Epidemic spreading in modular time-varying networks

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We investigate the effects of modular and temporal connectivity patterns on epidemic spreading. To this end, we introduce and analytically characterise a model of time-varying networks with tunable modularity. Within this framework, we study the epidemic size of Susceptible-Infected-Recovered, SIR, models and the epidemic threshold of Susceptible-Infected-Susceptible, SIS, models. Interestingly, we find that while the presence of tightly connected clusters inhibits SIR processes, it speeds up SIS phenomena. In this case, we observe that modular structures induce a reduction of the threshold with respect to time-varying networks without communities. We confirm the theoretical results by means of extensive numerical simulations both on synthetic graphs as well as on a real modular and temporal network.

Network thinking has become a prominent and convenient paradigm to unveil the properties of complex systems. Such a paradigm has been rapidly enriched, giving rise to variants that account for inherent features of real complex systems inferred by the availability of large, often time-resolved datasets1–4. Three are the main features that have captured the attention of researchers in the area. The first is heterogeneity in the statistical distributions of key topological properties such as the number of connections per node (degree) and the intensity of interactions (weight)3,4. This property is one of the hallmarks of complexity and is linked to a range of non-trivial dynamics3,5. For example, heterogeneity in the connectivity patterns makes networks extremely fragile to the spreading of infectious diseases and malicious attacks6,7. The second feature regards the presence of modules and communities8. Available datasets have highlighted that real networks are organized in modules, or communities, whereby the density of links within the community is much greater than the density of links between communities. On the one hand, communities can be treated as fairly independent entities within a large network, like the behaviour of different organs within the same body. On the other hand, from time to time, phenomena originating in a community may involve a huge portion, if not all, of the network. This is for example the case of pandemics originating from local outbreaks6,9–11. Nevertheless, the role played communities is still ambivalent. For example, the presence of communities might slow down or speed up the propagation of a disease and facilitate the spreading of social norms12–20. Detecting communities in a real system is not a trivial task, due to their fuzzy, vanishing, and overlapping nature. Also, in most of the available datasets communities are not explicitly labeled, so the validation of community detection algorithms cannot often be rigorously carried out21. Finally, networks are characterised by non trivial temporal dynamics22,23. The propensity of nodes to initiate and to attract interactions per unit time is typically heterogeneously distributed24,25. The same applies to the duration and the time interval between connections13,26. Furthermore, the creation or renewal of interactions might be correlated, and the dynamics driving the temporal evolution of networks are function of the time-scale considered27,28. However, the large majority of studies on dynamical processes unfolding on networks have been conducted under the hypothesis of time scale separation which effectively neglects all such features. In particular, the evolution of the process and the evolution of network are considered to take place at well-distinct time scales. Within this paradigm two opposite limits have been considered. In the first case, the dynamical process is assumed to be much faster than the evolution of the network. This is the limit of quenched/static networks29. Here, networks are fully characterised from an adjacency...
At the next time step $P$ be only slightly smaller than the extracted value. The empirical distribution of community sizes in the network is much smaller than the total number of nodes, thus the actual size of the last community can average value of last extraction we might have available only a fraction of the nodes necessary to fill the community. However, the one community. Very rarely, all nodes can be perfectly assigned to the extracted community sizes. Indeed, in the parameters. The assignment of nodes to communities is done as follow. A community of size $s$ is extracted from a heavy-tailed distribution, i.e. $F(s)$, where $s \in [\epsilon, 1]$ and $\epsilon = 10^{-5}$ to avoid divergence in the distribution. Furthermore, each node is assigned to only one group/community. To take into account empirical evidences, the size of each community is extracted from a heavy-tailed distribution, i.e. $P(s) \sim s^{-\omega}$ with $s \in [s_{\text{min}}, \sqrt{N}]$, $\omega \approx 8.54$. Therefore, we do not limit ourselves in studying a fixed number of modules, whilst their number is driven from the model's parameters. The assignment of nodes to communities is done as follow. A community of size $s$ is extracted. The ID of $s$ nodes is progressively assigned to the community. The processes is repeated until all $N$ nodes are assigned to one community. Very rarely, all nodes can be perfectly assigned to the extracted community sizes. Indeed, in the last extraction we might have available only a fraction of the nodes necessary to fill the community. However, the average value of $s$ is much smaller than the total number of nodes, thus the actual size of the last community can be only slightly smaller than the extracted value. The empirical distribution of community sizes in the network will then follow $P(s)$. Given these settings, a generative network model is defined by the following steps (see Fig. 1).

