epiworld

0.8.2

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1 Example: 00-hello-world	1
2 Benchmarking	3
3 Contributor Code of Conduct	5
4 ModelMeaslesMixing	7
4.1 Overview	7
4.2 Disease States	7
4.3 Key Features	8
4.3.1 Disease Progression	8
4.3.2 Population Mixing	8
4.3.3 Public Health Measures	8
4.4 Parameters	8
4.4.1 Disease Parameters	8
4.4.2 Detection and Isolation	9
4.4.3 Contact Tracing and Quarantine	9
4.4.4 Vaccination	9
4.5 Usage Example	9
4.6 Differences from Other Models	10
4.6.1 vs. ModelMeaslesSchool	10
4.6.2 vs. ModelSEIRMixingQuarantine	10
4.7 Testing	10
5 ModelMeaslesMixingRiskQuarantine	11
5.1 Overview	11
5.1.1 Key Features	11
5.2 Disease States	11
5.3 Risk Classification System	12
5.3.1 High Risk	12
5.3.2 Medium Risk	12
5.3.3 Low Risk	12
5.4 Enhanced Detection	13
5.5 Parameters	13
5.5.1 Disease Parameters	13
5.5.2 Detection and Isolation	13
5.5.3 Risk-based Quarantine	13
5.5.4 Contact Tracing	14
5.5.5 Vaccination	14
5.6 Usage Example	14
5.7 Differences from Other Models	14
5.7.1 vs. ModelMeaslesMixing	14
5.7.2 vs. ModelMeaslesSchool	15
5.8 Testing	15
	_

5.9 Applications	15
5.10 Implementation Notes	15
6 epiworld c++ template library	17
6.1 Main features	17
6.2 Algorithm	17
6.3 Hello world (C++)	18
6.4 Surveillance simulation	18
6.4.1 Preliminary results	19
6.4.2 Cases detected	20
7 General parameters	21
7.1 Compartmental Models	21
7.1.1 SIR Model	21
7.1.2 SEIR Model	22
7.2 Agent-Based Model Approach	22
7.2.1 Mathematical preliminaries	23
7.2.2 Simulation study	23
7.3 Comparing ABM with Compartmental Models	23
7.3.1 SIR	23
7.3.2 SEIR	24
7.3.3 Rates	24
8 MIT License	25
9 model1	27
10 Mixing probabilities in connected model	29
10.1 Case 1: No grouping	
10.2 Case 2: Grouping	
11 EPI Simulator	31
11.1 Disease dynamics	
11.2 Network dynamics	
11.3 Contagion dynamics	
11.4 Time dynamics	
11.5 Updating agent's status	
11.5.1 Other parameters	32
12 Fix for rewire_degseq Bug	33
12.1 Problem	
12.2 Root Cause	
12.3 Solution	
12.3.1 AdjList Version Changes	34
12.3.2 Agent Vector Version Changes	34

12.4 Testing	
12.4.1 Test Results	
12.5 Files Modified	
12.6 Notes	34
13 Namespace Index	35
13.1 Namespace List	35
14 Hierarchical Index	37
14.1 Class Hierarchy	
15 Class Index	41
15.1 Class List	41
16 File Index	45
16.1 File List	45
17 Namespace Documentation	47
17.1 epiworld::sampler Namespace Reference	47
17.1.1 Detailed Description	47
17.1.2 Function Documentation	47
17.1.2.1 make_sample_virus_neighbors()	47
17.1.2.2 make_update_susceptible()	48
17.1.2.3 sample_virus_single()	48
17.2 sampler Namespace Reference	50
17.2.1 Detailed Description	50
17.2.2 Function Documentation	50
17.2.2.1 make_sample_virus_neighbors()	50
17.2.2.2 make_update_susceptible()	51
17.2.2.3 sample_virus_single()	51
18 Class Documentation	55
18.1 AdjList Class Reference	
18.1.1 Constructor & Destructor Documentation	
18.1.1.1 AdjList()	
18.1.2 Member Function Documentation	56
18.1.2.1 read_edgelist()	56
18.2 epiworld::AdjList Class Reference	56
18.2.1 Constructor & Destructor Documentation	57
18.2.1.1 AdjList()	57
18.2.2 Member Function Documentation	57
18.2.2.1 read_edgelist()	57
18.3 Agent < TSeq > Class Template Reference	58
18.3.1 Detailed Description	

18.3.2 Member Function Documentation	60
18.3.2.1 operator()()	60
18.3.2.2 swap_neighbors()	62
18.3.3 Friends And Related Function Documentation	62
18.3.3.1 default_rm_entity	62
18.4 epiworld::Agent < TSeq > Class Template Reference	63
18.4.1 Detailed Description	65
18.4.2 Member Function Documentation	65
18.4.2.1 operator()()	65
18.4.2.2 swap_neighbors()	66
18.4.3 Friends And Related Function Documentation	66
18.4.3.1 default_rm_entity	66
18.5 AgentsSample < TSeq > Class Template Reference	66
18.5.1 Detailed Description	67
18.5.2 Constructor & Destructor Documentation	67
18.5.2.1 AgentsSample()	67
18.6 epiworld::AgentsSample < TSeq > Class Template Reference	68
18.6.1 Detailed Description	68
18.6.2 Constructor & Destructor Documentation	69
18.6.2.1 AgentsSample()	69
18.7 DataBase < TSeq > Class Template Reference	69
18.7.1 Detailed Description	71
18.7.2 Member Function Documentation	72
18.7.2.1 get_generation_time()	72
18.7.2.2 get_reproductive_number()	72
18.7.2.3 get_transition_probability()	73
18.7.2.4 get_transmissions()	73
18.7.2.5 operator==() [1/3]	73
18.7.2.6 operator==() [2/3]	74
<b>18.7.2.7</b> operator==() [3/3]	74
18.7.2.8 record_virus()	74
$18.8 \; epiworld:: DataBase < TSeq > Class \; Template \; Reference \qquad $	75
18.8.1 Detailed Description	77
18.8.2 Member Function Documentation	77
18.8.2.1 get_generation_time()	77
18.8.2.2 get_reproductive_number()	77
18.8.2.3 get_transition_probability()	78
18.8.2.4 get_transmissions()	78
18.8.2.5 operator==()	79
18.8.2.6 record_virus()	79
18.9 Entities < TSeq > Class Template Reference	79
18.9.1 Detailed Description	80

18.10 epiworld::Entities< TSeq > Class Template Reference	80
18.10.1 Detailed Description	81
18.11 Entities_const< TSeq > Class Template Reference	81
18.11.1 Detailed Description	81
18.12 epiworld::Entities_const< TSeq > Class Template Reference	82
18.12.1 Detailed Description	82
18.13 Entity < TSeq > Class Template Reference	83
18.13.1 Constructor & Destructor Documentation	83
18.13.1.1 Entity()	83
18.13.2 Friends And Related Function Documentation	84
18.13.2.1 default_rm_entity	84
18.14 epiworld::Entity< TSeq > Class Template Reference	84
18.14.1 Constructor & Destructor Documentation	85
18.14.1.1 Entity()	85
18.14.2 Friends And Related Function Documentation	85
18.14.2.1 default_rm_entity	86
18.15 epiworld::Event < TSeq > Struct Template Reference	86
18.15.1 Detailed Description	86
18.15.2 Constructor & Destructor Documentation	87
18.15.2.1 Event()	87
18.16 Event < TSeq > Struct Template Reference	87
18.16.1 Detailed Description	88
18.16.2 Constructor & Destructor Documentation	89
18.16.2.1 Event()	89
18.17 epiworld::GlobalEvent< TSeq > Class Template Reference	89
18.17.1 Detailed Description	90
18.17.2 Constructor & Destructor Documentation	90
18.17.2.1 GlobalEvent()	90
18.18 GlobalEvent< TSeq > Class Template Reference	91
18.18.1 Detailed Description	91
18.18.2 Constructor & Destructor Documentation	91
18.18.2.1 GlobalEvent()	91
18.19 epiworld::LFMCMC < TData > Class Template Reference	92
18.19.1 Detailed Description	93
18.20 LFMCMC< TData > Class Template Reference	93
18.20.1 Detailed Description	94
18.21 epiworld::Model < TSeq > Class Template Reference	95
18.21.1 Detailed Description	103
18.21.2 Member Function Documentation	103
18.21.2.1 add_globalevent()	103
18.21.2.2 clone_ptr()	104
18.21.2.3 draw()	104

18.21.2.4 events_add()	 104
18.21.2.5 events_run()	 105
18.21.2.6 load_agents_entities_ties()	 105
18.21.2.7 reset()	 106
18.21.2.8 run_multiple()	 106
18.21.2.9 set_agents_data()	 106
18.21.2.10 set_name()	 107
18.21.2.11 write_data()	 107
18.21.3 Member Data Documentation	 108
18.21.3.1 initial_states_fun	 108
18.21.3.2 rbinomd	 108
18.21.3.3 rexpd	 108
18.21.3.4 rgammad	 109
18.21.3.5 rgeomd	 109
18.21.3.6 rlognormald	 109
18.21.3.7 rnbinomd	 109
18.21.3.8 rnormd	 109
18.21.3.9 rpoissd	 110
18.21.3.10 runifd	 110
18.21.3.11 time_elapsed	 110
18.22 Model < TSeq > Class Template Reference	 110
18.22.1 Detailed Description	 118
18.22.2 Member Function Documentation	 119
18.22.2.1 add_globalevent()	 119
18.22.2.2 clone_ptr()	 119
18.22.2.3 draw()	 119
18.22.2.4 events_add()	 120
18.22.2.5 events_run()	 120
18.22.2.6 load_agents_entities_ties()	 122
18.22.2.7 reset()	 122
18.22.2.8 run_multiple()	 123
18.22.2.9 set_agents_data()	 123
18.22.2.10 set_name()	 123
18.22.2.11 write_data()	 124
18.22.3 Member Data Documentation	 124
18.22.3.1 initial_states_fun	 124
18.22.3.2 rbinomd	 124
18.22.3.3 rexpd	 125
18.22.3.4 rgammad	 125
18.22.3.5 rgeomd	 125
18.22.3.6 rlognormald	 125
18.22.3.7 rnbinomd	 125

18.22.3.8 rnormd	126
18.22.3.9 rpoissd	126
18.22.3.10 runifd	126
18.22.3.11 time_elapsed	126
18.23 epiworld::ModelDiagram Class Reference	127
18.24 ModelDiagram Class Reference	127
18.25 epiworld::epimodels::ModelDiffNet< TSeq > Class Template Reference	127
18.25.1 Detailed Description	128
18.26 ModelDiffNet< TSeq > Class Template Reference	129
18.26.1 Detailed Description	130
18.27 epiworld::epimodels::ModelMeaslesMixing< TSeq > Class Template Reference	130
18.27.1 Detailed Description	132
18.27.2 Constructor & Destructor Documentation	133
<b>18.27.2.1 ModelMeaslesMixing()</b> [1/2]	133
18.27.2.2 ModelMeaslesMixing() [2/2]	135
18.27.3 Member Function Documentation	136
18.27.3.1 clone_ptr()	136
18.27.3.2 get_agent_quarantine_triggered()	136
18.27.3.3 get_contact_matrix()	137
18.27.3.4 get_isolation_willingness()	137
18.27.3.5 get_quarantine_willingness()	137
18.27.3.6 initial_states()	137
18.27.3.7 run()	138
18.27.3.8 set_contact_matrix()	138
18.28 ModelMeaslesMixing < TSeq > Class Template Reference	139
18.28.1 Detailed Description	141
18.28.2 Constructor & Destructor Documentation	142
18.28.2.1 ModelMeaslesMixing() [1/2]	142
18.28.2.2 ModelMeaslesMixing() [2/2]	143
18.28.3 Member Function Documentation	144
18.28.3.1 clone_ptr()	144
18.28.3.2 get_agent_quarantine_triggered()	145
18.28.3.3 get_contact_matrix()	145
18.28.3.4 get_isolation_willingness()	145
18.28.3.5 get_quarantine_willingness()	146
18.28.3.6 initial_states()	146
18.28.3.7 run()	146
18.28.3.8 set_contact_matrix()	147
$18.29 \; epiworld:: epimodels:: Model Measles Mixing Risk Quarantine < TSeq > Class \; Template \; Reference  .$	147
18.29.1 Detailed Description	149
18.29.2 Constructor & Destructor Documentation	150
18 29 2 1 ModelMeaclesMixingRickOuaranting() [1/2]	150

18.29.2.2 ModelMeaslesMixingRiskQuarantine() [2/2]
18.29.3 Member Function Documentation
18.29.3.1 clone_ptr()
18.29.3.2 get_agent_quarantine_triggered()
18.29.3.3 get_contact_matrix()
18.29.3.4 get_isolation_willingness()
18.29.3.5 get_quarantine_risk_levels()
18.29.3.6 get_quarantine_willingness()
18.29.3.7 initial_states()
18.29.3.8 run()
18.29.3.9 set_contact_matrix()
18.30 ModelMeaslesMixingRiskQuarantine < TSeq > Class Template Reference
18.30.1 Detailed Description
18.30.2 Constructor & Destructor Documentation
18.30.2.1 ModelMeaslesMixingRiskQuarantine() [1/2]
18.30.2.2 ModelMeaslesMixingRiskQuarantine() [2/2]
18.30.3 Member Function Documentation
18.30.3.1 clone_ptr()
18.30.3.2 get_contact_matrix()
18.30.3.3 get_isolation_willingness()
18.30.3.4 get_quarantine_willingness()
18.30.3.5 initial_states()
18.30.3.6 run()
18.30.3.7 set_contact_matrix()
18.31 epiworld::epimodels::ModelMeaslesSchool < TSeq > Class Template Reference
18.31.1 Detailed Description
18.31.2 Constructor & Destructor Documentation
18.31.2.1 ModelMeaslesSchool()
18.31.3 Member Function Documentation
18.31.3.1 clone_ptr()
18.31.3.2 quarantine_agents()
18.31.3.3 reset()
18.32 ModelMeaslesSchool < TSeq > Class Template Reference
18.32.1 Detailed Description
18.32.2 Constructor & Destructor Documentation
18.32.2.1 ModelMeaslesSchool()
18.32.3 Member Function Documentation
18.32.3.1 clone_ptr()
18.32.3.2 quarantine_agents()
18.32.3.3 reset()
18.33 epiworld::epimodels::ModelSEIR< TSeq > Class Template Reference
18.33.1 Detailed Description

18.33.2 Member Function Documentation
18.33.2.1 initial_states()
18.33.3 Member Data Documentation
18.33.3.1 update_exposed_seir
18.33.3.2 update_infected_seir
18.34 ModelSEIR < TSeq > Class Template Reference
18.34.1 Detailed Description
18.34.2 Member Function Documentation
18.34.2.1 initial_states()
18.34.3 Member Data Documentation
18.34.3.1 update_exposed_seir
18.34.3.2 update_infected_seir
18.35 epiworld::epimodels::ModelSEIRCONN< TSeq > Class Template Reference
18.35.1 Constructor & Destructor Documentation
18.35.1.1 ModelSEIRCONN()
18.35.2 Member Function Documentation
18.35.2.1 clone_ptr()
18.35.2.2 initial_states()
18.35.2.3 reset()
18.36 ModelSEIRCONN < TSeq > Class Template Reference
18.36.1 Constructor & Destructor Documentation
18.36.1.1 ModelSEIRCONN()
18.36.2 Member Function Documentation
18.36.2.1 clone_ptr()
18.36.2.2 initial_states()
18.36.2.3 reset()
18.37 epiworld::epimodels::ModelSEIRD< TSeq > Class Template Reference
18.37.1 Detailed Description
18.37.2 Constructor & Destructor Documentation
18.37.2.1 ModelSEIRD() [1/2]
18.37.2.2 ModelSEIRD() [2/2]
18.37.3 Member Data Documentation
18.37.3.1 update_exposed_seir
18.38 ModelSEIRD< TSeq > Class Template Reference
18.38.1 Detailed Description
18.38.2 Constructor & Destructor Documentation
18.38.2.1 ModelSEIRD() [1/2]
18.38.2.2 ModelSEIRD() [2/2]
18.38.3 Member Data Documentation
18.38.3.1 update_exposed_seir
18.39 epiworld::epimodels::ModelSEIRDCONN< TSeq > Class Template Reference
18.39.1 Constructor & Destructor Documentation

18.39.1.1 ModelSEIRDCONN()	191
18.39.2 Member Function Documentation	192
18.39.2.1 clone_ptr()	192
18.39.2.2 initial_states()	192
18.39.2.3 reset()	192
18.40 ModelSEIRDCONN < TSeq > Class Template Reference	193
18.40.1 Constructor & Destructor Documentation	194
18.40.1.1 ModelSEIRDCONN()	194
18.40.2 Member Function Documentation	195
18.40.2.1 clone_ptr()	195
18.40.2.2 initial_states()	195
18.40.2.3 reset()	196
18.41 epiworld::epimodels::ModelSEIRMixing< TSeq > Class Template Reference	196
18.41.1 Constructor & Destructor Documentation	197
<b>18.41.1.1 ModelSEIRMixing()</b> [1/2]	198
<b>18.41.1.2 ModelSEIRMixing()</b> [2/2]	198
18.41.2 Member Function Documentation	199
18.41.2.1 clone_ptr()	199
18.41.2.2 initial_states()	199
18.41.2.3 reset()	200
18.42 ModelSEIRMixing < TSeq > Class Template Reference	200
18.42.1 Constructor & Destructor Documentation	201
<b>18.42.1.1 ModelSEIRMixing()</b> [1/2]	201
<b>18.42.1.2</b> ModelSEIRMixing() [2/2]	202
18.42.2 Member Function Documentation	202
18.42.2.1 clone_ptr()	203
18.42.2.2 initial_states()	203
18.42.2.3 reset()	203
$18.43\ epiworld:: epimodels:: Model SEIR Mixing Quarantine < TSeq > Class\ Template\ Reference \\ \ldots \\ \ldots \\ \vdots$	204
18.43.1 Detailed Description	206
18.43.2 Constructor & Destructor Documentation	206
18.43.2.1 ModelSEIRMixingQuarantine() [1/2]	207
18.43.2.2 ModelSEIRMixingQuarantine() [2/2]	208
18.43.3 Member Function Documentation	209
18.43.3.1 clone_ptr()	209
18.43.3.2 get_agent_quarantine_triggered()	209
18.43.3.3 get_contact_matrix()	210
18.43.3.4 get_isolation_willingness()	210
18.43.3.5 get_quarantine_willingness()	210
18.43.3.6 initial_states()	210
18.43.3.7 run()	
18.43.3.8 set_contact_matrix()	211

18.44 ModelSEIRMixingQuarantine < TSeq > Class Template Reference
18.44.1 Detailed Description
18.44.2 Constructor & Destructor Documentation
18.44.2.1 ModelSEIRMixingQuarantine() [1/2]
18.44.2.2 ModelSEIRMixingQuarantine() [2/2]
18.44.3 Member Function Documentation
18.44.3.1 clone_ptr()
18.44.3.2 get_agent_quarantine_triggered()
18.44.3.3 get_contact_matrix()
18.44.3.4 get_isolation_willingness()
18.44.3.5 get_quarantine_willingness()
18.44.3.6 initial_states()
18.44.3.7 run()
18.44.3.8 set_contact_matrix()
$18.45 \; epiworld:: epimodels:: Model SIR < TSeq > Class \; Template \; Reference \; . \; . \; . \; . \; . \; . \; . \; . \; . \; $
18.45.1 Detailed Description
18.45.2 Member Function Documentation
18.45.2.1 initial_states()
18.46 ModelSIR < TSeq > Class Template Reference
18.46.1 Detailed Description
18.46.2 Member Function Documentation
18.46.2.1 initial_states()
$18.47 \; epiworld:: epimodels:: Model SIRCONN < TS eq > Class \; Template \; Reference \; \dots \; \dots \; \dots \; 22441 = 1000 \; Model \; Mod$
18.47.1 Constructor & Destructor Documentation
18.47.1.1 ModelSIRCONN()
18.47.2 Member Function Documentation
18.47.2.1 clone_ptr()
18.47.2.2 get_n_infected()
18.47.2.3 initial_states()
18.47.2.4 reset()
18.48 ModelSIRCONN < TSeq > Class Template Reference
18.48.1 Constructor & Destructor Documentation
18.48.1.1 ModelSIRCONN()
18.48.2 Member Function Documentation
18.48.2.1 clone_ptr()
18.48.2.2 get_n_infected()
18.48.2.3 initial_states()
18.48.2.4 reset()
$18.49 \; epiworld:: epimodels:: Model SIRD < TSeq > Class \; Template \; Reference \; . \; . \; . \; . \; . \; . \; . \; . \; . \; $
18.49.1 Detailed Description
18.49.2 Constructor & Destructor Documentation
18.49.2.1 ModelSIRD()

18.49.3 Member Function Documentation	33
18.49.3.1 initial_states()	33
18.50 ModelSIRD < TSeq > Class Template Reference	33
18.50.1 Detailed Description	34
18.50.2 Constructor & Destructor Documentation	35
18.50.2.1 ModelSIRD()	35
18.50.3 Member Function Documentation	35
18.50.3.1 initial_states()	35
18.51 epiworld::epimodels::ModelSIRDCONN< TSeq > Class Template Reference	36
18.51.1 Constructor & Destructor Documentation	37
18.51.1.1 ModelSIRDCONN()	37
18.51.2 Member Function Documentation	38
18.51.2.1 clone_ptr()	38
18.51.2.2 reset()	38
18.52 ModelSIRDCONN< TSeq > Class Template Reference	39
18.52.1 Constructor & Destructor Documentation	40
18.52.1.1 ModelSIRDCONN()	40
18.52.2 Member Function Documentation	40
18.52.2.1 clone_ptr()	40
18.52.2.2 reset()	41
18.53 epiworld::epimodels::ModelSIRLogit< TSeq > Class Template Reference	41
18.53.1 Detailed Description	42
18.53.2 Constructor & Destructor Documentation	43
18.53.2.1 ModelSIRLogit()	43
18.53.3 Member Function Documentation	44
18.53.3.1 clone_ptr()	44
18.53.3.2 reset()	44
18.54 ModelSIRLogit < TSeq > Class Template Reference	45
18.54.1 Detailed Description	46
18.54.2 Constructor & Destructor Documentation	46
18.54.2.1 ModelSIRLogit()	46
18.54.3 Member Function Documentation	47
18.54.3.1 clone_ptr()	47
18.54.3.2 reset()	47
18.55 epiworld::epimodels::ModelSIRMixing < TSeq > Class Template Reference	48
18.55.1 Constructor & Destructor Documentation	49
18.55.1.1 ModelSIRMixing() [1/2]	49
18.55.1.2 ModelSIRMixing() [2/2]	50
18.55.2 Member Function Documentation	50
18.55.2.1 clone_ptr()	51
18.55.2.2 initial_states()	51
18.55.2.3 reset()	51

18.56 ModelSIRMixing < TSeq > Class Template Reference
18.56.1 Constructor & Destructor Documentation
18.56.1.1 ModelSIRMixing() [1/2]
18.56.1.2 ModelSIRMixing() [2/2]
18.56.2 Member Function Documentation
18.56.2.1 clone_ptr()
18.56.2.2 initial_states()
18.56.2.3 reset()
18.57 epiworld::epimodels::ModelSIS< TSeq > Class Template Reference
18.57.1 Detailed Description
18.58 ModelSIS< TSeq > Class Template Reference
18.58.1 Detailed Description
18.59 epiworld::epimodels::ModelSISD< TSeq > Class Template Reference
18.59.1 Detailed Description
18.60 ModelSISD< TSeq > Class Template Reference
18.60.1 Detailed Description
18.61 epiworld::epimodels::ModelSURV < TSeq > Class Template Reference
18.61.1 Member Function Documentation
18.61.1.1 reset()
18.62 ModelSURV < TSeq > Class Template Reference
18.62.1 Member Function Documentation
18.62.1.1 reset()
18.63 Network< Nettype, Nodetype, Edgetype > Class Template Reference
18.64 epiworld::PersonTools < TSeq > Class Template Reference
18.65 PersonTools < TSeq > Class Template Reference
18.66 epiworld::Progress Class Reference
18.66.1 Detailed Description
18.67 Progress Class Reference
18.67.1 Detailed Description
18.68 epiworld::Queue < TSeq > Class Template Reference
18.68.1 Detailed Description
18.69 Queue < TSeq > Class Template Reference
18.69.1 Detailed Description
18.70 RandGraph Class Reference
18.71 epiworld::SAMPLETYPE Class Reference
18.72 SAMPLETYPE Class Reference
18.73 SimpleAdjList Struct Reference
18.74 SimpleModel Struct Reference
18.75 epiworld::Tool < TSeq > Class Template Reference
18.75.1 Detailed Description
18.76 Tool < TSeq > Class Template Reference
18.76.1 Detailed Description 27

	$18.77 \; epiworld:: ToolFunctions < TSeq > Class \; Template \; Reference \; . \; . \; . \; . \; . \; . \; . \; . \; . \; $	275
	18.77.1 Detailed Description	275
	18.78 ToolFunctions < TSeq > Class Template Reference	276
	18.78.1 Detailed Description	276
	18.79 epiworld::Tools< TSeq > Class Template Reference	276
	18.79.1 Detailed Description	276
	18.80 Tools < TSeq > Class Template Reference	277
	18.80.1 Detailed Description	277
	18.81 epiworld::Tools_const< TSeq > Class Template Reference	278
	18.81.1 Detailed Description	278
	18.82 Tools_const < TSeq > Class Template Reference	278
	18.82.1 Detailed Description	279
	$18.83 \; epiworld:: User Data < TSeq > Class \; Template \; Reference \; \ldots \; $	279
	18.83.1 Detailed Description	280
	18.83.2 Constructor & Destructor Documentation	281
	18.83.2.1 UserData()	281
	18.84 UserData < TSeq > Class Template Reference	281
	18.84.1 Detailed Description	282
	18.84.2 Constructor & Destructor Documentation	282
	18.84.2.1 UserData()	282
	$18.85 \; epiworld:: vec Hasher < T > Struct \; Template \; Reference \qquad $	283
	18.85.1 Detailed Description	283
	18.86 vecHasher < T > Struct Template Reference	283
	18.86.1 Detailed Description	283
	18.87 epiworld::Virus < TSeq > Class Template Reference	284
	18.87.1 Detailed Description	285
	18.88 Virus < TSeq > Class Template Reference	286
	18.88.1 Detailed Description	288
	$18.89 \; epiworld:: Viruses < TSeq > Class \; Template \; Reference \; \dots $	288
	18.89.1 Detailed Description	288
	18.90 Viruses < TSeq > Class Template Reference	289
	18.90.1 Detailed Description	289
	18.91 epiworld::Viruses_const< TSeq > Class Template Reference	290
	18.91.1 Detailed Description	290
	18.92 Viruses_const< TSeq > Class Template Reference	290
	18.92.1 Detailed Description	291
	18.93 epiworld::VirusFunctions < TSeq > Class Template Reference	291
	18.94 VirusFunctions < TSeq > Class Template Reference	292
19	File Documentation	293
	19.1 include/epiworld/agent-meat-state.hpp File Reference	293
	19.1.1 Detailed Description	

19.2 include/epiworld/models/measlesmixing.hpp File Reference	295
19.2.1 Detailed Description	295
19.2.2 Macro Definition Documentation	295
19.2.2.1 GET_MODEL	296
19.2.2.2 SAMPLE_FROM_PROBS	296
19.3 include/epiworld/models/measlesmixingriskquarantine.hpp File Reference	296
19.3.1 Detailed Description	297
19.3.2 Macro Definition Documentation	297
19.3.2.1 GET_MODEL	297
19.3.2.2 SAMPLE_FROM_PROBS	297
19.4 include/epiworld/models/seirmixingquarantine.hpp File Reference	298
19.4.1 Detailed Description	298
19.4.2 Macro Definition Documentation	298
19.4.2.1 GET_MODEL	299
19.4.2.2 SAMPLE_FROM_PROBS	299
Index	301
HIUCA	30 I

# Example: 00-hello-world

#### Output from the program:

```
Running the model...
SIMULATION STUDY
Name of the model
Population size : 10000
Agents' data : (none)
Number of entities : 0
Days (duration) : 100 (of 100)
Number of viruses : 1
Last run elapsed t : 17.00ms
Last run speed : 56.33 million agents x day / second Rewiring : off
Rewiring Global events:
 (none)
Virus(es):
 - covid 19
Tool(s):
 - vaccine
Model parameters:
 (none)
Distribution of the population at time 100:
 - (0) Susceptible: 9950 -> 0
- (1) Exposed: 50 -> 0
- (2) Recovered: 0 -> 9399
- (3) Removed: 0 -> 601
Transition Probabilities:
- Susceptible 0.92 0.08 - - Exposed - 0.85 0.14 0.01 - Recovered - 1.00 - 1.00
 - Removed
```

# **Benchmarking**

Here we keep a list of scenarios where we compare epiworld with other ABM simulation engines. Although the comparison is made at the speed level, we also list features of capabilities and main differences between the engines.

4 Benchmarking

## **Contributor Code of Conduct**

As contributors and maintainers of this project, we pledge to respect all people who contribute through reporting issues, posting feature requests, updating documentation, submitting pull requests or patches, and other activities.

We are committed to making participation in this project a harassment-free experience for everyone, regardless of level of experience, gender, gender identity and expression, sexual orientation, disability, personal appearance, body size, race, ethnicity, age, or religion.

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# ModelMeaslesMixing

A comprehensive epidemiological model that combines measles-specific disease progression with population mixing and quarantine measures.

#### 4.1 Overview

ModelMeaslesMixing implements a measles transmission model that extends the SEIR framework with:

- $\bullet \ \ \text{Measles-specific disease progression} : \mathsf{Susceptible} \to \mathsf{Exposed} \to \mathsf{Prodromal} \to \mathsf{Rash} \to \mathsf{Recovered}$
- · Population mixing: Uses contact matrices to model heterogeneous mixing between population groups
- Infectious period: Agents are infectious during the Prodromal state
- · Detection and isolation: Detection occurs during the Rash state when symptoms become visible
- Contact tracing: Comprehensive contact tracing with configurable success rates
- Quarantine measures: Multiple quarantine states for exposed contacts
- · Vaccination: Vaccine distribution with configurable efficacy and recovery enhancement

#### 4.2 Disease States

The model includes 13 distinct states:

- 1. Susceptible Can become infected
- 2. Exposed Infected but not yet infectious (incubation period)
- 3. Prodromal Infectious individuals (replaces "Infected" in standard SEIR)
- 4. Rash Non-infectious with visible symptoms (detection occurs here)
- 5. Isolated Detected individuals in self-isolation
- 6. Isolated Recovered Recovered individuals still in isolation

8 ModelMeaslesMixing

- 7. Detected Hospitalized Hospitalized individuals who were contact-traced
- 8. Quarantined Exposed Exposed individuals in quarantine
- 9. Quarantined Susceptible Susceptible individuals in quarantine
- 10. Quarantined Prodromal Prodromal individuals in quarantine
- 11. Quarantined Recovered Recovered individuals in quarantine
- 12. Hospitalized Individuals requiring hospital care
- 13. Recovered Immune individuals

### 4.3 Key Features

#### 4.3.1 Disease Progression

- · Incubation Period: Time from exposure to becoming infectious
- · Prodromal Period: Duration of infectiousness before rash appears
- Rash Period: Duration of visible symptoms (detection window)
- Detection: Probabilistic detection during rash period
- Hospitalization: Some individuals require hospitalization

#### 4.3.2 Population Mixing

- · Contact Matrices: Define mixing patterns between population groups
- · Heterogeneous Mixing: Different contact rates between groups
- Scalable: Supports multiple population entities

#### 4.3.3 Public Health Measures

- · Contact Tracing: Trace contacts of detected individuals
- · Quarantine: Quarantine exposed contacts with configurable compliance
- · Isolation: Isolate detected cases with configurable willingness
- · Vaccination: Reduce susceptibility and enhance recovery

#### 4.4 Parameters

#### 4.4.1 Disease Parameters

- contact\_rate: Average number of contacts per day
- transmission\_rate: Probability of transmission per contact
- incubation\_period: Average incubation period (days)
- prodromal\_period: Average prodromal period (days)
- rash\_period: Average rash period (days)
- hospitalization\_rate: Probability of hospitalization
- hospitalization\_period: Average hospitalization duration (days)

4.5 Usage Example 9

#### 4.4.2 Detection and Isolation

- days\_undetected: Average time until detection during rash period
- isolation period: Duration of isolation for detected cases
- isolation\_willingness: Proportion willing to self-isolate

#### 4.4.3 Contact Tracing and Quarantine

- contact\_tracing\_success\_rate: Probability of successfully tracing a contact
- contact\_tracing\_days\_prior: Number of days to trace back
- quarantine\_period: Duration of quarantine for contacts
- quarantine\_willingness: Proportion willing to quarantine

#### 4.4.4 Vaccination

- prop\_vaccinated: Proportion of population vaccinated
- vax\_efficacy: Vaccine efficacy (reduction in susceptibility)
- vax\_reduction\_recovery\_rate: Recovery enhancement from vaccination

## 4.5 Usage Example

```
#include "epiworld.hpp"
// Contact matrix for single homogeneous population
std::vector<double> contact_matrix = {1.0};
// Create the model
epimodels::ModelMeaslesMixing<> model(
    "Measles",
                     // Virus name
// Population size
    1000.
                     // Initial prevalence
    0.005,
                     // Contact rate
    4.0,
    0.9,
                     // Transmission rate
    0.95,
                     // Vaccine efficacy
                      // Vaccine recovery enhancement
    0.5,
                      // Incubation period
    10.0.
                      // Prodromal period
    4.0,
                      // Rash period
    5.0,
    contact_matrix,
                     // Contact matrix
    0.1,
                     // Hospitalization rate
    7.0,
                      // Hospitalization period
                      // Days undetected
    2.0,
                     // Quarantine period
    14,
    0.8,
                     // Quarantine willingness
    0.9,
                     // Isolation willingness
                      // Isolation period
    10,
                      // Vaccination rate
    0.7,
                      // Contact tracing success rate
    0.8,
                      // Contact tracing days prior
    3u
);
// Add population entity
model.add_entity(Entity<>("Population", dist_factory<>(0, 1000)));
// Run simulation
model.run(60, 123);
model.print();
```

10 ModelMeaslesMixing

#### 4.6 Differences from Other Models

#### 4.6.1 vs. ModelMeaslesSchool

- Population Mixing: Adds contact matrices for heterogeneous mixing
- Contact Tracing: Enhanced contact tracing with configurable success rates
- · Scalability: Supports multiple population entities

#### 4.6.2 vs. ModelSEIRMixingQuarantine

- · Disease-Specific States: Uses Prodromal/Rash instead of generic Infected
- Detection Timing: Detection occurs during Rash state, not Infected state
- · Infectious Period: Only Prodromal individuals are infectious
- Vaccination: Includes vaccine distribution and efficacy

### 4.7 Testing

The model includes comprehensive tests that verify:

- · Correct transition probabilities between states
- · Proper detection and quarantine mechanics
- · Population mixing functionality
- · Reproductive number calculations
- · All quarantine states function correctly

See tests/19-measles-mixing.cpp for test examples.

# ModelMeaslesMixingRiskQuarantine

A comprehensive epidemiological model that extends measles transmission modeling with risk-stratified quarantine strategies.

#### 5.1 Overview

ModelMeaslesMixingRiskQuarantine implements a measles transmission model based on the ModelMeaslesMixing framework, with enhanced quarantine policies that vary based on exposure risk levels. This allows for targeted public health interventions that can optimize resource allocation and epidemic control.

#### 5.1.1 Key Features

- Measles-specific disease progression: Susceptible o Exposed o Prodromal o Rash o Recovered
- Population mixing: Uses contact matrices to model heterogeneous mixing between population groups
- · Risk-stratified quarantine: Three-tier system (high/medium/low risk) with customizable durations
- Enhanced detection: Detection rate parameter that activates during quarantine periods
- Contact tracing: Comprehensive contact tracing with configurable success rates
- · Entity-based risk assessment: Risk levels determined by shared entity membership
- · Vaccination effects: Vaccine distribution with configurable efficacy

#### 5.2 Disease States

The model includes 13 distinct states:

- 1. Susceptible Can become infected
- 2. Exposed Infected but not yet infectious (incubation period)

- 3. Prodromal Infectious individuals (replaces "Infected" in standard SEIR)
- 4. Rash Non-infectious with visible symptoms (detection occurs here)
- 5. **Isolated** Detected individuals in self-isolation
- 6. Isolated Recovered Recovered individuals still in isolation
- 7. Detected Hospitalized Hospitalized individuals who were contact-traced
- 8. Quarantined Exposed Exposed individuals in quarantine
- 9. Quarantined Susceptible Susceptible individuals in quarantine
- 10. Quarantined Prodromal Prodromal individuals in quarantine
- 11. Quarantined Recovered Recovered individuals in quarantine
- 12. Hospitalized Individuals requiring hospital care
- 13. Recovered Individuals who have recovered and gained immunity

## 5.3 Risk Classification System

When the quarantine process is triggered by a detected case, contacts are classified into three risk levels:

#### 5.3.1 High Risk

- Definition: Unvaccinated agents who share entity membership with the case that triggered quarantine
- Rationale: Highest transmission risk due to close, prolonged contact within same household/workplace/school
- Default quarantine duration: 21 days (configurable)

#### 5.3.2 Medium Risk

- · Definition: Unvaccinated agents who had contact with infected individuals but don't share entity membership
- · Rationale: Moderate transmission risk from community contact
- Default quarantine duration: 14 days (configurable)

#### 5.3.3 Low Risk

- Definition: All other unvaccinated agents
- Rationale: Lowest transmission risk, potential for community spread
- Default quarantine duration: 7 days (configurable)

**Note**: Vaccinated agents (those with tools) are not subject to quarantine regardless of contact patterns.

5.4 Enhanced Detection 13

#### 5.4 Enhanced Detection

The model includes an enhanced detection mechanism that operates during active quarantine periods:

• **Detection rate quarantine**: Additional parameter specifying detection probability for prodromal individuals during active quarantine

- **Mechanism**: When any quarantine process is active, prodromal individuals have an additional chance of being detected and moved to quarantined prodromal state
- Purpose: Models increased surveillance and testing during outbreak response

#### 5.5 Parameters

#### 5.5.1 Disease Parameters

- · Contact rate: Average number of contacts per step
- Transmission rate: Probability of transmission per contact
- Incubation period: Average time from exposure to becoming infectious
- Prodromal period: Average duration of infectious period
- Rash period: Average duration of rash symptoms
- Hospitalization rate: Probability of requiring hospitalization
- · Hospitalization period: Average duration of hospital stay

#### 5.5.2 Detection and Isolation

- Days undetected: Average time before rash cases are detected
- · Isolation period: Duration of isolation for detected cases
- Isolation willingness: Proportion willing to self-isolate when detected
- Detection rate quarantine: Detection rate during active quarantine periods

#### 5.5.3 Risk-based Quarantine

- Quarantine period high: Duration for high-risk contacts (days)
- Quarantine period medium: Duration for medium-risk contacts (days)
- Quarantine period low: Duration for low-risk contacts (days)
- Quarantine willingness: Proportion willing to comply with quarantine

#### 5.5.4 Contact Tracing

- · Contact tracing success rate: Probability of successfully identifying contacts
- · Contact tracing days prior: Number of days prior to detection for tracing

#### 5.5.5 Vaccination

- · Vaccination rate: Proportion of agents initially vaccinated
- · Vax efficacy: Vaccine effectiveness in preventing infection
- · Vax improved recovery: Enhanced recovery rate for vaccinated individuals

### 5.6 Usage Example

```
#include <epiworld/epiworld.hpp>
    Contact matrix for 3 population groups
std::vector<double> contact_matrix = {
       0.8, 0.1, 0.1, // Group 1 mixing
0.1, 0.8, 0.1, // Group 2 mixing
0.1, 0.1, 0.8 // Group 3 mixing
// Create model with risk-based quarantine
epimodels::ModelMeaslesMixingRiskQuarantine<> model(
       1000,
                          // Population size
       1000, // Population Size

0.01, // Initial prevalence

2.0, // Contact rate

0.2, // Transmission rate

0.9, // Vaccination efficacy

0.3, // Vaccine recovery enhancement
       7.0,
                             // Incubation period
       4.0,
                           // Prodromal period
      5.0, // Rash period
contact_matrix, // Contact matrix
0.2, // Hospitalization rate
7.0, // Hospitalization duration
3.0, // Days undetected
21, // Quarantine period high risk
14, // Quarantine period medium risk
7, // Quarantine period low risk
0.8, // Quarantine willingness
0.8, // Isolation willingness
4, // Isolation period
0.1, // Proportion vaccinated
0.15, // Detection rate during quarantine
1.0, // Contact tracing success rate
4u // Contact tracing days prior
                              // Rash period
       5.0.
                             // Contact tracing success rate
// Contact tracing days prior
       4u
// Add population entities
model.add_entity(Entity<>("Households", dist_factory<>(0, 400)));
model.add_entity(Entity<>("Schools", dist_factory<>(400, 700)))
model.add_entity(Entity<>("Workplaces", dist_factory<>(700, 1000)));
// Set initial conditions
model.initial_states({1.0, 0.0}); // All infected start as exposed
model.run(60, 123);
model.print();
```

#### 5.7 Differences from Other Models

#### 5.7.1 vs. ModelMeaslesMixing

- · Risk-stratified quarantine: Three different quarantine durations based on exposure risk
- Enhanced detection: Additional detection during active quarantine periods
- · Entity-based risk assessment: Risk levels determined by shared entity membership
- Flexible quarantine policies: Each risk level can have different quarantine duration or be disabled

5.8 Testing 15

#### 5.7.2 vs. ModelMeaslesSchool

- · Population mixing: Supports contact matrices for heterogeneous mixing
- · Risk stratification: Multiple quarantine strategies rather than uniform approach
- · Enhanced contact tracing: Risk-based contact management
- · Scalability: Supports multiple population entities with different mixing patterns

#### 5.8 Testing

The model includes comprehensive tests that verify:

- · Correct transition probabilities between all 13 states
- Proper risk level assignment (high/medium/low)
- Enhanced detection mechanism during quarantine periods
- · Different quarantine durations for different risk levels
- · Comparison of uniform vs. risk-stratified quarantine strategies
- · Population mixing functionality with multiple entities

See tests/20a-measles-mixing-risk-quarantine.cpp for test examples.

### 5.9 Applications

This model is particularly useful for:

- 1. Outbreak response planning: Comparing different quarantine strategies
- 2. Resource optimization: Allocating quarantine resources based on transmission risk
- 3. Policy evaluation: Assessing effectiveness of risk-stratified interventions
- 4. Contact tracing optimization: Understanding impact of enhanced detection during outbreaks
- 5. Vaccination strategy: Evaluating how vaccination coverage affects quarantine effectiveness

### 5.10 Implementation Notes

- · Risk levels are assigned dynamically when quarantine is triggered
- · Vaccinated agents are excluded from quarantine regardless of contact patterns
- Enhanced detection only operates when at least one quarantine process is active
- Quarantine durations can be set to -1 to disable quarantine for specific risk levels
- · The model maintains backward compatibility with uniform quarantine by setting all periods equal

# epiworld c++ template library

#### 6.1 Main features

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.

## 6.2 Algorithm

#### Setup

- · Create viruses.
- · Create tools (arbitrary).
- · Set model parameters (arbitrary).
- Create global events (e.g., surveillance).
- · Set up the population: small world network (default).
- Set up rewiring (optional).
- Set states (arbitrary number of them).

#### Run

- 1. Distribute the tool(s) and virus(es)
- 2. For each t in 1 -> Duration:
  - Update state for susceptible/infected/removed(?)
  - Mutate virus(es) (each individual)
  - Run Global events (e.g., surveillance)
  - · Run rewiring algorithm

#### Along update:

- · Contagion events are applied recorded.
- · New variants are recorded.
- · Optional user data is recorded.

### 6.3 Hello world (C++)

```
#include "include/epiworld/epiworld.hpp"
int main()
  // Creating a virus
 epiworld::Virus<> covid19("covid 19", .01, true);
 covid19.set_infectiousness(.8);
  // Creating a tool
 epiworld::Tool<> vax("vaccine", .5, true);
 vax.set_contagion_reduction(.95);
// Creating a model
  epiworld::Model<> model;
  // Adding the tool and virus
 model.add_virus(covid19);
 model.add_tool(vax);
  // Generating a random pop
 model.population_from_adjlist(
   epiworld::rgraph_smallworld(1000, 5, .2)
 // Initializing setting days and seed
model.init(60, 123123);
 // Running the model
model.run();
 model.print();
 return;
```

#### 6.4 Surveillance simulation

- Incubation time of the disease  $\sim~\text{Gamma}$  (3, ~1)
- Duration of the disease  $\sim$ Gamma (12, 1)
- · Probability of becoming symptomatic: 0.9
- Prob. of transmission: 1.0.
- · Vaccinated population: 25%
- · Vaccine efficacy: .9.
- · Vaccine reduction on transmission: 0.5.
- Surveillance program of x% of the population at random.
- · Individuals who test positive become isolated.

### 6.4.1 Preliminary results

```
# With low surveillance
pop_size <- 20e3
pop_seed <- pop_size * .01
s_levels <- c(0.0001, 0.002)
 system(sprintf("./07-surveillance.o %i %i 100 %.04f 2>&1", pop\_seed, pop\_size, s\_levels[1]), intern = TRUE) 
 cat(sep = "\n")
## Running the model...
##
##
##
## SIMULATION STUDY
##
                   : 20000
## Population size
## Days (duration)
                  : 200 (of 200)
## Number of variants : 1
## Last run elapsed t : 505.00ms
## Rewiring
                    : off
##
## Virus(es):
## - Covid19 (baseline prevalence: 100 seeds)
## Tool(s):
##
   - Vaccine (baseline prevalence: 25.00%)
##
## Model parameters:
## - Infect period
                          : 12.0000
## - Latent period
## - Latent period : 3.0000
## - Prob of symptoms : 0.7000
##
   - Prob of transmission : 1.0000
## - Prob. death
                          : 0.0010
## - Prob. reinfect
                         : 0.1000
## - Surveilance prob. : 1.0e-04
## - Vax efficacy : 0.9000
## - Vax redux transmision : 0.5000
##
\#\# Distribution of the population at time 200:
## - Total susceptible (S) : 19900 -> 2106
## - Total recovered (S)
                                           0 -> 17369
## - Total latent (I)
                                         100 -> 109
## - Total symptomatic (I)
                                          0 -> 155
                                          0 -> 2
## - Total symptomatic isolated (I) :
##
   - Total asymptomatic (I)
  - Total asymptomatic isolated (I) :
                                           0 -> 0
##
## - Total removed (R)
                                           0 -> 187
##
## (S): Susceptible, (I): Infected, (R): Recovered
## _
hist1 <- read.csv("07-surveillance_hist.txt", sep = " ")</pre>
surv1 <- read.csv("07-surveillance_user_data.txt", sep = " ")</pre>
# With high surveillance
system(sprintf("./07-surveillance.o %i %i 100 %.04f 2>&1", pop_seed, pop_size, s_levels[2]), intern = TRUE)
 cat(sep = "\n")
## Running the model...
##
##
##
## SIMULATION STUDY
## Population size : 20000 . 200 (duration)
## Days (duration)
                    : 200 (of 200)
## Number of variants : 1
## Last run elapsed t : 530.00ms
## Rewiring
##
## Virus(es):
```

```
## - Covid19 (baseline prevalence: 100 seeds)
## Tool(s):
## - Vaccine (baseline prevalence: 25.00%)
##
## Model parameters:
## - Infect period
                           : 12.0000
                          : 3.0000
##
   - Latent period
## - Prob of symptoms
                            : 0.7000
  - Prob of transmission : 1.0000
                           : 0.0010
   - Prob. death
##
   - Prob. reinfect
##
                            : 0.1000
  - Surveilance prob.
                           : 0.0020
## - Vax efficacy
                          : 0.9000
##
   - Vax redux transmision : 0.5000
##
\#\# Distribution of the population at time 200:
  - Total susceptible (S)
##
                                       : 19900 -> 2125
## - Total recovered (S)
                                              0 -> 17325
                                             100 -> 109
##
  - Total latent (I)
##
   - Total symptomatic (I)
                                              0 -> 155
## - Total symptomatic isolated (I)
                                               0 -> 8
                                      :
## - Total asymptomatic (I)
                                               0 -> 76
##
   - Total asymptomatic isolated (I) :
                                               0 -> 1
  - Total removed (R)
                                               0 -> 201
##
## (S): Susceptible, (I): Infected, (R): Recovered
## _
hist2 <- read.csv("07-surveillance_hist.txt", sep = " ")</pre>
surv2 <- read.csv("07-surveillance_user_data.txt", sep = " ")</pre>
hist_comb <- rbind(
 cbind(sim = as.character(s_levels[1]), hist1),
 cbind(sim = as.character(s_levels[2]), hist2)
qqplot(hist\_comb, aes(x = date, y = counts + 1, colour = state, linetype=sim)) +
 geom_line() +
 # scale_y_log10() + labs(y = "Counts (log)")
```

#### 6.4.2 Cases detected

```
survdat <- rbind(
  with(surv1, rbind()
   data.frame(Id = as.character(s_levels[1]), Date = date, Type = "N Sampled", n = nsampled),
   data.frame(Id = as.character(s_levels[1]), Date = date, Type = "N detected", n = ndetected),
   data.frame(Id = as.character(s_levels[1]), Date = date, Type = "N detected Asymp", n =
        ndetected_asympt),
   data.frame(Id = as.character(s_levels[1]), Date = date, Type = "N Asymp", n = nasymptomatic)
)),
  with(surv2, rbind(
   data.frame(Id = as.character(s_levels[2]), Date = date, Type = "N Sampled", n = nsampled),
   data.frame(Id = as.character(s_levels[2]), Date = date, Type = "N detected", n = ndetected),
   data.frame(Id = as.character(s_levels[2]), Date = date, Type = "N detected Asymp", n =
        ndetected_asympt),
   data.frame(Id = as.character(s_levels[2]), Date = date, Type = "N Asymp", n = nasymptomatic)
))
)
ggplot(survdat, aes(x = Date, y = n + 1, colour = Type)) +
   geom_line() +
   facet_wrap(~Id) +
   scale_y_log10() +
   labs(y = "Counts (log)")</pre>
```

## **General parameters**

The following are parameters used for both ABM and Compartmental models.

