Evolutionary dynamics of higher-order interactions in social networks

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Abstract

We live and cooperate in networks. However, links in networks only allow for pairwise interactions, thus making the framework suitable for dyadic games, but not for games that are played in groups of more than two players. Here, we study the evolutionary dynamics of a public goods game in social systems with higher-order interactions. First, we show that the game on uniform hypergraphs corresponds to the replicator dynamics in the well-mixed limit, providing a formal theoretical foundation to study cooperation in networked groups. Secondly, we unveil how the presence of hubs and the coexistence of interactions in groups of different sizes affects the evolution of cooperation. Finally, we apply the proposed framework to extract the actual dependence of the synergy factor on the size of a group from real-world collaboration data in science and technology. Our work provides a way to implement informed actions to boost cooperation in social groups.

Introduction

Cooperation among unrelated individuals distinguishes humans markedly from other mammals, and it is one of the central pillars of our evolutionary success [1]. Past research has emphasized that the structure of social interactions is crucial for the evolution of cooperation, but thus far predominantly in the realm of networks where links connect pairs of players [2, 3]. However, since cooperation often unfolds in groups, the need for a paradigm shift in the way we model social interactions is evident and indeed urgent. Regardless of the model that we use to describe human interactions, cooperation remains at odds with the fundamental principles of Darwinian evolution, and it is fascinating that we have succeeded in collectively holding off self-interest over most of the last two million years, ever since the genus Homo first emerged [4].

Given this puzzle, the search for reasons and mechanisms that may allow cooperation to evolve and proliferate is an evergreen and vibrant subject across the social and natural sciences [5, 6, 7, 8, 9, 10, 11]. Evolutionary game theory is long established as the theory of choice for addressing the puzzle mathematically [12, 13, 14], wherein social dilemmas constitute a particularly important class of games. Namely, social dilemmas capture the essence of the problem since defection is the individually optimal strategy, whilst cooperation is the optimal strategy for the highest social welfare [15]. An important mechanism for cooperation in social dilemmas is network reciprocity [15], which stands for the fact that a limited interaction range, as dictated by lattices or other types of networks, facilitates the formation of compact clusters of cooperators that are in this way protected against invading defectors. This basic mechanism could also be seen if the degree distribution of the interaction network is strongly heterogeneous [17, 18, 19], if there is set or community structure [20, 21], or
if the evolution unfolds on two or more network layers that mutually support cooperative clusters. The simplest remedy is to consider members of a group to be all the players that are pairwise-connected to a central player. However, since the other players are further connected in a pairwise manner, one would also need to consider all the groups in which the central player is a member but is not central. Evidently, classical networks do not provide a unique procedure for defining a group. Moreover, members of the same group are commonly not all directly connected with one another, which prevents strategy changes among them, either in terms of imitation, replication, or exploration. These facts posit a lack of common theoretical foundation for studying the evolution of cooperation in networked groups. Without knowing who is connected to whom in a group, it is also impossible to implement fundamental mechanisms that promote cooperation, such as reciprocity, image scoring, and reputation.

As a solution, we here introduce and study higher-order interactions in evolutionary games that are played in groups. The distinctive feature of higher-order interactions is that, unlike in classical networks, a link can connect more than just two individuals. Thus, higher-order networks naturally account for structured group interactions, wherein a group is simply made up of all players that are connected by a so-called hyperlink, which is the higher-order analogous of the link. As a paradigmatic example, we consider a standard public goods game on the higher-order analogous of a network, referred to as a hypergraph, see Figure 1. We first show that it corresponds exactly to the replicator dynamics in the well-mixed limit as long as no hyperdegree-hyperdegree correlations exist. As such, it thus provides a formal theoretical foundation to study cooperation in networked groups – effectively a null model – that is amenable to further upgrades. Next, we consider the public goods game on hypergraphs with heterogeneity either in their node hyperdegrees (number of hyperlinks a node is involved into) or in the order of their hyperlinks (number of nodes that form each hyperlink), which allow us to describe the dynamics induced by the presence of highly connected players and to consider scenarios in which the synergy factor depends on the group size in a systematic and consistent way. We show, for example, how synergy factors that are given by different powers of the group size lead to a critical scaling in the transition from defection to cooperation. Lastly, we also demonstrate how the proposed higher-order interaction framework can be used to determine the synergy factor as a function of the group size from empirical data on cooperation and collaborations. Under the assumption that the structure of the hypergraph is the outcome of an optimisation process of the game it hosts, we extract the game parameters from datasets describing collaborations in science and technology, showing that higher-order interactions induce diverse benefits and costs in different social domains.

The public goods game constitutes the fundamental example of a social
dilemma when multiple individuals interact simultaneously. It presents a situation where the gain or loss of an initial investment is shared symmetrically between the members of a group, even if the investment itself can be asymmetric. In other words, there is no correlation between the individual effort and the distribution of the reward, meaning that some players receive more than what they give or deserve, while some others receive less. Metaphorically, one would say that the game has no memory, in the sense that the payoff is assigned blindly to all the players as if the system had lost the information about the original contribution of each player. More formally, the public goods game describes a setting where $N$ players are requested to contribute to a common pool with a token of value $c$.

Cooperators do contribute, and defectors do not. The collected amount is then multiplied by the so-called synergy factor $R$, and the benefit is shared amongst all the members of the group. The payoff for the defectors and cooperators playing in a group of $g$ members is given by $\pi_D = Rcw_C/g$ and $\pi_C = Rcw_C/g - c$ respectively, with $w_C$ representing the number of cooperators in the group. Typically $c$ has a fixed value of $c = 1$, so that the behaviour of the system is determined by the synergy factor $R$, or the reduced synergy factor $r = R/g$. Besides, it is common to represent the state of the system by the fraction of players adopting each strategy, $x_C$ for the cooperators and $x_D$ for the defectors.

The evolutionary dynamics determines how the strategies of the players evolve with each iteration of the PGG, that is, how the fractions $x_C$ and $x_D$ change with time. Here, we implement the so-called fixed cost per game approach, where cooperators contribute with an entire token to each game they play. Individual updates constitute micro-steps of the dynamics, whereas a (global) time step corresponds to $N$ individual steps, so that all the players in the system have the chance to play the game and update their strategies. Players interact among them following the links of the network they are embedded in. As mentioned before, the standard network implementation \cite{35}, henceforth referred to as graph implementation (GI), is not able to account for the most general type of interaction in groups. One of the first proposals to overcome the limitation of a GI is Evolutionary Set Theory \cite{20}, that considers a structure of interaction in which the players are organised as the elements of a set. Yet, the game itself is pairwise, and thus different from the type of approach proposed here. However, it is worth pointing out that the set theory description is equivalent to the hypergraph formalism, and therefore, one should expect the same results when studying the same game on both structures. In this work we have opted for hypergraphs because, as a higher-order generalization of graphs, they inherit the whole family of graph tools with which evolutionary game theory scholars are more familiar with. A few years later, it was proposed to address higher-order interactions by bipartite graphs, having a set of nodes for the players and a second set for the groups \cite{48,49,50}. The authors adapted the PGG to the bipartite graph, in what we call the bipartite implementation (BI). In such a case the game is indeed polyadic, but the update process is still dyadic, and the constrains associated to the formalism do not make it suitable for an analytical
treatment. Here, we generalise the BI to a fully higher-order implementation and provide the theoretical foundation to study higher-order cooperative games in uniform and heterogeneous hypergraphs. Finally, we mention that in a more recent work [51], the authors have considered games played by agents belonging to subpopulations and whose interactions occur across and within the population, providing a useful methodology for situations in which one can get rid of the fine details of the individual connections.

Results

Game Implementation

In order to account for higher-order interactions, we use hypergraphs [46]. A hypergraph, $H(\mathcal{N}, \mathcal{L})$, is a mathematical object that consists of a set of $N$ nodes $\mathcal{N} = \{n_1 = 1, ..., n_N = N\}$ and a set of $L$ hyperlinks $\mathcal{L} = \{l_1, ..., l_L\}$. Each hyperlink is a subset of two or more elements of $\mathcal{N}$ and represents a group interaction. For instance, in Figure 1a, the hyperlink $l_1$ contains nodes $n_1$ and $n_3$, whereas the hyperlink $l_3$ is the subset made up by nodes $n_4$, $n_5$ and $n_6$. Furthermore, the cardinality of a subset, known as the order of the hyperlink, is the number of nodes in the group. In the previous example, $l_1$ has order 2 and $l_3$ has order 3. In a hypergraph, the hyperdegree, $k_i$, of a node $i$ represents the number of hyperlinks in which the node is involved into, thus, the number of groups of a specific order $g$ that contains $i$ can be denoted by $k^g_i$. Hence, the hyperdegree of $i$ is given as $k_i = \sum_{g=g^-}^{g^+} k^g_i$, where $g^-$ and $g^+$ account for the minimal and maximal orders in $\mathcal{L}$. For example, in Figure 1a, $k_4 = 3 = k^2_4 + k^3_4$, with $k^2_4 = 1$ (the hyperlink $l_4$) and $k^3_4 = 2$ (the hyperlinks $l_2$ and $l_3$). As $\langle k \rangle$ we indicate the average hyperdegree of node $i$, where the averages are evaluated over all the nodes in the system, i.e. $\langle k \rangle = \frac{1}{N} \sum_{i \in \mathcal{N}} k_i$.

