The contagious nature of imprisonment: an agent-based model to explain racial disparities in incarceration rates

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We build an agent-based model of incarceration based on the susceptible–infected–susceptible (SIS) model of infectious disease propagation. Our central hypothesis is that the observed racial disparities in incarceration rates between Black and White Americans can be explained as the result of differential sentencing between the two demographic groups. We demonstrate that if incarceration can be spread through a social influence network, then even relatively small differences in sentencing can result in large disparities in incarceration rates. Controlling for effects of transmissibility, susceptibility and influence network structure, our model reproduces the observed large disparities in incarceration rates given the differences in sentence lengths for White and Black drug offenders in the USA without extensive parameter tuning. We further establish the suitability of the SIS model as applied to incarceration by demonstrating that the observed structural patterns of recidivism are an emergent property of the model. In fact, our model shows a remarkably close correspondence with California incarceration data. This work advances efforts to combine the theories and methods of epidemiology and criminology.

1. Introduction

The rapid increase in the US incarceration rate over the last few decades has been described as an epidemic. According to the Bureau of Justice Statistics, the per capita rate of incarceration nearly quadrupled between 1978 and 2011, from 137 to 511 persons per 100 000 [1]. This prison boom has primarily affected Black Americans, especially Black males. By 2011, Black incarceration rates were over six times higher than White rates (3023 per 100 000 for Blacks, and 478 per 100 000 for Whites).

Racial disparities in incarceration rates have been studied extensively [2–6], and while these disparities are partially due to differences in criminal involvement [7], the increase in imprisonment for Black males since 1980 was not matched by a similar increase in Black-male criminality [8–10]. What then accounts for the racial disparities in incarceration? Scholars offer several explanations, including differential exposure to police surveillance [9], prosecutorial discrimination [11], the use of incarceration to deal with a ‘racial threat’ [12,13] or sentencing disparities between Blacks and Whites. Although studies reveal that racial sentencing disparities are reduced when legal factors [14–16] or social contexts [17,18] are considered, a recent meta-analysis reports that sentencing disparities remain even after controlling for these factors [4]. While the magnitude of the difference is small and variable, it is largest in cases involving discretionary powers and for drug offences. In fact, it has been shown that Blacks receive longer sentences than Whites for drug offences [3,9].

Careful study of patterns of incarceration reveals that incarceration behaves like a contagious disease in that the close associates of an incarcerated person have higher-than-average probabilities of being incarcerated. An individual’s incarceration can be ‘transmitted’ to others via several mechanisms. First, an individual’s incarceration can increase the family members’ emotional and
economic stress, and cumulative strains are related to crimi-
nal behaviour [19]. Specifically, it has been shown that the
children of incarcerated parents tend to display increased
levels of behaviour issues [20], including aggressive behav-
our [21], a predictor of criminality in later life. Close family
members, such as the inmates’ domestic partners, experience
the acute effects of their family member’s incarceration.
Decreased household income due to the inmate’s inability
to work while incarcerated and the inability of the inmate
to contribute to child care responsibilities put the remaining
family members at increased risk of work–family conflicts,
stress and depression [22,23]. Economic losses within the
household further percolate to the next generation [24]. Eco-


nonlinearities. Nonlinear processes such as infectious disease
may aptly characterize incarceration.

In addition, an individual’s incarceration can expose his
or her family and friends to a network of criminals, thereby
exposing them to criminal norms [26]. These factors may
increase the criminality of the incarcerated person’s family
and friends, thereby increasing the probability that these
family members and friends would themselves be incar-
cerated. Thus, an individual’s incarceration can be ‘transmitted’
to others. Alternatively, there is evidence that an individual’s
incarceration can create an ‘official bias’ against his or her
family and friends [27–29]. That is, once a person is incarcer-
ated, the police and the courts pay more attention to the
incarcerated person’s family and friends, thereby increasing
the probability they will be caught, prosecuted and processed
by the criminal justice system. Thus, even if criminal behaviour
is not transmitted from person to person, a similar pattern
emerges because a convicted person’s associates will be
called and convicted more frequently than will a non-
incarcerated person’s associates, even if their criminal
behaviours are the same. Regardless of the mechanisms
involved, the incarceration of one family member undoubt-
dedly increases the likelihood of other family members being incar-
cerated [20,21,30,31]. This suggests that models of contagion
may aptly characterize incarceration.

