Scaling Laws in Spatial Network Formation

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Geometric constraints impact the formation of a broad range of spatial networks, from amino acid chains folding to proteins structures to rearranging particle aggregates. How the network of interactions dynamically self-organizes in such systems is far from fully understood. Here, we analyze a class of spatial network formation processes by introducing a mapping from geometric to graph-theoretic constraints. Combining stochastic and mean field analyses yields an algebraic scaling law for the extent (graph diameter) of the resulting networks with system size, in contrast to logarithmic scaling known for networks without constraints. Intriguingly, the exponent falls between that of self-avoiding random walks and that of space filling arrangements, consistent with experimentally observed scaling (of the spatial radius of gyration) for protein tertiary structures.

Most networks forming in the real world are spatially extended and often geometrically constrained. Common examples include volume exclusion in the dynamics of polymers, chemical interactions in folding proteins and local electromagnetic forces in ferrofluidic aggregates \cite{1-6}. How geometric constraints impact the dynamic formation processes of spatial networks and thereby their function, is far from fully understood.

In many physical, chemical and biological systems, interaction structure and geometrical arrangement are equally important \cite{7}, in particular for their dynamics. Key examples include proteins folding into their tertiary structures \cite{8,9}. During the folding process, not only do amino acids interact with their neighbours along the chain but also with units that are far apart in the chain but close in space \cite{10,11}. On the level of abstract contact networks \cite{12,13}, the process of protein folding can thus be considered as adding interaction links to a network, akin to percolation \cite{15,16}, but spatially transforming the network at the same time.

In this Letter, we demonstrate that geometric constraints induce algebraic scaling laws in the formation of spatial networks, suggesting self-similar ('fractal') structures. We introduce a stochastic model that explicitly captures the essential impact of such geometric constraints on establishing spatial contacts and map them to constraints on graph-theoretic link additions. Combining probabilistic analysis with mean field calculations, we show that the extensions of the resulting networks exhibit an algebraic scaling law with system size. In stark contrast, network formation processes without such constraints exhibit logarithmic scaling \cite{18} such that geometric constraints qualitatively change the nature of the scaling law. Intriguingly, the algebraic scaling law \textit{per se} as well as its exponent are consistent with the scaling of the experimentally observed spatial radius of gyration with the chain length of protein tertiary structures.

\textit{Geometrically constrained network formation.} To understand basic principles underlying geometrically constrained network formation dynamics consider an initial chain of identical, spatially extended units, each in contact with its nearest neighbor units (as all units are identical, this is a special case of a \textit{coin graph} \cite{19}). For later analytic accessibility, we take the space to be two-dimensional and the chain to be closed to a single cycle such that initially the units are indistinguishable. The latter does not change the scaling behaviour, because folding an open chain results in a collection of closed cycles, as we will see below. This chain represents the original aggregate such as an unfolded protein where the units are amino acids or an initial contact sequence of ferrofluidic particles.

In a time-discrete network forming process (Fig.1), the units randomly come into contact with each other under the geometric constraints that in each step (i) no two units overlap and (ii) units in contact at some point in time stay in contact. The chain thus non-locally deforms each time a new contact forms (Fig.1). The sequence of connections models the emergence of pair-wise contacts between interacting units moving in space under the above constraints. In the model, new contacts keep

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure1}
\caption{(color online) \textbf{Mapping spatial structure formation onto network formation.} Units coming into spatial contact (green dashed lines) induce additional links on the network level. The network becomes more and more compact as links add, in two dimensions yielding a subgraph of the triangular grid. For illustration, panels show networks of $N=11$ units for time steps $t \in \{1, 2, \ldots, 8\}$.}
\end{figure}
FIG. 2. (color online) Mapping constraints from spatial geometry to network topology. Links in the contact graph form when two randomly chosen units come into spatial contact, subject to geometric constraints (a)-(d) specified in the text. Process 1: adding a link (green dashed line) is allowed because all conditions (a)-(d) are satisfied. Process 2: adding a link (red dashed line) is forbidden due to condition (a) to avoid overlapping units. Process 3: adding a link on the outer face is forbidden due to condition (b) to avoid the possibility that units (here the one shaded yellow) may with later links (red dotted line) be enclosed by less than six other units during a subsequent step (e.g., red dotted line).

FIG. 3. (color online) Algebraic scaling laws in spatial network formation. (a) Scaling of chain lengths of experimentally analyzed proteins vs. their radius of gyration (Eq. 1) (37162 data points from [20] log-binned, with error bars indicating standard deviations). Best fits suggest algebraic (red) rather than logarithmic (gray) scaling (b) Algebraic scaling of graph diameter $D_{\text{final}}(N)$ as derived in this Letter (orange line), plotted vs. the chain lengths $N$. Black dots indicate 450 stochastic realizations of network formation processes (uniformly sampled on a logarithmic chain length scale, binned and evaluated as in (a)) indicating the diameter of the original graph with best algebraic fit (red line). The algebraic scaling law with the (inverse) scaling dimension $\nu$ lying between that of self avoiding random walk $\nu_{\text{RW}}$ (green dashed lines) and that of space filling aggregates (blue dotted lines) is consistent with biological data but inconsistent with logarithmic scaling as expected from network formation processes without geometric constraints.