Although hypergraphs are not the only possible representation of group interactions, they allow exploiting the analogy between the links representing pairwise interactions in contact networks and hyperlinks, which are based on higher-order, group interactions. As we will show next, the differences between these two approaches lead to fundamentally distinct outcomes of the PGG evolutionary dynamics. To see how the evolutionary dynamics evolves in hypergraphs, let us consider the first step of a standard graph implementation of the PGG. When a node $n_i$ and one of its neighbours $n_j$ are selected on a graph, it is equivalent to say that a node and one of its links are selected. Such a procedure can be easily generalised to group interactions of more than $g = 2$ individuals, see Figure 1b. Note that if we choose more neighbours of $n_i$ to generate higher-order interactions, such an extension would still be based on dyadic ones. Instead, we propose a hypergraph implementation (HI) of the game that consists of selecting one of the hyperlinks of $n_i$. That is, in the HI setup, we select at random with uniform probability a node $n_i$ in the hypergraph and one of its hyperlinks, $l_i$. Then, all the members of the hyperlink $l_i$ play a game for each of the hyperlinks they are part of, as illustrated in Figure 1. Finally, as it
is customary, the nodes accumulate the payoffs of all the rounds they play, and we normalise this quantity by the total number of played games, such that each node’s performance is represented by its average payoff.

The second part of each micro-step of the evolutionary dynamics of the game involves updating the strategy of node $n_i$. To this end, we normalise the discrete replicator dynamics for the case of higher-order interactions. We propose to compare the payoff $\pi_i$ of a node $n_i$ with the maximal payoff of the selected hyperlink $l_i$. Under this rule, $n_i$ will adopt the strategy of the node with the maximal payoff with a probability $\frac{1}{\Delta}(\max_{j} \pi_j - \pi_i)$, where $\Delta$, whose precise definition is provided in Equation (9), accounts for the maximal payoff difference, and is employed to guarantee that the probability is normalised. The rationale behind the choice of this expression is that node $i$ will compare its payoff to that of the node with the largest payoff in hyperlink $l_i$. Note that the previous expression reduces to the standard one of the GI when $g = 2$.

Summing up, the HI accounts for a more realistic update than that in the BI, since the player inspiring a strategy change is the one with the highest payoff of the group, and not a randomly chosen one.

Uniform Hypergraphs

To get some insights into the dynamics of the system in a simple configuration, we first studied the PGG on uniform random hypergraphs (URH) with hyper-links of order equal to $g = 2$, 3, 4 and 5 (see Methods for details on how to generate URH). Numerical simulations have been carried out for hypergraphs with $N = 1000$ nodes (players), and the game has been iterated for $T = 10^4$ time steps. The results obtained are reported in Fig. 2. Panel (a) shows the final fraction of cooperators as a function of the reduced synergy factor $r$. In each case, the simulations refer to hypergraphs with $L = L_c$ hyperlinks, where $L_c$ accounts for the minimal number of hyperlinks that guarantees the connectedness of the hypergraph. As it can be seen in the figure, there is a value of $r$ beyond which cooperation emerges. We define this critical value of the reduced synergy factor, $r_c$, which depends on $g$, as the lowest value of $r$ for which the fraction of cooperators is nonzero.

The results show that $r_c$ decreases when the order $g$ of the hyperlinks of the hypergraph increases. This is equivalent to say that $r_c$ decreases when the same number of $N = 1000$ individuals play in larger groups. We believe that this observation is important, since determining how $r$ varies with the size of the group, allows us to get more realistic insights. Admittedly, the well-mixed limit of population-size groups is rarely applicable in reality, thus, the study of the impact of having large groups inside large populations, as allowed by our higher-order framework, is key. The panel (b) of Fig. 2 displays how the value of $r_c$ depends on the number of hyperlinks $L$ in the hypergraphs. For each value of $g$, we observe an increase of $r_c$ with $L$, and a tendency, for large hypergraph densities, to the value $r_c = 1$, which corresponds to the well-mixed replicator approximation [52]. The replicator equation approximation relies on the indistinguishability of the nodes, and as such, it is exact when the hypergraph is fully
Figure 1: Higher-order vs pairwise interactions in a Public Goods Game (PGG). Comparison of the proposed hypergraph implementation (HI) with a standard graph implementation (GI) of the game based on pairwise interactions only. (a) In the HI implementation, a node, \( n_2 \), and one of its hyperlinks, \( l_2 \), are randomly selected. All the nodes in \( l_2 \), namely node \( n_2 \), and the two nodes highlighted in red \( n_3 \) and \( n_4 \), play all the games they are involved in, corresponding, in this example, to PGG defined for the subset of nodes of the hyperlinks \( l_1, l_2, l_3 \) and \( l_4 \). Then, the strategy of \( n_2 \) is updated by comparing its payoff with that of the node with the highest accumulated payoff of the hyperlink \( l_2 \). This is not equivalent to play the PGG in the graph generated by projecting the interactions of the hypergraph, which is shown in (b). In the standard GI implementation, a neighbour of \( n_2 \), let us say \( n_3 \) —highlighted in red— is randomly selected. The two nodes \( n_2 \) and \( n_3 \) then play all the games of the groups they are part of, that is, of the groups made up by the subsets of nodes \( \{n_1, n_3\} \), \( \{n_2, n_3, n_4\} \), \( \{n_1, n_2, n_3, n_4\} \) and \( \{n_2, n_3, n_4, n_5, n_6, n_7\} \). These subsets, coloured as indicated in the figure, could be represented by a different set of hyperlinks \( \bar{l}_1, \bar{l}_2, \bar{l}_3 \) and \( \bar{l}_4 \), respectively, which are different from the set of hyperlinks of the original hypergraph. Finally, the strategy of \( n_2 \) is updated by comparing its accumulated payoff to that of node \( n_3 \).

connected, i.e. contains all the possible hyperlinks. However, we show that the approximation is good also for sparse hypergraphs, with a number of hyperlinks of the order of the critical value for ensuring a giant component. Therefore it is natural that the higher the value of \( L \), the closer \( r_c \) is to 1. The same argument can be used to explain the results in (a). The ratio \( L_c / C_N^g \), which represents the fraction between the critical number of hyperlinks \( L_c \) and the total possible number of hyperlinks, given by the binomial coefficient \( C_N^g \), decreases with \( g \). This implies that, if two hypergraphs have \( L = L_c \), but different values of \( g \), the one with lower \( g \) will be denser, and thus will exhibit a critical point closer to the analytic prediction. Therefore, we can say that at fixed reduced synergy factor \( r \), large groups are better to foster cooperation in sparse hypergraphs, as the
Figure 2: Public Goods Game with higher-order interactions in uniform random hypergraphs. Numerical simulation of the Hypergraph Implementation of the game on uniform random hypergraphs of $N = 1000$ players and different orders $g$. (a) Fraction of cooperators, $x_C$, as a function of the synergy factor, $r$, for hypergraphs with hyperdegree $\langle k \rangle = k_c$, or total number of hyperlinks $L = L_c$, where $k_c$ and $L_c$ stand for the critical hyperdegree or number of hyperlinks guaranteeing a connected hypergraph. (b) Critical value of the synergy factor, $r_c$, as a function of the ratio between the number of hyperlinks $L$ and the critical value $L_c$ in hypergraphs of different density. (c) Relaxation times as a function of the synergy factor, $r$, for hypergraphs with hyperdegree $\langle k \rangle = 5k_c$. In all plots, triangles correspond to numerical simulations, while the solid lines are the results of our theoretical predictions.

number of hyperlinks required for connecting all the players represents a smaller fraction of the total number of hyperlinks. Finally, the value of $r$ also influences how long it takes for the system to converge to the stationary solution. This is illustrated in panel (c), where we report the measured relaxation time $T$ from an initial configuration with $x_D = x_C = 0.5$, in a hypergraph with $L = 5L_c$. These results are obtained by running the simulations up to a maximum of $10^4$ steps. Furthermore, for the replicator approximation, the value of $T$ can be analytically computed as $T = \frac{\ln(N-1)}{Q}$, with $Q = (1 - r)/\Delta$ (see Equations (10) and (11) for the details of the calculation). As it can be seen in the figure, the agreement between the theoretical predictions and the numerical results is not only qualitatively but also quantitatively very good. The absorbing state, either full cooperation or full defection, emerges when the system is at equilibrium, a condition that can only be reached if enough iterations have occurred. On the other hand, real-world social interactions that can be modelled as games usually take place over a limited time interval $\tau$. Hence, the relation between the relaxation time $T$, which depends on the synergy factor $r$, and $\tau$ is crucial to determine if the system does or does not reach the equilibrium, and consequently, if the replicator dynamics can or cannot predict the numerically computed fraction of cooperators. All these results indicate that the dynamics of the PGG on uniform random hypergraphs corresponds to the replicator dynamics in the well-mixed limit. In order to test the robustness of these findings with respect to the implementation selection, we have also carried out numerical simulations in the BI implementation (see Figure 6).
Hyperdegree-Heterogeneous Hypergraphs

The previous section addressed the simplest scenario in which the individuals of a population are assumed to be indistinguishable (URH). However, such an assumption can be too oversimplified to describe real situations as it is well known that social systems are heterogeneous. Think of your friends at college. It is likely that a minority of those are involved in considerably more activities, and therefore social circles, than the rest. Such heterogeneity is typically characterised by a non-exponential degree distribution, allowing the presence of hubs, or highly connected individuals [45]. Hence, we consider here two families of hyperdegree-heterogeneous hypergraphs that we name power random hypergraphs (PRH) and scale-free random hypergraphs (SRH). The algorithms we have used to generate these hypergraphs are reported in the Methods section, and their properties have been studied in Figure 7 and Figure 8. Scale-free hypergraphs are characterised by a power-law distribution, and represent the most hyperdegree-heterogeneous family of hypergraph considered here. For this reason, these hypergraphs display a hierarchy between the nodes, as a few of them are involved in most of the hyperlinks and thus have a dominant position in the dynamics of the system. In contrast, power random hypergraphs stay in between uniform and scale-free hypergraphs, as their hyperdegree distribution combines exponential and non-exponential functions.

