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Hierarchical team structure and multidimensional localization (or siloing) on networks

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Abstract

Knowledge silos emerge when structural properties of organizational interaction networks limit the diffusion of information. These structural barriers are known to take many forms at different scales-hubs in otherwise sparse organizations, large dense teams, or global core-periphery structure—but we lack an understanding of how these different structures interact and shape dynamics. Here we take a first theoretical step in bridging the gap between the mathematical literature on localization of spreading dynamics and the more applied literature on knowledge silos in organizational interaction networks. To do so, we introduce a new model that considers a layered structure of teams to unveil a new form of hierarchical localization (i.e. the localization of information at the top or center of an organization) and study its interplay with known phenomena of mesoscopic localization (i.e. the localization of information in large groups), k-core localization (i.e. around denser subgraphs) and hub localization (i.e. around high degree stars). We also include a complex contagion mechanism by considering a general infection kernel which can depend on hierarchical level (influence), degree (popularity), infectious neighbors (social reinforcement) or team size (importance). This very general model allows us to explore the multifaceted phenomenon of information siloing in complex organizational interaction networks and opens the door to new optimization problems to promote or hinder the emergence of different localization regimes.

1. Introduction

Organizational interaction networks, from those that follow the structures of private companies and social media to public institutions and academia, have long been known to influence the spread of information and dynamics among their agents [1–3]. Open source projects can falter or thrive through the interplay of maintainers and newcomers [4-6] much like how social media network structures hasten or hinder the development of echo chambers around specific groups [7, 8]. The collaborative structure in science can both create knowledge silos or help diffuse new ideas [9], just as the hierarchical structure of an organization can shape culture [10] and individual engagement [2, 11]. The strength of connections between individuals further shape these effects. Ideas and emotions spread over weak organizational ties [12–14]. Stronger ties and repeated exposures amplify these effects [15, 16], facilitating the spread of socialized behaviors and actions [10, 17]. At a theoretical level, all of these phenomena are related to the idea of dynamic localization: dictating whether specific parts of a system contribute alone to a dynamical process (localized), or whether the system as a whole undergoes collective activity (delocalized).

'Knowledge silos' (or *thought worlds* [18]) that constrain information or activity within specific parts of an organization are neither nor good nor bad *per se*. They can be detrimental to organizations if they limit adoption of norms or practices and hinder the spread of innovation [19]. They can also be desired when they protect confidential information or foster social reinforcement within teams [20]. In any case, knowing about the potential for different types of silos can be important. Through a better understanding of the different theoretical mechanisms under which silos emerge would help us control and design systems with information localization in mind. We therefore aim to provide a broad theoretical model for localization (the mathematical phenomenon) in order to enable future work around the design and control of silos (the empirical manifestation).

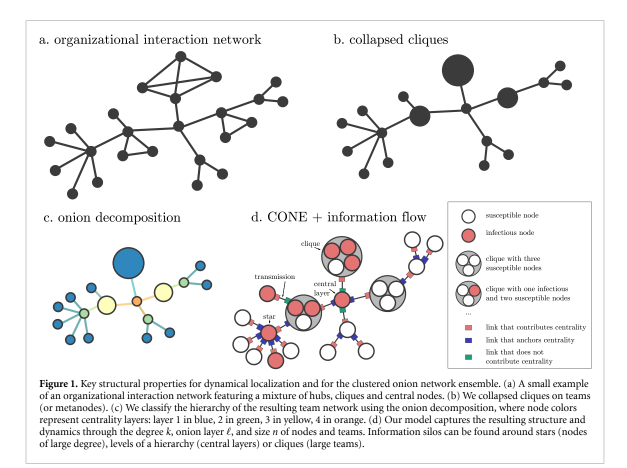
In an organizational interaction network of nodes (agents) and edges (information pathways), dynamical localization can be defined as a phase transition or an exponentially increasing level of activity across node categories with different levels of connectivity [21, 22]. For instance, if we consider only degree heterogeneity, simple models generally assume a node with *k* neighbors to be exposed *k* times more to some dynamical process than a node with a single neighbor and therefore to be *k* times more active (or less, if saturation occurs) [23]. However, for sufficiently heterogeneous networks, more detailed descriptions show that activity scales exponentially with the level of connectivity of a node [24]. The subsets of the networks where dynamics localize can then be identified in simulations with ad hoc tools or, as we will do, predicted analytically by demonstrating the exponential relationship.

Several approaches exist to identify specific types of localization, resulting in a rich and varied literature. By modeling epidemic spread and using a spectral approach [25], Goltsev *et al* showed in 2012 how a dynamical process can localize around the hubs of a network [26]. In particular, when studying a simple contagion process which diffuses at a normalized rate λ and dissipates at a unit rate, one can define a local activation threshold of a star subgraph (or star motif) with *k* neighbors as $\lambda_i \sim 1/\sqrt{k_i}$. Of course, in a larger complex network, the local activation threshold would be decreased by other active nodes in the system. Still, it remains that in heterogeneous networks, the node with the largest degree, k_{max} , can maintain activity on its own if the transmission rate lies around $1/\sqrt{k_{\text{max}}}$, which is often much smaller than the rate necessary for the collective activation of the whole network. This phenomenon can also be captured numerically by studying the participation ratios of individual nodes to either the dynamical steady state or the eigenvector centrality of the whole system [21].

The description of the localization phenomenon was extended beyond degree (a microscopic metric of connectivity) to k-core centrality (a macroscopic metric of connectivity) in 2018 by Pastor–Satorras and Castellano [27]. Based on the k-core decomposition—which prunes the network to identify the nested maximal subsets (or k-cores) of nodes who have at least k connections among each other—this version of localization lets the denser core of a network maintain a spreading process on its own with subcritical spillovers in the periphery of the network. More recently, St-Onge et al investigated the possibility of mesoscopic localization around specific groups rather than nodes or cores [22, 28]. Finding that a heterogeneous distribution of group size and weak (finite) coupling could allow large enough groups to maintain a diffusion process, again with subcritical spillover into smaller groups. These results all show that diverse sets of subgraphs with high enough connectivity can localize and maintain dynamics independently, despite the lack of collective activation across the entire network. Dynamical localization is thus analogous to how information traps are used in other areas of network analysis, such as community detection using compression of random walks patterns [29], and therefore has the potential to help detect important network properties beyond simple modules. We draw an example of the structural network properties relevant to organization in figure 1, illustrating how the interplay of degree, centrality, and group density, can be hard to quantify even in simple examples.

