

Implementation of $O(nm \log n)$ Weighted Matchings in General Graphs: The Power of Data Structures

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We describe the implementation of an algorithm which solves the weighted matching problem in general graphs with n vertices and m edges in time $O(nm \log n)$. Our algorithm is a variant of the algorithm of Galil, Micali and Gabow [Galil et al. 1986] and extensively uses sophisticated data structures, in particular *concatenable priority queues*, so as to reduce the time needed to perform dual adjustments and to find tight edges in Edmonds' blossom-shrinking algorithm.

We compare our implementation to the experimentally fastest implementation, named *Blossom IV*, due to Cook and Rohe [Cook and Rohe 1997]. Blossom IV requires only very simple data structures and has an asymptotic running time of $O(n^2 m)$. Our experiments show that our new implementation is superior to Blossom IV. A closer inspection reveals that the running time of Edmonds' blossom-shrinking algorithm in practice heavily depends on the time spent to perform dual adjustments and to find tight edges. Therefore, optimizing these operations, as is done in our implementation, indeed speeds-up the practical performance of implementations of Edmonds' algorithm.

1. INTRODUCTION

We consider the *weighted matching problem* in general graphs. That is, given an undirected graph $G = (V, E)$ we wish to find a matching of maximum weight. A *matching* M in G is a set of edges no two of which share an endpoint. The edges of G have weights associated with them and the *weight* of a matching is simply the

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sum of the weights of the edges in the matching. M may further be restricted to being *perfect*, which constitutes the *weighted perfect matching problem*; a matching is *perfect* if every vertex of the graph is matched.

In 1965, Edmonds [Edmonds 1965b] invented the famous *blossom-shrinking algorithm*, which solves the weighted perfect matching problem in polynomial time. A straightforward implementation of the blossom-shrinking algorithm, as originally proposed by Edmonds himself, requires time $O(n^2m)$, where n and m are the number of vertices and edges of G , respectively. Since then, the worst-case complexity of the blossom-shrinking algorithm has been improved successively: both Lawler [Lawler 1976] and Gabow [Gabow 1974] achieved a running time of $O(n^3)$, Galil, Micali and Gabow [Galil et al. 1986] improved the running time to $O(nm \log n)$ and finally Gabow [Gabow 1990] achieved a running time of $O(n(m + n \log n))$. Somewhat better asymptotic running times are known for integral edge weights.

Mainly, the improved worst-case complexity is achieved by using sophisticated data structures which reduce the time needed to perform dual adjustments and to find tight edges. For example, the algorithm of Galil, Micali and Gabow requires *concatenable priority queues*, in which the priorities of certain subgroups of vertices can be changed by a single operation. As a consequence, a dual adjustment can be performed in time $O(1)$ as opposed to $O(n)$ as in the algorithms of Edmonds [Edmonds 1965b], Lawler [Lawler 1976] and Gabow [Gabow 1974].

The currently most efficient codes implement variants of Edmonds' blossom-shrinking algorithm and are based on either the algorithm of Edmonds [Edmonds 1965b], Lawler [Lawler 1976] or Gabow [Gabow 1974]. All these algorithms only use very simple data structures. The fastest implementation currently available, named *Blossom IV*, is due to Cook and Rohe [Cook and Rohe 1997]. Their work is based on earlier work by Applegate and Cook [Applegate and Cook 1993]. Blossom IV is known to be highly efficient in practice.

It has been an open question, and one explicitly posed in [Applegate and Cook 1993] and [Cook and Rohe 1997], whether or not the use of sophisticated data structures will help to improve the performance of weighted matching algorithms in practice. We will answer this question in the affirmative: our implementation is a variant of the algorithm of Galil, Micali and Gabow [Galil et al. 1986] and, as the experiments show, superior to Blossom IV on most of the tested instances. Our implementation can be asked to compute either a maximum-weight perfect matching or a maximum-weight matching. Previous implementations were restricted to compute optimal perfect matchings.¹

No previous implementation guaranteed a worst-case performance of $O(nm \log n)$. As mentioned above, the improved asymptotics results from improved time bounds for dual adjustments and finding tight edges. Our experiments show that the time spend for dual adjustments and finding tight edges is not only the theoretical bottleneck for weighted matchings; Blossom IV actually spends most of its time in

¹One may argue that the maximum-weight matching problem can be reduced to the maximum-weight perfect matching problem: the original graph is doubled and zero weight edges are inserted from each original vertex to the corresponding doubled vertex. However, for an original graph with n vertices and m edges the reduction doubles n and increases m by $m + n$. Thus, for large instances the reduction becomes infeasible.

these operations.

The preflow-push method for maximum network flow is another example of an algorithm whose asymptotic running time improves dramatically through the use of sophisticated data structures. With the highest level selection rule and only simple data structures the worst-case running time is $O(n^2 \sqrt{m})$. It improves to $\tilde{O}(nm)$ with the use of dynamic trees. None of the known efficient implementations [Cherkassky and Goldberg ; Mehlhorn and Näher 1999] uses sophisticated data structures and attempts to use them produced far inferior implementations [Humble 1996]. The reason here is that the scenario in which dynamic trees help rarely occurs in practical problems.

Our implementation is based on (and part of) the *Library of Efficient Data Types and Algorithms* (LEDA). It may seem astonishing that an algorithm based on an all-purpose library such as LEDA is superior to an algorithm which was developed from scratch with the only purpose to efficiently solve the weighted matching problem. We see two reasons: (1) the use of LEDA allowed us to implement the asymptotically faster algorithm and, as argued above, the theoretical bottleneck is also the actual bottleneck in the case of weighted matching and (2) LEDA aims for zero-overhead library design and comes close to this goal in the area of graph algorithms.

This paper is organized as follows: in Section 2 we review Edmonds' blossom-shrinking algorithm, in Section 3 we describe our implementation, in Section 4 we report our experimental findings and in Section 5 we offer a short conclusion.

2. EDMONDS' BLOSSOM-SHRINKING ALGORITHM

Edmonds' blossom-shrinking algorithm is a primal-dual method based on a linear programming formulation of the maximum-weight perfect matching problem. The details of the algorithm depend on the underlying formulation. We will first give the formulation we use and then present all details of the resulting blossom-shrinking algorithm.

2.1 LP Formulations

Let $G = (V, E)$ be a general graph with edge weights $w : E \rightarrow \mathbb{R}$. An incidence vector x is associated with the edges of G : x_e is set to 1 if e belongs to the matching; otherwise, x_e is set to 0. For any subset $S \subseteq E$, we define $x(S) = \sum_{e \in S} x_e$. The edges of G having both endpoints in $S \subseteq V$ are denoted by $\gamma(S) = \{uv \in E : u \in S \text{ and } v \in S\}$, and the set of all edges having exactly one endpoint in S is referred to by $\delta(S) = \{uv \in E : u \in S \text{ and } v \notin S\}$. (For a singleton set $S = \{u\}$ we use $\delta(u)$ for short.) Moreover, let \mathcal{O} consist of all non-singleton odd cardinality subsets of V : $\mathcal{O} = \{B \subseteq V : |B| \text{ is odd and } |B| \geq 3\}$.