It is well known that some models of contagion exhibit
nonlinearities. Nonlinear processes such as infectious disease
outbreaks are capable of amplifying small differences in par-


2. The susceptible – incarcerated – susceptible
model

The hypothesized ‘transmissibility’ of incarceration suggests
that an SIS model in which incarceration is modelled as
though it was an infectious disease is appropriate. An
agent-based simulation of the SIS model requires three
main components: a contact network through which individ-
uals stochastically transmit the disease, transmission
probabilities that dictate the rate at which agents transmit
to each other and a period of infectivity [32]. Network ties
represent opportunities for transmission, and the structure
of the network through which the disease spreads affects
the dynamics of the outbreak. In a disease model, network
ties between agents typically denote physical contact or
close proximity; in the case of incarceration, they denote
the existence of a familial relationship or close friendship, i.e.
the existence of a strong influence between the agents.

In the disease modelling paradigm, transmission probabil-
ities may be a function of characteristics of the individual
agents (e.g. elderly individuals have a higher probability of
contracting a disease) or the relationship between the agents
(e.g. transmission rates are higher between parents and small
children). In our model, incarcerated people are considered
‘infectious’ to those who are most profoundly affected by
their absence. Incarcerated people ‘transmit’ the incarceration
‘disease’ to their network members with a probability that is
a function of the relationship type (e.g. an individual’s
incarceration has greater effect on his or her child than on a
friend) and personal characteristics (e.g. males are more
susceptible than females). We denote the probability that
infected agent i transmits to susceptible agent j by p(i → j).

In modelling the spread of disease using the SIS model,
the period of infectivity, s, is the duration of time during
which the infected individual is contagious. In our model,
s is the length of the individual’s prison sentence, as that is
the time during which the inmate’s family and close friends
are acutely affected. Here, we do not explicitly model
increased risk of incarceration due to an inmate’s difficulty
re-integrating into society. In our model, individuals released
from prison cease to be ‘infectious’ and return to a ‘suscep-
tible’ state from which they may become re-infected. The
source of the new infection will be incarcerated friends and
relations, introducing positive feedback into the system. Given the infectious period, the probability that agent \(i\) transmits the disease to agent \(j\) over the whole course of the infectious period is given by \(p_i(t \rightarrow j) = 1 - (1 - p_i(t \rightarrow j))^s\), i.e. the complement of the probability that agent \(i\) does not transmit to agent \(j\) in any of the \(s\) iterations during which it is infectious. Thus, for fixed \(p_i(t \rightarrow j)\), a longer infectious period results in a higher transmission probability (figure 1) and a greater chance that the outbreak becomes widespread.

3. Simulation model

3.1. Overview

Simulating the described contagious process requires three main components: a synthetic population through which the ‘disease’ will be passed, transmission probabilities and durations of infectivity. In our simulation, we synthesize a realistic multi-generational population of agents for which all family and friendship ties are known. All parameters involved in creating this population are based on recent, high-quality data. For example, distributions for the sex, lifespan and the number of children of each agent are taken from the US Census, the Centers for Disease Control and Prevention and the Social Security Administration, respectively. Transmission probabilities are derived directly from the survey of prison inmates presented in [33]. This survey provides the probability that an inmate’s mother, father, sister, brother or adult child are also incarcerated by inmate gender. That is, \(p_i(t \rightarrow j)\) is taken directly from the literature. In our simulation, we treat close friends as siblings in terms of transmission probabilities. We focus on the crime of drug possession and use data from the US Census, the Centers for Disease Control and Prevention and ‘Black’ simulations using a distribution of sentence lengths corresponding to those received by White Americans for the same crime, as described.

In order to test whether differential sentencing alone can explain racial disparities in incarceration rates, we use the same transmission probabilities and the same network to represent both Black and White populations. We acknowledge that there may be differences between the two communities in terms of demographics that affect the structure of the networks (number of children, for example); however, in order to isolate the effect of differential sentencing, we hold all else constant—the network included. Indeed, the ability to isolate one mechanism and disentangle its effect from all others is one of the main advantages of taking a simulation approach.

