

# Engineering fast almost optimal algorithms for bipartite graph matching

Ioannis Panagiotas, Bora Uçar

### ▶ To cite this version:

Ioannis Panagiotas, Bora Uçar. Engineering fast almost optimal algorithms for bipartite graph matching. ESA 2020 - European Symposium on Algorithms, Sep 2020, Pisa, Italy. hal-02463717v3

## HAL Id: hal-02463717 https://hal.inria.fr/hal-02463717v3

Submitted on 1 Jul2020

**HAL** is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers. L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.

# Engineering fast almost optimal algorithms for bipartite graph matching

#### 3 Ioannis Panagiotas 💿

- 4 ENS Lyon, France
- 5 ioannis.panagiotas@ens-lyon.fr
- 6 Bora Uçar 💿
- 7 CNRS and LIP, ENS Lyon, France
- 8 bora.ucar@ens-lyon.fr

#### <sup>9</sup> — Abstract

We consider the maximum cardinality matching problem in bipartite graphs. There are a number 10 of exact, deterministic algorithms for this purpose, whose complexities are high in practice. There 11 are randomized approaches for special classes of bipartite graphs. Random 2-out bipartite graphs, 12 where each vertex chooses two neighbors at random from the other side, form one class for which 13 there is an  $O(m + n \log n)$ -time Monte Carlo algorithm. Regular bipartite graphs, where all vertices 14 have the same degree, form another class for which there is an expected  $O(m + n \log n)$ -time Las 15 Vegas algorithm. We investigate these two algorithms and turn them into practical heuristics with 16 randomization. Experimental results show that the heuristics are fast and obtain near optimal 17 matchings. They are also more robust than the state of the art heuristics used in the cardinality 18 matching algorithms, and are generally more useful as initialization routines. 19

 $_{20}$  2012 ACM Subject Classification Theory of computation  $\rightarrow$  Design and analysis of algorithms

- 21 Keywords and phrases bipartite graphs, matching, randomized algorithm
- 22 Supplement Material https://gitlab.inria.fr/bora-ucar/fast-matching

#### <sup>23</sup> 1 Introduction

A matching in a graph is a set of edges, such that no two of them share a common vertex. 24 We consider the maximum cardinality problem in bipartite graphs which asks for a matching 25 with maximum cardinality. There are a number of exact algorithms for this problem. The 26 best known algorithms [21] run in  $O(m\sqrt{n})$  time for a graph with n vertices and m edges. 27 Such complexity can be prohibiting for large instances. For this reason, there is significant 28 interest in algorithms which can find large matchings in linear or near linear time [37]. The 29 practical use of approximate matchings in applications [33] and as an initialization to exact 30 algorithms [30] are well known. 31

We investigate two randomized algorithms by Karp et al. [22] and Goel et al. [18], both 32 of which run in  $O(m + n \log n)$  time. The former algorithm finds, almost surely, maximum 33 cardinality matchings on random graphs formed by allowing each vertex to select two 34 vertices from the other side uniformly at random. The latter algorithm finds maximum 35 cardinality matchings in regular bipartite graphs, where all vertices have equal degree. In 36 both of these classes of graphs, the bipartite graphs have equal number of vertices in each 37 part, and the maximum cardinality matchings cover all vertices (such matchings are called 38 perfect). We investigate these two theoretical algorithms for very special cases of bipartite 39 graphs and convert them to efficient heuristics for general bipartite graphs. We discuss 40 our implementations and investigate the performance of the resulting heuristics in terms of 41 run time and the matching cardinality. Both heuristics run in near linear time and obtain 42 matchings whose cardinality is more than 0.99 of the maximum, even in cases where the 43 current state of the art approaches have difficulties. 44

The rest of the paper is organized as follows. In Section 2, we give the necessary background. In Sections 3.1 and 3.2 we review the existing randomized algorithms and then discuss how we adapt them. Section 4 contains the experimental results, and Section 5 concludes the paper. Appendices A–D provide some additional results and discussion.

#### <sup>49</sup> **2** Background and notation

Let  $G = (R \cup C, E)$  be a bipartite graph, where R and C are two disjoint set of vertices, and E is the set of edges. The bipartite graph G can be represented with a matrix  $\mathbf{A}_G$ . The vertex  $r_i \in R$  corresponds to the *i*th row, and the vertex  $c_j \in C$  corresponds to the *j*th column, so that  $\mathbf{A}_G(i, j) = 1$  if and only if  $(r_i, c_j) \in E$ . We will refer to vertices of R as rows and to those of C as columns from this point on, and use  $\mathbf{A}$  to refer to  $\mathbf{A}_G$ .

Let  $\mathcal{M}$  be a matching. For  $(u, v) \in \mathcal{M}$ , the vertices u and v are matched, and they are each other's mate. A vertex is called free if it is not matched by  $\mathcal{M}$ . If there are no free vertices in R or in C, then  $\mathcal{M}$  is called perfect. An augmenting path with respect to  $\mathcal{M}$  is a path which starts with a free vertex and ends at another free vertex, where every second edge is in  $\mathcal{M}$ . A matching is maximum if and only if there are no augmenting paths [7].

A square matrix is called doubly stochastic if the sum of entries in each row and column is 60 equal to one. An  $n \times n$  matrix **A** has support if there is a perfect matching in the associated 61 bipartite graph G. A is said to have total support if each edge in G is used in a perfect 62 matching. A square matrix is fully indecomposable, if it has total support and cannot be 63 permuted into a block diagonal matrix. Any nonnegative matrix  $\mathbf{A}$  with total support can be 64 scaled with two positive diagonal matrices  $D_R$  and  $D_C$  such that  $A_S = D_R A D_C$  is doubly 65 stochastic, and if  $\mathbf{A}$  is fully indecomposable, then the matrices  $\mathbf{D}_{\mathbf{R}}$  and  $\mathbf{D}_{\mathbf{C}}$  are unique. The 66 Sinkhorn–Knopp algorithm [38] is a well-known method for finding such  $\mathbf{D}_{\mathbf{R}}$  and  $\mathbf{D}_{\mathbf{C}}$  for a 67 given matrix. This is an iterative algorithm, where at each iteration each row is normalized to 68 have unit length, and then each column is normalized to have unit length. If a given matrix 69 A has total support, then Sinkhorn–Knopp algorithm finds the unique scaling matrices. If A 70 has support but not total support, then entries that cannot be put into a perfect matching 71 tend to zero. The method converges with an asymptotical convergence rate depending on 72 the second singular value of the final doubly stochastic matrix. There are other iterative, 73 faster converging methods [1, 10, 28], whose iterations are more sophisticated than that of 74 Sinkhorn-Knopp's. 75

A k-out subgraph  $G_k$  of a host graph G is defined by allowing each vertex in G to 76 randomly select uniformly k of its neighbors, and the union of all selections forms the edge 77 set of  $G_k$ . Walkup [40] shows that in the pure random k-out setting, where the host graph is 78 the complete bipartite graph, the resulting  $G_k$  has a perfect matching with high probability 79 for  $k \geq 2$ . We do not know any general result about properties of  $G_2$  sampled from any 80 arbitrary host graph. Frieze and Johansson [17] investigate some other properties of  $G_k$ s on 81 host graphs where the minimum degree of a vertex is at least n/2. Dufossé et al. [16] propose 82 using the doubly stochastic matrix  $\mathbf{A}_{\mathbf{S}}$  (scaled version of the matrix representation) for 83 sampling and show an approximation result for  $G_1$ , when A has total support. We give some 84 experiments in which  $G_{2s}$  generated using the same probabilities have perfect matchings in 85 majority of the cases. 86

Two popular classes of randomized algorithms are *Las Vegas* and *Monte Carlo* algorithms. *Las Vegas* algorithms always return a correct answer, but their run time can depend on random choices, whereas Monte Carlo algorithms can fail with small probability, but their complexity is independent of the random choices made (see for example [34, p. 70]).

There are a number of heuristics for the cardinality matching problem [30, 37] (see Appendix A for a relevant discussion). Among those, that by Karp and Sipser [23] is very well known and widely used. This heuristic eliminates vertices of degree at most two in the following way. It matches any degree-1 vertices with their neighbors (and discards both), or merges the neighbors of a degree-2 vertex (which is then discarded) to a single node, and removes any parallel edges that occur. If neither operation can be done, it matches a pair of vertices randomly.

#### **3** Two heuristics

We describe the original Monte Carlo algorithm [22] for finding perfect matchings in 2-out 99 bipartite graphs in Section 3.1 and the original Las Vegas algorithm [18] for finding perfect 100 matchings in *d*-regular bipartite graphs in Section 3.2. These two algorithms are based on 101 uniform sampling. We generalize these two algorithms to general bipartite graphs within a 102 common framework. The framework we propose scales the adjacency matrix of the input 103 bipartite graph and uses the nonzero values of the scaled matrix for sampling. We also 104 identify and fix an oversight in the description of the Monte Carlo algorithm, and describe 105 efficient implementations of the two heuristics. 106

#### **3.1** 2OUTMC: Monte Carlo on 2-out graphs

#### <sup>108</sup> 3.1.1 Description of the algorithm

The Monte Carlo algorithm by Karp et al. [22] finds a perfect matching, with high probability, 109 in a random 2-out bipartite graph, sampled from the complete bipartite graph. A random 110 2-out bipartite graph  $B_{2o}$  is constructed by selecting uniformly at random two row vertices 111 for each column, and two column vertices for each row. These selections form the edges 112 of  $B_{2o}$ . Given the edges of  $B_{2o}$ , Karp et al. define two multigraphs. The Column-Graph 113 (CG) is the multigraph whose vertices are the rows, and whose edges are the choices of the 114 columns. That is, there is an edge in CG for a column vertex in  $B_{2o}$ . Parallel edges occur 115 if two columns select the same rows. The Row-Graph (RG) is defined similarly. The main 116 idea to show that  $B_{2o}$  has a perfect matching is the following. In a component of CG that 117 contains a cycle, it is possible to match all rows (vertices in CG) with one of the columns 118 that have selected them (edges in CG). On the other hand in a tree component of CG, in 119 any matching (pairing of edges with vertices) there will always be a free row vertex. As a 120 consequence, when one or more trees appear in CG, the choices of the columns alone do 121 not suffice to find a perfect matching, and those of the rows must be used. The algorithm 122 thus keeps track of the tree components of CG and tries to identify one row vertex per tree 123 component whose selections should be taken into account. The columns selected by such a 124 row could be used for a set of rows belonging in tree components. Thus one should go back 125 and forth identifying trees in CG and analyzing components in RG. Karp et al.'s algorithm, 126 which is described in Algorithm 1, formalizes this approach. 127

The algorithm operates on  $H_1$ , a copy of CG, and  $H_2$ , a copy of RG initially devoid of edges. It furthermore uses two arrays **checked** for columns and **marked** for rows. These two arrays together signal whether a vertex will be matched with one of its two selections or not. More specifically, if a row vertex r is marked (i.e., **marked**[r]=true), then the algorithm will match r with one of its two selections. On the other hand, if a column c is checked (i.e., **checked**[c]=true), then the algorithm will match c with one of the marked row vertices that have selected it.