### 7.1 Compartmental Models

### 7.1.1 SIR Model

```
library(deSolve)
library(ggplot2)
library(data.table)
# Code from
# Chapter 2: SIR
# Book "Epidemics: Models and Data using R."
# By: Ottar N. Bjørnstad
sirmod <- function(t, y, parms) {
    # Pull state variables from the vector y</pre>
       S = y[1]
       R = y[3]
       # Pull parameter values from parms vector
       beta = parms["beta"]
mu = parms["mu"]
       gamma = parms["gamma"]
N = parms["N"]
       # Define equations
       dS = mu * (N - S) - beta * S * I/N
dI = beta * S * I/N - (mu + gamma) * I
dR = gamma * I - mu * R
       res = c(dS, dI, dR)
       # Return list of gradients
       list(res)
times <- seq(0, EPI_NDAYS, by = 1)

parms <- c(mu = 0, N = EPI_N, beta = EPI_BETA, gamma = EPI_GAMMA)

start <- c(S = EPI_N * (1 - EPI_0), I = EPI_N * EPI_0, R = 0)

out <- ode(y = start, times = times, func = sirmod, parms= parms)
out <- as.data.frame(out)
out <- rbind(
```

22 **General parameters** 

```
with (out, data.table(date = time, state = "Susceptible", counts = S)),
     with (out, data.table(date = time, state = "Infected", counts = I)), with (out, data.table(date = time, state = "Recovered", counts = R))
Now we visualize the model
ggplot(out, aes(x = date, y = counts)) +
    geom_line(aes(colour = state)) +
     labs(title = "Compartmental SIR")
7.1.2 SEIR Model
# Code adapted from
  Chapter 2: SIR
```

```
# Book "Epidemics: Models and Data using R"
# By: Ottar N. Bjørnstad
seirmod <- function(t, y, parms) {
    # Pull state variables from y vector</pre>
     S = y[1]
     E = y[2]
     I = y[3]
     R = y[4]
     # Pull parameter values from parms vector
     beta = parms["beta"]
mu = parms["mu"]
     alpha = parms["alpha"
     gamma = parms["gamma"]
             = parms["N"]
     # Define equations
     dS = mu * (N - S) - beta * S * I/N - mu * S
         = beta * S * I/N - (mu + alpha) * E
     dI = alpha * E - (mu + gamma) * I
dR = gamma * I - mu * R
     res = c(dS, dE, dI, dR)
     # Return list of gradients
# Initial parameters
parms <- c(

mu = 0, N = EPI_N, beta = EPI_BETA,
     alpha = 1/EPI_LATENCY, gamma = EPI_GAMMA
start <- c(S = EPI_N * (1 - EPI_0), E = EPI_N * EPI_0, I = 0, R = 0)
out_seir <- ode(y = start, times = times, func = seirmod, parms = parms)</pre>
out_seir <- as.data.frame(out_seir)</pre>
out_seir <- rbind(
     with(out_seir, data.table(date = time, state = "Susceptible", counts = S)),
     with (out_seir, data.table(date = time, state = "Exposed", counts = E)), with (out_seir, data.table(date = time, state = "Infected", counts = I)), with (out_seir, data.table(date = time, state = "Recovered", counts = I)), with (out_seir, data.table(date = time, state = "Recovered", counts = R))
Now we visualize the model
```

```
ggplot(out\_seir, aes(x = date, y = counts)) + geom\_line(aes(colour = state)) +
      labs(title = "Compartmental SEIR")
```

### **Agent-Based Model Approach**

Calculation of the expected number of days in state \$a\$ when prob of changing state equals \$\alpha\$ is \$1\alpha\$

```
set.seed(712)
a <- .3
R \leftarrow matrix(runif(2e5 * 50), ncol = 50)
dat <- apply(R, 1, \setminus(x) {
   which.max(x < a)
mean(dat) - 1 / a
[1] -0.01049333
```

### 7.2.1 Mathematical preliminaries

That agent \$i\$ becomes infected can be computed as follows:

At the same time, the probability of not becoming infected equals to the probability of no infected agent transmitting the infection. The probability that agent \$i\\$ infects \$i\\$ equals

In this case, \$\beta\$ is parametrized such that its values are within \$(0,1)\$. Since transmission from the \$1\$ infected agents happens independently, we finally have the following:

With the above equation, we can now calculate the change in the number of susceptible agents. In this case, it equals the expected number of new infections:

With the same parametrization in the canonical SIR model (Kermack and McKendrick), the instantaneous change in the number of susceptible agents equals  $\frac{s}{d} = -S \beta$  is. Given \$S\$ and \$I\$, we can show that, as  $\frac{s}{d} = -S \beta$  to the same number. Formally:

The same can be shown for the change in the number recovered.

### 7.2.2 Simulation study

ggplot(epiworld, aes(x = date, y = counts)) +
 geom\_line(aes(colour = state)) +
 labs(title = "ABM SEIR")

### 7.3 Comparing ABM with Compartmental Models

To this end, we will compare the results of the first run of the Compartmental model with 100 runs of the ABM, compute the confidence interval, and see how likely is the compartmental model to fall within the trajectory of the ABM simulation.

#### 7.3.1 SIR

```
system("./09-sir-connected.o -n $EPI N -b $EPI BETA -d $EPI NDAYS -p $EPI 0 -r $EPI GAMMA -i 1.0 -s555599 -e
        100")
library(ggplot2)
library(data.table)
epiworld <- data.table::fread("09-sir-connected-experiments.csv")</pre>
epiworld <- epiworld[, .(
    min = quantile(counts, probs = .025),
    mean = mean(counts),
    max = quantile(counts, probs = .975)), by = .(date, state)]
# Merging Compartmental
epiworld <- merge(
     epiworld,
    out[, .(date = date, state = state, compartmental = counts)], by = c("date", "state")
setorder(epiworld, state, date)
ggplot(epiworld, aes(x = date, y = mean)) +
     geom_ribbon(aes(ymin = min, ymax = max, colour = state), alpha = 0.1) +
     \texttt{geom\_line}(\texttt{aes}\,(\texttt{x}\,=\,\texttt{date},\,\,\texttt{y}\,=\,\texttt{compartmental},\,\,\texttt{colour}\,=\,\texttt{sprintf}\,(\texttt{"\$s}\,\,\,(\texttt{compt})\,\texttt{"},\,\,\texttt{state})))
```

It seems that, although both yield the same equilibria, compartmental models reach the highest point of the simulation earlier. This makes sense as within a single day of the ABM simulation, compartmental models have more events taking place. Nonetheless, as predicted, as \$\beta\downarrow 0\\$, the differences become lesser. Furthermore, we could use the fact that the transition rates are known to compute an adjustment.

24 General parameters

### 7.3.2 **SEIR**

### 7.3.3 Rates

```
S <- 1000
rate_comp <- function(I,B) S * B * I
rate_abm <- function(I,B) S * (1 - (1 - B)^I)
op <- par(mfrow = c(3, 2))
for (i in c(1, 10, 100)) {
        curve(rate_comp(i, x), from = .01, to = 0.05)
        curve(rate_abm(i, x), from = .01, to = 0.05, add = FALSE, lty = 2)
}
par(op)</pre>
```

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### model1

The dynamics of the simulation process are:

- 1. Discrete Markov process.
- 2. The simulation has the following parameters:
  - a. New variant emergence at rate X. b. For each variant k:
    - Unvaccinated individuals become sick rate C(k),
    - Mortality rate D (k),
    - Recovery rate H(k),
    - Vaccines have an efficaccy rate  $\mathbb{E}\left(v,k\right)$  and pseudo vaccines (recovered) have efficacy rate  $\mathbb{E}\left(r,k\right)$   $< \mathbb{E}\left(v,k\right)$ . In general, the probability of i acquiring the disease k from j will be equal to

```
``` P(i gets the disease from j | their states) = C(k) * (1 - E(i,k)) * (1 - E(j, k)) ```
```

where (i,j) in (u,v,r). Efficacy rate for unvaccinated is zero.

- Vaccinated individuals have a reduced mortality rate D(k,v) > D(k), and recovered individuals D(k,r) in (D(k,v), D(k)]
- Vaccinated individuals have an increased recovery rate H (k, v) > H (k), whereas recovered's rate H (k, r) in [H(k), H(k, v)).

The sum of mortality and recovery rates is less than one since the difference represents no change.

- c. Each country vaccinates citizens at rate V function of A (availability) and B (citizens' acceptance rate.) d. In each country i, the entire population N(i) distributes between the following states:
  - Healthy unvaccinated (N (i,t,u)),
  - Healthy vaccinated ( $\mathbb{N}(i,t,v)$ ),
  - Deceased (N(i,t,d)),
  - Recovered (N(i,t,r)),
  - Unvaccinated and sick with variant (N(i,t,s,k|u))k., and
  - Vaccinated and sick with variant (N (i,t,s,k|v)) k .

```
Total sick are N(i,t,k,s) = sum(g in \{u,v\}) N(i,t,k,s|g)
```

Globally, we keep track of the prevalence of new variants. Variants can disappear if no more individuals port the variant, i.e., the prevalence rate P(k,t) = sum(i) N(i,s,k) equals zero.

d. Vaccines are manufactured at each country at rates M(i) and uniformly shared with other countries at rate S(i). c. Population flows between each country pair (i,j) at a rate F(i,j). Flows between countries do not change Population and are symmetric.

28 model1

- 3. The simulation process is as follows:
  - (a) Countries are initialized with a total population N(i).
  - (b) Variant zero initializes at a random location i, with an initial prevalence P(k,t) = N(i,t,k).
  - (c) For time t in (0,T) do:
    - a. Unvaccinated individuals can become sick of variant  ${\bf k}$  with probability:
    - ```  $Pr(h->s|i,t,k,u) \sim sum(g in \{u,v\}) (N(i,t-1,s,k|g) + sum(j != i) F(i,j) * N(j,t-1,s,k|g)) * C(k) / (N(i) + sum(j != i) N(j)) ```$
    - b. Vaccinated individuals can become sick of variant k with probability:  $\Pr(v->s|i,t,k,v) \sim \Pr(h->s|i,t,k) * (1 E(v,k))$ .
    - b. Recovered individuals can become sick of variant k with probability:  $\Pr(v->s|i,t,k,r) \sim \Pr(h->s|i,t,k) * (1 E(r,k))$ .
    - c. Sick individuals with variant k die with probability D(k) or recover with probability H(k), otherwise they stay infected; with the rates depending on their vaccination status v or n.
    - d. Unvaccinated individuals vaccinate in country i with probability  $P(u->v) \sim V(A(i,t), B(i))$ .
    - e. The country vaccine supply changes.

## Mixing probabilities in connected model

George G. Vega Yon, Ph.D. 2024-04-25

### 10.1 Case 1: No grouping

We will look into the probability of drawing infected individuals to simplify the algorithm. There are \$1\$ infected individuals at any time in the simulation; thus, instead of drawing from \$Bern(c/N, N)\$, we will be drawing from \$\infty\$ Bern(c/N, I)\$. The next step is to check which infected individuals should be drawn. Let's compare the distributions using the hypergeometric as an example:

```
set.seed(132)
nsims <- 1e4
N <- 400
rate <- 5
p <- rate/N
I <- 10
sim_complex <- parallel::mclapply(1:nsims, \(i) {</pre>
 nsamples <- rbinom(N, N, p)
sum(rbinom(N, size = nsamples, prob = I/N) > 0)
}, mc.cores = 4L) |> unlist()
sim_simple <- parallel::mclapply(1:nsims, \(i) {</pre>
sum(rbinom(N, I, p) > 0)
}, mc.cores = 4L) |> unlist()
op <- par(mfrow = c(1,2))
MASS::truehist(sim_complex)
MASS::truehist(sim_simple)
par(op)
quantile(sim_complex)
  0% 25% 50% 75% 100%
  27 43 47 51
quantile(sim_simple)
  0% 25% 50% 75% 100%
  23 43 47 51 71
```

These two approaches are equivalent, but the second one is more efficient from the computational perspective.

plotter(sim\_complex, sim\_simple)

### 10.2 Case 2: Grouping

This explores the case when we have mixing across groups. The question is if we can replicate the effect at the group level.

set.seed(123133)

```
ngroups <- 3
mixing <- matrix(
  c(0.1, 0.2, 0.3, 0.2, 0.1, 0.2, 0.3, 0.2, 0.1),
  nrow = ngroups,
  ncol = ngroups
mixing <- mixing/rowSums(mixing)</pre>
mixing
             [,1]
                         [,2]
[1,] 0.1666667 0.3333333 0.5000000
[2,] 0.4000000 0.2000000 0.4000000
[3,] 0.5000000 0.3333333 0.1666667
N <- 500
sizes <- c(100, 150, 250) rate <- 5
p <- rate/N
I <- c(10, 30, 20)
ids <- rep.int(1:ngroups, times = sizes)</pre>
nsims <- 1e4
sim_complex <- parallel::mclapply(1:nsims, \(i) {</pre>
  # Sampling group first
  # How many each individual will sample from the groups
    ans <- rbinom(
     n = N, size = sizes[g], prob = mixing[ids,][,g] * p
      ) |> sum()
    # Sampling with replacement rbinom(ans, size = 1, prob = I[g]/sizes[g]) |> sum()
  }) |> sum()
, mc.cores = 4L) > unlist()
Using the alternative method in which we directly weight the probabilities:
sim_simple <- parallel::mclapply(1:nsims, \(i) {
    # Sampling group first</pre>
  sapply(1:ngroups, \(g) {
    rbinom(
      n = N, size = I[g], prob = mixing[cbind(ids,g)] * p
 ) |> sum()
}) |> sum()
}, mc.cores = 4L) |> unlist()
op <- par(mfrow = c(1,2))</pre>
MASS::truehist(sim_complex)
MASS::truehist(sim_simple)
par(op)
quantile(sim_complex)
  0% 25% 50% 75% 100%
              94 101 131
       88
quantile(sim_simple)
  0% 25% 50% 75% 100%
  58 87 94 101 135
plotter(sim_complex, sim_simple)
```

### **EPI Simulator**

### 11.1 Disease dynamics

Diseases continuously evolve in time. Changes in their genetic sequence make them more or less resistant to the particular version of the vaccine. Mutations also affect the transmissibility level and mortality rate of the disease. Using this approach allows making vaccination efficacy a function of compatibility between the variant and the vaccine.

When an individual becomes infected, the disease accumulates mutations in the new host. Ultimately, there is no single version of the disease present in the model, but rather an infinite number of them, each slightly different from the other.

### 11.2 Network dynamics

We can assume that the Population is organized in fully connected blocks for the first version of the model. Block sizes and the number of connections between blocks are Poisson random variables. Individuals interact with all the members of their blocks, and bridging individuals allow the disease to move across blocks.

### 11.3 Contagion dynamics

The transmission of the disease will be governed by the number of vaccinated, infected, and recovered within each block. Transmission between blocks will be treated in the same way, although individuals bridging the block will only interact with others within the block and their direct connections across the blocks.

### 11.4 Time dynamics

Time dynamics has two components, how biology evolves and how agents react.

The model develops as a continuous-time Markov process. Each block of individuals takes action at rates  $\mathbb{L}\left(\frac{1}{N}\right)$  function of the local number of infections. This way, if

32 EPI Simulator

### 11.5 Updating agent's status

Like most other components, updating agents' states can be personalized. A naive approach allows agents to get infected with a single virus or stay as-is. The probability of this event is conditional on acquiring at most one virus. Since these are independent events, the conditional probability is computed as follows:

### Where

```
P(only variant k) = P(k) * Prod(m!=v) (1 - P(m))

P(at most 1) = P(None) + Sum(v in variants) P(v) * Prod(m != v) (1 - P(m))

P(None) = Prod(v in variants) (1 - P(v))
```

Furthermore, the (Variant, Person) pairs are treated independently.

### 11.5.1 Other parameters

- · Who did you get the infection from.
- · Omicron is 1.5 more infectious than delta.
- · Surveillance:
  - Pull people to be tested at random.
  - Or at symptoms.
  - A mix of the two.
- Define a class for passing extra functions and datasets, for example, testing surveillance.
- · Exposed people become infectious after k days.
- Network changesthe can be a function of an ERGM. Apply K steps throughout time.
- · Add progress bar.

## Fix for rewire\_degseq Bug

### 12.1 Problem

The rewire\_degseq function in randgraph.hpp was incorrectly swapping edge weights instead of actually rewiring the network connections. This meant that:

- The network structure (who connects to whom) remained unchanged
- · Only the weight values stored in the adjacency maps were being swapped

### 12.2 Root Cause

```
In the AdjList version:
// OLD CODE (incorrect):
std::swap(p0[id01], p1[id11]); // This only swaps the weights!
```

The code was swapping the VALUES in the map (the weights), not the KEYS (the neighbor IDs).

### 12.3 Solution

The fix properly rewires edges by:

- 1. Saving the edge weights
- 2. Removing the old edges from the adjacency lists
- 3. Adding new edges with the swapped connections
- 4. For undirected graphs, also updating the reverse edges
- 5. Checking for duplicate edges before swapping to preserve the degree sequence

### 12.3.1 AdjList Version Changes

- Replace std::swap(p0[id01], p1[id11]) with proper edge removal and addition
- · Add duplicate edge detection to skip rewires that would create parallel edges
- · Ensure both directions are updated for undirected graphs

### 12.3.2 Agent Vector Version Changes

- Fixed bug where model->get\_agents()[id0] should have been model->get\_agents()[non←\_isolates[id0]]
- Added duplicate edge detection before calling swap\_neighbors()
- The swap\_neighbors () function itself was already correctly implemented

### 12.4 Testing

A standalone test (test-rewire-simple.cpp) was created to verify:

- 1. Degree preservation: Each node maintains its original number of connections
- 2. Network structure change: The rewiring actually changes who connects to whom
- 3. Graph symmetry: For undirected graphs, if A connects to B, then B connects to A

### 12.4.1 Test Results

```
=== Testing rewire_degseq fix ===
Degrees preserved: YES
Nodes with changed neighbors: 15 out of 20
Graph symmetric: YES
PASS: All tests successful!
```

### 12.5 Files Modified

- include/epiworld/randgraph.hpp: Fixed both rewire\_degseq implementations
- tests/test-rewire-simple.cpp: Standalone test to verify the fix

### **12.6 Notes**

- · The main test suite has pre-existing compilation errors in unrelated models that prevent it from compiling
- The standalone test validates the core fix without requiring the full test infrastructure
- The fix is minimal and surgical, changing only what's necessary to correct the rewiring logic

# Namespace Index

### 13.1 Namespace List

Here is a list of all documented namespaces with brief descriptions:

epiworld	::sampler					
	Functions for sampling viruses	 	 	 	 	47
sampler						
	Functions for sampling viruses					50

36 Namespace Index

## **Hierarchical Index**

### 14.1 Class Hierarchy

This inheritance list is sorted roughly, but not completely, alphabetically:

AdjList
epiworld::AdjList
Agent < TSeq >
epiworld::Agent < TSeq >
Agent < EPI_DEFAULT_TSEQ >
AgentsSample < TSeq >
epiworld::AgentsSample < TSeq >
DataBase < TSeq >
epiworld::DataBase < TSeq >
Entities < TSeq >
epiworld::Entities < TSeq >
Entities_const< TSeq >
epiworld::Entities_const< TSeq >
Entity < TSeq >
epiworld::Entity< TSeq >
Entity < EPI_DEFAULT_TSEQ >
epiworld::Event < TSeq >
Event< TSeq >
epiworld::GlobalEvent< TSeq >
GlobalEvent< TSeq >
epiworld::LFMCMC< TData >
LFMCMC< TData >
epiworld::Model < TSeq >
Model < TSeq >
ModelMeaslesSchool < TSeq >
epiworld::Model < EPI_DEFAULT_TSEQ >
ModelDiffNet < TSeq >
ModelMeaslesMixing < TSeq >
ModelMeaslesMixingRiskQuarantine < TSeq >
ModelSEIR < TSeq >
ModelSEIRCONN < TSeq >
ModelSEIRD < TSeq >
ModelSEIRDCONN < TSeq >
ModelSEIRMixing < TSeq >
ModelSEIRMixingQuarantine < TSeq >

38 Hierarchical Index

$ModelSIR{}\dots\dots\dots\dots\dots\dots\dots\dots\dots\dots\dots\dots\dots\dots\dots\dots\dots\dots\dots\dots\dots\dots\dots\dots\dots\dots\dots\dots\dots\dots\dots\dots\dots\dots\dots$	. 221
ModelSIRCONN< TSeq >	
ModelSIRD< TSeq >	
ModelSIRDCONN< TSeq >	. 239
ModelSIRLogit < TSeq >	
ModelSIRMixing< TSeq >	
ModelSIS< TSeq >	
ModelSISD< TSeq >	
ModelSURV < TSeq >	
$epiworld:: epimodels:: Model Diff Net < TSeq > \dots $	
$epiworld:: epimodels:: Model Measles Mixing < TS eq > \dots $	
$epiworld:: epimodels:: Model Measles Mixing Risk Quarantine < TS eq > \dots $	
$epiworld::epimodels::ModelMeaslesSchool < TSeq > \dots $	
epiworld::epimodels::ModelSEIR< TSeq >	
epiworld::epimodels::ModelSEIRCONN< TSeq >	
epiworld::epimodels::ModelSEIRD< TSeq >	
$epiworld:: epimodels:: Model SEIRDCONN < TS eq > \dots $	
epiworld::epimodels::ModelSEIRMixing< TSeq >	
$epiworld:: epimodels:: Model SEIR Mixing Quarantine < TS eq > \dots $	
epiworld::epimodels::ModelSIR< TSeq >	
$epiworld::epimodels::ModelSIRCONN < TSeq > \dots $	
epiworld::epimodels::ModelSIRD< TSeq >	
$epiworld:: epimodels:: Model SIRDCONN < TS eq > \dots $	
epiworld::epimodels::ModelSIRLogit< TSeq >	
epiworld::epimodels::ModelSIRMixing< TSeq >	
epiworld::epimodels::ModelSIS< TSeq >	
epiworld::epimodels::ModelSISD< TSeq >	
epiworld::epimodels::ModelSURV< TSeq >	. 262
epiworld::ModelDiagram	127
ModelDiagram	127
${\sf Network} {<} \ {\sf Nettype}, \ {\sf Nodetype}, \ {\sf Edgetype} {>} \ \ldots \ $	267
${\sf epiworld::PersonTools} {< \sf TSeq >  \dots \dots$	
${\sf PersonTools}{<{\sf TSeq}>} \ \dots $	268
epiworld::Progress	
Progress	
epiworld::Queue < TSeq >	
Queue < TSeq >	
RandGraph	270
epiworld::SAMPLETYPE	271
SAMPLETYPE	271
SimpleAdjList	271
SimpleModel	272
epiworld::Tool< TSeq >	272
Tool< TSeq >	273
epiworld::ToolFunctions < TSeq >	275
ToolFunctions < TSeq >	276
epiworld::Tools< TSeq >	276
Tools < TSeq >	277
epiworld::Tools_const< TSeq >	278
Tools_const< TSeq >	278
epiworld::UserData < TSeq >	279
UserData < TSeq >	281
epiworld::vecHasher< T >	283
vecHasher< T >	283
epiworld::Virus< TSeq >	284
Virus < TSeq >	286
epiworld::Viruses< TSeq >	288
Viruses< TSeq >	289

14.1 Class Hierarch	y 39
---------------------	------

epiworld::Viruses_const< TSeq >	290
Viruses_const< TSeq >	290
epiworld::VirusFunctions< TSeq >	291
VirusFunctions < TSeq >	292

40 Hierarchical Index

## **Class Index**

### 15.1 Class List

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

AdjList	55
epiworld::AdjList	56
Agent < TSeq >	
Agent (agents)	58
epiworld::Agent< TSeq >	
Agent (agents)	63
AgentsSample < TSeq >	
Sample of agents	66
epiworld::AgentsSample < TSeq >	
Sample of agents	68
DataBase < TSeq >	
Statistical data about the process	69
epiworld::DataBase< TSeq >	
Statistical data about the process	75
Entities < TSeq >	
Set of Entities (useful for building iterators)	79
epiworld::Entities< TSeq >	
Set of Entities (useful for building iterators)	80
Entities_const< TSeq >	
Set of Entities (const) (useful for iterators)	81
epiworld::Entities_const< TSeq >	
Set of Entities (const) (useful for iterators)	82
Entity < TSeq >	83
epiworld::Entity< TSeq >	84
epiworld::Event < TSeq >	
Event data for update an agent	86
Event< TSeq >	
Event data for update an agent	87
epiworld::GlobalEvent< TSeq >	
Template for a Global Event	89
GlobalEvent < TSeq >	
Template for a Global Event	91
epiworld::LFMCMC< TData >	
Likelihood-Free Markov Chain Monte Carlo	92
LFMCMC< TData >	
Likelihood-Free Markov Chain Monte Carlo	93

42 Class Index

epiworld::Model < TSeq >	
Core class of epiworld	95
Model < TSeq >	
Core class of epiworld	110
epiworld::ModelDiagram	127
ModelDiagram	. 127
epiworld::epimodels::ModelDiffNet< TSeq >	
Template for a Network Diffusion Model	127
ModelDiffNet< TSeq >	
Template for a Network Diffusion Model	129
epiworld::epimodels::ModelMeaslesMixing< TSeq >	
Measles model with population mixing, quarantine, and contact tracing	130
ModelMeaslesMixing < TSeq >	
Measles model with population mixing, quarantine, and contact tracing	139
epiworld::epimodels::ModelMeaslesMixingRiskQuarantine< TSeq >	100
Measles model with population mixing and risk-based quarantine strategies	. 147
ModelMeaslesMixingRiskQuarantine < TSeq >	147
	150
Measles model with population mixing and risk-based quarantine strategies	. 156
epiworld::epimodels::ModelMeaslesSchool< TSeq >	400
Template for a Measles model with quarantine	163
ModelMeaslesSchool < TSeq >	400
Template for a Measles model with quarantine	168
epiworld::epimodels::ModelSEIR< TSeq >	
Template for a Susceptible-Exposed-Infected-Removed (SEIR) model	172
ModelSEIR < TSeq >	
Template for a Susceptible-Exposed-Infected-Removed (SEIR) model	
${\sf epiworld::epimodels::ModelSEIRCONN} < {\sf TSeq} > \ \dots \$	
ModelSEIRCONN< TSeq >	181
epiworld::epimodels::ModelSEIRD< TSeq >	
Template for a Susceptible-Exposed-Infected-Removed-Deceased (SEIRD) model	184
ModelSEIRD< TSeq >	
Template for a Susceptible-Exposed-Infected-Removed-Deceased (SEIRD) model	
${\sf epiworld::epimodels::ModelSEIRDCONN{< TSeq > \dots $	
ModelSEIRDCONN < TSeq >	
${\it epiworld::epimodels::} Model SEIR Mixing < TSeq > \dots $	196
${\sf ModelSEIRMixing}{<}  {\sf TSeq} {>}  \dots $	200
epiworld::epimodels::ModelSEIRMixingQuarantine< TSeq >	
SEIR model with mixing, quarantine, and contact tracing	204
ModelSEIRMixingQuarantine < TSeq >	
SEIR model with mixing, quarantine, and contact tracing	212
epiworld::epimodels::ModelSIR< TSeq >	
Template for a Susceptible-Infected-Removed (SIR) model	220
ModelSIR < TSeq >	
Template for a Susceptible-Infected-Removed (SIR) model	221
epiworld::epimodels::ModelSIRCONN< TSeq >	224
ModelSIRCONN < TSeq >	227
epiworld::epimodels::ModelSIRD< TSeq >	
Template for a Susceptible-Infected-Removed-Deceased (SIRD) model	230
ModelSIRD < TSeq >	
Template for a Susceptible-Infected-Removed-Deceased (SIRD) model	233
epiworld::epimodels::ModelSIRDCONN< TSeq >	
ModelSIRDCONN < TSeq >	
epiworld::epimodels::ModelSIRLogit< TSeq >	
Template for a Susceptible-Infected-Removed (SIR) model	241
ModelSIRLogit < TSeq >	
Template for a Susceptible-Infected-Removed (SIR) model	245
epiworld::epimodels::ModelSIRMixing< TSeq >	
the contract of the commence of the contract o	270
ModelSIRMixing < TSeq >	

15.1 Class List 43

aniwaylduanimadalauMadalQlQ < TQaa >	
epiworld::epimodels::ModelSIS< TSeq > Template for a Susceptible-Infected-Susceptible (SIS) model	255
ModelSIS < TSeq >	
Template for a Susceptible-Infected-Susceptible (SIS) model	258
epiworld::epimodels::ModelSISD< TSeq >	
	259
ModelSISD< TSeq >	
	261
	262
	265
Network< Nettype, Nodetype, Edgetype >	267
	267
PersonTools < TSeq >	268
epiworld::Progress	
A simple progress bar	268
Progress	
A simple progress bar	268
	269
Queue < TSeq >	203
	269
	209 270
·	270 271
•	271 271
	271 271
	272
epiworld::Tool< TSeq >	070
	272
Tool< TSeq >	070
	273
epiworld::ToolFunctions< TSeq >	
Helper class to store the functions avoiding multiple shared_pointers (we have only one for the	
,	275
ToolFunctions < TSeq >	
Helper class to store the functions avoiding multiple shared_pointers (we have only one for the	
,	276
epiworld::Tools< TSeq >	
Set of tools (useful for building iterators)	276
Tools< TSeq >	
Set of tools (useful for building iterators)	277
epiworld::Tools_const< TSeq >	
Set of Tools (const) (useful for iterators)	278
Tools_const< TSeq >	
Set of Tools (const) (useful for iterators)	278
epiworld::UserData< TSeq >	
Personalized data by the user	279
UserData < TSeq >	
Personalized data by the user	281
epiworld::vecHasher< T >	
Vector hasher	283
vecHasher< T >	
Vector hasher	283
epiworld::Virus< TSeq >	
	284
Virus< TSeq >	
·	286
epiworld::Viruses< TSeq >	
Set of viruses (useful for building iterators)	288

44 Class Index

/iruses< TSeq >	
Set of viruses (useful for building iterators)	289
piworld::Viruses_const< TSeq >	
Set of Viruses (const) (useful for iterators)	290
/iruses_const< TSeq >	
Set of Viruses (const) (useful for iterators)	290
piworld::VirusFunctions< TSeq >	291
· /irusFunctions< TSeg >	

## File Index

### 16.1 File List

Here is a list of all documented files with brief descriptions:

epiworld.hpp
include/epiworld/adjlist-bones.hpp
include/epiworld/adjlist-meat.hpp
include/epiworld/agent-bones.hpp
include/epiworld/agent-events-meat.hpp??
include/epiworld/agent-meat-state.hpp
Sampling functions are getting big, so we keep them in a separate file
include/epiworld/agent-meat-virus-sampling.hpp??
include/epiworld/agent-meat.hpp
include/epiworld/agentssample-bones.hpp
include/epiworld/config.hpp
include/epiworld/database-bones.hpp
include/epiworld/database-meat.hpp??
include/epiworld/entities-bones.hpp
include/epiworld/entity-bones.hpp
include/epiworld/entity-distribute-meat.hpp??
include/epiworld/entity-meat.hpp
include/epiworld/epiworld-macros.hpp
include/epiworld/epiworld.hpp
include/epiworld/ <b>globalevent-bones.hpp</b>
include/epiworld/ <b>globalevent-meat.hpp</b>
include/epiworld/misc.hpp
include/epiworld/model-bones.hpp
include/epiworld/model-meat-print.hpp
include/epiworld/model-meat.hpp
include/epiworld/modeldiagram-bones.hpp
include/epiworld/modeldiagram-meat.hpp
include/epiworld/ <b>network-bones.hpp</b>
include/epiworld/ <b>progress.hpp</b>
include/epiworld/queue-bones.hpp
include/epiworld/ <b>randgraph.hpp</b>
include/epiworld/ <b>random_graph.hpp</b>
include/epiworld/seq_processing.hpp
include/epiworld/tool-bones.hpp
include/epiworld/tool-distribute-meat.hpp

46 File Index

include/epiworld/tool-meat.hpp	??
include/epiworld/tools-bones.hpp	??
include/epiworld/ <b>userdata-bones.hpp</b>	??
include/epiworld/ <b>userdata-meat.hpp</b>	??
include/epiworld/virus-bones.hpp	??
include/epiworld/virus-distribute-meat.hpp	??
include/epiworld/virus-meat.hpp	??
include/epiworld/viruses-bones.hpp	??
include/epiworld/math/distributions.hpp	??
include/epiworld/math/ <b>Ifmcmc.hpp</b>	??
include/epiworld/math/lfmcmc-bones.hpp	??
include/epiworld/math/lfmcmc/lfmcmc-meat-print.hpp	??
include/epiworld/math/lfmcmc/lfmcmc-meat.hpp	??
include/epiworld/models/diffnet.hpp	??
include/epiworld/models/ <b>globalevents.hpp</b>	??
include/epiworld/models/ <b>init-functions.hpp</b>	??
include/epiworld/models/measlesmixing.hpp	
Template for a Measles model with population mixing, quarantine, and contact tracing	295
include/epiworld/models/measlesmixingriskquarantine.hpp	
Template for a Measles model with population mixing and risk-based quarantine	296
include/epiworld/models/ <b>measlesquarantine.hpp</b>	??
include/epiworld/models/ <b>models.hpp</b>	??
include/epiworld/models/ <b>seir.hpp</b>	??
include/epiworld/models/seirconnected.hpp	??
include/epiworld/models/ <b>seird.hpp</b>	??
include/epiworld/models/seirdconnected.hpp	??
include/epiworld/models/ <b>seirmixing.hpp</b>	??
include/epiworld/models/seirmixingquarantine.hpp	
Template for a Susceptible-Exposed-Infected-Removed (SEIR) model with mixing, quarantine,	
and contact tracing	298
include/epiworld/models/ <b>sir.hpp</b>	??
include/epiworld/models/ <b>sirconnected.hpp</b>	??
include/epiworld/models/ <b>sird.hpp</b>	??
include/epiworld/models/ <b>sirdconnected.hpp</b>	??
include/epiworld/models/ <b>sirlogit.hpp</b>	??
include/epiworld/models/ <b>sirmixing.hpp</b>	??
include/epiworld/models/ <b>sis.hpp</b>	??
include/epiworld/models/ <b>sisd.hpp</b>	??
include/epiworld/models/ <b>surveillance.hpp</b>	??
tooto/tooto hnn	22

## **Namespace Documentation**

### 17.1 epiworld::sampler Namespace Reference

Functions for sampling viruses.

### **Functions**

```
    template<typename TSeq = EPI_DEFAULT_TSEQ>
    std::function< void(Agent< TSeq > *, Model< TSeq > *)> make_update_susceptible (std::vector<
    epiworld_fast_uint > exclude={})
```

Make a function to sample from neighbors.

template<typename TSeq = EPI\_DEFAULT\_TSEQ>
 std::function< Virus< TSeq > \*(Agent< TSeq > \*, Model< TSeq > \*)> make\_sample\_virus\_neighbors
 (std::vector< epiworld\_fast\_uint > exclude={})

Make a function to sample from neighbors.

Sample from neighbors pool of viruses (at most one)

template<typename TSeq = EPI\_DEFAULT\_TSEQ>
 Virus< TSeq > \* sample\_virus\_single (Agent< TSeq > \*p, Model< TSeq > \*m)

### 17.1.1 Detailed Description

Functions for sampling viruses.

#### 17.1.2 Function Documentation

### 17.1.2.1 make\_sample\_virus\_neighbors()

Make a function to sample from neighbors.

This is akin to the function default\_update\_susceptible, with the difference that it will create a function that supports excluding states from the sampling frame. For example, individuals who have acquired a virus can be excluded if in incubation state.

### **Template Parameters**

#### **Parameters**

exclude	unsigned vector of states that need to be excluded from the sampling
---------	----------------------------------------------------------------------

#### Returns

Virus<TSeq>\* of the selected virus. If none selected (or none available,) returns a nullptr;

### 17.1.2.2 make\_update\_susceptible()

Make a function to sample from neighbors.

This is akin to the function default\_update\_susceptible, with the difference that it will create a function that supports excluding states from the sampling frame. For example, individuals who have acquired a virus can be excluded if in incubation state.

#### **Template Parameters**

TC-	
i Seq	

### **Parameters**

exclude	unsigned vector of states that need to be excluded from the sampling
CACIGGE	and great vector of states that need to be excluded from the sampling

### Returns

Virus<TSeq>\* of the selected virus. If none selected (or none available,) returns a nullptr;

### 17.1.2.3 sample\_virus\_single()

Sample from neighbors pool of viruses (at most one)

This function samples at most one virus from the pool of viruses from its neighbors. If no virus is selected, the function returns a nullptr, otherwise it returns a pointer to the selected virus.

This can be used to build a new update function (EPI\_NEW\_UPDATEFUN.)

### **Template Parameters**

TSeq	

#### **Parameters**

р	Pointer to person	
m	Pointer to the model	

#### Returns

Virus<TSeg>\* of the selected virus. If none selected (or none available,) returns a nullptr;

### 17.2 sampler Namespace Reference

Functions for sampling viruses.

### **Functions**

```
    template<typename TSeq = EPI_DEFAULT_TSEQ>
    std::function< void(Agent< TSeq > *, Model< TSeq > *)> make_update_susceptible (std::vector<
    epiworld_fast_uint > exclude={})
```

Make a function to sample from neighbors.

template<typename TSeq = EPI\_DEFAULT\_TSEQ>
 std::function
 Virus< TSeq > \*(Agent< TSeq > \*, Model< TSeq > \*)> make\_sample\_virus\_neighbors
 (std::vector< epiworld\_fast\_uint > exclude={})

Make a function to sample from neighbors.

```
• template<typename TSeq = EPI_DEFAULT_TSEQ> 
    Virus< TSeq > * sample_virus_single (Agent< TSeq > *p, Model< TSeq > *m)
```

Sample from neighbors pool of viruses (at most one)

### 17.2.1 Detailed Description

Functions for sampling viruses.

### 17.2.2 Function Documentation

### 17.2.2.1 make\_sample\_virus\_neighbors()

Make a function to sample from neighbors.

This is akin to the function default\_update\_susceptible, with the difference that it will create a function that supports excluding states from the sampling frame. For example, individuals who have acquired a virus can be excluded if in incubation state.

#### **Template Parameters**

#### **Parameters**

exclude	unsigned vector of states that need to be excluded from the sampling
---------	----------------------------------------------------------------------

#### Returns

Virus<TSeq>\* of the selected virus. If none selected (or none available,) returns a nullptr;

### 17.2.2.2 make\_update\_susceptible()

Make a function to sample from neighbors.

This is akin to the function default\_update\_susceptible, with the difference that it will create a function that supports excluding states from the sampling frame. For example, individuals who have acquired a virus can be excluded if in incubation state.

#### **Template Parameters**

TSeq	

### **Parameters**

exclude	unsigned vector of states that need to be excluded from the sampling
---------	----------------------------------------------------------------------

### Returns

Virus<TSeq>\* of the selected virus. If none selected (or none available,) returns a nullptr;

### 17.2.2.3 sample\_virus\_single()

Sample from neighbors pool of viruses (at most one)

This function samples at most one virus from the pool of viruses from its neighbors. If no virus is selected, the function returns a nullptr, otherwise it returns a pointer to the selected virus.

This can be used to build a new update function (EPI\_NEW\_UPDATEFUN.)

Tem	nlate	Parar	neters
10111	viaic	ı araı	Hetelo

### **Parameters**

p	Pointer to person	
m	Pointer to the model	

### Returns

 $\label{thm:constraint} \mbox{Virus$<$TSeq$>*$ of the selected virus. If none selected (or none available,) returns a nullptr;}$ 

# **Chapter 18**

# **Class Documentation**

# 18.1 AdjList Class Reference

# **Public Member Functions**

- AdjList (const std::vector < int > &source, const std::vector < int > &target, int size, bool directed)
   Construct a new Adj List object.
- AdjList (AdjList &&a)
- AdjList (const AdjList &a)
- AdjList & operator= (const AdjList &a)
- void read\_edgelist (std::string fn, int size, int skip=0, bool directed=true)

Read an edgelist.

- std::map< int, int > operator() (epiworld\_fast\_uint i) const
- void **print** (epiworld\_fast\_uint limit=20u) const
- size t vcount () const

Number of vertices/nodes in the network.

• size\_t ecount () const

Number of edges/arcs/ties in the network.

- std::vector< std::map< int, int > > & get\_dat ()
- bool is\_directed () const

true if the network is directed.

# 18.1.1 Constructor & Destructor Documentation

# 18.1.1.1 AdjList()

Construct a new Adj List object.

Ids in the network are assume to range from 0 to size - 1.

#### **Parameters**

source	Unsigned int vector with the source
target	Unsigned int vector with the target
size	Number of vertices in the network.
directed	Bool true if the network is directed

# 18.1.2 Member Function Documentation

# 18.1.2.1 read\_edgelist()

```
void AdjList::read_edgelist (
    std::string fn,
    int size,
    int skip = 0,
    bool directed = true ) [inline]
```

Read an edgelist.

Ids in the network are assume to range from 0 to size - 1.

#### **Parameters**

fn	Path to the file
skip	Number of lines to skip (e.g., 1 if there's a header)
directed	true if the network is directed
size	Number of vertices in the network.

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

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

# 18.2 epiworld::AdjList Class Reference

# **Public Member Functions**

- AdjList (const std::vector< int > &source, const std::vector< int > &target, int size, bool directed)
   Construct a new Adj List object.
- AdjList (AdjList &&a)
- AdjList (const AdjList &a)
- AdjList & operator= (const AdjList &a)
- void read\_edgelist (std::string fn, int size, int skip=0, bool directed=true)

Read an edgelist.

- std::map< int, int > operator() (epiworld\_fast\_uint i) const
- void **print** (epiworld\_fast\_uint limit=20u) const
- size\_t vcount () const

Number of vertices/nodes in the network.

size\_t ecount () const

Number of edges/arcs/ties in the network.

- std::vector< std::map< int, int > > & get\_dat ()
- bool is\_directed () const

true if the network is directed.

# 18.2.1 Constructor & Destructor Documentation

# 18.2.1.1 AdjList()

Construct a new Adj List object.

lds in the network are assume to range from 0 to size - 1.

### **Parameters**

source	Unsigned int vector with the source
target	Unsigned int vector with the target
size	Number of vertices in the network.
directed	Bool true if the network is directed

# 18.2.2 Member Function Documentation

# 18.2.2.1 read\_edgelist()

```
void AdjList::read_edgelist (
    std::string fn,
    int size,
    int skip = 0,
    bool directed = true ) [inline]
```

Read an edgelist.

lds in the network are assume to range from 0 to size - 1.

#### **Parameters**

fn	Path to the file	
skip	Number of lines to skip (e.g., 1 if there's a header)	
directed	cted true if the network is directed	
size	Number of vertices in the network.	

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

· epiworld.hpp

# 18.3 Agent < TSeq > Class Template Reference

Agent (agents)

#include <agent-bones.hpp>

#### **Public Member Functions**

- Agent (Agent < TSeq > &&p)
- Agent (const Agent < TSeq > &p)
- Agent < TSeq > & operator= (const Agent < TSeq > &other\_agent)
- int get\_id () const

Id of the individual.

- VirusPtr< TSeq > & get\_virus ()
- const VirusPtr< TSeq > &  $get\_virus$  () const
- ToolPtr< TSeq > & get\_tool (int i)
- Tools < TSeq > get\_tools ()
- const Tools\_const< TSeq > get\_tools () const
- size t get n tools () const noexcept
- void mutate\_virus ()
- void add\_neighbor (Agent < TSeq > &p, bool check\_source=true, bool check\_target=true)
- void swap\_neighbors (Agent < TSeq > &other, size\_t n\_this, size\_t n\_other)

Swaps neighbors between the current agent and agent other

- std::vector< Agent< TSeq > \* > get\_neighbors ()
- size t get n neighbors () const
- void change\_state (Model < TSeq > \*model, epiworld\_fast\_uint new\_state, epiworld\_fast\_int queue=0)
- const unsigned int & get\_state () const
- · void reset ()
- bool has\_tool (epiworld\_fast\_uint t) const
- bool has\_tool (std::string name) const
- bool  ${\sf has\_tool}$  (const  ${\sf Tool}{<}$   ${\sf TSeq} > {\&t}$ ) const
- · bool has\_virus (epiworld\_fast\_uint t) const
- · bool has\_virus (std::string name) const
- bool has\_virus (const Virus< TSeq > &v) const
- bool has\_entity (epiworld\_fast\_uint t) const
- · bool has\_entity (std::string name) const
- · void print (bool compressed=false) const
- Entities < TSeq > get\_entities ()
- const Entities\_const< TSeq > get\_entities () const

- const Entity < TSeq > & get\_entity (size\_t i) const
- Entity< TSeq > & get\_entity (size\_t i)
- · size t get n entities () const
- bool operator== (const Agent < TSeq > & other) const
- bool operator!= (const Agent < TSeq > & other) const

#### Add/Remove Virus/Tool

Any of these is ultimately reflected at the end of the iteration.

#### **Parameters**

tool	Tool to add
virus	Virus to add
state_new	state after the change
queue	

- void add\_tool (ToolPtr< TSeq > &tool, Model< TSeq > \*model, epiworld\_fast\_int state\_new=-99, epiworld\_fast\_int queue=-99)
- void add\_tool (const Tool < TSeq > &tool, Model < TSeq > \*model, epiworld\_fast\_int state\_new=-99, epiworld fast int queue=-99)
- void set\_virus (VirusPtr< TSeq > &virus, Model< TSeq > \*model, epiworld\_fast\_int state\_new=-99, epiworld\_fast\_int queue=-99)
- void set\_virus (const Virus < TSeq > &virus, Model < TSeq > \*model, epiworld\_fast\_int state\_new=-99, epiworld\_fast\_int queue=-99)
- void add\_entity (Entity < TSeq > &entity, Model < TSeq > \*model, epiworld\_fast\_int state\_new=-99, epiworld\_fast\_int queue=-99)
- void rm\_tool (epiworld\_fast\_uint tool\_idx, Model < TSeq > \*model, epiworld\_fast\_int state\_new=-99, epiworld\_fast\_int queue=-99)
- void rm\_tool (ToolPtr< TSeq > &tool, Model< TSeq > \*model, epiworld\_fast\_int state\_new=-99, epiworld\_fast\_int queue=-99)
- void rm\_virus (Model < TSeq > \*model, epiworld\_fast\_int state\_new=-99, epiworld\_fast\_int queue=-99)
- void rm\_entity (epiworld\_fast\_uint entity\_idx, Model < TSeq > \*model, epiworld\_fast\_int state\_new=-99, epiworld fast int queue=-99)
- void rm\_entity (Entity < TSeq > &entity, Model < TSeq > \*model, epiworld\_fast\_int state\_new=-99, epiworld fast int gueue=-99)
- void rm\_agent\_by\_virus (Model < TSeq > \*model)
   Agent removed by virus.

# Get the rates (multipliers) for the agent

# Parameters

v A pointer to a virus.

#### Returns

epiworld\_double

- epiworld\_double get\_transmission\_reduction (VirusPtr< TSeq > v, Model< TSeq > \*model)
- epiworld\_double get\_recovery\_enhancer (VirusPtr< TSeq > v, Model< TSeq > \*model)
- epiworld\_double get\_death\_reduction (VirusPtr< TSeq > v, Model< TSeq > \*model)

```
    double & operator() (size_t j)
```

Access the j-th column of the agent.

- double & operator[] (size\_t j)
- double **operator()** (size\_t j) const
- double operator[] (size\_t j) const

#### **Friends**

- class Model < TSeq >
- class Virus < TSeq >
- class Tool < TSeq >
- class Tools < TSeq >
- class Tools const< TSeq >
- class Queue < TSeq >
- class Entities < TSeq >
- class AgentsSample < TSeq >
- void default\_add\_virus (Event< TSeq > &a, Model< TSeq > \*m)
- void default\_add\_tool (Event< TSeq > &a, Model< TSeq > \*m)
- void default add entity (Event< TSeq > &a, Model< TSeq > \*m)
- void default\_rm\_virus (Event< TSeq > &a, Model< TSeq > \*m)
- void default\_rm\_tool (Event< TSeq > &a, Model< TSeq > \*m)
- void default rm entity (Event< TSeq > &a, Model< TSeq > \*m)
- void default\_change\_state (Event< TSeq > &a, Model< TSeq > \*m)

# 18.3.1 Detailed Description

```
template<typename TSeq>class Agent< TSeq>
```

Agent (agents)

**Template Parameters** 

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

# 18.3.2 Member Function Documentation

# 18.3.2.1 operator()()

Access the j-th column of the agent.

If an external array has been specified, then these two functions can be used to access additional agent's features not included in the model.

The operator[] method is with no boundary check, whereas the operator() method checks boundaries. The former can result in a segfault.

#### **Parameters**



# Returns

double&

# 18.3.2.2 swap\_neighbors()

Swaps neighbors between the current agent and agent other

#### **Parameters**

other	
n_this	
n_other	

# 18.3.3 Friends And Related Function Documentation

# 18.3.3.1 default\_rm\_entity

- < Last entity of the agent
- < Last agent of the entity

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

- include/epiworld/agent-bones.hpp
- include/epiworld/agent-meat.hpp

# 18.4 epiworld::Agent < TSeq > Class Template Reference

# Agent (agents)

#include <epiworld.hpp>

#### **Public Member Functions**

- Agent (Agent < TSeq > &&p)
- Agent (const Agent < TSeq > &p)
- Agent < TSeq > & operator= (const Agent < TSeq > &other\_agent)
- int get id () const

Id of the individual.

- VirusPtr< TSeq > & get\_virus ()
- const VirusPtr< TSeq > & get\_virus () const
- ToolPtr< TSeq > & get\_tool (int i)
- Tools < TSeq > get\_tools ()
- const Tools\_const < TSeq > get\_tools () const
- size\_t get\_n\_tools () const noexcept
- void mutate\_virus ()
- void add\_neighbor (Agent < TSeq > &p, bool check\_source=true, bool check\_target=true)
- void swap\_neighbors (Agent < TSeq > &other, size\_t n\_this, size\_t n\_other)

Swaps neighbors between the current agent and agent other

- std::vector< Agent< TSeq > \* > get\_neighbors ()
- size\_t **get\_n\_neighbors** () const
- void **change\_state** (Model < TSeq > \*model, epiworld\_fast\_uint new\_state, epiworld\_fast\_int queue=0)
- const unsigned int & get\_state () const
- · void reset ()
- · bool has tool (epiworld fast uint t) const
- · bool has\_tool (std::string name) const
- bool has\_tool (const Tool < TSeq > &t) const
- bool has\_virus (epiworld\_fast\_uint t) const
- bool has\_virus (std::string name) const
- bool has\_virus (const Virus < TSeq > &v) const
- bool has\_entity (epiworld\_fast\_uint t) const
- bool has\_entity (std::string name) const
- void print (bool compressed=false) const
- Entities < TSeq > get\_entities ()
- const Entities\_const< TSeq > get\_entities () const
- const Entity < TSeq > & get\_entity (size\_t i) const
- Entity < TSeq > & get\_entity (size\_t i)
- size\_t get\_n\_entities () const
- bool operator== (const Agent < TSeq > &other) const
- bool operator!= (const Agent < TSeq > & other) const

# Add/Remove Virus/Tool

Any of these is ultimately reflected at the end of the iteration.

### **Parameters**

tool virus	Tool to add  Virus to add
viius	VIIUS IO AUU
state_new	state after the change
Gளு <b>டிரு</b> by Doxygen	

void add\_tool (ToolPtr< TSeq > &tool, Model< TSeq > \*model, epiworld\_fast\_int state\_new=-99, epiworld\_fast\_int queue=-99)

- void add\_tool (const Tool < TSeq > &tool, Model < TSeq > \*model, epiworld\_fast\_int state\_new=-99, epiworld\_fast\_int queue=-99)
- void set\_virus (VirusPtr< TSeq > &virus, Model< TSeq > \*model, epiworld\_fast\_int state\_new=-99, epiworld fast int queue=-99)
- void set\_virus (const Virus < TSeq > &virus, Model < TSeq > \*model, epiworld\_fast\_int state\_new=-99, epiworld\_fast\_int queue=-99)
- void add\_entity (Entity < TSeq > &entity, Model < TSeq > \*model, epiworld\_fast\_int state\_new=-99, epiworld fast int gueue=-99)
- void rm\_tool (epiworld\_fast\_uint tool\_idx, Model < TSeq > \*model, epiworld\_fast\_int state\_new=-99, epiworld fast int gueue=-99)
- void rm\_tool (ToolPtr< TSeq > &tool, Model< TSeq > \*model, epiworld\_fast\_int state\_new=-99, epiworld fast int gueue=-99)
- void rm\_virus (Model < TSeq > \*model, epiworld\_fast\_int state\_new=-99, epiworld\_fast\_int queue=-99)
- void rm\_entity (epiworld\_fast\_uint entity\_idx, Model < TSeq > \*model, epiworld\_fast\_int state\_new=-99, epiworld fast int gueue=-99)
- void rm\_entity (Entity < TSeq > &entity, Model < TSeq > \*model, epiworld\_fast\_int state\_new=-99, epiworld\_fast\_int queue=-99)
- void rm agent by virus (Model < TSeq > \*model)

Agent removed by virus.

#### Get the rates (multipliers) for the agent

#### **Parameters**

v A pointer to a virus.

### Returns

epiworld double

- epiworld\_double get\_transmission\_reduction (VirusPtr< TSeq > v, Model< TSeq > \*model)
- epiworld\_double get\_recovery\_enhancer (VirusPtr< TSeq > v, Model< TSeq > \*model)
- epiworld\_double get\_death\_reduction (VirusPtr< TSeq > v, Model< TSeq > \*model)
- double & operator() (size\_t j)

Access the j-th column of the agent.

- double & operator[] (size\_t j)
- double operator() (size\_t j) const
- double operator[] (size\_t j) const

# **Friends**

- class Model < TSeq >
- class Virus < TSeq >
- class Tool < TSeq >
- class Tools< TSeq >
- class Tools\_const< TSeq >
- class Queue < TSeq >
- class Entities < TSeq >
- class AgentsSample < TSeq >

- void default\_add\_virus (Event< TSeq > &a, Model< TSeq > \*m)
- void default\_add\_tool (Event< TSeq > &a, Model< TSeq > \*m)
- void default\_add\_entity (Event< TSeq > &a, Model< TSeq > \*m)
- void default\_rm\_virus (Event< TSeq > &a, Model< TSeq > \*m)
- void default\_rm\_tool (Event< TSeq > &a, Model< TSeq > \*m)
- void default\_rm\_entity (Event< TSeq > &a, Model< TSeq > \*m)
- void default\_change\_state (Event< TSeq > &a, Model< TSeq > \*m)

# 18.4.1 Detailed Description

