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Moment-Closure Approximations for Discrete Adaptive Networks

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Abstract

Moment-closure approximations are an important tool in the analysis of the dynamics on both static and adaptive networks. Here, we provide a broad survey over different approximation schemes by applying each of them to the adaptive voter model. While already the simplest schemes provide reasonable qualitative results, even very complex and sophisticated approximations fail to capture the dynamics quantitatively. We then perform a detailed analysis that identifies the emergence of specific correlations as the reason for the failure of established approaches, before presenting a simple approximation scheme that works best in the parameter range where all other approaches fail. By combining a focused review of published results with new analysis and illustrations, we seek to build up an intuition regarding the situations when existing approaches work, when they fail, and how new approaches can be tailored to specific problems.

Keywords: adaptive network, moment-closure approximation, adaptive voter model, fragmentation transition, state correlations

1. INTRODUCTION

Complex networks have been ubiquitously used to model problems from various disciplines \cite{1-6}. Treating a complex system as a network, a set of discrete nodes and links, leads to a conceptual simplification that often allows subsequent analytical insight that provides a deep understanding.

For many questions the networks of interest are not static entities but change in time due to the dynamics of and on the network. In the dynamics of networks, the network itself is regarded as a dynamical system. Prominent examples are network growth models leading to specific topologies such as scale-free \cite{7} and small-world networks \cite{8}. The dynamics on networks concerns dynamical processes such as epidemic spreading \cite{9} that occur on a given fixed network, where each node carries a state which evolves through interactions with its neighbors.

If the dynamics on and of networks occur simultaneously and interdependently then the network topology coevolves with the states of the nodes and an adaptive network is formed \cite{10,11}. Adaptive networks have been used to model problems of opinion formation \cite{12,18}, epidemic spreading \cite{13,27}, evolution of cooperation \cite{28,40}, synchronization \cite{41,45}, neuronal activity \cite{48,54}, collective motion \cite{53,56}, cartelisation of markets \cite{57}, and particle diffusion \cite{58} among others.

Network models in general and adaptive networks in particular provide a powerful framework to model, analyze, and eventually understand a wide range of self-organization phenomena.

For instance Tomita et al. \cite{59} showed that a very simple adaptive network model can be used to produce a huge variety of different self-organizing structures including a self-replicating Turing machine.

While specific models can be studied by agent-based simulation, the numerical performance scales badly with the complexity of update rules in the model, which makes exploration of a wider range of models difficult. In particular those models where the update of the state or neighborhood of nodes depends on the states of multiple other neighboring nodes pose strong numerical demands. Additionally the general bad data locality of network simulations precludes efficient parallelization. This defines a strong need for analytical approaches, and, based on recent successes, highlights the exploration of dynamic networks with complex update rules as an area where analytical work could outpace and guide numerical exploration.

A direct microscopic description of dynamical networks generally constitutes a very high-dimensional dynamical system. While in some cases exact analytical results were nevertheless obtained (e.g. \cite{53,60,61}), there are presently no approaches that are generally applicable. Much of the theoretical progress therefore relies on the derivation of reasonably low-dimensional coarse-grained approximations to the full microscopic model.

For networks in which the node can only assume states from a (small) discrete set of possibilities, approximation schemes are well established. These schemes are deeply rooted in physics and can be traced back to early work on the Ising model \cite{62,63}. In the networks literature there is presently a veritable zoo of different approximation schemes that build on similar principles but take different information into account. In the following we refer to these approaches as moment-closure approximations. The common idea in all of these approaches is to derive evolution equations for the abundance of certain subgraphs in

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the network. One starts with writing an evolution equation for small subgraphs, such as single nodes, before writing equations for larger motifs – a process that is reminiscent of classical moment expansions. The system of equations that is thus obtained generally depends on the abundance of other, typically larger, subgraphs that are not captured, and thus needs to be closed by estimating the abundance of these subgraphs – the actual moment-closure approximation.

Despite their underlying similarity moment-closure approximations proposed in the recent literature differ widely by the type and number of the subgraphs they capture. Generally speaking, capturing the dynamics of more subgraphs, leads to better approximations at the cost of having to deal with a larger system of equations (see [13, 23, 64, 67]). In practice some recently proposed schemes are successfully applied which only capture one or two subgraphs, while others capture thousands or millions of subgraphs.

For the analysis of adaptive networks, but also certain types of dynamics on static networks, moment-closure approximations are presently the most commonly applied theoretical tool. In adaptive networks they were used for instance to study epidemics [19, 24, 66, 27, 68, 72], collective motion [55, 56], evolution of cooperation [36, 23, 74], and social opinion formation [13, 15, 75, 76].

Despite the abundance of examples there is so far little intuition on when particular approximation schemes work and when they fail. This is most notable when considering the adaptive SIS model [19] and the adaptive voter model [13, 15]. Both of these models are adaptive network models of similar complexity, and, depending on personal taste, either can be considered as the most simple non-trivial adaptive network. However, for the adaptive SIS model, the dynamics can be faithfully captured already by simple approximation schemes [19], with more sophisticated approaches leading expectedly to a further improvement [23, 70, 71]. By contrast, for the adaptive voter model, simple approximation schemes only provide unsatisfactory results [13, 15] and, as we show here, more sophisticated approaches can actually perform worse.

In the present paper, we aim to offer an in-depth analysis of the performance and the failure of different approximation schemes. For the purpose of illustration we focus on the adaptive voter model as it provides a mathematically simple, yet challenging example system. To this system we apply the major approximation schemes proposed in the recent literature. Thereby, we build up an intuition of the advantages and disadvantages of the respective schemes.

A second goal of the present paper is to provide researchers entering the field with “worked examples” for the major approximation schemes. We present these examples in strongly abbreviated form in the main text, while providing all calculations in full detail in the appendices.

The paper is structured as follows: We start in Sec. 2 by introducing the adaptive voter model. In Sec. 3 we study the so-called homogeneous approximations, which result in relatively low dimensional equation systems. For these we explore in particular the effect of the order of approximation. Then, in Sec. 4 we discuss different heterogeneous approximations, which yield high-dimensional equation systems, but surprisingly do not significantly improve the performance of the approximation. Finally, in Sec. 5 we introduce a slightly different expansion that captures very similar information but works exactly in those parameter ranges where other approximations fail. A summary and discussion in Sec. 6 concludes the paper.

2. ADAPTIVE VOTER MODEL

The voter model considers the competition of equally attractive and mutually exclusive opinions (say A and B) in a population of interacting agents. The agents are represented by nodes that have an internal binary state variable, indicating the opinion held by the corresponding agents. The state is updated dynamically in time due to social interactions, occurring between linked agents.

In the original non-adaptive voter model [77] the underlying interaction topology is static. At each time step, a pair of nodes connected by a link is selected. If they share the same state, nothing happens. Otherwise, one of the two adopts the other’s state. This model has been explored in many subsequent works and in particular the dependence of the time needed to reach consensus on the underlying topology and details of the update rule is well understood [78, 86].

![Figure 1: (Color online) Parabola of active states. Shown are five representative trajectories for each of three parameter values and three different update rules. The long-term behavior depends on the value of the rewiring rate p. If this rate exceeds a threshold \( p^* \) then the system quickly approaches a fragmented state in which the number of active links \( |AB| \) vanishes, while the density of a given opinion (say, \( \langle A \rangle \)) remains almost constant. If \( p \) is below the threshold rewiring rate then the model approaches a parabola of meta-stable active states on which it remains for a long time while undergoing a random walk in the opinion density that eventually leads to consensus. Parameters: \( N = 10^5 \) and \( \langle k \rangle = 4 \).](image)

The adaptive variants of the model [12–15] additionally incorporates a phenomenon known as social segregation, the tendency of humans to link preferentially to others holding similar opinions and distance themselves from those holding opposite opinions. Starting from a suitable initial condition, a pair of nodes connected by a link is selected at each time step. If the link is inert, connecting nodes in the same state, then nothing...
happens. Otherwise, the link is active and either (with probability $p$) one of the nodes rewires the link by detaching it from the opinion, or one of the nodes copies the other node’s state.

In the spirit of discrete event simulations, using the Gillespie algorithm, the probability $p$ can be thought of as an effective parameter capturing the rate of rewiring events, normalized by the total rate of update events happening in the network. We therefore call $p$ the rewiring rate.

Previous investigations [12–16, 87] have shown that there is a critical rewiring rate, $p^*$, at which the behavior of the model changes qualitatively. If $p > p^*$ then the system quickly approaches a fragmented state in which the network splits into two disconnected components that hold opinion A and B, respectively. Since no active links survive the dynamics freezes in this fragmented state. By contrast, if $p < p^*$ then the system remains active for a long time before eventually reaching a complete consensus on one of the opinions, which is likewise an absorbing state (see Fig. 1).

We note many variants of the adaptive voter model using different update rules have been investigated in the literature [12–16, 75, 88–93]. Apart from models with more than two states[12, 18, 94] and models with directed links[16], the most prominent difference is the precise procedure for selecting the link that is updated and the direction of the update (e.g. which node adopts the others opinion) on this link. One distinguishes between direct node update, reverse node update, and link update rules. In the two node update rules, one selects a random node (node X) then a random neighbor (node Y). Selecting the nodes in this way creates a slight bias in the selected nodes regarding the degree, i.e. the number of neighbors of the respective nodes. Essentially, a node of higher degree has a proportionally higher probability of being found by following a link, and conversely the nodes found in this way have a higher degree. Therefore, the average degree of node Y is higher than the average degree of node X [13]. In models using direct node update, X retains the link in rewiring events and adopts Y’s state in update events, whereas in models using reverse node update Y retains the link in rewiring events and copies X’s opinion in update events. By contrast, in the link update rule one randomly selects a link, such that both selected nodes have identical statistical properties. While the precise update rule can have a significant impact on convergence times [14], the qualitative features of the models, described above remain unchanged (see Fig. 1).

In the remainder of this paper we apply different moment-closure approximations for capturing the dynamics of the voter model. Specifically, we compare the ability of the different approximation schemes to predict the density of active links in the state when both opinions are equally abundant.

The specific model that we consider throughout most of the paper is as follows: We start with an Erdős-Renyi random graph with $N$ nodes and $L$ links, such that the mean degree is $(k) = 2L/N$. Initial opinions are assigned randomly with equal probability. Therefore, the initial abundances obey $N_A = N_B = N/2$. The network is then evolved by a link update rule: At each time step an active link is selected at random. With probability $p$ one of the nodes rewires the link and connects to a random node of same state. Otherwise, that is with the complementary probability $1 − p$, one of the nodes adopts the other’s state. The respective node that retains the link or adopts the other’s opinion is selected randomly with equal probability. The model is simulated according to these rules until either fragmentation occurs or an active state is reached where the density of active links remains approximately constant over an intermediate time scale.

3. HOMOGENEOUS APPROXIMATIONS

For classifying the different approximation schemes that have been proposed, it is useful to distinguish between homogeneous and heterogeneous approximations. While all approximations attempt to capture the dynamics of certain subgraphs, they differ in the way in which subgraphs are identified. Homogeneous approximations {13–15, 19, 20, 36, 68, 74, 84, 95} classify subgraphs according to states of the nodes and the internal topology in the subgraph, whereas heterogeneous approximations additionally take the degree of the nodes in the subgraph into account {9, 17–19, 23, 72, 78, 74, 96–100}.

