Evolving Networks through Deletion and Duplication

Nadia Farid∗ and Kim Christensen†

Blackett Laboratory, Imperial College London,
Prince Consort Road, London SW7 2BW, United Kingdom

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We introduce a minimalistic model based on dynamic node deletion and node duplication with heterodimerisation. The model is intended to capture the essential features of the evolution of protein interaction networks. We derive an exact two-step rate equation to describe the evolution of the degree distribution. We present results for the case of a fixed-size network. The results are based on the exact numerical solution to the rate equation which are consistent with Monte Carlo simulations of the model’s dynamics. Power-law degree distributions with apparent exponents < 1 were observed for generic parameter choices. However, a proper finite-size scaling analysis revealed that the actual critical exponent in such cases is equal to 1. We present a mean-field argument to determine the asymptotic value of the average degree, illustrating the existence of an attractive fixed point, and corroborate this result with numerical simulations of the first moment of the degree distribution as described by the two-step rate equation. Using the above results, we show that the apparent exponent is determined by the heterodimerisation probability. Our preliminary results are consistent with empirical data for a wide range of organisms, and we believe that through implementing some of the suggested modifications, the model could be well-suited to other types of biological and non-biological networks.

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I. INTRODUCTION

In recent years, models of networks growing via single-node duplication, divergence and mutation of links, considered in isolation and combination, have assumed prominence in the literature on complex networks. In a series of independent studies it was suggested that these duplication-divergence-mutation models (hereafter called ‘duplication models’ for brevity) are good candidates to describe the evolution and large-scale topological features of real protein-protein interaction networks (PINs) in several organisms such as S.Cerevisiae and H.Pylori [1–6].

In duplication models, proteins are represented by nodes, and a pairwise interaction between any two proteins is represented by an undirected link between the associated nodes, assumed to be fully operative at all times. In a duplication event, a mother node is chosen uniformly at random (u.a.r.) and each of its links are copied to a newly created daughter node. Divergence refers to the subsequent loss of links from the daughter node [2, 4–7], and/or the mother node [8], or a shared neighbour of both the daughter and mother node [3]. Often, for simplicity, it is assumed that only the daughter node diverges. In a mutation event, new links are added between the daughter node and all other nodes in the network which are not already connected to the mother node. Typically, one duplication-divergence event occurs at each update step, and mutations, when considered, are modelled at a rate much less than the divergence rate – typically, one new link per update step is added [4–6].

The idea of evolution through gene duplication is taken from biology [9–12], where it was popularised in the 1970’s by Ohno who conjectured that single and whole genome duplications could provide the raw material for evolutionary diversification [13]. Whilst there is mounting evidence that duplicate genes do occur in genomes [1, 12, 14–17], it is widely acknowledged that little is known about the details of the process of duplication itself, such as the frequency of duplication events, the fate of duplicate genes, and the frequency with which duplicate genes acquire novel functions [1, 9, 11, 18]. In fact, whilst the microscopic parameters are not yet known exactly, it has been suggested by Berg et al. [19] that the rate of gain and loss of interactions through mutations is at least an order of magnitude higher than the growth rate due to duplications; as a result, the link dynamics act as the dominant evolutionary force shaping the large-scale statistical structure of the network, and not the gene duplications; the slower gene duplication events only really affect the size of the network. Thus, it is still unclear if, and to what extent, gene duplication is the dominant mechanism responsible for the observed statistical features of PINs. Moreover, gene deletion and rearrangement are
known to play important roles in the long-term evolution of genomes [10, 17, 20] but have received comparatively little attention in the literature on networks addressing PINs [20, 21].

In this paper, in a move to go beyond the duplication models and to expand upon the emerging literature on network models of PINs including gene deletion, we present a four-parameter model addressing the scenario of network evolution through dynamic total node removal in conjunction with growth by node duplication and heterodimerisation. We refer to our network model as a deletion-duplication-divergence-heterodimerisation (DDDHD) model [22]. Although the model is primarily aimed at describing the evolution of PINs, its description is kept general enough so as to be applicable to a diverse range of complex systems where components are added and removed throughout the system’s evolution.

In the wider literature on complex networks, previous studies which have considered node deletion have generally regarded it as a perturbation effect, used to test the tolerance of a network to random and targeted attack [3, 25–27]. More recently, the mechanism of dynamic node removal in conjunction with growth by preferential attachment has been explored in independent studies by Chung and Lu [28], Cooper and Frieze [29], and Wang [30]. They refer to such models as growth-deletion models. Since we consider growth by duplication as opposed to growth by preferential attachment, our model therefore also contributes to the literature on growth-deletion models.

We focus our analysis on the degree distribution, \(P(k)\), characterising the probability for a node to have exactly \(k\) links [31–33]. The degree distribution is the simplest topological feature to measure and as a result, it has attracted and received the most attention in the literature on complex networks. It has been shown [2–4, 6–8] that the degree distribution of networks generated by duplication models exhibits a power-law tail, \(P(k) \sim k^{-\gamma}\), for \(k \gg 1\), where the critical exponent \(\gamma\) can be tuned such that it is in agreement with observed exponents which are found to be in the range \(1 < \gamma < 3\) [2, 34]. Importantly, it has also been shown that the degree distribution is a robust and generic property of PINs common across different data sets – an important consideration given that current experimental techniques are notorious for suffering from a high rate of false positives and false negatives [19, 34].

