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FAST TRACK COMMUNICATION

Degree-based graph construction

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Abstract

Degree-based graph construction is a ubiquitous problem in network modelling (Newman *et al* 2006 *The Structure and Dynamics of Networks* (*Princeton Studies in Complexity*) (Princeton, NJ: Princeton University Press), Boccaletti *et al* 2006 *Phys. Rep.* **424** 175), ranging from social sciences to chemical compounds and biochemical reaction networks in the cell. This problem includes existence, enumeration, exhaustive construction and sampling questions with aspects that are still open today. Here we give necessary and sufficient conditions for a sequence of nonnegative integers to be realized as a simple graph's degree sequence, such that a given (but otherwise arbitrary) set of connections from an arbitrarily given node is avoided. We then use this result to present a swap-free algorithm that builds *all* simple graphs realizing a given degree sequence. In a wider context, we show that our result provides a greedy construction method to build all the *f*-factor subgraphs (Tutte 1952 *Can. J. Math.* **4** 314) embedded within $K_n \setminus S_k$, where K_n is the complete graph and S_k is a star graph centred on one of the nodes.

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1. Introduction and summary

In network modelling of complex systems [1, 2], one usually defines a graph with components of the system, being represented by the nodes, and the interactions amongst the components being represented as the links (edges) of this graph. This graph is usually inferred from empirical observations of the system and it is uniquely determined if one can specify all the connections in the graph. Occasionally, however, the data available from the system are incomplete, and one cannot uniquely determine this graph. In this case there will be a *set* \mathcal{G} of graphs satisfying the data, and one is faced with the following problems: (1) construct a graph

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from \mathcal{G} , (2) count the number of elements (graphs) in \mathcal{G} , (3) construct all graphs from \mathcal{G} and (4) construct a *typical* element of \mathcal{G} , often interpreted as a uniform random sample taken from \mathcal{G} . Problems (1), (3) and (4) are construction type problems, whereas (2) is an enumeration type problem. In this communication, we restrict ourselves to simple, undirected graphs, that is, any edge connects a single pair of distinct nodes (no hypergraph, no self-loops) and there is at most one edge incident on any pair of nodes (no parallel or multiple edges).

A rather important and typical situation is when the empirical data specify only the degrees of the nodes, in the form of a sequence $d = \{d_1, d_2, \dots, d_n\}$ of positive integers, $d_i \ge 1$, $1 \le i \le n$. (We exclude zero degree nodes, since they represent isolated points.) In the following, we will refer to such cases as 'degree-based' construction (enumeration) problems. There are numerous examples, we will mention only a few here. In epidemic studies of sexually transmitted diseases [4] the data collected are from anonymous surveys, where the individuals specify the *number* of different partners they have had in a given period of time, without revealing their identity. In this case, the epidemiologist is faced with constructing the most typical contact graph obeying the empirical degree sequence. Another example comes from chemistry where the task is to determine the total number of structural isomers of chemical compounds, such as alkanes. In this case, nodes represent chemical elements (atoms) in the compound and a link represents a chemical bond. In the case of alkanes the bond can be interpreted as a single link in the corresponding graph (no double bonds). Since the valence of an atom is fixed, the formula of an alkane such as C₄H₁₀ (butane) will specify only the degree sequence. Knowing all the possible graphs with this degree sequence provides a starting point from which the feasible structures can be inferred. In particular, butane has 2, octane (C₈H₁₈) has 18, C₂₀H₄₂ has 366 319 isomers, etc. Degree-based graph construction is also found in many other network modelling problems, such as communications (the internet, WWW, peer-to-peer networks), biology (metabolic networks, gene transcription, etc), ecology (food webs) and social networks.