```
\label{template} \begin{tabular}{ll} template < typename TSeq > \\ class epiworld::Agent < TSeq > \\ \end{tabular}
```

Agent (agents)

#### **Template Parameters**

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

# 18.4.2 Member Function Documentation

# 18.4.2.1 operator()()

Access the j-th column of the agent.

If an external array has been specified, then these two functions can be used to access additional agent's features not included in the model.

The operator[] method is with no boundary check, whereas the operator() method checks boundaries. The former can result in a segfault.

#### **Parameters**



#### Returns

double&

# 18.4.2.2 swap\_neighbors()

Swaps neighbors between the current agent and agent other

#### **Parameters**

other	
n_this	
n_other	

# 18.4.3 Friends And Related Function Documentation

#### 18.4.3.1 default\_rm\_entity

- < Last entity of the agent
- < Last agent of the entity

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

· epiworld.hpp

# 18.5 AgentsSample < TSeq > Class Template Reference

Sample of agents.

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

#### **Public Member Functions**

• AgentsSample ()=delete

Default constructor.

AgentsSample (const AgentsSample < TSeq > &a)=delete

Copy constructor.

AgentsSample (AgentsSample < TSeq > &&a)=delete

Move constructor.

- **AgentsSample** (Model < TSeq > &model\_, size\_t n, std::vector < size\_t > states\_={}, bool truncate=false)
- AgentsSample (Model < TSeq > \*model, Entity < TSeq > &entity\_, size\_t n, std::vector < size\_t > states ←
   \_={}, bool truncate=false)
- AgentsSample (Model < TSeq > \*model, Agent < TSeq > &agent\_, size\_t n, std::vector < size\_t > states ←
   \_={}, bool truncate=false)

Sample from the agent's entities.

- std::vector < Agent < TSeq > \* >::iterator begin ()
- std::vector < Agent < TSeq > \* >::iterator end ()
- Agent < TSeq > \* operator[] (size\_t n)
- Agent< TSeq > \* operator() (size\_t n)
- size\_t size () const noexcept

# 18.5.1 Detailed Description

```
template<typename TSeq = EPI_DEFAULT_TSEQ> class AgentsSample< TSeq >
```

Sample of agents.

This class allows sampling agents from Entity<TSeq> and Model<TSeq>.

**Template Parameters** 

TSeq	

#### 18.5.2 Constructor & Destructor Documentation

#### 18.5.2.1 AgentsSample()

Sample from the agent's entities.

For example, how many individuals the agent contacts in a given point in time.

# **Template Parameters**

TSeq
------

#### **Parameters**

agent⊷	
_	
n	Sample size
truncate	If the agent has fewer than $n$ connections, then truncate = true will automatically reduce the number of possible samples. Otherwise, if false, then it returns an error.

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

• include/epiworld/agentssample-bones.hpp

# 18.6 epiworld::AgentsSample < TSeq > Class Template Reference

Sample of agents.

#include <epiworld.hpp>

#### **Public Member Functions**

• AgentsSample ()=delete

Default constructor.

AgentsSample (const AgentsSample < TSeq > &a)=delete

Copy constructor.

AgentsSample (AgentsSample < TSeq > &&a)=delete

Move constructor.

- **AgentsSample** (Model < TSeq > &model\_, size\_t n, std::vector < size\_t > states\_={}, bool truncate=false)
- AgentsSample (Model < TSeq > \*model, Entity < TSeq > &entity\_, size\_t n, std::vector < size\_t > states ←
   \_={}, bool truncate=false)
- AgentsSample (Model < TSeq > \*model, Agent < TSeq > &agent\_, size\_t n, std::vector < size\_t > states ←
   \_={}, bool truncate=false)

Sample from the agent's entities.

- std::vector< Agent< TSeq > \* >::iterator begin ()
- std::vector < Agent < TSeq > \* >::iterator end ()
- Agent< TSeq > \* operator[] (size\_t n)
- Agent< TSeq > \* operator() (size\_t n)
- size\_t size () const noexcept

# 18.6.1 Detailed Description

template<typename TSeq = EPI\_DEFAULT\_TSEQ> class epiworld::AgentsSample< TSeq >

Sample of agents.

This class allows sampling agents from Entity<TSeq> and Model<TSeq>.

**Template Parameters** 

# 18.6.2 Constructor & Destructor Documentation

### 18.6.2.1 AgentsSample()

Sample from the agent's entities.

For example, how many individuals the agent contacts in a given point in time.

# **Template Parameters**

|--|

# **Parameters**

agent⊷	
_	
n	Sample size
truncate	If the agent has fewer than $n$ connections, then truncate = true will automatically reduce the number of possible samples. Otherwise, if false, then it returns an error.

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

· epiworld.hpp

# 18.7 DataBase < TSeq > Class Template Reference

Statistical data about the process.

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

#### **Public Member Functions**

- DataBase (Model < TSeq > &m)
- DataBase (const DataBase < TSeq > &db)
- void record virus (Virus < TSeq > &v)

Registering a new variant.

- void record\_tool (Tool < TSeq > &t)
- void set\_seq\_hasher (std::function< std::vector< int >(TSeq)> fun)
- · void reset ()
- 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\_virus\_info, std::string fn\_virus\_hist, std::string fn\_tool\_info, std::string fn\_tool\_hist, std::string fn\_transmission, std::string fn\_transmiss
- void record\_transmission (int i, int j, int virus, int i\_expo\_date)
- size\_t get\_n\_viruses () const

Get the number of viruses.

size\_t get\_n\_tools () const

Get the number of tools.

- void set\_user\_data (std::vector< std::string > names)
- void add\_user\_data (std::vector< epiworld\_double > x)
- void add user data (epiworld fast uint j, epiworld double x)
- UserData < TSeq > & get user data ()
- std::vector< epiworld\_double > get\_transition\_probability (bool print=true, bool normalize=true) const

Calculates the transition probabilities.

- bool operator== (const DataBase< TSeg > &other) const
- bool operator!= (const DataBase < TSeq > &other) const
- bool operator== (const DataBase< std::vector< int >> &other) const
- bool operator== (const DataBase< std::vector< int >> &other) const

# Get recorded information from the model

#### **Parameters**

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

#### Returns

```
In get_today_total, the current counts of what.
In get_today_virus, the current counts of what for each virus.
In get_hist_total, the time series of what
In get_hist_virus, the time series of what for each virus.
In get_hist_total_date and get_hist_virus_date the corresponding date
```

- int get today total (const std::string &what) const
- · int get today total (const epiworld fast uint &what) const
- $\bullet \ \ \ \ \text{void } \textbf{get\_today\_total} \ (\text{std::vector} < \ \text{std::string} > * \ \text{state=nullptr}, \ \text{std::vector} < \ \text{int} > * \ \text{counts=nullptr}) \ \text{const}$
- void get\_today\_virus (std::vector< std::string > &state, std::vector< int > &id, std::vector< int > &counts) const
- void get\_today\_transition\_matrix (std::vector< int > &counts) const
- void get\_hist\_total (std::vector< int > \*date, std::vector< std::string > \*state, std::vector< int > \*counts) const

- void get\_hist\_virus (std::vector< int > &date, std::vector< int > &id, std::vector< std::string > &state, std::vector< int > &counts) const
- void get\_hist\_tool (std::vector< int > &date, std::vector< int > &id, std::vector< std::string > &state, std::vector< int > &counts) const
- void get\_hist\_transition\_matrix (std::vector< std::string > &state\_from, std::vector< std::string > &state\_to, std::vector< int > &date, std::vector< int > &counts, bool skip\_zeros) const
- $\hbox{$\bullet$ void $\tt get\_transmissions (std::vector< int> \&date, std::vector< int> \&source, std::vector< int> \&target, std::vector< int> \&virus, std::vector< int> \&source\_exposure\_date) const } \\$

Get the transmissions object.

- void **get\_transmissions** (int \*date, int \*source, int \*target, int \*virus, int \*source\_exposure\_date) const
- MapVec\_type< int, int > get\_reproductive\_number () const
- Computes the reproductive number of each case.

   void **get\_reproductive\_number** (std::string fn) const
- void get\_generation\_time (std::vector< int > &agent\_id, std::vector< int > &virus\_id, std::vector< int > &time, std::vector< int > &gentime) const

Get the generation time.

• void get\_generation\_time (std::string fn) const

Write the generation time to a file.

### **Friends**

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

# 18.7.1 Detailed Description

template<typename TSeq> class DataBase< TSeq >

Statistical data about the process.

**Template Parameters** 

TSeq	

# 18.7.2 Member Function Documentation

# 18.7.2.1 get\_generation\_time()

```
template<typename TSeq >
void DataBase< TSeq >::get_generation_time (
    std::vector< int > & agent_id,
    std::vector< int > & virus_id,
    std::vector< int > & time,
    std::vector< int > & gentime ) const [inline]
```

Get the generation time.

Calculates the generating time

**Parameters** 

ent_id,virus_id,time,gentin	vectors where to save the values
-----------------------------	----------------------------------

The generation time is the time between the infection of the source and the infection of the target.

# 18.7.2.2 get\_reproductive\_number()

```
template<typename TSeq >
MapVec_type< int, int > DataBase< TSeq >::get_reproductive_number [inline]
```

Computes the reproductive number of each case.

By definition, whereas it computes R0 (basic reproductive number) or Rt/R (the effective reproductive number) will depend on whether the virus is allowed to circulate naïvely or not, respectively.

# **Parameters**

fn File where to write out the reproductive number.

In the case of MapVec\_type<int, int>, the key is a vector of 3 integers:

- Virus id
- Source id
- · Date when the source was infected

# 18.7.2.3 get\_transition\_probability()

Calculates the transition probabilities.

#### **Parameters**

print	Print the transition matrix.
normalize	Normalize the transition matrix. Otherwise, it returns raw counts.

The transition matrix is the matrix of the counts of transitions from one state to another. So the ij-th element of the matrix is the number of transitions from state i to state j (when not normalized), or the probability of transitioning from state i to state j (when normalized).

#### Returns

```
std::vector< epiworld_double >
```

# 18.7.2.4 get\_transmissions()

```
template<typename TSeq >
void DataBase< TSeq >::get_transmissions (
    std::vector< int > & date,
    std::vector< int > & source,
    std::vector< int > & target,
    std::vector< int > & virus,
    std::vector< int > & source_exposure_date ) const [inline]
```

Get the transmissions object.

#### **Parameters**

date	
source	
target	
virus	
source_exposure_date	

# **18.7.2.5** operator==() [1/3]

```
< Date of the transmission eve,
< Id of the sour,
< Id of the targ,
< Id of the varia,
< Date when the source acquired the varia,
18.7.2.6 operator==() [2/3]
bool DataBase< std::vector< int > >::operator== (
              const DataBase< std::vector< int >> & other ) const [inline]
< Date of the transmission eve,
< Id of the sour,
< Id of the targ,
< Id of the varia,
< Date when the source acquired the varia,
18.7.2.7 operator==() [3/3]
template<typename TSeq >
bool DataBase< TSeq >::operator== (
             const DataBase< TSeq > & other ) const [inline]
< Date of the transmission eve
< Id of the sour
< Id of the targ
< Id of the varia
< Date when the source acquired the varia
```

# 18.7.2.8 record\_virus()

Registering a new variant.

#### **Parameters**

*v* Pointer to the new virus. Since viruses are originated in the agent, the numbers simply move around. From the parent virus to the new virus. 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

# 18.8 epiworld::DataBase < TSeq > Class Template Reference

Statistical data about the process.

```
#include <epiworld.hpp>
```

#### **Public Member Functions**

- DataBase (Model < TSeq > &m)
- DataBase (const DataBase < TSeq > &db)
- void record\_virus (Virus < TSeq > &v)

Registering a new variant.

- void record\_tool (Tool < TSeq > &t)
- void set seg hasher (std::function< std::vector< int >(TSeg)> fun)
- · void reset ()
- 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\_virus\_info, std::string fn\_virus\_hist, std::string fn\_tool\_info, std::string fn\_tool\_hist, std::string fn\_total\_hist, std::string fn\_transmission, std::string fn\_transmissio
- void record\_transmission (int i, int j, int virus, int i\_expo\_date)
- size\_t get\_n\_viruses () const

Get the number of viruses.

· size\_t get\_n\_tools () const

Get the number of tools.

- void set\_user\_data (std::vector< std::string > names)
- void add\_user\_data (std::vector< epiworld\_double > x)
- void add user data (epiworld fast uint j, epiworld double x)
- UserData < TSeq > & get\_user\_data ()
- std::vector< epiworld\_double > get\_transition\_probability (bool print=true, bool normalize=true) const Calculates the transition probabilities.
- bool operator== (const DataBase < TSeq > &other) const
- bool operator!= (const DataBase< TSeq > &other) const

# Get recorded information from the model

#### **Parameters**

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

#### Returns

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

In get\_today\_virus, the current counts of what for each virus.

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

In get\_hist\_virus, the time series of what for each virus.

In get\_hist\_total\_date and get\_hist\_virus\_date the corresponding date

- int get\_today\_total (const std::string &what) const
- int get\_today\_total (const epiworld\_fast\_uint &what) const
- void get\_today\_total (std::vector< std::string > \*state=nullptr, std::vector< int > \*counts=nullptr) const
- void get\_today\_virus (std::vector< std::string > &state, std::vector< int > &id, std::vector< int > &counts) const
- void get\_today\_transition\_matrix (std::vector< int > &counts) const
- void get\_hist\_total (std::vector< int > \*date, std::vector< std::string > \*state, std::vector< int > \*counts) const
- void get\_hist\_virus (std::vector< int > &date, std::vector< int > &id, std::vector< std::string > &state, std::vector< int > &counts) const
- void get\_hist\_tool (std::vector< int > &date, std::vector< int > &id, std::vector< std::string > &state, std::vector< int > &counts) const
- void get\_hist\_transition\_matrix (std::vector< std::string > &state\_from, std::vector< std::string > &state\_to, std::vector< int > &date, std::vector< int > &counts, bool skip\_zeros) const
- void get\_transmissions (std::vector< int > &date, std::vector< int > &source, std::vector< int > &target, std::vector< int > &virus, std::vector< int > &source\_exposure\_date) const

Get the transmissions object.

- void get transmissions (int \*date, int \*source, int \*target, int \*virus, int \*source exposure date) const
- MapVec\_type < int, int > get\_reproductive\_number () const
   Computes the reproductive number of each case.
- · void get\_reproductive\_number (std::string fn) const
- void get\_generation\_time (std::vector< int > &agent\_id, std::vector< int > &virus\_id, std::vector< int > &time, std::vector< int > &gentime) const

Get the generation time.

· void get\_generation\_time (std::string fn) const

Write the generation time to a file.

#### **Friends**

- class Model < TSeq >
- void default\_add\_virus (Event< TSeq > &a, Model< TSeq > \*m)
- void default add tool (Event< TSeq > &a, Model< TSeq > \*m)
- void default\_rm\_virus (Event< TSeq > &a, Model< TSeq > \*m)
- void default\_rm\_tool (Event< TSeq > &a, Model< TSeq > \*m)
- void default\_change\_state (Event< TSeq > &a, Model< TSeq > \*m)

# 18.8.1 Detailed Description

```
\label{template} \begin{tabular}{ll} template < typename TSeq > \\ class epiworld::DataBase < TSeq > \\ \end{tabular}
```

Statistical data about the process.

**Template Parameters** 

# 18.8.2 Member Function Documentation

# 18.8.2.1 get\_generation\_time()

Get the generation time.

Calculates the generating time

**Parameters** 

```
agent_id,virus_id,time,gentime vectors where to save the values
```

The generation time is the time between the infection of the source and the infection of the target.

# 18.8.2.2 get\_reproductive\_number()

```
template<typename TSeq >
MapVec_type< int, int > DataBase< TSeq >::get_reproductive_number [inline]
```

Computes the reproductive number of each case.

By definition, whereas it computes R0 (basic reproductive number) or Rt/R (the effective reproductive number) will depend on whether the virus is allowed to circulate naïvely or not, respectively.

#### **Parameters**

```
fn File where to write out the reproductive number.
```

In the case of MapVec\_type<int, int>, the key is a vector of 3 integers:

- · Virus id
- · Source id
- · Date when the source was infected

# 18.8.2.3 get\_transition\_probability()

Calculates the transition probabilities.

#### **Parameters**

print	Print the transition matrix.
normalize	Normalize the transition matrix. Otherwise, it returns raw counts.

The transition matrix is the matrix of the counts of transitions from one state to another. So the ij-th element of the matrix is the number of transitions from state i to state j (when not normalized), or the probability of transitioning from state i to state j (when normalized).

### Returns

```
std::vector< epiworld_double >
```

# 18.8.2.4 get\_transmissions()

```
template<typename TSeq >
void DataBase< TSeq >::get_transmissions (
    std::vector< int > & date,
    std::vector< int > & source,
    std::vector< int > & target,
    std::vector< int > & virus,
    std::vector< int > & source_exposure_date ) const [inline]
```

Get the transmissions object.

#### **Parameters**

date	
source	
target	
virus	
source_exposure_date	

#### 18.8.2.5 operator==()

< Date when the source acquired the varia

# 18.8.2.6 record\_virus()

Registering a new variant.

#### **Parameters**

Pointer to the new virus. Since viruses are originated in the agent, the numbers simply move around. From the parent virus to the new virus. And the total number of infected does not change.

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

· epiworld.hpp

# 18.9 Entities < TSeq > Class Template Reference

Set of Entities (useful for building iterators)

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

# **Public Member Functions**

```
    Entities (Agent< TSeq > &p)
    std::vector< Entity< TSeq > * >::iterator begin ()
    std::vector< Entity< TSeq > * >::iterator end ()
```

• Entity< TSeq > & operator() (size\_t i)

- Entity < TSeq > & operator[] (size\_t i)
- size\_t size () const noexcept
- bool operator== (const Entities < TSeq > &other) const

#### **Friends**

- class Entity< TSeq >
- class Agent < TSeq >

# 18.9.1 Detailed Description

```
template<typename TSeq> class Entities< TSeq>
```

Set of Entities (useful for building iterators)

**Template Parameters** 



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

- include/epiworld/agent-bones.hpp
- · include/epiworld/entities-bones.hpp

# 18.10 epiworld::Entities < TSeq > Class Template Reference

Set of Entities (useful for building iterators)

```
#include <epiworld.hpp>
```

#### **Public Member Functions**

- Entities (Agent < TSeq > &p)
- std::vector< Entity< TSeq > \* >::iterator begin ()
- std::vector< Entity< TSeq > \* >::iterator end ()
- Entity < TSeq > & operator() (size\_t i)
- Entity< TSeq > & operator[] (size t i)
- size\_t size () const noexcept
- bool operator== (const Entities < TSeq > &other) const

#### **Friends**

- class Entity< TSeq >
- class Agent < TSeq >

# 18.10.1 Detailed Description

 $\label{template} \begin{tabular}{ll} template < typename TSeq > \\ class epiworld::Entities < TSeq > \\ \end{tabular}$ 

Set of Entities (useful for building iterators)

**Template Parameters** 



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

· epiworld.hpp

# 18.11 Entities\_const < TSeq > Class Template Reference

Set of Entities (const) (useful for iterators)

#include <entities-bones.hpp>

# **Public Member Functions**

- Entities\_const (const Agent < TSeq > &p)
- std::vector< Entity< TSeq > \* >::const\_iterator begin ()
- std::vector< Entity< TSeq > \* >::const\_iterator end ()
- const Entity < TSeq > & operator() (size\_t i)
- const Entity < TSeq > & operator[] (size\_t i)
- size\_t size () const noexcept
- bool operator== (const Entities\_const< TSeq > &other) const

# **Friends**

- class Virus < TSeq >
- class Agent < TSeq >

# 18.11.1 Detailed Description

template<typename TSeq> class Entities\_const< TSeq >

Set of Entities (const) (useful for iterators)

# **Template Parameters**

TSeq	
,	

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

• include/epiworld/entities-bones.hpp

# 18.12 epiworld::Entities\_const< TSeq > Class Template Reference

Set of Entities (const) (useful for iterators)

```
#include <epiworld.hpp>
```

#### **Public Member Functions**

- Entities\_const (const Agent < TSeq > &p)
- std::vector< Entity< TSeq > \* >::const\_iterator begin ()
- std::vector< Entity< TSeq > \* >::const\_iterator end ()
- const Entity < TSeq > & operator() (size t i)
- const Entity < TSeq > & operator[] (size\_t i)
- size\_t size () const noexcept
- bool **operator==** (const Entities\_const< TSeq > &other) const

# **Friends**

- class Virus < TSeq >
- class Agent < TSeq >

# 18.12.1 Detailed Description

template<typename TSeq>
class epiworld::Entities\_const< TSeq>

Set of Entities (const) (useful for iterators)

**Template Parameters** 

TSeq	

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

· epiworld.hpp

# 18.13 Entity < TSeq > Class Template Reference

#### **Public Member Functions**

- Entity (std::string name, EntityToAgentFun< TSeq > fun=nullptr)
   Constructs an Entity object.
- void add\_agent (Agent < TSeq > &p, Model < TSeq > \*model)
- void add agent (Agent < TSeq > \*p, Model < TSeq > \*model)
- void rm\_agent (size\_t idx, Model < TSeq > \*model)
- size\_t size () const noexcept
- void set\_location (std::vector< epiworld\_double > loc)
- std::vector< epiworld double > & get location ()
- std::vector< size\_t >::iterator begin ()
- std::vector< size\_t >::iterator end ()
- std::vector< size t >::const iterator begin () const
- std::vector< size t >::const iterator end () const
- size\_t operator[] (size\_t i)
- int get\_id () const noexcept
- · const std::string & get\_name () const noexcept
- void **set\_state** (epiworld\_fast\_int init, epiworld\_fast\_int post)
- void set\_queue (epiworld\_fast\_int init, epiworld\_fast\_int post)
- void get state (epiworld fast int \*init, epiworld fast int \*post)
- void get\_queue (epiworld\_fast\_int \*init, epiworld\_fast\_int \*post)
- · void reset ()
- bool operator== (const Entity < TSeq > &other) const
- bool operator!= (const Entity < TSeq > & other) const

#### **Entity distribution**

These functions are used for distributing agents among entities. The idea is to have a flexible way of distributing agents among entities.

- void distribute (Model < TSeq > \*model)
- std::vector< size\_t > & get\_agents ()
- · void print () const
- void set\_distribution (EntityToAgentFun < TSeq > fun)

# **Friends**

- class Agent < TSeq >
- class AgentsSample < TSeq >
- class Model < TSeq >
- void default\_add\_entity (Event< TSeq > &a, Model< TSeq > \*m)
- void default\_rm\_entity (Event< TSeq > &a, Model< TSeq > \*m)

# 18.13.1 Constructor & Destructor Documentation

#### 18.13.1.1 Entity()

Constructs an Entity object.

This constructor initializes an Entity object with the specified parameters.

#### **Parameters**

name	The name of the entity.
fun	A function pointer to a function that maps the entity to an agent.

# 18.13.2 Friends And Related Function Documentation

#### 18.13.2.1 default rm entity

- < Last agent of the entity
- < Last entity of the agent
- < Last agent of the entity

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

- include/epiworld/agent-bones.hpp
- include/epiworld/entity-bones.hpp
- include/epiworld/entity-meat.hpp

# 18.14 epiworld::Entity < TSeq > Class Template Reference

# **Public Member Functions**

```
    Entity (std::string name, EntityToAgentFun < TSeq > fun=nullptr)
    Constructs an Entity object.
```

```
    void add_agent (Agent < TSeq > &p, Model < TSeq > *model)
```

- void add\_agent (Agent < TSeq > \*p, Model < TSeq > \*model)
- void rm\_agent (size\_t idx, Model < TSeq > \*model)
- size\_t size () const noexcept
- void set\_location (std::vector< epiworld\_double > loc)
- std::vector< epiworld\_double > & get\_location ()
- std::vector< size\_t >::iterator begin ()
- std::vector< size\_t >::iterator end ()
- std::vector< size\_t >::const\_iterator begin () const
- std::vector< size\_t >::const\_iterator end () const
- size\_t operator[] (size\_t i)
- int get\_id () const noexcept
- const std::string & get\_name () const noexcept

- void **set\_state** (epiworld\_fast\_int init, epiworld\_fast\_int post)
- void **set\_queue** (epiworld\_fast\_int init, epiworld\_fast\_int post)
- void get\_state (epiworld fast int \*init, epiworld fast int \*post)
- void **get\_queue** (epiworld\_fast\_int \*init, epiworld\_fast\_int \*post)
- · void reset ()
- bool operator== (const Entity < TSeq > &other) const
- bool operator!= (const Entity < TSeq > &other) const

# **Entity distribution**

These functions are used for distributing agents among entities. The idea is to have a flexible way of distributing agents among entities.

- void distribute (Model < TSeq > \*model)
- std::vector< size t > & get\_agents ()
- void print () const
- void set distribution (EntityToAgentFun< TSeq > fun)

#### **Friends**

- class Agent < TSeq >
- class AgentsSample < TSeq >
- class Model < TSeq >
- void default\_add\_entity (Event< TSeq > &a, Model< TSeq > \*m)
- void default\_rm\_entity (Event< TSeq > &a, Model< TSeq > \*m)

# 18.14.1 Constructor & Destructor Documentation

#### 18.14.1.1 Entity()

Constructs an Entity object.

This constructor initializes an Entity object with the specified parameters.

#### Parameters

name	The name of the entity.
fun	A function pointer to a function that maps the entity to an agent.

# 18.14.2 Friends And Related Function Documentation

#### 18.14.2.1 default\_rm\_entity

- < Last entity of the agent
- < Last agent of the entity

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

· epiworld.hpp

# 18.15 epiworld::Event < TSeq > Struct Template Reference

Event data for update an agent.

```
#include <epiworld.hpp>
```

### **Public Member Functions**

Event (Agent < TSeq > \*agent\_, VirusPtr < TSeq > &virus\_, ToolPtr < TSeq > &tool\_, Entity < TSeq > \*entity\_, epiworld\_fast\_int new\_state\_, epiworld\_fast\_int queue\_, EventFun < TSeq > &call\_, int idx\_agent ← \_\_, int idx\_object\_)

Construct a new Event object.

# **Public Attributes**

- Agent < TSeq > \* agent
- VirusPtr< TSeq > virus
- ToolPtr< TSeq > tool
- Entity < TSeq > \* entity
- epiworld\_fast\_int new\_state
- · epiworld\_fast\_int queue
- EventFun< TSeq> call
- int idx\_agent
- · int idx\_object

# 18.15.1 Detailed Description

```
template<typename TSeq = EPI_DEFAULT_TSEQ> struct epiworld::Event< TSeq >
```

Event data for update an agent.

# **Template Parameters**

#### 18.15.2 Constructor & Destructor Documentation

### 18.15.2.1 Event()

Construct a new Event object.

All the parameters are rather optional.

#### **Parameters**

agent_	Agent over who the action will happen
virus_	Virus to add
tool_	Tool to add
virus_idx	Index of virus to be removed (if needed)
tool_idx	Index of tool to be removed (if needed)
new_←	Next state
state_	
queue_	Efect on the queue
call_	The action call (if needed)
idx_⊷	Location of agent in object.
agent_	
idx_⊷	Location of object in agent.
object_	

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

· epiworld.hpp

# 18.16 Event < TSeq > Struct Template Reference

Event data for update an agent.

#include <config.hpp>

Collaboration diagram for Event < TSeq >:



# **Public Member Functions**

Event (Agent < TSeq > \*agent\_, VirusPtr < TSeq > &virus\_, ToolPtr < TSeq > &tool\_, Entity < TSeq > \*entity\_, epiworld\_fast\_int new\_state\_, epiworld\_fast\_int queue\_, EventFun < TSeq > &call\_, int idx\_agent ← \_\_, int idx\_object\_)

Construct a new Event object.

#### **Public Attributes**

- Agent < TSeq > \* agent
- VirusPtr< TSeq > virus
- ToolPtr< TSeq> tool
- Entity < TSeq > \* entity
- epiworld\_fast\_int new\_state
- epiworld\_fast\_int queue
- EventFun< TSeq > call
- int idx\_agent
- int idx\_object

# 18.16.1 Detailed Description

template<typename TSeq = EPI\_DEFAULT\_TSEQ> struct Event< TSeq >

Event data for update an agent.

**Template Parameters** 

TSeq	

# 18.16.2 Constructor & Destructor Documentation

# 18.16.2.1 Event()

Construct a new **Event** object.

All the parameters are rather optional.

#### **Parameters**

agent_	Agent over who the action will happen
virus_	Virus to add
tool_	Tool to add
virus_idx	Index of virus to be removed (if needed)
tool_idx	Index of tool to be removed (if needed)
new_←	Next state
state_	
queue_	Efect on the queue
call_	The action call (if needed)
idx_←	Location of agent in object.
agent_	
idx_←	Location of object in agent.
object_	

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

- · include/epiworld/agent-bones.hpp
- include/epiworld/config.hpp

# 18.17 epiworld::GlobalEvent< TSeq > Class Template Reference

Template for a Global Event.

```
#include <epiworld.hpp>
```

# **Public Member Functions**

- GlobalEvent (GlobalFun < TSeq > fun, std::string name, int day=-99)
   Construct a new Global Event object.
- void operator() (Model < TSeq > \*m, int day)
- void set\_name (std::string name)
- std::string get\_name () const
- void set\_day (int day)
- int get\_day () const
- · void print () const
- bool operator== (const GlobalEvent < TSeq > &other) const
- bool operator!= (const GlobalEvent < TSeq > &other) const

# 18.17.1 Detailed Description

```
\label{template} \begin{tabular}{ll} template < typename TSeq > \\ class epiworld::GlobalEvent < TSeq > \\ \end{tabular}
```

Template for a Global Event.

Global events are functions that Model<TSeq> executes at the end of a day.

# 18.17.2 Constructor & Destructor Documentation

#### 18.17.2.1 GlobalEvent()

Construct a new Global Event object.

#### **Parameters**

fun	A function that takes a Model $<$ TSeq $> *$ as argument and returns void.
name	A descriptive name for the action.
day	The day when the action will be executed. If negative, it will be executed every day.

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

· epiworld.hpp

# 18.18 GlobalEvent < TSeq > Class Template Reference

Template for a Global Event.

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

#### **Public Member Functions**

- GlobalEvent (GlobalFun < TSeq > fun, std::string name, int day=-99)
   Construct a new Global Event object.
- void operator() (Model < TSeq > \*m, int day)
- void set\_name (std::string name)
- std::string get\_name () const
- void set\_day (int day)
- int get\_day () const
- · void print () const
- bool operator== (const GlobalEvent < TSeq > &other) const
- bool operator!= (const GlobalEvent< TSeq > &other) const

### 18.18.1 Detailed Description

```
template<typename TSeq> class GlobalEvent< TSeq>
```

Template for a Global Event.

Global events are functions that Model<TSeq> executes at the end of a day.

### 18.18.2 Constructor & Destructor Documentation

### 18.18.2.1 GlobalEvent()

Construct a new Global Event object.

#### **Parameters**

fun	A function that takes a Model <tseq> * as argument and returns void.</tseq>
name	A descriptive name for the action.
day	The day when the action will be executed. If negative, it will be executed every day.

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

- include/epiworld/globalevent-bones.hpp
- include/epiworld/globalevent-meat.hpp

## 18.19 epiworld::LFMCMC< TData > Class Template Reference

Likelihood-Free Markov Chain Monte Carlo.

```
#include <epiworld.hpp>
```

### **Public Member Functions**

- void run (std::vector< epiworld\_double > params\_init\_, size\_t n\_samples\_, epiworld\_double epsilon\_, int seed=-1)
- LFMCMC (const TData &observed\_data\_)
- void set\_observed\_data (const TData &observed\_data\_)
- void set proposal fun (LFMCMCProposalFun < TData > fun)
- void set\_simulation\_fun (LFMCMCSimFun < TData > fun)
- void set\_summary\_fun (LFMCMCSummaryFun < TData > fun)
- void set kernel fun (LFMCMCKernelFun < TData > fun)
- void set\_params\_names (std::vector< std::string > names)
- void set\_stats\_names (std::vector< std::string > names)
- size\_t get\_n\_samples () const
- size\_t get\_n\_stats () const
- size\_t get\_n\_params () const
- epiworld\_double get\_epsilon () const
- const std::vector< epiworld\_double > & get\_initial\_params () const
- const std::vector< epiworld\_double > & get\_current\_proposed\_params () const
- const std::vector< epiworld\_double > &  $get\_current\_accepted\_params$  () const
- const std::vector< epiworld\_double > & <code>get\_current\_proposed\_stats</code> () const
- const std::vector< epiworld double > & get\_current\_accepted\_stats () const
- const std::vector< epiworld\_double > & get\_observed\_stats () const
- const std::vector< epiworld\_double > & get\_all\_sample\_params () const
- const std::vector< epiworld\_double > & get\_all\_sample\_stats () const
- const std::vector< bool > & get\_all\_sample\_acceptance () const
- const std::vector< epiworld\_double > & get\_all\_sample\_drawn\_prob () const
- const std::vector< epiworld\_double > & get\_all\_sample\_kernel\_scores () const
- const std::vector< epiworld\_double > & get\_all\_accepted\_params () const
- const std::vector< epiworld\_double > &  ${\tt get\_all\_accepted\_stats}$  () const
- const std::vector< epiworld double > & get all accepted kernel scores () const
- std::vector< TData > \* get simulated data () const
- std::vector< epiworld double > get\_mean\_params ()
- std::vector< epiworld\_double > get\_mean\_stats ()
- LFMCMC< TData > & verbose\_off ()
- LFMCMC< TData > & verbose\_on ()
- void print (size\_t burnin=0u) const

### Random number generation

#### **Parameters**

eng

- void set\_rand\_engine (std::shared\_ptr< std::mt19937 > &eng)
- std::shared\_ptr< std::mt19937 > & get\_rand\_endgine ()
- void seed (epiworld\_fast\_uint s)
- void **set\_rand\_gamma** (epiworld\_double alpha, epiworld\_double beta)
- epiworld double runif ()
- epiworld double rnorm ()
- epiworld double rgamma ()
- epiworld double runif (epiworld double lb, epiworld double ub)
- epiworld double rnorm (epiworld double mean, epiworld double sd)
- epiworld\_double **rgamma** (epiworld\_double alpha, epiworld\_double beta)

### 18.19.1 Detailed Description

template<typename TData> class epiworld::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:

• epiworld.hpp

# 18.20 LFMCMC< TData > Class Template Reference

Likelihood-Free Markov Chain Monte Carlo.

#include <1fmcmc-bones.hpp>

### **Public Member Functions**

- void run (std::vector< epiworld\_double > params\_init\_, size\_t n\_samples\_, epiworld\_double epsilon\_, int seed=-1)
- LFMCMC (const TData & observed data )
- void set\_observed\_data (const TData &observed\_data\_)
- void set\_proposal\_fun (LFMCMCProposalFun < TData > fun)
- void set\_simulation\_fun (LFMCMCSimFun < TData > fun)
- void set\_summary\_fun (LFMCMCSummaryFun< TData > fun)
- void set\_kernel\_fun (LFMCMCKernelFun < TData > fun)
- void set\_params\_names (std::vector< std::string > names)
- void set\_stats\_names (std::vector< std::string > names)
- size\_t get\_n\_samples () const

- size\_t get\_n\_stats () const
- · size\_t get\_n\_params () const
- · epiworld double get\_epsilon () const
- const std::vector< epiworld double > & get\_initial\_params () const
- const std::vector< epiworld\_double > & get\_current\_proposed\_params () const
- const std::vector< epiworld\_double > & get\_current\_accepted\_params () const
- const std::vector< epiworld\_double > & get\_current\_proposed\_stats () const
- const std::vector< epiworld double > & get current\_accepted\_stats () const
- const std::vector< epiworld double > & get observed stats () const
- const std::vector< epiworld double > & get all sample params () const
- const std::vector< epiworld double > & get all sample stats () const
- const std::vector< bool > & get\_all\_sample\_acceptance () const
- const std::vector< epiworld\_double > & get\_all\_sample\_drawn\_prob () const
- const std::vector< epiworld\_double > & get\_all\_sample\_kernel\_scores () const
- const std::vector< epiworld double > & get all\_accepted\_params () const
- const std::vector< epiworld double > & get all accepted stats () const
- const std::vector< epiworld\_double > & get\_all\_accepted\_kernel\_scores () const
- std::vector< TData > \* get simulated data () const
- std::vector< epiworld\_double > get\_mean\_params ()
- std::vector< epiworld double > get mean stats ()
- LFMCMC< TData > & verbose off ()
- LFMCMC< TData > & verbose\_on ()
- · void print (size\_t burnin=0u) const

#### Random number generation

#### **Parameters**

eng

- void set\_rand\_engine (std::shared\_ptr< std::mt19937 > &eng)
- std::shared ptr< std::mt19937 > & get\_rand\_endgine ()
- void seed (epiworld\_fast\_uint s)
- void **set\_rand\_gamma** (epiworld\_double alpha, epiworld\_double beta)
- epiworld double runif ()
- epiworld\_double rnorm ()
- epiworld\_double rgamma ()
- epiworld double runif (epiworld double lb, epiworld double ub)
- epiworld double **rnorm** (epiworld double mean, epiworld double sd)
- epiworld double **rgamma** (epiworld double alpha, epiworld double beta)

### 18.20.1 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 files:

- include/epiworld/math/lfmcmc/lfmcmc-bones.hpp
- include/epiworld/math/lfmcmc/lfmcmc-meat-print.hpp
- include/epiworld/math/lfmcmc/lfmcmc-meat.hpp

## 18.21 epiworld::Model < TSeq > Class Template Reference

Core class of epiworld.

```
#include <epiworld.hpp>
```

Collaboration diagram for epiworld::Model < TSeq >:



### **Public Member Functions**

- DataBase < TSeq > & get\_db ()
- const DataBase< TSeq > & get\_db () const
- epiworld\_double & operator() (std::string pname)
- size\_t size () const
- void load\_agents\_entities\_ties (std::string fn, int skip)

Associate agents-entities from a file.

- void load\_agents\_entities\_ties (const std::vector< int > &agents\_ids, const std::vector< int > &entities\_ids)
   Associate agents-entities from data.
- void load\_agents\_entities\_ties (const int \*agents\_id, const int \*entities\_id, size\_t n)
- size\_t get\_n\_viruses () const

Number of viruses in the model.

• size\_t get\_n\_tools () const

Number of tools in the model.

- epiworld\_fast\_uint get\_ndays () const
- · epiworld\_fast\_uint get\_n\_replicates () const
- size\_t get\_n\_entities () const
- void set\_ndays (epiworld\_fast\_uint ndays)
- bool get\_verbose () const
- Model < TSeq > & verbose\_off ()
- Model < TSeq > & verbose\_on ()

· int today () const

The current time of the model.

void write\_data (std::string fn\_virus\_info, std::string fn\_virus\_hist, std::string fn\_tool\_info, std::string fn\_tool\_hist, std::string fn\_transmission, std::string fn\_transmiss

Wrapper of DataBase::write\_data

- std::map< std::string, epiworld\_double > & params ()
- virtual void reset ()

Reset the model.

- const Model < TSeq > & print (bool lite=false) 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\_globalevent (std::function< void(Model< TSeq > \*)> fun, std::string name="A global action", int date=-99)

Set a global action.

- void add\_globalevent (GlobalEvent < TSeq > action)
- GlobalEvent < TSeq > & get\_globalevent (std::string name)

Retrieve a global action by name.

GlobalEvent < TSeq > & get globalevent (size t i)

Retrieve a global action by index.

void rm\_globalevent (std::string name)

Remove a global action by name.

· void rm globalevent (size ti)

Remove a global action by index.

- void run globalevents ()
- void clear\_state\_set ()
- const std::vector< VirusPtr< TSeq > > & get\_viruses () const
- const std::vector< ToolPtr< TSeq > > & get\_tools () const
- Virus < TSeq > & get\_virus (size\_t id)
- Tool < TSeq > & get\_tool (size\_t id)
- void set\_agents\_data (double \*data\_, size\_t ncols\_)

Set the agents data object.

- double \* get\_agents\_data ()
- size\_t get\_agents\_data\_ncols () const
- void set\_name (std::string name)

Set the name object.

- std::string get\_name () const
- bool operator== (const Model < TSeq > &other) const
- bool operator!= (const Model < TSeq > &other) const
- void events\_run ()

Executes the stored action.

• void <a href="draw">draw</a> (DiagramType diagram\_type=DiagramType::Mermaid, const std::string &fn\_output="", bool self=false)

Draws a mermaid diagram of the model.

### 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 ()

### Random number generation

#### **Parameters**

eng	Random number generator
S	Seed

- void set\_rand\_engine (std::shared\_ptr< std::mt19937 > &eng)
- std::shared\_ptr< std::mt19937 > & get\_rand\_endgine ()
- void seed (size\_t s)
- void set rand norm (epiworld double mean, epiworld double sd)
- void set rand unif (epiworld double a, epiworld double b)
- void set rand exp (epiworld double lambda)
- void set rand gamma (epiworld double alpha, epiworld double beta)
- void set rand lognormal (epiworld double mean, epiworld double shape)
- void set\_rand\_binom (int n, epiworld\_double p)
- void set\_rand\_nbinom (int n, epiworld\_double p)
- void set\_rand\_geom (epiworld\_double p)
- void set rand poiss (epiworld double lambda)
- epiworld\_double runif ()
- epiworld\_double runif (epiworld\_double a, epiworld\_double b)
- epiworld double rnorm ()
- epiworld double **rnorm** (epiworld double mean, epiworld double sd)
- epiworld double rgamma ()
- epiworld double rgamma (epiworld double alpha, epiworld double beta)
- epiworld double rexp ()
- epiworld double rexp (epiworld double lambda)
- epiworld\_double rlognormal ()
- epiworld double **riognormal** (epiworld double mean, epiworld double shape)
- int rbinom ()
- int **rbinom** (int n, epiworld\_double p)
- int rnbinom ()
- int **rnbinom** (int n, epiworld\_double p)
- int rgeom ()
- int **rgeom** (epiworld double p)
- int rpoiss ()
- · int rpoiss (epiworld double lambda)

### Add Virus/Tool to the model

This is done before the model has been initialized.

### Parameters

٧	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)
- void add\_tool (Tool < TSeq > &t)
- void add\_entity (Entity < TSeq > e)
- void rm\_virus (size\_t virus\_pos)
- void rm tool (size t tool pos)
- void rm\_entity (size\_t entity\_id)

#### 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.
size	Size of the network.
al	AdjList to read into the model.

- void agents from adilist (std::string fn, int size, int skip=0, bool directed=false)
- void agents\_from\_edgelist (const std::vector< int > &source, const std::vector< int > &target, int size, bool directed)
- · void agents from adjlist (AdjList al)
- · bool is directed () const
- std::vector< Agent< TSeq > > & get\_agents ()

Returns a reference to the vector of agents.

- Agent < TSeq > & get\_agent (size\_t i)
- std::vector< epiworld\_fast\_uint > get\_agents\_states () const

Returns a vector with the states of the agents.

std::vector< Viruses\_const< TSeq > > get\_agents\_viruses () const

Returns a const vector with the viruses of the agents.

std::vector< Viruses< TSeq > > get agents viruses ()

Returns a vector with the viruses of the agents.

- std::vector< Entity< TSeq > > & get entities ()
- Entity < TSeg > & get entity (size t entity id, int \*entity pos=nullptr)
- Model < TSeq > & agents\_smallworld (epiworld\_fast\_uint n=1000, epiworld\_fast\_uint k=5, bool d=false, epiworld\_double p=.01)
- void agents\_empty\_graph (epiworld\_fast\_uint n=1000)

#### 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 update\_state ()
- void mutate\_virus ()
- void next ()
- virtual Model < TSeq > & run (epiworld\_fast\_uint ndays, int seed=-1)

Runs the simulation (after initialization)

void run\_multiple (epiworld\_fast\_uint ndays, epiworld\_fast\_uint nexperiments, int seed\_=-1, std::function 
 void(size\_t, Model < TSeq > \*) > fun=make\_save\_run < TSeq >(), bool reset=true, bool verbose=true, int nthreads=1)

### 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< Agent< 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< int > &source, std::vector< int > &target) const

### Manage state (states) in the model

The functions get\_state return the current values for the states included in the model.

#### **Parameters**

lab std::s	string <b>Name of the state</b> .
------------	-----------------------------------

### Returns

add\_state\* returns nothing.
get\_state\_\* returns a vector of pairs with the states and their labels.

- void add\_state (std::string lab, UpdateFun< TSeq > fun=nullptr)
- const std::vector< std::string > & get\_states () const
- size\_t get\_n\_states () const
- const std::vector< UpdateFun< TSeq > > & get\_state\_fun () const
- void print\_state\_codes () const

### Initial states

These functions are called before the simulation starts.

### **Parameters**

proportions↔	Vector of proportions for each state.
queue_	Vector of queue for each state.

virtual Model < TSeq > & initial\_states (std::vector < double >, std::vector < int >)

### 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 epiworld\_fast\_uint 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 par() function members are aliases for get\_param().

In the case of the function read\_params, users can pass a file listing parameters to be included in the model. Each line in the file should have the following structure:

```
[name of parameter 1]: [value in double]
[name of parameter 2]: [value in double]
```

The only condition for parameter names is that these do not include a colon.

#### **Parameters**

initial_val	
pname	Name of the parameter to add or to fetch
fn	Path to the file containing parameters

#### Returns

The current value of the parameter in the model.

- epiworld double add param (epiworld double initial val, std::string pname, bool overwrite=false)
- Model < TSeq > & read\_params (std::string fn, bool overwrite=false)
- epiworld double **get param** (epiworld fast uint k)
- epiworld double **get param** (std::string pname)
- void **set\_param** (std::string pname, epiworld\_double val)
- epiworld\_double par (std::string pname) const

### Set the user data object

### **Parameters**

names	string vector with the names of the variables.
-------	------------------------------------------------

- void set\_user\_data (std::vector< std::string > names)
- void add user data (epiworld fast uint 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.)

Model < TSeq > & 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**



#### Returns

epiworld double

- 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)

### **Protected Member Functions**

- · void dist\_tools ()
- void dist\_virus ()
- void dist\_entities ()
- · void chrono\_start ()
- void chrono\_end ()

Construct a new Event object.

### **Protected Attributes**

```
std::string name = ""
```

Name of the model.

- DataBase< TSeq > db = DataBase<TSeq>(\*this)
- std::vector< Agent< TSeq >> population = {}
- bool using\_backup = true
- std::vector< Agent< TSeq > > population\_backup = {}
- bool directed = false
- std::vector< VirusPtr< TSeq >> viruses = {}
- $std::vector < ToolPtr < TSeq > > tools = {}$
- std::vector< Entity< TSeq >> entities = {}
- std::vector< Entity< TSeq > >  $entities_backup$  = {}
- $std::shared\_ptr < std::mt19937 > engine = std::make\_shared < std::mt19937 > ()$
- std::uniform\_real\_distribution runifd
- · std::normal distribution rnormd
- std::gamma\_distribution rgammad
- · std::lognormal\_distribution rlognormald
- · std::exponential\_distribution rexpd
- · std::binomial distribution rbinomd
- std::negative\_binomial\_distribution rnbinomd
- std::geometric\_distribution rgeomd
- std::poisson\_distribution rpoissd
- std::function< void(std::vector< Agent< TSeq >> \*, Model< TSeq > \*, epiworld\_double)> rewire\_fun
- epiworld\_double rewire\_prop = 0.0
- std::map< std::string, epiworld\_double > parameters
- epiworld\_fast\_uint **ndays** = 0
- · Progress pb

```
    std::vector< UpdateFun< TSeq > > state_fun = {}

     Functions to update states.
std::vector< std::string > states labels = {}
     Labels of the states.

    std::function< void(Model< TSeq > *)> initial_states_fun

• epiworld fast uint nstates = 0u
• bool verbose = true
• int current date = 0

    std::chrono::time_point< std::chrono::steady_clock > time_start

    std::chrono::time_point< std::chrono::steady_clock > time_end

    std::chrono::duration< epiworld_double, std::micro > time_elapsed

• epiworld_fast_uint n_replicates = 0u

    std::vector< GlobalEvent< TSeq >> globalevents

    Queue < TSeq > queue

• bool use_queuing = true
std::vector< Event< TSeq >> events = {}
```

Variables used to keep track of the events to be made regarding viruses.

# Auxiliary variables for AgentsSample<TSeq> iterators

These variables+objects are used by the AgentsSample<TSeq> class for building efficient iterators over agents. The idea is to reduce the memory allocation, so only during the first call of AgentsSample<TSeq>::Agents←Sample(Model<TSeq>) these vectors are allocated.

- std::vector < Agent < TSeq > \* > sampled\_population
- size\_t sampled\_population\_n = 0u
- std::vector< size\_t > population\_left
- size\_t population\_left\_n = 0u

• epiworld fast uint nactions = 0u

### Agents features

Optionally, a model can include an external data source pointing to agents information. The data can then be access through the Agent::operator() method.

- double \* agents\_data = nullptr
- size\_t agents\_data\_ncols = 0u

### **Friends**

- class Agent < TSeq >
- class AgentsSample< TSeq >
- class DataBase< TSeq >
- class Queue < TSeq >

### **Tool Mixers**

These functions combine the effects tools have to deliver a single effect. For example, wearing a mask, been vaccinated, and the immune system combine together to jointly reduce the susceptibility for a given virus.

- MixerFun< TSeq > susceptibility\_reduction\_mixer = susceptibility\_reduction\_mixer\_default<TSeq>
- MixerFun< TSeq > transmission reduction mixer = transmission reduction mixer default<TSeq>
- MixerFun< TSeq > recovery\_enhancer\_mixer = recovery\_enhancer\_mixer\_default<TSeq>

- MixerFun< TSeq > death\_reduction\_mixer = death\_reduction\_mixer\_default<TSeq>
- std::vector< epiworld\_double > array\_double\_tmp
- std::vector< Virus< TSeq > \* > array\_virus\_tmp
- virtual Model < TSeq > \* clone ptr ()

Advanced usage: Makes a copy of data and returns it as undeleted pointer.

- · Model ()
- Model (const Model < TSeq > &m)
- Model (Model < TSeq > &m)
- Model (Model < TSeq > &&m)
- Model < TSeq > & operator= (const Model < TSeq > &m)
- virtual ∼Model ()

### 18.21.1 Detailed Description

```
template<typename TSeq> class epiworld::Model< TSeq >
```

Core class of epiworld.

The model class provides the wrapper that puts together Agent, 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.)
```

#### 18.21.2 Member Function Documentation

#### 18.21.2.1 add globalevent()

Set a global action.

#### **Parameters**

fun	A function to be called on the prescribed date
name	Name of the action.
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.

### 18.21.2.2 clone\_ptr()

```
template<typename TSeq >
Model< TSeq > * Model< TSeq >::clone_ptr [inline], [protected], [virtual]
```

Advanced usage: Makes a copy of data and returns it as undeleted pointer.

#### **Parameters**

сору

Reimplemented in ModelSIRMixing < TSeq >, ModelSIRLogit < TSeq >, ModelSIRDCONN < TSeq >, ModelSIRCONN < TSeq >, ModelSIRDCONN < TSeq >, ModelMeaslesMixingRiskQuarantine < TSeq >, ModelMeaslesMixing < TSeq >, epiworld::epimodels::ModelMeaslesMixing < TSeq >, epiworld::epimodels::ModelSIRMixingQuarantine < TSeq >, epiworld::epimodels::ModelSIRMixing < TSeq >, epiworld::epimodels::ModelSIRMixing < TSeq >, epiworld::epimodels::ModelSIRLogit < TSeq >, epiworld::epimodels::ModelSIRDCONN < TSeq >, epiworld::epimodels::ModelSIRCONN < TSeq >.

#### 18.21.2.3 draw()

Draws a mermaid diagram of the model.

### **Parameters**

model	The model to draw.
fn_output	The name of the file to write the diagram. If empty, the diagram will be printed to the standard output.
self	Whether to allow self-transitions.

### 18.21.2.4 events\_add()

```
int idx_agent_,
int idx_object_ ) [inline], [protected]
```

Construct a new Event object.

#### **Parameters**

agent_	Agent over which the action will be called
virus_	Virus pointer included in the action
tool_	Tool pointer included in the action
entity_	Entity pointer included in the action
new_←	New state of the agent
state_	
call_	Function the action will call
queue_	Change in the queue
idx_⊷	Location of agent in object.
agent_	
idx_⊷	Location of object in agent.
object_	

### 18.21.2.5 events\_run()

```
template<typename TSeq >
void Model< TSeq >::events_run [inline]
```

Executes the stored action.

