Successive whole genome duplications have recently been firmly established in all major eukaryote kingdoms. It is not clear, however, how such dramatic evolutionary processes have contributed to the large scale topology of protein-protein interaction (PPI) networks. We propose and analyze a generic model of PPI network evolution under successive whole genome duplications. This demonstrates that the observed scale-free degree distributions and conserved multi-protein complexes may have concomitantly emerged from (i) intrinsic exponential dynamics of PPI network evolution and (ii) asymmetric divergence of gene duplicates. This requirement of asymmetric divergence is in fact "spontaneously" fulfilled at the level of protein-binding domains. In addition, domain shuffling of multi-domain proteins is shown to provide a powerful combinatorial source of PPI network innovation, while preserving essential structures of the underlying single-domain interaction network. Finally, large scale features of PPI networks reflecting the "combinatorial logic" behind direct and indirect protein interactions are well reproduced numerically with only two adjusted parameters of clear biological significance.

Gene duplication is considered the main evolutionary source of new protein functions. Although long suspected, whole genome duplications have only been recently confirmed through large scale comparisons of complete genomes.

Whole genome duplications are rare evolutionary transitions followed by random nonfunctionalization of most gene duplicates on time scales of about 100MY (with large variations between genes, see discussion). Whole genome duplications presumably provide unique opportunities to evolve many new functional genes at once through accretion of functional domains from contiguous pseudogenes (or redundant genes) and may also promote speciation events by preventing genetic recombinations between close descendants with different random deletion patterns.

Recent whole genome duplications (WGDs) within the last 500MY (about 15% of life history) have now been firmly established in all major eukaryote kingdoms. For instance, there are 4 consecutive WGDs between the sea squirt Ciona intestinalis and the common carp Cyprinus carpio, with most tetrapods (including mammals) in between at +2WGDs from sea squirt and −2WGDs from carp and most bony fish at +3WGDs from sea squirt and −1WGDs from carp (a pseudotetraploid bony fish duplicated about 10MY ago). There are also 3 consecutive WGDs in the recent evolution of the flowering plant Arabidopsis thaliana and at least 3 consecutive WGDs for the protist Paramecium tetraurelia (Patrick Wincker, personal communication). Extrapolating these 500MY old records, one roughly expects a few tens consecutive WGDs (or equivalent "doubling events") since the origin of life. These rare but dramatic evolutionary transitions must have had major consequences on the evolution of large biological networks, such as protein-protein interaction (PPI) networks.

From a theoretical point of view, we also expect that alternating whole genome duplications and extensive gene deletions lead to exponential dynamics of PPI network evolution. In the long time limit, this should outweigh all time-linear dynamics that have been assumed in PPI network evolution models under local structure changes (see discussion). In fact, the intrinsic exponential dynamics of genome evolution is already transparently reduced from the wide distribution of genome sizes and proliferation of repetitive elements: it is hard to imagine that the 10^4-fold span in lengths of eukaryote genomes could have solely arisen through time-linear increases (and decreases) in genome sizes.

**Modelling PPI network evolution by whole genome duplication**

We propose a simple model of PPI network evolution focussing on whole genome duplications (extensions to local or partial genome duplication are presented in ref. and confirm the conclusions of this paper). Each time step corresponds to a whole genome duplication and leads to a complete duplication of the PPI network, whereby each node is duplicated (×2) and each interaction quadrupled (×4) as depicted in Fig.1. Links from the duplicated network are then kept with different probabilities γ (0 ≤ γ ≤ 1) reflecting symmetric or asymmetric divergences between protein or link copies.

The interaction network is characterized at each step n by its number of nodes with k neighbours N(n) and its total number of links L(n) = ∑k≥1 kN(n)/2. As stochastic differences exist between network realizations, we study the evolution of typical networks by introducing a generating function averaged over all network realizations,

\[ F^{(n)}(x) = \sum_{k=0}^{\infty} (N^{(n)}(k)x^k). \]  

This use of generating functions can in fact be generalized to other, possibly non local features of interest (e.g. the average connectivity of first neighbors, \( g_{first} \) is introduced below).

In the following, we discuss a general model of PPI network evolution through whole genome duplication with asymmetric divergence of duplicated genes (Figs.1&2A). We compare it, first, to an alternative model with symmetric protein divergence but random link "complementation" (Fig.1B), and also to direct physical interactions from Yeast PPI network data (Fig. 2B&C). We then redefine this initial asymmetric divergence model (Fig. 1) in terms of protein-binding domains (Figs. 3A&B) to account for indirect protein-protein interaction within multi-protein complexes (Figs. 3A&C).

