Eindhoven University of Technology Molecular machine learning group of F. Grisoni

2019-2021

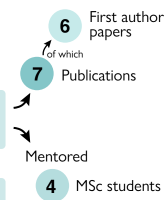
MSc, Bioinformatics

Wageningen University & Research Average grade of 8.5, MSc thesis 9.5 out of 10

2015-2019

BSc, Biomedical science

Avans University of Applied Sciences, Breda Average grade of 8.0, BSc thesis 9.0 out of 10



BSc students

Skills

Scientific & domain knowledge

Artificial intelligence Statistics Pharmacology Molecular (Bio)chemistry biology

Molecular machine learning

Cheminformatics Molecular **Bioinformatics** docking Uncertainty Generative estimation models

Graph neural networks

Other experience

2021

Internship - Radboud Medical Hospital

Developed a computational framework for 3D modeling of peptide:MHC complexes, aimed at immunotherapy design.

2018-2019

Internship - Crossfire Oncology

Predicted drug response on an in house panel of cancer cell lines using multimodal machine learning.

Most exciting projects I've spearheaded:

Molecular property prediction with active learning.

A computational study for active learning in low-data regimes. Published in Nature Computational Science 4 (10), 786-796

Molecular deep learning at the edge of chemical space.

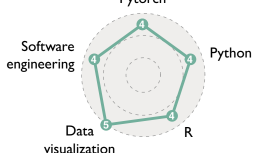
I introduced a new method to estimate model reliability and applied my method prospectively in the lab, discovering multiple novel bioactive molecules across two protein targets. Under review

Nanoparticle design with active learning.

I combined active learning with microfluidics and high-content imaging to design nanoparticles for drug delivery in cancer cells. Published in Digital Discovery 3 (7), 1280-1291

Al-guided evolution of immune cell-specific nanobiologics. I designed nanobody binders using generative protein language models. A collaboration with A. Hokke and K. de Bruin. Manuscript in preparation

Data science / programming skills **Pytorch**



Soft skills

Problem solving Collaboration Self-management & teamwork Communication Communication (written) (verbal)

> Levels of competence are scored according to the Dreyfus model:

1. Novice, 2. Advanced beginner,

3. Competent, 4. Proficient, 5. Expert

List of first author journal articles

Cancers in Agreement? Exploring the Cross-Talk of Cancer Metabolomic and Transcriptomic Landscapes Using Publicly Available Data.

Derek van Tilborg and Edoardo Saccenti. Cancers 13 (3), 393, 2021.

Exposing the Limitations of Molecular Machine Learning with Activity Cliffs.

Derek van Tilborg et al. Journal of Chemical Information and Modeling 62 (23), 5938-5951, 2022.

Structure-based Drug discovery with Deep Learning. Rıza Özçelik, Derek van Tilborg, et al. ChemBioChem, e202200776, 2023.

Machine learning-guided high throughput nanoparticle design.

Ana Ortiz-Perez, Derek van Tilborg, et al. Digital Discovery 3 (7), 1280-1291, 2024.

Deep learning for low-data drug discovery: Hurdles and opportunities.

Derek van Tilborg et al.

Current Opinion in Structural Biology 86, 102818, 2024.

Traversing chemical space with active deep learning for low-data drug discovery.

Derek van Tilborg and Francesca Grisoni. Nature Computational Science 4 (10), 786-796, 2024.

Molecular deep learning at the edge of chemical space.

Derek van Tilborg et al. ChemRxiv, 2025 (preprint).

Selected conference contributions

Poster presentation (best poster prize)

Modeller Forum, online, February 2022

MoleculeACE – a benchmark for molecular machine learning with activity cliffs
Molecular Graphics and Modelling Society - Young

Invited talk

Exposing the limitations of molecular machine learning with activity cliffs.

Valence labs, Molecular Modeling and Drug Design, online, May 2022.

Poster presentation

MoleculeACE: a benchmark for machine learning with activity cliffs.

International Conference on Chemical Structures, Noordwijkerhout, The Netherlands, June 2022.

Poster presentation

Self-supervised learning with graph neural networks for drug discovery.

NWO CHAINS, Veldhoven, The Netherlands, September 2022.

Poster presentation

Traversing Chemical Space with Active Learning. Institute of Complex Molecular Systems - Annual Symposium, Eindhoven, The Netherlands, March 2023.

Poster presentation (best poster prize)

Traversing Chemical Space with Active Learning. 20th IEEE Conference on Computational Intelligence in Bioinformatics and Computational Biology, Eindhoven, The Netherlands, August 2023.

Plenary lecture

Traversing Chemical Space with Active Learning, International Symposium on Bioinformatics and Computer-Aided Drug Discovery, online, September 2023.

Invited talk

Traversing Chemical Space with Active Learning. Figon Dutch medicine days, Oss, The Netherlands, September 2023.

Poster presentation

Machine learning-guided high throughput design. ELLIS machine learning for molecules, online, December 2023.

Poster presentation

Traversing Chemical Space with Active Learning. Institute of Complex Molecular Systems - Annual Symposium, Eindhoven, The Netherlands, March 2024.

Oral presentation

Drug discovery at the edge of chemical space. NWO CHAINS, Veldhoven, The Netherlands, December 2024.

Invited talk

Molecular deep learning at the edge of chemical space. École Polytechnique Fédérale de Lausanne, online, April 2025.

Poster presentation (best poster prize)

Molecular deep learning at the edge of chemical space. Institute of Complex Molecular Systems - Annual Symposium, Eindhoven, The Netherlands, March 2025.