The maximum-weight perfect matching problem of G with weight function w can then be formulated as a linear program:

$$\begin{aligned}
 \text{(WPM)} \quad & \text{maximize} && w^T x \\
 & \text{subject to} && x(\delta(u)) = 1 && \text{for all } u \in V, && (1) \\
 & && x(\gamma(B)) \leq \lfloor |B|/2 \rfloor && \text{for all } B \in \mathcal{O}, && (2) \\
 & && x_e \geq 0 && \text{for all } e \in E. && (3)
 \end{aligned}$$

(WPM)(1) states that each vertex u of G must be matched and (WPM)(2)–(3) assure that x_e is set either to 0 or 1.

Consider the dual linear program of (WPM). A *potential* y_u and $z_{\mathcal{B}}$ is assigned to each vertex u and non-singleton odd cardinality set \mathcal{B} , respectively.

$$\begin{aligned}
 (\overline{\text{WPM}}) \quad & \text{minimize} && \sum_{u \in V} y_u + \sum_{\mathcal{B} \in \mathcal{O}} \lfloor |\mathcal{B}|/2 \rfloor z_{\mathcal{B}} \\
 & \text{subject to} && z_{\mathcal{B}} \geq 0 \quad \text{for all } \mathcal{B} \in \mathcal{O}, \quad (1) \\
 & && y_u + y_v + \sum_{\substack{\mathcal{B} \in \mathcal{O} \\ uv \in \gamma(\mathcal{B})}} z_{\mathcal{B}} \geq w_{uv} \quad \text{for all } uv \in E. \quad (2)
 \end{aligned}$$

The *reduced cost* π_{uv} of an edge uv with respect to a dual solution (y, z) of $(\overline{\text{WPM}})$ is defined as given below. We will say that an edge uv is *tight* if its reduced cost π_{uv} equals 0.

$$\pi_{uv} = y_u + y_v - w_{uv} + \sum_{\substack{\mathcal{B} \in \mathcal{O} \\ uv \in \gamma(\mathcal{B})}} z_{\mathcal{B}}. \quad (1)$$

Edmonds' blossom-shrinking algorithm is a primal-dual method that keeps a primal (not necessarily feasible) solution x of (WPM) and also a dual feasible solution (y, z) of $(\overline{\text{WPM}})$; the primal solution may violate (WPM)(1). The solutions are adjusted successively until they are recognized to be optimal. The optimality of x and (y, z) will be assured by the feasibility of x and (y, z) and the validity of the complementary slackness conditions (CS)(1)–(2):

$$\begin{aligned}
 (\text{CS}) \quad & x_{uv} > 0 \implies \pi_{uv} = 0 \quad \text{for all } uv \in E, \quad (1) \\
 & z_{\mathcal{B}} > 0 \implies x(\gamma(\mathcal{B})) = \lfloor |\mathcal{B}|/2 \rfloor \quad \text{for all } \mathcal{B} \in \mathcal{O}. \quad (2)
 \end{aligned}$$

(CS)(1) states that matching edges must be tight and (CS)(2) states that non-singleton sets \mathcal{B} with a positive potential are *full*, i.e., a maximum number of edges in \mathcal{B} are matched.

There exists an alternative formulation, in which the second constraint (WPM)(2) is replaced by “ $x(\delta(\mathcal{B})) = 1$ for all $\mathcal{B} \in \mathcal{O}$ ”. The details of an algorithm based on the alternative formulation differ from those which will be presented below.² We will outline the main differences below. Both the implementation of Applegate and Cook [Applegate and Cook 1993] and the implementation of Cook and Rohe [Cook and Rohe 1997] use the alternative formulation. The formulation above is used by Galil, Micali and Gabow [Galil et al. 1986] and seems to be more suitable to achieve a running time of $O(nm \log n)$. It has the additional advantage that changing the constraint (WPM)(1) to “ $x(\delta(u)) \leq 1$ for all $u \in V$ ” gives a formulation of the non-perfect maximum-weight matching problem.

2.2 Blossom-Shrinking Algorithm

2.2.1 Initialization. The algorithm starts with an arbitrary matching M , which satisfies (WPM)(2)–(3).³ Each vertex u has a potential y_u associated with it and

²On the one hand, performing a dual adjustment becomes slightly simpler, but, on the other hand, the computation of the reduced cost of an edge is computationally more expensive.

³We will often use the concept of a matching M and its incidence vector x interchangeably.

all z_B 's are (implicitly) set to 0. The potentials are chosen such that (y, z) is a dual feasible solution of $(\overline{\text{WPM}})$ and, moreover, satisfies $(\text{CS})(1)–(2)$ with respect to M . The algorithm operates in phases.

2.2.2 A Phase. In each phase, two additional vertices become matched and hence do no longer violate $(\text{WPM})(1)$. After $O(n)$ phases, either all vertices will satisfy $(\text{WPM})(1)$ and thus the computed matching is optimal, or it has been discovered that no perfect matching exists.

The algorithm attempts to match free vertices by growing *alternating trees* rooted at free vertices.⁴ Each alternating tree T_r is rooted at a free vertex r . The edges in T_r are tight and alternately matched and unmatched with respect to the current matching M . We further assume a labeling for the vertices of T_r : a vertex $u \in T_r$ is labeled *even* or *odd*, when the path from u to r is of even or odd length, respectively. Vertices that are not part of any alternating tree are matched and said to be *unlabeled*. We will use u^+ , u^- or u^\emptyset to denote an even, odd or unlabeled vertex.

An alternating tree is extended, or *grown*, from even labeled tree vertices $u^+ \in T_r$: when a tight edge uv from u to a non-tree vertex v^\emptyset exists, the edge uv and also the matching edge vw of v (which must exist, since v is unlabeled) is added to T_r . Here, v and w are labeled odd and even, respectively.

When a tight edge uv with $u^+ \in T_r$ and $v^+ \in T_{r'}$ exists, with $T_r \neq T_{r'}$, an *augmenting path* from r to r' has been discovered. A path p from a free vertex r to another free vertex r' is called *augmenting*, if the edges along p are alternately in M and not in M (the first and last edge are unmatched). Let p_r denote the tree path in T_r from u to r and, correspondingly, $p_{r'}$ the tree path in $T_{r'}$ from v to r' . By $\overline{p_r}$ we denote the path p_r in reversed order. The current matching M is augmented by $P = (\overline{p_r}, uv, p_{r'})$, i.e., all non-matching edges along P become matching edges and vice versa. After that, all vertices in T_r and $T_{r'}$ will be matched; therefore, we can destroy T_r and $T_{r'}$ and unlabeled all their vertices.