Initially, all row vertices are unmarked and all column vertices are unchecked. The 135 algorithm at each step picks a tree from  $H_1$  and marks one of its vertices x. This signifies 136 that x can only be matched with one of its choices. Then, the edge of x is inserted in  $H_2$ . 137 The algorithm then finds the component  $Q_x$  in  $H_2$  containing the edge x, and selects an 138 unchecked column y from  $Q_x$ . Column y is checked, which means that it can only be matched 139 with a marked vertex. As y's choices are rendered useless now, the corresponding edge is 140 removed from  $H_1$  upon which new trees can arise. For each tree vertex x identified in  $H_1$ , one 141 should be able to find a vertex in the associated component  $Q_x$ , so that x can be matched in 142 that component. Otherwise,  $Q_x$  has more edges than vertices, and any matching of vertices 143 with edges in  $Q_x$  will hence leave some edges unpaired. In other words, Algorithm 1 has 144 decided that all columns that correspond to edges in  $Q_x$  should be matched with one of their 145 two selections. However, the union of the rows denoted by these selections has cardinality 146 strictly smaler than the number of such columns, and that is why a column is always left 147 unmatched by the algorithm if this scenario occurs. The algorithm returns failure upon 148 detecting this case (Line 10). The algorithm terminates successfully if all trees have a marked 149 vertex. If this happens, each component in  $H_1$  will have as many edges as unmarked vertices. 150 Likewise, each component in  $H_2$  will have as many edges as checked vertices. It is therefore 151 possible to orient the edges in either  $H_1$  or  $H_2$  such that each vertex (excluding marked rows 152 or unchecked columns) is matched with a unique adjacent edge. This gives a perfect matching 153 in  $B_{2o}$ , which can be found by the Karp–Sipser heuristic in linear time. Algorithm 1 finds a 154 perfect matching with probability  $1 - O(n^{-\alpha})$ , where  $\alpha$  is a positive constant. 155

**Algorithm 1** 20UTMC: Monte Carlo on 2-out graphs

1:  $H_1 \leftarrow CG, H_2 \leftarrow$  empty graph with columns as vertices;

2: All vertices in 
$$H_1$$
 are unmarked, all vertices in  $H_2$  are unchecked;

3: CORE  $\leftarrow$  edges in cycles of CG

4: while there exists a tree T in  $H_1$  with no marked vertex do

- 5: Let x be a random vertex of  $T \rightarrow x$  is a column vertex
- 6:  $marked[x] \leftarrow true \qquad \blacktriangleright x$  must be matched with one of its choices
- 7: Add the edge of x in  $H_2$
- 8: Let  $Q_x$  be the component in  $H_2$  containing the edge of x
- 9: **if**  $Q_x$  has no unchecked vertices **then**
- 10: Return Fail  $\blacktriangleright Q_x$  has more edges than vertices (no 1-1 pairing possible)
- 11: else
- 12: Select an unchecked vertex y of  $Q_x$ . In case of ties, prefer one from CORE
- 13:  $checked[y] \leftarrow true \rightarrow y$  will be matched with a row that selected it

14: delete y in  $H_1$   $\blacktriangleright$  The algorithm forgets y's choices

15: Create  $B'_{2o}$  from  $B_{2o}$  by keeping only edges between marked rows and checked columns (edges in  $H_2$ ) or unmarked rows and unchecked columns (edges in  $H_1$ )

16: Apply Karp–Sipser on  $B'_{2o}$  to find a perfect matchin

The authors then describe how to efficiently implement the algorithm such that it runs in  $O(n \log n)$  worst case time. They identify two main tasks:

**Task A**: Keep track of the tree components during edge deletions in  $H_1$ .

<sup>159</sup> **Task B**: Keep track of the connected components during edge insertions in  $H_2$ , and the single unchecked vertex in each component.

Task B can be efficiently done in amortized near linear time (over the course of the algorithm) by using a union-find data-structure and keeping the identity of the single unchecked vertex in a component of  $H_2$  at the root of the component. For Task A, Karp et al. propose the following. In the beginning, the edges of CG are labeled as  $\mathcal{F}$ , if their



**Figure 1** Algorithm 1 does not recognize new trees, if another edge is deleted after (u, v).

deletion creates a tree;  $\mathcal{T}$ , if they belong to a tree component; and  $\mathcal{C}$  otherwise. Let c-degree of a vertex v be the number of  $\mathcal{C}$  edges incident on v. During deleting the edge (u, v) from  $H_1$ , one of the following is performed depending on the label of (u, v).

**Case 1:** (u, v) is C: The c-degrees of u and v are decreased by one. Then, while there is a vertex with a single C edge; its C edge is relabeled as  $\mathcal{F}$ .

**Case 2:** (u, v) is  $\mathcal{F}$ : Using a dove-tailed depth-first search, where depth-first searches from *u* and *v* are interleaved, the tree component created can be found in time proportional to its size. One then changes the labels of all edges in this tree from  $\mathcal{F}$  to  $\mathcal{T}$ .

**Case 3:** (u, v) is  $\mathcal{T}$ : Deleting (u, v) creates two trees. As in the previous case, a dovetailed DFS is used to find these two trees in time proportional to the size of the smaller one. The new trees are to be examined by the algorithm.

<sup>176</sup> We identify an oversight in this procedure, where the algorithm fails to keep track of some <sup>177</sup> trees in  $H_1$ . We demonstrate this by an example. In Figure 1, if the edge between vertices u<sup>178</sup> and v gets deleted, then the connected component is split into two triangles. The **c-degree** of <sup>179</sup> both u and v decreases to two, and as both are greater to one, the deletion procedure stops <sup>180</sup> without any action. However, both triangles are unicylic. If an edge is deleted from either <sup>181</sup> triangle, then Case-1 does not recognize that the remaining edges should be relabeled as  $\mathcal{T}$ <sup>182</sup> not  $\mathcal{F}$ .

If Algorithm 1 is not able to keep track of all the trees in  $H_1$ , then it can exit the loop of Line 4 prematurely. As a consequence Karp–Sipser in Line 16 will return a suboptimal matching. We propose a fix for this oversight in Lemma 1.

**Lemma 1.** Let u be an endpoint of a deleted edge (u, v) with label C. Apply the procedure of Case-1 until we arrive at a vertex p with  $c-degree[p] \neq 1$ . If c-degree[p] = 0, then u's component has become a tree.

**Proof.** We claim that if c-degree [p] = 0, then p and v are the same vertex. Each vertex on 189 the path from u to p had its c-degree affected twice (from 2 to 0), except p. Hence for p to 190 become 0, its c-degree must have been equal to 1. If  $p \neq v$ , then p should had its C edge 191 relabeled during another deletion process. Therefore, prior to the deletion of (u, v), there was 192 a cycle on  $H_1$  with all vertices having c-degree equal to 2, and both their C edges participated 193 in the cycle. Any outgoing edges from vertices of the cycle therefore were labeled  $\mathcal{F}$  and by 194 definition, their deletion led to a tree being formed. The component was hence unicyclic 195 before. 196

<sup>197</sup> Case 1-continuation is therefore as follows:

<sup>198</sup> Once there are no vertices with **c-degree** equal to 1, take the last vertex v whose **c-degree** <sup>199</sup> was reduced. If **c-degree**[v] = 0, then relabel all edges in vs component from  $\mathcal{F}$  to  $\mathcal{T}$ . <sup>200</sup> This addition has overall O(n) cost, because each edge can change label at most twice.

#### <sup>201</sup> 3.1.2 Conversion to an efficient general heuristic

Algorithm 1 works well when the random 2-out graph is sampled from  $K_{n,n}$ . However, in the case of an arbitrary host graph, the underlying theory is not shown to hold, and the algorithm

can make erroneous decisions. Here we discuss how to turn Algorithm 1 into a general 204 heuristic. Apart from the aim of obtaining a practical heuristic for bipartite matching, there 205 is another reason to investigate the matching problem in 2-out bipartite graphs. We show in 206 Appendix D that an O(f(n,m)) time algorithm to find a maximum cardinality matching in 207 a 2-out bipartite graph can be used to find a maximum cardinality matching in any bipartite 208 graph with m edges in O(f(m, m)) time, where f is a function on the number of vertices n 209 and edges m. Such a reduction is important because it shows that an algorithm for finding 210 maximum cardinality matchings in 2-out graphs with similar complexity to 20UTMC can be 211 used to obtain an  $O(m \log m)$  algorithm for matchings in general bipartite graphs. 212

If the algorithm reaches Line 10 during execution, it quits immediately before examining 213 all trees in  $H_1$ . We instead propose to continue with the execution of the algorithm to make 214 the returned matching as large as possible. To achieve this efficiently, we keep for each tree T215 a list  $L_T$  of unmarked vertices. At Line 5 we randomly sample x from  $L_T$  and discard it from 216  $L_T$ . Contrary to Algorithm 1, we neither mark x nor insert it in  $H_2$  yet. Instead, we examine 217 first whether the component in  $H_2$  of either of the two choices of x has an unchecked column 218 y. If y exists, we mark x, insert it to  $H_2$  and continue by deleting y from  $H_1$ . Otherwise, 219 we perform the same set of actions with another randomly sampled vertex from  $L_T$ . If  $L_T$ 220 becomes empty, and no vertex was marked, we abandon T and proceed to another tree. Each 221 such tree in the final state of  $H_1$  decreases the cardinality of the returned matching by one, 222 as a row is left free. If T is split into two trees, the lists of unmarked vertices for the new 223 trees contain only those vertices still inside  $L_T$  at the moment of splitting. This is necessary 224 to avoid sampling vertices more than once. 225

The overall algorithm 20UTMC is as follows. It takes the matrix representation of the 226 given bipartite graph and scales it with a few steps of the Sinkhorn–Knopp algorithm to 227 obtain  $A_{S}$ . It then chooses two random neighbors for each column and row using their 228 respective probability distributions in the corresponding row and column of  $A_S$ , which are 229 given as input to Algorithm 1. Then, the auxiliary graph  $B_{2\rho}$  is constructed and Karp–Sipser 230 is run on this graph to retrieve a maximum cardinality matching in  $B_{2o}$ . If one allows vertices 231 to choose neighbors uniformly, then there are no guarantees on the maximum cardinality of 232 a matching in  $B_{2o}$ . As an example, consider the graph where the *i*th row and *i*th column 233 are connected for  $i = 1, \ldots, n$ , and additionally the first  $\ell$  rows and columns are connected 234 with every vertex on the opposite side. Then, in expectation  $O(\frac{\ell-1}{\ell+1} \cdot n)$  rows (resp. columns) 235 make both choices from the first  $\ell$  columns (resp. rows), such that in the generated  $B_{2o}$  the 236 maximum cardinality matching is of size  $O(\frac{n}{\ell} + \ell)$ . Using **A**<sub>**S**</sub>'s values to perform the random 237 choices spreads the choices so that the maximum cardinality of the matching in the subgraph 238 increases (see Theorem 2 and Lemmas 6–8 in [16] that examines the 1-out subgraph model). 239

In Appendix B we describe two heuristics for 20UTMC which can lead to an increase in the cardinality of the returned matching. The main idea of both heuristics is to reduce the chance that an edge deletion in  $H_1$  creates a new tree.

