Coupling Epidemiological Models with Social Dynamics

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Abstract

In this work we study a Susceptible-Infected-Susceptible model coupled with a continuous opinion dynamics model. We assume that each individual can take measures to reduce the probability of contagion, and the level of effort each agent applies can change due to social interactions. We propose simple rules to model the propagation of behaviors that modify the level of effort, and analyze their impact on the dynamics of the disease. We derive a two dimensional set of ordinary differential equations describing the dynamic of the proportion of the number of infected individuals and the mean value of the effort parameter, and analyze the equilibria of the system. The stability of the endemic phase and disease free equilibria depends only on the mean value of the levels of efforts, and not on the initial distribution of levels of effort.

Keywords
Epidemic models · Social interactions · Reproduction number · Opinion dynamic

Mathematics Subject Classification 91C20 · 82B21 · 92D25

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1 Introduction

In the last few years an increasing interest emerged on epidemic models involving changes in socio-cultural norms and behaviors, partly motivated by the emergence of anti-vaccine movements in different countries and the challenge they pose to the control of epidemic outbreaks.

Several authors divided the susceptible population in three (or more) different groups, the vaccinated, the non-vaccinated individuals, and the ones that decided to remain non-vaccinated, see for instance (Bauch 2005; Bauch and Earn 2004; Pires and Crokidakis 2017). Also, related models considered two groups of susceptible agents, the ones who are aware and those who are not aware of the threat of an infection (Clancy 2018; Misra et al. 2011; Yang et al. 2017). In those cases, each group has its own rate of contagion. Groups with different degrees of susceptibility, such as age-dependent groups, and a class of exposed agents were considered. The difficulty to analyze the corresponding models increases with the amount of classes and groups since they lead to coupled systems of differential equations involving many parameters describing the transitions between different groups.

Usually, agents can move from one group to another due to the mechanisms of the disease (for instance, a susceptible agent was exposed and becomes infected, or an infected agent recovers after the infection), or due to a personal decision based on its own beliefs or the influence of other agents. So, the transitions involve a set of ordinary differential equations modelling the disease dynamics coupled with different social contact processes like the voter model, Axelrod’s cultural transmission model (Axelrod 1997), imitation dynamics (Bauch 2005), or other discrete opinion dynamics models like Galam’s, Sznajd’s, or Ochrombel’s (Galam and Zucker 2000; Ochronbel 2001; Slanina and Lavicka 2003; Sznajd-Weron and Sznajd 2000). Now, the possibility of an epidemic outbreak, and its long-term impact, will depend on the size of those groups, the mechanisms of transmission of the disease, the social contact process, and also on the clustering or other properties of the network modelling the social structure of the population, see for instance (Dorso et al. 2017; da Silva et al. 2019; Su et al. 2018; Tyson et al. 2020; Zhou et al. 2019).

In particular, when vaccines are available, opinion-based models were used to analyze the transition from susceptible to vaccinated states, as a result of social interactions, see (Dorso et al. 2017; Pires and Crokidakis 2017). However, the decision of being vaccinated depends on other factors like their cost, risks, and perceived advantages, see for instance the Bauch’s work (Bauch 2005), where a replicator-type equation was added to a classical SIR model, and the payoff of being vaccinated changes depending on the risk of contagion.

However, there are many infectious diseases that can not be analyzed with these kind of models because their prevention depends on a sustained effort over time through preventive measures, and there are no vaccines available. As a paradigmatic examples we can consider the actual Covid-19 pandemic, the 2009 flu pandemic caused by the H1N1 virus, or the outbreaks of Dengue, Zika, and Chikungunya (Fauci and Morens 2016). In those cases it is possible to analyze the spread of behaviors (and misbehaviors) related to infectious diseases using continuous opinion dynamics models, like the ones considered by Bellomo, Deffuant, Toscani, Weisbuch and their collaborators,
Coupling Epidemiological Models with Social Dynamics

So, we present here a variant of a classic Susceptible-Infected-Susceptible (SIS) model coupled with a continuous opinion dynamics model. Briefly, the state of any agent \( j \) in the society is characterized by the pair \((a_j, p_j)\), where \( a_j \in \{I, S\} \) means that the agent is infected or susceptible, respectively, and \( p_j \in (0, 1) \) is a factor reducing the transmission of the disease; \( p_j \) is related to its level of effort to avoid the infection, the lower is \( p_j \), the greater are the prevention measures taken by the agent to avoid contagion.

Let us mention a related model by Funk, Gilad, Watkins and Jansen (Funk et al. 2009), where the discrete set of positive integers \( \{k\}_{k\geq 0} \) represents different levels of awareness or effort (being 0 the maximum level of awareness, which implies immunity against the disease). The spreading of awareness on the population depends on several mechanisms: information transmission due to interaction among agents, a decay term since individuals forget the acquired information, and a reset term since each infected agent becomes fully informed and goes to 0. A Susceptible-Infected-Recovered (SIR) model was considered, which always converges to an equilibrium without infected people. When an infected agent interacts with a susceptible agent at level \( k \), the probability of infection is proportional to \((1 - \rho^k)\) for some fixed \( \rho \in [0, 1] \). The authors studied this model using agent-based simulations, and derived a SIR system coupled with an equation for the information diffusion which is non-closed since it depends on the location of agents. Let us observe that long range jumps of agents awareness levels are allowed, since infected agents go directly to zero. This introduces some kind of non-locality to the equations.

In our model, when two agents interact, they change their states as a consequence of both the epidemic and opinion dynamics as detailed in Sect. 2. Informally, although the rates of contact \( \beta \) and recovery \( \gamma \) remain unchanged, a susceptible agent \( i \) is infected with probability \( p_i \beta \) after a contact with an infected agent \( j \). Then, both agents \( i \) and \( j \) increase or decrease their levels of effort depending on the existence or not of contagion after the interaction. We also examine the possibility where the contagion probability depends on the infected agent, \( p_j \beta \), or depends on both agents measures, \( p_i p_j \beta \).

On the other hand, if two susceptible or two infected agents interact, each one moves its own effort level toward the other agents value as in continuous opinion dynamics models (Aletti et al. 2007; Toscani 2006), depending on a third parameter which does not enter in the coefficients of the system of equations. It plays a role only on the formation of consensus, see (Pinasco et al. 2017), since the mean value of the level of effort remains unchanged after this kind of interaction.

We are mainly interested here in the extinction or not of the disease. To this end, in Sect. 3 we derive a system of ordinary differential equations describing the proportion of susceptible agents and the mean value of the level of efforts of the population, \( \langle p \rangle \), in the mean field approximation.

Although the system depends on \( \langle p_S \rangle \), the mean value of the level of efforts of the susceptible population, simulations show that \( \langle p \rangle \) and \( \langle p_S \rangle \) are very similar, and coincide when the parameter in the social interaction goes to zero. So, assuming \( \langle p \rangle = \langle p_S \rangle \), we study the system of ordinary differential equations satisfied by \( \langle p \rangle \).
and the proportion $S$ of susceptible agents. We find the fixed points and classify them according to their stability in Sect. 4.