The paper is organised as follows. In Sec. II, we present the formulation of the dynamics which describe the rules of evolution of the network, defining the parameters and interpreting the rates. In Sec. III, we present and discuss the exact two-step rate equation for the evolution of the degree distribution. In Sec. IV, we present results obtained from Monte Carlo (MC) simulations of the model and the exact numerical solution of the rate equation for a generic choice of parameter values. We discuss our results in Sec. V and end with a conclusion in Sec. VI.

II. DDDHD MODEL

We consider undirected networks where loops and multiple links are forbidden. We start with a network of known size, \(N\), and degree distribution, \(P(k, t = 0)\), and allow it to evolve under the following rules (see Fig. 1):

![FIG. 1: (Colour online) A schematic representation of a network evolving through deletion and duplication. (i) A node is chosen u.a.r. and deleted with probability \(p_{\text{del}}\) (grey). (ii) A mother node is chosen u.a.r. and duplicated with probability \(p_{\text{dupl}}\) (green \(\rightarrow\) red). The links are retained with probability \(p\) (dashed red line), and a further link is established between the daughter and mother node with probability \(\theta\) (dotted blue line).](image)

1. Deletion. With probability \(p_{\text{del}}\), a node is chosen u.a.r. This node and all of its links are deleted from the network.

2. Duplication-Divergence-Heterodimerisation. With probability \(p_{\text{dupl}}\), a mother node is chosen u.a.r. and duplicated. This entails a new daughter node being added to the network and linking to each of the neighbours of the mother node with probability \(p\). A further link is established between the daughter and mother node with heterodimerisation probability \(\theta\).

The evolution of the network is thus governed by four parameters: \(p_{\text{del}}, p_{\text{dupl}}, p, \theta\). In Sec. III, we cast these rules into a concrete mathematical framework. First, however, it is instructive to discuss the motivation behind the choice of dynamics and its implications.

The mechanism of growth by duplication is preferred to the mechanism of growth by preferential attachment in PIN network models as well as many other network models since it reproduces the effects of preferential attachment...
without having to artificially put the mechanism in. In other words, it arises naturally from the dynamics: nodes that have a large number of links are more likely to be neighbours of a duplicating node, and hence are more likely to gain a link to the newly created node. The DDDH model preserves this effect and introduces another one: implicit preferential detachment. Nodes that have a large number of links are more likely to lose a link each time a node is deleted. It is interesting to note that these two effects do not cancel out each other, as one is deleted. It is interesting to note that these might intuitively expect. The inclusion of heterodimerisation, \( \theta > 0 \), means that we do not consider mutations in the traditional sense (as described above), but rather we restrict the addition of new links to only occur between the mother and the daughter node. Heterodimerisation is preferred over mutations as it has been noted that the former increases the likelihood of clique-formation – a feature observed in real PINs – whilst the latter, in order to form the observed number of triads and higher cliques, would require a prohibitively/physically unrealistic high rate \( p \). Moreover, in duplication models, whilst a lack of random linking (mutations) has been found to destroy fine structure such as the self-averaging and existence of a smooth degree distribution \( p \), the large-scale statistical features of the final network do not depend on the existence of mutations [3–5].

From the general definition of the model's dynamics given above, one can already determine specific features of the network, namely its overall size, by focusing on particular choices of the model's parameter values. For example, for a network that remains, on average, fixed in size, \( p_{\text{del}} = p_{\text{dupl}} \). We will refer to this case as a 'fixed-size' network. For a network that is monotonically growing, on average, \( p_{\text{dupl}} > p_{\text{del}} \); if \( p_{\text{del}} = 0 \) no nodes are removed from the network. Moreover, if we fix \( \theta = 0 \) and \( 0 < p < 1 \) and \( p_{\text{dupl}} = 1 \) for this case, the DDDH model is equivalent to the duplication-divergence model [5, 6, 8, 35]. For \( p_{\text{del}} = 0, p_{\text{dupl}} = 1, \theta = 0 \) and \( p = 1 \) the duplication-divergence model is equivalent to Polya's Urn [2, 8, 36]. For an on average monotonically shrinking network, \( p_{\text{dupl}} < p_{\text{del}} \); if \( p_{\text{dupl}} = 0 \) no new nodes or links are added to the network [37]. For a network fluctuating in size the values of \( p_{\text{del}} \) and \( p_{\text{dupl}} \) would be stochastic variables chosen anew at each update step from a suitably chosen distribution [38]. If \( p = 1 \), the daughter node inherits all of the links; this is the case of perfect, or full duplication. If \( 0 < p < 1 \), the daughter node inherits only some of the links: this represents imperfect, or partial duplication. If \( p = 0 \), the daughter node inherits none of the links, that is, an isolated node is added to the network.

In this paper, we study the case of fixed-size networks, \( p_{\text{del}} = p_{\text{dupl}} = 1 \), with heterodimerisation \( \theta > 0 \) and perfect duplication \( p = 1 \), unless otherwise stated (in which case, similar to Refs. [2, 4, 6] we assume that only the daughter node diverges) [40].

