epiworld

0.0-1

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# **Chapter 1**

# Main Page

# 1.1 epiworld

This C++ template-header-only library provides a general framework for epidemiologic simulation. The main features of the library are:

- 1. Four key classes: Model, Person, Tool, and Virus.
- 2. The model features a social networks of Persons.
- 3. Persons can have multiple Tools as a defense system.
- 4. Tools can reduce contagion rate, transmissibility, death rates, and improve recovery rates.
- 5. Viruses can mutate (generating new variants).
- 6. Models can feature multiple states, e.g., HEALTHY, SUSCEPTIBLE, etc.
- 7. Models can have an arbitrary number of parameters.
- 8. **REALLY FAST** About 6.5 Million person/day simulations per second.

# 1.2 Hello world

# Here is a simple SIRS model implemented with

```
#include "../include/epiworld/epiworld.hpp"
using namespace epiworld;
int main()
{
          // Creating a model
          Model<> model;
          // Adding the tool and virus
          Virus<> virus("covid 19");
          virus.set_post_immunity(1.0);
          model.add_virus_n(virus, 5);

          Tool<> tool("vaccine");
          model.add_tool(tool, .5);
          // Generating a random pop
          model.population_smallworld(100000);
          // Initializing setting days and seed
          model.init(100, 123);
```

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```
// Running the model
model.run();
model.print();
```

And you should get something like the following:

Running the model...

```
SIMULATION STUDY
Population size
                 : 100000
Days (duration)
                 : 100 (of 100)
Number of variants : 1
Last run elapsed t : 280.00ms
Rewiring
                : off
Virus(es):
 - covid 19 (baseline prevalence: 5 seeds)
Tool(s):
  vaccine (baseline prevalence: 50.00%)
Model parameters:
Distribution of the population at time 100:
- Total healthy (S) : 99995 -> 97390
- Total recovered (S) : 0 -> 2554
- Total infected (I)
                    :
                            5 -> 56
 - Total removed (R)
                            0 -> 0
(S): Susceptible, (I): Infected, (R): Recovered
```

Which took about 0.280 seconds.

### 1.2.1 **Tools**

# 1.2.2 Contagion

Susceptible individuals can acquire a virus from any of their infected connections. The probability that susceptible individual i gets the virus v from individual j depends on how three things:

- 1. The transmissibility of the virus, Pv in [0,1],
- 2. The contagion reduction factor of i, Cr in [0,1], and
- 3. The host's transmission reduction factor, Tr [0,1].

The last two are computed from i and j's tools. Ultimately, the probability of i getting virus v from j equals:  $P(Virus \ v) = Pv \ \star \ (1 - Cr) \ \star (1 - Tr)$ 

Nonetheless, the default behavior of the simulation model is to assume that individuals can acquire only one disease at a time, if any. This way, the actual probability is:

```
P(Virus \ v| \ at \ most \ one \ virus) = Prcond(i,v,j)
```

```
The latter is calculated using Bayes' rule
```

This way, viruses with higher transmissibility will be more likely to be acquired when competing with other variants.

# Chapter 2

# **Class Index**

# 2.1 Class List

Here are the classes, structs, unions and interfaces with brief descriptions:

Action < TSeq >	 	. 5
AdjList		
DataBase < TSeq >		
Statistical data about the process		. 7
LFMCMC< TData >		
Likelihood-Free Markov Chain Monte Carlo		. 8
Location < TSeq >		. 10
Model < TSeq >		
Core class of epiworld		. 10
Person < TSeq >		
Person (agents)		. 18
PersonTools < TSeq >		
List of tools available for the individual to		20
PersonViruses < TSeq >		
Set of viruses in host		. 21
Progress		
A simple progress bar	 	. 22
Queue < TSeq >		
Controls which agents are verified at each step	 	. 22
RandGraph	 	. 23
Tool< TSeq >		
Tools for defending the host against the virus	 	23
UserData < TSeq >		
Personalized data by the user		. 24
vecHasher< T >		
Vector hasher		. 26
Virus< TSeq >		
Virus	 	. 27

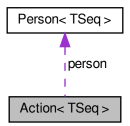
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# **Chapter 3**

# **Class Documentation**

# 3.1 Action < TSeq > Struct Template Reference

Collaboration diagram for Action< TSeq >:



# **Public Attributes**

- Person< TSeq > \* person
- epiworld\_fast\_uint new\_status
- ActionFun< TSeq > call
- · epiworld\_fast\_int queue

The documentation for this struct was generated from the following file:

• include/epiworld/config.hpp

# 3.2 AdjList Class Reference

# **Public Member Functions**

AdjList (const std::vector< unsigned int > &source, const std::vector< unsigned int > &target, bool directed, int min\_id=-1, int max\_id=-1)

Construct a new Adj List object.

- void read\_edgelist (std::string fn, int skip=0, bool directed=true, int min\_id=-1, int max\_id=-1)
- std::map< unsigned int, unsigned int > operator() (unsigned int i) const
- void print (unsigned int limit=20u) const
- · unsigned int get\_id\_max () const
- unsigned int get\_id\_min () const
- size\_t vcount () const
- size\_t ecount () const
- std::map< unsigned int, std::map< unsigned int, unsigned int > > & get\_dat ()
- · bool is directed () const

# 3.2.1 Constructor & Destructor Documentation

# 3.2.1.1 AdjList()

Construct a new Adj List object.

It will create an adjacency list object with maxid - minid + 1 nodes. If min\_id and max\_id are not specified (both < 0), then the program will try to figure them out automatically by looking at the range of the observed ids.

#### **Parameters**

source	Unsigned int vector with the source
target	Unsigned int vector with the target
directed	Bool true if the network is directed
min_id	int min id.
max_id	int max id.

The documentation for this class was generated from the following files:

- include/epiworld/adjlist-bones.hpp
- · include/epiworld/adjlist-meat.hpp

# 3.3 DataBase < TSeq > Class Template Reference

Statistical data about the process.

#include <database-bones.hpp>

### **Public Member Functions**

- DataBase (int freq=1)
- void record\_variant (VirusPtr< TSeq > v)

Registering a new variant.

- void record\_tool (ToolPtr< TSeq > t)
- void set\_seq\_hasher (std::function< std::vector< int >(TSeq)> fun)
- void set\_model (Model < TSeq > &m)
- Model < TSeq > \* get\_model ()
- · void record ()
- const std::vector< TSeq > & get\_sequence () const
- const std::vector< int > & get\_nexposed () const
- size\_t size () const
- void write\_data (std::string fn\_variant\_info, std::string fn\_variant\_hist, std::string fn\_tool\_info, std::string fn\_tool\_hist, std::string fn\_transmission, std::string fn\_transition) const
- void record\_transmission (int i, int j, int variant)
- size\_t get\_nvariants () const
- · void reset ()
- void set\_user\_data (std::vector< std::string > names)
- void add\_user\_data (std::vector< epiworld\_double > x)
- void **add\_user\_data** (unsigned int j, epiworld\_double x)
- UserData < TSeq > & get\_user\_data ()

# Get recorded information from the model

#### **Parameters**

what std::string, The status, e.g., 0, 1, 2, ...

#### Returns

In get\_today\_total, the current counts of what.

In get\_today\_variant, the current counts of what for each variant.

In get\_hist\_total, the time series of what

In get\_hist\_variant, the time series of what for each variant.

In get\_hist\_total\_date and get\_hist\_variant\_date the corresponding dates

- int get\_today\_total (std::string what) const
- int get\_today\_total (epiworld\_fast\_uint what) const
- void get today total (std::vector < std::string > \*status=nullptr, std::vector < int > \*counts=nullptr) const
- void get\_today\_variant (std::vector< std::string > &status, std::vector< int > &id, std::vector< int > &counts) const
- void get\_hist\_total (std::vector< int > \*date, std::vector< std::string > \*status, std::vector< int > \*counts) const
- void get\_hist\_variant (std::vector< int > &date, std::vector< int > &id, std::vector< std::string > &status, std::vector< int > &counts) const

### **Friends**

- class Model < TSeq >
- void default\_add\_virus (Action< TSeq > &a, Model< TSeq > \*m)
- void default\_add\_tool (Action < TSeq > &a, Model < TSeq > \*m)
- void default\_rm\_virus (Action < TSeq > &a, Model < TSeq > \*m)
- void default\_rm\_tool (Action < TSeq > &a, Model < TSeq > \*m)

# 3.3.1 Detailed Description

```
template<typename TSeq> class DataBase< TSeq>
```

Statistical data about the process.

**Template Parameters** 



## 3.3.2 Member Function Documentation

### 3.3.2.1 record\_variant()

Registering a new variant.

## **Parameters**

*v* Pointer to the new variant. Since variants are originated in the host, the numbers simply move around. From the parent variant to the new variant. And the total number of infected does not change.

The documentation for this class was generated from the following files:

- · include/epiworld/database-bones.hpp
- include/epiworld/database-meat.hpp

# 3.4 LFMCMC< TData > Class Template Reference

Likelihood-Free Markov Chain Monte Carlo.