# 3.2 TRUNCRW: Truncated random walk with nonuniform sampling 3.2.1 Description of the algorithm for regular bipartite graphs

Goel et al. [18] propose a randomized algorithm (of the Las Vegas type) that finds a perfect matching in a *d*-regular bipartite graph with *n* vertices in each side in  $O(n \log n)$  time in expectation. This algorithm starts a random walk from a randomly chosen free column-vertex. At a column vertex *c*, the algorithm selects uniformly at random one of the row-vertices that are not matched to *c*, and goes to the chosen row vertex *r*. If *r* is free, then an augmenting path is obtained by removing possible loops from the walk. If *r* is matched, then the random

$$\begin{pmatrix} \sqrt{2} & & & \\ & \frac{1}{\sqrt{2}} & & \\ & & \frac{1}{\sqrt{8}} & & \\ & & & \frac{1}{\sqrt{8}} & \\ & & & & \frac{1}{\sqrt{8}} \end{pmatrix} \begin{pmatrix} 1 & 1 & 0 & 0 \\ 1 & 1 & 1 & 0 \\ 1 & 1 & 1 & 1 \end{pmatrix} \begin{pmatrix} \frac{1}{\sqrt{8}} & & & \\ & & \frac{1}{\sqrt{8}} & & \\ & & & \frac{1}{\sqrt{2}} & \\ & & & & \sqrt{2} \end{pmatrix} = \begin{pmatrix} 1/2 & 1/2 & 0 & 0 \\ 1/4 & 1/4 & 1/2 & 0 \\ 1/8 & 1/8 & 1/4 & 1/2 \\ 1/8 & 1/8 & 1/4 & 1/2 \end{pmatrix}$$

**Figure 2** The matrix **A** associated with a  $4 \times 4$  Hessenberg matrix, the scaling matrices **D**<sub>R</sub> and **D**<sub>C</sub>, and the resulting doubly stochastic matrix  $\mathbf{A}_{\mathbf{S}} = \mathbf{D}_{\mathbf{R}} \mathbf{A} \mathbf{D}_{\mathbf{C}}$ . In general,  $\mathbf{A}_{\mathbf{S}}(n, 1) = 1/2^{n-1}$ .

walk goes to the mate of r. Goel et al. show that the total length of the random walks is  $O(n \log n)$  in expectation, and thus the algorithm obtains a perfect matching in the stated time [18, Theorem 4]. They also show that one can obtain a Monte Carlo-type algorithm by truncating the random walks. The expected length of an augmenting path with respect to a given matching of cardinality j is 2(4 + 2n/(n - j)), and the random walks could be truncated at this length to obtain near optimal matchings in  $O(n \log n)$  time.

A random walk is easy to implement for d-regular bipartite graphs. At a column vertex  $c_{i}$ 257 one can create a random number between 1 and d in O(1) time and choose the neighbor at 258 that position, and repeat the experiment if the mate of c is chosen. This will take O(1) time 259 in expectation for each step of the walk, and the run time bound of  $O(n \log n)$  is maintained. 260 Goel et al. show that the random-walk based algorithm will work for finding perfect 261 matchings in the bipartite graph representation of a doubly stochastic matrix. They also 262 suggest using an existing data structure [20] when the row and column sums are constant 263 with nonnegative integer entries bounded by a polynomial in n, to attain an  $O(n \log n)$  run 264 time bound. A more recent paper [32] removes the restriction on the entries, and obtains 265 an expected constant time per update and sampling. Further investigations and a careful 266 implementation are necessary to apply the mentioned sampling approaches in our context. 267 Instead, for general doubly stochastic matrices without any bound on the entries, Goel et 268 al. propose an augmented binary search tree with which each selection step of the random 269 walk can be implemented in  $O(\log n)$  time, and obtain a run time of  $O(m + n \log^2 n)$  in 270 expectation, with a total of O(m) preprocessing time. 271

#### **3.2.2** Conversion to an efficient general heuristic

Let c be a free column vertex with respect to a given matching of cardinality j. Assuming there is a perfect matching, one can find an augmenting path to match c, and a random walk can find it. The  $O(\frac{n}{n-j})$  bound on the expected length of such a path will not hold if the bipartite graph is not regular. One may perform more than m steps, which is the worst case time complexity of deterministically finding an augmenting path starting from a free vertex. We propose two methods to make the random walks more useful and to sample efficiently in a random walk. We also discuss an efficient implementation of the whole approach.

The first proposed method is to scale the matrix representation  $\mathbf{A}$  of a given bipartite 280 graph to obtain a doubly stochastic matrix  $\mathbf{A}_{\mathbf{S}}$  for random selections. The expected length 281 of a random walk to find an augmenting path holds when  $A_S$  has bounded nonzero entries. 282 In general, ones does not have any bound on the entries of  $A_{\mathbf{S}}$ . Consider the matrix  $\mathbf{A}$ 283 associated with an upper Hessenberg matrix of size n. A has a full lower triangular part, 284 and additional n-1 entries  $\mathbf{A}(i-1,i) = 1$  for  $i = 2, \ldots, n$ , and fully indecomposable. The 285  $4 \times 4$  example along with its unique scaling matrices are shown in Fig. 2. In the resulting 286 scaled matrix  $\mathbf{A}_{\mathbf{S}}(n,1) = 1/2^{n-1}$  whose inverse is not bounded polynomially in n. 287

As highlighted at the end of Section 3.2, one needs an  $O(\log n)$  time algorithm to select a row vertex randomly from a given column vertex. The second proposed method is a simple yet efficient algorithm for this purpose, rather than a sophisticated augmented tree. The main components of the proposed sampling method are as follows. For each column vertex c, with  $d_c$  neighbors, we have:

- <sup>293</sup> =  $\operatorname{adj}_{c}[1, \ldots, d_{c}]$ : an array keeping the neighbors of c.
- wghts<sub>c</sub>[1,...,d<sub>c</sub>]: the weight of the edges incident on c. This array is parallel to the first one so that the weight of the edge  $(c, \mathsf{adj}_c[i])$  is wghts<sub>c</sub>[i].
- medge[c]: the position of the mate of c in the array  $adj_c$ , or -1 if c is not matched.
- At the beginning, we compute the prefix sum of  $\operatorname{wghts}_c[1,\ldots,d_c]$ . After this operation, the total weight of the edges incident on c is  $\operatorname{wghts}_c[d_c]$ , and the weight of the edge  $(c, \operatorname{mate}[c])$ is  $\operatorname{wghts}_c[\operatorname{medge}[c]] - \operatorname{wghts}_c[\operatorname{medge}[c] - 1]$ , assuming that  $\operatorname{wghts}_c[0]$  signifies zero.
- Given the prefix sums in  $\mathsf{wghts}_c[1,\ldots,d_c]$ , the position of the mate of c at  $\mathsf{medge}[c]$ , we 300 can choose a random neighbor (which is not equal to mate[c]) as shown in Algorithm 2. We 301 use a binary search function, binSearch, which takes an array, the array's start and end 302 positions, a target value, and returns the smallest index of an array element which is larger 303 than the given value with binary search (we skip the details of this search function). At 304 Line 5, since c does not have a mate, we search in the whole list. At Line 8, since the prefix 305 sum just before medge[c] is larger than the target value, we search in the first part of wghts<sub>c</sub> 306 until the current mate located at medge[c]. At Line 10, we search on the right of medge[c], 307 by a modified target value. This last part is the gist of the algorithm's efficiency as it avoids 308 updating the prefix sums when the mate changes. 309
  - **Algorithm 2** Sampling a random neighbor of the column vertex c with  $d_c$  neighbors.

```
Require: \operatorname{adj}_{c}[1,\ldots,d_{c}], wghts<sub>c</sub>[1,\ldots,d_{c}], and medge[c]
 1: mwght \leftarrow wghts<sub>c</sub>[medge[c]] - wghts<sub>c</sub>[medge[c] - 1] if medge[c] \neq -1, otherwise 0
                                                 ▶ The total weight of the edges that can be sampled
 2: totalW \leftarrow wghts<sub>c</sub>[d<sub>c</sub>] - mwght
 3: create a random value rv between 0 and totalW
 4: if medge<sub>c</sub> = -1 then
        return binarySearch(wghts<sub>c</sub>[1,...,d<sub>c</sub>], rv)
 5:
 6: else
 7:
           wghts_c[medge_c] - mwght \ge rv  then
        if
           return binSearch(wghts<sub>c</sub>[1,...,medge[c] - 1], rv)
 8:
 9:
        else
10:
           return binSearch(wghts<sub>c</sub>[medge[c] + 1, ..., d_c], rv + mwght) + medge<sub>c</sub>
```

The sampling algorithm returns the index of the neighbor in  $\operatorname{adj}_c$  different from the current mate in time  $O(\log d_c)$ , independent of the values of the edges. It thus respects the required run time bound. If we were to apply the rejection sampling (as discussed before for the regular bipartite graphs), the run time would depend on the value of the matching edge that we want to avoid. This could of course lead to an expected run time of more than O(n).

