Pathogen.jl: Infectious Disease Transmission Network Modeling with Julia

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Abstract

We introduce Pathogen.jl for simulation and inference of transmission network individual level models (TN-ILMs) of infectious disease spread in continuous time. TN-ILMs can be used to jointly infer transmission networks, event times, and model parameters within a Bayesian framework via Markov chain Monte Carlo (MCMC). We detail our specific strategies for conducting MCMC for TN-ILMs, and our implementation of these strategies in the Julia package, Pathogen.jl, which leverages key features of the Julia language. We provide an example using Pathogen.jl to simulate an epidemic following a susceptible-infectious-removed (SIR) TN-ILM, and then perform inference using observations that were generated from that epidemic. We also demonstrate the functionality of Pathogen.jl with an application of TN-ILMs to data from a measles outbreak that occurred in Hagelloch, Germany, in 1861 (Pfeilsticker 1863; Oesterle 1992).

Keywords: epidemic, individual level models, transmission networks, Bayesian, Julia.

1. Introduction

The availability of data to support the use of highly detailed epidemic models is an increasing reality. Individual level models (ILMs) are a framework that accommodates individual specific risk factor information to describe infectious disease dynamics (Deardon, Brooks, Grenfell, Keeling, Tildesley, Savill, Shaw, and Woolhouse 2010). In accounting for population heterogeneity with ILMs, disease dynamics can be captured with more realistic models, and in turn, control strategies can be evaluated using these models. As the number of individuals and the associated data for these individuals increase, the computational requirements for simulation and inference of ILMs are magnified. With epidemic modeling in support of decision making in ongoing epidemics, time, whether for development or computation, is at a premium. Time constraints aside, the epidemiologists that utilize these models may not have the training to support development of a high performance implementation of a specialized ILM in a low level language.
Julia is a high level, high performance language that has been motivated by the needs of modern scientific computing (Bezanson, Chen, Chung, Karpinski, Shah, Vitek, and Zoubiritzky 2018). Previously, the use of lower level compiled languages, such as Fortran or C++, has typically been required to implement high performance features of packages in high level scientific computing languages (Bezanson et al. 2018), such as R (R Core Team 2022). This two language problem is insidious, and generates an unnecessary divide between those that can use software, and those that can review or advance its development. We use Julia to avoid this problem. As a high level and high performance language, Julia is well situated to address some of the ongoing challenges and opportunities in developing and deploying specialized ILMs.

To date, there have been no packages specific for epidemic modeling published on the Julia General Registry (https://github.com/JuliaRegistries/General). There are Julia packages, however, that could be used for this purpose. DifferentialEquations.jl is a comprehensive, general purpose package for differential equations (Rackauckas and Nie 2017b), and is well suited for population level epidemic modeling through its discrete stochastic differential equation functionality (Rackauckas and Nie 2017a). Use of these kinds of models can be complementary to the use of ILMs (Webb 2017). DifferentialEquations.jl can simulate from population level epidemic models, and provides several methods of Bayesian parameter estimation through DiffEqBayes.jl. The BioSimulator.jl package provides simulation methods for interacting populations (Landeros, Stutz, Keys, Alekseyenko, Sinsheimer, Lange, and Sehl 2018), and could also be used for population level epidemic models. While there are multiple simulation algorithms and full flexibility into the interaction network of the simulated entities in BioSimulator.jl, there are no inference methods provided.

Modeling of heterogeneous populations can be done with Agents.jl, which provides a general purpose package for stochastic simulation of agents on grid systems in discrete time (Vahdati 2019). Agent based models (ABMs) can incorporate agent movements and complex behaviors into the epidemics they generate. They prove to be powerful tools in the consideration of individual decision making, especially in the context of learning and behavioral shifts by individuals during an epidemic, and relating those individual actions to higher level disease dynamics, such as seen in Abdulkareem, Augustijn, Filatova, Musial, and Mustafa (2020). Such dynamics are not readily modeled by other methods, but ABMs are also limited in their ability to be fit with traditional inference procedures. Approximate Bayesian computation (ABC) is one avenue for ABM parameterization, where prior beliefs on realistic parameter values are refined using measures of similarity between flexibly summarized ABM simulations and experimental or observational data (Ross, Baker, Parker, Ford, Mort, and Yates 2017). Which is to say, ABMs, like ILMs, and stochastic differential equation models all have their place in epidemics research.

While not registered on the Julia General Registry at this time, Shoukat, Wells, Langley, Singer, Galvani, and Moghadas (2020) use Julia in the construction of an agent based model specific to the application of predicting hospital and ICU capacity in Canada during the COVID-19 pandemic. Using a population of 10000 agents, they perform simulations to assess different scenarios in regards to resulting proportions of the population requiring care in a hospital or ICU.

Outside of Julia, there are several packages for epidemic modeling on R’s Comprehensive R Archive Network (CRAN) at https://CRAN.R-project.org/. Amongst these epidemic modeling packages, there are two for working with ILMs specifically: EpiILM (Warriyar K. V., Almutiry, and Deardon 2020) and EpiILMCT (Almutiri, Deardon, and Warriyar K. V. 2021).
EpiILMCT is a package for network and spatial continuous time ILMs. This package offers a high performance low level implementation in Fortran for performing MCMC for continuous time ILMs. The functions that describe disease state transition rates in this package are limited in form, and may only contain a coefficient and a power-parameter for each risk factor. This package offers data augmentation for epidemics in which event times are assumed to be unknown. EpiILM, on the other hand, supports network and spatial ILMs in discrete time, and does not have data augmentation functionality. EpiILM similarly has its core functionality programmed in Fortran for performance reasons.

Beyond the R epidemic modeling packages that utilize the ILM framework of Deardon et al. (2010), there is also package epinet which provides tools for modeling epidemics that spread over contact networks described by exponential-family random graph models (Groendyke and Welch 2018). This capable package can perform simulations and inference with optional event time augmentation. Its core functionality is implemented in C. Another package is EpiModel, which provides functionality to build and simulate individual based epidemic models over stochastic and dynamic contact networks (Jenness, Goodreau, and Morris 2018). The R package surveillance is also notable for its implementation of several spatio-temporal epidemic models, including individual level epidemic models (Meyer, Held, and Höhle 2017). However, the individual level models in surveillance do not impute transmission networks or event times.

The remainder of the paper is organized as follows. In Section 2, we describe ILMs (Deardon et al. 2010), and then an extension of ILMs, the transmission network individual level model (TN-ILM). Section 3 presents methods for simulating epidemics and fitting TN-ILMs. In Section 4, we introduce and present in detail our high performance package, Pathogen.jl, for implementing the simulation, model fitting, and inference of TN-ILMs with data augmentation capability in continuous time. Section 5 demonstrates how to use Pathogen.jl to simulate epidemics and how to fit TN-ILMs via MCMC to simulated data sets through an example. An application to the measles epidemic in Hagelloch, Germany in 1861 (Pfeilsticker 1863; Oesterle 1992) is then provided in Section 6. The paper concludes in Section 7 with a brief discussion of future work.

### 2. Individual level models

In an ILM, each individual is considered to be in one of several disease states at any time, and the rates governing their transition through the disease states are a function of both individual specific risk factors and the disease states of other individuals in the population at that time. As a whole, the disease state transitions, and subsequent disease state transition rate updates constitute a time-heterogeneous Poisson process.