3.2. Synthetic population

3.2.1. Agent attributes

We begin with a seed population of \(n = 1500\) individuals, \(\{a_1, a_2, \ldots, a_n\}\), from which all members of the population will be descended. To initialize each agent, it is assigned several attributes. The \(i\)th agent, \(a_i\), is assigned a gender from the distribution \(g_i \sim \text{Bernoulli}(0.5)\), a birth year \(b_i \sim \text{Uniform}([0, 15])\) and a spatial location in the unit square \((x_i, y_i) \sim \text{Uniform}(0, 1))\). The location may be thought of geographically, as a physical location in a city or as a preference space. Regardless of how one prefers to conceptualize the spatial location, it serves the function of creating communities in the network, as friends and spouses are selected with respect to these locations. To simulate a realistic distribution of life durations, the age at death is sampled according to the 2009 life tables released by the Social Security Administration.\(^2\) In these tables, the probability of death in the next year at each age (from 0 to 119) is given by sex. We treat these as the probabilities of death at each age throughout the simulation. If individual \(i\) is female and born in year \(b_i\), we select a life duration, \(l_i\), at random from the distribution given in the female life tables (\(l_i \sim \text{Multinomial}(p_{\text{female}})\)). The iteration of death is then given by \(d_i = b_i + l_i\).

Female agents are assigned an age at first birth attribute. Age at first birth is based upon a figure released by the Centers for Disease Control, which lists the mean age at first childbirth for women in 2011 as 25.6 years.\(^3\) We specify \(h_i\), the age at first birth, to be drawn from the distribution \(h_i = 15 + r_i\), where \(r_i \sim \text{Poisson}(10.6)\), which has mean 25.6.

Each female agent is also assigned the number of children she will have throughout her lifetime. The US Census provides the distribution of the number of children that women in the age bracket 40–44 have had and lists the total fertility rate for several years between 1980 and 2008, which ranges from about 1.84 to 2.1. To prevent our simulated population from dying out (fertility rate < 2), we adjust the raw distribution given for 40- to 44-year-old women to be consistent with historical fertility rates. Under this adjusted distribution, the expected fertility rate is 2.07 children per woman.

3.2.2. Network ties

In addition to personal characteristics, each agent is endowed with relationships with other agents. These are represented as edges in a network. In our simulation, all parent–child, sibling, spouse and close friend relationships are represented. When an agent reaches its 10th ‘birthday’, the agent forms friendship ties. In order to select the number of close friends assigned to each agent, we use data from the 2004 General Social Survey,\(^4\) in which respondents indicate the number
of individuals with whom they discuss important matters and their relationship to up to five of these people. Because many familial relations are already accounted for in our simulation, we count only those people listed who are not parents, children, spouses or siblings to calculate the probability of selecting each possible number of friends, \( f_i \).

Conditional on \( f_i \), we select the specific agents that will be designated as friends. We consider potential friends to be non-siblings between the ages of 9 and 11. This age range is based on information obtained from the National Longitudinal Study of Adolescent Health, in which children were asked to list several of their friends. Of the friends listed, nearly 95% were within one grade of the student (75% shared the same grade). Because friends within 1 year of the students made up the majority of close childhood friendships, we also restrict to this range. From this set of potential friends, we select the \( f_i \) agents that are closest in Euclidean space to the given agent.

In the iteration in which a female actor’s first child is born, she is assigned a partner. In this hypothetical population, we only model opposite-sex spousal ties. The algorithm first finds all potential partners—unrelated and non-friend male agents whose current age is between the female agent’s age and 9 years older. We selected this scheme based upon data from the 1999 US Census that shows that for about 80% of marriages, the husband’s age minus the wife’s age falls into the range \( [9, 11] \). Because friends within 1 year of the students made up the majority of close childhood friendships, we also restrict to this range. From this set of potential friends, we select the \( f_i \) agents that are closest in Euclidean space to the given agent.

At this point, the children of the couple (or single mother) are initialized. The first child is assigned to be born in the current iteration. Subsequent children are assigned birth years as the current iteration plus independent and identically distributed draws from a Poisson distribution with mean parameter \( \lambda = 4.5 \). The mean parameter, \( \lambda \), is again based on the Centers for Disease Control’s data used to calibrate the age at first birth. Child locations are set to be half-way between the mother’s and father’s location along the x- and y-axis, respectively.

We run our algorithm for 200 iterations, resulting in a total population of 13,826 individuals. We discard the first 150 iterations as a burn-in period, reducing dependence on our initial conditions. Those agents that are part of our simulation (i.e. those that are alive at any point beyond the 150th iteration) are retained. The total population consists of 8856 agents, with 61,376 family and friendship ties. This total population size is not pre-specified, as it is the result of the random birth and death process initialized with 1500 agents. Figure 2a shows an example family tree generated by our algorithm. This individual family, of course, does not exist in isolation of the rest of the population. The population-wide network is shown in figure 2b.

