



Robrecht Cannoodt

SOFTWARE ENGINEER · DATA SCIENTIST

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Hello! This is me.

- I'm a software engineer specialised in data science.
- I proudly advocate open research and good software development practices.
- I prefer honesty and multi-perspective thinking over pride and success.

Education

VIB – Ghent University

PHD IN COMPUTER SCIENCE

Ghent, Belgium

Sep 2013 - Dec 2019

- Dissertation: "Modelling single-cell dynamics with trajectories and gene regulatory networks". [Download]
- Science Communication skills: Academic Writing, Presenting, and Data Visualisation
- Bioinformatics skills: Single-Cell Omics, Gene Regulatory Network Inference, Trajectory Inference,
- Data Science skills: Data Analysis, Data Wrangling, Benchmarking, (Co-)authored 24 R Software Packages
- Soft skills: Mentored 6 Master Thesis Students, Teaching Assistant 6 times, Intense collaborations with PhD students

Ghent University

MASTER OF SCIENCE IN COMPUTER SCIENCE ENGINEERING: SOFTWARE ENGINEERING

Ghent, Belgium

Sep 2011 - Jun 2013

- Master Thesis: "Network inference by integrating biclustering and feature selection"
- Programming courses: Programming Languages, Functional Programming
- Engineering courses: Software Architecture, Advanced Computer Architecture, Distributed Software, High Performance Computing
- Other courses: Philosophy and Science, Robotics, Computational Biology

Ghent University

BACHELOR OF SCIENCE IN INFORMATICS

Ghent, Belgium

Sep 2008 - Jun 2011

- Engineering courses: Software Development, Programming, Scripting, Operating Systems, Computer Architecture, Network Management
- Computer Science courses: Machine Learning, Algorithms and Data Structures, Formal Logic
- Math courses: Statistics, Linear Algebra, Geometry, Discrete Math, Calculus

International School of Berne

INTERNATIONAL BACCALAUREATE

Berne, Switzerland

Sep 2003 - Jul 2007

- Taught in English
- Main courses: Math, Physics, Philosophy, Visual Arts

Skills

Research Domains	Data Science, Machine Learning, Software Engineering, Bioinformatics
Communication	Effective Data Visualisation, Academic Writing, Presentations (Technical and Non-technical)
Programming	R, Scala, Java, C++, bash, awk, sed, MySQL, RMarkdown, \LaTeX
Software Development	Git, Docker, Singularity, Travis CI, UML, Agile/Scrum
Languages	Dutch (Native), English (Fluent), French (If-need-be), Hindi (None at all)

Software

Below is a selection of software projects. Visit cannoodt.dev/software for a full list of software I've (co-)authored.

1. Watanabe K and **Cannoodt R**. proxyC: Computes Proximity in Large Sparse Matrices. Role: Contributor. <https://CRAN.R-project.org/package=proxyC>
2. Hastie T, Weingessel A, Hornik K, Bengtsson H and **Cannoodt R**. princurve: Fits a Principal Curve in Arbitrary Dimension. Role: Maintainer. <https://CRAN.R-project.org/package=princurve>
3. **Cannoodt R** and Saelens W. GillespieSSA2: Gillespie's Stochastic Simulation Algorithm for Impatient People. Role: Lead Developer. <https://CRAN.R-project.org/package=GillespieSSA2>
4. **Cannoodt R** and Saelens W. dyno: Inferring, interpreting and visualising trajectories using a streamlined set of packages. Role: Lead Developer. <https://github.com/dynverse/dyno>
5. **Cannoodt R** and Saelens W. lmds: Landmark Multi-Dimensional Scaling. Role: Lead Developer. <https://CRAN.R-project.org/package=lmds>
6. **Cannoodt R** and Saelens W. SCORPIUS: Inferring Developmental Chronologies from Single-Cell RNA Sequencing Data. Role: Lead Developer. <https://CRAN.R-project.org/package=SCORPIUS>
7. **Cannoodt R** and Saelens W. babelwhale: Talking to Docker and Singularity Containers. Role: Lead Developer. <https://CRAN.R-project.org/package=babelwhale>

Publications

Below is a selection of publications. Visit cannoodt.dev/publication or my ORCID profile for a full list of publications.

1. Saelens W*, **Cannoodt R***, Todorov H, Saeys Y. A comparison of single-cell trajectory inference methods. *Nature Biotechnology*. 2019 Apr 1;37(5):547–54. doi:10.1038/s41587-019-0071-9
2. Todorov H, **Cannoodt R**, Saelens W, Saeys Y. Network Inference from Single-Cell Transcriptomic Data. *Gene Regulatory Networks*. 2018 Dec 14;235–49. doi:10.1007/978-1-4939-8882-2_10
3. Weber LM, Saelens W, **Cannoodt R**, Sonesson C, Hapfelmeier A, Gardner PP, et al. Essential guidelines for computational method benchmarking. *Genome Biology*. 2019 Jun 20;20(1). doi:10.1186/s13059-019-1738-8
4. Saelens W, **Cannoodt R**, Saeys Y. A comprehensive evaluation of module detection methods for gene expression data. *Nature Communications*. 2018 Mar 15;9(1). doi:10.1038/s41467-018-03424-4
5. **Cannoodt R**, Saelens W, Sichien D, Tavernier S, Janssens S, Guilliams M, et al. SCORPIUS improves trajectory inference and identifies novel modules in dendritic cell development. *bioRxiv*; 2016 Oct 6; doi:10.1101/079509
6. **Cannoodt R***, Saelens W*, Saeys Y. Computational methods for trajectory inference from single-cell transcriptomics. *European Journal of Immunology*. 2016 Oct 19;46(11):2496–506. doi:10.1002/eji.201646347