**Asymmetric divergence of duplicated proteins**

The case of asymmetric divergence between duplicated genes corresponds to the following evolution scenario: while duplicated proteins are initially equivalent and experience, at first, the same functional constraints, their divergence becomes eventually asymmetric (see discussion). This presumably occurs...
Once one duplicate copy has lost an essential interaction and thus function, which has then to be fulfilled entirely by the other duplicate. The evolution of this latter duplicate is, from then on, more constrained to retain “old” interactions, while the former duplicate is left largely free to accumulate more neutral mutations with the likely outcome to become nonfunctional and eventually deleted unless some “new”, duplication-derived interactions are selected; “Old” duplicates, on the other hand, are more constrained to conserve “old” interactions already present before duplication. The duplicated network with quadruplicated links is graphically rearranged for convenience into old and new network copies (e.g. 2 and 2′ duplicated nodes are swapped here). Links from the duplicated network are then kept with different probabilities e.g. δ,o new and δ,o old each link emerging from each node of the duplicated graph. Hence, the generating function recurrence for PPI network evolution as follows: since each node is initially duplicated, \( p^{(n+1)}(x) \) is the sum of two \( p^{(n)}(x) \) where \( x \) is first replaced by \( x^2 \) (since each node degree can at most double) and then substituted as \( x \rightarrow \gamma x + \delta \), where \( \gamma_i \) [resp. \( \delta_i = 1 - \gamma_i \)] corresponds to the probability to keep [resp. delete] each link emerging from each node of the duplicated graph. Hence, the generating function recurrence for PPI network evolution with asymmetric divergence of duplicated proteins yields:

\[
F^{(n+1)}(x) = p^{(n)}\left((\gamma x + \delta)(\gamma_0 x + \delta_0)\right) + F^{(n)}\left((\gamma x + \delta)(\gamma_0 x + \delta_0)\right).
\]

where \( \gamma, \gamma_0 \) and \( \delta, \delta_0 \) stand for \( \gamma_{\text{cross}}, \gamma_{\text{new}} \) and \( \gamma_{\text{old}}, \delta_{\text{cross}}, \delta_{\text{new}} \) and \( \delta_{\text{old}} \) in Fig. 1 (see supporting information for proof details).

The overall graph dynamics through successive global duplications is clearly exponential as anticipated; in particular, the total number of nodes grows as \( F^{(n)}(1) = A 2^n \), where \( A \) is the initial number of nodes, and the number of links scales as \( \langle L^{(n)} \rangle \propto (2\gamma + \gamma_0) A 2^n \). We remove permanently disconnected nodes from the list of relevant nodes, assuming that they correspond to proteins that have in fact lost their function and are eventually eliminated from the genome. To this end, we redefine the graph size as, \( \langle N^{(n)} \rangle = \sum_{k \geq 1} \langle N^{(n)}_k \rangle \) and introduce a normalized generating function \( p^{(n)}(x) \) for the mean degree distribution,

\[
p^{(n)}(x) = \sum_{k \geq 1} p^{(n)}(x)^k, \quad \text{where} \quad p^{(n)}(x) = \frac{\langle N^{(n)}_k \rangle}{\langle N^{(n)} \rangle}.
\]

Absolute and relative generating functions are related through,

\[
F^{(n)}(x) = p^{(n)}(x) \langle N^{(n)} \rangle + \langle N^{(n)}_0 \rangle.
\]

Inserting this expression 4 in recurrence 3 gives a closed relation between successive \( p^{(n)}(x) \),

\[
p^{(n+1)}(x) = 1 - 2 - p^{(n)}\left((\gamma x + \delta)(\gamma_0 x + \delta_0)\right) - p^{(n)}\left((\gamma x + \delta)(\gamma_0 x + \delta_0)\right),
\]

where \( \Delta(n) \) is the ratio between consecutive numbers of connected nodes, \( \Delta(n) = \langle N^{(n+1)} \rangle / \langle N^{(n)} \rangle = 2 - p(n) / (\delta_0 / p(n) - p(n) / \delta_0) \leq 2. \)

The evolution of the mean degree is obtained by taking the first derivative of 5 at \( x = 1 \):

\[
\partial_x p^{(n+1)}(1) = \frac{\Gamma_n + \Gamma_0}{\Delta(n)} \partial_x p^{(n)}(1),
\]

where \( \Gamma_n = \gamma + \gamma_0 \) and \( \Gamma_0 = \gamma + \gamma_0 \) hereafter.

We will limit the discussion here to degree distributions approaching a stationary regimes \( p^{(n)}(x) \rightarrow p(x) \) with a finite mean degree \( 1 \leq p(x) < \infty. \) This seems to cover the most biologically relevant networks; for completeness, other cases are discussed elsewhere.