### 3.3. Generating sentences

Data released by the Bureau of Justice Statistics indicate that the duration of sentence served for the same crime varies by race. In particular, the Bureau of Justice Statistics states that for drug possession, the mean sentence for Whites is 14 months with a median of 10 months. For Blacks, the mean sentence served is 17 months with a median of 10 months. We use a negative binomial distribution to generate sentence lengths that are consistent with the specified summary statistics. A comparison of the sentence distributions is shown in figure 3.

### 3.4. Transmission probabilities

Dallaire [33] presents the results of a survey of incarcerated individuals. In this survey, each inmate is asked which of
Table 1. Derived monthly transmission probabilities, \( p(i \rightarrow j) \).

<table>
<thead>
<tr>
<th></th>
<th>women</th>
<th>men</th>
</tr>
</thead>
<tbody>
<tr>
<td>mother</td>
<td>0.001</td>
<td>0.003</td>
</tr>
<tr>
<td>father</td>
<td>0.011</td>
<td>0.011</td>
</tr>
<tr>
<td>sister</td>
<td>0.008</td>
<td>0.004</td>
</tr>
<tr>
<td>brother</td>
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<td>0.030</td>
</tr>
<tr>
<td>spouse</td>
<td>0.004</td>
<td>0.001</td>
</tr>
<tr>
<td>adult</td>
<td>0.017</td>
<td>0.006</td>
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</tbody>
</table>

their relations are also incarcerated. The proportion of inmates whose relations are incarcerated are reported in these data by sex. We use these probabilities to derive our transmission probabilities, \( p(i \rightarrow j) \). We have noted that if agent \( i \) has probability \( p(i \rightarrow j) \) of infecting agent \( j \) each month it is incarcerated and its sentence is \( s \) months, then the probability of transmission over the course of its incarceration, \( p_{\text{sentence}}(i \rightarrow j) \), is given by \( p_{\text{sentence}}(i \rightarrow j) = 1 - (1 - p(i \rightarrow j))^s \). One can easily solve for \( p(i \rightarrow j) = 1 - (1 - p_{\text{sentence}}(i \rightarrow j))^{1/s} \). We set \( s = 14 \) to approximately calibrate to a value between the Blacks and Whites. The derived monthly transmission probabilities used in this simulation are given in table 1.

The derived monthly transmission probabilities are most usefully understood in the context of the probability of transmission averaging over sentence length, i.e. the marginal transmission probabilities,

\[
p_{\text{race}}(i \rightarrow j) = \sum_{s=0}^{\infty} p_{\text{sentence}}(i \rightarrow j) \pi_{\text{race}}(s),
\]

where \( p_{\text{sentence}} \) is defined above and \( \pi_{\text{race}}(s) \) is the distribution of sentence lengths for each race. Marginal probabilities by race, sex of inmate and relation are given in table 2, along with the original probabilities listed in [33]. These were calculated using the Monte Carlo method. We note that our marginal probabilities for the White sentences tend to be slightly higher than that given. For Black sentences, the marginal probability of transmission tends to be slightly lower than that given. This is, of course, unsurprising as the sentences tend to be shorter for Whites and they thus have fewer opportunities for transmission. Recall that the monthly transmission rates under the two scenarios are precisely the same, so one should not interpret the differences in marginal probabilities to mean that this model implies that Blacks are more susceptible than Whites under the same conditions. The only differences that exist here are due to the discrepancies in sentencing. We find that the probability of transmission for Blacks under these parameters tends to be about 20% greater than the probability of transmission for Whites.

The assumption that one’s infectivity is constant over the duration of the sentence and non-existent upon release is likely flawed. In reality, the degree of infectivity likely varies over the course of the sentence and slowly decays after releases. Ex-inmates may even have heightened infectivity immediately after release, particularly as they try to reintegrate into their families and communities and find work. However, we find it plausible that the duration of infectivity is at the very least positively correlated with sentence length. If we were to assume that the infectious period were proportional to the sentence length and calibrate the transmission probabilities to this increased period of infectivity, the eventual results of our simulation would be quite similar, as the marginal probability of transmission over the course of the infectious period would be similar. Parsimony is an important quality in an agent-based model, and we have endeavoured to adhere to this principle whenever possible, particularly in cases where there are no data to inform more detailed parameters. In this case, we have no information about the duration and rate of post-release reacclimatization. Thus, we assume that infections only take place during the sentence, a period about which we have concrete data.