ILMs may be continuous or discrete with respect to time. In continuous time ILMs, we think of the temporal data structure as consisting of variable time periods, and these representing the length of time between events; i.e., inter-event periods. Here, events are considered to be the disease state transitions in the population. A discrete time ILM approximates its continuous time counterpart, and generally uses time periods that are equal in duration. The likelihoods and strategies for inference for these models are different, however the disease state transition rates each uses can be defined in the same way.

There is flexibility in the disease states that are considered, with susceptible and infectious states being the only necessary states. In the following, the structure of a more complex framework, the susceptible-exposed-infectious-removed (SEIR) ILM, will be described.
In an $\mathcal{SEIR}$ ILM, the rate that an individual, $i$, transitions from the susceptible state to the exposed state during the $t$-th time period is given as:

$$\lambda_{SE}(i, t) = \left[ \Omega_S(i) \sum_{k \in \mathcal{I}(t)} \Omega_T(k) \kappa(i, k) \right] + \epsilon(i) \quad \text{for } i \in \mathcal{S}(t),$$

where,

- $\mathcal{I}(t)$ is the set of infectious individuals during the $t$-th time period,
- $\mathcal{S}(t)$ is the set of susceptible individuals during the $t$-th time period,
- $\Omega_S(i)$ is a function of risk factors associated with the risk of susceptible individual $i$ contracting the disease (susceptibility),
- $\Omega_T(k)$ is a function of risk factors associated with the risk of infection transmission from the $k$-th individual (transmissibility),
- $\kappa(i, k)$ is an infection kernel, a function of risk factors involving both the $i$-th and $k$-th individuals, which often describes the connectivity between these individuals, and,
- $\epsilon(i)$ is a function of risk factors associated with exposure to the $i$-th individual that the model otherwise fails to explain. Typically this refers to exposure from a non-specified source outside of the observed population. This is also referred to as the *sparks function*.

Transition between exposed and infectious states for the $j$-th individual during the $t$-th time period occurs with rate:

$$\lambda_{EI}(j, t) = \Omega_L(j) \quad \text{for } j \in \mathcal{E}(t)$$

where,

- $\mathcal{E}(t)$ is the set of exposed individuals during the $t$-th time period, and
- $\Omega_L(j)$ is a function of risk factors associated with the latent period, the length of time between exposure and onset of infectiousness, in the $j$-th individual.

Lastly, the $k$-th individual transitions from the infectious to the removed state during the $t$-th time period with rate:

$$\lambda_{IR}(k, t) = \Omega_R(k) \quad \text{for } k \in \mathcal{I}(t)$$

where,

- $\Omega_R(k)$ is a function of risk factors associated with the removal of the $k$-th individual from an infectious state. Removal can refer to recovery with acquired immunity, death, quarantine, etc.
In combination, the rates $\lambda_{SE}(i, t)$, $\lambda_{EI}(j, t)$, and $\lambda_{IR}(k, t)$ describe the spread of disease in an SEIR ILM.

In a continuous time SEIR ILM, no change occurs to the sets $S(t)$, $E(t)$, $I(t)$, and $R(t)$ during an inter-event period. It follows that the rates $\lambda_{SE}(i, t)$, $\lambda_{EI}(j, t)$, and $\lambda_{IR}(k, t)$ remain constant during each of these time periods, and that the occurrence of any single event (disease state transition) marks the end of the $t$-th inter-event time period. Movement of an individual into or out of a modeled population may be represented through a deterministic change of state membership of that individual to or from $R(t)$, followed by the appropriate rate updates. For the remainder of this paper we focus exclusively on continuous time ILMs.

2.1. Transmission network ILM extension

In ILMs, a susceptible individual’s risk of being infected by an infectious disease is based on the culmination of various risk factors, such that the influence of specific sources of exposure are masked. We introduce transmission network ILMs (TN-ILMs), that instead are explicit with respect to exposure sources. In an SEIR TN-ILM, a set of competing transition rates for each susceptible individual, $i$, to the exposed state are defined as

$$
\lambda^*_SE(i, k, t) = \Omega_S(i)\Omega_T(k)\kappa(i, k)
$$

for $i \in S(t)$, $k \in I(t)$

describing transition rates specific to each infectious individual, $k$, and with,

$$
\lambda^*_SE(i, t) = \epsilon^*(i)
$$

describing the transition rate specific to any exogenous exposure source. While $\epsilon(i)$ in an ILM is not necessarily specific to exogenous sources, that assumption is made with $\epsilon^*(i)$ in TN-ILMs.

3. Methods

This section starts with a description of simulation methods for continuous time ILMs and TN-ILMs in Section 3.1. In Section 3.2, the TN-ILM likelihood function is presented, again drawing comparisons to continuous time ILMs. Lastly, in Section 3.3, an approach is detailed for Bayesian inference of TN-ILMs via Markov chain Monte Carlo (MCMC).

3.1. Continuous time-to-event simulation

As a Poisson process, an epidemic following a continuous time ILM can be stochastically simulated using the Gillespie (1977) algorithm. Here, the inter-event period is the minimum of competing exponential random variables, with each of these exponential random variables representing a possible event given the disease status of the population. The minimum, and the inter-event time, is generated directly from an exponential distribution with rate

$$
v(t) = \sum_{i \in S(t)} \lambda_{SE}(i, t) + \sum_{j \in E(t)} \lambda_{EI}(j, t) + \sum_{k \in I(t)} \lambda_{IR}(k, t).
$$

The specific event that occurs at this time is generated from a multinomial distribution with probability vector

$$
\pi_1(t) = \left[ \frac{\lambda_{SE}(1, t)}{v(t)}, \ldots, \frac{\lambda_{SE}(N, t)}{v(t)}, \frac{\lambda_{EI}(1, t)}{v(t)}, \ldots, \frac{\lambda_{EI}(N, t)}{v(t)}, \frac{\lambda_{IR}(1, t)}{v(t)}, \ldots, \frac{\lambda_{IR}(N, t)}{v(t)} \right]^T,
$$
for a population of size \( N \). Which is to say, the probability of a specific event being the first event to occur amongst competing events is proportional to its contribution to \( v(t) \). If the generated event is the transition of a susceptible individual to the exposed state, and a TN-ILM is used, the transmission source is then generated from a multinomial distribution, with probability vector

\[
\pi_2(t) = \left[ \frac{\lambda_{SE}(i,1,t)}{\sum_j \lambda_{SE}(j,t)}, \ldots, \frac{\lambda_{SE}(i,N,t)}{\sum_j \lambda_{SE}(j,t)}, \ldots, \frac{\lambda_{SE}(i,t)}{\sum_j \lambda_{SE}(j,t)} \right]^T, \tag{2}
\]

following the same logic. With the generation of an inter-event interval, and generation of the specific event occurrence, the population is then updated (i.e., membership in the sets of \( S_{(t+1)}, E_{(t+1)}, I_{(t+1)}, \) and \( R_{(t+1)} \)), and the process repeats, until either no further events are possible (i.e., \( \pi_1(t) \) is a vector of zeros), or until some earlier stop condition is met.