From 6 and the condition of finite mean degree, we readily obtain that \( \Delta(n) \rightarrow \Gamma_n + \Gamma_0, \) which implies that the network evolution is asymptotically equivalent in terms of connected nodes and links,

\[
\langle N^{(n+1)} \rangle / \langle N^{(n)} \rangle \rightarrow \langle L^{(n+1)} \rangle / \langle L^{(n)} \rangle = \Gamma_n + \Gamma_0 \leq 2,
\]

The stationary degree distribution is then solution of the functional equation,

\[
p(x) = 1 - 2 - p\left((\gamma x + \delta)(\gamma_0 x + \delta_0)\right) - p\left((\gamma x + \delta)(\gamma_0 x + \delta_0)\right),
\]

which can be differentiated \( k \) times to express the \( k \)th derivative in terms of lower derivatives,

\[
\partial_x^k p(1) \left[ 1 - \frac{\Gamma_n + \Gamma_0}{\Gamma_n + \Gamma_0} \right] = \sum_{m=[k/2]}^k \alpha_m (\gamma_n, \gamma_0, \gamma) \partial_x^m p(1)
\]

where the coefficients \( \alpha_m \) are all positive from the definition 8.

The finite or infinite nature of \( \partial_x^k p(1) \) depends on the two parameters \( \Gamma_n \) and \( \Gamma_0 \) and defines the form of the limit degree distribution. The phase diagram Fig. 2A summarizes in the plane \( (\Gamma_n + \Gamma_0, \Gamma_0 - \Gamma_n) \) the different regimes for the asymptotic degree distribution \( p_k \). \( \Gamma_0 + \Gamma_n \) is the global growth rate of the network.
(Γ_o+Γ_n > 1 to ensure a growing network) and Γ_o−Γ_n corresponds to the divergence asymmetry between duplicated proteins. We now discuss the two main stationary regimes for p_k in the case of Γ_n ≤ Γ_o (the case Γ_n ≥ Γ_o is deduced by permutating indices):

- **Exponential, non-conservative regime.** If both Γ_o < 1 and Γ_n < 1,

  \[ \Gamma_n^k + \Gamma_o^k < \Gamma_n + \Gamma_o, \quad \text{for all } k \geq 2 \quad (10) \]

  and the factor in front of \( \partial_k^r p(1) \) in [B] is always strictly positive, which implies that all derivatives of the limit degree distribution are finite. Hence, in this case, the limit degree distribution decreases more rapidly than any power law (see explicit asymptotic development in [B]). Note that this “exponential” regime occurs when the links emerging from each node (Fig. 1) are more likely lost than duplicated at each round of global duplication (as Γ_i = γ + γ_i < 1 is equivalent to δ_i > γ_i). This implies that most nodes eventually disappear, and with them all traces of network evolution, after just a few rounds of global duplication. The network topology is not conserved, but instead continuously renewed from duplication of the (few) most connected nodes.

- **Scale-free, conservative regime.** If Γ_o > 1 > Γ_n, the factor in front of \( \partial_k^r p(1) \) in [B] can become negative. However, since the generating function should have all its derivatives positive, a negative value for one of them means that it simply does not exist. In fact, for Γ_o ln Γ_o + Γ_o ln ln Γ_o ≥ 0 (red line in Fig. 2A and [B]), there is an integer \( r \geq 1 \) such that

  \[ \Gamma_n^r + \Gamma_o^r \leq \Gamma_n + \Gamma_o < \Gamma_n^{r+1} + \Gamma_o^{r+1}, \quad (11) \]

  implying that all derivatives \( \partial_k^r p(1) \) are finite up to the \( r \)th order, while \( \partial_k^r p(1) \) is infinite. This justifies the following asymptotic expansion of \( p(x) \) in the vicinity of \( x = 1 \),

  \[ p(x) = 1 - A_1 (1-x) + \ldots + (-1)^r A_r (1-x)^r - A_r (1-x)^{\alpha-1} + \ldots, \]

  for some appropriate \( r < \alpha < r + 1 \). This ansatz is then inserted in [B] using \( (\gamma x + \delta)((\gamma_n, x + \delta_n, x) = 1 = \Gamma_n, (1-x) + \gamma_n, (1-x)^2 \) to obtain an equation on the coefficients \( A_1, \ldots, A_n \). The term \( A_n \) does not mix with previous terms and gives the following equation for \( \alpha \),

  \[ \Gamma_n^\alpha + \Gamma_o^\alpha = \Gamma_n + \Gamma_o. \]

  (13)

  The limit degree distribution follows a power law in this case,

  \[ p_k \propto k^{-\alpha-1}, \]

  (14)

  (see red and blue “exponent” lines in Fig. 2A for \( \alpha = 1 = 2, 3, 4, \ldots \)).

  Note that scale-free degree distributions emerge under successive, global network duplications only if the “old” node copy has its links more likely duplicated than lost at each round of global duplication (as \( \Gamma_n = \gamma + \gamma_o > 1 \) is equivalent to \( \gamma + \gamma_o > \delta \)). Thus, “old” nodes statistically keep on increasing their connectivity once they have emerged as “new” nodes by duplication. This implies that most nodes and their surrounding links are conserved throughout the evolution process, thereby ensuring that local topologies of previous networks remain embedded in subsequent networks.

