

Episimlab: a Python package for modeling epidemics

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Summary

Computational models play a crucial role in our scientific understanding of, and preparedness for, infectious disease epidemics. These *in silico* disease models simulate real-world transmission dynamics, and are thereby well-suited for such tasks as early detection of novel pandemics, improving situational awareness during periods of high prevalence, and estimating efficacy of intervention strategies. These modeling approaches have proven valuable in responding to emergent epidemics such as the H1N1 flu pandemic, the Ebola epidemic, the Zika virus pandemic, and the recent COVID-19 pandemic (ref). For example, during the early waves of the COVID-19 pandemic, compartmental disease models were instrumental for projecting case counts and hospitalizations (K. Pierce et al., 2020; K. A. Pierce et al., 2020). As more data on case incidence, hospitalizations, and viral genomics became available, disease modelers are able to simulate increasingly complex disease dynamics. Developing such complex compartmental models is time-consuming, however, and few implementations share a common software framework or application flow. Therefore, there is a urgent need for more robust cyberinfrastructure in the field of epidemic modeling.

Episimlab is a software development kit (SDK) written in Python that seeks to address this problem by providing a flexible framework for developing compartmental disease models. Models in Episimlab are collections of modular components, known as processes, which can be added, removed, or replaced to modify the dynamics of the simulated disease. In practice, this means that Episimlab supports development of models that:

- 1. Have many parameters, often with multiple, varying dimensions
- 2. Use any compartment structure that can be represented as a graph
- 3. Simulate interventions dynamically, such as administering vaccines only when case incidence exceeds a threshold.

The package is designed to be approachable; it includes several pre-packaged models that the user can run in a few lines of code. When the user chooses to add data streams or more complex transmission dynamics, they can easily do so by adding or replacing processes in the model. In addition, Episimlab provides a scalable and performant runtime environment for model execution, thanks to integration with packages in the PyData stack such as Dask (Dask Development Team, 2016), Xarray (Hoyer & Hamman, 2017), and xarray-simlab (Bovy et al., 2021). Finally, Episimlab provides a standard for packaging, versioning, and sharing models, using Python's built-in class attributes.

Statement of Need

Episimlab is a Python package that provides a common framework for rapid development of complex compartmental disease models. It specifically supports models of epidemics that:

- 1. Are Markovian
- 2. Are compartmental
- 3. Run in discrete time
- 4. Are either continuous or stochastic (Gillespie dynamics, see Gillespie (1977))

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Software

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Episimlab provides sufficient boilerplate such that the user can quickly instantiate a basic compartmental model. Simple models such as the SIR model are not unique to Episimlab; they have a long history of use in disease modeling since their origin in the early 20th century (Kermack & McKendrick, 1927; Ross, 1916; Ross & Hudson, 1917a, 1917b). More recently, various implementations of compartmental models have been made publicly available and easily usable as open-source software packages (refs). These packages are valuable because they simplify execution of many simple compartmental models, but their usage is limited to the discrete handful of model structures that are published with the package. In addition, such projects rarely support complex models with more than 5 or 6 compartments, presumably because complex models are difficult to develop and reproduce.

While inspired by these previous works, Episimlab aims to support development of models with arbitrary complexity. It gives the user flexibility to define key components of their compartmental model, such as the dimensionality of the Markov state matrix, the number of compartments, and custom stochastic behavior. Of note, the comparable Python package epydemic (Simon Dobson, 2021) also supports a generic model of compartmental disease. It does not, however, support arbitrary dimensionality in input variables or in the Markov state matrix.

Episimlab was designed with epidemiological use cases in mind, via collaboration with data scientists and epidemiologists in the UT Austin COVID-19 Modeling Consortium. Specifically, prototypes of Episimlab were used in studies forecasting hospital burden due to the COVID-19 epidemic in Austin, Texas (K. Pierce et al., 2020; K. A. Pierce et al., 2020). Although the package was originally intended for use by epidemiologists, Episimlab is useful for anyone developing Markovian models of disease spread. The package is useful for students because it provides a minimal, approachable boilerplate for developing basic models in pure Python. It introduces and reinforces best practices in object-oriented software development, such as modularity and reproducibility. For disease modeling experts, Episimlab provides a platform that supports a wide variety of modeling use cases. Simple models can be easily adapted into more complex ones, encouraging a model development approach that is rapid, iterative, and organic. Under the hood, Episimlab leverages concurrency in xarray-simlab, dataset chunking in Dask, and accelerated matrix math in xarray, so models are performant even when using large datasets.

