

# Meghan M. Thommes

meghan.thommes@gmail.com | (617) 702-4989 | Somerville, MA

[linkedin.com/in/meghanthommes](https://www.linkedin.com/in/meghanthommes) | [github.com/megthommes](https://github.com/megthommes) | [meghanthommes.com](https://meghanthommes.com)

## Summary

---

Experienced computational biologist with strong engineering capabilities seeking bioinformatics and engineering opportunities. Looking for roles that nurture growth, foster collaboration, and with tangible impact.

## Skills

---

<b>Languages:</b>	Python, SQL, Bash/Shell
<b>Python Packages:</b>	pandas, sqlalchemy, pydantic, sklearn, statsmodels, numpy, scipy, seaborn, matplotlib
<b>Techniques:</b>	Regression (multivariate, linear, logistic), Random Forest, Clustering/Dimensionality Reduction (PCA, t-SNE, k-means), Data Modeling (conceptual)
<b>Tools:</b>	GitHub, VSCode, Jupyter Notebook, Google Colab, Linux, High-Performance Computing

## Education

---

<b>PhD Biomedical Engineering</b> , Boston University	<b>May 2020</b>
Thesis: <i>Strategies for Engineering Microbial Communities</i>	
<b>BS Biomedical Engineering</b> , Case Western Reserve University	<b>May 2013</b>

## Experience

---

Dyno Therapeutics	Watertown, MA
<b>Scientist I, Computational Biology</b>	<b>Oct. 2021 – March 2024</b>
<ul style="list-style-type: none"><li>Built data analysis pipelines of NGS data in Python using Pydantic models and Matplotlib figures to derive insights on engineered AAV capsids</li><li>Collaborated with wet-lab scientists to design experiments, develop new approaches to data generation, and generate reports for internal stakeholders</li><li>Collaborated with software engineers to construct and improve analysis frameworks used by all computational biologists</li></ul>	
Joyn Bio	Boston, MA
<b>Data Scientist</b>	<b>June 2020 – Sept. 2021</b>
<ul style="list-style-type: none"><li>Evaluated and predicted translation from <i>in vitro</i> to <i>in planta</i> biological performance of engineered microbes by developing modeling frameworks (linear regression, random forest) in Python and R</li><li>Built data integration pipelines and packages in Python and R to clean, standardize, and consolidate data across departments; developed interactive dashboards in Tableau</li></ul>	

Joyn Bio, *continued*

Boston, MA

**Data Scientist**

**June 2020 – Sept. 2021**

- Created conceptual data models to inform modeling efforts, create data templates, and organize data in a centralized location so they can easily be queried and imported into databases
- Collaborated with scientists to design schemas for LIMS and ELN implementation in Benchling; tested API in Python with requests to import existing data; queried PostgreSQL database to generate example dashboards

Insight, *Health Data Science*

Boston, MA

**Fellow**

**Jan. 2020 – April 2020**

- Created “NoveList”, a web app that predicts a Goodreads user’s want-to-read book ratings, enabling them to prioritize what to read next
- Built, trained, and evaluated a collaborative filtering recommender system in Jupyter Notebooks with scikit-surprise, pandas, numpy, and seaborn
- Deployed “NoveList” using streamlit and Heroku

## **Publications and Presentations**

---

**Meghan Thommes**, Taiyao Wang, Qi Zhao, Ioannis Ch. Paschalidis, Daniel Segrè. Designing metabolic division of labor in microbial communities. *mSystems* (2019). doi: [10.1128/mSystems.00263-18](https://doi.org/10.1128/mSystems.00263-18) (Editor’s Choice)

Konrad Herbst, Elena J. Forchielli, Taiyao Wang, **Meghan Thommes**, Ioannis Ch. Paschalidis, Daniel Segrè. Multi-Attribute Subset Selection enables prediction of representative phenotypes across microbial populations. *Communications Biology* (2024). doi: [10.1038/s42003-024-06093-w](https://doi.org/10.1038/s42003-024-06093-w)

**Meghan Thommes**, Taiyao Wang, Qi Zhao, Joshua Goldford, Ioannis Ch. Paschalidis, Daniel Segrè. Microbes distributing metabolism: How cross-feeding interactions support metabolic division of labor. 2018 Boston University Microbiome Day. Boston, MA. February 14, 2018. ([Won Best Poster Presentation](#))

**Meghan Thommes\***, Taiyao Wang, Qi Zhao, Joshua Goldford, Ioannis Ch. Paschalidis, Daniel Segrè. Computational design of metabolic division of labor for synthetic microbial communities. 2017 Metabolic Pathway Analysis Conference. Boseman, MT. July 24-28, 2017. Abstract Book, page 83. ([Won Best Poster Presentation](#))