Homogeneous moment-closure approximations have been used in the past two decades to explain dynamics on static networks [64, 65, 95, 101, 102] and have more recently been applied to adaptive models of epidemic spreading [13, 15, 22, 74, 103], opinion formation [13–15, 73, 74], cooperation games [36, 73, 74], and collective motion [55, 56].

The central idea of homogeneous approximations is to capture the dynamics of the network by writing balance equations for the density or abundance of a certain set of labeled subgraphs (motifs), which are called network moments. The system of differential equations of network moments constitutes the moment expansion which is then closed by the so-called moment-closure approximation.

3.1. Moment Expansion

Writing the rate equation for a network moment necessitates calculating the rates of all possible processes that result in either the formation or destruction of the respective subgraph. For the adaptive voter model, the moment expansion for small subgraphs (nodes and links) has already been developed in Vazquez et al. [13] and independently for an identical model by Kimura and Hayakawa [15]:

\[
\frac{d}{dt}[A] = 0,
\]

\[
\frac{d}{dt}[AB] = \frac{1}{2}([AB] + \frac{(1 − p)}{2}(2[ABA] − [AAB]),
\]

\[
\frac{d}{dt}[BB] = \frac{1}{2}([AB] + \frac{(1 − p)}{2}(2[BAB] − [ABB]).
\] (1)

where $[X]$ denotes the density of nodes with state $X$, $[XY]$ denotes the density of links between nodes of state $X$ and $Y$, and $[XYZ]$ denotes the density of triplets constituted by a node of state $Y$ in the center and its two neighbors of state $X$ and $Z$, with $X, Y, Z \in \{A,B\}$. The densities are normalized with respect to
the number of nodes $N$, i.e. $|\Omega| = N_\Omega/N$ where we used $\Omega$ as a placeholder for an arbitrary subgraph (e.g. A-node, AB-link, AAB-triplet), such that $N_\Omega$ is the total abundance of that subgraph in the network.

The first equation in Eq. (1) states that the density of nodes of state A (and equivalently B) is conserved in the thermodynamic limit. This is peculiar to the voter model and is a direct consequence of symmetry in state adoption. In an update step, the A-node adopts state B with probability $(1 - p)/2$ and equivalently the B-node adopts state A with the same probability $(1 - p)/2$ leading to a vanishing net drift in the deterministic limit.

The second equation captures the change in the density of AA-links [AA]. If a B-node adopts state A on an AB-link (at rate $(1 - p)[AB]/2$) or an A-node rewire an AB-link and forms a link to another A-node (at rate $p[AB]/2$), an AA-link is directly created in the update. The total rate of this direct creation of AA-Links is thus $[AB]/2$. Additionally, AA-links can be created indirectly. Consider a B-node that is connected to two A-neighbors such that the three nodes form an ABA-triplet. When this B-node adopts state A, instead of one, two AA-links are formed. While the rate of adoption events creating A-nodes is $(1 - p)[AB]/2$ the expected additional links created indirectly in such adoption events are $2/[ABA]/[AB]$. Thus the total rate for the indirect creation of AA-links is $(1 - p)[ABA]$. Finally, AA-links can be destroyed indirectly if one of the A-nodes forming an AA-link adopts state B due to an interaction with another B-node, which happens at rate $(1 - p)[AAB]/2$. The rate equation for $[BB]$ is obtained by interchanging A and B in the second equation.

The system of equations (1) is not closed as it contains the (unknown) densities of larger subgraphs. This is a general property of moment expansions. For a system where the update rules directly affect subgraphs of diameter $d$ (here, 1) the evolution equation for a subgraph of diameter $l$ generally contains subgraphs of diameter up to $d + l$ due to indirect effects.

For a more precise discussion it is useful to define the order of a network moment as the number of links contained in the corresponding subgraph. For instance, the moments $\langle X \rangle$, $\langle XY \rangle$ and $\langle XYZ \rangle$ have order zero, one and two respectively. Accordingly, one can define the order of a network model as the largest order of subgraphs used in definition of the network update process, e.g. the voter model has order one since it involves a link at most in the definition.

Let us denote the model order as $o_M$, the set of node states as $S$ ($S = \{A, B\}$ in the model), and the set of network moments of order $o$ as $[\Omega_o]$, e.g. $[\Omega_0] = \{\langle X \rangle : X \in S\}$, $[\Omega_1] = \{\langle XY \rangle, \langle X\rangle, \langle Y \rangle : X, Y \in S\}$, and $[\Omega_2] = \{\langle XYZ \rangle, \langle XY \rangle, \langle X \rangle, \langle Y \rangle, \langle Z \rangle : X, Y, Z \in S\}$. Then,

$$d/dt[\omega_o] = f_{o_M}([\Omega_0], [\Omega_1], \ldots, [\Omega_{o_M}]),$$

where $[\omega_o] \in [\Omega_o]$ and $f_{o_M}$ is a $\mathbb{R}$-valued function that is moment specific.

We note that there are variants of moment expansions that differ in the definition of the set $\Omega_o$. One intuitive approach is to include all possible subgraphs that contain $o$ links. However, more appropriate basis of subgraphs can be constructed by considering the specific properties of the model. For instance, sparse random graphs tend to be locally tree-like. If a model does not have any particular update rules that actively create small cycles, cyclic motifs are exceedingly rare and can thus in general be safely ignored.

![Figure 2](image-url) (Color online) Validity of the moment expansion. The quasi-stationary density $[AB]$ at $A = [B] = 1/2$ from agent-based simulations is compared with $[AB]$ computed analytically from the moment expansion (Eq. (3) for link and reverse node update [Appendix D], and see [Appendix E] for direct node update). In this plot, the moment-closure approximation is avoided and the analytical results use the triplet densities $[AAB]$ and $[ABA]$ measured in the numerical simulation. Parameters: $N = 10^3$, $\langle k \rangle = 4$.

In order to test the validity of the moment expansion Eq. (1) for the adaptive voter model, we solve for the density $[AB]$ in the same equation at the steady state, which yields

$$[AB] = (1 - p)([AAB] - 2[AAB]).$$

(3)

To assess the validity of the expansion independently of the subsequent closure approximation we compute the expected number of active links for the respective triplet densities $[AAB]$ and $[ABA]$ observed in an agent-based simulation. The resulting diagram, Fig. 2, shows an almost perfect match for the active link density $[AB]$ with the numerical results for the model with link update. The match for node update rules is less good, in particular for low rewiring rates, but becomes better as we approach the transition point where the active links vanish and the network fragments.

Let us discuss the bad performance of the expansion for the node update rules in a little more detail. The moment expansion can only be exactly valid in the thermodynamic limit. However, the link update model shows that for the network sizes considered here finite size effects can be neglected. The only other
explanation is that the simulated system contains some correlations that are not picked up by the moment expansion. Because the effect of long-ranged correlations should have been captured. However, in the expansion we are implicitly assuming that the triplets are statistically distributed within the network, which is approximately true for the link update rule at low rewiring rates. By contrast in the node update rules the bias can induce additional correlations. This effect is counteracted by the rewiring as this mixes the network and thus improves the prediction.

As a side-note, let us remark that the beneficial effect of rewiring is a general property. In a given static network model a specific topological feature, such as a node of particular high degree or a densely clustered region, might exist that biases the system constantly in a specific direction. In an adaptive network such topological features may emerge for a limited time before being destroyed by topological dynamics. In the long run the effect of unlikely local configurations often averages out. In this sense adaptive networks are ensembles of themselves and can thus often be very well approximated unless correlations arise systematically that are not captured by the approximation scheme used.

3.2. Moment-Closure Approximation

In the previous section we have seen that moment expansions lead to an infinite hierarchy of equations. So far we truncated this expansion after the first order and used numerical values for the densities of larger subgraphs. Because we generally develop the expansion to obtain an analytical solution, we cannot rely on numerics, but must estimate the density of large subgraphs using a suitable approximation, known as moment-closure approximations.

For illustration let us approximate the second order moment \([ABA]\) in terms of the zeroth order moment \([B]\) and the first order moment \([AB]\) by the so-called pair approximation. An ABA-triplet comprises two adjacent AB-links sharing a common B-node. Therefore we first note that the density of single AB-links is \([AB]\). Next, we compute the probability that an additional A-node is connected to the B-node in this link.

Since we reached the B-node by following a link, we can expect it to have a higher-than-average degree. Specifically, the degree of the B-node in a randomly selected AB-link follows the distribution \(Q^B_k = kP^B_k/\langle[B]\rangle_{kB}\), where \(P^B_k\) denotes the probability that a randomly selected B-node has degree \(k\) and \(\langle k \rangle_B\) is the mean degree of a randomly selected B-node. The expected number of additional links of a B-node in an AB-link is thus \(\langle q_B \rangle = \sum_k (k-1)Q^B_k\), which is also known as the mean excess degree \([2]\) (of B-nodes).

A key assumption that we have to make at this point is that the AB-links are uncorrelated, except for the effect of the node degree described above. Using this assumption, each of the additional links of the B-node is an AB-link with probability \([AB]/\langle[AB] + 2[BB]\rangle\). Using \([AB] + 2[BB] = \langle k_B \rangle_B\), we find

\[
2[ABA] = \kappa_B \frac{\langle AB \rangle^2}{\langle B \rangle}, \tag{4}
\]

where \(\kappa_B = \langle q_B \rangle / \langle k_B \rangle\).

Similarly,

\[
[ABB] \approx \kappa_B \frac{2[BB][AB]}{[B]} \quad \text{and} \quad [BBB] \approx \kappa_B \frac{2[BB]^2}{[B]}.
\tag{5}
\]

The parameter \(\kappa_B\) that appears in these equations is generally not known because the degree distribution is reshaped by the rewiring process (although see \([16]\) for an exception). Governing \(\kappa_B\) are two counteracting effects, on the one hand we know that the degree of a node that is reached via a link is on average greater than the mean degree and on the other hand we have to subtract 1 from this increased degree, because we are only interested in the number of additional links. On Erdős-Rényi random graphs, these effects cancel exactly such that \(\kappa_B = 1\). Although the value of \(\kappa_B\) can be significantly higher in networks with broad degree distributions, assuming \(\kappa_B = 1\) has yielded good results for models with a fairly wide degree distribution \([19]\).

Substituting Eqs. (4, 5) into Eq. (1) and using the random-graph approximation \(\kappa_B = 1\) yields

\[
\frac{d}{dt}[A] = 0,
\]

\[
\frac{d}{dt}[AA] = \frac{1}{2}[AB] + \frac{(1-p)}{2} \left[\frac{[AB]^2}{[B]} - \frac{2[AA][AB]}{[A]}\right],
\]

\[
\frac{d}{dt}[BB] = \frac{1}{2}[AB] + \frac{(1-p)}{2} \left[\frac{[AB]^2}{[A]} - \frac{2[BB][AB]}{[B]}\right].
\tag{6}
\]

In order to test the performance of approximation we solve Eq. (6) for the stationary density of active links

\[
[AB] = \left(\frac{(\kappa_B)(1-p) - 1}{(1 - \kappa_B)}\right)[A](1 - [A]), \tag{7}
\]
which nicely captures the parabola shape of the states, shown in Fig. 1. Furthermore, considering the tip of the parabola \( [A] = 1/2 \), we find

\[
[AB] = \left( \frac{\langle k \rangle (1 - p) - 1}{4(1 - p)} \right).
\]

A comparison of the Eq. (8) with numerical results is shown in Fig. 3. The comparison shows that the approximation captures qualitative features of the model. The highest density of active links is found for \( p = 0 \), then as \( p \) is increased the density of active links declines and finally reaches zero at a finite rewiring event \( p^* \). However, the quantitative correspondence between the analytical and numerical results is very bad. In particular the pair approximation significantly overestimates the rewiring rate at which fragmentation occurs.