It is easy to see that not all integer sequences can be realized as the degrees of a simple graph (the existence problem). For example, while $\{2,1,1\}$ and $\{2,2,2\}$ are the degree sequences of a path $(P_3, \bullet - \bullet - \bullet)$ and a triangle, there is no simple graph with degree sequence $\{3,2,1\}$ or $\{1,1,1\}$ or $\{4,4,2,1,1\}$. Let G(V,E) denote a simple graph where $V=\{v_1,v_2,\ldots,v_n\}$ denotes the set of nodes and E the set of edges. Consider a sequence of positive integers $d=\{d_1,d_2,\ldots,d_n\}$ arranged in decreasing order, $d_1 \geq d_2 \geq \cdots \geq d_n$ (for convenience reasons, only). If there is a simple graph G(V,E) with degree sequence d, then we call the sequence d a graphical sequence and in this case we also say that d realizes d. A second observation is that given a graphical d (and thus, we know that a simple graph d exists with this degree sequence), careless connections of pairs of nodes may not result in a simple graph. For example, consider the sequence $\{2,2,2,2\}$ which is graphical d (4-cycle). Making the connections d (d (d (d), d), d), d (d), d (d

Degree-based graph construction

Given a sequence of integers $d = \{d_1, d_2, \dots, d_n\}, d_1 \geqslant \dots \geqslant d_n \geqslant 1$,

- (A) Is there a simple graph G(V, E) on *n*-nodes realizing d?
- (B) If the answer to (A) is yes, how can we build such a graph?
- (C) Can we build all such graphs?
- (D) Let $\mathcal{G}(d)$ be the set of all such graphs. How can we sample at uniform from $\mathcal{G}(d)$?

There are two well-known theorems that answer question (A) above, namely the Erdős–Gallai theorem [12] and the Havel–Hakimi theorem [13, 14], the latter also giving a

construction algorithm for a graph with degree sequence d and thus answering question (B) as well (see section 2). In principle, problem (C) can be resolved via the method of edge swaps starting from the graph produced by the Havel-Hakimi procedure from (B) (called an HH-graph from now on) and book-keeping the swaps (which gets rather involved). Given two edges (v_1, v_2) and (v_3, v_4) , they can be swapped into (v_1, v_3) and (v_2, v_4) , or (v_1, v_4) and (v_2, v_3) leaving the degree sequence unchanged. Due to a theorem by Ryser [5], if G_1 and G_2 are two simple graphs with identical degree sequences, then there is a sequence of edge swaps that transforms one into another [6, 7]. Edge swapping is also at the basis of all sampling algorithms attempting to answer (D), using a Markov chain Monte Carlo approach, the literature of which is too extensive to be reviewed here. The basic idea is to keep swapping edges until the memory of the initial condition (HH-graph) is lost and one produces a (pseudo)-random instance. This sampling method is approximative and it is not well controlled in general (except for some specific sequences). A simple and direct (swap-free) construction method to produce a uniformly sampled random graph from $\mathcal{G}(d)$ was presented by Molloy and Reed (M-R) [8] (see section 5) and subsequently used to generate graphs with given degree sequences [9], including those described by power-law degree distributions [10]. The problem with the M-R algorithm is that it can become very slow due to rejections caused by self-loops and parallel edges (see section 5).

Here we present a new approach to degree-based graph construction. First we prove our main result that gives the sufficient and necessary conditions for a sequence of integers to be graphical such that a given (but otherwise arbitrary) set of connections from a given (but otherwise arbitrary) node is avoided. We then show how to use this result to present an algorithm that builds *all graphs* from $\mathcal{G}(d)$ (question (C)). It is important to emphasize that our algorithm does not use edge swaps, it is a direct construction method. Lastly, (section 5) we show how our result improves on the M–R method of uniform sampling, allowing us to reject some of the samples without getting to the point where the multi-edge conflicts would actually occur (see section 5). We also make a connection with the renowned Tutte's f-factor theorem [3, 11], showing that our result provides a greedy algorithm for constructing all f-factors in the case of $K_n \setminus S_k$, where K_n is the complete graph on n nodes and S_k is a star graph with k leaves centred on some arbitrary node.