Unused Text

Unlike past compartmental modeling software, models in Episimlab are modular. Episim lab models are essentially unique sets of discrete components. These components, formally known as processes in the API, can be modified, exchanged, reused, and shared between models. Commonly used processes, such as calculating the force of infection (FOI), ship with Episimlab, and can be modified using class inheritance in Python.

Dependencies

xarray xarray-simlab dask networkx matplotlib

Related Packages

epydemic

epydemic is a Python package that provides a common framework for building models of epidemic processes (Simon Dobson, 2021). It supports simulations that are discrete-time synchronous or continuous-time stochastic (Gillespie). Like Episimlab, it supports a generic model for compartmental disease, allowing for flexibitlity in the compartmental model structure. In addition, it ships with several basic compartmental models such as SIR, SIS, and SEIR.



EoN (Epidemics on Networks)

Epidemics on Networks (EoN) is a Python package that simulates disease dynamics for SIR and SIS models (Miller & Ting, 2019). The package includes numerical solutions for 20 different differential equation models, and supports complex contagions using the Gillespie algorithm (Gillespie, 1977).

Eir

Eir is a Python package that simulates epidemics using compartmental models. It includes 4 distinct models with different mobility dynamics (Jacob, 2021). In addition, it provides utilities for inspecting transmission chains, analyzing state histories, and visualizing simulation results.

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References

- Bovy, B., McBain, G. D., Gailleton, B., & Lange, R. (2021). *Benbovy/xarray-simlab: 0.5.0* (Version 0.5.0) [Computer software]. Zenodo. https://doi.org/10.5281/zenodo.4469813
- Dask Development Team. (2016). Dask: Library for dynamic task scheduling. https://dask.org
- Gillespie, D. T. (1977). Exact stochastic simulation of coupled chemical reactions. *The Journal of Physical Chemistry*, 81(25), 2340–2361. https://doi.org/10.1021/j100540a008
- Hoyer, S., & Hamman, J. (2017). Xarray: N-D labeled arrays and datasets in Python. *Journal of Open Research Software*, 5(1). https://doi.org/10.5334/jors.148
- Jacob, M. (2021). Eir: A python package for epidemic simulation. *Journal of Open Source Software*, 6(62), 3247. https://doi.org/10.21105/joss.03247
- Kermack, W. O., & McKendrick, A. G. (1927). A contribution to the mathematical theory of epidemics. Proceedings of the Royal Society of London. Series A, Containing Papers of a Mathematical and Physical Character, 115(772), 700–721. https://doi.org/10.1098/rspa. 1927.0118
- Miller, J. C., & Ting, T. (2019). EoN (epidemics on networks): A fast, flexible python package for simulation, analytic approximation, and analysis of epidemics on networks. *Journal of Open Source Software*, 4(44), 1731. https://doi.org/10.21105/joss.01731
- Pierce, K. A., Ho, E., Wang, X., Pasco, R., Du, Z., Zynda, G., Song, J., Wells, G., Fox, S. J., & Meyers, L. A. (2020). Early COVID-19 pandemic modeling: Three compartmental model case studies from texas, USA. *Computing in Science & Engineering*, 23(1), 25–34. https://doi.org/10.1109/MCSE.2020.3037033
- Pierce, K., Ho, E., Wang, X., Pasco, R., Du, Z., Fox, S., Zynda, G., Song, J., & Meyers, L. A. (2020). COVID-19 healthcare demand projections: Beaumont-port arthur MSA texas. *UT COVID-19 Model. Consortium.* https://doi.org/10.13140/RG.2.2.14066.53443
- Ross, R. (1916). An application of the theory of probabilities to the study of a priori pathometry. & #x2014; part i. Proceedings of the Royal Society of London. Series A, Con-



- taining Papers of a Mathematical and Physical Character, 92(638), 204–230. https://doi.org/10.1098/rspa.1916.0007
- Ross, R., & Hudson, H. P. (1917a). An application of the theory of probabilities to the study of a priori pathometry.—part II. *Proceedings of the Royal Society of London. Series A, Containing Papers of a Mathematical and Physical Character*, *93*(650), 212–225. https://doi.org/10.1098/rspa.1917.0014
- Ross, R., & Hudson, H. P. (1917b). An application of the theory of probabilities to the study of a priori pathometry. \$\precept \times 2014; part III. Proceedings of the Royal Society of London. Series A, Containing Papers of a Mathematical and Physical Character, 93(650), 225–240. https://doi.org/10.1098/rspa.1917.0015
- Simon Dobson. (2021). *Epydemic* (Version 1.7.2) [Computer software]. https://pyepydemic.readthedocs.io/