Let us investigate the reason of the bad performance of the approximation in more detail. We have already argued above that the random-graph approximation \( \kappa_3 = 1 \) is probably harmless. This can be confirmed by comparing the degree distribution observed in simulations to the Poissonian distribution of a random graph (Fig. 4). The comparison shows that the degree distribution stays very close to the Poissonian distribution. For such a close match the random-graph approximation is almost exact and cannot be the source of the major discrepancy observed in the results.

Accepting the validity of the moment expansion and ruling out the random-graph approximation as a source of errors leaves us with only two further sources of errors: We have assumed that a) the actual density of large motifs can be replaced with its expectation value and b) that correlations between active links can be neglected when computing this expectation value.

Let us first consider assumption a), which we call truncation assumption. Approximating any system by a lower-dimensional system is only possible if there is a time scale separation between slow low-order moments and fast higher-order moments \([68]\). The system then quickly converges to the slow manifold, characterized by the slow variables only \([104]\). Therefore, dynamics of moments higher than some order are enslaved to the dynamics of lower ones and they can be expressed as algebraic functions of low order slow moments. In our moment expansion the higher-order moments are disproportionately more likely to be affected by updates. For instance a single rewiring event affects one link, but approximately \( 2(\kappa_B(k))^2 \) triplets. While a more detailed investigation of this point would probably be fruitful, we conclude that assumption a) is probably not the main source of error in the present approximation scheme.

Accepting that the actual dynamical densities of higher-order moments can be replaced by their static expectation values, leaves us with the task of capturing the corresponding slow manifolds in a suitable functional form. Above, we derived such a functional form based on the assumption b), the absence of longer-ranged correlations. We can test this assumption by comparing the numbers of triplets observed in simulations to the expected values based on the observed number of nodes and links. This comparison, shown in Fig. 5, indicates that the expectation for ABB triplets diverges as the system approaches the fragmentation point \( p^* \). The sketch in the inset explains this failure. Close to fragmentation many active links are created by very few nodes that are in the wrong cluster. This induces a very high correlation between active links which is not captured by the pair approximation. Parameters: \( N = 10^5, \langle k \rangle = 4 \).

Figure 4: (Color online) Validity of the random-graph approximation. Comparison of the degree distribution of B-nodes, \( P^B_k \), in agent-based simulations with a Poisson distribution \( P_k = e^{-\langle k \rangle} \langle k \rangle^k / k! \). The comparison shows that the degree distribution of the adaptive network remains almost exactly Poissonian, and hence the random-graph approximation is valid to very good approximation. Parameters: \( N = 10^5, \langle k \rangle = 4 \).

Figure 5: (Color online) Test of the pair approximation. Shown is the ratio of the observed number of triplets in agent-based simulations and the expected number based on the observed number of nodes and links. The pair approximation is approximately valid for ABB triplets, whereas the error in the approximation of ABA triplets diverges as the system approaches the fragmentation point \( p^* \). The sketch in the inset explains this failure. Close to fragmentation many active links are created by very few nodes that are in the wrong cluster. This induces a very high correlation between active links which is not captured by the pair approximation. Parameters: \( N = 10^5, \langle k \rangle = 4 \).
nodes have sorted into two large clusters, connected by a few remaining links. On those links occasionally opinion adoption events take place which introduce some “wrong” nodes into the clusters and thus create many active links. The majority of active links is thus located on few nodes. This creates both a dis-

Figure 6: (Color online) Emergence of ABA-correlations close to fragmentation. Shown is \( T^{m}_{ABA} \), the fraction of B-nodes with \( m \) A-neighbors. Averages from agent-based simulations are compared with the expectation of an Erdős-Rényi random graph without any second neighbor correlations (see Appendix F). Close to the fragmentation transition, the A-neighbor distribution of ABA-triplets deviate from the pair approximation. Parameters: \( N = 10^7 \), \( (k) = 4 \).

3.3. Better homogeneous approximations

The reasoning presented above identifies the actual closure approximation, essentially assuming the absence of longer-ranged correlations as the reason for the failure of the pair approximation close to the fragmentation transition. Let us therefore discuss ways in which the approximation can be improved.

In principle every moment expansion should converge to the correct result if the order of the approximation is increased. Clearly despite its shortcomings the pair approximation is much better than the zeroth-order node approximation that ignores all, even link level, correlations and assumes that the network is well-mixed in terms of node states \([14]\). This is analogous to the molecular field approximation of the Ising model \([105]\].

The first-order approximation we used above is called the pair approximation \([13, 15, 19, 20, 36, 76, 80]\), which accounts for nearest-neighbor correlations but neglects higher order ones. This is analogous to the Bethe-Peierls approximation \([62, 63]\).

Increasing the order of the approximation is expected to yield better results \([15, 62, 63]\). An enormous number of subgraphs including exceedingly rare ones, which clearly cannot be of importance. Consider further-

Approximations that make the closure at higher orders are analogous to high order Kikuchi approximations \([109]\).

The performance of higher level-closures is shown in Fig. 2. Although the triplet-level closures perform better than the pair-level closure, the prediction is still very bad close to the bifurcation point. We can explain this result by considering Fig. 6 again. In the triplet-level closure we have to estimate the den-

sity of four-node subgraphs. In this estimation we use the as-

sumption that the four node subgraphs are uncorrelated. How-

ever, from the numerical results we know that many active links connect to nodes that have ten or more such links, which im-

plies also a high correlation at the four-node-level.

The reasoning above suggests that the order of the approx-

imation will have to raised beyond the mean degree of the sys-

tem to achieve a faithful result. Because of the technical dif-

ficulties described above this is clearly infeasible. However, note that by raising the expansion to this point we would be captur-

ing much information that is clearly not important. Considering that the number of subgraphs rises combinatorially with theor-

of equation-free modeling \([110]\) to extract a proper closure term for an epidemic model automatically from very short bursts of

in statistical physics. When the density of cycles is non-

negligible, i.e. when the network has high clustering, the pair approximation is extended in order to account for the cyclic motifs, corresponding to Bethe-Kirkwood type approximations \([67, 108]\). Approximations that make the closure at higher or-

of cycle is well-mixed in terms of node states \([14]\). This is analogous
to the Bethe-Peierls approximation \([62, 63]\).

A promising alternative approach is to use a relatively low order moment expansion, but use a more intelligent closure. For instance, Gross and Kevrekidis \([68]\) implements the approach of equation-free modeling \([110]\) to extract a proper closure term for an epidemic model automatically from very short bursts of
simulation runs. This enabled a semi-analytical investigation where continuation software was used to explore the dynamics of the system.

Another approach proposed in a recent paper [111] is to generate closures from a maximum entropy principle. This approach solves at least the problem of non-uniqueness of higher-order closures. However, in many cases the approach provides only implicit equations for the closure that do not seem to have a closed-form solution. This can be seen as an indication that explicit fully-consistent closure approximations for moment-expansions beyond the pair level might not exist at all.

4. HETEROGENEOUS APPROXIMATIONS

Presently it is widely believed that quite universally better results can be obtained by heterogeneous approximations that capture information on the degree of the nodes. Indeed, such approaches have yielded an improvement in several example systems [17, 23, 98, 102].

In this section, we investigate two prominent heterogeneous moment-closure approximations. In the heterogeneous pair approximation [98], links are grouped according to the state and the degree of the nodes at their ends. In the active neighborhood approximation [23, 99, 100, 112], nodes are placed in compartments according to their state, degree and number of neighbors in a given state.

Although the heterogeneous approaches can capture effects resulting from the heterogeneity of the degree distribution, they do not specifically address the complications identified above. Here, we test the performance of these approaches for the voter model, which reveals that they do not perform better than the homogeneous approximations, in this context.

4.1. Heterogeneous Pair Approximation

The heterogeneous pair approximation is based on writing a set of coupled rate equations for the density of active links \([AB]_{k,k'}\) between a node of degree \(k\) and a node of degree \(k'\). In networks with narrow degree distribution we expect these densities to be independent of \(k\) and \(k'\), whereas the same is not true for networks with broad degree distribution [98]. Here we follow an approach which is an extension of the one developed in [81, 98] for the case of adaptive networks. We consider the direct node update rule, since the derivation is easier in this case.

We start by writing the active link density as \([AB]_{k,k'} = [A_k B_{k'}] + [B_k A_{k'}]\), where, for instance, \([A_k B_{k'}]\) denotes the density of links connecting a node of state A and degree \(k\) with a node of state B and degree \(k'\). Assuming a node with state A (node \(i\)) is chosen in an update event, then

\[
d[A_k B_{k'}](t) = \sum_{j=1}^{N} \frac{[A_j]}{N} \sum_{n=0}^{N} \sum_{n=0}^{N} \sum_{n=0}^{N} \sum_{n=0}^{N} \sum_{n=0}^{N} \sum_{n=0}^{N} \frac{\Delta [A_i B_{k'}]}{N}
\]

\[
\times M(N_1, \ldots, N_6, l) \sum_{m=1}^{\infty} B(m, N_m)
\]

\[
\times \left\{ \begin{array}{l}
\frac{n}{l} (1-p) \left[ (N_i - n_k) \delta_{i,k'} - n_{k'} \delta_{i,k} \right] - \frac{n_{k'}}{l} p \delta_{i,k'} \\
+ \frac{p}{l} [n_{k'}] N(A_{k'} \mid B_{i} + A_{k'}) - n_{k'} N(A_{k'} \mid B_{i} + A_{k'}) \\
+ \frac{n}{l} \left[ (A_{k'} - 1) N(B_{k'} \mid A_{k'} - 1) - (A_{k'} - 1) N(B_{k'} \mid A_{k'}) \right] \right\} \frac{1}{N}.
\]

\[(9)\]
With probability \( [A_k] \), node \( i \) belongs to class \((A, l)\). The product \( M(N_1, ..., N_{S_l}) \prod_{m=1}^{S_l} B(n_m; N_m) \) expresses the probability that the configuration around node \( i \) consists of \( N_m \) links to neighbors of the degree class \( m \) (\( m = 1, ..., S_l \), with \( k \) the maximum degree) and \( n_m \) of these neighbors have the opposite state \( B \) (\( n_m = 0, ..., N_m \)). Here \( B(n_m; N_m) \) stands for the probability that \( n_m \) of the \( N_m \) links to neighbors of class \( m \) are active.

We distinguish again between direct and indirect changes in \([A_k B_{k'}]\). A direct change takes place when node \( i \) is in either class \( A_k \) or \( A_k' \), giving the first two terms inside the brackets of Eq. (2), respectively. Node \( i \) adopts state \( B \) when it copies the state of a randomly chosen B-neighbor, which happens with probability \( (1 - p)n/k \), where \( n = \sum_{m=1}^{S_l} n_m \) is the number of active links. In these events, the corresponding changes in the density \([A_k B_{k'}]\) are \( \Delta[A_k B_{k'}] = (N_k - n_k)/N \) and \( \Delta[A_k B_{k'}] = -n_k / N \), respectively (see Fig. 7-a).

An indirect change occurs due to an update on a neighboring node. The third term inside the brackets of Eq. (2) corresponds to the rewiring of an active link connected to a neighbor \( j \) of class \((B, k')\) (with probability \( pn_k/k \)), that results in the loss of an \( A_k B_{k'} \) link.