### 3.2. Continuous time ILM likelihood

The likelihood of the continuous time-to-event \( S\bar{E}IR \) ILM is the product of likelihoods at time periods indexed by \( t = 1, \ldots, T - 1 \), where \( T \) is the total number of events that have occurred. With the length of time since the beginning of the \( t \)-th time period denoted as \( \Delta_t \), the likelihood function for the associated parameters, \( \theta \), is given as:

\[
L(\theta) = \prod_{t=1}^{T-1} \psi(t) v(t) \exp \left\{ -v(t) \Delta_t \right\},
\]

where

\[
\psi(t) = \begin{cases} 
\frac{\lambda_{SE}(i,t)}{v(t)} & \text{if } i \in (S_t \cap E_{(t+1)}), \text{ i.e., } i \text{ transitioned from susceptible to exposed}, \\
\frac{\lambda_{EI}(j,t)}{v(t)} & \text{if } j \in (E_t \cap I_{(t+1)}), \\
\frac{\lambda_{IR}(k,t)}{v(t)} & \text{if } k \in (I_t \cap R_{(t+1)}), 
\end{cases}
\]

and,

\[
N = |S_t| + |E_t| + |I_t| + |R_t| \quad \forall t.
\]

Modification is required for TN-ILMs to account for specific transmissions sources. With the TN-ILM, the likelihood function is given as:

\[
L(\theta) = \prod_{t=1}^{T-1} \psi^*(t) v(t) \exp \left\{ -v(t) \Delta_t \right\},
\]

where

\[
\psi^*(t) = \begin{cases} 
\frac{\lambda_{SE}^*(i,k,t)}{v(t)} & \text{if } i \in (S_t \cap E_{(t+1)}) \text{ by endogenous exposure from individual } k, \\
\frac{\lambda_{SE}^*(i,k,t)}{v(t)} & \text{if } i \in (S_t \cap E_{(t+1)}) \text{ by exogenous exposure}, \\
\frac{\lambda_{EI}(j,t)}{v(t)} & \text{if } j \in (E_t \cap I_{(t+1)}), \\
\frac{\lambda_{IR}(k,t)}{v(t)} & \text{if } k \in (I_t \cap R_{(t+1)}).
\end{cases}
\]

This likelihood mirrors the simulation process described in Section 3.1. That is, inter-event periods follow exponential distributions, which are paired with multinomial distributions for
the occurrence of specific events. The product of the likelihoods of these distributions compose the likelihood of the epidemic as a whole.

3.3. Bayesian inference via Markov chain Monte Carlo

In the Bayesian framework, beliefs about a parameter are described by a posterior distribution. The posterior distribution is a probability distribution for model parameter values, $\theta$, conditioned on observational data $D$. From a posterior distribution, credible intervals and point estimates for parameters can be obtained, and hypothesis testing can be conducted. Monte Carlo methods, such as Markov chain Monte Carlo (MCMC) methods, are typically used to generate a sufficient quantity of samples from the posterior distribution, such that the properties of the posterior distribution can be estimated through the generated samples. The Metropolis-Hastings (M-H) MCMC algorithm is used to perform Bayesian parameter estimation for continuous time ILMs. The M-H algorithm generates a Markov chain consisting of a sequence of samples within the parameter space, the distribution of which converges to the target distribution – the posterior distribution of the parameters in this case. To generate such a Markov chain, some initial values, here denoted as $\theta_1$, must be first selected or generated. Following initialization of the Markov chain, new samples are proposed with a transition kernel. For a symmetric transition kernel, a sample proposed for the $w$-th iteration, $\theta_w'$, is accepted with probability

$$
\alpha(\theta_w') = P(\theta_w = \theta_w') = \min \left(1, \frac{L(\theta_w')P(\theta_w')}{L(\theta_w-1)P(\theta_w-1)} \right).
$$

If the proposal $\theta_w'$ is rejected, $\theta_w$ remains at $\theta_w-1$ (Hastings 1970; Robert and Casella 2013).

Sampling from the posterior distribution of continuous time ILMs using the M-H algorithm is straightforward when an epidemic is fully observed; i.e., with event times that are known exactly with certainty. While for certain epidemics complete or nearly complete case identification may occur, exact event times are rarely known. Here, unknown event times are imputed using a data augmentation process, in which they are treated as additional parameters to be estimated. This tends to result in a massive increase in the dimensionality of the parameter space, and with that computational challenges arise. With TN-ILMs these challenges are exacerbated by the transmission network, which is latent and requires imputation. A compatibility must be enforced in the generation of MCMC proposals between the transmission network and event times. Without this incorporated into the proposal mechanism, the rejection rate using M-H would be unreasonably high (i.e., we must avoid transmission network proposals that are impossible given the order of infection of individuals, as dictated by the event time set). As such, a specialized MCMC algorithm is required for inference of TN-ILMs.

**Initialization strategy**

Initialization is an important step to facilitate efficient MCMC in these models. High dimensionality and event time interdependence results in vast areas of the parameter space having near-zero posterior density, which is approximated to zero in computation. With MCMC, it may take a long time to move into and sample from areas of the parameter space that have non-zero computationally approximated posterior mass. This also impacts adaptive tuning of transition kernel variance and further contributes to bottlenecks in MCMC convergence.
Our strategy is to generate many potential sets of initial values, and to select the set with the highest posterior density. Multiple Markov chains, and a higher number of initialization generations per Markov chain are recommended for higher dimensional applications of TN-ILMs, as is common when implementing MCMC in general (Robert and Casella 2013). These independently initialized Markov chains allow for a more substantiated assessment of Markov chain convergence, in comparison to assessing the behavior of a single Markov chain in isolation.

The initialization of each Markov chain begins with sampling a set of model parameters from their corresponding prior distributions. A set of event times is also generated for observation delays, as well as latent periods if applicable. Since the transmission network can be generated directly from its conditional distribution with TN-ILMs, we marginalize the TN-ILM likelihood over all possible transmission networks, simplifying it to the standard ILM likelihood for the purposes of Markov chain initialization. The ILM likelihood is calculated by iterating through each event in the generated set of event times. While this calculation is computationally intensive, conveniently, it can be stopped early when its running result drops below some threshold. This convenience is leveraged in our initialization generation process such that computational cost does not increase linearly with an increase in the number of initialization attempts.

**Iteration strategy**

Each MCMC iteration can be broken down into several sub-steps: event time sampling, parameter sampling, and transmission network sampling.

**Event times.** A random walk M-H sampling procedure is used to sample event times from the TN-ILM posterior distribution. Specifically a bounded normal distribution is used as the transition kernel, with its mean set to the event time in the previous iteration, and with a pre-specified variance. These bounds are determined by the times of the events that are dependent through the transmission network or model structure.

The event times relevant to determining the bounds of event time proposal distributions include: the times of other state transitions by the involved individual (e.g., an individual may not transition to a removed state prior to being in an infected state); the infection and, if applicable, removal times of their transmission source (e.g., a susceptible individual may only transition to the exposed state while their transmission source is in an infected state). Similarly, if an individual transmits the disease themselves, they must be in the infected state prior to their earliest transmission, and must not be removed until after their final transmission. This ensures the transmission network remains compatible with the set of event times.

