## **Algorithms for the Steiner Problem in Networks**

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### Abstract

The Steiner problem in networks is the problem of connecting a set of required vertices in a weighted graph at minimum cost. It is a classical  $\mathcal{NP}$ -hard problem with many important applications. For this problem we develop, implement and test several new techniques. On the side of lower bounds, we present a hierarchy of linear relaxations and class of new relaxations that are the currently strongest polynomially solvable linear relaxations. On the side of preprocessing techniques, we improve some known reduction tests and introduce powerful new ones. For upper bounds we introduce the successful concept of heuristic reductions. Finally, we integrate these blocks into an exact algorithm. For the exact algorithm and for the different components we present very good computational results on the large benchmark library SteinLib.

## Kurzzusammenfassung

Das Steiner Problem in Netzwerken ist das Problem, eine Menge von Basisknoten in einem gewichteten Graphen kostenminimal zu verbinden. Es ist ein klassisches  $\mathcal{NP}$ -schweres Problem mit vielen Anwendungen. Für dieses Problem entwickeln, implementieren und bewerten wir einige neue Techniken. Bezüglich unterer Schranken stellen wir eine Hierarchie von Linearen Relaxationen auf und entwickeln eine Klasse von Relaxationen, die die zur Zeit stärksten polynomiell lösbaren Linearen Relaxationen darstellen. Im Bereich des Preprocessing verbessern wir einige bekannte Reduktionstests und entwickeln einige starke neue Tests. Für obere Schranken führen wir das erfolgreiche Konzept der "heuristischen Reduktionen" ein. Schließlich werden diese Bausteine zu einem exakten Algorithmus zusammengesetzt. Sowohl der exakte Algorithmus, als auch die einzelnen Komponenten erzielen sehr gute experimentelle Resultate auf der umfassenden Benchmarkbibliothek SteinLib.

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# Chapter 1

# Introduction

#### **1.1 About This Work**

The Steiner problem in networks is the problem of connecting a set of required vertices in a weighted graph at minimum cost. This is a classical  $\mathcal{NP}$ -hard problem with many important applications in network design in general and VLSI layout in particular. The primary goal of our research has been the development of empirically successful algorithms. This means we designed and implemented algorithms that

- 1. generate Steiner trees of low cost in reasonable running times (upper bounds),
- 2. prove the quality of a Steiner tree by providing a lower bound on the optimal value (lower bounds),
- 3. or find an optimal Steiner tree (exact algorithms).
- 4. As an important prerequisite for the first three tasks, we used preprocessing techniques to reduce the size of the original problem without changing the optimal solution (reduction tests).

The value of our algorithms is measured by comparing our results to those of other research groups on the huge and well-established benchmark library for Steiner tree problems [SteinLib, KMV01]. In the case of exact algorithms, this measure is the best one can get because of the  $\mathcal{NP}$ -hardness result. For upper and lower bounds, the extensive experimental evaluation gives much sharper and much more relevant information than a worst-case analysis.

Why is one interested in solving Steiner tree problems? First of all, there are a lot of practical problems that can be modeled as Steiner tree problems. The question remains why an optimal solution is relevant. One could argue that in practical applications the computation of a heuristic Steiner tree and a lower bound that proves a gap of less than one percent is sufficient, because typically there are bigger inaccuracies in the model anyway. Still, the development of exact algorithms has a solid practical justification. The best algorithms for upper and lower bounds rely heavily on the computation of very sharp bounds for subproblems of the original problem, and the best results are achieved when an optimal solution of these subproblems can be found. In addition, the empirical success of our exact algorithms (even large benchmark instances with 30000 vertices can be solved in a matter of minutes) puts into question the justification of heuristic upper bound algorithms.

Our work is also motivated from the theoretical side: Classical  $\mathcal{NP}$ -hard combinatorial optimization problems like the "Traveling Salesman Problem" (TSP), scheduling problems, and the Steiner tree problem, have attracted researchers as test environments for their methods for decades. As a consequence, all important techniques in the fields of Operations Research and Combinatorial Optimization have been tried on these problems, leading to a respected competition for the best implementation. The relevance of this is twofold: On the one hand, the current state of affairs (which problems can be solved and what are the best known upper and lower bounds) reflects to some extent the state of the art in solving combinatorial optimization problems. On the other hand, these classical problems have always been "creation engines" for new methods and techniques that are also applicable to other problems. The outcome has been such general techniques as the cutting plane approach, which was invented in the TSP context. Likewise in this work, we present appealing approaches (e.g., partitioning as reduction technique, reduction-based heuristics) that may be applicable to other problems.

Similar to many other elaborate optimization packages, our package for the Steiner tree problem consists of a large collection of different components that interact extensively. In fact, our best programs for generating upper bounds, lower bounds, and exact solution all use essentially the same code, and just arrange the use of the components in different ways. Therefore, it is not possible to give a concise description of "how to produce a good upper bound" in some dozen lines of pseudo-code. Hence, we have to give a bottom-up description: We will first describe the different building blocks separately and give pointers to the necessary connections of the blocks elsewhere. Still, we cannot provide a close grained picture of our program. This becomes obvious given the fact that merely printing the code without any further explanation requires roughly 1000 pages. Therefore, we describe the algorithms on a rather abstract level, and give only pointers to the description of standard techniques used.

This work summarizes the current state of our research, which has already been presented at conferences (ESA, APPROX, Combinatorial Optimization) and has been published in journals (Discrete Applied Mathematics, Operations Research Letters). It has already received considerable recognition in research on the Steiner tree problem, visible in citations in recently published literature [RUW02, PW02, UdAR99, Uch01, KMV01, CT01] and university lectures [JKP<sup>+</sup>02].

In the following, we will give a list of our main contributions:

#### Lower Bounds:

- There are many (mixed) integer programming formulations of the Steiner problem. The corresponding linear programming relaxations are of great interest particularly, but not exclusively, for computing lower bounds, but not much was known about the relative quality of these relaxations. We compare the linear relaxations of all classical, frequently cited integer programming formulations of this problem from a theoretical point of view with respect to their optimal values. We present several new results, establishing very clear relations between relaxations, which have often been treated as unrelated or incomparable, forming a **hierarchy of relaxations**.
- We introduce a **collection of new relaxations** that are stronger than any known relaxation that can be solved in polynomial time, and place the new relaxations into our hierarchy. Further, we show how such a relaxation can be used in practical algorithms. Except for the flow-balance constraints introduced by [KM98], this is the first successful attempt to use a relaxation that is stronger than Wong's directed cut relaxation from 1984 [Won84].

#### **Preprocessing/Reduction techniques:**

- For some of the classical reduction tests, which would have been too time-consuming for large instances in their original form, we design **efficient realizations**, improving the worst-case running time to  $O(m + n \log n)$  in many cases. Furthermore, we design new tests, filling some of the gaps left by the classical tests.
- Previous reduction tests were either alternative based or bound based. That means to simplify the problem they either argued with the existence of alternative solutions, or they used some constrained lower bound and upper bound. We develop a framework for **extended reduction tests**, which extends the scope of inspection of reduction tests to larger patterns and combines for the first time alternative-based and bound-based approaches effectively.
- We introduce the new concept of **partitioning-based reduction techniques**, which has a significant impact on the reduction results in some cases.
- We integrate all tests into a **reduction packet**, which performs stronger reductions than any other package we are aware of. Additionally, the reduction results of other packages can be achieved typically in a fraction of the running time. ("The result reported in Polzin

and Daneshmand is by far the best result in terms of both number of instances solved to optimality and computational times factoring into the difference in cpu speed."[CT01].)

#### **Upper Bounds:**

- We present variants of known path heuristics, including an empirically fast variant with a fast worst-case running time of  $O(m + n \log n)$ . The previous running time for this kind of path heuristic was  $O(rm \log n)$ .
- We introduce a new meta-heuristic, **reduction-based heuristics**. On the basis of this concept, we develop heuristics that achieve typically sharper upper bounds than the strongest known heuristics for this problem despite running times that are smaller by orders of magnitude.

#### **Exact Algorithm:**

We integrate the previously mentioned building blocks into an exact algorithm that achieves very good running times.

- For most benchmark instances the program computes the exact solution in running times that are shorter than the running times of other authors by orders of magnitude. ("Computational times reported in [PV01c] are far better than the rest." [CT01])
- There are 73 instances in SteinLib that have not been solved by any other research group. We have been able to solve 32 of them.
- For geometric Steiner problems, our algorithm for general networks is (together with a preprocessing phase [WWZ01] that exploits some of the geometric properties) the fastest algorithm and beats the specially tailored MSTH approach [WWZ00], which has received much attention.

Additionally, we present a procedure that uses the fixed-parameter tractability of the Steiner problem for subgraphs of small width.

In Chapter 2, we study some relaxations of the problem and methods for computing lower bounds using them; they are also frequently used in the following chapters. In Chapter 3, reduction techniques are discussed, which play a central role in our approach. These techniques are also the basis of the reduction-based heuristics, which we introduce in Chapter 4 on upper bounds. In Chapter 5, the building blocks from the previous chapters are integrated into an exact algorithm, which is shown to be empirically successful.

Some comparison of our empirical results to those of other authors is presented in Section 3.8 for reduction techniques, in Section 4.6 for upper bounds, and in Section 5.4 for the exact algorithm. Detailed results for the exact algorithm are given in the appendix (see page 105).

Most of the background information relevant to this work can be found in Hwang, Richards and Winter [HRW92]; we have tried to keep the notation compatible with that book. The basic definitions are repeated in Section 1.3.

The implementation and most of the results presented were produced jointly with Siavash Vahdati Daneshmand [PV00, PV01a, PV01c, PV01e, PV02a, PV02b, PV03]. I declare that my contribution to these results constituted at least half of the work.

#### **1.2** About Experimental Results in this Work

In each of the following sections, we will report on the experimental behaviour of certain algorithms. We do not claim that algorithms can be evaluated beyond doubt by running them on a set of test instances. But when considering (exact) algorithms for an  $\mathcal{NP}$ -hard problem, there is no fully satisfactory alternative. Proving guaranteed performance ratios for certain components (like heuristics for computing upper bounds) cannot be a complete substitute, because such results are often too pessimistic due to their worst case character or lack of better proof techniques. From a comparative point of view, a much sharper differentiation is necessary; particularly in the context of exact algorithms, where even marginal differences (small fractions of a percent) in the value of the bounds can have a major impact on the behaviour of the algorithm.

In addition, we consider the comparability of results a critical issue, which strongly suggests using benchmark instances. There are two major benchmarks for the Steiner problem in networks: the collection in the OR-Library [Bea90] and SteinLib [SteinLib]. The instances of the OR-Library are much older, with the advantage that more comparative results exist on them. On the other hand, only one type of instance is represented (sparse and random). The library SteinLib is much more extensive, containing instances of all common types. But giving experimental results for all these instances in each section would make the work unreasonably long, so we have chosen a compromise option: For the intermediary results (for example concerning upper bounds or reductions), we give average results on each group of the problem instances from SteinLib (if the gap to the optimal solution is measured, we restrict ourselves to those groups where all optimal values are known). For the final results of the complete algorithm, however, we additionally give results for all instances in SteinLib. We leave aside some very small and easy instance groups.

In Table 1.1, we give a brief description of the instance classes of SteinLib. For more comprehensive information, see [SteinLib]. The column "Instances" gives the number of instances in the group. The column "Status" shows whether all instances of this column have been solved by other authors and by us ("solved"), or only by us ("*solved*" in italics), or if there are some "unsolved" instances.

Also it must be mentioned that for actual tests, we did not always implement the data structures and algorithms with the best known (worst-case) time bound, especially if the extra work did not seem to pay off. So, statements concerning worst-case time bounds for a component merely mean the possibility of implementation of that component with that bound.

All results in this work (except for Section 5.4.1) were produced single-threaded on a Sunfire 15000 with 900 MHz SPARC III+ CPUs, using the operating system SunOS 5.9. We always used the GNU g++ 2.95.3 compiler with the -O4 flag. As it is a multi-processor machine with shared memory, it is slower than a single processor system with the same processor. A comparison of the running times in Section 5.4.1 and in the appendix shows that the machine is approximately half as fast as a PC with an AMD Athlon XP 1800+ (1.53 GHz) processor, which was used in Section 5.4.1.

We used ExpLab [HPKS02] and CVS to address the issue of reproduceability of experiments.

#### **1.3 Definitions and Notations**

For any undirected graph G = (V, E), we define n := |V|, m := |E|, and assume that  $(v_i, v_j)$  and  $(v_j, v_i)$  denote the same (undirected) edge  $\{v_i, v_j\}$ . A network is a weighted graph (V, E, c) with an edge weight function  $c : E \to \mathbb{R}$ . We sometimes refer to networks simply as graphs. For each edge  $(v_i, v_j)$ , we use terms like cost, weight, length, etc. of  $(v_i, v_j)$  interchangeably to denote  $c((v_i, v_j))$  (also denoted by  $c(v_i, v_j)$ ). For any directed network  $\vec{G} = (V, A, c)$ , we use  $[v_i, v_j]$  to denote

Class Name	Instances	V	Status	Description
D	20	1000	solved	sparse random with varying graph param-
E	20	2500	solved	∫ eters, OR-Library
Х	3	52-666	solved	complete with Euclidean weights
ES1000FST	15	2532–2984	solved	rectilinear, derived with geosteiner
ES10000FST	1	27019	solved	[WWZ01] from 1000 (rsp. 10000) random
TSPFST	76	89–17127	solved	points in the plane, rsp. instances from TSPLIB [Rei91], only other algorithm for solving these instances uses additional in-
1080	100	80	solved	formation computed in the geometric pre- processing phase (see Section 3.6.1)
I160	100	160	solved	incidence networks, constructed with
I320	100	320	solved	the aim of being difficult for known
I320 I640	100	640	unsolved	techniques, introduced by Duin [Dui93].
MC	6	97-400	solved	constructed difficult instances
PUC	50	64-4096	unsolved	constructed difficult instances: hypercubes,
				from code covering and bipartite graph [RdAR <sup>+</sup> 01].
SP	8	6–3997	unsolved	constructed instances, combination of odd wheels and odd circles, difficult for Linear
				Programming approaches
VLSI	116	90–36711	solved	grid graph with holes (not metric) from VLSI
				design, SteinLib instance groups alue,
				alut, diw, dmxa, gap, msm, and taq
LIN	37	53-38418	solved	grid graph with holes (not metric) from VLSI
				design
WRP3	63	84–3168	solved	) wire routing problems from industry
WRP4	62	110–1898	solved	∫[ZR00]
1 <b>R</b>	27	1250	solved	2D cross grid graph [Fre97]
2R	27	2000	solved	3D cross grid graph [Fre97]

Table 1.1: Classes of Problem Instances in [SteinLib]

the directed edge, or arc, from  $v_i$  to  $v_j$ ; and define a := |A|.

The **degree** of a vertex  $v_i \in V$  is the number of incident edges of  $v_i$ . For any network G, c(G) denotes the sum of the edge weights of G.

The Steiner problem in networks (NSP) can be formulated as follows: Given a network G = (V, E, c) and a non-empty set  $R, R \subseteq V$ , of required vertices (or terminals), find a subnetwork  $T_G(R)$  of G that contains a path between every pair of terminals and minimizes  $\sum_{(v_i, v_j) \in T_G(R)} c_{ij}$ .

We define r := |R|. For ease of notation we assume  $R = \{v_1, \ldots, v_r\}$ . If we want to stress that  $v_i$  is a terminal, we will write  $z_i$  instead of  $v_i$ . The vertices in  $V \setminus R$  are called **non-terminals**. Without loss of generality, we assume that the edge weights are positive and that G (and  $T_G(R)$ ) are connected. Now  $T_G(R)$  is a tree, called **Steiner minimal tree** (for historical reasons). A **Steiner tree** is an acyclic, connected subnetwork of G, spanning (a superset of) R. We call non-terminals in a Steiner tree its **Steiner nodes**.

The directed version of this problem (also called the Steiner arborescence problem) is defined similarly (see [HRW92]): In addition to G and R, a root  $z_1 \in V$  is given and it is required that the solution contains a path from  $z_1$  to every terminal in R. Every instance of the undirected version can be transformed into an instance of the directed version in the corresponding bidirected network, by fixing a terminal  $z_1$  as the root. We define:  $R^{z_1} := R \setminus \{z_1\}$ .

With  $d(v_i, v_j), d_{ij}$  we denote the length of a shortest path between  $v_i$  and  $v_j$ .

For a given network G = (V, E, c) and  $W \subseteq V$ , the corresponding distance network is defined as  $D_G(W) = (W, W \times W, d)$ .

For each terminal  $z_i$ , one can define a neighborhood  $N(z_i)$  as the set of vertices that are not closer to any other terminal. More precisely, a partition of V is defined:

$$V = \bigcup_{z_i \in R} N(z_i) \text{ with } v_j \in N(z_i) \Rightarrow d(v_j, z_i) \le d(v_j, z_k) \quad \text{ (for all } z_k \in R)$$

If  $v_j \in N(z_i)$ , we call  $z_i$  the **base** of  $v_j$  (written  $base(v_j)$ ). In accordance with the parlance of algorithmic geometry, we call  $N(z_i)$  the **Voronoi region** of z. We consider two terminals  $z_i$  and  $z_j$  as neighbors if there is an edge  $(v_k, v_l)$  with  $v_k \in N(z_i)$  and  $v_l \in N(z_j)$ . Given G and R, the Voronoi regions can be computed in time  $O(m + n \log n)$ . Using them, a minimum spanning tree for the corresponding distance network  $D_G(R)$  (we denote this tree by  $T'_D(R)$ ) can be computed in the same time [Meh88].

#### **1.4 Applications and Background**

#### 1.4.1 History

The Steiner problem in networks is the combinatorial variant of the much older Euclidean Steiner problem, which asks for the minimal tree that connects a given set of points in the plane. A special case of this problem has already been discussed before 1640 by Fermat:

Given three points in the plain, find a point that minimizes the sum of the distances to the given points.

Jacob Steiner (1796-1863) considered a generalization of this problem for r points (the **generalized Fermat problem**), but not the (Euclidean) Steiner problem. These two problems are identical only in the case r = 3. The "Steiner" problem is believed to have been presented first by Gauß (see [Aro96]). The first (terminating) algorithm for the Euclidean Steiner problem was given by Melzak

#### 1.4. APPLICATIONS AND BACKGROUND

[Mel61]. More information on the Euclidean Steiner problem and its history is contained in Hwang et al. [HRW92]. Its relation to the network variant will be discussed later.

The Steiner problem in networks was explicitly formulated for the first time by Hakimi [Hak71] and Levin [Lev71]. Since then hundreds of articles have been published concerning different variants of this problem. A good (although not fully up-to-date) overview is given in Hwang et al. [HRW92].

#### 1.4.2 Related Problems

#### **Euclidean Steiner Problem (ESP)**

**Definition 1** Given a finite set R of points in the (Euclidean) plane, find a point set S together with a minimal spanning tree T for  $R \cup S$ , such that T has minimal length concerning the  $L_2$ -norm (Euclidean distance).

In comparison to the ESP, the Steiner problem in networks (NSP) is in some sense more general: The cost function can be general and the network does not need to have geometric properties. On the other hand, in the NSP the set of potential Steiner nodes is finite, whereas in the ESP every point in the plain can be part of a feasible solution. But it was proven that a minimal Steiner tree for an ESP instance has at most r - 2 (r = |R|) Steiner points (nonterminals of degree at least 3) and that the number of possible topologies is finite [HRW92].

Furthermore it holds for every ESP instance I and for every  $\epsilon > 0$  that it can be approximated by an NSP instance  $I_{\epsilon}$  with fewer than  $const \cdot \frac{r^2}{\epsilon^2}$  vertices (i.e., the quotient of the lengths of the optimal solutions of  $I_{\epsilon}$  and I is at most  $1 + \epsilon$ ). The basic idea is to put a grid with appropriate granularity on the convex hull of the given points and to use the grid points as possible Steiner points (for details see [HRW92]). Admittedly, the direct application of this method is not practical, as for a guarantee of 1% one needs a complete network with up to  $const \cdot 10^4 \cdot r^2$  vertices.

#### **Rectilinear Steiner Problem (RSP)**

**Definition 2** Given a finite set R of points in the (Euclidean) plane, find a point set S together with a minimal spanning tree T for  $R \cup S$ , such that T has minimal length concerning the  $L_1$ -norm (Manhattan/rectilinear distance).

The similarity between the RSP and the ESP is obvious, but an observation by Hanan [Han66] showed that the RSP is a special case of the NSP:

Draw horizontal and vertical lines through the given points. Define a network G = (V, E, c), with V as the set of all line crossings and E corresponding to the line segments. Hanan showed that an optimal solution in G is an optimal solution for the original RSP instance.