In this paper, we pursue a holistic view of dynamical localization in networks. Are there other mechanisms for localization which have been so far ignored? In particular, we are interested in demonstrating the mathematical potential for hierarchical localization, which is known in applied organizational research [2, 10, 11] but not yet clearly connected to dynamical localization. Colloquially, the CEO of a company should be more likely to be active in important discussions than a lower-level manager, even if both have the same number of direct subordinates. In other words, local connectivity alone does not fully characterize the potential for localization. And we are then also interested in how different structural features interact and how different types of localization might combine and amplify each other. How much more robust is localization around a central team than it is around a central star or a peripheral clique?

To this end, we develop an analytical description of organizational interaction network structure based on clique size distribution, the external degree distribution of cliques (inter-clique connectivity), and the



hierarchical structure of the clique network as described by a refined *k*-core decomposition called onion decomposition [30]. We then describe nonlinear dynamics on these networks using an approximate master equation (AME) framework [22, 28, 31] which allows us to analytically demonstrate hierarchical localization and numerically investigate its interplay with other types of dynamical localization.

2. Construction of the clustered onion network ensemble (CONE)

We look at networks of teams (higher order networks which can reduce to usual networks for teams of size one) and describe them analytically using the hierarchy and connectivity patterns between teams. The layered structure of the hierarchy is specified by the onion decomposition [30], a refined version of the *k*-core decomposition. The onion decomposition follows the same pruning algorithm but keeps track not only of the cores in which a node is found, but also of the layers in which they are removed in the pruning process. Starting with an entire network as the 0-core, nodes of degree 0 are removed to define the zeroth layer of the onion decomposition and leave us with the 1-core of the network. We then remove nodes of degree 1 in layer 1, and we remove nodes that are now of degree 1 (if any) in layer 2, and so on until we are left with the 2-core of the network. The process then repeats, starting with nodes of degree 2 and always increasing the layer count. In the ℓ th layer of this pruning process, one can think of the links remaining around a node to be removed in layer $\ell + 1$ as links contributing to its centrality, either by anchoring it to layer ℓ or by connecting it to more central layers. This structure can therefore be captured in our model using connection rules detailed in the onion network ensemble (ONE) [30] and described mathematically as a layered & correlated configuration model [32].

Each node in the ONE is then assumed to represent a team of arbitrary size, modeled as a clique, and thereby producing a CONE, see figure 1. Therein, the structural role of each team is specified by a size n, a degree k (connections with other teams) and an onion layer ℓ corresponding to core $c(\ell)$ in the k-core decomposition (team centrality). We denote N(x), N(x,y) and N(x,y,z) the distributions of teams where x, y, and z can represent their size n, degree k, or layer ℓ (and any combination thereof) over the three-dimensional space (n, k, ℓ) . Links between teams are assigned uniformly at random between the members of the respective teams, but the teams themselves must respect the underlying onion structure as shown in figure 1. Teams of

size 1 are individual nodes, teams of size 2 are simple edges but can be defined as a team to avoid imposing a hierarchical relationship between the two nodes involved, teams of size 3 are triangles, and so on.

We can further describe the connection rules suggested above, which will allow the model to respect a hierarchy specified as an onion decomposition. Teams of degree k in layer ℓ must have exactly $c(\ell)$ links to layers $\ell' \ge \ell$ if they are in the first layer of their core, and otherwise at least $c(\ell) + 1$ links to layers $\ell' \ge \ell - 1$ and at most $c(\ell)$ links to layers $\ell' \ge \ell$ [30]. As mentioned above, one can think of these links as those remaining once a node is reached by the pruning process of the k-core decomposition. In fact, we call degrees connecting layer ℓ to layer $\ell' < \ell - 1$ green or peripheral stubs (and these do not contribute to their position in the k-core decomposition), degrees connecting them to layer $\ell' = \ell - 1$ blue or anchor stubs (as they potentially anchor them to their layer and can contribute to their position of the k-core decomposition), and degrees connecting them to layer $\ell' \ge \ell$ red or central stubs (which always contribute to the position of the k-core decomposition). The color naming scheme for stub types is used for visualization purposes, but colors always map to a concrete structural role: green stubs connect to more peripheral neighbors, blue stubs anchor nodes to the previous layer, and red stubs connect towards the core and contribute to the onion centrality. These different links across layers are specified by a joint node-type connection matrix $L(k,\ell,k',\ell')$ counting the number of stubs starting from nodes of connectivity (ℓ,k) and leading to nodes in (ℓ', k') . From there, we simply assume that connections are made at random under the constraints of the connection matrix $L(k, \ell, k', \ell')$ and of the ONE: blue and green stubs only connect to red stubs while red stubs connect to any stubs of any color. The entire structure of this model is represented in figure 1, starting from the conceptualization of a network in teams and centrality to the layer structure of the CONE.

3. AMEs

We describe the spread of information (ideas, norms, etc) as a susceptible-infectious-susceptible (SIS) contagion model using an AME framework. This model is not meant to capture any specific spreading mechanisms, but simply describe how infectious nodes can transmit to their susceptible neighbors at a given rate λ and recover at a fixed unit rate to become susceptible again. In the context of organizational network, one can think of the SIS model as describing how up-to-date nodes (infectious) that possess all information relevant to the organization can update their neighbors who are behind or uninformed (transmission to susceptible neighbors) but also fall behind themselves (recovery). After recovery, nodes still have the potential to be exposed to new information and update themselves again. This 'reinfection' mechanism is key to SIS dynamics and leads to a dynamical equilibrium where the system can reach an active steady-state where infection and recovery events are balanced. This steady-state allows us to study the stability of the dynamics. Thus, regardless of the exact conceptualization, the SIS dynamics is a minimal tractable model that captures how information can emerge, spread, and remain active on a given network structure.