Assume a tight edge uv connecting two even tree vertices $u^+ \in T_r$ and $v^+ \in T_r$ exists (in the same tree T_r). We follow the tree paths from u and v towards the root until the lowest common ancestor vertex lca has been found. lca must be even by construction of T_r and the simple cycle $C = (lca, \dots, u, v, \dots, lca)$ is full. The subset $B \subseteq V$ of vertices on C are said to form a *blossom*, as introduced by Edmonds [Edmonds 1965b], and uv will be called a *blossom forming edge*. A key observation is that one can *shrink* blossoms into an even labeled pseudo-vertex, say lca , and continue the growth of the alternating trees in the resulting graph.⁵ To shrink a cycle C means to collapse all vertices of B into a single pseudo-vertex lca . All edges uv between vertices of B , i.e., $uv \in \gamma(B)$, become non-existent and all edges uv having exactly one endpoint v in B , i.e., $uv \in \delta(B)$, are replaced by an edge from u to lca .

However, we will regard these vertices to be conceptually shrunk into a new

⁴We concentrate on a *multiple search tree* approach, where alternating trees are grown from all free vertices simultaneously. The *single search tree* approach, where just one alternating tree is grown at a time, is slightly simpler to describe, gives the same asymptotic running time, but leads to an inferior implementation.

⁵The crucial point is that any augmenting path in the resulting graph can be lifted to an augmenting path in the original graph.

pseudo-vertex only. Since pseudo-vertices might get shrunk into other pseudo-vertices, the following view is appropriate: the current graph is partitioned into a *nested family* of odd cardinality subsets of V . Each odd cardinality subset is called a blossom. A blossom might contain other blossoms, called *subblossoms*. A *trivial* blossom corresponds to a single vertex of G . A blossom \mathcal{B} is said to be a *surface blossom*, if \mathcal{B} is not contained in any other blossom. All edges lying completely in some blossom \mathcal{B} are *dead* and will not be considered by the algorithm; all other edges are *alive*.⁶

2.2.3 Dual Adjustment. The algorithm might come to a halt due to the lack of further tight edges. Then, a *dual adjustment* is performed: the dual solution (y, z) is adjusted such that the objective value of $(\overline{\text{WPM}})$ decreases. However, the adjustment will be of the kind such that (y, z) stays dual feasible and moreover preserves (CS)(1)–(2). The potentials y_u of *all* vertices $u \in V$ and the potential $z_{\mathcal{B}}$ of each non-trivial *surface* blossom \mathcal{B} are updated by some $\delta > 0$:

$$y'_u = y_u + \sigma\delta, \quad \text{and} \quad z'_{\mathcal{B}} = z_{\mathcal{B}} - 2\sigma\delta.^7$$

The *status indicator* σ equals -1 or 1 for even or odd tree blossoms (trivial or non-trivial), respectively, and equals 0 , otherwise. (The status of a vertex is the status of the surface blossom containing it.) Let T_{r_1}, \dots, T_{r_k} denote the current alternating trees. The value of δ must be chosen as $\delta = \min\{\delta_2, \delta_3, \delta_4\}$, with

$$\delta_2 = \min_{uv \in E} \{\pi_{uv} : u^+ \in T_{r_i} \text{ and } v^\varnothing \text{ not in any tree}\},$$

$$\delta_3 = \min_{uv \in E} \{\pi_{uv}/2 : u^+ \in T_{r_i} \text{ and } v^+ \in T_{r_j}\},$$

$$\delta_4 = \min_{\mathcal{B} \in \mathcal{O}} \{z_{\mathcal{B}}/2 : \mathcal{B}^- \in T_{r_i}\},$$

where T_{r_i} and T_{r_j} denote any alternating tree, with $1 \leq i, j \leq k$.⁸ The minimum of an empty set is defined to be ∞ . When $\delta = \infty$, the dual linear program $(\overline{\text{WPM}})$ is unbounded and thus no optimal solution to (WPM) exists (by weak duality).

If δ is chosen as δ_4 , the potential of an odd tree blossom, say $\mathcal{B}^- \in T_r$, will drop to 0 after the dual adjustment. Due to $(\overline{\text{WPM}})(1)$, \mathcal{B} must be prevented from participating in further dual adjustments. The action to be taken is to *expand* \mathcal{B} , i.e., the defining subblossoms $\mathcal{B}_1, \dots, \mathcal{B}_{2k+1}$ of \mathcal{B} are lifted to the surface and \mathcal{B} is abandoned. Since \mathcal{B} is odd, there exists a matching tree edge ub and a non-matching tree edge dv . The vertices b and d in \mathcal{B} are called the *base* and *discovery*

⁶Note that during the course of the algorithm the reduced costs of alive edges have to be computed frequently. If the alternative LP formulation is used this is a quite tedious task:

$$\pi_{uv} = y_u + y_v - w_{uv} + \sum_{\substack{\mathcal{B} \in \mathcal{O} \\ uv \in \delta(\mathcal{B})}} z_{\mathcal{B}}.$$

That is, not only the vertex potentials of the endpoints but also the potentials of all blossoms containing exactly one endpoint must be taken into account.

⁷For the alternative LP formulation, the potential updates reduce to

$$y'_u = y_u + \sigma\delta, \quad \text{and} \quad z'_{\mathcal{B}} = z_{\mathcal{B}} + \sigma\delta$$

for each *surface* vertex u and *surface* blossom \mathcal{B} .

⁸Notice that T_{r_i} and T_{r_j} need not necessarily to be different in the definition of δ_3 .

vertex of \mathcal{B} . Assume \mathcal{B}_i and \mathcal{B}_j correspond to the subblossoms containing b and d , respectively. Let p denote the even length alternating (now alive) path from \mathcal{B}_j to \mathcal{B}_i with its edges lying exclusively in $\gamma(\mathcal{B})$. The path p and all subblossoms along p are added to T_r and are labeled accordingly. All remaining subblossoms are unlabeled and leave T_r .

2.2.4 Running Time of Different Realizations. As mentioned above, the algorithm terminates after $O(n)$ phases. The number of dual adjustments is bounded by $O(n)$ per phase.⁹ A *union-find* data structure that additionally supports a *split* operation is sufficient to maintain the surface graph in time $O(n \log n)$ per phase.¹⁰ The existing realizations of the blossom-shrinking algorithm mainly differ in the way they determine the value of δ , perform dual adjustments and find tight edges.

The most trivial realization inspects all edges and explicitly updates the vertex and blossom potentials and thus needs $O(m + n)$ time per dual adjustment. The resulting $O(n^2 m)$, or equivalently $O(n^4)$, approach is essentially the one which was suggested first by Edmonds [Edmonds 1965a].

Lawler [Lawler 1976] and Gabow [Gabow 1974] improved the asymptotic running time to $O(n^3)$ by reducing the time spent to determine the value of δ and to find tight edges to $O(n)$. The idea is to keep for each non-tree vertex v° a *best edge*, i.e., the edge having minimum reduced cost, to an even labeled tree vertex. The same data is kept for each odd tree vertex $v^- \in T_r$. (The necessity for the latter is due to the expansion of blossoms). Additionally, each even tree blossom knows its best edges to other even tree blossoms. Then, only all best edges have to be considered in order to determine the value of δ and to find tight edges. Moreover, the best edge data can be maintained in time $O(n^2)$ per phase.