Although it is possible to translate every RSP instance into an NSP instance, this approach has the obvious disadvantage that one loses the knowledge of the special structure of the RSP instance. A refined scheme that uses some geometric based preprocessing before the translation to an NSP instance still loses the geometric information. Nevertheless, in combination with the NSP algorithms presented in this work, it is in many cases the fastest approach for solving an RSP instance (see Section 5.4.1).

We do not discuss the RSP in more detail in this work. For an overview of the very extensive literature for the RSP, see [HRW92, Zac01]. For recent results, see [WWZ00, PV03].

#### 1.4.3 Applications

The different variants of the NSP are among those problems in Combinatorial Optimization that have most applications. It is easy to imagine possible applications. A voluminous book [CD01] is between to Steiner tree applications. As examples, we describe some applications of the NSP from different fields:

#### **Routing in Computer Networks**

In multiple destination routing (MDR) we are given a network G = (V, E, c) with a source  $s \in V$ and a set of destinations  $D \subset V$  ( $s \notin D$ ). The cost function c can be a complicated function with arguments like delay of a channel, fees, and so on. Two well-known special cases are |D| = 1 (single destination routing) and |D| = n - 1 (broadcasting). A routing tree T is a subnetwork of G, containing all vertices of  $D \cup \{s\}$  and can be viewed as a directed tree rooted at s with leaves from D. An NC (network cost) optimal routing asks for a routing tree with minimal total cost.

We present a possible approach for NC optimal routing (for details see [BKJ83]): First, a (minimal) Steiner tree T for G with  $R = D \cup \{s\}$  is calculated. The source sends a copy of the message to each of its neighbors in T, together with information on the respective subtree. Each vertex that receives a message repeats this procedure for its subtree.

For a similar application of the NSP in directed networks, see [BD93]. For a distributed variant that uses local information, see [NK94].

#### **VLSI Layout**

Several variants of the NSP have numerous applications in VLSI layout. A simple scenario is to find a connection for a set of points on a chip that should carry the same signal.

For some (classical) problems in VLSI layout it may be more appropriate to formulate them as RSP, in many cases also additional constraints (e.g., delay) have to be satisfied or multiple disjoint trees have to be found. But in many applications a formulation as NSP is indicated, e.g., because there are obstacles on the chip (see for example Koch and Martin [KM96]).

For an overview of applications of the Steiner problem in VLSI layout see Korte, Pr'omel and Steger [KPS90] or Lengauer [Len90].

#### Phylogeny

Phylogeny is the study of the evolution of life forms. A central problem in Phylogeny is the task of reconstructing an evolutionary tree for a set of (biological) species. One typical variant is the following:

Each given species is represented by some segment of its DNA code. Each DNA sequence is identified with a sequence of m letters of a finite alphabet A (i.e., a vector from  $A^m$ ). Then we have a (complete) network G = (V, E, c) with  $V = A^m$ , where the cost function c represents the "distance" between two sequences; in the simplest case this is the Hamming distance of the vectors.

The task is now to find a (minimal) Steiner tree in G, where the set of given species corresponds to the set of terminals.

The literature in this area is very voluminous and deals with a number of very heterogeneous definitions and aims. An overview is given in [HRW92].

#### 1.4.4 Some Efficiently Solvable Special Cases

Some important special cases of the NSP can be solved very efficiently. This forms the basis for many algorithms for the NSP.

- r = n: The solution is a minimal spanning tree (MST) of G. It can be obtained in time  $O(m + n \log n)$ .
- r = 1: The solution is the given terminal.
- r = 2: The solution is a shortest path between the two given terminals. It can be obtained in time  $O(m + n \log n)$ .
- r = 3: Let  $R = \{z_1, z_2, z_3\}$ . It is obvious that every minimal Steiner tree has at most one vertex with degree three. Thus, we need to find a vertex v that minimizes  $d(v, z_1) + d(v, z_2) + d(v, z_3)$ . Let  $v^*$  be such a vertex. For the solution calculate shortest paths trees with roots  $z_1, z_2$  and  $z_3$ . This can be done in time  $O(m + n \log n)$ .

Note that for some of the tasks there are algorithms with even better running time guarantees [PR00, KKT95, Tho97], but they are considered to be impractical because of the constant factors hidden in the *O*-notation.

#### **1.5 Reformulations**

Reformulations of a problem are typically used in two cases: Either they are used for simpler proofs of properties of the original problem, or they are used for calculating lower bounds based on linear programming relaxations.

#### **1.5.1** Complete Networks

If it is desirable (e.g., for some simpler proofs), every network G can be transferred to a (with respect to the Steiner problem) equivalent, complete network  $G^*$  by introducing new edges with a weight > c(G) between each pair of non-adjacent vertices.

#### 1.5.2 Distance Networks

**Lemma 1** Let  $D := D_G(V)$  be the distance network of G. It holds that  $c(T_G(R)) = c(T_D(R))$ .

#### **Proof:**

I)  $c(T_G(R)) = c(T_{G^*}(R)) \ge c(T_D(R))$ , because no edge in D is longer than the corresponding edge in  $G^*$ .

II)  $c(T_D(R)) \ge c(T_G(R))$ , because every edge in  $T_D(R)$  can be replaced by a corresponding shortest path in G.

**Corollary 1.1** For each NSP instance there is an equivalent instance that fulfills the triangle inequality. If the preprocessing time for the calculation of all pairs shortest paths (e.g.,  $O(n^3)$ ) is acceptable, one can always assume metric graphs.

An explicit transformation of the instance is not advisable in most cases, as one loses many structural properties that could be exploited by algorithms. Furthermore, the necessary time for the calculation of all pairs shortest paths can be very large.

#### 1.5.3 Directed Networks

**Definition 3** Let  $\vec{G} = (V, A, c)$  be a directed network, and  $R \subseteq V$  a nonempty set of terminals with a special root vertex  $z_1 \in R$ . The Steiner problem in  $\vec{G}$  is to find a subnetwork  $T_{\vec{G}}(R)$  of  $\vec{G}$ , such that

1. in  $T_{\vec{C}}(R)$  there is a path from  $z_1$  to every other terminal,

2.  $c(T_{\vec{C}}(R))$  is minimal.

Every instance of the Steiner problem in an undirected network G can be translated into an instance of the Steiner problem in a directed Network  $\vec{G}$ : Replace every undirected edge  $(v_i, v_j)$ , by two directed arcs  $[v_i, v_j]$  and  $[v_j, v_i]$ , each with weight  $c(v_i, v_j)$ . The root vertex can be chosen arbitrarily from R. Each solution  $T_{\vec{G}}(R)$  yields a solution  $T_G(R)$  by replacing each directed arc by the corresponding undirected edge.

#### 1.5.4 Minimal Spanning Trees with Degree Constraints

Let G = (V, E, c) with terminals R be an NSP instance. Introduce a new vertex  $v_0$  and connect it with every vertex from  $V \setminus R$  and one arbitrary, but fixed  $z_1 \in R$  by zero edges. Let  $G_0 = (V_0, E_0, c_0)$ be the network resulting from this transformation. We consider a minimal spanning tree (MST)  $T_0 = \overline{T}_{G_0}(V_0)$  with the additional constraint that in  $T_0$  every vertex from  $V \setminus R$  that is adjacent to  $v_0$  has degree 1.

**Lemma 2** Remove  $v_0$  and all incident edges from  $T_0$ . The resulting tree  $\hat{T}$  is a minimal Steiner tree for R in G.

#### **Proof:**

I)  $c(\hat{T}) \ge c(T_G(R))$ , because  $\hat{T}$  is a tree and connects all terminals. II)  $c(\hat{T}) \le c(T_G(R))$ : Add  $v_0$  to  $T_G(R)$  and connect  $v_0$  with zero edges to  $z_1$  and all vertices  $v_i \notin$ 

 $T_G(R)$ . This is a spanning tree for  $G_0$  that satisfies the degree constraints and has cost  $c(T_G(R))$ .  $\Box$ A similar reformulation is also possible for the directed version. Let  $\vec{G} = (V, A, c)$  be a directed

A similar reformulation is also possible for the directed version. Let G = (V, A, c) be a directed network with terminals R and a root vertex  $z_1 \in R$ . We introduce an additional vertex  $v_0$  with zero arcs  $[v_0, v_i]$  (for all  $v_i \in V \setminus R$ ) and  $[v_0, z_1]$ . Let  $\vec{G}_0 = (V_0, A_0, c_0)$  denote this extended network. With a similar argument as in the last proof, it follows that finding  $T_{\vec{G}}(R)$  is essentially the same as finding a directed spanning tree (arborescence)  $\vec{T}_0$  with root  $v_0$  in  $\vec{G}_0$ , such that in  $\vec{T}_0$  every descendant of  $v_0$  has degree 1.

### **1.6 Complexity Results**

#### **1.6.1 General Propositions**

The decision variant of the Steiner problem in networks (with  $c : E \to \mathbb{N}$ ) is strongly  $\mathcal{NP}$ -complete (Karp [Kar72]). The optimization variant is  $\mathcal{NP}$ -hard.

It is still  $\mathcal{NP}$ -hard to solve the Steiner problem for many important metrics, for example:

- shortest distance in networks (direct consequence of 1.5.2),
- Euclidean distance (Garey, Graham and Johnson [GGJ77]),
- Manhattan distance (Garey and Johnson [GJ77]),
- Hamming distance (Foulds and Graham [FG82]).

#### 1.6.2 Special Networks

#### $\mathcal{NP}$ -hard cases

The Steiner problem is still  $\mathcal{NP}$ -hard for most important classes of graphs, for example:

**bipartite networks,** are between R and  $V \setminus R$  and have weight 1. The proof is a simple reduction from EXACT-COVER BY 3-SETS, see for example [HRW92].

planar networks (Garey and Johnson [GJ77]), the special case for edge weights 1 is open.

complete networks with edge weights {1,2} (Bern and Plassman [BP89]).

A detailed overview of this is given in Johnson [Joh85].

#### **Polynomially Solvable Cases**

As we deal with the general Steiner problem, considering polynomially solvable cases could seem only weakly motivated. But this is not the case. A simple scheme is provided by the previously mentioned cases r = 2, r = 3 and r = n. The very efficient algorithms for these cases are used in nearly every algorithm for the Steiner problem. Other schemes are more pretentious: It could be possible that the transformations of the original instance performed by reduction techniques yield subinstances that are polynomially solvable special cases. For most known polynomially solvable cases (trees, series parallel networks, Halin networks, ...) we found these approaches to be not helpful, but the special case of constant pathwidth could be used fruitfully, as we show in Section 5.2.

For a more comprehensive list of polynomially solvable cases, see [HRW92, Chapter 5].

#### 1.6.3 Approximability

The Steiner problem is  $\mathcal{APX}$ -complete (Bern and Plassman [BP89]), even if edge weights are only from  $\{1, 2\}$ . This means there exists  $\epsilon > 0$ , such that it is  $\mathcal{NP}$ -hard to find a  $(1 + \epsilon)$ -approximation for this problem (Arora [Aro94]).

There are a large number of approximation algorithms with constant approximation ratio. Most of the classical heuristics for generating upper bounds guarantee a ratio of close to two.

Better ratios were obtained by a series of algorithms that improved the ratio step by step. The fastest of these algorithm has a ratio of approximately 1.84 and a running time of  $O(r(m + n \log n + rn))$  [Zel93, DV97]. But the empirical results of this algorithm were only mediocre and do not justify the comparably long running times.

The most recent result is a ratio of  $1 + \ln(3)/2 \approx 1.55$  [RZ00], but the polynomial describing the running time has a k in the exponent that has to go to  $\infty$  to reach this ratio. Thus, the result is hardly of practical relevance. For a recent survey on approximation results, see [GHNP01].

At this point we mention that Arora [Aro96] (and, independently, Mitchell [Mit96]) developed a polynomial time approximation scheme for the Euclidean Steiner problem in  $\mathbb{R}^2$  (as well as for a huge number of other  $\mathcal{NP}$ -hard geometric problems). The methods used are quite general and the proofs hold for all geometric norms  $L_p$  ( $p \ge 1$ ).

Chapter 2

# Theoretical and Practical Aspects of Linear Programming Relaxations for the Steiner Problem

### 2.1 Introduction

This chapter deals with theoretical and practical aspects of linear programming approaches for the Steiner problem. The basic idea is to reformulate the given problem as an integer linear program. Then, the linear relaxation of these reformulation can be solved (or approximated). This is useful, not only for computing lower bounds, but also as the basis of many empirically successful heuristics for computing upper bounds and sophisticated reduction techniques, culminating in an exact algorithm, which achieves very good empirical results.

There are many (mixed) integer programming formulations of the Steiner problem in networks. Although the strength of the corresponding linear relaxation has great impact on the practical effectiveness of the algorithms that are based on it, not much was known about the relative quality of these relaxations. In Sections 2.3 to 2.8, we compare the linear relaxations of all classical, frequently cited and some modified or new integer programming formulations of this problem from a theoretical point of view with respect to their optimal values. We present several new results, establishing clear relations between relaxations, which have often been treated as unrelated or incomparable.

In Section 2.7, we introduce a collection of new relaxations that are stronger than any relaxation we are aware of, and place the new relaxations into our hierarchy. Further, we show how one relaxation of the collection can be used in practical algorithms. Except for the flow-balance constraints introduced to the Steiner problem by [KM98], this is the first successful attempt to use a relaxation that is stronger than Wong's directed cut relaxation (Section 2.3) from 1984 [Won84].

In Section 2.9, we report on our experimental study of some of these relaxations. Their previously mentioned algorithmic application for upper bound calculation, reduction techniques, and in the exact algorithm will be explained in detail in the corresponding chapters.

#### 2.2 Additional Definitions for Lower Bounds

Here we recall two reformulations of the Steiner problem from Section 1.5, because they are used in some relaxations. One uses the directed version: Given G = (V, E, c) and R, find a minimum weight arborescence in  $\vec{G} = (V, A, c)$  ( $A := \{[v_i, v_j], [v_j, v_i] \mid (v_i, v_j) \in E\}$ , c defined accordingly) with a terminal (say  $z_1$ ) as the root that spans  $R^{z_1} := R \setminus \{z_1\}$ .

The problem can also be stated as finding a degree-constrained minimum spanning tree  $T_0$  in a modified network  $G_0 = (V_0, E_0, c_0)$ , produced by adding a new vertex  $v_0$  and connecting it through zero cost edges to all vertices in  $V \setminus R$  and to a fixed terminal (say  $z_1$ ). The problem is now equivalent to finding a minimum spanning tree  $T_0$  in  $G_0$  with the additional restriction that in  $T_0$  every vertex in  $V \setminus R$  adjacent to  $v_0$  must have degree one. For more details on this reformulation, see [BP87, Bea89]. Again, a similar directed version for a network  $\vec{G}_0$  can be defined, this time by adding zero cost arcs  $[v_0, v_i]$  (for all  $v_i \in V \setminus R$ ) and  $[v_0, z_1]$  to  $\vec{G}$ .

We introduce some additional definitions that make the notation of linear relaxations easier to understand.

A cut in  $\vec{G} = (V, A, c)$  (or in G = (V, E, c)) is defined as a partition  $C = \{\overline{W}, W\}$  of V $(\emptyset \subset W \subset V; V = W \cup \overline{W})$ . We use  $\delta^-(W)$  to denote the set of arcs  $[v_i, v_j] \in A$  with  $v_i \in \overline{W}$  and  $v_j \in W$ . For simplicity, we write  $\delta^-(v_i)$  instead of  $\delta^-(\{v_i\})$ . The sets  $\delta^+(W)$  and, for the undirected version,  $\delta(W)$  are defined similarly. A cut  $C = \{\overline{W}, W\}$  is called a **Steiner cut** if  $z_1 \in \overline{W}$  and  $R^{z_1} \cap W \neq \emptyset$  (for the undirected version:  $R \cap W \neq \emptyset$  and  $R \cap \overline{W} \neq \emptyset$ ).

In the integer programming formulations we use (binary) variables  $x_{ij}$  for each arc  $[v_i, v_j] \in A$ (respectively  $X_{ij}$  for each edge  $(v_i, v_j) \in E$ ), indicating whether an arc is part of the solution  $(x_{ij} =$  1) or not  $(x_{ij} = 0)$ . Thus, the cost of the solution can be calculated by the dot product  $c \cdot x$ , where c is the cost vector. For any  $B \subseteq A$ , x(B) is short for  $\sum_{a \in B} x_a$ , and A(W) denotes  $\{[v_i, v_j] \in A | v_i, v_j \in W\}$  for any  $W \subseteq V$ . For example,  $x(\delta^-(W))$  is short for  $\sum_{[v_i, v_j] \in A, v_i \notin W, v_j \in W} x_{ij}$ .

Let  $P_1$  be an integer linear program. The corresponding linear relaxation is denoted by  $LP_1$ . The dual of such a relaxation is denoted by  $DLP_1$  and a Lagrangian relaxation by  $LaLP_1$ . The value of an optimal solution of the integer programming formulation (for given  $\vec{G}$  and R), denoted by  $v(P_1)$ , is of course the value of an optimal solution of the corresponding Steiner arborescence problem in  $\vec{G}$ . Thus, in this context we are interested in the optimal value  $v(LP_1)$  of the corresponding linear relaxation, which can differ from  $v(P_1)$ . The notations  $P_1$  (or  $LP_1$ ) always denote an integer (or linear) program corresponding to an arbitrary, but fixed instance (G, R) of the Steiner problem (with G replaced by  $\vec{G}$ ,  $G_0$  or  $\vec{G}_0$  when appropriate).

For a linear program  $P_x$ , the identifier x describes the program (see Table 2.1 for further explanation of the abbreviations for linear programs).

C	cut based	Section 2.3.1
F	flow based	Section 2.3.2
$F^R$	common-flow formulation	Section 2.7.3
$F^{j_1, j_2}$	restricted version of $F^R$ (multiple roots)	Section 2.7.4
$F^2$	restricted version of $F^R$ (only one root)	Section 2.7.4
2T	two-terminal formulation	Section 2.3.2
Т	tree based	Section 2.4
mT	multiple tree based	Section 2.6
$T_0$	degree constraint spanning tree based (with $v_0$ vertex)	Section 2.4.1
$\vec{T}$	directed version of $T$	
U	undirected version	
X + FB	with added flow-balance constraints	Section 2.6.2
X-	weaker version of X	
X + +	aggregated version of $X$	
X'	modified version of $X$	

Table 2.1: Abbreviations for linear programs and their meaning.

We compare relaxations using the predicates **equivalent** and (**strictly**) **stronger**: We call a relaxation  $R_1$  stronger than a relaxation  $R_2$  if the optimal value of  $R_1$  is no less than that of  $R_2$  for all instances of the problem. If  $R_2$  is also stronger than  $R_1$ , we call them equivalent, otherwise we say that  $R_1$  is strictly stronger than  $R_2$ . If neither is stronger than the other, they are **incomparable**.

### 2.3 Cut and Flow Formulations

In this section, we state the basic flow and cut-based formulations of the Steiner problem. There are some well-known observations concerning these formulations, which we cite without proof.

#### 2.3. CUT AND FLOW FORMULATIONS

#### 2.3.1 Cut Formulations

The directed cut (or dicut) formulation was stated in [Won84].

$$\frac{P_C}{x(\delta^-(W))} \geq 1 \quad (z_1 \notin W, R \cap W \neq \emptyset),$$
(1.1)

$$(W)) \geq 1 \quad (z_1 \notin W, R \cap W \neq \emptyset), \tag{1.1}$$

$$x \in \{0,1\}^{|A|}. \tag{1.2}$$

The constraints (1.1) are called Steiner cut constraints. They guarantee that in any arc set corresponding to a feasible solution, there is a path from  $z_1$  to any other terminal.

A formulation for the undirected version was stated in [Ane80]:

$$\begin{array}{cccc}
\underline{P_{UC}} & c \cdot X & \to & \min, \\
\hline X(\delta(W)) & \geq & 1 & (W \cap R \neq R, W \cap R \neq \emptyset), \\
& X & \in & \{0,1\}^{|E|}.
\end{array}$$
(2.1)
(2.2)

**Lemma 3**  $LP_C$  is strictly stronger than  $LP_{UC}$ ; and  $\sup\left\{\frac{v(LP_C)}{v(LP_{UC})}\right\} = 2$  [CR94a, Dui93].

We just mention here that  $\frac{v(P_{UC})}{v(LP_{UC})} \leq 2$  [GB93]; and that when applied to undirected instances, the value  $v(LP_C)$  is independent of the choice of the root [GM93]. For much more information on  $LP_C$ ,  $LP_{UC}$  and their relationship, see [CR94a]. Also, many related results are discussed in [MW95].