We follow the dynamics on a networked, hierarchical, organization of teams. We denote $C_{i|n,k,\ell}$ the fraction of these teams which have a given structural role (size *n*, connectivity *k*, and onion centrality ℓ) and have *i* nodes currently 'infectious' or active. We use SIS dynamics with a general infection kernel $\lambda(i, n, k, \ell)$ that can depend on the number of infectious nodes in a team (e.g. social reinforcement if $\lambda \propto i$), or on the implicit authority of a node (e.g. structural features *k* and ℓ). This general process can then be followed through the following set of AMEs

$$\begin{split} \dot{C}_{i|n,k,\ell} &= -(n-i) \left[i\lambda(i,n,k,\ell) + S^{\rm r}_{n,k,\ell} + S^{\rm g}_{n,k,\ell} + S^{\rm b}_{n,k,\ell} \right] C_{i|n,k,\ell} - iC_{i|n,k,\ell} \\ &+ (n-i+1) \left[(i-1)\lambda(i-1,n,k,\ell) + S^{\rm r}_{n,k,\ell} + S^{\rm g}_{n,k,\ell} + S^{\rm b}_{n,k,\ell} \right] C_{i-1|n,k,\ell} + (i+1)C_{i+1|n,k,\ell} \end{split}$$
(1)

where the $S_{n,k,\ell}^x$ are the mean-field quantities coupling different groups through stubs of a given color. On average, any stub of a node of degree *k* and layer ℓ will be red with probability

$$p_{k,\ell}^{\mathrm{r}} = \frac{\sum_{k',\ell' \ge \ell} L(k,\ell,k',\ell')}{c(\ell)N(k,\ell)}$$
(2a)

where the denominator counts the number of stubs that contribute to the centrality of nodes of degree k and layer ℓ (since they each have c contributing stubs) while the numerator counts the fraction of those that are classified as red (since they reach inward in the onion structure). Similarly, the remaining stubs will be green with probability

$$p_{k,\ell}^{g} = \frac{\sum_{k',\ell' < \ell-1} L(k,\ell,k',\ell')}{[k-c(\ell)]N(k,\ell)}$$
(2b)

where the denominator now counts non-contributing stubs and the numerator counts those that would be classified as green. With these frequencies of stubs of different colors, we can calculate the average degree of each color for a node with a given connectivity (k, ℓ) . This can be done using the following multinomial process with extra constraints enforced by the Kronecker deltas,

$$\langle k^{\mathbf{r}} \rangle_{k,\ell} = c(\ell) p^{\mathbf{r}}_{k,\ell} \tag{3a}$$

$$\langle k^{g} \rangle_{k,\ell} = \begin{cases} (k - c(\ell)) p_{k,\ell}^{g} - \delta_{c(\ell),c(\ell-1)} [p_{k,\ell}^{r}]^{c(\ell)} (k - c(\ell)) p_{k,\ell}^{g} & \text{if } k - c(\ell) \leqslant 1 \\ (k - c(\ell)) p_{k,\ell}^{g} & \text{otherwise} \end{cases}$$
(3b)

$$\langle k^{\mathbf{b}} \rangle_{k,\ell} = \begin{cases} c(\ell)(1 - p_{k,\ell}^{\mathbf{g}}) + (k - c(\ell))(1 - p_{k,\ell}^{\mathbf{g}}) \\ + \delta_{c(\ell),c(\ell-1)}[p_{k,\ell}^{\mathbf{r}}]^{c(\ell)}(k - c(\ell))p_{k,\ell}^{\mathbf{g}} & \text{if } k - c(\ell) \leq 1 \\ c(\ell)(1 - p_{k,\ell}^{\mathbf{r}}) + (k - c(\ell))(1 - p_{k,\ell}^{\mathbf{g}}) & \text{otherwise.} \end{cases}$$
(3c)

The details of this calculation are available in our previous treatment of the onion network structure [32]. Altogether, this colored stub matching scheme allows us to evaluate the mean-field couplings for a dynamical process, one color of stub at a time. First, for green stubs, we write

$$S_{n,k,\ell}^{g} = \frac{\langle k^{g} \rangle_{k,\ell}}{n} \sum_{\ell' < \ell - 1, n', k', i'} \lambda(1, n', k', \ell') \frac{L(k, \ell, k', \ell')}{\sum_{k'', \ell'' < \ell - 1} L(k, \ell, k'', \ell'')} \frac{i'}{n'} C_{i'|n', k', \ell'} , \qquad (4a)$$

which is constructed as follows: expected number of green stubs per node times the infection kernel of potential neighbors summed over all possible layers and cliques for these neighbors (biased by the layer–layer connection matrix *L* and their expected number of red stubs). The infection kernel is given by the rate of infection from a neighbor of a given type (k', ℓ') , times the probability that a green stub from (k, ℓ) reaches (k', ℓ') times the probability that the neighbor is actually infectious. For blue stubs, we write

$$S_{n,k,\ell}^{\rm b} = \frac{\langle k^{\rm b} \rangle_{k,\ell}}{n} \sum_{\ell'=\ell-1,n',k',i'} \lambda(1,n',k',\ell') \frac{L(k,\ell,k',\ell')}{\sum_{k'',\ell''=\ell-1} L(k,\ell,k'',\ell'')} \frac{i'}{n'} C_{i'|n',k',\ell'} , \qquad (4b)$$

and for red stubs,

$$S_{n,k,\ell}^{\mathrm{r}} = \frac{\langle k^{\mathrm{r}} \rangle_{k,\ell}}{n} \sum_{\ell' \ge \ell, n', k', i'} \lambda(1, n', k', \ell') \frac{L(k, \ell, k', \ell')}{\sum_{k'', \ell'' \ge \ell} L(k, \ell, k'', \ell'')} \frac{i'}{n'} C_{i'|n', k', \ell'} . \tag{4c}$$

Equations (1)–(4) provide a closed system of equations to follow generalized SIS dynamics on the CONE. This model describes network based on a unique combination of degree heterogeneity, clique structure and centrality pattern. However, the model does not capture correlations between neighboring nodes and teams; destroyed by the annealed structure of the master equations which averages over all nodes with a given structural description (n, k, ℓ) . This is where AMEs can deviate from stochastic simulations. While the master equation is exact for a disconnected group, the mean-field coupling for external connections assume an *annealed* network structure. One can think of this annealed structure as one where all nodes or teams of a given type are shuffled much faster than the dynamical process, such that their connectivity and hierarchical position is preserved but the identity of their neighbors lost in the shuffle. Conceptually, it is unclear whether reality is best described by a static network or an annealed structure, but it is important to know that the dynamical correlations present in simulations on static network can help reinforce localization phenomena [28].

We also note that the CONE framework is somewhat agnostic to the exact details of the dynamical process, such that similar systems of equations could be written to follow other types of binary social dynamics. Now that we have modeled the network structure in detail, we could write a more general dynamical process where the rate at which one node transitions from one state to another is a general function of its type and of the state of its team. This has been done for less detailed AME framework before [33, 34].