```
#include <lfmcmc.hpp>
```

### **Public Member Functions**

- void run (VEC(epiworld\_double) param\_init, size\_t n\_samples\_, epiworld\_double epsilon\_)
- · LFMCMC (TData &observed data )
- · void set observed data (TData &observed data )
- void **set\_proposal\_fun** (FUN< void(VEC(epiworld\_double)&, LFMCMC< TData > \*)> fun)
- void set simulation fun (FUN< TData(VEC(epiworld double)&, LFMCMC< TData > \*)> fun)
- void set summary fun (FUN< VEC(epiworld double)(TData &, LFMCMC< TData > \*)> fun)
- void set\_kernel\_fun (FUN< epiworld\_double(VEC(epiworld\_double)&, epiworld\_double, LFMCMC< TData > \*)> fun)
- const size t get n samples ()
- const size\_t get\_n\_statistics ()
- const size\_t get\_n\_parameters ()
- const epiworld double get epsilon ()
- const VEC (epiworld\_double) &get\_params\_now()
- const VEC (epiworld double) &get params prev()
- const VEC (epiworld double) &get params init()
- const VEC (epiworld\_double) &get\_statistics\_obs()
- const VEC (epiworld\_double) &get\_statistics\_hist()
- · const VEC (bool) &get\_statistics\_accepted()
- const VEC (epiworld double) &get posterior If prob()
- const VEC (epiworld double) &get acceptance prob()
- const VEC (epiworld\_double) &get\_drawn\_prob()
- VEC (TData) \*get sampled data()

### Random number generation

#### **Parameters**

eng

- void set rand engine (std::mt19937 &eng)
- std::mt19937 \* get\_rand\_endgine ()
- void seed (unsigned int s)
- void **set\_rand\_gamma** (epiworld\_double alpha, epiworld\_double beta)
- epiworld\_double runif ()
- epiworld\_double rnorm ()
- epiworld\_double **rnorm** (epiworld\_double mean, epiworld\_double sd)
- epiworld double rgamma ()
- epiworld\_double rgamma (epiworld\_double alpha, epiworld\_double beta)

# 3.4.1 Detailed Description

template < typename TData > class LFMCMC < TData >

Likelihood-Free Markov Chain Monte Carlo.

### **Template Parameters**

TData Type of data that is generated

The documentation for this class was generated from the following file:

• include/epiworld/math/lfmcmc.hpp

# 3.5 Location < TSeq > Class Template Reference

## **Public Member Functions**

```
    add_person (Person< TSeq > &p)
    add_person (Person< TSeq > *p)
```

- size\_t count () const
- · void reset ()

The documentation for this class was generated from the following file:

· include/epiworld/location-bones.hpp

# 3.6 Model < TSeq > Class Template Reference

Core class of epiworld.

```
#include <model-bones.hpp>
```

# **Public Member Functions**

- Model (const Model < TSeq > &m)
- Model (Model < TSeq > &&m)
- Model < TSeq > & operator= (const Model < TSeq > &m)
- void **clone\_population** (std::vector< Person< TSeq >> &p, std::map< int, int > &p\_ids, bool &d, Model< TSeq > \*m=nullptr) const
- void clone\_population (const Model < TSeq > &m)
- DataBase < TSeq > & get\_db ()
- epiworld\_double & operator() (std::string pname)
- size\_t size () const
- void record\_variant (VirusPtr< TSeq > v)
- int get\_nvariants () const
- unsigned int get\_ndays () const
- unsigned int **get\_n\_replicates** () const
- void set\_ndays (unsigned int ndays)
- bool **get\_verbose** () const
- void verbose\_off ()
- void verbose\_on ()
- · int today () const

The current time of the model.

void write\_data (std::string fn\_variant\_info, std::string fn\_variant\_hist, std::string fn\_tool\_info, std::string fn tool hist, std::string fn total hist, std::string fn transmission, std::string fn transition) const