### **Parameters**

model⊷	Model over which it will be executed.
_	

### 18.21.2.6 load\_agents\_entities\_ties()

Associate agents-entities from a file.

The structure of the file should be two columns separated by space. The first column indexing between 0 and nagents-1, and the second column between 0 and nentities - 1.

### **Parameters**

fn	Path to the file.
skip	How many rows to skip.

Generated by Doxygen

#### 18.21.2.7 reset()

```
template<typename TSeq >
void Model< TSeq >::reset [inline], [virtual]
```

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

Reimplemented in ModelSURV < TSeq >, ModelSIRMixing < TSeq >, ModelSIRLogit < TSeq >, ModelSIRDCONN < TSeq >, ModelSIRDCONN < TSeq >, ModelSIRDCONN < TSeq >, ModelSEIRMixingQuarantine < TSeq >, ModelSEIRMixing < TSeq >, ModelSEIRDCONN < TSeq >, ModelSEIRDCONN < TSeq >, ModelSEIRCONN < TSeq >, ModelMeaslesMixingRiskQuarantine < TSeq >, ModelMeaslesMixing < TSeq >, epiworld::epimodels::ModelMeaslesMixingRiskQuarantine < TSeq >, epiworld::epimodels::ModelMeaslesMixing < TSeq >, epiworld::epimodels::ModelMeaslesSchool < TSeq >, epiworld::epimodels::ModelSIRMixing < TSeq >, epiworld::epimodels::ModelSIRMixing < TSeq >, epiworld::epimodels::ModelSIRMixing < TSeq >, epiworld::epimodels::ModelSIRDCONN < TSeq >, epiworld::epimodels::ModelSIRDCONN < TSeq >, epiworld::epimodels::ModelSIRDCONN < TSeq >, epiworld::epimodels::ModelSIRDCONN < TSeq >, and epiworld::epimodels::ModelSURV < TSeq >.

### 18.21.2.8 run\_multiple()

#### **Parameters**

ndays | Multiple runs of the simulation

#### 18.21.2.9 set\_agents\_data()

template<typename TSeq >

Set the agents data object.

The data should be an array with the data stored in a column major order, i.e., by column.

#### Parameters 4 8 1

data⊷	Pointer to the first element of an array of size size() *
_	ncols
ncols⇔	Number of features included in the data.
_	

### 18.21.2.10 set\_name()

Set the name object.

### **Parameters**

name

### 18.21.2.11 write\_data()

Wrapper of DataBase::write\_data

#### **Parameters**

fn_virus_info	Filename. Information about the virus.
fn_virus_hist	Filename. History of the virus.
fn_tool_info	Filename. Information about the tool.

#### **Parameters**

fn_tool_hist	Filename. History of the tool.
fn_total_hist	Filename. Aggregated history (state)
fn_transmission	Filename. Transmission history.
fn_transition	Filename. Markov transition history.
fn_reproductive_number	Filename. Case by case reproductive number

### 18.21.3 Member Data Documentation

### 18.21.3.1 initial\_states\_fun

```
= [] (Model<TSeq> * )
     -> void {}
```

Function to distribute states. Goes along with the function

#### 18.21.3.2 rbinomd

```
template<typename TSeq >
std::binomial_distribution epiworld::Model< TSeq >::rbinomd [protected]
```

### Initial value:

```
std::binomial_distribution<>()
```

### 18.21.3.3 rexpd

```
template<typename TSeq >
std::exponential_distribution epiworld::Model< TSeq >::rexpd [protected]
```

### Initial value:

```
std::exponential_distribution<>()
```

#### 18.21.3.4 rgammad

```
template<typename TSeq >
std::gamma_distribution epiworld::Model< TSeq >::rgammad [protected]
Initial value:
       std::gamma_distribution<>()
18.21.3.5 rgeomd
template<typename TSeq >
std::geometric_distribution epiworld::Model< TSeq >::rgeomd [protected]
Initial value:
       std::geometric_distribution<>()
18.21.3.6 rlognormald
template<typename TSeq >
std::lognormal_distribution epiworld::Model < TSeq >::rlognormald [protected]
Initial value:
       std::lognormal_distribution<>()
18.21.3.7 rnbinomd
template<typename TSeq >
std::negative_binomial_distribution epiworld::Model< TSeq >::rnbinomd [protected]
Initial value:
       std::negative_binomial_distribution<>()
18.21.3.8 rnormd
template<typename TSeq >
std::normal_distribution epiworld::Model< TSeq >::rnormd [protected]
```

std::normal\_distribution<>(0.0)

Initial value:

#### 18.21.3.9 rpoissd

#### 18.21.3.10 runifd

### 18.21.3.11 time\_elapsed

```
template<typename TSeq >
std::chrono::duration<epiworld_double,std::micro> epiworld::Model< TSeq >::time_elapsed [protected]
```

#### Initial value:

std::chrono::duration<epiworld\_double,std::micro>::zero()

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

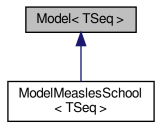
· epiworld.hpp

# **18.22** Model < TSeq > Class Template Reference

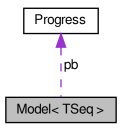
Core class of epiworld.

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

Inheritance diagram for Model < TSeq >:



Collaboration diagram for Model < TSeq >:



### **Public Member Functions**

- DataBase < TSeq > & get\_db ()
- const DataBase< TSeq > & get\_db () const
- epiworld double & operator() (std::string pname)
- · size t size () const
- void load\_agents\_entities\_ties (std::string fn, int skip)

Associate agents-entities from a file.

- void load\_agents\_entities\_ties (const std::vector< int > &agents\_ids, const std::vector< int > &entities\_ids)

  Associate agents-entities from data.
- void load\_agents\_entities\_ties (const int \*agents\_id, const int \*entities\_id, size\_t n)
- size\_t get\_n\_viruses () const

Number of viruses in the model.

• size\_t get\_n\_tools () const

Number of tools in the model.

- epiworld\_fast\_uint get\_ndays () const
- epiworld\_fast\_uint get\_n\_replicates () const
- size\_t get\_n\_entities () const
- void set\_ndays (epiworld\_fast\_uint ndays)
- bool get\_verbose () const
- Model < TSeq > & verbose\_off ()
- Model < TSeq > & verbose\_on ()
- · int today () const

The current time of the model.

void write\_data (std::string fn\_virus\_info, std::string fn\_virus\_hist, std::string fn\_tool\_info, std::string fn—tool\_hist, std::string fn\_total\_hist, std::string fn\_transmission, std::string fn\_transition, std::string fn\_erroductive\_number, std::string fn\_generation\_time) const

Wrapper of DataBase::write\_data

- std::map< std::string, epiworld\_double > & params ()
- virtual void reset ()

Reset the model.

- const  ${\sf Model}{<}$  TSeq > &  ${\sf print}$  (bool lite=false) 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\_globalevent (std::function< void(Model< TSeq > \*)> fun, std::string name="A global action", int date=-99)

Set a global action.

- void add\_globalevent (GlobalEvent < TSeq > action)
- GlobalEvent < TSeq > & get\_globalevent (std::string name)

Retrieve a global action by name.

GlobalEvent < TSeq > & get\_globalevent (size\_t i)

Retrieve a global action by index.

void rm globalevent (std::string name)

Remove a global action by name.

· void rm\_globalevent (size\_t i)

Remove a global action by index.

- void run\_globalevents ()
- · void clear state set ()
- const std::vector< VirusPtr< TSeq > > & get\_viruses () const
- const std::vector< ToolPtr< TSeq > > & get\_tools () const
- Virus< TSeq > & get\_virus (size\_t id)
- Tool < TSeq > & get\_tool (size t id)
- void set\_agents\_data (double \*data\_, size\_t ncols\_)

Set the agents data object.

- double \* get\_agents\_data ()
- · size t get agents data ncols () const
- void set name (std::string name)

Set the name object.

- std::string get\_name () const
- bool operator== (const Model < TSeq > &other) const
- bool operator!= (const Model < TSeq > &other) const
- void events run ()

Executes the stored action.

 void draw (DiagramType diagram\_type=DiagramType::Mermaid, const std::string &fn\_output="", bool self=false)

Draws a mermaid diagram of the model.

### 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 ()

#### Random number generation

#### **Parameters**

eng	Random number generator
S	Seed

- void set\_rand\_engine (std::shared\_ptr< std::mt19937 > &eng)
- std::shared\_ptr< std::mt19937 > & get\_rand\_endgine ()
- void seed (size ts)
- void set\_rand\_norm (epiworld\_double mean, epiworld\_double sd)
- void **set\_rand\_unif** (epiworld\_double a, epiworld\_double b)
- void set\_rand\_exp (epiworld\_double lambda)
- void **set\_rand\_gamma** (epiworld\_double alpha, epiworld\_double beta)
- · void set rand lognormal (epiworld double mean, epiworld double shape)
- void **set\_rand\_binom** (int n, epiworld\_double p)

- void set\_rand\_nbinom (int n, epiworld\_double p)
- void **set\_rand\_geom** (epiworld\_double p)
- void set\_rand\_poiss (epiworld\_double lambda)
- epiworld double runif ()
- epiworld\_double runif (epiworld\_double a, epiworld\_double b)
- epiworld\_double rnorm ()
- epiworld double **rnorm** (epiworld double mean, epiworld double sd)
- epiworld double rgamma ()
- epiworld\_double rgamma (epiworld\_double alpha, epiworld\_double beta)
- epiworld double rexp ()
- epiworld\_double rexp (epiworld\_double lambda)
- epiworld\_double rlognormal ()
- epiworld\_double **rlognormal** (epiworld\_double mean, epiworld\_double shape)
- int rbinom ()
- int **rbinom** (int n, epiworld\_double p)
- int rnbinom ()
- int **rnbinom** (int n, epiworld double p)
- int rgeom ()
- int **rgeom** (epiworld\_double p)
- int rpoiss ()
- int rpoiss (epiworld\_double lambda)

#### 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)
- void add\_tool (Tool < TSeq > &t)
- void add\_entity (Entity < TSeq > e)
- void rm\_virus (size\_t virus\_pos)
- void rm\_tool (size\_t tool\_pos)
- void rm\_entity (size\_t entity\_id)

#### 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.
size	Size of the network.
al	AdjList to read into the model.

- void agents\_from\_adjlist (std::string fn, int size, int skip=0, bool directed=false)
- void agents\_from\_edgelist (const std::vector< int > &source, const std::vector< int > &target, int size, bool directed)
- void agents\_from\_adjlist (AdjList al)
- bool is\_directed () const
- std::vector< Agent< TSeq > > & get\_agents ()

Returns a reference to the vector of agents.

- Agent < TSeq > & get\_agent (size\_t i)
- std::vector< epiworld\_fast\_uint > get\_agents\_states () const

Returns a vector with the states of the agents.

-  $std::vector < Viruses\_const < TSeq > > get\_agents\_viruses$  () const

Returns a const vector with the viruses of the agents.

std::vector< Viruses< TSeq >> get\_agents\_viruses ()

Returns a vector with the viruses of the agents.

- std::vector< Entity< TSeq > > & get entities ()
- Entity < TSeq > & get\_entity (size t entity id, int \*entity pos=nullptr)
- Model < TSeq > & agents\_smallworld (epiworld\_fast\_uint n=1000, epiworld\_fast\_uint k=5, bool d=false, epiworld\_double p=.01)
- void agents empty graph (epiworld fast uint n=1000)

#### 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 update\_state ()
- void mutate\_virus ()
- void next ()
- virtual Model < TSeq > & run (epiworld\_fast\_uint ndays, int seed=-1)
   Runs the simulation (after initialization)
- void run\_multiple (epiworld\_fast\_uint ndays, epiworld\_fast\_uint nexperiments, int seed\_=-1, std::function 
   void(size\_t, Model < TSeq > \*) > fun=make\_save\_run < TSeq >(), bool reset=true, bool verbose=true, int nthreads=1)

#### 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< Agent< 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< int > &source, std::vector< int > &target) const

#### Manage state (states) in the model

The functions get\_state return the current values for the states included in the model.

#### **Parameters**

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

#### Returns

add\_state\* returns nothing.

get state \* returns a vector of pairs with the states and their labels.

- void add\_state (std::string lab, UpdateFun< TSeq > fun=nullptr)
- const std::vector< std::string > & get\_states () const
- size t get n states () const
- const std::vector< UpdateFun< TSeq > > & get\_state\_fun () const
- · void print\_state\_codes () const

#### **Initial states**

These functions are called before the simulation starts.

#### **Parameters**

proportions ←	Vector of proportions for each state.
queue_	Vector of queue for each state.

virtual Model < TSeq > & initial\_states (std::vector < double >, std::vector < int >)

#### 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 epiworld\_fast\_uint 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 par() function members are aliases for get\_param().

In the case of the function read\_params, users can pass a file listing parameters to be included in the model. Each line in the file should have the following structure:

```
[name of parameter 1]: [value in double]
[name of parameter 2]: [value in double]
```

The only condition for parameter names is that these do not include a colon.

#### **Parameters**

initial_val	
pname	Name of the parameter to add or to fetch
fn	Path to the file containing parameters

#### Returns

The current value of the parameter in the model.

- epiworld\_double add\_param (epiworld\_double initial\_val, std::string pname, bool overwrite=false)
- Model < TSeq > & read\_params (std::string fn, bool overwrite=false)
- epiworld\_double **get\_param** (epiworld\_fast\_uint k)
- epiworld\_double **get\_param** (std::string pname)
- void **set\_param** (std::string pname, epiworld\_double val)
- epiworld double par (std::string pname) const

#### Set the user data object

#### **Parameters**

names

string vector with the names of the variables.

- void set\_user\_data (std::vector< std::string > names)
- void add user data (epiworld fast uint 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.)

Model < TSeq > & 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**



### Returns

epiworld\_double

- 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)

### **Protected Member Functions**

- void dist\_tools()
- · void dist\_virus ()

- void dist\_entities ()
- void chrono\_start ()
- void chrono\_end ()
- void events add (Agent < TSeq > \*agent , VirusPtr < TSeq > virus , ToolPtr < TSeq > tool , Entity < TSeq</li> > \*entity\_, epiworld\_fast\_int new\_state\_, epiworld\_fast\_int queue\_, EventFun< TSeq > call\_, int idx\_~ agent\_, int idx\_object\_)

Construct a new Event object.

```
Protected Attributes
    std::string name = ""
         Name of the model.

    DataBase< TSeq > db = DataBase<TSeq>(*this)

    std::vector< Agent< TSeq >> population = {}
    • bool using backup = true

    std::vector< Agent< TSeq > > population_backup = {}

    bool directed = false

    std::vector< VirusPtr< TSeq >> viruses = {}
    std::vector< ToolPtr< TSeq > > tools = {}
    std::vector< Entity< TSeq >> entities = {}

    std::vector< Entity< TSeq > > entities_backup = {}

    std::shared_ptr< std::mt19937 > engine = std::make_shared< std::mt19937 >()
    · std::uniform_real_distribution runifd
    • std::normal_distribution rnormd
    · std::gamma distribution rgammad
    · std::lognormal_distribution rlognormald

    std::exponential distribution rexpd

    · std::binomial distribution rbinomd
    • std::negative_binomial_distribution rnbinomd

    std::geometric_distribution rgeomd

    · std::poisson distribution rpoissd

    std::function< void(std::vector< Agent< TSeq >> *, Model< TSeq > *, epiworld_double)> rewire_fun

    • epiworld_double rewire_prop = 0.0

    std::map< std::string, epiworld_double > parameters

    • epiworld_fast_uint ndays = 0
    · Progress pb

    std::vector< UpdateFun< TSeq > > state fun = {}

         Functions to update states.
    std::vector< std::string > states_labels = {}
         Labels of the states.

    std::function< void(Model< TSeq > *)> initial_states_fun

    • epiworld fast uint nstates = 0u
    • bool verbose = true
    • int current date = 0

    std::chrono::time_point< std::chrono::steady_clock > time_start

    - std::chrono::time\_point < std::chrono::steady\_clock > time\_end

    std::chrono::duration< epiworld_double, std::micro > time_elapsed

    • epiworld_fast_uint n_replicates = 0u
```

 Queue < TSeq > queue • bool use\_queuing = true

std::vector < GlobalEvent < TSeq > > globalevents

Variables used to keep track of the events to be made regarding viruses.

std::vector< Event< TSeq > > events = {}

• epiworld\_fast\_uint nactions = 0u

#### Auxiliary variables for AgentsSample<TSeq> iterators

These variables+objects are used by the AgentsSample<TSeq> class for building efficient iterators over agents. The idea is to reduce the memory allocation, so only during the first call of AgentsSample<TSeq>::Agents←Sample(Model<TSeq>) these vectors are allocated.

```
    std::vector< Agent< TSeq > * > sampled_population
```

- size t sampled population n = 0u
- std::vector< size\_t > population\_left
- size t population left n = 0u

### Agents features

Optionally, a model can include an external data source pointing to agents information. The data can then be access through the Agent::operator() method.

```
double * agents_data = nullptr
```

• size\_t agents\_data\_ncols = 0u

#### **Friends**

```
class Agent < TSeq >
```

- class AgentsSample < TSeq >
- class DataBase < TSeq >
- class Queue < TSeq >

#### **Tool Mixers**

These functions combine the effects tools have to deliver a single effect. For example, wearing a mask, been vaccinated, and the immune system combine together to jointly reduce the susceptibility for a given virus.

```
    MixerFun< TSeq > susceptibility reduction mixer = susceptibility reduction mixer default<TSeq>
```

- MixerFun< TSeq > transmission\_reduction\_mixer = transmission\_reduction\_mixer\_default<TSeq>
- MixerFun< TSeq > recovery\_enhancer\_mixer = recovery\_enhancer\_mixer\_default<TSeq>
- MixerFun< TSeq > death\_reduction\_mixer = death\_reduction\_mixer\_default<TSeq>
- std::vector< epiworld double > array double tmp
- std::vector< Virus< TSeq > \* > array virus tmp
- virtual Model < TSeq > \* clone\_ptr ()

Advanced usage: Makes a copy of data and returns it as undeleted pointer.

- · Model ()
- Model (const Model < TSeq > &m)
- Model (Model < TSeq > &m)
- Model (Model < TSeq > &&m)
- Model < TSeq > & operator= (const Model < TSeq > &m)
- virtual ∼Model ()

### 18.22.1 Detailed Description

```
template<typename TSeq> class Model< TSeq >
```

Core class of epiworld.

The model class provides the wrapper that puts together Agent, 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.)	

### 18.22.2 Member Function Documentation

### 18.22.2.1 add\_globalevent()

Set a global action.

#### **Parameters**

fun	A function to be called on the prescribed date	
name	e Name of the action.	
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.

### 18.22.2.2 clone\_ptr()

```
template<typename TSeq >
Model< TSeq > * Model< TSeq >::clone_ptr [inline], [protected], [virtual]
```

Advanced usage: Makes a copy of data and returns it as undeleted pointer.

### **Parameters**

сору

Reimplemented in ModelMeaslesSchool < TSeq >.

### 18.22.2.3 draw()

```
template<typename TSeq >
void Model< TSeq >::draw (
```

```
DiagramType diagram_type = DiagramType::Mermaid,
const std::string & fn_output = "",
bool self = false ) [inline]
```

Draws a mermaid diagram of the model.

#### **Parameters**

model	The model to draw.
fn_output The name of the file to write the diagram. If empty, the diagram will be printed to the stand	
self	Whether to allow self-transitions.

### 18.22.2.4 events\_add()

Construct a new Event object.

### **Parameters**

Agent over which the action will be called
Virus pointer included in the action
Tool pointer included in the action
Entity pointer included in the action
New state of the agent
Function the action will call
Change in the queue
Location of agent in object.
Location of object in agent.

### 18.22.2.5 events\_run()

```
template<typename TSeq >
void Model< TSeq >::events_run [inline]
```

Executes the stored action.

#### **Parameters**

model⊷	Model over which it will be executed.	1

### 18.22.2.6 load\_agents\_entities\_ties()

Associate agents-entities from a file.

The structure of the file should be two columns separated by space. The first column indexing between 0 and nagents-1, and the second column between 0 and nentities - 1.

#### **Parameters**

fn	Path to the file.
skip	How many rows to skip.

### 18.22.2.7 reset()

```
template<typename TSeq >
void Model< TSeq >::reset [inline], [virtual]
```

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

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#### 18.22.2.8 run\_multiple()

#### **Parameters**

```
ndays | Multiple runs of the simulation
```

### 18.22.2.9 set\_agents\_data()

Set the agents data object.

The data should be an array with the data stored in a column major order, i.e., by column.

#### **Parameters**

data⊷	Pointer to the first element of an array of size size () *	
_	ncols	
ncols⇔	cols  Our Number of features included in the data.	
_		

### 18.22.2.10 set\_name()

```
template<typename TSeq >
void Model< TSeq >::set_name (
          std::string name) [inline]
```

Set the name object.

#### **Parameters**

name

#### 18.22.2.11 write data()

Wrapper of DataBase::write\_data

#### **Parameters**

fn_virus_info	Filename. Information about the virus.
fn_virus_hist	Filename. History of the virus.
fn_tool_info	Filename. Information about the tool.
fn_tool_hist	Filename. History of the tool.
fn_total_hist	Filename. Aggregated history (state)
fn_transmission	Filename. Transmission history.
fn_transition	Filename. Markov transition history.
fn_reproductive_number	Filename. Case by case reproductive number

### 18.22.3 Member Data Documentation

### 18.22.3.1 initial\_states\_fun

Function to distribute states. Goes along with the function

### 18.22.3.2 rbinomd

Generated by Doxygen

#### 18.22.3.3 rexpd

```
template<typename TSeq >
std::exponential_distribution Model< TSeq >::rexpd [protected]
Initial value:
       std::exponential_distribution<>()
18.22.3.4 rgammad
template<typename TSeq >
std::gamma_distribution Model< TSeq >::rgammad [protected]
Initial value:
       std::gamma_distribution<>()
18.22.3.5 rgeomd
template<typename TSeq >
std::geometric_distribution Model< TSeq >::rgeomd [protected]
Initial value:
       std::geometric_distribution<>()
18.22.3.6 rlognormald
template<typename TSeq >
std::lognormal_distribution Model< TSeq >::rlognormald [protected]
Initial value:
       std::lognormal_distribution<>()
18.22.3.7 rnbinomd
template<typename TSeq >
```

std::negative\_binomial\_distribution Model< TSeq >::rnbinomd [protected]

std::negative\_binomial\_distribution<>()

# Generated by Doxygen

Initial value:

#### 18.22.3.8 rnormd

#### 18.22.3.9 rpoissd

std::poisson\_distribution<>()

### 18.22.3.10 runifd

```
template<typename TSeq >
std::uniform_real_distribution Model< TSeq >::runifd [protected]
Initial value:
```

std::uniform\_real\_distribution<> (0.0, 1.0)

### 18.22.3.11 time\_elapsed

```
template<typename TSeq >
std::chrono::duration<epiworld_double,std::micro> Model< TSeq >::time_elapsed [protected]
```

### Initial value:

```
std::chrono::duration<epiworld_double,std::micro>::zero()
```

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

- include/epiworld/agent-bones.hpp
- include/epiworld/model-bones.hpp
- include/epiworld/model-meat-print.hpp
- include/epiworld/model-meat.hpp

## 18.23 epiworld::ModelDiagram Class Reference

## **Public Member Functions**

- void draw\_from\_data (DiagramType diagram\_type, const std::vector< std::string > &states, const std
   ::vector< epiworld\_double > &tprob, const std::string &fn\_output="", bool self=false)
- void draw\_from\_file (DiagramType diagram\_type, const std::string &fn\_transition, const std::string &fn\_← output="", bool self=false)
- void **draw\_from\_files** (DiagramType diagram\_type, const std::vector< std::string > &fns\_transition, const std::string &fn\_output="", bool self=false)

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

· epiworld.hpp

## 18.24 ModelDiagram Class Reference

## **Public Member Functions**

- void draw\_from\_data (DiagramType diagram\_type, const std::vector< std::string > &states, const std
   ::vector< epiworld\_double > &tprob, const std::string &fn\_output="", bool self=false)
- void draw\_from\_file (DiagramType diagram\_type, const std::string &fn\_transition, const std::string &fn\_← output="", bool self=false)
- void **draw\_from\_files** (DiagramType diagram\_type, const std::vector< std::string > &fns\_transition, const std::string &fn\_output="", bool self=false)

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

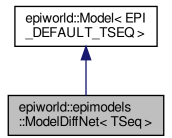
- include/epiworld/modeldiagram-bones.hpp
- include/epiworld/modeldiagram-meat.hpp

## 18.25 epiworld::epimodels::ModelDiffNet< TSeq > Class Template Reference

Template for a Network Diffusion Model.

#include <epiworld.hpp>

 $Inheritance\ diagram\ for\ epiworld:: epimodels:: Model DiffNet < TSeq >:$ 



Collaboration diagram for epiworld::epimodels::ModelDiffNet< TSeq >:



## **Public Member Functions**

- **ModelDiffNet** (ModelDiffNet< TSeq > &model, const std::string &innovation\_name, epiworld\_double prevalence, epiworld\_double prob\_adopt, bool normalize\_exposure=true, double \*agents\_data=nullptr, size\_← t data\_ncols=0u, std::vector< size\_t > data\_cols={}, std::vector< double > params={})
- ModelDiffNet (const std::string &innovation\_name, epiworld\_double prevalence, epiworld\_double prob
   \_adopt, bool normalize\_exposure=true, double \*agents\_data=nullptr, size\_t data\_ncols=0u, std::vector <
   size\_t > data\_cols={}, std::vector < double > params={})

## **Public Attributes**

- bool normalize\_exposure = true
- std::vector< size t > data\_cols
- std::vector< double > params

## **Static Public Attributes**

- static const int NONADOPTER = 0
- static const int ADOPTER = 1

## **Additional Inherited Members**

## 18.25.1 Detailed Description

template<typename TSeq = EPI\_DEFAULT\_TSEQ> class epiworld::epimodels::ModelDiffNet< TSeq >

Template for a Network Diffusion Model.

model	A Model <tseq> object where to set up the SIR.</tseq>	
vname	std::string Name of the virus	Generated by Doxygen
initial_prevalence	epiworld_double Initial prevalence	deficiated by Boxygen
initial_efficacy	epiworld_double Initial susceptibility_reduction of the immune system	
initial_recovery	epiworld_double Initial recovery rate of the immune system	

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

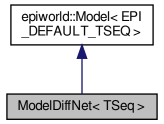
· epiworld.hpp

## 18.26 ModelDiffNet< TSeq > Class Template Reference

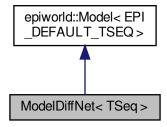
Template for a Network Diffusion Model.

```
#include <diffnet.hpp>
```

Inheritance diagram for ModelDiffNet< TSeq >:



Collaboration diagram for ModelDiffNet< TSeq >:



## **Public Member Functions**

- **ModelDiffNet** (ModelDiffNet< TSeq > &model, const std::string &innovation\_name, epiworld\_double prevalence, epiworld\_double prob\_adopt, bool normalize\_exposure=true, double \*agents\_data=nullptr, size\_
  t data\_ncols=0u, std::vector< size\_t > data\_cols={}, std::vector< double > params={})
- ModelDiffNet (const std::string &innovation\_name, epiworld\_double prevalence, epiworld\_double prob
   \_adopt, bool normalize\_exposure=true, double \*agents\_data=nullptr, size\_t data\_ncols=0u, std::vector <
   size\_t > data\_cols={}, std::vector < double > params={})

## **Public Attributes**

- bool normalize\_exposure = true
- $std::vector < size_t > data_cols$
- std::vector< double > params

## **Static Public Attributes**

- static const int **NONADOPTER** = 0
- static const int ADOPTER = 1

## **Additional Inherited Members**

## 18.26.1 Detailed Description

```
template<typename TSeq = EPI_DEFAULT_TSEQ> class ModelDiffNet< TSeq >
```

Template for a Network Diffusion Model.

#### **Parameters**

model	A Model <tseq> object where to set up the SIR.</tseq>
vname	std::string Name of the virus
initial_prevalence	epiworld_double Initial prevalence
initial_efficacy	epiworld_double Initial susceptibility_reduction of the immune system
initial_recovery	epiworld_double Initial recovery rate of the immune system

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

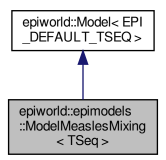
• include/epiworld/models/diffnet.hpp

# 18.27 epiworld::epimodels::ModelMeaslesMixing< TSeq > Class Template Reference

Measles model with population mixing, quarantine, and contact tracing.

#include <epiworld.hpp>

Inheritance diagram for epiworld::epimodels::ModelMeaslesMixing < TSeq >:



Collaboration diagram for epiworld::epimodels::ModelMeaslesMixing < TSeq >:



## **Public Member Functions**

• ModelMeaslesMixing (ModelMeaslesMixing< TSeq > &model, epiworld\_fast\_uint n, epiworld\_double prevalence, epiworld\_double contact\_rate, epiworld\_double transmission\_rate, epiworld\_double vax\_\top efficacy, epiworld\_double vax\_reduction\_recovery\_rate, epiworld\_double incubation\_period, epiworld\_double prodromal\_period, epiworld\_double rash\_period, std::vector< double > contact\_matrix, epiworld\_double hospitalization\_rate, epiworld\_double hospitalization\_period, epiworld\_double days\_undetected, epiworld\top fast\_int quarantine\_period, epiworld\_double quarantine\_willingness, epiworld\_double isolation\_willingness, epiworld\_fast\_int isolation\_period, epiworld\_double prop\_vaccinated, epiworld\_double contact\_tracing\_\top success\_rate=1.0, epiworld\_fast\_uint contact\_tracing\_days\_prior=4u)

Constructs a ModelMeaslesMixing object.

ModelMeaslesMixing (epiworld\_fast\_uint n, epiworld\_double prevalence, epiworld\_double contact\_rate, epiworld\_double transmission\_rate, epiworld\_double vax\_efficacy, epiworld\_double vax\_reduction\_
 recovery\_rate, epiworld\_double incubation\_period, epiworld\_double prodromal\_period, epiworld\_double rash\_period, std::vector< double > contact\_matrix, epiworld\_double hospitalization\_rate, epiworld\_double hospitalization\_period, epiworld\_double days\_undetected, epiworld\_fast\_int quarantine\_period, epiworld
 \_double quarantine\_willingness, epiworld\_double isolation\_willingness, epiworld\_fast\_int isolation\_period,

epiworld\_double prop\_vaccinated, epiworld\_double contact\_tracing\_success\_rate=1.0, epiworld\_fast\_uint contact\_tracing\_days\_prior=4u)

Constructs a ModelMeaslesMixing object.

ModelMeaslesMixing < TSeq > & run (epiworld fast uint ndays, int seed=-1)

Run the model simulation.

· void reset ()

Reset the model to initial state.

Model < TSeq > \* clone ptr ()

Create a clone of this model.

ModelMeaslesMixing< TSeq > & initial\_states (std::vector< double > proportions\_, std::vector< int > queue\_={})

Set the initial states of the model.

void set contact matrix (std::vector< double > cmat)

Set the contact matrix for population mixing.

std::vector< double > get\_contact\_matrix () const

Get the current contact matrix.

• std::vector< size\_t > get\_agent\_quarantine\_triggered () const

Get the quarantine trigger status for all agents.

std::vector< bool > get quarantine willingness () const

Get the quarantine willingness for all agents.

std::vector< bool > get\_isolation\_willingness () const

Get the isolation willingness for all agents.

## **Static Public Attributes**

- static const int SUSCEPTIBLE = 0
- static const int EXPOSED = 1
- static const int PRODROMAL = 2
- static const int RASH = 3
- static const int ISOLATED = 4
- static const int ISOLATED RECOVERED = 5
- static const int **DETECTED\_HOSPITALIZED** = 6
- static const int **QUARANTINED\_EXPOSED** = 7
- static const int QUARANTINED\_SUSCEPTIBLE = 8
- static const int **QUARANTINED\_PRODROMAL** = 9
- static const int **QUARANTINED\_RECOVERED** = 10
- static const int HOSPITALIZED = 11
- static const int **RECOVERED** = 12
- static const size t QUARANTINE PROCESS INACTIVE = 0u
- static const size t QUARANTINE PROCESS ACTIVE = 1u
- static const size\_t QUARANTINE\_PROCESS\_DONE = 2u

## **Additional Inherited Members**

## 18.27.1 Detailed Description

template<typename TSeq = EPI\_DEFAULT\_TSEQ>
class epiworld::epimodels::ModelMeaslesMixing< TSeq >

Measles model with population mixing, quarantine, and contact tracing.

This class implements a Measles epidemiological model based on the SEIR framework with additional features including:

- · Population mixing based on contact matrices
- Measles-specific disease progression: Susceptible o Exposed o Prodromal o Rash
- Prodromal individuals are infectious (replace the "Infected" state in SEIR)
- · Rash individuals are no longer infectious but can be detected for isolation
- · Quarantine measures for exposed contacts during contact tracing
- · Isolation policies for detected individuals during the rash state
- · Contact tracing with configurable success rates
- · Hospitalization of severe cases
- · Individual willingness to comply with public health measures

The model supports 13 distinct states:

- · Susceptible: Individuals who can become infected
- Exposed: Infected but not yet infectious (incubation period)
- Prodromal: Infectious individuals in the community (replaces "Infected" in SEIR)
- Rash: Non-infectious individuals with visible symptoms (detection occurs here)
- · Isolated: Detected individuals in self-isolation
- · Isolated Recovered: Recovered individuals still in isolation
- · Detected Hospitalized: Hospitalized individuals who were contact-traced
- · Quarantined Exposed: Exposed individuals in quarantine due to contact tracing
- · Quarantined Susceptible: Susceptible individuals in guarantine due to contact tracing
- · Quarantined Prodromal: Prodromal individuals in quarantine due to contact tracing
- · Quarantined Recovered: Recovered individuals in quarantine due to contact tracing
- · Hospitalized: Individuals requiring hospital care
- · Recovered: Individuals who have recovered and gained immunity

## **Template Parameters**

```
TSeq Type for genetic sequences (default: EPI_DEFAULT_TSEQ)
```

## 18.27.2 Constructor & Destructor Documentation

## 18.27.2.1 ModelMeaslesMixing() [1/2]

```
template<typename TSeq >
ModelMeaslesMixing< TSeq >::ModelMeaslesMixing (
```

```
ModelMeaslesMixing< TSeq > & model,
epiworld_fast_uint n,
epiworld_double prevalence,
epiworld_double contact_rate,
epiworld_double transmission_rate,
epiworld_double vax_efficacy,
epiworld_double vax_reduction_recovery_rate,
epiworld_double incubation_period,
epiworld_double prodromal_period,
epiworld_double rash_period,
std::vector< double > contact_matrix,
epiworld_double hospitalization_rate,
epiworld_double hospitalization_period,
epiworld_double days_undetected,
epiworld_fast_int quarantine_period,
epiworld_double quarantine_willingness,
epiworld_double isolation_willingness,
epiworld_fast_int isolation_period,
epiworld_double prop_vaccinated,
epiworld_double contact_tracing_success_rate = 1.0,
epiworld_fast_uint contact_tracing_days_prior = 4u ) [inline]
```

## Constructs a ModelMeaslesMixing object.

Template for a Measles model with population mixing, quarantine, and contact tracing.

A reference to an existing ModelMeaslesMixing object.
The number of entities in the model.
The initial prevalence of the disease in the model.
The contact rate between entities in the model.
The transmission rate of the disease in the model.
The efficacy of the vaccine.
The reduction in recovery rate due to the vaccine.
The incubation period of the disease in the model.
The prodromal period of the disease in the model.
The rash period of the disease in the model.
The contact matrix between entities in the model. Specified in column-major
order.
The rate at which infected individuals are hospitalized.
The average duration of hospitalization in days.
The average number of days an infected individual remains undetected.
The duration of quarantine in days for exposed contacts.
The proportion of individuals willing to comply with quarantine measures.
The proportion of individuals willing to self-isolate when detected.
The duration of isolation in days for detected infected individuals.
The proportion of vaccinated agents.
The probability of successfully identifying and tracing contacts (default: 1.0).
The number of days prior to detection for which contacts are traced (default:
4).
A ModelMeaslesMixing <tseq> object where to set up the model.</tseq>
Number of agents in the population
Initial prevalence (proportion of infected individuals)

#### **Parameters**

contact_rate	Average number of contacts (interactions) per step
transmission_rate	Probability of transmission per contact
vax_efficacy	The efficacy of the vaccine
vax_reduction_recovery_rate	The reduction in recovery rate due to the vaccine
incubation_period	Average incubation period in days
prodromal_period	Average prodromal period in days
rash_period	Average rash period in days
contact_matrix	Contact matrix specifying mixing patterns between population groups
hospitalization_rate	Rate at which infected individuals are hospitalized
hospitalization_period	Average duration of hospitalization in days
days_undetected	Average number of days an infected individual remains undetected
quarantine_period	Duration of quarantine in days for exposed contacts
quarantine_willingness	Proportion of individuals willing to comply with quarantine
isolation_willingness	Proportion of individuals willing to self-isolate when detected
isolation_period	Duration of isolation in days for detected infected individuals
prop_vaccinated	Proportion of vaccinated agents
contact_tracing_success_rate	Probability of successfully identifying contacts during tracing
contact_tracing_days_prior	Number of days prior to detection for contact tracing

## 18.27.2.2 ModelMeaslesMixing() [2/2]

```
template<typename TSeq >
ModelMeaslesMixing< TSeq >::ModelMeaslesMixing (
             epiworld_fast_uint n,
             epiworld_double prevalence,
             epiworld_double contact_rate,
             epiworld_double transmission_rate,
             epiworld_double vax_efficacy,
             epiworld_double vax_reduction_recovery_rate,
             epiworld_double incubation_period,
             epiworld_double prodromal_period,
             epiworld_double rash_period,
             std::vector< double > contact_matrix,
             epiworld_double hospitalization_rate,
             epiworld_double hospitalization_period,
             epiworld_double days_undetected,
             epiworld_fast_int quarantine_period,
             epiworld_double quarantine_willingness,
             epiworld_double isolation_willingness,
             epiworld_fast_int isolation_period,
             epiworld_double prop_vaccinated,
             epiworld_double contact_tracing_success_rate = 1.0,
             {\tt epiworld\_fast\_uint} \ \ contact\_tracing\_days\_prior = 4u \ ) \quad [{\tt inline}]
```

Constructs a ModelMeaslesMixing object.

#### **Parameters**

n	The number of entities in the model.
prevalence	The initial prevalence of the disease in the model.
contact_rate	The contact rate between entities in the model.
transmission_rate	The transmission rate of the disease in the model.
vax_efficacy	The efficacy of the vaccine.
vax_reduction_recovery_rate	The reduction in recovery rate due to the vaccine.
incubation_period	The incubation period of the disease in the model.
prodromal_period	The prodromal period of the disease in the model.
rash_period	The rash period of the disease in the model.
contact_matrix	The contact matrix between entities in the model.
hospitalization_rate	The rate at which infected individuals are hospitalized.
hospitalization_period	The average duration of hospitalization in days.
days_undetected	The average number of days an infected individual remains undetected.
quarantine_period	The duration of quarantine in days for exposed contacts.
quarantine_willingness	The proportion of individuals willing to comply with quarantine measures.
isolation_willingness	The proportion of individuals willing to self-isolate when detected.
isolation_period	The duration of isolation in days for detected infected individuals.
prop_vaccinated	The proportion of vaccinated agents.
contact_tracing_success_rate	The probability of successfully identifying and tracing contacts (default: 1.0).
contact_tracing_days_prior	The number of days prior to detection for which contacts are traced (default: 4).

## 18.27.3 Member Function Documentation

## 18.27.3.1 clone\_ptr()

```
template<typename TSeq >
Model< TSeq > * ModelMeaslesMixing< TSeq >::clone_ptr [inline], [virtual]
```

Create a clone of this model.

#### Returns

Pointer to a new model instance with the same configuration

Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

## 18.27.3.2 get\_agent\_quarantine\_triggered()

```
template<typename TSeq = EPI_DEFAULT_TSEQ>
std::vector< size_t > epiworld::epimodels::ModelMeaslesMixing< TSeq >::get_agent_quarantine
_triggered ( ) const [inline]
```

Get the quarantine trigger status for all agents.

## Returns

Vector indicating quarantine process status for each agent

#### 18.27.3.3 get\_contact\_matrix()

```
template<typename TSeq = EPI_DEFAULT_TSEQ>
std::vector< double > epiworld::epimodels::ModelMeaslesMixing< TSeq >::get_contact_matrix ( )
const [inline]
```

Get the current contact matrix.

#### Returns

Vector representing the contact matrix

## 18.27.3.4 get\_isolation\_willingness()

```
template<typename TSeq = EPI_DEFAULT_TSEQ>
std::vector< bool > epiworld::epimodels::ModelMeaslesMixing< TSeq >::get_isolation_willingness
( ) const [inline]
```

Get the isolation willingness for all agents.

#### Returns

Vector of boolean values indicating each agent's willingness to self-isolate

## 18.27.3.5 get\_quarantine\_willingness()

```
template<typename TSeq = EPI_DEFAULT_TSEQ>
std::vector< bool > epiworld::epimodels::ModelMeaslesMixing< TSeq >::get_quarantine_willingness
( ) const [inline]
```

Get the quarantine willingness for all agents.

## Returns

Vector of boolean values indicating each agent's willingness to quarantine

## 18.27.3.6 initial\_states()

```
template<typename TSeq >
ModelMeaslesMixing< TSeq > & ModelMeaslesMixing< TSeq >::initial_states (
    std::vector< double > proportions_,
    std::vector< int > queue_ = {} ) [inline], [virtual]
```

Set the initial states of the model.

#### **Parameters**

proportions↔	Double vector with two elements:
_	• [0]: The proportion of initially infected individuals who start in the exposed state.
	• [1]: The proportion of initially non-infected individuals who have recovered (immune).
queue_	Optional vector for queuing specifications (default: empty).

Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

## 18.27.3.7 run()

Run the model simulation.

#### **Parameters**

ndays	Number of days to simulate
seed	Random seed for reproducibility (default: -1 for random seed)

## Returns

Reference to this model instance

 $\label{eq:local_problem} \mbox{Reimplemented from epiworld::} \mbox{Model} < \mbox{EPI\_DEFAULT\_TSEQ} >.$ 

## 18.27.3.8 set\_contact\_matrix()

Set the contact matrix for population mixing.

## **Parameters**

cmat   Contact matrix specifying interaction rates between groups
-------------------------------------------------------------------

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

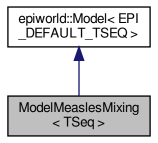
epiworld.hpp

## 18.28 ModelMeaslesMixing < TSeq > Class Template Reference

Measles model with population mixing, quarantine, and contact tracing.

#include <measlesmixing.hpp>

Inheritance diagram for ModelMeaslesMixing < TSeq >:



Collaboration diagram for ModelMeaslesMixing < TSeq >:



## **Public Member Functions**

Constructs a ModelMeaslesMixing object.

ModelMeaslesMixing (epiworld\_fast\_uint n, epiworld\_double prevalence, epiworld\_double contact\_rate, epiworld\_double transmission\_rate, epiworld\_double vax\_efficacy, epiworld\_double vax\_reduction\_← recovery\_rate, epiworld\_double incubation\_period, epiworld\_double prodromal\_period, epiworld\_double rash\_period, std::vector< double > contact\_matrix, epiworld\_double hospitalization\_rate, epiworld\_double hospitalization\_period, epiworld\_double days\_undetected, epiworld\_fast\_int quarantine\_period, epiworld← \_double quarantine\_willingness, epiworld\_double isolation\_willingness, epiworld\_fast\_int isolation\_period, epiworld\_double prop\_vaccinated, epiworld\_double contact\_tracing\_success\_rate=1.0, epiworld\_fast\_uint contact\_tracing\_days\_prior=4u)

Constructs a ModelMeaslesMixing object.

ModelMeaslesMixing < TSeq > & run (epiworld\_fast\_uint ndays, int seed=-1)

Run the model simulation.

• void reset ()

Reset the model to initial state.

Model < TSeq > \* clone\_ptr ()

Create a clone of this model.

ModelMeaslesMixing< TSeq > & initial\_states (std::vector< double > proportions\_, std::vector< int > queue\_={})

Set the initial states of the model.

void set contact matrix (std::vector< double > cmat)

Set the contact matrix for population mixing.

std::vector< double > get\_contact\_matrix () const

Get the current contact matrix.

std::vector< size\_t > get\_agent\_quarantine\_triggered () const

Get the quarantine trigger status for all agents.

std::vector< bool > get\_quarantine\_willingness () const

Get the quarantine willingness for all agents.

std::vector< bool > get isolation willingness () const

Get the isolation willingness for all agents.

#### Static Public Attributes

- static const int SUSCEPTIBLE = 0
- static const int **EXPOSED** = 1
- static const int **PRODROMAL** = 2
- static const int RASH = 3
- static const int ISOLATED = 4
- static const int ISOLATED\_RECOVERED = 5
- static const int **DETECTED\_HOSPITALIZED** = 6
- static const int **QUARANTINED\_EXPOSED** = 7
- static const int QUARANTINED\_SUSCEPTIBLE = 8
- static const int QUARANTINED PRODROMAL = 9
- static const int QUARANTINED\_RECOVERED = 10
- static const int HOSPITALIZED = 11
- static const int **RECOVERED** = 12
- static const size t QUARANTINE PROCESS INACTIVE = 0u
- static const size t QUARANTINE PROCESS ACTIVE = 1u
- static const size t QUARANTINE PROCESS DONE = 2u

## **Additional Inherited Members**

## 18.28.1 Detailed Description

template<typename TSeq = EPI\_DEFAULT\_TSEQ> class ModelMeaslesMixing< TSeq >

Measles model with population mixing, quarantine, and contact tracing.

This class implements a Measles epidemiological model based on the SEIR framework with additional features including:

- · Population mixing based on contact matrices
- $\bullet \ \ \mathsf{Measles\text{-}specific \ disease \ progression:} \ \mathsf{Susceptible} \to \mathsf{Exposed} \to \mathsf{Prodromal} \to \mathsf{Rash}$
- · Prodromal individuals are infectious (replace the "Infected" state in SEIR)
- · Rash individuals are no longer infectious but can be detected for isolation
- · Quarantine measures for exposed contacts during contact tracing
- · Isolation policies for detected individuals during the rash state
- · Contact tracing with configurable success rates
- · Hospitalization of severe cases
- · Individual willingness to comply with public health measures

The model supports 13 distinct states:

- Susceptible: Individuals who can become infected
- Exposed: Infected but not yet infectious (incubation period)
- Prodromal: Infectious individuals in the community (replaces "Infected" in SEIR)
- · Rash: Non-infectious individuals with visible symptoms (detection occurs here)
- · Isolated: Detected individuals in self-isolation
- · Isolated Recovered: Recovered individuals still in isolation
- · Detected Hospitalized: Hospitalized individuals who were contact-traced
- · Quarantined Exposed: Exposed individuals in quarantine due to contact tracing
- · Quarantined Susceptible: Susceptible individuals in quarantine due to contact tracing
- · Quarantined Prodromal: Prodromal individuals in quarantine due to contact tracing
- · Quarantined Recovered: Recovered individuals in quarantine due to contact tracing
- · Hospitalized: Individuals requiring hospital care
- · Recovered: Individuals who have recovered and gained immunity

## **Template Parameters**

TSeq Type for genetic sequences (default: EPI\_DEFAULT\_TSEQ)

## 18.28.2 Constructor & Destructor Documentation

## 18.28.2.1 ModelMeaslesMixing() [1/2]

```
{\tt template}{<}{\tt typename}~{\tt TSeq}~{>}
ModelMeaslesMixing< TSeq >::ModelMeaslesMixing (
             ModelMeaslesMixing< TSeq > & model,
              epiworld_fast_uint n,
              epiworld_double prevalence,
              epiworld_double contact_rate,
              epiworld_double transmission_rate,
              epiworld_double vax_efficacy,
              epiworld_double vax_reduction_recovery_rate,
              epiworld_double incubation_period,
              epiworld_double prodromal_period,
              epiworld_double rash_period,
              std::vector< double > contact_matrix,
              epiworld_double hospitalization_rate,
              epiworld_double hospitalization_period,
              epiworld_double days_undetected,
              epiworld_fast_int quarantine_period,
              epiworld_double quarantine_willingness,
              epiworld_double isolation_willingness,
              epiworld_fast_int isolation_period,
              epiworld_double prop_vaccinated,
              epiworld_double contact_tracing_success_rate = 1.0,
              {\tt epiworld\_fast\_uint} \ \ contact\_tracing\_days\_prior = 4u \ ) \quad [{\tt inline}]
```

## Constructs a ModelMeaslesMixing object.

Template for a Measles model with population mixing, quarantine, and contact tracing.

model	A reference to an existing ModelMeaslesMixing object.
п	The number of entities in the model.
prevalence	The initial prevalence of the disease in the model.
contact_rate	The contact rate between entities in the model.
transmission_rate	The transmission rate of the disease in the model.
vax_efficacy	The efficacy of the vaccine.
vax_reduction_recovery_rate	The reduction in recovery rate due to the vaccine.
incubation_period	The incubation period of the disease in the model.
prodromal_period	The prodromal period of the disease in the model.
rash_period	The rash period of the disease in the model.
contact_matrix	The contact matrix between entities in the model. Specified in column-major order.

## **Parameters**

hospitalization_rate	The rate at which infected individuals are hospitalized.
hospitalization_period	The average duration of hospitalization in days.
days_undetected	The average number of days an infected individual remains undetected.
quarantine_period	The duration of quarantine in days for exposed contacts.
quarantine_willingness	The proportion of individuals willing to comply with quarantine measures.
isolation_willingness	The proportion of individuals willing to self-isolate when detected.
isolation_period	The duration of isolation in days for detected infected individuals.
prop_vaccinated	The proportion of vaccinated agents.
contact_tracing_success_rate	The probability of successfully identifying and tracing contacts (default: 1.0).
contact_tracing_days_prior	The number of days prior to detection for which contacts are traced (default: 4).
model	A ModelMeaslesMixing <tseq> object where to set up the model.</tseq>
n	Number of agents in the population
prevalence	Initial prevalence (proportion of infected individuals)
contact_rate	Average number of contacts (interactions) per step
transmission_rate	Probability of transmission per contact
vax_efficacy	The efficacy of the vaccine
vax_reduction_recovery_rate	The reduction in recovery rate due to the vaccine
incubation_period	Average incubation period in days
prodromal_period	Average prodromal period in days
rash_period	Average rash period in days
contact_matrix	Contact matrix specifying mixing patterns between population groups
hospitalization_rate	Rate at which infected individuals are hospitalized
hospitalization_period	Average duration of hospitalization in days
days_undetected	Average number of days an infected individual remains undetected
quarantine_period	Duration of quarantine in days for exposed contacts
quarantine_willingness	Proportion of individuals willing to comply with quarantine
isolation_willingness	Proportion of individuals willing to self-isolate when detected
isolation_period	Duration of isolation in days for detected infected individuals
prop_vaccinated	Proportion of vaccinated agents
contact_tracing_success_rate	Probability of successfully identifying contacts during tracing
contact_tracing_days_prior	Number of days prior to detection for contact tracing
	1

## 18.28.2.2 ModelMeaslesMixing() [2/2]

```
epiworld_double rash_period,
std::vector< double > contact_matrix,
epiworld_double hospitalization_rate,
epiworld_double hospitalization_period,
epiworld_double days_undetected,
epiworld_fast_int quarantine_period,
epiworld_double quarantine_willingness,
epiworld_double isolation_willingness,
epiworld_fast_int isolation_period,
epiworld_double prop_vaccinated,
epiworld_double contact_tracing_success_rate = 1.0,
epiworld_fast_uint contact_tracing_days_prior = 4u ) [inline]
```

## Constructs a ModelMeaslesMixing object.

## **Parameters**

п	The number of entities in the model.
prevalence	The initial prevalence of the disease in the model.
contact_rate	The contact rate between entities in the model.
transmission_rate	The transmission rate of the disease in the model.
vax_efficacy	The efficacy of the vaccine.
vax_reduction_recovery_rate	The reduction in recovery rate due to the vaccine.
incubation_period	The incubation period of the disease in the model.
prodromal_period	The prodromal period of the disease in the model.
rash_period	The rash period of the disease in the model.
contact_matrix	The contact matrix between entities in the model.
hospitalization_rate	The rate at which infected individuals are hospitalized.
hospitalization_period	The average duration of hospitalization in days.
days_undetected	The average number of days an infected individual remains undetected.
quarantine_period	The duration of quarantine in days for exposed contacts.
quarantine_willingness	The proportion of individuals willing to comply with quarantine measures.
isolation_willingness	The proportion of individuals willing to self-isolate when detected.
isolation_period	The duration of isolation in days for detected infected individuals.
prop_vaccinated	The proportion of vaccinated agents.
contact_tracing_success_rate	The probability of successfully identifying and tracing contacts (default: 1.0).
contact_tracing_days_prior	The number of days prior to detection for which contacts are traced (default: 4).