In Sec. III, we apply the rate equation approach [4, 41] to study the evolution of the expected number of nodes with \( k \) links at time \( t \), \( f(k, t) = N_t P(k, t) \) where \( N_t \) is the total number of nodes at time \( t \).

### III. DDDH MODEL: RATE EQUATION

In this section, we present a two-step rate equation for the general DDDH model, and then explain in detail the origin of each term.

The two-step rate equation for the DDDH model is given by,

\[
f(k, t+1) = f(k, t) - p_{\text{del}} \frac{f(k, t)}{N_t} - p_{\text{del}} \frac{k f(k, t)}{N_t} + p_{\text{del}} \frac{(k+1) f(k+1, t)}{N_t}, \tag{1a}
\]

\[
f(k, t+2) = f(k, t+1) - p_{\text{dupl}} p \frac{k f(k, t+1)}{N_{t+1}} - p_{\text{dupl}} \theta \frac{f(k, t+1)}{N_{t+1}}
+ p_{\text{dupl}} \left( \frac{(k-1) f(k-1, t+1)}{N_{t+1}} + \frac{f(k-1, t+1)}{N_{t+1}} \right)
+ p_{\text{dupl}} \theta \sum_{j \geq (k-1)} \binom{j}{k-1} p^{(k-1)}(1-p)^{(k-1)} \frac{f(j, t+1)}{N_{t+1}}
+ p_{\text{dupl}}(1-\theta) \sum_{j \geq k} \binom{j}{k} p^{k}(1-p)^{j-k} \frac{f(j, t+1)}{N_{t+1}}. \tag{1b}
\]
Equation (1) is **exact** and there are no approximations in its derivation. It describes the DDDH process for all parameters, \( p_{\text{del}}, p_{\text{dupl}}, p, \theta \). It holds for all \( k \geq 0 \), with \( f(-1, t) = 0 \) and \( f(k > k_{\text{max}}, t) = 0 \) for all \( t \). Moreover, \( f(k, t) \) satisfies the following normalisation conditions:

\[
\begin{align*}
\sum_{k=0}^{\infty} f(k, t) &= N_t, \quad (2a) \\
\sum_{k=0}^{\infty} k f(k, t) &= 2L_t, \quad (2b)
\end{align*}
\]

where \( L_t \) is the total number of links at time \( t \) in the network.

The distinct feature of the DDDH rate equation compared to rate equations for duplication models is that it is defined in two steps, and not one, reflecting the fact that we now include a node deletion in addition to node duplication. To keep the notation simple, we have written \( t \), to keep notation simple, we have written \( t \) instead of \( t+\frac{1}{2} \) and \( t+1 \) to indicate that we only observe the network after both the deletion and the duplication steps have been completed.

Moreover, each of these actions is executed sequentially, highlighting the fact that we clearly also make the distinction between this process and the process of adding nodes at an ‘effective’ duplication rate, \( p_{\text{dupl}} - p_{\text{del}} \), or merging nodes as is Refs. [23, 24], or substituting nodes where the duplication of a node automatically implies the deletion of some other node as in Ref. [20]. We note that the exact correspondence between time, \( t \), and real biological timescales is unclear, however, at this stage.

In Eq. (1a) we consider the effects on the network of the deletion of a node and the removal of its links. The probability a deletion event occurs is given by \( p_{\text{del}} \). The terms on the right-hand side (RHS) are interpreted as follows. A **loss** in the number of nodes with degree \( k \) at time \( t+1 \) from deletion will occur either if the node deleted is of degree \( k \) at time \( t \), or if a neighbour (of arbitrary degree) of a \( k \)-node is chosen for deletion, as the \( k \)-node will lose a link and become a node of degree \( k-1 \). Since every node has an equal probability of being deleted in a given time step, a node of degree \( k \) is chosen with probability \( f(k, t)/N_t \) (second term); the probability that a neighbour of a \( k \)-node is chosen for deletion is \( kf(k, t)/N_t \) (third term). The final term on the RHS represents a **gain** in the number of nodes with degree \( k \) at time \( t \). This can occur if the neighbour of a node with degree \( k+1 \) is deleted. Given that a node of degree \( k+1 \) has \( k+1 \) neighbours, the probability a neighbour is chosen for deletion is \((k+1)f(k+1, t)/N_t\).

In Eq. (1b) we consider the effects of node duplication and subsequent heterodimerisation. The probability a duplication event occurs is given by \( p_{\text{dupl}} \), and the probability that subsequent heterodimerisation occurs is given by \( \theta \). All but the two final terms on the RHS represent changes a duplication-heterodimerisation event has on the existing nodes in the network; the last two terms represent the daughter node’s contribution.

A **loss** in the number of nodes with degree \( k \) at time \( t+2 \) from duplication and heterodimerisation can occur in one of two ways. If one of the neighbours (of arbitrary degree) of a node of degree \( k \) at time \( t + 1 \) is duplicated, the \( k \)-node will, with probability \( p \), gain a link from duplication thus becoming a \( k+1 \) node at time \( t+2 \) (second term). Alternatively, if the mother node to be duplicated is already of degree \( k \) at time \( t + 1 \), it will become a node with degree \( k+1 \) at time \( t+2 \) by gaining a link to the daughter node via heterodimerisation. The probability a node of degree \( k \) is chosen for duplication is given by \( f(k, t+1)/N_{t+1} \), and the probability of heterodimerisation is \( \theta \) (third term).