Other indirect changes take place when node \( i \) is in a generic class \( A_k \), and one of its links to a neighbor \( j \) is rewired, affecting the class of the links to the node \( j \), and the class of a-b links, where \( a \) is the node that receives the rewired link and node \( b \) is a neighbor of node \( a \) (see Figs. 7-b and c). In Fig. 7-b we describe the situation in which node \( j \) is in class \((B, k' + 1)\) (with probability \( n_k/k \)). Given that node \( j \) loses one link, it changes to class \((B, k')\), thus there is a gain of \( N(A_kB_{k'+1}A_k) \) links, represented in the fourth term of Eq. (3). The fifth term (see Fig. 7-c) corresponds to a loss in a similar update, when node \( j \) is in class \((B, k')\).

Finally, the last gain and loss terms belong to the case where any of the active links of node \( i \) is rewired (with probability \( pn_k/k \)) to a node \( a \) in class \((A, k - 1)\) (see Fig. 7-b), or in class \((A, k)\) (see Fig. 7-c), respectively. Given that the link is rewired to an \( A \)-node chosen at random, the probabilities for these events are \([A_{k-1}]/[A] \) and \([A]/[A] \), respectively. In the former event, the \( N(B_k'A_{k-1}) \) links of type \( A_{k-1}B_{k'} \) attached to node \( a \) change to type \( A_k B_{k'} \), while in the latter event the \( N(B_k'A_k) \) links of type \( A_k B_{k'-1} \) attached to node \( a \) change to type \( A_k B_{k'} \).

Given that \([A_k B_{k'}]\) may also change when a \( B \)-node is chosen, the evolution of \([A_k B_{k'}]\) is given by

\[
\frac{d[A_k B_{k'}]}{dt} = \left[ \frac{d[A_k B_{k'}]}{dt} \right]_A + \left[ \frac{d[A_k B_{k'}]}{dt} \right]_B.
\]

The second term on the right hand side can be obtained from Eq. 3 by interchanging \( A \) and \( k \) by \( B \) and \( k' \), respectively. By carrying out the summations, we arrive at the rate equation for the evolution of \([AB]_{k,k'}\) (see Appendix G) for the derivation.

\[
\frac{d[AB]_{k,k'}}{dt} = (1 - p) \left( Q_k \frac{k - 1}{k} \frac{[AB]_{k'}}{[AB]_{k}'} + \frac{k' - 1}{k'} \frac{[AB]_{k'}}{[AB]_{k}'} \right) - \frac{(1 - p)}{2\langle k \rangle [A] [B]} \left[ \frac{k - 1}{k} \frac{[AB]_{k}}{Q_k} + \frac{k' - 1}{k'} \frac{[AB]_{k'}}{Q_{k'}} \right] + \frac{1}{k} \frac{1}{k'} \frac{[AB]_{k,k'}}{[AB]_{k}'} + \frac{p}{4\langle k \rangle [A] [B]} \left( k'[AB]_{k,k'+1} [AB]_{k+1} \right.
\]

\[
\left. + \frac{k[AB]_{k+1,k} [AB]_{k+1} \right)}{Q_k} + \frac{\left( k' - 1 \right) \langle AB \rangle_{k'}}{Q_{k'}} + \frac{(k - 1) \langle AB \rangle_{k'}}{Q_k} \right) \frac{[AB]_{k,k'}}{[AB]_{k}'} + 2p \left\{ [AB]_{k-1,k} + [AB]_{k,k'-1} - 2[AB]_{k,k'} \right\} [AB],
\]

where \( Q_k = kP_k/(k) \) is the excess degree distribution, \( [A_k B] = \sum_{l=1}^{S} [A_k B_l]/[A] \), \( [AB] = [A] [B] \), \( [AB] = \sum_{l=1}^{S} ([A_k B_l] + [B_k A_l]), \) and \( [AB] = \sum_{l=1}^{S} [A_k B_l] \).

Equation (10) together with the consistency conditions \([AB]_{k} = \sum_{l=1}^{S} [A_k B_l]/[A] \) and \( [AB]_{k} = \sum_{l=1}^{S} [A_k B_l]/[A] \) form a closed system of coupled ordinary differential equations. Since the system is too complex to solve directly, we obtain the stationary solution by numerical integration starting from equiprobable \([A]_0 = [B]_0 = 0.5 \) initial states in a random graph with \( P_k = e^{-2/k} \).

In Fig. 8, we plot the global stationary density of active links \([AB] = \sum_{k=1}^{S} [AB]_{k} \) as a function of rewiring rate \( p \), and compare it with results from agent-based simulations. The heterogeneous pair approximation is in good agreement with simula-
tions for low $p$, but discrepancies become very large as $p$ increases. The overall performance is surprisingly even worse than the homogeneous pair approximation. This might be due to the accruing of discrepancies in individual terms $[A_i B_i]$. While the heterogeneous approximation certainly provides a more accurate description of networks with wide degree distribution, it does not suitably capture the correlations arising in the fragmentation transition.

4.2. Active Neighborhood Approach

Recently, an alternative heterogeneous moment-closure approach, the active neighborhood approach [23 99], was proposed to study the dynamics of the SIS model on adaptive networks [19]. Here, nodes are grouped by their compartment, defined according to their state, total degree and the number of active links. Therefore, not only the heterogeneity of the network is taken into account, but also the state correlations between nearest neighbors and associated heterogeneities. In this formalism, there is no need of estimating the neighborhood of a node, because this information is already contained in its class.

The active neighborhood approach has been applied to other epidemic systems [103, 112, 113], Glauber dynamics [100], and a voter-like model where the rewiring is state independent [17]. The approximation was found to reproduce the time evolution of both the states and the structure of the network with remarkable accuracy. A moment-generating function approach has been applied to mitigate the computational cost of the numerical integration [70, 71].

We now follow the active neighborhood approach for the adaptive voter model. We place nodes with state A (B), degree $k$, and $n$ ($n = 0, \ldots, k$) neighbors in the opposite state B (A), into the compartment labeled as $[A, k, n]$ ($[B, k, n]$). For reasons of simplicity, we assume that the time is continuous, thus that opinion adoption and rewiring processes take place in parallel. Every node in the network transmits its state to its neighbors at rate $\beta$, and rewires the connection from each neighbor in the opposite state to a random node with the same state at rate $\gamma$. That is, in a small time interval $dt$ all links are updated with the same probability $(\beta + \gamma)dt$. This “link homogeneous” dynamics is equivalent to the link update dynamics, in which links are chosen and updated with probability $1/L$, where $L$ is the number of links in the network. Therefore, stationary states obtained from numerical simulations of both dynamics are similar, as shown in Fig[10].

The evolution of $A_{k,n}$ is governed by the rate equation,

$$\frac{dA_{k,n}}{dt} = \beta \left[ (k-n)B_{k,n} - nA_{k,n} \right] + \beta \left[ A_{AB} \right] \frac{A_{AB}}{A_A} \left[ (k-n+1)B_{k,n-1} - (k-n)A_{k,n} \right] + \beta \frac{B_{AB}}{B_A} \left[ (n+1)B_{k,n+1} - nA_{k,n} \right] + \gamma \left[ (n+1)A_{k,n+1} - nA_{k,n} \right] + \gamma \left[ A_{k-1,n} - A_{k,n} \right] + \gamma \left[ (n+1)A_{k+1,n+1} - nA_{k,n} \right], \quad (11)$$

with the zeroth-order moments

$$A \equiv \sum_{k,n} A_{k,n} \quad \text{and} \quad B \equiv \sum_{k,n} B_{k,n}, \quad (12)$$

first-order moments

$$A_A \equiv \sum_{k,n} (k-n)A_{k,n} \quad \text{and} \quad A_B \equiv \sum_{k,n} nA_{k,n}, \quad (13)$$

and second-order moments

$$A_{AB} \equiv \sum_{k,n} n(k-n)A_{k,n} \quad \text{and} \quad B_{AA} \equiv \sum_{k,n} n^2B_{k,n}, \quad (14)$$

$$B_{BA} \equiv \sum_{k,n} n(k-n)B_{k,n} \quad \text{and} \quad A_{BB} \equiv \sum_{k,n} n^2A_{k,n}.$$}

These moments are related to the moments we defined in Sec.3 by $A = [A]$, $A_A = 2[A]$, $A_B = [AB]$, $A_{AB} = [AAB]$ and $A_{BB} = [AB] + 2[BAB]$.

Figure 9: Schematic representation of the possible update events in node states (a), (b) and (c) and links (d), (e) and (f), for the active neighborhood approach to the voter dynamics. Open and filled circles represent nodes in state A and B, respectively.

In Fig.9 we illustrate the six possible transitions of nodes from compartment $[A, k, n]$ to other compartments (we denote the reference node in compartment $[A, k, n]$ as node i), which correspond to the six loss terms in brackets of Eq. (11). The first of these describes the transition of node i from compartment $[A, k, n]$ to compartment $[B, k, k - n]$ at rate $\beta n$, when it adopts state B from an active neighbor (see Fig.9(a)). The second loss term describes the change of one of the $k - n$ A-neighbors of node i to B that happens at rate $\beta A_{AB}/A_A$, where $A_{AB}/A_A$ is the estimated number of the B-neighbors (see Fig.9(b)). This yields the transition of i to $[A, k, n + 1]$. The third term is analogous to the second term, but with the change $B \rightarrow A$ of one of the $n$ B-neighbors of node i at rate $\beta B_{BA}/B_A$ (see Fig.9(c)), that brings i to compartment $[A, k, n - 1]$. The fourth term represents the
replaced by an A-node at rewiring rate \( \gamma \) (see Fig. 9d), thus node \( i \) moves to \([A, k, n - 1]\). Node \( i \) gains a link coming from an A-node due to a rewiring event which occurs at rate \( \gamma A_B / A \) (see Fig. 9e). Node \( i \) switches accordingly to \([A, k + 1, n]\), represented by the fifth term. Finally, node \( i \) switches to the compartment \([A, k - 1, n - 1]\) when it loses a link due to the disconnection of one of its B-neighbors at rate \( \gamma n \) (see Fig. 9f). The gain terms can be explained analogously.

\[
\frac{\partial P_k}{\partial t} = \sum_{\ell} \left( p_{\ell k} A^\ell_k \right)
\]

\[\gamma A_B / A \]

We obtain the equation for the density \([B, k, n]\) by exchanging \( A \) and \( B \) in Eq. \(11\) using the symmetry of the model. This closes the system of equations. Numerical integration of the closed system of equations by standard numerical integration algorithms gives the values of the fractions \( A_{k,n} \) for a given time, and therefore, it allows one to obtain the time evolution of macroscopic variables, such as the densities of active links. We numerically solve Eq. \(11\) with initial conditions

\[A_{k,n}(0) = A_0 P_k \left( \frac{k}{n} \right) (1 - A_0)^n A_0^{n-k}\]

\[B_{k,n}(0) = B_0 P_k \left( \frac{k}{n} \right) B_0^n (1 - A_0)^{n-k}\]

where \( A_0 = B_0 = 1/2 \) and \( P_k = e^{-k^2} k! \). We determine the asymptotic values of the density of \( AB \)-pairs \( A_B = \sum_{k,n} n A_{k,n} \) for different values of the ratio between the rewiring and state adoption dynamics \( \gamma/\beta = p/(1 - p) \), where \( p \) is the rewiring probability.

Results from the active neighborhood approximation are compared with agent-based simulations in Fig. 10. The agreement between the analytical approach and simulations is good for small values of \( p \), but discrepancy increases with increasing \( p \), such that also the active neighborhood approach fails to capture the fragmentation transition faithfully.