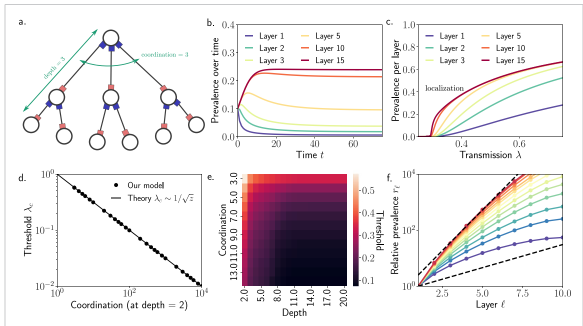


Figure 2. Localization on a Cayley tree. (a) Parameterization of finite Cayley trees, and their stub structure in the CONE. (b) Prevalence of contagion over time for individual layers on a Cayley tree of 20 layers with coordination number 4 and a transmission rate $\lambda = 0.33$. This shows how the dynamics can be supercritical in some layers with minimal spillover in others. (c) We fully explore the steady-state prevalence per layer (i.e. as time goes to infinity) in the same Cayley tree of 20 layers with coordination number 4. We see how the global threshold $\lambda_c \simeq 0.28$ is driven by the central layers only. (d) Global activation threshold λ_c of a star motif, i.e. a tree of depth $\ell_{max} = 2$, with varying coordination number (or degree). (e) Global activation threshold λ_c on the CONE from Cayley trees with varying depth and coordination number. This threshold is related to the largest eigenvalue of the tree structure and analyzed in appendix. (f) We demonstrate hierarchical localization through the relative prevalence r_ℓ of layer ℓ versus layer 1, evaluated at the global activation threshold λ_c for Cayley trees of depth 10 and coordination number between 3 (bottom blue curve) and 15 (top red curve). The two dashed lines show the exponential factor from equation (17), parameterized to match the bottom and top curves and therefore matching their main behavior observed in the middle layers where additional factors are less important.

4. Hierarchical localization

We confirm that the CONE can capture dynamical localization around central nodes, at both the microscopic (degree) and macroscopic (onion centrality) levels in figure 2. As a first experiment, we simulate the SIS dynamics using equations (1)–(4) on regular Cayley trees parameterized by their depth (denoted ℓ_{max})) and coordination number (the degree of non-leaf nodes, denoted *z*) as shown in figure 2(a). Note that this section completely ignores both the team structure of the CONE, i.e. we fix n = 1, as well as the complex contagion mechanisms, i.e. we fix $\lambda(i, n, k, \ell) = \lambda$.

Figure 2(b) shows the temporal evolution of the activity level (i.e. the *prevalence* or expected fraction of infectious nodes at a given layer) starting from a uniform initial condition in a tree with depth 20 and coordination number 4. While all layers $\ell > 1$ have the same connectivity, we see very different time series across different layers driven by differences in hierarchical position: With activity relaxing to very low levels in the peripheral layers while more central layers are orders of magnitude more active. Figure 2(c) then shows the expected steady state prevalence (or infinite time limit) in the different layers of the tree as we vary the transmission rate. These results provide a first glimpse of hierarchical localization since the activation of the system around the global $\lambda_c = 0.3$ is driven only by some of the core layers of the tree. Conversely, layer 3 for example appears to activate only around 0.35 if we use the maximum derivative of its prevalence curve with respect to λ as a proxy for its peak susceptibility [35].

Figure 2(d) shows the value of the global threshold at a fixed depth of two and therefore looking at star motifs with different coordination number or degree. The CONE captures the previous key theoretical result from spectral analysis [26]: the threshold for microscopic localization or activation around a star motif goes as the inverse of the square root of the star degree. More generally, figure 2(e) shows the global threshold λ_c as a function of the depth and coordination number of the trees; illustrating the relative symmetry of the system and how adding hierarchical layers and local degree is somewhat equivalent.

We can investigate these previous results analytically by considering a Cayley tree of depth *L* with coordination number z = d + 1 where *d* is the number of descendants or children of non-leaf nodes. Given

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the one-to-one mapping between degree and layer, we can directly write the CONE equation for the fraction of infected nodes I_{ℓ} in the layer ℓ as

$$I_{\ell} = -I_{l} + (1 - I_{\ell}) \left[\lambda (d + \delta_{\ell, \ell_{\max}}) I_{\ell-1} + \lambda I_{\ell+1} \right],$$
(5)

where it is implicit that $I_0 = I_{\ell_{max}+1} = 0$. In the stationary state and near the epidemic threshold (where $I_{\ell}^* \to 0$ for all ℓ , see appendix), we can write the following recursive equation

$$I_{\ell}^{*} = \frac{1}{\lambda} I_{\ell-1}^{*} - dI_{\ell-2}^{*} , \qquad (6)$$

which is valid for $\ell \in \{3, L\}$, where we define $L \equiv \ell_{\max} - 1$. We also have the boundary conditions $I_2^* = I_1^* / \lambda$ and $I_{\ell_{\max}}^* = \lambda(d+1)I_L^*$. Since $I_\ell \to 0$, we are in fact interested in the relative quantity $r_\ell \equiv I_\ell^* / I_1^*$, which also respect the recursion

$$r_{\ell} = \frac{1}{\lambda} r_{\ell-1} - dr_{\ell-2} , \qquad (7)$$

with boundary conditions $r_1 = 1$, $r_2 = 1/\lambda$ and $r_{\ell_{max}} = \lambda(d+1)r_L$. We define the following generating function

$$G(x) = \sum_{\ell=1}^{\infty} r_{\ell} x^{\ell}$$
(8)

but are only interested in its first *L* terms. By multiplying equation (7) by x^{ℓ} and summing from $\ell = 3$ to $\ell \to \infty$,⁶ we obtain

$$G(x) - r_1 x - r_2 x^2 = \frac{x}{\lambda} \left[G(x) - r_1 x \right] - dx^2 G(x) .$$
(9)

Rearranging the terms and using the boundary conditions to simplify the equation, we obtain

$$G(x) = \frac{x}{1 - \frac{x}{\lambda} + dx^2} = \frac{x}{d(x - x_+)(x - x_-)},$$
(10)

where

$$x_{\pm} = \frac{1 \pm \Delta}{2d\lambda}, \quad \Delta = \sqrt{1 - 4\lambda^2 d}.$$
 (11)