We arrive at the fourth and fifth terms which describe the **gain** in the number of nodes with degree \( k \) at time \( t+2 \) by similar considerations. If one of the neighbours of a node of degree \( k-1 \) at time \( t + 1 \) is chosen for duplication, the \( k-1 \) node will, with probability \( p \), gain a link from duplication, thus becoming a \( k \) node at time \( t+2 \) (fourth term). Alternatively, if the mother node to be duplicated is of degree \( k-1 \) at time \( t + 1 \), it will become a node with degree \( k \) at time \( t+2 \) by gaining a link to the daughter node via heterodimerisation (fifth term).

The final two terms on the RHS of Eq. (1b) account for the daughter node’s contribution. The sixth term is to account for the contribution of the daughter node in the event it does establish a link, with probability \( \theta \), to the mother node. The seventh term is to account for the case where it does not, which happens with probability \((1 - \theta)\). Note the lower limits on these sums are not identical. This is because if a link is established via heterodimerisation, in order to become a node of degree \( k \), the daughter node is restricted to copying \( k-1 \) out of \( j \) links, each with probability \( p \). However, if such a link is not established, the daughter node is restricted to copying \( k \) out of \( j \) links of the mother node.

Since the exact analytical solution to the rate equation in Eq. (1) is not tractable at present,
in Sec. IV we present results obtained from the numerical solution of the exact two-step rate equation and compare them with MC simulations of the model. Unless otherwise stated, our analysis is based on the case of a fixed-size network, \( p_{\text{del}} = p_{\text{dupl}} = 1 \), evolving through perfect duplication, \( p = 1 \), with heterodimerisation, \( \theta > 0 \).

IV. DDDH MODEL: RESULTS

1. Comparison between exact numerical solution of the rate equation and MC simulations

Figure 2 displays the evolution of the degree distribution obtained from the exact numerical solution of the rate equation compared to MC simulations of the model. The stationary regime is defined by \( P(k, t) = P(k, t + 2) \).

We start with an initial network of \( N = 400 \) nodes, with a random degree distribution centred around \( k_{\text{init}} = 100 \), and iterate the rules with the following parameter settings: \( p_{\text{del}} = p_{\text{dupl}} = 1 \) (fixed-size network), \( p = 1 \) (perfect duplication), with heterodimerisation \( \theta = 0.1 \). The value for \( \theta \) was chosen as such as it is believed that heterodimerisation occurs at a rate not greater than 0.1 [8]. In Fig. 2, we show two snapshots of the network in the transient regime when \( t = 1000, 5000 \), respectively, and one in the stationary regime for \( t = 10^6 \). The MC simulations are averaged over \( 10^5 \) realisations for \( t = 1000, 5000 \) and \( 3 \times 10^5 \) realisations for \( t = 10^6 \).

There is excellent agreement between the exact numerical solution and MC results, lending support to our statement earlier that there are no approximations involved in our derivation of the rate equation. We have verified through extensive simulations that the agreement holds true over a range of \( p_{\text{del}}, p_{\text{dupl}}, p \), and \( \theta \) values. Hence, all remaining figures are generated using data obtained from the exact numerical solution of the rate equation only, hereafter referred to as ‘exact numerical results’. The interesting feature to note is that even for the simplest realisation of the DDDH model which we have presented in Fig. 2, fat-tailed degree distributions are obtained in the stationary regime (\( t = 10^6 \) curve). This is in stark contrast to the duplication models where only the case of imperfect duplication leads to a power law [2]. In the following section, we describe quantitatively the exact form of the stationary degree distribution.

FIG. 2: Exact numerical (solid line) and MC (open circles) results of the degree distribution of a fixed-size network, \( p_{\text{del}} = p_{\text{dupl}} = 1 \), with \( N = 400 \) nodes, evolving under perfect duplication, \( p = 1 \), with heterodimerisation \( \theta = 0.1 \). Snapshots were taken at times \( t = 1000, 5000 \) (transient regime) and \( t = 10^6 \) (stationary regime). The MC simulations are averaged over \( 10^5 \) realisations for \( t = 1000, 5000 \), and \( 3 \times 10^5 \) realisations for \( t = 10^6 \). The exact numerical results show excellent agreement with the MC simulations. The distribution in the stationary regime is well approximated by a power-law decay, \( P(k) \sim k^{-\gamma} \) with an apparent exponent \( \gamma = 0.8 \) and sharp cut-off at \( k_{\text{cut-off}} = 399 \). Note that the maximum degree a node can attain in networks of the type we consider is \( N - 1 \).

2. Scaling for \( p_{\text{del}} = p_{\text{dupl}} = p = 1, 0 < \theta \leq 1/2 \)

We are interested in quantifying the form of the degree distribution, in the stationary regime, as a function of the model’s parameters. Given that we are, for the moment, investigating fixed-size networks evolving under perfect duplication, \( p_{\text{del}}, p_{\text{dupl}} \) and \( p \) are all fixed. This reduces the number of variables to just one: \( \theta \). However, given that we are observing the degree distribution for specific fixed network sizes, we have \( N \) as another variable in the problem. Hence, we would like to know how \( P(k) \) depends on \( \theta \) and \( N \). In order to investigate this, we have performed numerical simulations for the following two cases: (i) Fixed \( \theta \), varying \( N \), and (ii) Fixed \( N \), varying \( \theta \). We discuss (i) in this subsection, and (ii) in Sec. IV. 3.