To study the emergence of cooperators in hyperdegree-heterogeneous hypergraphs, we have run $T = 10^4$ time steps of the game on ensembles of hypergraphs with $N = 1000$ nodes and orders $g = 2, 3, 4, 5$, respectively sampled from PRH and SRH. In order to compare the simulations with those reported in Fig 2a, we have fixed the total number of hyperlinks to $L = L_c$. When, for high heterogeneity, some of the nodes (a minimal fraction of the total) do not belong to the main component, we have neglected their contribution to the fraction of cooperators.

The results reported in Fig. 3 show an important difference between PRH and SRH. In the case of PRH (top panels) the position of the transition does not depend on the heterogeneity of the node hyperdegree distribution, tuned by parameter $\mu$, (see Methods for the precise definition of $\mu$), and the critical point is the same as that obtained in URH. Conversely, the simulation of the game on SRH (bottom panels) shows that, the larger the heterogeneity in the hyperdegree distribution (larger values of $\mu$), the more the solution deviates from that of URH, and the closer the critical point gets to $r = 1$. This indicates that hierarchically structured systems inhibit cooperation in the PGG with higher-order interactions at variance with numerical simulations obtained on traditional networks under the same evolutionary dynamics.

In order to be able to explain these results we need to consider a refinement of the replicator approximation that takes into account the possible presence of correlations between the hyperdegrees of nodes belonging to the same hyperlink. Let $K$ be the set of all possible hyperdegrees a node can have, and let $k \in K$ be the hyperdegree of a randomly chosen node. We now denote as $p(k' | k)$ the conditional probability that the node of hyperdegree $k$ is part of a hyperlink
where the remaining $g - 1$ nodes have hyperdegrees $k'' = \{k_1, k_2, \ldots, k_{g-1}\}$, where $k'' \in K^{g-1}$ is a vector whose $g - 1$ components are elements of $K$. We have been able to show that the system will fulfil the replicator approximation as long as the conditional probability $p(k''|k)$ does not depend on $k$ (see Equation 12 and the section below it for the detailed analysis). This is true for the case of the PRH. Conversely, in the case of the SRH, increasing heterogeneity while maintaining the total number of hyperlinks in the hypergraph requires reducing the number of effective nodes. This induces non-trivial correlations in the model between the hyperdegrees of nodes belonging to the same hyperlink, and has a similar effect of driving the system closer to the $r_c = 1$ threshold, as that we have observed when we increase the hyperlink density in the uniform case (URH). Intuitively this can be explained by the notion of locality. When the density is low, or when no large hubs are present in the system, there is a non-negligible probability that cooperator bubbles emerge below the critical threshold, because there may be regions of the hypergraph that are semi-isolated, and therefore protected from defectors, even if they belong to the same component. However, either increasing the density or introducing hubs will reduce the probability of finding these isolated groups of nodes, and therefore will inhibit the formation of cooperator bubbles below $r = 1$. 

Figure 3: Public Goods Game with higher-order interactions in hyperdegree-heterogeneous random hypergraphs. Numerical simulation for the fraction of cooperators $x_C$ as a function of the reduced synergy factor $r$ on hyperdegree-heterogeneous random hypergraphs of $N = 1000$ players and different orders $g$. Top and bottom panels refer respectively to PRHs and scale-free random hypergraphs Random Hypergraphs (SRH), while the four different columns (a)-(e), (b)-(f), (c)-(g) and (d)-(h) correspond to the values of $g = 2, 3, 4, 5$. Legends in the PRH and SRH plots denote the value of $\mu$ characterising the hyperdegree distribution, where larger values $\mu$ imply higher heterogeneity.
Order-Heterogeneous Hypergraphs

Heterogeneity can also arise in the order of the hyperlinks. Indeed, the proposed HI of the PGG allows studying the more general, realistic and interesting case of hypergraphs where not all the hyperlinks have the same order. Important examples of such systems include teams of different sizes working for a common goal or one-to-many communication via apps like WhatsApp, where users can create and belong to several groups of different sizes. In what follows, we consider order-heterogeneous random hypergraphs with an assigned distribution of hyperlinks. Such hypergraphs are characterised by their total number of hyperlinks $L$ and by a probability vector $\mathbf{p} = \{p_g\}_{g=g_-}^{g=},$ whose entry $p_g = k^g/k$ specifies how likely it is, on average, that the hyperdegree $k$ of the node contains $k^g$ groups of order $g$. $\mathbf{p}$ is normalised such that $\sum_{g=g_-}^{g=} p_g = 1$. Considering groups of different orders in the same hypergraph allows us to focus on another important aspect of the PGG on higher-order structures, namely, the possible dependence of the rescaled synergy factor $r$ on the order of the group. This is important for practical purposes, given the increasing interest in understanding how the size of a group impacts its performance. As it has been shown recently $^{53}$, large and small teams play different roles in science and technology ecosystems. Thus, it is natural to assume that the synergy factor of a group depends on its size. This is particularly true in scientific publications, where it has been shown that the larger the group, the more citations a produced publication is likely to attract $^{54, 55}$. Therefore, as a general form for such a dependence we assume that the synergy factor $R$ is an increasing power-law function of $g$, namely:

$$R(g) = \alpha g^\beta$$

with parameter $\alpha > 0$ and exponent $\beta \geq 0$. The value of the exponent allows to tune the benefit that the players are able to produce when working as a group. In particular, adopting a superlinear scaling $\beta > 1$, means considering a synergistic effect of a group that goes beyond the sum of the individual contributions $^{56, 57}$. Notice, however, that the assumed dependence in Eq. (1) is only a first approximation as it neglects saturation effects or even possible disadvantages due to difficulties in coordinating large groups, which, as we will see later on, appear in real systems. Under this assumption, the average payoff difference between cooperation and defection can be written as:

$$\pi_D - \pi_C = \sum_{g=g_-}^{g=} p_g (1 - \alpha g^\beta - 1)$$

where $g_-$ and $g_+$ are again the minimal and maximal orders of hyperlinks, respectively. The relaxation time is again given by $T = \ln(N - 1)/|Q|$, where $Q = (\pi_D - \pi_C)/\Delta$ (see the Appendix for the definition of $\Delta$ in the general case and for explicit calculations). It is then possible to derive the critical value of the parameter $\alpha$ as a function of the exponent $\beta$ as:

$$\alpha_c(\beta) = \frac{1}{\sum_{g=g_-}^{g=} p_g g^\beta - 1} = \frac{1}{K_\beta},$$

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where, for simplicity, we have defined $K_\beta \equiv \sum_{g=2}^{\infty} p^g g^{\beta-1}$. We remark here that $\alpha = \alpha_c$ for a fixed value of $\beta$ is the critical point separating the defection and cooperation phases. This means that when $\alpha < \alpha_c$ the system will converge to full defection, while for $\alpha > \alpha_c$ it will converge to full cooperation.

To explore how the dynamics evolves in order-heterogeneous random hypergraphs, we have performed numerical simulations of the PGG considering four different values of $g = 2, 3, 4$ and 5 and allowing the values of $p^g$ to take only multiples of 0.25. This leads to 35 possible hypergraphs, one for each of all conceivable convex sums of \{2, 3, 4, 5\} with the previous constraints. This means that the hypergraphs we consider are composed by hyperlinks of different orders, where each order $g$ takes $Lp^g$ hyperlinks out of the total number $L$. For instance, on a hypergraph with $L = 100$ and order probabilities $(0, 0.25, 0.25, 0.5)$, on average we would expect 25 hyperlinks of order $g = 3$, another 25 of order $g = 4$ and the remaining 50 of order $g = 5$. Results are reported in Fig. 4 for four different values of the power exponent $\beta$, namely, $\beta = 0, 1, 2, 3$, shown with different colours. Notice that the case $\beta = 1$ corresponds to the underlying linear assumption of the standard PGG: in this case, $\alpha$ plays the role of the reduced synergy factor $r$. Panels a) through d) plot the colour-coded fraction of cooperators as a function of the parameter $\alpha$ in the definition of the synergy factor. The hypergraphs $H_i$ have $\langle k \rangle = 2k_c$ and are displayed according to their value of $K_\beta$, i.e., the value of the critical point $\alpha_c(\beta)$. As for the case of uniform random hypergraphs, we find that although the critical point is slightly overestimated for low densities by the analytical approximation, there is still a good agreement between the theoretical predictions of the well-mixed replicator approximation and the numerical simulations. We next explore the behaviour of the relaxation time. Panels e) through h) show results obtained for order-heterogeneous hypergraphs with $\langle k \rangle = 5k_c$. As it was done for the homogeneous scenario, we follow the dynamics of the system up to a maximum of $T = 10^4$ time steps. The plots show that the relaxation times depend on $\alpha$ for all values of $\beta \neq 1$, albeit rather differently with respect to the dependence of the critical value $\alpha_c$ for $\beta < 1$ and $\beta > 1$. In order to further explore this relationship, we analysed how the average relaxation time varies as a function of the critical point $\alpha_c$. Results shown in panels i) to l) reveal that the dependence is always linear. However, when the synergy factor increases superlinearly, there appear different curves, each one corresponding to a distinct family of hypergraphs and characterised by a different linear relation between the average relaxation time and the critical value. This behaviour introduces an additional degree of freedom that can turn very useful, since the degeneracy that is observed for $\beta \leq 1$ is broken for $\beta > 1$, and therefore one can independently set a critical point and a relaxation time by opportunely choosing the corresponding hypergraph. We remind the reader that cases with $\beta > 1$ are those in which the synergy factor $r(g)$ has a superlinear dependence on the order $g$. Those values of $\beta$ are a priori the most interesting ones to study, and the ones more likely to be found in real situations. Therefore, our results about the relaxation are particularly relevant, because in this case one can potentially turn an unstable system into a stable one, and the opposite, by changing the
order of the hyperlinks, while still respecting the value of the critical point.