#### 2.3.2 Flow Formulations

Viewing the Steiner problem as a multicommodity flow problem leads to the following formulation (see [Won84]).

$$\frac{P_F}{v_i^t(\delta^-(v_i))} = y^t(\delta^+(v_i)) - \begin{cases} 1 & (z_t \in R^{z_1}; v_i = z_t), \\ 0 & (z_t \in R^{z_1}; v_i \in V \setminus \{z_1, z_t\}), \end{cases} (3.1)$$

$$y^t \leq x \quad (z_t \in R^{z_1}), \tag{3.2}$$

$$y^t \ge 0 \quad (z_t \in R^{z_1}),$$
 (3.3)

$$x \in \{0,1\}^{|A|}. \tag{3.4}$$

Each variable  $y_{ii}^t$  denotes the quantity of the commodity t flowing through  $[v_i, v_i]$ . Constraints (3.1) guarantee that for each terminal  $z_t \in R^{z_1}$ , there is a flow of one unit of commodity t from  $z_1$  to  $z_t$ . Together with (3.2), they guarantee that in any arc set corresponding to a feasible solution, there is a path from  $z_1$  to any other terminal.

#### **Lemma 4** $LP_C$ is equivalent to $LP_F$ [Won84].

The correspondence is even stronger: Every feasible solution x for  $LP_C$  corresponds to a feasible solution (x, y) for  $LP_F$ .

The straightforward translation of  $P_F$  for the undirected version leads to  $LP_{UF}$  with  $v(LP_{UF}) =$  $v(LP_{UC})$  (see [GM93]). There are other undirected formulations (see [GM93]), leading to relaxations that are all equivalent to  $LP_F$ ; so we use the notation  $LP_{FU}$  for all of them.

Of course, there is no need for different commodities in  $P_F$ . In an aggregated version, which we call  $P_{F++}$ , one unit of a single commodity flows from  $z_1$  to each terminal  $z_t \in \mathbb{R}^{z_1}$  (see [Mac87]). This program has only  $\Theta(a)$  variables and constraints, which is asymptotically minimal. But the corresponding linear relaxation  $LP_{F++}$  is not a strong one:

$$\frac{P_{F++}}{V(\delta^{-}(v_{i}))} = Y(\delta^{+}(v_{i})) - \begin{cases} 1 & (v_{i} \in R^{z_{1}}), \\ 0 & (v_{i} \in V \setminus R), \end{cases}$$
(4.1)

$$(r-1)x \geq Y,$$

$$Y \geq 0,$$

$$(4.2)$$

$$(4.3)$$

$$1 \ge 0,$$
 (4.3)

 $x \in \{0,1\}^{|A|}.$ (4.4)

The variables Y describe a flow of one unit from  $z_1$  to each terminal in  $\mathbb{R}^{z_1}$ .

**Lemma 5**  $LP_F$  is strictly stronger than  $LP_{F++}$ . The worst-case ratio  $\frac{v(LP_F)}{v(LP_{F++})}$  is r-1 [Mac87, Dui93].

In [Liu90], the two-terminal formulation was stated:

$$\begin{array}{cccc}
P_{2T} & c \cdot x & \to & \min, \\
\check{y}^{kl}(\delta^{-}(v_{i})) - \check{y}^{kl}(\delta^{+}(v_{i})) & \geq & \begin{cases} -1 & (\{z_{k}, z_{l}\} \subseteq R^{z_{1}}; v_{i} = z_{1}), \\
0 & (\{z_{k}, z_{l}\} \subseteq R^{z_{1}}; v_{i} \in V \setminus \{z_{1}\}), \end{cases} (5.1) \\
\begin{pmatrix} (\delta^{-}(v_{i})) & (\check{z}^{kl} + \check{z}^{kl})(\delta^{+}(v_{i})) & \int 1 & (\{z_{k}, z_{l}\} \subseteq R^{z_{1}}; v_{i} = z_{k}), \\
\end{pmatrix} (5.2)$$

$$(\check{y}^{kl} + \check{y}^{kl})(\delta^{-}(v_i)) - (\check{y}^{kl} + \check{y}^{kl})(\delta^{+}(v_i)) = \begin{cases} 1 & (\{z_k, z_l\} \subseteq R^{z_1}; v_i = z_k), \\ 0 & (\{z_k, z_l\} \subseteq R^{z_1}; v_i \in V \setminus \{z_1, z_k\}), \end{cases}$$
(5.2)

$$(\check{y}^{kl} + \acute{y}^{kl})(\delta^{-}(v_i)) - (\check{y}^{kl} + \acute{y}^{kl})(\delta^{+}(v_i)) = \begin{cases} 1 & (\{z_k, z_l\} \subseteq R^{z_1} ; v_i = z_l), \\ 0 & (\{z_k, z_l\} \subseteq R^{z_1} ; v_i \in V \setminus \{z_1, z_l\}), \end{cases}$$
(5.3)

$$\check{y}^{kl} + \check{y}^{kl} + \check{y}^{kl} \leq x \quad (\{z_k, z_l\} \subseteq R^{z_1}),$$
(5.4)

$$x \in \{0,1\}^{|A|}.$$
(5.6)

The formulation  $P_F$  is based on the flow formulation of the shortest path problem (the special case of the Steiner problem with  $|R^{z_1}| = 1$ ). The formulation  $P_{2T}$  is based on the special case with  $|R^{z_1}| = 2$ , namely the two-terminal (2T) Steiner arborescence problem. In a Steiner tree, for any two terminals  $z_k, z_l \in R^{z_1}$ , there is a two-terminal tree consisting of a path from  $z_1$  to a splitter node  $v_s$  and two paths from  $v_s$  to  $z_k$  and  $z_l$  ( $v_s$  can belong to  $\{z_1, z_k, z_l\}$ ). In  $P_{2T}$ ,  $\check{y}$ ,  $\check{y}$  and  $\check{y}$  describe flows from  $z_1$ to  $v_s$ , from  $v_s$  to  $z_k$  and from  $v_s$  to  $z_l$ . Note that the flow described by  $\check{y}$  can have an excess at some vertices (because of the inequality in (5.1)), this excess is carried by the flows described by  $\hat{y}$  and  $\hat{y}$  to  $z_k$  and  $z_l$  (because of (5.2) and (5.3)).

**Lemma 6**  $LP_{2T}$  is strictly stronger than  $LP_F$  [Liu90].

 $\check{y}'$ 

#### 2.4**Tree Formulations**

In this section, we state the basic tree-based formulations and prove that the corresponding linear relaxations are all equivalent. We also discuss some variants from the literature, which we prove to be weaker.

#### 2.4.1 Degree-Constrained Tree Formulations

In [Bea89], the following program was suggested, which is a translation of the degree-constrained minimum spanning tree problem in  $G_0$ .

$$\begin{array}{cccc} \hline P_{T_0} & c \cdot X & \to & \min, \\ \{(v_i, v_j) \mid X_{ij} = 1\} & : & \text{builds a spanning tree for } G_0, \\ & X_{0k} + X_{ki} & \leq & 1 & (v_k \in V \setminus R \ ; \ (v_k, v_i) \in \delta(v_k)), \end{array}$$

$$(6.2)$$

$$X \in \{0,1\}^{|E_0|}. \tag{6.3}$$

The requirement (6.1) can be stated by linear constraints. In the following, we assume that (6.1) is replaced by the following constraints.

$$X(E_0) = n, (6.4)$$

$$X(E_0(W)) \leq |W| - 1 \quad (\emptyset \neq W \subset V_0). \tag{6.5}$$

The constraints (6.4) and (6.5), together with the nonnegativity of X, define a polyhedron whose extreme points are the incidence vectors of spanning trees in  $G_0$  (see [Edm71, MW95]). Thus, no other set of linear constraints replacing (6.1) can lead to a stronger linear relaxation.

A directed version can be stated as follows.

$$\begin{array}{cccc}
P_{\vec{T}_0} & c \cdot x & \to & \min, \\
& & x(\delta^-(v_i)) &= 1 \quad (v_i \in V), \\
\end{array}$$
(7.1)

$$x(A_0(W)) \leq |W| - 1 \quad (\emptyset \neq W \subseteq V_0), \tag{7.2}$$

$$x_{0i} + x_{ij} + x_{ji} \leq 1 \quad (v_i \in V \setminus R; [v_i, v_j] \in \delta^+(v_i)),$$
 (7.3)

$$x \in \{0,1\}^{|A_0|}. \tag{7.4}$$

Again, the constraints (7.1) and (7.2), together with the nonnegativity of x, define a polyhedron whose extreme points are the incidence vectors of spanning arborescences with root  $v_0$  (see [MW95]). Note that  $\delta^-(v_0) = \emptyset$  by the construction of  $\vec{G}_0$ .

In the literature on the Steiner problem, one usually finds a directed variant  $P_{\vec{T}_0-}$  that uses

$$x_{0i} + x_{ij} \leq 1$$
  $(v_i \in V \setminus R; [v_i, v_j] \in \delta^+(v_i))$ 

instead of the constraints (7.3) (see for example [HRW92]). Obviously  $v(P_{\vec{T}_0-}) = v(P_{\vec{T}_0})$ , and  $v(LP_{\vec{T}_0-}) \leq v(LP_{\vec{T}_0})$ . The following example shows that  $LP_{\vec{T}_0}$  is strictly stronger than the version in the literature.

**Example 1** Figure 2.1 shows the network  $\vec{G}$  with  $R = \{z_1, z_2\}, \gamma \ge 100$  and the network  $\vec{G}_0$ . The minimum Steiner arborescence has the value  $\gamma + 10$ .

The following  $\dot{x}$  is feasible (and optimal) for  $LP_{\vec{T}_0-}$  and gives the value 11:  $\dot{x}_{01} = 1$ ,  $\dot{x}_{03} = \dot{x}_{04} = \dot{x}_{34} = \dot{x}_{43} = \dot{x}_{32} = \dot{x}_{42} = \frac{1}{2}$  and  $\dot{x}_{ij} = 0$  (for all other arcs). But for  $LP_{\vec{T}_0}$ ,  $\dot{x}$  is infeasible. The optimal value here is:  $v(LP_{\vec{T}_0}) = \frac{\gamma}{3} + 14$  (this value is reached for example by  $\hat{x}$  with  $\hat{x}_{01} = 1$ ,  $\hat{x}_{03} = \dot{x}_{04} = \hat{x}_{13} = \hat{x}_{23} = \hat{x}_{32} = \frac{1}{3}$ ,  $\hat{x}_{42} = \hat{x}_{34} = \frac{2}{3}$  and  $\hat{x}_{ij} = 0$  (for all other arcs)). So the ratio  $v(LP_{\vec{T}_0-})/v(LP_{\vec{T}_0})$  can be arbitrarily close to 0.



Figure 2.1: Example with  $v(LP_{\vec{T}_0-}) \ll v(LP_{\vec{T}_0}) = v(LP_{T_0}) \ll v(P_{T_0})$ .

#### 2.4.2 Rooted Tree Formulation

The rooted tree formulation is stated, for example, in [KPH93]:

$$P_{\vec{T}}$$
  $c \cdot x \rightarrow \min,$ 

$$x(\delta^{-}(v_i)) = 1 \quad (v_i \in R^{z_1}),$$
 (8.1)

$$x(\delta^{-}(v_i) \setminus \{[v_j, v_i]\}) \geq x_{ij} \quad (v_i \in V \setminus R ; [v_i, v_j] \in \delta^+(v_i)),$$

$$(8.2)$$

$$x(A(W)) \leq |W| - 1 \quad (\emptyset \neq W \subseteq V), \tag{8.3}$$

$$x \in \{0,1\}^{|A|}. \tag{8.4}$$

To get rid of the exponential number of constraints for avoiding cycles, many authors have considered replacing (8.3) by the subtour elimination constraints introduced in the TSP-context (known as the Miller-Tucker-Zemlin constraints [MTZ60]), allowing additional variables  $t_i$  for all  $v_i \in V$ :

$$t_i - t_j + nx_{ij} \le n - 1$$
 ([ $v_i, v_j$ ]  $\in A$ ). (8.5)

This leads to the program  $P_{\vec{T}-}$  with  $\Theta(a)$  variables and constraints, which is asymptotically minimal. The linear relaxation  $LP_{\vec{T}-}$  was used by [KP95]. We will now prove the intuitive guess that  $LP_{\vec{T}}$  is stronger than  $LP_{\vec{T}-}$ . Indeed, the ratio  $\frac{v(LP_{\vec{T}-})}{v(LP_{\vec{T}})}$  can be arbitrarily close to 0 (see Figure 2.2 on page 30).

Lemma 7  $v(LP_{\vec{T}-}) \leq v(LP_{\vec{T}}).$ 

**Proof:** Let  $\hat{x}$  denote an (optimal) solution for  $LP_{\vec{T}}$ . Obviously  $\hat{x}$  satisfies the constraints (8.1) and (8.2). We now show that it is possible to construct  $\hat{t}$  such that  $(\hat{x}, \hat{t})$  satisfies (8.5), too.

We start with an arbitrary  $\hat{t}$  (e.g.,  $\hat{t}_i = 0$  (for all  $v_i \in V$ )). We define for every arc  $[v_i, v_j] \in A$ :  $s_{ij} := (n-1) - (\hat{t}_i - \hat{t}_j + n\hat{x}_{ij})$ ; and call an arc  $[v_i, v_j]$  good, if  $s_{ij} \ge 0$ ; used, if  $s_{ij} \le 0$ ; and bad, if  $s_{ij} < 0$ . Suppose  $[v_i, v_j]$  is a bad arc (if no bad arcs exist,  $(\hat{x}, \hat{t})$  satisfies (8.5)).

We now show how  $\hat{t}_j$  (and perhaps some other  $\hat{t}_p$ ) can be increased in a way that  $[v_i, v_j]$  becomes good, but no good arc becomes bad. By repeating this procedure we can make all arcs good and prove the lemma.

In each step we denote by  $W_j$  the set of vertices  $v_k \in V$  that can be reached from  $v_j$  through paths with only used arcs. We define  $\Delta$  as  $\min\{s_{kl} \mid [v_k, v_l] \in \delta^+(W_j)\}$ , if this set is nonempty, and  $\infty$  otherwise. Now we increase for all vertices  $v_p \in W_j$  the variables  $\hat{t}_p$  by  $\min\{-s_{ij}, \Delta\}$  (these values can change in every step). By doing this, no arc of  $\delta^+(W_j)$  becomes bad. For arcs  $[v_p, v_q]$  with  $v_p, v_q \in W_j$  or  $v_p, v_q \notin W_j$  the value of  $s_{pq}$  does not change; and for arcs  $[v_q, v_p] \in \delta^-(W_j) s_{qp}$  does not decrease.

Because  $\hat{t}_j$  is increased in every step, there is only one situation that could prevent that  $[v_i, v_j]$  becomes good: In one step  $v_i$  is absorbed by  $W_j$ . But then, according to the definition of  $W_j$ , there exists a path  $v_j \rightsquigarrow v_i$  with only used arcs. Thus, there exists a cycle  $C := (v_i, v_j = v_{k_1}, \dots, v_{k_l} = v_i)$ , with  $s_{k_lk_1} < 0$  and  $s_{k_{t-1}k_t} \leq 0$  (for all  $t \in \{2, \dots, l\}$ ). Summation of the inequalities for arcs on the cycle C leads to:  $n\hat{x}(C) > l(n-1)$ . On the other hand, since  $\hat{x}$  satisfies the constraints (8.3),  $\hat{x}(C) \leq l-1$ . The consequence,  $\frac{l-1}{l} > \frac{n-1}{n}$ , is a contradiction.

#### 2.4.3 Equivalence of Tree-Class Relaxations

We now show the equivalence of the tree-based relaxations  $LP_{T_0}, LP_{\vec{T}_0}$ , and  $LP_{\vec{T}}$ .

**Lemma 8**  $v(LP_{\vec{T}_0}) = v(LP_{T_0}).$ 

#### **Proof:**

I)  $v(LP_{\vec{T}_0}) \ge v(LP_{T_0})$ : Let x denote an (optimal) solution for  $LP_{\vec{T}_0}$ . Define X with  $X_{ij} := x_{ij} + x_{ji}$ (for all  $(v_i, v_j) \in E$ ),  $X_{0i} := x_{0i}$  (for all  $v_i \in V \setminus R$ ) and  $X_{01} := x_{01}$ . It is easy to check that X satisfies all constraints of  $LP_{T_0}$  and yields the same value as  $v(LP_{\vec{T}_0})$ .

II)  $v(LP_{T_0}) \ge v(LP_{\vec{T_0}})$ : Now let X denote an (optimal) solution for  $LP_{T_0}$ . Define  $\Delta$  with  $\Delta_{ij} \in [0, 1]$ arbitrarily (for all  $(v_i, v_j) \in E$ ) and set x to  $x_{ij} := \Delta_{ij}X_{ij}, x_{ji} := (1 - \Delta_{ij})X_{ij}$  (for all  $(v_i, v_j) \in E$ ),  $x_{0i} := X_{0i}$  (for all  $v_i \in V \setminus R$ ) and  $x_{01} := X_{01}$ . Again, it is easy to validate that x satisfies the constraints (7.2) and (7.3) and yields the same value as  $v(LP_{T_0})$ .

The only question is, whether there is a  $\Delta$  such that x satisfies the constraints (7.1), too. This question can be stated in the following way:

Is it possible to distribute the "supply"  $X_{ij}$  of each edge  $(v_i, v_j)$  in such a way to its end-vertices that every vertex  $v_i \in V$  gets one unit at the end?

It is known that this problem can be viewed as a flow problem: Construct a flow network with source s, sink t, and vertices  $u_{ij}$  (for all  $(v_i, v_j) \in E_0$ ) and  $u_i$  (for all  $v_i \in V_0$ ). Every  $u_{ij}$  is connected with  $u_i$  and  $u_j$  through arcs  $[u_{ij}, u_i]$  and  $[u_{ij}, u_j]$  with capacity  $\infty$ . Furthermore, there are arcs  $[s, u_{ij}]$  with capacity  $X_{ij}$  and arcs  $[u_i, t]$  with capacity 1 (or 0, if i = 0). The question above is equivalent to the question, whether a flow from s to t with value n can be constructed. The max-flow min-cut theorem says that this is possible if and only if there is no cut  $C = \{U, \overline{U}\}$  (with  $s \in U$  and  $t \notin U$ ) with capacity less than n (Obviously  $U = \{s\}$  and  $U = V \setminus \{t\}$  correspond to cuts with capacity n).

Suppose that U corresponds to a cut C with minimum capacity. Define  $W := \{v_i \in V_0 \mid u_i \in U\}$ ,  $E_W := \{(v_i, v_j) \in E_0 \mid v_i, v_j \in W\}$ , and  $E_U := \{(v_i, v_j) \in E_0 \mid u_{ij} \in U\}$ . For every  $[v_i, v_j] \in E_U$   $(u_{ij} \in U)$ ,  $u_i$  and  $u_j$  must belong to  $U([v_i, v_j] \in E_W)$ , because otherwise the capacity of C would be  $\infty$ , which is not minimal. It follows that:  $E_U \subseteq E_W$ .

The capacity of C is:

$$|W \setminus \{v_0\}| + X(E_0 \setminus E_U) \geq |W \setminus \{v_0\}| + X(E_0 \setminus E_W) \quad (\text{since } E_U \subseteq E_W)$$
  
$$\geq |W| - 1 + X(E_0) - X(E_W)$$
  
$$= |W| - 1 + n - X(E_W) \quad (\text{because of } 6.4)$$
  
$$\geq n. \quad (\text{because of } 6.5)$$

It follows that the minimal cut has capacity of n.

Lemma 9  $v(LP_{\vec{T}}) = v(LP_{\vec{T}_0}).$ 

**Proof:** 

I)  $v(LP_{\vec{T}_0}) \geq v(LP_{\vec{T}})$ : Let  $\hat{x}$  denote an (optimal) solution for  $LP_{\vec{T}_0}$ . Define  $\tilde{x}$  with  $\tilde{x}_{ij} := \hat{x}_{ij}$  (for all  $[v_i, v_j] \in A$ ). Because  $\hat{x}$  satisfies the constraints (7.1) and in  $\vec{G}_0$  only arcs in A are incident with terminals in  $R^{z_1}$ ,  $\tilde{x}$  satisfies the constraints (8.1).

Furthermore,  $\tilde{x}$  satisfies the constraints (8.2), because for every arc  $[v_i, v_j] \in A$  with  $v_i \in V \setminus R$  it holds that:

$$\begin{split} \tilde{x}(\delta^{-}(v_i) \setminus \{[v_j, v_i]\}) &= \tilde{x}(\delta^{-}(v_i)) - \tilde{x}_{ji} & (\delta \text{ in } \vec{G}) \\ &= \hat{x}(\delta^{-}(v_i)) - \hat{x}_{0i} - \hat{x}_{ji} & (\delta \text{ in } \vec{G}_0) \\ &= 1 - \hat{x}_{0i} - \hat{x}_{ji} & (because \text{ of } (7.1)) \\ &\geq \hat{x}_{ij} & (because \text{ of } (7.3)) \\ &= \tilde{x}_{ij}. \end{split}$$

Finally  $\tilde{x}$  satisfies (8.3), because  $\hat{x}$  satisfies (7.2).