Figure 3(a) shows the exact numerical results of the degree distribution, \( P(k; N) \) versus \( k \) on a double logarithmic plot in the stationary regime for \( \theta = 0.1 \) and networks of increasing \( N \), specifically, \( N = 50, 100, 200, 400, 1000 \) nodes. There are clear power-law fluctuations in the node degrees present in the network, im-
We can say that \( k_{\text{max}} = k_{\text{local}} + k_{\text{swap}} \), where \( k_{\text{local}} \) is the number of nodes a node can contact within its local (and hence, unseen) region, and \( k_{\text{swap}} \) is the number of nodes a node can contact via a swap operation, which is equivalent to the maximum possible degree a node in the network can acquire. This implies that \( k_{\text{max}} \) increases linearly with increasing network size, \( N \), hence, in the limit of \( N \rightarrow \infty \), the characteristic scale diverges and a pure power law is recovered, as expected. In the region, \( 1 \leq k \ll k_{\text{max}} \), we find that the gradient of the lines in Fig. 3(a) are well-approximated by an ‘apparent’ exponent, \( \gamma = 0.8 \) and we will shortly demonstrate that \( \gamma = 1 - 2\theta \). Generally speaking, an exponent less than 1 is unusual, and seems to contradict certain known results about scaling functions. However, we will be able to resolve this apparent contradiction in Sec. IV. 3.

With the above discussion in mind, we propose the following general ansatz for \( P(k;N) \),

\[
P(k;N) \propto \begin{cases} 
  k^{-\gamma} & 1 \leq k \ll k_{\text{max}} \\
  \text{sharp cut-off} & k \gg k_{\text{max}}.
\end{cases}
\]  

(3)

From simulations, we find that \( k_{\text{max}} = N - 1 \) which is equivalent to the maximum possible degree that a node in the network can acquire. This implies that \( k_{\text{max}} \) increases linearly with increasing network size, \( N \), hence, in the limit of \( N \rightarrow \infty \), the characteristic scale diverges and a pure power law is recovered, as expected. In the region, \( 1 \leq k \ll k_{\text{max}} \), we find that the gradient of the lines in Fig. 3(a) are well-approximated by an ‘apparent’ exponent, \( \gamma = 0.8 \) and we will shortly demonstrate that \( \gamma = 1 - 2\theta \). Generally speaking, an exponent less than 1 is unusual, and seems to contradict certain known results about scaling functions. However, we will be able to resolve this apparent contradiction in Sec. IV. 3.

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\end{cases}
\]  

(4)

and, assuming for simplicity that \( k_{\text{max}} \) is approximated by \( N \), the equation is valid for \( 1 \leq k \leq N \), where \( N \gg 1 \) and \( \gamma < 1 \). In Eq. (4), \( a(N) \) is a prefactor, dependent on the network size, and \( G(x) \) is the cut-off function, dependent on the rescaled variable \( k/N \). \( G(x) \) is required to fall-off fast enough to ensure \( P(k;N) \) is finite and integrable. From Fig. 3(a) it seems reasonable to assume that \( G(x) \) is constant for \( x \ll 1 \) and decays abruptly for \( x \gg 1 \).

Note that Eq. (4) is not a finite-size scaling (FSS) ansatz since the prefactor, rather than being a constant (as is typical), is \( N \)-dependent. This difference turns out to be important as it leads to an interesting result about the critical exponent which we will demonstrate below.

In theory, we can use the fact that the probability density function must be properly normalised,

\[
\int_{1}^{\infty} P(k;N)dk = 1,
\]  

(5)

to derive an expression for \( a(N) \). However, without knowing \textit{a priori} the correct scaling for \( P(k;N) \), we can only guess at the form of \( G(x) \).

For simplicity, we assume that the cut-off function, \( G(x) \), is of the form,

\[
G(x) = \begin{cases} 
  1 - k/N & 1 \leq k \leq N \\
  0 & \text{otherwise}
\end{cases}
\]  

(6)

(which is in-keeping with our stated requirements for the form of \( G(x) \) and is an excellent
Therefore, we expect the curves to collapse onto the curve
\[ \tilde{G}(k/N) = \left(\frac{1}{\Gamma(1-\gamma)\Gamma(1+\gamma)}\right) (k/N)^{1-\gamma}(1-k/N)^{\gamma}, \]
with the gradient of the slope to be equal to \( 1-\gamma \). As shown in Fig. 3(b), a convincing data collapse is obtained, with all curves collapsing onto the scaling function described by \( \tilde{G}(x) \) with \( \gamma = 1 - 2\theta \). We have repeated the data collapse for different values of \( \theta \), and observed a convincing data collapse in all cases, with all curves collapsing onto the scaling function described by \( \tilde{G}(x) \) with \( \gamma = 1 - 2\theta \).