- At each time $t$, the network, $G_t$, starts with $N$ disconnected nodes.
- With probability $a \Delta t$ each vertex $i$ is active and willing to create $m$ connections.
- Each link being generated points with probability $\mu$ within the node's community, and with probability $1 - \mu$ to one of any other groups. In both cases, the target node $j$ of the link is randomly selected in the target community.
- At the next time step $t + \Delta t$ all the edges in $G_t$ are deleted.

All the interactions have a constant duration $\Delta t$. In the model, neither self-loops nor multiple edges are allowed. In the following, without loss of generality, we fix $\Delta t = 1$. Furthermore, we consider the case $m = 1$.

Given an heterogeneous distribution of activity, at each time step, the model generates a random, structureless network in which few nodes are active. The modular features of the network emerge integrating connections in time. Such time-integrated properties, at different time regimes, can be computed analytically.
we will report the results for the evolution of the average number of connections of each node \( \langle k_i(t) \rangle \) (average degree) and the overall degree distribution \( \rho(k) \). The complete set of results is shown in the Supplementary Information (SI).

To solve the average degree’s dynamics, let us introduce the effective activity \( \bar{a} = a + \langle a \rangle \) (where \( \langle a \rangle \) is the average value of the activity distribution) and the mixing parameter \( \mu' = 1 - \mu \). We refer to the degree of node \( i \) at time \( t \) as \( k_i(a_i, s, t) \), where \( s \) is the node’s community size. By defining an activity class as the group of nodes featuring similar activity values \( a \), we set the average in-community degree \( \langle k_c(a, s, t) \rangle \) to be the average number of connections that nodes belonging to the activity class \( a \) and falling in communities of size \( s \) have toward nodes of their same community. The latter grows as

\[
\tau(a, s) = \frac{ka s t}{\mu' \tau(a, s)} \sim \frac{ka s t}{\mu' \tau(a, s)} \quad (1)
\]

where \( \tau(a, s) \) is the characteristic time that it takes for the degree \( k_c(a, s, t) \) of nodes of activity \( a \) belonging to a community of size \( s \) to be \( \sim (s-1) \), being \( s-1 \) the maximum value of the in-community degree (see the Supplementary Information for the evaluation of \( \tau(a, s) \)).

Similarly, we can define the average out-community degree \( \langle k_o(a, t) \rangle \) as the number of connections that nodes of activity class \( a \) have outside of their communities at time \( t \). We expect this quantity to be independent of the nodes’ community size \( s \) so that, for large networks, we can write:

\[
\langle k_o(a, t) \rangle = \mu' \bar{a} t \quad (2)
\]

The average total degree \( \langle k(a, s, t) \rangle \) can be computed as the simple sum between the two previous equations, obtaining

\[
\langle k(a, s, t) \rangle = \langle k_c(a, s, t) \rangle + \langle k_o(a, t) \rangle \cong \begin{cases} \bar{a} t \quad t \ll \tau(a, s) \\ \mu' \bar{a} t + (s-1) t \sim \tau(a, s) \\ \mu' \bar{a} t \quad t \gg \tau(a, s) \end{cases} \quad (3a) \quad (3b) \quad (3c)
\]

Three regimes are readily identified: an initial growth in which both the in-community and the out-community degrees are growing linearly in time, followed by the slowing down of the in-community degree, which saturates to \( s-1 \), and then a further linear regime driven only by the out-community degree growth. Figure 2 shows that the numerical simulations perfectly match with the theoretical formulas (see the SI for details).