The communication is organized as follows. Section 2 recalls known fundamental theorems for graph construction, and via a simple counter-example it shows that HH is not sufficient to build all graphs from $\mathcal{G}(d)$; section 3 presents our main theorem with its proof and section 4 describes the algorithm for building all graphs in $\mathcal{G}(d)$; section 5 is devoted to discussions.

2. Previous results

For simplicity of the notation we will identify node v_i by the integer i. There are two well-known necessary and sufficient conditions for a sequence of nonnegative integers to be graphical: one was given independently by Havel [13] and Hakimi [14], while the other is due to Erdős and Gallai [12]. We now announce these results for later reference, however, without proof, those can be found in the corresponding references.

Theorem 1 (Hakimi–Havel, HH). There exists a simple graph with degree sequence $d_1 > 0$, $d_2 \ge \cdots \ge d_n > 0$ if and only if there exists one with degree sequence $d_2 - 1, \ldots, d_{d_1+1} - 1, d_{d_1+2}, \ldots, d_n$.

Theorem 2 (Erdős–Gallai, EG). Let $d_1 \ge d_2 \ge \cdots \ge d_n > 0$ be integers. Then they are the degree sequence of a simple graph if and only if



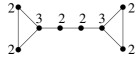


Figure 1. This graph cannot be obtained by the Havel–Hakimi procedure. The integers indicate node degrees.

(i) $d_1 + \cdots + d_n$ is even;

(ii) for all k = 1, ..., n - 1 we have

$$\sum_{i=1}^{k} d_i \leqslant k(k-1) + \sum_{i=k+1}^{n} \min\{k, d_i\}.$$
 (1)

Note that theorem 1 provides a greedy algorithm (the HH-algorithm) to generate an actual graph with the given degree sequence d while theorem 2 is an existence result. Tripathi and Vijay have recently shown [15] that it is enough to check the inequalities (1) for $1 \le k \le s$, where s is determined by $d_s \ge s$, $d_{s+1} < s+1$, that is only as many times as many distinct terms are in the degree sequence.

In the following, we will imagine the given degree sequence as a collection of *stubs*: at each vertex i there are d_i stubs ('half-edges'), anchored at the vertex, but the other ends are free. Connecting two stubs at two distinct nodes will form an edge between those nodes. We will call *the residual degree* the number of current stubs of a node.

The HH-algorithm for constructing a graph realizing a graphical sequence d proceeds as follows: connect all stubs of a node to nodes that have the largest residual degrees and repeat until no stubs are left. It is important to emphasize that one can choose any node to connect its stubs, as long as we connect all its stubs to the other nodes with the largest residual degrees. Clearly, if we always choose a high degree node (from the residual sequence) to connect its stubs, the HH-algorithm will create a graph in which high degree nodes tend to be connected to other high degree nodes, called assortative property [16]. However, if we always pick a node with a low (residual) degree to connect, we will likely obtain a graph with dissassortative property [17]. The HH-theorem is also a consequence (as a corollary) of our main result, see section 3. Nevertheless, this is still not enough to produce all graphs realizing a graphical node to connect is a node with degree 3, then the HH-algorithm connects it to the other node with degree 3 (highest degree). If the first node to connect has degree 2, then the HH-algorithm connects both its stubs to nodes with degree 3. However, the graph in figure 1 does not have any of the connections just mentioned (a 3-3 or 3-2-3 connection), and thus it cannot be constructed with the HH-algorithm. In the following section, we introduce a theorem that allows us to construct all labelled graphs with a given degree sequence.