```
Wrapper of DataBase::write_data
```

std::map< std::string, epiworld\_double > & params ()

void reset ()

Reset the model.

- · void print () const
- Model < TSeq > && clone () const
- void **get\_elapsed** (std::string unit="auto", epiworld\_double \*last\_elapsed=nullptr, epiworld\_double \*total\_
   elapsed=nullptr, std::string \*unit\_abbr=nullptr, bool print=true) const
- void add\_global\_action (std::function < void(Model < TSeq > \*) > fun, int date)
   Set a global action.
- void run\_global\_actions ()
- void clear\_status\_set ()

#### Set the backup object

backup can be used to restore the entire object after a run. This can be useful if the user wishes to have individuals start with the same network from the beginning.

- void set backup ()
- void restore\_backup ()

## Random number generation

#### **Parameters**

eng	Random number generator
s	Seed

- void set\_rand\_engine (std::mt19937 &eng)
- std::mt19937 \* get\_rand\_endgine ()
- void **seed** (unsigned int s)
- void set\_rand\_gamma (epiworld\_double alpha, epiworld\_double beta)
- epiworld double runif ()
- epiworld double rnorm ()
- epiworld\_double  ${\bf rnorm}$  (epiworld\_double mean, epiworld\_double sd)
- epiworld\_double rgamma ()
- epiworld\_double **rgamma** (epiworld\_double alpha, epiworld\_double beta)

#### Add Virus/Tool to the model

This is done before the model has been initialized.

## Parameters

V	Virus to be added
t	Tool to be added
preval	Initial prevalence (initial state.) It can be specified as a proportion (between zero and one,) or an integer indicating number of individuals.

- void add\_virus (Virus < TSeq > v, epiworld\_double preval)
- void add\_virus\_n (Virus< TSeq > v, unsigned int preval)
- void  ${\it add\_tool}$  ( ${\it Tool}{<}$  TSeq > t, epiworld\_double preval)
- void add\_tool\_n (Tool < TSeq > t, unsigned int preval)

## Accessing population of the model

#### **Parameters**

fn	std::string Filename of the edgelist file.
skip	int Number of lines to skip in fn.
directed	bool Whether the graph is directed or not.
min_id	int Minimum id number (if negative, the program will try to guess from the data.)
max_id	int Maximum id number (if negative, the program will try to guess from the data.)
al	AdjList to read into the model.

- void population\_from\_adjlist (std::string fn, int skip=0, bool directed=false, int min\_id=-1, int max\_id=-1)
- void population\_from\_adjlist (AdjList al)
- bool is\_directed () const
- std::vector< Person< TSeq > > \* get\_population ()
- void population\_smallworld (unsigned int n=1000, unsigned int k=5, bool d=false, epiworld\_double p=. ← 01)

#### Functions to run the model

#### **Parameters**

seed	Seed to be used for Pseudo-RNG.
ndays	Number of days (steps) of the simulation.
fun	In the case of run_multiple, a function that is called after each experiment.

- · void init (unsigned int ndays, unsigned int seed)
- void update\_status ()
- void mutate\_variant ()
- void next ()
- void run ()

Runs the simulation (after initialization)

 void run\_multiple (unsigned int nexperiments, std::function< void(Model< TSeq > \*)> fun, bool reset, bool verbose)

### Rewire the network preserving the degree sequence.

This implementation assumes an undirected network, thus if  $\{(i,j), (k,l)\} \rightarrow \{(i,l), (k,j)\}$ , the reciprocal is also true, i.e.,  $\{(j,i), (l,k)\} \rightarrow \{(j,k), (l,i)\}$ .

### **Parameters**

proportion	Proportion of ties to be rewired.
------------	-----------------------------------

### Returns

A rewired version of the network.

- void set\_rewire\_fun (std::function< void(std::vector< Person< TSeq >> \*, Model< TSeq > \*, epiworld\_double)> fun)
- void set\_rewire\_prop (epiworld\_double prop)
- epiworld\_double get\_rewire\_prop () const
- void rewire ()

# Export the network data in edgelist form

#### **Parameters**

fn	std::string. File name.
source	Integer vector
target	Integer vector

When passing the source and target, the function will write the edgelist on those.

- · void write edgelist (std::string fn) const
- void write\_edgelist (std::vector< unsigned int > &source, std::vector< unsigned int > &target) const

### Manage status (states) in the model

The functions get\_status return the current values for the statuses included in the model.

#### **Parameters**