In the long-time limit, when the increase or decrease of the level of effort depends on the same parameter, the value $\langle p \rangle$ converges to $\langle p \rangle_\infty = (1 + \beta)^{-1}$, independently of the initial distribution of values of $p$ in the population. If they depend on different parameters, we get two values $\langle p \rangle_\pm$, depending on which one is greater, and analogous results hold. We show, in particular, the existence of a threshold depending only on the contact rate $\beta$ and on the recovery rate $\gamma$, and not in the initial value of $\langle p \rangle$, such that the disease does not become endemic if and only if

$$R_m := \langle p \rangle_\infty R_0 = \frac{2\beta}{\gamma(1 + \beta)} \leq 1.$$  

Notice that $2\beta$ is the rate of contagion in this model, and that $R_0 = 2\beta/\gamma$ is the basic reproduction number of the standard SIS model. We call this modified reproduction number $R_m$ the behaviorally reduced reproduction number, since the factor $1 + \beta$ appears due to the social process and helps to reduce the propagation of the disease, decreasing the basic reproduction number $R_0$ of the standard SIS model. Notice that this modified reproduction number $R_m$ is obtained asymptotically when both the SIS and the social dynamics evolve. A discussion of the significance of $R_m$ is given in Sect. 5. Agent-based simulations of the dynamics strongly agree with the theoretical results.

We conclude in Sect. 6, and we add a short “Appendix” with the derivation of the equation for the evolution of the proportion of susceptible agents, and the corresponding equations for other contagion rules.

2 The Model

Let us assume that we have a population of $n$ agents, each one characterized by a pair $(a_i, p_i)$, where $a_i$ represents the state of agent $i$, namely $a_i = S$ if it is susceptible and $a_i = I$ if it is infected, and $p_i \in [0, 1]$ denotes its level of effort to avoid infection.

The purely epidemiological parameters are the recovery rate $\gamma$, and the contact rate $\beta$. Both are assumed to be constant.

When two agents $i, j$ interact, they will change their parameters from $(a_i, p_i)$ to $(a_i^*, p_i^*)$, and from $(a_j, p_j)$ to $(a_j^*, p_j^*)$ respectively. The recovery dynamic is similar to the classical SIS model: an infected agent becomes susceptible in a unit of time with probability $\gamma$, without interactions. On the other hand the dynamics of infection depends on the social parameter: if a susceptible agent $i$ interacts with an infected agent, it becomes infected with probability $p_i \beta$.

**Remark 2.1** We are assuming here that only the level of effort of the susceptible agent matters, as in Funk et al. (2009). Alternative rules can be imposed here. For instance, the contagion dynamic could depend on $p_i, p_j$, or both. These rules can be studied in the exact same way. We present the corresponding equations below, and their derivation can be found in the “Appendix”.
However, drastic changes in effort levels, such as those that occur in Funk et al. (2009), need a different approach, since this model includes long jumps in their values of awareness.

When two agents $i$ and $j$, either both susceptible or both infected, interact, their level of measures of prevention $p_i$ and $p_j$ will change following a social dynamic rule similar to the one introduced by Deffuant and Weisbuch (see (Deffuant et al. 2000; Pérez-Llanos et al. 2018; Toscani 2006)): each agent moves its parameter $p$ toward the other agent’s value. More precisely, following (Deffuant et al. 2000), we introduce a positive fixed parameter $h \in [0, 1]$, the *compromise* parameter, which measures the ability of an agent to adapt its own level of effort in social interactions. After an interaction with agent $j$, agent $i$ will change its level to

$$p_i^* = p_i + h(p_j - p_i).$$

When $h = 0$ no change occurs, while for $h = 1/2$ both agents choose the mean value of their levels of effort. This compromise parameter $h$ represents the susceptibility to persuasion, the extent to which an agent is willing to modify its behavior due to the influence of other agents. Different theories from social sciences support this behavior: Asch’s social pressure theory (Asch 1955), where an agent tries to adapt its opinion to others agents opinions, and the persuasive argument theory of Burnstein and Vinokur (1977), where agents exchange arguments and modify their positions after analyzing them.

On the other hand, if a susceptible agent interacts with an infected agent, both will increase (respectively, decrease) their levels of effort whenever the agent remains susceptible (resp., becomes infected). We can think this rule as if both agents have a social interaction with a third hypothetical agent who has the maximum (resp., minimum) level of effort $p = 0$ (resp., $p = 1$), with two different compromise parameters $h_1, h_2 \in [0, 1/2]$. Hence, we have

$$p_i^* = p_i - h_1 p_i, \quad \text{or} \quad p_i^* = p_i + h_2 (1 - p_i)$$

if contagion occurs or not.

Let us state precisely the rules of the dynamics, together with their interpretation:

- **Persuasion or imitation:** a susceptible (respectively, infected) agent $i$ remains susceptible if he interacts with another susceptible (resp., infected) agent $j$. Both change their levels of effort to

$$p_i^* = p_i + h(p_j - p_i),$$

$$p_j^* = p_j + h(p_i - p_j),$$

that is, each one adopts an effort level intermediate between its own value and the one of the other agent.

- **Contagion and Fear:** a susceptible agent $i$ becomes infected with probability $p_i \beta$ after an interaction with an infected agent $j$. In this case, $i$ and $j$ feel that their
efforts are not enough, and change their $p$ parameters to

$$
p^*_i = p_i - h_1 p_i,
p^*_j = p_j - h_1 p_j.
$$

- **Confidence**: a susceptible agent $i$ remains susceptible with probability $1 - p_i \beta$ after an interaction with an infected agent $j$. In this case, $i$ and $j$ feel that their efforts are excessive and change them to

$$
p^*_i = p_i + h_2 (1 - p_i),
p^*_j = p_j + h_2 (1 - p_j).
$$

- **Recovery**: a random agent $i$ is selected and becomes susceptible with probability $\gamma$ if it was infected, and no change of $p_i$ occur.

Let us observe that both dynamics are coupled, in the sense that the same contact can produce a new infection and a change in the levels of effort of both agents. It is easy to add a separate dynamic which only impacts on the effort of agents.

### 3 Theoretical Analysis

In this section we derive an ordinary differential equation for $\langle p \rangle$, the mean value of agents levels of effort, and another one for $S$, the proportion of susceptible agents.

Let us mention that if $h_1 \neq h_2$, the system is not closed, since it involves the second moment $\langle p^2 \rangle$, and we need an extra equation for it. However, this new equation depends on $\langle p^3 \rangle$, and so on. This way we get an infinite hierarchy of coupled equations.

There are several ways to deal with this problem. In statistical mechanics and many other problems, under appropriate hypotheses, a closure of moments is introduced to truncate this kind of hierarchies. Another option is to introduce a measure valued function $\mu_t$ for the distribution of agents on the levels of effort and it is possible to obtain a more detailed description through a Boltzmann-like equation coupled with a SIS system describing the infection dynamics. We prefer to defer the technical details and the mathematical proofs of existence, uniqueness and stability of such equations to another work.

In the next section we consider two simpler approaches.

Let us first introduce the following notation which will be used below. We assume that there are finitely many agents, say $n$, and denote $k(t)$ the expected number of susceptible agents at time $t$. The proportions $S(t)$ and $I(t)$ of susceptible and infected agents are then given by

$$
S(t) = \frac{k(t)}{n}, \quad I(t) = 1 - S(t).
$$
The mean values $\langle p \rangle$, $\langle p_S \rangle$ and $\langle p^2_S \rangle$ of the $p$ parameter in the susceptible population and the mean of $p$ in the whole population are

$$P_S = \sum_{i \in \text{Sus}} p_i, \quad \langle p_S \rangle = \frac{P_S}{k(t)},$$

$$\langle p^2_S \rangle = \frac{1}{k(t)} \sum_{i \in \text{Sus}} p_i^2, \quad \langle p \rangle = \frac{1}{n} \sum_i p_i,$$  

where we denote by Sus and Inf the (time dependent) subsets of susceptible and infected agents respectively.