  In summary, whole genome duplication with asymmetric divergence of duplicated proteins leads to the emergence of two classes of PPI networks with finite asymptotic degree distributions: i) PPI networks with an exponential degree distribution and without conserved topology and ii) PPI networks with a scale-free limit degree distribution and at least local topology conservation. All other evolution scenarios, which do not lead to finite asymptotic degree distributions, are unlikely to model biologically relevant cases; they correspond either to an exponential disappearance of the whole PPI network (i.e. if \( \Gamma_n + \Gamma_o < 1 \)) or to an exponential shift of all proteins towards higher and higher connectivities (i.e. dense regime in Fig. 2A for \( \Gamma_n, \Gamma_o > 1 \)).
Symmetric divergence of duplicates with link “complementation”

Another model of interest is the so-called “duplication-mutation-complementation” model initially proposed in the context of protein network evolution through successive local duplications. This model can be easily adapted to the context of PPI network evolution through whole genome duplication, Fig. S1. After each global duplication step, the probability to keep an instance of each interaction is now distributed randomly over the four equivalent links with out reference to particular protein duplicates, unlike in the previous model. The complementation step (which ensures that at least one instance of each previous link is retained) can be enforced here through the “old” link copy ($\gamma = 1$) with $\gamma_n$ corresponding to the “new” interaction sharing no node with $\gamma_n$, while $\gamma$ still pertains to the last two equivalent cross links. This model is thus effectively symmetric from the protein point of view and readily yields the following recurrence for the generating function of the network degree distribution.

$$F^{(n+1)}(x) = 2F^{(n)}((\gamma x + \delta)(\gamma_n x + \delta_n)),$$

where $\gamma_n = (\gamma_n + \gamma_0)/2$ and $\delta_n = (\delta_n + \delta)/2$ are effective average probabilities to retain or delete old and new links (see supporting information for proof details). Hence, the model of PPI network evolution with link complementation is in fact equivalent to the case of a symmetric divergence of duplicated proteins in the previous general model. Such symmetric divergence of duplicated proteins yields either a stationary exponential regime ($\Gamma_n = \Gamma_\alpha < 2$, Fig.2A) or a non-stationary dense regime ($\Gamma_n > 2$, Fig.2A).

Hence, the “duplication-mutation-complementation” model cannot lead to scale-free degree distributions, and thus to locally conserved network topology, in the context of whole genome duplication evolution, by contrast to the same model applied to local duplication with time-linear evolution.

Fitting PPI network data with a one-parameter model

Scale-free degree distributions have been widely reported for large biological networks and other exponentially growing networks like the WWW. We showed in the previous discussion that scale-free limit degree distributions require an asymmetric divergence of duplicated proteins ($\Gamma_n - \Gamma_\alpha = \gamma_n - \gamma_\alpha > 0$) which corresponds to the probability difference between conservation of old interactions ($\gamma_0$) and coevolution of new binding sites ($\gamma_n$). The expected range of parameters for actual biological networks is $1 \approx \gamma_0 \gg \gamma > \gamma_n \approx 0$; In particular, the most conservative ($\gamma_0 = 1$) and least correlated ($\gamma_n = 0$) evolution scenario corresponds to the strongest divergence asymmetry between duplicated proteins ($\Gamma_n - \Gamma_\alpha = 1$, upper border on Fig.2A). The condition $\gamma_0 = 1$ ensures that not only local but also global topologies of all previous networks remain embedded in all subsequent networks. This model is effectively a one-parameter model (\gamma) for PPI network evolution through whole genome duplication. It converges towards a stationary scale-free limit degree distribution $p_k \sim k^{-\gamma}$ with $0 < \alpha < 2$ for $0 < \gamma < (1/\sqrt{5} - 1)/2$ and generates non-stationary dense networks for $(1/\sqrt{5} - 1)/2 < \gamma < 1$.

We used this one-parameter model to fit both the degree distribution (Fig.2B) and the average connectivity of first neighbors (Fig.2C) for direct physical interaction data of S. cerevisiae taken from two databases, BIND and hand curated MIPS (with presumably fewer nonspecific spurious interactions).

The predicted asymptotic regime is in fact approached for $k \leq 20$ due to the finite size of yeast PPI network. The fitting parameter $\gamma = 0.26$ corresponds to a fixed growth rate of $1 + 2\gamma = 1.52$ (i.e. the number of links and nodes increases by 52% at each global duplication). Adding and removing up to 30% of links randomly, or drawing $\gamma$ from a uniform distribution between 0 and 0.52 (with average $\tilde{\gamma} = 0.26$) yield remarkably similar fits (not shown) to the experimental data. This reveals a large insensitivity to false-positive and negative noises and fluctuations in $\gamma$ (as long as the non-stationary dense regime is avoided, Fig.2A). The fixed (or averaged) growth rate of 52% at each round of global duplication is enough to generate networks of the size of S. cerevisiae starting from a few interacting “seeds” after about 20 global duplications (i.e. 1.52$^{20} = 4334$ times more nodes with an average of one global duplication per 200MY for 4BY). Such scenario is not a priori incompatible with experimental data, as we only have clear records on global duplications dating back up to 400-500MY ago (i.e. only 10 to 20% of life history). Yet, these records suggest that “recent” whole genome duplications might be more frequent (every 100-150MY) and more selective (growth rates between 10 and 25%).