If $\Delta \neq 0$, we can use the following partial fraction decomposition

$$G(x) = \frac{\lambda x}{\Delta} \left[\frac{1}{x - x_+} - \frac{1}{x - x_-} \right] , \qquad (12)$$

and develop both terms using their Maclaurin series, which results in

$$G(x) = \frac{\lambda}{\Delta} \left[\sum_{\ell=1}^{\infty} \left(\frac{x^{\ell}}{x_{-}^{\ell}} \right) - \sum_{\ell=1}^{\infty} \left(\frac{x^{\ell}}{x_{+}^{\ell}} \right) \right].$$
(13)

Therefore, for all $\ell \in \{1, L\}$,

$$r_{\ell} = \frac{\lambda}{\Delta} \left[\left(\frac{1}{x_{-}} \right)^{\ell} - \left(\frac{1}{x_{+}} \right)^{\ell} \right] \,, \tag{14}$$

which is rewritten as

$$r_{\ell} = \frac{\lambda}{\Delta} \left(\frac{1}{2\lambda}\right)^{\ell} \left[(1+\Delta)^{\ell} - (1-\Delta)^{\ell} \right] \,. \tag{15}$$

⁶ We are extending the validity of the recursion beyond $\ell = L$ without consequences as we will then only extract the first *L* terms of the generating function.

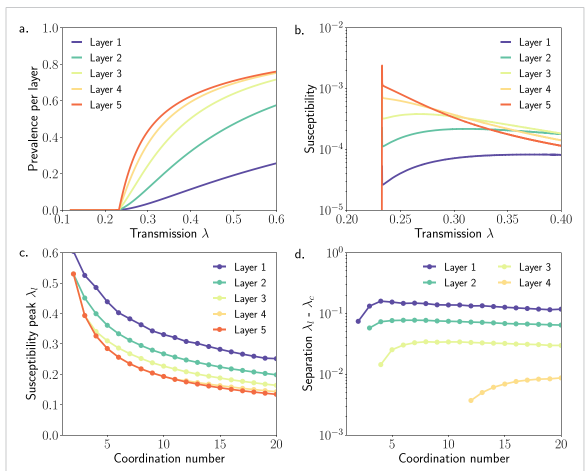


Figure 3. Analysis of hierarchical localization. (a) Prevalence per layer in a Cayley tree of 5 layers with coordination number 7. (b) Susceptibility, approximated by a numerical derivative over the prevalence, at each layer of the tree as a function of transmission rate λ . Peaks in susceptibility indicate important phase transitions: first all curves peak at the global threshold of the dynamical process (with numerical noise filtered out before the threshold), then, some curves show a second broad peak can indicate a local activation threshold for a given layer. In this example, the dynamics are initially localized around layer 5 and 4, then grow to layer 3 as it activates around $\lambda = 0.25$, then to layer 2 around $\lambda = 0.3$ and finally to layer 1 close to $\lambda = 0.4$ (c) Activation threshold per layer, identified as the final local maximum of each susceptibility curve, as we vary the coordination number of the Cayley tree with 5 layers. Hierarchical localization around specific layers requires a sufficiently large coordination number. For example, the dynamics can localize exclusively in layers 4 and 5 only if the coordination number is at least 4, and localized solely on the root only if the coordination number is at least 12. This can be seen by the separation of susceptibility peaks. Any contagion with transmission rate between the two most extreme curves can therefore exhibit some level of hierarchical localization; whereas contagions fail to survive under the lower bound and collective delocalized dynamics occur above the upper bound. (d) Plot of the differences between the local susceptibility peak λ_{ℓ} and the global threshold λ_c , showing the phase transitions behind the emergence of specific localization patterns.

Note that in the limit $\Delta \rightarrow 0 \ (4\lambda^2 d \rightarrow 1)$, the preceding equation is well-defined

$$\lim_{\Delta \to 0} r_{\ell} = 2\lambda \ell \left(\frac{1}{2\lambda}\right)^{\ell} \iff \lim_{\Delta \to 0} r_{\ell} = 2\lambda \ell d^{\ell/2} , \qquad (16)$$

even though the partial fraction decomposition should have been computed differently. If Δ is imaginary $(4\lambda^2 d > 1)$, which should be the case in general given that the threshold of a Cayley tree is well approximated by the threshold of the root $\lambda_c \sim 1/\sqrt{z}$ (see the appendix for a more accurate approximation), we can rewrite the solution as

$$r_{\ell} = \frac{2\lambda}{||\Delta||} d^{\ell/2} \sin(\theta \ell) , \quad \theta = \arctan(||\Delta||) .$$
(17)

Therefore, there is a regime close to the epidemic threshold where we find an exponential relationship between a node's depth in the network (layer ℓ) and its level of activity: Hierarchical localization. The phenomenon is confirmed in the last panel of figure 2.

We further explore hierarchical localization numerically in figure 3. Borrowing tools from statistical physics and the study of phase transitions [36, 37], we can use the derivative of local prevalence per layer as a

proxy for the susceptibility (here used in the statistical mechanics sense: the response of a system to changes in parameters). This quantity gives us a tool to identify local activation and further study the localization phenomenon [35]. Figure 3(a) presents the prevalence diagram on a tree with only five layers. In panel (b), we then show the discrete derivative which should have a unified, infinite, peak at phase transitions. Here, these curves show a global maximum at the global activation threshold of the system (a classic phase transition), peaking at a finite value because we approximate susceptibility through a numerical derivative. Still, we also find local maxima at the local activation thresholds of unique layers that do not participate in the global activation. Before a local peak of layer 2, for example, we can assume that the dynamics are localized in layers at least greater than 2 in order to explain the double peak: The first peak is caused by the localized activation with subcritical spillover of activity to lower layers like layer 2, whereas the second peak identifies the point where nodes in the second layer reach their local critical point and can maintain activity. If layer 2 participated in a global activation, as in classic phase transitions, then there could not be a second local maxima of susceptibility as that layer would have already reached its peak response from inactive to active. This result is similar to double percolation transitions that have been previously observed in highly clustered networks [36] and core-periphery structure [38].

In figure 3(c) we automate the detection of susceptibility peaks and tune the coordination number of the five-layer tree. Because of the interplay between microscopic and hierarchical localization, we find that certain localization patterns require a minimal number of layers and degree. For example, localizing the dynamics solely on the root of a five-layer tree requires a coordination number of at least 12. These specific localization patterns can emerge discontinuously, as shown in figure 3(d). As we tune the structure of a network (through discrete changes because of the discrete nature of networks), we can create explosive localization patterns: new susceptibility peaks can appear far from the global activation threshold. All of our results show the need to consider dynamical localization as a multidimensional phenomenon with multiple causes at different scales, like local connectivity and global layer structure, all of which can interact in interesting ways.