Noticeably, the long-time evolution of the node degree is linear in time and proportional to its activity class \( a \), so that we find the asymptotic degree distribution of the system to feature the same functional form of \( F(a) \propto a^{-\nu} \), as found in non-modular activity driven networks

\[
F(a) \propto a^{-\nu} \quad (4)
\]

In Fig. 3, we integrate the network for \( T = \) and we plot the three degree distributions. As expected, the out-community \( \rho(k_o) \) and the total \( \rho(k) \) degree distributions fall as power laws with exponent \( -\nu \). On the other
hand, the in-community degree $\rho(k_c)$ saturates to the community size distribution $P(s)$, as all the nodes reach their maximum in-community degree value ($s-1$), being that the modules’ size is far smaller than the network size $N_s = sN \max$. On the contrary, the out-community degree takes longer times to saturate to its maximum value $N_s/s$.

It is worth stressing that the results presented in this section apply to the networks obtained integrating links over time. A process unfolding on such networks, in general, will be affected by the time-aggregated features of the graph. The extent to which this is true, is function of the interplay between the time-scale describing its evolution, $\tau_p$, and the various $\tau(a, s)$. In the limit $\tau_p \ll \tau(a, s)$ the process would effectively evolve on the instantaneous, annealed networks that are characterised by a small average degree and modularity. In the opposite limit instead, the process would effectively unfold on static networks obtained integrating links over longer time characterised by high average degree and low modularity. Indeed, the average degree in this regime will be dominated by out-community links that make the connections between different communities increasingly stronger, thus increasingly destroying the identity of communities. In the limit $\tau_p \sim \tau(a, s)$ the process would effectively evolve on maximally modular networks (for a given set of parameters). Arguably, this is the most interesting regime that we will consider in the following.

**Epidemic processes on modular activity driven networks.** Let us turn our attention on the dynamical properties of SIR and SIS processes (see the Methods section for a detailed definition of the two) unfolding on the proposed model. Although similar, the two processes are intrinsically different33,57-59. Indeed, SIR processes are always characterised by the so called disease-free equilibrium, provided $d, N > 0$. The illness eventually disappear, i.e., $I = 0$ for $t \rightarrow \infty$. SIS models instead allow the existence of an endemic state where a finite and constant fraction of infected individuals permanently colonise the population, i.e., $I > 0$ for $t \rightarrow \infty$. Here, we focus on a central concept of contagion phenomena: the epidemic threshold. This quantity defines the conditions necessary for the spreading of the illness. In annealed networks, the threshold is determined by the moments of the degree distribution $\rho(k)$. In static graphs the expression is given by the principle eigenvalue of the adjacency matrix57,60,61. In time-varying networks instead, the threshold is determined by the interplay between the time-scales of the contagion and network evolution processes24,41,45,50,62-69. In the case of SIR models, we also consider another important...
quantity: the epidemic size $R_\infty$, which is defined as the final ratio of recovered nodes. This describes the fraction of nodes affected by the disease.

To develop a deeper understanding, let us derive the mean-field level dynamical equations describing the contagion process in modular activity driven networks. We define the activity block variables $S_{a,s}$, $I_{a,s}$, and $R_{a,s}$ as the number of susceptible, infected and recovered individuals, respectively, in the class of activity $a$ and community of size $s$ at time $t$ (to enhance readability, we omit to notate the dependence on time). This allows us to write the mean-field equation of the number of infected individuals for a SIR process, in each group of nodes with activity $a$ as:

$$dI_a = -\gamma I_a + \lambda s (\sum_{s'} a s' + (1 - \mu) \mu I_{a,s} \frac{S_s}{s} + (1 - \mu) I_{a,s} \frac{S_s}{N},$$

where $I$ and $I$ are the number of infected in communities of size $s$ and in the whole network, respectively. The first term in the r.h.s. accounts for the recovery of infected individuals. The other four terms account for the probability that a Susceptible node in a community of size $s$ connects to an Infected node inside (first) or outside (second) its community acquiring the infection, and for the probability that an Infected node of class $a'$ connects to a Susceptible node inside (third) or outside (forth) a community of size $s$, contracting the disease. For simplicity, we consider that $N - s \sim N$ and, at least initially, $I - I \sim I$. Summing over all the activities and community sizes, and considering only the first order terms in $a, I_{a,s}$, $R_{a,s}$, and their products, we obtain

$$dI = -\gamma I + \lambda (a) I + \lambda \Theta + \lambda \mu \sum_s (\langle a \rangle_s - \langle a \rangle) I_s,$$