The above results are naturally placed in the larger context of Tutte's famous f-factor theorem [3]. Given an integer function $f:V\to\mathbb{N}\cup\{0\}$, the f-factor of a given simple graph G(V,E) is a subgraph H of G such that $d_H(v)=f(v)$ for all $v\in V$. Here $d_H(v)$ is the degree of v within H. Tutte gave sufficient and necessary conditions for the existence of an f-factor for G [3] and later connected this to the problem of finding perfect matchings in bipartite graphs [11]. It is not hard to see that taking $G=K_n$, that is the complete graph on n-nodes, the f-factor problem is exactly question (A) of the degree-based construction problem with $d=\{f(v_1),\ldots,f(v_n)\}$. In this sense, the HH-algorithm is a greedy method for constructing an f-factor on K_n .



3. Graphical sequences with constraints from a single node

Before we can announce and prove our main result, we need to introduce a number of definitions and observations.

Definition 1. Let A(i) be an increasingly ordered set of d_i distinct nodes associated with node i: $A(i) = \{a_k \mid a_k \in V, \ a_k \neq i, \ \forall k, 1 \leq k \leq d_i\}$.

Usually, this set will represent the set of nodes adjacent to node i in some graph G, therefore we will refer to A(i) as an *adjacency set* of i.

Definition 2. If for two adjacency sets $A(i) = \{..., a_k, ...\}$ and $B(i) = \{..., b_k, ...\}$ we have $b_k \le a_k$ for all $1 \le k \le d_i$, we say that $B(i) \le A(i)$.

In this case we also say that B(i) is 'to the left' of A(i).

Definition 3. Let $d_1 \ge d_2 \ge \cdots \ge d_n \ge 1$ be a graphical sequence, and let A(i) be an adjacency set of node i. The degree sequence reduced by A(i) is defined as

$$d'_k|_{A(i)} = \begin{cases} d_k - 1 & \text{if } k \in A(i) \\ d_k & \text{if } k \in [1, n] \setminus (A(i) \cup \{i\}) \\ 0 & \text{if } k = i. \end{cases}$$
 (2)

In other words, if A(i) is the set of adjacent nodes to i in the graph G, then the reduced degree sequence $d'|_{A(i)}$ is obtained after removing node i with all its edges from G.

Lemma 3. Let $\{d_1, \ldots, d_j, \ldots, d_k, \ldots, d_n\}$ be a non-increasing graphical sequence and assume $d_j > d_k$. Then the sequence $\{d_1, \ldots, d_j - 1, \ldots, d_k + 1, \ldots, d_n\}$ is also graphical (not necessarily ordered).

Proof. Since $d_j > d_k$, there exists a node m connected to node j, but not connected to node k. Let us cut the edge (m, j) and remove the disconnected stub of j. If we add one more stub to k and connect this new stub to the disconnected stub of m, then we can see that the new graph is also simple with degree sequence $\{d_1, d_2, \ldots, d_j - 1, \ldots, d_k + 1, \ldots, d_n\}$.

Lemma 4. Let $d = \{d_1, d_2, \dots, d_n\}$ be a non-increasing graphical sequence, and let A(i), B(i) be two adjacency sets for some node $i \in V$, such that $B(i) \leq A(i)$. If the degree sequence reduced by A(i) (that is $d'|_{A(i)}$) is graphical, then the degree sequence reduced by B(i) (that is $d'|_{B(i)}$) is also graphical.