Due to event time interdependence, event time proposals are generated one at a time. The order in which event times are updated is randomized at each iteration. A decision to accept or reject proposed event times can be conducted in batches such that fewer likelihood calculations occur. Successful tuning of the event time batch size and transition kernel variance results in effective sampling of the event time posterior distribution while minimizing computational costs. To achieve this, experimental tuning is required for each TN-ILM application by the user.
**TN-ILM parameters.** Following updates to the event times, a new set of TN-ILM parameters is proposed, and the proposed values are subject to the acceptance rule for M-H MCMC algorithms shown in Equation 3. For these proposals, a multivariate normal transition kernel is used. The covariance matrix of the transition kernel can be automatically tuned during sampling, using the adaptive sampling method of Roberts and Rosenthal (2007).

**Transmission network.** Finally, the transmission network is updated using the Gibbs sampler. Each transmission source is sampled from a multinomial distribution corresponding to its conditional distribution. The probability vector of each multinomial distribution is generated following Equation 2, for each applicable time period using the current set of event times and model parameters.

## 4. Software implementation with Julia

We have implemented simulation and inference methods for TN-ILMs as described in Section 3 in Julia with the Pathogen.jl package. Pathogen.jl is open source and released with an MIT (Expat) license. Our description of Pathogen.jl is consistent with its v0.4.12 release.

Package Pathogen.jl leverages several features of Julia. One of the more impactful features of Pathogen.jl is the ability for the user to define functions for disease state transition rates with a high level of flexibility, without inherently sacrificing performance. This high flexibility-high performance quality of Pathogen.jl is owed to Julia code being just-in-time (JIT) compiled to highly optimized machine code. While Julia is a compiled language, it is still interactive and can be used with a read-evaluate-process-loop (REPL) interface that is familiar to those working with other high level interactive scientific computing languages. In a set of benchmarks performed by Bezanson, Edelman, Karpinski, and Shah (2017), Julia was typically within a factor of two in regards to computation time of equivalent C code.

In Julia, new user-defined types are implemented without performance penalty in comparison to the basic types provided by the language. This is made no clearer than by the fact that much of Julia’s features, are written in Julia itself. Julia’s *multiple dispatch* enables both generalist and specialist function methods that are invoked based on argument types. The type system results in code that can often be common across a set of types. Pathogen.jl provides types to support \textit{SEIR}, \textit{SEI}, \textit{SIR}, and \textit{SI} TN-ILMs. With only some basic methods defined for these types or across unions of these types, higher level code such as that involved in epidemic simulation, likelihood calculations, and performing MCMC is able to be kept common across all disease model classes. This same functionality in theory would reduce the development time required to implement additional disease model classes. The types implemented by Pathogen.jl, and their hierarchy are shown in Figure 1.

Distributed computing functionality is highly relevant, if not essential for large data applications. For TN-ILMs, this would be for modeling large populations and/or higher complexity models. Julia has been designed for simple, but powerful distributed computing. Some of this is leveraged in Pathogen.jl for conducting MCMC for TN-ILMs. Independent Markov chains are easily initialized, and ran across multiple cores, or on a high performance cluster in parallel.
Figure 1: The hierarchy of the types provided by Pathogen.jl are shown above. Type names followed by \{T\} indicate that that type is parametric – in all cases the type is parameterized by a subtype of ‘EpidemicModel’. Subscripts are used to indicate the kind of type: \(p\)primitive: elementary data representations; \(a\)abstract: types used for organization purposes or to invoke specific function methods but do not contain data; \(s\)struct: types that have fields containing other types.
4.1. Installing Pathogen.jl

Pathogen.jl v0.4.12 is available for Julia version 1.1 and higher, and is listed on Julia’s General Registry (https://github.com/JuliaRegistries/General). Julia has a built-in package manager, Pkg. Pkg provides a special REPL that is entered from the Julia REPL by typing a single ] . From the package REPL, Pathogen.jl v0.4.12 is installed with add Pathogen@v0.4.12. Users also have the option to install Plots.jl to make use of custom plotting tools provided by Pathogen.jl. This can be done with add Plots from the package REPL. Once installed, Pathogen.jl can be immediately used with using Pathogen from back in the Julia REPL. This entire process takes only a few seconds.

4.2. Pathogen.jl basics

Whether using Pathogen.jl for simulation or inference of TN-ILM epidemics, a population must be first defined. We represent populations in our package with a ‘Population’ type. More specifically, our ‘Population’ type is a struct. In Julia, a struct is a type that is composed of other types – perhaps other structs, or primitive types such as Float64 or Bool. To construct a ‘Population’, a ‘DataFrame’ containing individual specific risk factors must be specified. ‘DataFrame’ is a type provided by the DataFrames.jl package, which is a dependency of Pathogen.jl. In this ‘DataFrame’, each row will represent an individual. There is also an option to provide a distance matrix when describing a population. Distance measures are common components of infectivity kernels. The distance matrix can be used to avoid repeated calculation of these distances from individual specific risk factor information. ‘Population’ in Pathogen.jl is declared as

```julia
struct Population
    risks::DataFrame
    distances::Union{Nothing, AbstractArray}
    individuals::Int64
end
```

The structure of TN-ILMs is described with ‘RiskFunctions{T}’, which is a collection of functions that calculate individual specific disease state transition rates. ‘RiskFunctions{T}’ is a parametric struct, declared as shown below, with slightly different construction and behavior for different values of T.

```julia
struct RiskFunctions{T<:EpidemicModel}
    sparks::Union{Nothing, Function}
    susceptibility::Union{Nothing, Function}
    infectivity::Union{Nothing, Function}
    transmissibility::Union{Nothing, Function}
    latency::Union{Nothing, Function}
    removal::Union{Nothing, Function}
end
```

Parametric structs are used throughout Pathogen.jl to provide specialization to different disease model classes. This allows for modified functionality where it is needed for the various model class implementations in otherwise common code. For instance, construction of
‘RiskFunctions{SIR}’ when compared to ‘RiskFunctions{SEIR}’ does not involve specification of a function describing the transition rate between exposed and infectious classes (i.e., $\Omega_L$ in Equation 1). There is full flexibility in the form of TN-ILM risk functions used in ‘RiskFunctions{T}’, as long as these risk functions follow an expected signature for their arguments. Each risk function must accept a ‘Population’, a parameter vector (Vector{Float64}), and an Int64 individual identifier, as arguments. Infectivity kernels are an exception to this, and must accept two Int64 identifiers – for infection source and newly infected individual. Each risk function should return a Float64. A complete example of constructing ‘RiskFunctions{SIR}’, including the construction of the risk functions that compose it, is provided in Section 5.

The parameterization of TN-ILM risk functions is represented by ‘RiskParameters{T}’, a separate type. For each of the risk functions required by a TN-ILM, a parameter vector must be provided in ‘RiskParameters{T}’, which has been declared as:

```julia
struct RiskParameters{T<: EpidemicModel} <: AbstractRisk{T}
    sparks::Union{Nothing, AbstractVector}
    susceptibility::Union{Nothing, AbstractVector}
    infectivity::Union{Nothing, AbstractVector}
    transmissibility::Union{Nothing, AbstractVector}
    latency::Union{Nothing, AbstractVector}
    removal::Union{Nothing, AbstractVector}
end
```

### 4.3. Simulation with Pathogen.jl

To simplify the simulation interface, we have provided with Pathogen.jl a ‘Simulation{T}’ struct. ‘Simulation{T}’ objects include all of the information required to iterate and track the progression of an epidemic. Once constructed, the `simulate!` function is used to run the simulation, updating event times, individual disease states, and the transmission network within the ‘Simulation{T}’ object as appropriate. The `simulate!` function will iterate until a specified stop condition is met (processing time, simulation time, and/or number of iterations) or if there are no further events possible. ‘Simulation{T}’ is a parametric mutable struct, which is like a parametric struct, except it allows for its values to be changed or updated. For instance, the Int64 value for `iterations`, shown in the type declaration below, is incremented at each iteration, which is possible because of this mutability. Explicitness about struct mutability in Julia allows for certain optimizations by the compiler when dealing with structs that do not change in composition.

```julia
mutable struct Simulation{T <: EpidemicModel}
    time::Float64
    iterations::Int64
    population::Population
    risk_functions::RiskFunctions{T}
    risk_parameters::RiskParameters{T}
    disease_states::Vector{DiseaseState}
    transmission_rates::TransmissionRates
    event_rates::EventRates{T}
end
```
events::Events{T}
transmission_network::TransmissionNetwork
end

The use of \{T\} throughout the declaration of ‘Simulation\{T\}’ ensures matching type parameterizations for all of the types that compose it. This also means that the parameterization of ‘Simulation\{T\}’ can be inferred during its construction.