Together, Eq. (4) and the success of the data collapse in Fig. 3(b) demonstrate that the degree distribution for any network size, \( N \), is determined by the scaling function \( \tilde{G}(x) \). This means that we can deduce the degree distribution for any network size, \( N \), without having to actually perform the numerical simulation itself. Hence, our results are applicable to networks larger than those which we have demonstrated directly, that is for \( N > 10^3 \). This is in contrast to the duplication models considered in [8] where the networks generated do not attain power-law degree distributions even for very large networks.

Thus far, we have not yet justified the relation given between the apparent exponent and the parameter \( \theta \). In the following section, we derive a result for the average degree, \( \langle k \rangle \). We then demonstrate how we can use this result to find an expression for the apparent exponent \( \gamma \) in terms of the parameter, \( \theta \).

### 3. Mean-field Equation for the Average Degree

The average degree, \( \langle k \rangle \), can be determined in various different ways. Ideally, one would be able to calculate it directly from the two-step rate equation for the evolution of the degree distribution in Eq. (1). This would be achieved by taking the first moment of the normalised degree distribution, \( P(k; t) = f(k, t)/N_t \), according to,

\[ \langle k \rangle_t = \int_1^\infty k P(k, t) dk. \]

Strictly speaking, where we write integration signs we should have sums, as we are dealing with a discrete probability distribution. Either way, the solution we are interested in is

\[ \lim_{t \to -\infty} \langle k \rangle_t = \langle k \rangle, \]

which is analytically intractable. In order to get around this problem we have calculated this quantity numerically, and compared this result to the asymptotic value of \( \langle k \rangle \) obtained as a solution to a mean-field rate equation approach (described below).

Figure 4 illustrates the exact numerical results for the time-evolution of \( \langle k \rangle \) for \( \theta \in [0.01, 0.5] \) and clearly illustrates the existence of a stationary asymptotic \( \langle k \rangle \). We now describe our mean-field argument to determine the asymptotic value of the average degree, \( \langle k \rangle \).

At each time step, for each node we delete we lose on average, \( \langle k \rangle_t \) links, and for each node...
Thus, both the mean-field calculation and the exact numerical results for the first moment of the degree distribution. As shown in Fig. 4, there is exact agreement between the asymptotic value of the average degree determined from the mean-field calculation and the exact numerical results (for $t > 10^3$). Thus, both the mean-field calculation and the exact numerical results, as illustrated in Fig. 4, demonstrate the existence of an attractive fixed point in the average degree, $\langle k \rangle$. This is clearly related to the existence of a stationary degree distribution, that is, $P(k, t) = P(k, t + 2)$.

Equation (13) highlights the importance of accounting for the mechanism of heterodimerisation for finite-sized networks, as it shows that the network self-organises to a stationary state where the average degree $\langle k \rangle$ is constant, and determined by the system size, $N$ and the probability for heterodimerisation, $\theta$. Since real PINs are of finite size, typically with no more than $10^4$ nodes (see Table I), the point of information regarding the role of $\theta$ in finite-sized networks is of significance.

We have also verified through further simulations varying $p$ such that $p < 1$ for constant $\theta$ and $N$ clearly dramatically reduces $\langle k \rangle$, as one would expect, although the precise nature of this effect has not yet been quantified and seems to be non-trivial. Thus, Eq. (13) actually gives an upper bound for $\langle k \rangle$.

An alternative method for calculating $\langle k \rangle$ analytically, is to calculate the first moment of the degree distribution as expressed in Eq. (10), $\langle k \rangle_{SF}$. This turns out to be very useful as far as determining an expression for the apparent exponent, $\gamma$, in terms of $\theta$. We find that,

$$\langle k \rangle_{SF} = \int_{1}^{N} k \, dP(k; N)$$

$$= \int_{1}^{N} \frac{1}{\Gamma(1-\gamma)\Gamma(1+\gamma)} \left( \frac{k}{N} \right)^{2-\gamma} \left( 1 - \frac{k}{N} \right)^{\gamma} \, dk$$

$$= \frac{N \Gamma(2-\gamma)}{2 \Gamma(1-\gamma)} \text{ for } N \to \infty$$

$$= \frac{N}{2(1-\gamma)}. \quad (14)$$

Since we already know that $\langle k \rangle_{MF} = \theta(N-1)$ from Eq. (13), the RHS of Eq. (14) must be equivalent to Eq. (13), hence,

$$\gamma = 1 - \theta. \quad (15)$$

This justifies our previous finding in Sec. IV. 2 that the apparent exponent is $\gamma = 1 - 2\theta$. In the following section, we investigate the effect on the degree distribution of varying $\theta \in (0, 0.5)$ for fixed $N$, completing the analysis of the two scenarios outlined at the beginning of Sec. IV. 2.

4. Effects of varying $\theta$

Figure 5 illustrates the topological effect of varying the probability to heterodimerise in the range $0 < \theta \leq 0.5$, in a fixed-size network ($N = 200$), evolving through perfect duplication.

We find that the degree distribution exhibits a power law with a slope that varies with $\theta$ according to $\gamma = 1 - 2\theta$, reaching a uniform distribution at a value of $\theta = 0.5$. We have repeated the above simulations for a range of network sizes, up to $N = 1000$, and confirmed this behaviour. Hence, the result for $\gamma$ is consistent with our previous findings in Sec. IV. 2.