4. Results

We run our simulation 250 times using each sentence-length distribution, resulting in 250 ‘Black’ epidemics and 250 ‘White’ epidemics. The populations are initialized to have approximately 1% of individuals incarcerated at the beginning of the simulation. Although White and Black incarceration rates, in reality, have been disparate throughout history, we initialize the simulations equivalently under the two scenarios to rule out the possibility that the resultant disparities are due to initial conditions alone. Our first analysis looks at the effects of differential sentencing after 50 years, a duration slightly longer than the time since the term ‘War on Drugs’ was coined in the USA. Figure 4a shows the mean epidemic curves and corresponding confidence intervals by race; figure 4b shows several example trajectories. While the prison epidemic takes off under Black sentence lengths, reaching just under 3% incarcerated on average after the 50 years in our simulation, under the model using White sentence lengths, the prison population first declines and then increases at a much slower rate, reaching about 0.725% over the same time period. Eventually, the Black and White simulations level off (or increase at a negligibly slow rate) at around 7% and 1%, respectively.

We have shown that under a largely arbitrary simulation period and initial values, large racial disparities in incarceration rates are an emergent property of this model. We now test whether this conclusion holds under realistic time periods and initial values. As a comparison with real data, we initialize the Black and White simulations to be near the real incarceration rates in California in the mid-1980s when mandatory drug sentencing became law there (1% and 0.15% for Blacks and Whites, respectively). The results of this are shown in figure 5. From 1986 to 2010, the incarceration rate for Blacks in California climbed from about 1 to 2.18%, whereas the rate for Whites rose much more modestly from 0.15 to 0.277%. In our simulations, the Black and White simulations increase to 2.13% and 0.33%, respectively, over the same time period. For the Black trajectory, our mean trajectory deviates slightly though the ultimate results after 25 years are remarkably similar to reality; however, we note that the real trajectory in California is within the range of values that are typical under our simulation, as it falls well within the cloud of trajectories shown. The White trajectory in California is quite similar to the mean trajectory in our simulation.

Agent-based models are often evaluated with respect to the model’s ability to reproduce a set of ‘stylized facts’ as opposed to specific values [34,35]. These facts are often...
structural or qualitative in nature; examples include patterns of persistence or high autocorrelation in financial time-series [34], spatial clustering of crime [36] or habitation sites [37] and racial segregation in housing [38]. Thus, it is not strictly necessary that a generative model be able to perfectly match observed statistics to establish the plausibility of the generative mechanism; rather, it is necessary that the agent-based model be able to reproduce observed structural properties at a qualitative level. For example, in Schelling’s foundational work in describing a generative mechanism for racial segregation in housing, it is enough to show that simple decision rules produce clusters of like agents [38]; one need not establish an exact correspondence between clustering coefficients in these simulations and those seen in reality.

In that spirit, here we establish the ability of our model to reproduce structural properties of the incarceration epidemic with respect to the patterns found in recidivism data. In figure 6, we compare the recidivism rates derived from the simulation model to those released by the state of California. The plots to the right show the same statistics from a variety of other states. The chosen states vary because not all states release the same statistics; we chose California as our primary point of comparison because its report shows recidivism rates disaggregated by the most factors and at the highest level of discretization. Remarkably, our model reproduces the structural properties of recidivism very accurately. For example, in both our model and the data from California, recidivism rates increase with the number of times an inmate has been incarcerated, with the largest increase occurring between the first and second incarceration. This effect emerges as a structural property of the SIS model without including an increased probability of incarceration for those who have previously been incarcerated. An agent’s incarceration affects its local network enough that, upon release, it has a higher probability of return. In all cases, including our simulation results, the recidivism rate is lower for those who are released at an advanced age. In fact, our simulation even reproduces a subtle demographic bulge in rates that is apparent in the California report. This is again an emergent property of the model—as an agent ages, it tends to have fewer (and different types of) contacts who generate positive feedback for incarceration. The rate by months since release (for those who recidivated within 3 years) closely matches the structure of the data from all states. These results suggest that our model is reflecting the underlying regularities in the system.