We assume that interactions between agents occur following a Poisson process with constant rate, that can be assumed equal to 1 without loss of generality, rescaling time if necessary. In particular the probability of an interaction in a small time interval $[t, t + \Delta t]$ is $\Delta t + o(\Delta t)$, no interaction occurs with probability $1 - \Delta t + o(\Delta t)$, and two or more interactions occur with a negligible probability $o(\Delta t)$. In what follows we will neglect the $o(\Delta t)$. When an interaction occurs, two agents are selected at random (uniformly across all agents), and they interact following the rules described in Sect. 2.

Our main objective in this section is to derive the differential equations describing the evolution of the expected values of $S(t)$ and $\langle p \rangle$.

### 3.1 Evolution of the effort level of a single agent.

Let us fix a single agent $i$, and study the evolution of its effort level at time $t$, i.e. the expected change $p_i(t + \Delta t) - p_i(t)$ in a small time window $[t, t + \Delta t]$. We have the following master equation for $p_i(t + \Delta t)$:

$$p_i(t + \Delta t) = (1 - \Delta t) p_i(t) + \Delta t p^*_i(t),$$  \hspace{1cm} (1)

where the first term is the probability that no interaction occurs and $p_i$ remains the same, and, in the second one, we have the expected value of $p^*_i$ when a single interaction occurs. In order to compute $p^*_i$ we need the possible values it can take and their probabilities, which depend on the state of agent $i$.

The probability that agents $i$ and $j$ are selected is $\frac{2}{n(n-1)}$. Notice that the factor 2 comes from the fact that agent $i$ can be selected as the first or second agent in the interaction. If we derive a classical SIS model under this assumption, we get $R_0 = 2\beta/\gamma$, a fact that must be remembered in order to compare it to the modified reproduction number when agents behaviors are considered.

Let us assume first that $i$ is a susceptible agent at time $t$. The new value $p^*_i$ will change depending on the following two things: that an infection occurs if the other agent is infected, and the specific level of effort of agent $j$ if it is susceptible. We have

- $p^*_i = p_i - h_1 p_i$, when $i$ becomes infected after interacting with one of the $n - k$ infected agents. This event occurs with probability $\frac{2}{n} \frac{n-k}{n-1} \beta p_i$;
\(- p^*_i = p_i + h_2(1 - p_i)\) when \(i\) remains susceptible after interacting with one of the \(n - k\) infected agents, which occurs with probability \(\frac{2(n-k)}{n} (1 - \beta p_i)\);

\(- p^*_i = p_i - h(p_i - p_j)\) when \(i\) interacts with another susceptible agent \(j\), which occurs with probability \(\frac{2}{n(n-1)}\), for any susceptible agent \(j\) we can choose.

After replacing in Eq. (1) and rearranging terms, we get

\[
p_i(t + \Delta t) - p_i(t) = \frac{2\Delta t}{n(n-1)} \left[ -h_1(n-k)\beta p_i^2 + h_2(1 - p_i)(n-k)(1 - \beta p_i) \right. \\
- \left. h \sum_{j \in \text{Sus}} (p_i - p_j) \right].
\]  

(2)

**Remark 3.1** Let us remark that in the last term of the sum all the susceptible agents appear since the expected level of effort depends on the particular agent we choose, while in the first and second terms we can select any of the \(n - k\) infected agents and their parameters do not enter in the computation.

If the probability of contagion depends on the parameter of the infected agent, or on the parameters of both agents, we must change the probability of contagion and the selection of the involved agents accordingly, see Sect. 3.3 below.

Now we consider the case that \(i\) is an infected agent. The value \(p^*_i\) will depend on the specific parameter of the susceptible agent \(j\) we choose, and whether or not contagion occurs. The probabilities that \(i\) infects or not \(j\) are

\[
\text{Prob}(i \text{ infects } j) = \frac{2}{n(n-1)} \beta p_j,
\]

\[
\text{Prob}(i \text{ does not infect } j) = \frac{2}{n(n-1)} (1 - \beta p_j).
\]

We write again the expected change of \(p^*_i\) as a conditional probability on the effort of the particular susceptible agent:

\(- p^*_i = p_i - h_1 p_i\) when the other agent becomes infected, and this occurs with probability \(\frac{2\beta}{n(n-1)}\) for each susceptible agent \(j\);

\(- p^*_i = p_i + h_2(1 - p_i)\) when the other agent remains susceptible, and this occurs with probability \(\frac{2\beta p_j}{n(n-1)}\) for each susceptible agent \(j\);

\(- p^*_i = p_i - h(p_i - p_j)\) when \(i\) interacts with any other infected agent \(j\), which occurs with probability \(\frac{2(1 - \beta p_j)}{n(n-1)}\).

Hence, after replacing in the master Eq. (1) we obtain

\[
p_i(t + \Delta t) - p_i(t) = \frac{2\Delta t}{n(n-1)} \left[ -h_1 p_i \sum_{j \in \text{Sus}} p_j \beta + h_2(1 - p_i) \sum_{j \in \text{Sus}} (1 - p_j \beta) \right.
\]

\[
\left. - h \sum_{j \in \text{Sus}} (p_i - p_j) \right].
\]
\[-h \sum_{j \in \text{Inf}} (p_i - p_j)\].

We can write the last equation in a more compact way noticing that

\[
\sum_{j \in \text{Sus}} p_j = P_S, \quad \sum_{j \in \text{Sus}} (1 - p_j \beta) = k - \beta P_S,
\]

and we obtain

\[
p_i(t + \Delta t) - p_i(t) = \frac{2 \Delta t}{n(n-1)} \left[ -h_1 \beta p_i P_S + h_2 (1 - p_i) (k - \beta P_S) \right. \\
- \left. h \sum_{j \in \text{Inf}} (p_i - p_j) \right]. \tag{3}
\]

Finally, dividing by \(\Delta t\) in (2) and (3), and taking the limit as \(\Delta t \to 0\), we obtain an ordinary differential equation for \(p_i\) when \(i\) is a susceptible or infected agent,

\[
\frac{d}{dt} p_i(t) = \frac{2}{n(n-1)} \left[ -h_1 (n-k) \beta p_i^2 + h_2 (1 - p_i) (n-k) (1 - \beta p_i) - h \sum_{j \in \text{Inf}} (p_i - p_j) \right]. \tag{4}
\]

\[
\frac{d}{dt} p_i(t) = \frac{2}{n(n-1)} \left[ -h_1 \beta p_i P_S + h_2 (1 - p_i) (k - \beta P_S) - h \sum_{j \in \text{Inf}} (p_i - p_j) \right]. \tag{5}
\]

3.2 Derivation of the mean-field equations.

We can now obtain the equation for \(\langle p \rangle = \frac{1}{n} \sum_{i=1}^{n} p_i\), the mean value of \(p\). Splitting the sum as \(\sum_{i=1}^{n} p_i = \sum_{i \in \text{Sus}} p_i + \sum_{i \in \text{Inf}} p_i\), and using (4) and (5), we obtain

\[
\frac{n^2(n-1)}{2} \frac{d}{dt} (p(t)) = \sum_{i \in \text{Sus}} \left[ -h_1 (n-k) \beta p_i^2 + h_2 (1 - p_i) (n-k) (1 - \beta p_i) - h \sum_{j \in \text{Sus}} (p_i - p_j) \right. \\
- \left. h \sum_{j \in \text{Inf}} (p_i - p_j) \right] \\
+ \sum_{i \in \text{Inf}} \left[ -h_1 \beta p_i P_S + h_2 (1 - p_i) (k - \beta P_S) - h \sum_{j \in \text{Inf}} (p_i - p_j) \right].
\]

Noticing that

\[
\sum_{i \in \text{Sus}} \sum_{j \in \text{Sus}} (p_i - p_j) = \sum_{i \in \text{Inf}} \sum_{j \in \text{Inf}} (p_i - p_j) = 0.
\]
we observe that the parameter $h$ has no influence in $\langle p \rangle$ since the levels of effort of both agents get closer to each other and the mean does not change. Hence, the competition between fear and confidence drives the dynamics.