There are two key components of Algorithm 2. The first one is the prefix sum, which 315 is computed once before the random walks start and does not change. The second one is 316 medge[c], the position of mate[c] in  $adj_c$ . The value medge[c] changes and needs to be updated 317 when we perform an augmentation. We handle this update as follows. We keep the random 318 walk in a stack by storing only the column vertices, as the row vertices direct the walk to 319 their mate, or terminate the walk if not matched. We discard the cycles from the random 320 walk as soon as they arise—this way we only store a path on the stack, and its length can 321 be at most n. Storing a path also enables keeping the  $medge[\cdot]$  up-to-date. Every time we 322 sample an outgoing edge from a column vertex c, we assign the location of the sampled 323 row vertex in  $adj_c$  to a variable nmedge[c]. When we find a free row, the stack contains the 324 column vertices of the corresponding augmenting path, whose new mates' locations are in 325  $\mathsf{nmedge}[\cdot]$  and thus can be used to update  $\mathsf{medge}[\cdot]$ . 326

The described procedure will work gracefully in expected  $O(m + n \log n)$  time for regular

bipartite graphs and for doubly stochastic matrices where the nonzero values do not differ by 328 large. On the other hand, when there are large differences in edge weights, a random walk can 320 get stuck in a cycle. That is why truncating the long walks is necessary to make the algorithm 330 work for any given doubly stochastic matrix. Furthermore, such a truncation is necessary 331 with the proposed matrix scaling approach for defining random choices. For the overall 332 approach to be practical, we should not apply the scaling algorithms until convergence. As 333 in the previous approaches [15, 16], we allot a linear time of O(m+n) for scaling. Applying 334 Sinkhorn–Knopp algorithm for a few iterations will thus be allowable. The known convergence 335 bounds for the Sinkhorn–Knopp algorithm [27, Thm. 4.5] apply asymptotically, therefore 336 we do not have any bounds on the error after a few iterations; it can be large. That is why 337 truncation makes the random walk based augmenting path search practical. 338

The overall algorithm TRUNCRW is thus as follows. It takes the matrix representation of the given bipartite graph and scales it with a few steps of the Sinkhorn-Knopp algorithm. Then for j = 0 to n - 1, it uniformly at random picks a free column vertex, and starts a random walk starting from that column, for at most 2(4 + 2n/(n - j)) steps, after which the walk is truncated. Some follow discussion and experiments with different parameters for TRUNCRW may be found in Appendix C.

#### 345 **4** Experiments

We implemented 20UTMC and TRUNCRW in C/C++, and the codes are accessible from 346 https://gitlab.inria.fr/bora-ucar/fast-matching. The codes, all are sequential, were 347 compiled with "-O3" and run on a machine with 2 x Intel Xeon CPU Gold 6136 CPUs and 187 348 GB RAM. We evaluate 20UTMC and TRUNCRW both on real-life and synthetic bipartite 349 graphs with equal number of vertices in each side. We compared the two algorithms 350 against KASI, the widely used version of Karp–Sipser which applies degree-1 reduction 351 (own implementation), and KASI2, the original version of Karp–Sipser with both reduction 352 rules. We use a publicly available implementation of KASI2 (https://gitlab.inria.fr/ 353 bora-ucar/karp--sipser-reduction) which is the fastest of recent implementations [26, 354 29]. We note that there are other heuristics (a short summary and further references are 355 in Appendix A) which deliver very good results in practice. For most of these heuristics, 356 especially for those based on vertex degree, there are known worst case upper bounds close 357 to 1/2. We therefore restrict the focus on KASI and KASI2, which are efficient and very 358 effective in practice [14, 25, 30]. We also investigated if random 2-out bipartite graphs of a 359 general host graph have perfect matchings if rows and columns select neighbors with the 360 probabilities in the scaled matrix representation. The quality of a matching refers to the 361 ratio of the cardinality of the matching to the maximum cardinality of a matching in a given 362 graph. The practical version of Sinkhorn-Knopp is referred to as SK-t, where t is the number 363 of allowed iterations. All run times are reported in seconds. 364

#### **4.1** Investigation of perfect matchings in 2-out graphs

Here, we investigate the claim that  $G_2$  will likely have a perfect matching for G, if created with the probabilities in the scaled matrix. We used a set of 39 large sparse square matrices from the SuiteSparse Collection [12], whose bipartite graphs have perfect matchings. These matrices are automatically selected from all square matrices available at the collection with  $10^6 \le n \le 28 \times 10^6$ , and with at least two nonzeros per row or column.

<sup>371</sup> We consider two different models to create  $G_2$ . In the model  $M_1$ , row choices are <sup>372</sup> independent of the column choices. Under this model, a row and a column can select each <sup>373</sup> other resulting in parallel edges—only one of them is kept. The model  $M_2$  tries to avoid

$\frac{m}{n}$	[	0,10)	[] []	10,20)	[4	20,30)	[3	30,40)	[4	10,50)
#Instances		27		5		5		1		1
	# PM	deficiency	#PM	deficiency	#PM	deficiency	#PM	deficiency	#PM	deficiency
Model M <sub>1</sub>	0	223	0	8	1	20	0	2	1	0
Model $M_2$	27	0	3	3	1	10	0	1	0	1

**Table 1** We divide the real-life graphs into five groups. The *i*th group consists of graphs whose  $\frac{m}{n}$  ratio is between 10(i-1) and 10i. For each group, we give the number of instances in which a 2-out graph built using the models M<sub>1</sub> and M<sub>2</sub> has a perfect matching and the largest difference from the maximum cardinality of a matching.

			20UT	мс	Trunc	RW
h	KaSi	KAS12	Uniform	SK-5	Uniform	SK-5
2	0.93	1.00	0.78	0.99	0.88	0.99
8	0.80	0.85	0.59	0.99	0.91	0.99
32	0.69	0.72	0.52	0.99	0.83	0.99
128	0.64	0.65	0.51	0.99	0.78	0.99
512	0.61	0.63	0.52	0.99	0.76	0.99

**Table 2** Average quality of the matchings found by the algorithms on graphs from the synthetic family  $\mathcal{I}$  for n = 30000 and various values of h.

parallel edges. In this model, all columns perform their selections. Then, each row r attempts 374 to randomly choose two columns, only from those that did not select r. These selections again 375 are based on the scaled matrix. In this model, parallel edges can arise (and be discarded) 376 only when a vertex v is connected in the 2-out graph with all of its neighbors in G, because 377 it is impossible for v to select otherwise. We experimented three times with each real-life 378 graph.  $M_i$ 's result is the maximum of those three experiments. In each test, we first created 379 the choices of all columns. Then we allowed the two models to generate the choices of the 380 rows accordingly. 381

The results are shown in Table 1 for the 39 real-life graphs and are with SK-5. As seen in this table, the random  $G_2$  graphs generated with the model  $M_1$  have near perfect matchings, but they do not contain perfect matchings in most cases. In contrast, the random  $G_2$  graphs generated by  $M_2$  in many cases contain a perfect matching. In only a few graphs this does not hold true, and in these cases the deficiency is no more than 10.

#### **387** 4.2 On synthetic graphs

In Table 2, we give results with a synthetic family  $\mathcal{I}$  of graphs from literature [16], whose matrix representations do not have total support. To create a member of  $\mathcal{I}$ , we separate the vertex set R into  $R_1 = \{r_1, \ldots, r_{n/2}\}$  and  $R_2 = \{r_{n/2+1}, \ldots, r_n\}$  and likewise for C. All vertices of  $R_1$  are connected to all vertices of  $C_1$ . Edges  $(r_i, c_{n/2+i})$  and  $(r_{n/2+i}, c_i)$  for  $i = 1, \ldots, n/2$  are added to introduce a perfect matching. A parameter h is used to connect h vertices from  $R_1$ , and h vertices from  $C_1$  to every vertex on the opposite side.

As seen in Table 2, KASI and KASI2 have more and more difficulty with increasing h. 394 The matching quality drops over 30% between h = 2 and h = 512 for KASI and almost 40% 395 for KAS12. On the contrary, 20UTMC and TRUNCRW both obtain a near perfect matching, 396 with SK-5. Even though the matrices associated with the graphs of  $\mathcal{I}$  lack total support, 397 SK-5 sufficed to obtain near optimal matchings. We notice the effect of scaling: if vertices 398 select without scaling (Uniform), the matching quality reduces. This is particularly true 399 for 20UTMC, which exhibits the worst overall performance with uniform selection. Family 400  $\mathcal I$  shows the importance of scaling, and more importantly highlights the robustness of the 401 proposed methods. An adversary can create graphs which make degree-based randomized 402

	KaSi	KaSi2	2	outMC		Τı	RUNCRW	7
n	quality	quality	uniform	SK-5	SK-20	uniform	SK-5	SK-20
10000	0.76	0.84	0.81	0.92	0.95	0.97	0.97	0.97
20000	0.73	0.83	0.81	0.92	0.95	0.97	0.97	0.97
30000	0.73	0.83	0.81	0.92	0.95	0.97	0.97	0.97

**Table 3** Average quality of the matchings found by the algorithms on graphs from the synthetic family  $\mathcal{J}$  for  $n \in \{10000, 20000, 30000\}$ .

<sup>403</sup> approaches lose quality—some of those heuristics are briefly mentioned in Appendix A, and <sup>404</sup> the full details including negative results on KASI2 can be found elsewhere [9]. On the other <sup>405</sup> hand, the use of scaling helps to avoid such cases for 20UTMC and TRUNCRW.

We now discuss another synthetic family of graphs  $\mathcal J$  in which the proposed approaches 406 obtain matchings of much higher quality than KASI and KASI2. A bipartite graph with n407 vertices per side belonging to  $\mathcal{J}$  contains the following edges:  $(r_i, c_j)$  for all  $i \leq j$ ;  $(r_2, c_1)$ , 408  $(r_n, c_{n-1}); (r_3, c_1), (r_3, c_2), (r_n, c_{n-2});$  and  $(r_{n-1}, c_{n-2})$ . The graphs in  $\mathcal{J}$  are hard for 409 Karp–Sipser-based heuristics because only few of the edges participate in a perfect matching, 410 the deterministic rules do not apply, and hence they resort to multiple suboptimal random 411 decisions. Likewise, due to the large number of entries without support in the matrix 412 representation, Sinkhorn–Knopp will take many iterations to properly scale the matrix. 413

In Table 3, we give results of the algorithms for a few graphs from this family. In the 414 table, we also show the effects of scaling on 20UTMC and TRUNCRW by showing results 415 without scaling (under column "uniform", in which a column vertex chooses a neighbor 416 uniformly at random), with SK-5, and with SK-20. As can be seen, despite the lack of 417 total support, both 20UTMC and TRUNCRW obtain matchings whose cardinality is more 418 than 0.92 of the maximum, when SK-5 or SK-20 is used. TRUNCRW in particular is nearly 419 optimal. These results are always better than that of KASI and KASI2, with the difference 420 in matching quality being about 20-25% for the former, and 10-15% for the latter. With 421 increased iterations of Sinkhorn-Knopp, 20UTMC increases the cardinality of its matchings 422 by 3%. If we do not use scaling ("uniform"), while there's no noticeable effect on TRUNCRW's 423 matchings, 20UTMC matchings decrease by roughly 10%. Even so, its results remain better 424 than KASI's and on par with those of KASI2. 425

#### 426 4.3 On real-life graphs

We compared TRUNCRW and 20UTMC with KASI and KASI2 on all 39 real-life graphs from 427 Section 4.1. Figure 3a and Figure 3b present the high level picture. For the experiments, we 428 did not permute the matrices randomly, which generally increases the experimentation time. 429 The results for matching quality can be seen in Figure 3a, where we plot the ratio of the 430 cardinality of the matchings found by different algorithms to the maximum cardinality of 431 the matching. The graphs are indexed in nondecreasing number of edges. 20UTMC and 432 TRUNCRW use SK-3 for scaling. As can be observed, both 20UTMC and TRUNCRW obtain 433 near perfect matchings. The average matching quality obtained by 20UTMC is 0.9979 and 434 that obtained by TRUNCRW is 0.9984. Both algorithms never drop below 0.9900 in any of 435 the 39 cases. 436

Figure 3a also shows the matching quality of KASI2 and KASI. KASI obtains matchings
of quality 0.9862 on average, with always smaller cardinality than TRUNCRW and 20UTMC.
KASI2 fares better and its average quality is 0.9968. Even so, in the majority of cases, it
obtains matchings that are inferior quality-wise to both TRUNCRW and 20UTMC.