$$d\Theta = -\gamma \Theta + \lambda (a^2) I + \lambda (a) \Theta + \lambda \mu \sum_s [\langle (a^2) \rangle_s - \langle a \rangle^2 \langle a \rangle \Theta] - \langle \Theta \rangle - \langle \Theta \rangle \langle \Theta \rangle \Theta,$$

where we defined $\Theta = \sum_s a I_s$ and $\Theta = \sum_s a I_s$. The term $\langle a^2 \rangle_s = \sum_s N_s a^2/s$ describes the moments of the activity distribution in any community of size $s$. The second, auxiliary, equation is obtained from the first by multiplying both sides by $a$ and summing over all $a$ and $a$. The epidemic threshold, in principle, can be derived evaluating the principle eigenvalue of the Jacobian matrix of the system of differential equations in $I$ and $\Theta$. In general, a closed expression for the threshold does not exist. However, we can point out some interesting observations.

First of all, the terms associated to $R_{a,s}$ vanish, implying that, at the first order, the thresholds of both SIR and SIS are equal. Furthermore, the terms in $\mu$ weight a comparison between the moments of the activity distribution in the network with the corresponding quantities evaluated inside each community. If fluctuations of these terms are negligible, due for example to very large community sizes or to narrow distribution of activity, the equations become equivalent to the case $\mu = 0$. In the limit $\mu \rightarrow 0$ the network has no modular structure, and the threshold, for both SIR and SIS, becomes $\beta/\gamma \geq 2/(1 + \sqrt{\gamma})$ as derived with different approaches in refs. The spreading condition is determined by the interplay between the time-scale of the contagion process and the time-scale of the network. Furthermore, the threshold is significantly larger with respect to the case in which the disease would spread in static or annealed networks generated integrating connections over time. Indeed, the concurrency of contacts in the two time scale separation regimes drastically facilitates the spreading of diseases. It is important to notice how the threshold is not function of $m$ even in the case of $m > 1$. This is due to the fact that we absorbed the contact rates in the definition of $\beta$. As detailed in the methods section, this is defined as the per capita rate of infection. By adopting such definition, we are able to estimate the spreading power of a disease independently of possible differences in contacts rates. The expression could be easily changed to explicitly account this aspect obtaining $\lambda/\gamma \geq 1/(N a (1 + \sqrt{\gamma})$. In the opposite limit $\mu \rightarrow 1$ networks are extremely modular, fluctuations become important and the symmetry between SIR and SIS breaks. In order to understand this limit, let us consider first a SIR process started from a single infected node in a community of size $s$. The large majority of connections are towards a small number of vertices in the same group. As soon as the disease start to spread within the community the number of infected and then recovered nodes grows, thus the probability of links $I - I$ and $I - R$ increases. Such connections cannot help the spreading of the disease. In fact they hamper the contagion process. In case instead of a network characterised by smaller values of modularity the growth of infected and recovered nodes has a much smaller effect. Indeed, the connectivity between communities would guarantee access to larger pool of susceptible nodes to sustain the spreading. From these simple observations we can expect that SIR processes are inhibited by highly modular connectivity patterns. Except for few exceptions in particular topologies, this is the case on static and annealed graphs. As we will show below, the same arguments hold also in time-varying connectivity patterns. On the other hand, in case of SIS processes, the repetition of contacts does not lead to such “pair annihilation”: contacts between infected nodes do not help the spreading of the disease, but they are only temporary (eventually, all infected nodes become susceptible again). Thus, we can expect that modularity plays a different role in SIS dynamics. This is what has been found also in the case of one annealed network model. Below we will show how this applies also in the case of time-varying networks. In order to numerically characterise SIR models, we study the epidemic size, $R_\infty$, as a function of $\beta/\gamma$. This quantity
acts as the order parameter of a second-order phase transition. For SIS processes instead, the order parameter is the final fraction of infected individuals, $I_\infty$. The numerical estimation of this quantity is challenging, since it requires the precise determination of endemic states. For these reasons, we follow ref. 74, measuring the lifetime of the disease, $L$, that acts as the susceptibility in phase transitions. This quantity is defined as the average time it takes for the disease to either die out or reach a macroscopic fraction, $Y$, of the populations. Without loss of generality, we start our simulations by setting 1% of randomly selected nodes as initial infected seed. Other parameters are set as: $\gamma = 0.01$, $m = 1$, $\nu = 2.1$, $\omega = 2.1$, $N = 10^5$ and $Y = 0.5$ (see SI for similar plots obtained fixing $\omega = 1.5$).