Letting $g := h_1 - h_2$ i.e. $h_1 = h_2 + g$, after rearranging terms we have

$$
\frac{n^2(n-1)}{2} \frac{d}{dr} \langle p(t) \rangle = h_2(n-k) \sum_{i \in \text{Sus}} (1 - (1+\beta) p_i) + h_2 \sum_{i \in \text{Inf}} (k - \beta P_S - p_i k) - \sum_{i \in \text{Sus}} g(n-k) \beta p_i^2 - \sum_{i \in \text{Inf}} g \beta p_i P_S \\
= h_2(n-k)[2k - (1 + 2\beta) P_S] - h_2 k \sum_{i \in \text{Inf}} p_i - \sum_{i \in \text{Sus}} g(n-k) \beta p_i^2 - \sum_{i \in \text{Inf}} g \beta p_i P_S.
$$

Writing

$$
\sum_{i \in \text{Inf}} p_i = \sum_{i=1}^n p_i - \sum_{i \in \text{Sus}} p_i = n \langle p \rangle - k \langle p_S \rangle = n(\langle p \rangle - \langle p_S \rangle) + (n-k) \langle p_S \rangle,
$$

we obtain

$$
\frac{n^2(n-1)}{2} \frac{d}{dr} \langle p(t) \rangle = 2 h_2(n-k) k[1 - (1+\beta) \langle p_S \rangle] - h_2 k n(\langle p \rangle - \langle p_S \rangle) \\
- g(n-k) \beta k [\langle p_S^2 \rangle - \sum_{i \in \text{Sus}} p_i] = h_2(n-k)[2k - (1 + 2\beta) P_S] - h_2 k \sum_{i \in \text{Inf}} p_i - \sum_{i \in \text{Sus}} g(n-k) \beta p_i^2 - \sum_{i \in \text{Inf}} g \beta p_i P_S \\
= 2 h_2 S(1-S)[1 - (1+\beta) \langle p_S(t) \rangle] - h_2 S(\langle p \rangle - \langle p_S \rangle) \\
- g \beta S \left( (1-S) \langle p_S^2 \rangle + \sum_{i \in \text{Inf}} p_i \right) \left( (\langle p \rangle - \langle p_S \rangle) + (1-S) \langle p_S \rangle \right).
$$

Finally, we can rescale time to get rid off the factor $n - 1$. Rearranging the terms we obtain the equation

$$
\frac{d}{dr} \langle p \rangle = 4 h_2 S(1-S)[1 - (1+\beta) \langle p_S \rangle] - 2 h_2 S(\langle p \rangle - \langle p_S \rangle) \\
- 2 g \beta S \left[(1-S) \left( \langle p_S^2 \rangle + \langle p_S \rangle \right) \right] \left( (\langle p \rangle - \langle p_S \rangle) \right). \tag{6}
$$

**Remark 3.2** Let us recall that $h_1 - h_2 = g$, so the last term is not present if $h_1 = h_2$.

**Remark 3.3** In much the same way, we get the equation

$$
\frac{d}{dr} S = (1-S)(\gamma - 2 \langle p_S \rangle \beta S). \tag{7}
$$
Its derivation can be found in the “Appendix”. It is similar to the usual derivation of the SIS model. The main differences with classical models are the mean susceptibility $\langle p_S \rangle \beta$, and the factor 2, which appears since both selected agents can infect or get infected (as opposed to the derivation of the classical SIS equation where only one agent, usually selected first, can get infected).

### 3.3 Other contagion rules

It is possible to derive the mean field equations for other contagion rules in a similar way. To this end, let us call

$$\langle p_I \rangle = \frac{1}{n-k} \sum_{j \in \text{Inf}} p_j, \quad \langle p_I^2 \rangle = \frac{1}{n-k} \sum_{j \in \text{Inf}} p_j^2. \quad (8)$$

Suppose that when a susceptible agent $i$ interacts with an infected agent $j$, contagion occurs with probability $p_j \beta$. In this case, after changing the probability of the different events accordingly, we get the following equations:

$$\frac{d}{dt} \langle p \rangle = 2h_2 S (1 - S) (2 - 2\beta \langle p_I \rangle - \langle p_S \rangle - \langle p_I \rangle) + 2(h_2 - h_1) \beta S (1 - S) (\langle p_I \rangle \langle p_S \rangle + \langle p_I^2 \rangle), \quad (8)$$

$$\frac{d}{dt} S = (1 - S) (\gamma - 2 \langle p_I \rangle \beta S). \quad (9)$$

On the other hand, if the probability of contagion depends on both effort levels, that is, contagion occurs with probability $p_I p_j \beta$, we get

$$\frac{d}{dt} \langle p \rangle = 2h_2 S (1 - S) (2 - \langle p_I \rangle - \langle p_S \rangle - 2\beta \langle p_S \rangle \langle p_I \rangle) + 2(h_2 - h_1) \beta (\langle p_S^2 \rangle \langle p_I \rangle + \langle p_S \rangle \langle p_I^2 \rangle), \quad (10)$$

$$\frac{d}{dt} S = (1 - S) (\gamma - 2 \langle p_I \rangle \langle p_S \rangle \beta S). \quad (11)$$

See the “Appendix” for details.

### 4 Qualitative study of the dynamic

Let us note that the definition of the microscopic interaction rules given in Sect. 2 concerning the level of effort implies that both susceptible and infected agents react in the same way. This suggests that their mean level of effort would be the same. The agent simulations in the left panel of Fig. 1 confirm this intuition, showing that the difference between $\langle p_S \rangle$ and $\langle p \rangle$ is of order $h$. Thus, taking $h \ll 1$, we will assume from now on that $\langle p_S \rangle = \langle p \rangle$. 