```
lab std::string Name of the status.
```

#### Returns

```
add_status* returns nothing.
get_status_* returns a vector of pairs with the statuses and their labels.
```

- void add\_status (std::string lab, UpdateFun< TSeq > fun)
- const std::vector< std::string > & get status () const
- const std::vector< UpdateFun< TSeq > > & get\_status\_fun () const
- void print\_status\_codes () const

## Set the user data object

### Parameters

name	s	string vector with the names of the variables.

- void set\_user\_data (std::vector< std::string > names)
   [@
- void add\_user\_data (unsigned int j, epiworld\_double x)
- void add\_user\_data (std::vector< epiworld\_double > x)
- UserData< TSeq > & get\_user\_data ()

## **Queuing system**

When queueing is on, the model will keep track of which agents are either in risk of exposure or exposed. This then is used at each step to act only on the aforementioned agents.

void queuing\_on ()

Activates the queuing system (default.)

void queuing\_off ()

Deactivates the queuing system.

bool is\_queuing\_on () const

Query if the queuing system is on.

Queue < TSeq > & get\_queue ()

Retrieve the Queue object.

### Get the susceptibility reduction object

#### **Parameters**

V

#### Returns

epiworld double

- epiworld\_double get\_susceptibility\_reduction (VirusPtr< TSeq > v)
- epiworld double get\_transmission\_reduction (VirusPtr< TSeq > v)
- epiworld\_double get\_recovery\_enhancer (VirusPtr< TSeq > v)
- epiworld\_double get\_death\_reduction (VirusPtr< TSeq > v)
- void set susceptibility reduction mixer (MixerFun < TSeq > fun)
- void set transmission reduction mixer (MixerFun < TSeg > fun)
- void set recovery enhancer mixer (MixerFun< TSeq > fun)
- void set\_death\_reduction\_mixer (MixerFun< TSeq > fun)

# **Public Attributes**

- std::vector< epiworld\_double > array\_double\_tmp
- std::vector< VirusPtr< TSeq >> array\_virus\_tmp

#### **Friends**

- class Person < TSeq >
- class DataBase< TSeq >
- class  $\mathbf{Queue} < \mathbf{TSeq} >$

# Setting and accessing parameters from the model

Tools can incorporate parameters included in the model. Internally, parameters in the tool are stored as pointers to an std::map<> of parameters in the model. Using the unsigned int method directly fetches the parameters in the order these were added to the tool. Accessing parameters via the std::string method involves searching the parameter directly in the std::map<> member of the model (so it is not recommended.)

The function set\_param() can be used when the parameter already exists in the model.

The par() function members are aliases for get\_param().

# **Parameters**

initial_val	
pname	Name of the parameter to add or to fetch

### Returns

The current value of the parameter in the model.

- epiworld\_double \* p0
- epiworld double \* p1
- epiworld\_double \* p2

- epiworld\_double \* p3
- epiworld\_double \* p4
- epiworld\_double \* p5
- epiworld double \* p6
- epiworld double \* p7
- epiworld\_double \* p8
- epiworld\_double \* p9
- epiworld\_double \* p10
- epiworld\_double \* p11
- epiworld\_double \* p12
- epiworld double \* p13
- epiworld\_double \* p14
- epiworld double \* p15
- epiworld\_double \* p16
- epiworld double \* p17
- epiworld\_double \* p18
- epiworld double \* p19
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- epiworld\_double \* p29
- epiworld\_double \* p30
- epiworld\_double \* p31
- epiworld\_double \* p31epiworld\_double \* p32
- epiworld double \* p33
- epiworld\_double \* p34
- epiworld\_double \* p35
- epiworld\_double \* p36
- epiworld\_double \* p37
- epiworld\_double \* p38
- epiworld\_double \* p39
- unsigned int npar\_used = 0u
- epiworld\_double add\_param (epiworld\_double initial\_val, std::string pname)
- epiworld\_double set\_param (std::string pname)
- epiworld\_double **get\_param** (unsigned int k)
- epiworld double **get\_param** (std::string pname)
- epiworld\_double **par** (unsigned int k)
- epiworld double par (std::string pname)

# 3.6.1 Detailed Description

template<typename TSeq = bool> class Model< TSeq >

Core class of epiworld.

The model class provides the wrapper that puts together Person, Virus, and Tools.

# **Template Parameters**

TSeq

Type of sequence. In principle, users can build models in which virus and human sequence is represented as numeric vectors (if needed.)

# 3.6.2 Member Function Documentation

# 3.6.2.1 add\_global\_action()

```
template<typename TSeq = bool>
void Model< TSeq >::add_global_action (
            std::function< void(Model< TSeq > *)> fun,
            int date )
```

Set a global action.

#### **Parameters**

fun	A function to be called on the prescribed dates
date	Integer indicating when the function is called (see details)

When date is less than zero, then the function is called at the end of every day. Otherwise, the function will be called only at the end of the indicated date.

### 3.6.2.2 reset()

```
template<typename TSeq = bool>
void Model< TSeq >::reset ( )
```

Reset the model.

Resetting the model will:

- · clear the database
- restore the population (if set\_backup() was called before)
- · re-distribute tools
- · re-distribute viruses
- set the date to 0

# 3.6.2.3 run\_multiple()

#### **Parameters**

nexperiments	Multiple runs of the simulation
--------------	---------------------------------

# 3.6.2.4 write\_data()

Wrapper of DataBase::write\_data

### **Parameters**

fn_variant_info	Filename. Information about the variant.
fn_variant_hist	Filename. History of the variant.
fn_tool_info	Filename. Information about the tool.
fn_tool_hist	Filename. History of the tool.
fn_total_hist	Filename. Aggregated history (status)
fn_transmission	Filename. Transmission history.
fn_transition	Filename. Markov transition history.

The documentation for this class was generated from the following files:

- include/epiworld/config.hpp
- include/epiworld/model-bones.hpp

# 3.7 Person < TSeq > Class Template Reference

# Person (agents)

```
#include <person-bones.hpp>
```

# **Public Member Functions**

- int get\_id () const
   Id of the individual.
- unsigned int <a href="mailto:get\_index">get\_index</a> () const