5. ACTIVE MOTIF APPROACH

In conventional moment expansions of previous sections, moments are taken as densities of regular subgraphs, where subgraphs were characterized by a given number of links and prescribed node states, and degrees in case of heterogeneous moments. While such basis provide reasonable general purpose approaches, they do not take into account the specific dynamics of the system. The considerations presented in Sec. 3 and the failure of the heterogeneous approximations in Sec. 4 convey a clear message: To capture the fragmentation transition faithfully it is essential to capture the very heterogeneous distribution of active links that appears close to the fragmentation transition. Even the very complex and sophisticated active neighborhood approximation is in essence only a first order approximation and thus fails to pick up the correlations in the active links.

We can conclude that capturing the fragmentation transition faithfully requires tracking subgraphs of an order roughly up to the mean degree of the network. While the task of tracking all such subgraphs would be of enormous difficulty, it is greatly simplified, by tailoring the subgraph basis to the problem by using only those subgraphs that capture much information on the specific system. Our analysis above has shown that properties such as the density of ABB-triplets and even the degree distribution conforms very well to statistical expectations. By contrast the number of ABA-triplets and larger subgraphs comprising a number of active links attached to the same node defies statistical expectations close to the fragmentation point.

The reasoning above suggests that we should use a basis consisting of subgraphs that contain different numbers of active links attached to the same node. Two such active motif bases were proposed in B"ohme and Gross \[87\]. In application to the adaptive voter model, it was shown that both bases allowed a precise prediction of the transition point \[87\]. The approach was subsequently extended to multi-state voter models \[18\], where it likewise yielded good results. In the following, we briefly explain the approach and illustrate it by application to the adaptive voter model. Going beyond the previous works, we extend this approach to the estimation of active link densities, which allows a direct comparison with the approaches discussed above.

For computing the fragmentation point \( p^* \) we consider the situation where two communities holding opposite opinions have formed that are only connected by few active links. In analogy to the approaches discussed so far, considering the possible updates affecting subgraphs in the basis we derive a system of differential equations, governing the evolution of the subgraphs in time. However, because we are only concerned with subgraphs containing active links and such links are rare close to the fragmentation point, we arrive at a linear system of equations. The time evolution is thus fully captured by the eigenval-
The corresponding Jacobian is

\[ J = \begin{pmatrix} -1 & 2 \\ 1 - p & -1 - p \end{pmatrix}, \]

and the condition \( \lambda(p, 3) = 0 \) yields \( p^* = 1/3 \) for the transition point. The described procedure can be generalized to arbitrary \( k \). For \( k = 4 \) (the degree considered here), the predicted transition point is in good agreement with the value \( (p^*) \approx 0.445 \) from agent-based simulations which violates the assumption of degree regularity.

In order to account for a degree heterogeneous network, a basis set of \( \{m, l\} \)-spiders is used: a spider motif consists of one central base node which is connected to \( m \) nodes of its own opinion and \( l \) nodes of opposing opinion. The dynamical evolution equation for spider motifs is provided in [Appendix 1] and leads to a very close approximation of the true transition point.

In the estimation above for the calculation of the transition point, we assumed that in a \( q \)-fan motif, all neighbors of the fringe nodes (except for the base node) hold the same opinion as the fringe nodes. This assumption is valid for vanishing active link density at the symmetric state \( \sigma = |AB|/(|k|A) | \to 0 \). Now we consider the case \( p < p^* \), where there is a finite density of active links. In our equations we therefore have to include the possibility that a fringe node already holds outer active links (apart from the one connecting to the base node), which become inert when the fringe node adopts the state of the base node. By utilizing the observation that active links tend to gather, rather than distribute homogeneously over the whole system, we assume that whenever a new \( k-1 \)-fan is created, this fan is either active, with probability \( (1-\sigma) \), or inactive with probability \( \sigma \).

In Fig. 12 an example for the degree regular case \( (k = 3) \) is illustrated, where we include the active link density in the transition probabilities. As before, this can be summarized in a system of evolution equations for the motif densities

\[ \frac{d[1]}{dt} = -(1) + 2[2], \]
\[ \frac{d[2]}{dt} = -2[2] + (1 - p)[1] + (1 - p)[2]. \]  

Note that for \( \sigma = 0 \) we recover Eq. 16.

The corresponding Jacobian now depends on \( p \) and \( \sigma \). Solving \( \lambda(p, \sigma) = 0 \) yields

\[ \sigma(p) = \frac{4(1 - 2p) - (1 - p)^2}{8(1 - p) - (1 - p)^2}. \]
even the evolution of problematic models can be captured if with other long-ranged state correlations [30, 119, 120].

networks, approximately bipartite nets, or complex topologies self-organize into specific topologies such as leader-follower failure close to the fragmentation point because some specific correlations appear. These correlations can be generally expected to arise in fragmentation transitions regardless of the specific situation, whereas for small rewiring rates it becomes very bad as active links are increasingly well mixed.

6. SUMMARY AND DISCUSSION

In this paper we investigated the performance of moment-closure approximations for discrete adaptive networks. In particular we used the adaptive voter model as a benchmark model to assess different approaches. The comparison with agent-based simulations revealed that both homogeneous and heterogeneous moment-closure approximations capture qualitative properties of the fragmentation transition, but fail to provide good quantitative estimates close to the fragmentation point. Remarkably, even very sophisticated heterogeneous approaches can produce results that are worse than those from simple homogeneous schemes. Finally, we identified the active motif approximations as a class of approximations that were able to capture the behavior close to fragmentation point quantitatively.

The present results are likely to hold in a much larger class of models. In the adaptive voter model conventional approaches fail close to the fragmentation point because some specific correlations appear. These correlations can be generally expected to arise in fragmentation transitions regardless of the specific model under consideration and should thus occur in a large variety of models [12, 15, 17, 18, 73, 88, 114–118]. We further expect that similar correlations could arise in networks that self-organize into specific topologies such as leader-follower networks, approximately bipartite nets, or complex topologies with other long-ranged state correlations [30, 113, 120].

Perhaps the main message from the current work is that even the evolution of problematic models can be captured if an expansion is used that is tailored specifically to the system at hand. In this paper we have used extensive numerical and analytical investigations to identify the problematic correlations. However, in retrospect just considering a sketch of the situation close to fragmentation, such as the inset in Fig. 5 could have pointed us to these correlations and hence to a suitable approximation scheme.

Anticipating the structures that are likely to emerge prominently in a given model should generally allow to identify a suitable approximation scheme. Using the spider or fan motifs of the active motif approximation will improve predictions in models close to fragmentation or related de-mixing transitions. By contrast, including such motifs in the subgraph basis of an approximation scheme could be cumbersome and even have an adverse effect in models that remain well-mixed. Models that are prone to evolve degree-state correlations or strong clustering will require specific approaches such as [64, 65, 101, 121–124] (see [102] for a review). By contrast, in models with strong random rewiring and sufficiently low degree, small cycles should be rare and thus subgraphs containing cycles can (and should) be ignored when selecting the approximation scheme. Clearly, for all models that tend to evolve very heterogeneous degree distributions, heterogeneous approximations such as the heterogeneous pair approximation or the active neighborhood approximation are required. However, even in the case when degree distributions become broad but not exceedingly broad, such as in Gross et al. [19], it may be worth to consider homogeneous approximations as they may still provide relatively good results, at a significantly lower cost than the heterogeneous approximations.

Finally, we note even for the deceptively simple adaptive voter model, we have not yet identified an approximation that works well over the whole parameter range. While the active motif approximation yields faithful results in the ordered states close to fragmentation it fails in well-mixed systems. Conversely, all other approaches studied here work reasonably well when far from fragmentation, but fail at the fragmentation point. In the future a scheme that works over the whole range of rewiring rates may be found, either as a reasonable interpolation between the existing approaches, or by the informed design of a suitable approximation. We hope that the information gathered in the present survey will contribute to reaching this goal.

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Appendix A. The origin of second-order terms in the first-order moment expansion

Here, we derive the triplet density terms in the moment expansion Eq. 1 from local selection events and combinatorics. We illustrate it on $Q(AB|A)$ which denotes the number of additional A-neighbors of the B-end of a randomly selected AB-link.
The quantity \( Q(ABA) \) can be written as \( Q(ABA) = P(ABA) \langle q_B \rangle \), where \( P(ABA) \) denotes the probability that a random neighbor of the B-end of a randomly selected AB-link has state A and \( \langle q_B \rangle \) denotes the mean excess degree, as derived in the main text. The probability \( P(ABA) \) is expressed as a ratio of the corresponding triplet densities, i.e. \( P(ABA) = 2[ABA]/(2[ABA] + [ABB]) \), where \([ABA] = N_{ABA}/N \) (\( N_{ABA} \) is the number of ABA-triplets) and analogously \([ABB] \equiv N_{ABB}/N \).

Let \( k_i \) and \( n_i \) be the number of all (A and B) neighbors and A-neighbors of a B-node selected at random (node i) respectively. The number of triplets is obtained from a summation over all B-nodes: \( N_{ABA} = \sum_{i \in B} n_i(n_i - 1)/2 \) and \( N_{ABB} = \sum_{i \in B} n_i k_i \), where \( B \) is the set of B-nodes. We thus obtain \( 2N_{ABA} + N_{ABB} = \sum_{i \in B} n_i(k_i - 1) \). Now, we proceed with the summation

\[
\sum_{i \in B} n_i(k_i - 1) = \sum_{i \in B} n_i k_i - \sum_{i \in B} n_i = \sum_{j \in A} k_i - N[AB] = N[AB]\left(\sum_k k \frac{kP^R}{[B][k_B]} - 1\right)
\]

\[
= N[AB] \sum_k (k - 1) kP^R / ([B][k_B]) = N[AB][q_B]
\]

and obtain \( 2[ABA] + [ABB] = (2N_{ABA} + N_{ABB})/N = [AB]\langle q_B \rangle \).

By this, we reach

\[
Q(ABA) = \frac{2[ABA]}{[AB]}
\]

Appendix B. Second order moment expansion for the link update rule

Since the right-hand side of Eq. [1] involves second-order moments, the first-order expansion is not closed. We now treat the second-order moments as dynamical variables and derive the corresponding rate equations. In Fig. B.14, we illustrate the second-order moments as dynamical variables and derive the update rule

\[
\frac{d}{dt}[A] = 0,
\]

\[
\frac{d}{dt}[AA] = \frac{1 - p}{2}([AB] + 2[ABA] - [AAB]) + \frac{p}{2}[AB],
\]

\[
\frac{d}{dt}[BB] = \frac{1 - p}{2}([AB] + 2[BAB] - [ABB]) + \frac{p}{2}[AB],
\]

\[
\frac{d}{dt}[AAA] = \frac{1 - p}{2}(2[ABA] + [AAB] + [ABB]) + \frac{p}{2}[ABAB] + [ABA]
\]

\[
+ 3[(B)AAA] - [AAB] - [(A)BAAB] + \frac{p}{2}[ABA] + \frac{1}{2} [AAB] B_{6}^{AAB} - 2[ABA] - 2[(A)BAB] + \frac{p}{2}[ABA] + \frac{1}{2} [AAB] B_{6}^{ABA} - 3[(B)AAA] - 4p[ABA],
\]

(B.1)

where \([XYZW]\) denotes the density of chain-quadruplets constituted by a node of state X (node 1), a Y-neighbor of node 1 (node 2), a Z-neighbor of node 2 (node 3), and a W-neighbor of node 3 (node 4), and \([(X)Y\ldots W]\) denotes the density of star-quadruplets constituted by a node of state X at the center and its three neighbors of state Y, Z, and W with X, Y, Z, W \(\in \{A,B\} \).