While all algorithms obtain matchings of high quality, the absolute different is remarkable in some cases. For example, the largest difference observed between the matching cardinalities



(a) Quality results

443

(b) Run time results

**Figure 3** Quality (left) and run time (right) results for all 39 graphs from Section 4.1.

obtained by 20UTMC and KASI was 346577, in favor of 20UTMC.

Figure 3b shows the run time of all examined heuristics, where the graphs are again indexed in nondecreasing number of edges. KASI is in general the fastest of these four algorithms when there are not too many edges. TRUNCRW and 20UTMC are close run-time wise to KASI and in some instances faster than it. This is especially true in instances with many edges because KASI depends more on *m*. KASI2 has the slowest performance overall.

For a detailed study, we show results on the five largest graphs from the mentioned dataset and Circuit5M, which was identified as a challenging instance in earlier work [25]. Degree-1 vertices from Circuit5M are removed by applying Rule-1 of KAS12 as a preprocessing step—this is without loss of generality of the heuristics. For each graph we relabeled its row-vertices randomly and executed five tests with each algorithm.

Table 4 shows the matching quality and the run time of the four heuristics. 20UTMC and TRUNCRW used SK-3 for this set of experiments for speed. For each graph, we give the minimum, maximum, and averages over five runs. As already discussed, all heuristics obtain high quality matchings. On a closer look, we see that TRUNCRW, on average, matched 158410 more edges than KASI, and 50847 more edges than KASI2. Similarly 20UTMC matched 139220 more edges than KASI on average, and 31652 more edges than KASI2. Interestingly, on graph Channel-500 TRUNCRW was able to find the maximum matching.

Concerning run time, as KASI is a linear time heuristic it is expected to be the fastest. 461 Surprisingly, TRUNCRW even with the scaling time added is faster than KASI in three 462 instances. This is due to the fact that each iteration of the scaling algorithm takes linear time 463 with small constants. As an algorithm on its own (without scaling time), TRUNCRW becomes 464 the fastest one, thanks to its run time not depending on m after the initialization. 20UTMC, 465 though slower, also exhibits good behavior, except in nlpkkt240. KASI2 has the worst run 466 time overall. Its initialization takes more time, and its implementation is more involved. 467 SK-3 is fast except for nlpkt240 where it requires about 30 seconds. The reason that SK-3 468 requires 30 seconds for this particular graph is due to the random permutation of its rows, 469 which is not cache-friendly (if SK-3 is run on nlpkkt240 using the initial ordering of rows, it 470 finishes in less than 10 seconds). In the other cases and despite the large size of the graphs, 471 scaling finishes in less than seven seconds. Table 4 additionally shows that TRUNCRW and 472 20UTMC's run time performance does not seem to be affected by their random decisions. 473 The largest difference between the result of the minimum, and the maximum run is no more 474 than two seconds for both of these algorithms. 475

			KA	Sı	KA	S12	SK-3	20ut	MC	TRUN	CRW
name	n	statistics	quality	time	quality	time	time	quality	time	quality	time
		min.	0.99	12.67	0.99	26.89	4.59	0.99	8.82	0.99	8.27
cage15	5.15	avg.	0.99	12.81	0.99	27.08	4.68	0.99	8.88	0.99	9.32
		max.	0.99	13.17	0.99	27.27	4.83	0.99	8.96	0.99	10.23
		min.	0.99	10.12	0.99	20.63	2.74	0.99	7.63	1.00	3.86
Channel-500	4.80	avg.	0.99	10.16	0.99	20.94	2.75	0.99	7.66	1.00	4.48
		max.	0.99	10.18	0.99	21.87	2.75	0.99	7.70	1.00	5.11
		min.	0.99	6.57	0.99	24.74	2.45	0.99	4.40	0.99	2.07
Circuit5M	5.55	avg.	0.99	6.76	0.99	24.93	2.84	0.99	4.56	0.99	2.19
		max.	0.99	7.03	0.99	25.33	4.16	0.99	4.81	0.99	2.35
		min.	0.99	11.58	0.99	65.97	4.32	0.99	23.34	0.99	11.21
Delaunay_24	16.00	avg.	0.99	11.61	0.99	68.30	4.44	0.99	23.58	0.99	11.31
		max.	0.99	11.66	0.99	72.47	4.48	0.99	24.38	0.99	11.37
		min.	0.99	14.97	0.99	91.42	6.26	0.99	30.96	0.99	14.25
Hugebub-20	21.19	avg.	0.99	15.04	0.99	97.77	6.29	0.99	31.28	0.99	14.38
		max.	0.99	15.15	0.99	106.78	6.31	0.99	31.59	0.99	14.57
		min.	0.98	98.58	0.99	182.08	29.77	0.99	52.34	0.99	34.34
nlpkkt240	27.99	avg.	0.98	98.66	0.99	183.10	29.92	0.99	52.53	0.99	34.50
		max.	0.98	98.76	0.99	186.08	30.27	0.99	52.76	0.99	34.70

**Table 4** Full run time comparisons with heuristics for the graphs of Section 4.3. The run time of SK-3 should be added to TRUNCRW and 20UTMC. For each instance we give the minimum, the average, and the maximum of five runs for all columns regarding the quality and the run time. The number of vertices n per side is in the order of millions. Hugebub-20 stands for Hugebubbles-0020.

476 Combined with the results in the previous section, we conclude thus that (i) 20UTMC
477 and TRUNCRW always obtain near perfect matchings, while KASI and KASI2 are not as
478 robust; (ii) 20UTMC and TRUNCRW are nearly as fast as the linear time algorithm KASI,
479 and are much faster than KASI2.

Next, we consider the impact of our heuristics as initialization to an exact algorithm for 480 finding a maximum cardinality matching. We first run the heuristics to obtain an initial 481 matching, then call an exact algorithm to augment the initial matchings for maximum 482 cardinality. We consider three different exact algorithms MC21, PR, and PF+ for the 483 augmentation steps. MC21 [13] from mmaker [14, 25] visits free vertices one by one and 484 tries to match the visited vertex with a depth-first search, and hence is closely related 485 to TRUNCRW. In this setting, differences among the qualities of initial matchings should 486 be observable while computing an exact matching. PR [25] is based on the Push-Relabel 487 method [19], and PF+ which is a depth-first search based method [14, 36]. The last two 488 algorithms are more elaborate than MC21, and the cardinality difference between two different 489 initial matchings does not necessarily correlate with the run time. 490

The statistics of five runs with MC21 are given in Table 5. In this table, the time 491 spent in augmentations is given in column "augment.". The overall time to compute a 492 maximum cardinality matching is given in column "overall', which includes the time spent in 493 heuristics—in case of 20UTMC and TRUNCRW it includes the scaling time as well. The 494 runs on nlpkkt240 did not finish within an hour and are not presented. As seen in the 495 table, the overall time to obtain a maximum cardinality matching is always the smallest 496 with TRUNCRW initialization. 20UTMC is usually competitive with the faster of KAS12 497 and KASI, without a clear winner. It is also interesting to note that in all graphs the worst 498 behavior of TRUNCRW is better than the best behavior of KASI2 and KASI and in some 499 cases (see cage15 or Channel-500) significantly so. The same is almost true for 20UTMC as 500 well except for graphs Delaunay\_24 and Hugebbubles-0020 where 20UTMC's worst result 501 is only a few seconds slower than KASI's best result, or cage15 versus KASI2. 502

		KaSi		KaSi2		20utMC		TRUNCRW	
name	statistics	augment	overall.	augment	overall.	augment	overall.	augment	overall
	min.	133.85	146.52	7.42	34.47	27.29	40.75	0.22	14.07
cage15	avg.	140.13	152.94	8.81	35.90	31.44	45.00	1.85	15.84
	max.	144.42	157.28	10.70	37.59	37.84	51.47	2.46	16.84
	min.	64.29	74.46	9.15	29.81	12.18	22.62	0.04	6.65
Channel-500	avg.	71.61	81.76	10.93	31.86	15.28	25.68	0.14	7.36
	max.	78.81	88.98	11.71	33.58	18.84	29.25	0.25	8.11
	min.	14.33	20.94	10.51	35.32	4.38	12.21	0.50	5.02
Circuit5M	avg.	15.26	22.01	13.11	38.04	5.70	13.09	0.77	5.80
	max.	16.00	22.72	14.42	39.43	6.81	13.68	1.31	7.79
	min.	49.95	61.54	26.93	94.02	35.10	63.71	26.77	42.49
Delaunay_24	avg.	54.79	66.40	29.99	98.29	36.68	64.70	31.06	46.81
	max.	61.23	72.81	32.70	104.13	40.30	68.11	34.09	49.77
Hugebub-20	min.	68.17	83.14	55.79	148.64	44.83	82.31	42.02	62.56
	avg.	73.15	88.20	58.95	156.72	50.65	88.21	44.54	65.21
	max.	75.99	91.10	61.18	166.98	54.35	91.60	47.11	67.68

**Table 5** Detailed run times when MC21 is used for augmentations on the graphs described in Section 4.3. The quality of heuristics are in Table 4. We have omitted graph nlpkkt240 for which MC21 did not finish within a reasonable amount of time. For each instance we give the minimum, the average, and the maximum run time of five runs. Hugebub-20 stands for Hugebubbles-0020.