Figure 4: Public Goods Game with higher-order interactions in order-heterogeneous random hypergraphs. We assume that the synergy factor grows according to Eq. (1) and consider the set of hypergraphs \( \mathcal{H}_i \) that contain hyperlinks of orders \( g = \{2, 3, 4, 5\} \) with probabilities \( p_g \) taking values in the set \( \{0, 0.25, 0.5, 0.75, 1\} \) —there are 35 possible such hypergraphs. (a-d) Fraction of cooperators as a function of \( \alpha \) for each of the 35 hypergraphs \( \mathcal{H}_i \) and several values of \( \beta \). The hypergraphs are ordered according to their value of \( K_\beta \). Simulations have been carried out up to \( T = 10^4 \) time steps for hypergraphs with \( \langle k \rangle = 2k_c \), and triangles correspond to the theoretical predictions in the replicator approximation (see the Appendix for details). (e-h) Relaxation time as a function of \( \alpha \) for the set of hypergraphs \( \mathcal{H}_i \). Now hypergraphs have \( \langle k \rangle = 5k_c \). (i-l) Predictions for the critical value \( \alpha_c \) as a function of the average relaxation time, calculated for each hypergraph in \( \mathcal{H}_i \) by averaging over the intervals of \( \alpha \) \([0, 0.8] \), \([0, 2] \), \([0, 1]\) and \([0, 0.5]\) for \( \beta = 0, 1, 2, 3 \) respectively.

**Synergy factor of real games**

From the previous results, a natural question arises: is it possible to determine the value of the synergy factor for a real PGG for each of the possible group sizes? A plausible answer to this question can be obtained under the assumption that the very same structure of the hypergraph is the result of an evolutionary process in which nodes select the groups they belong to. We hypothesise that
each individual tries to optimise the ideal number of groups of each order, based on the perceived dependence of the synergy factor on the group size. In this way, each real-world hypergraph would be the optimal structure that supports the game it hosts. We could then extract the functional form $R(g)$ directly from the hyperdegree distribution of the hypergraph. More precisely, the goal would be to use the information in the vector $p$ of the hypergraphs on which the PGG occurs to determine the functional form, $R(g)$, of the synergy factor by imposing two conditions. The first condition comes from the assumption that the unknown reduced synergy factor $r(g)$ is proportional to $p_g$. This originates in the intuition that the distribution of the hyperdegree of a generic player should be aligned with the potential benefit that each player expects to obtain for each higher-order interaction. The second condition imposes that the average payoff of cooperators is equal to the average payoff of defectors. This implies that the system is at equilibrium and guarantees the coexistence of cooperators and defectors. Thus, given that these two conditions are satisfied, it is possible to extract the curves of $r(g)$ and $R(g)$ from empirical data on higher-order interactions.

![Figure 5: Synergy factors of scientific collaborations. (a) Empirical synergy factors extracted from the structure of hypergraphs describing co-authorships from publications in the American Physical Society journals. Solid lines are fit of the empirical dots according to Eq. (6). Different symbols and colours refer to different journals, i.e. to different scientific communities. (b) Journal (labelled from 1 to 13) positions in the costs-benefits diagram. The synergy factors are factorized as a function of two competing terms, one modeling the benefits of cooperation, which is dominant for small group size, and another one accounting for costs associated to an excessively large number of co-authors, which describes for the exponential decays observed in the first two panels.](image)

In order to show how the above-mentioned procedure works in practice in real cases, we have studied collaboration in science and technology. We believe that this could constitute an example in which the benefit of a group depends on its size and at the same time, all group members do not contribute the same to the collective task, which essentially leads to a PGG dynamics. Although there is not a single way of classifying in a binary manner (either cooperator or defector) the authors of a scientific publication, one can think of two type of players mimicking cooperators and defectors. A cooperator can be considered as anyone...
that has contributed at least a “fair” amount of work. The reverse applies to defectors, which can be considered those that put less effort in producing a teamwork than the average or the “fair” amount of work. Note that whatever the effort of the team members is, they all receive the same benefit, for instance, in terms of citations (the citation is to the paper, not to the individual). Thus, given that there are cooperators and defectors, what is the optimal collaboration (group) size? And that of the synergy factor?

In particular, we have considered a large data set of all the scientific articles published in the last century in thirteen journals of the American Physical Society (APS). For each journal, we have constructed a hypergraph whose nodes and hyperlinks represent respectively scientists and co-authored publications (see Table 1 for further details). The reduced synergy factors have then been obtained from information on the number of authors in each publication (see Methods). From the plots of $r(g)$ vs $g$ reported in Fig. 5a we notice the existence of a maximum value of $r$ at intermediate group orders $g$. This indicates that there is an optimal trade-off between the positive and negative effects of increasing the group size. The optimal value of $g$ depends on the specific scientific community, as it varies from journal to journal. In the case of PhysRevLett the maximum of $r(g)$ is located at $g = 3$. Different journals are associated with other optimal collaboration sizes. For instance, for PhysRevApplied $r(g)$ is maximum at $g = 5$, indicating that larger collaborations are more beneficial in applied topics, such as device physics, electronics and industrial physics. For almost all journals, the synergy factor is low for $g = 1$, showcasing the difficulty of publishing alone in physics, a research area where teamwork has been becoming increasingly important in the last decades [58]. Interestingly, a paradigmatic case is the one of PhysRevSeriesI, the very first journal published by the APS in the early 1900s, for which a sharp peak is located at $g = 1$, showing how most publications were produced by single scientists, in contrast with current trends. In order to shed light on this result we have factorised the synergy factor as the product of an increasing function of $g$ times a decreasing function of $g$, and we have performed a numerical fit to extract the benefit exponent $\beta$ and the so-called cost parameter $\gamma$ (see Methods). This enables us to interpret the synergy factor as a combination of two contrary effects of the higher-order interactions in this particular dataset.

Fig. 5b reports the values of $\beta$ and $\gamma$ obtained for each journal of the APS, and it allows us to classify the different scientific communities in terms of benefits and costs of higher-order interactions. These results provide a game-theoretic interpretation of the APS dataset. Specifically, in the context of this bibliographic dataset, hidden benefits and costs that conform to the synergy factor can be associated with several aspects of the task of producing a publication. Benefits (an increase of the synergy factor with increasing $g$) would correspond to the potential reinforcement of the amount and quality of the ideas and the potential increase in the outreach of the work with the number of co-authors involved. On the contrary, the costs (decrease with increasing $g$) would be the additional organisational effort in the process of arriving at a consensus and carrying out the tasks for publishing a paper. Experimental communities, such as that of
nuclear physicists publishing in PhysRevC, tend to have low costs. These ideas are aligned with recent studies about the creation and production of research ideas [59] and the role, group dynamics and success of teams [53, 54, 55]. Our formalism allows for a quantitative analysis of these phenomena and could be used in future applications to design ways to foster higher-order cooperation.

Discussion

Summing up, we have introduced higher-order interactions in evolutionary games to study cooperation in groups. Since higher-order interactions allow for a single link to connect more than just two individuals, they are naturally suitable to define groups in networks. In doing so, higher-order interactions thus do away with the arbitrary definitions of groups in classical networks, and they provide a formal theoretical foundation to study cooperation in networked groups. We have shown that the public goods game on a hypergraph is effectively a null model that agrees exactly with the replicator dynamics in the well-mixed limit as long as no hyperdegree-hyperdegree correlations exist. As such, it can be used in future research towards upgrades that add additional layers of reality in models of human cooperation, either by means of strategic complexity [10], or by means of more complex interaction networks [50].

Towards the latter effect, we have also studied the public goods game on hyperdegree-heterogeneous and on order-heterogeneous hypergraphs, where we study the effects of the presence of highly connected individuals and of hyperlinks of different orders respectively. Due to the exact definition of a group in the proposed framework, we have been able to systematically and consistently consider synergy factors that are dependent on group size. Indeed, the framework allows us to unveil the effects of group size on cooperation in its most general form. As an example, we have considered synergy factors that are given by different powers of the group size, showing a critical scaling in the transition from defection to cooperation. In this case too, we have observed a substantial agreement between the simulations and the analytical predictions of the model. Interestingly, we found that hierarchically structured hypergraphs could hinder cooperation in a structured population. Our framework enables analysis of real systems, as we have shown for the APS publications dataset, providing insights regarding the positive and negative effects associated to higher-order interactions and the nature of group dynamics. However, even if our framework includes diverse forms of higher-order interactions, we recognize that a current limitation of this representation of human interactions is given by the constraints imposed by the available data. Admittedly, the identification of interactions in social networks beyond the traditional pairwise relationships constitutes nowadays an important challenge. Interestingly enough, this also represents an opportunity from an experimental point of view. It is also worth mentioning that the application of our results to scientific publications is based on the hypothesis that the interaction structure is the outcome of an optimization process, where the average distribution of groups that each node is part of coincides with the
synergy factor, such that the system is in a stationary state of the dynamics of the PGG. This hypothesis, which constitutes a limitation of our method to extract synergy factors from real data, could potentially be either validated or refuted by models considering the dynamics within the topology of interactions on top of the PGG. Moreover, the PGG imposes all cooperators to contribute with the same amount, making this contribution a boolean variable in practice. And therefore, an additional limitation arises when adapting real systems to the rigid formalization of the PGG, as the role of cooperators and defectors cannot be unequivocally defined when the contributions are not only 0 or 1.