The algorithm of Galil, Micali and Gabow [Galil et al. 1986] considerably improves the time needed to perform a dual adjustment to $O(1)$. The determining of δ and the finding of tight edges reduces to a priority queue operation and takes time $O(\log n)$. The maintenance of the underlying priority queues requires time $O(m \log n)$ per phase. The total running time is therefore $O(nm \log n)$. This algorithm will be discussed in more detail below.

Gabow [Gabow 1990] further improved the running time to $O(n(m + n \log n))$. The main difficulty here is to deal with the (potentially m many) blossom forming edges. The rough idea is to replace each blossom forming edge uv by two back edges to the lowest common ancestor of u and v and to partition these back edges into packets. However, the underlying data structures are complex and we doubt that an efficient implementation would be possible.

⁹Whenever $\delta = \delta_2, \delta_3$, at least one (formerly non-even labeled) vertex becomes an even tree vertex, or a phase terminates. An even tree vertex stays even and resides in its tree until the end of the phase. Thus, $\delta = \delta_2, \delta_3$ occurs $O(n)$ times per phase. The maximum cardinality of a blossom is n and therefore $\delta = \delta_4$ occurs $O(n)$ times per phase.

¹⁰Each vertex knows the *name* of its surface blossom. In a *shrink* or an *expand* step all vertices of the smaller group are renamed.

3. IMPLEMENTATION

3.1 Blossom IV

The implementation, named Blossom IV, of Cook and Rohe [Cook and Rohe 1997] is the most efficient code for weighted perfect matchings in general graphs currently available. Cook and Rohe do not claim any worst-case guarantee. Blossom IV only uses very simple data structures. In particular, no priority queues are used to determine the value of δ or to find tight edges and the potential updates are made explicitly, i.e., take time $O(n)$ per dual adjustment. Thus, the worst-case guarantee cannot be better than $O(n^3)$. Since no *best edge* data is kept, we even believe that it is $O(n^2m)$. The algorithm is implemented in C.

The comparison to other implementations is made in two papers: (1) In [Cook and Rohe 1997] Blossom IV is compared to the implementation of Applegate and Cook [Applegate and Cook 1993]. It is shown that Blossom IV is substantially faster. (2) In [Applegate and Cook 1993] the implementation of Applegate and Cook is compared to other implementations. The authors show that their code is superior to all other codes.

In Blossom IV one can choose between three approaches: a single search tree approach, a multiple search tree approach and a refinement of the multiple search tree approach, called the *variable δ approach*. In the variable δ approach, each alternating tree T_{r_i} chooses its own dual adjustment value δ_{r_i} so as to maximize the decrease in the dual objective value. A heuristic is used to make the choices (an exact computation would be too costly). The variable δ approach does not only lead to a faster decrease of the dual objective value, it, typically, also creates tight edges faster. The experiments in [Cook and Rohe 1997] show that the variable δ approach is superior to the other approaches in practice. We make all our comparisons to Blossom IV with the variable δ approach.

3.1.1 Price-and-Repair Strategy for Geometric Instances. Blossom IV provides a price-and-repair heuristic which allows it to run on implicitly defined complete geometric graphs (edge weight = Euclidean distance). The running time of Blossom IV on these instances is significantly improved if the price-and-repair heuristic is used. We have not yet implemented such a heuristic for our algorithm.

The idea is simple. A minimum-weight matching (it is now more natural to talk about minimum-weight matchings) has a natural tendency of avoiding large weight edges; this suggests to compute a minimum-weight matching iteratively. One starts with a sparse subgraph of light edges and computes an optimal matching. Once the optimal matching is computed, one checks optimality with respect to the full graph. This is what is called *pricing*. Some of the edges having negative reduced cost are added to the current graph, with the matching and the potentials being modified such that all preconditions of the matching algorithm are satisfied. The algorithm is resumed so as to *repair* the matching for the current graph. This process is repeated until the obtained matching is optimal for the full graph. Derigs and Metz [Derigs and Metz 1991; Applegate and Cook 1993; Cook and Rohe 1997] discuss the repair step in detail.

There are several natural strategies for selecting the sparse subgraph. One can, e.g., take the lightest d edges incident to any vertex. For complete graphs induced by a set of points in the plane and with edge weights equal to the Euclidean distance,

the Delaunay diagram of the points is a good choice.

3.2 Our Implementation

We come to our implementation. It has a worst-case running time of $O(nm \log n)$, extensively uses (concatenable) priority queues and is able to compute a non-perfect or a perfect maximum-weight matching. It is based on the *Library of Efficient Data Types and Algorithms* (LEDA, [Mehlhorn and Näher 1999]) and the implementation language is C++. The implementation can run either a single search tree approach or a multiple search tree approach. As the experiments will show, the additional programming expenditure for the multiple search tree approach is well worth the effort regarding the efficiency in practice. Comparisons of our multiple search tree algorithm to the variable δ approach of Blossom IV will be given in Section 4.

The underlying strategies are similar to or have been evolved from the ideas of Galil, Micali and Gabow [Galil et al. 1986]. The time required to perform a dual adjustment is considerably improved to $O(1)$. The determining of δ and the finding of tight edges reduces to a priority queue operation and takes time $O(\log n)$. Moreover, the maintenance of the underlying priority queue data structures takes time $O(m \log n)$ per phase. However, our approach differs with regard to the maintenance of the varying potentials and reduced costs. Galil, Micali and Gabow handle these varying values within the priority queues, i.e., by means of an operation that changes all priorities in a priority queue by the same amount, whereas we establish a series of formulae that enable us to compute the values on demand.

Next, we will briefly sketch the key ideas of our implementation. As above, we concentrate on the description of the multiple search tree approach.

3.2.1 Determination of δ — towards a Priority Queue Approach. For each δ_2, δ_3 and δ_4 we keep a priority queue *delta2*, *delta3* and *delta4*, respectively. The priorities stored in each priority queue correspond to the value of interest, i.e., to (one half of) the reduced cost of edges for *delta2* and *delta3* and to blossom potentials for *delta4*. The difficulty that these priorities decrease by δ with each dual adjustment is overcome as follows. We keep track of the amount $\Delta = \sum \delta_i$ of all dual adjustments and compute the *actual* priority \tilde{p} of any element in the priority queues by taking Δ into consideration: $\tilde{p} = p - \Delta$. A dual adjustment by δ then reduces to an increase of Δ by δ , i.e., the time required by a dual adjustment is reduced to $O(1)$.