II)  $v(LP_{\vec{T}}) \geq v(LP_{\vec{T}_0})$ : Let  $\tilde{x}$  denote an (optimal) solution for  $LP_{\vec{T}}$ . Define  $\hat{x}$  with  $\hat{x}_{ij} := \tilde{x}_{ij}$  (for all  $[v_i, v_j] \in A$ ) and  $\hat{x}_{0i} := 1 - \tilde{x}(\delta^-(v_i))$  (for all  $v_i \in V \setminus R^{z_1}$ ). Notice that for an optimal  $\tilde{x}$ ,  $\tilde{x}(\delta^-(v_i)) > 1$  could only be forced by (8.2) for some arc  $[v_i, v_l]$  with  $\tilde{x}(\delta^-(v_i) \setminus \{[v_j, v_i]\}) = \tilde{x}_{il}$ , and it would follow that  $1 < \tilde{x}(\delta^-(v_i)) = \tilde{x}_{li} + \tilde{x}_{il}$ , but this is excluded by (8.3) (for  $W = \{v_i, v_l\}$ ). So  $\hat{x}$  satisfies (7.1) in a trivial way.

The constraints (7.2) are satisfied by  $\hat{x}$  for every  $W \subseteq V$ , because  $\tilde{x}$  satisfies (8.3). For  $W \subseteq V_0$  with  $v_0 \in W$  it holds that:

$$\begin{aligned} \hat{x}(A_0(W)) &\leq \sum_{v_i \in W \setminus \{v_0\}} \hat{x}(\delta^-(v_i)) & \text{ (in } \vec{G}_0) \\ &= \sum_{v_i \in W \setminus \{v_0\}} 1 & \text{ (because of (7.1) )} \\ &= |W| - 1. \end{aligned}$$

Finally for every  $[v_i, v_j] \in A$  with  $v_i \in V \setminus R$ :

$$\hat{x}_{0i} + \hat{x}_{ij} + \hat{x}_{ji} = 1 - \tilde{x}(\delta^{-}(v_i)) + \tilde{x}_{ij} + \tilde{x}_{ji} \quad (\text{in } G) \\
= 1 - \tilde{x}(\delta^{-}(v_i) \setminus \{[v_j, v_i]\}) + \tilde{x}_{ij} \\
\leq 1. \quad (\text{because of } (8.2))$$

Thus,  $\hat{x}$  also satisfies the constraints (7.3).

#### 2.5 Relationship between the Two Classes

In this section, we settle the question of the relationship between flow and tree-based relaxations by proving that  $LP_C$  is strictly stronger than  $LP_{\vec{T}}$ . Our proofs also show that  $LP_C$  cannot be strengthened by adding constraints that are present in  $LP_{\vec{T}_0}$  or  $LP_{\vec{T}}$ . First, we show that every (optimal) solution  $\hat{x}$  of  $LP_C$  has certain properties:

**Lemma 10** For every (optimal) solution  $\hat{x}$  of  $LP_C$ ,  $W \subseteq V \setminus \{z_1\}$  and  $v_k \in W$  the following holds:

$$\hat{x}(\delta^{-}(W)) \ge \hat{x}(\delta^{-}(v_k)).$$

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#### 2.5. RELATIONSHIP BETWEEN THE TWO CLASSES

**Proof:** Suppose that  $\hat{x}$  violates the inequality for some W and  $v_k$ . Among all such inequalities, choose one for which |W| is minimal. For this inequality to be violated, there must be an arc  $[v_l, v_k] \in \delta^-(v_k) \setminus \delta^-(W)$  with  $\hat{x}_{lk} > 0$ . Because of the optimality of  $\hat{x}, \hat{x}_{lk}$  cannot be decreased without violating a Steiner cut constraint, so there is a  $U \subset V$  with  $z_1 \notin U, U \cap R \neq \emptyset, [v_l, v_k] \in \delta^-(U)$ , and  $\hat{x}(\delta^-(U)) = 1$ . Now one has the inequality  $\dagger$ :

$$\begin{aligned} \hat{x}(\delta^{-}(U)) + \hat{x}(\delta^{-}(W)) &= \hat{x}(\delta^{-}(U \cup W)) + \hat{x}(\delta^{-}(U \cap W)) + \\ \hat{x}(\{[v_i, v_j] \in A \mid v_i \in W \backslash U, v_j \in U \backslash W) + \\ \hat{x}(\{[v_j, v_i] \in A \mid v_i \in W \backslash U, v_j \in U \backslash W) \\ &\geq \hat{x}(\delta^{-}(U \cup W)) + \hat{x}(\delta^{-}(U \cap W)) \end{aligned}$$

Since  $z_1 \notin U \cup W$  and  $(U \cup W) \cap R \neq \emptyset$ ,  $U \cup W$  corresponds to a Steiner cut, and  $\hat{x}(\delta^-(U \cup W)) \ge 1 = \hat{x}(\delta^-(U))$ . Using  $\dagger$ , one obtains:  $\hat{x}(\delta^-(W)) \ge \hat{x}(\delta^-(U \cap W))$ . This implies that  $\hat{x}$  also violates the lemma for  $U \cap W$  and  $v_k$ . Since  $v_l \in W \setminus U$ , we have  $|U \cap W| < |W|$ , and this contradicts the minimality of W.<sup>1</sup>

**Lemma 11** For every (optimal) solution  $\hat{x}$  of  $LP_C$  and  $v_k \in V \setminus \{z_1\}$  the following holds:

$$\hat{x}(\delta^-(v_k)) \le 1.$$

**Proof:** Suppose  $\hat{x}$  violates the inequality for  $v_k$ . There is an arc  $[v_l, v_k] \in \delta^-(v_k)$  with  $\hat{x}_{lk} > 0$ . Because of the optimality of  $\hat{x}$ ,  $\hat{x}_{lk}$  cannot be decreased without violating a Steiner cut constraint, so there is a  $W \subset V$  with  $z_1 \notin W$ ,  $W \cap R \neq \emptyset$ ,  $[v_l, v_k] \in \delta^-(W)$ , and  $\hat{x}(\delta^-(W)) = 1$ . Together with Lemma 10 (for  $v_k$  and W), one gets a contradiction.

**Lemma 12** For every (optimal) solution  $\hat{x}$  of  $LP_C$ ,  $v_l \in V \setminus \{z_1\}$ , and  $[v_l, v_k] \in A$  the following holds:

$$\hat{x}(\delta^{-}(v_l) \setminus \{[v_k, v_l]\}) \ge \hat{x}_{lk}.$$

**Proof:** This follows directly from Lemma 10 (for  $v_k$  and  $W = \{v_l, v_k\}$ ) by subtracting  $\hat{x}(\delta^-(v_l) \setminus \{[v_k, v_l]\})$  from both sides. Note that the special case  $v_k = z_1$  is trivial, because  $\hat{x}_{l1} = 0$  in every optimal solution.

**Theorem 13**  $v(LP_{\vec{T}}) \leq v(LP_C)$ .

**Proof:** Let  $\hat{x}$  be an (optimal) solution for  $LP_C$ . We will show that  $\hat{x}$  is feasible for  $LP_{\vec{T}}$ : Because  $\{v_i\}$  corresponds to a Steiner cut for  $v_i \in R^{z_1}$ , through the use of Lemma 11,  $\hat{x}$  satisfies (8.1).

Because of Lemma 12,  $\hat{x}$  satisfies (8.2).

Let  $W \subseteq V$  be a nonempty set. If  $z_1 \in W$ :

$$\begin{split} \hat{x}(A(W)) &\leq \sum_{v_i \in W} \hat{x}(\delta^-(v_i)) \\ &= \sum_{v_i \in W \setminus \{z_1\}} \hat{x}(\delta^-(v_i)) \qquad \text{(optimality of } \hat{x}) \\ &\leq \sum_{v_i \in W \setminus \{z_1\}} 1 \qquad \text{(Lemma 11)} \\ &= |W| - 1. \end{split}$$

<sup>1</sup>In a different context this argumentation was used in [GM93].

Now we assume  $z_1 \notin W$  and define  $\Delta := \hat{x}(\delta^-(W))$ . There are two cases: I)  $\Delta \ge 1$ :

$$\begin{split} \hat{x}(A(W)) &= \sum_{v_i \in W} \hat{x}(\delta^-(v_i)) - \hat{x}(\delta^-(W)) \\ &\leq \sum_{v_i \in W} \hat{x}(\delta^-(v_i)) - 1 & (\Delta \ge 1) \\ &\leq \sum_{v_i \in W} 1 - 1 & (\text{Lemma 11}) \\ &= |W| - 1. \end{split}$$

II)  $\Delta < 1$  :

$$\begin{split} \hat{x}(A(W)) &= \sum_{v_i \in W} \hat{x}(\delta^-(v_i)) - \hat{x}(\delta^-(W)) \\ &\leq \sum_{v_i \in W} \hat{x}(\delta^-(W)) - \hat{x}(\delta^-(W)) \qquad \text{(Lemma 10)} \\ &= (|W| - 1)\hat{x}(\delta^-(W)) \\ &< |W| - 1. \qquad (\Delta < 1) \end{split}$$

It follows that  $\hat{x}$  satisfies (8.3) too.

**Corollary 13.1** The proof shows that adding constraints of  $LP_{\vec{T}}$  to  $LP_C$  cannot improve  $v(LP_C)$ .

**Corollary 13.2** Because the proofs of the equivalence of the tree relaxations require the optimality only in one step of Lemma 9 to show that  $\tilde{x}(\delta^-(v_i)) \leq 1$ , which is forced by Lemma 11 for each (optimal) solution of  $LP_C$ , adding constraints of  $LP_{\vec{T}_0}$  to  $LP_C$  cannot improve  $v(LP_C)$  either.

To show that  $LP_F$  and  $LP_C$  are strictly stronger than the tree-based relaxations  $LP_{T_0}, LP_{\vec{T}_0}$ , and  $LP_{\vec{T}}$ , it is sufficient to give the following example.



Figure 2.2: Example for  $v(LP_{\vec{T}-}) \ll v(LP_{\vec{T}}) \ll v(LP_F) = v(P_F)$ .

**Example 2** For the network G (or in the directed view  $\vec{G}$ ) in Figure 2.2 set  $\alpha \gg 1$  and  $\gamma \gg \alpha$ . Obviously,  $v(P_F) = v(LP_F) = \gamma$ . For  $LP_{\vec{T}}$  is  $\hat{x}$  with  $\hat{x}_{23} = \hat{x}_{34} = \hat{x}_{42} = \frac{2}{3}$ ,  $\hat{x}_{25} = \hat{x}_{56} = \hat{x}_{62} = \frac{1}{3}$ , and  $\hat{x}_{ij} = 0$  (otherwise) feasible, even optimal, and gives the value  $v(LP_{\vec{T}}) = \alpha + 2$ . Thus, there is

no positive lower bound for the ratio  $\frac{v(LP_{\vec{T}})}{v(LP_F)}$ .

With respect to  $LP_{\vec{T}-}$  and  $LP_{\vec{T}-}$  one observes that  $(\dot{x}, \dot{t})$  with  $\dot{t}_i = 0$  (for all  $v_i \in V$ ),  $\dot{x}_{23} = \dot{x}_{32} =$  $\dot{x}_{34} = \dot{x}_{43} = \dot{x}_{24} = \dot{x}_{42} = \frac{1}{2}$ , and  $\dot{x}_{ij} = 0$  (otherwise) is an (optimal) solution for  $LP_{\vec{T}-}$  with the value 3. So, there is no positive lower bound for the ratio  $\frac{v(LP_{\vec{T}-})}{v(LP_{\vec{\tau}})}$ .

#### 2.6 Multiple Trees and the Relation to the Flow Model

In this section, we consider a relaxation based on multiple trees and prove its equivalence to an augmented flow relaxation. We also discuss some variants of the former relaxation.

#### 2.6.1 **Multiple Trees Formulation**

In [KPH93], a variant of  $P_{\vec{T}}$  was stated, using the idea that an undirected Steiner tree can be viewed as |R| different Steiner arborescences with different roots.

$$\mathcal{K}(\delta(v_i)) \geq 1 \qquad (v_i \in R), \tag{9.1}$$

$$(v_i)) \geq 2s_i \qquad (v_i \in V \setminus R), \tag{9.2}$$

$$s_i \geq X_{ij} \qquad (v_i \in V \setminus R ; (v_i, v_j) \in \delta(v_i)), \tag{9.3}$$

$$x + x_{ji}^{\kappa} = X_{ij} \qquad (v_k \in R \; ; \; (v_i, v_j) \in E),$$
(9.4)

$$x^{k}(\delta^{-}(v_{i})) = \begin{cases} 1 & (v_{k} \in R ; v_{i} \in R \setminus \{v_{k}\}), \\ 0 & (v_{k} \in R ; v_{i} = v_{k}), \end{cases}$$
(9.5)

$$x^{k}(\delta^{-}(v_{i})) = s_{i} \qquad (v_{k} \in R \; ; \; v_{i} \in V \setminus R), \tag{9.6}$$

$$\{[v_i, v_j] \mid x_{ij}^k = 1\} \quad : \quad \text{contains no cycles} \quad (v_k \in R), \tag{9.7}$$

$$X \in \{0,1\}^{|E|}, \tag{9.8}$$

$$x^{k} \in \{0,1\}^{|A|} \quad (v_{k} \in R),$$
(9.9)

$$s_i \in \{0,1\} \qquad (v_i \in V \setminus R). \tag{9.10}$$

In any feasible solution for  $P_{m\vec{T}}$ , each group of variables  $x^k$  describes an arborescence (with root  $z_k$ ) spanning all terminals. The variables s describe the set of the other vertices used by these arborescences.

We will relate this formulation to the flow formulations. First, we have to present an improvement of  $LP_F$ .

#### 2.6.2 Flow-Balance Constraints and an Augmented Flow Formulation

There is a group of constraints (see for example [KM98]) that can be used to make  $LP_F$  stronger. We call them flow-balance constraints:

$$x(\delta^{-}(v_i)) \le x(\delta^{+}(v_i)) \qquad (v_i \in V \setminus R).$$
(10.1)

We denote the linear program that consists of  $LP_F$  and (10.1) by  $LP_{F+FB}$ . It is obvious that  $LP_{F+FB}$ is stronger than  $LP_F$ . The following example shows that it is even strictly stronger.



Figure 2.3: Example with  $v(LP_F) < v(LP_{F+FB}) = v(P_{F+FB})$ .

**Example 3** The network  $\vec{G}$  in Figure 2.3 with  $z_1$  as the root and  $R^{z_1} = \{z_2, z_3\}$  gives an example for  $v(LP_F) < v(LP_{F+FB})$ :  $v(P_{F+FB}) = v(LP_{F+FB}) = 6$ ,  $v(LP_F) = 5\frac{1}{2}$ .

Now consider the following formulation:

$$\begin{array}{cccc} \hline P_{F'+FB} & c \cdot X & \to & \min, \\ & & x_{ij} + x_{ji} & = & X_{ij} & & ((v_i, v_j) \in E), \end{array}$$
(11.1)

$$(x,y)$$
 : is feasible for  $P_{F+FB}$ . (11.2)

**Lemma 14** If (X, x, y) is an (optimal) solution for  $LP_{F'+FB}$  with root terminal  $z_a$ , then there exists an (optimal) solution  $(X, \check{x}, \check{y})$  for  $LP_{F'+FB}$  for any other root terminal  $z_b \in R \setminus \{z_a\}$ .

**Proof:** One can verify that  $(X, \check{x}, \check{y})$  with  $\check{x}_{ij} := x_{ij} + y_{ji}^b - y_{ij}^b$ ,  $\check{y}_{ij}^t := \max\{0, y_{ij}^t - y_{ij}^b\} + \max\{0, y_{ji}^b - y_{ji}^t\}, \check{y}_{ij}^a := y_{ji}^b$  (for all  $[v_i, v_j] \in A$ ,  $z_t \in R \setminus \{z_a, z_b\}$ ) satisfies (11.1) and (3.2). Because of  $\sum_{[v_j, v_i] \in \delta^-(v_i)} (\check{y}_{ji}^t - \check{y}_{ij}^t) = \sum_{[v_j, v_i] \in \delta^-(v_i)} (\max\{0, y_{ji}^t - y_{ji}^b\} + \max\{0, y_{ij}^b - y_{ij}^t\} + \min\{0, -y_{ij}^t + y_{ij}^b\}) = \sum_{[v_j, v_i] \in \delta^-(v_i)} y_{ji}^t - y_{ji}^b + y_{ij}^b - y_{ij}^t$  (for all  $v_i \in V$ ,  $z_t \in R \setminus \{z_a, z_b\}$ ) the constraints (3.1) are satisfied, too. From (3.1) for  $y^b$ , it follows that  $x(\delta^-(v_i)) = \check{x}(\delta^-(v_i))$  and  $x(\delta^+(v_i)) = \check{x}(\delta^+(v_i))$  for all  $v_i \in V \setminus R$ ; therefore  $\check{x}$  satisfies the flow-balance constraints (10.1). Because this translation could also be performed from any (optimal) solution with root terminal  $z_b$  to

a feasible solution with root terminal  $z_a$ , the value  $v(LP_{F'+FB})$  is independent of the choice of the root terminal and  $(X, \check{x}, \check{y})$  is an (optimal) solution.

It follows immediately that  $LP_{F'+FB}$  is equivalent to  $LP_{F+FB}$ .

#### 2.6.3 Relationship between the two Models

We will now show that the linear relaxation  $LP_{m\vec{T}}$  (where (9.7) is replaced by linear constraints of the form (8.3)) is equivalent to  $LP_{F+FB}$ .

**Lemma 15**  $v(LP_{m\vec{T}}) = v(LP_{F'+FB}).$ 

#### **Proof:**

I)  $v(LP_{m\vec{T}}) \geq v(LP_{F'+FB})$ : Let  $(\hat{X}, \hat{x}, \hat{s})$  denote an (optimal) solution for  $LP_{m\vec{T}}$ . Define x with  $x := \hat{x}^1$ , and y with  $y^t := \max\{\hat{x}^1 - \hat{x}^t, 0\}$  (for all  $z_t \in R^{z_1}$ ). Because of (9.4) and the definition of  $y, y_{ij}^t = 0$  if  $y_{ji}^t > 0$  (for all  $[v_i, v_j] \in A$  and  $z_t \in R^{z_1}$ ).

#### 2.7. A COLLECTION OF NEW FORMULATIONS

For all  $z_t \in R^{z_1}, v_i \in V \setminus \{z_1, z_t\}$  it holds that:

$$\begin{split} y^t(\delta^-(v_i)) - y^t(\delta^+(v_i)) &= (\hat{x}^1 - \hat{x}^t)(\{[v_j, v_i] \in \delta^-(v_i) | \hat{x}_{ji}^1 > \hat{x}_{ji}^t\}) - \\ &\quad (\hat{x}^1 - \hat{x}^t)(\{[v_i, v_j] \in \delta^+(v_i) | \hat{x}_{ij}^1 > \hat{x}_{ij}^t\}) \\ &= (\hat{x}^1 - \hat{x}^t)(\{[v_j, v_i] \in \delta^-(v_i) | \hat{x}_{ji}^1 > \hat{x}_{ji}^t\}) + \\ &\quad (\hat{x}^1 - \hat{x}^t)(\{[v_j, v_i] \in \delta^-(v_i) | \hat{x}_{ji}^1 < \hat{x}_{ji}^t\}) &\quad (\text{because of (9.4) }) \\ &= (\hat{x}^1 - \hat{x}^t)(\delta^-(v_i)) &= 0 &\quad (\text{because of (9.5) or (9.6) }). \end{split}$$

With the same argumentation adapted to  $v_i = z_t$ , it follows that y satisfies (3.1). The other constraints (3.2) are satisfied in a trivial way. A substitution of (9.4) and (9.6) into (9.2) gives the flow-balance constraints (10.1). Thus, (X, x, y) is feasible for  $LP_{F'+FB}$ . II)  $v(LP_{m\vec{T}}) \leq v(LP_{F'+FB})$ : Let (X, x, y) denote an (optimal) solution for  $LP_{F'+FB}$ . From lemma 14 we know that there is an (optimal) solution  $(X, \hat{x}^r, \hat{y}^r)$  for each choice of the root vertex  $z_r \in R$ , with the property that  $\hat{s}_i := \hat{x}^t (\delta^-(v_i))$  (for any  $v_i \in V \setminus R$ ) has the same value for any choice of  $z_t \in R$ . With the argumentation of Theorem 13 it follows that  $(X, \hat{x}, \hat{s})$  is feasible for  $LP_{m\vec{T}}$ .