Proof. Let $A(i) = \{\ldots, a_k, \ldots\}$ and $B(i) = \{\ldots, b_k, \ldots\}$, $k = 1, \ldots, d_i$. Consider the adjacency set $B^1(i) = \{b_1, a_2, a_3, \ldots, a_{d_i}\}$ (we replaced node a_1 by node $b_1 \leq a_1$). If $b_1 = a_1$ then there is nothing to do, we move on (see below). If $b_1 < a_1$ then the conditions in lemma 3 are fulfilled. Namely, $b_1 < a_1$ implies $d_{b_1} \geq d_{a_1} > d_{a_1} - 1$ and we know that the sequence $d'|_{A(i)} = \{\ldots, d_{b_1}, \ldots, d_{a_1} - 1, \ldots, d_{a_2} - 1, \ldots\}$ is graphical by assumption. Thus, according to lemma 3, the sequence $\{\ldots, d_{b_1} - 1, \ldots, d_{a_1}, \ldots, d_{a_2} - 1, \ldots\}$ is also graphical, that is the one reduced by the set $B^1(i)$. Next, we will proceed by induction. Consider the adjacency set $B^m(i) = \{b_1, \ldots, b_m, a_{m+1}, a_{m+2}, \ldots, a_{d_i}\}$ and assume that the degree sequence reduced by it (from d) is graphical. Now, consider the adjacency set $B^{m+1}(i) = \{b_1, \ldots, b_{m+1}, a_{m+2}, a_{m+3}, \ldots, a_{d_i}\}$ (replaced a_{m+1} by a_{m+1}). If $a_{m+1} < a_{m+1}$, lemma 3 can be applied again since $a_{m+1} < a_{m+1}$ implies $a_{m+1} > a_{m+1} > a_{m+1} - 1$, showing that the sequence reduced by $a_{m+1} > a_{m+1} < a_{m+1}$ implies $a_{m+1} > a_{m+1} > a_{m+1} < a_{m+1} < a_{m+1}$ finishes the proof.

Definition 4. Let $d = \{d_1, d_2, \ldots, d_n\}$ be a decreasing graphical sequence and consider an arbitrary node $i \in V$, and an arbitrarily fixed integer m with $0 \le m \le n-1-d_i$. Let us fix a set of nodes $X(i) = \{j_1, \ldots, j_m\} \subset V \setminus \{i\}$ and consider the set $L(i) = \{l_1, \ldots, l_{d_i}\}$ containing the d_i lowest index nodes not in X(i) and different from i. We call L(i) the leftmost adjacency set of i restricted by X(i). Accordingly, we call the set of nodes X(i) the set of forbidden connections for i.

Lemma 5. If $d = \{d_1, d_2, ..., d_n\}$ is a decreasing graphical sequence, and $Y(i) = \{y_1, ..., y_{d_i}\}$ is an adjacency set disjoint from $X(i) \cup \{i\}$, then $L(i) \leq Y(i)$.

Proof. This is immediate, since by definition 4, $l_j \leq y_j$, for all $j = \{1, ..., d_i\}$.

We are now ready for the main theorem.

Theorem 6 (Star-constrained graphical sequences). Let $d_1 \ge d_2 \ge \cdots d_n \ge 1$ be a sequence of integers. For an arbitrary node $i \in V$ define a set $X(i) = \{j_1, \ldots, j_m\} \subset V \setminus \{i\}$ with $m \le n-1-d_i$, and consider L(i), the leftmost adjacency set of i restricted by X(i). Then the degree sequence $\mathbf{d} = \{d_1, \ldots, d_n\}$ can be realized by a simple graph G(V, E) in which $(i, j) \notin E$, for all $j \in X(i)$, if and only if the degree sequence reduced by L(i) is graphical.

Proof. ' \Leftarrow ' is straightforward: add node i to the reduced set of nodes, then connect it with edges to the nodes of L(i). Thus we obtained a graphical realization of d in which there are no connections between i and any node in X. ' \Longrightarrow ' In this case, d is graphical with no links between i and X(i), and we have to show that the sequence obtained from d by reduction via L(i) is also graphical. However, d graphical means that there is an adjacency set A(i) (with $A(i) \cap X(i) = \emptyset$) containing all the nodes that i is connected to in G. Thus, according to lemma S, we must have $L(i) \leq A(i)$. Then, by lemma S, the sequence reduced by S is graphical.