Spatial scaling of the network. The spatial extension of an aggregate is often measured by its radius of gyration

$$R_g = N^{-1} \left( \frac{1}{2} \sum_{i,j} (r_i - r_j)^2 \right)^{1/2} \sim N^{\nu},$$

(1)

quantifying the average distance between any pair out of $N$ units. Here, $r_i$ is the spatial position of unit $i \in \{1, \ldots, N\}$ and $\frac{1}{2}$ is the scaling dimension. Real three-dimensional protein structures indeed exhibit an algebraic scaling law (Fig 3b) with an exponent $\nu \approx 0.42 \pm 0.04$ above a lower bound $\nu_{\text{SF}} = 1/3$ implied by compact space filling aggregates [20][23] and below an upper bound $\nu_{\text{RW}} = 3/5$ resulting from self-avoiding random walks in three dimensions without further restrictions [24][26], together yielding:

$$\nu_{\text{SF}} < \nu < \nu_{\text{RW}}.$$  

(2)

For spatially embedded networks where each unit occupies space of the same order of magnitude we expect the diameter $D$ to increase linearly with spatial extension. Direct numerical simulations of the model processes for various system sizes indicate an algebraic scaling law

$$D_{\text{final}} \sim N^{\nu}.$$  

(3)
as found for biological protein tertiary structures, see Fig. 3. Specifically, the obtained scaling exponent \( \nu \approx 0.62 \pm 0.04 \) moreover satisfies the same types of upper and lower bounds (Eq. 2) as experiments on proteins suggest, between space filling configurations (in two dimensions \( \nu_{\text{SR}} = 1/2 \)) and that of self-avoiding random walks (\( \nu_{\text{RW}} = 3/4 \)).

**FIG. 4.** (color online) **Diameter path, diameter graph and end cycles.** A diameter path is a sequence of cycles of maximum length (here \( D_t = 7 \)) indicated by the dashed red line. For large graphs with defined average cycle length, \( D_t \) is proportional to the diameter of the original graph (black dots, black solid lines, pink solid lines indicate diameter). The diameter graph is the union of all such diameter paths (all shaded regions). \( V_t \) denotes the number of cycles on the diameter graph (here \( V_t = 12 \)) and \( E_t \) the number of end cycles (with only one neighbour) on any diameter path (here \( E_t = 5 \), shaded light rose).

**Network formation integrating constraints.** To understand the emergence of this scaling law and estimate its exponent, we mathematically analyze the network formation in the simplified network model with graph-theoretic constraints (a)-(d) inherited from the geometric ones (i) and (ii).

Consider at time \( t = 1 \) an initial graph consisting of one cycle of \( N \) units that evolves in a process in discrete time \( t \in \{1, 2, \ldots \} \), with exactly one link adding at a time. Each new link divides one cycle into two smaller cycles. Such a process exclusively generates networks that are planar graphs consisting of cycles.

How does the above scaling emerge? How do the constraints impact the structure formation process on the network level? The graph-theoretical diameter of the dual graph of a given network serves as a natural quantity representing a longest sequence of neighboring cycles in the original graph. At time \( t \), the diameter \( D_t \) of the dual therefore equals the length of (one of) its longest paths, representing a longest sequence of neighboring cycles in the original graph. We call such a sequence a **diameter path**. The union of all diameter paths (all sequences of cycles of the same (largest) length) in the original graph is called the **diameter graph**.

For small times \( t \), the cycles are typically of different lengths, for larger times become similar and eventually all become triangles. Thus, for sufficiently large times \( t \), the diameter of the network is proportional to that of the dual (Fig. 3). We thus take a mean field view and simply talk about the diameter, also when analyzing the scaling of the the diameter of the dual. Since no two cycles share more than one link, and no unit of the original network becomes enclosed in any path (due to condition (b)), the resulting dual graph stays a tree at all times. The diameter graph thus is the union of all paths of cycles of length \( D_t \). We note that the total number of cycles present at that time \( t \) equals \( t \).