There are several ways to construct a ‘Simulation\{T\}’ object. The simplest construction method requires only specification of the following for the TN-ILM: a ‘Population’, ‘RiskFunctions\{T\}’, and ‘RiskParameters\{T\}’. In this case, an entirely susceptible population is assumed for a starting time of 0.0 time units. The internal code of this basic ‘Simulation\{T\}’ construction method is shown below. Individuals can start the simulation from other specified ‘DiseaseState’ s and/or the simulation may have a different start time using the other construction methods (not shown).

function Simulation(pop::Population,
rf::RiskFunctions{T},
rp::RiskParameters{T}) where T <: EpidemicModel
states = fill(State_S, pop.individuals)
tr = initialize(TransmissionRates, states, pop, rf, rp)
rates = initialize(EventRates, tr, states, pop, rf, rp)
events = Events{T}(pop.individuals)
net = TransmissionNetwork(pop.individuals)
return new{T}(0.0, 0, pop, rf, rp, states, tr, rates, events, net)
end

Finally, observational data can be generated from a completed ‘Simulation’ using the provided observe() function, with statistical distributions specified for observation delays, in the form of ‘UnivariateDistribution’s from the Distributions.jl package (Besançon, Anthoff, Arslan, Byrne, Lin, Papamarkou, and Pearson 2021), another dependency of Pathogen.jl.

4.4. Inference with Pathogen.jl

As with simulation in Pathogen.jl, performing parameter estimation for TN-ILMs via MCMC requires the definition of a ‘Population’ and of ‘RiskFunctions\{T\}’: ‘RiskParameters’ are now unknown, and will be sampled from the posterior distribution through MCMC. In order to do this, specification of prior distributions for each risk parameter is first required. These priors are structured through the ‘RiskPriors\{T\}’ type. This parametric struct has the same form as ‘RiskParameters’, but in place of each parameter value, a ‘UnivariateDistribution’ must be provided.

We also must specify priors for the event time data augmentation process. Currently uniform prior distributions are supported and applied broadly through an ‘EventExtents\{T\}’ struct. These are upper bounds on the length of time between observations of infectiousness, and the actual onset of infection, as well as the length of time between removal observations and actual removal times. For TN-ILMs that incorporate an exposed disease state, a bound on the length of the latent period must also be specified with ‘EventExtents\{T\}’.
Finally, an ‘MCMC{T}’ struct is constructed, containing all of the information required for performing MCMC, common across individual Markov chains (e.g., observational data, population data, prior distributions), as well as the individual Markov chains that sample from the TN-ILM posterior distribution. The composition of ‘MCMC{T}’ is:

```julia
mutable struct MCMC{T <: EpidemicModel}
    event_observations::EventObservations{T}
    event_extents::EventExtents{T}
    population::Population
    starting_states::Vector{DiseaseState}
    risk_functions::RiskFunctions{T}
    risk_priors::RiskPriors{T}
    transmission_network_prior::Union{Nothing, TNDistribution}
    markov_chains::Vector{MarkovChain{T}}
end
```

‘MarkovChain{T}’ includes vectors of event times from data augmentation, networks, as well as TN-ILM parameters, and is declared as:

```julia
mutable struct MarkovChain{T <: EpidemicModel}
    iterations::Int64
    events::Vector{Events{T}}
    transmission_network::Vector{TransmissionNetwork}
    risk_parameters::Vector{RiskParameters{T}}
    log_posterior::Vector{Float64}
    cov::OnlineStats.CovMatrix
end
```

To construct an ‘MCMC{T}’ object to perform inference, all of the following need to be specified: ‘EventObservations{T}’, ‘EventExtents{T}’, ‘Population’, ‘RiskFunctions{T}’, and ‘RiskPriors{T}’. Specification of a ‘TNDistribution’ as a prior distribution for the transmission network is optional. If a prior for the transmission network is not specified, a flat uniform prior is used. After construction of an ‘MCMC’ object, it is initialized using the `start!` function. During initialization the user must specify the number of chains to initialize, and the number of initialization attempts per chain. Each chain can be initialized on different cores. After initialization, MCMC can proceed using the `iterate!` function. With this function the number of iterations are specified, as well as the transition kernel variance for event time data augmentation. MCMC for each ‘MarkovChain{T}’ can also be ran in parallel. An example of the process of performing MCMC with `Pathogen.jl` is provided in Section 5.
5. Simulated example

In the following we present a full example using Pathogen.jl to:

- Generate an epidemic population;
- Simulate from an SIR TN-ILM;
- Simulate observations from the epidemic; and
- Use the observations to estimate event times, transmission network, and TN-ILM parameters via MCMC.

The source code to replicate this example with Julia v1.4.2 is included in section-5/code.jl. Changes and improvements to random number generation can occur between Julia releases (The Julia Project 2022). Using a release other than Julia v1.4.2 may lead to different results than those presented for this epidemic simulation. We start by loading the various publicly available Julia packages used in the example, and setting the seed of the random number generator:

```julia
julia> using Distances, Pathogen, Random, Plots, Plots.PlotMeasures;
julia> Random.seed!(11235);
```

We generate risk factor data for a population containing 100 individuals. A location (x and y coordinates over a $15 \times 30$ unit area), and an arbitrary Gamma($\alpha = 1, \beta = 1$) distributed risk factor:

```julia
julia> n = 100
julia> risks = DataFrame(x = rand(Uniform(0, 15), n),
                     y = rand(Uniform(0, 30), n), riskfactor1 = rand(Gamma(), n))
```

We pre-calculate Euclidean distances between individuals in a distance matrix, which is used in the specification of a 'Population' object:

```julia
julia> dists = [euclidean([risks[i, :x]; risks[j, :y]], [risks[j, :x]; risks[j, :y]]) for i = 1:n, j = 1:n]
julia> pop = Population(risks, dists)
```

Next, several functions of risk factors are defined with the signature expected by Pathogen.jl, and these structured into a 'RiskFunctions{SIR}' object:

```julia
julia> function _constant(params::Vector{Float64}, pop::Population, i::Int64)
    return params[1]
end
julia> function _one(params::Vector{Float64}, pop::Population, i::Int64)
    return 1.0
end
julia> function _linear(params::Vector{Float64}, pop::Population, i::Int64)
    return params[1] * pop.risks[i, :riskfactor1]
end
```
In the following call the first argument specifies the sparks function, the second the susceptibility function, the third the infectivity kernel, the forth the transmissibility function and the fifth and last the removal function:

```julia
julia> rf = RiskFunctions{SIR}(_constant, _one, _powerlaw, _one, _linear)
```