Direct vs indirect protein-protein interactions

The protein-protein interactions we have considered so far correspond to direct physical contact between protein pairs derived, for instance, from two-hybrid expression assays. However, we expect from the proposed scale-free fit of the degree distribution (Fig. 2B) that the underlying PPI network has conserved not only pairwise interactions during evolution but also some level of network topology (see above). The emergence of locally conserved topology in PPI network evolution leads “naturally” to conserved associations or “modules” between multiple proteins and, beyond, to recurrent “motifs” across different types of biological networks.

In fact, many biological functions are known to rely on multiple direct and indirect interactions within protein complexes. Moreover, the combinatorial complexity of multiple-protein interactions is likely responsible for the remarkable diversity amongst living organisms, despite their rather limited and largely shared genetic background (i.e. a few (ten) thousand genes built from a few hundreds families of homologous protein domains).

High-throughput studies using affinity precipitation methods coupled to mass spectroscopy have proposed some 80,000 direct and indirect protein interactions for S. cerevisiae (raw data) and similar data are now becoming available for several other species.

Yet, from a theoretical point of view, the evolution of indirect interactions is expected to depend not only on locally conserved network topology but also on the actual “combinatorial logic” between direct interactions. This cannot be readily defined on traditional PPI network representation (e.g. Fig. 1) and requires a somewhat more elaborated model as we now discuss.

Redefining PPI network evolution in terms of protein domains

Indirect protein interactions reflect the occurrence of simultaneous direct interactions within protein complexes. This requires that some proteins have more than one binding sites to simultaneously interact with several protein partners. Indeed, proteins with a single protein-binding site can only bind to one partner at a time, underlying a simple “XOR”-like combinatorial logic. By contrast, proteins with several protein-binding sites (which are usually multi-domain proteins) greatly increase the combinatorial complexity of biological processes (like gene regulation or cell signaling) by adding “AND” operators to the computational logic between multiple direct interactions. Multi-domain proteins also provide a versatile support for protein evolution through accretion or deletion of individual domains.

In addition, we note that binding sites on specific protein domains are likely the primary source of asymmetric divergence in PPI network evolution, as binding site mutations necessarily affect interactions with all binding partners (Fig. 1) and not just a random subset of them (Fig. S1). Hence, asymmetric divergence of protein duplicates “naturally” originates from “spontaneous symmetry breaking” of their equivalent protein-binding sites (or domains).

We propose to highlight this central role of protein domains in the evolution of PPI networks by simply redefining our initial asymmet-
Generating function depletion events leads to virtually identical results for the large scale elaborate model of protein evolution detailing domain accretion and are typically observed across living kingdoms [14]). Besides, a more multi-domain proteins (indeed, different multi-domain combinations lines) from cumulative, indirect interactions (“AND”) within multi-protein complexes (red dashed lines). B&C. Comparison with protein direct & indirect interaction data for Yeast from BIND [53] database (B&C filled symbols, indirect interactions from [54, 55]) and Ref [56] (C open symbols, see supporting information). Data are statistically averaged as in Fig. 2B&C to account for gaps in connectivities for large k ≥ 20, due to the finite size of Yeast PPI network. B. Two-parameter fit of both direct connectivity distribution $p_k$ and average direct connectivity of first neighbor proteins $g_1$ [24] (see Fig. 2C and text). Numerical predictions are averaged over 1,000 network realizations (central green and blue lines). Numerical averages plus or minus two standard deviations are also displayed to show the predicted dispersions (upper and lower green and blue lines). The two adjusted parameters ($\gamma = 0.1$ and $\lambda = 0.3$) correspond to a network growth rate of 20% and an average of 1.5 protein-binding sites (domains) per protein. The connectivity distribution of the underlying single-domain network (corresponding to $\gamma = 0.1$ and $\lambda = 0.0$) is also displayed (brown line) to illustrate its relation to the full multi-domain protein network (see text). C. Two-parameter fit of both direct & indirect “matrix” connectivity distribution $p_k$ and average direct & indirect “matrix” connectivity of first neighbor proteins $g_1$ [24] (see text). Same two adjusted parameters ($\gamma = 0.1$ and $\lambda = 0.3$) as in B while a selection of indirect interactions is added up to a total of 28,000 direct & indirect interactions (see supporting information).

Combining whole genome duplication and domain shuffling.