It is also worth mentioning that in his essay titled Innate Social Aptitudes of Man, W. D. Hamilton wrote, “There may be reasons to be glad that human life is a many-person game and not just a disjointed collection of two-person games”. He was referring to the fact that social enforcement works better in groups with more than two members, which can offer at least a partial cure for the problems with reciprocation in larger groups [38]. We note that the theoretical framework of higher-order interactions also invites to re-examine other fundamental mechanisms that may promote cooperation, such as image scoring [39, 40, 41], rewarding [61], and punishment [62, 63, 64, 65].

Given the fundamental differences between pairwise and higher-order interactions, it would also be of interest to revisit the role of specific network properties and their role in the evolution of cooperation. In this regard, the role of community structure [21], as well as two or more network layers [22, 23, 24, 25, 26, 27, 28, 29], promise to be fruitful ground for future explorations on how interaction structure impacts cooperation. Overall, we believe that the introduction of higher-order interactions to evolutionary games has the potential to improve our understanding of the evolution of cooperation and other social processes in networks.

Methods

Uniform random hypergraphs (URH): We detail here the procedure we have adopted to sample $g$-uniform hypergraphs, i.e. hypergraphs with all hyperlinks of the same order $g$. A URH of order $g$ can be constructed by assigning a uniform probability $p$ to each $g$-tuple of $N$. For each of them, a random number in the $[0,1)$ interval is generated, and if this number is lower than $p$, the hyperlink containing the $g$-tuple is created. However, this method scales badly with $g$ since the number of $g$-tuples to be considered is equal to the binomial coefficient $C_g^N = \binom{N}{g}$, which grows fast as a function of $g$. A more efficient procedure is to fix the total number of hyperlinks, $L$, and generate a random integer in the $[1, C_g^N]$ interval. One has to provide an ordering for the set of all possible hyperlinks, so that each of the random integers corresponds to a hyperlink. The hyperlinks selected through this process are then added to the hypergraph. The hyperlink ordering is based on the following combinatorial
that allows us to partition set $\mathcal{L}$ of all the possible hyperlinks of a $g$-uniform hypergraph in terms of disjoint hypergraphs, each one of them containing the hyperlinks that form the corresponding $g$-star hypergraph \[66\]. This holds true in general, which enables us to apply the same argument recursively, such that we can order all the possible hyperlinks univocally, and even more, the probability for having a specific node in a hyperlink is equal for all the nodes. These properties arise from the combinatorial probabilities \[ \frac{C_N}{C_g} \] for \( i = 1, \ldots, N - (g - 1) \) i.e. the normalised weights of each of the terms in the summation. We have empirically found a distribution that can be used as an approximation to \( c_i = \sum_{j=1}^{i} d_j \), the cumulative distribution of \( d_i \), namely given by \( 1 - (1 - x)^g \), where \( x = i/(N - (g - 1)) \). See Figure 7 for a numerical proof of the convergence between both expressions.

For the purpose of studying the stationary condition of a game, we are interested in having a connected hypergraph. The critical thresholds for the number of hyperlinks, \( L_c \), and the hyperdegree, \( k_c \), are equal to \( L_c = \frac{N}{g} \ln N \) and \( k_c = \ln N \). Hence, when \( L \) is larger than \( L_c \), there is a high probability that the resultant hypergraph is connected.

**Power random hypergraphs (PRH):** We have seen that using the combinatorial probabilities \( d_i \) allows us to create uniform random hypergraphs. Therefore, increasing the value of the exponent \( g \) to \( g' \) in \( c_i \), such that \( g' > g \), will increase the probability of sampling the hyperlinks belonging to the g-star hypergraphs of low index nodes, and therefore introduce heterogeneities in the degree distribution. The control parameter that we use in the simulations in the manuscript is \( \mu \in [0, 1] \). In terms of \( \mu \), one can obtain the power to use in the cumulative distribution \( g' \) as \( g' = (1 + \mu)g \). In order to sample hyperlinks of order \( g \) according to the new distribution, we transform the random number \( r \) to a different random \( r' \)

\[
r' = c_{i-1}(0) + \left[ r - c_{i-1}(\mu) \right] \frac{d_i(0)}{d_i(\mu)}
\]

Here \( i \) is the g-star to which the hyperlink would belong if it was sampled according to \( r \). In this expression \( d_i(\mu) \) and \( c_i(\mu) \) account for the distributions using the value of \( g' \) as a function of \( \mu \). Accordingly, \( d_i(0) \) and \( c_i(0) \) are simply the distributions of the uniform case. See Figure 8 for the analysis of degree distribution emerging from the PRH.

**Scale-free random hypergraphs (SRH):** The standard indicator of heterogeneity in graphs is the power-law decay of the degree distribution. Here we employ the static scale-free algorithm \[67\] to generate such a profile. We use the same control parameter as in the PRH, \( \mu \in [0, 1] \), which in this case results
in a power-law $p_k \sim k^{-\lambda}$ where the power $\lambda$ is $\lambda = 1 + 1/\mu$. See Figure ?? for the degree distribution of the hypergraphs generated with the SRH.

**Extracting synergy factors from real data:** We show here how the dependence of the reduced synergy factor $r(g)$ on group size $g$ can be derived for real systems, based on the assumption that this information is encoded in the very same structure of a hypergraph. In particular, we have considered a data set of scientific publications and we have used it to investigate how benefits change with the size of groups in scientific collaborations. The data set consists of 577,886 papers published in the period from 1904 to 2015 in the collection of all the journals of the American Physical Society (APS) [68]. We have constructed the 13 hypergraphs corresponding to different journals, such as Physical Review, Physical Review Letters, etc. of the APS. The nodes and hyperlinks of these hypergraphs represent scientists and publications respectively. The order of a hyperlink is equal to the number of authors of the corresponding publication. For each hypergraph, we have extracted the number $L^g$ of hyperlinks of a given order $g$, which we used to compute the average number $k^g = gL^g/N$ of hyperlinks of order $g$ a node is involved in. The reduced synergy factor $r(g)$ can then be extracted from the proportion $p^g = k^g/k$ of hyperlinks of order $g$ of a node, by assuming that $r(g) = zp^g$ and using the critical point relation:

$$ \sum_{g=g^-}^{g=g^+} p^g (1 - r(g)) = 0$$

(5)

to calculate the proportionality constant $z$.

**Cost-benefit factorisation of the synergy factor:** In scientific collaborations across all journals of the APS, an optimal team size is associated with a maximum in the synergy factor, suggesting that an excessively large number of co-authors might lead to disadvantages in cooperation. In order to account for these effects, we have modelled the synergy factor extracted from real-world collaboration data as the following function of $g$:

$$f(g, \alpha, \beta, \gamma) = \alpha g^\beta e^{-\gamma(g-1)}.$$  \hspace{1cm} (6)

ruled by the three parameters, $\alpha, \beta$ and $\gamma$. The first parameter, $\alpha$, introduced in Eq. (1), is determined by the critical point condition. The remaining two parameters account respectively for the benefits and costs of the higher-order interactions. Benefits are modelled as a power-law of the group size $g$ with an exponent $\beta$. Costs are described by an exponential decrease in the group size tuned by the cost parameter $\gamma$. Different functions of $g$ might also provide a satisfactory fit of the data. Here we have opted for this expression because it enables to factorize the group size dependence into two different contributions, benefits and costs, that can be interpreted in terms of behaviors of the players. The benefits grow as $g^\beta$, where $\beta$ captures the synergistic effect of group interactions. The term due to the cost associated to task organization in groups has
its maximum at $g = 1$, and the exponential dependence has been adopted to avoid possible singularities of other functional forms at $g = 1$. In conclusion, Eq. (6) has a maximum at $g = \beta/\gamma$, which summarizes the result in a compressed expression. To extract the pair of parameters $(\beta, \gamma)$ for each journal, we have explored the parameter space and performed an optimisation in order to reproduce the empirical points correctly. For each considered pair $(\beta, \gamma)$, we have computed the normalised distance between the synergy factor inferred analytically and the one associated to the data (see Equation (13) for further details on the procedure). The pairs with the smallest distance are selected as the outcome of the optimisation process and are those reported in Fig. 5c.

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Author contributions
U. A.-R, F. B. and V. L. conceived the study with contributions from G. F. A, M. P and Y. M. U. A.-R performed the calculations, U. A.-R, F. B., G. F. A, M. P, Y. M. and V. L. analysed the data and discussed the results. U. A.-R, F. B., G. F. A, M. P, Y. M. and V. L. wrote the paper.

Competing interests
The authors declare no competing interests.

Data availability
The APS dataset is provided by the APS at their website: https://journals.aps.org/datasets.

Code availability
Custom code that supports the findings of this study is available from the corresponding author upon request.
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Appendix

Game Implementations

Graph Implementation In the original graph implementation, or GI, the core idea is to derive the higher-order structure from the network of players. In the GI every node constructs its own focal hyperlink, whose constituents are the node itself and its first neighbours. Therefore, for a system of \( N \) players the GI imposes \( N \) different groups, each of them of \( g = k_i^2 + 1 \) members. A micro-step in the game is played by following a sequence of steps: Firstly a node, and one of its neighbours, say \( n_i \) and \( n_j \), are randomly selected from \( N \). Secondly, the nodes play a round of the game for all the groups they belong to, accumulate their payoffs, \( \pi_i \) and \( \pi_j \), and normalize them with the number of games they have played. As there is a group for each player every node will play \( k_i^2 + 1 \) rounds. The third step is the update of the strategy, and for that purpose we employ the replicator update rule. According to this, \( n_i \) will adopt the strategy of \( n_j \) with probability \( \frac{1}{\Delta} (\pi_j - \pi_i) \theta(\pi_j - \pi_i) \), where \( \Delta \) is the absolute value of the maximal payoff difference for all the possible strategies. It is noteworthy to mention that the replicator update rule is not the only update mechanism, but it is the one that we have selected because it provides an effective and noisy payoff oriented update rule by means of a simple analytic expression.