These risk functions are then parameterized:

```julia
julia> rparams = RiskParameters{SIR}([0.0001], Float64[], [4.0], Float64[], [0.1])
```

Starting states for each of the 100 individuals are then set, using the DiseaseState primitive type that is provided by Pathogen.jl. State_S, State_I, and State_R are valid states for the SIR model. At the start of the epidemic, individual 1 is infectious and the remaining 99 individuals are susceptible. The starting states and risk factors of the population, in addition to the parameterized risk functions of the TN-ILM, are organized into a ‘Simulation’ struct. With this, the simulation is finally run. The simulation ends once 200 time units have elapsed in the epidemic.

```julia
julia> starting_states = append!([State_I], fill(State_S, n-1))
julia> sim = Simulation(pop, starting_states, rf, rparams)
julia> simulate!(sim, tmax = 200.0)
```

The simulated epidemic can be explored visually. For our example, plots are generated in Julia using Plots.jl (Breloff 2015) with the GR plotting back-end (Heinen et al. 2020), with plot recipes provided in Pathogen.jl. An epidemic curve can be generated from an ‘Events{T}’ object, such as the one within our ‘Simulation{T}’, with

```julia
julia> p1 = plot(sim.events)
```

and transmission network plots showing the disease states of individuals at specified times, with

```julia
julia> p2 = plot(sim.transmission_network, sim.population, sim.events, 0.0, title = "Time = 0")
[...]
julia> p6 = plot(sim.transmission_network, sim.population, sim.events, 50.0, title = "Time = 50")
```

These plots can then be combined in a layout, to obtain Figure 2 with
Figure 2: An epidemic simulated from a TN-ILM using the code provided in Section 5. The top plot shows the overall number of individuals in each of the represented disease states over the length of the epidemic. The bottom row of plots shows individual disease state information along with the state of the transmission network at 5 time points in the epidemic.

```julia
julia> l = @layout [a; b c d e f]
julia> plot(p1, p2, p3, p4, p5, p6, layout = l)
```

From the simulated epidemic, observations can be generated, with statistical distributions used for the generation of observation delay. The optional `force = true` keyword argument is used to bound the infection observation delay such that an infection observation is guaranteed (i.e., an individual cannot move to the removed state undetected).

```julia
julia> obs = observe(sim, Uniform(0.5, 2.5), Uniform(0.5, 2.5),
force = true)
```

With observational data, inference for a specified model can be conducted. For our example, we will assume the model structure is known, and reuse the set of risk functions we declared for the epidemic simulation. For each parameter value, a prior distribution must be specified before commencing MCMC. We must also specify priors for our event times through `EventExtents`.

```julia
julia> rpriors = RiskPriors(SIR)([Exponential(0.0001)],
UnivariateDistribution[], [Uniform(1.0, 7.0)],
UnivariateDistribution[], [Uniform(0.0, 1.0)])
julia> ee = EventExtents(SIR)(5.0, 5.0)
```
Figure 3: Here we show several plots for visualizing MCMC results for a TN-ILM. The top plot shows the values of three TN-ILM parameters over 50k iterations. The bottom row of plots shows the epidemic curve posterior distributions in blue (posterior density estimated with every 20th iteration from iteration number 10k to 50k), with the true epidemic curves indicated with black.

MCMC will now be initialized, following the initialization strategy detailed in Section 3.3, with:

\begin{verbatim}
julia> mcmc = MCMC(obs, ee, pop, rf, rpriors)
julia> start!(mcmc, attempts = 50000)
\end{verbatim}

We then perform 50k iterations (second positional argument) for the initialized Markov chain using the `iterate!` function. Here, we elect to batch event time data augmentation into 10 sets (specified with the `event_batches` keyword argument), with a transition kernel variance of 1.0 for each event time (third positional argument). We also condition event time augmentation on the previous transmission network (specified with the `condition_on_network` keyword argument). 50k iterations in this manner require approximately 18.5 minutes on a computer with an Intel 2.7 GHz i7-3740QM processor, using Julia 1.4.2. For applications to real world data, or for simulations aiming at real world application, longer runs by multiple chains is advised in order to validate convergence.

\begin{verbatim}
julia> iterate!(mcmc, 50000, 1.0, condition_on_network = true, event_batches = 10)
\end{verbatim}

We provide convenient plotting functions for visualizing MCMC and posterior distributions,
after MCMC is complete, yielding the plots seen in Figures 3 and 4. The trace plot in Figure 3 is generated with:

```
 julia> p1 = plot(1:20:50001, mcmc.markov_chains[1].risk_parameters,
                yscale = :log10, title = "TN-ILM parameters")
```

For the remainder of this example we take every 20th iteration from iteration 10000 through 50000 as being representative samples from the TN-ILM posterior distribution. With this, the epidemic curve posterior distributions are visualized with

```
 julia> p2 = plot(mcmc.markov_chains[1].events[10000], State_S,
                linealpha = 0.01, title = "S")
 julia> for i = 10020:20:50000
        plot!(p2, mcmc.markov_chains[1].events[i], State_S,
              linealpha = 0.01)
    end
 julia> plot!(p2, sim.events, State_S, linecolor = :black)
```

for each state. Partially transparent epidemic curves are repeatedly plotted to show the posterior density. We also overlay the true event times in black. A plot layout is specified, and the plots are combined to form Figure 3.
### Table 1: The summary() function outputs a ‘DataFrame’ containing estimates of posterior mean, posterior variance, as well as credible intervals for each parameter in a TN-ILM, when applied to ‘MCMC’ objects.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Mean</th>
<th>Variance</th>
<th>95% credible interval</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\zeta$</td>
<td>$1.033 \times 10^{-4}$</td>
<td>$9.861 \times 10^{-9}$</td>
<td>$(2.869 \times 10^{-6}, 3.686 \times 10^{-4})$</td>
</tr>
<tr>
<td>$\beta$</td>
<td>$3.967$</td>
<td>$1.794 \times 10^{-2}$</td>
<td>$(3.715, 4.241)$</td>
</tr>
<tr>
<td>$\eta$</td>
<td>$8.168 \times 10^{-2}$</td>
<td>$8.457 \times 10^{-5}$</td>
<td>$(6.520 \times 10^{-2}, 1.001 \times 10^{-1})$</td>
</tr>
</tbody>
</table>

6. Application: 1861 Hagelloch measles outbreak

We apply the methods of Pathogen.jl to fit an $\text{SEIR}$ TN-ILM to the 1861 Hagelloch measles data set, originally published by Pfieflsticker (1863). This data set includes detailed observations on 188 children in Hagelloch, Germany. The data were analyzed by Oesterle (1992), who expanded the individual level detail of the data set, and predicted the transmission source for each case. The 1861 Hagelloch measles data have since been used to demonstrate numerous new analysis methods and new software in the area of individual level epidemic modeling and transmission network models. Neal and Roberts (2004) demonstrated reversible jump MCMC for nested epidemic models with event time data augmentation. Also using reversible jump MCMC for Bayesian model selection, Groendyke, Welch, and Hunter (2012) demonstrated...
exponential family random graphs that model the contact network over which the disease was transmitted, while considering the covariates in the Hagelloch data set, as implemented in package epinet. In Groendyke et al. (2012), infection and removal times are treated as known, and exposure/transmission times are inferred. Meyer et al. (2017) also used the outbreak data as an example application for the TwinSIR individual level model in the surveillance package. Here, we apply an SEIR TN-ILM which considers the available risk factor information and assumes the transmission network, as well as exposure, infection, and removal times to be all unknown, and to be imputed. The purpose of this analysis is to exhibit the functionality of Pathogen.jl using this exceptional data set, rather than to extend our understanding of the outbreak itself. The code to recreate this analysis with Julia v1.4.2 is contained in the supplementary material in section-6/code.jl.