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**Coupling Epidemiological Models with Social Dynamics**

Page 11 of 23
Fig. 1 Agent based simulations with \( n = 10000 \) agents, \( \beta = 0.7, \gamma = 0.3 \), and \( p \) initially uniformly distributed in \([0.7, 0.9]\). Left panel: plot of \( \langle p_S \rangle - \langle p \rangle \) for \( h_1 = h_2 = h = 0.1, 0.01, 0.001, \) and \( 0.0001 \). Right panel: plot of the variance \( \text{Var}(p) = \langle p^2 \rangle - \langle p \rangle^2 \) for \( h_1 = h_2 = h = 0.01, 0.001, \) and \( 0.0001 \)

Notice the Eq. (6) for \( \langle p \rangle \) involves \( \langle p^2 \rangle \), and an equation for \( \langle p^2 \rangle \) would involve \( \langle p^3 \rangle \), and so on, thus resulting in an infinite hierarchy of equations. In order to avoid the infinite hierarchy of moments, we consider two options. The first and simplest one is to limit ourselves to the case \( h_1 = h_2 \). Indeed in that case, \( g = 0 \) and, recalling we assumed \( \langle p_S \rangle = \langle p \rangle \), Eq. (6) reduces to

\[
\frac{d}{dt} \langle p \rangle = 4h_2S(1 - S)[1 - (1 + \beta)\langle p \rangle].
\] (12)

The second one comes from the fact that the variance \( \text{Var}(p) = \langle p^2 \rangle - \langle p \rangle^2 \) goes to zero, so \( \langle p^2 \rangle \sim \langle p \rangle^2 \), and this enables us to deal also with the case \( h_1 \neq h_2 \). We show in the right panel in Fig. 1 the evolution of \( \text{Var}(p) \) for three values of \( h \), and \( h = h_1 = h_2 \). We observe in the simulations that \( \text{Var}(p) = o(h) \).

Moreover, let us note that \( h \) disappeared in the limit equation since \( \langle p \rangle \) is invariant under this interaction. Hence, by considering the approximation \( h \gg h_1, h_2 \), a consensus is reached in a shorter time scale, see for instance (Pinasco et al. 2017).

4.1 The case \( h_1 = h_2 \)

Since \( g = 0 \), and recalling we assumed \( \langle p_S \rangle = \langle p \rangle \), Eqs. (6) and (7) for \( \langle p \rangle \) and \( S \) become

\[
\frac{d}{dt} \langle p \rangle = 4h_2(1 - S)S[1 - (1 + \beta)\langle p \rangle],
\] (13)

\[
\frac{d}{dt} S = (1 - S)\left(\gamma - 2\langle p \rangle\beta S\right).
\] (14)

Notice that the square \([0, 1] \times [0, 1]\) is clearly invariant for the system, and that its fixed points in \([0, 1] \times [0, 1]\) are the line \( S = 1 \) and the point \( (\langle p \rangle, S) = \left(\frac{1}{1 + \beta}, \frac{\gamma(1 + \beta)}{2\beta}\right) \) with \( \frac{\gamma(1 + \beta)}{2\beta} \leq 1 \).

Let us study the asymptotic behaviour of a solution starting from a point \( (\langle p \rangle(0), S(0)) \in [0, 1] \times [0, 1] \). Since \( S(t) < 1 \) for any \( t \) and noticing that \( \frac{d}{dt} \langle p \rangle \)
has same sign as $1 - (1 + \beta)\langle p \rangle$, we see that

$$\lim_{t \to +\infty} \langle p \rangle = p_\infty = \frac{1}{1 + \beta}.$$  

We can then rewrite (14) as

$$\frac{1}{2\beta} \frac{d}{dt} S = \langle p \rangle (1 - S) \left( \frac{\gamma}{2\beta \langle p \rangle} - S \right) = \langle p \rangle (1 - S) \left( R_m^{-1} + \varepsilon(t) - S \right),$$

where $R_m^{-1} := \frac{\gamma(1+\beta)}{2\beta}$ and $\varepsilon(t) \xrightarrow{t \to +\infty} 0$. We now distinguish three cases.

If $R_m^{-1} > 1$ then there exists $T > 0$ such that for any $t \geq T$, $R_m^{-1} + \varepsilon(t) \geq 1$. It follows that for $t \geq T$ we have $\frac{d}{dt} S \geq 0$ with equality only when $S = 1$. We conclude that $\lim_{t \to +\infty} S(t) = 1$.

If $R_m^{-1} < 1$ then for any $a, b \in [0, 1), a < R_m^{-1} < b$, there exists $T' > 0$ such that for $t \geq T'$ we have

$$\langle p \rangle (1 - s) \left( R_m^{-1} + \varepsilon(t) - s \right) > 0 \quad \forall s \in [0, a],$$

$$\langle p \rangle (1 - s) \left( R_m^{-1} + \varepsilon(t) - s \right) < 0 \quad \forall s \in [b, 1)$$

Thus $S$ enters the interval $[a, b]$ at some time $T \geq T'$ and stays there forever. Since this holds for any $a, b \in [0, 1)$ with $a < R_m^{-1} < b$, we deduce that $\lim_{t \to +\infty} S(t) = R_m^{-1}$.

If $R_m = 1$ then for any $\delta \in (0, 1)$, there exists $T' > 0$ such that for $t \geq T'$, $|\varepsilon(t)| < \delta/2$ so that

$$\langle p \rangle (1 - s) \left( R_m^{-1} + \varepsilon(t) - s \right) > 0 \quad \forall s \in [0, 1 - \delta].$$

Thus if $S(0) < 1 - \delta$, then $S(t)$ must enter the interval $[1 - \delta, 1]$ at some time $T \geq T'$ and stay there forever. Since this holds for any $\delta > 0$, we deduce that $\lim_{t \to +\infty} S(t) = 1$.

We can summarize the previous discussion in the following Theorem.

**Theorem 4.1** For any initial condition $(\langle p \rangle(0), S(0)) \in [0, 1] \times [0, 1]$, the solution $(\langle p \rangle, S)$ of (13)-(14) satisfies

$$\lim_{t \to +\infty} \langle p \rangle = p_\infty = \frac{1}{1 + \beta}.$$  

If $S(0) = 1$ then $S(t) = 1$ for any $t \geq 0$, and if $S(0) < 1$ then

$$\lim_{t \to +\infty} S(t) = \begin{cases} 1 & \text{if } R_m \leq 1, \\ R_m^{-1} & \text{if } R_m > 1, \end{cases}$$
where $R_m = \frac{2\beta}{\gamma(1+\beta)}$.

### 4.2 The case $h_1 \neq h_2$

Assuming that $h \gg h_1, h_2$, or $\langle p^2 \rangle \sim \langle p \rangle^2$, Eqs. (6) and (7) for $\langle p \rangle$ and $S$ read now

\[
\frac{d}{dt} \langle p \rangle = 4h_2S(1 - S)[1 - (1 + \beta)\langle p \rangle + (1 - h_1/h_2)\beta\langle p \rangle^2],
\]

\[
\frac{d}{dt} S = (1 - S)(\gamma - 2\langle p \rangle\beta S).
\]

We consider first the case $h_1 > h_2$ where fear dominates confidence. It is easily seen that there exists $p^-_\infty \in (0, (1 + \beta)^{-1})$ such that

\[
\frac{d}{dt} \langle p \rangle \begin{cases} > 0 & \text{if } \langle p \rangle \in [0, p^-_\infty) \\ < 0 & \text{if } \langle p \rangle \in [p^-_\infty, (1 + \beta)^{-1}) \end{cases}
\]

Thus $\langle p \rangle \to p^-_\infty$. Hence, there exists a unique positive equilibrium $p^-_\infty$ for the first equation, which is stable, and $0 < p^-_\infty < (1 + \beta)^{-1}$. The rest of the analysis follows as before, replacing $R_m$ by $R^-_m = p^-_\infty R_0$.

In the same way, when $h_1 < h_2$, agents relax their protection measures, and there exists a unique positive equilibrium $p^+_\infty$ to Eq. (15), which is stable and $p^+_\infty > (1 + \beta)^{-1}$. Again, we can repeat the rest of the analysis as before, obtaining $R^+_m = p^+_\infty R_0$.