**Corollary 15.1** The constraints (9.1), (9.3), and (9.7) are useless with respect to the value of the linear relaxation  $LP_{m\vec{T}}$ .

**Corollary 15.2** The linear program  $LP_{m\vec{T}-}$  (with the same objective function as  $LP_{m\vec{T}}$ ) that contains only the equations (9.4), (9.5), and (9.6) is equivalent to  $LP_F$ .

### 2.7 A Collection Of New Formulations

In [PV01a] we introduced the common flow formulation  $LP_{F^2}$ , the strongest linear relaxation of polynomial size known so far. Concerning this relaxation we have frequently been asked three questions: How we developed the new relaxation, whether the result can be strengthened (the notation  $LP_{F^2}$ looks as if there could be an  $LP_{F^3}$ ) and how a relaxation of this kind could be used in a practical algorithm. We will address all three issues in the following.

#### 2.7.1 Properties of the Flow/Cut Relaxations



Figure 2.4: Example with  $v(LP_C) < v(P_C)$ .

A network with a deviation between the integral and the linear solution is shown in Figure 2.4. Choosing  $z_1$  as root yields the value 7.5 as  $v(LP_F)$  (setting all x-variables in the direction away from  $z_1$  to 0.5 leads to an optimal solution), while an optimal Steiner tree (and  $v(P_F)$ ) has value 8. Thus the integrality gap is at least  $\frac{8}{7.5}$ .

Goemans [Goe98] extends the example in Figure 2.4 and constructs networks  $G_k$  whose integrality gap comes arbitraily close to  $\frac{8}{7}$ . The network  $G_k$  consists essentially of  $\binom{k}{2}$  copies of the network in Figure 2.4. It has k + 1 terminals  $a_0, a_1, ..., a_k$ , and  $k^2$  non-terminals  $b_1, ..., b_k, c_{12}, ..., c_{ij}, ..., c_{k-1,k}, d_{12}, ..., d_{ij}, ..., d_{k-1,k}$ . For any i and j,  $1 \le i < j \le k$  one includes the network of Figure 2.4 with the following labeling:  $z_1$  is  $a_0, z_2$  is  $a_i, z_3$  is  $a_j, v_4$  and  $v_5$  are  $c_{ij}$  and  $d_{ij}$ , and  $v_6$  and  $v_7$  are  $b_i$  and  $b_j$ . An optimal Steiner tree has value 4k and the optimal  $LP_C$  solution is obtained by setting all x-variables in the direction away from  $a_0$  to  $\frac{1}{k}$ , yielding the value  $\frac{1}{k}(4k + 7\binom{k}{2}) = 3.5k + 0.5$ . Thus, the integrality gap approaches  $\frac{8}{7}$  with increasing k. This is the largest integrality gap known<sup>2</sup>.



Figure 2.5: *y*-variables and decomposition into trees.

Figure 2.5 shows how the y-variables of the  $LP_F$  solution for Figure 2.4 can be viewed as a linear combination of incidence vectors of paths in some trees. The upper-left network shows the  $y^2$  variables, each arc  $[v_i, v_j]$  corresponds to  $y_{ij}^2 = 0.5$ . The upper-right figure shows the same for  $y^3$ . The y values can be decomposed into two rooted trees  $T_1$  and  $T_2$  that are depicted in the two figures at the bottom. For any  $z_t \in R^{z_1}$ ,  $y^t$  is a linear combination of the incidence vectors of the paths from

<sup>&</sup>lt;sup>2</sup>Independently, we constructed other examples with the same asymptotic integrality gap. For example, the edges between  $a_i$  and  $b_i$  can be deleted for every *i* without deteriorating the asymptotic result.

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 $z_1$  to  $z_t$  in  $T_1$  and  $T_2$  (in this example both incidence vectors have weight 0.5).

If the x-variables were a linear combination of the incidence vectors of these trees, the optimal value of the linear relaxation would be the same as of the integral formulation. In this example, we see that the arc  $[v_4, v_5]$  causes the problem: The linear combination of the incidence vectors of the trees yields 1 for  $[v_4, v_5]$ , while  $x_{45}$  has the value 0.5. Looking at the flow-variables  $y^2$  and  $y^3$  one can see that at vertex  $v_4$  flows of the different commodities enter from different arcs  $(y_{74}^2 = y_{64}^3 = 0.5)$ , but depart together on arc  $[v_4, v_5]$ . We denote this situation as a rejoining. Formally, a rejoining of flows of the commodities  $\{z_{i_1}, z_{i_2}, \ldots, z_{i_k}\} =: B$  at vertex  $v_i \in V$  is defined by the condition  $\sum_{e \in \delta^-(v_i)} \max\{y_e^t | z_t \in B\} > \sum_{e \in \delta^+(v_i)} \max\{y_e^t | z_t \in B\}$ . In Section 2.7.3 we will show how to attack rejoining of flow in an extended linear program.

This situation is typical. In fact, we do not know of any example for a network with an integrality gap for  $LP_C$  that does not include a variant of Figure 2.4. Of course, there may be additional edges, edges replaced by paths, other edge weights and non-terminals replaced by terminals, but still Figure 2.4 is a minor of the network (a minor is a subgraph obtained after a sequence of contractions and deletions, for the relation of minors to linear relaxations for the Steiner problem see [CR94a]). Furthermore, the rejoining of flows of different commodities is a common property of all examples known to us, although more than two flows can rejoin (as we will show in Figure 2.6) and it may need a change of the root to see a rejoining. Figure 2.4 can be used to illustrate the latter: Turn  $v_4$  into a terminal  $z_4$  and choose  $z_4$  as root, the optimal linear solution value stays the same (setting all x-variables in the direction away from  $z_4$  to 0.5 leads to an optimal solution). Again, the y-variables can be decomposed into the same trees  $T_1$  and  $T_2$  (just with a reorientation of the arcs), but now there is no rejoining of flows.

#### 2.7.2 Common Flow

As we have seen in the last section, the typical problematic situation for  $LP_F$  can be described as the rejoining of flows of different commodities. We capture this condition in linear constraints with the help of additional variables  $y^B$  (for all subsets  $B \subseteq R^{z_1}$ ) that describe the amount of flow going from the root to any terminal  $z_t \in B$ :  $y^B \ge y^t$ , for all  $z_t \in B$ . From Figure 2.5 it is easy to see the flow to  $z_2$  and  $z_3$  ( $y^{\{z_2,z_3\}}$ ) incoming at  $v_4$  is greater than the flow outgoing of  $v_4$ . This cannot be the case if the *x*-variables correspond to a feasible Steiner tree. Thus, we can introduce additional constraints that assure that for each common flow the outgoing flow at any vertex has at least the value of the incoming flow.<sup>3</sup>

As already noted in Section 2.7.1, the same network can or cannot have rejoining of flow, depending on the choice of the root. To detect all rejoining we have to consider all possible roots. Thus, the variables that will be used to describe the common flow formulation will have the shape  $y^{r,B}$  and they will describe a flow from  $z_r$  to all terminals  $z_t \in B \subseteq R^{\{z_t\}}$  (we will skip the chosen root in the notation if the root is clear from the context).

#### 2.7.3 A Collection of Common Flow Formulations

We will first describe the full common flow formulation that has an exponential number of constraints and variables, and describe afterwards how to reduce it to different formulations (e.g., of only polynomial size)

<sup>&</sup>lt;sup>3</sup>In [PV01a] we used a semantically inverse, but mathematically equivalent definition of the common fbw. Meanwhile, we find the new definition more intuitive and compact.

$$P_{F^R}$$
  $c \cdot X \to \min,$ 

$$y^{r,\{z_t\}}(\delta^{-}(z_r)) = y^{r,\{z_t\}}(\delta^{+}(z_r)) - 1 \quad (z_r \in R; z_t \in R^{z_r}),$$

$$(12.1)$$

$$y^{r_{1}(z_{1})} - y^{r_{1}(z_{2})} \leq y^{r_{1}(z_{2})} \quad (\{z_{r}, z_{s}, z_{t}\} \in R),$$

$$(12.2)$$

$$y_{ji}^{r,B}(\delta^{-}(v_{i})) \leq y_{ij}^{r,B}(\delta^{+}(v_{i})) \quad (z_{r} \in R; B \subseteq R^{z_{r}}; v_{i} \in V \setminus (B \cup \{z_{r}\}),$$
(12.3)  
$$y^{r,B}(\delta^{-}(v_{i})) \leq y^{r,B}(\delta^{+}(v_{i})) \quad (z_{r} \in R; B \subseteq R^{z_{r}}; v_{i} \in V \setminus (B \cup \{z_{r}\}),$$
(12.4)

$$r^{r,B}(\delta^{-}(v_i)) \leq y^{r,C}(\delta^{-}(v_i)) \quad (z_r \in R; B \subseteq R^{*r}; v_i \in V \setminus (B \cup \{z_r\}), \quad (12.4)$$

$$y_{ij}^{r,R^{z_r}} + y_{ii}^{r,R^{z_r}} \leq X_{ij} \quad (z_r \in R; (i,j) \in E),$$
(12.6)

$$y^{r,\{z_r\}} = 0 \quad (z_r \in R), \tag{12.7}$$

$$y \ge 0,$$
 (12.8)  
 $y \ge 0,$ 

$$X \in \{0,1\}^{|E|}.$$
 (12.9)

It follows from the proof of Lemma 14 that the constraints (12.2) and (12.3) redirect the flows from root  $z_r$  to flows from any other root  $z_s \in R$  to each terminal  $z_t \in R$ . In particular, from these constraints in combination with (12.7) it follows that  $y_{ji}^{t,\{z_r\}} = y_{ij}^{r,\{z_t\}}$  for all  $z_r, z_t \in R$ . For each flow from a root  $z_r$  to a terminal  $z_t$ , described by the variables  $y^{r,\{z_t\}}$ , the constraints (12.1) guarantee an outflow of 1 out of  $z_r$ , and (together with the previous observation) an inflow of 1 in  $z_t$ . Together with the constraints (12.4) (with  $B = \{z_t\}$ ) and (12.8) this guarantees a flow of one unit from  $z_r$  to each terminal  $z_t \in R^{z_r}$ .

The constraints (12.4) also guarantee for any root  $z_r$  and set of terminals B that there is no rejoining at any vertex  $v_i$  (to be more precise, there is a penalty for rejoined flows in form of increased  $y^{r,B}$ values). The constraints (12.5) set each common flow from a root  $z_r$  to a set of terminal C to at least the maximum of all flows from  $z_r$  to a subset  $B \subset C$  of the terminals.

Finally the constraints (12.6) build the edgewise maximum over all flows from all terminals in the edge variables X.

We do not know of any network where this relaxation has an integrality gap. Unfortunately, we are neither able to prove that there is no integrality gap, nor to do a really broad experimental study to support this (see Section 2.9.3). Note that due to the exponential number of constraints *and* variables the possibility that there is no integrality gap is not ruled out by complexity arguments.

#### 2.7.4 Polynomial Variants

We will now show how to derive a polynomial-size relaxation from  $LP_{F^R}$ : We simply limit the terminal sets B to a polynomial size, e.g., by choosing two constants  $k_1 \ge 1$  and  $k_2 \ge 1$   $(k_1 + k_2 < |R|)$ , and using only those variables  $y^{r,B}$  with |B| either at most  $k_1$  or at least  $|R| - k_2$ . We denote the remaining relaxation with  $LP_{F^{k_1,k_2}}$ . It has  $O(r^{1+\max\{k_1,k_2-1\}}a)$  variables and  $O(r^{2\max\{k_1,k_2-1\}}a)$ constraints. For example, if we choose  $k_1 = 2$  and  $k_2 = 1$  and also use only one fixed root  $z_r$  we get a relaxation that is equivalent to  $LP_{F^2}$  presented in [PV01a]. Here, we state the relaxation in the form as it is derives from  $P_{F^R}$ , using the variables  $y^{r,B}$  for a fixed  $z_r$  and for  $B \in \mathcal{B} = \{C \subseteq R^{z_r} \mid |C| \in \{1, 2, |R| - 1\}\}$ 

$$\begin{array}{|c|c|c|c|}\hline P_{F^2} & c \cdot x & \to & \min, \\ \hline r \left( r \right) \left( r \right)$$

$$y^{r,\{z_t\}}(\delta^-(z_r)) = y^{r,\{z_t\}}(\delta^+(z_r)) - 1 \quad (z_t \in \mathbb{R}^{z_r}),$$
(13.1)

$$y^{r,B}(\delta^{-}(v_i)) \leq y^{r,B}(\delta^{+}(v_i)) \quad (B \in \mathcal{B}; v_i \in V \setminus (B \cup \{z_r\}),$$
(13.2)

$$y^{r,B} \leq y^{r,C} \quad (B \subset C \in \mathcal{B}), \tag{13.3}$$
$$y^{r,R^{z_r}} = x, (13.4)$$

$$y \ge 0, (13.5)$$
  

$$x \in \{0,1\}^{|A|}. (13.6)$$



Figure 2.6:  $v(LP_F) = 14\frac{1}{3} < v(LP_{F^{2,1}}) = 14\frac{2}{3} < v(LP_{F^R}) = 15 = v(P_{F^R}).$ 

The drawback of this limitation is that not all rejoinings will be detected by the limited relaxation, as the following example shows: In Figure 2.6, using  $LP_F$  with  $z_1$  as root, there is a rejoining of 3 different flows  $(y^2, y^3 \text{ and } y^4)$  at the central vertex. As  $LP_{F^2}$  captures only rejoining of 2 flows, there will still be some integrality gap. The figure also shows how  $k_2$  weakens the relaxation: Only if the common flow  $y^{r,B}$  to a set of terminals B is considered where  $z_2, z_3$  and  $z_4$  are in B, but  $z_5$  is not in B, the relaxation gives the optimal integer value. Thus, for  $k_1 \ge 3$  or  $k_2 \ge 2$  there is no integrality gap.

Note that the example in Figure 2.6 can be generalized to a network  $G[k_1, k_2]$ , which produces an integrality gap for any common flow relaxation that is limited by some constants  $k_1$  and  $k_2$ : Around a center vertex  $v_0$  put  $k_1 + 2$  edges  $[v_0, v_1], [v_0, v_2], \ldots, [v_0, v_{k_1+2}]$  of cost 1. Additionally use  $k_1 + 2$  terminals  $(a_1, a_2, \ldots, a_{k_1+2})$ . Each of the terminals  $z_i \in R$  is connected to all vertices  $v_j, j \neq 0, j \neq i$  with edges of cost  $k_1 + 1$ . Then connect the center vertex to  $k_2$  different terminals  $(b_1, b_2, \ldots, b_{k_2})$  with edges of cost 1. An integer optimal solution has the cost  $k_2 + (k_1 + 1)(k_1 + 2) + 2$ , but the following X-variables are feasible for the  $k_1, k_2$  limited common flow relaxation: Set X-variables of edges connected to  $b_i, 1 \leq i \leq k_2$  to 1, the X-variable for  $(v_0, v_1)$  to  $1 - (k_1 + 1)^{-1}$  and all other X variables to  $(k_1+1)^{-1}$ . Thus, the optimal value of the relaxation is at most  $k_2 + (k_1+1)(k_1+2)+2-(k_1+1)^{-1}$ . Figure 2.6 shows the network G[2, 1].

As a consequence, we have derived a collection of polynomial relaxations  $LP_{F^{k_1,k_2}}$ , where  $LP_{F^{k_1,k_2}}$  can be related to  $LP_{F^{j_1,j_2}}$ . We assume that  $k_1 + k_2 \leq j_1 + j_2$ .

- $k_1 \leq j_1, k_2 \leq j_2$ : The relaxation  $LP_{F^{j_1,j_2}}$  is stronger than  $LP_{F^{k_1,k_2}}$  (since it contains a superset of the constraints and variables). If additionally  $k_1 < j_1$  or  $k_2 < j_2$ , the relaxation  $LP_{F^{j_1,j_2}}$  is even strictly stronger. Consider the network  $G[\min\{k_1, j_1\}, \min\{k_2, j_2\}]$ ,  $LP_{F^{j_1,j_2}}$  gives the optimal solution, while  $LP_{F^{k_1,k_2}}$  has an integrality gap.
- **otherwise:** The relaxations are incomparable. To see this, consider the networks  $G[\min\{k_1, j_1\}, \max\{k_2, j_2\}]$  and  $G[\max\{k_1, j_1\}, \min\{k_2, j_2\}]$ .

Now, we show why considering all roots is important. The only difference between the relaxations  $LP_{F^{2,1}}$  and  $LP_{F^2}$  is that  $LP_{F^{2,1}}$  considers all terminals as root.

**Lemma 16** The relaxation  $LP_{F^{2,1}}$  is strictly stronger than  $LP_{F^2}$ .

**Proof:** The relaxation  $LP_{F^{2,1}}$  is stronger than  $LP_{F^2}$ , as it contains a superset of the constraints and variables. The network G[1,1] can be used to show that  $LP_{F^{2,1}}$  is even strictly stronger than  $LP_{F^2}$ . Here,  $LP_{F^{2,1}}$  gives the optimal value, but for  $LP_{F^2}$  it depends on the choice of the root. Using any of the vertices  $a_1, a_2$ , or  $a_3$  as root gives the integer optimal value, while with  $b_1$  as root there is an integrality gap.

#### 2.7.5 Restricted Version

We tested different restricted versions of the common flow relaxation, but did not come to a definitive conclusion which approach is the best. Nevertheless, we can present a very restricted, but still very useful version.

We observed that in many cases the tracing of the common flow to all |R| - 1 terminals ( $k_2 = 1$ ) catches most of the problematic situations except for insightfully constructed pathological instances. As depicted in Figure 2.6, to fool the relaxation with  $k_2 = 1$ , there typically has to be a terminal directly connected to the vertex at which the rejoining happens. In most of the cases where this happens, the terminal is the vertex itself. To attack those situations without the need to introduce a quadratic number of constraints or variables, we restrict the common flow relaxation by  $k_1 = k_2 = 1$  on non-terminals, and to  $k_2 = 2$  on terminals, but only considering the set  $R \setminus \{z_r, z_i\}$  at a terminal  $z_i$  (with respect to root  $z_r$ ). Additionally, we restrict the relaxation to one fixed root  $z_r$ , and because we do not need the X variables for different roots anymore, we can use the  $y^{R^{z_r}}$  as x variables.

In a branch-cut-price framework it is highly desirable to introduce as few variables as possible. Working with flow-based relaxations makes this difficult, because if a flow variable on one edge is necessary, all variables and all flow conservation constraints have to be inserted to get something useful. Therefore the cut-based relaxations are more suitable in this context. In analogy to  $P_C$ , we can use constraints of the form  $y_i^{\{z\}}(\delta^-(W)) \ge 1$  for all  $W \cap \{z_r, z_i\} = z_i$ . Finally, we eliminate all variables  $y^{\{z_i\}}$  and  $y_e^{R \setminus \{z_r, z_i\}}$  if e is not in  $\delta^-(z_i)$  in the following way:

Finally, we eliminate all variables  $y^{\{z_i\}}$  and  $y_e^{\mathbb{R}\setminus\{z_r,z_i\}}$  if e is not in  $\delta^-(z_i)$  in the following way: Unwanted  $y^{\{z_i\}}$  variables are replaced by  $y^{\mathbb{R}\setminus\{z_r,z_i\}}$  if the replacement is possible using the constraints (12.5). Similarly, unwanted  $y^{\mathbb{R}\setminus\{z_r,z_i\}}$  variables are replaced by x if possible. All constraints that still use unwanted variables are deleted.

Now, we have the following formulation:

$$\begin{array}{rcl}
P_{C'} & c \cdot x & \to & \min, \\
x(\delta^{-}(W)) &\geq & 1 & (z_1 \notin W, R \cap W \neq \emptyset), \quad (14.1) \\
y^{R \setminus \{z_r, z_i\}}(\delta^{-}(W) \cap \delta^{-}(z_i)) + x(\delta^{-}(W) \setminus \delta^{-}(z_i)) &\geq & 1 & (z_r \notin W, R^{z_i} \cap W \neq \emptyset), \quad (14.2) \\
& & x(\delta^{-}(v_i)) &\leq & x(\delta^{+}(v_i)) \quad (v_i \in V \setminus R), \quad (14.3) \\
& & y^{R \setminus \{z_r, z_i\}}(\delta^{-}(z_i)) &\leq & x(\delta^{+}(z_i)) \quad (z_i \in R^{z_r}), \quad (14.4) \\
& & y \geq & 0, \quad (14.5)
\end{array}$$

$$x \in \{0,1\}^{|A|}.$$
 (14.6)

This new formulation has fewer than 2|A| variables. Although it has an exponential number of constraints, solving the linear relaxation is relatively easy. This will be described in Section 2.9.3.