```
Location (0, ..., n-1).
```

- std::mt19937 \* get\_rand\_endgine ()
- Model < TSeq > \* get\_model ()
- VirusPtr< TSeq > & get\_virus (int i)
- std::vector< VirusPtr< TSeq > > & get\_viruses ()
- · size\_t get\_n\_viruses () const noexcept
- ToolPtr< TSeq > & get\_tool (int i)
- std::vector< ToolPtr< TSeq > > & get\_tools ()
- size t get n tools () const noexcept
- void mutate\_variant ()
- void add\_neighbor (Person < TSeq > \*p, bool check\_source=true, bool check\_target=true)
- std::vector < Person < TSeq > \* > & get\_neighbors ()
- void change\_status (epiworld\_fast\_uint new\_status, epiworld\_fast\_int queue=0)
- const epiworld\_fast\_uint & get\_status () const
- · void reset ()
- · bool has tool (unsigned int t) const
- · bool has tool (std::string name) const
- · bool has\_virus (unsigned int t) const
- · bool has virus (std::string name) const

#### Add/Remove Virus/Tool

Calling any of these functions will lock the agent (person) until the action is applied at the end of the iteration. Calling any of this functions when the agent is locked will cause an error.

#### **Parameters**

tool	Tool to add
virus	Virus to add
status_new	Status after the change
queue	

- void add\_tool (ToolPtr< TSeq > tool, epiworld\_fast\_int status\_new=-1, epiworld\_fast\_int queue=0)
- void **add\_virus** (VirusPtr< TSeq > virus, epiworld\_fast\_int status\_new=-1, epiworld\_fast\_int queue=0)
- void rm\_tool (epiworld\_fast\_uint tool\_idx, epiworld\_fast\_int status\_new=-1, epiworld\_fast\_int queue=0)
- void rm\_virus (epiworld\_fast\_uint virus\_idx, epiworld\_fast\_int status\_new=-1, epiworld\_fast\_int queue=0)

# Get the rates (multipliers) for the agent

## **Parameters**

v A pointer to a virus.

## Returns

epiworld double

- epiworld\_double get\_susceptibility\_reduction (VirusPtr< TSeq > v)
- epiworld double get\_transmission\_reduction (VirusPtr< TSeq > v)
- epiworld\_double get\_recovery\_enhancer (VirusPtr< TSeq > v)
- epiworld\_double get\_death\_reduction (VirusPtr< TSeq > v)

#### **Friends**

class Model < TSeq >

- class Tool < TSeq >
- class Queue < TSeq >
- void default add virus (Action < TSeq > &a, Model < TSeq > \*m)
- void default\_add\_tool (Action < TSeq > &a, Model < TSeq > \*m)
- void default\_rm\_virus (Action < TSeq > &a, Model < TSeq > \*m)
- void default\_rm\_tool (Action < TSeq > &a, Model < TSeq > \*m)

# 3.7.1 Detailed Description

```
template<typename TSeq = bool> class Person< TSeq >
```

Person (agents)

**Template Parameters** 

TSeq | Sequence type (should match TSeq across the model)

The documentation for this class was generated from the following files:

- include/epiworld/config.hpp
- include/epiworld/person-bones.hpp

# 3.8 PersonTools < TSeq > Class Template Reference

List of tools available for the individual to.

#include <persontools-bones.hpp>

## **Public Member Functions**

- void add\_tool (int date, Tool< TSeq > tool)
- epiworld\_double get\_susceptibility\_reduction (VirusPtr< TSeq > v)
- epiworld\_double get\_transmission\_reduction (VirusPtr< TSeq > v)
- epiworld\_double get\_recovery\_enhancer (VirusPtr< TSeq > v)
- epiworld double get\_death\_reduction (VirusPtr< TSeq > v)
- void set\_susceptibility\_reduction\_mixer (MixerFun < TSeq > fun)
- void set\_transmission\_reduction\_mixer (MixerFun < TSeq > fun)
- void set\_recovery\_enhancer\_mixer (MixerFun < TSeq > fun)
- void set\_death\_reduction\_mixer (MixerFun< TSeq > fun)
- · size t size () const
- Tool < TSeq > & operator() (int i)
- Person< TSeq > \* get\_person ()
- Model < TSeq > \* get\_model ()
- void reset ()
- bool has\_tool (unsigned int t) const
- bool has\_tool (std::string name) const

### **Friends**

- class Person < TSeq >
- class Model < TSeq >

# 3.8.1 Detailed Description

template<typename TSeq = bool> class PersonTools< TSeq >

List of tools available for the individual to.

**Template Parameters** 



The documentation for this class was generated from the following files:

- include/epiworld/config.hpp
- · include/epiworld/persontools-bones.hpp
- include/epiworld/persontools-meat.hpp

# 3.9 PersonViruses < TSeq > Class Template Reference

Set of viruses in host.

#include <personviruses-bones.hpp>

#### **Public Member Functions**

- void add\_virus (epiworld\_fast\_uint new\_status, Virus < TSeq > v)
- size\_t size () const
- int size\_active () const
- Virus < TSeq > & operator() (int i)
- void mutate ()
- void reset ()
- void deactivate (Virus < TSeq > &v)
- Person< TSeq > \* get\_host ()
- bool has\_virus (unsigned int v) const
- bool has\_virus (std::string vname) const

## **Friends**

- class Person < TSeq >
- class Model < TSeq >

# 3.9.1 Detailed Description

template<typename TSeq = bool> class PersonViruses< TSeq >

Set of viruses in host.

# **Template Parameters**

TSeq Type of sequence	
-----------------------	--

The documentation for this class was generated from the following files:

- include/epiworld/person-bones.hpp
- include/epiworld/personviruses-bones.hpp
- include/epiworld/personviruses-meat.hpp

# 3.10 Progress Class Reference

A simple progress bar.