In Table 6, we observe the behavior of the heuristics when used for initializing the PF+ 503 algorithm. The table shows the minimum, average, and maximum time over the five runs. As 504 can be observed, TRUNCRW exhibits the best overall behavior. TRUNCRW has the fastest 505 performance in four out of six instances, and in the remaining two instances it is very close 506 to KASI. The largest difference between the two can be observed in nlpkkt240 where KASI 507 is overall almost 50 seconds slower. The total run time with KAS12 is never better than that 508 with TRUNCRW. It roughly takes the same amount of time for PF+ to augment 20UTMC's 500 initial matching, as it takes for it to augment the matching of TRUNCRW. Therefore, when 510 20UTMC has a run time similar to TRUNCRW their overall run times are similar. In the 511 largest of instances 20UTMC's and TRUNCRW's performance diverge, but 20UTMC's overall 512 behavior is superior to KASI2 and competitive with that of KASI. 513

In Table 7, we observe the behavior of the heuristics when used for initializing the PR algorithm. The behavior of KASI in Circuit5M demonstrates the robustness of our approaches. The average behavior of PR initialized with KASI is 339 seconds with the maximum run time exceeding 500 seconds. In stark contrast, PR with TRUNCRW's input never needs more than 25 seconds, whereas with 20UTMC it never surpasses 150 seconds. In the remaining instances, the proposed algorithms are competitive with KASI or even faster.

In summary, the effects of the proposed methods as an initialization routine are more 520 observable with MC21 on all instances. With PF+, we see that the augmentations take 521 less time on average with 20UTMC and TRUNCRW, but the overall time with KASI can 522 be sometimes better than that of TRUNCRW slightly thanks to KASI being faster. When 523 PR is used, the augmentations take less time with KASI in three instances compared to 524 TRUNCRW; and in four instances compared to 20UTMC. When 20UTMC and TRUNCRW 525 serve better than KASI as an initialization to PR, the difference is more significant. The 526 above results with three different algorithms demonstrate the merits of the two proposed 527 algorithms for use as initialization routines in exact matching algorithms. 528

		KaSi		KAS12		20UTMC		TRUNCRW	
name	statistics	augment.	overall	augment.	overall	augment.	overall	augment.	overall
	min.	2.19	14.89	2.11	29.18	1.90	15.46	0.73	14.22
cage15	avg.	2.51	15.33	2.59	29.67	1.97	15.53	1.16	15.15
	max.	2.98	16.15	3.16	30.43	2.01	15.69	1.55	15.63
	min.	1.70	11.84	1.82	22.50	1.19	11.60	0.04	6.66
Channel-500	avg.	1.91	12.06	2.07	23.01	1.30	11.71	0.04	7.27
	max.	2.60	12.77	2.89	23.69	1.40	11.84	0.05	7.90
	min.	0.63	7.20	0.45	25.28	0.45	7.34	0.48	5.01
Circuit5M	avg.	0.77	7.53	0.62	25.55	0.53	7.93	0.58	5.61
	max.	0.97	7.97	0.90	25.92	0.67	9.55	0.64	7.04
	min.	18.47	30.06	13.88	80.75	14.24	42.05	14.20	29.92
Delaunay_24	avg.	20.83	32.44	14.89	83.19	15.47	43.49	17.67	33.41
	max.	22.33	33.91	16.17	86.35	17.12	44.98	20.40	36.09
	min.	23.09	38.09	14.99	106.41	23.27	60.75	21.97	42.54
Hugebub-20	avg.	28.13	43.17	19.63	117.40	26.97	64.53	24.49	45.16
	max.	34.11	49.26	23.00	127.49	30.38	68.17	29.65	50.53
	min.	27.01	125.69	28.19	210.27	14.91	97.26	13.76	77.87
nlpkkt240	avg.	27.09	125.76	29.63	212.73	17.56	100.01	13.96	78.38
-	max.	27.24	125.83	30.27	216.15	20.99	103.47	14.09	79.06

**Table 6** Detailed run times when PF+ is used for augmentations on the graphs described in Section 4.3. The quality of heuristics are in Table 4. For each instance we give the minimum, the average, and the maximum run time of five runs. Hugebub-20 stands for Hugebubbles-0020.

		KaSi		KaSi2		20utMC		TRUNCRW	
name	statistics	augment.	overall	augment.	overall	augment	overall	augment	overall
	min.	2.15	14.85	3.63	30.52	1.19	14.67	1.10	14.03
cage15	avg.	2.41	15.22	3.80	30.88	1.39	14.95	1.28	15.28
	max.	2.68	15.85	4.01	31.08	1.69	15.32	1.69	16.59
	min.	1.57	11.75	2.83	23.47	1.63	12.03	0.04	6.68
Channel-500	avg.	1.66	11.81	2.92	23.86	1.75	12.16	0.06	7.28
	max.	1.70	11.85	3.01	24.88	2.02	12.44	0.08	7.92
	min.	116.67	123.24	107.51	132.34	2.02	8.89	0.74	5.26
Circuit5M	avg.	332.29	339.05	235.54	260.47	37.11	44.51	5.37	10.40
	max.	559.09	566.09	378.31	403.12	139.61	148.58	18.30	24.78
	min.	40.52	52.15	32.09	98.89	41.66	69.52	48.63	64.32
Delaunay_24	avg.	45.48	57.09	36.90	105.20	46.94	74.96	52.48	68.23
	max.	52.47	64.06	43.74	110.18	53.19	81.04	58.07	73.91
	min.	41.01	56.16	55.22	146.78	44.71	81.96	49.46	70.34
Hugebub-20	avg.	47.53	62.58	58.56	156.33	51.59	89.15	53.16	73.84
	max.	52.59	67.56	61.17	166.57	58.54	96.15	54.82	75.36
	min.	13.98	112.59	22.87	205.18	15.49	97.63	19.74	84.26
nlpkkt240	avg.	14.13	112.80	24.17	207.27	17.34	99.79	28.70	93.13
	max.	14.51	113.27	25.77	211.10	19.01	101.46	47.31	112.28

**Table 7** Detailed run times when PR is used for augmentations on the graphs described in Section 4.3. The quality of heuristics are in Table 4. For each instance we give the minimum, the average, and the maximum run time of five runs. Hugebub-20 stands for Hugebubbles-0020.

#### 529 **5** Conclusions

We have examined two randomized algorithms for the maximum cardinality matching problem in bipartite graphs. These algorithms originally were designed for two very special classes of bipartite graphs. We have discussed how to convert them into efficient and effective heuristics. Our experimental results show that these approaches obtain near perfect matchings in real-life and synthetic instances and have a near linear time run time. The two approaches are also shown to be more robust than the state of the art heuristics used in the cardinality matching algorithms, and are generally more useful as initialization routines.

Our adaptation of 20UTMC is based on the premise that 2-out graphs sampled from a host graph have perfect matchings, assuming that the matrix representation of the host graph have total support. We showed evidence that this may be true and even if not, the sampled graphs have close to perfect matchings. A proof or the disproof of such 2-out graphs having perfect matchings is certainly welcome. Furthermore, this was the first attempt to implement 20UTMC, and there is room for improved performance.

#### <sup>543</sup> **A** Other heuristics for bipartite matching and recent work

In the main text, we compared the proposed heuristics with KASI and KASI2. There are a few other effective heuristics, which we briefly review here (see a recent survey [37]).

Hopcroft and Karp's original algorithm [21] proceeds in phases. At each phase, it finds 546 shortest augmenting paths, and augments the current matching along a maximal set of 547 disjoint such paths, where each phase runs in O(n+m) time. Stopping when the shortest 548 augmenting paths is of length 2k + 1 at a phase no larger than k results in an 1 - 1/(k + 1)549 approximate matching in O(k(m+n)) time in the worst case. Greedy [39] chooses a random 550 edge and matches the two endpoints and discards both vertices and the edges incident on 551 them. Modified Greedy [39] chooses a free vertex and then randomly matches it to one of 552 the available neighbors. MinGreedy [39] (see also Magun [31] and Langguth et al. [30] for 553 related algorithms) improves upon Modified Greedy by selecting a random vertex with the 554 minimum degree at the first step. The Greedy-like algorithms obtain maximal matchings 555 and therefore are 1/2 approximate. Slight improvements in the form of  $1/2 + \varepsilon$  are shown 556 for these algorithms [2, 35], but there are theoretical bounds in the same vicinity [9]. Duff et 557 al. [14] and Langguth et al. [25, 30] compare these algorithms for initialization in maximum 558 cardinality matching algorithms and suggest using KASI as initialization for general problems 559 especially with the push-relabel based algorithms. 560

Another class of heuristics use randomization for breaking the 1/2 barrier. RANKING [24] algorithm achieves an approximation ratio of 1 - 1/e, where e is the base of the natural logarithm. The same approximation ratio is also achieved by a very simple parallel algorithm [16] whose most involved step is the application of a matrix scaling algorithm. This last paper also proposes an algorithm based on sampling 1-out subgraphs of a general bipartite graph (as we did in this paper) to obtain matchings of size about 0.86 times the maximum cardinality.

Matching has stirred some recent interest in the theoretical computer science community, with works focusing on parallel and distributed settings [4, 5, 11, 3] or on the fully dynamic version [6, 8] among others. Among the recent work, a method by Assadi et al. [4] shares similarities with the 20UTMC algorithm. Their approach similarly sparsifies a given graph G to produce a subgraph with some approximation guarantees for the maximum cardinality matching. A detailed experimentation with this sparsification approach will reveal useful.

#### <sup>574</sup> **B** Further comments on 20UTMC

As demonstrated in the experiments in Section 4, 20UTMC obtains matchings of very high cardinality. We can improve its matching quality by the following two heuristics. These two heuristics are not used in the given experiments. We plan to improve their run time.

#### 578 B.1 Heuristic 1: Delayed tree vertex selection during Line 5

The ideal case at Line 5 of Algorithm 1 is to select an x such that x's insertion as an edge to  $H_2$  does not lead to a new tree in  $H_1$  after the deletion of the edge corresponding to the unchecked vertex of the connected component  $Q_x$ . This is only possible if  $Q_x$  contains an unchecked column labeled as C in  $H_1$ . Otherwise, a new tree will be created in  $H_1$ , and the algorithm will have to process it in a future step. For the first heuristic, we greedily select an x such that, if possible, the creation of a tree in  $H_1$  is avoided.

We replace  $L_T$  is with two lists  $L_T^1$  and  $L_T^2$ . The lists  $L_T^1$  contains those unmarked vertices of T whose insertion in  $H_2$  leads to a new tree;  $L_T^2$  contains all other  $L_T$  vertices that have not been tried yet. At first, we sample x from  $L_T^2$  and see whether the components of x's choices in  $H_2$  have an unchecked vertex of type C in  $H_1$ . If they have, x is marked and inserted to  $H_2$ . Otherwise, x is inserted in  $L_T^1$ , and we consider another random vertex of  $L_T^2$ . If  $L_T^2$  becomes empty, we start sampling from  $L_T^1$ .

With the union-find data structure, this heuristic requires constant amortized time per sample and each vertex can be sampled at most twice. Therefore the overhead associated with this heuristic is almost linear in *n*.