5. Exploration of multidimensional localization

Localization around stars or hubs has been a very active topic of research for the last decade, and here we have illustrated how it is a subset of hierarchical localization around trees of depth 2. Other structural mechanisms for localization (e.g. cliques) can be thought of as independent dimensions of the localization phenomenon. Using simple *in silico* experiments and an empirical case study, we now investigate how hierarchical localization can be amplified or hampered by localization around other network features.

5.1. Synthetic experiments

First, we can use the clique formulation of the CONE to include elements of mesoscopic localization. This allows us to look at the interplay of macroscopic hierarchy and mesoscopic cliques by using the CONE to study dynamics on a tree of cliques (or a hierarchy of teams). Organizations tend to have a pyramid structures as a result of power centralization [39], which suggest a strong potential for hierarchical localization. However, in these pyramids, team size can be either (1) positively correlated with onion layers such that teams higher in the hierarchy are also larger, (2) negatively correlated with layers such that peripheral teams tend to be larger, or (3) uncorrelated such that all teams have an average size regardless of their position in the hierarchy. Note that correlations between team size and centrality changes the density of the resulting network, but with surprising impacts on the dynamics. We show these results in figure 4. Our results illustrate how cliques can amplify hierarchical localization through positive correlations, as we find a larger separation of susceptibility in scenario 1 compared to 3. Conversely, negative correlations can complete delocalize the dynamics, as seen in scenario 2. Interestingly, despite having more links in the network, scenario 3 delocalizes the dynamics at the price of having a higher global activation threshold than scenario 1.

We then deviate from trees and leverage the ability of the CONE to produce networks with a fixed *k*-core structure. In figure 5, we introduce a *k*-core at the top of the hierarchy. This of course amplifies the localization phenomenon by allowing the *k*-core of root nodes to self-activate. Interestingly, while this self-activation lowers the global threshold and spills over into the penultimate onion layers, it barely affects the lower layers of the hierarchy. The local activation threshold of nodes in layers 1 or remains essentially unchanged as we grow the inner *k*-core of the network. In figure 6 we do the opposite and introduce a *k*-core among leaves in the periphery of the initial structure. This process is akin to team building exercise or

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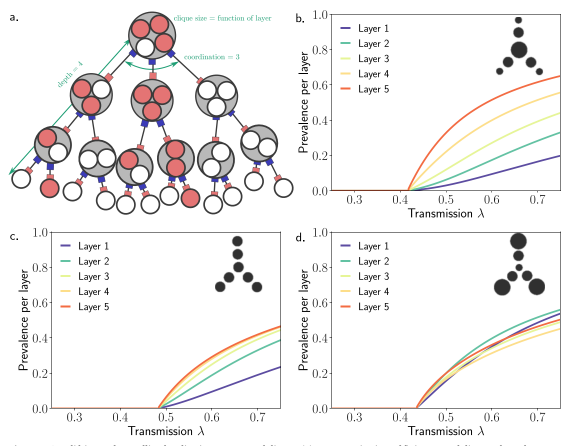


Figure 4. Amplifying and cancelling localization on a tree of cliques. (a) Parameterization of finite trees of cliques where the tree structure is enforced by the stub structure of the CONE and the sizes of cliques are specified as a function of their layer. (b) We reinforce localization by setting clique size *n* equal to layer ℓ : teams at the top of the hierarchy are larger. We use depth $\ell_{\text{max}} = 5$ and coordination number z = 3. (c) We use the same tree and set all cliques size equal to 3. (d) Again using the same tree, we delocalize the dynamics by setting clique size *n* inversely proportional to layer: $n = \ell_{\text{max}} + 1 - \ell$ such that teams at the bottom of the hierarchy are larger.

socialization meant to disseminate information from one branch of an organization to another. We find that this process allows an organization to progressively delocalize the dynamics. Yet, unlike the results of figure 4, socialization across branches also simultaneously decreases the global activation threshold of the whole network. In other words, this type of structure allows easier and more homogeneous spread of information.

As a last synthetic experiment, we briefly explore the generality of the CONE beyond network structure by using the $\lambda(i, n, k, \ell)$ function. This complex transmission rate can allow us model classic complex contagion where groups with more activity (i.e. higher *i*) diffusion information more effectively by setting $\lambda(i, n, k, \ell) \propto i$. This scenario has recently been well studied in the context of dynamics on higher-order networks [40]. Such dynamics can lead to interesting transitions where activity emerges first in larger teams before diffusing to smaller ones. Using AMEs formulation in terms of group structure, we recently showed how this modifies classic notions of network influence [41]: in non-linear contagions, influence becomes a group or team property, rather than an individual property of network hubs.

One unique feature of the CONE is that we can now also model situations where teams or individuals with privileged network structure (larger size *n*, connectivity *k* or centrality ℓ) have access to more or less information, by setting for example $\lambda(i, n, k, \ell) \propto \ell^{\gamma}$ as we do in figure 7. (Note that our analysis presented in appendix can be generalized to this type of complex contagion.) When γ is positive, central nodes receive more power and amplify the localization: we can saturate the dynamics around central layers while the periphery remains virtually inactive. More interestingly, we can also localize the dynamics around central layers by opposing power and network positions when γ is negative. In this case, the periphery might perceive a higher transmission rate but remains at a structural disadvantage while the root sees a lower transmission rate despite its advantageous network position. In between, certain layers can strike the right balance and outperform both the center and periphery of the hierarchy. While weakly negative γ values