The Hagelloch Measles data are included in the R package surveillance. These data are further modified for ease of use with Pathogen.jl, and we include them as an example data set in our own package.

Our analysis of the outbreak begins by loading our required packages, and setting a seed for replicability:

```julia
julia> using CSV, DelimitedFiles, Distances, Random, Pathogen, Plots,
        Plots.PlotMeasures, DataFrames
julia> Random.seed!(4321);
```

Our first step with Pathogen.jl is to construct a Population. We do this by loading our risk factor data, and pre-calculating a matrix of Euclidean distances between the homes of all children. We also allow transmission rates to depend on whether children were in the same household, or the same class at school. If individuals were listed at the same location, we set their distance to Inf, such that household transmission would be excluded from the Euclidean distance-based element of the infectivity kernel. For brevity we reserve the Julia code for construction of this distance matrix to the supplementary material. With the risk factors and distance matrix, a Population is then constructed with:

```julia
julia> risks = CSV.read("data/measles_hagelloch_1861_risk_factors.csv",
        DataFrame)
julia> pop = Population(risks, dist)
```

We specify observations for model fitting using the dates of prodrome, rash, and death. The date of prodrome is assumed to occur after the true onset of infectiousness. If an individual recovered, a removal time observation of the date of rash plus four days was assumed. This implies the true removal of that individual from the infectious state must have occurred before four days after displaying a rash, which is the typical maximum extent of infectiousness (Public Health Agency of Canada 2020). If an individual perished before this assumed removal time, their date of death would instead be taken as their removal time. This data processing step is again reserved for the supplementary material. We can construct an EventObservationsSEIR once we have vectors of infection and removal observations with:

```julia
julia> obs = EventObservations{SEIR}(infected, removed)
```

We do not exclude any cases, as has been done in some past analyses due to an isolated case following the primary outbreak. Groendyke, Welch, and Hunter (2011) found inclusion of
this particular case to affect parameter estimates of the random network model they present. In the application of a TN-ILM to these data, we expect that this case would be attributed to an external transmission source and not adversely impact model estimates, so its removal is not required. Specification of an initial infection is also not required with the TN-ILM due to it allowing for external transmission sources.

In our TN-ILM, individual disease state transitions are described by:

\[
\begin{align*}
\epsilon^*(i) &= \zeta, \\
\Omega_S(i) &= 1.0, \\
\Omega_T(j) &= 1.0, \\
\kappa(i, j) &= \alpha d_{i,j}^\beta + \tau I_{class}(i, j) + \rho I_{house}(i, j), \\
\Omega_L(i) &= \gamma, \\
\Omega_R(i) &= \eta,
\end{align*}
\]  

where \(d_{i,j}\) is the Euclidean distance between individuals \(i\) and \(j\), \(I_{class}(i, j)\) and \(I_{house}(i, j)\) are indicator functions for whether \(i\) and \(j\) are members of the same classroom, and same household, respectively. This model structure is specified as

```julia
julia> function _constant(params::Vector{Float64}, pop::Population, i::Int64)
    return params[1]
end
julia> function _one(params::Vector{Float64}, pop::Population, i::Int64)
    return 1.0
end
julia> function _powerlaw_plus(params::Vector{Float64}, pop::Population, i::Int64, k::Int64)
    return params[1] * pop.distances[k, i][1]^(-params[2]) +
        params[3] * pop.distances[k, i][2] +
        params[4] * pop.distances[k, i][3]
end
julia> rf = RiskFunctions{SEIR}(_constant, # sparks function
      _one, # susceptibility function
      _powerlaw_plus, # infectivity function
      _one, # transmissibility function
      _constant, # latency function
      _constant) # removal function
```

Weakly informative priors that restrain parameters to positive values were selected for all variables for the purpose of fitting this model using MCMC, with:

\[
\begin{align*}
\zeta &\sim \text{Uniform}(0.0, 0.1), \\
\alpha &\sim \text{Uniform}(0.0, 7.0), \\
\beta &\sim \text{Uniform}(0.0, 7.0), \\
\tau &\sim \text{Uniform}(0.0, 1.0), \\
\rho &\sim \text{Uniform}(0.0, 1.0), \\
\gamma &\sim \text{Uniform}(0.0, 1.0), \\
\eta &\sim \text{Uniform}(0.0, 1.0).
\end{align*}
\]

Uniform prior distributions were specified for the delay between true infection and removal times and infection and removal observations of Uniform(0.0, 3.0) and Uniform(0.0, 2.0), respectively. The prior distribution for exposure times is specified relative to the latent infection time as having occurred Uniform(5.0, 14.0) days earlier. A flat uniform prior was assumed for the transmission network.

Using Pathogen.jl, we specify these prior distributions with:
Figure 5: The trace plot after 200k iterations of a TN-ILM with 7 parameters applied to the 1861 Hagelloch measles data. Convergence was assumed to occur by 100k iterations, with a thinning rate of 50 used in selection of samples to estimate the joint posterior distribution of the TN-ILM. For display purposes, only every 20th iteration is shown in this trace plot.

```
julia> rpriors = RiskPriors{SEIR}([Uniform(0.0, 0.1)], UnivariateDistribution[], [Uniform(0.0, 7.0); Uniform(0.0, 7.0); Uniform(0.0, 1.0); Uniform(0.0, 1.0)], UnivariateDistribution[], [Uniform(0.0, 1.0), [Uniform(0.0, 1.0)]])
julia> ee = EventExtents{SEIR}((5.0, 14.0), 3.0, 2.0)
```

We next construct an ‘MCMC{SEIR}’ object, and initialize it by selecting the best parameter set from 100k initialization attempts. From the selected initial parameter values, we then perform 200k MCMC iterations. For this example, event time data augmentation is completed in 10 blocks, with a transition kernel variance set to 1.0 for each event time.

```
julia> mcmc = MCMC(obs, ee, pop, rf, rpriors)
julia> start!(mcmc, attempts = 100000)
julia> iterate!(mcmc, 200000, 1.0, condition_on_network = true, event_batches = 10)
```