We associate a *concatenable priority queue* P_B with each surface blossom B . A concatenable priority queue supports all standard priority queue operations and, in addition, a *concat* and *split* operation. Moreover, the elements are regarded to form a sequence. *concat* concatenates the sequences of two priority queues and, conversely, *split* splits the sequence of a priority queue at a given item into two priority queues. It can be achieved that both operations run in time $O(\log n)$, see, e.g., [Mehlhorn 1984, Section III.5.3] and [Aho et al. 1974, Section 4.12]. Our implementation is based on (2, 16)-trees.

P_B contains exactly one element $\langle p, u \rangle$ for each vertex u contained in B . Generally, p represents the reduced cost of the best edge of u to an even labeled tree vertex. More precisely, let T_{r_1}, \dots, T_{r_k} denote the alternating trees at some stage of the blossom-shrinking algorithm. Every vertex u is associated with a series of (at most k) incident edges uv_1, \dots, uv_k and their reduced costs $\pi_{uv_1}, \dots, \pi_{uv_k}$ (maintained

by a standard priority queue). Each edge uv_i represents the best edge from u to an even tree vertex $v_i^+ \in T_{r_i}$. The reduced cost $\pi_{uv_i^*}$ of u 's best edge uv_i^* (along all best edges uv_i associated with u) is the priority stored with the element $\langle p, u \rangle$ in P_B .

When a new blossom B is formed by $B_1, B_2, \dots, B_{2k+1}$ the priority queues $P_{B_1}, P_{B_2}, \dots, P_{B_{2k+1}}$ are concatenated one after another and the resulting priority queue P_B is assigned to B . Thereby, we keep track of each t_i , $1 \leq i \leq 2k+1$, the last item in B_i . Later, when B is expanded the priority queues of $B_1, B_2, \dots, B_{2k+1}$ can easily be recovered by splitting the priority queue of B at each item t_i , $1 \leq i \leq 2k+1$.

Some details for the maintenance of *delta2* are given next. Whenever a blossom B becomes unlabeled, it sends its best edge and the reduced cost of this edge to *delta2*. Moreover, when the best edge of B changes, the appropriate element in *delta2* is adjusted. If a non-tree blossom B^\emptyset becomes a tree blossom its element in *delta2* is deleted. These insertions, adjustments and deletions on *delta2* contribute $O(n \log n)$ time per phase.

We come to *delta3*. Each tree T_{r_i} maintains its own priority queue *delta3_{r_i}* containing all *blossom forming* edges, i.e., the edges connecting two even vertices in T_{r_i} . The priority of each element corresponds to one half of the reduced cost; note, however, that the actual reduced cost of each element is computed as stated above. The edges inserted into *delta3_{r_i}* are assured to be alive. However, during the course of the algorithm some edges in *delta3_{r_i}* might become dead. We use a *lazy-deletion* strategy for these edges: dead edges are simply discarded when they occur as the minimum element of *delta3_{r_i}*. The minimum element of each *delta3_{r_i}* is sent to *delta3*. When a tree T_{r_i} is destroyed, its representative is deleted from *delta3* and *delta3_{r_i}* is freed. Moreover, each even labeled blossom $B^+ \in T_{r_i}$ sends its best edge uv with $u^+ \in T_{r_i}$ and $v^+ \in T_{r_j}$, $T_{r_i} \neq T_{r_j}$, and the reduced cost of this edge to *delta3*. Since $m \leq n^2$, the time needed for the maintenance of all priority queues responsible for *delta3* is $O(m \log n)$ per phase.

Handling *delta4* is trivial. Each non-trivial odd surface blossom $B^- \in T_{r_i}$ sends an element to *delta4*. The priority corresponds to one half of the potential z_B .

3.2.2 Varying Potentials and Reduced Costs. What remains to be shown is how to treat the varying blossom and vertex potentials as well as the reduced cost of all edges associated with the vertices. The crux is that with each dual adjustment all these values uniformly change by some multiple of δ .

For example, consider an even surface blossom $B^+ \in T_{r_i}$. If a dual adjustment by δ is performed, the potential of B changes by $+2\delta$, the potential of each vertex $u \in B$ changes by $-\delta$ and the reduced costs of all edges associated with each $u \in B$ change by -2δ . Taking advantage of this fact, the actual value of interest can be computed by taking Δ and some additional information into consideration. The idea is as follows.

Each surface blossom B has an offset *offset_B* assigned to it. This offset is initially set to 0 and will be adjusted whenever B changes its status. The formulae to compute the actual potential \tilde{z}_B for a (non-trivial) surface blossom B , the actual potential \tilde{y}_u for a vertex u (with surface blossom B) and the actual reduced cost

$\tilde{\pi}_{uv_i}$ of an edge uv_i associated with u (and surface blossom \mathcal{B}) are given below:

$$\tilde{z}_{\mathcal{B}} = z_{\mathcal{B}} - 2\text{offset}_{\mathcal{B}} - 2\sigma\Delta, \quad (2)$$

$$\tilde{y}_u = y_u + \text{offset}_{\mathcal{B}} + \sigma\Delta, \quad (3)$$

$$\tilde{\pi}_{uv_i} = \pi_{uv_i} + \text{offset}_{\mathcal{B}} + (\sigma - 1)\Delta. \quad (4)$$

Here, σ and Δ are defined as above. A status change for a surface blossom \mathcal{B} then reduces to an update of its offset value:

$$\text{offset}_{\mathcal{B}} = \text{offset}_{\mathcal{B}} + (\sigma - \sigma')\Delta, \quad (5)$$

which is necessary at the point of time, when \mathcal{B} changes its status indicator from σ to σ' . The update of a blossom offset takes time $O(1)$. We conclude, that each value of interest can be computed (if required) in time $O(1)$ by the formulae (2)–(4).

It is an easy matter to handle the blossom offsets if a blossom \mathcal{B} is expanded: $\text{offset}_{\mathcal{B}}$ is assigned to each offset of the defining subblossoms of \mathcal{B} .

However, it is not so obvious how to handle the possibly different blossom offsets in a shrink step. Let \mathcal{B} denote the blossom that is about to be formed by the defining subblossoms $\mathcal{B}_1, \mathcal{B}_2, \dots, \mathcal{B}_{2k+1}$. Each odd labeled subblossom \mathcal{B}_i is made even by adjusting its offset as in (5). The corresponding offsets $\text{offset}_{\mathcal{B}_1}, \text{offset}_{\mathcal{B}_2}, \dots, \text{offset}_{\mathcal{B}_{2k+1}}$ may have different values. However, we want to determine a common offset value $\text{offset}_{\mathcal{B}}$ such that the actual potential and the actual reduced costs associated with each vertex $u \in \mathcal{B}_i$ can be computed with respect to $\text{offset}_{\mathcal{B}}$. The following strategy assures that the offsets of all defining subblossoms \mathcal{B}_i , $1 \leq i \leq 2k+1$, are set to 0 and thus the desired result is achieved by the common offset $\text{offset}_{\mathcal{B}} = 0$.