*: Equal contribution

Teaching

TEACHING ASSISTANT

2014	Bioinformatics: lessons in R	<i>Biomedical sciences</i>
2015	Bioinformatics: lessons in R	<i>Biomedical sciences</i>
2016	Bioinformatics: lessons in R	<i>Biomedical sciences</i>
2017	Bioinformatics: lessons in R	<i>Biomedical sciences</i>
2018	Bioinformatics: lessons in R	<i>Biomedical sciences</i>
2019	Bioinformatics: lessons in R	<i>Biomedical sciences</i>

MASTER THESIS GUIDANCE

2013-2014	Identificatie van nieuwe kankergenen voor neuroblastoomonderzoek met machine learning	<i>Leen De Baets</i>
2013-2014	Locale cel-type specifieke genexpressie in het myeloïde transcriptoom	<i>Wouter Saelens</i>
2015-2016	Quantitative evaluation of network inference methods for single-cell cancer regulomes	<i>Charlotte De Vogelaere</i>
2015-2016	Comparative review of dimensionality reduction methods for high-throughput single-cell transcriptomics	<i>Sofie Veys</i>
2016-2017	Het afleiden van dynamische grafen op basis van snapshot data	<i>Chloë Guidi</i>
2018-2019	Inferentie van cellontwikkelingstrajecten met machine learning	<i>Jarre Knockaert</i>

Conferences and workshops

ACCEPTED TALKS

2013	Network inference by integrating biclustering and feature selection, BeNeLux Bioinformatics Conference	<i>Brussels, Belgium</i>
2015	Differential module analysis in neuroblastoma regulatory networks, Oncopoint	<i>Ghent, Belgium</i>
2015	Inferring developmental chronologies from single cell RNA, BeNeLux Bioinformatics Conference	<i>Antwerp, Belgium</i>
2016	SCORPIUS: Inferring trajectories along dynamic processes from single-cell RNA-seq data, Single Cell Biology Workshop	<i>Ghent, Belgium</i>
2016	Improving marker gene discovery from high-dimensional single-cell snapshot data, CYTO	<i>Seattle, USA</i>
2016	Unbiased modelling of dynamic processes with single-cell RNA-sequencing, BENELEARN	<i>Kortrijk, Belgium</i>
2017	Unbiased modelling of dynamic processes identifies novel modules in dendritic cell development, VIB Seminar	<i>Veldhoven, Netherlands</i>
2019	Automated building and unit testing, Docker and Singularity, VIB Developers Meeting	<i>Ghent, Belgium</i>
2019	Assessing Accuracy, Robustness and Usability of Single-Cell Trajectory Inference methods, Keystone Symposia: Single Cell Biology	<i>Colorado, USA</i>

ACCEPTED POSTERS

2014	Network Inference by Integrating Biclustering and Feature Selection, N2N Annual Symposium	<i>Ghent, Belgium</i>
2014	Differential Module Analysis in Neuroblastoma Regulatory Networks, BeNeLux Bioinformatics Conference	<i>Luxembourg, Belgium</i>
2015	Differential Module Analysis in Neuroblastoma Regulatory Networks, BIG N2N symposium	<i>Ghent, Belgium</i>
2016	Unbiased modelling of dynamic processes with single-cell RNA-sequencing, Single Cell Biology	<i>Hinxton, United Kingdom</i>
2016	Unbiased modelling of dynamic processes with single-cell RNA-sequencing, BIG N2N symposium	<i>Ghent, Belgium</i>
2016	Improving marker gene discovery from high-dimensional single-cell snapshot data, CYTO	<i>Seattle, USA</i>
2016	Unbiased modelling of dynamic processes with single-cell RNA-sequencing, BENELEARN	<i>Kortrijk, Belgium</i>
2016	Unbiased modelling of dynamic processes with single-cell RNA-sequencing, Single Cell Genomics	<i>Hinxton, United Kingdom</i>
2017	Unbiased modelling of dynamic processes with SCORPIUS identifies novel modules in dendritic cell development, VIB Seminar	<i>Veldhoven, Netherlands</i>
2017	True single cell network inference: Modelling gene regulation of individual cells, Keystone Symposia: Single Cell Omics	<i>Stockholm, Sweden</i>
2017	Generalised framework for and comparison of 24 trajectory inference methods, BeNeLux Bioinformatics Conference	<i>Louvain, Belgium</i>
2018	A comparison of single-cell trajectory inference methods: towards more accurate and robust tools, Single Cell Biology	<i>Hinxton, United Kingdom</i>
2019	Assessing Accuracy, Robustness and Usability of Single-Cell Trajectory Inference methods, Keystone Symposia: Single Cell Biology	<i>Colorado, USA</i>