## 18.28.3 Member Function Documentation

## 18.28.3.1 clone\_ptr()

```
template<typename TSeq >
Model< TSeq > * ModelMeaslesMixing< TSeq >::clone_ptr [inline], [virtual]
```

Create a clone of this model.

#### Returns

Pointer to a new model instance with the same configuration

Reimplemented from epiworld::Model < EPI DEFAULT TSEQ >.

## 18.28.3.2 get\_agent\_quarantine\_triggered()

```
template<typename TSeq = EPI_DEFAULT_TSEQ>
std::vector< size_t > ModelMeaslesMixing< TSeq >::get_agent_quarantine_triggered ( ) const
[inline]
```

Get the quarantine trigger status for all agents.

## Returns

Vector indicating quarantine process status for each agent

## 18.28.3.3 get\_contact\_matrix()

```
template<typename TSeq = EPI_DEFAULT_TSEQ>
std::vector< double > ModelMeaslesMixing< TSeq >::get_contact_matrix ( ) const [inline]
```

Get the current contact matrix.

## Returns

Vector representing the contact matrix

## 18.28.3.4 get\_isolation\_willingness()

```
template<typename TSeq = EPI_DEFAULT_TSEQ>
std::vector< bool > ModelMeaslesMixing< TSeq >::get_isolation_willingness ( ) const [inline]
```

Get the isolation willingness for all agents.

## Returns

Vector of boolean values indicating each agent's willingness to self-isolate

## 18.28.3.5 get\_quarantine\_willingness()

```
template<typename TSeq = EPI_DEFAULT_TSEQ>
std::vector< bool > ModelMeaslesMixing< TSeq >::get_quarantine_willingness ( ) const [inline]
```

Get the quarantine willingness for all agents.

#### Returns

Vector of boolean values indicating each agent's willingness to quarantine

## 18.28.3.6 initial\_states()

Set the initial states of the model.

#### **Parameters**

proportions⊷	Double vector with two elements:	]
_	• [0]: The proportion of initially infected individuals who start in the exposed state.	
	• [1]: The proportion of initially non-infected individuals who have recovered (immune).	
queue_	Optional vector for queuing specifications (default: empty).	1

Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

## 18.28.3.7 run()

Run the model simulation.

ndays	Number of days to simulate
seed	Random seed for reproducibility (default: -1 for random seed)

#### Returns

Reference to this model instance

Reimplemented from epiworld::Model < EPI DEFAULT TSEQ >.

## 18.28.3.8 set\_contact\_matrix()

Set the contact matrix for population mixing.

#### **Parameters**

cmat Contact matrix specifying interaction rates between groups

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

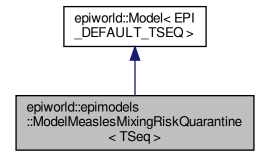
• include/epiworld/models/measlesmixing.hpp

# 18.29 epiworld::epimodels::ModelMeaslesMixingRiskQuarantine< TSeq > Class Template Reference

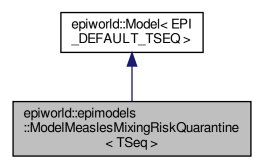
Measles model with population mixing and risk-based quarantine strategies.

```
#include <epiworld.hpp>
```

 $Inheritance\ diagram\ for\ epiworld:: epimodels:: Model Measles Mixing Risk Quarantine < TSeq>:$ 



Collaboration diagram for epiworld::epimodels::ModelMeaslesMixingRiskQuarantine < TSeq >:



#### **Public Member Functions**

ModelMeaslesMixingRiskQuarantine (ModelMeaslesMixingRiskQuarantine < TSeq > &model, epiworld\_ ⇔ fast\_uint n, epiworld\_double prevalence, epiworld\_double contact\_rate, epiworld\_double transmission ← \_ rate, epiworld\_double vax\_efficacy, epiworld\_double vax\_reduction\_recovery\_rate, epiworld\_double incubation\_period, epiworld\_double prodromal\_period, epiworld\_double rash\_period, std::vector < double > contact\_matrix, epiworld\_double hospitalization\_rate, epiworld\_double hospitalization\_period, epiworld\_ ⇔ double days\_undetected, epiworld\_fast\_int quarantine\_period\_high, epiworld\_fast\_int quarantine\_period\_ ⇔ medium, epiworld\_fast\_int quarantine\_period\_low, epiworld\_double quarantine\_willingness, epiworld\_double isolation\_willingness, epiworld\_fast\_int isolation\_period, epiworld\_double prop\_vaccinated, epiworld\_ ⇔ double detection\_rate\_quarantine, epiworld\_double contact\_tracing\_success\_rate=1.0, epiworld\_fast\_uint contact\_tracing\_days\_prior=4u)

Constructs a ModelMeaslesMixingRiskQuarantine object.

ModelMeaslesMixingRiskQuarantine (epiworld\_fast\_uint n, epiworld\_double prevalence, epiworld\_double contact\_rate, epiworld\_double transmission\_rate, epiworld\_double vax\_efficacy, epiworld\_double vax\_ereduction\_recovery\_rate, epiworld\_double incubation\_period, epiworld\_double prodromal\_period, epiworld← \_\_double rash\_period, std::vector< double > contact\_matrix, epiworld\_double hospitalization\_rate, epiworld← \_\_double hospitalization\_period, epiworld\_double days\_undetected, epiworld\_fast\_int quarantine\_period← \_\_high, epiworld\_fast\_int quarantine\_period\_medium, epiworld\_fast\_int quarantine\_period\_low, epiworld← \_\_double quarantine\_willingness, epiworld\_double isolation\_willingness, epiworld\_fast\_int isolation\_period, epiworld\_double prop\_vaccinated, epiworld\_double detection\_rate\_quarantine, epiworld\_double contact\_← tracing success rate=1.0, epiworld fast uint contact tracing days prior=4u)

Constructs a ModelMeaslesMixingRiskQuarantine object.

ModelMeaslesMixingRiskQuarantine< TSeq > & run (epiworld\_fast\_uint ndays, int seed=-1)

Run the model simulation.

· void reset ()

Reset the model to initial state.

Model < TSeq > \* clone\_ptr ()

Create a clone of this model.

ModelMeaslesMixingRiskQuarantine < TSeq > & initial\_states (std::vector < double > proportions\_, std ← ::vector < int > queue\_={})

Set the initial states of the model.

void set contact matrix (std::vector< double > cmat)

Set the contact matrix for population mixing.

• std::vector< double > get\_contact\_matrix () const

Get the current contact matrix.

• std::vector< size\_t > get\_agent\_quarantine\_triggered () const

Get the quarantine trigger status for all agents.

- bool get\_system\_quarantine\_triggered () const
- std::vector< bool > get\_quarantine\_willingness () const

Get the quarantine willingness for all agents.

std::vector< bool > get isolation willingness () const

Get the isolation willingness for all agents.

std::vector< int > get quarantine risk levels () const

Get the risk levels assigned to all agents.

## **Static Public Attributes**

- static constexpr int SUSCEPTIBLE = 0
- static constexpr int **EXPOSED** = 1
- static constexpr int **PRODROMAL** = 2
- static constexpr int **RASH** = 3
- static constexpr int ISOLATED = 4
- static constexpr int ISOLATED\_RECOVERED = 5
- static constexpr int **DETECTED\_HOSPITALIZED** = 6
- static constexpr int QUARANTINED\_EXPOSED = 7
- static constexpr int QUARANTINED SUSCEPTIBLE = 8
- static constexpr int QUARANTINED PRODROMAL = 9
- static constexpr int QUARANTINED RECOVERED = 10
- static constexpr int HOSPITALIZED = 11
- static constexpr int **RECOVERED** = 12
- static constexpr size t QUARANTINE PROCESS INACTIVE = 0u
- static constexpr size t QUARANTINE PROCESS ACTIVE = 1u
- static constexpr size\_t QUARANTINE\_PROCESS\_DONE = 2u
- static constexpr int RISK\_LOW = 0
- static constexpr int RISK MEDIUM = 1
- static constexpr int RISK\_HIGH = 2

## **Additional Inherited Members**

## 18.29.1 Detailed Description

```
template<typename TSeq = EPI_DEFAULT_TSEQ>
class epiworld::epimodels::ModelMeaslesMixingRiskQuarantine< TSeq >
```

Measles model with population mixing and risk-based quarantine strategies.

This class extends the Measles epidemiological model to support different quarantine strategies based on exposure risk levels:

- · High Risk: Unvaccinated agents who share entity membership with the case
- · Medium Risk: Unvaccinated agents who contacted an infected individual but don't share entity membership
- · Low Risk: Other unvaccinated agents

Each risk level can have different quarantine durations, allowing for targeted public health interventions. The model also includes enhanced detection during active quarantine periods.

Disease progression follows the same states as ModelMeaslesMixing: Susceptible  $\to$  Exposed  $\to$  Prodromal  $\to$  Rash  $\to$  Recovered

## **Template Parameters**

TSeq Type for genetic sequences (default: EPI\_DEFAULT\_TSEQ)

## 18.29.2 Constructor & Destructor Documentation

#### 18.29.2.1 ModelMeaslesMixingRiskQuarantine() [1/2]

```
{\tt template}{<}{\tt typename}~{\tt TSeq}~{>}
{\tt ModelMeaslesMixingRiskQuarantine} < {\tt TSeq} >: : {\tt ModelMeaslesMixingRiskQuarantine} \quad (
             ModelMeaslesMixingRiskQuarantine< TSeq > & model,
             epiworld_fast_uint n,
             epiworld_double prevalence,
             epiworld_double contact_rate,
             epiworld_double transmission_rate,
             epiworld_double vax_efficacy,
             epiworld_double vax_reduction_recovery_rate,
             epiworld_double incubation_period,
             epiworld_double prodromal_period,
             epiworld_double rash_period,
              std::vector< double > contact_matrix,
             epiworld_double hospitalization_rate,
             epiworld_double hospitalization_period,
             epiworld_double days_undetected,
             epiworld_fast_int quarantine_period_high,
             epiworld_fast_int quarantine_period_medium,
             epiworld_fast_int quarantine_period_low,
             epiworld_double quarantine_willingness,
              epiworld_double isolation_willingness,
             epiworld_fast_int isolation_period,
             epiworld_double prop_vaccinated,
             epiworld_double detection_rate_quarantine,
             epiworld_double contact_tracing_success_rate = 1.0,
             epiworld_fast_uint contact_tracing_days_prior = 4u ) [inline]
```

Constructs a ModelMeaslesMixingRiskQuarantine object.

Template for a Measles model with population mixing and risk-based quarantine.

model	A reference to an existing ModelMeaslesMixingRiskQuarantine object.
n	The number of entities in the model.
prevalence	The initial prevalence of the disease in the model.
contact_rate	The contact rate between entities in the model.
transmission_rate	The transmission rate of the disease in the model.
vax_efficacy	The efficacy of the vaccine.
vax_reduction_recovery_rate	The reduction in recovery rate due to the vaccine.
incubation_period	The incubation period of the disease in the model.
prodromal_period	The prodromal period of the disease in the model.

rash_period	The rash period of the disease in the model.
contact_matrix	The contact matrix between entities in the model. Specified in column-major order.
hospitalization_rate	The rate at which infected individuals are hospitalized.
hospitalization_period	The average duration of hospitalization in days.
days_undetected	The average number of days an infected individual remains undetected.
quarantine_period_high	The duration of quarantine in days for high-risk contacts.
quarantine_period_medium	The duration of quarantine in days for medium-risk contacts.
quarantine_period_low	The duration of quarantine in days for low-risk contacts.
quarantine_willingness	The proportion of individuals willing to comply with quarantine measures.
isolation_willingness	The proportion of individuals willing to self-isolate when detected.
isolation_period	The duration of isolation in days for detected infected individuals.
prop_vaccinated	The proportion of vaccinated agents.
detection_rate_quarantine	The detection rate during active quarantine periods.
contact_tracing_success_rate	The probability of successfully identifying and tracing contacts (default: 1.0).
contact_tracing_days_prior	The number of days prior to detection for which contacts are traced (default: 4).
model	A ModelMeaslesMixingRiskQuarantine <tseq> object where to set up the model.</tseq>
n	Number of agents in the population
prevalence	Initial prevalence (proportion of infected individuals)
contact_rate	Average number of contacts (interactions) per step
transmission_rate	Probability of transmission per contact
vax_efficacy	The efficacy of the vaccine
vax_reduction_recovery_rate	The reduction in recovery rate due to the vaccine
incubation_period	Average incubation period in days
prodromal_period	Average prodromal period in days
rash_period	Average rash period in days
contact_matrix	Contact matrix specifying mixing patterns between population groups
hospitalization_rate	Rate at which infected individuals are hospitalized
hospitalization_period	Average duration of hospitalization in days
days_undetected	Average number of days an infected individual remains undetected
quarantine_period_high	Duration of quarantine in days for high-risk contacts
quarantine_period_medium	Duration of quarantine in days for medium-risk contacts
quarantine_period_low	Duration of quarantine in days for low-risk contacts
quarantine_willingness	Proportion of individuals willing to comply with quarantine
isolation_willingness	Proportion of individuals willing to self-isolate when detected
isolation_period	Duration of isolation in days for detected infected individuals
prop_vaccinated	Proportion of vaccinated agents
detection_rate_quarantine	Detection rate during active quarantine periods
contact_tracing_success_rate	Probability of successfully identifying contacts during tracing
contact_tracing_days_prior	Number of days prior to detection for contact tracing

## 18.29.2.2 ModelMeaslesMixingRiskQuarantine() [2/2]

```
template<typename TSeq >
{\tt Model Measles Mixing Risk Quarantine < TSeq >:: \tt Model Measles Mixing Risk Quarantine (IN the Second Foundation of the Model Measles Mixing Risk Quarantine (IN the Model Measles Mixing Risk Quara
  epiworld_fast_uint n,
  epiworld_double prevalence,
  epiworld_double contact_rate,
  epiworld_double transmission_rate,
  epiworld_double vax_efficacy,
  epiworld_double vax_reduction_recovery_rate,
  epiworld_double incubation_period,
  epiworld_double prodromal_period,
  epiworld_double rash_period,
  std::vector< double > contact_matrix,
  epiworld_double hospitalization_rate,
  epiworld_double hospitalization_period,
  epiworld_double days_undetected,
  epiworld_fast_int quarantine_period_high,
  epiworld_fast_int quarantine_period_medium,
  epiworld_fast_int quarantine_period_low,
  epiworld_double quarantine_willingness,
  epiworld_double isolation_willingness,
  {\tt epiworld\_fast\_int} \ isolation\_period,
  epiworld_double prop_vaccinated,
  epiworld_double detection_rate_quarantine,
  epiworld_double contact_tracing_success_rate = 1.0,
  epiworld_fast_uint contact_tracing_days_prior = 4u) [inline]
```

## Constructs a ModelMeaslesMixingRiskQuarantine object.

n	The number of entities in the model.
prevalence	The initial prevalence of the disease in the model.
contact_rate	The contact rate between entities in the model.
transmission_rate	The transmission rate of the disease in the model.
vax_efficacy	The efficacy of the vaccine.
vax_reduction_recovery_rate	The reduction in recovery rate due to the vaccine.
incubation_period	The incubation period of the disease in the model.
prodromal_period	The prodromal period of the disease in the model.
rash_period	The rash period of the disease in the model.
contact_matrix	The contact matrix between entities in the model.
hospitalization_rate	The rate at which infected individuals are hospitalized.
hospitalization_period	The average duration of hospitalization in days.
days_undetected	The average number of days an infected individual remains undetected.
quarantine_period_high	The duration of quarantine in days for high-risk contacts.
quarantine_period_medium	The duration of quarantine in days for medium-risk contacts.
quarantine_period_low	The duration of quarantine in days for low-risk contacts.
quarantine_willingness	The proportion of individuals willing to comply with quarantine measures.
isolation_willingness	The proportion of individuals willing to self-isolate when detected.
isolation_period	The duration of isolation in days for detected infected individuals.
prop_vaccinated	The proportion of vaccinated agents.
detection_rate_quarantine	The detection rate during active quarantine periods.
contact_tracing_success_rate	The probability of successfully identifying and tracing contacts (default: 1.0).
contact_tracing_days_prior	The number of days prior to detection for which contacts are traced (default: 4).
	-

## 18.29.3 Member Function Documentation

## 18.29.3.1 clone ptr()

```
template<typename TSeq >
Model< TSeq > * ModelMeaslesMixingRiskQuarantine< TSeq >::clone_ptr [inline], [virtual]
```

Create a clone of this model.

#### Returns

Pointer to a new model instance with the same configuration

Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

## 18.29.3.2 get\_agent\_quarantine\_triggered()

```
template<typename TSeq = EPI_DEFAULT_TSEQ>
std::vector< size_t > epiworld::epimodels::ModelMeaslesMixingRiskQuarantine< TSeq >::get_
agent_quarantine_triggered ( ) const [inline]
```

Get the quarantine trigger status for all agents.

## Returns

Vector indicating quarantine process status for each agent

#### 18.29.3.3 get\_contact\_matrix()

```
template<typename TSeq = EPI_DEFAULT_TSEQ>
std::vector< double > epiworld::epimodels::ModelMeaslesMixingRiskQuarantine< TSeq >::get_
contact_matrix ( ) const [inline]
```

Get the current contact matrix.

## Returns

Vector representing the contact matrix

## 18.29.3.4 get\_isolation\_willingness()

```
template<typename TSeq = EPI_DEFAULT_TSEQ>
std::vector< bool > epiworld::epimodels::ModelMeaslesMixingRiskQuarantine< TSeq >::get_\(\cup \)
isolation_willingness ( ) const [inline]
```

Get the isolation willingness for all agents.

#### Returns

Vector of boolean values indicating each agent's willingness to self-isolate

## 18.29.3.5 get\_quarantine\_risk\_levels()

```
template<typename TSeq = EPI_DEFAULT_TSEQ>
std::vector< int > epiworld::epimodels::ModelMeaslesMixingRiskQuarantine< TSeq >::get_
quarantine_risk_levels ( ) const [inline]
```

Get the risk levels assigned to all agents.

## Returns

Vector of integers indicating each agent's risk level (0=low, 1=medium, 2=high)

## 18.29.3.6 get\_quarantine\_willingness()

```
template<typename TSeq = EPI_DEFAULT_TSEQ>
std::vector< bool > epiworld::epimodels::ModelMeaslesMixingRiskQuarantine< TSeq >::get_←
quarantine_willingness ( ) const [inline]
```

Get the quarantine willingness for all agents.

## Returns

Vector of boolean values indicating each agent's willingness to quarantine

## 18.29.3.7 initial\_states()

Set the initial states of the model.

#### **Parameters**

proportions↔	Double vector with two elements:
_	• [0]: The proportion of initially infected individuals who start in the exposed state.
	• [1]: The proportion of initially non-infected individuals who have recovered (immune).
queue_	Optional vector for queuing specifications (default: empty).

Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

#### 18.29.3.8 run()

Run the model simulation.

#### **Parameters**

ndays	Number of days to simulate
seed	Random seed for reproducibility (default: -1 for random seed)

## Returns

Reference to this model instance

 $\label{eq:local_problem} \mbox{Reimplemented from epiworld::} \mbox{Model} < \mbox{EPI\_DEFAULT\_TSEQ} >.$ 

## 18.29.3.9 set\_contact\_matrix()

Set the contact matrix for population mixing.

## **Parameters**

cmat	Contact matrix specifying interaction rates between groups

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

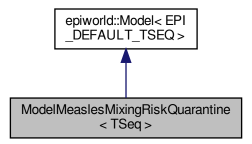
· epiworld.hpp

## 18.30 ModelMeaslesMixingRiskQuarantine< TSeq > Class Template Reference

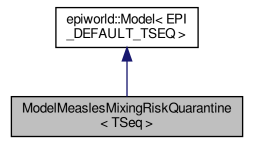
Measles model with population mixing and risk-based quarantine strategies.

#include <measlesmixingriskquarantine.hpp>

Inheritance diagram for ModelMeaslesMixingRiskQuarantine < TSeq >:



Collaboration diagram for ModelMeaslesMixingRiskQuarantine < TSeq >:



## **Public Member Functions**

ModelMeaslesMixingRiskQuarantine (ModelMeaslesMixingRiskQuarantine < TSeq > &model, epiworld\_←
fast\_uint n, epiworld\_double prevalence, epiworld\_double contact\_rate, epiworld\_double transmission←
\_rate, epiworld\_double vax\_efficacy, epiworld\_double vax\_reduction\_recovery\_rate, epiworld\_double
incubation\_period, epiworld\_double prodromal\_period, epiworld\_double rash\_period, std::vector < double >
contact\_matrix, epiworld\_double hospitalization\_rate, epiworld\_double hospitalization\_period, epiworld\_←
double days\_undetected, epiworld\_fast\_int quarantine\_period\_high, epiworld\_fast\_int quarantine\_period\_←
medium, epiworld\_fast\_int quarantine\_period\_low, epiworld\_double quarantine\_willingness, epiworld\_double
isolation\_willingness, epiworld\_fast\_int isolation\_period, epiworld\_double prop\_vaccinated, epiworld\_←
double detection\_rate\_quarantine, epiworld\_double contact\_tracing\_success\_rate=1.0, epiworld\_fast\_uint
contact\_tracing\_days\_prior=4u)

Constructs a ModelMeaslesMixingRiskQuarantine object.

ModelMeaslesMixingRiskQuarantine (epiworld\_fast\_uint n, epiworld\_double prevalence, epiworld\_double contact\_rate, epiworld\_double transmission\_rate, epiworld\_double vax\_efficacy, epiworld\_double vax\_ereduction\_recovery\_rate, epiworld\_double incubation\_period, epiworld\_double prodromal\_period, epiworld← \_\_double rash\_period, std::vector< double > contact\_matrix, epiworld\_double hospitalization\_rate, epiworld← \_\_double hospitalization\_period, epiworld\_double days\_undetected, epiworld\_fast\_int quarantine\_period← \_\_high, epiworld\_fast\_int quarantine\_period\_medium, epiworld\_fast\_int quarantine\_period\_low, epiworld← \_\_double quarantine\_willingness, epiworld\_double isolation\_willingness, epiworld\_fast\_int isolation\_period, epiworld\_double prop\_vaccinated, epiworld\_double detection\_rate\_quarantine, epiworld\_double contact\_← tracing\_success\_rate=1.0, epiworld\_fast\_uint contact\_tracing\_days\_prior=4u)

Constructs a ModelMeaslesMixingRiskQuarantine object.

ModelMeaslesMixingRiskQuarantine < TSeq > & run (epiworld fast uint ndays, int seed=-1)

Run the model simulation.

· void reset ()

Reset the model to initial state.

Model < TSeq > \* clone\_ptr ()

Create a clone of this model.

ModelMeaslesMixingRiskQuarantine < TSeq > & initial\_states (std::vector < double > proportions\_, std
 ::vector < int > queue\_={})

Set the initial states of the model.

void set contact matrix (std::vector< double > cmat)

Set the contact matrix for population mixing.

std::vector< double > get\_contact\_matrix () const

Get the current contact matrix.

std::vector< bool > get\_quarantine\_willingness () const

Get the quarantine willingness for all agents.

std::vector< bool > get isolation willingness () const

Get the isolation willingness for all agents.

## **Static Public Attributes**

- static constexpr int SUSCEPTIBLE = 0
- static constexpr int EXPOSED = 1
- static constexpr int **PRODROMAL** = 2
- static constexpr int RASH = 3
- static constexpr int ISOLATED = 4
- static constexpr int ISOLATED\_RECOVERED = 5
- static constexpr int **DETECTED HOSPITALIZED** = 6
- static constexpr int QUARANTINED\_EXPOSED = 7
- static constexpr int **QUARANTINED\_SUSCEPTIBLE** = 8
- static constexpr int **QUARANTINED\_PRODROMAL** = 9
- static constexpr int **QUARANTINED\_RECOVERED** = 10
- static constexpr int HOSPITALIZED = 11
- static constexpr int **RECOVERED** = 12
- static constexpr size\_t QUARANTINE\_PROCESS\_INACTIVE = 0u
- static constexpr size t QUARANTINE PROCESS ACTIVE = 1u
- static constexpr size t QUARANTINE PROCESS\_DONE = 2u
- static constexpr int RISK LOW = 0
- static constexpr int RISK\_MEDIUM = 1
- static constexpr int RISK\_HIGH = 2

## **Additional Inherited Members**

## 18.30.1 Detailed Description

```
template<typename TSeq = EPI_DEFAULT_TSEQ> class ModelMeaslesMixingRiskQuarantine< TSeq >
```

Measles model with population mixing and risk-based quarantine strategies.

This class extends the Measles epidemiological model to support different quarantine strategies based on exposure risk levels:

- · High Risk: Unvaccinated agents who share entity membership with the case
- Medium Risk: Unvaccinated agents who contacted an infected individual but don't share entity membership
- · Low Risk: Other unvaccinated agents

Each risk level can have different quarantine durations, allowing for targeted public health interventions. The model also includes enhanced detection during active quarantine periods.

Disease progression follows the same states as ModelMeaslesMixing: Susceptible  $\to$  Exposed  $\to$  Prodromal  $\to$  Rash  $\to$  Recovered

**Template Parameters** 

```
TSeq Type for genetic sequences (default: EPI_DEFAULT_TSEQ)
```

## 18.30.2 Constructor & Destructor Documentation

#### 18.30.2.1 ModelMeaslesMixingRiskQuarantine() [1/2]

```
epiworld_double days_undetected,
epiworld_fast_int quarantine_period_high,
epiworld_fast_int quarantine_period_medium,
epiworld_fast_int quarantine_period_low,
epiworld_double quarantine_willingness,
epiworld_double isolation_willingness,
epiworld_fast_int isolation_period,
epiworld_double prop_vaccinated,
epiworld_double detection_rate_quarantine,
epiworld_double contact_tracing_success_rate = 1.0,
epiworld_fast_uint contact_tracing_days_prior = 4u )
```

## Constructs a ModelMeaslesMixingRiskQuarantine object.

## **Parameters**

model	A reference to an existing ModelMeaslesMixingRiskQuarantine object.
n	The number of entities in the model.
prevalence	The initial prevalence of the disease in the model.
contact_rate	The contact rate between entities in the model.
transmission_rate	The transmission rate of the disease in the model.
vax_efficacy	The efficacy of the vaccine.
vax_reduction_recovery_rate	The reduction in recovery rate due to the vaccine.
incubation_period	The incubation period of the disease in the model.
prodromal_period	The prodromal period of the disease in the model.
rash_period	The rash period of the disease in the model.
contact_matrix	The contact matrix between entities in the model. Specified in column-major
	order.
hospitalization_rate	The rate at which infected individuals are hospitalized.
hospitalization_period	The average duration of hospitalization in days.
days_undetected	The average number of days an infected individual remains undetected.
quarantine_period_high	The duration of quarantine in days for high-risk contacts.
quarantine_period_medium	The duration of quarantine in days for medium-risk contacts.
quarantine_period_low	The duration of quarantine in days for low-risk contacts.
quarantine_willingness	The proportion of individuals willing to comply with quarantine measures.
isolation_willingness	The proportion of individuals willing to self-isolate when detected.
isolation_period	The duration of isolation in days for detected infected individuals.
prop_vaccinated	The proportion of vaccinated agents.
detection_rate_quarantine	The detection rate during active quarantine periods.
contact_tracing_success_rate	The probability of successfully identifying and tracing contacts (default: 1.0).
contact_tracing_days_prior	The number of days prior to detection for which contacts are traced (default:
	4).

## 18.30.2.2 ModelMeaslesMixingRiskQuarantine() [2/2]

```
epiworld_double contact_rate,
epiworld_double transmission_rate,
epiworld_double vax_efficacy,
epiworld_double vax_reduction_recovery_rate,
epiworld_double incubation_period,
epiworld_double prodromal_period,
epiworld_double rash_period,
std::vector< double > contact_matrix,
epiworld_double hospitalization_rate,
epiworld_double hospitalization_period,
epiworld_double days_undetected,
epiworld_fast_int quarantine_period_high,
epiworld_fast_int quarantine_period_medium,
epiworld_fast_int quarantine_period_low,
epiworld_double quarantine_willingness,
epiworld_double isolation_willingness,
epiworld_fast_int isolation_period,
epiworld_double prop_vaccinated,
epiworld_double detection_rate_quarantine,
epiworld_double contact_tracing_success_rate = 1.0,
epiworld_fast_uint contact_tracing_days_prior = 4u )
```

## Constructs a ModelMeaslesMixingRiskQuarantine object.

n	The number of entities in the model.
prevalence	The initial prevalence of the disease in the model.
contact_rate	The contact rate between entities in the model.
transmission_rate	The transmission rate of the disease in the model.
vax_efficacy	The efficacy of the vaccine.
vax_reduction_recovery_rate	The reduction in recovery rate due to the vaccine.
incubation_period	The incubation period of the disease in the model.
prodromal_period	The prodromal period of the disease in the model.
rash_period	The rash period of the disease in the model.
contact_matrix	The contact matrix between entities in the model.
hospitalization_rate	The rate at which infected individuals are hospitalized.
hospitalization_period	The average duration of hospitalization in days.
days_undetected	The average number of days an infected individual remains undetected.
quarantine_period_high	The duration of quarantine in days for high-risk contacts.
quarantine_period_medium	The duration of quarantine in days for medium-risk contacts.
quarantine_period_low	The duration of quarantine in days for low-risk contacts.
quarantine_willingness	The proportion of individuals willing to comply with quarantine measures.
isolation_willingness	The proportion of individuals willing to self-isolate when detected.
isolation_period	The duration of isolation in days for detected infected individuals.
prop_vaccinated	The proportion of vaccinated agents.
detection_rate_quarantine	The detection rate during active quarantine periods.
contact_tracing_success_rate	The probability of successfully identifying and tracing contacts (default: 1.0).
contact_tracing_days_prior	The number of days prior to detection for which contacts are traced (default: 4).

## 18.30.3 Member Function Documentation

## 18.30.3.1 clone ptr()

```
template<typename TSeq = EPI_DEFAULT_TSEQ>
Model<TSeq>* ModelMeaslesMixingRiskQuarantine< TSeq >::clone_ptr ( ) [virtual]
```

Create a clone of this model.

## Returns

Pointer to a new model instance with the same configuration

Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

## 18.30.3.2 get\_contact\_matrix()

```
template<typename TSeq = EPI_DEFAULT_TSEQ>
std::vector< double > ModelMeaslesMixingRiskQuarantine< TSeq >::get_contact_matrix ( ) const
[inline]
```

Get the current contact matrix.

## Returns

Vector representing the contact matrix

## 18.30.3.3 get\_isolation\_willingness()

```
template<typename TSeq = EPI_DEFAULT_TSEQ>
std::vector< bool > ModelMeaslesMixingRiskQuarantine< TSeq >::get_isolation_willingness ( )
const [inline]
```

Get the isolation willingness for all agents.

## Returns

Vector of boolean values indicating each agent's willingness to self-isolate

## 18.30.3.4 get\_quarantine\_willingness()

```
template<typename TSeq = EPI_DEFAULT_TSEQ>
std::vector< bool > ModelMeaslesMixingRiskQuarantine< TSeq >::get_quarantine_willingness ( )
const [inline]
```

Get the quarantine willingness for all agents.

## Returns

Vector of boolean values indicating each agent's willingness to quarantine

## 18.30.3.5 initial\_states()

Set the initial states of the model.

## **Parameters**

proportions⇔	Double vector with two elements:
_	• [0]: The proportion of initially infected individuals who start in the exposed state.
	• [1]: The proportion of initially non-infected individuals who have recovered (immune).
queue_	Optional vector for queuing specifications (default: empty).

 $\label{eq:control_equation} \mbox{Reimplemented from epiworld::} \mbox{Model} < \mbox{EPI\_DEFAULT\_TSEQ} >.$ 

## 18.30.3.6 run()

Run the model simulation.

ndays	Number of days to simulate
seed	Random seed for reproducibility (default: -1 for random seed)

#### Returns

Reference to this model instance

Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

# 18.30.3.7 set\_contact\_matrix()

Set the contact matrix for population mixing.

#### **Parameters**

cmat Contact matrix specifying interaction rates between groups

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

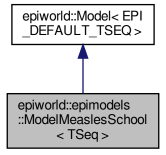
• include/epiworld/models/measlesmixingriskquarantine.hpp

# 18.31 epiworld::epimodels::ModelMeaslesSchool < TSeq > Class Template Reference

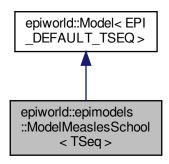
Template for a Measles model with quarantine.

```
#include <epiworld.hpp>
```

 $Inheritance\ diagram\ for\ epiworld::epimodels::ModelMeaslesSchool<\ TSeq>:$ 



Collaboration diagram for epiworld::epimodels::ModelMeaslesSchool < TSeq >:



#### **Public Member Functions**

• void quarantine agents ()

Quarantine agents that are in the system.

· void reset ()

Reset the model.

- void update\_infectious ()
- Model < TSeq > \* clone\_ptr ()

Advanced usage: Makes a copy of data and returns it as undeleted pointer.

- ModelMeaslesSchool (ModelMeaslesSchool < TSeq > &model, epiworld\_fast\_uint n, epiworld\_fast\_uint n\_exposed, epiworld\_double contact\_rate, epiworld\_double transmission\_rate, epiworld\_double vax
   \_efficacy, epiworld\_double vax\_reduction\_recovery\_rate, epiworld\_double incubation\_period, epiworld\_double prodromal\_period, epiworld\_double rash\_period, epiworld\_double days\_undetected, epiworld\_double hospitalization\_rate, epiworld\_double hospitalization\_period, epiworld\_double prop\_vaccinated, epiworld\_double fast int quarantine period, epiworld double quarantine willingness, epiworld fast int isolation period)
- ModelMeaslesSchool (epiworld\_fast\_uint n, epiworld\_fast\_uint n\_exposed, epiworld\_double contact\_
   rate, epiworld\_double transmission\_rate, epiworld\_double vax\_efficacy, epiworld\_double vax\_reduction
   \_recovery\_rate, epiworld\_double incubation\_period, epiworld\_double prodromal\_period, epiworld\_double
   rash\_period, epiworld\_double days\_undetected, epiworld\_double hospitalization\_rate, epiworld\_double
   hospitalization\_period, epiworld\_double prop\_vaccinated, epiworld\_fast\_int quarantine\_period, epiworld\_
   double quarantine\_willingness, epiworld\_fast\_int isolation\_period)

# **Public Attributes**

std::vector < Agent < TSeq > \* > infectious

Agents infectious for contact.

- bool system\_quarantine\_triggered = false
- std::vector< int > day\_flagged

Either detected or started quarantine.

std::vector< int > day\_rash\_onset

Day of rash onset.

#### **Static Public Attributes**

- static constexpr epiworld\_fast\_uint SUSCEPTIBLE = 0u
- static constexpr epiworld\_fast\_uint EXPOSED = 1u
- static constexpr epiworld fast uint PRODROMAL = 2u
- static constexpr epiworld\_fast\_uint RASH = 3u
- static constexpr epiworld fast uint ISOLATED = 4u
- static constexpr epiworld\_fast\_uint ISOLATED\_RECOVERED = 5u
- static constexpr epiworld\_fast\_uint **DETECTED\_HOSPITALIZED** = 6u
- static constexpr epiworld fast uint QUARANTINED EXPOSED = 7u
- static constexpr epiworld\_fast\_uint QUARANTINED\_SUSCEPTIBLE = 8u
- static constexpr epiworld fast uint QUARANTINED PRODROMAL = 9u
- static constexpr epiworld\_fast\_uint QUARANTINED\_RECOVERED = 10u
- static constexpr epiworld\_fast\_uint HOSPITALIZED = 11u
- static constexpr epiworld\_fast\_uint RECOVERED = 12u

#### **Additional Inherited Members**

# 18.31.1 Detailed Description

```
template<typename TSeq = EPI_DEFAULT_TSEQ> class epiworld::epimodels::ModelMeaslesSchool< TSeq >
```

Template for a Measles model with quarantine.

#### **Parameters**

TSeq	The type of the sequence to be used.
------	--------------------------------------

This model can be described as a SEIHR model with isolation and quarantine. The infectious state is divided into prodromal and rash phases. Furthermore, the quarantine state includes exposed, susceptible, prodromal, and recovered states.

The quarantine process is triggered any time that an agent with rash is detected. The agent is then isolated and all agents who are unvaccinated are quarantined. Isolated agents then may be moved out of the isolation in isolation ← \_period days.

# 18.31.2 Constructor & Destructor Documentation

#### 18.31.2.1 ModelMeaslesSchool()

```
epiworld_double transmission_rate,
epiworld_double vax_efficacy,
epiworld_double vax_reduction_recovery_rate,
epiworld_double incubation_period,
epiworld_double prodromal_period,
epiworld_double days_undetected,
epiworld_double hospitalization_rate,
epiworld_double hospitalization_period,
epiworld_double prop_vaccinated,
epiworld_fast_int quarantine_period,
epiworld_double quarantine_willingness,
epiworld_fast_int isolation_period) [inline]
```

#### **Parameters**

n	The number of agents in the system.
n_exposed	The number of exposed agents in the system.
contact_rate	The rate of contact between agents.
transmission_rate	The rate of transmission of the virus.
vax_efficacy	The efficacy of the vaccine.
vax_reduction_recovery_rate	The reduction in recovery rate due to the vaccine.
incubation_period	The incubation period of the virus.
prodromal_period	The prodromal period of the virus.
rash_period	The rash period of the virus.
days_undetected	The number of days the virus goes undetected.
hospitalization_rate	The rate of hospitalization.
hospitalization_period	The duration of hospitalization.
prop_vaccinated	The proportion of vaccinated agents.
quarantine_period	The number of days for quarantine.
quarantine_willingness	The willingness to be quarantined.
isolation_period	The number of days for isolation.

# 18.31.3 Member Function Documentation

# 18.31.3.1 clone\_ptr()

```
template<typename TSeq >
Model< TSeq > * ModelMeaslesSchool< TSeq >::clone_ptr [inline], [virtual]
```

Advanced usage: Makes a copy of data and returns it as undeleted pointer.

#### **Parameters**



Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

#### 18.31.3.2 quarantine agents()

```
template<typename TSeq >
void ModelMeaslesSchool< TSeq >::quarantine_agents [inline]
```

Quarantine agents that are in the system.

The flow should be:

- The function only runs if the quarantine status is active.
- · Agents who are in quarantine, isolation, removed, or hospitalized are ignored.
- · Agents who are in the RASH state are isolated.
- Vaccinated agents are ignored.
- Susceptible, Exposed, and Prodromal agents are moved to the QUARANTINED\_\* state.
- At the end of the function, the quarantine status is set false.

#### 18.31.3.3 reset()

```
template<typename TSeq >
void ModelMeaslesSchool< TSeq >::reset [inline], [virtual]
```

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

Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

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

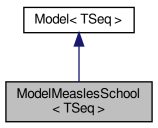
· epiworld.hpp

# 18.32 ModelMeaslesSchool < TSeq > Class Template Reference

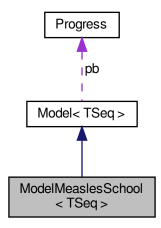
Template for a Measles model with quarantine.

#include <measlesquarantine.hpp>

Inheritance diagram for ModelMeaslesSchool < TSeq >:



 $\label{local_continuous_continuous_continuous_continuous} Collaboration \ diagram \ for \ ModelMeaslesSchool < TSeq >:$ 



# **Public Member Functions**

• void quarantine\_agents ()

Quarantine agents that are in the system.

· void reset ()

Reset the model.

• void update\_infectious ()

Model < TSeq > \* clone\_ptr ()

Advanced usage: Makes a copy of data and returns it as undeleted pointer.

- ModelMeaslesSchool (ModelMeaslesSchool < TSeq > &model, epiworld\_fast\_uint n, epiworld\_fast\_uint n\_exposed, epiworld\_double contact\_rate, epiworld\_double transmission\_rate, epiworld\_double vax
   \_efficacy, epiworld\_double vax\_reduction\_recovery\_rate, epiworld\_double incubation\_period, epiworld\_←
   double prodromal\_period, epiworld\_double rash\_period, epiworld\_double days\_undetected, epiworld\_double hospitalization\_rate, epiworld\_double hospitalization\_period, epiworld\_double prop\_vaccinated, epiworld\_←
   fast int quarantine period, epiworld double quarantine willingness, epiworld fast int isolation period)
- ModelMeaslesSchool (epiworld\_fast\_uint n, epiworld\_fast\_uint n\_exposed, epiworld\_double contact\_
   rate, epiworld\_double transmission\_rate, epiworld\_double vax\_efficacy, epiworld\_double vax\_reduction
   \_recovery\_rate, epiworld\_double incubation\_period, epiworld\_double prodromal\_period, epiworld\_double
   rash\_period, epiworld\_double days\_undetected, epiworld\_double hospitalization\_rate, epiworld\_double
   hospitalization\_period, epiworld\_double prop\_vaccinated, epiworld\_fast\_int quarantine\_period, epiworld\_
   double quarantine\_willingness, epiworld\_fast\_int isolation\_period)

#### **Public Attributes**

- std::vector< Agent< TSeq > \* > infectious
  - Agents infectious for contact.
- bool system\_quarantine\_triggered = false
- std::vector< int > day\_flagged

Either detected or started quarantine.

std::vector< int > day\_rash\_onset

Day of rash onset.

# **Static Public Attributes**

- static constexpr epiworld fast uint SUSCEPTIBLE = 0u
- static constexpr epiworld\_fast\_uint EXPOSED = 1u
- static constexpr epiworld\_fast\_uint PRODROMAL = 2u
- static constexpr epiworld\_fast\_uint RASH = 3u
- static constexpr epiworld\_fast\_uint ISOLATED = 4u
- static constexpr epiworld fast uint ISOLATED\_RECOVERED = 5u
- static constexpr epiworld\_fast\_uint DETECTED\_HOSPITALIZED = 6u
- static constexpr epiworld\_fast\_uint QUARANTINED\_EXPOSED = 7u
- static constexpr epiworld\_fast\_uint **QUARANTINED\_SUSCEPTIBLE** = 8u
- static constexpr epiworld\_fast\_uint **QUARANTINED\_PRODROMAL** = 9u
- static constexpr epiworld fast uint QUARANTINED\_RECOVERED = 10u
- static constexpr epiworld\_fast\_uint HOSPITALIZED = 11u
- static constexpr epiworld fast uint RECOVERED = 12u

### **Additional Inherited Members**

# 18.32.1 Detailed Description

template<typename TSeq = EPI\_DEFAULT\_TSEQ> class ModelMeaslesSchool< TSeq >

Template for a Measles model with quarantine.

#### **Parameters**

TSeq The type of the sequence to be used.

This model can be described as a SEIHR model with isolation and quarantine. The infectious state is divided into prodromal and rash phases. Furthermore, the quarantine state includes exposed, susceptible, prodromal, and recovered states.

The quarantine process is triggered any time that an agent with rash is detected. The agent is then isolated and all agents who are unvaccinated are quarantined. Isolated agents then may be moved out of the isolation in isolation period days.

# 18.32.2 Constructor & Destructor Documentation

# 18.32.2.1 ModelMeaslesSchool()

```
template<typename TSeq >
ModelMeaslesSchool < TSeq >::ModelMeaslesSchool (
             ModelMeaslesSchool< TSeq > & model,
             epiworld_fast_uint n,
             epiworld_fast_uint n_exposed,
             epiworld_double contact_rate,
             epiworld_double transmission_rate,
             epiworld_double vax_efficacy,
             epiworld_double vax_reduction_recovery_rate,
             epiworld_double incubation_period,
             epiworld_double prodromal_period,
             epiworld_double rash_period,
             epiworld_double days_undetected,
             epiworld_double hospitalization_rate,
             epiworld_double hospitalization_period,
             epiworld_double prop_vaccinated,
             epiworld_fast_int quarantine_period,
             epiworld_double quarantine_willingness,
             epiworld_fast_int isolation_period ) [inline]
```

#### **Parameters**

n	The number of agents in the system.
n_exposed	The number of exposed agents in the system.
contact_rate	The rate of contact between agents.
transmission_rate	The rate of transmission of the virus.
vax_efficacy	The efficacy of the vaccine.
vax_reduction_recovery_rate	The reduction in recovery rate due to the vaccine.
incubation_period	The incubation period of the virus.
prodromal_period	The prodromal period of the virus.
rash_period	The rash period of the virus.
days_undetected	The number of days the virus goes undetected.
hospitalization_rate	The rate of hospitalization.

#### **Parameters**

hospitalization_period	The duration of hospitalization.
prop_vaccinated	The proportion of vaccinated agents.
quarantine_period	The number of days for quarantine.
quarantine_willingness	The willingness to be quarantined.
isolation_period	The number of days for isolation.

# 18.32.3 Member Function Documentation

#### 18.32.3.1 clone ptr()

```
template<typename TSeq >
Model< TSeq > * ModelMeaslesSchool< TSeq >::clone_ptr [inline], [virtual]
```

Advanced usage: Makes a copy of data and returns it as undeleted pointer.

#### **Parameters**

сору

Reimplemented from Model < TSeq >.

#### 18.32.3.2 quarantine agents()

```
template<typename TSeq >
void ModelMeaslesSchool< TSeq >::quarantine_agents [inline]
```

Quarantine agents that are in the system.

The flow should be:

- The function only runs if the quarantine status is active.
- · Agents who are in quarantine, isolation, removed, or hospitalized are ignored.
- · Agents who are in the RASH state are isolated.
- · Vaccinated agents are ignored.
- Susceptible, Exposed, and Prodromal agents are moved to the QUARANTINED\_\* state.
- At the end of the function, the quarantine status is set false.

#### 18.32.3.3 reset()

```
template<typename TSeq >
void ModelMeaslesSchool< TSeq >::reset [inline], [virtual]
```

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

Reimplemented from Model < TSeq >.

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

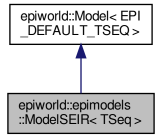
• include/epiworld/models/measlesquarantine.hpp

# 18.33 epiworld::epimodels::ModelSEIR< TSeq > Class Template Reference

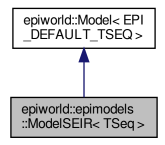
Template for a Susceptible-Exposed-Infected-Removed (SEIR) model.

```
#include <epiworld.hpp>
```

Inheritance diagram for epiworld::epimodels::ModelSEIR < TSeq >:



Collaboration diagram for epiworld::epimodels::ModelSEIR< TSeq >:



#### **Public Member Functions**

- **ModelSEIR** (ModelSEIR< TSeq > &model, const std::string &vname, epiworld\_double prevalence, epiworld\_double transmission\_rate, epiworld\_double avg\_incubation\_days, epiworld\_double recovery\_← rate)
- **ModelSEIR** (const std::string &vname, epiworld\_double prevalence, epiworld\_double transmission\_rate, epiworld\_double avg\_incubation\_days, epiworld\_double recovery\_rate)
- ModelSEIR < TSeq > & initial\_states (std::vector < double > proportions\_, std::vector < int > queue\_={})
   Set up the initial states of the model.

# **Public Attributes**

- epiworld::UpdateFun< TSeq > update exposed seir
- epiworld::UpdateFun< TSeq > update\_infected\_seir

# **Static Public Attributes**

- static const int SUSCEPTIBLE = 0
- static const int **EXPOSED** = 1
- static const int **INFECTED** = 2
- static const int **REMOVED** = 3

#### **Additional Inherited Members**

# 18.33.1 Detailed Description

template<typename TSeq = EPI\_DEFAULT\_TSEQ> class epiworld::epimodels::ModelSEIR< TSeq >

Template for a Susceptible-Exposed-Infected-Removed (SEIR) model.

#### **Parameters**

model	A Model <tseq> object where to set up the SIR.</tseq>
vname	std::string Name of the virus
prevalence	epiworld_double Initial prevalence the immune system
transmission_rate	epiworld_double Transmission rate of the virus
avg_incubation_days	epiworld_double Average incubation days of the virus.
recovery_rate	epiworld_double Recovery rate of the virus.

# 18.33.2 Member Function Documentation

# 18.33.2.1 initial\_states()

Set up the initial states of the model.

#### **Parameters**

proportions⊷	Double vector with the following values:
_	0: Proportion of non-infected agents who are removed.
	1: Proportion of exposed agents to be set as infected.

Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

# 18.33.3 Member Data Documentation

# 18.33.3.1 update\_exposed\_seir

#### 18.33.3.2 update\_infected\_seir

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

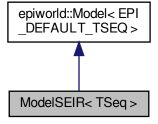
· epiworld.hpp

# 18.34 ModelSEIR < TSeq > Class Template Reference

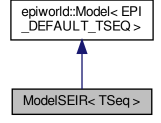
Template for a Susceptible-Exposed-Infected-Removed (SEIR) model.

```
#include <seir.hpp>
```

Inheritance diagram for ModelSEIR< TSeq >:



Collaboration diagram for ModelSEIR< TSeq >:



# **Public Member Functions**

• **ModelSEIR** (ModelSEIR< TSeq > &model, const std::string &vname, epiworld\_double prevalence, epiworld\_double transmission\_rate, epiworld\_double avg\_incubation\_days, epiworld\_double recovery\_← rate)

- **ModelSEIR** (const std::string &vname, epiworld\_double prevalence, epiworld\_double transmission\_rate, epiworld double avg incubation days, epiworld double recovery rate)
- ModelSEIR < TSeq > & initial\_states (std::vector < double > proportions\_, std::vector < int > queue\_={})
   Set up the initial states of the model.

# **Public Attributes**

- epiworld::UpdateFun< TSeq > update\_exposed\_seir
- epiworld::UpdateFun< TSeq > update infected seir

# **Static Public Attributes**

- static const int SUSCEPTIBLE = 0
- static const int EXPOSED = 1
- static const int INFECTED = 2
- static const int **REMOVED** = 3

#### **Additional Inherited Members**

# 18.34.1 Detailed Description

 $\label{template} \mbox{template} < \mbox{typename TSeq} = \mbox{EPI\_DEFAULT\_TSEQ} > \\ \mbox{class ModelSEIR} < \mbox{TSeq} > \\ \mbox{}$ 

Template for a Susceptible-Exposed-Infected-Removed (SEIR) model.

#### **Parameters**

model	A Model <tseq> object where to set up the SIR.</tseq>
vname	std::string Name of the virus
prevalence	epiworld_double Initial prevalence the immune system
transmission_rate	epiworld_double Transmission rate of the virus
avg_incubation_days	epiworld_double Average incubation days of the virus.
recovery_rate	epiworld_double Recovery rate of the virus.

# 18.34.2 Member Function Documentation

#### 18.34.2.1 initial\_states()

Set up the initial states of the model.

#### **Parameters**

# proportions↔ Double vector with the following values: 0: Proportion of non-infected agents who are removed. 1: Proportion of exposed agents to be set as infected.

Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

# 18.34.3 Member Data Documentation

#### 18.34.3.1 update\_exposed\_seir

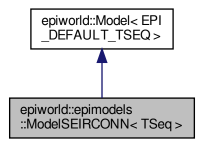
# 18.34.3.2 update\_infected\_seir

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

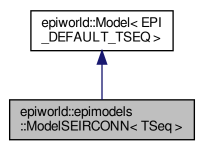
• include/epiworld/models/seir.hpp

# 18.35 epiworld::epimodels::ModelSEIRCONN< TSeq > Class Template Reference

Inheritance diagram for epiworld::epimodels::ModelSEIRCONN < TSeq >:



Collaboration diagram for epiworld::epimodels::ModelSEIRCONN< TSeq >:



# **Public Member Functions**

ModelSEIRCONN (ModelSEIRCONN
 TSeq > &model, const std::string &vname, epiworld\_fast\_uint n, epiworld\_double prevalence, epiworld\_double contact\_rate, epiworld\_double transmission\_rate, epiworld
 \_double avg\_incubation\_days, epiworld\_double recovery\_rate)

Template for a Susceptible-Exposed-Infected-Removed (SEIR) model.