#### <sup>594</sup> B.2 Heuristic 2: Online creation of the RG multigraph

In this heuristic, the decisions of the rows are not given as input, but are instead defined during the course of the algorithm. Similar to the previous idea, this heuristic aims to reduce the possibility that a tree in  $H_1$  gets created following an edge insertion into  $H_2$ .

More specifically, consider a vertex x randomly chosen at Line 5. In this heuristic, x598 has not picked its two choices yet, and we let x choose them at this point, in the way that 599 benefits the algorithm the most. This is done as follows. Initially, we iterate over all of x's 600 neighbors in the host graph G. Let c be one of x's neighbors and  $c^*$  be the sole unchecked 601 vertex in c's connected component in  $H_2$ , or  $c^* = -1$  if no unchecked vertices exist. We 602 assign values to x's neighbors to classify them. If  $c^*$  is equal to -1, c's value is 0. If  $c^*$  has 603 label  $\mathcal{F}$  or  $\mathcal{T}$  in  $H_1$ , c's value is 1. Otherwise, c's value is 2. Based on these assigned values, 604 we partition the neighbors of x in G into three disjoint sets  $C_0$ ,  $C_1$  and  $C_2$  such that  $C_i$ 605 contains all neighbors of x with value equal to i. Selecting columns from  $C_2$  is preferred, as 606 they can avoid creating a tree in  $H_1$ . Vertex x will attempt to sample first from  $C_2$ , and if 607 needed from  $C_1$  or  $C_0$ , with a preference for  $C_1$  over  $C_0$ . The sets  $C_0$ ,  $C_1$  and  $C_2$  are kept 608 implicitly, and each vertex x requires amortized  $O(d_x)$  to make its choices, where  $d_x$  is its 609 degree. Hence, the overhead associated with this heuristic is almost linear in m. 610

#### 611 **B.3 Comparison with** 20UTMC

Here, we briefly discuss the effects that the above two heuristics have on the performance of the
20UTMC algorithm. Since 20UTMC obtains high quality results, the two heuristics can only
yield a relatively small improvement. When they are enabled and used with SK-5 20UTMC
finds matchings with average quality of 0.9997 for the real-world graphs from Section 4.3 for

which 20UTMC obtained matchings of quality 0.9983. This difference corresponds to about 616 13113 additionally matched edges, and hence signals that 13113 augmentations are avoided. 617 It is also interesting to consider the effects that these heuristics can have on cases where 618 20UTMC did not deliver near-optimal matchings. As an example, we consider the synthetic 619 family  $\mathcal{J}$  from Section 4.2. When scaling was not enabled, 20UTMC found matchings of 620 average cardinality 0.80 - 0.81% of the maximum. If however one uses the two heuristics 621 proposed in this section, then there is a significant improvement in performance, and 20UTMC 622 finds matchings of cardinality 0.89 of the maximum. 623

#### <sup>624</sup> **C** Further comments on TRUNCRW

We incorporated a known heuristic called look-ahead [13, 14] for speeding up the augmenting 625 path search in practice. All our experiments with TRUNCRW in Section 4 were with the 626 look-ahead approach. In this heuristic, before sampling an arbitrary row-vertex from a 627 column-vertex c, we check if there is a free row vertex in the adjacency list of c. If so, such a 628 row is returned, and the random walk terminates. The implementation of this heuristic has 629 a total overhead of O(m) for the whole course of the algorithm [13, 14]. We note that the 630 look-ahead technique trades the quality of TRUNCRW with run time. In our experiments, the 631 look-ahead heuristic reduced the run time significantly; it interferes with the randomization 632 though. 633

We can easily apply TRUNCRW to bipartite graphs with different number of vertices in each side. This is based on the fact that we can scale a rectangular  $n_1 \times n_2$  matrix (say  $n_1 \ge n_2$ ) so that all columns have sum of 1, and all rows have equal sum of  $n_2/n_1$ , if there is matching covering all columns, and all entries can be put in such a matching. Then, all components of TRUNCRW work without any change.

If there is no total support, then Sinkhorn–Knopp works in such a way that the entries 639 that cannot put into a perfect matching tend to zero. This is helpful in TRUNCRW's context, 640 as the corresponding edges will not likely be selected in a random walk. If there is no perfect 641 matching, then little is known about scaling. It is our experience that the Sinkhorn–Knopp 642 iterations tend to zero out entries that cannot be put into a maximum cardinality matching. 643 Therefore, in this case again, scaling, random selection, and truncation should help. We 644 present some experiments to support this observation and leave the question of showing this 645 theoretically as an open problem. 646

We experimented with bipartite graphs without total support which correspond to square 647  $(10000 \times 10000)$  and rectangular matrices  $(12000 \times 10000)$  with a uniform nonzero distribution. 648 These matrices are generated with sprand command of Maltab and have about  $d \times 10000$ 649 nonzeros for d = 2, 3, 4, 5. The matrix representation of the bipartite graphs were scaled with 650 10 iterations of SK. For each d, we created five random matrices and ran TRUNCRW on the 651 corresponding five instances. We report the worst quality of the five instances in Table 8. As 652 seen in this table, TRUNCRW works just fine for this case. We did not report in the table 653 but with increased SK iterations, the results improve, which is in accordance with earlier 654 work [16]. 655

#### 656 C.1 Engineering TRUNCRW

<sup>657</sup> The experiments here are on real-life instances from Subsection 4.3 and with SK-5.

Recall that TRUNCRW tries to find an augmenting path starting from a column vertex a certain number of times before giving up and moving to the next column vertex. When we allowed TRUNCRW just a single attempt, it was unable to find a perfect matching in any

	1000	$0 \times 10000$	$12000\times10000$			
d	sprank	TRUNCRW	sprank	TRUNCRW		
2	7787	0.9888	8724	0.9919		
3	9266	0.9697	9667	0.9958		
4	9761	0.9828	9899	0.9995		
5	9918	0.9922	9973	1.0000		

**Table 8** The quality of TRUNCRW on bipartite graphs without perfect matchings.

of the cases, and its average matching quality was 0.9984. When we allowed five attempts,
TRUNCRW found a perfect matching for 13 graphs, and its average matching quality was
0.9999. With 10 attempts, it managed to find a perfect matching in 5 additional graphs.
This verifies that allowing more attempts indeed improves the performance of the algorithm.
The drawback, however, was the increased run time, which we did not think worth. That is
why our implementation of TRUNCRW starts a random walk from a vertex only once.

We also test the effects of the look-ahead mechanism. Let us define the walk efficiency of 667 TRUNCRW as the ratio of the cardinality of the matching found to the total length of the 668 random walks. The higher this ratio, the more useful the random walks are. We evaluate 669 the walk efficiency on a set of seven instances (real-life instances having at most 10000000 670 edges). We test both with and without scaling and report the results of the 14 tests. In 13 671 cases, the look-ahead mechanism improved the walk efficiency. The geometric mean (of 14 672 cases) of the ratios of walk efficiencies with look-ahead to that of without was 1.37. In the 673 case where the look-ahead did not help (ratio was 0.71 in an instance named Hamrle3), the 674 maximum deviation of a row or column sum from one after SK-5 was 0.28, which is high. 675 We conclude that the look-ahead mechanism is very helpful. 676

Finally we test the effects that the length of the augmenting walk has on TRUNCRW. We doubled the allowed length of a random walk to 4(4 + 2n/(n-j)). On average, the matching quality rose from 0.9984 to 0.9998. This modification was not able to find a perfect matching in any of the 39 instances. This led to an increase in the run time, which we deemed too large. We therefore keep 2(4 + 2n/(n-j)) as the truncation length.

#### **D** Reducing bipartite graph matching to matching on 2-out graphs

Here, we prove our claim in Section 3.1 that bipartite matching can be reduced to matching on a 2-out bipartite graph. Let  $G = (V_G, E_G)$ , with be a graph with minimum degree at least two. If G's minimum degree is one, we can apply the first deterministic rule of Karp–Sipser to match degree-1 vertices with their neighbors and consider as G the resulting graph.

We produce a new graph G' from G in the following way. For any edge  $e = (a, b) \in E$ we add edges  $e' = (a, a_e), e'' = (a_e, b_e)$ , and  $e''' = (b_e, b)$  to G'. We hence introduce two new vertices  $a_e, b_e$  s.t  $d_{G'}(a_e) = d_{G'} = 2$  for each edge  $e \in E_G$ . The degree of nodes in  $V_G$ remains unchanged in G'.

<sup>691</sup> ► Lemma 2. Let H be a random 2-out subgraph G'. Then H = G'.

<sup>692</sup> **Proof.** The added vertices  $a_e, b_e$  have degree two and will select both neighbors, hence no <sup>693</sup> edge will remain unpicked.

<sup>694</sup> In what follows, we refer to the second reduction rule of Karp–Sipser which merges the <sup>695</sup> neighbors of a degree-2 vertex, which is then discarded, as a degree-2 reduction.

**Lemma 3.** It is possible to obtain G by doing only degree-2 reductions on G'.

**Proof.** Let  $a_e$  be a vertex of G', introduced due to the edge e = (a, b). Since  $d_{G'}(a_e) = 2$  we can apply a degree-2 reduction which will merge a with  $b_e$  to create a single node  $ab_e$ . As a consequence of this merge, the edge  $(ab_e, b)$  will be created and edges  $(a, a_e), (a_e, b_e), (b_e, b)$ will be erased. We simply relabel  $ab_e$  to a again. The proof then follows similarly by applying degree-2 reduction for all  $a_e$  corresponding to  $e \in E$  until we obtain G.

Now we show that maximum matchings in G' are related to those on G and vice versa.

**Lemma 4.** Any maximum cardinality matching M' on G' corresponds to a maximum cardinality matching M on G.

- <sup>705</sup> **Proof.** Let M' be a maximum cardinality matching on G'. A matching M for G can be
- <sup>706</sup> generated in the following way: If both  $(a, a_e)$  and  $(b_e, b)$  appear in M', e is added to M. <sup>707</sup> Hence it suffices to show that any maximum cardinality matching M' in G' necessarily
- contains |M| pair of matched edges  $(a, a_e)$  and  $(b, b_e)$ .
- First, we have that  $|M'| = |E_G| + |M|$ . To see this, note that per Lemma 2 we perform
- $|E_G|$  degree-2 reductions, and result in G. Each of this reductions corresponds with a matched redge in M'. Then, we only need to find the maximum cardinality on G which is |M|.

Let  $S_a$  contain all indices e such that  $(a, a_e)$  is in M' and  $(b_e, b)$  is not in M'. Set  $S_b$  is defined similarly. Set  $S_{\emptyset}$  contains all indices e such that  $(a_e, b_e)$  appears in M'. Finally,  $S_{ab}$ contains all indices e such that  $(a, a_e)$  and  $(b, b_e)$  are matched together in M'. Then, since M' is a maximum cardinality matching we have

$$|S_a| + |S_b| + |S_{\emptyset}| + 2 \cdot |S_{ab}| = |E_G| + |M| .$$

This is true because of the fact that for each edge e exactly one matched edge appears in M'in case  $e \in S_a \cup S_b \cup S_{\emptyset}$  and two edges are added if  $e \in S_{ab}$ .