Note that the forbidden set of connections form a star graph $S_{|X(i)|}$ centred on node i. Also note that considering the empty set as the set of forbidden nodes, $X(i) = \emptyset$, we obtain the Havel–Hakimi theorem 1 as corollary. Informally, theorem 6 can be announced as follows:

Let $\mathbf{d} = \{d_1, d_2, \dots, d_n\}$, be a decreasing graphical sequence and let i be a fixed, but an arbitrary vertex. Assume we are given a set of forbidden connections in V incident on i. Then there exists a realization of the degree sequence avoiding all forbidden connections if and only if there also exists a realization where i is connected with vertices of highest degree among the non-forbidden ones.

Since the forbidden connections emanating from a node i form a star graph S_k , k = |X(i)|, theorem 6 provides sufficient and necessary conditions for the existence of an f-factor for $G = K_n \setminus S_k$. More importantly, it gives a *greedy algorithm* for finding such an f-factor.

4. Building all graphs from $\mathcal{G}(d)$

As we show next, theorem 6 provides us with a procedure that allows for the construction of *all* graphs realizing the same degree sequence.

Consider a graphical degree sequence d on n nodes. Certainly, we can produce all graphs realizing this sequence by connecting all the stubs of a chosen node first, before moving on to another node with stubs to connect (that is we finish with a node, before moving on). In this vein, now choose a node i and connect one of its stubs to some other node j_1 . Is the remaining degree sequence $d' = \{d_1, \ldots, d_i - 1, \ldots, d_{j_1} - 1, \ldots, d_n\}$ still graphical such

that nodes i and j_1 avoid another connection in subsequent connections of the other stubs? Certainly, as a necessary condition, d' has to be graphical as a sequence, since all subgraphs of a simple graph are simple, and thus if G is a simple graph realizing d with $(i, j_1) \in E$, then after removing this edge, the remaining graph is still simple. However, after making some connections from a node, it is not sufficient that the residual sequence is graphical. One might still be forced to make multiple edges, as illustrated by the following example. Consider the graphical sequence $\{2, 2, 1, 1\}$ (the path P_4 , $\bullet - \bullet - \bullet - \bullet$) as the degrees of the set of nodes $V = \{u, v, x, y\}$ ($d_u = d_v = 2, d_x = d_y = 1$). Connect nodes u and v. We certainly have not broken the graphical character yet, since we could still finish the path by connecting next u to u (or to u) and u to u (or to u). The remaining sequence u in the path by connecting next u to u (or to u) and u to u (or to u). Next, connect node u to node u. The remaining (residual) sequence is u (emanating from node u and u, respectively), graphical on its own, however, we can no longer connect nodes u and u, because the very first connection is already there. Thus, after making one, or more connections from a node u, how can we check that the next connection from u will not break the graphical character?

Theorem 6 answers this question if we think of the connections already made from node i as forbidden connections. That is, after the first connection of i to j_1 take d' as d in theorem 6 and $X(i) = \{j_1\}$. Then, to test whether the sequence reduced by the corresponding L(i) is graphical we can employ for example the Erdős-Gallai theorem 2, checking all the inequalities, or the Havel-Hakimi theorem 1. If the test fails on the reduced sequence, one must disconnect i from j_1 and reconnect it somewhere else. The graphical character of the original sequence guarantees that there is always j_1 where the test will not fail. If, however, the remaining degree sequence is graphical with the constraint imposed by X(i), we connect another stub of i to some other node j_2 (different from j_1), adding an element to the forbidden set of connections X(i). To check whether after the second connection the remaining sequence is still graphical with the constraint imposed by the new set $X(i) = \{j_1, j_2\}$, we proceed in exactly the same way, using theorem 6, repeating the procedure until all the stubs of node i are connected away into edges. After this we can move on to some other node i (arbitrary) from the remaining set of nodes and repeat the procedure. Note that this procedure is not a real procedure in the sense that it does not prescribe which stubs to connect. It only tells us whether the connection we just made (by whatever process) has broken the graphical character. Since every element from $\mathcal{G}(d)$ can be realized by some sequence of connections, it is clear that if we specify a systematic way of going through all the possible connections while employing theorem 6, we will realize all elements of $\mathcal{G}(d)$. However, taking all possible connections would be very inefficient. Next we present a version of a more economical algorithm that constructs every labelled graph with degree sequence d, and only once. For simplicity of the notation we will call the test for the graphical character via theorem 6, the 'CG test' (constrained graphicality test). The algorithm also exploits lemma 4, which guarantees preservation of graphicality for all adjacency sets to the left of a graphical one, thus avoiding costlier checks with EG or HH theorems for those adjacency sets. Clearly, a labelled graph can be characterized by the sequence of its adjacency sets $G = \{A(1), \ldots, A(n)\}$. This algorithm creates all the possible adjacency sets for node 1, then for each one of those repeats the same procedure on the reduced sequence by that adjacency set (in sense of definition 3) of at most n-1 nodes.