For illustration, we derive the corresponding contributions to \(d[ABA]/dt\) for processes (e) and (h) in Fig. B.14.

In the process (e), an ABA-triplet is created per each AB-link connected through its B-node to the B-end of an AB-link on which B-node adopts state A. We denote the A (B) end of a randomly selected AB-link as node X (Y) and a random excess neighbor of node Y as node Z. The expected number of AB-links connected through its B-node to node Y is \(Q(ABA) = P(ABA)K_{BA}\), where \(P(ABA)\) is the probability that node Z has state B and a random neighbor of node Z has state A and \(K_{BA}\) is the expected number of excess triplets attached to node Y.

Node Y has degree k with probability \(kP^R/(\langle B(k_B) \rangle)\). Assuming neutral mixing by degree, node Z has degree \(k'\) with probability \(k'P^R/(\langle k' \rangle)\). Thus, \(K_{BA} = \langle \sum_k (k - 1) kP^R / \langle B(k_B) \rangle \rangle \times \langle \sum_k (k' - 1) kP^R / \langle k' \rangle \rangle = \langle q_B \rangle \langle q \rangle^2 \).

Analogous to the expression of \(P(ABA)\) in Appendix A, we write the conditional probability \(P(AB|BA)\) as a fraction of appropriate network moments, i.e. \(P(AB|BA) = [2[ABA] + [ABBB] + [ABAB] + [ABAA]]/[2[ABA] + [ABB] + [AAB] + [ABA]]\). In order to approximate the denominator, we use the second-order approximation for chain-quadruplets, which is derived in Appendix C.
following this approximation,

\[ 2[ABBA] + [ABB] + [ABAB] + [BAAA] \approx [AB][[AB] + 2[BBB]] = \frac{2[BB]}{[AB]} = \langle q \rangle [AB] + 2[ABA] \]

Thus, \( P(ABA) = 2[ABBA]/\langle (q)^2 \rangle [AB] \). Since \( K_{AB} = \langle q \rangle^2 \), we reach

\[ Q(AB|BA) = 2[ABBA]/[AB]. \tag{B.2} \]

Similarly,

\[ Q(AA|AB) = \frac{[AAAB]}{[AB]}, \]

\[ Q(AB|AB) = \frac{[ABAB]}{[AB]}, \]

\[ Q(BA|AB) = 2[BAAB]/[AB], \]

\[ Q(BB|AB) = \frac{[BBAB]}{[AB]}. \tag{B.3} \]

Furthermore, in the process (h), an ABA-triplet becomes an AAA-triplet per every two A-neighbors of the B-end of a randomly selected AB-link. We call such subgraphs constituted by a node and its three neighbors as stars. We again denote the A (B) end of a randomly selected AB-link as node X (Y). We denote the expected number of the configurations where node Y is connected to two additional A-nodes as \( Q(A, A|BA) = P(A, A|BA)S_B \), where \( P(A, A|BA) \) is the probability that two random neighbors of node Y both have state A, and \( S_B \) is the average number of two-combinations of excess neighbors of node Y.

Node Y has degree \( k \) with probability \( kP_k/B(k_b) \). The term \( S_B \) is obtained from \( S_B = \sum_k (k - 2)(k - 1)kP_k/(2B(k_b)) \). In terms of statistical moments of the degree distribution,

\[ S_B = \frac{\langle k_B^2 \rangle - 3\langle k_B \rangle + 2\langle k_B \rangle}{2B(k_b)}. \]

We write the conditional probability \( P(A, A|BA) \) as a fraction of appropriate moments as before such that \( P(A, A|BA) = 3[(B)AAA]/[(B)AAA] + 2[(B)AB] + [(B)ABB] \). The density \( [(B)AAA] \) is defined as \( [(B)AAA] = N_{(B)AAA}/N \), where \( N_{(B)AAA} \) is the total number of (B)AAA-stars (central B-node, three A-neighbors). The densities \( [(B)AB] = N_{(B)AB}/N \) and \( [(B)ABB] = N_{(B)ABB}/N \) are defined analogously. The number of stars is obtained from a summation over B-nodes such that \( N_{(B)AAA} = \sum_{i\in[B]} n_i(n_i - 1)(n_i - 2)/6 \), \( N_{(B)AB} = \sum_{i\in[B]} n_i(n_i - 1)(k_i - n_i)/2 \), and \( N_{(B)ABB} = \sum_{i\in[B]} n_i(k_i - n_i)(k_i - n_i - 1)/2 \), where \( k_i \) and \( n_i \) are the number of all neighbors and A-neighbors of node i respectively. We reach the expression \( 3N_{(B)AAA} + 2N_{(B)AB} + N_{(B)ABB} = \sum_{i\in[B]} n_i(k_i^2 - 3k_i + 2)/2 \) and proceed with the summation

\[ \sum_{i\in[B]} n_i(k_i^2 - 3k_i + 2) = \sum_{i\in[B]} \sum_{j\in[A]} (k_j^2 - 3k_j + 2) \]

and obtain \( 3[(B)AAA] + 2[(B)AB] + [(B)ABB] = [AB]S_B \). We reach

\[ Q(A, A|BA) = 3[(B)AAA]/[AB]. \tag{B.4} \]

Similarly,

\[ Q(A, A|AB) = \frac{[(A)ABA]}{[AB]}, \]

\[ Q(A, B|AB) = 2[(A)BBA]/[AB], \]

\[ Q(B, B|AB) = 3[(A)BBA]/[AB]. \tag{B.5} \]

Appendix C. Second-order moment closure approximation

In order to close the expansion Eq. [B.1], we need to express third-order moments in terms of lower order ones.

We first derive the second-order approximation for the chain quadruplet density [ABBA]. We start with the BB-link at the
center, which has density \([BB]\), and estimate the number of the A-nodes connected to the two ends, between which no correlation exists according to the second-order approximation. Each B-end has \([ABB]/2[BB]\) A-neighbors on average, as obtained in Appendix A. We reach the following second-order approximation: \(2[ABB]/2[BB]\). We note that 2 appears on both sides, because we can't distinguish between the two B-nodes in the BB-link and the two ABB-triplets.

We derive the remaining terms for chain quadruplets analogously and obtain
\[
\begin{align*}
[AAAB] & \approx \frac{[AAA][AAB]}{[AA]}, \\
[ABAA] & \approx 2 \frac{[ABA][AAB]}{[AB]}, \\
[ABAB] & \approx 4 \frac{[ABA][BAB]}{[AB]}, \\
[ABBA] & \approx \frac{[ABB]^2}{4[BB]}.
\end{align*}
\]

We now illustrate the second-order approximation for the star motif density \([AB][BA]\), which is the density of subgraphs formed by an A-node in the center and its three (two B and one A) neighbors. We consider the star motif as an assembly of two triplets originating from the same end of a shared link. We start with locating an AB-link that appears at density \([AB]\). Each excess neighbor of the A-node has state A with probability \([ABB]/q[AB]\) and state B with probability \(2[BAB]/q[AB]\). The expected number of star-quadruplets originating from one end of a random link is \(1/2(k^3 - 3(k^2) + 2(k))/k\), where \((k^n)\) is the \(n^{th}\) moment of the degree distribution. By using it, we obtain \(2[(AB)BA] = \langle k\rangle((k^3 - 3(k^2) + 2(k))/k) - \langle k\rangle^2 \langle 2[BAB][AAB]/[AB]\rangle). If we assume a Poissonian distribution, the second moment \(\langle k^2\rangle = \langle k\rangle^2 + \langle k\rangle\) and third moment \(\langle k^3\rangle = \langle k\rangle^3 + 3\langle k^2\rangle + \langle k\rangle\). Thence, we reach the approximation
\[
[(AB)BA] \approx \frac{[AAB][BAB]}{[AB]}.
\]

However, approximations for star-quadruplet densities are not unique. It depends on which pair is considered to be shared by the two constituent triplets. Above we took the shared link to be an AB-link. We could also select the AA-link instead as the shared link which leads to a different approximation. As considering such alternative ways of motif construction, star quadruplets can be approximated as
\[
\begin{align*}
[(AB)AA] & \approx \frac{[AAB]^2}{2[AB]} \text{ or } \frac{[AAB][AAA]}{2[AA]}, \\
[(AB)BA] & \approx \frac{[AAB]^2}{4[AA]} \text{ or } \frac{[AAB][BAB]}{[AB]}, \\
[(AB)BB] & \approx \frac{[BAB]^2}{3[AB]}.
\end{align*}
\]

Appendix D. Reverse node update rule

Moment expansion equations for the reverse node update rule are very similar to those of the link update rule with the following changes in rates.

An A-node adopts state B at rate \((1 - p)[A][AB]/\langle k_A\rangle\) and an AB-link is replaced by an BB-link through rewiring at rate \(p[A][AB]/\langle k_B\rangle\). Analogously, a B-node adopts state A at rate \((1 - p)[B][AB]/\langle k_B\rangle\) and an AB-link is replaced by an AA-link through rewiring at rate \(p[B][AB]/\langle k_A\rangle\).

The degree distributions of the nodes selected at an update event depend on the update rule. In the link update rule, both of the nodes at the two ends of a selected link have degree \(k\) with the same probability \(kP_k/k\). In the reverse node update rule, the first selected node (node X) has degree \(k'\) with probability \(P_{k'}\) and its random neighbor (node Y) has degree \(k\) with probability \(kP_k/k\). The degree distribution of node X does not show up in the first-order equation and the degree distribution of node Y is the same as in the link update rule. Therefore, the first-order equation for the reverse node update rule differs from that of the link update rule due to the changes of rates of events only
\[
\begin{align*}
\frac{d}{dt}[A] & = (1 - p)\left[\frac{[A][AB]}{\langle k_A\rangle} - \frac{[AB]}{\langle k_B\rangle}\right], \\
\frac{d}{dt}[AA] & = (1 - p)\left[\frac{[A][AB]}{\langle k_A\rangle} + \frac{[AB]^2}{\langle k_A\rangle} - \frac{2[AB][BB]}{\langle k_B\rangle}\right] \\
& \quad + p\left[\frac{[A][AB]}{\langle k_A\rangle} - \frac{[AB]}{\langle k_B\rangle}\right], \\
\frac{d}{dt}[BB] & = (1 - p)\left[\frac{[A][AB]}{\langle k_B\rangle} + \frac{[AB]^2}{\langle k_A\rangle} - \frac{2[AB][AA]}{\langle k_A\rangle}\right] \\
& \quad + p\left[\frac{[A][AB]}{\langle k_A\rangle} - \frac{[AB]}{\langle k_B\rangle}\right].
\end{align*}
\]

At the symmetric state \([A] = [B] = 1/2\), Eq. (D.1) is equivalent to Eq. (6) with the rescaled time \(t' = \langle k\rangle t\).

Appendix E. Direct node update rule

An A-node adopts state A at rate \((1 - p)[A][AB]/\langle k_A\rangle\) and an AB-link is replaced by an AA-link through rewiring at rate \(p[A][AB]/\langle k_B\rangle\). Similarly, a B-node adopts state A at rate \((1 - p)[B][AB]/\langle k_B\rangle\) and an AB-link is replaced by a BB-link at rate \(p[B][AB]/\langle k_A\rangle\).