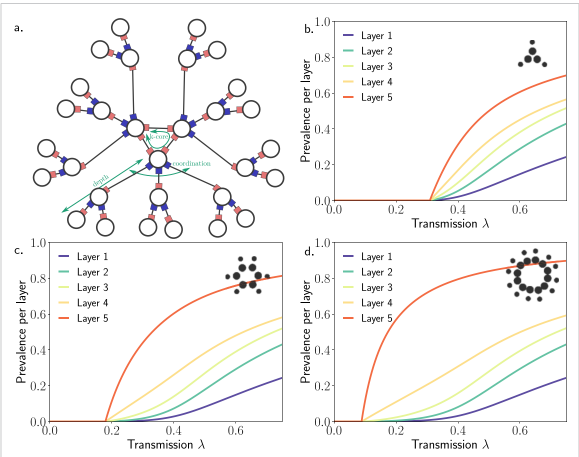


Figure 5. Amplifying localization on a core with treelike periphery structure. (a) Parameterization of a uniform *k*-core where all nodes are roots of finite trees, all enforced by the stub structure of the CONE. In panels (b)–(d), we use a central group of 20 nodes that form a 2-, 5-, or 11-core respectively. All nodes in that core are the roots of their own finite trees of depth 5 and coordination number 3, resulting in a 5 layers of Onion Decomposition with a bimodal *k*-core structure (core vs periphery). As the *k*-core grows denser, network structure amplifies the localization phenomenon through a synergy between core localization around the roots and hierarchical localization in the periphery. Importantly, while the number of peripheral trees depends on the density of the core, the prevalence observed on each tree is almost independent of the prevalence in the core as inner layers provide a buffer zone (i.e. the blue curves barely change).

hinder localization (figure 7(a)), for sufficiently negative γ (figure 7(a)) we find a phase transition where activity emerges in a strongly localized manner within these balanced central layers, before weakly diffusing asymmetrically towards the periphery and root. The most influential nodes are thus those that have the right balance of network centrality and dynamical activity. This nontrivial interplay of structure and dynamics define the *influential layers* of the network, i.e. those able to sustain the dynamics on their own.

5.2. Empirical case study and simulations

Finally, we illustrate how the entire conceptual pipeline of the CONE can be integrated in a single software mapping any organizational network to a bifurcation diagram like the ones visualized in the last experiments. The software takes a simple network edgelist as input, as in figure 1(a). It then finds cliques in the original network and merges them into teams, prioritizing larger cliques and merging cliques of the same size in an arbitrary order. This steps gives us a network of teams as in figure 1(b). The software then runs the onion decomposition on this network of teams, as in figure 1(c), and outputs a CONE parametrization as illustrated in figure 1(d). The software for this pipeline is open and available [42].

To showcase this pipeline, we use friendship data collected as part of previous research on network effects in a manufacturing firm [43]. This previous study used an established survey tool for constructing social networks of workplace friendship ties [44–46]. The survey asked participants to specify individuals they consider to be friends from a complete list of company employees. Of the firm's 142 manufacturing floor employees, 75 responded (52.8%) and reported 4.55 friendships on average with a 4.60 sample standard deviation. This yielded a network of directed edges from individuals who claim friendships and toward

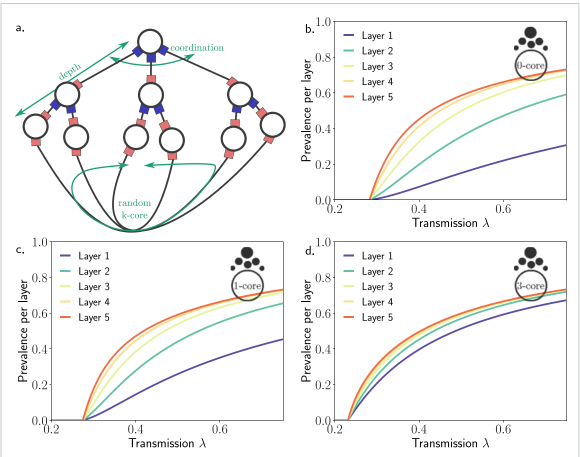


Figure 6. Delocalizing dynamics on a tree through *k*-core structure in the periphery. (a) We start with a finite tree and connect all leaves as a uniform *k*-core. The resulting networks has a number of onion layers equal to the depth of the original tree and the root preserves its centrality but with all nodes in the (k + 1)-core, as long as k + 1 is smaller than the coordination number of the original tree. In panels (b)–(d), we fix the depth and coordination number of the original tree to 5, and modify the peripheral *k*-core from 0 (control) to a 1-core and a 3-core. As the peripheral *k*-core grows denser, we lower the threshold of the whole system and also delocalize the dynamics away from the inner layers.

individuals who are considered friends by those individuals, which we then map to a simple, undirected, friendship network. The resulting network features significant degree heterogeneity given its size with a few smaller cliques dispersed throughout its core-periphery structure.

The raw network data is illustrated in figure 8(a) and used as input to our computational pipeline. The output of the computation is either a time series of prevalence per node type for a given transmission rate or a steady state prevalence per node time across transmission rates. We show the latter in figure 8(b). We compare the results of our analytical framework with the distribution of prevalence observed in exact, stochastic, simulations on the fixed network structure [47]. We only simulate after the epidemic threshold of the analytical system to limit finite size effects. Within these simulations, we separate nodes based on the different node types characterized by the CONE and plot the mean prevalence and the entire range of prevalence within each type. In doing so, we find that our analytical results match and explain the spread of activity observed throughout the real network.

Interestingly, we find that hubs and cliques deep in the core-periphery structure both drive the activation of the system around its global threshold. Core cliques reach a higher prevalence than core hubs, despite having lower connectivity with the rest of the network. This computational pipeline can therefore be used to investigate how well different parts of the network can spread information relative to each other. Insights from our previous experiments could then be used to suggest network interventions and fine tune the output of the network.

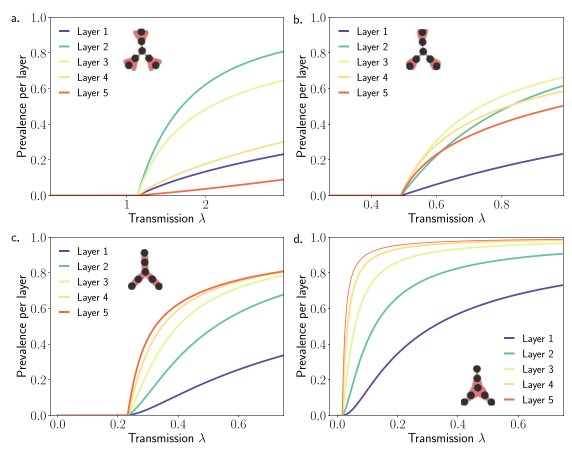


Figure 7. Modifying hierarchical localization through a complex transmission function. We use a finite tree of 5 layers with coordination number 7, as in figure 3 and implement a transmission function that varies with layer: $\lambda(l) \equiv \lambda \times \ell^{\gamma}$. This function exogeneously models the fact that certain position in the hierarchical organization might have more power and or sensitivity to the contagion process. (a) and (b) We amplify the transmission rate in the periphery with γ equal to -3.0 and -1.0, respectively. (c) Classic contagion case with $\gamma = 0$. (d) We amplify the power of core layers with $\gamma = 2.0$, lowering the global threshold and increase the extent of hierarchical localization.