Results obtained from SIR models are represented in Fig. 4A,B, whilst results from SIS models are shown in Fig. 5A,B. In Figs 4B and 5B we study different community structure, either by considering a constant community size (dashed curves) or by drawing community sizes directly from the community size distribution $P(s)$. In solid curves, we draw community sizes as equal to the average value of $P(s)$ for all communities. The 95% confidence interval is in grey. Each point is an average of $10^2$ independent simulations.

Figure 4. Panel (A) $R_\infty$ as a function of $\beta/\gamma$, for selected values of $\mu$ and $s_{\text{min}} = 10$. Vertical black line represents the theoretical value of the epidemic threshold for $\mu = 0$ as derived in refs 24,71. Panel (B) $R_{\text{max}}$, i.e. the max value of $R_\infty$, as a function of $\mu$. In red curves we set $s_{\text{min}} = 100$, in blue curves $s_{\text{min}} = 10$. In solid curves, we draw community sizes directly from the community size distribution $P(s)$. In dashed curves, we fix the community sizes as equal to the average value of $P(s)$ for all communities. The 95% confidence interval is in grey. Each point is an average of $10^2$ independent simulations.

Figure 5. Panel (A) Lifetime of the disease $L$ as a function of $\beta/\gamma$, for selected values of $\mu$ and when $s_{\text{min}} = 10$. Vertical lines are the epidemic threshold. Panel (B) Ratio $\xi_{\text{SIS}} = \beta/\gamma$ in correspondence of $L_{\text{max}}$, as a function of $\mu$. In red curves we set $s_{\text{min}} = 100$, blue curves $s_{\text{min}} = 10$. Each point is an average of $10^2$ independent simulations. Note that we avoid to simulate $\mu = 1$ because the criterion we follow for the estimation of the threshold does not hold for a network with many connected components.
For SIR models, Fig. 4A tells us that, as expected, the higher $\beta/\gamma$ the higher the epidemic size. The figure also confirms the intuitions about the threshold. Indeed, we observe a dependence on $\mu$: the higher $\mu$ the higher the threshold. However, it is important to notice how such dependence is weak especially when compared with the SIS case (see below). Moreover, the higher the fraction of links created between pair of nodes sharing the same community (i.e. the higher $\mu$), the lower the epidemic size. This second observation is confirmed studying different community structures, as done in Fig. 4B, in which we plot the maximum epidemic size (corresponding to the largest value of $\beta/\mu$ in our settings), $R_{\text{max}}$, as a function of $\mu$. In the limit $\mu \to 0$, we observe that the disease impact is the same: the networks behave as if no community structure was present. Instead, when $\mu \to 1$, the modular structure influences the spread of the disease. As mentioned before, repeating contacts within communities significantly narrows the chances of having new infected individuals. Indeed, in SIR models, once a node recovers, it cannot be infected again. Repeating contacts with nodes already recovered does not favour the spread of the disease. Overall, the main observations are four. (i) Increasing the modularity reduces the epidemic size. (ii) A network with, on average, larger modules is likely to yield a higher epidemic size. (iii) The larger the modules the weaker the dependence on $\mu$ of the epidemic size. (iv) In case of small modules, the distribution of community sizes seems to influence the spreading of the disease. In particular, a network organised in small groups of constant sizes leads to smaller epidemic size respect to a network in which the average community size is the same, but individual sizes are extracted from a power-law distribution.