Let us note that $p^-_\infty < (1 + \beta)^{-1} < p^+_\infty$ implies that, for $h_1 > h_2$, people attain a level of effort that helps to mitigate the propagation of the disease better than the corresponding one for $h_1 = h_2$ - the opposite occurs when $h_1 < h_2$.

### 4.3 Other contagion rules

We omit the analysis of the alternative rules presented before in Sect. 3.3 since it is similar. Let us mention only that, for $h_1 = h_2$, and assuming $\langle p_I \rangle = \langle p_S \rangle = \langle p \rangle$, we get

\[
\frac{d}{dt} \langle p \rangle = 4h_2S(1 - S)(1 - (1 + \beta)\langle p \rangle),
\]

\[
\frac{d}{dt} S = (1 - S)(\gamma - 2\langle p \rangle\beta S).
\]

when the contagion depends on the level of effort of infected individuals. The equilibria and their stability are the same as before.

If the contagion depends on the level of effort of both agents, we obtain

\[
\frac{d}{dt} \langle p \rangle = 4h_2S(1 - S)(1 - \langle p \rangle - \beta\langle p \rangle^2),
\]

\[
\frac{d}{dt} S = (1 - S)(\gamma - 2\langle p \rangle^2\beta S).
\]
The first equation has a single positive equilibria

$$\langle p \rangle_\infty = \frac{\sqrt{1+4\beta} - 1}{2\beta} \in (0, 1),$$

and $\langle p \rangle \to \langle p \rangle_\infty$ when $t \to +\infty$ if initially $S(0) \in (0, 1)$. In order to compare $\langle p \rangle_\infty$ and the limit number of Susceptible individuals in the endemic equilibria, namely $\frac{\gamma}{2 \langle p \rangle_\infty^2 \beta}$, with what happens when contagion depends on the susceptible individual only, notice that

$$\frac{\sqrt{1+4\beta} - 1}{2\beta} > \frac{1}{1 + \beta}, \quad \text{and} \quad \left(\frac{\sqrt{1+4\beta} - 1}{2\beta}\right)^2 < \frac{1}{1 + \beta}.$$ 

It follows that if contagion depends on the level of effort of both agents, they asymptotically implement more relaxed measures of protection, and the combined efforts contribute lead to a bigger number of limit susceptible individuals.

### 5 The behaviourally reduced reproduction number $R_m$ and simulations

A critical parameter in compartmental epidemic models is $R_0$, the basic reproduction number, which gives the expected number of contagions associated to a single infected individual who enters in a susceptible population. In the classical SIS model,

$$R_0 = \frac{2\beta}{\gamma},$$

see for instance (Anderson and May 1992) (recall that the factor 2 appears since both the first or second agent can become infected). Then, as time evolves, the effective reproduction number $R_t = R_0 S(t)$ is a better descriptor of the dynamics since it takes into account that there are less susceptible agents. In the SIS model, $R_t$ stabilizes at $R_t = 1$ in the endemic phase, and an equilibrium between recovered and new infected individuals appears, reaching a stable number of infected agents.

In the model studied in this paper, we observe from Eq. (14) that at time $t$ we have an effective reproduction number

$$R_t = \langle p(t) \rangle \frac{2\beta}{\gamma} S(t).$$

Since $\langle p(t) \rangle \to p_\infty$ for $h_1 = h_2$, and to $p_\infty^\pm$ if $h_1 \neq h_2$, we introduce the behaviourally reduced reproduction numbers

$$R_m = p_\infty \frac{2\beta}{\gamma} \quad \text{and} \quad R_m^\pm = p_\infty^\pm \frac{2\beta}{\gamma}.$$
Fig. 2 Plot of the proportion of Infected agents at the stable equilibria, for $\gamma = 0.6$ and varying $\beta$. The dotted blue curve represent the equilibria for the classical SIS dynamics, the black one are the theoretical equilibria computed in Theorem 4.1, and the red stars are agent-based simulations of the dynamics with 10000 agents and $p$ uniformly distributed in $[0.7, 0.9]$

It follows from the previous results that given an epidemic outbreak, i.e. $S(0) < 1$, then the disease tends to disappear if $R_m \leq 1$ whereas it becomes endemic if $R_m > 1$ - the same is valid for $R_m^\pm$. In Fig. 2, for $\gamma = 0.6$, we compare the bifurcation diagram of the classical SIS model with the one presented in this work. Varying $\beta$, we observe the bifurcation at $R_0 = 2\beta/\gamma$ of the classical SIS model (blue dotted curve). In contrast, the role played by the social interactions in our model delays the bifurcation until $R_m = R_0/(1 + \beta)$, and effectively lowers the final number of infected agents in the endemic state (black line). The red stars in the diagram correspond to agent-based simulations of the dynamics. We observe a very good agreement with the predicted value of infected agents at equilibria given by Theorem 4.1.

To further appreciate the agreement between the theoretical results of Theorem 4.1 and the agent-based simulations, we present in Fig. 3 a heat map for the proportion of susceptible agents at equilibria obtained in Theorem 4.1 (left panel), and performing agent-based simulations (right panel). We observe in the left panel that the region in the space of parameters where the disease is endemic is smaller than the corresponding one for a classical SIS model, delimited with the yellow line $R_0 = 1$. Also, the agent-based simulations in the right panel confirm that the ODE system (13)–(14) accurately models the dynamics.

6 Conclusions and Final Remarks

We derived an epidemic model coupled with a continuous opinion dynamics model. We assumed that each individual can take measures to reduce the probability of con-
Fig. 3 Heat map of the proportion of susceptible agents at equilibria. Upper panel: theoretical values from Theorem 4.1. Lower panel: agent based simulation of the dynamics with 10000 agents and $p$ initially uniformly distributed in $[0.7, 0.9]$. The yellow line correspond to $R_0 = 1$ in a classical SIS model.

tagion, and the level of effort that each agent applies change due to social interactions. We model few mechanisms, fear to contagion, confidence after a contact without contagion, and persuasion, as the main reasons for behavioral change, and we studied their impact on the dynamics of the disease.

We obtained a system of two ordinary differential equations, one for the proportion of infected people, and another one for the mean value of the effort parameter. The mean value of the effort enters as a coefficient in the original SIS system which modifies the rate of contagion. At the same time it changes as a consequence of epidemic events,
since it decreases or increases when a contagion occurs or not. Hence, the behavioral dynamics modify the parameters of the SIS system. We can compare the system with other behavioral epidemic models, where an additional equation in a SIS or SIR model as in Bauch (2005) represent new compartment, and the behavioral dynamics drags the susceptible population into the vaccinated compartment. Different factors like imitation, costs of vaccination, or perceived risk due to the infected population, appears in the coefficients of this additional equation, which is a replicator equation with varying payoffs since depends on the proportion of infected agents.

We study the asymptotic behaviour of the solutions, and we obtained a generalization of the basic reproduction number $R_0$, denoted $R_m$, and given by

$$R_m = \frac{2\beta}{\gamma(1 + \beta)}$$

when $h_1 = h_2$, and where $\beta$ and $\gamma$ are the contact and recovery rate of the disease. We proved the existence of similar critical values $R_{m}^{\pm}$ when $h_1 \neq h_2$. Let us recall that the factor 2 in $R_m$ appears since both the first and second agents can become infected after an interaction. We prefer to keep this for simplicity and symmetry in the formulation of the problem. Notice however that comparison with the $R_0$ of classical models must include the factor 2 (that is, if the rate of contagion of a disease studied with our model is $\beta$, this is equivalent to a rate $2\beta$ when only the first agent in some encounter can be infected).