- ModelSEIRCONN (const std::string &vname, epiworld\_fast\_uint n, epiworld\_double prevalence, epiworld
   —double contact\_rate, epiworld\_double transmission\_rate, epiworld\_double avg\_incubation\_days, epiworld
   —double recovery\_rate)
- ModelSEIRCONN
   TSeq > & run (epiworld\_fast\_uint ndays, int seed=-1)

Runs the simulation (after initialization)

• void reset ()

Reset the model.

Model < TSeq > \* clone\_ptr ()

Advanced usage: Makes a copy of data and returns it as undeleted pointer.

ModelSEIRCONN< TSeq > & initial\_states (std::vector< double > proportions\_, std::vector< int > queue ← \_={})

Set the initial states of the model.

- size\_t get\_n\_infected () const
- std::vector< double > generation\_time\_expected (int max\_days=200, int max\_contacts=200) const

#### **Static Public Attributes**

- static const int SUSCEPTIBLE = 0
- static const int **EXPOSED** = 1
- static const int INFECTED = 2
- static const int RECOVERED = 3

#### **Additional Inherited Members**

#### 18.35.1 Constructor & Destructor Documentation

#### 18.35.1.1 ModelSEIRCONN()

Template for a Susceptible-Exposed-Infected-Removed (SEIR) model.

#### **Parameters**

model	A Model <tseq> object where to set up the SIR.</tseq>
vname	std::string Name of the virus
prevalence	Initial prevalence (proportion)
contact_rate	Average number of contacts (interactions) per step.
transmission_rate	Probability of transmission
recovery_rate	Probability of recovery

# 18.35.2 Member Function Documentation

#### 18.35.2.1 clone\_ptr()

```
template<typename TSeq >
Model< TSeq > * ModelSEIRCONN< TSeq >::clone_ptr [inline], [virtual]
```

Advanced usage: Makes a copy of data and returns it as undeleted pointer.

#### **Parameters**

```
сору
```

Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

# 18.35.2.2 initial\_states()

Set the initial states of the model.

#### **Parameters**

proportions↔	Double vector with a single element:
_	The proportion of non-infected individuals who have recovered.

 $\label{eq:control_equation} \mbox{Reimplemented from epiworld::} \mbox{Model} < \mbox{EPI\_DEFAULT\_TSEQ} >.$ 

#### 18.35.2.3 reset()

```
template<typename TSeq >
void ModelSEIRCONN< TSeq >::reset [inline], [virtual]
```

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

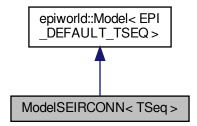
Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

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

epiworld.hpp

# 18.36 ModelSEIRCONN< TSeq > Class Template Reference

Inheritance diagram for ModelSEIRCONN< TSeq >:



Collaboration diagram for ModelSEIRCONN< TSeq >:



# **Public Member Functions**

ModelSEIRCONN (ModelSEIRCONN 
 TSeq > &model, const std::string &vname, epiworld\_fast\_uint n, epiworld\_double prevalence, epiworld\_double contact\_rate, epiworld\_double transmission\_rate, epiworld—double avg\_incubation\_days, epiworld\_double recovery\_rate)

Template for a Susceptible-Exposed-Infected-Removed (SEIR) model.

- ModelSEIRCONN (const std::string &vname, epiworld\_fast\_uint n, epiworld\_double prevalence, epiworld
   — double contact\_rate, epiworld\_double transmission\_rate, epiworld\_double avg\_incubation\_days, epiworld
   — double recovery\_rate)
- ModelSEIRCONN < TSeq > & run (epiworld\_fast\_uint ndays, int seed=-1)

Runs the simulation (after initialization)

· void reset ()

Reset the model.

Model < TSeq > \* clone ptr ()

Advanced usage: Makes a copy of data and returns it as undeleted pointer.

ModelSEIRCONN< TSeq > & initial\_states (std::vector< double > proportions\_, std::vector< int > queue ← \_={})

Set the initial states of the model.

- size\_t get\_n\_infected () const
- std::vector< double > generation\_time\_expected (int max\_days=200, int max\_contacts=200) const

#### **Static Public Attributes**

- static const int SUSCEPTIBLE = 0
- static const int **EXPOSED** = 1
- static const int INFECTED = 2
- static const int **RECOVERED** = 3

#### **Additional Inherited Members**

#### 18.36.1 Constructor & Destructor Documentation

#### 18.36.1.1 ModelSEIRCONN()

Template for a Susceptible-Exposed-Infected-Removed (SEIR) model.

# **Parameters**

model	A Model <tseq> object where to set up the SIR.</tseq>
vname	std::string Name of the virus
prevalence	Initial prevalence (proportion)
contact_rate	Average number of contacts (interactions) per step.
transmission_rate	Probability of transmission
recovery_rate	Probability of recovery

#### 18.36.2 Member Function Documentation

#### 18.36.2.1 clone\_ptr()

```
template<typename TSeq >
Model< TSeq > * ModelSEIRCONN< TSeq >::clone_ptr [inline], [virtual]
```

Advanced usage: Makes a copy of data and returns it as undeleted pointer.

#### **Parameters**

```
сору
```

Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

# 18.36.2.2 initial\_states()

Set the initial states of the model.

#### **Parameters**

proportions⊷	Double vector with a single element:
_	The proportion of non-infected individuals who have recovered.

 $\label{eq:control_equation} \mbox{Reimplemented from epiworld::} \mbox{Model} < \mbox{EPI\_DEFAULT\_TSEQ} >.$ 

#### 18.36.2.3 reset()

```
template<typename TSeq >
void ModelSEIRCONN< TSeq >::reset [inline], [virtual]
```

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

Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

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

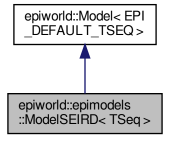
• include/epiworld/models/seirconnected.hpp

# 18.37 epiworld::epimodels::ModelSEIRD< TSeq > Class Template Reference

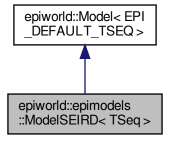
Template for a Susceptible-Exposed-Infected-Removed-Deceased (SEIRD) model.

#include <epiworld.hpp>

Inheritance diagram for epiworld::epimodels::ModelSEIRD< TSeq >:



Collaboration diagram for epiworld::epimodels::ModelSEIRD< TSeq >:



# **Public Member Functions**

• ModelSEIRD (ModelSEIRD< TSeq > &model, const std::string &vname, epiworld\_double prevalence, epiworld\_double transmission\_rate, epiworld\_double avg\_incubation\_days, epiworld\_double recovery\_rate, epiworld\_double death\_rate)

Constructor for the SEIRD model.

• ModelSEIRD (const std::string &vname, epiworld\_double prevalence, epiworld\_double transmission\_rate, epiworld\_double avg\_incubation\_days, epiworld\_double recovery\_rate, epiworld\_double death\_rate)

Constructor for the SEIRD model.

• ModelSEIRD< TSeq > & initial\_states (std::vector< double > proportions\_, std::vector< int > queue\_={})

# **Public Attributes**

- epiworld::UpdateFun< TSeq > update\_exposed\_seir
- epiworld::UpdateFun< TSeq > update\_infected

#### **Static Public Attributes**

- static const int SUSCEPTIBLE = 0
- static const int EXPOSED = 1
- static const int INFECTED = 2
- static const int **REMOVED** = 3
- static const int **DECEASED** = 4

# **Additional Inherited Members**

# 18.37.1 Detailed Description

```
template<typename TSeq = EPI_DEFAULT_TSEQ> class epiworld::epimodels::ModelSEIRD< TSeq >
```

Template for a Susceptible-Exposed-Infected-Removed-Deceased (SEIRD) model.

# 18.37.2 Constructor & Destructor Documentation

### 18.37.2.1 ModelSEIRD() [1/2]

Constructor for the SEIRD model.

# **Template Parameters**

TSeq	Type of the sequence used in the model.

#### **Parameters**

model	Reference to the SEIRD model.
vname	Name of the model.

#### **Parameters**

prevalence	ence Prevalence of the disease.	
transmission_rate Transmission rate of the disease.		
avg_incubation_days	Average incubation period of the disease.	
recovery_rate	Recovery rate of the disease.	
death_rate	Death rate of the disease.	

# 18.37.2.2 ModelSEIRD() [2/2]

#### Constructor for the SEIRD model.

#### **Parameters**

vname	name Name of the model.	
prevalence	Initial prevalence of the disease.	
transmission_rate	Transmission rate of the disease.	
avg_incubation_days	Average incubation period of the disease.	
recovery_rate Recovery rate of the disease.		
death_rate	Death rate of the disease.	

#### 18.37.3 Member Data Documentation

# 18.37.3.1 update\_exposed\_seir

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

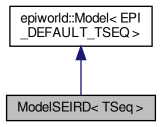
· epiworld.hpp

# **18.38** ModelSEIRD< TSeq > Class Template Reference

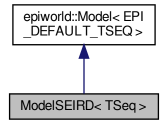
Template for a Susceptible-Exposed-Infected-Removed-Deceased (SEIRD) model.

```
#include <seird.hpp>
```

Inheritance diagram for ModelSEIRD< TSeq >:



Collaboration diagram for ModelSEIRD< TSeq >:



# **Public Member Functions**

• ModelSEIRD (ModelSEIRD< TSeq > &model, const std::string &vname, epiworld\_double prevalence, epiworld\_double transmission\_rate, epiworld\_double avg\_incubation\_days, epiworld\_double recovery\_rate, epiworld double death rate)

Constructor for the SEIRD model.

 ModelSEIRD (const std::string &vname, epiworld\_double prevalence, epiworld\_double transmission\_rate, epiworld\_double avg\_incubation\_days, epiworld\_double recovery\_rate, epiworld\_double death\_rate)

Constructor for the SEIRD model.

• ModelSEIRD< TSeq > & initial\_states (std::vector< double > proportions\_, std::vector< int > queue\_={})

# **Public Attributes**

- epiworld::UpdateFun< TSeq > update\_exposed\_seir
- epiworld::UpdateFun< TSeq > update\_infected

#### **Static Public Attributes**

- static const int SUSCEPTIBLE = 0
- static const int **EXPOSED** = 1
- static const int INFECTED = 2
- static const int **REMOVED** = 3
- static const int **DECEASED** = 4

# **Additional Inherited Members**

# 18.38.1 Detailed Description

```
template<typename TSeq = EPI_DEFAULT_TSEQ> class ModelSEIRD< TSeq >
```

Template for a Susceptible-Exposed-Infected-Removed-Deceased (SEIRD) model.

# 18.38.2 Constructor & Destructor Documentation

### 18.38.2.1 ModelSEIRD() [1/2]

Constructor for the SEIRD model.

# **Template Parameters**

TSeq Type of the sequence used in the model.
----------------------------------------------

#### **Parameters**

model	Reference to the SEIRD model.
vname	Name of the model.

#### **Parameters**

prevalence	ence Prevalence of the disease.	
transmission_rate Transmission rate of the disease.		
avg_incubation_days	Average incubation period of the disease.	
recovery_rate	Recovery rate of the disease.	
death_rate	Death rate of the disease.	

# 18.38.2.2 ModelSEIRD() [2/2]

#### Constructor for the SEIRD model.

#### **Parameters**

vname	name Name of the model.	
prevalence	Initial prevalence of the disease.	
transmission_rate	Transmission rate of the disease.	
avg_incubation_days	Average incubation period of the disease.	
recovery_rate Recovery rate of the disease.		
death_rate	Death rate of the disease.	

# 18.38.3 Member Data Documentation

# 18.38.3.1 update\_exposed\_seir

```
template<typename TSeq = EPI_DEFAULT_TSEQ>
epiworld::UpdateFun<TSeq> ModelSEIRD< TSeq >::update_exposed_seir

Initial value:
= [](
    epiworld::Agent<TSeq> * p,
    epiworld::Model<TSeq> * m
) -> void {
    auto v = p->get_virus();

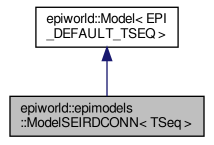
    if (m->runif() < 1.0/(v->get_incubation(m)))
        p->change_state(m, ModelSEIRD<TSeq>::INFECTED);
    return;
```

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

· include/epiworld/models/seird.hpp

# 18.39 epiworld::epimodels::ModelSEIRDCONN< TSeq > Class Template Reference

Inheritance diagram for epiworld::epimodels::ModelSEIRDCONN< TSeq >:



Collaboration diagram for epiworld::epimodels::ModelSEIRDCONN < TSeq >:



#### **Public Member Functions**

ModelSEIRDCONN (ModelSEIRDCONN < TSeq > &model, const std::string &vname, epiworld\_fast\_uint n, epiworld\_double prevalence, epiworld\_double contact\_rate, epiworld\_double transmission\_rate, epiworld\_double avg\_incubation\_days, epiworld\_double recovery\_rate, epiworld\_double death\_rate)

Template for a Susceptible-Exposed-Infected-Removed (SEIR) model.

- ModelSEIRDCONN (const std::string &vname, epiworld\_fast\_uint n, epiworld\_double prevalence, epiworld
   —double contact\_rate, epiworld\_double transmission\_rate, epiworld\_double avg\_incubation\_days, epiworld
   —double recovery\_rate, epiworld\_double death\_rate)
- ModelSEIRDCONN< TSeq > & run (epiworld\_fast\_uint ndays, int seed=-1)

Runs the simulation (after initialization)

· void reset ()

Reset the model.

Model < TSeq > \* clone\_ptr ()

Advanced usage: Makes a copy of data and returns it as undeleted pointer.

ModelSEIRDCONN< TSeq > & initial\_states (std::vector< double > proportions\_, std::vector< int > queue\_={})

Set up the initial states of the model.

• size\_t get\_n\_infected () const

#### Static Public Attributes

- static const int SUSCEPTIBLE = 0
- static const int **EXPOSED** = 1
- static const int INFECTED = 2
- static const int REMOVED = 3
- static const int **DECEASED** = 4

# **Additional Inherited Members**

# 18.39.1 Constructor & Destructor Documentation

# 18.39.1.1 ModelSEIRDCONN()

Template for a Susceptible-Exposed-Infected-Removed (SEIR) model.

#### **Parameters**

model	A Model <tseq> object where to set up the SIR.</tseq>
vname	std::string Name of the virus
prevalence	Initial prevalence (proportion)
contact_rate	Average number of contacts (interactions) per step.
transmission_rate	Probability of transmission
recovery_rate	Probability of recovery
death_rate	Probability of death

# 18.39.2 Member Function Documentation

# 18.39.2.1 clone\_ptr()

```
template<typename TSeq >
Model< TSeq > * ModelSEIRDCONN< TSeq >::clone_ptr [inline], [virtual]
```

Advanced usage: Makes a copy of data and returns it as undeleted pointer.

#### **Parameters**

сору

Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

# 18.39.2.2 initial\_states()

Set up the initial states of the model.

# **Parameters**

proportions → Double vector with the following values:
 0: Proportion of non-infected agents who are removed.
 1: Proportion of exposed agents to be set as infected.

Reimplemented from epiworld::Model < EPI DEFAULT TSEQ >.

#### 18.39.2.3 reset()

```
template<typename TSeq >
void ModelSEIRDCONN< TSeq >::reset [inline], [virtual]
```

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

Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

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

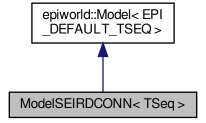
· epiworld.hpp

# 18.40 ModelSEIRDCONN< TSeq > Class Template Reference

Inheritance diagram for ModelSEIRDCONN< TSeq >:



Collaboration diagram for ModelSEIRDCONN< TSeq >:



#### **Public Member Functions**

ModelSEIRDCONN (ModelSEIRDCONN < TSeq > &model, const std::string &vname, epiworld\_fast\_uint n, epiworld\_double prevalence, epiworld\_double contact\_rate, epiworld\_double transmission\_rate, epiworld\_← double avg\_incubation\_days, epiworld\_double recovery\_rate, epiworld\_double death\_rate)

Template for a Susceptible-Exposed-Infected-Removed (SEIR) model.

- ModelSEIRDCONN (const std::string &vname, epiworld\_fast\_uint n, epiworld\_double prevalence, epiworld
   — double contact\_rate, epiworld\_double transmission\_rate, epiworld\_double avg\_incubation\_days, epiworld
   — double recovery\_rate, epiworld\_double death\_rate)
- ModelSEIRDCONN< TSeq > & run (epiworld\_fast\_uint ndays, int seed=-1)

Runs the simulation (after initialization)

· void reset ()

Reset the model.

Model < TSeq > \* clone\_ptr ()

Advanced usage: Makes a copy of data and returns it as undeleted pointer.

ModelSEIRDCONN< TSeq > & initial\_states (std::vector< double > proportions\_, std::vector< int > queue\_={})

Set up the initial states of the model.

· size t get n infected () const

# **Static Public Attributes**

- static const int SUSCEPTIBLE = 0
- static const int EXPOSED = 1
- static const int INFECTED = 2
- static const int **REMOVED** = 3
- static const int **DECEASED** = 4

# **Additional Inherited Members**

# 18.40.1 Constructor & Destructor Documentation

# 18.40.1.1 ModelSEIRDCONN()

Template for a Susceptible-Exposed-Infected-Removed (SEIR) model.

#### **Parameters**

model	A Model <tseq> object where to set up the SIR.</tseq>
vname	std::string Name of the virus
prevalence	Initial prevalence (proportion)
contact_rate	Average number of contacts (interactions) per step.
transmission_rate	Probability of transmission
recovery_rate	Probability of recovery
death_rate	Probability of death

# 18.40.2 Member Function Documentation

#### 18.40.2.1 clone\_ptr()

```
template<typename TSeq >
Model< TSeq > * ModelSEIRDCONN< TSeq >::clone_ptr [inline], [virtual]
```

Advanced usage: Makes a copy of data and returns it as undeleted pointer.

#### **Parameters**

сору

Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

# 18.40.2.2 initial\_states()

Set up the initial states of the model.

# **Parameters**

proportions←	Double vector with the following values:
_	0: Proportion of non-infected agents who are removed.
	• 1: Proportion of exposed agents to be set as infected.

 $\label{lem:lemented_lemented} \textbf{Reimplemented from epiworld::} \\ \textbf{Model} < \textbf{EPI\_DEFAULT\_TSEQ} >. \\$ 

#### 18.40.2.3 reset()

```
template<typename TSeq >
void ModelSEIRDCONN< TSeq >::reset [inline], [virtual]
```

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

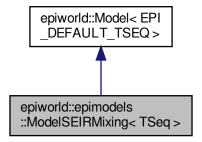
Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

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

• include/epiworld/models/seirdconnected.hpp

# 18.41 epiworld::epimodels::ModelSEIRMixing< TSeq > Class Template Reference

 $Inheritance\ diagram\ for\ epiworld::epimodels::ModelSEIRMixing < TSeq >:$ 



Collaboration diagram for epiworld::epimodels::ModelSEIRMixing< TSeq >:



#### **Public Member Functions**

ModelSEIRMixing (ModelSEIRMixing < TSeq > &model, const std::string &vname, epiworld\_fast\_uint n, epiworld\_double prevalence, epiworld\_double contact\_rate, epiworld\_double transmission\_rate, epiworld—double avg\_incubation\_days, epiworld\_double recovery\_rate, std::vector< double > contact\_matrix)

Constructs a ModelSEIRMixing object.

ModelSEIRMixing (const std::string &vname, epiworld\_fast\_uint n, epiworld\_double prevalence, epiworld\_
 double contact\_rate, epiworld\_double transmission\_rate, epiworld\_double avg\_incubation\_days, epiworld
 double recovery\_rate, std::vector< double > contact\_matrix)

Constructs a ModelSEIRMixing object.

ModelSEIRMixing < TSeq > & run (epiworld\_fast\_uint ndays, int seed=-1)

Runs the simulation (after initialization)

• void reset ()

Reset the model.

Model < TSeq > \* clone\_ptr ()

Advanced usage: Makes a copy of data and returns it as undeleted pointer.

ModelSEIRMixing< TSeq > & initial\_states (std::vector< double > proportions\_, std::vector< int > queue ← \_={})

Set the initial states of the model.

void set\_contact\_matrix (std::vector< double > cmat)

#### **Static Public Attributes**

- static const int SUSCEPTIBLE = 0
- static const int **EXPOSED** = 1
- static const int INFECTED = 2
- static const int **RECOVERED** = 3

# **Additional Inherited Members**

# 18.41.1 Constructor & Destructor Documentation

# 18.41.1.1 ModelSEIRMixing() [1/2]

Constructs a ModelSEIRMixing object.

Template for a Susceptible-Exposed-Infected-Removed (SEIR) model.

#### **Parameters**

model	A reference to an existing ModelSEIRMixing object.
vname	The name of the ModelSEIRMixing object.
п	The number of entities in the model.
prevalence	The initial prevalence of the disease in the model.
contact_rate	The contact rate between entities in the model.
transmission_rate	The transmission rate of the disease in the model.
avg_incubation_days	The average incubation period of the disease in the model.
recovery_rate	The recovery rate of the disease in the model.
contact_matrix	The contact matrix between entities in the model. Specified in column-major order.
model	A Model <tseq> object where to set up the SIR.</tseq>
vname	std::string Name of the virus
prevalence	Initial prevalence (proportion)
contact_rate	Average number of contacts (interactions) per step.
transmission_rate	Probability of transmission
recovery_rate	Probability of recovery

# 18.41.1.2 ModelSEIRMixing() [2/2]

Constructs a ModelSEIRMixing object.

#### **Parameters**

vname	The name of the ModelSEIRMixing object.
n	The number of entities in the model.
prevalence	The initial prevalence of the disease in the model.
contact_rate	The contact rate between entities in the model.
transmission_rate	The transmission rate of the disease in the model.
avg_incubation_days	The average incubation period of the disease in the model.
recovery_rate	The recovery rate of the disease in the model.
contact_matrix	The contact matrix between entities in the model.

## 18.41.2 Member Function Documentation

## 18.41.2.1 clone\_ptr()

```
template<typename TSeq >
Model< TSeq > * ModelSEIRMixing< TSeq >::clone_ptr [inline], [virtual]
```

Advanced usage: Makes a copy of data and returns it as undeleted pointer.

#### **Parameters**

сору

Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

## 18.41.2.2 initial\_states()

Set the initial states of the model.

## **Parameters**

proportions↔	Double vector with a single element:
_	The proportion of non-infected individuals who have recovered.

Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

#### 18.41.2.3 reset()

template<typename TSeq >
void ModelSEIRMixing< TSeq >::reset [inline], [virtual]

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

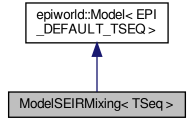
Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

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

· epiworld.hpp

## 18.42 ModelSEIRMixing < TSeq > Class Template Reference

Inheritance diagram for ModelSEIRMixing< TSeq >:



Collaboration diagram for ModelSEIRMixing < TSeq >:



#### **Public Member Functions**

ModelSEIRMixing (ModelSEIRMixing < TSeq > &model, const std::string &vname, epiworld\_fast\_uint n, epiworld\_double prevalence, epiworld\_double contact\_rate, epiworld\_double transmission\_rate, epiworld—double avg\_incubation\_days, epiworld\_double recovery\_rate, std::vector < double > contact\_matrix)

Constructs a ModelSEIRMixing object.

ModelSEIRMixing (const std::string &vname, epiworld\_fast\_uint n, epiworld\_double prevalence, epiworld\_
 double contact\_rate, epiworld\_double transmission\_rate, epiworld\_double avg\_incubation\_days, epiworld
 double recovery\_rate, std::vector< double > contact\_matrix)

Constructs a ModelSEIRMixing object.

ModelSEIRMixing< TSeq > & run (epiworld\_fast\_uint ndays, int seed=-1)

Runs the simulation (after initialization)

· void reset ()

Reset the model.

Model < TSeq > \* clone\_ptr ()

Advanced usage: Makes a copy of data and returns it as undeleted pointer.

ModelSEIRMixing < TSeq > & initial\_states (std::vector < double > proportions\_, std::vector < int > queue ← \_={})

Set the initial states of the model.

void set\_contact\_matrix (std::vector< double > cmat)

#### Static Public Attributes

- static const int SUSCEPTIBLE = 0
- static const int EXPOSED = 1
- static const int INFECTED = 2
- static const int **RECOVERED** = 3

## **Additional Inherited Members**

#### 18.42.1 Constructor & Destructor Documentation

## 18.42.1.1 ModelSEIRMixing() [1/2]

Constructs a ModelSEIRMixing object.

Template for a Susceptible-Exposed-Infected-Removed (SEIR) model.

#### **Parameters**

model	A reference to an existing ModelSEIRMixing object.
vname	The name of the ModelSEIRMixing object.
п	The number of entities in the model.
prevalence	The initial prevalence of the disease in the model.
contact_rate	The contact rate between entities in the model.
transmission_rate	The transmission rate of the disease in the model.
avg_incubation_days	The average incubation period of the disease in the model.
recovery_rate	The recovery rate of the disease in the model.
contact_matrix	The contact matrix between entities in the model. Specified in column-major order.
model	A Model <tseq> object where to set up the SIR.</tseq>
vname	std::string Name of the virus
prevalence	Initial prevalence (proportion)
contact_rate	Average number of contacts (interactions) per step.
transmission_rate	Probability of transmission
recovery_rate	Probability of recovery

## 18.42.1.2 ModelSEIRMixing() [2/2]

## Constructs a ModelSEIRMixing object.

## **Parameters**

vname	The name of the ModelSEIRMixing object.
n	The number of entities in the model.
prevalence	The initial prevalence of the disease in the model.
contact_rate	The contact rate between entities in the model.
transmission_rate	The transmission rate of the disease in the model.
avg_incubation_days	The average incubation period of the disease in the model.
recovery_rate	The recovery rate of the disease in the model.
contact_matrix	The contact matrix between entities in the model.

## 18.42.2 Member Function Documentation

#### 18.42.2.1 clone\_ptr()

```
template<typename TSeq >
Model< TSeq > * ModelSEIRMixing< TSeq >::clone_ptr [inline], [virtual]
```

Advanced usage: Makes a copy of data and returns it as undeleted pointer.

#### **Parameters**

```
сору
```

Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

## 18.42.2.2 initial\_states()

Set the initial states of the model.

#### **Parameters**

proportions⊷	Double vector with a single element:
_	The proportion of non-infected individuals who have recovered.

 $\label{eq:control_loss} \mbox{Reimplemented from epiworld::} \mbox{Model} < \mbox{EPI\_DEFAULT\_TSEQ} >.$ 

#### 18.42.2.3 reset()

```
template<typename TSeq >
void ModelSEIRMixing< TSeq >::reset [inline], [virtual]
```

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

Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

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

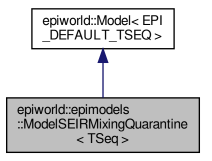
include/epiworld/models/seirmixing.hpp

# 18.43 epiworld::epimodels::ModelSEIRMixingQuarantine< TSeq > Class Template Reference

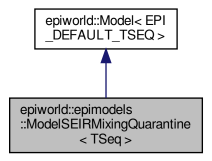
SEIR model with mixing, quarantine, and contact tracing.

#include <epiworld.hpp>

Inheritance diagram for epiworld::epimodels::ModelSEIRMixingQuarantine< TSeg >:



Collaboration diagram for epiworld::epimodels::ModelSEIRMixingQuarantine < TSeq >:



## **Public Member Functions**

ModelSEIRMixingQuarantine (ModelSEIRMixingQuarantine < TSeq > &model, const std::string &vname, epiworld\_fast\_uint n, epiworld\_double prevalence, epiworld\_double contact\_rate, epiworld\_double transmission\_rate, epiworld\_double avg\_incubation\_days, epiworld\_double recovery\_rate, std::vector < double > contact\_matrix, epiworld\_double hospitalization\_rate, epiworld\_double hospitalization\_period, epiworld\_double days\_undetected, epiworld\_fast\_int quarantine\_period, epiworld\_double quarantine = \_\_willingness, epiworld\_double isolation\_willingness, epiworld\_fast\_int isolation\_period, epiworld\_double contact\_tracing\_success\_rate=1.0, epiworld\_fast\_uint contact\_tracing\_days\_prior=4u)

Constructs a ModelSEIRMixingQuarantine object.

ModelSEIRMixingQuarantine (const std::string &vname, epiworld\_fast\_uint n, epiworld\_double prevalence, epiworld\_double contact\_rate, epiworld\_double transmission\_rate, epiworld\_double avg\_incubation\_days, epiworld\_double recovery\_rate, std::vector< double > contact\_matrix, epiworld\_double hospitalization\_rate, epiworld\_double hospitalization\_period, epiworld\_double days\_undetected, epiworld\_fast\_int quarantine — \_\_period, epiworld\_double quarantine\_willingness, epiworld\_double isolation\_willingness, epiworld\_fast\_int isolation\_period, epiworld\_double contact\_tracing\_success\_rate=1.0, epiworld\_fast\_uint contact\_tracing\_cdays prior=4u)

Constructs a ModelSEIRMixingQuarantine object.

ModelSEIRMixingQuarantine< TSeq > & run (epiworld\_fast\_uint ndays, int seed=-1)

Run the model simulation.

· void reset ()

Reset the model to initial state.

Model < TSeq > \* clone\_ptr ()

Create a clone of this model.

ModelSEIRMixingQuarantine < TSeq > & initial\_states (std::vector < double > proportions\_, std::vector < int > queue ={})

Set the initial states of the model.

void set contact matrix (std::vector< double > cmat)

Set the contact matrix for population mixing.

std::vector< double > get\_contact\_matrix () const

Get the current contact matrix.

• std::vector< size\_t > get\_agent\_quarantine\_triggered () const

Get the quarantine trigger status for all agents.

std::vector< bool > get\_quarantine\_willingness () const

Get the quarantine willingness for all agents.

• std::vector< bool > get isolation willingness () const

Get the isolation willingness for all agents.

#### **Static Public Attributes**

- static const int SUSCEPTIBLE = 0
- static const int EXPOSED = 1
- static const int **INFECTED** = 2
- static const int **ISOLATED** = 3
- static const int **DETECTED\_HOSPITALIZED** = 4
- static const int QUARANTINED\_SUSCEPTIBLE = 5
- static const int QUARANTINED\_EXPOSED = 6
- static const int ISOLATED\_RECOVERED = 7
- static const int **HOSPITALIZED** = 8
- static const int **RECOVERED** = 9
- static const size\_t QUARANTINE\_PROCESS\_INACTIVE = 0u
- static const size\_t QUARANTINE\_PROCESS\_ACTIVE = 1u
- static const size\_t **QUARANTINE\_PROCESS\_DONE** = 2u

## **Additional Inherited Members**

## 18.43.1 Detailed Description

template<typename TSeq = EPI\_DEFAULT\_TSEQ>
class epiworld::epimodels::ModelSEIRMixingQuarantine< TSeq>

SEIR model with mixing, quarantine, and contact tracing.

This class implements a Susceptible-Exposed-Infected-Removed (SEIR) epidemiological model with additional features including:

- · Population mixing based on contact matrices
- · Quarantine measures for exposed contacts
- · Isolation policies for detected infected individuals
- · Contact tracing with configurable success rates
- · Hospitalization of severe cases
- · Individual willingness to comply with public health measures

The model supports 10 distinct states:

- · Susceptible: Individuals who can become infected
- Exposed: Infected but not yet infectious (incubation period)
- · Infected: Infectious individuals in the community
- Isolated: Detected infected individuals in self-isolation
- Detected Hospitalized: Hospitalized individuals who were contact-traced
- · Quarantined Susceptible: Susceptible individuals in guarantine due to contact tracing
- · Quarantined Exposed: Exposed individuals in quarantine due to contact tracing
- · Isolated Recovered: Recovered individuals still in isolation
- · Hospitalized: Individuals requiring hospital care
- · Recovered: Individuals who have recovered and gained immunity

## **Template Parameters**

TSeq Type for genetic sequences (default: EPI\_DEFAULT\_TSEQ)

## 18.43.2 Constructor & Destructor Documentation

## 18.43.2.1 ModelSEIRMixingQuarantine() [1/2]

```
template<typename TSeq >
{\tt ModelSEIRMixingQuarantine<\ TSeq\ >::} ModelSEIRMixingQuarantine\ (
             ModelSEIRMixingQuarantine< TSeq > & model,
             const std::string & vname,
             epiworld_fast_uint n,
             epiworld_double prevalence,
             epiworld_double contact_rate,
             epiworld_double transmission_rate,
             epiworld_double avg_incubation_days,
             epiworld_double recovery_rate,
             std::vector< double > contact_matrix,
             epiworld_double hospitalization_rate,
             epiworld_double hospitalization_period,
             epiworld_double days_undetected,
             epiworld_fast_int quarantine_period,
             epiworld_double quarantine_willingness,
             epiworld_double isolation_willingness,
             epiworld_fast_int isolation_period,
             epiworld_double contact_tracing_success_rate = 1.0,
             epiworld_fast_uint contact_tracing_days_prior = 4u ) [inline]
```

Constructs a ModelSEIRMixingQuarantine object.

Template for a Susceptible-Exposed-Infected-Removed (SEIR) model with mixing, quarantine, and contact tracing.

#### **Parameters**

madal	A reference to an existing MadelCEIDMixingOverenting chiest
model	A reference to an existing ModelSEIRMixingQuarantine object.
vname	The name of the ModelSEIRMixingQuarantine object.
n	The number of entities in the model.
prevalence	The initial prevalence of the disease in the model.
contact_rate	The contact rate between entities in the model.
transmission_rate	The transmission rate of the disease in the model.
avg_incubation_days	The average incubation period of the disease in the model.
recovery_rate	The recovery rate of the disease in the model.
contact_matrix	The contact matrix between entities in the model. Specified in column-major order.
hospitalization_rate	The rate at which infected individuals are hospitalized.
hospitalization_period	The average duration of hospitalization in days.
days_undetected	The average number of days an infected individual remains undetected.
quarantine_period	The duration of quarantine in days for exposed contacts.
quarantine_willingness	The proportion of individuals willing to comply with quarantine measures.
isolation_willingness	The proportion of individuals willing to self-isolate when detected.
isolation_period	The duration of isolation in days for detected infected individuals.
contact_tracing_success_rate	The probability of successfully identifying and tracing contacts (default: 1.0).
contact_tracing_days_prior	The number of days prior to detection for which contacts are traced (default: 4).
model	A ModelSEIRMixingQuarantine <tseq> object where to set up the SEIR model.</tseq>
vname	Name of the virus
n	Number of agents in the population
prevalence	Initial prevalence (proportion of infected individuals)

#### **Parameters**

contact_rate	Average number of contacts (interactions) per step
transmission_rate	Probability of transmission per contact
avg_incubation_days	Average incubation period in days
recovery_rate	Probability of recovery per day
contact_matrix	Contact matrix specifying mixing patterns between population groups
hospitalization_rate	Rate at which infected individuals are hospitalized
hospitalization_period	Average duration of hospitalization in days
days_undetected	Average number of days an infected individual remains undetected
quarantine_period	Duration of quarantine in days for exposed contacts
quarantine_willingness	Proportion of individuals willing to comply with quarantine
isolation_willingness	Proportion of individuals willing to self-isolate when detected
isolation_period	Duration of isolation in days for detected infected individuals
contact_tracing_success_rate	Probability of successfully identifying contacts during tracing
contact_tracing_days_prior	Number of days prior to detection for contact tracing

## 18.43.2.2 ModelSEIRMixingQuarantine() [2/2]

```
template<typename TSeq >
{\tt ModelSEIRMixingQuarantine<\ TSeq\ >::} ModelSEIRMixingQuarantine\ (
            const std::string & vname,
             epiworld_fast_uint n,
             epiworld_double prevalence,
             epiworld_double contact_rate,
             epiworld_double transmission_rate,
             epiworld_double avg_incubation_days,
             epiworld_double recovery_rate,
             std::vector< double > contact_matrix,
             epiworld_double hospitalization_rate,
             epiworld_double hospitalization_period,
             epiworld_double days_undetected,
             epiworld_fast_int quarantine_period,
             epiworld_double quarantine_willingness,
             epiworld_double isolation_willingness,
             epiworld_fast_int isolation_period,
             epiworld_double contact_tracing_success_rate = 1.0,
             epiworld_fast_uint contact_tracing_days_prior = 4u ) [inline]
```

## Constructs a ModelSEIRMixingQuarantine object.

#### **Parameters**

vname	The name of the ModelSEIRMixingQuarantine object.
п	The number of entities in the model.
prevalence	The initial prevalence of the disease in the model.
contact_rate	The contact rate between entities in the model.
transmission_rate	The transmission rate of the disease in the model.
avg_incubation_days	The average incubation period of the disease in the model.

#### **Parameters**

recovery_rate	The recovery rate of the disease in the model.
contact_matrix	The contact matrix between entities in the model.
hospitalization_rate	The rate at which infected individuals are hospitalized.
hospitalization_period	The average duration of hospitalization in days.
days_undetected	The average number of days an infected individual remains undetected.
quarantine_period	The duration of quarantine in days for exposed contacts.
quarantine_willingness	The proportion of individuals willing to comply with quarantine measures.
isolation_willingness	The proportion of individuals willing to self-isolate when detected.
isolation_period	The duration of isolation in days for detected infected individuals.
contact_tracing_success_rate	The probability of successfully identifying and tracing contacts (default: 1.0).
contact_tracing_days_prior	The number of days prior to detection for which contacts are traced (default: 4).

## 18.43.3 Member Function Documentation

## 18.43.3.1 clone\_ptr()

```
template<typename TSeq >
Model< TSeq > * ModelSEIRMixingQuarantine< TSeq >::clone_ptr [inline], [virtual]
```

Create a clone of this model.

## Returns

Pointer to a new model instance with the same configuration

Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

## 18.43.3.2 get\_agent\_quarantine\_triggered()

```
template<typename TSeq = EPI_DEFAULT_TSEQ>
std::vector< size_t > epiworld::epimodels::ModelSEIRMixingQuarantine< TSeq >::get_agent_
quarantine_triggered ( ) const [inline]
```

Get the quarantine trigger status for all agents.

## Returns

Vector indicating quarantine process status for each agent

#### 18.43.3.3 get\_contact\_matrix()

```
template<typename TSeq = EPI_DEFAULT_TSEQ>
std::vector< double > epiworld::epimodels::ModelSEIRMixingQuarantine< TSeq >::get_contact_
matrix ( ) const [inline]
```

Get the current contact matrix.

#### Returns

Vector representing the contact matrix

#### 18.43.3.4 get\_isolation\_willingness()

```
template<typename TSeq = EPI_DEFAULT_TSEQ>
std::vector< bool > epiworld::epimodels::ModelSEIRMixingQuarantine< TSeq >::get_isolation_
willingness ( ) const [inline]
```

Get the isolation willingness for all agents.

#### Returns

Vector of boolean values indicating each agent's willingness to self-isolate

## 18.43.3.5 get\_quarantine\_willingness()

```
template<typename TSeq = EPI_DEFAULT_TSEQ>
std::vector< bool > epiworld::epimodels::ModelSEIRMixingQuarantine< TSeq >::get_quarantine_
willingness ( ) const [inline]
```

Get the quarantine willingness for all agents.

#### Returns

Vector of boolean values indicating each agent's willingness to quarantine

#### 18.43.3.6 initial\_states()

Set the initial states of the model.

#### **Parameters**

$proportions \leftarrow$	Double vector with two elements:	
_	• [0]: The proportion of initially infected individuals who start in the exposed state.	
	• [1]: The proportion of initially non-infected individuals who have recovered (immune).	
queue_	Optional vector for queuing specifications (default: empty).	

Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

#### 18.43.3.7 run()

Run the model simulation.

#### **Parameters**

ndays	Number of days to simulate
seed	Random seed for reproducibility (default: -1 for random seed)

#### Returns

Reference to this model instance

 $\label{eq:local_problem} \mbox{Reimplemented from epiworld::} \mbox{Model} < \mbox{EPI\_DEFAULT\_TSEQ} >.$ 

## 18.43.3.8 set\_contact\_matrix()

Set the contact matrix for population mixing.

#### **Parameters**

_		
	cmat	Contact matrix specifying interaction rates between groups

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

epiworld.hpp

## 18.44 ModelSEIRMixingQuarantine < TSeq > Class Template Reference

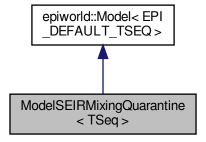
SEIR model with mixing, quarantine, and contact tracing.

#include <seirmixingquarantine.hpp>

Inheritance diagram for ModelSEIRMixingQuarantine< TSeq >:



Collaboration diagram for ModelSEIRMixingQuarantine < TSeq >:



#### **Public Member Functions**

ModelSEIRMixingQuarantine (ModelSEIRMixingQuarantine < TSeq > &model, const std::string &vname, epiworld\_fast\_uint n, epiworld\_double prevalence, epiworld\_double contact\_rate, epiworld\_double transmission\_rate, epiworld\_double avg\_incubation\_days, epiworld\_double recovery\_rate, std::vector < double > contact\_matrix, epiworld\_double hospitalization\_rate, epiworld\_double hospitalization\_period, epiworld\_double days\_undetected, epiworld\_fast\_int quarantine\_period, epiworld\_double quarantine = \_\_willingness, epiworld\_double isolation\_willingness, epiworld\_fast\_int isolation\_period, epiworld\_double contact\_tracing\_success\_rate=1.0, epiworld\_fast\_uint contact\_tracing\_days\_prior=4u)

Constructs a ModelSEIRMixingQuarantine object.

ModelSEIRMixingQuarantine (const std::string &vname, epiworld\_fast\_uint n, epiworld\_double prevalence, epiworld\_double contact\_rate, epiworld\_double transmission\_rate, epiworld\_double avg\_incubation\_days, epiworld\_double recovery\_rate, std::vector< double > contact\_matrix, epiworld\_double hospitalization\_rate, epiworld\_double hospitalization\_period, epiworld\_double days\_undetected, epiworld\_fast\_int quarantine — \_\_period, epiworld\_double quarantine\_willingness, epiworld\_double isolation\_willingness, epiworld\_fast\_int isolation\_period, epiworld\_double contact\_tracing\_success\_rate=1.0, epiworld\_fast\_uint contact\_tracing\_days\_prior=4u)

Constructs a ModelSEIRMixingQuarantine object.

• ModelSEIRMixingQuarantine < TSeq > & run (epiworld fast uint ndays, int seed=-1)

Run the model simulation.

· void reset ()

Reset the model to initial state.

Model < TSeq > \* clone\_ptr ()

Create a clone of this model.

ModelSEIRMixingQuarantine< TSeq > & initial\_states (std::vector< double > proportions\_, std::vector< int > queue\_={})

Set the initial states of the model.

void set contact matrix (std::vector< double > cmat)

Set the contact matrix for population mixing.

std::vector< double > get contact matrix () const

Get the current contact matrix.

std::vector< size\_t > get\_agent\_quarantine\_triggered () const

Get the quarantine trigger status for all agents.

std::vector< bool > get\_quarantine\_willingness () const

Get the quarantine willingness for all agents.

std::vector< bool > get\_isolation\_willingness () const

Get the isolation willingness for all agents.

#### **Static Public Attributes**

- static const int SUSCEPTIBLE = 0
- static const int EXPOSED = 1
- static const int INFECTED = 2
- static const int ISOLATED = 3
- static const int **DETECTED HOSPITALIZED** = 4
- static const int QUARANTINED\_SUSCEPTIBLE = 5
- static const int QUARANTINED\_EXPOSED = 6
- static const int ISOLATED\_RECOVERED = 7
- static const int **HOSPITALIZED** = 8
- static const int **RECOVERED** = 9
- static const size\_t QUARANTINE\_PROCESS\_INACTIVE = 0u
- static const size t QUARANTINE PROCESS ACTIVE = 1u
- static const size\_t QUARANTINE\_PROCESS\_DONE = 2u

## **Additional Inherited Members**

## 18.44.1 Detailed Description

template<typename TSeq = EPI\_DEFAULT\_TSEQ> class ModelSEIRMixingQuarantine< TSeq >

SEIR model with mixing, quarantine, and contact tracing.

This class implements a Susceptible-Exposed-Infected-Removed (SEIR) epidemiological model with additional features including:

- · Population mixing based on contact matrices
- · Quarantine measures for exposed contacts
- · Isolation policies for detected infected individuals
- · Contact tracing with configurable success rates
- · Hospitalization of severe cases
- · Individual willingness to comply with public health measures

The model supports 10 distinct states:

- · Susceptible: Individuals who can become infected
- Exposed: Infected but not yet infectious (incubation period)
- · Infected: Infectious individuals in the community
- Isolated: Detected infected individuals in self-isolation
- Detected Hospitalized: Hospitalized individuals who were contact-traced
- · Quarantined Susceptible: Susceptible individuals in quarantine due to contact tracing
- · Quarantined Exposed: Exposed individuals in quarantine due to contact tracing
- · Isolated Recovered: Recovered individuals still in isolation
- · Hospitalized: Individuals requiring hospital care
- · Recovered: Individuals who have recovered and gained immunity

## **Template Parameters**

TSeq Type for genetic sequences (default: EPI\_DEFAULT\_TSEQ)

## 18.44.2 Constructor & Destructor Documentation

## 18.44.2.1 ModelSEIRMixingQuarantine() [1/2]

```
template<typename TSeq >
{\tt ModelSEIRMixingQuarantine<\ TSeq\ >::} ModelSEIRMixingQuarantine\ (
             ModelSEIRMixingQuarantine< TSeq > & model,
             const std::string & vname,
             epiworld_fast_uint n,
             epiworld_double prevalence,
             epiworld_double contact_rate,
             epiworld_double transmission_rate,
             epiworld_double avg_incubation_days,
             epiworld_double recovery_rate,
             std::vector< double > contact_matrix,
             epiworld_double hospitalization_rate,
             epiworld_double hospitalization_period,
             epiworld_double days_undetected,
             epiworld_fast_int quarantine_period,
             epiworld_double quarantine_willingness,
             epiworld_double isolation_willingness,
             epiworld_fast_int isolation_period,
             epiworld_double contact_tracing_success_rate = 1.0,
             epiworld_fast_uint contact_tracing_days_prior = 4u ) [inline]
```

Constructs a ModelSEIRMixingQuarantine object.

Template for a Susceptible-Exposed-Infected-Removed (SEIR) model with mixing, quarantine, and contact tracing.

#### **Parameters**

model	A reference to an existing ModelSEIRMixingQuarantine object.
vname	The name of the ModelSEIRMixingQuarantine object.
n	The number of entities in the model.
prevalence	The initial prevalence of the disease in the model.
contact_rate	The contact rate between entities in the model.
transmission_rate	The transmission rate of the disease in the model.
avg_incubation_days	The average incubation period of the disease in the model.
recovery_rate	The recovery rate of the disease in the model.
contact_matrix	The contact matrix between entities in the model. Specified in column-major order.
hospitalization_rate	The rate at which infected individuals are hospitalized.
hospitalization_period	The average duration of hospitalization in days.
days_undetected	The average number of days an infected individual remains undetected.
quarantine_period	The duration of quarantine in days for exposed contacts.
quarantine_willingness	The proportion of individuals willing to comply with quarantine measures.
isolation_willingness	The proportion of individuals willing to self-isolate when detected.
isolation_period	The duration of isolation in days for detected infected individuals.
contact_tracing_success_rate	The probability of successfully identifying and tracing contacts (default: 1.0).
contact_tracing_days_prior	The number of days prior to detection for which contacts are traced (default: 4).
model	A ModelSEIRMixingQuarantine <tseq> object where to set up the SEIR model.</tseq>
vname	Name of the virus
n	Number of agents in the population
prevalence	Initial prevalence (proportion of infected individuals)

#### **Parameters**

contact_rate	Average number of contacts (interactions) per step
transmission_rate	Probability of transmission per contact
avg_incubation_days	Average incubation period in days
recovery_rate	Probability of recovery per day
contact_matrix	Contact matrix specifying mixing patterns between population groups
hospitalization_rate	Rate at which infected individuals are hospitalized
hospitalization_period	Average duration of hospitalization in days
days_undetected	Average number of days an infected individual remains undetected
quarantine_period	Duration of quarantine in days for exposed contacts
quarantine_willingness	Proportion of individuals willing to comply with quarantine
isolation_willingness	Proportion of individuals willing to self-isolate when detected
isolation_period	Duration of isolation in days for detected infected individuals
contact_tracing_success_rate	Probability of successfully identifying contacts during tracing
contact_tracing_days_prior	Number of days prior to detection for contact tracing

## 18.44.2.2 ModelSEIRMixingQuarantine() [2/2]

```
template<typename TSeq >
{\tt ModelSEIRMixingQuarantine<\ TSeq\ >::} ModelSEIRMixingQuarantine\ (
            const std::string & vname,
             epiworld_fast_uint n,
             epiworld_double prevalence,
             epiworld_double contact_rate,
             epiworld_double transmission_rate,
             epiworld_double avg_incubation_days,
             epiworld_double recovery_rate,
             std::vector< double > contact_matrix,
             epiworld_double hospitalization_rate,
             epiworld_double hospitalization_period,
             epiworld_double days_undetected,
             epiworld_fast_int quarantine_period,
             epiworld_double quarantine_willingness,
             epiworld_double isolation_willingness,
             epiworld_fast_int isolation_period,
             epiworld_double contact_tracing_success_rate = 1.0,
             epiworld_fast_uint contact_tracing_days_prior = 4u ) [inline]
```

## Constructs a ModelSEIRMixingQuarantine object.

#### **Parameters**

vname	The name of the ModelSEIRMixingQuarantine object.
п	The number of entities in the model.
prevalence	The initial prevalence of the disease in the model.
contact_rate	The contact rate between entities in the model.
transmission_rate	The transmission rate of the disease in the model.
avg_incubation_days	The average incubation period of the disease in the model.

#### **Parameters**

recovery_rate	The recovery rate of the disease in the model.
contact_matrix	The contact matrix between entities in the model.
hospitalization_rate	The rate at which infected individuals are hospitalized.
hospitalization_period	The average duration of hospitalization in days.
days_undetected	The average number of days an infected individual remains undetected.
quarantine_period	The duration of quarantine in days for exposed contacts.
quarantine_willingness	The proportion of individuals willing to comply with quarantine measures.
isolation_willingness	The proportion of individuals willing to self-isolate when detected.
isolation_period	The duration of isolation in days for detected infected individuals.
contact_tracing_success_rate	The probability of successfully identifying and tracing contacts (default: 1.0).
contact_tracing_days_prior	The number of days prior to detection for which contacts are traced (default: 4).

## 18.44.3 Member Function Documentation

## 18.44.3.1 clone\_ptr()

```
template<typename TSeq >
Model< TSeq > * ModelSEIRMixingQuarantine< TSeq >::clone_ptr [inline], [virtual]
```

Create a clone of this model.

## Returns

Pointer to a new model instance with the same configuration

Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

## 18.44.3.2 get\_agent\_quarantine\_triggered()

```
template<typename TSeq = EPI_DEFAULT_TSEQ>
std::vector< size_t > ModelSEIRMixingQuarantine< TSeq >::get_agent_quarantine_triggered ( )
const [inline]
```

Get the quarantine trigger status for all agents.

## Returns

Vector indicating quarantine process status for each agent

## 18.44.3.3 get\_contact\_matrix()

```
template<typename TSeq = EPI_DEFAULT_TSEQ>
std::vector< double > ModelSEIRMixingQuarantine< TSeq >::get_contact_matrix ( ) const [inline]
```

Get the current contact matrix.

#### Returns

Vector representing the contact matrix

## 18.44.3.4 get\_isolation\_willingness()

```
template<typename TSeq = EPI_DEFAULT_TSEQ>
std::vector< bool > ModelSEIRMixingQuarantine< TSeq >::get_isolation_willingness ( ) const
[inline]
```

Get the isolation willingness for all agents.

#### Returns

Vector of boolean values indicating each agent's willingness to self-isolate

## 18.44.3.5 get\_quarantine\_willingness()

```
template<typename TSeq = EPI_DEFAULT_TSEQ>
std::vector< bool > ModelSEIRMixingQuarantine< TSeq >::get_quarantine_willingness ( ) const
[inline]
```

Get the quarantine willingness for all agents.

## Returns

Vector of boolean values indicating each agent's willingness to quarantine

#### 18.44.3.6 initial\_states()

Set the initial states of the model.

#### **Parameters**

proportions↔	Double vector with two elements:
_	• [0]: The proportion of initially infected individuals who start in the exposed state.
	• [1]: The proportion of initially non-infected individuals who have recovered (immune).
queue_	Optional vector for queuing specifications (default: empty).

Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

#### 18.44.3.7 run()

Run the model simulation.