A progress bar is displayed during MCMC initialization and iteration. It took approximately 15 minutes, and 3 hours 53 minutes to run those processes, respectively, on a computer with an Intel 2.7 GHz i7-3740QM processor, using Julia 1.4.2. Once completed, we can visually assess a trace plot of the TN-ILM parameters with

```
julia> plot(1:20:200001, mcmc.markov_chains[1].risk_parameters)
```

which shows every 20th set of parameter values, and generates the plot seen in Figure 5. The TN-ILM parameters appear to converge quickly to the posterior distribution.
Table 2: Estimates of posterior mean, posterior variance, and 95% credible intervals for the 7 parameters of the TN-ILM that we applied to the Hagelloch measles data.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Mean</th>
<th>Variance</th>
<th>95% credible interval</th>
</tr>
</thead>
<tbody>
<tr>
<td>ζ</td>
<td>0.001261</td>
<td>4.107 × 10^{-7}</td>
<td>(0.0003319, 0.002840)</td>
</tr>
<tr>
<td>α</td>
<td>4.138</td>
<td>3.376</td>
<td>(0.6007, 6.872)</td>
</tr>
<tr>
<td>β</td>
<td>1.963</td>
<td>0.03318</td>
<td>(1.478, 2.227)</td>
</tr>
<tr>
<td>τ</td>
<td>0.02006</td>
<td>1.147 × 10^{-5}</td>
<td>(0.01398, 0.02711)</td>
</tr>
<tr>
<td>ρ</td>
<td>0.1952</td>
<td>0.001172</td>
<td>(0.1324, 0.2699)</td>
</tr>
<tr>
<td>γ</td>
<td>0.1259</td>
<td>8.735 × 10^{-5}</td>
<td>(0.1084, 0.1452)</td>
</tr>
<tr>
<td>η</td>
<td>0.1210</td>
<td>8.102 × 10^{-5}</td>
<td>(0.1039, 0.1391)</td>
</tr>
</tbody>
</table>

Which yields the posterior estimates shown in Table 2. We can also visually explore the posterior distribution of event times and overall epidemic curves, as well as the transmission network as we did in the simulated example in Section 5. At the top of Figure 6, we contrast the posterior distribution of the transmission network from our TN-ILM, to that suggested by Oesterle (1992). We add a small amount of variation to individual locations for display purposes only, to enable viewing of multiple individuals in the same household. The epidemic curves at the bottom of Figure 6 show credible bands for aggregate infection and removal times. The black lines on the infection and removal time plots indicate the observed infection and removal times. The code to recreate Figure 6 is included in the supplementary material.

When comparing our transmission network posterior distribution to Oesterle’s analysis, we note that Oesterle suggests individual 45 to be the source of 30 further infections, while the transmission network posterior distribution from our TN-ILM indicates an out degree of only 11.11 for this individual. Oesterle may have been able to incorporate more contextual information into their transmission network reconstruction, resulting in this higher estimate. Our estimated out degree considers probabilities for specific transmissions, while Oesterle’s transmission network reconstruction does not represent uncertainty, which also may account for some of this difference. Measles is a highly infectious airborne disease, with a basic reproductive number, the average number of secondary cases from an index case in a susceptible population, usually cited as in the range of 12–18 (Guerra, Bolotin, Lim, Heffernan, Deeks, Li, and Crowcroft 2017). Variation in the number of secondary cases from individual exposure sources during an outbreak is expected. Both the number of secondary transmissions for individual 45 in our transmission network reconstruction, and in Oesterle’s reconstruction, are within a plausible range for this disease. Individual 141 is the case that has been removed in some previous analyses. Our approach yields a marginal posterior probability of this individual having an external transmission source of approximately 1.0.

Overall, we note that the posterior mode transmission network from our TN-ILM is in agreement with Oesterle on 103 of the 188 transmission sources. More informative prior distributions for event times, and for exposure time especially, would likely affect our transmission network reconstruction. Consideration of other risk factors, or consideration of the presently
Figure 6: Top: The marginal posterior distribution of the transmission network is shown in contrast to the transmission network presented by Oesterle (1992). Bottom: The marginal posterior distributions of the count of individuals in each disease state through time are shown, with observation of infectiousness (onset of prodrome), and observation of removal (assumed to be 4 days following onset of rash), overlaid in black.

used risk factors in different forms, would also likely impact the transmission network posterior distribution.

The epidemic curves show narrow bands for infection and removal times in the population, with higher uncertainty indicated for the exposure time. The posterior mean estimate for $\gamma$ implies a posterior mean for the latent period of 7.95 days. The posterior mean of the incubation period is 9.42 days. An analysis of measles data from households with two cases by Klinkenberg and Nishiura (2011) put the mean incubation period for this disease to be slightly higher, between 11 and 12 days (Klinkenberg and Nishiura 2011). The posterior mean for $\eta$ implies a mean infectious period of 8.27 days.

The remaining parameters speak to the relative sources of infectious pressure. The infectious pressure from an infectious family member was 9.73 times that of an infectious classmate. An infectious individual residing 15 meters away applied infectious pressure approximately equal to that of an infectious classmate. With 30 meters between residences, the infectious pressure from an infectious individual drops to approximately 25% of that of an infectious classmate. While it was less common for an infection to have an external source, an estimate of a constant rate of $1.26 \times 10^{-3}$ was produced for $\zeta$. Our transmission network posterior
distribution indicated an average of 7.70 transmissions as coming from outside the observed population of 188 children in the data set.

While modeling approaches have varied, we find these results are generally in agreement with the conclusions of other authors, being that household, then classroom transmission, were of primary importance in this outbreak.

7. Conclusion

We have introduced Pathogen.jl, a Julia package, with flexible simulation and inference capabilities for TN-ILMs. Pathogen.jl, currently supports SEIR, SIR, SEI and SI TN-ILMs. Functions of risk factors for disease state transitions within each of these model classes are effectively unrestricted, and this generality does not come at an implicit performance cost due to JIT compilation in Julia. With TN-ILM simulations, observational data can be generated such that realistic simulation studies can be conducted. Simulated observations such as these or real data can be fit to TN-ILMs with Pathogen.jl, which performs MCMC with event time and transmission network data augmentation. For several of the types that are introduced by Pathogen.jl, visualization methods are also provided. These visualization methods include epidemic curves, transmission network visuals, and trace plots for MCMC.

In the future, we hope to increase the feature set of Pathogen.jl to include TN-ILMs incorporating time-varying disease state transition rates, and to provide an API for individual movement into and out of populations. The flexibility of Pathogen.jl could be explored further through the use of artificial neural networks for one or more rate-describing functions, similar to what has been done with neural differential equations with DiffEqFlux.jl (Rackauckas, Innes, Ma, Bettencourt, White, and Dixit 2019). Providing simulation and inference capabilities for phylodynamic ILMs, which incorporate densely sampled pathogen genetic sequence data, is also of particular interest. Such data sets have only relatively recently become feasible to compile with advances to sequencing technology (Leventhal, Günthard, Bonhoeffer, and Stadler 2014). Pathogen genetic data can help identify likely transmission paths, and improve our understanding of key connectivity factors in disease transmission. Additional inference methods may also be added to Pathogen.jl, such as approximate Bayesian computation. Beyond these, as the Julia package ecosystem continues to mature, integration with other Julia packages will be prioritized. While not only reducing the code base of Pathogen.jl, this process will present the opportunity to add features, increase generality, improve performance, and improve usability of the package. Notably, there are several maturing probabilistic programming language packages in Julia including Turing.jl (Ge, Xu, and Ghahramani 2018), Soss.jl (Scherrer and Zhao 2020), and Gen.jl (Cusumano-Towner, Saad, Lew, and Mansinghka 2019). The use of one of these packages could be leveraged in providing additional algorithms for fitting TN-ILMs, such as Hamiltonian Monte Carlo, variational inference, or particle filtering based approaches.

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