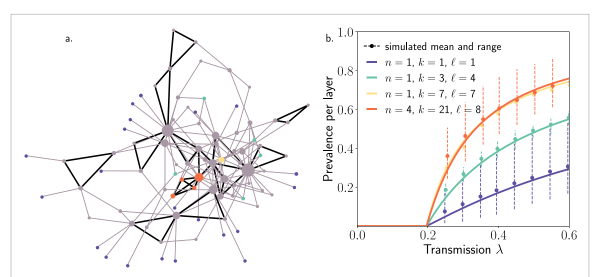


Figure 8. Application of the CONE framework to network data from a manufacturing firm [43]. (a) Our computational pipeline uses raw network data as input. It first collapses cliques, a few of which are highlighted with thick black edges. It then runs the onion decomposition to identify team centrality. And finally, the pipeline uses the resulting structure to parametrize and run the CONE equations. All nodes are assigned to a particular type, and just a few are highlighted in color on the figure. (b) From the CONE, we obtain a bifurcation diagram exploring the potential of different types of nodes to remain active through local or global activation. We compare the predictions of the CONE for the highlighted node types using simulations on the real network data. At different value of transmission rate, we calculate the prevalence of each node (i.e. the fraction of time it spends in an active infectious state) and plot the mean and full range of prevalence for nodes in a given node type under the CONE framework. The orange node type contains a single node. Our equations match and explain the observed spread of simulations.

6. Conclusion

In this paper, we have made both concrete and forward-looking steps towards a better theoretical understanding of how dynamical processes unfold in organizational interaction networks. First, we provided a mathematical demonstration of hierarchical localization; showing how key layers of centrality can potentially support a dynamical process on their own. Second, we explored the multidimensional interplay of different localization mechanisms by varying degree distributions, centrality patterns, team sizes, core structure, and transmission functions. Through a series of experiments, we have highlighted how carefully designing organizational networks can either amplify or hinder dynamical localization.

This exploratory analysis paves the way for future work on multiple fronts. Future theoretical studies could attempt to optimize network structure to either enhance or limit siloing of information as desired by a given organization. More applied work could leverage the CONE to quantify the localization potential of real organizational structures through formal organizational charts, network surveys, or email archives. This could be done, for example, by measuring the standard deviation around the peak susceptibility of different teams based on their size, connectivity and centrality in the organization; i.e. the standard deviation over the values shown for a given network in figure 3(c). Using peaks of susceptibility means that localization and siloing potential are not measured against a specific dynamical process with fixed parameters, but simply against a spreading mechanism regardless of parameter values. This might provide a parameter-agnostic tool for the comparison of organizational networks. Similarly, the localization properties identified by the CONE could be used to compress network structure similarly to community detection [29], and help automatically identify key network properties that combine hubs, groups, and hierarchy.

This line of enquiry could eventually contribute to important questions in organizational research: How do structures affect organizations' abilities to share different kinds of information? And, as others also ask [19, 48, 49], how do network properties moderate the role of factors like process, context, information content, and identity? Moreover, while researchers increasingly understand *how* and *why* knowledge embeds in organizational units, organizational scholarship still faces critical challenges. How can we limit the spread of stress-related disorders [50] without hindering the dissemination of knowledge? How can we prevent the diffusion of maladaptive behaviors, like corruption [16], while promoting inclusive practices [10] through socialization? The answer could lie in the complex interplay of information dynamics and network structure; by amplifying the transmission of certain types of information in key regions of an interaction network, one can intentionally localize or delocalize their spread. Our work introduced a general network model that could allow us to explore solutions to these important questions. The CONE provides a mathematical sandbox to develop a greater understanding of localization within subsets of a network, and of diffusion throughout organizational structures.

Data availability statement

The codes that produced the findings of this study are openly available [42].

The data that support the findings of this study are openly available at the following URL/DOI: https://zenodo.org/record/7930362.

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Appendix . Epidemic threshold for the modified Cayley tree

Let us consider a modified Cayley tree where the root node is of degree d and each descendent of this node are of degree d + 1, except for the leaves which are of degree 1. This is also called a perfect d-ary tree. The CONE equations are then

$$\dot{I}_{\ell} = -I_{\ell} + (1 - I_{\ell}) \left[\lambda dI_{\ell-1} + \lambda I_{\ell+1} \right] \,. \tag{18}$$

(19)

Near the epidemic threshold, we can linearize the preceding equation and write it under matrix form

 $\frac{\mathrm{d}\boldsymbol{I}}{\mathrm{d}\boldsymbol{t}} = \boldsymbol{M}\boldsymbol{I}\,,$

where

$$\boldsymbol{M} = \begin{bmatrix} -1 & \lambda & 0 & \dots & 0\\ \lambda d & -1 & \lambda & & \\ 0 & \lambda d & \ddots & \ddots & \\ \vdots & & \ddots & \ddots & \lambda\\ 0 & & & \lambda d & -1 \end{bmatrix}$$
(20)

is a tridiagonal Toeplitz matrix. The eigenvalues $(\phi_j)_{i=1}^{\ell_{\text{max}}}$ of **M** are given by [51]

$$\phi_j = -1 + 2\lambda \sqrt{d} \cos\left(\frac{j\pi}{\ell_{\max} + 1}\right) \,. \tag{21}$$

The critical point is reached when the largest eigenvalue

$$\Phi = -1 + 2\lambda\sqrt{d}\cos\left(\frac{\pi}{\ell_{\max} + 1}\right)$$
(22)

equals zero. Therefore, the epidemic threshold is

$$\lambda_{\rm c} = \frac{1}{2\sqrt{d}\cos\left(\frac{\pi}{\ell_{\rm max}+1}\right)} \,. \tag{23}$$

This expression is equivalent to known results for the largest eigenvalues of Bethe trees [52]. For ℓ_{max} sufficiently large, we also recover $\lambda_c \simeq 1/(2\sqrt{d})$ which is a known result from the study of dynamics on networks [26]. Since the degree of the root node for the standard Cayley tree is larger, we expect the epidemic threshold of the Cayley tree to be upper-bounded by the expression above, i.e. $\lambda_c \leq 1/(2\sqrt{d})$.

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