However,  $|S_a| + |S_b| + |S_{\emptyset}| + |S_{ab}| = |E_G|$ , since each edge *e* must appear in one of those rss sets and there exist exactly  $|E_G|$  of them.

Hence,  $|S_{ab}| = |M|$  necessarily. As they define a matching in G and their cardinality is |M|, the matching is maximum.

718

<sup>719</sup> Using the above lemma, we can prove Theorem 5 below.

**Theorem 5.** Assume there is an algorithm ALG working in O(f(n,m)) time for finding a maximum cardinality matching in a 2-out graph. Then we can find a maximum cardinality matching in O(f(m,m)) time for any given graph.

**Proof.** Let G be any bipartite graph without degree-1 vertices and  $m = |E_G|$ . In O(m) time we generate G'. By Lemma 2, the 2-out subgraph of G' corresponds to G' itself. In addition  $|E_{G'}|, |V_{G'}| \in O(m)$ . Using ALG, we can find a maximum cardinality M' for G' in O(f(m, m))time. By Lemma 4 then, we can convert M' to a maximum cardinality matching M for G in O(m) time.

As a byproduct of Lemma 4, we observe that the transformation of G to G' also eliminates 728 the need to perform SK as a preprocessing step. We briefly experimented with this method 729 on the real-world graphs of Section 4.3. For each graph G of the test-set, we generated 730 its extension G' and executed the 20UTMC algorithm on 2-out graphs sampled from G', 731 with uniform selections. The behavior of 20UTMC was similar with that of the previous 732 experiments. It was not able to obtain a perfect matching in G' (and consequently G), but 733 it always returned near-optimal matchings of quality over 0.99. These matchings, when 734 converted into matchings of G (following the idea in Lemma 4) yielded also near-optimal 735 matchings with quality over 0.99. 736

-

737		References
738	1	Z. Allen-Zhu, Y. Li, R. Mendes de Oliveira, and A. Wigderson. Much faster algorithms for
739		matrix scaling. In 58th IEEE Annual Symposium on Foundations of Computer Science, FOCS,
740		pages 890–901, Berkeley, CA, USA, October 2017.
741	2	J. Aronson, M. Dyer, A. Frieze, and S. Suen. Randomized greedy matching II. Random
742		Structures & Algorithms, 6(1):55–73, 1995.
743	3	S. Assadi, M. Bateni, A. Bernstein, V. Mirrokni, and C. Stein. Coresets meet edcs: algorithms
744		for matching and vertex cover on massive graphs. In Proceedings of the Thirtieth Annual
745		ACM-SIAM Symposium on Discrete Algorithms, pages 1616–1635. SIAM, 2019.
746	4	S. Assadi and A. Bernstein. Towards a unified theory of sparsification for matching problems.
747		arXiv preprint arXiv:1811.02009, 2018.
748	5	S. Behnezhad, S. Brandt, M. Derakhshan, M. Fischer, M. Hajiaghayi, R.M. Karp, and J. Uitto.
749		Massively parallel computation of matching and mis in sparse graphs. In <i>Proceedings of the</i>
750		2019 ACM Symposium on Principles of Distributed Computing, pages 481–490, 2019.
751	6	S. Behnezhad, J. Łącki, and V. Mirrokni. Fully dynamic matching: Beating 2-approximation in
752		$\delta^{\epsilon}$ update time. In Proceedings of the Fourteenth Annual ACM-SIAM Symposium on Discrete
753		Algorithms, pages 2492–2508. SIAM, 2020.
754	7	C. Berge. Two theorems in graph theory. Proceedings of the National Academy of Sciences of
755		the $USA$ , $43:842-844$ , $1957$ .
756	8	A. Bernstein and C. Stein. Fully dynamic matching in bipartite graphs. In International
757		Colloquium on Automata, Languages, and Programming, pages 167–179. Springer, 2015.
758	9	B. Besser and M. Poloczek. Greedy matching: Guarantees and limitations. <i>Algorithmica</i> , 77(1):201–234, 2017
759	10	M.B. Cohon, A. Madry, D. Tsipres, and A. Vlady. Matrix scaling and balancing via how
760	10	constrained newton's method and interior point methods. In 58th IEEE Annual Symposium
762		on Foundations of Computer Science, FOCS, pages 902–913, Berkeley, CA, USA, October
763		2017.
764	11	A. Czumaj, J. Łacki, A. Madry, S. Mitrovic, K. Onak, and P. Sankowski. Round compression
765		for parallel matching algorithms. In Proceedings of the 50th Annual ACM SIGACT Symposium
766		on Theory of Computing, pages 471–484. Association for Computing Machinery, 2018.
767	12	T. A. Davis and Y. Hu. The University of Florida sparse matrix collection. ACM Transactions
768		on Mathematical Software, 38(1):1:1–1:25, 2011.
769	13	I. S. Duff. On algorithms for obtaining a maximum transversal. ACM Transactions on
770		Mathematical Software, 7(3):315–330, 1981.
771	14	I. S. Duff, K. Kaya, and B. Uçar. Design, implementation, and analysis of maximum transversal
772		algorithms. ACM Transactions on Mathematical Software, 38:13:1–13:31, 2011.
773	15	F. Dufossé, K. Kaya, I. Panagiotas, and B. Uçar. Approximation algorithms for maximum
774		matchings in undirected graphs. In 2018 Proceedings of the Seventh SIAM Workshop on
775		Combinatorial Scientific Computing, pages 56–65, 2018.
776	16	F. Dufossé, K. Kaya, and B. Uçar. Two approximation algorithms for bipartite matching on
777		multicore architectures. Journal of Parallel and Distributed Computing, 85:62–78, 2015.
778	17	A. Frieze and T. Johansson. On random k-out subgraphs of large graphs. Random Structures
779	10	@ Algorithms, 50(2):143–157, 2017.
780	10	A. Goel, M. Kapralov, and S. Khanna. Perfect matchings in $O(n \log n)$ time in regular bipartite
781	10	graphs. SIAM Journal on Computing, 42(5):1592–1404, 2015.
782	13	A. v. Goldberg and R. E. rarjan. A new approach to the maximum-now problem. J. $ACM$ , $35(A) \cdot 021 = 0.40$ 1088
783	20	T Hagorup K Mohlhorn and I I Munro Maintaining discrete probability distributions
795	20	ontimally In A Lingas B Karlsson and S Carlsson editors 20th International Colloquium
786		on Automata, Languages, and Programming (ICALP), pages 253–264. Berlin, Heidelberg 1993
787		Springer Berlin Heidelberg.

- <sup>788</sup> **21** J. E. Hopcroft and R. M. Karp. An  $n^{5/2}$  algorithm for maximum matchings in bipartite <sup>789</sup> graphs. *SIAM Journal on Computing*, 2(4):225–231, 1973.
- R. M. Karp, A. H. G. Rinnooy Kan, and R. V. Vohra. Average case analysis of a heuristic for the assignment problem. *Mathematics of Operations Research*, 19(3):513–522, 1994.
- R. M. Karp and M. Sipser. Maximum matching in sparse random graphs. In 22nd Annual IEEE Symposium on Foundations of Computer Science (FOCS), pages 364–375, Los Alamitos, CA, USA, 1981. IEEE Computer Society.
- R. M. Karp, U. V. Vazirani, and V. V. Vazirani. An optimal algorithm for on-line bipartite matching. In *Proceedings of the twenty-second annual ACM symposium on Theory of computing*, STOC '90, pages 352–358, New York, NY, USA, 1990. ACM.
- K. Kaya, J. Langguth, F. Manne, and B. Uçar. Push-relabel based algorithms for the maximum transversal problem. *Computers & Operations Research*, 40(5):1266–1275, 2013.
- K. Kaya, J. Langguth, I. Panagiotas, and B. Uçar. Karp–Sipser based kernels for bipartite
   graph matching. In SIAM Symposium on Algorithm Engineering and Experiments (ALENEX),
   pages 134–145, Salt Lake City, Utah, US, January 2020.
- P. A. Knight. The Sinkhorn-Knopp algorithm: Convergence and applications. SIAM Journal
   on Matrix Analysis and Applications, 30(1):261–275, 2008.
- P. A. Knight and D. Ruiz. A fast algorithm for matrix balancing. *IMA Journal of Numerical* Analysis, 33(3):1029–1047, 2013.
- V. Korenwein, A. Nichterlein, R. Niedermeier, and P. Zschoche. Data reduction for maximum matching on real-world graphs: Theory and experiments. In 26th Annual European Symposium on Algorithms (ESA 2018), volume 112, pages 53:1–53:13, Dagstuhl, Germany, 2018.
- 30 J. Langguth, F. Manne, and P. Sanders. Heuristic initialization for bipartite matching problems.
   Journal of Experimental Algorithmics (JEA), 15:1–22, 2010.
- J. Magun. Greedy matching algorithms, an experimental study. Journal of Experimental Algorithmics, 3:6, 1998.
- Y. Matias, J. S. Vitter, and W.-C. Ni. Dynamic generation of discrete random variates. *Theory* of Computing Systems, 36(4):329–358, 2003.
- 33 N. McKeown. The iSLIP scheduling algorithm for input-queued switches. *IEEE/ACM* Transactions on Networking, 7:188–201, 1999.
- 34 M. Mitzenmacher and E. Upfal. Probability and computing: Randomized algorithms and probabilistic analysis. Cambridge University Press, 1st edition, 2005.
- M. Poloczek and M. Szegedy. Randomized greedy algorithms for the maximum matching
   problem with new analysis. In *Foundations of Computer Science (FOCS)*, 2012 IEEE 53rd
   Annual Symposium on, pages 708–717, 2012.
- A. Pothen and C.-J. Fan. Computing the block triangular form of a sparse matrix. ACM
   Transactions on Mathematical Software, 16(4):303-324, 1990.
- A. Pothen, S. M. Ferdous, and F. Manne. Approximation algorithms in combinatorial scientific computing. *Acta Numerica*, 28:541–633, 2019.
- R. Sinkhorn and P. Knopp. Concerning nonnegative matrices and doubly stochastic matrices.
   *Pacific Journal of Mathematics*, 21(2):343–348, 1967.
- 39 G. Tinhofer. A probabilistic analysis of some greedy cardinality matching algorithms. Annals
   of Operations Research, 1(3):239–254, 1984.
- 40 D. Walkup. Matchings in random regular bipartite digraphs. *Discrete Mathematics*, 31(1):59–64, 1980.