Bipartite Implementation The Bipartite Implementation (BI) lies at an intermediate point between the GI and the HI. On the one hand, and like in the HI and unlike in the GI, each round of BI occurs in a hyperlink of a hypergraph, which makes the implementation consistent with the structure of higher-order interactions present in the system. On the other hand, the update process does not benefit from the simultaneous interaction with multiple players, as each node updates its strategy by comparing its payoff with a randomly selected neighbour. The algorithm would be as follows: at each step of the game a node, one of its hyperlinks and a neighbour belonging to this hyperlink are randomly selected. Both nodes play a round of the game for each hyperlink they are part of. The original node updates its strategy by comparing its payoff with the one of its randomly selected neighbour.

We have carried out a series of numerical simulations in an ensemble of hypergraphs with 1000 nodes for different values of the group size \( G \), see Figure 6. We first observe that the critical point is the same as in the HI, meaning that the asymptotic emergence of the defectors or cooperators is the same in both cases. However we do observe a difference in the relaxation time: even if HI and BI are comparable, proving the robustness of the procedure, BI is slower, and it exceeds the limit established by the replicator approach.
Figure 6: Public Goods Game in the Bipartite Implementation. (a) We depict the fraction of cooperators as a function of the synergy factor for a system with $N = 1000$ nodes, and $L = L_C$ hyperlinks evolving during $T = 10^4$ time steps. (b) We show the relaxation time of the system as a function of the synergy factor for a hypergraph with $N = 1000$ nodes and $L = 5L_C$ hyperlinks.

Uniform Random Hypergraphs

Creation Algorithm

We provide here an example of the hyperlink counting process. Let us suppose that we have a system with $N = 5$ and $g = 3$, meaning that we could create a hypergraph of a potential number of $C_5^3 = 10$ hyperlinks, given by

$$L = \{123, 124, 125, 134, 135, 145, 234, 235, 245, 345\}$$

The mechanism to assign a random natural number in $[1, C_5^N]$ to each hyperlink is to make use of the star decomposition. Following with our example, we have $C_3^5 = C_3^2 + C_2^3 + C_3^2$, which enables us to make a first partition on the g-star hypergraphs: $\{123, 124, 125\}, \{134, 135\}, \{145\}, \{234, 235, 245\}$ and $\{345\}$. We can recursively apply our procedure, and make another partition in the subgraphs that we have here, for instance $\{123, 124, 125\}, \{134, 135\}$ and $\{145\}$ conform the first set of the previous step. These lists of combinatorial numbers may be approximated with a continuous function for better handling them. As discussed in the methods section, we propose to use $c_i \sim 1 - (1 - x)^g$ as an approximation for the cumulative probability, which gives $d_i \propto g(1 - x)^{g-1}$ for the ordinary probability. In Figure 7, we show the convergence between the continuous approximation and the real discrete function. These formulas are crucial for carrying out the PRH algorithm.

Replicator Dynamics for URH

Let us introduce now the replicator model for predicting the dynamics of the system. This technique is based on the indistinguishability between different
Figure 7: Combinatorial Approximation. The coloured lines show (a) the probability, and (b) the cumulative probability of the combinatorial summatory. The shaded area corresponds to the continuous approximation. The variable $x$ is the continuous equivalent of the discrete $i$ which allows us to associate each term in the summatory to a rational number in $[0, 1]$.

nodes, and therefore it should describe the system as long as the hypergraph is connected and the hyperlinks are uniformly distributed among the nodes. We are working with URH, and therefore the more connected the system is, the better it fulfills the uniform requirement, and the closer it gets to the replicator description.

In this simplified analysis we will describe the system in terms the fraction of defectors $x_D$ and the fraction of cooperators $x_C$. We compute the average normalized payoff of defectors and cooperators $\pi_D$ and $\pi_C$, as a function of their fractions, $x_D$ and $x_C$, in a hyperlink of order $g$. In order to do so, we sum the payoff of all the possible configurations of $g - 1$ nodes. The sum is taken over $g - 1$ because one of the nodes is already occupied by a defector, in $\pi_D$, or a cooperator in $\pi_C$. The payoff is calculated with the reduced synergy factor $r$.

$$\pi_D = \sum_{i=0}^{g-1} \binom{g-1}{i} x_D^{g-1-i} x_C^i r$$
$$\pi_C = \sum_{i=0}^{g-1} \binom{g-1}{i} x_D^{g-1-i} x_C^i ((i + 1)r - 1)$$

The total average payoffs, are obtained by multiplying the ones of Equation (7) with the average hyperdegree. But, we also normalize the total payoff with the hyperdegree, so the terms that go into the dynamical equation are indeed $\pi_D$ and $\pi_C$. One can observe that the expression for $\pi_D - \pi_C$ does not contain an
explicit dependence on \( g \), and therefore the same formula is valid for any order.

\[
\pi_D - \pi_C = \sum_{i=0}^{g-1} \binom{g-1}{i} x_D^{g-1-i} x_C^i [ir - ((i + 1)r - 1)]
\]

\[
= (1 - r) \sum_{i=0}^{g-1} \binom{g-1}{i} x_D^{g-1-i} x_C^i.
\]

\[
= (1 - r)(x_D + x_C)^{g-1} = 1 - r
\]  

(8)

We now compute \( \Delta \), the maximal payoff difference to be used in the replicators update rule. The \( \pi^+ \) and \( \pi^- \) indicate the maximal and minimal payoff values from all the possible configuration of strategies.

\[
\Delta \equiv \begin{cases} 
\pi_D^+ - \pi_C^- & \text{if } \pi_D^+ - \pi_C^- > \pi_C^+ - \pi_D^- \\
\pi_C^+ - \pi_D^- & \text{if } \pi_D^+ - \pi_C^- < \pi_C^+ - \pi_D^-
\end{cases}
\]

\( \Delta \) is only expressed for comparisons of payoffs of different strategies, since the strategies are copied from the neighbours, and therefore the terms \( \pi_i^+ - \pi_i^- \), where two nodes share the same strategy, are irrelevant. For any order \( g \), \( \Delta \) is given by

\[
\Delta = \begin{cases} 
r(g - 2) + 1 & \text{if } r < 1 \\
gr - 1 & \text{if } r > 1
\end{cases}
\]  

(9)

We now work with \( Q \), the normalized payoff difference, \( Q \equiv (\pi_D - \pi_C)/\Delta \).

The system equation is derived as the sum of all the channels via which the strategy of a node can change. These channels correspond to the different configurations of strategies of hyperlinks of order \( g \), weighted with the probabilities of the nodes having a particular strategy. For each configuration, one has to compute the probability of having a defector-cooperator pair, \( w_D x_C / g(g - 1) \) and the probability that this pair results in a strategy change. Notice that the values of \( w_D \) and \( w_C \), the number of defectors and cooperators in the group, are precisely the exponents of \( x_C \) and \( x_D \). We employ the replicator update, which in this context means that the flip probability is computed as \(-Q\theta(\pi_C - \pi_D)\) when a defector changes into a cooperator and as \(Q\theta(\pi_D - \pi_C)\) in the opposite case. This sign corresponds to the calculation of the fraction of defectors, the contrary holds true for the fraction of cooperators.
\[ \partial_t x_D = \sum_{i=0}^{G-2} \left( \frac{g}{1+i} \right) x_D^{g-1-i} x_C^{1+i} \left[ -\left( -Q\theta(\pi_C - \pi_D) \right) + Q\theta(\pi_D - \pi_C) \right] \]

\[ = \sum_{i=0}^{G-2} \left( \frac{g}{1+i} \right) x_D^{g-1-i} x_C^{1+i} \frac{Q(g - 1 - i)(1 + i)}{g(g - 1)} \]

\[ = \sum_{i=0}^{G-2} Qx_D^{g-1-i} x_C^{1+i} \frac{g!(g - 1 - i)(1 + i)}{(g - 1 - i)!(1 + i)!g(g - 1)} \]

\[ = \sum_{i=0}^{G-2} Qx_D^{g-1-i} x_C^{1+i} \frac{(g - 2)!}{(g - 2 - i)!!} \]

\[ = Qx_D x_C \sum_{i=0}^{G-2} \left( \frac{g - 2}{i} \right) x_D^{g-2-i} x_C^i = Qx_D x_C(x_D + x_C)^{g-2} = Qx_D x_C \]

This is precisely the replicator equation which provides a very useful tool for describing the system since its structure does not explicitly depend on \( g \), even if the evolution itself does, since \( Q \) is a function of \( g \).

\[ \partial_t x_D = Qx_D x_C \]

\[ \partial_t x_C = -Qx_D x_C \quad (10) \]

The stationary condition yields three possible outcomes, \( x_D = 0, x_C = 0 \) and \( Q = 0 \). The first two are the trivial phases of the system, and the third one denotes the critical point, that corresponds to \( r = 1 \). Notice that this condition is reduced to finding the zeros of Equation (8), and is therefore common for all systems, and independent of their order. This implies that for random uniform hypergraphs cooperators only emerge when \( R > g \).