Whenever a surface blossom \mathcal{B}' (trivial or non-trivial) becomes an even tree blossom, its offset $\text{offset}_{\mathcal{B}'}$ is set to 0. Consequently, in order to preserve the validity of (3) and (2) for the computation of the actual potential \tilde{y}_u of each vertex $u \in \mathcal{B}'$ and of the actual potential $\tilde{z}_{\mathcal{B}'}$ of \mathcal{B}' itself (if \mathcal{B}' is non-trivial only), the following adjustments have to be performed:

$$\begin{aligned} y'_u &= y_u + \text{offset}_{\mathcal{B}'}, \\ z'_{\mathcal{B}'} &= z_{\mathcal{B}'} - 2\text{offset}_{\mathcal{B}'}. \end{aligned}$$

Moreover, the stored reduced cost π_{uv_i} of each edge uv_i associated with each vertex $u \in \mathcal{B}'$ is subject to correction:

$$\pi'_{uv_i} = \pi_{uv_i} + \text{offset}_{\mathcal{B}'}.$$

Observe that the adjustments are performed at most once per phase for a fixed vertex. Thus, the time required for the potential adjustments is $O(n)$ per phase. On the other hand, the corrections of the reduced costs contributes total time $O(m \log n)$ per phase.

In summary, we have established a convenient way to handle the varying potentials as well as the reduced costs of edges associated with a vertex. The values of interest can be computed on demand by the developed formulae. It has been shown that the additional overhead produced by the offset maintenance consumes $O(m \log n)$ time per phase.

This concludes the description of our approach (for a more extensive discussion the reader is referred to [Schäfer 2000]).

3.3 Construction of Initial Solutions

Both Blossom IV and our implementation support two strategies for finding initial solutions; both of them are well known and also used in previous codes.

3.3.1 Greedy Heuristic. The *greedy heuristic* first sets the vertex potentials: the potential y_u of a vertex u is set to one half of the weight of the heaviest incident edge. This guarantees that all edges have non-negative reduced cost. It then chooses a matching within the tight edges in a greedy fashion.

3.3.2 Fractional Matching Heuristic. The *fractional matching heuristic* [Derigs and Metz 1986] first solves the fractional matching problem (consisting of constraints (WPM)(1) and (WPM)(3) only). In the solution all variables are half-integral and the edges with value $1/2$ form odd length cycles. The initial matching consists of the edges with value 1 and of $\lfloor |C|/2 \rfloor$ edges from every odd cycle. Applegate and Cook [Applegate and Cook 1993] describe how to solve the fractional matching problem.

Our fractional matching algorithm uses priority queues and similar strategies to the ones described above. The priority queue based approach appears to be highly efficient.¹¹

3.4 Correctness

Blossom IV and our program does not only compute an optimal matching M but also an optimal dual solution. This makes it easy to verify the correctness of a solution. One only has to check that M is a (perfect) matching, that the dual solution is feasible (in particular, all edges must have non-negative reduced costs), and that the complementary slackness conditions are satisfied.

4. EXPERIMENTAL RESULTS

4.1 Experimental Setting

We experimented with three kinds of instances: triangulation instances (including Delaunay triangulations), sparse and dense random instances containing a perfect matching and complete geometric instances.

The triangulation instances were constructed as follows. First, n random points were chosen from a $2^{20} \times 2^{20}$ square. Then, a triangulation of the point set was computed. We experimented with two kinds of triangulations: Delaunay triangulations and triangulations computed by a sweep-line algorithm (called sweep-line triangulations henceforth). Both triangulations were constructed using the appropriate functions of LEDA. Finally, each edge in the triangulation was assigned a weight equal to the Euclidean distance of the endpoints (rounded down). Delaunay triangulations are known to contain perfect matchings [Dillencourt 1990].

For the random instances we chose random graphs with n vertices. The number of edges for sparse graphs was chosen as $m = \alpha n$ for small values of α , $\alpha \leq 10$. For dense graphs the density is approximately 20%, 40% and 60% of the density of

¹¹Experimental results (not presented in this paper) showed that the priority queue approach is substantially faster than the specialized algorithm of LEDA to compute a maximum-weight (perfect) matching in a bipartite graph. A closer inspection revealed that this superiority is due to a heuristic improvement which is described in detail in [Mehlhorn and Schäfer 2001].

Table 1. MS_{SST} vs. MS_{MST} on sweep-line triangulations. $-$, $+$ or $*$ indicates usage of no, the greedy or the fractional matching heuristic. The time needed by the greedy or the fractional matching heuristic is shown in the columns GY and FM.

n	MS_{SST}^-	MS_{MST}^-	MS_{SST}^+	MS_{MST}^+	GY	MS_{SST}^*	MS_{MST}^*	FM	t
10000	37.01	6.27	24.05	4.91	0.13	5.79	3.20	0.40	5
20000	142.93	14.81	89.55	11.67	0.24	18.54	8.00	0.83	5
40000	593.58	31.53	367.37	25.51	0.64	76.73	17.41	1.78	5

Table 2. $B4^*$, $B4_{var}^*$ vs. MS^* on Delaunay triangulations.

n	$B4^*$	$B4_{var}^*$	MS^*	t
10000	30.00	2.69	4.88	20
20000	101.96	6.00	10.12	20
40000	374.67	17.60	21.68	20
80000	—	64.32	46.07	20
160000	—	138.29	98.13	20

a complete graph. Random weights out of the range $[1, \dots, 2^{16})$ were assigned to the edges. We checked for perfect matchings with the LEDA cardinality matching implementation.

Complete geometric instances were induced by n random points in an $n \times n$ square and their Euclidean distances.

All our running times are in seconds and are the average of $t = 5$ runs, unless stated otherwise. All experiments were performed on a Sun Ultra Sparc, 333 Mhz.

4.2 Experiments

4.2.1 Initial Solution, Single vs. Multiple Search Tree Strategies. Table 1 compares the usage of different heuristics in combination with the single search tree (MS_{SST}) and the multiple search tree (MS_{MST}) approach.

The fractional matching heuristic is computational more intensive than the greedy heuristic, but leads to overall improvements of the running time. The multiple search tree strategy is superior to the single search tree strategy with both heuristics. We therefore take the multiple search tree strategy with the fractional matching heuristic as our canonical implementation; we refer to this implementation as MS^* . Blossom IV also uses the fractional matching heuristic for constructing an initial solution. We remark that the difference between the two heuristics is more pronounced for the single search tree approach.

4.2.2 Triangulation Instances. Table 2 compares Blossom IV with the fractional matching heuristic, multiple search trees without ($B4^*$) and with ($B4_{var}^*$) the variable δ approach with MS^* . We used Delaunay triangulations.

The table shows that the variable δ approach $B4_{var}^*$ makes a tremendous difference for Blossom IV and that MS^* is competitive to $B4_{var}^*$.

It seems that Delaunay triangulations constructed as described above are rather simple instances. In Table 3 we present experiments on sweep-line triangulations. MS^* is significantly faster than $B4_{var}^*$.