As noted in the introduction, whole-genome duplications promote efficient shuffling of multi-domain proteins by enabling many accretion and deletion events of functional domains after each genome doubling. We will assume in the following that this shuffling of multi-domain proteins is so efficient that protein domains encoded along the genome evolve independently from their inclusion in single- or multi-domain proteins (indeed, different multi-domain combinations are typically observed across living kingdoms [14]). Besides, a more elaborate model of protein evolution detailing domain accretion and deletion events leads to virtually identical results for the large scale topological features of PPI network (not shown). The asymptotic generating function $\hat{p}(x)$ for multi-domain protein networks with independent domain evolution can be deduced a posteriori as,

$$\hat{p}(x) = (1 - \lambda)p(x)(1 + \lambda p(x) + \lambda^2 p^2(x) + \ldots) = \frac{(1 - \lambda)p(x)}{1 - \lambda p(x)}$$

where $\lambda$ is the probability of covalent connection between successive protein domains encoded along the genome. This leads to an exponential distribution of multi-domain proteins, in agreement with actual distributions [58, 59], with an average of $1/(1 - \lambda)$ protein-binding sites per protein. While $p(x)$ now reflects the independent evolution of single protein-binding domains according to Eqs. [81, 12] it also controls the asymptotic properties of the derived multi-domain networks $\tilde{p}(x)$; in particular, for $\Gamma_o > 1 > \Gamma_n$, we obtain from Eq. [12] the following asymptotic expansion in the vicinity of $x = 1$,

$$\tilde{p}(x) = 1 - \frac{1 - p(x)}{1 - \lambda p(x)} \sim 1 - \ldots - \frac{\Lambda_o}{1 - \lambda}(1 - x)^\alpha - \ldots$$

which implies that degree distributions of multi-domain protein networks $\tilde{p}_k$ increase with respect to the underlying single-domain interaction network $p_k$ as $\tilde{p}_k \sim p_k/(1 - \lambda)$ for large $k$, while the fraction of proteins with a single binding partner $p_1$ decreases at the same time as $\tilde{p}_1 = \tilde{p}(0) = (1 - \lambda)p(0) = (1 - \lambda)p_1$ (see Fig. 3B). Note that the scale-free degree distribution of such multi-domain protein networks results from an asymmetric divergence of individual binding sites (or domains) rather than asymmetric divergence of global protein architectures. This has also consequences for the functionalization of duplicated genes (see supporting information). In particular, random (symmetric) “subfunctionalization” between protein duplicates at the level of protein domains does not prevent the emergence of scale-free networks with locally conserved topology, by contrast to random link “complementation” at the level of individual interactions (Fig. S1) which leads to exponential networks without conserved topology (as discussed above).

Hence, domain shuffling of multi-domain proteins provides a powerful, yet non-disruptive source of combinatorial innovation, as it preserves essential topological features inherited from the underlying protein-domain interaction network evolution.

Finally, comparison with experimental data sets including indirect protein-protein interactions [53, 54, 55] is made by adopting a statistical implementation of the “combinatorial logic” discussed above (see supporting information). It is based on a Dijkstra algorithm that estimates the relative importance of all possible indirect interactions between multi-domain (and single-domain) proteins for each PPI network realization. Figs. 3B&C show rather good fits of experimental
data sets corresponding to an estimated 30% to 60% coverage of actual PPI networks (see, however, supporting information). The two adjusted parameters, \( \gamma = 0.1 \) and \( \lambda = 0.3 \), correspond to a network growth rate of 20% (i.e. \( 1 + 2\gamma \)) and an average of 1.5 (i.e. \( 1/(1 - \lambda) \)) protein-binding sites (domains) per protein in agreement with broad estimates for these biological parameters (see above § and [58, 59]). This also confirms that the properties of PPI networks we have predicted from first principles (i.e. \( i \) exponential dynamics and \( ii \) symmetry breaking) are already transparent from partial data sets.

**Discussion**

Beyond whole genome duplications, local genome rearrangements such as small segmental duplications, rearrangements and horizontal transfers might well have been critical for the emergence and proliferation of living organisms. Moreover, we note that local duplications/deletions may also lead to exponential dynamics of PPI network evolution if they are selected independently in parallel (exponential models of local or partial genome duplication are presented in ref[25]). Yet, recent records (<500MY) from various eukaryote kingdoms (from protists to animals and plants) suggest that the majority of duplicates may still have arisen from successive whole genome duplications (although this will need to be confirmed as more fully sequenced eukaryote genomes will become available).

One possible origin for this less efficient selection of local duplications might be the dosage imbalance they initially induce, thereby raising the odds for their rapid nonfunctionalization (unless proved beneficial under concomitant environmental changes[25]). By contrast, rapid nonfunctionalization of duplicates following a whole genome duplication should be opposed by dosage effect. This is because whole genome duplications initially preserve correct relative dosage between expressed genes, while subsequent random nonfunctionalizations disrupt this initial dosage balance. Preventing rapid asymmetric divergence between duplicates from recent whole genome duplications appears, in the end, to increase their chance of neo- or subfunctionalization by favoring longer (symmetric) genetic drift rather than early (asymmetric) functional loss.

**Conclusion**

Large scale topological features of PPI networks emerge “spontaneously” in the course of evolution under simple duplication/deletion events [25]. regardless of the specific evolutionary advantages individual proteins might have been selected for. Yet, the intrinsic exponential dynamics of PPI network evolution by whole genome duplications (or independent local duplications selected in parallel[25]) requires an asymmetric divergence of protein duplicates. Such asymmetric divergence arises “naturally” at the level of protein-binding sites or domains (through “spontaneous symmetry breaking”) and is robust to extensive domain shuffling of multi-domain proteins.