If the initial condition is given by a uniform distribution of strategies, \( x_D = x_C = 0.5 \), the time evolution reads

\[ x_D(t) = \frac{1}{1 + e^{Qt}} \]

\[ x_C(t) = \frac{1}{1 + e^{-Qt}} \]

Even if the relaxation time is infinite, we can get the time to arrive the neighbourhood of the asymptotic state by imposing the condition \( x_i = 1/N \) or \( x_i = (N - 1)/N \) that yield

\[ T = \frac{\ln(N - 1)}{|Q|} \quad (11) \]

**Hyperdegree-Heterogeneous Random Hypergraphs**

**PRH and SRH**

In this subsection we report a statistical analysis about the average hyperdegree distribution obtained when creating the hypergraph using the PRH and SRH.
algorithms explained in the Methods section of the main text. We supply numerical evidence about the nature of the PRH and SRH, see Figure 8. For the former the hyperdegree distribution has two clear phases, sharp decay and an intermediate flatter regime. This combination enables the existence of hubs with a maximum hyperdegree of $1.5$ orders of magnitude larger than in the uniform case. For the latter the hyperdegree distribution follows a power law, visualized as a straight line in the log-log scale. The hubs created using the SRH have a hyperdegree $3$ orders of magnitude larger than those in the uniform case.

Figure 8: Hyperdegree-Heterogeneous Distributions. We show the value of $p_k$ the probability of having hyperdegree $k$ as a function of the value of the hyperdegree $k$ in a logarithmic scale. In the first and second row we plot the results for Power Random Hypergraphs and Scale-Free Random Hypergraphs respectively. Each of the columns is associated to $g = 2, 3, 4, 5$. In each of the subplots we plot the distributions obtained for $\mu = 0, 0.25, 0.5, 0.75, 1$. The simulations have been performed over ensembles of Hypergraphs with $N = 1000$ nodes and $L = L_c$ hyperlinks.

Replicator Dynamics for Hyperdegree-Heterogeneous Hypergraphs

In this subsection we provide theoretical arguments that clarify the results of the numerical simulations. More precisely, we are looking for the conditions that allow hypergraphs with a heterogeneous hyperdegree distribution to behave according to the dynamics derived for the uniform case. In order to do so, we are going to repeat all the steps in the formalization of the mean-field dynamics introducing an explicit dependence on the hyperdegree, such that the fraction of defectors and cooperators are expressed in terms of the corresponding fractions
conditioned to the nodes hyperdegree

\[ x_D = \sum_{k \in K} p(k)p(D|k) \]

\[ x_C = \sum_{k \in K} p(k)p(C|k) \] (12)

Here \( K \) denotes the set of all possible hyperdegrees and \( k \) an element of that set.

We first notice that the value of \( \Delta \) is the same as in the uniform case, as \( \Delta \) is a function of the maximal and minimal possible payoffs, which are the same. We continue with the proposal of the dynamical equation. The time evolution of Equation (12) can be expressed as a sum of each of the \( p(D|k) \) terms, that for clarity we relabel as \( x_{Dk} \). There are two possible transitions in which the fraction of defectors with hyperdegree \( k \), \( x_{Dk} \) can change: a cooperator with hyperdegree \( k \) changes into a defector of hyperhyperdegree \( k \), and a defector of hyperdegree \( k \) changes into a cooperator of hyperdegree \( k \). The dynamics accounts for these two contributions as follows:

\[
\frac{d}{dt} x_{Dk} = -x_{Dk} \sum_{k' \in K} p(k'|k)p(C|k') \sum_{k'' \in K^{G-2}} p(k''|kk') \sum_{x \in X^{G-2}} p(x|k'') \times \\
\frac{(\pi_{Ck'} - \pi_{Dk'})\theta(\pi_{Ck'} - \pi_{Dk'})}{\Delta} \\
+ x_{Ck} \sum_{k' \in K} p(k'|k)p(D|k') \sum_{k'' \in K^{G-2}} p(k''|kk') \sum_{x \in X^{G-2}} p(x|k'') \times \\
\frac{(\pi_{Dk'} - \pi_{Ck})\theta(\pi_{Dk'} - \pi_{Ck})}{\Delta}
\]

The variable and first summatory in both terms account for the probability of having the defector and cooperator pair that will take part in the strategy update. In order to ensure that both are in the same hyperlink we introduce the hyperdegree-hyperdegree correlation to account for the probability that the hyperdegree of the player inspiring the strategy update is \( k' \), given that the player changing is strategy has hyperdegree \( k \). The next term corresponds to the remaining \( g-2 \) nodes of the hyperlink, and thus, \( k'' \) represents \( G-2 \) elements of \( K \). Again the hyperdegree-hyperdegree correlation provides the information about the conditional probability of having \( g-2 \) nodes with hyperdegrees \( k'' \) given that the remaining two nodes have hyperdegrees \( k \) and \( k' \). In the next term, the summation in \( x \), \( x \) accounts for \( G-2 \) elements of \( X \), where \( X \) is the set of all the possible strategies, either \( D \) or \( C \). The last term contains the comparison of the payoffs, that will either increase or decrease the amount of defectors. Following the same strategy as in the uniform case, we will have a closer look at the payoff comparison. The \( \Delta \) contribution is solved already, so
we focus on the numerator.

\[
\pi_{Dk} = \sum_{k'' \in K^{G-1}} p(k'' | k) \sum_{x'' \in X^{G-1}} p(x'' | k'')(nr)
\]

\[
\pi_{Ck'} = \sum_{k'' \in K^{G-1}} p(k'' | k') \sum_{x'' \in X^{G-1}} p(x'' | k'')((n+1)r - 1)
\]

Here \( n \) is the number of cooperators of each particular case of the summatory, which is of course a function of \( x'' \).

The important terms are \( p(k'' | k) \) and \( p(k'' | k') \), those that account for the hyperdegree-hyperdegree distribution, i.e., the probability that a node with a given hyperdegree \( k \) or \( k' \) is part of a hyperlink with nodes of hyperdegrees given by \( k'' \). For hypergraphs in which the neighbouring hyperdegree distribution \( p(k'' | k) \) is given by the total hyperdegree distribution \( p(k'') \), this expression can be reduced to

\[
\pi_{Dk} = \sum_{k'' \in K^{G-1}} p(k'') \sum_{x'' \in X^{G-1}} p(x'' | k'')(nr)
\]

\[
\pi_{Ck'} = \sum_{k'' \in K^{G-1}} p(k'') \sum_{x'' \in X^{G-1}} p(x'' | k'')((n+1)r - 1)
\]

and then

\[
\pi_{Dk} - \pi_{Ck'} = \sum_{k'' \in K^{G-1}} p(k'') \sum_{x'' \in X^{G-1}} p(x'' | k'')((nr) - ((n+1)r - 1))
\]

\[
= \sum_{k'' \in K^{G-1}} p(k'') \sum_{x'' \in X^{G-1}} p(x'' | k'')(1 - r)
\]

\[
= (1 - r)
\]

This is indeed the same result as in the uniform case, \( Q = \frac{1 - r}{x} \), an expression that we can now use in the main derivation to simplify the equation

\[
\frac{d}{dt} x_{Dk} = -x_{Dk}(-Q\theta(r - 1)) \sum_{k'' \in K} p(k' | k) p(C | k') \sum_{k''' \in K^{G-2}} p(k'' | kk') \sum_{x'' \in X^{G-2}} p(x | k'')
\]

\[
+ x_{Ck} Q\theta(1 - r) \sum_{k'' \in K} p(k' | k) p(D | k') \sum_{k'''' \in K^{G-2}} p(k''' | kk') \sum_{x'' \in X^{G-2}} p(x | k''')
\]

\[
= Q\theta(r - 1)x_{Dk}x_{C} + Q\theta(1 - r)x_{Ck}x_{D}
\]

In the second step we have make use of the assumption that there are not hyperdegree-hyperdegree correlations, \( p(k' | k) = p(k') \). We then insert this equation into Equation (12) which yields

\[
\frac{d}{dt} x_{D} = Q\theta(r - 1)x_{D}x_{C} + Q\theta(1 - r)x_{C}x_{D} = Qx_{D}x_{C}
\]

The dynamics is equivalent to the one derived in the uniform case. The interpretation is the following: when the nodes are indistinguishable in the sense that their neighbourhoods are equivalent the mean-field dynamics is still valid even if the hypergraphs are heterogeneous.
Order-Heterogeneous Random Hypergraphs

Replicator Dynamics for Order-Heterogeneous Hypergraphs

The first thing to notice is that the average payoff difference, is nothing but the sum of the average payoff differences for each of the orders, weighted with the $p^g$. The sum goes from $g^-$, the minimal value of $g$ to $g^+$, the maximal one.

$$\pi_D - \pi_C = \sum_{g=g^-}^{g^+} p^g (1 - r^g)$$

We first derive the normalization factor $\Delta$, from Equation (9), via the explicit expression for the reduced synergy factor $r^g = \alpha g^{\beta - 1}$. We consider $\alpha, \beta \geq 0$.

The extreme point of $r^g$ at $\beta = 1$ divides the analysis in two different intervals, $\beta < 1$ and $\beta \geq 1$.