In Table 4 we give some detailed information about the number of operations performed by the algorithms on sweep-line triangulations. Observe that the run-

Table 3. $B4^*$, $B4_{\text{var}}^*$ vs. MS^* on sweep-line triangulations.

n	$B4^*$	$B4_{\text{var}}^*$	MS^*	t
10000	26.44	9.89	5.16	20
20000	98.69	24.82	11.06	20
40000	509.07	95.96	24.09	20
80000	—	346.51	52.46	20
160000	—	2373.18	121.99	20

Table 4. $B4_{\text{var}}^*$ vs. MS^* on sweep-line triangulations with $n = 40000$ vertices. *heur* and *free* denotes the time needed and the number of vertices left unmatched by the fractional matching heuristic, respectively. *adj*, *shrinks*, *expands* and *updates* states the number of dual adjustments, shrinks, expands and potentials being updated. Note that the value *updates* for MS^* corresponds to hypothetical updates, since, as opposed to $B4_{\text{var}}^*$, our implementation does not update the potentials explicitly. (The values of *updates* differ so drastically since different LP formulations are used.) *time* refers to the time needed to match the additional *free* many vertices and *total* denotes the overall running time of the algorithm.

Alg.	<i>heur</i>	<i>free</i>	<i>adj</i>	<i>shrinks</i>	<i>expands</i>	<i>updates</i>	<i>time</i>	<i>total</i>
$B4_{\text{var}}^*$	1.45	1408	19532	8575	653	6467296	166.07	167.52
MS^*	1.81	1408	21044	10273	2024	(239707800)	21.89	23.70
$B4_{\text{var}}^*$	0.98	1408	12466	4913	433	1179070	22.43	23.41
MS^*	1.81	1408	15812	7058	1766	(147217992)	20.17	21.98
$B4_{\text{var}}^*$	0.96	1428	15790	5953	747	2886144	61.90	62.86
MS^*	1.87	1428	17710	7768	2365	(176127169)	21.59	23.46
$B4_{\text{var}}^*$	0.98	1446	13996	5667	495	1764564	34.13	35.11
MS^*	1.81	1446	16322	7402	1777	(160005292)	20.99	22.80
$B4_{\text{var}}^*$	1.20	1440	18176	7852	764	8214055	182.03	183.23
MS^*	1.84	1440	21007	9841	2565	(263387831)	23.31	25.15
$B4_{\text{var}}^*$	0.94	1464	14316	5529	538	2166822	46.55	47.49
MS^*	1.77	1464	17114	7506	2280	(181317577)	22.37	24.14
$B4_{\text{var}}^*$	1.71	1444	21294	10205	713	10621905	252.71	254.42
MS^*	1.79	1444	24687	12530	2475	(394330879)	24.33	26.12
$B4_{\text{var}}^*$	1.06	1474	17230	6431	756	3670495	87.46	88.52
MS^*	1.85	1474	18408	8509	2237	(202292903)	22.82	24.67
$B4_{\text{var}}^*$	1.68	1474	22488	10585	967	11353087	278.25	279.93
MS^*	1.78	1474	24505	12140	2573	(367147596)	24.27	26.05
$B4_{\text{var}}^*$	1.00	1416	11180	4079	338	472361	9.62	10.62
MS^*	1.79	1416	14231	6070	1764	(126532110)	19.27	21.06

ning time of $B4_{\text{var}}^*$ varies between 10.62 and 279.93 seconds. The running time of our implementation varies only slightly. In general, MS^* performs more dual adjustments, shrink and expand operations than $B4_{\text{var}}^*$. The running time of $B4_{\text{var}}^*$ heavily depends on the number of potential updates; observe the strong correlation between columns *updates* and *time* for $B4_{\text{var}}^*$. This dependence suggests the assumption that Blossom IV spends most of its time to perform dual adjustments. To verify our conjecture, we profiled the code of Blossom IV with the GNU profiler **gprof**: Blossom IV indeed spends more than 50% of its computation time in the functions which determine the dual adjustment value δ (33.63%) and perform the potential updates (20%).

4.2.3 Influence of Edge Weights. Table 5 shows the influence of the edge weights on the running time. We took random graphs with $m = 4n$ edges and random edge

Table 5. B4*, B4*_{var} vs. MS* on random instances with variable weight range.

n	α	b	B4*	B4* _{var}	MS*	t
10000	4	1	3.98	3.99	0.85	1
10000	4	10	2.49	3.03	2.31	1
10000	4	100	3.09	3.10	2.58	1
10000	4	1000	17.41	8.40	2.91	1
10000	4	10000	13.69	11.91	2.78	1
10000	4	100000	12.06	11.20	2.69	1

Table 6. B4*, B4*_{var} vs. MS* on sparse random instances with $m = 6n$.

n	α	B4*	B4* _{var}	MS*	t
10000	6	20.94	18.03	3.51	5
20000	6	82.96	53.87	9.97	5
40000	6	194.48	177.28	29.05	5

weights in the range $[1, \dots, b]$ for different values of b . For $b = 1$, the problem is unweighted.

The running time of B4* and B4*_{var} depends significantly on the size of the range, the running time of MS* depends only weakly (except for the unweighted case which is simpler). MS* is superior to B4*_{var}.

We try an explanation. When the range of edge weights is small, a dual adjustment is more likely to make more than one edge tight. Also it seems to take fewer dual adjustments until an augmenting path is found. Since dual adjustments are cheaper in our implementation ($O(m \log n)$ for all adjustments in a phase of our implementation versus $O(n)$ for a single adjustment in Blossom IV), our implementation is less harmed by large numbers of adjustments.

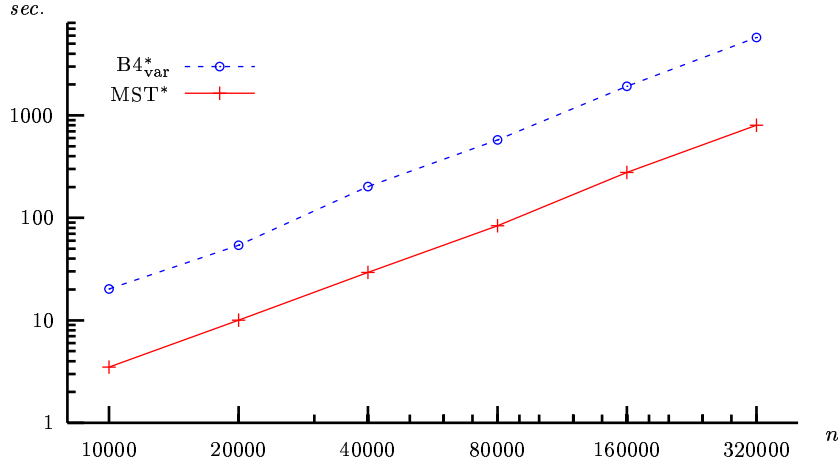
4.2.4 Asymptotics. Tables 6 and 7 give some information about the asymptotics. For Table 6 we have fixed the graph density at $m = 6n$ and varied n .