#### 2.7.6 Relation to Other Relaxations

**Lemma 17**  $v(LP_{F^2}) \ge v(LP_{2T}).$ 

**Proof:** Let (x, y) be an (optimal) solution of  $LP_{F^2}$ . For all  $\{z_k, z_l\} \subseteq R^{z_r}$  and k < l define  $\hat{y}^{kl} := y^{r,\{z_k\}} - y^{r,\{z_k,z_l\}}, \hat{y}^{kl} := y^{r,\{z_k,z_l\}}, \text{ and } \check{y}^{kl} := y^{r,\{z_k,z_l\}}$ . The variables  $(x,\check{y},\check{y},\check{y},\check{y})$  satisfy the constraints of  $LP_{2T}$ .

Because  $LP_{F^2}$  contains the flow-balance constraints and is stronger than  $LP_{2T}$ , it is stronger than  $LP_{2T+FB}$  (constructed by adding (10.1) to  $LP_{2T}$ ). It follows directly that  $LP_{F^2}$  is also stronger than  $LP_{F+FB}$ . The following example shows that it is even strictly stronger than the other stated relaxations.



Figure 2.7: Example with  $v(LP_{2T}) < v(LP_{F+FB}) = v(LP_{F^2}) = v(P_{F^2})$ .

**Example 4** Here is an example for  $v(LP_{2T}) < v(LP_{F^2}) = v(P_{F^2})$ : Setting all x-variables to 0.5 leads to a feasible (and optimal) solution for  $LP_{2T}$  with the value 13.5. An optimal solution for  $LP_{F^2}$  is  $x_{13} = x_{35} = x_{56} = x_{62} = x_{64} = 1$ , which forms a Steiner tree with value 14. Notice that this is also an example with  $v(LP_{2T}) < v(LP_{F+FB})$ . On the other hand, if  $v_5$  is moved to R,  $v(LP_{F+FB}) = v(LP_F) = 12 < v(LP_{2T}) = v(LP_{2T+FB}) = 13.5 < v(LP_{F^2}) = v(P_{F^2})$  (The optimal value for  $LP_{F+FB}$  is reached by  $\hat{x}$  with  $\hat{x}_{12} = \hat{x}_{13} = \hat{x}_{14} = \hat{x}_{25} = \hat{x}_{35} = \hat{x}_{45} = 1/3, \hat{x}_{56} = \hat{x}_{62} = \hat{x}_{63} = \hat{x}_{64} = 2/3$ ). Thus,  $LP_{F+FB}$  and  $LP_{2T}$  are incomparable.

This example has been chosen because it is especially instructive. For  $v(LP_{2T+FB}) < v(LP_{F^2})$ , as for all other statements in this work that one relaxation is *strictly* stronger than another, we also know (originally) undirected instances as examples.

The relaxation  $LP_{F^2}$ , as well as  $LP_{2T}$ , makes it difficult for flows to two different terminals to split up and rejoin by increasing the x-variables on arcs with rejoined flow. One could say that rejoining has to be "payed" for. To get an intuitive impression why  $LP_{F^2}$  is strictly stronger than  $LP_{2T}$  (or even  $LP_{2T+FB}$ ), notice that in  $LP_{F^2}$ , there is one flow to each terminal and rejoining of each pair of these flows has to be payed for; while in  $LP_{2T}$ , it is just required that for each pair of terminals there are two flows and rejoining them has to be payed for. The latter task is easier; for example it is possible (for given x-values) that for each pair of terminals there are two flows that do not rejoin, but there are not  $|R^{z_1}|$  flows to all terminals in  $R^{z_1}$  that do not rejoin pairwise; this is the case in example 2.7 (setting all x-variables to 0.5).

Still, we have to prove two more relations to relate the common flow relaxations to the other relaxations.

**Lemma 18** The relaxation  $LP_{F+FB}$  is equivalent to  $LP_{F^{1,1}}$ .

**Proof:** The relaxations are nearly the same. The only difference is that  $LP_{F^{1,1}}$  considers all possible roots. But as we have seen in Lemma 14, the choice of the root does not change the value of the

solution and the transformation of a solution for one root into a solution for another is covered exactly by the constraints (12.2) and (12.3).  $\Box$ 

**Lemma 19** The relaxation  $LP_{F^2}$  is strictly stronger than  $LP_{C'}$  and  $LP_{C'}$  is strictly stronger than  $LP_{F^{1,1}}$ .

**Proof:** It is obvious that  $LP_{F^2}$  is stronger than  $LP_{C'}$ , as  $LP_{C'}$  is a restricted and aggregated version of  $LP_{F^2}$ . Similarly,  $LP_{C'}$  contains a superset of variables and constraints of  $LP_{C+FB}$ , which is equivalent to  $LP_{F^{1,1}}$ . To see that the relations are also strictly stronger, consider the networks G[1, 1]. If we choose  $a_1, a_2$ , or  $a_3$  as root,  $LP_{C'}$  has an integrality gap,  $LP_{F^2}$  not. And if we contract the edge  $(v_0, b_1), LP_{C'}$  gives the integer optimal value, while  $LP_{F^{1,1}}$  still has an integrality gap.  $\Box$ 

# 2.8 A Hierarchy of Relaxations

## 2.8.1 Summary of the Relations

The following Figure 2.8 summarizes the relations stated before. All relaxations in the same box are equivalent. A line between two boxes means that the relaxations in the upper box are strictly stronger than those in the lower box. Notice that the "strictly stronger" relation is transitive. For an overview of the meaning of the abbreviations, see Table 2.1 on page 22.



Figure 2.8: Hierarchy of relaxations.

#### 2.8.2 Extensions to Polyhedral Results

It should be mentioned that some of the stated results on the relationship between the optimal values of linear relaxations extend directly to polyhedral results concerning the corresponding feasible sets. This is always the case if optimality is not used in the proofs (e.g., in Lemmas 7 or 8); and hence the feasible set of one relaxation (projected into the x-space) is mapped into the corresponding set of some other. The situation is different in the other cases (e.g., the proofs of Lemma 9 or Theorem 13). Here the assumption of optimality of x can obviously be replaced by the assumption of minimality of x (a feasible x is minimal if there is no feasible  $x' \neq x$  with  $x' \leq x$ ). In such cases, the presented results extend directly to polyhedral results in the sense of inclusions between the dominants of the corresponding polyhedra (projected into the x-space). (The dominant of Q is  $\{x' \mid x' \geq x \in Q\}$ .)

Note also that polyhedral results concerning the facets of the Steiner tree polyhedron (like those in [CR94a, CR94b]) fall into a different category. Our line of approach has been studying linear relaxations of general, explicitly given (and frequently used) integer formulations; not methods for describing facet defining inequalities. Applying such descriptions is typically possible only if the graph has certain properties (e.g., that it contains a special substructure) and involves separation problems that are believed to be difficult.

## 2.9 Using Relaxations

To actually exploit the relaxations for computing lower bounds, two factors are of more or less equal importance: How good the optimal values of the corresponding linear programs are, and how fast these values can be determined or sufficiently approximated. In the following, we investigate both questions for the stated relaxations.

#### 2.9.1 The Spanning Tree Formulation and Lagrangian Relaxation

A Lagrangian relaxation  $LaP_{T_0}$  of the tree formulation  $P_{T_0}$  is described in [Bea89], relaxing the degree constraints (6.2). After this, a subgradient optimization of the Lagrangian multiplier problem can be used, which involves calculating a minimum spanning tree in each iteration. Using this approach, the value  $v(LP_{T_0})$  can be approximated fairly fast (this relaxation has the integrality property). The problem here is the value  $v(LP_{T_0})$  itself. Theorem 13 already indicates theoretically that  $LP_{T_0}$  is not a generally tight relaxation. Empirically, we observed that usually the bound  $v(LP_{T_0})$  is only satisfactory for instances where the average distance between terminals is not too high in comparison to the average edge length (e.g., random networks with many terminals). A bad situation for this relaxation typically arises from instances modeling points in the plane with respect to a given metric. For instances with Euclidean distances or grid instances with few terminals, gaps of more than 50% are not exceptional. Nevertheless, we have further investigated the mentioned Lagrangian relaxation, since it can be useful for some instances.

We obtained a minor improvement in the speed of the subgradient optimization by applying a sensitivity analysis for the Lagrangian multipliers. Using data structures for efficient handling of tree bottlenecks and alternative chords (see [Tar79, VJ83]) allows fast calculation of the quantities by which each multiplier can be changed without affecting the validity of the calculated minimum spanning tree. Modifying the multipliers by these quantities improves the lower bound immediately.

In [DV87], some modifications for this relaxation are suggested, for example adding (and relaxing) further constraints and using another structure for  $G_0$ . In our experiments, these modifications did not

improve the overall results of the lower bound calculation: In situations where  $LaP_{T_0}$  leads to a substantial gap, no decisive improvements could be achieved using these modifications.

In [BL98], a relaxation constructed by adding the Steiner cut (and some other) constraints to  $LP_{T_0}$  is used. This indeed leads to a stronger relaxation than  $LP_{T_0}$ . However, as we have proved in Corollary 13.1 in Section 2.5,  $LP_C$  cannot be strengthened (i.e.,  $v(LP_C)$  does not change) by adding constraints like those present in  $LP_{T_0}$ ; this motivates concentrating on  $LP_C$  itself.

#### 2.9.2 The Cut Formulation, Dual Ascent and Row Generating

Considering the relaxation  $LP_C$ , the situation is to some degree the converse of the case  $LP_{T_0}$ . It is known that  $v(LP_C)$  does not deviate from the optimum by more than 50% [GB93]. All our experimental investigations strongly suggest that  $LP_F$  is a very tight relaxation. As an example, for all D-instances of the OR-Library  $v(LP_C)$  is equal to  $v(P_C)$ . Even for the instances where there is a gap, the knowledge of a solution of  $LP_C$  has been usually sufficient to solve the instance exactly (without branching) through bound-based reduction techniques (Section 3.4). So, the really interesting problem is how to calculate (or sufficiently approximate) a solution for  $LP_C$ .

The direct approach of solving the complete linear program using a standard LP-solver is not practical, even for the equivalent multicommodity flow relaxation [Won84], which has approximately ra variables and r(a + n) constraints: This is still too much for moderate and large instances; and the resulting linear programs are often highly degenerated.

We present those two approaches that have been proven to be very effective, dual ascent and a row generating approach.

**Dual Ascent:** A fast dual ascent algorithm that generally provides fairly good lower bounds was described in [Won84] for the equivalent multicommodity flow relaxation. Below, we give an alternative description of it as a dual ascent algorithm for  $LP_C$ , which we call **DUAL-ASCENT**. First, we state the dual linear program  $DLP_C$ , that uses dual variables  $u_W$  for each Steiner cut W ( $z_1 \in \overline{W}$  and  $R^{z_1} \cap W \neq \emptyset$ ):

$$\frac{\overline{DLP_C}}{\sum_{W,a\in\delta^{-}(W)}} \sum_{W_W} d_W \quad \forall max, \qquad (15.1)$$

$$u \geq 0. \tag{15.2}$$

Now, we can describe the algorithm:

- Initialize the reduced costs ( $\tilde{c} := c$ ), the lower bound (*lower* := 0) and assume all dual variables u have been set to zero.
- In each iteration, choose a terminal  $z_t \in R^{z_1}$  not reachable from the root by edges of zero reduced cost. Let  $W, z_t \in W$ , be the smallest set such that  $\{\overline{W}, W\}$  is a Steiner cut and  $\tilde{c}_{ij} > 0$  for all  $[v_i, v_j] \in \delta^-(W)$ . Set the dual variable  $u_W$  to  $\Delta := \min\{\tilde{c}_{ij} \mid [v_i, v_j] \in \delta^-(W)\}$  and let  $lower := lower + \Delta$  and  $\tilde{c}_{ij} := \tilde{c}_{ij} \Delta$  (for all  $[v_i, v_j] \in \delta^-(W)$ ).
- Repeat until no such terminal is left.

A good implementation of this algorithm has running time  $O(a \cdot \min\{a, rn\})$  (see for example [Dui93]). Empirically, it is usually faster than this time bound would suggest.

#### 2.9. USING RELAXATIONS

The algorithm DUAL-ASCENT already achieves very good results. Out of the 20 D-instances of the OR-Library, a DUAL-ASCENT run yields the optimum (i.e.,  $v(P_C)$ ) for 12 instances. The average gap between lower bound and optimum is 0.4% and the average running time is 0.02 seconds.

A critical point in this algorithm is the choice of  $z_t$  in each iteration and, for the undirected version, the choice of the root  $z_1$ . Although it has been shown [GM93] that  $v(LP_C)$  is independent of the choice of the root vertex, the lower bound calculated by DUAL-ASCENT is not. For this reason we start DUAL-ASCENT with different roots if a strengthening of the bound is necessary. Again, for the D-instances, considering up to ten different roots improves the average gap to 0.1%; achieving the optimum for 16 instances. The average running time for this lower bound calculation is 0.08 seconds. Further improvements can be achieved by using the reductions and upper bound calculations, which are done in combination with DUAL-ASCENT (see Sections 3.4.2 and 4.4). Some results on this can be found in [PV01c].

Even using this amplification, there are still cases where DUAL-ASCENT does not reach the value  $v(LP_C)$ . We tested different criteria for the choice of  $z_t$  in each iteration. Our standard criterion is: Choose  $z_t$  such that W is smallest. We had some success with the following idea that tries to guide DUAL-ASCENT with the help of a heuristically constructed Steiner tree: Assume that the upper bound is already optimal. DUAL-ASCENT can reach the optimum only if in each set  $\delta^-(W)$  there is exactly one edge of the corresponding Steiner tree. Of course this criterion can not always be realized, especially if the best known Steiner tree is not optimal or  $v(LP_C) < v(P_C)$ . Nevertheless, it is a heuristic criterion that in many cases leads to better lower (and, indirectly, upper) bounds.

A natural alternative for a better approximation of  $v(LP_C)$  builds upon a Lagrangian relaxation of the multicommodity flow formulation; an approach already used in in [Bea84] (but with the much weaker undirected relaxation; see also [HRW92]). Relaxing the constraints that bind edge and flow variables together, the problem decomposes into (mainly) r - 1 single pair shortest path problems, which can be solved in time  $O(r(m + n \log n))$ . This relaxation has the integrality property, and can be used in combination with subgradient optimization to approximate  $v(LP_C)$ . In [PV97], we have investigated this approach and presented some improvements, particularly in combination with the algorithm DUAL-ASCENT and with sophisticated reduction techniques. Although this approach is quite effective in many cases, for large instances with many terminals it tends to be too slow. So, it is not used here and is replaced by the following approach.

**Row Generating:** To get an optimal solution for  $LP_C$ , one can begin with a subset of constraints of  $LP_C$  as the initial program, and successively solve the current program, find Steiner cut inequalities violated by the current solution x, add them to the program, and iterate this process by reoptimizing the program, until no Steiner cut inequality is violated anymore. This is an approach already used by many authors (see for example [CGR92, BL98, KM98]).

In order to find violated Steiner cut inequalities (or to establish that no such inequality exists), one can compute a minimum capacity cut in each of the r-1 flow networks constructed from G by choosing the root  $(z_1)$  as the source, a terminal  $z_t \in R^{z_1}$  as the sink and the current  $x_{ij}$ -value as the capacity of the arc  $[v_i, v_j]$ . Although there are other (heuristical) ways to find such violated inequalities, using those corresponding to minimum cuts usually leads to better overall results. Indeed, it is even very advantageous to find in each case a minimum capacity cut with a minimum number of cut edges, an idea already used in [KM98]. This can be realized by adding a small  $\epsilon$  to the capacity of each edge before solving the minimum cut problem. Although this leads to much denser flow networks, the linear programs obtained are easier to (re-)optimize (and the corresponding constraints seem to be much stronger). As a consequence, the overall results (especially the total number of necessary reoptimizations) are clearly superior. It must be mentioned that in our implementation, the time for finding all the r - 1 minimum cuts is dominated by the time for reoptimizing the linear

programs.

For computing minimum cuts, we implemented the highest-label preflow-push algorithm with several auxiliary heuristics, including the global and the gap relabeling heuristics [CG97]. Although no better time bound than  $O(n^2\sqrt{m})$  can be given for this algorithm, using the heuristics mentioned the empirical running times were much better described by  $O(n^{1.5})$ . As long as only minimum cuts from the sink-side are to be computed, only the first stage of the algorithm has to be performed. Besides, in this context several additional heuristics can be used to improve the empirical times further; for example, sinks which are reachable from the root (or another terminal) by paths of capacity no less than 1 need not be considered.

For (re-)optimizing the linear programs, we use the dual simplex routine in the callable library of CPLEX 8.0. Here, the warm-start ability of the simplex algorithm can be particularly utilized.

We have achieved considerable speedups by inserting the cuts generated by the algorithm DUAL-ASCENT into the initial linear program. In this case the lower bound provided by DUAL-ASCENT (which is often very close to  $v(LP_C)$ ) is already reached in the first iteration; and the number of necessary reoptimizations and the time needed per reoptimization are comparable to the case without these cuts *after* reaching this lower bound value. As a consequence, the overall times are clearly improved.

As mentioned earlier, the flow balance constraints (10.1) can used to strengthen the linear programs. Empirically, we found it advantageous in terms of running times to insert all the flow-balance inequalities into the initial program. Although the other additional constraints used in [KM98] cannot enhance the value of the relaxation (see the corollaries of Theorem 13), a group of them can speed the process up if its violated members are added to the current program. These are constraints of the form of equations (8.2) of  $LP_{\vec{T}}$ .

To save time and space, we do some garbage collection every ten iterations, purging the constraints that have had large positive slack values in all the iterations since the last garbage collection. Further we make sure that no constraint is present in a linear program more than once.

Another idea, which is promising at first sight, is pricing: To achieve further speedups one can begin with a subset of variables as active variables and at certain stages (especially when a correct lower bound is necessary) add variables that do not price out correctly (have negative reduced costs) to the program (activate them); a correct lower bound is given when all non-active variables have non-negative reduced costs with respect to the current dual solution. We have tried several schemes for using this idea, but could not achieve decisive *additional* improvements through these schemes. The main reason is that because of our massive usage of reduction techniques (see the next chapter), most variables that could be priced out are eliminated anyway. It seems that the information provided by the linear relaxations (like reduced costs) are more effectively used in bound-based reduction techniques (see Section 3.4).

#### 2.9.3 Using Tighter Relaxations

In this section, we will outline approaches for actually using tighter relaxations than  $LP_C$  algorithmically.

The complete common flow relaxation  $LP_{FR}$  is not a good starting point for building algorithms, because its exponential size makes it difficult (if not impossible) to come up with a practical algorithm using it. Simply writing down the linear program and starting an LP-solver fails already on problems of toy size. But also a branch-cut-price approach fails, because too many variables and constraints have to be included and each iteration of the column and row generation method can take too much time without making any substantial progress.

#### 2.10. SOME EXPERIMENTAL RESULTS

Although a more efficient realization cannot be ruled out, turning to the restricted version  $LP_{C'}$  (see Section 2.7.5) of the common flow relaxation is a by far more appealing approach. Here one can start with the *x* variables and the flow-balance constraints (14.3), and then use a column and row generation approach. We work in three levels. In each iteration, we process a level only if the previous levels could not find a new row or column.

- 1. We look for violated Steiner cut constraints (14.1) as described in the previous section.
- 2. For those  $y^{R \setminus \{z_r, z_i\}}$  that have already been inserted, we look for violated Steiner cut constraints (14.2), again with the same procedure as previously described.
- 3. Now, we are looking for violated constraints (14.4) of the form y<sup>R\{z<sub>r</sub>,z<sub>i</sub>}</sup>(δ<sup>-</sup>(z<sub>i</sub>)) ≤ x(δ<sup>+</sup>(z<sub>i</sub>)) for some z<sub>i</sub>. Here, one has to assume that all y<sup>R\{z<sub>r</sub>,z<sub>i</sub>}</sup> that are not already included in the linear program are equal to x if x(δ<sup>+</sup>(z<sub>i</sub>)) > 0. Otherwise, a lot of unnecessary y-variables would have to be included because of negative reduced costs. If we find violated constraints (14.4), we also include the necessary variables y<sup>R\{z<sub>r</sub>,z<sub>i</sub>}</sup>.

With this procedure the linear program will reach the value of  $v(LP_{C'})$ , which is in some cases significantly above the value of  $v(LP_{F+FB})$  (see next section). Some other practical approaches for improving the lower bound are mentioned in Section 2.11.

# 2.10 Some Experimental Results

In this section, we present summarized results of some different approaches for computing lower bounds on bechmark instances from [SteinLib], see Table 2.2, (see Section 1.2 for a description of the instance groups). We measured running times and the average gap between the lower bound (rounded up to the next integer) and the known optimum. We give results for all groups of instances from SteinLib where all optimal values are known. A stroke in the table means that we were not able to compute the lower bound for some of the instances of this class, because of running time constraints.