For SIS models, the lower $\mu$, the lower the life time $\tau$ (see Fig. 5). Inter-community links speed up the disease spreading and an endemic state, i.e. $Y = 0.5$, is reached faster. Moreover, the higher $\mu$, the lower the epidemic threshold. This last observation, which implies that increasing values of modularity favour the survival of the disease, is confirmed in Fig. 5B where we also test the effects of different community structures. In the limit $\mu \to 0$, there is no community structure and the curves converge to the same epidemic threshold. On the contrary, when $\mu \to 1$, the community structure becomes increasingly important and influences the spreading. Qualitatively, higher levels of modularity diminish the epidemic threshold. This is due to the repetition of the same contacts within a community which becomes increasingly more likely. Indeed, in SIS models, reinfection is allowed and nodes can become infected many times: communities act as a reservoir for the disease and favour the contagion process pushing the epidemic threshold to smaller values. Besides this last point, there are two main observations. (i) A network with larger modules is likely to have a higher epidemic threshold. (ii) In case of communities with smaller average sizes and high values of modularity, having the community size extracted from a power-law seems to slightly increase the threshold. Thus, the disease is able to spread more easily in modular networks with communities of similar or equal sizes. With the exception of one data point, this observed for $\mu > 0.5$ (see the dashed blue line in Fig. 5B).

### Real networks.

Although the modelling framework presented captures realistic activity and community size distributions of real networks, it neglects other important features such as burstiness\(^\text{70–80}\), and more complex temporal/structural correlations\(^\text{81–85}\). It is then crucial grounding the picture emerging from synthetic models with a real-world system. To this extent, we consider a temporal and modular network about scientific collaborations in the American Physical Society (APS). We study 96940 scholars connected by 692667 links (see the Supplementary Information for more details)\(^\text{86}\). We focus on ten years of data (January 1997–December 2006) coarse-grained at the temporal/structural correlations\(^\text{81–85}\). It is then crucial grounding the picture emerging from synthetic models with a real effect of modularity on SIS processes in this case. The temporal features of networks have been found to either slow down or facilitate the propagation of a disease\(^\text{6}\). Moreover, the presence of communities in static/annealed graphs might either significantly narrow the chances of having new infected individuals. Indeed, in SIR models, once a node recovers, it cannot be infected again. Repeating contacts with nodes already recovered does not favour the spread of the disease. Overall, the main observations are four. (i) Increasing the modularity reduces the epidemic size. (ii) A network with, on average, larger modules is likely to yield a higher epidemic size. (iii) The larger the modules the weaker the dependence on $\mu$ of the epidemic size. (iv) In case of small modules, the distribution of community sizes seems to influence the spreading of the disease. In particular, a network organised in small groups of constant sizes leads to smaller epidemic size respect to a network in which the average community size is the same, but individual sizes are extracted from a power-law distribution.