```
#include cprogress.hpp>
```

### **Public Member Functions**

- Progress (int n\_, int width\_)
- void start ()
- · void next ()
- · void end ()

# 3.10.1 Detailed Description

A simple progress bar.

The documentation for this class was generated from the following file:

• include/epiworld/progress.hpp

# 3.11 Queue < TSeq > Class Template Reference

Controls which agents are verified at each step.

```
#include <queue-bones.hpp>
```

### **Public Member Functions**

- void operator+= (Person< TSeq > \*p)
- void operator-= (Person < TSeq > \*p)
- epiworld\_fast\_int operator[] (unsigned int i) const
- void set\_model (Model < TSeq > \*m)
- void update ()

# 3.11.1 Detailed Description

```
template<typename TSeq = bool> class Queue< TSeq >
```

Controls which agents are verified at each step.

The idea is that only agents who are either in an infected state or have an infected neighbor should be checked. Otherwise it makes no sense (no chance to recover or capture the disease).

**Template Parameters** 

TSea	
1004	

The documentation for this class was generated from the following files:

- include/epiworld/model-bones.hpp
- · include/epiworld/queue-bones.hpp

# 3.12 RandGraph Class Reference

# **Public Member Functions**

- RandGraph (int N\_)
- · void init (int s)
- void set\_rand\_engine (std::mt19937 &e)
- epiworld\_double runif ()

The documentation for this class was generated from the following file:

• include/epiworld/random\_graph.hpp

# 3.13 Tool < TSeq > Class Template Reference

Tools for defending the host against the virus.

```
#include <tools-bones.hpp>
```

### **Public Member Functions**

- Tool (std::string name="unknown tool")
- void set\_sequence (TSeq d)
- void set\_sequence\_unique (TSeq d)
- void  $set\_sequence$  (std::shared\_ptr< TSeq > d)
- std::shared ptr< TSeq > get\_sequence ()
- TSeq & get\_sequence\_unique ()
- void set\_name (std::string name)
- std::string get\_name () const
- Person < TSeq > \* get\_person ()
- unsigned int get\_id () const
- void set\_id (int id)
- void set date (int d)
- void set\_status (epiworld\_fast\_int init, epiworld\_fast\_int end)
- void set\_queue (epiworld\_fast\_int init, epiworld\_fast\_int end)

#### Get and set the tool functions

#### **Parameters**

٧	The virus over which to operate
fun	the function to be used

### Returns

epiworld\_double

- epiworld double get\_susceptibility\_reduction (VirusPtr< TSeq > v)
- epiworld double get\_transmission\_reduction (VirusPtr< TSeq > v)
- epiworld double get recovery enhancer (VirusPtr< TSeg > v)
- epiworld\_double get\_death\_reduction (VirusPtr< TSeq > v)
- void set\_susceptibility\_reduction\_fun (ToolFun < TSeq > fun)
- void set\_transmission\_reduction\_fun (ToolFun < TSeq > fun)
- void set\_recovery\_enhancer\_fun (ToolFun < TSeq > fun)
- void set\_death\_reduction\_fun (ToolFun < TSeq > fun)
- void set\_susceptibility\_reduction (epiworld\_double \*prob)
- void set\_transmission\_reduction (epiworld\_double \*prob)
- void set\_recovery\_enhancer (epiworld\_double \*prob)
- void set death reduction (epiworld double \*prob)
- void set susceptibility reduction (epiworld double prob)
- void set transmission reduction (epiworld double prob)
- void set recovery enhancer (epiworld double prob)
- void set\_death\_reduction (epiworld\_double prob)

# **Friends**

- class PersonTools < TSeq >
- class Person < TSeq >
- class Model < TSeq >

### 3.13.1 Detailed Description

template < typename TSeq = bool> class Tool< TSeq >

Tools for defending the host against the virus.

**Template Parameters** 

TSeq Type of sequence

The documentation for this class was generated from the following files:

- · include/epiworld/config.hpp
- include/epiworld/tools-bones.hpp
- include/epiworld/tools-meat.hpp

# 3.14 UserData < TSeq > Class Template Reference

Personalized data by the user.

#include <userdata-bones.hpp>

# **Public Member Functions**

UserData (std::vector< std::string > names)

Construct a new User Data object.

- std::vector< std::string > & get\_names ()
- std::vector< int > & get\_dates ()
- std::vector< epiworld\_double > & get\_data ()
- void **get\_all** (std::vector< std::string > \*names=nullptr, std::vector< int > \*date=nullptr, std::vector<</li>
   epiworld\_double > \*data=nullptr)
- · unsigned int nrow () const
- unsigned int ncol () const
- void write (std::string fn)
- · void print () const

# Append data

### **Parameters**

Х	A vector of length ncol() (if vector), otherwise a epiworld_double.
j	Index of the data point, from 0 to ncol () - 1.

- void add (std::vector< epiworld\_double > x)
- void add (unsigned int j, epiworld\_double x)

### Access data

#### **Parameters**

i	Row (0 through ndays - 1.)
j	Column (0 through ncols()).

# Returns

epiworld\_double&

- epiworld\_double & operator() (unsigned int i, unsigned int j)
- epiworld\_double & operator() (unsigned int i, std::string name)

# **Friends**

- class Model < TSeq >
- class DataBase< TSeq >

# 3.14.1 Detailed Description

template<typename TSeq> class UserData< TSeq>

Personalized data by the user.

# **Template Parameters**

# 3.14.2 Constructor & Destructor Documentation

## 3.14.2.1 UserData()

Construct a new User Data object.

#### **Parameters**

names A vector of names. The length of the vector sets the number of columns to record.

The documentation for this class was generated from the following files:

- include/epiworld/database-bones.hpp
- include/epiworld/userdata-bones.hpp
- include/epiworld/userdata-meat.hpp

# 3.15 vecHasher< T > Struct Template Reference

Vector hasher.