We now derive a recurrence relation for the average diameter \( \langle D_t \rangle \) to then estimate how the final diameter scales with the chain length. Let \( \langle V_t \rangle \) be the expected number of cycles on the diameter graph and let \( \langle E_t \rangle \) be the number of end cycles (degree-one vertices of the dual) on any diameter path, as shown in Fig. 4. The average diameter \( \langle D_t \rangle \) evolves with time in three different ways. First, if a new link divides a cycle that is not part of the diameter graph, the diameter \( \langle D_t \rangle \) stays unchanged. Second, if a new link divides an end cycle of the diameter graph (Fig. 4), which in mean field approximation occurs with probability \( \langle E_t \rangle / t \), \( \langle D_t \rangle \) grows by one. Finally, if in the diameter graph a new link divides a cycle that is not an end, which analogously occurs with probability \( \langle V_t \rangle - \langle E_t \rangle \rangle / t \), \( \langle D_t \rangle \) grows by one if the splitting is transverse to a diameter path, which in turn occurs with some probability \( P_t^+ \); otherwise, if the splitting is parallel to the diameter path, \( \langle D_t \rangle \) also remains unchanged, compare Fig. 5. We thus obtain the recurrence relation

\[
\langle D_{t+1} \rangle = \langle D_t \rangle + \frac{1}{t} \left( \langle E_t \rangle + \langle (V_t) - \langle E_t \rangle \rangle P_t^+ \right)
\]

for the expectation value of the diameter. It remains to estimate \( P_t^+ \), \( \langle E_t \rangle \) and \( \langle V_t \rangle \) and then to iterate the recurrence relation in time to obtain the diameter of the final network.

**Approximating \( P_t^+ \).** To find \( P_t^+ \), we first compute the probability \( P_t(D_t \text{ increases}) / \ell \) of the diameter increasing given that a link adds in a cycle of length \( \ell \) on the diameter path. There are two ways such a link can add, see Fig. 5. If adding a link splits the cycle parallel to the diameter path, the newly created cycle becomes a side arm of the path, leaving the diameter unchanged. Alternatively, if the new link splits the cycle transversally to the direction of the path the diameter extends by one. Let \( h_1 \) and \( h_2 (= \ell - h_1) \) be the numbers of units in the two fractions transversal to the diameter path (Fig. 5). Increasing the diameter thus requires to connect one of the \( h_1 \) units to one of the other \( h_2 \) units. Then \( P(D_t \text{ increases}) / \ell, h_1) = 2 h_1 (\ell - h_1) / (\ell (\ell - 3)) \), because there are \( \ell (\ell - 3)/2 \) ways of connecting any two units in the cycle and \( h_1 (\ell - h_1) \) ways of forming a transversal connection, the term “−2” taking care of the two links that already exist between the two fractions of the original cycle. As every splitting of the cycle into two parts is
The number of cycles in the diameter graph are exponentially suppressed, the second term is negligible (see Supplemental Material for more details).

We now approximate the detailed dynamics \( \langle V_t \rangle \), by its rigorous lower bound given by Jensen’s inequality.

Thus even without the approximation of the dynamics \( P_t \), (see SM), indicates that any estimate of the time average \( \nu_{\text{theory}} = \langle \bar{D} \rangle \) must lie within an interval \( \nu_{\text{theory}} \in [\nu_{\text{min}}, \nu_{\text{max}}] \), where \( \nu_{\text{max}} < 2/3 \) and \( \nu_{\text{min}} > 1/2 \). Thus even without the approximation of the dynamics \( P_t \), an algebraic scaling is guaranteed and its exponent is above that for space filling aggregates, \( \nu_{\text{theory}} > \nu_{\text{SF}} \).

The scaling law intrinsically results from the geometric constraints: without such constraints the process analyzed above exactly reduces to the formation of Watts-Strogatz small-world networks with new links randomly adding to a circular graph \([28–30]\); for sufficiently many links, the diameter of such networks exhibits logarithmic scaling that is thus inconsistent with the algebraic scaling we found. Roughly speaking, due to the geometric constraints, any new link between two units drastically increases the probability of creating further links in these units’ respective neighborhoods. As a consequence, the structures cannot be arbitrarily compact. Our numerical results as well as analytic derivations above indicate that the spatial extent is modified qualitatively, changing a logarithmic to an algebraic scaling law.

**Conclusion and outlook.** Taken together, we uncovered an algebraic scaling law for network formation processes under geometric constraints. We have analyzed a spatial network formation model by mapping geometric constraints in space to purely graph-theoretical constraints on the topological changes of a network. Direct numerical simulations as well as analytic mean field calculations strongly indicate a scaling law with the graph diameter growing algebraically with system size, representing spatially self-similar (‘fractal’) networks. This algebraic law is largely independent of the details of the model setup and clearly induced by geometric constraints. Even without the time-averaging approximation of the dynamics \( P_t \), an algebraic scaling is guaranteed, exhibiting an exponent larger than that of a space filling aggregate, \( \nu_{\text{theory}} > \nu_{\text{SF}} \).
\( \nu > \nu_{5\ell}, \) thus indicating self-similar features. Both the algebraic scaling per se and its exponent are consistent with experimentally observed scaling of protein tertiary structures in real space \([20, 21, 23]\). More generally, our results may suggest that geometric constraints generically induce algebraic (rather than logarithmic) scaling laws of networks forming in space.

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