Figure 4. (a) The mean incarceration prevalence over time by race. Log p-values shown below indicate that at all but the first time point, the mean prevalence is significantly different between the two populations. (b) Several example epidemic curves.

Table 2. Probabilities given in [33] and marginal transmission probabilities for Whites and Blacks.

<table>
<thead>
<tr>
<th></th>
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</thead>
<tbody>
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<td></td>
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<tr>
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reflects the trend as a function of sentence length. The discrepancy between the real data and the simulated data offer suggestions for refinements to the model that could improve its accuracy. Such refinements, however, come at the expense of parsimony. Given how well our extremely parsimonious model is able to reproduce many facets of the incarceration epidemic, we anticipate that the marginal returns for incorporating the described refinements will be negligible. That is, although we may be able to more precisely match exact figures from the real data with a more complicated and highly parameterized model, realizing small improvements to the accuracy of the incarceration and recidivism rates in our simulation will do little to further establish the plausibility of our hypotheses. In its current form, our model clearly demonstrates the plausibility of our hypothesis that incarceration may be spread in a manner similar to a contagious disease and that sentencing disparities may be a major cause of racial differences in incarceration rates.

4.1. A non-contagious model

It is important to test whether the causal mechanism of interest (in this case, contagion) is a necessary component of the model [36,37]. If it is possible to reproduce an equivalent fit to the structural properties of the observed process without the mechanism, then, under an appeal to parsimony, it is not sensible to put forth the mechanism as a plausible candidate for the true causal mechanism. We address this issue by running a simulation in which incarceration is not contagious. In these simulations, the mechanism of interest is ‘turned off’. Here, there is only spontaneous infection—at each iteration, each agent has equal probability, \( p \), of becoming infected independent of any relationship with an incarcerated agent. In order to make a fair comparison, in this simulation we tune the probability of spontaneous infection such that, at the end of the simulation, the Black population’s prevalence is roughly 3% as in the SIS simulations. We again initialize both populations to have a 1% incarceration rate at the outset and use the appropriate sentence-length distributions for Black and White simulations. Figure 8 shows the results of this experiment. Under this model, we are not able to reproduce the large difference in incarceration rates between the Black and White simulations. Here, the curves are only about 20% different, whereas in the simulations that incorporate the contagious mechanism, as well as in reality, the incarceration rate of Black agents is many times that of the White agents. The trajectory of the
epidemic under this scenario also fails to resemble that of the real incarceration epidemic—it takes off very quickly and within approximately 5 years reaches equilibrium (figure 8). Recidivism statistics that pertain to the non-contagious simulation appear in figure 9. Under the non-contagious model, we find a very poor correspondence to reality. Under this model, the rate of recidivism with respect to the number of times previously incarcerated is decreasing, whereas in the contagious simulation and the real data it is increasing. Here, the cumulative rate of recidivism by months since release shows a convex pattern where both data generated by the contagious model as well as the real data exhibit a concave shape. The probability of recidivism is essentially flat with respect to the length of the sentence under a non-contagious model; real data and the contagious model both show that longer sentences result in a decreased probability of recidivism. Recidivism by age is the only metric by which the non-contagious model roughly reproduces the same results as seen in the California data, though even here the shape of the curve appears markedly rounder than in the California data and lacks the characteristic demographic bulge.

Based on the lack of correspondence between reality and recidivism data generated under a non-contagious model coupled with the close correspondence between reality and

Figure 6. A comparison of the output of our simulation model with recidivism rates in California and several other states. Results from ‘Black’ simulations are shown with black bars, and those from the ‘White’ simulations are shown with lighter grey bars. In the case where differences between the results from the two sets of simulations are visually indistinguishable, we present only the ‘Black’ results.
We conclude that sentence-length disparities in the absence of network effects do not account for the observed difference in incarceration rates. It is feedback through the local network coupled with sentencing disparities that causes such large differences in incarceration rates.

4.2. An ordinary differential equations approach

Under assumptions of random mixing and homogeneity of transmission rate, the SIS model can be written as a set of ordinary differential equations (ODEs). In this case, the number of people infected—the prevalence—when an outbreak reaches a steady state, $I$, is determined by the expected number of transmissions per infected person, which in turn is given by the product $ps$ of transmission rate and the duration of infectivity. The presence of a positive feedback loop makes the relation between prevalence and number of transmissions highly nonlinear. In particular, ignoring births and deaths, $I = 0$ if $s < p^{-1}$ and $1 - 1/ps$ otherwise. Thus, given the transmission rate $p$, we can define a critical duration of infectivity $s_c = p^{-1}$ such that for $s < s_c$ the outbreak dies out, while for $s > s_c$ it achieves a non-zero steady-state prevalence. Near $s_c$, small differences in sentencing (i.e. duration of infectivity) can cause large differences in incarceration (i.e. disease) prevalence.