To estimate the indirect contribution term, we use the quantity \(Q(A|A_1B_2)\) that represents the average number of A-neighbors of an A-node given that it already has a B-neighbor. Indices 1 and 2 indicate that the A-node was chosen first and the B-node was chosen after that. Given that node X is chosen at random in the direct node update rule, it has degree \(k\) with probability \(P_k\). We also know that X has a B-neighbor, thus the probability that each of the \(k - 1\) remaining neighbors has state A can be estimated as \([AAB]/\langle q_A\rangle[AB]\). Averaging over the entire network we obtain
\[
Q(A|A_1B_2) = \sum_k (k - 1)P_k \frac{[AAB]}{\langle q_A\rangle[AB]} = \frac{\langle k_A\rangle - 1}{\langle q_A\rangle} \frac{[AAB]}{[AB]}.
\]

Then, the following set of equations is reached for the direct node update rule:
in terms of triplet density in Eq. E.1

\[
\begin{align*}
\frac{d}{dt}[A] &= (1 - p) \left( \frac{[AB]}{\langle kb \rangle} - \frac{[AB]}{\langle kA \rangle} \right),
\end{align*}
\]

\[
\begin{align*}
\frac{d}{dt}[AA] &= (1 - p) \left( \frac{[AB]}{\langle kb \rangle} + \frac{2(\langle kA \rangle - 1)[AB]}{\langle kA \rangle^2} \right)
- \frac{(\langle kA \rangle - 1)[AA]}{\langle kA \rangle} + p \frac{[AB]}{\langle kA \rangle},
\end{align*}
\]

\[
\begin{align*}
\frac{d}{dt}[BB] &= (1 - p) \left( \frac{[AB]}{\langle kb \rangle} + \frac{2(\langle kB \rangle - 1)[BAB]}{\langle kA \rangle^2} \right)
- \frac{(\langle kB \rangle - 1)[BBA]}{\langle kB \rangle} + p \frac{[AB]}{\langle kB \rangle}.
\end{align*}
\]

(E.1)

At the steady-state, the link density \([AB]\) can be expressed in terms of triplet density in Eq. E.1

\[
[AB] = \frac{(1 - p)(\langle k \rangle - 1)}{\kappa(k)} ([AA] - 2[ABA]).
\]

(E.2)

However, this requires numerical values of the triplet densities that should be measured from the agent-based simulations, which is undesirable as explained in the main text. Instead, we develop a first-order moment closure approximation. Since node X is selected without resorting to the information of node Y, node Y should be treated as a sample from the neighbors of node X. An alternative solution to this problem is to ignore the information from node Y and use a pair approximation. Then, any neighbor of node X has state A with probability \(2[AA]/[A(k)]\). The pair approximation formulation for \(Q(A|A_1B_2)\) is

\[
Q(A|A_1B_2) \approx Q(A|A_1) = \sum_k P_k(k - 1) \frac{2[AA]}{[A(k)]} = \frac{2(\langle kA \rangle - 1)[AA]}{\langle kA \rangle}.
\]

This is equivalent to using the pair approximation for the density of triplets, such as

\[
[AA] \approx 2\kappa(k) \frac{[AA][AB]}{[A]}. \quad \text{(E.1)}
\]

Replacing this expression for \([AA]\) and the analogous expressions for the other densities of triplets in Eq. E.1, we arrive at

\[
\begin{align*}
\frac{d}{dt}[A] &= (1 - p) \left( \frac{[AB]}{\langle kb \rangle} - \frac{[AB]}{\langle kA \rangle} \right),
\end{align*}
\]

\[
\begin{align*}
\frac{d}{dt}[AA] &= (1 - p) \left( \frac{[AB]}{\langle kb \rangle} + \frac{2(\langle kA \rangle - 1)[AB]}{\langle kA \rangle^2} \right)
- \frac{2(\langle kA \rangle - 1)[AA][AB]}{\langle kA \rangle^2} + p \frac{[AB]}{\langle kA \rangle},
\end{align*}
\]

\[
\begin{align*}
\frac{d}{dt}[BB] &= (1 - p) \left( \frac{[AB]}{\langle kb \rangle} + \frac{2(\langle kB \rangle - 1)[BAB]}{\langle kA \rangle^2} \right)
- \frac{2(\langle kB \rangle - 1)[BBA][AB]}{\langle kB \rangle^2} + p \frac{[AB]}{\langle kB \rangle}.
\end{align*}
\]

(E.3)

Appendix F. Derivation of pair approximation for the fraction of active triplets

Let’s denote \(B_k^m\) as the fraction of B-nodes with k total (A and B) and \(m\) A neighbors with the normalization \(\sum_k \sigma_k^m = 1\). The fraction \(B_k^m\) is expressed as \(B_k^m = P_k^m P(m,k)/|B|\), where \(P_k^m / |B|\) is the probability of the B-nodes with degree k and \(P(m,k)\) is the probability that \(m\) links out of \(k\) are active. For an infinite Erdős-Rényi random graph, \(P_k^m / |B|\) is Poisson distributed, i.e. \(P_k^m / |B| = (k)^k e^{-k}/k!\). If we neglect state correlations to second nearest-neighbors, then every link connected to a B-node is active with probability \(\sigma_{p,(k)} = [AB]_{p,(k)} / |B(k)| = 2[AB]_{p,(k)} / (k)\), at \(|B| = 1/2\), where we measure the average active link density \([AB]_{p,(k)}^v\) at a specific p and (k). When second nearest-neighbor correlations are ignored, \(P(m,k)\) is a binomial distribution, i.e. \(P(m,k) = (m!(k-m))!k! e^{m(k)}(1 - \sigma_{p,(k)})^{k-m}\). Therefore, in the uncorrelated infinite-size Erdős-Rényi case \(B_k^m\) obeys

\[
\frac{d[B_k^m]}{dt} = \frac{m!}{k^2} e^{(k-m)(1 - \sigma_{p,(k)})^{k-m}}. \quad \text{(F.1)}
\]

The quantity \(T_{ABA}^m\) is the fraction of ABA-triplets that have a B-node with \(m\) A-neighbors. By definition,

\[
T_{ABA}^m = \sum_{k=2}^{\infty} \frac{1}{2m(m-1)} B_k^m. \quad \text{(F.2)}
\]

Appendix G. Derivation of the heterogeneous pair approximation

Here, we first derive the rate equation for the average change in the density of active links \([A_kB_k]\) when a node with state A is chosen. We rewrite Eq. (9) as

\[
\frac{d[A_kB_k]}{dt} = \sum_{i=1}^{\infty} \sum_{j=0}^{\infty} \sum_{l=0}^{\infty} \sum_{m=1}^{\infty} \text{S}(N_1, \ldots, N_\tau) B(n_m,N_m), \quad \text{G.1}
\]

where

\[
\text{S}(N_1, \ldots, N_\tau) = \sum_{n_1=0}^{N_1} \cdots \sum_{n_\tau=0}^{N_\tau} \prod_{m=1}^{\tau} B(n_m, N_m)
\]

\[
\left\{ \begin{array}{l}
(1 - p) \left[ N_k n_k - n_k^2 + (N_k - n_k) \sum_{m+k} n_m \delta_{i,k} \right] 
- (1 - p) n_k^2 + n_k \sum_{m+k} n_m \delta_{i,k} - n_n \delta_{i,k} 
+ \nu n_k \sum_{m+k} n_m \delta_{i,k} - n_n \delta_{i,k} 
= (1 - \sigma_{p,(k)})^k - \sigma_{p,(k)} M(A_{k-1},A_k) - \sigma_{p,(k)} M(A_k,A_{k-1}) 
\end{array} \right. \left\} \right. \times \sum_{m=1}^{\tau} n_m \cdots \}
\]

(G.2)

We have also replaced the number of active links \(n\) around the chosen node by \(\sum_{m=1}^{\tau} n_m\). To carry out the summations in
Eqs. (G.1) and (G.2), we assume that the network has no degree correlations, thus the probability that a given node has a neighbor of degree \( m \) is \( Q_m = m P_m / \langle k \rangle \). Then, the probabilities \( M \) and \( B \) become the multinomial and binomial distributions, respectively,

\[
M(N_1, \ldots, N_{r'}; l; Q_1, \ldots, Q_r) = \begin{cases} \binom{N}{N_1} \cdots \binom{N}{N_r} & \text{when } \sum_m l_m = l; \\ 0 & \text{otherwise}, \end{cases}
\]

and

\[
B(n_m; N_m) = \frac{N_m!}{n_m!(N_m - n_m)!} (1 - q_{ml})^{N_m - n_m}.
\]

Here, \( q_{ml} = P(B_l m; A, l) \approx [A B_m] [l Q_m] [A] \) is the conditional probability that a neighbor of a node in class \((A, l)\) that has degree \( m \) is in state \( B \). This probability is estimated as

\[
\frac{N(A, l \to B, m)/N(A, l \to m)}{N(A, l \to B, m) - N(A, l \to m) / [A] | N Q_m} \approx N(A, l \to m) / [A] | N Q_m
\]

from the number of nodes of degree \( m \) and density \( m \) in state \( A \). In the following, we denote the number of nodes of degree \( m \) and density \( l \) in state \( A \) by \( n_m = \sum_m B(n_m; N_m) n_m \) and \( n_m^2 = \sum_m B(n_m; N_m) n_m^2 \), respectively, obtaining

\[
S(N_1, \ldots, N_{r'}) = (1 - p) \sum_{m} S_m \sum_{l} \sum_{m} t_{ml} \delta_{l,l'}
\]

defined in Eq. (G.4). Now, inserting expression (G.3) for \( S \) in Eq. (G.1) we obtain

\[
\frac{d[A_k B_{l'}]}{dt} \bigg|_A = \sum_{l} \sum_{m} t_{ml} \left[ \left( 1 - p \right) \sum_{l'} \left( 1 - q_{l'l'} \right) q_{l'l'} \langle N_{l'} \rangle \right. \\
\left. + \sum_{m} q_{ml} \langle N_{m} \rangle \right] \delta_{l,l'}
\]

and

\[
\frac{d[A_{k'}]}{dt} \bigg|_A = \sum_{l} \sum_{m} t_{ml} \left( 1 - p \right) \left( 1 - q_{l'l'} \right) q_{l'l'} \langle N_{m} \rangle
\]

where we have used the expression for the moments \( \langle n_m \rangle = q_{ml} N_m \) and \( \langle n_m^2 \rangle = q_{ml} N_m + q_{ml}^2 N_m (N_m - 1) \) of the binomial

\[
\langle N_{l'} \rangle \equiv \sum_{l' \neq l} M(N_1, \ldots, N_{r'}; l) N_{l'} = Q_{l'-l}
\]

and

\[
\langle N_{m} \rangle \equiv \sum_{N_m} M(N_1, \ldots, N_{r'}; l) N_{m} = Q_{m}
\]

where we have used the expression for the moments \( \langle n_m \rangle = q_{ml} N_m \) and \( \langle n_m^2 \rangle = q_{ml} N_m + q_{ml}^2 N_m (N_m - 1) \) of the binomials

\[
\langle N_{l'} \rangle \equiv \sum_{l' \neq l} M(N_1, \ldots, N_{r'}; l) N_{l'} = Q_{l'-l}
\]

and

\[
\langle N_{m} \rangle \equiv \sum_{N_m} M(N_1, \ldots, N_{r'}; l) N_{m} = Q_{m}
\]
Adding the two contributions leads to we reach the final equation Eq. (10).