Algorithm 1 (All graphs). *Given a graphical sequence* $d_1 \ge d_2 \ge \cdots \ge d_n \ge 1$,

- (I) Create the rightmost adjacency set $A_R(1)$ for node 1: connect node 1 to n (this never breaks graphicality). Let k = n 1.
 - (I.1) Connect another stub of 1 to k. Run the CG test.
 - (I.2) If it fails, make k = k 1. Repeat I.1.



- (I.3) If it passes, keep (save) the connection, make k = k 1, and if i has stubs left, repeat from I.1.
- (II) Create the set $A(\mathbf{d})$ of all adjacency sets of node 1 that are colexicographically smaller than $A_R(1)$ and preserve graphicality:

$$A(d) = \{A(1) = \{a_1, \dots, a_{d_1}\}, a_i \in V | A(1) <_{CL} A_R(1), d'|_{A(1)} - graphical \}.$$

(III) For every $A(1) \in \mathcal{A}(d)$ create all graphs from the corresponding $\mathcal{G}(d'|_{A(1)})$ using this algorithm, where $d'|_{A(1)}$ is the sequence reduced by A(1).

For simplicity of the notation, we will drop the (1) from A(1), tacitly assuming that it refers to the leftmost node 1. Observe that the ordering relation '<' in definition 2 is a partial order, while the colexicographic order '< $_{\text{CL}}$ ' is a total order over all adjacency sets, however, '<' implies '< $_{\text{CL}}$ '. It is not hard to see that A_R is colexicographically the largest ('rightmost') sequence which still preserves graphicality. When constructing $\mathcal{A}(d)$, checking graphicality with the EG or HH theorems is only needed for those adjacency sets, which are incomparable by the '<' relationship to any of the current elements of $\mathcal{A}(d)$, while for the rest graphicality is guaranteed by lemma 4.

5. Discussion and outlook

Algorithm 1 proceeds by attempting to connect all stubs of the largest degree node as much to the right as possible. Depending on the degree sequence, it might happen that the CG test fails many times at step I, until it finds A_R . However, in that case, A_R is located more towards the higher degree nodes (towards the left) and thus the number of adjacency sets that preserve graphicality, namely $|\mathcal{A}|$ is smaller and accordingly, the algorithm has fewer cases to run through in subsequent steps. The more heterogeneous a degree sequence is, the more likely this will happen. Of course, it only makes sense to produce all graphs from $\mathcal{G}(d)$ for small graphs (chemistry), or graphical sequences that do not admit too many solutions. An interesting question would then be finding the conditions on the sequence d that would guarantee a given upper bound C on the size of $\mathcal{G}(d)$, $|\mathcal{G}(d)| \leq C$. A possible starting point in this direction could be Koren's [18] characterization of sequences uniquely realizable by a simple graph. Sequences that admit only a small number of realizations (labelled simple graphs) would likely be 'close' in some sense to these special sequences.