Current estimates from empirical data for yeast, fly and human PINs indicate that $\theta$ never
for realistic values of the duplication model considered in Ref. [8], 200 nodes, and $k_{\text{initial}} = 50$. The network evolved through perfect duplication and increasing $\theta = 0.01, 0.025, 0.05, 0.07, 0.1, 0.3, 0.5$ (marked with lines of decreasing dash-length). The value $\theta = 0.5$ marks a change in behaviour from a power-law decay with a negative exponent to a uniform distribution.

FIG. 5: Exact numerical results for the degree distribution in the stationary regime for a fixed-size network, $p_{\text{del}} = p_{\text{dupl}} = 1$, with $N = 200$ nodes, and $k_{\text{initial}} = 50$. The network evolved through perfect duplication and increasing $\theta = 0.01, 0.025, 0.05, 0.07, 0.1, 0.3, 0.5$ (marked with lines of decreasing dash-length). The value $\theta = 0.5$ marks a change in behaviour from a power-law decay with a negative exponent to a uniform distribution.

exceeds 0.1 [8]. Since we observe power-law degree distributions in the range $0 < \theta < 0.5$, a value of $\theta < 0.1$ in our model is consistent with empirical data. The fact that $\gamma = 1 - 2\theta$ might go some way towards explaining why in the duplication model considered in Ref. [8], for realistic values of $\theta < 0.1$ their results were not affected.

5. Effects of varying $p$

Up until now, we have been investigating the effects of varying $\theta$ and $N$, for fixed $p_{\text{del}} = p_{\text{dupl}} = p = 1$, on the degree distribution. We now report our findings for a third possible scenario: the effect of varying $p$, for fixed $\theta$ and $N$ (keeping $p_{\text{del}} = p_{\text{dupl}} = 1$, as before).

Figure 6 illustrates the topological effect of varying $p$ in a fixed-size network with heterodimerisation $\theta = 0.1$. There is clearly a marked difference between the curve for $p = 1$ and the family of curves for $p < 1$. We established in Sec. IV. 2-4 that for $p = 1$ and $\theta \in (0, 0.5)$, the degree distribution is well approximated by a power-law decay. We now see that for $p < 1$, the power-law behaviour is no longer observed and a characteristic degree is present. We have confirmed this result for a range of network sizes, specifically, $N = 50, 100, 400, 1000$. Moreover, we have found that the second moment $\langle k^2 \rangle$ does not diverge with increasing network size as one would expect if, in the limit of $N \to \infty$, the degree distribution were indeed described by a power-law decay.

FIG. 6: Exact numerical results for the degree distribution in the stationary regime for a fixed-size network, $p_{\text{del}} = p_{\text{dupl}} = 1$, with $N = 200$ nodes, for increasing duplication probabilities $p = 0.1, 0.2, 0.4, 0.6, 0.8, 0.9, 0.95, 0.975, 0.99, 1$ (marked with lines of decreasing dash-length) and heterodimerisation $\theta = 0.1$. There is a marked difference between the curves generated with $p < 1$ and with $p = 1$. Whereas the latter is described by a power-law decay, the former are not and a characteristic degree is present.

We can offer a simple heuristic argument to account for the difference between the case $p = 1$ and $p < 1$. We believe that it is directly related to the choices we have made for the remaining parameters of the model, namely $p_{\text{del}} = p_{\text{dupl}} = 1$ and $\theta > 0$. For $p < 1$ at each duplication event only some of the links of the mother node are copied by the daughter node, whereas for $p = 1$, all of the mother node’s links are copied. Given that $p_{\text{del}} = 1$ we delete a node and all of its links at each time step, and thus at each duplication event, if $p < 1$, we do not compensate for the loss of links incurred through the deletion process. Hence, the repeated application of duplication events with $p < 1$, given that $p_{\text{del}} = 1$ accounts for the fact that the observed degree distribution does not follow a power-law decay but rather has a characteristic degree present.

6. Comment on relation to biological data

We can test the suitability of the DDDH model as a representation of the evolution of PINs by comparing our results against empirical data. Using data cited in Ref. [8] we can obtain values of the number of proteins in the network and estimates of the average degree of the PINs for yeast, fly, and human. From these, we can derive estimates for $\theta$ using Eq. (13) and
the apparent scaling exponent, $\gamma = 1 - 2\theta$. The results are tabulated in Table I.

| Data       | N   | $\langle k \rangle$ | $\theta$ | $\gamma$ |
|------------|-----|----------------------|---------|---------|
| Yeast (I)  | 4873 | 6.6                  | 0.0014  | 1.0     |
| Yeast (II)| 5397 | 29.2                 | 0.0054  | 1.0     |
| Fly        | 6954 | 5.9                  | 0.00085 | 1.0     |
| Human      | 5275 | 5.7                  | 0.0011  | 1.0     |

**Table I:** Comparison between empirical data [8, 42] and the fixed-size, perfect duplication DDDH model. Values of $\theta$ and $\gamma$ are quoted to 2 decimal places.

For all data sets, the calculated value of $\theta$ is of the order of $10^{-3}$ which agrees well with the fact that it is believed heterodimerisation occurs at a rate not greater than 0.1 [8]. Moreover, the corresponding apparent scaling exponent extracted in all cases is found to be $\gamma = 1.0$. This is consistent with our FSS analysis and corroborates Ref. [8] where a power law with degree exponent $\gamma \approx 1.1$ is given for the data for yeast derived from [42].