We take an agent-based simulation approach to modelling the incarceration epidemic because neither the assumption of uniform mixing nor the assumption of homogeneity is met. Indeed, the network of family and friends plays a crucial role in our hypothesis. Furthermore, the data show that transmission rates depend on the nature of the relationship between the infectious and susceptible people, and any particular susceptible may simultaneously have several types of relationships with different infectious people (e.g. mother and sister and daughter). It is easier to capture this heterogeneity in transmission rates in an agent-based simulation than in a set of ODEs.

Moreover, using the output of our simulation, we can generate a population-wide mean transmission rate $p$ to calibrate an ODE model. The result is $p \approx 0.0612$ transmissions per infected person per month, or $s_c \approx 16.3$ months. Thus, the mean sentence lengths (17 and 14) are on opposite sides of the critical point, illustrated in figure 10. Under this model,
the incarceration epidemic in the White population would eventually die out. However, the Black population’s incarceration rate would reach a steady state of about 3.9%. As this model does not account for spontaneous infections, this result is consistent with those of the agent-based model. This approach, however, does not allow us to assess the validity of the model or comment on its ability to reproduce structural properties of the epidemic using other withheld sources of information, such as the recidivism data we have shown.

5. Future directions
Incarceration as a contagion is a complex process that co-evolves with many other complex social processes, e.g. poverty. Our model by no means accounts for all of the complexities in the process; however, we view the simplicity of our model as one of its major strengths. Even without accounting for the details, our model is able to capture many of the facets (or ‘stylized facts’) observed in real incarceration data.

Extensions to this model could nonetheless prove fruitful. For example, one could investigate the extent to which differences in network structure between Black and White communities might encourage or discourage the spread of incarceration. In particular, increasing the fertility rate in certain communities would raise the average degree in the network, giving more opportunities for transmission. One might investigate whether the differences between Black and White communities in fertility rate plausibly play a significant role in the spread of incarceration. Black and White communities do not exist completely separately; real networks include both Black and White individuals. One might simulate a more realistic mixed-race influence network to investigate the extent to which the rate with which the two communities are commingled might affect the epidemic, even under inequitable sentencing for Blacks. In this work, we have ignored the residual effects of incarceration once the agent is released. Future work might seek to model an extended infectious period during which the ex-inmate is trying to reintegrate into their home life and find work. Last, one very interesting extension would be to simulate the mechanisms by which incarceration is spread. People with low income are more likely to become incarcerated, and incarceration decreases one’s expected lifetime earnings. These effects are passed on through the generations, as social mobility data indicate the children of low income parents are themselves likely to have low income, and thus are at increased risk of incarceration. It has also been documented that incarceration increases the likelihood of relationship dissolution and prevents young men from forming stable long-term relationships. A lack of stable family life is a risk factor for incarceration, increasing the likelihood of re-incarceration for the ex-inmates as well as for their children. Understanding and quantifying the relationship between poverty and incarceration could have a potentially large impact, as the insights gained could be used to help drive policy decisions.
The plausibility of social transmission suggests that the incarceration epidemic might be ameliorated by reducing the spread of incarceration through contact networks. In a model of contagion, reducing the overall probability of transmission reduces the steady-state prevalence of the epidemic. Transmission probabilities can be reduced by decreasing the time-rate of transmission or shortening the infectious period. Our model provides a framework for assessing the possible long-term effects of policies targeted at reducing transmission probabilities, but it is critical to first determine empirically the effectiveness of various policies. To this end, studies of the effect of different sentence lengths, alternative punishments, targeted interventions and retraining/re-integration programmes on ‘transmissibility’ are urgently needed. These data, in conjunction with models of contagion, would provide a substantially improved framework for analysis of incarceration-related policies and their cascading effects.

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Endnotes
1 Simulation code is available at http://ndsl.vbi.vt.edu/synthetic-data/.
3 http://www3.norc.org/gss+website/.
4 http://www.cpc.unc.edu/projects/addhealth; data not publicly available.

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