\[ Q_{\ell} \approx kP_{\ell}/(k), \quad [A_{\ell}B] \equiv \sum_{m=1}^{\infty} [A_{\ell}B_m], \quad \{AB\}_{\ell} \equiv \sum_{m=1}^{\infty} [A_mB_m]/l, \quad \text{and} \quad \{AB\} \equiv \sum_{m=1}^{\infty} [A_mB_m]/l. \]

Given that \([A_{\ell}B_{\ell}]\) may also change when a B-node is chosen, the evolution of \([A_{\ell}B_{\ell}]\) is given by

\[ \frac{d[A_{\ell}B_{\ell}]}{dt} = \frac{d[A_{\ell}B_{\ell}]}{dt} \bigg|_A + \frac{d[A_{\ell}B_{\ell}]}{dt} \bigg|_B. \]

The second term on the right hand side can be obtained from Eq. (G.9), by interchanging A and k by B and k, respectively. Adding the two contributions leads to

\[ \frac{d[A_{\ell}B_{\ell}]}{dt} = (1-p) \left( Q_{\ell} \frac{k-1}{k} [AB_{\ell}] + Q_{\ell} \frac{k-1}{k'} [AK_{\ell}B] \right. \]

\[ \left. - \left. \left[ \frac{[AB_{\ell}] k-1}{k^2} + \frac{[A_{\ell}B]}{[A_{\ell}k]} \right. \frac{k'-1}{k} \right] [A_{\ell}B_{\ell}] \right) \]

\[ + \left. \left( \frac{1}{k} + \frac{1}{k'} \right) [A_{\ell}B_{\ell}] \right) 

+ \left. \left( \frac{k}{k'} \left[ [A_{\ell}B_{\ell-1}] [AB_{\ell+1}] \right] + \frac{k}{k'} \left[ [A_{\ell}B_{\ell}] [A_{\ell}B_{\ell-1}] \right] \right) \right) \]

\[ \frac{d[A_{\ell}B_{\ell}]}{dt} = \beta \left( [AAB] + [BBA] - 2[ABA] - 2[ABA] \right) - 2[ABA] - 2[ABA] \]

\[ \frac{d[A_{\ell}B_{\ell}]}{dt} = \beta \left( [AAB] + [BBA] - 2[ABA] - 2[ABA] \right) - 2[ABA] - 2[ABA] \]

\[ \frac{d[A_{\ell}B_{\ell}]}{dt} = \beta \left( [AAB] + [BBA] - 2[ABA] - 2[ABA] \right) - 2[ABA] - 2[ABA] \]

\[ \frac{d[A_{\ell}B_{\ell}]}{dt} = \beta \left( [AAB] + [BBA] - 2[ABA] - 2[ABA] \right) - 2[ABA] - 2[ABA] \]

\[ \frac{d[A_{\ell}B_{\ell}]}{dt} = \beta \left( [AAB] + [BBA] - 2[ABA] - 2[ABA] \right) - 2[ABA] - 2[ABA] \]

\[ \frac{d[A_{\ell}B_{\ell}]}{dt} = \beta \left( [AAB] + [BBA] - 2[ABA] - 2[ABA] \right) - 2[ABA] - 2[ABA] \]

\[ \frac{d[A_{\ell}B_{\ell}]}{dt} = \beta \left( [AAB] + [BBA] - 2[ABA] - 2[ABA] \right) - 2[ABA] - 2[ABA] \]

\[ \frac{d[A_{\ell}B_{\ell}]}{dt} = \beta \left( [AAB] + [BBA] - 2[ABA] - 2[ABA] \right) - 2[ABA] - 2[ABA] \]

\[ \frac{d[A_{\ell}B_{\ell}]}{dt} = \beta \left( [AAB] + [BBA] - 2[ABA] - 2[ABA] \right) - 2[ABA] - 2[ABA] \]

\[ \frac{d[A_{\ell}B_{\ell}]}{dt} = \beta \left( [AAB] + [BBA] - 2[ABA] - 2[ABA] \right) - 2[ABA] - 2[ABA] \]

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\[ \frac{d[A_{\ell}B_{\ell}]}{dt} = \beta \left( [AAB] + [BBA] - 2[ABA] - 2[ABA] \right) - 2[ABA] - 2[ABA] \]

Finally, given that \([AB]_{\ell} = [A_{\ell}B_{\ell}] + [B_{\ell}A_{\ell}]\), and using the symmetry between A and B states and the fact that in a time of order unity, a quasi-stationary state is established, in which the fraction of nodes in different degree classes \([A_k]/P_k\) \((B_k)/P_k\)) reach the value corresponding to the global density \([A]) ([B]), we reach the final equation Eq. (10).
The equations look similar for reverse node update.

\[
\begin{align*}
&{\frac{d}{dt}} [1, l] = - l [1, l]
\end{align*}
\]

\[+(1-p)m[l,m] \sum_{g} P_{\langle g \rangle}(g+1) \beta_{m,l}(g)\]

\[+(p+1)[m,l+1] \sum_{g} P_{\langle g \rangle}(g+1) \alpha_{m,l}(g).\]

Where \( \alpha_{m,l}(g) = (l+m)/(l+m+g+1) \) and \( \beta_{m,l}(g) = (g+1)/(l+m+g+1) \).

The equations look similar for reverse node update.
A. Levina, J. M. Herrmann, T. Geisel, Phase transitions towards criticality in a neural system with adaptive interactions, Phys. Rev. Lett. 102 (2007) 118110.

J. Jost, K. M. Kolwankar, Evolution of network structure by temporal learning, Physica A 388 (2009) 1959–1966.

C. Meisel, T. Gross, Adaptive self-organization in a realistic neural network model, Phys. Rev. E 80 (2009) 061917.

R. Ren, K. M. Kolwankar, A. Samal, J. Jost, Stdp-driven networks and the c. elegans neuronal network, Physica A 389 (2010) 3900–3914.

C. Meisel, A. Storch, S. Hallmeyer-Eigner, E. Bullmore, T. Gross, Failure of adaptive self-organized criticality due to epileptic seizure attacks, PLoS Comput. Biol. 8(1) (2012) e1002312.

F. Droste, A.-L. Do, T. Gross, Analytical investigation of self-organized criticality in neural networks, J. R. Soc. Interface (2012).

C. Haupe, G. Zschaler, A.-L. Do, T. Gross, Adaptive-network models of swarm dynamics, New J. Phys. 13 (2011) 073022.

I. D. Couzin, C. C. Ioannou, G. Demirel, T. Gross, C. J. Torney, A. Hartnett, L. Conradt, S. A. Levin, N. E. Leonard, Uninformed individuals promote democratic consensus in animal groups, Science 334 (2011) 1578–1580.

T. P. Peixoto, S.Bornholdt, No need for conspiracy: self-organized cartel formation in a modified trust game, Phys. Rev. Lett. 108 (2012) 218702.

S. W. Kim, J. D. Noh, Instability in a network coevolving with a particle system, Phys. Rev. Lett. 100 (2008) 118702.

K. Tomita, H. Kurokawa, S. Muarata, Graph rewiring automata as a natural extension of cellular automata, in: T. Gross, H. Sayama (Eds.), Adaptive Networks. Theory, Models and Applications, Springer Verlag, New York, 2009, pp. 291–310.

F. M. Atay, J. Jost, Delays, connection topology, and synchronization of coupled chaotic maps, Phys. Rev. Lett. 92 (2004) 144101.

A.-L. Do, S. Boccaletti, T. Gross, Graphical notation reveals topological stability criteria for collective dynamics in complex networks, Phys. Rev. Lett. 108 (2012) 194102.

H. A. Bethe, Statistical theory of superlattices, Proc. R. Soc. B 150 (1935) 552–575.

R. Pastor-Satorras, C. Castellano, A. Vespignani, Incomplete ordering of the voter model dynamics on annealed complex heterogeneous networks, Europhys. Lett. 88 (2009) 58004–58006.

S. Jolad, W. Liu, B. Schmittmann, R. K. P. Zia, Epidemic spreading on preferred degree adaptive networks, 2011. ArXiv:1109.5440.

J. P. Gleeson, S. Melnik, J. Ward, M. A. Porter, P. J. Mucha, Accuracy of mean-field theory for dynamics on real-world networks, Phys. Rev. E 85 (2012) 026106.

J. P. Gleeson, S. Melnik, J. Ward, M. A. Porter, P. J. Mucha, Accuracy of mean-field theory for dynamics on real-world networks, Phys. Rev. E 85 (2012) 026106.

J. Yedidia, W. Freeman, Y. Weiss, Constructing free energy approximations for disparate systems and the voter model, Ann. Probab. 3 (1975) 643–663.
imations and generalized belief propagation algorithms, IEEE Trans. Inform. Theory 51 (2005) 2282–2312.
[108] R. Law, D. J. Murrell, U. Dieckmann, Population growth in space and time: spatial logistic equations, Ecology 84 (2003) 252–262.
[109] R. Kikuchi, A theory of cooperative phenomena, Phys. Rev. 81 (1951) 908–1003.
[110] I. G. Kevrekidis, C. W. Gear, G. Hummer, Equation-free: The computer-aided analysis of complex multiscale systems, AIChE Journal 50 (2004) 1346–1355.
[111] T. Rogers, Maximum-entropy moment-closure for stochastic systems on networks, J. Stat. Mech. (2011) P05007.
[112] J. Lindquist, J. Ma, P. van den Driessche, F. H. Willeboordse, Effective degree network disease models, J. Math. Biol. 62 (2011) 143–164.
[113] M. Taylor, T. J. Taylor, I. Z. Kiss, Epidemic threshold and control in a dynamic network, Phys. Rev. E 85 (2012) 016103.
[114] F. Vazquez, J. C. González-Avella, V. M. Eguiluz, M. S. Miguel, Time scale competition leading to fragmentation and recombination transition in the coevolution of network and states, Phys. Rev. E 76 (2007) 046120.
[115] B. Kozma, A. Barrat, Consensus formation on adaptive networks, Phys. Rev. E 77 (2008) 016102.
[116] B. Kozma, A. Barrat, Consensus formation on coevolving networks: groups’ formation and structure, J. Phys. A: Math. Theor. 41 (2008) 224020.
[117] G. Iñiguez, J. Kertész, K. K. Kaski, R. A. Barrio, Opinion and community formation in coevolving networks, Phys. Rev. E 80 (2009) 066119.
[118] J. Bryden, S. Funk, N. Geard, S. Bullock, V. A. A. Jansen, Stability in flux: community structure in dynamic networks, J. R. Soc. Interface 8 (2011) 1031–1040.
[119] P. Holme, G. Ghoshal, Dynamics of networking agents competing for high centrality and low degree, Phys. Rev. Lett. 96 (2006) 098701.
[120] M. G. Zimmermann, V. M. Eguiluz, Cooperation, social networks and the emergence of leadership in a prisoners dilemma with adaptive local interactions, Phys. Rev. E 72 (2005) 056118.
[121] K. T. D. Eames, M. J. Keeling, Modeling dynamic and network heterogeneities in the spread of sexually transmitted diseases, Proc. Natl. Acad. Sci. USA 99 (2002) 13330–13335.
[122] S. Bansal, B. T. Grenfell, L. A. Meyers, When individual behaviour matters: homogeneous and network models in epidemiology, J. R. Soc. Interface 4 (2007) 879–891.
[123] E. Volz, L. A. Meyers, Susceptibleinfectedrecovered epidemics in dynamic contact networks, Proc. R. Soc. B 274 (2007) 2925–2934.
[124] E. Volz, L. A. Meyers, Epidemic thresholds in dynamic contact networks, J. R. Soc. Interface 6 (2009) 233–241.