There are several questions of practical interest, particularly in this moment with the Covid-19 pandemic active. To be clear, we are not claiming the direct applicability of our specific rules of social interaction, although trends in the population can be detected (like the degree of use of masks, sanitizer products, or social distance), thus giving a personal probability of contagion $\beta_i$, which can be studied as the product $p_i\beta$ in our model. Of course, different questions are relevant if we consider Covid-19, since a SEIR model (including exposed and removed agents) seems to be a better one to analyze its evolution.

We have chosen the same scale of time for both the epidemic and social dynamics, scaling only the number of agents, and keeping the parameter $h_2$ in the equation for $\langle p \rangle$. It is possible to change the frequency of interactions, by separating the ones related to the disease transmission, to the social ones ones which only affect the level of effort. This is specially important in SIR type dynamics, since the interactions at the start of the epidemic are critical for its evolution.

Also, fundamental agents, like e.g. governments, media, health organizations, were not considered here, though their role as social agents interacting with all the population cannot be neglected. On the negative side of social interactions, there are groups of people proposing innocuous and even harmful measures, like anti-vaccines movements, and we can see them today violating the social distance, or without masks, and trying to convince other people to imitate them. They act as zealots or stubborn agents and in a forthcoming paper we will study their role and their influence on the stability of equilibria.

**Author Contributions** All the authors contributed equally.
Declaration

Conflict of interest The authors declare that they have no conflict of interest.

Appendix

We briefly present the derivation of the mean-field Eq. (7), and Eqs. (8)–(11). We closely follow the derivation presented in Sects. 3.1 and 3.2.

Contagion depending on the level of effort of the infected agent.

If agent \( i \) is susceptible then

\[
\frac{n(n-1)}{2} p_i(t + \Delta t) - p_i(t) \Delta t = -h_1 p_i \sum_{j \in \text{Inf}} \beta \ p_j + h_2 (1-p_i) \sum_{j \in \text{Sus}} (1 - \beta p_j) + h \sum_{j \in \text{Sus}} (p_j - p_i)
\]

\[
= -h_2 p_i (n-k) \langle p_I \rangle + h_2 (n-k)(1-p_i)(1 - \beta \langle p_I \rangle) + h \sum_{j \in \text{Sus}} (p_j - p_i).
\]

Next if agent \( i \) is infected,

\[
\frac{n(n-1)}{2} p_i(t + \Delta t) - p_i(t) \Delta t = -h_1 p_i \sum_{j \in \text{Sus}} \beta p_i + h_2 (1-p_i) \sum_{j \in \text{Sus}} (1 - \beta p_i) + h \sum_{j \in \text{Inf}} (p_j - p_i)
\]

\[
= -h_1 \beta p_i^2 k + h_2 k (1-p_i)(1 - \beta p_i) + h \sum_{j \in \text{Inf}} (p_j - p_i).
\]

Summing over \( i \), by using that \( \sum_{i,j \in \text{Sus}} (p_j - p_i) = \sum_{i,j \in \text{Inf}} (p_j - p_i) = 0 \), after taking the limit \( \Delta t \to 0 \), we obtain

\[
\frac{n^2(n-1)}{2} \frac{d}{dt} \langle p \rangle = -h_1 \beta (n-k) \langle p_I \rangle \sum_{i \in \text{Sus}} p_i + h_2 (n-k) \sum_{i \in \text{Sus}} (1-p_i)(1 - \beta \langle p_I \rangle)
\]

\[
- h_1 \beta k \sum_{i \in \text{Inf}} p_i^2 + h_2 k \sum_{i \in \text{Inf}} (1-p_i)(1 - \beta p_i),
\]

i.e.

\[
\frac{n^2(n-1)}{2} \frac{d}{dt} \langle p \rangle = -h_1 \beta k (n-k)(\langle p_I \rangle \langle p_S \rangle + \langle p_I^2 \rangle)
\]

\[
+ h_2 k (n-k)(2 - (1+2\beta)\langle p_I \rangle - \langle p_S \rangle + \beta \langle p_I \rangle \langle p_S \rangle + \beta \langle p_I^2 \rangle).
\]
Dividing by $n^2$ gives

$$\frac{n - 1}{2} \frac{d}{dt} \langle p \rangle = (h_2 - h_1) \beta S (1 - S) \left( \langle p_I \rangle \langle p_S \rangle + \langle p_I^2 \rangle \right) + h_2 S (1 - S) \left( 2 - (1 + 2\beta) \langle p_I \rangle - \langle p_S \rangle \right),$$

and after rearranging and rescaling $t$ we get Eq. (8).

**Contagion depending on the level of effort of both agents.**

As before, if agent $i$ is susceptible then

$$\frac{n(n - 1)}{2} p_i (t + \Delta t) - p_i (t) \Delta t = -h_1 p_i \sum_{j \in \text{Inf}} \beta p_i p_j + h_2 (1 - p_i) \sum_{j \in \text{Inf}} (1 - \beta p_i p_j) + h \sum_{j \in \text{Sus}} (p_j - p_i)$$

$$= -h_1 \beta p_i^2 (n - k) \langle p_I \rangle + h_2 (n - k) (1 - p_i) (1 - \beta p_i \langle p_I \rangle) + h \sum_{j \in \text{Sus}} (p_j - p_i).$$

Now, if agent $i$ is infected,

$$\frac{n(n - 1)}{2} p_i (t + \Delta t) - p_i (t) \Delta t = -h_1 p_i \sum_{j \in \text{Sus}} \beta p_i p_j + h_2 (1 - p_i) \sum_{j \in \text{Sus}} (1 - \beta p_i p_j) + h \sum_{j \in \text{Inf}} (p_j - p_i)$$

$$= -h_1 \beta p_i^2 k \langle p_S \rangle + h_2 k (1 - p_i) (1 - \beta p_i \langle p_S \rangle) + h \sum_{j \in \text{Inf}} (p_j - p_i).$$

Summing over $i$, by using that $\sum_{i,j \in \text{Sus}} (p_j - p_i) = \sum_{i,j \in \text{Inf}} (p_j - p_i) = 0$, we obtain

$$\frac{n^2 (n - 1)}{2} \frac{d}{dt} \langle p \rangle = -h_1 \beta (n - k) \langle p_I \rangle \sum_{i \in \text{Sus}} p_i^2 + h_2 (n - k) \sum_{i \in \text{Sus}} (1 - p_i) (1 - \beta p_i \langle p_I \rangle)$$

$$- h_1 \beta k \langle p_S \rangle \sum_{i \in \text{Inf}} p_i^2 + h_2 k \sum_{i \in \text{Inf}} (1 - p_i) (1 - \beta p_i \langle p_S \rangle),$$

i.e.

$$\frac{n^2 (n - 1)}{2} \frac{d}{dt} \langle p \rangle = -h_1 \beta k (n - k) (\langle p_I \rangle \langle p_S^2 \rangle + \langle p_I^2 \rangle \langle p_S \rangle)$$

$$+ h_2 k (n - k) (2 - 2 \beta \langle p_S \rangle \langle p_I \rangle - \langle p_S \rangle - \langle p_I \rangle + \beta \langle p_S^2 \rangle \langle p_I \rangle + \beta \langle p_S \rangle \langle p_I^2 \rangle).$$
Dividing by $n^2$ gives

\[
\frac{n - 1}{2} \frac{d}{dr} \langle p \rangle = (h_2 - h_1)\beta S(1 - S) \left( \langle p_I \rangle \langle p_S \rangle^2 + \langle p_I \rangle \langle p_S \rangle \right) \\
+ h_2 S(1 - S) \left( 2 - 2\beta \langle p_S \rangle \langle p_I \rangle - \langle p_S \rangle - \langle p_I \rangle \right),
\]

and after rearranging and rescaling $t$ we get Eq. (10).