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### Discussion

Real networks are characterised by heterogeneous statistical distributions of crucial topological features; they are organised in modules/communities; they are subject to non-trivial temporal dynamics\(^\text{3–5,8,22,23}\). It has long been acknowledged that such attributes have critical effects on contagion processes evolving on systems’ fabric\(^\text{6}\). In particular, the heterogeneity in the connectivity patterns makes static/annealed networks extremely fragile to the spreading of infectious diseases\(^\text{6}\). Moreover, the presence of communities in static/annealed graphs might either slow down or facilitate the propagation of a disease\(^\text{12–18}\). In particular, community structures are most likely to inhibit SIR-like processes\(^\text{12–17}\). However, in particular networks, such as Autonomous System Graphs, peculiar topological properties might have the opposite effect\(^\text{17}\). The study of SIS processes in the context of modular networks has received much less attention. However, at least in the case of one artificial network model, modularity has been found to help the spreading\(^\text{18}\). A note of caution is however important. Indeed, in this paper, modular networks are compared with random graphs of different degree distribution. Thus, it is hard to disentangle the real effect of modularity on SIS processes in this case. The temporal features of networks have been found to either facilitate or hamper the spreading of contagion processes. We refer the reader to ref.\(^\text{90}\) for a recent compendium of epidemic spreading on time-varying networks. The study of the effects of modularity on epidemic spreading...
unfolding on time-varying networks has been very limited. At the best of our knowledge only one paper so far tackled directly this issue. In this work, an activity-driven network organised in two communities of equal size has been considered. In these settings, the threshold is not function of the modularity as each community is a good representation of the full network, thus the critical behaviour is the same independently of the number of connections between the two modules. Furthermore, the authors found a weak correlation between modularity and epidemic size. Starting from all these results, here we aimed to characterise the interplay between modularity and temporal dynamics of networks considering realistic community structures. To this end, we proposed a model of temporal networks with tunable modularity and heterogeneous activity distributions. We provided an analytical description of time-aggregated properties of such networks, and studied the impact of modular and temporal features on epidemic spreading processes. In synthetic networks, we found that modularity reduces the epidemic size and threshold in SIR models, slowing down the spreading process. The effect of modularity on the threshold is weak and appreciable only for large values of it. In SIS models, modularity reduces the epidemic threshold making the system more prone to disease spreading. The dependence on modularity is in this case much stronger. The repetition of the same contacts between nodes belonging to the same community acts as a reservoir for SIS-like diseases and allows the pathogen to reach an endemic state more easily. Our work is not exempt by caveats. The most important limitation, perhaps, is represented by the adoption of Poissonian activation dynamics, which has been shown to be unrealistic in empirical temporal networks which are characterised by bursty behaviours (heterogenous inter-event time distributions). Furthermore, not only links are dynamical entities being created and terminated, but also nodes may appear and disappear during the dynamics. Our model of modular activity-driven networks does not capture all these aspects of real temporal networks. Thus, we confirmed the picture emerging in synthetic graphs by numerical simulations of SIR and SIS models on real network, characterised by both modular and temporal features. We selected a co-authorship network in which nodes describe authors and links between them capture scientific collaborations. Clearly, such network is not of direct epidemiological relevance. Unfortunately, datasets more suitable for the study of the behaviour of spreading processes on temporal networks, such as face-to-face interactions, are scarce and quite small in size. Thus, they are not the optimal choice to study how temporal and topological properties affect epidemic thresholds which are analytically defined in the limit of $N \rightarrow \infty$. However, it is important to notice that social networks, in many different contexts, are characterised by the wide range of features missing in our simple model. Thus, in the spirit of a qualitative comparison, studying the behaviour of contagion processes on co-authorship networks is a meaningful exercise. In conclusion, our findings show that on time-varying networks modularity can have opposite effects on different classes of spreading processes. Dynamical processes unfolding on real networks have been show to exhibit a rich and complex phenomenology, depending on the topological and temporal properties of the underlying substrate as well as on the characteristics of the process under investigation. Such complexity may be addressed by both the definition of proper generative models, that allows to control the desired features, and the study of the processes behaviour in real-world network. Our work is set within this framework, and it contributes to shed light on the impact of modular and temporal properties of real networks on epidemic spreading dynamics.

**Methods**

**SIS and SIR models.** In both processes nodes are divided in different classes according to their disease status. In SIR models nodes are either Susceptible (S), Infected (I) or Recovered (R). Susceptible nodes describe healthy individuals. Infected nodes contract the disease and are infectious. Recovered nodes are no longer

![Figure 6. Panel (A) $R_0$ as a function of $\beta/\gamma$ for SIR processes diffusing on APS (cyan circles) and on the randomized APS dataset (green circles). Panel (B) $L$ as a function of $\beta/\gamma$ for a SIS models evolving on the same two networks. Each point is the average of $10^5$ independent simulations started from 1% of random seeds. We fix $\gamma = 0.05$.](image-url)
infected and acquire complete immunity to the illness. The model is fully characterized by two transitions: $S + I \xrightarrow{\beta} 2I$ and $I \xrightarrow{\gamma} R$. The first describes the infection propagation and $\beta$ is the capita infection rate. This quantity is defined by the average contacts per node ($k$) and by the per contact probability of transmission $\lambda$, i.e. $\beta = \lambda k$. The second transition describes the recovery process. Infected individuals recover spontaneously and permanently with rate $\gamma$. In SIS models instead we have just Susceptible and Infected nodes. While the contagion process is equivalent to the SIR case, the recovery is different and described by the following transition: $I \xrightarrow{\gamma} S$. Infected nodes spontaneously return in the susceptible compartment with rate $\gamma$.

References


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79. Author Contributions N.P. conceived the research, M.N. and K.S. conducted the numerical simulations, E.U. developed the analytical calculations, all authors analysed the results, wrote and reviewed the manuscript.
Additional Information

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