#### **Parameters**

ndays	Number of days to simulate
seed	Random seed for reproducibility (default: -1 for random seed)

#### Returns

Reference to this model instance

 $\label{eq:control_equation} \mbox{Reimplemented from epiworld::} \mbox{Model} < \mbox{EPI\_DEFAULT\_TSEQ} >.$ 

## 18.44.3.8 set\_contact\_matrix()

Set the contact matrix for population mixing.

#### **Parameters**

_		
	cmat	Contact matrix specifying interaction rates between groups

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

include/epiworld/models/seirmixingquarantine.hpp

# 18.45 epiworld::epimodels::ModelSIR< TSeq > Class Template Reference

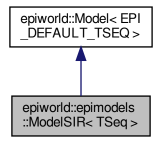
Template for a Susceptible-Infected-Removed (SIR) model.

#include <epiworld.hpp>

Inheritance diagram for epiworld::epimodels::ModelSIR < TSeq >:



Collaboration diagram for epiworld::epimodels::ModelSIR< TSeq >:



## **Public Member Functions**

- ModelSIR (ModelSIR < TSeq > &model, const std::string &vname, epiworld\_double prevalence, epiworld
   —double transmission\_rate, epiworld\_double recovery\_rate)
- **ModelSIR** (const std::string &vname, epiworld\_double prevalence, epiworld\_double transmission\_rate, epiworld\_double recovery\_rate)
- ModelSIR< TSeq > & initial\_states (std::vector< double > proportions\_, std::vector< int > queue\_={})

  Set the initial states of the model.

## **Additional Inherited Members**

## 18.45.1 Detailed Description

```
template<typename TSeq = EPI_DEFAULT_TSEQ> class epiworld::epimodels::ModelSIR< TSeq >
```

Template for a Susceptible-Infected-Removed (SIR) model.

#### **Parameters**

model	A Model <tseq> object where to set up the SIR.</tseq>
vname	std::string Name of the virus
initial_prevalence	epiworld_double Initial prevalence
initial_efficacy	epiworld_double Initial susceptibility_reduction of the immune system
initial_recovery	epiworld_double Initial recovery_rate rate of the immune system

## 18.45.2 Member Function Documentation

## 18.45.2.1 initial\_states()

Set the initial states of the model.

#### **Parameters**

proportions↔	Double vector with a single element:
_	The proportion of non-infected individuals who have recovered.

Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

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

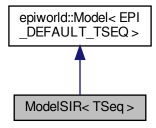
• epiworld.hpp

## 18.46 ModelSIR < TSeq > Class Template Reference

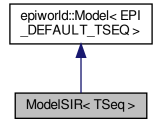
Template for a Susceptible-Infected-Removed (SIR) model.

```
#include <sir.hpp>
```

Inheritance diagram for ModelSIR < TSeq >:



 $\label{local_continuity} \mbox{Collaboration diagram for ModelSIR} < \mbox{TSeq} >:$ 



## **Public Member Functions**

- ModelSIR (ModelSIR < TSeq > &model, const std::string &vname, epiworld\_double prevalence, epiworld
   —double transmission\_rate, epiworld\_double recovery\_rate)
- **ModelSIR** (const std::string &vname, epiworld\_double prevalence, epiworld\_double transmission\_rate, epiworld\_double recovery\_rate)
- ModelSIR < TSeq > & initial\_states (std::vector < double > proportions\_, std::vector < int > queue\_={})
   Set the initial states of the model.

## **Additional Inherited Members**

## 18.46.1 Detailed Description

template<typename TSeq = EPI\_DEFAULT\_TSEQ> class ModelSIR< TSeq >

Template for a Susceptible-Infected-Removed (SIR) model.

#### **Parameters**

model	A Model <tseq> object where to set up the SIR.</tseq>
vname	std::string Name of the virus
initial_prevalence	epiworld_double Initial prevalence
initial_efficacy	epiworld_double Initial susceptibility_reduction of the immune system
initial_recovery	epiworld_double Initial recovery_rate rate of the immune system

## 18.46.2 Member Function Documentation

## 18.46.2.1 initial\_states()

Set the initial states of the model.

#### **Parameters**

proportions⊷	Double vector with a single element:	
_	The proportion of non-infected individuals who have recovered.	

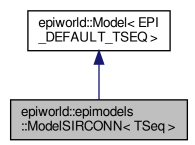
Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

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

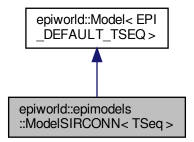
• include/epiworld/models/sir.hpp

# 18.47 epiworld::epimodels::ModelSIRCONN< TSeq > Class Template Reference

Inheritance diagram for epiworld::epimodels::ModelSIRCONN< TSeq >:



Collaboration diagram for epiworld::epimodels::ModelSIRCONN< TSeq >:



## **Public Member Functions**

ModelSIRCONN (ModelSIRCONN TSeq > &model, const std::string &vname, epiworld\_fast\_uint n, epiworld\_double prevalence, epiworld\_double contact\_rate, epiworld\_double transmission\_rate, epiworld
 —double recovery\_rate)

Template for a Susceptible-Infected-Removed (SIR) model.

- **ModelSIRCONN** (const std::string &vname, epiworld\_fast\_uint n, epiworld\_double prevalence, epiworld\_← double contact\_rate, epiworld\_double transmission\_rate, epiworld\_double recovery\_rate)
- ModelSIRCONN
   TSeq > & run (epiworld\_fast\_uint ndays, int seed=-1)

Runs the simulation (after initialization)

· void reset ()

Reset the model.

Model < TSeq > \* clone\_ptr ()

Advanced usage: Makes a copy of data and returns it as undeleted pointer.

ModelSIRCONN< TSeq > & initial\_states (std::vector< double > proportions\_, std::vector< int > queue ← \_={})

Set the initial states of the model.

• size\_t get\_n\_infected () const

Get the infected individuals.

• std::vector< double > generation\_time\_expected (int max\_days=200, int max\_contacts=200) const

#### **Static Public Attributes**

- static const int **SUSCEPTIBLE** = 0
- static const int INFECTED = 1
- static const int **RECOVERED** = 2

## **Additional Inherited Members**

## 18.47.1 Constructor & Destructor Documentation

## 18.47.1.1 ModelSIRCONN()

Template for a Susceptible-Infected-Removed (SIR) model.

## **Parameters**

model	A Model <tseq> object where to set up the SIR.</tseq>
vname	std::string Name of the virus
prevalence	Initial prevalence (proportion)
contact_rate	Average number of contacts (interactions) per step.
transmission_rate	Probability of transmission
recovery_rate	Probability of recovery

## 18.47.2 Member Function Documentation

#### 18.47.2.1 clone\_ptr()

```
template<typename TSeq >
Model< TSeq > * ModelSIRCONN< TSeq >::clone_ptr [inline], [virtual]
```

Advanced usage: Makes a copy of data and returns it as undeleted pointer.

#### **Parameters**

```
сору
```

Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

## 18.47.2.2 get\_n\_infected()

```
template<typename TSeq = EPI_DEFAULT_TSEQ>
size_t epiworld::epimodels::ModelSIRCONN< TSeq >::get_n_infected ( ) const [inline]
```

Get the infected individuals.

#### Returns

```
std::vector< epiworld::Agent<TSeq> * >
```

#### 18.47.2.3 initial states()

Set the initial states of the model.

## **Parameters**

proportions↔	Double vector with a single element:
_	The proportion of non-infected individuals who have recovered.

Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

## 18.47.2.4 reset()

```
template<typename TSeq >
void ModelSIRCONN< TSeq >::reset [inline], [virtual]
```

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

Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

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

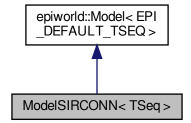
· epiworld.hpp

## 18.48 ModelSIRCONN< TSeq > Class Template Reference

Inheritance diagram for ModelSIRCONN < TSeq >:



Collaboration diagram for ModelSIRCONN < TSeq >:



#### **Public Member Functions**

ModelSIRCONN (ModelSIRCONN
 TSeq > &model, const std::string &vname, epiworld\_fast\_uint n, epiworld\_double prevalence, epiworld\_double contact\_rate, epiworld\_double transmission\_rate, epiworld← \_double recovery\_rate)

Template for a Susceptible-Infected-Removed (SIR) model.

- ModelSIRCONN (const std::string &vname, epiworld\_fast\_uint n, epiworld\_double prevalence, epiworld\_← double contact\_rate, epiworld\_double transmission\_rate, epiworld\_double recovery\_rate)
- ModelSIRCONN
   TSeq > & run (epiworld fast uint ndays, int seed=-1)

Runs the simulation (after initialization)

· void reset ()

Reset the model.

Model < TSeq > \* clone\_ptr ()

Advanced usage: Makes a copy of data and returns it as undeleted pointer.

ModelSIRCONN< TSeq > & initial\_states (std::vector< double > proportions\_, std::vector< int > queue ← \_={})

Set the initial states of the model.

• size t get n infected () const

Get the infected individuals.

std::vector< double > generation\_time\_expected (int max\_days=200, int max\_contacts=200) const

## **Static Public Attributes**

- static const int **SUSCEPTIBLE** = 0
- static const int INFECTED = 1
- static const int **RECOVERED** = 2

## **Additional Inherited Members**

## 18.48.1 Constructor & Destructor Documentation

## 18.48.1.1 ModelSIRCONN()

Template for a Susceptible-Infected-Removed (SIR) model.

#### **Parameters**

model	A Model <tseq> object where to set up the SIR.</tseq>	
vname	std::string Name of the virus	
prevalence	Initial prevalence (proportion)	Generated by Doxygen
contact_rate	Average number of contacts (interactions) per step.	
transmission_rate	Probability of transmission	
recovery rate	Probability of recovery	

## 18.48.2 Member Function Documentation

## 18.48.2.1 clone\_ptr()

```
template<typename TSeq >
Model< TSeq > * ModelSIRCONN< TSeq >::clone_ptr [inline], [virtual]
```

Advanced usage: Makes a copy of data and returns it as undeleted pointer.

#### **Parameters**

сору

Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

## 18.48.2.2 get\_n\_infected()

```
template<typename TSeq = EPI_DEFAULT_TSEQ>
size_t ModelSIRCONN< TSeq >::get_n_infected ( ) const [inline]
```

Get the infected individuals.

## Returns

std::vector< epiworld::Agent<TSeq> \* >

## 18.48.2.3 initial\_states()

Set the initial states of the model.

## **Parameters**

proportions⊷	Double vector with a single element:
_	The proportion of non-infected individuals who have recovered.

 $\label{lem:lemented_lemented} \textbf{Reimplemented from epiworld::} \\ \textbf{Model} < \textbf{EPI\_DEFAULT\_TSEQ} >. \\$ 

#### 18.48.2.4 reset()

```
template<typename TSeq >
void ModelSIRCONN< TSeq >::reset [inline], [virtual]
```

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

Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

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

· include/epiworld/models/sirconnected.hpp

# 18.49 epiworld::epimodels::ModelSIRD< TSeq > Class Template Reference

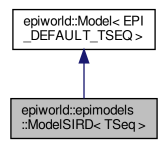
Template for a Susceptible-Infected-Removed-Deceased (SIRD) model.

```
#include <epiworld.hpp>
```

Inheritance diagram for epiworld::epimodels::ModelSIRD < TSeq >:



Collaboration diagram for epiworld::epimodels::ModelSIRD< TSeq >:



#### **Public Member Functions**

- ModelSIRD < TSeq > & initial\_states (std::vector < double > proportions\_, std::vector < int > queue\_={})
   Set the initial states of the model.
- ModelSIRD (ModelSIRD< TSeq > &model, const std::string &vname, epiworld\_double prevalence, epiworld\_double transmission\_rate, epiworld\_double recovery\_rate, epiworld\_double death\_rate)
   Constructs a new SIRD model with the given parameters.
- **ModelSIRD** (const std::string &vname, epiworld\_double prevalence, epiworld\_double transmission\_rate, epiworld\_double recovery\_rate, epiworld\_double death\_rate)

#### **Additional Inherited Members**

## 18.49.1 Detailed Description

template<typename TSeq = EPI\_DEFAULT\_TSEQ> class epiworld::epimodels::ModelSIRD< TSeq >

Template for a Susceptible-Infected-Removed-Deceased (SIRD) model.

## 18.49.2 Constructor & Destructor Documentation

## 18.49.2.1 ModelSIRD()

Constructs a new SIRD model with the given parameters.

#### **Parameters**

model	The SIRD model to copy from.
vname	The name of the vertex associated with this model.
prevalence	The initial prevalence of the disease in the population.
transmission_rate	The rate at which the disease spreads from infected to susceptible individuals.
recovery_rate	The rate at which infected individuals recover and become immune.
death_rate	The rate at which infected individuals die.

#### 18.49.3 Member Function Documentation

## 18.49.3.1 initial\_states()

Set the initial states of the model.

#### **Parameters**

proportions←	Double vector with two elements:
_	The proportion of non-infected individuals who have recovered.
	The proportion of non-infected individuals who have died.

Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

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

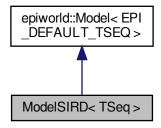
· epiworld.hpp

# 18.50 ModelSIRD< TSeq> Class Template Reference

Template for a Susceptible-Infected-Removed-Deceased (SIRD) model.

```
#include <sird.hpp>
```

Inheritance diagram for ModelSIRD< TSeq >:



Collaboration diagram for ModelSIRD < TSeq >:



## **Public Member Functions**

- ModelSIRD < TSeq > & initial\_states (std::vector < double > proportions\_, std::vector < int > queue\_={})
   Set the initial states of the model.
- ModelSIRD (ModelSIRD< TSeq > &model, const std::string &vname, epiworld\_double prevalence, epiworld\_double transmission\_rate, epiworld\_double recovery\_rate, epiworld\_double death\_rate)
   Constructs a new SIRD model with the given parameters.
- **ModelSIRD** (const std::string &vname, epiworld\_double prevalence, epiworld\_double transmission\_rate, epiworld\_double recovery\_rate, epiworld\_double death\_rate)

## **Additional Inherited Members**

## 18.50.1 Detailed Description

template<typename TSeq = EPI\_DEFAULT\_TSEQ> class ModelSIRD< TSeq >

Template for a Susceptible-Infected-Removed-Deceased (SIRD) model.

# 18.50.2 Constructor & Destructor Documentation

## 18.50.2.1 ModelSIRD()

Constructs a new SIRD model with the given parameters.

#### **Parameters**

model	The SIRD model to copy from.
vname	The name of the vertex associated with this model.
prevalence	The initial prevalence of the disease in the population.
transmission_rate	The rate at which the disease spreads from infected to susceptible individuals.
recovery_rate	The rate at which infected individuals recover and become immune.
death_rate	The rate at which infected individuals die.

# 18.50.3 Member Function Documentation

#### 18.50.3.1 initial\_states()

Set the initial states of the model.

#### **Parameters**

proportions⇔	Double vector with two elements:
_	The proportion of non-infected individuals who have recovered.
	The proportion of non-infected individuals who have died.

Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

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

• include/epiworld/models/sird.hpp

# 18.51 epiworld::epimodels::ModelSIRDCONN< TSeq > Class Template Reference

Inheritance diagram for epiworld::epimodels::ModelSIRDCONN< TSeq >:



Collaboration diagram for epiworld::epimodels::ModelSIRDCONN< TSeq >:



# **Public Member Functions**

ModelSIRDCONN (ModelSIRDCONN
 TSeq > &model, const std::string &vname, epiworld\_fast\_uint n, epiworld\_double prevalence, epiworld\_double contact\_rate, epiworld\_double transmission\_rate, epiworld\_← double recovery\_rate, epiworld\_double death\_rate)

Template for a Susceptible-Infected-Removed (SIR) model.

- ModelSIRDCONN (const std::string &vname, epiworld\_fast\_uint n, epiworld\_double prevalence, epiworld
   \_double contact\_rate, epiworld\_double transmission\_rate, epiworld\_double recovery\_rate, epiworld\_double death\_rate)
- ModelSIRDCONN< TSeq > & run (epiworld\_fast\_uint ndays, int seed=-1)

Runs the simulation (after initialization)

· void reset ()

Reset the model.

Model < TSeq > \* clone\_ptr ()

Advanced usage: Makes a copy of data and returns it as undeleted pointer.

### **Static Public Attributes**

- static const int SUSCEPTIBLE = 0
- static const int INFECTED = 1
- static const int RECOVERED = 2
- static const int **DECEASED** = 3

## **Additional Inherited Members**

# 18.51.1 Constructor & Destructor Documentation

## 18.51.1.1 ModelSIRDCONN()

Template for a Susceptible-Infected-Removed (SIR) model.

#### **Parameters**

model	A Model <tseq> object where to set up the SIR.</tseq>
vname	std::string Name of the virus
prevalence	Initial prevalence (proportion)
contact_rate	Average number of contacts (interactions) per step.
transmission_rate	Probability of transmission
recovery_rate	Probability of recovery
death_rate	Probability of death

# 18.51.2 Member Function Documentation

# 18.51.2.1 clone\_ptr()

```
template<typename TSeq >
Model< TSeq > * ModelSIRDCONN< TSeq >::clone_ptr [inline], [virtual]
```

Advanced usage: Makes a copy of data and returns it as undeleted pointer.

**Parameters** 

сору

Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

### 18.51.2.2 reset()

```
template<typename TSeq >
void ModelSIRDCONN< TSeq >::reset [inline], [virtual]
```

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

Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

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

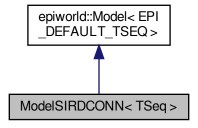
· epiworld.hpp

# 18.52 ModelSIRDCONN< TSeq > Class Template Reference

Inheritance diagram for ModelSIRDCONN< TSeq >:



Collaboration diagram for ModelSIRDCONN< TSeq >:



# **Public Member Functions**

ModelSIRDCONN (ModelSIRDCONN < TSeq > &model, const std::string &vname, epiworld\_fast\_uint n, epiworld\_double prevalence, epiworld\_double contact\_rate, epiworld\_double transmission\_rate, epiworld\_← double recovery\_rate, epiworld\_double death\_rate)

Template for a Susceptible-Infected-Removed (SIR) model.

- ModelSIRDCONN (const std::string &vname, epiworld\_fast\_uint n, epiworld\_double prevalence, epiworld
   —double contact\_rate, epiworld\_double transmission\_rate, epiworld\_double recovery\_rate, epiworld\_double death\_rate)
- ModelSIRDCONN< TSeq > & run (epiworld\_fast\_uint ndays, int seed=-1)

Runs the simulation (after initialization)

· void reset ()

Reset the model.

Model < TSeq > \* clone ptr ()

Advanced usage: Makes a copy of data and returns it as undeleted pointer.

### **Static Public Attributes**

- static const int **SUSCEPTIBLE** = 0
- static const int INFECTED = 1
- static const int **RECOVERED** = 2
- static const int **DECEASED** = 3

# **Additional Inherited Members**

#### 18.52.1 Constructor & Destructor Documentation

### 18.52.1.1 ModelSIRDCONN()

Template for a Susceptible-Infected-Removed (SIR) model.

# Parameters

model	A Model <tseq> object where to set up the SIR.</tseq>
vname	std::string Name of the virus
prevalence	Initial prevalence (proportion)
contact_rate	Average number of contacts (interactions) per step.
transmission_rate	Probability of transmission
recovery_rate	Probability of recovery
death_rate	Probability of death

### 18.52.2 Member Function Documentation

# 18.52.2.1 clone\_ptr()

```
template<typename TSeq >
Model< TSeq > * ModelSIRDCONN< TSeq >::clone_ptr [inline], [virtual]
```

Advanced usage: Makes a copy of data and returns it as undeleted pointer.

#### **Parameters**

сору

Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

#### 18.52.2.2 reset()

```
template<typename TSeq >
void ModelSIRDCONN< TSeq >::reset [inline], [virtual]
```

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

Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

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

• include/epiworld/models/sirdconnected.hpp

# 18.53 epiworld::epimodels::ModelSIRLogit< TSeq > Class Template Reference

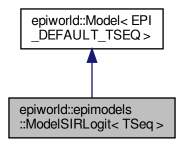
Template for a Susceptible-Infected-Removed (SIR) model.

```
#include <epiworld.hpp>
```

 $Inheritance\ diagram\ for\ epiworld::epimodels::ModelSIRLogit < TSeq >:$ 



Collaboration diagram for epiworld::epimodels::ModelSIRLogit< TSeq >:



#### **Public Member Functions**

ModelSIRLogit (ModelSIRLogit < TSeq > &model, const std::string &vname, double \*data, size\_t ncols, std::vector < double > coefs\_infect, std::vector < double > coefs\_recover, std::vector < size\_t > coef\_← infect\_cols, std::vector < size\_t > coef\_recover\_cols, epiworld\_double transmission\_rate, epiworld\_double recovery\_rate, epiworld\_double prevalence)

Template for a Susceptible-Infected-Removed (SIR) model.

- ModelSIRLogit (const std::string &vname, double \*data, size\_t ncols, std::vector< double > coefs\_infect, std::vector< double > coefs\_recover, std::vector< size\_t > coef\_infect\_cols, std::vector< size\_t > coef← \_recover\_cols, epiworld\_double transmission\_rate, epiworld\_double recovery\_rate, epiworld\_double prevalence)
- ModelSIRLogit < TSeq > & run (epiworld\_fast\_uint ndays, int seed=-1)

Runs the simulation (after initialization)

Model < TSeq > \* clone ptr ()

Advanced usage: Makes a copy of data and returns it as undeleted pointer.

· void reset ()

Reset the model.

# **Public Attributes**

- std::vector< double > coefs infect
- std::vector< double > coefs\_recover
- std::vector< size t > coef\_infect\_cols
- std::vector< size\_t > coef\_recover\_cols

# **Additional Inherited Members**

# 18.53.1 Detailed Description

 $\label{template} \mbox{typename TSeq = EPI_DEFAULT_TSEQ>} \\ \mbox{class epiworld::epimodels::ModelSIRLogit< TSeq>} \\$ 

Template for a Susceptible-Infected-Removed (SIR) model.

In this model, infection and recoveru probabilities are computed using a logit model. Particularly, the probability of infection is computed as:

$$\frac{1}{1 + \exp\left(-\left(\beta_0 E_i + \sum_{i=1}^n \beta_i x_i\right)\right)}$$

where  $\beta_0$  is the exposure coefficient and  $E_i$  is the exposure number,  $\beta_i$  are the coefficients for the features  $x_i$  of the agents, and n is the number of features. The probability of recovery is computed as:

$$\frac{1}{1 + \exp\left(-\left(\sum_{i=1}^{n} \beta_i x_i\right)\right)}$$

where  $\beta_i$  are the coefficients for the features  $x_i$  of the agents, and n is the number of features.

#### **Parameters**

TSeq Type of the seq	uence (e.g. std::vector, std::deque)
----------------------	--------------------------------------

# 18.53.2 Constructor & Destructor Documentation

# 18.53.2.1 ModelSIRLogit()

Template for a Susceptible-Infected-Removed (SIR) model.

#### **Parameters**

vname	Name of the virus.
coefs_infect	Double ptr. Infection coefficients.
coefs_recover	Double ptr. Recovery coefficients.
ncoef_infect	Unsigned int. Number of infection coefficients.
ncoef_recover	Unsigned int. Number of recovery coefficients.
coef_infect_cols	Vector <unsigned int="">. Ids of infection vars.</unsigned>
coef_recover_cols	Vector <unsigned int="">. Ids of recover vars.</unsigned>

#### **Parameters**

model	A Model <tseq> object where to set up the SIR.</tseq>
vname	std::string Name of the virus
prevalence	Initial prevalence (proportion)
contact_rate	Average number of contacts (interactions) per step.
prob_transmission	Probability of transmission
prob_recovery	Probability of recovery

# 18.53.3 Member Function Documentation

#### 18.53.3.1 clone\_ptr()

```
template<typename TSeq >
Model< TSeq > * ModelSIRLogit< TSeq >::clone_ptr [inline], [virtual]
```

Advanced usage: Makes a copy of data and returns it as undeleted pointer.

#### **Parameters**

сору

 $\label{eq:control_equation} \textbf{Reimplemented from epiworld::} \\ \textbf{Model} < \textbf{EPI\_DEFAULT\_TSEQ} >.$ 

# 18.53.3.2 reset()

```
template<typename TSeq >
void ModelSIRLogit< TSeq >::reset [inline], [virtual]
```

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

Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

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

· epiworld.hpp

# 18.54 ModelSIRLogit < TSeq > Class Template Reference

Template for a Susceptible-Infected-Removed (SIR) model.

#include <sirlogit.hpp>

Inheritance diagram for ModelSIRLogit< TSeg >:



Collaboration diagram for ModelSIRLogit < TSeq >:



# **Public Member Functions**

ModelSIRLogit (ModelSIRLogit < TSeq > &model, const std::string &vname, double \*data, size\_t ncols, std::vector < double > coefs\_infect, std::vector < double > coefs\_recover, std::vector < size\_t > coef\_ ← infect\_cols, std::vector < size\_t > coef\_recover\_cols, epiworld\_double transmission\_rate, epiworld\_double recovery\_rate, epiworld\_double prevalence)

Template for a Susceptible-Infected-Removed (SIR) model.

- ModelSIRLogit (const std::string &vname, double \*data, size\_t ncols, std::vector< double > coefs\_infect, std::vector< double > coefs\_recover, std::vector< size\_t > coef\_infect\_cols, std::vector< size\_t > coef← \_recover\_cols, epiworld\_double transmission\_rate, epiworld\_double recovery\_rate, epiworld\_double prevalence)
- ModelSIRLogit < TSeq > & run (epiworld\_fast\_uint ndays, int seed=-1)

Runs the simulation (after initialization)

Model < TSeq > \* clone\_ptr ()

Advanced usage: Makes a copy of data and returns it as undeleted pointer.

· void reset ()

Reset the model.

### **Public Attributes**

- $std::vector < double > coefs_infect$
- std::vector< double > coefs\_recover
- std::vector< size t > coef\_infect\_cols
- std::vector< size\_t > coef\_recover\_cols

#### **Additional Inherited Members**

# 18.54.1 Detailed Description

```
template<typename TSeq = EPI_DEFAULT_TSEQ> class ModelSIRLogit < TSeq >
```

Template for a Susceptible-Infected-Removed (SIR) model.

In this model, infection and recoveru probabilities are computed using a logit model. Particularly, the probability of infection is computed as:

$$\frac{1}{1 + \exp\left(-\left(\beta_0 E_i + \sum_{i=1}^n \beta_i x_i\right)\right)}$$

where  $\beta_0$  is the exposure coefficient and  $E_i$  is the exposure number,  $\beta_i$  are the coefficients for the features  $x_i$  of the agents, and n is the number of features. The probability of recovery is computed as:

$$\frac{1}{1 + \exp\left(-\left(\sum_{i=1}^{n} \beta_i x_i\right)\right)}$$

where  $\beta_i$  are the coefficients for the features  $x_i$  of the agents, and n is the number of features.

#### **Parameters**

```
TSeq Type of the sequence (e.g. std::vector, std::deque)
```

#### 18.54.2 Constructor & Destructor Documentation

## 18.54.2.1 ModelSIRLogit()

```
std::vector< double > coefs_infect,
std::vector< double > coefs_recover,
std::vector< size_t > coef_infect_cols,
std::vector< size_t > coef_recover_cols,
epiworld_double transmission_rate,
epiworld_double recovery_rate,
epiworld_double prevalence ) [inline]
```

Template for a Susceptible-Infected-Removed (SIR) model.

#### **Parameters**

vname	Name of the virus.
coefs_infect	Double ptr. Infection coefficients.
coefs_recover	Double ptr. Recovery coefficients.
ncoef_infect	Unsigned int. Number of infection coefficients.
ncoef_recover	Unsigned int. Number of recovery coefficients.
coef_infect_cols	Vector <unsigned int="">. Ids of infection vars.</unsigned>
coef_recover_cols	Vector <unsigned int="">. Ids of recover vars.</unsigned>
model	A Model <tseq> object where to set up the SIR.</tseq>
vname	std::string Name of the virus
prevalence	Initial prevalence (proportion)
contact_rate	Average number of contacts (interactions) per step.
prob_transmission	Probability of transmission
prob_recovery	Probability of recovery

# 18.54.3 Member Function Documentation

#### 18.54.3.1 clone\_ptr()

```
template<typename TSeq >
Model< TSeq > * ModelSIRLogit< TSeq >::clone_ptr [inline], [virtual]
```

Advanced usage: Makes a copy of data and returns it as undeleted pointer.

#### **Parameters**

сору

Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

#### 18.54.3.2 reset()

```
template<typename TSeq >
void ModelSIRLogit< TSeq >::reset [inline], [virtual]
```

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

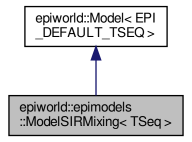
Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

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

• include/epiworld/models/sirlogit.hpp

# 18.55 epiworld::epimodels::ModelSIRMixing< TSeq > Class Template Reference

Inheritance diagram for epiworld::epimodels::ModelSIRMixing< TSeq >:



 $\label{localized} \mbox{Collaboration diagram for epiworld::epimodels::} \mbox{ModelSIRMixing} < \mbox{TSeq} > :$ 



#### **Public Member Functions**

ModelSIRMixing (ModelSIRMixing < TSeq > &model, const std::string &vname, epiworld\_fast\_uint n, epiworld\_double prevalence, epiworld\_double contact\_rate, epiworld\_double transmission\_rate, epiworld
 — double recovery\_rate, std::vector < double > contact\_matrix)

Constructs a ModelSIRMixing object.

ModelSIRMixing (const std::string &vname, epiworld\_fast\_uint n, epiworld\_double prevalence, epiworld\_
 double contact\_rate, epiworld\_double transmission\_rate, epiworld\_double recovery\_rate, std::vector < double > contact\_matrix)

Constructs a ModelSIRMixing object.

ModelSIRMixing < TSeq > & run (epiworld fast uint ndays, int seed=-1)

Runs the simulation (after initialization)

· void reset ()

Reset the model.

Model < TSeq > \* clone\_ptr ()

Advanced usage: Makes a copy of data and returns it as undeleted pointer.

ModelSIRMixing< TSeq > & initial\_states (std::vector< double > proportions\_, std::vector< int > queue ← \_={})

Set the initial states of the model.

- size\_t get\_n\_infected (size\_t group) const
- void set contact matrix (std::vector< double > cmat)

#### Static Public Attributes

- static const int SUSCEPTIBLE = 0
- static const int INFECTED = 1
- static const int **RECOVERED** = 2

#### **Additional Inherited Members**

#### 18.55.1 Constructor & Destructor Documentation

### 18.55.1.1 ModelSIRMixing() [1/2]

Constructs a ModelSIRMixing object.

Template for a Susceptible-Exposed-Infected-Removed (SEIR) model.

#### **Parameters**

model	A reference to an existing ModelSIRMixing object.
vname	The name of the ModelSIRMixing object.
n	The number of entities in the model.
prevalence	The initial prevalence of the disease in the model.
contact_rate	The contact rate between entities in the model.
transmission_rate	The transmission rate of the disease in the model.
recovery_rate	The recovery rate of the disease in the model.
contact_matrix	The contact matrix between entities in the model.
model	A Model <tseq> object where to set up the SIR.</tseq>
vname	std::string Name of the virus
prevalence	Initial prevalence (proportion)
contact_rate	Average number of contacts (interactions) per step.
transmission_rate	Probability of transmission
recovery_rate	Probability of recovery

# 18.55.1.2 ModelSIRMixing() [2/2]

# Constructs a ModelSIRMixing object.

#### **Parameters**

vname	The name of the ModelSIRMixing object.
n	The number of entities in the model.
prevalence	The initial prevalence of the disease in the model.
contact_rate	The contact rate between entities in the model.
transmission_rate	The transmission rate of the disease in the model.
recovery_rate	The recovery rate of the disease in the model.
contact_matrix	The contact matrix between entities in the model.

# 18.55.2 Member Function Documentation

#### 18.55.2.1 clone\_ptr()

```
template<typename TSeq >
Model< TSeq > * ModelSIRMixing< TSeq >::clone_ptr [inline], [virtual]
```

Advanced usage: Makes a copy of data and returns it as undeleted pointer.

#### **Parameters**

```
сору
```

Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

### 18.55.2.2 initial\_states()

Set the initial states of the model.

#### **Parameters**

proportions↔	Double vector with a single element:
_	The proportion of non-infected individuals who have recovered.

 $\label{eq:control_equation} \mbox{Reimplemented from epiworld::} \mbox{Model} < \mbox{EPI\_DEFAULT\_TSEQ} >.$ 

#### 18.55.2.3 reset()

```
template<typename TSeq >
void ModelSIRMixing< TSeq >::reset [inline], [virtual]
```

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

Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

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

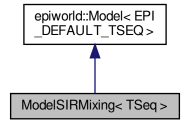
· epiworld.hpp

# 18.56 ModelSIRMixing < TSeq > Class Template Reference

Inheritance diagram for ModelSIRMixing < TSeq >:



Collaboration diagram for ModelSIRMixing < TSeq >:



#### **Public Member Functions**

ModelSIRMixing (ModelSIRMixing < TSeq > &model, const std::string &vname, epiworld\_fast\_uint n, epiworld\_double prevalence, epiworld\_double contact\_rate, epiworld\_double transmission\_rate, epiworld← \_double recovery\_rate, std::vector < double > contact\_matrix)

Constructs a ModelSIRMixing object.

ModelSIRMixing (const std::string &vname, epiworld\_fast\_uint n, epiworld\_double prevalence, epiworld\_
 double contact\_rate, epiworld\_double transmission\_rate, epiworld\_double recovery\_rate, std::vector < double > contact\_matrix)

Constructs a ModelSIRMixing object.

ModelSIRMixing< TSeq > & run (epiworld\_fast\_uint ndays, int seed=-1)

Runs the simulation (after initialization)

· void reset ()

Reset the model.

Model < TSeq > \* clone\_ptr ()

Advanced usage: Makes a copy of data and returns it as undeleted pointer.

ModelSIRMixing< TSeq > & initial\_states (std::vector< double > proportions\_, std::vector< int > queue ← \_={})

Set the initial states of the model.

- size\_t get\_n\_infected (size\_t group) const
- void set\_contact\_matrix (std::vector< double > cmat)

#### **Static Public Attributes**

- static const int SUSCEPTIBLE = 0
- static const int INFECTED = 1
- static const int **RECOVERED** = 2

### **Additional Inherited Members**

#### 18.56.1 Constructor & Destructor Documentation

### 18.56.1.1 ModelSIRMixing() [1/2]

### Constructs a ModelSIRMixing object.

Template for a Susceptible-Exposed-Infected-Removed (SEIR) model.

#### **Parameters**

A reference to an existing ModelSIRMixing object.
The name of the ModelSIRMixing object.
The number of entities in the model.
The initial prevalence of the disease in the model.
The contact rate between entities in the model.
The transmission rate of the disease in the model.
The recovery rate of the disease in the model.
The contact matrix between entities in the model.
A Model <tseq> object where to set up the SIR.</tseq>
std::string Name of the virus
Initial prevalence (proportion)
Average number of contacts (interactions) per step.
Probability of transmission
Probability of recovery

### 18.56.1.2 ModelSIRMixing() [2/2]

Constructs a ModelSIRMixing object.

#### **Parameters**

vname	The name of the ModelSIRMixing object.
n	The number of entities in the model.
prevalence	The initial prevalence of the disease in the model.
contact_rate	The contact rate between entities in the model.
transmission_rate	The transmission rate of the disease in the model.
recovery_rate	The recovery rate of the disease in the model.
contact_matrix	The contact matrix between entities in the model.

### 18.56.2 Member Function Documentation

## 18.56.2.1 clone\_ptr()

```
template<typename TSeq >
Model< TSeq > * ModelSIRMixing< TSeq >::clone_ptr [inline], [virtual]
```

Advanced usage: Makes a copy of data and returns it as undeleted pointer.

#### **Parameters**



 $\label{eq:local_problem} \mbox{Reimplemented from epiworld::} \mbox{Model} < \mbox{EPI\_DEFAULT\_TSEQ} >.$ 

### 18.56.2.2 initial\_states()

```
template<typename TSeq >
ModelSIRMixing< TSeq > & ModelSIRMixing< TSeq >::initial_states (
```

```
std::vector< double > proportions_,
std::vector< int > queue_ = {} ) [inline], [virtual]
```

Set the initial states of the model.

#### **Parameters**

proportions↔	Double vector with a single element:
_	The proportion of non-infected individuals who have recovered.

Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

#### 18.56.2.3 reset()

```
template<typename TSeq >
void ModelSIRMixing< TSeq >::reset [inline], [virtual]
```

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

Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

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

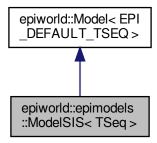
• include/epiworld/models/sirmixing.hpp

# 18.57 epiworld::epimodels::ModelSIS< TSeq > Class Template Reference

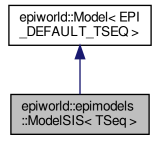
Template for a Susceptible-Infected-Susceptible (SIS) model.

```
#include <epiworld.hpp>
```

Inheritance diagram for epiworld::epimodels::ModelSIS < TSeq >:



Collaboration diagram for epiworld::epimodels::ModelSIS< TSeq >:



# **Public Member Functions**

- ModelSIS (ModelSIS < TSeq > &model, const std::string &vname, epiworld\_double prevalence, epiworld
   —double transmission\_rate, epiworld\_double recovery\_rate)
- **ModelSIS** (const std::string &vname, epiworld\_double prevalence, epiworld\_double transmission\_rate, epiworld\_double recovery\_rate)

# **Static Public Attributes**

- static const int **SUSCEPTIBLE** = 0
- static const int INFECTED = 1

# **Additional Inherited Members**

# 18.57.1 Detailed Description

 $\label{template} \begin{tabular}{ll} template < typename TSeq = EPI\_DEFAULT\_TSEQ > \\ class epiworld::epimodels::ModelSIS < TSeq > \\ \end{tabular}$ 

Template for a Susceptible-Infected-Susceptible (SIS) model.

#### **Parameters**

vname	std::string Name of the virus
initial_prevalence	epiworld_double Initial prevalence
initial_efficacy	epiworld_double Initial susceptibility_reduction of the immune system
initial_recovery	epiworld_double Initial recovery_rate rate of the immune system

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

· epiworld.hpp

# **18.58** ModelSIS< TSeq > Class Template Reference

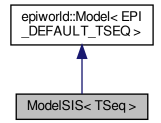
Template for a Susceptible-Infected-Susceptible (SIS) model.

#include <sis.hpp>

Inheritance diagram for ModelSIS < TSeq >:



Collaboration diagram for ModelSIS < TSeq >:



### **Public Member Functions**

- ModelSIS (ModelSIS < TSeq > &model, const std::string &vname, epiworld\_double prevalence, epiworld
   \_double transmission\_rate, epiworld\_double recovery\_rate)
- **ModelSIS** (const std::string &vname, epiworld\_double prevalence, epiworld\_double transmission\_rate, epiworld\_double recovery\_rate)

### **Static Public Attributes**

- static const int **SUSCEPTIBLE** = 0
- static const int INFECTED = 1

### **Additional Inherited Members**

### 18.58.1 Detailed Description

```
template<typename TSeq = EPI_DEFAULT_TSEQ> class ModelSIS< TSeq >
```

Template for a Susceptible-Infected-Susceptible (SIS) model.

#### **Parameters**

vname	std::string Name of the virus
initial_prevalence	epiworld_double Initial prevalence
initial_efficacy	epiworld_double Initial susceptibility_reduction of the immune system
initial_recovery	epiworld_double Initial recovery_rate rate of the immune system

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

• include/epiworld/models/sis.hpp

# 18.59 epiworld::epimodels::ModelSISD< TSeq > Class Template Reference

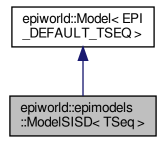
Template for a Susceptible-Infected-Susceptible-Deceased (SISD) model.

```
#include <epiworld.hpp>
```

Inheritance diagram for epiworld::epimodels::ModelSISD< TSeq >:



Collaboration diagram for epiworld::epimodels::ModelSISD< TSeq >:



### **Public Member Functions**

- **ModelSISD** (ModelSISD< TSeq > &model, const std::string &vname, epiworld\_double prevalence, epiworld double transmission rate, epiworld double recovery rate, epiworld double death rate)
- **ModelSISD** (const std::string &vname, epiworld\_double prevalence, epiworld\_double transmission\_rate, epiworld\_double recovery\_rate, epiworld\_double death\_rate)

### **Additional Inherited Members**

## 18.59.1 Detailed Description

template<typename TSeq = EPI\_DEFAULT\_TSEQ> class epiworld::epimodels::ModelSISD< TSeq >

Template for a Susceptible-Infected-Susceptible-Deceased (SISD) model.

#### **Parameters**

vname	std::string Name of the virus
initial_prevalence	epiworld_double Initial prevalence
initial_efficacy	epiworld_double Initial susceptibility_reduction of the immune system
initial_recovery	epiworld_double Initial recovery_rate rate of the immune system
inital_death	epiworld_double Initial death_rate of the immune system

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

· epiworld.hpp

# 18.60 ModelSISD< TSeq > Class Template Reference

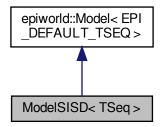
Template for a Susceptible-Infected-Susceptible-Deceased (SISD) model.

#include <sisd.hpp>

Inheritance diagram for ModelSISD< TSeq >:



Collaboration diagram for ModelSISD < TSeq >:



### **Public Member Functions**

• **ModelSISD** (ModelSISD< TSeq > &model, const std::string &vname, epiworld\_double prevalence, epiworld\_double transmission\_rate, epiworld\_double recovery\_rate, epiworld\_double death\_rate)

• **ModelSISD** (const std::string &vname, epiworld\_double prevalence, epiworld\_double transmission\_rate, epiworld\_double recovery\_rate, epiworld\_double death\_rate)

### **Additional Inherited Members**

# 18.60.1 Detailed Description

```
template<typename TSeq = EPI_DEFAULT_TSEQ> class ModelSISD< TSeq >
```

Template for a Susceptible-Infected-Susceptible-Deceased (SISD) model.

#### **Parameters**

vname	std::string Name of the virus
initial_prevalence	epiworld_double Initial prevalence
initial_efficacy	epiworld_double Initial susceptibility_reduction of the immune system
initial_recovery	epiworld_double Initial recovery_rate rate of the immune system
inital_death	epiworld_double Initial death_rate of the immune system

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

• include/epiworld/models/sisd.hpp

# 18.61 epiworld::epimodels::ModelSURV< TSeq > Class Template Reference

Inheritance diagram for epiworld::epimodels::ModelSURV< TSeq >:



Collaboration diagram for epiworld::epimodels::ModelSURV < TSeq >:



# **Public Member Functions**

• void reset ()

Reset the model.

# Construct a new ModelSURV object

The ModelSURV class simulates a survaillence model where agents can be isolated, even if asyptomatic.

#### **Parameters**

vname	String. Name of the virus
prevalence	Integer. Number of initial cases of the virus.
efficacy_vax	Double. Efficacy of the vaccine (1 - P(acquire the disease)).
latent_period	Double. Shape parameter of a Gamma (latent_period, 1) distribution. This coincides with the expected number of latent days.
infect_period	Double. Shape parameter of a Gamma (infected_period, 1) distribution. This coincides with the expected number of infectious days.
prob_symptoms	Double. Probability of generating symptoms.
prop_vaccinated	Double. Probability of vaccination. Coincides with the initial prevalence of vaccinated individuals.
prop_vax_redux_transm	Double. Factor by which the vaccine reduces transmissibility.
prop_vax_redux_infect	Double. Factor by which the vaccine reduces the chances of becoming infected.
surveillance_prob	Double. Probability of testing an agent.
prob_transmission	Double. Raw transmission probability.
prob_death	Double. Raw probability of death for symptomatic individuals.
prob_noreinfect	Double. Probability of no re-infection.

This model features the following states:

- Susceptible
- Latent
- Symptomatic
- · Symptomatic isolated

- · Asymptomatic
- · Asymptomatic isolated
- Recovered
- Removed

#### Returns

An object of class epiworld\_surv

- ModelSURV ()
- ModelSURV (ModelSURV < TSeq > &model, const std::string &vname, epiworld\_fast\_uint prevalence=50, epiworld\_double efficacy\_vax=0.9, epiworld\_double latent\_period=3u, epiworld\_double infect⇔ \_period=6u, epiworld\_double prob\_symptoms=0.6, epiworld\_double prop\_vaccinated=0.25, epiworld⇔ \_double prop\_vax\_redux\_transm=0.5, epiworld\_double prop\_vax\_redux\_infect=0.5, epiworld\_double surveillance\_prob=0.001, epiworld\_double prob\_transmission=1.0, epiworld\_double prob\_death=0.001, epiworld\_double prob noreinfect=0.9)
- ModelSURV (const std::string &vname, epiworld\_fast\_uint prevalence=50, epiworld\_double efficacy\_
   vax=0.9, epiworld\_double latent\_period=3u, epiworld\_double infect\_period=6u, epiworld\_double prob
   \_symptoms=0.6, epiworld\_double prop\_vaccinated=0.25, epiworld\_double prop\_vax\_redux\_transm=0.5,
   epiworld\_double prop\_vax\_redux\_infect=0.5, epiworld\_double surveillance\_prob=0.001, epiworld\_double
   prob\_transmission=1.0, epiworld\_double prob\_death=0.001, epiworld\_double prob\_noreinfect=0.9)

#### **Public Attributes**

std::vector< epiworld\_double > days\_latent\_and\_infectious
 Vector of days spent in latent and infectious states A row-major matrix.

### **Additional Inherited Members**

#### 18.61.1 Member Function Documentation

### 18.61.1.1 reset()

```
template<typename TSeq >
void ModelSURV< TSeq >::reset [inline], [virtual]
```

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

 $\label{eq:local_problem} \mbox{Reimplemented from epiworld::} \mbox{Model} < \mbox{EPI\_DEFAULT\_TSEQ} >.$ 

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

epiworld.hpp

# 18.62 ModelSURV < TSeq > Class Template Reference

Inheritance diagram for ModelSURV < TSeq >:



Collaboration diagram for ModelSURV < TSeq >:



# **Public Member Functions**

• void reset ()

Reset the model.

# Construct a new ModelSURV object

The ModelSURV class simulates a survaillence model where agents can be isolated, even if asyptomatic.

#### **Parameters**

vname	String. Name of the virus
prevalence	Integer. Number of initial cases of the virus.
efficacy_vax	Double. Efficacy of the vaccine (1 - P(acquire the disease)).
latent_period	Double. Shape parameter of a Gamma (latent_period, 1) distribution.  This coincides with the expected number of latent days.

#### **Parameters**

infect_period	Double. Shape parameter of a Gamma (infected_period, 1) distribution. This coincides with the expected number of infectious days.
prob_symptoms	Double. Probability of generating symptoms.
prop_vaccinated	Double. Probability of vaccination. Coincides with the initial prevalence of vaccinated individuals.
prop_vax_redux_transm	Double. Factor by which the vaccine reduces transmissibility.
prop_vax_redux_infect	Double. Factor by which the vaccine reduces the chances of becoming infected.
surveillance_prob	Double. Probability of testing an agent.
prob_transmission	Double. Raw transmission probability.
prob_death	Double. Raw probability of death for symptomatic individuals.
prob_noreinfect	Double. Probability of no re-infection.

This model features the following states:

- Susceptible
- · Latent
- Symptomatic
- · Symptomatic isolated
- · Asymptomatic
- · Asymptomatic isolated
- · Recovered
- · Removed

#### Returns

An object of class epiworld\_surv

- ModelSURV ()
- ModelSURV (ModelSURV < TSeq > &model, const std::string &vname, epiworld\_fast\_uint prevalence=50, epiworld\_double efficacy\_vax=0.9, epiworld\_double latent\_period=3u, epiworld\_double infect⇔ \_period=6u, epiworld\_double prob\_symptoms=0.6, epiworld\_double prop\_vaccinated=0.25, epiworld⇔ \_double prop\_vax\_redux\_transm=0.5, epiworld\_double prop\_vax\_redux\_infect=0.5, epiworld\_double surveillance\_prob=0.001, epiworld\_double prob\_transmission=1.0, epiworld\_double prob\_death=0.001, epiworld\_double prob\_noreinfect=0.9)
- ModelSURV (const std::string &vname, epiworld\_fast\_uint prevalence=50, epiworld\_double efficacy\_ 
  vax=0.9, epiworld\_double latent\_period=3u, epiworld\_double infect\_period=6u, epiworld\_double prob
  \_symptoms=0.6, epiworld\_double prop\_vaccinated=0.25, epiworld\_double prop\_vax\_redux\_transm=0.5,
  epiworld\_double prop\_vax\_redux\_infect=0.5, epiworld\_double surveillance\_prob=0.001, epiworld\_double
  prob\_transmission=1.0, epiworld\_double prob\_death=0.001, epiworld\_double prob\_noreinfect=0.9)

#### **Public Attributes**

std::vector< epiworld\_double > days\_latent\_and\_infectious
 Vector of days spent in latent and infectious states A row-major matrix.

#### **Additional Inherited Members**

## 18.62.1 Member Function Documentation

#### 18.62.1.1 reset()

```
template<typename TSeq >
void ModelSURV< TSeq >::reset [inline], [virtual]
```

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

Reimplemented from epiworld::Model < EPI\_DEFAULT\_TSEQ >.

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

• include/epiworld/models/surveillance.hpp

# 18.63 Network< Nettype, Nodetype, Edgetype > Class Template Reference

# **Public Member Functions**

- NType ()
- Edgetype operator() (int i, int j)
- bool is\_directed () const
- size\_t vcount () const
- size t ecount () const
- · void add edge (int i, int j)
- void **rm\_edge** (int i, int j)

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

• include/epiworld/network-bones.hpp

# 18.64 epiworld::PersonTools < TSeq > Class Template Reference

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

· epiworld.hpp

# 18.65 PersonTools < TSeq > Class Template Reference

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

· include/epiworld/config.hpp

# 18.66 epiworld::Progress Class Reference

A simple progress bar.

```
#include <epiworld.hpp>
```

#### **Public Member Functions**

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

# 18.66.1 Detailed Description

A simple progress bar.

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

· epiworld.hpp

# 18.67 Progress Class Reference

A simple progress bar.

```
#include cpress.hpp>
```

### **Public Member Functions**

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

# 18.67.1 Detailed Description

A simple progress bar.

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

• include/epiworld/progress.hpp

# 18.68 epiworld::Queue < TSeq > Class Template Reference

Controls which agents are verified at each step.

#include <epiworld.hpp>

#### **Public Member Functions**

- void operator+= (Agent < TSeq > \*p)
- void operator-= (Agent < TSeq > \*p)
- epiworld\_fast\_int & operator[] (epiworld\_fast\_uint i)
- · void reset ()
- bool operator== (const Queue < TSeq > &other) const
- bool operator!= (const Queue < TSeq > &other) const

### **Static Public Attributes**

- static const int NoOne = 0
- static const int OnlySelf = 1
- static const int **Everyone** = 2

#### **Friends**

class Model < TSeq >

### 18.68.1 Detailed Description

template<typename TSeq> class epiworld::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** 



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

· epiworld.hpp

# 18.69 Queue < TSeq > Class Template Reference

Controls which agents are verified at each step.