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**SUPPORTING INFORMATION**

I. Supplementary Figure.

[Diagram showing the relationship between genome and P-P Interaction Network, with symbols indicating duplication and deletion processes.]

**FIG. S1:** Alternative Model of PPI network evolution through whole genome duplication with symmetric divergence of duplicated proteins and random link “complementation” [13, 21].

II. Proof of Recurrence Relations for Generating Functions (Eq.2 and Eq.15).

After each whole genome duplication, each node has at most doubled its number of neighbors counted through powers of \( x \) in the generating function. Hence, a given PPI network realization with \( N_k \) nodes of connectivity \( k (k \geq 0) \) will contribute to the next duplicated ensemble of PPI networks as,

\[
N_k x^k \rightarrow N_k x^{2k} \quad (16)
\]

After link deletion with probability \( \delta \) or \( \delta_i = \delta_o, \delta_n, \) it contributes to the \( x^m \) terms of the generating function (with \( m = 0, \ldots, 2k \)) as,

\[
N_k x^{2k} \rightarrow N_k \left( \sum_{\ell=0}^{k} \left( \frac{k}{\ell} \right) (\gamma x)^{\ell} \delta^{k-\ell} \right) \left( \sum_{\ell=0}^{k} \left( \frac{k}{\ell} \right) (\gamma o x)^{\ell} \delta^{k-\ell} \right) = N_k \left( (\gamma x + \delta)(\gamma o x + \delta) \right)^k \quad (17)
\]

for the asymmetric divergence model (Fig. 1, Eq. 2) and as,

\[
N_k x^{2k} \rightarrow N_k \left( \sum_{\ell=0}^{k} \left( \frac{k}{\ell} \right) (\gamma x)^{\ell} \delta^{k-\ell} \right) \left[ \sum_{j=0}^{k} \left( \frac{k}{j} \right) \frac{1}{2} \sum_{\ell=0}^{j} \left( \frac{j}{\ell} \right) (\gamma o x)^{\ell} \delta^{j-\ell} \right] \left( \sum_{\ell_n=0}^{k-j} \frac{1}{2k-j} \left( \frac{k-j}{\ell_n} \right) (\gamma o x)^{\ell_n} \delta^{k-j-\ell_n} \right) \]


with \( \gamma_e = (\gamma_0 + \gamma_n)/2 \) and \( \delta_e = (\delta_0 + \delta_n)/2 \) for the symmetric divergence model with link “complementation” \([18, 27]\) (Fig. S1, Eq. 15).

III. Gene functionalization patterns in different models of PPI network evolution through whole genome duplication.

The initial model depicted on Fig. 1 with asymmetric divergence of duplicated proteins leads typically to “neofunctionalization” of “new” duplicates, while “old” duplicates retain most initial interactions (if not all for \( \gamma_0 = 1 \)).

By contrast, the alternative model depicted on Fig. S1 with symmetric divergence of duplicated proteins and random link “complementation” \([19, 27]\) leads typically to random “subfunctionalization” between protein duplicates at the level of individual interactions. However, this eventually leads to exponential degree distributions with no topology conservation of the PPI network (see main text), whereas scale-free degree distributions with at least local topology conservation of the PPI network indeed emerge under the initial asymmetric model, Fig. 1.

Yet, as discussed in the main text, the necessary asymmetric divergence of protein duplicates occurs “spontaneously” at the level of protein-binding sites rather than of the entire (multi-domain) proteins, as assumed in Fig. 1. This motivates the redefinition of the initial model in terms of protein-binding domains (Fig. 3A) to capture the asymmetric divergence of protein duplicates at the level of protein-binding sites and allow, at the same time, for extensive domain shuffling events of multidomain proteins (see main text).

This more elaborate model of PPI network evolution by whole genome duplication and domain shuffling encompasses both “neofunctionalization” and “subfunctionalization” of gene duplicates at the level of protein domains, in agreement with the suggestion that gene/protein evolution should be analyzed in terms of domains rather than entire protein chains \([11, 12, 13, 26, 41]\). In addition, this combined model of PPI network evolution also provides a theoretical framework to describe the evolution of the “combinatorial logic” behind indirect interactions within multi-protein complexes (see Fig. 3A and main text).

IV. Statistical weighting of indirect interactions from protein complexes.

We use a statistical implementation of the “combinatorial logic” underlying indirect protein interactions. Indirect interactions between protein pairs are weighted by the product of binding site “availabilities” along the shortest weighted path of intermediate direct interactions connecting them. The “availability” \( a_i \) of a binding site \( i \) is defined as the relative expression level \( e_i \) with respect to its first neighbor binding partners \( j \) of connectivity \( d_{ij} \),

\[
\frac{a_i}{e_i + \sum_{j \in \{j \} \cap (i)} \frac{e_j}{d_{ij}}} < 1
\]

(19)

Where expression level \( e_j \) can be distributed with specific statistics, such as randomly, uniformly or according to characteristic power laws, as reported experimentally \([25, 43, 54, 55, 56]\). Yet, in practice, we found that the predicted large scale topological features of PPI networks depend only weakly on the specific distribution of expression levels (for reasonable distribution range).

