An Agent-Based Model of COVID-19 Transmission

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Introduction

This model is a simple agent-based model of COVID-19 transmission in a population connected by social networks. The transmission of COVID-19 is modeled using a compartmental SEIR model in which individual agents progress through disease stages of susceptible, exposed, infected or recovered. Agents are connected to one another through non-spatial social networks which are modeled as Watts-Strogatz small world networks (?) or Barab'asi-Albert scale-invariant networks (??). The disease may be transmitted through these network links.

Model Overview

Purpose

The purpose of this model is to use agent-based modeling to simulate the spread of COVID-19 in communities. The model is used to simulate the effects of different types of network connectivity (including changes to the network connections through interventions such as "social isolation") on the spread of infections.

The model does not attempt to provide reliable predictions of the future spread of disease, but focuses on making comparisons between different patterns of social network connections.

Entities, State Variables, and Scales

There are two kinds of entities in this model: agents and links.

Agents

Agents represent individual people. Each agent is initialized with the following characteristics:

- age, which is represented as both a number and as a bracket.
- *sex*, a binary variable representing male or female sex phenotype. This model does not currently treat intersex individuals.
- *comorbidity*, a binary variable that represents whether the indivisual has a comorbidity that could complicate susceptibility to new infections.
- *symptomatic*, a binary variable that controls whether the individaul will be symptomatic if they are infected. This is important, because symptomatic individuals are reported to shed greater amounts of virus than asymptomatic ones. Future variations on this model may also account for behavioral differences (symptomatic individuals may be more likely to self-quarantine or seek medical attention, whereas asymptomatic ones may continue to interact with others and thus spread the disease)

- health status, an ordinal factor that represents the status of the individual within a four-compartment SEIR model, where S represents susceptible individuals, E represents exposed individuals who are incubating an infection but are not yet infectious, I represents infectious individuals who can spread the disease, and R represents recovered individuals who are no longer contagious and who have acquired immunity against being reinfected.
- ticks, an integer representing the number of time steps since the individual entered the current *health status*. This affects the progression from E to I and from *I* to *R*.
- *shedding factor*, a real number (positive or negative) that represents the relative degree of viral shedding (e.g., symptomatic individuals are reported to have higher shedding factors than asymptomatic ones).
- susceptibility factor, a real number (positive or negative) that represents the relative susceptibility of an individual to contracting an infection when exposed to the virus.
- EI scale, EI shape, IR scale, and IR shape, real numbers that parameterize progress of the disease. See further details under Submodels

Links

*Links** represent social connections. These can be *household* connections (people who live together), social connections of people who live in different households, but are connected through friendship, church, community groups, etc., and see one another regularly, and work connections.

Links have parameters:

- contact frequency, represents the mean frequency of this kind of contact, in number of meetings per time step.
- contact intensity, represents the intensity of close interaction (e.g., visits to a health-care professional will generally be more intense than interactions with cashiers at a store)

The characteristics of the network is described in greater detail under *collectives*, below.

Spatiotemporal scales

This model does not explicitly represent space, and agents interact through social networks rather than spatial proximity.

The time step represents one day. The number of ticks in a model run can be specified by the user.

Environment

Initially, we will not model the environment. We will have agents connected by social networks with no representation of the physical environments they inhabit. Later versions will incorporate infrastructure to represent connections via transportation, and spatial distributions of housing units and workplaces.

Process Overview and Scheduling

At each time step, two things happen:

- I. Each agent in the E or I state progresses stochastically toward the next state (I or R, respectively). The probability of progression may depend on the number of time steps the agent has been in this status. See the disease progression submodel for details.
- 2. Every agent in the I state stochastically infects its immediate neighbors on the network (those with whom it shares a link) who are in the S status. See the infect submodel for details.

Design Concepts

Basic Principles

The transmission of COVID-19 is represented using a four-compartment SEIR model. The SEIR model categorizes agents into susceptible, exposed, infectious or recovered.

Infection (transition from susceptible to exposed status) occurs stochastically. At each time step, every susceptible agent that is connected to an infectious agent by a network link has a probability of transitioning to an exposed state. The probability depends on the characteristics of the two agents and of the network link.

Agents are heterogeneous, so two agents with the same age, sex, etc. can have different shedding and susceptibility factors. These factors are drawn at random from probability distributions that are parameterized by the agent characteristics (age, sex, comorbidities, and symptomatic status)

Exposed and infectious agents stochastically transition to the next stage (infectious and recovered, respectively), with time-dependent probabilities that follow gamma or Weibull distributions. The parameters of these distributions may depend on the agent's age, sex, comorbidities, etc.

Emergence

The spread of the disease emerges from individual interactions on the network.

Adaptation

Currently, the agents do not adapt their behavior to changing conditions. Future versions may allow agents to change their social interactions when they get sick or in response to public policies, such as stay-at-home orders.

Objectives

The agents do not pursue objectives.

Learning

The agents do not learn.

Prediction

The agents do not engage in prediction.

Sensing

The agents do not currently use sensing. In the future, they may sense aspects of their own or other agents' health.

Interaction

Agents interact through links. These are how infections are transmitted.

Stochasticity

Agent initialization is stochastic with characteristics (age, sex, comorbidities, future symptomatic response to infection, shedding intensity, and susceptibility) drawn from distributions that can be specified at run-time.

Disease progress (E to I and I to R) are stochastic, with probabilities that vary with the amount of time an agent has been in that status.

Disease transmission is stochastic. Disease is transmitted across links that connect infectious to susceptible agents. The probability of transmission depends on the shedding intensity of the infectious agent, the susceptibility of the susceptible agent, and the contact characteristics of the link.

Collectives

Multiple overlapping social networks (household, social, and work) connect agents. These networks can have different topologies that are specified at runtime when the agents are initialized.

Currently available topologies are Strogatz-Watts *small-world* (?) and Barabasi Albert *preferential attachment* (??). The big difference between these is that the degree of connection is fairly uniformly distributed in the Strogatz Watts model, but is very unequally distributed in Barab'asi-Albert networks, with a few highly connected nodes that may be able to simulate super-spreaders.

Observation

At each time step we record the number of agents in each health status (S, E, I, or R).

Details

Initialization

Input Data

Submodels