In the first one, $\beta < 1$, the maximal payoffs $\pi^+$ are obtained for the lowest order $g^-$, and the minimal payoffs $\pi^-$ for the highest order $g^+$.

$$\pi_D^+ = \alpha g^{\beta - 1} (g^- - 1), \quad \pi_D^- = 0, \quad \pi_C^+ = \alpha g^{\beta} - 1, \quad \pi_C^- = \alpha g^{\beta - 1} - 1$$

Therefore, we get

$$\Delta(\beta < 1) = \begin{cases} \alpha g^{\beta - 1} (g^- - 1) - \alpha g^{\beta - 1}_+ + 1 & \text{if } \alpha \leq \frac{2}{g^- + g^+_+} \\ \alpha g^{\beta - 1}_+ - 1 & \text{if } \alpha > \frac{2}{g^- + g^+_+} \end{cases}$$

In the second interval, $\beta \geq 1$, the maximal payoffs are obtained for the maximal orders, and the minimal payoffs for the minimal orders.

$$\pi_D^+ = \alpha g^{\beta - 1}_+ (g^+ - 1), \quad \pi_D^- = 0, \quad \pi_C^+ = \alpha g^{\beta}_+ - 1, \quad \pi_C^- = \alpha g^{\beta - 1}_+ - 1$$

which yields,

$$\Delta(\beta \geq 1) = \begin{cases} \alpha g^{\beta - 1}_+ (g^+ - 1) - \alpha g^{\beta - 1}_- + 1 & \text{if } \alpha \leq \frac{2}{g^- + g^+_+} \\ \alpha g^{\beta}_+ - 1 & \text{if } \alpha > \frac{2}{g^- + g^+_+} \end{cases}$$

With $\pi_D - \pi_C$ and $\Delta$, we can obtain $Q$, as in the uniform case. And in terms of $Q$ the differential equation for the time evolution is exactly Equation (10). Therefore, the relaxation time is again given by Equation (11), and the critical point is again obtained by making $\pi_D - \pi_C = 0$, which yields

$$\alpha_c = \frac{1}{\sum_{g=g^-}^{g^+} p^g g^{\beta - 1}}$$
Hypergraphs describing real-world collaborations

**IETF Dataset.** In this last section, we provide an additional practical example of the procedure for extracting the synergy factor from real data, based on the publication records of the Internet Engineering Task Force (IETF). This is a large open international community of network designers, operators, vendors, and researchers concerned with the evolution and the smooth operation of the Internet. Similarly to the APS journals, the system is described by a hypergraph where authors who participated together in a project are part of the same hyperlink. For the IETF dataset, we have 34280 articles. In Figure 9 we report the histogram of the number of articles with a given number of authors. The normalized version of this histogram is precisely the hyperlink distribution as a function of the order of the hyperlink. Next, we retrieve the hyperdegree-order distribution by assuming that the hyperlinks are equally distributed amongst the nodes, meaning that for each order \( g \) the hyperdegree can be deduced from the number of hyperlinks as \( k_g = gL^g/N \). We assume that the reduced synergy factor is proportional to the hyperdegree distribution. This assumption means that we need an additional parameter to provide the full expression of the reduced synergy factor (and not the normalized one). This parameter can be obtained by introducing the critical point condition \( \sum_{g=g^+}^g \rho^g (1 - r(g)) = 0 \). Notice that \( \rho^g \) is the \( g \)th component of the hyperdegree, and that \( r(g) \) is the synergy factor we are looking for, which can be expressed as \( z \rho^g \) under the aforementioned assumption. This provides a simple equation to obtain \( z \), \( z \sum (\rho^g)^2 = 1 \), which completes the procedure for extracting \( r(g) \).

For this particular dataset, for completeness we fit the experimental data with different analytical expressions. We numerically find those parameters that minimize the distance with either \( R(g) \) or \( r(g) \). To validate the fit, the measure we have selected is the distance between the normalized points of the analytical and the experimental synergy factor, which is bounded to the \([0, 1] \) interval. Therefore, values close to 0 account for a good estimation of the data, otherwise the opposite is true. To extract the parameters we have performed a search process by limiting the possible values to meaningful regions of the parameter space, and by discretizing this region into a finite set values. Then, we iterated over all the sets, one for each parameter, and calculated the normalized distance between the analytical expression and the experimental one for each iteration. The best results that we have obtained in Figure 9 are given by

\[
\begin{align*}
e_1 : r(g) &= \alpha g^\beta e^{-\gamma (g-1)} \quad (\beta, \gamma) = (1.382, 0.696) \quad (d_r, d_R) = (0.054, 0.090) \\
e_2 : R(g) &= \alpha g^\beta e^{-\gamma (g-1)} \quad (\beta, \gamma) = (2.89, 0.865) \quad (d_r, d_R) = (0.065, 0.082) \\
e_3 : r(g) &= \alpha g^\beta e^{-\gamma (g-1)^\delta} \quad (\beta, \gamma, \delta) = (0.5, 0.12, 1.7) \quad (d_r, d_R) = (0.039, 0.079) \\
e_4 : R(g) &= \alpha g^\beta e^{-\gamma (g-1)^\delta} \quad (\beta, \gamma, \delta) = (1.51, 0.13, 1.65) \quad (d_r, d_R) = (0.041, 0.077)
\end{align*}
\]

Here \( e_i \) is associated to the expressions as reported in the figure, \( \alpha \) is obtained analytically by applying the critical point condition and \( d_r \) and \( d_R \) the distance...
between the approximation and the points inferred from the data. Notice that the \((e_1, e_2)\) and \((e_3, e_4)\) pairs of equations have the same structure, but their parameters have been obtained by minimizing the distance with \(r(g)\) and \(R(g)\), respectively. As before, \(\beta\) and \(\gamma\) are the benefits and costs fit parameter, that respectively grow as a power law and decrease exponentially.

Figure 9: Synergy factors of the Internet Engineering Task Force dataset. We extract the synergy factor as a function of the group size for a bibliographic dataset in technology that contains the number of publications as a function of the number of authors \(g\). We infer the distribution in the order of the hyper-degrees from the original dataset by assuming that the hyperlinks are evenly distributed amongst the nodes. We then impose the critical point condition and extract the value of the synergy factor under the hypothesis that \(r(g)\) is proportional to \(p\). Finally, the synergy factors \(r(g)\) in (a) and \(R(g)\) in (b) are factorized into two analytical expressions that respectively account for the benefits of large cooperations, and the costs associated to saturation effects.

**APS Dataset.** Here we provide some additional details about the dataset of scientific collaborations from publications in the journals of the American Physical Society described in the main text. In Table 1 we report the main properties of the hypergraph describing each journal.

We clarify here the optimization method for \(\beta\) and \(\gamma\). Once we have selected

\[
f(g, \alpha, \beta, \gamma) = \alpha g^\beta e^{-\gamma(g-1)}
\]

as the base equation to fit the data, we run a first coarse-grained analysis on the possible minimal and maximal limits of \(\beta\) and \(\gamma\). These limits account for the values outside which the approximation does not resemble the data. We then discretize the domain of \(\beta\) and \(\gamma\), and carry out a brute-force optimization, namely, we evaluate a distance \(\sigma(r, f)\) for all the possible combinations of \(\beta\) and \(\gamma\) and select those with the smallest value. The function \(\sigma(r, f)\) is bounded to the \([0, 1]\) interval, and measures the distance between the normalized synergy factor.
and the normalized approximation $f$, calculated as $\sigma = 1 - \sum_g \text{min}(r(g), f(g))$.

We also provide here the benefit and cost parameters associated to each journal, as well as the normalized error between the maximum synergy factor extracted directly from the data and from the analytical estimate. In Figure 10 we show the experimental and approximated reduced synergy factors $r(g)$ as a function of all hyperlink orders $g$ contained in the publications dataset. This figure is an extended version of Fig.4(a) in the main dataset.

| Journal           | $L$  | $<g>$ | $g_{(\text{max} r)}$ | $\beta$ | $\gamma$ | $d_r$ |
|-------------------|------|-------|----------------------|---------|----------|-------|
| PhysRev           | 47313| 1.95  | 2                    | 2.936   | 1.573    | 0.033 |
| PhysRevA          | 70502| 3.07  | 3                    | 2.679   | 0.986    | 0.067 |
| PhysRevB          | 171268| 3.75  | 3                    | 1.531   | 0.49     | 0.05  |
| PhysRevC          | 36290| 5.98  | 3                    | 0.02    | 0.075    | 0.146 |
| PhysRevD          | 74715| 3.02  | 2                    | 2.178   | 0.941    | 0.206 |
| PhysRevE          | 50988| 2.93  | 3                    | 3.84    | 1.41     | 0.048 |
| PhysRevApplied    | 327  | 5.39  | 5                    | 3.356   | 0.62     | 0.09  |
| PhysRevLett       | 113674| 4.57  | 3                    | 0.848   | 0.33     | 0.175 |
| PhysRevSeriesI    | 1240 | 1.21  | 1                    | 2.691   | 2.831    | 0.019 |
| PhysRevSTAB       | 2393 | 5.52  | 4                    | 0.566   | 0.173    | 0.127 |
| PhysRevSTPER      | 484  | 2.42  | 3                    | 2.75    | 1.21     | 0.078 |
| PhysRevX          | 611  | 5.28  | 5                    | 1.85    | 0.416    | 0.127 |
| RevModPhys        | 3153 | 2.05  | 2                    | 1.19    | 0.79     | 0.112 |

Table 1: American Physical Society Dataset. For each journal we report the total number of hyperlinks $L$, the average hyperlink order $<g>$, the order associated to the maximal synergy factor $g_{(\text{max} r)}$, the benefit parameter $\beta$ and the cost parameter $\gamma$ (from the analytical approximation of $r(g)$), and the normalized distance between the reduced synergy factor extracted from the data empirically and after the fit, that we indicate as $d_r$.  

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Figure 10: Synergy factors of the American Physical Society dataset. Value of the reduced synergy factor $r(g)$ for all the values of the group size $g$ available in the dataset. These plots are an extended version of Fig.4 shown in the main manuscript.