The statistical probability of an (intermediate) direct interaction between domains \( i \) and \( j \) is then proportional to \( a_i a_j \), which we use in a Dijkstra-like algorithm \([23]\) for additive distance minimization assigning \( d_{ij}^{\pi} = -\ln(a_i a_j) > 0 \) weights between interacting domains \( i \) and \( j \). Because of the presence of both covalent peptide bonds and direct, noncovalent interactions between protein domains (Fig. 3A), indirect protein-protein interactions correspond to alternating paths of noncovalent and covalent interactions with no successive noncovalent interactions which are forbidden by the shared binding site constraint (i.e. a binding site can only interact with one binding partner at a time). We describe below an algorithm which performs a simultaneous minimization for paths starting with a covalent bond \( (e_i, j) \) and paths starting with a direct, noncovalent interaction \( (d_{ij}) \). An additional variable for second node \( v_{ij} \) on the path is also needed to avoid non-physical “covalent loops”.

The initialization of distances between protein domains is:

\[
\begin{align*}
\delta_{ij} &= d_{ij} = \max & \forall (i,j) \text{ pairs, and} \\
\delta_{ij} &= d_{ij} = -\ln(a_i a_j) & \text{for direct, noncovalent interactions,} \\
\delta_{ij} &= d_{ij} = \max & \text{for covalent bonds,} \\
\delta_{ij} &= d_{ij} = \max & \text{otherwise.}
\end{align*}
\]

We then iterate until convergence (after \( N^2 \times \lambda \) (longest path) operations):

\[
\begin{align*}
d_{ij} &= \min(d_{ij}, \min_{k \in \{l\} \cap d_{ik}} (\delta_{ik} + c_{kj})) \\
\delta_{ij} &= \min(c_{ij}, \min_{k \in \{l\} \cap v_{kj} \neq i} (\delta_{ik} + \min(d_{kj}, c_{kj})))) \\
v_{ij} &= \{k \in \{l\} | v_{kj} \neq i, \min(\delta_{ik} + \min(d_{kj}, c_{kj})))
\end{align*}
\]

and remove eventually the minimum paths starting with a covalent bond (to avoid double counting of indirect interactions for multidomain proteins below):

\[
d_{ij} = \max & \text{ if } d_{ij} \geq \min(c_{ij}, c_{ji})
\]

(20)

Hence, the probabilities to observe a single indirect interactions within protein complexes is given by:

\[
\begin{align*}
w_{ij} &= 0 & \text{if } d_{ij} = \max \\
\beta w_{ij} &= \beta \exp(-d_{ij}) & \text{otherwise,}
\end{align*}
\]

with the normalization condition \( \Sigma_{i<j} w_{ij} = 1 \), which gives \( 1/\beta = \Sigma_{i<j} \exp(-d_{ij}) \). \( w_{ij} \) is thus the normalized product of availabilities \( a_k \) along the shortest weighted path between \( i \) and \( j \).

Finally, the individual probabilities \( p_{ij} \) to observe a total of \( M \) indirect interactions within protein complexes are given by:

\[
p_{ij} = 1 - (1 - w_{ij})^n
\]

(21)

where \( n \) is solution of \( \Sigma_{i<j} p_{ij} = M \).

Given the number \( M \) of indirect interactions in various data sets \([53, 54, 55]\), we have assessed their expected contribution to the large scale topology of Yeast PPI network from the two-parameter \( \gamma - \lambda \) model described in the main text. \( M \approx 28,000 \) corresponds to the sum of about 9,000 direct physical interactions from the BIND database \([33]\) (Fig. 2B&C filled symbols) and about 19,000 “matrix” interactions from \([53, 54]\) between 2,100 proteins already involved in direct physical interactions (out of 4,576 proteins in the BIND database, Fig. 3C filled symbols). “Matrix” interactions from
ref. 55 (Fig. 3C open symbols) are “reconstructed” from supplementary information files of 55 as follows: “matrix” interactions are included for (each complex core)×(each associated “module”) and (each complex core)×(each associated “attachment” = one protein). This reconstructed dataset should therefore be considered as incomplete, since “matrix” interactions between compatible modules and/or attachments associated to a given core are not taken into account (information not given in 55).

Numerical fits (γ = 0.1, λ = 0.3) are displayed on Fig. 3C (for direct and indirect interactions) for both connectivity distribution (green) and average connectivity of first neighbors (blue). They correspond to the same adjusted values (γ = 0.1, λ = 0.3) as in Fig. 3B (for direct interactions only).

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