### The SIS model with a single agent effort

Let us note that after an interaction the value of $S$ can remain the same or change by an amount of $\pm \frac{1}{N}$. The transition $S \rightarrow S + \frac{1}{N}$ happens when an infected agent is cured, and occurs with probability

\[
\text{Prob}(S \rightarrow S + \frac{1}{N}) = \sum_{i \in \text{Inf}} \frac{\gamma}{N} = \gamma(1 - S).
\]

On the other hand the transition $S \rightarrow S - \frac{1}{N}$ happens when a susceptible agent becomes infected upon contact with an infected one, and this occurs with the following probabilities:

- when the contagion depends on the level of effort of the susceptible agent,

\[
\text{Prob}(S \rightarrow S - \frac{1}{N}) = \sum_{i \in \text{Sus}} \sum_{j \in \text{Inf}} \frac{2}{n(n - 1)} p_i \beta = \frac{2k(n - k)}{n(n - 1)} \beta \langle p_S \rangle \\
\approx 2\beta S(1 - S)(p_S);
\]

- when the contagion depends on the level of effort of the infected agent,

\[
\text{Prob}(S \rightarrow S - \frac{1}{N}) = \sum_{i \in \text{Sus}} \sum_{j \in \text{Inf}} \frac{2}{n(n - 1)} p_j \beta = \frac{2k(n - k)}{n(n - 1)} \beta \langle p_I \rangle \\
\approx 2\beta S(1 - S)(p_I);
\]

- when the contagion depends on the levels of effort of both agents,

\[
\text{Prob}(S \rightarrow S - \frac{1}{N}) = \sum_{i \in \text{Sus}} \sum_{j \in \text{Inf}} \frac{2}{n(n - 1)} p_i p_j \beta = \frac{2k(n - k)}{n(n - 1)} \beta \langle p_S \rangle \langle p_I \rangle \\
\approx 2\beta S(1 - S)(p_S)(p_I).
\]

We deduce the equation for the variation of $S$ for the first case:

\[
\frac{d}{dr} S = \frac{1}{N} (1 - S)(\gamma - 2\beta S(p_S)).
\]
and in the other two the factor $\langle p_S \rangle$ changes to $\langle p_I \rangle$ and $\langle p_S \rangle \langle p_I \rangle$ respectively.

Rescaling time to absorb the factor $1/N$ so that $N$ interactions are done in one time unit (which is asymptotically the time rescaling we did in the final step of the derivation of the equation for $\langle p \rangle$), yields

$$\frac{d}{dt} S = (1 - S)(\gamma - 2\beta \langle p_S \rangle),$$

and the corresponding ones in the other two cases.

References

Aletti G, Naldi G, Toscani G (2007) First-order continuous models of opinion formation. SIAM J Appl Math 67(3):837–853
Anderson RM, May RM (1992) Infectious diseases of humans: dynamics and control. Oxford University Press
Asch SE (1955) Opinions and social pressure. Sci Am 193(5):31–35
Axelrod R (1997) The dissemination of culture: a model with local convergence and global polarization. J Conflict Resolut 41(2):203–226
Bauch CT (2005) Imitation dynamics predict vaccinating behaviour. Proc R Soc B Biol Sci 272(1573):1669–1675
Bauch CT, Earn DJ (2004) Vaccination and the theory of games. Proc Nat Acad Sci 101(36):13391–13394
Bellomo N (2008) Modeling complex living systems: a kinetic theory and stochastic game approach. Springer
Bellomo N, Marsan GA, Tosin A (2013) Complex systems and society: modeling and simulation. Springer
Burnstein E, Vinokur A (1977) Persuasive argumentation and social comparison as determinants of attitude polarization. J Exp Soc Psychol 13(4):315–332
Clancy D (2018) Precise estimates of persistence time for sis infections in heterogeneous populations. Bull Math Biol 80(11):2871–2896
Defuant G, Neau D, Amblard F, Weisbuch G (2000) Mixing beliefs among interacting agents. Adv Complex Syst 3(01n04):87–98
Dorso CO, Medus A, Balenzuela P (2017) Vaccination and public trust: a model for the dissemination of vaccination behaviour with external intervention. Physica A 482:433–443
Fauci AS, Morens DM (2016) Zika virus in the Americas—yet another arbovirus threat. N Engl J Med 374(7):601–604
Funk S, Gilad E, Watkins C, Jansen VA (2009) The spread of awareness and its impact on epidemic outbreaks. Proc Nat Acad Sci 106(16):6872–6877
Galam S, Zucker JD (2000) From individual choice to group decision-making. Physica A 287(3–4):644–659
Misra AK, Sharma A, Shukla JB (2011) Modeling and analysis of effects of awareness programs by media on the spread of infectious diseases. Math Comput Modell 53(5–6):1221–1228
Ochombok R et al (2001) Simulation of Sznaid sociophysics model with convincing single opinions. Int J Mod Phys C 12(7):1091–1092
Pedraza L, Pinasco JP, Saintier N (2020) Measure-valued opinion dynamics. Math Models Methods Appl Sci 30(02):225–260
Pérez-Llanos M, Pinasco JP, Saintier N, Silva A (2018) Opinion formation models with heterogeneous persuasion and zealotry. SIAM J Math Anal 50(5):4812–4837
Pinasco JP, Semeshenko V, Balenzuela P (2017) Modeling opinion dynamics: theoretical analysis and continuous approximation. Chaos Solitons Fractals 98:210–215
Pires MA, Crokidakis N (2017) Dynamics of epidemic spreading with vaccination: impact of social pressure and engagement. Physica A 467:167–179
da Silva PCV, Velásquez-Rojas F, Connaughton C, Vazquez F, Moreno Y, Rodrigues FA (2019) Epidemic spreading with awareness and different timescales in multiplex networks. Phys Rev E 100(3):032313
Slanina F, Lavicka H (2003) Analytical results for the sznaid model of opinion formation. Eur Phys J B Condens Matter Complex Syst 35(2):279–288

Springer
Su Z, Wang W, Li L, Stanley HE, Braunstein LA (2018) Optimal community structure for social contagions. New J Phys 20(5):053053
Sznajd-Weron K, Sznajd J (2000) Opinion evolution in closed community. Int J Mod Phys C 11(06):1157–1165
Toscani G et al (2006) Kinetic models of opinion formation. Commun Math Sci 4(3):481–496
Tyson RC, Hamilton SD, Lo AS, Baumgaertner BO, Krone SM (2020) The timing and nature of behavioural responses affect the course of an epidemic. Bull Math Biol 82(14):1–28
Yang C, Wang X, Gao D, Wang J (2017) Impact of awareness programs on cholera dynamics: two modeling approaches. Bull Math Biol 79(9):2109–2131
Zhou Y, Zhou J, Chen G, Stanley HE (2019) Effective degree theory for awareness and epidemic spreading on multiplex networks. New J Phys 21(3):035002

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