**V. DISCUSSION**

The results in Sec. IV indicate that the degree distribution of fixed-size networks where the rate of node deletion is equal to the rate of node duplication, $p_{del} = p_{dupl} = 1$, is dependent on several features. We summarise these as follows. (a) In order to observe a power-law degree distribution it is necessary to have $p = 1$, that is, perfect duplication, and $0 < \theta < 0.5$. (b) No power-law is observed in the degree distribution for $p < 1$, or for $p = 1$ and $\theta \geq 0.5$. (c) In all cases, it is necessary for $\theta$ to be non-zero in order for a positive average degree to be obtained in the stationary regime since for $\theta = 0$, $\langle k \rangle = 0$. (d) In cases where a positive average degree is obtained, it is notable that the network self-organises into a stationary state. We see this as an advantageous feature of this model and comment that this is in contrast to some other network models, such as in Ref. [19], where the average degree is a fixed parameter in the model, or the duplication models where the average degree scales with the network size. (e) Our FSS analysis indicates that for fixed $\theta \in (0, 0.5)$, the scaling exponent of the associated (proper) FSS ansatz is fixed and equal to 1. The new result is obtained through the inclusion of a system-size dependency in the prefactor, $a(N)$, necessary when the apparent exponent $\gamma < 1$, which results in the scaling function being recast in such a way that the only relevant scale parameterising the system is determined by the cut-off, given by $N$ in our case. This example illustrates that it is necessary to be cautious when doing a FSS ansatz in systems where the apparent exponent appears to be less than 1 [43]. (f) Estimates of $\theta$ and $\gamma$ for networks for yeast, fly and human are consistent with estimates from empirical data and our FSS analysis.

If we accept the fact that the fixed-size version of the DDDH model in spite of its simplicity is able to reproduce certain observed topological features of PINs, this in turn would require us to revise the idea that the protein repertoire has evolved over millions of years from a small set of genes to the genomes we observed today in multi-cellular organisms which are typically composed of tens of thousands of genes since the two are not compatible. Clearly, this is a rather drastic measure. Rather than accept such a state of affairs, perhaps all that the results of the fixed-size DDDH model indicate thus far is that we should exercise caution when interpreting minimalistic network models, as attractive as they are. Since we can conjure up many varied and simple network models, with and without growth, which are capable of reproducing observed features of complex systems perhaps the only recourse when trying to pick one network model over the other is to carefully use our knowledge of the essence of the original real system [44].

**A. Extensions**

As a first step in investigating the behaviour of the DDDH model, it seems reasonable to keep things as simple as possible, as we have done here. We have reported on the case of fixed-size networks, and are currently investigating how the topology is affected by varying the relative rates of node deletion and duplication. Moreover, sensitivity to initial conditions is being probed further; we believe that for the fixed-size case, the network features are independent of initial conditions.

However, beyond the steps we have mentioned, there are obvious extension of this model which further investigations could benefit from including. For example, the model is based on an undirected network – it would be interesting to see how best to incorporate dynamics based on directed links and what affect this would have on the in-degree distribution and out-degree distribution. Incorporating this feature would make the model well-suited to describing genetic regulatory networks, for exam-
Moreover, the model assumes that the rate of node deletion and duplication are independent of one another, and independent of any feature of the network such as the size; one could imagine the scenario where this is not the case. Moreover, we consider single-node deletion and single-node duplication – an interesting variation would be to consider multiple-node deletion or duplication, or even duplication of whole modules (motifs) as in Ref. [45], for example.

Finally, the only cause of an increase or decrease in the number of links is either due to a node deletion or node duplication event: links are not added or deleted through any other mechanism. The scenario where (directed) links are stochastically added or removed between already existing nodes would be an interesting amendment to investigate, particularly with regards to the resulting effect on the degree distribution and its corresponding exponent [28, 30, 46].

A final example is that there is no fitness parameter in the model, nor any rule based on selection – our results are independent of both of these features at the gene/protein level, and at the network level, yet it is widely believed that both features are driving forces in the evolution of most, if not all, biological systems. Including these features in a meaningful way would be a highly relevant step towards understanding some of the thornier questions in modern biology today.

VI. CONCLUSIONS

We have introduced and discussed a minimalistic model governed by four parameters, based on dynamic node deletion and node duplication with heterodimerisation. The model is intended to capture some basic features in the evolution of protein interaction networks but we believe that it is also suited to other types of networks in light of the suggested modifications.

Power-law degree distributions were observed for generic parameter values, and a novel finite-size scaling effect was observed for the case of fixed-size networks evolving through perfect duplication and $\theta \in (0, 0.5)$. The existence of an attractive fixed point in the average degree was derived based on mean-field arguments, and corroborated with numerical simulations of the first moment of the degree distribution as described by the two-step rate equation. The above results were then used to to derive a relation for the apparent exponent, $\gamma = 1 - 2\theta$.

Our results thus far indicate consistency with empirical data. Further investigations are required to fully explore and understand the wider phase-space inhabited by this model, and several suggestions have been made to this end.

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