#include <queue-bones.hpp>

### **Public Member Functions**

- void operator+= (Agent < TSeq > \*p)
- void operator-= (Agent < TSeq > \*p)
- epiworld fast int & operator[] (epiworld fast uint i)
- void reset ()
- bool operator== (const Queue < TSeq > &other) const
- bool operator!= (const Queue < TSeq > &other) const

### **Static Public Attributes**

- static const int NoOne = 0
- static const int OnlySelf = 1
- static const int **Everyone** = 2

### **Friends**

class Model < TSeq >

# 18.69.1 Detailed Description

template<typename TSeq>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** 



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

- include/epiworld/agent-bones.hpp
- include/epiworld/queue-bones.hpp

# 18.70 RandGraph Class Reference

## **Public Member Functions**

- RandGraph (int N )
- · void init (int s)
- void  $set\_rand\_engine$  (std::shared\_ptr< std::mt19937 > &e)
- epiworld\_double runif ()

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

• include/epiworld/random\_graph.hpp

## 18.71 epiworld::SAMPLETYPE Class Reference

## **Static Public Attributes**

- static const int MODEL = 0
- static const int **ENTITY** = 1
- static const int AGENT = 2

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

· epiworld.hpp

## 18.72 SAMPLETYPE Class Reference

#### **Static Public Attributes**

- static const int MODEL = 0
- static const int **ENTITY** = 1
- static const int AGENT = 2

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

• include/epiworld/agentssample-bones.hpp

## 18.73 SimpleAdjList Struct Reference

## **Public Member Functions**

- SimpleAdjList (const std::vector< int > &source, const std::vector< int > &target, int n, bool dir)
- std::vector< std::map< int, int > > & get\_dat ()
- · bool is directed () const
- size\_t vcount () const

#### **Public Attributes**

- std::vector< std::map< int, int > > dat
- · bool directed

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

· tests/test-rewire-simple.cpp

## 18.74 SimpleModel Struct Reference

## **Public Member Functions**

- · void seed (unsigned int s)
- · double runif ()
- · bool is\_directed () const

#### **Public Attributes**

std::mt19937 gen

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

· tests/test-rewire-simple.cpp

# 18.75 epiworld::Tool < TSeq > Class Template Reference

Tools for defending the agent against the virus.

```
#include <epiworld.hpp>
```

#### **Public Member Functions**

- Tool (std::string name="unknown tool")
- Tool (std::string name, epiworld\_double prevalence, bool as\_proportion)
- void set\_sequence (TSeq d)
- void set\_sequence (std::shared\_ptr< TSeq > d)
- EPI\_TYPENAME\_TRAITS (TSeq, int) get\_sequence()
- void set\_name (std::string name)
- std::string get\_name () const
- Agent < TSeq > \* get\_agent ()
- int get\_id () const
- void set\_id (int id)
- void set\_date (int d)
- int get\_date () const
- void set\_state (epiworld\_fast\_int init, epiworld\_fast\_int post)
- void set\_queue (epiworld\_fast\_int init, epiworld\_fast\_int post)
- void get\_state (epiworld\_fast\_int \*init, epiworld\_fast\_int \*post)
- void get\_queue (epiworld\_fast\_int \*init, epiworld\_fast\_int \*post)
- bool operator== (const Tool < TSeq > &other) const
- bool operator!= (const Tool < TSeq > &other) const
- · void print () const
- void distribute (Model < TSeq > \*model)
- void set\_distribution (ToolToAgentFun< TSeq > fun)

## 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, Model< TSeq > \*model)
- epiworld double get transmission reduction (VirusPtr< TSeq > v, Model< TSeq > \*model)
- epiworld\_double get\_recovery\_enhancer (VirusPtr< TSeq > v, Model< TSeq > \*model)
- epiworld double get\_death\_reduction (VirusPtr< TSeq > v, Model< TSeq > \*model)
- 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 Agent < TSeq >
- class Model < TSeq >
- void default\_add\_tool (Event< TSeq > &a, Model< TSeq > \*m)
- void default\_rm\_tool (Event< TSeq > &a, Model< TSeq > \*m)

## 18.75.1 Detailed Description

template<typename TSeq> class epiworld::Tool< TSeq>

Tools for defending the agent against the virus.

**Template Parameters** 

TSeq Type of sequence

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

· epiworld.hpp

## 18.76 Tool < TSeq > Class Template Reference

Tools for defending the agent against the virus.

#include <tool-bones.hpp>

#### **Public Member Functions**

- Tool (std::string name="unknown tool")
- Tool (std::string name, epiworld double prevalence, bool as proportion)
- void set sequence (TSeq d)
- void set sequence (std::shared ptr< TSeq > d)
- EPI TYPENAME TRAITS (TSeq, int) get sequence()
- void set name (std::string name)
- std::string get\_name () const
- Agent < TSeq > \* get agent ()
- int get id () const
- · void set id (int id)
- · void set date (int d)
- · int get date () const
- void set\_state (epiworld\_fast\_int init, epiworld\_fast\_int post)
- void set\_queue (epiworld\_fast\_int init, epiworld\_fast\_int post)
- void get state (epiworld fast int \*init, epiworld fast int \*post)
- void get\_queue (epiworld\_fast\_int \*init, epiworld\_fast\_int \*post)
- bool operator== (const Tool < TSeq > &other) const
- bool operator!= (const Tool < TSeq > &other) const
- · void print () const
- void distribute (Model < TSeq > \*model)
- void set\_distribution (ToolToAgentFun< TSeq > fun)
- void set\_sequence (int d)
- bool operator== (const Tool < std::vector < int >> &other) const
- void set\_sequence (int d)
- bool operator== (const Tool< std::vector< int >> &other) const

#### 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, Model < TSeq > \*model)
- epiworld double get transmission reduction (VirusPtr< TSeq > v, Model< TSeq > \*model)
- epiworld double get recovery enhancer (VirusPtr< TSeq > v, Model< TSeq > \*model)
- epiworld\_double get\_death\_reduction (VirusPtr< TSeq > v, Model< TSeq > \*model)
- void set\_susceptibility\_reduction\_fun (ToolFun < TSeq > fun)
- void  ${\bf set\_transmission\_reduction\_fun}$  (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 Agent < TSeq >
- class Model < TSeq >
- void default\_add\_tool (Event< TSeq > &a, Model< TSeq > \*m)
- void default\_rm\_tool (Event< TSeq > &a, Model< TSeq > \*m)

## 18.76.1 Detailed Description

```
template < typename TSeq > class Tool < TSeq >
```

Tools for defending the agent against the virus.

**Template Parameters** 

```
TSeq Type of sequence
```

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

- · include/epiworld/agent-bones.hpp
- include/epiworld/tool-bones.hpp
- · include/epiworld/tool-meat.hpp

# 18.77 epiworld::ToolFunctions < TSeq > Class Template Reference

Helper class to store the functions avoiding multiple shared\_pointers (we have only one for the four of these)

```
#include <epiworld.hpp>
```

## **Public Attributes**

- ToolFun< TSeq > susceptibility\_reduction = nullptr
- ToolFun< TSeq> transmission\_reduction = nullptr
- ToolFun< TSeq> recovery\_enhancer = nullptr
- $\bullet \quad \mathsf{ToolFun} \! < \mathsf{TSeq} > \mathbf{death\_reduction} = \mathsf{nullptr}$
- ToolToAgentFun< TSeq > dist = nullptr

## 18.77.1 Detailed Description

```
\label{template} \mbox{template} < \mbox{typename TSeq} > \\ \mbox{class epiworld::ToolFunctions} < \mbox{TSeq} > \\
```

Helper class to store the functions avoiding multiple shared\_pointers (we have only one for the four of these)

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

· epiworld.hpp

## 18.78 ToolFunctions < TSeq > Class Template Reference

Helper class to store the functions avoiding multiple shared\_pointers (we have only one for the four of these)

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

#### **Public Attributes**

- ToolFun< TSeq > susceptibility\_reduction = nullptr
- ToolFun< TSeq > transmission\_reduction = nullptr
- ToolFun< TSeq > recovery\_enhancer = nullptr
- ToolFun< TSeq > death\_reduction = nullptr
- ToolToAgentFun< TSeq > dist = nullptr

## 18.78.1 Detailed Description

```
template<typename TSeq> class ToolFunctions< TSeq >
```

Helper class to store the functions avoiding multiple shared\_pointers (we have only one for the four of these)

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

include/epiworld/tool-bones.hpp

# 18.79 epiworld::Tools < TSeq > Class Template Reference

Set of tools (useful for building iterators)

```
#include <epiworld.hpp>
```

#### **Public Member Functions**

- Tools (Agent < TSeq > &p)
- std::vector< ToolPtr< TSeq > >::iterator begin ()
- std::vector< ToolPtr< TSeq > >::iterator end ()
- ToolPtr< TSeq > & operator() (size t i)
- ToolPtr< TSeq > & operator[] (size\_t i)
- size\_t size () const noexcept
- · void print () const noexcept

## Friends

- class Tool < TSeq >
- class Agent < TSeq >

## 18.79.1 Detailed Description

```
\label{template} \mbox{typename TSeq} > \\ \mbox{class epiworld::Tools} < \mbox{TSeq} > \\
```

Set of tools (useful for building iterators)

**Template Parameters** 

TSeq	
,	

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

· epiworld.hpp

# 18.80 Tools < TSeq > Class Template Reference

Set of tools (useful for building iterators)

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

## **Public Member Functions**

- Tools (Agent < TSeq > &p)
- std::vector< ToolPtr< TSeq > >::iterator **begin** ()
- std::vector < ToolPtr < TSeq > >::iterator end ()
- ToolPtr< TSeq > & operator() (size t i)
- ToolPtr< TSeq > & operator[] (size\_t i)
- size\_t size () const noexcept
- · void print () const noexcept

## **Friends**

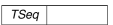
- class Tool < TSeq >
- class Agent < TSeq >

## 18.80.1 Detailed Description

template<typename TSeq> class Tools< TSeq>

Set of tools (useful for building iterators)

**Template Parameters** 



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

- · include/epiworld/agent-bones.hpp
- include/epiworld/tools-bones.hpp

# 18.81 epiworld::Tools\_const< TSeq > Class Template Reference

Set of Tools (const) (useful for iterators)

```
#include <epiworld.hpp>
```

#### **Public Member Functions**

- Tools const (const Agent < TSeq > &p)
- std::vector< ToolPtr< TSeq > >::const\_iterator begin () const
- $std::vector < ToolPtr < TSeq > >::const_iterator end () const$
- const ToolPtr < TSeq > & operator() (size\_t i)
- const ToolPtr< TSeq > & operator[] (size\_t i)
- size\_t size () const noexcept
- · void print () const noexcept

## **Friends**

- class Tool < TSeq >
- class Agent < TSeq >

## 18.81.1 Detailed Description

template<typename TSeq>
class epiworld::Tools\_const< TSeq >

Set of Tools (const) (useful for iterators)

**Template Parameters** 



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

· epiworld.hpp

# ${\bf 18.82 \quad Tools\_const} < {\bf TSeq} > {\bf Class\ Template\ Reference}$

Set of Tools (const) (useful for iterators)

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

#### **Public Member Functions**

- Tools\_const (const Agent < TSeq > &p)
- std::vector< ToolPtr< TSeq > >::const\_iterator begin () const
- std::vector< ToolPtr< TSeq > >::const\_iterator end () const
- const ToolPtr< TSeq > & operator() (size\_t i)
- const ToolPtr< TSeq > & operator[] (size\_t i)
- · size\_t size () const noexcept
- · void print () const noexcept

#### **Friends**

- class Tool < TSeq >
- class Agent < TSeq >

## 18.82.1 Detailed Description

template<typename TSeq> class Tools\_const< TSeq>

Set of Tools (const) (useful for iterators)

**Template Parameters** 



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

- · include/epiworld/agent-bones.hpp
- include/epiworld/tools-bones.hpp

# 18.83 epiworld::UserData < TSeq > Class Template Reference

Personalized data by the user.

```
#include <epiworld.hpp>
```

#### **Public Member Functions**

- UserData (Model < TSeq > &m)
- UserData (Model < TSeq > \*m)
- 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< epiworld\_double > \*data=nullptr)

- epiworld\_fast\_uint nrow () const
- epiworld\_fast\_uint 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** (epiworld\_fast\_uint j, epiworld\_double x)

#### Access data

#### **Parameters**

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

#### Returns

epiworld double&

- epiworld\_double & operator() (epiworld\_fast\_uint i, epiworld\_fast\_uint j)
- epiworld double & operator() (epiworld fast uint i, std::string name)

## **Friends**

- class Model < TSeq >
- class  ${\bf DataBase}{<}{\,{\sf TSeq}}{\,>}$

## 18.83.1 Detailed Description

template<typename TSeq> class epiworld::UserData< TSeq>

Personalized data by the user.

**Template Parameters** 

TSeq	

## 18.83.2 Constructor & Destructor Documentation

#### 18.83.2.1 UserData()

Construct a new User Data object.

#### **Parameters**

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

· epiworld.hpp

## 18.84 UserData < TSeq > Class Template Reference

Personalized data by the user.

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

## **Public Member Functions**

- UserData (Model < TSeq > &m)
- UserData (Model < TSeq > \*m)
- 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< epiworld\_double > \*data=nullptr)
- · epiworld\_fast\_uint nrow () const
- epiworld\_fast\_uint **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 (epiworld\_fast\_uint j, epiworld\_double x)

#### Access data

#### **Parameters**

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

## Returns

epiworld\_double&

- epiworld\_double & operator() (epiworld\_fast\_uint i, epiworld\_fast\_uint j)
- epiworld\_double & **operator()** (epiworld\_fast\_uint i, std::string name)

## **Friends**

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

## 18.84.1 Detailed Description

```
template<typename TSeq> class UserData< TSeq>
```

Personalized data by the user.

**Template Parameters** 

```
TSeq
```

## 18.84.2 Constructor & Destructor Documentation

## 18.84.2.1 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

# 18.85 epiworld::vecHasher< T > Struct Template Reference

Vector hasher.
#include <epiworld.hpp>

## **Public Member Functions**

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

## 18.85.1 Detailed Description

 $\label{eq:typename} \begin{array}{l} \text{template}{<} \text{typename T}{>} \\ \text{struct epiworld::vecHasher}{<} \text{ T}{>} \\ \end{array}$ 

Vector hasher.

**Template Parameters** 



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

· epiworld.hpp

# **18.86** vecHasher< T > Struct Template Reference

Vector hasher.

#include <misc.hpp>

## **Public Member Functions**

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

## 18.86.1 Detailed Description

$$\label{template} \begin{split} \text{template} &< \text{typename T}> \\ \text{struct vecHasher} &< \text{T}> \end{split}$$

Vector hasher.

## **Template Parameters**



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

· include/epiworld/misc.hpp

# 18.87 epiworld::Virus < TSeq > Class Template Reference

#### Virus.

```
#include <epiworld.hpp>
```

## **Public Member Functions**

- Virus (std::string name="unknown virus")
- Virus (std::string name, epiworld\_double prevalence, bool as\_proportion)
- void mutate (Model < TSeq > \*model)
- void set\_mutation (MutFun< TSeq > fun)
- EPI\_TYPENAME\_TRAITS (TSeq, int) get\_sequence()
- void **set\_sequence** (TSeq sequence)
- Agent < TSeq > \* get\_agent ()
- void set\_agent (Agent < TSeq > \*p)
- 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
- bool **operator==** (const Virus< TSeq > &other) const
- bool operator!= (const Virus < TSeq > &other) const
- void **print** () const

#### 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 (Model< TSeq > \*model)
- epiworld\_double get\_prob\_recovery (Model < TSeq > \*model)
- epiworld\_double get\_prob\_death (Model < TSeq > \*model)
- epiworld\_double get\_incubation (Model< TSeq > \*model)
- void post\_recovery (Model < TSeq > \*model)

- 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 incubation fun (VirusFun< TSeq > fun)
- void set\_prob\_infecting (const epiworld\_double \*prob)
- void set prob recovery (const epiworld double \*prob)
- void set prob death (const epiworld double \*prob)
- void set\_incubation (const epiworld\_double \*prob)
- void set\_prob\_infecting (epiworld\_double prob)
- void set\_prob\_recovery (epiworld\_double prob)
- void set\_prob\_death (epiworld\_double prob)
- void set\_incubation (epiworld\_double prob)

#### Get and set the state and queue

After applied, viruses can change the state and affect the queue of agents. These function sets the default values, which are retrieved when adding or removing a virus does not specify a change in state or in queue.

#### Parameters

init	After the virus/tool is added to the agent.
end	After the virus/tool is removed.
removed	After the agent (Agent) is removed.

- void set state (epiworld fast int init, epiworld fast int end, epiworld fast int removed=-99)
- void set\_queue (epiworld\_fast\_int init, epiworld\_fast\_int end, epiworld\_fast\_int removed=-99)
- void get\_state (epiworld\_fast\_int \*init, epiworld\_fast\_int \*end, epiworld\_fast\_int \*removed=nullptr)
- void **get\_queue** (epiworld\_fast\_int \*init, epiworld\_fast\_int \*end, epiworld\_fast\_int \*removed=nullptr)
- void distribute (Model < TSeq > \*model)

Get information about the prevalence of the virus.

void set\_distribution (VirusToAgentFun < TSeq > fun)

#### **Friends**

- class Agent < TSeq >
- class  $\mathbf{Model} < \mathbf{TSeq} >$
- class DataBase < TSeq >
- void default\_add\_virus (Event< TSeq > &a, Model< TSeq > \*m)
- void default\_rm\_virus (Event< TSeq > &a, Model< TSeq > \*m)

## 18.87.1 Detailed Description

template<typename TSeq> class epiworld::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 agent.

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

· epiworld.hpp

# 18.88 Virus < TSeq > Class Template Reference

#### Virus.

#include <virus-bones.hpp>

#### **Public Member Functions**

- Virus (std::string name="unknown virus")
- Virus (std::string name, epiworld\_double prevalence, bool as\_proportion)
- void mutate (Model < TSeq > \*model)
- void set\_mutation (MutFun< TSeq > fun)
- EPI\_TYPENAME\_TRAITS (TSeq, int) get\_sequence()
- void **set\_sequence** (TSeq sequence)
- Agent < TSeq > \* get\_agent ()
- void set\_agent (Agent < TSeq > \*p)
- 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
- bool operator== (const Virus< TSeq > &other) const
- bool operator!= (const Virus < TSeq > &other) const
- · void print () const
- void **set\_sequence** (int sequence)
- bool operator== (const Virus< std::vector< int >> &other) const
- void set\_sequence (int sequence)
- bool operator== (const Virus< std::vector< int >> &other) const

## 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 (Model< TSeq > \*model)
- epiworld\_double get\_prob\_recovery (Model < TSeq > \*model)
- epiworld\_double get\_prob\_death (Model < TSeq > \*model)
- epiworld\_double get\_incubation (Model < TSeq > \*model)
- void post\_recovery (Model < TSeq > \*model)
- 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\_incubation\_fun (VirusFun < TSeq > fun)
- void set\_prob\_infecting (const epiworld\_double \*prob)
- void set\_prob\_recovery (const epiworld\_double \*prob)
- void set\_prob\_death (const epiworld\_double \*prob)
- void set\_incubation (const epiworld\_double \*prob)
- void set\_prob\_infecting (epiworld\_double prob)
- void set\_prob\_recovery (epiworld\_double prob)
- void set\_prob\_death (epiworld\_double prob)
- void set\_incubation (epiworld\_double prob)

#### Get and set the state and queue

After applied, viruses can change the state and affect the queue of agents. These function sets the default values, which are retrieved when adding or removing a virus does not specify a change in state or in queue.

#### **Parameters**

init	After the virus/tool is added to the agent.
end	After the virus/tool is removed.
removed	After the agent (Agent) is removed.

- void set\_state (epiworld\_fast\_int init, epiworld\_fast\_int end, epiworld\_fast\_int removed=-99)
- · void set queue (epiworld fast int init, epiworld fast int end, epiworld fast int removed=-99)
- void get\_state (epiworld\_fast\_int \*init, epiworld\_fast\_int \*end, epiworld\_fast\_int \*removed=nullptr)
- void get\_queue (epiworld\_fast\_int \*init, epiworld\_fast\_int \*end, epiworld\_fast\_int \*removed=nullptr)
- void distribute (Model < TSeq > \*model)

Get information about the prevalence of the virus.

• void **set distribution** (VirusToAgentFun< TSeq > fun)

#### **Friends**

- class Agent < TSeq >
- class Model < TSeq >
- class DataBase < TSeq >
- void default\_add\_virus (Event< TSeq > &a, Model< TSeq > \*m)
- void default\_rm\_virus (Event< TSeq > &a, Model< TSeq > \*m)

## 18.88.1 Detailed Description

template<typename TSeq> class Virus< TSeq>

#### Virus.

#### **Template Parameters**



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 agent.

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

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

## 18.89 epiworld::Viruses < TSeq > Class Template Reference

Set of viruses (useful for building iterators)

#include <epiworld.hpp>

#### **Public Member Functions**

- Viruses (Agent < TSeq > &p)
- std::vector< VirusPtr< TSeq > >::iterator begin ()
- std::vector< VirusPtr< TSeq > >::iterator end ()
- VirusPtr< TSeq > & operator() (size\_t i)
- VirusPtr< TSeq > & operator[] (size\_t i)
- size\_t size () const noexcept
- · void print () const noexcept

## **Friends**

- class Virus < TSeq >
- class Agent < TSeq >

## 18.89.1 Detailed Description

template<typename TSeq> class epiworld::Viruses< TSeq>

Set of viruses (useful for building iterators)

**Template Parameters** 

TSeq	
,	

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

· epiworld.hpp

# 18.90 Viruses < TSeq > Class Template Reference

Set of viruses (useful for building iterators)

#include <viruses-bones.hpp>

## **Public Member Functions**

- Viruses (Agent < TSeq > &p)
- std::vector< VirusPtr< TSeq > >::iterator begin ()
- std::vector< VirusPtr< TSeq > >::iterator end ()
- VirusPtr< TSeq > & operator() (size t i)
- VirusPtr< TSeq > & operator[] (size\_t i)
- size\_t size () const noexcept
- · void print () const noexcept

## **Friends**

- class Virus < TSeq >
- class Agent < TSeq >

## 18.90.1 Detailed Description

template<typename TSeq> class Viruses< TSeq>

Set of viruses (useful for building iterators)

**Template Parameters** 



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

- include/epiworld/agent-bones.hpp
- include/epiworld/viruses-bones.hpp

# 18.91 epiworld::Viruses\_const< TSeq > Class Template Reference

Set of Viruses (const) (useful for iterators)

```
#include <epiworld.hpp>
```

#### **Public Member Functions**

- Viruses\_const (const Agent < TSeq > &p)
- std::vector< VirusPtr< TSeq > >::const\_iterator begin () const
- std::vector< VirusPtr< TSeq >>::const\_iterator end () const
- const VirusPtr< TSeq > & operator() (size\_t i)
- const VirusPtr< TSeq > & operator[] (size\_t i)
- size\_t size () const noexcept
- · void print () const noexcept

## **Friends**

- class Virus < TSeq >
- class Agent < TSeq >

## 18.91.1 Detailed Description

template<typename TSeq>
class epiworld::Viruses\_const< TSeq>

Set of Viruses (const) (useful for iterators)

**Template Parameters** 



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

· epiworld.hpp

# 18.92 Viruses\_const < TSeq > Class Template Reference

Set of Viruses (const) (useful for iterators)

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

#### **Public Member Functions**

- Viruses\_const (const Agent < TSeq > &p)
- std::vector< VirusPtr< TSeq > >::const\_iterator begin () const
- std::vector< VirusPtr< TSeq > >::const\_iterator end () const
- const VirusPtr< TSeq > & operator() (size\_t i)
- const VirusPtr< TSeq > & operator[] (size\_t i)
- · size\_t size () const noexcept
- · void print () const noexcept

#### **Friends**

- class Virus < TSeq >
- class Agent < TSeq >

## 18.92.1 Detailed Description

template<typename TSeq> class Viruses\_const< TSeq >

Set of Viruses (const) (useful for iterators)

**Template Parameters** 



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

- · include/epiworld/agent-bones.hpp
- · include/epiworld/viruses-bones.hpp

# 18.93 epiworld::VirusFunctions < TSeq > Class Template Reference

## **Public Attributes**

- MutFun< TSeq > mutation = nullptr
- PostRecoveryFun< TSeq > post\_recovery = nullptr
- VirusFun< TSeq > probability\_of\_infecting = nullptr
- $\bullet \quad \text{VirusFun} < \text{TSeq} > \textbf{probability\_of\_recovery} = \text{nullptr}$
- VirusFun< TSeq > probability\_of\_death = nullptr
- VirusFun< TSeq > incubation = nullptr
- VirusToAgentFun< TSeq > dist = nullptr

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

· epiworld.hpp

# 18.94 VirusFunctions < TSeq > Class Template Reference

## **Public Attributes**

- MutFun< TSeq > mutation = nullptr
- PostRecoveryFun< TSeq > post\_recovery = nullptr
- VirusFun< TSeq > probability\_of\_infecting = nullptr
- VirusFun< TSeq > probability\_of\_recovery = nullptr
- VirusFun< TSeq > probability\_of\_death = nullptr
- VirusFun< TSeq > incubation = nullptr
- VirusToAgentFun< TSeq > dist = nullptr

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

• include/epiworld/virus-bones.hpp

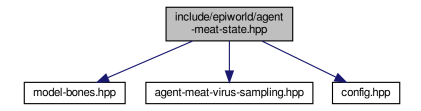
# **Chapter 19**

# **File Documentation**

# 19.1 include/epiworld/agent-meat-state.hpp File Reference

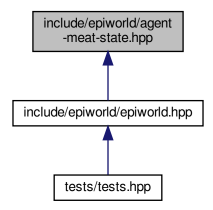
Sampling functions are getting big, so we keep them in a separate file.

```
#include "model-bones.hpp"
#include "agent-meat-virus-sampling.hpp"
#include "config.hpp"
Include dependency graph for agent-meat-state.hpp:
```



294 File Documentation

This graph shows which files directly or indirectly include this file:



## **Functions**

- template<typename TSeq = EPI\_DEFAULT\_TSEQ>
   void default\_update\_susceptible (Agent< TSeq > \*p, Model< TSeq > \*m)
- template<typename TSeq = EPI\_DEFAULT\_TSEQ> void **default\_update\_exposed** (Agent< TSeq > \*p, Model< TSeq > \*m)

## 19.1.1 Detailed Description

Sampling functions are getting big, so we keep them in a separate file.

**Author** 

George G. Vega Yon (g.vegayon en gmail)

Version

0.1

Date

2022-06-15

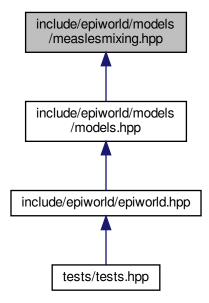
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# 19.2 include/epiworld/models/measlesmixing.hpp File Reference

Template for a Measles model with population mixing, quarantine, and contact tracing.

This graph shows which files directly or indirectly include this file:



#### Classes

class ModelMeaslesMixing
 TSeq >

Measles model with population mixing, quarantine, and contact tracing.

#### **Macros**

- #define MM(i, j, n) j \* n + i
- #define **GET\_MODEL**(model, output)
- #define SAMPLE\_FROM\_PROBS(n, ans)

## 19.2.1 Detailed Description

Template for a Measles model with population mixing, quarantine, and contact tracing.

## 19.2.2 Macro Definition Documentation

296 File Documentation

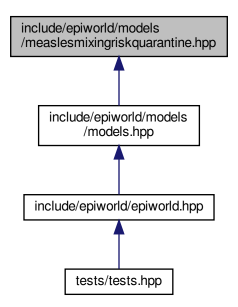
## 19.2.2.1 GET\_MODEL

## 19.2.2.2 SAMPLE\_FROM\_PROBS

# 19.3 include/epiworld/models/measlesmixingriskquarantine.hpp File Reference

Template for a Measles model with population mixing and risk-based quarantine.

This graph shows which files directly or indirectly include this file:



## **Classes**

class ModelMeaslesMixingRiskQuarantine< TSeq >

Measles model with population mixing and risk-based quarantine strategies.

## **Macros**

- #define MM(i, j, n) j \* n + i
- #define **GET\_MODEL**(model, output)
- #define SAMPLE\_FROM\_PROBS(n, ans)

Macro to sample from a list of probabilities.

## 19.3.1 Detailed Description

Template for a Measles model with population mixing and risk-based quarantine.

## 19.3.2 Macro Definition Documentation

## 19.3.2.1 GET\_MODEL

#### 19.3.2.2 SAMPLE\_FROM\_PROBS

#define SAMPLE\_FROM\_PROBS(

Macro to sample from a list of probabilities.

## Returns

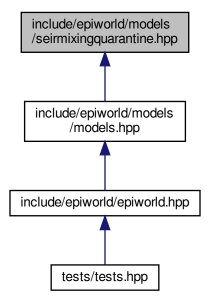
The index of the sampled probability; and the total length if none is found, returns n.

298 File Documentation

# 19.4 include/epiworld/models/seirmixingquarantine.hpp File Reference

Template for a Susceptible-Exposed-Infected-Removed (SEIR) model with mixing, quarantine, and contact tracing.

This graph shows which files directly or indirectly include this file:



#### **Classes**

 class ModelSEIRMixingQuarantine < TSeq >
 SEIR model with mixing, quarantine, and contact tracing.

#### **Macros**

- #define MM(i, j, n) j \* n + i
- #define **GET\_MODEL**(model, output)
- #define SAMPLE\_FROM\_PROBS(n, ans)

## 19.4.1 Detailed Description

Template for a Susceptible-Exposed-Infected-Removed (SEIR) model with mixing, quarantine, and contact tracing.

## 19.4.2 Macro Definition Documentation

## 19.4.2.1 GET\_MODEL

## 19.4.2.2 SAMPLE\_FROM\_PROBS

300 File Documentation

# Index

```
add globalevent
  ModelSIRCONN < TSeq >, 229
    epiworld::Model < TSeq >, 103
  ModelSIRDCONN < TSeq >, 240
    Model < TSeq >, 119
  ModelSIRLogit < TSeq >, 247
AdjList, 55
  ModelSIRMixing < TSeq >, 254
    AdjList, 55
  DataBase < TSeq >, 69
    epiworld::AdjList, 57
  get generation time, 72
    read edgelist, 56
  get_reproductive_number, 72
Agent < TSeq >, 58
  get_transition_probability, 72
    default_rm_entity, 62
  get transmissions, 73
    operator(), 60
  operator==, 73, 74
    swap neighbors, 62
  record virus, 74
AgentsSample
  default rm entity
    AgentsSample < TSeq >, 67
  Agent < TSeq >, 62
    epiworld::AgentsSample < TSeq >, 69
  Entity < TSeq >, 84
AgentsSample < TSeq >, 66
  epiworld::Agent < TSeq >, 66
    AgentsSample, 67
  epiworld::Entity< TSeq >, 85
clone ptr
  draw
    epiworld::epimodels::ModelMeaslesMixing< TSeq
  epiworld::Model < TSeq >, 104
         >, 136
  Model < TSeq >, 119
    epiworld::epimodels::ModelMeaslesMixingRiskQuarantine<
  Entities < TSeq >, 79
         TSeq >, 153
   Entities_const< TSeq >, 81
    epiworld::epimodels::ModelMeaslesSchool< TSeq
   Entity
         >, 166
   Entity < TSeq >, 83
    epiworld::epimodels::ModelSEIRCONN< TSeq >,
  epiworld::Entity< TSeq >, 85
  Entity < TSeq >, 83
    epiworld::epimodels::ModelSEIRDCONN< TSeq
  default rm entity, 84
         >, 192
  Entity, 83
    epiworld::epimodels::ModelSEIRMixing< TSeq >,
  epiworld::AdjList, 56
         199
    epiworld::epimodels::ModelSEIRMixingQuarantine<
  AdjList, 57
  read edgelist, 57
         TSeq >, 209
  epiworld::Agent < TSeq >, 63
    epiworld::epimodels::ModelSIRCONN< TSeq >,
  default_rm_entity, 66
         225
  operator(), 65
    epiworld::epimodels::ModelSIRDCONN< TSeq >,
  swap neighbors, 65
   epiworld::AgentsSample < TSeq >, 68
    epiworld::epimodels::ModelSIRLogit< TSeq
  AgentsSample, 69
   epiworld::DataBase< TSeq >, 75
    epiworld::epimodels::ModelSIRMixing< TSeq >,
  get generation time, 77
         250
  get_reproductive_number, 77
    epiworld::Model < TSeq >, 103
    Model < TSeq >, 119
  get_transition_probability, 78
  get transmissions, 78
    ModelMeaslesMixing< TSeq >, 144
  operator==, 79
    ModelMeaslesMixingRiskQuarantine< TSeq >,
  record_virus, 79
  epiworld::Entities < TSeq >, 80
    ModelMeaslesSchool < TSeq >, 171
  epiworld::Entities const< TSeq >, 82
    ModelSEIRCONN < TSeq >, 182
  epiworld::Entity< TSeq >, 84
    ModelSEIRDCONN < TSeq >, 195
  default_rm_entity, 85
    ModelSEIRMixing < TSeq >, 202
  Entity, 85
    ModelSEIRMixingQuarantine < TSeq >, 217
```

epiworld::epimodels::ModelDiffNet< TSeq >, 127 epiworld::epimodels::ModelMeaslesMixing< TSeq >,	get_isolation_willingness, 210 get_quarantine_willingness, 210
130	initial states, 210
clone_ptr, 136	ModelSEIRMixingQuarantine, 206, 208
get_agent_quarantine_triggered, 136	run, 211
get_contact_matrix, 136	set_contact_matrix, 211
get_isolation_willingness, 137	epiworld::epimodels::ModelSIR< TSeq >, 220
get_quarantine_willingness, 137	initial_states, 221
initial_states, 137	epiworld::epimodels::ModelSIRCONN< TSeq >, 224
ModelMeaslesMixing, 133, 135	clone_ptr, 225
run, 138	get_n_infected, 226
set_contact_matrix, 138	initial_states, 226
epiworld::epimodels::ModelMeaslesMixingRiskQuarantine	
TSeq >, 147	reset, 226
clone_ptr, 153	epiworld::epimodels::ModelSIRD< TSeq >, 230
get_agent_quarantine_triggered, 153	initial_states, 233
get_contact_matrix, 153	ModelSIRD, 231
get_isolation_willingness, 153	epiworld::epimodels::ModelSIRDCONN< TSeq >, 236
get_quarantine_risk_levels, 154	clone ptr, 238
get quarantine willingness, 154	ModelSIRDCONN, 237
initial states, 154	reset, 238
ModelMeaslesMixingRiskQuarantine, 150, 151	epiworld::epimodels::ModelSIRLogit< TSeq >, 241
run, 155	clone_ptr, 244
set_contact_matrix, 155	ModelSIRLogit, 243
epiworld::epimodels::ModelMeaslesSchool< TSeq >,	reset, 244
163	epiworld::epimodels::ModelSIRMixing< TSeq >, 248
clone_ptr, 166	clone_ptr, 250
ModelMeaslesSchool, 165	initial_states, 251
quarantine_agents, 167	ModelSIRMixing, 249, 250
reset, 167	reset, 251
epiworld::epimodels::ModelSEIR< TSeq >, 172	epiworld::epimodels::ModelSIS< TSeq >, 255
initial_states, 174	epiworld::epimodels::ModelSISD< TSeq >, 259
update_exposed_seir, 174	epiworld::epimodels::ModelSURV< TSeq >, 262
update_infected_seir, 174	reset, 264
epiworld::epimodels::ModelSEIRCONN< TSeq >, 178	epiworld::Event< TSeq >, 86
clone_ptr, 179	Event, 87
initial_states, 180	epiworld::GlobalEvent< TSeq >, 89
ModelSEIRCONN, 179	GlobalEvent, 90
reset, 180	epiworld::LFMCMC< TData >, 92
epiworld::epimodels::ModelSEIRD< TSeq >, 184	epiworld::Model< TSeq >, 95
ModelSEIRD, 185, 186	add globalevent, 103
update_exposed_seir, 186	clone_ptr, 103
epiworld::epimodels::ModelSEIRDCONN< TSeq >,	draw, 104
190	events_add, 104
clone_ptr, 192	events_run, 105
initial_states, 192	initial_states_fun, 108
ModelSEIRDCONN, 191	load_agents_entities_ties, 105
reset, 192	rbinomd, 108
epiworld::epimodels::ModelSEIRMixing< TSeq >, 196	reset, 106
clone_ptr, 199	rexpd, 108
initial_states, 199	rgammad, 108
ModelSEIRMixing, 197, 198	rgeomd, 109
reset, 199	rlognormald, 109
epiworld::epimodels::ModelSEIRMixingQuarantine<	rnbinomd, 109
TSeq >, 204	rnormd, 109
clone_ptr, 209	rpoissd, 109
get_agent_quarantine_triggered, 209	run_multiple, 106
get_contact_matrix, 209	runifd, 110

set_agents_data, 106	get_isolation_willingness
set_name, 107	epiworld::epimodels::ModelMeaslesMixing< TSeq
time_elapsed, 110	>, 137
write_data, 107	epiworld::epimodels::ModelMeaslesMixingRiskQuarantine<
epiworld::ModelDiagram, 127	TSeq >, 153
epiworld::PersonTools< TSeq >, 267	epiworld::epimodels::ModelSEIRMixingQuarantine<
epiworld::Progress, 268	TSeq >, 210
epiworld::Queue< TSeq >, 269	ModelMeaslesMixing< TSeq >, 145
epiworld::sampler, 47	ModelMeaslesMixingRiskQuarantine< TSeq >,
make_sample_virus_neighbors, 47	161
make_update_susceptible, 48	ModelSEIRMixingQuarantine< TSeq >, 218
sample_virus_single, 48	GET_MODEL
epiworld::SAMPLETYPE, 271	measlesmixing.hpp, 295
epiworld::Tool< TSeq >, 272	measlesmixingriskquarantine.hpp, 297
epiworld::ToolFunctions< TSeq >, 275	seirmixingquarantine.hpp, 298
epiworld::Tools< TSeq >, 276	get_n_infected
epiworld::Tools_const< TSeq >, 278	• — —
epiworld::UserData < TSeq >, 279	epiworld::epimodels::ModelSIRCONN< TSeq >, 226
·	ModelSIRCONN< TSeq >, 229
UserData, 281	• •
epiworld::vecHasher< T >, 283	get_quarantine_risk_levels
epiworld::Virus< TSeq >, 284	epiworld::epimodels::ModelMeaslesMixingRiskQuarantine<
epiworld::Viruses< TSeq >, 288	TSeq >, 154
epiworld::Viruses_const< TSeq >, 290	get_quarantine_willingness
epiworld::VirusFunctions< TSeq >, 291	epiworld::epimodels::ModelMeaslesMixing< TSeq
Event	>, 137
epiworld::Event< TSeq >, 87	epiworld::epimodels::ModelMeaslesMixingRiskQuarantine<
Event< TSeq >, 89	TSeq >, 154
Event< TSeq >, 87	epiworld::epimodels::ModelSEIRMixingQuarantine<
Event, 89	TSeq >, 210
events_add	ModelMeaslesMixing < TSeq >, 145
epiworld::Model < TSeq >, 104	ModelMeaslesMixingRiskQuarantine< TSeq >,
Model < TSeq >, 120	161
events_run	ModelSEIRMixingQuarantine < TSeq >, 218
epiworld::Model< TSeq >, 105	get_reproductive_number
Model < TSeq >, 120	DataBase< TSeq >, 72
get agent guarantine triggered	epiworld::DataBase< TSeq >, 77
get_agent_quarantine_triggered	get_transition_probability
epiworld::epimodels::ModelMeaslesMixing< TSeq	DataBase < TSeq >, 72
>, 136	epiworld::DataBase< TSeq >, 78
epiworld::epimodels::ModelMeaslesMixingRiskQuara	
TSeq >, 153	DataBase< TSeq >, 73
epiworld::epimodels::ModelSEIRMixingQuarantine<	epiworld::DataBase< TSeq >, 78
TSeq >, 209	GlobalEvent
ModelMeaslesMixing< TSeq >, 145	epiworld::GlobalEvent $<$ TSeq $>$ , 90
ModelSEIRMixingQuarantine < TSeq >, 217	GlobalEvent< TSeq >, 91
get_contact_matrix	GlobalEvent< TSeq >, 91
epiworld::epimodels::ModelMeaslesMixing< TSeq >, 136	GlobalEvent, 91
epiworld::epimodels::ModelMeaslesMixingRiskQuara	• • •
TSeq >, 153	include/epiworld/models/measlesmixing.hpp, 295
epiworld::epimodels::ModelSEIRMixingQuarantine< TSeq >, 209	include/epiworld/models/measlesmixingriskquarantine.hpp,
ModelMeaslesMixing< TSeq >, 145	include/epiworld/models/seirmixingquarantine.hpp, 298
ModelMeaslesMixingRiskQuarantine< TSeq >,	initial_states
161	epiworld::epimodels::ModelMeaslesMixing< TSeq
ModelSEIRMixingQuarantine< TSeq >, 217	>, 137
get_generation_time	epiworld::epimodels::ModelMeaslesMixingRiskQuarantine<
DataBase< TSeq >, 72	TSeq >, 154
epiworld::DataBase< TSeq >, 77	epiworld::epimodels::ModelSEIR< TSeq >, 174

epiworld::epimodels::ModelSEIRCONN< TSeq >,	rgeomd, 125 rlognormald, 125
epiworld::epimodels::ModelSEIRDCONN< TSeq	rnbinomd, 125
>, 192	rnormd, 125
epiworld::epimodels::ModelSEIRMixing< TSeq >,	rpoissd, 126
199	run_multiple, 122
epiworld::epimodels::ModelSEIRMixingQuarantine<	runifd, 126
TSeq >, 210	set_agents_data, 123
epiworld::epimodels::ModelSIR< TSeq >, 221	set_name, 123
epiworld::epimodels::ModelSIRCONN< TSeq >,	time elapsed, 126
226	write_data, 124
epiworld::epimodels::ModelSIRD< TSeq >, 233	ModelDiagram, 127
epiworld::epimodels::ModelSIRMixing< TSeq >,	ModelDiffNet< TSeq >, 129
251	ModelMeaslesMixing
ModelMeaslesMixing < TSeq >, 146	_
ModelMeaslesMixingRiskQuarantine< TSeq >,	epiworld::epimodels::ModelMeaslesMixing< TSeq
162	>, 133, 135
	ModelMeaslesMixing < TSeq >, 142, 143
ModelSEIR < TSeq >, 176	ModelMeaslesMixing < TSeq >, 139
ModelSEIRCONN < TSeq >, 183	clone_ptr, 144
ModelSEIRDCONN < TSeq >, 195	get_agent_quarantine_triggered, 145
ModelSEIRMixing < TSeq >, 203	get_contact_matrix, 145
ModelSEIRMixingQuarantine < TSeq >, 218	get_isolation_willingness, 145
ModelSIR< TSeq >, 223	get_quarantine_willingness, 145
ModelSIRCONN< TSeq >, 229	initial_states, 146
ModelSIRD< TSeq >, 235	ModelMeaslesMixing, 142, 143
ModelSIRMixing< TSeq >, 254	run, 146
initial_states_fun	set_contact_matrix, 147
epiworld::Model < TSeq >, 108	ModelMeaslesMixingRiskQuarantine
Model < TSeq >, 124	epiworld::epimodels::ModelMeaslesMixingRiskQuarantine< TSeq >, 150, 151
LFMCMC< TData >, 93	ModelMeaslesMixingRiskQuarantine< TSeq >,
load_agents_entities_ties	158, 159
epiworld::Model < TSeq >, 105	ModelMeaslesMixingRiskQuarantine< TSeq >, 156
Model < TSeq >, 122	clone_ptr, 161
	get_contact_matrix, 161
make_sample_virus_neighbors	get_isolation_willingness, 161
epiworld::sampler, 47	get_quarantine_willingness, 161
sampler, 50	initial_states, 162
make_update_susceptible	ModelMeaslesMixingRiskQuarantine, 158, 159
epiworld::sampler, 48	run, 162
sampler, 51	set_contact_matrix, 163
measlesmixing.hpp	ModelMeaslesSchool
GET_MODEL, 295	epiworld::epimodels::ModelMeaslesSchool< TSeq
SAMPLE_FROM_PROBS, 296	>, 165
measlesmixingriskquarantine.hpp	ModelMeaslesSchool< TSeq >, 170
GET MODEL, 297	ModelMeaslesSchool < TSeq >, 170  ModelMeaslesSchool < TSeq >, 168
SAMPLE_FROM_PROBS, 297	•
Model < TSeq >, 110	clone_ptr, 171
add globalevent, 119	ModelMeaslesSchool, 170
clone_ptr, 119	quarantine_agents, 171
draw, 119	reset, 171
events_add, 120	ModelSEIR< TSeq >, 175
events_run, 120	initial_states, 176
initial_states_fun, 124	update_exposed_seir, 177
load_agents_entities_ties, 122	update_infected_seir, 177
_ <del>-</del>	ModelSEIRCONN
rbinomd, 124	epiworld::epimodels::ModelSEIRCONN< TSeq >,
reset, 122	179
rexpd, 124	ModelSEIRCONN< TSeq >, 182
rgammad, 125	

ModelSEIRCONN< TSeq >, 181	ModelSIRD< TSeq >, 235
clone_ptr, 182	ModelSIRD< TSeq >, 233
initial_states, 183	initial_states, 235
ModelSEIRCONN, 182	ModelSIRD, 235
reset, 183	ModelSIRDCONN
ModelSEIRD	epiworld::epimodels::ModelSIRDCONN< TSeq >,
epiworld::epimodels::ModelSEIRD< TSeq >, 185,	237
186	ModelSIRDCONN< TSeq >, 240
ModelSEIRD< TSeq >, 188, 189	ModelSIRDCONN< TSeq >, 239
ModelSEIRD< TSeq >, 187	clone ptr, 240
ModelSEIRD, 188, 189	ModelSIRDCONN, 240
update_exposed_seir, 189	reset, 241
ModelSEIRDCONN	ModelSIRLogit
epiworld::epimodels::ModelSEIRDCONN< TSeq	· II · II M LIGIDI II · TO
>, 191	epiworia::epimodels::ModelSIRLogit< 1Seq >,
ModelSEIRDCONN< TSeq >, 194	
•	ModelSIRLogit < TSeq >, 246
ModelSEIRDCONN < TSeq >, 193	ModelSIRLogit < TSeq >, 245
clone_ptr, 195	clone_ptr, 247
initial_states, 195	ModelSIRLogit, 246
ModelSEIRDCONN, 194	reset, 247
reset, 196	ModelSIRMixing
ModelSEIRMixing	epiworld::epimodels::ModelSIRMixing< TSeq >,
epiworld::epimodels::ModelSEIRMixing< TSeq >,	249, 250
197, 198	ModelSIRMixing< TSeq >, 253, 254
ModelSEIRMixing < TSeq >, 201, 202	ModelSIRMixing < TSeq >, 252
ModelSEIRMixing < TSeq >, 200	clone_ptr, 254
clone_ptr, 202	initial_states, 254
initial_states, 203	ModelSIRMixing, 253, 254
ModelSEIRMixing, 201, 202	reset, 255
reset, 203	ModelSIS < TSeq >, 258
ModelSEIRMixingQuarantine	ModelSISD< TSeq >, 261
epiworld::epimodels::ModelSEIRMixingQuarantine<	
TSeq >, 206, 208	reset, 266
ModelSEIRMixingQuarantine < TSeq >, 214, 216	
ModelSEIRMixingQuarantine < TSeq >, 212	Network< Nettype, Nodetype, Edgetype >, 267
clone_ptr, 217	
<del>_</del>	operator()
get_agent_quarantine_triggered, 217	Agent< TSeq >, 60
get_contact_matrix, 217	epiworld::Agent < TSeq >, 65
get_isolation_willingness, 218	operator==
get_quarantine_willingness, 218	DataBase < TSeq >, 73, 74
initial_states, 218	epiworld::DataBase< TSeq >, 79
ModelSEIRMixingQuarantine, 214, 216	opo
run, 219	PersonTools < TSeq >, 268
set_contact_matrix, 219	Progress, 268
ModelSIR< TSeq >, 221	
initial_states, 223	quarantine_agents
ModelSIRCONN	epiworld::epimodels::ModelMeaslesSchool< TSeq
epiworld::epimodels::ModelSIRCONN< TSeq >,	>, 167
225	ModelMeaslesSchool< TSeq >, 171
ModelSIRCONN< TSeq >, 228	Queue< TSeq >, 269
ModelSIRCONN< TSeq >, 227	44646 ( 1864 ) , 266
clone_ptr, 229	RandGraph, 270
get_n_infected, 229	rbinomd
initial_states, 229	epiworld::Model < TSeq >, 108
ModelSIRCONN, 228	Model < TSeq >, 124
reset, 230	read_edgelist
ModelSIRD	AdjList, 56
epiworld::epimodels::ModelSIRD< TSeq >, 231	epiworld::AdjList, 57
	epiwonaaujust, 37

record_virus	epiworld::epimodels::ModelSEIRMixingQuarantine<
DataBase < TSeq >, 74	TSeq >, 211
epiworld::DataBase< TSeq >, 79	ModelMeaslesMixing< TSeq >, 146
reset	${\sf ModelMeaslesMixingRiskQuarantine} < {\sf TSeq} >,$
epiworld::epimodels::ModelMeaslesSchool< TSeq	162
>, 167	ModelSEIRMixingQuarantine< TSeq >, 219
epiworld::epimodels::ModelSEIRCONN< TSeq >,	run_multiple
180	epiworld::Model < TSeq >, 106
epiworld::epimodels::ModelSEIRDCONN< TSeq	Model < TSeq >, 122
>, 192	runifd
epiworld::epimodels::ModelSEIRMixing< TSeq >, 199	epiworld::Model< TSeq >, 110 Model< TSeq >, 126
epiworld::epimodels::ModelSIRCONN< TSeq >,	,
226	SAMPLE_FROM_PROBS
epiworld::epimodels::ModelSIRDCONN< TSeq >,	measlesmixing.hpp, 296
238	measlesmixingriskquarantine.hpp, 297
epiworld::epimodels::ModelSIRLogit< TSeq >,	seirmixingquarantine.hpp, 299
244	sample_virus_single
epiworld::epimodels::ModelSIRMixing< TSeq >,	epiworld::sampler, 48
251	sampler, 51
epiworld::epimodels::ModelSURV< TSeq >, 264	sampler, 50
epiworld::Model < TSeq >, 106	make_sample_virus_neighbors, 50
Model < TSeq >, 122	make_update_susceptible, 51
ModelMeaslesSchool < TSeq >, 171	sample_virus_single, 51
ModelSEIRCONN< TSeq >, 183	SAMPLETYPE, 271
ModelSEIRDCONN < TSeq >, 196	seirmixingquarantine.hpp
ModelSEIRMixing < TSeq >, 203	GET_MODEL, 298 SAMPLE_FROM_PROBS, 299
ModelSIRCONN < TSeq >, 230	set_agents_data
ModelSIRDCONN< TSeq >, 241	epiworld::Model < TSeq >, 106
ModelSIRLogit < TSeq >, 247	Model < TSeq >, 123
ModelSTRMixing < TSeq >, 255	set_contact_matrix
ModelSURV < TSeq >, 266	epiworld::epimodels::ModelMeaslesMixing< TSeq
expd epiworld::Model < TSeq >, 108	>, 138
Model < TSeq >, 124	epiworld::epimodels::ModelMeaslesMixingRiskQuarantine<
gammad	TSeq >, 155
epiworld::Model < TSeq >, 108	epiworld::epimodels::ModelSEIRMixingQuarantine<
Model < TSeq >, 125	TSeq >, 211
geomd	ModelMeaslesMixing< TSeq >, 147
epiworld::Model< TSeq >, 109	ModelMeaslesMixingRiskQuarantine< TSeq >,
Model < TSeq >, 125	163
lognormald	ModelSEIRMixingQuarantine< TSeq >, 219
epiworld::Model < TSeq >, 109	set_name
Model < TSeq >, 125	epiworld::Model < TSeq >, 107
nbinomd	Model < TSeq >, 123
epiworld::Model < TSeq >, 109	SimpleAdjList, 271
Model < TSeq >, 125	SimpleModel, 272
normd	swap_neighbors
epiworld::Model < TSeq >, 109	Agent < TSeq >, 62
Model < TSeq >, 125	epiworld::Agent< TSeq >, 65
poissd	time planted
epiworld::Model < TSeq >, 109	time_elapsed epiworld::Model < TSeq >, 110
Model < TSeq >, 126	Model < TSeq >, 110
run	Tool < TSeq >, 120
epiworld::epimodels::ModelMeaslesMixing< TSeq	ToolFunctions < TSeq >, 276
>, 138	
epiworld::epimodels::ModelMeaslesMixingRiskQuara TSeq >, 155	Tools_const< TSeq >, 278

```
update_exposed_seir
     epiworld::epimodels::ModelSEIR< TSeq >, 174
     epiworld::epimodels::ModelSEIRD< TSeq>, 186
     \mathsf{ModelSEIR} {<\mathsf{TSeq}>},\, \mathsf{177}
     ModelSEIRD < TSeq >, 189
update infected seir
     epiworld::epimodels::ModelSEIR < TSeq >, 174
     ModelSEIR < TSeq >, 177
UserData
     epiworld::UserData < TSeq >, 281
     {\sf UserData} {< \mathsf{TSeq}>, \textcolor{red}{\textbf{282}}}
UserData < TSeq >, 281
     UserData, 282
vecHasher< T >, 283
Virus < TSeq >, 286
Viruses < TSeq >, {\color{red} 289}
Viruses_const< TSeq >, 290
VirusFunctions < TSeq >, 292
write_data
     epiworld::Model < TSeq >, 107
     Model < TSeq >, 124
```