```
#include <misc.hpp>
```

# **Public Member Functions**

• std::size\_t **operator()** (std::vector< T > const &dat) const noexcept

# 3.15.1 Detailed Description

```
template<typename T> struct vecHasher< T>
```

Vector hasher.

### **Template Parameters**



The documentation for this struct was generated from the following file:

· include/epiworld/misc.hpp

# 3.16 Virus < TSeq > Class Template Reference

### Virus.

```
#include <virus-bones.hpp>
```

# **Public Member Functions**

- Virus (std::string name="unknown virus")
- · void mutate ()
- void set\_mutation (MutFun < TSeq > fun)
- const TSeq \* get\_sequence ()
- void **set\_sequence** (TSeq sequence)
- Person< TSeq > \* get\_host ()
- Model < TSeq > \* get\_model ()
- void set\_date (int d)
- int get\_date () const
- void set\_id (int idx)
- int get\_id () const
- void set\_name (std::string name)
- std::string get\_name () const
- std::vector< epiworld\_double > & get\_data ()
- void set\_status (epiworld\_fast\_int init, epiworld\_fast\_int end)

## Get and set the tool functions

#### **Parameters**

V	The virus over which to operate
fun	the function to be used

#### Returns

epiworld\_double

- epiworld double get prob infecting ()
- epiworld\_double get\_prob\_recovery ()
- epiworld double get prob death ()
- void post recovery ()
- void set\_post\_recovery (PostRecoveryFun< TSeq > fun)
- void **set\_post\_immunity** (epiworld\_double prob)
- void set\_post\_immunity (epiworld\_double \*prob)

- void set\_prob\_infecting\_fun (VirusFun < TSeq > fun)
- void set\_prob\_recovery\_fun (VirusFun < TSeq > fun)
- void set\_prob\_death\_fun (VirusFun < TSeq > fun)
- void set\_prob\_infecting (epiworld\_double \*prob)
- void set\_prob\_recovery (epiworld\_double \*prob)
- void set\_prob\_death (epiworld\_double \*prob)
- void set prob infecting (epiworld double prob)
- void set\_prob\_recovery (epiworld\_double prob)
- void set\_prob\_death (epiworld\_double prob)

# **Friends**

- class Person < TSeq >
- class Model < TSeq >
- class PersonViruses < TSeq >
- class DataBase< TSeq >

# 3.16.1 Detailed Description

template<typename TSeq = bool> class Virus< TSeq >

# Virus.

**Template Parameters** 

TSeq	

Raw transmisibility of a virus should be a function of its genetic sequence. Nonetheless, transmisibility can be reduced as a result of having one or more tools to fight the virus. Because of this, transmisibility should be a function of the host.

The documentation for this class was generated from the following files:

- · include/epiworld/config.hpp
- include/epiworld/virus-bones.hpp
- include/epiworld/virus-meat.hpp

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