The main purpose of this table is to provide a rough overview of the power of the different approaches. Additionally, a comparison of our implementation of the different approaches with those of others is now possible.

To ensure comparability we did not use reductions before starting the lower bound computation. Note that the computation of a lower bound on an unreduced instance is a very artificial setting in our context. Obviously, it cannot reveal the true value of a technique as part of exact algorithm or upper bound computation. Usually, one would first try all reduction techniques (even those using lower bound calculations themselves) before starting time consuming lower bound calculations. Reduction techniques influence the lower bound calculation in different ways:

- 1. As the graph may be smaller, the computation may be faster. Also the structure of the graph may change: Typically, the graph becomes sparser, which may be an advantage or a disadvantage.
- 2. In the case of approximative approaches for the solution of a relaxation (e.g., DUAL-ASCENT), the quality of the approximation of the lower bound is typically improved.
- 3. Additionally, the value of the optimal solution of the relaxation can improve.

On the other hand, measuring running times and lower bound gaps on reduced instances would make a comparison with other implementations impossible.

instance	DUAL-ASCENT		DUAL-ASCENT, 10 roots		$LP_C$ , row generation	
group	time (s)	gap (%)	time (s)	gap (%)	time (s)	gap (%)
1R	0.01	3.49	0.02	2.03	_	_
2R	0.01	5.87	0.04	4.41		
D	0.02	0.40	0.08	0.10	662.29	0.00
E	0.12	0.26	0.62	0.25	5714.26	0.00
ES10000FST	257.30	1.19	2434.40	1.18	—	
ES1000FST	0.31	1.20	2.58	1.16	264.53	0.008
I080	0.01	1.25	0.05	0.76	10.81	0.14
I160	0.04	1.10	0.31	0.83	4726.18	0.22
I320	0.17	1.24	1.55	1.05		
LIN	0.39	2.49	3.39	1.88	—	_
MC	0.01	2.51	0.04	2.32	1941.06	0.54
TSPFST	1.72	0.67	16.95	0.60	5897.02	0.007
VLSI	0.23	2.00	2.16	1.51	_	_
WRP3	0.03	0.0006	0.20	0.0005		
WRP4	0.01	0.0008	0.10	0.0006	929.24	0.000003
Х	1.02	0.06	6.04	0.06	—	—

Table 2.2: Average results for lower bound calculations on unreduced instances.

We have not included values for  $LP_{F+FB}$  and  $LP_{C'}$  in this table. The reason for this is that on those instance groups, where the improvements due to  $LP_{F+FB}$  and  $LP_{C'}$  can show their strength, the computation of  $v(LP_C)$  is already too time consuming on the weakly reduced instances. The only exception are the ES1000FST instances, where  $LP_{C'}$  reduces the average gap from 0.008% of  $LP_C$ to 0.001%, which is a drastic improvement in the context of an exact algorithm.

# 2.11 Concluding Remarks on Lower Bounds

We have established a hierarchy of relaxations, providing very clear relations between relaxations. Furthermore, we introduced a collection of very tight linear relaxations for the Steiner problem. There are some polynomial variants in this collection that are stronger than any previously known relaxation of polynomial size. One of these relaxations can even be applied in practical algorithms and improve the performance of the current best approach for problem instances of the types mentioned.

Future work can be done in different directions: A theoretically important open question is whether a better worst-case guarantee for the integrality gap for any relaxation is possible. Practically, an improved scheme for using the presented formulations could lead to faster solutions of problem instances of SteinLib, where a small gap in linear relaxation is the crucial problem.

Also a constraint generation approach that is not based on some integer formulation, but on the "local cuts" approach as presented for the TSP by Applegate, Bixby, Chvátal, and Cook [ABCC01], could be applicable in this context. We made some steps in this direction by deriving and implementing some exact and heuristic projection methods, but we could apply them successfully only in rare cases. Our current version is able to improve the value of  $v(LP_{C'})$  in the cases it does not reach  $v(P_{C'})$ . But either the problem instances could be solved faster by using bound-based reductions or branching, or neither of the approaches is able to solve the instance (this holds for those problems that are insightfully constructed to fool the currently known methods). We conjecture that one reason why we could improve our results only rarely with the "local cuts" approach is our partitioning-based reduction techniques (see Section 3.6) that exploit locality even more effectively than the "local cuts" approach.

Recently, we have developed a new approach for improving the lower bound for the Steiner problem that works very well in practice. The main idea is to expand the network such that the value of the

#### 2.11. CONCLUDING REMARKS ON LOWER BOUNDS

linear relaxation is improved. This idea is inspired by the column replacement techniques that were introduced by Balas and Padberg [BP75] and generalized by Haus et. al. [HKW01] and Gentile et. al. [GHK<sup>+</sup>02]. In these and other papers a general technique for solving integer programs is developed. However, these techniques are mainly viewed as primal algorithms, and extensions for combinatorial optimization problems are presented for the Stable Set problem only. Furthermore, these extensions are not yet part of a practical algorithm (the general integer programming techniques have been applied successfully). Thus, we are the first to apply this basic idea in a practical algorithm for a concrete combinatorial optimization problem.



Figure 2.9: Splitting of vertex  $v_j$ . The filled circles are terminals,  $z_1$  is the root, all arcs have cost 1. An optimal Steiner arborescence has value 6 in each network. In the left network  $v(LP_{C+FB})$  is 5.5 (set the x-values of the dashed arcs to 0.5 and of  $[v_j, z_4]$  to 1), but 6 in the right network (again, set the x-values of the dashed arcs and of  $[v_j^a, z_4]$  and  $[v_j^b, z_4]$  to 0.5). The difference is that in the left network, there is a situation that is called "rejoining of flows": Flows from  $z_1$  to  $z_2$  and from  $z_1$  to  $z_3$  enter  $v_j$  on different arcs, but leave on the same arc, so they are accounted in the x variables only once. Before splitting, the x-value corresponding to the arc  $[v_j, v_c]$  is 0.5, after splitting the corresponding x-values sum up to 1.

As the algorithm is still under development, we will only sketch the basic operation here. This operation is the splitting of a vertex as depicted in Figure 2.9. A simplified version of the splitting operation is described by the following pseudocode:

```
\begin{array}{lll} SPLIT-VERTEX(G,v_j): & (\text{assuming } v_j \in V \setminus R) \\ 1 & \text{forall } [v_i,v_j] \in \delta^-(v_j): \\ 2 & insert \ a \ new \ vertex \ v_j^i \ into \ G \\ 3 & insert \ an \ arc \ [v_i,v_j^i] \ with \ cost \ c(v_i,v_j) \ into \ G \\ 4 & \text{forall } [v_j,v_k] \in \delta^+(v_j): \\ 5 & insert \ an \ arc \ [v_j^i,v_k] \ with \ cost \ c(v_j,v_k) \ into \ G \\ 6 & delete \ v_i \end{array}
```

The transformation is valid, as any optimal Steiner tree of the original network can be transformed into a feasible Steiner tree in the new network with the same cost and vice versa. The reasoning behind this transformation is that in an optimal directed Steiner tree, each vertex has at most indegree 1.

Details on the two methods are presented in a technical report [APV03].

Another area for further development are the polynomial approximation algorithms for  $LP_F$  and  $LP_C$  by Garg and K onemann [GK98] and Garg and Khandekar [GK02]. We did some preliminary tests with these methods, but we could not make them as effective as other approaches.

**Chapter 3** 

# **Simplifying Problem Instances Using Reduction Techniques**

# 3.1 Introduction

Intuitively spoken, reduction techniques reduce the size of an original problem instance without changing the optimal solution. It has been known for some time that reductions can play an important role as a preprocessing step when one wants to deal with  $\mathcal{NP}$ -hard problems algorithmically. In particular, the importance of reduction techniques for the Steiner problem has been widely recognized and a large number of techniques were developed, with the PhD thesis of Cees Duin [Dui93] being a milestone.

We will not give a formal definition of what a reduction is, an intuitive understanding is sufficient:

- 1. Some *reduction condition* of a *reduction test* is satisfied (e.g., an edge  $(v_i, v_j)$  is longer than the length of a shortest path between its endpoints  $v_i$  and  $v_j$ ).
- 2. By this, we get some knowledge about the optimal solutions of the problem. In the example, we know that an optimal solution exists that does not contain  $(v_i, v_j)$ , in fact no optimal solution contains  $(v_i, v_j)$ .
- 3. We transform the problem instance into one that is believed to be simpler, but we know that there is some efficient way to transform any optimal solution of the reduced problem instance into an optimal solution of the original instance. In the example, we can simply delete the edge  $(v_i, v_j)$ , a transformation procedure is not necessary.

We distinguish some major classes of reduction tests:

- The **alternative-based** tests (Section 3.3) use the existence of alternative solutions. For example in case of exclusion tests, it is shown that for any solution containing a certain part of the graph (e.g., a vertex or an edge) there is an alternative solution of no greater cost without this part; the inclusion tests use the converse argument.
- The **bound-based** tests (Section 3.4) use a lower bound for the value of an optimal solution with the additional constraint that a certain part of the graph is contained (in case of exclusion tests) or is not contained (in case of inclusion tests) in the solution; these tests can reduce the problem if such a lower bound exceeds a known upper bound. This can also be interpreted as a kind of an implicit branch-and-bound approach.
- The **extended** tests (Section 3.5) use backtracking to extend the scope of inspection of the reduction tests, here bound-based and alternative-based approaches can be combined.
- The **partitioning-based** tests (Section 3.6) exploit a partitioning of the network to derive reductions using the (optimal) solutions of subnetworks.

Our work in this field is threefold:

- 1. We design efficient realizations of some classical reduction tests, which would have been too time-consuming for large instances in their original form, especially for application in heuristics.
- 2. We design some new tests, filling some of the gaps the classical tests had left. Notice that each test is specially effective on a certain type of instances, having less (or even no) effect on others, so it is important to have a large arsenal of tests at one's disposal.

3. We integrated these tests into a program packet. It should be emphasized that the most impressive achievements of reductions are mainly due to the interaction of different tests, achieving results that are incomparable to those each single test could achieve on the same instance on its own.

# **3.2** Additional Definitions for Reductions

The **bottleneck distance**  $b(v_i, v_j)$  between  $v_i$  and  $v_j$  is the length of the shortest edge among the longest edges taken over all paths from  $v_i$  to  $v_j$  in G. The **restricted bottleneck distance**  $\bar{b}(v_i, v_j)$  between  $v_i$  and  $v_j$  is the bottleneck distance between  $v_i$  and  $v_j$  in G excluding  $(v_i, v_j)$ . An **elementary path** is a path that contains terminals only at the endpoints. Any path P between two vertices  $v_i$  and  $v_j$  in a network G can be broken at inner terminals into one or more elementary paths. The **Steiner distance** between  $v_i$  and  $v_j$  along P is the length of the longest elementary path in P. The **bottleneck Steiner distance** (sometimes also called "special distance")  $s(v_i, v_j)$  between  $v_i$  and  $v_j$  is the minimum Steiner distance taken over all paths between  $v_i$  and  $v_j$  in G. The **restricted bottleneck Steiner distance**  $\bar{s}(v_i, v_j)$  between  $v_i$  and  $v_j$  is the bottleneck Steiner distance between  $v_i$  and  $v_j$  is the bottleneck Steiner distance taken over all paths between  $v_i$  and  $v_j$  in G. The **restricted bottleneck Steiner distance**  $\bar{s}(v_i, v_j)$  between  $v_i$  and  $v_j$  is the bottleneck Steiner distance between  $v_i$  and  $v_j$  in G excluding  $(v_i, v_j)$ . Note that if R = V, then  $b_{ij} = s_{ij}$  and  $\bar{b}_{ij} = \bar{s}_{ij}$ .

A key node in a tree T is a node that is either a nonterminal of degree at least 3 or a terminal. The unique path in T between two vertices  $v_i \in T$  and  $v_j \in T$  is called the **fundamental path**. A **tree bottleneck** between  $v_i \in T$  and  $v_j \in T$  is a longest subpath on the fundamental path between  $v_i$  and  $v_j$  in T in which only the endpoints may be key nodes; and  $t_{ij}$  denotes the length of such a tree bottleneck.

# **3.3** Alternative-based Reductions

In this subsection we present a collection of alternative-based tests, including some new versions of classical tests and some new tests, which can all be realized in time  $O(m + n \log n)$ .

### **3.3.1 PT**<sub>m</sub> and Related Tests

The test  $PT_m$  (Paths with many Terminals) was introduced in [DV89]:

**PT**<sub>m</sub> test: Every edge  $(v_i, v_j)$  with  $c(v_i, v_j) > s(v_i, v_j)$  can be removed from G. [DV89]

**Proof:** Suppose all Steiner minimal trees contain an edge  $(v_i, v_j)$  with  $c_{ij} > s_{ij}$ . Let T be such a tree. Removing  $(v_i, v_j)$  from T divides it into two components. Let P be a path between  $v_i$  and  $v_j$  with the Steiner distance  $s_{ij}$ . On this path there is at least one elementary path P' that connects the two components of T. The tree T becomes shorter, if  $(v_i, v_j)$  is replaced by P', a contradiction.  $\Box$ 

The  $PT_m$  test is one of the most effective classical exclusion tests, but in its original form it is too time-consuming for large instances. Here we consider a fast realization of this test, which also uses inaccurate information. The modifications follow the same principal ideas as in [Dui93]. Later we will simply refer to this modified version as the  $PT_m$  test. Experimentally, one generally observes only a marginal difference in the effectiveness of the original test and its modified version.

For two terminals  $z_i$  and  $z_j$ , one observes that the bottleneck Steiner distance  $s(z_i, z_j)$  can be computed by determining a bottleneck on the fundamental path between  $z_i$  and  $z_j$  in the spanning tree  $T'_D(R)$ , which can be constructed in time  $O(m + n \log n)$  [Meh88]. Each such bottleneck can be trivially computed in time O(r), leading to a total time O(qr) for q inquiries  $(q \in O(\min\{m, r^2\}))$ . Observing that one actually has a static-tree variant of the bottleneck problem, one can use a strategy



Figure 3.1: Example for the  $PT_m$  test.

based on depth-first search (as described in [VJ83]) to achieve time  $\Theta(r^2)$  for all inquiries. One can go further and solve the problem as an off-line variant for all q inquiries in time  $O(q\alpha(m, r))$  using the Eval-Link-Update data structure [Tar79]. But this data structure is rather complex and leads to relatively large constant factors, and this bound is dominated by the worst-case time of other test operations anyway. So we suggest another method to achieve the desired worst-case time  $O(m+n \log n)$ for the whole test: One can sort the edges of  $T'_D(R)$  and then process them as links in increasing cost order, building a binary tree (whose internal nodes represent the edges of  $T'_D(R)$ ) using a suitable auxiliary union-find data structure. This transforms the problem to an instance of the off-line nearestcommon-ancestor problem, which is solvable, for example, in O(q) using a depth-first search strategy [Tar79]. This leads to a total time  $O(q + r \log r)$  for all q inquiries.

For non-terminals  $v_i$  and  $v_j$ , one can use an upper bound for the bottleneck Steiner distance  $s(v_i, v_j)$  considering only paths of the form  $v_i - z_{i,a} - z_{j,b} - v_j$ , where  $z_{i,a}$  and  $z_{i,b}$  are the a-th respectively b-th nearest terminals to  $v_i$  and  $v_j$ . The k (k constant, say 3) nearest terminals to all non-terminals (forbidding intermediary terminals on the corresponding paths) can be computed using a modification of the algorithm of Dijkstra in time O(m + $n \log n$ , as described in [Dui93]. After that, one works with the upper bound  $\hat{s}(v_i, v_j) :=$  $\min_{a,b\in\{1,...,k\}}\{\max\{\ d(v_i, z_{i,a}),\ s(z_{i,a}, z_{j,b}),\ d(z_{j,b}, v_j)\}\}$  instead of  $s(v_i, v_j)$ . But we do not precompute the  $\hat{s}$ -values, because very often not all the  $k^2$  combinations have to be checked; for example if the test condition turns out to be already satisfied during the computation (or, of course, if  $v_i$ or  $v_i$  were a terminal). More importantly, many additional observations can be used to do without  $\hat{s}(v_i, v_j)$  altogether. For example the lower bound  $\tilde{s}(v_i, v_j) := \max\{d(v_i, base(v_i)), d(v_j, base(v_j))\}$ (which is readily available) is often helpful: If both vertices  $v_i$  and  $v_j$  belong to the same Voronoi region, then we simply have  $\tilde{s}(v_i, v_j) = \hat{s}(v_i, v_j)$ . If  $v_i$  and  $v_j$  belong to different Voronoi regions and  $c(v_i, v_j) < \tilde{s}(v_i, v_j)$ , then the test cannot be successful for  $(v_i, v_j)$ . Furthermore, precomputing the  $\hat{s}$ -values (which can need time  $\Theta(n^2)$ ) would destroy the total time  $O(m + n \log n)$  for performing this test on all edges.

An additional lemma leads to a simple, very fast test, which is sometimes very powerful:

**Lemma 20** Let  $\hat{S}$  be the length of the longest edge in  $T'_D(R)$ . Every edge  $(v_i, v_j)$  with  $c(v_i, v_j) > \hat{S}$  can be removed from the network.

**Proof:** Suppose there is a Steiner minimal tree T containing an edge  $(v_i, v_j)$  with  $c_{ij} > \tilde{S}$ . Removing this edge from T divides it into two components:  $C_i$  containing  $v_i$  and  $C_j$  containing  $v_j$ . In each component, there is at least one terminal. Let  $z_k$  and  $z_l$  be two arbitrary terminals in  $C_i$  respectively  $C_j$ . In G, there is a path between  $z_k$  and  $z_l$ , corresponding to the fundamental path in  $T'_D(R)$ , with

Steiner distance at most  $\hat{S}$ . This path contains an elementary path P connecting  $C_i$  and  $C_j$ , whose length is at most  $\hat{S}$ . Reconnecting  $C_i$  and  $C_j$  by P yields a graph spanning all terminals and shorter than T, a contradiction.

Note that using this test, one can eliminate some edges that could not be eliminated by the  $PT_m$  test (even in its original form).

Since the tests above only consider paths with at least one terminal, they miss some of the edges the simple test LE (Long Edges) [Bea84, HRW92] would eliminate. On the other hand, after execution of other tests the graph is often sparse. So a weakened version of LE, which simply searches for shorter paths from both ends of an edge, can be effective. With the additional restriction that during the examination of each edge not more than a constant number of edges are visited in search for an alternative path, one gets the total time  $\Theta(m)$  for this modified test, which we call **Triangle**. This test is sometimes a nice complement to the PT<sub>m</sub> test (as described above), especially if the proportion of terminals to all vertices is not high.

### Improving the Test in the Case of Equality



Figure 3.2: Improvement for  $\bar{s}_{ij} = c_{ij}$ .

The test  $PT_m$  can actually be extended to the case of equality with the restricted bottleneck Steiner distance: An edge  $(v_i, v_j)$  can be removed from G if  $c_{ij} \ge \bar{s}_{ij}$  (remember that  $\bar{s}_{ij}$  is  $s_{ij}$  in G after removing  $(v_i, v_j)$ ). Figure 3.2 shows an example of edges that can be removed using this stronger test condition.

But removing edges with this test condition can change the (restricted) bottleneck Steiner distances, which makes a recalculation of these distances after each deletion necessary. In Figure 3.3, this case is shown: The edge  $(z_2, v_4)$  can be removed, because the restricted bottleneck Steiner distance is  $c(z_2, v_4) = \bar{s}(z_2, v_4) = s(z_2, v_4) = 3$ . Similarly, the edge  $(v_4, z_5)$  can be removed, because  $c(v_4, z_5) = \bar{s}(v_4, z_5)$ . But after removing the edge  $(z_2, v_4)$  the restricted bottleneck Steiner distances change, in particular  $\bar{s}(v_4, z_5) = \infty$ . Thus, only one of the two edges  $(z_2, v_4)$  and  $(v_4, z_5)$  can be deleted. These problems were the reason why in the literature on the Steiner problem it was assumed that the test  $PT_m$  cannot be performed for the case of equality without recalculation of the necessary information.

We have observed that the problematic cases can be efficiently identified, such that in all other cases the test actions can be performed even in case of equality (without recalculation) and using the approximated Steiner bottleneck distances  $\hat{s}$  (which are computed in advance) in the test condition  $c_{ij} \geq \hat{s}_{ij}$ . First, we list the problematic cases for  $\hat{s}_{ij} = c_{ij}$  and show how to deal with them. After that, we prove that other problematic cases cannot exist.



Figure 3.3: Critical cases with  $\bar{s}_{ij} = c_{ij}$ .