Algorithm 1 also provides a way to computationally enumerate all the labelled graphs $|\mathcal{G}(d)|$ realizing a degree sequence d (problem (B) of section 1). Naturally, the following recursion holds: $|\mathcal{G}(d)| = \sum_{A \in \mathcal{A}(d)} |\mathcal{G}(d'|_A)|$. Our graph construction process can be thought of as happening along the branches of a tree $\mathcal{T}(d)$ of depth at most n-1: the internal nodes of this tree on the kth level are all the allowed adjacency sets (from the corresponding \mathcal{A} set) of the node with the largest residual degree (the leftmost node). The reason this tree is of depth at most n-1 is because some other nodes (other than the one with the largest residual degree) in the process might loose all their stubs. A directed path towards a leaf of this tree corresponds to a graphical realization of d, because we end up specifying all the adjacency sets along this path. Based on this, during the realization of the graph, if we choose uniformly at random at every level of the tree within the children of a node, from the corresponding set \mathcal{A} , the probability of a final realization G in this process will be given by the product:

$$P(G) = \prod_{k=0} |\mathcal{A}(\mathbf{d}'|_{A^{(k)}})|^{-1},\tag{3}$$

where $A^{(k)}$ is the randomly chosen adjacency set of the node with the largest residual degree, on the kth level of $\mathcal{T}(d)$. By convention $A^{(0)} = \emptyset$, and $d'|_{A^{(0)}} = d$. It is not hard to convince

ourselves that the distribution in (3) is not uniform, and thus, this algorithm cannot be used in this form to produce uniform samples from $\mathcal{G}(d)$. However, theorem 6 can be used to improve on a well-known, direct uniform sampling process, the Molloy-Reed (M-R) algorithm [8]. In this process, one chooses between all the stubs uniformly at random, irrespective of what node they belong to. This is repeated until either a self-loop, or a double edge is created, or a simple graph is finished. When there is a self-loop or double edge, the process is stopped, and the algorithm starts from the very beginning. The CG test can be used along the way to test for graphicality after every connection just made. In particular, assume that we just connected node i with node j. We then run the CG test centred on node i (adding the (i, j)connection to the forbidden set from i). If it passes, we run the CG test on node j as well. If it fails either on i or j, we can stop the process, before actually running into a conflict later. Running into conflict usually happens towards the end, and this test can save us from possibly many unnecessary calls to the random number generator, speeding up the sampling algorithm. It is important to note that if the CG test passes on both i and j, we actually do not know whether graphicality is broken or still preserved at that stage! If the CG test fails, however, we know that graphicality was broken. The reason is because theorem 6 gives us the sufficient and necessary conditions for the sequence to be graphical such that no multiple edges will be made with the already existing connections emanating from the same node. It gives no such information for connections already made elsewhere! Of course, for degree sequences d for which there is only a small number of labelled graphs realizing it (compared to the total number $\prod_i d_i!$ of graphs that it produces), the M-R algorithm would search for needles in a haystack, and other (MCMC) methods will be necessary.

In summary, we have given necessary and sufficient conditions for a sequence of integers $d_1 \geqslant \cdots \geqslant d_n \geqslant 1$ to be graphical (realizable by simple, undirected graphs) avoiding multiple edges with an arbitrary star graph (the forbidden graph) $S_k(j)$, $0 \leqslant k < n - d_j$, centred on a node j. In a more general context, our result gives for the first time a greedy construction for Tutte's f-factor subgraphs within $K_n \backslash S_k(j)$. It would be desirable if such a *greedy* construction existed for an arbitrary forbidden graph F, not just for star graphs, however, at this point this still seems to be a rather difficult problem. Such an algorithm (if greedy) would further speed up the M-R sampling, because it would induce early rejections, as soon as they are made. Our main theorem also led to a direct and systematic construction algorithm that builds *all graphs* realizing a given degree sequence d. And finally, we mention that these studies can be extended to simple directed graphs as well (there are two degree sequences in this case), however, the computations are considerably more involved, and they will be presented separately.

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