The running times of B4*_{var} and MS* seem to grow less than quadratically (with B4*_{var} taking about six times as long as MS*). Table 7 gives more detailed information. We varied n and α .

A log-log plot indicating the asymptotics of B4*_{var} and our MS* algorithm on random instances ($\alpha = 6$) is depicted in Fig. 1.

Table 7. B4*, B4*_{var} vs. MS* on sparse random instances with $m = \alpha n$.

n	α	B4*	B4* _{var}	MS*	t
10000	6	20.90	20.22	3.49	5
10000	8	48.50	22.83	5.18	5
10000	10	37.49	30.78	5.41	5
20000	6	96.34	54.08	10.04	5
20000	8	175.55	89.75	12.20	5
20000	10	264.80	102.53	15.06	5
40000	6	209.84	202.51	29.27	5
40000	8	250.51	249.83	36.18	5
40000	10	710.08	310.76	46.57	5

Fig. 1. Asymptotics of $B4^*_{\text{var}}$ and MS^* algorithm on sparse random instances ($\alpha = 6$).

n	α	$B4^*_{\text{var}}$			MS^*			t
		best	worst	average	best	worst	average	
10000	6	16.88	20.03	18.83	3.34	4.22	3.78	5
20000	6	49.02	60.74	55.15	9.93	11.09	10.30	5
40000	6	162.91	198.11	180.88	25.13	32.24	29.09	5

4.2.5 Variance in Running Time. Table 8 gives information about the variance in running time. We give the best, worst and average time of five instances. The fluctuation is about the same for both implementations.

4.2.6 Dense Random Instances and Price-and-Repair. Our experiments suggest that MS^* is superior to $B4^*_{\text{var}}$ on sparse instances. Table 9 shows the running time on dense random instances. Our algorithm is superior to Blossom IV even on these instances.

As mentioned above, Blossom IV provides a price-and-repair heuristic for complete geometric instances. The running time on these instances is significantly

Table 9. $B4^*$, $B4^*_{\text{var}}$ vs. MS^* on dense random instances. The density is approximately 20%, 40% and 60%.

n	m	$B4^*$	$B4^*_{\text{var}}$	MS^*	t
1000	100000	6.97	5.84	1.76	5
1000	200000	16.61	11.35	3.88	5
1000	300000	18.91	18.88	5.79	5
2000	200000	46.71	38.86	8.69	5
2000	400000	70.52	70.13	16.37	5
2000	600000	118.07	115.66	23.46	5
4000	400000	233.16	229.51	42.32	5
4000	800000	473.51	410.43	92.55	5
4000	1200000	523.40	522.52	157.00	5

Table 10. Blossom IV variable δ without ($B4_{\text{var}}^*$) and with ($B4_{\text{var}}^{**}$) price-and-repair heuristic vs. MS^* on complete geometric instances. The Delaunay graph of the point set was chosen as the sparse subgraph for $B4_{\text{var}}^{**}$.

n	$B4_{\text{var}}^*$	$B4_{\text{var}}^{**}$	MS^*	t
1000	37.01	0.43	24.05	5
2000	225.93	1.10	104.51	5
4000	1789.44	4.33	548.19	5

Table 11. $B4_{\text{var}}^*$ vs. MS^* on chains.

$2n$	$B4_{\text{var}}^*$	MS^*	t
10000	94.75	0.25	1
20000	466.86	0.64	1
40000	2151.33	2.08	1

improved for $B4_{\text{var}}^*$ using the price-and-repair heuristic as can be seen in Table 10.

4.2.7 ‘Worse-Case’ Instances for Blossom IV. We wish to conclude the experiments with two ‘worse-case’ instances that demonstrate the superiority of our implementation to Blossom IV.

The first ‘worse-case’ instance for Blossom IV is simply a chain. We constructed a chain having $2n$ vertices and $2n - 1$ edges. The edge weights along the chain were alternately set to 0 and 2 (the edge weight of the first and last edge equal 0). $B4_{\text{var}}^*$ and our MS^* algorithm were asked to compute a maximum-weight perfect matching. Note that the fractional matching heuristic will always compute an optimal solution on instances of this kind. Table 11 shows the results.

The running time of $B4_{\text{var}}^*$ grows more than quadratically (as a function of n), whereas the running time of our MS^* algorithm grows about linearly with n . We present our argument as to why this is to be expected. First of all, the greedy heuristic will match all edges having weight 2; the two outer vertices remain unmatched. Each algorithm will then have to perform $O(n)$ dual adjustments so as to obtain the optimal matching. A dual adjustment takes time $O(n)$ for Blossom IV (each potential is explicitly updated), whereas it takes $O(1)$ for our MS^* algorithm. Thus, Blossom IV will need time $O(n^2)$ for all these adjustments and, on the other hand, the time required by our MS^* algorithm will be $O(n)$.

Another ‘worse-case’ instance for Blossom IV occurred in VLSI-Design having $n = 151780$ vertices and $m = 881317$ edges. Kindly, André Rohe made this instance available to us. We compared $B4^*$ and $B4_{\text{var}}^*$ to our MS algorithm. We ran our algorithm with the greedy heuristic (MS^+) as well as with the fractional matching heuristic (MS^*). The results are given in Table 12.

The second row states the times that were needed by the heuristics. Observe that both Blossom IV algorithms need more than two days to compute an optimal matching, whereas our algorithm solves the same instance in less than an hour.

Table 12. $B4^*$, $B4_{\text{var}}^*$ vs. MS on **boese.edg** instance.

n	m	$B4^*$	$B4_{\text{var}}^*$	MS^+	MS^*	t
151780	881317	200019.74 (332.01)	200810.35 (350.18)	3172.70 (5.66)	5993.61 (3030.35)	1

For our MS algorithm the fractional matching heuristic did not help at all on this instance: to compute a fractional matching took almost as long as computing an optimal matching for the original instance (using the greedy heuristic).

5. CONCLUSION

We have described an implementation of an $O(nm \log n)$ algorithm for the weighted matching problem. The experiments showed that our implementation is superior to the most efficient code Blossom IV of Cook and Rohe [Cook and Rohe 1997] on almost all instances. Only on complete geometric instances, for which Blossom IV additionally supports a highly effective price-and-repair strategy, our code was inferior. We can thus provide an affirmative answer to the question whether or not sophisticated data structures, such as concatenable priority queues, help to improve the performance of weighted matching algorithms in practice. Moreover, we think that the presented results give support to our belief that it is of a great practical value to have an all-purpose library such as LEDA, since the implementation of sophisticated data structures and algorithms from scratch would not be economical.

Our research raises several questions. (1) Is it possible to integrate the variable δ approach into an $O(nm \log n)$ algorithm? (2) A generator of instances forcing the implementations into their worst-case would be useful. (3) In order to handle complete geometric instances more efficiently the effect of a price-and-repair strategy is worth being considered; most likely, providing such a mechanism for MS* will improve its worst-case behaviour on these instances tremendously.

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