I) Edges from  $T'_D(R)$ : Obviously,  $\hat{s}_{ij} = c_{ij}$  holds for every edge  $(z_i, z_j) \in E$ ,  $z_i \in R$ ,  $z_j \in R$  that is contained in the tree  $T'_D(R)$ . In this case, we do not want the edge to be deleted, as we do not know the value of  $\bar{s}_{ij}$ . Thus, edges  $(z_i, z_j) \in E$  with both endpoints being terminals will only be deleted if  $(c_{ij} > \hat{s}_{ij}) \lor (c_{ij} = \hat{s}_{ij} \land (z_i, z_j) \notin T'_D(R))$ .

As an example take the edges  $(z_2, z_5)$  and  $(z_3, z_5)$  in Figure 3.3. Although  $c_{ij} \ge \hat{s}_{ij}$ , the edges will not be deleted, as they are contained in  $T'_D(R)$ .

**II**) Edges between a terminal and a non-terminal: For an edge  $(z_i, v_j) \in E, z_i \in R, v_j \in V \setminus R$ ,  $\hat{s}_{ij}$  can be equal to  $c_{ij}$ , arguing with the path consisting of the edge  $(z_i, v_j)$ . In Figure 3.3, this is shown for the edge  $(z_1, v_4)$ .

Therefore, an edge  $(z_i, v_j)$  will only be deleted, if  $c_{ij} < \hat{s}_{ij}$  or if  $c_{ij} = \hat{s}_{ij}$  and the path from  $v_j$  to  $z_i$  with Steiner distance  $\hat{s}_{ij}$  that was used in the calculation of  $\hat{s}_{ij}$  does not contain the edge. The last condition can be checked easily by maintaining "parent pointers" in the shortest path calculation.

III) Multiple edges to terminals with equal length: If two edges  $(v_i, z_j), (v_i, z_k) \in E$  have the same length and the bottleneck Steiner distance between  $z_j$  and  $z_k$  is not longer than the length of the edges, then any of the edges may be deleted (arguing with a path containing the other edge). But both edges may not be deleted, because after the deletion of the first edge, the argument for the deletion of the other is no longer valid. In Figure 3.3, this holds for the edges  $(v_4, z_2)$  and  $(v_4, z_5)$  as described above.

We have solved this problem in the following way: If an edge is deleted under the circumstances described (e.g.,  $(v_4, z_2)$ ), the other edge (in this case  $(v_4, z_5)$ ) is marked. A marked edge  $(v_i, v_j)$  will only be deleted if  $c_{ij} > \hat{s}_{ij}$ .

Now, we look at the problem from a different point of view. Under which circumstances does the following situation arise: An edge  $e_1 \in E$  has been deleted arguing with a path  $P_1, e_1 \notin P_1$ , with Steiner distance  $\hat{s}(e_1) \leq c(e_1)$ . Then, an edge  $e_2 \in E, e_2 \neq e_1$  is about to be deleted, arguing with a path  $P_2$  of the original network with Steiner distance  $\hat{s}(e_2) \leq c(e_2)$ . But  $e_2$  may not be deleted, because some part of  $P_2$  has been deleted and it is not possible to find a replacement for it in the current network  $G_2$ .

**Lemma 21** The three cases mentioned – denoted by the numbers I), II), and III) – are the only cases where  $c(e_2) \ge \hat{s}(e_2)$  holds for some edge  $e_2$ , but in the current network  $G_2$  it holds  $c(e_2) < \bar{s}(e_2)$ .

**Proof:** We have to distinguish several cases:

1.:  $e_2 \in P_2$ . This is either case I) or II).

2.:  $e_1, e_2 \notin P_2$ . As  $P_2$  is in  $G_2, \hat{s}(e_2) \geq \bar{s}(e_2)$  holds, a contradiction.

 $3: e_2 \notin P_2, e_1 \in P_2 \qquad (\Rightarrow c(e_1) \le \hat{s}(e_2)).$ 

3.1.:  $e_2 \notin P_1$ . We build a new path  $P'_2$  with the same Steiner distance  $\hat{s}(e_2)$  that is contained in  $G_2$ , by replacing  $e_1$  by the path  $P_1$ . It follows that  $\hat{s}(e_2) \geq \bar{s}(e_2)$ , a contradiction.

 $3.2.: e_2 \in P_1 \qquad (\Rightarrow c(e_2) \le \hat{s}(e_1)).$ 

3.2.1.:  $c(e_1) < \hat{s}(e_2)$ . It follows that  $c(e_1) < \hat{s}(e_2) \le c(e_2) \le \hat{s}(e_1) \le c(e_1)$ , a contradiction.

3.2.2.:  $c(e_1) = \hat{s}(e_2)$ . It follows that  $c(e_1) = \hat{s}(e_1) = c(e_2) = \hat{s}(e_2)$ .

3.2.2.1.: Both endpoints of  $e_1$  are terminals. Together with  $e_1 \in P_2$  it follows that  $e_1 \in T'_D(R)$ . This is case I) and  $e_1$  has not been deleted.

3.2.2.2.: Exactly one endpoint of  $e_1$  is a terminal. As  $e_1 \in P_2$  and  $P_2$  has Steiner distance  $\hat{s}(e_2) = c(e_1)$ , one endpoint of  $e_2$  must be the non-terminal endpoint of  $e_1$ . This is case III).

3.2.2.3.: No endpoint of  $e_1$  is a terminal. As  $e_2 \neq e_1$ , the path  $P_2$  must consist of more edges than just  $e_1$ . But then  $P_2$  must have a Steiner distance greater than  $c(e_1)$ , a contradiction.

It is easy to see that this proof also extends to the cases that no edge or more than one edge have been deleted before  $e_2$  is inspected.

It must be mentioned that this strengthening of the reduction test has a greater impact than one would assume, because in some cases the reduction process is blocked in face of many alternatives with equal weights.

## **3.3.2** NTD<sub>k</sub>

The test  $NTD_k$  (Non-Terminals of Degree k) was introduced in [DV89]:

**NTD**<sub>k</sub> test: A non-terminal  $v_i$  has degree at most 2 in at least one Steiner minimal tree if for each set  $\Delta$ ,  $|\Delta| \geq 3$ , of vertices adjacent to  $v_i$  the following holds: The sum of the lengths of the edges between  $v_i$  and vertices in  $\Delta$  is not less than the weight of a minimum spanning tree for the network  $(\Delta, \Delta \times \Delta, s)$ .

If this condition is satisfied, one can remove  $v_i$  and incident edges, introducing for each two vertices  $v_j$  and  $v_k$  adjacent to  $v_i$  an edge  $(v_j, v_k)$  with length  $c_{ij} + c_{ik}$  (and keeping only the shortest edge between each two vertices).

The special cases with k (degree of  $v_i$ ) in  $\{1, 2\}$  can be implemented with total time O(n) (for examination of all non-terminals). For  $k \in \{3, ..., 7\}$  we use the  $\hat{s}$ -values instead of the exact bottleneck Steiner distances, as described in Section 3.3.1. Again, empirically only a marginal difference in effectiveness is observed between the original and the modified version. As before, we do not precompute the  $\hat{s}$ -values, so the modified version has total time  $O(m + n \log n)$ .

Because the addition of new edges can be a nontrivial matter and the necessary  $\hat{s}$ -values are already available, it is a good idea to check whether each new edge could be eliminated using the PT<sub>m</sub> test. In this case it need not be inserted in the first place.

#### **3.3.3** NV and Related Tests

The test NV (Nearest Vertex) is a classical inclusion test [Bea84, HRW92]:

**NV test:** Let  $z_i$  be a terminal with degree at least 2, and let  $(z_i, v'_i)$  and  $(z_i, v''_i)$  be the shortest and second shortest edges incident to  $z_i$ . The edge  $(z_i, v'_i)$  belongs to at least one Steiner minimal tree, if there is a terminal  $z_i, z_i \neq z_i$ , with  $c(z_i, v''_i) \ge c(z_i, v'_i) + d(v'_i, z_i)$ .

#### 3.3. ALTERNATIVE-BASED REDUCTIONS

The original version of the test NV requires the computation of distances, which is too timeconsuming for large instances. But one can accelerate this test without making it less powerful, using the lemma given below. For this purpose, we use Voronoi regions again, saving some extra information while computing the regions. Let  $distance(z_i)$  be the length of a shortest path from  $z_i$ to another terminal  $z_j$  over the edge  $(z_i, v'_i)$ , computed as follows: Each time an edge  $(v_i, v_j)$  with  $v_i \in N(z_i), v_j \in N(z_j), z_j \neq z_i$  is visited, it is checked whether  $v_i$  is a successor of  $v'_i$  in the shortest paths tree with root  $z_i$  (simply done through marking the successors of  $v'_i$ ). In such a case  $distance(z_i)$ is updated to min{ $distance(z_i), d(z_i, v_i) + c(v_i, v_j) + d(v_j, z_j)$ }. Now we have:

**Lemma 22** The condition of the test NV is satisfied if and only if:  $c(z_i, v_i'') \ge c(z_i, v_i') + d(v_i', base(v_i'))$ , if  $v_i' \notin N(z_i)$ , and  $c(z_i, v_i'') \ge distance(z_i)$ , if  $v_i' \in N(z_i)$ .

**Proof:** Assume the condition formulated in the lemma is satisfied for a vertex  $z_i$ : If  $v'_i \notin N(z_i)$ , the NV test condition is satisfied for  $z_j = base(v'_i)$ . If  $v'_i \in N(z_i)$ , then a terminal  $z_j$  exists with  $c(z_i, v'_i) + d(v'_i, z_j) = distance(z_i) \leq c(z_i, v''_i)$ . Hence, the NV test condition is satisfied.

Now assume that the condition of the test NV,  $c(z_i, v_i'') \ge c(z_i, v_i') + d(v_i', z_j)$ , is satisfied: If  $v_i' \notin N(z_i)$ , it follows from  $d(v_i', z_j) \ge d(v_i', base(v_i'))$  that  $c(z_i, v_i'') \ge c(z_i, v_i') + d(v_i', base(v_i'))$ . If  $v_i' \in N(z_i)$ , we could get  $c(z_i, v_i') + d(v_i', z_j) \ge distance(z_i)$ , assuming that  $v_i'$  is on a shortest path between  $z_i$  and  $z_j$ . But the latter must be true, because otherwise we have  $c(z_i, v_i'') \ge c(z_i, v_i') + d(v_i', z_j) \ge d(v_i', z_j) \ge d(v_i', z_j) \ge d(z_i, z_j) \ge c(z_i, v_i'')$ , a contradiction.

Using this lemma, the test NV can be performed for all terminals in time  $O(m + n \log n)$ . Note that in inclusion tests, each included edge is contracted into a terminal.

The Voronoi regions can also be used to perform a related inclusion test, which we call **SL** (standing for Short Links):

**Lemma 23** Let  $z_i$  be a terminal, and  $(v_1, v'_1)$  and  $(v_2, v'_2)$  the shortest and second shortest edges that leave the Voronoi region of  $z_i$   $(v_1, v_2 \in N(z_i), v'_1, v'_2 \notin N(z_i)$ ; we call such edges **links**). The edge  $(v_1, v'_1)$  belongs to at least one Steiner minimal tree, if  $c(v_2, v'_2) \ge d(z_i, v_1) + c(v_1, v'_1) + d(v'_1, z_j)$ , where  $z_j = base(v'_1)$ .

**Proof:** Suppose that the edge  $(v_1, v'_1)$  is not in any Steiner minimal tree. Consider such a tree T and the path between  $z_i$  and  $z_j$  in T. An edge on this path must leave the Voronoi region of  $z_i$ . Removing this edge and inserting  $(v_1, v'_1)$  and two shortest paths to  $z_i$  and to  $z_j$ , we get a subgraph that includes  $(v_1, v'_1)$ , spans all terminals and is no longer than T, a contradiction.

This test can also be performed for all terminals in total time  $O(m + n \log n)$ .

The classical test SE (Short Edges) [DV89, HRW92] is a more powerful inclusion test. We have observed that even this test can be implemented with time  $O(m + n \log n)$ . But although this test is more effective than NV and SL in a single application, the difference almost vanishes when the reduction tests are iterated. Therefore, we only use the much simpler, empirically faster tests NV and SL in our actual implementations.

## **3.3.4** Path Substitution (PS)

We have designed another alternative-based reduction test that is more general than the previous tests in two ways: The test PS examines several edges along a path, instead of examining elementary graph objects (like single edges and vertices). If the test is successful, some of these edges can be deleted at the same time. The other more general aspect is a consequence of the first: Searching for alternatives



Figure 3.4: Example for the application of the NV- and SL-tests.

for a path, it is no longer sufficient to find *one* alternative, because the edges of the path can be involved in many different ways in a Steiner tree. As a consequence, such a test can only be efficient if it has a rather restricted test condition.

The basic idea is to start with a single edge as the path and then try to find alternative paths for the vertices adjacent to those on the path. If this is not possible for exactly one adjacent vertex, the path is extended by the edge to this vertex and the search for alternative paths is restarted. Such successive extensions could finally lead to the desired situation.

We describe the lemma that leads to the formal specification of the test in a simplified way: We give only the description for deleting *one* edge of the path and define it only for the special case that the starting vertex  $v_0$  has degree 3. The extensions to deleting many edges on the path and to vertices with higher degree are not too complicated. (For  $degree(v_0) > 3$  the additional condition  $d_P(v_0, v_i) \ge d_0(v_0^{k_0}, v_i^k) + c(v_i, v_i^k)$  is required in the last line of the definition of the test condition.)

**Lemma 24** Let P be a path  $(v_0, \ldots, v_l)$  with  $degree(v_0) = 3$  and  $v_i \in V \setminus R$  for all  $i \in \{0, \ldots, l\}$ . We denote by  $v_i^1, v_i^2, \ldots$  the vertices adjacent to each  $v_i$  on P that are not contained in P. Let  $d_0(v_i, v_j)$ be the length of a shortest path between  $v_i$  and  $v_j$  that does not contain  $(v_0, v_1)$ , and  $d_P(v_i, v_j)$  (for  $v_i$ and  $v_i$  in P) the length of the subpath of P between  $v_i$  and  $v_j$ .

The edge  $(v_0, v_1)$  can be deleted if for all  $i \in \{1, \ldots, l\}$  there are functions  $f^i$  and  $g^i$  such that: I) for all  $v_i^k$  adjacent to  $v_i$  and for  $k_0 = f^i(k)$ :  $d_P(v_0, v_i) \ge d_0(v_0^{k_0}, v_i^k)$ , II) for all  $v_0^{k_0}$  adjacent to  $v_0$  and for  $k = g^i(k_0)$ :  $d_P(v_0, v_i) \ge d_0(v_0^{k_0}, v_i^k)$ ,  $c(v_0, v_0^{k_0}) \ge c(v_i, v_i^k)$ .

**Proof:** Suppose all Steiner minimal trees contain the edge  $(v_0, v_1)$ . Consider such a tree T, and let  $t \geq 1$  be the smallest index such that there is an edge  $(v_t^k, v_t)$  in T. Notice that the degree of  $v_0$  in T must be greater than 1 and that all edges between  $v_0$  and  $v_t$  on P must be in T. There are two cases: 1) In T,  $v_0$  has degree three. Choose k such that  $(v_t^k, v_t)$  is in T. Let  $k_0 = f^t(k)$ . Remove the edges on the path  $(v_0, v_1, \ldots, v_{t-1}, v_t)$  from T. The resulting components can be reconnected without reinserting  $(v_0, v_1)$  by a path between  $v_0^{k_0}$  and  $v_t^k$  which is not longer.

2) In T,  $v_0$  has degree two. Choose  $k_0$  such that  $(v_0^{k_0}, v_0)$  is in T. Let  $k = g^t(k_0)$ . Remove the edges on the path  $(v_0^{k_0}, v_0, v_1, \dots, v_{t-1}, v_t)$  from T. The resulting components can be reconnected without reinserting  $(v_0, v_1)$  by a path between  $v_0^{k_0}$  and  $v_t^k$  and the edge  $(v_t^k, v_t)$ . Again the inserted edges together are not longer than the removed edges.

In both cases, we have a subgraph that does not contain  $(v_0, v_1)$ , spans all terminals and is not longer than T, a contradiction.  $\square$ 

One problem for an efficient implementation of this test is the calculation of the distances  $d_0(v_i, v_j)$ . Since we do not want to have running times like  $\Theta(n^3)$  for the calculation of shortest paths, we work with a weakened version: To determine an upper bound for  $d_0(v_0^{k_0}, v_t^k)$ , we examine only those paths that contain only vertices in  $\{v_{t'}^{k'} \mid 0 \leq t' \leq t\}$ . This makes it easy to maintain shortest paths trees for each  $v_0^i, i \in \{1, \ldots, degree(v_0) - 1\}$ , during the successive extensions of P. It is also possible to determine up to which vertex  $v_s$  in P the edge  $(v_s, v_{s+1})$  can be deleted, under the assumption that all edges between  $v_0$  and  $v_s$  have been deleted. If finally a situation is reached in which – according to the lemma above –  $(v_0, v_1)$  can be deleted, then all edges of P between  $v_0$  and  $v_{s+1}$  can be removed. Our implementation assures that each edge is considered as a part of P not more than twice (once in each direction). We have observed that if the test is successful, all involved vertices have low degrees. If one fixes a small constant g, e.g. g = 10, and aborts the successive extension of P each time a vertex with degree larger than g is visited, a total running time (for the whole network) of O(m) can be guaranteed, without impeding its reduction potential noticeably.

This version of the test is usually effective only for some sparse graphs (including some VLSIinstances). For such instances, 5-10% of edges could frequently be removed using this test alone.

## **3.4 Bound-based Reductions**

Since one cannot expect to solve all instances of an  $\mathcal{NP}$ -hard problem like the Steiner problem only through reduction tests with a (low order) polynomial worst-case time (like the tests in the previous subsection), the computation of (sharp) lower bounds is a common part of the standard algorithms for the exact solution of such a problem. But the information gained during such computations can be used to reduce the instance further; and sometimes small worst-case running times can be guaranteed even for this kind of tests.

#### 3.4.1 Using Voronoi Regions

The Voronoi regions can be used to determine a lower bound for the value of an optimal solution with additional constraints (for example, that the solution contains a certain non-terminal). For any terminal z, we define radius(z) as the length of a shortest path from z leaving its Voronoi region N(z). These values can be easily determined while computing the Voronoi regions. For convenience, we assume here that the terminals are numbered according to non-decreasing *radius*-values. For each non-terminal  $v_i$ , let  $z_{i,1}, z_{i,2}$  and  $z_{i,3}$  be the three terminals next to  $v_i$ , as described in Section 3.3.1. The following lemma can be used to eliminate a non-terminal.

**Lemma 25** Let T be a Steiner minimal tree and assume that  $v_i$  is a Steiner node in T. Then  $d(v_i, z_{i,1}) + d(v_i, z_{i,2}) + \sum_{t=1}^{r-2} radius(z_t)$  is a lower bound for the weight of T.

**Proof:** For each terminal  $z_l$ , we denote the path between  $z_l$  and  $v_i$  in T with  $P_l$ . Among such paths, there must be at least two (edge-)disjoint ones. For any path P, define  $\Delta(P)$  as the number of edges on P that have their vertices in two different Voronoi regions. Let  $P_j$  and  $P_k$  be two disjoint paths such that  $\Delta(P_j) + \Delta(P_k)$  is minimal. Note that no path  $P_l$  can have edges in common with both  $P_j$  and  $P_k$ . For each terminal  $z_l \notin \{z_j, z_k\}$ , let  $P'_l$  be the part of  $P_l$  from  $z_l$  up to the first vertex not in  $N(z_l)$ ;  $P'_l$  is well-defined, because otherwise  $P_l$  would be the only path with  $\Delta(P_l) = 0$  (namely for  $z_l = base(v_i)$ ) and would have been chosen as  $P_j$  or  $P_k$ . Obviously, all  $P'_l$  are disjoint. Now suppose that  $P_j$  has an edge in common with some  $P'_l$ . Let  $v_l$  be a vertex of this edge with  $v_l \in N(z_l)$ . The part of  $P_j$  between  $z_j$  and  $v_l$  contains an edge with only one vertex in  $N(z_l)$ , so  $\Delta(P_l) < \Delta(P_j)$ ,

which contradicts the choice of  $P_j$ . So  $P_j$  (or, similarly,  $P_k$ ) has no edge in common with a path  $P'_l$ . Since  $P_j$ ,  $P_k$  and the r-2 paths  $P'_l$  are all disjoint, the sum of their lengths cannot be larger than the weight of T. The sum of the lengths of  $P_j$  and  $P_k$  is at least  $d(v_i, z_{i,1}) + d(v_i, z_{i,2})$ . The sum of the lengths of the r-2 paths  $P'_l$  is at least  $\sum_{t=1}^{r-2} radius(z_t)$ .

A non-terminal  $v_i$  can be eliminated if this lower bound exceeds a known upper bound. This method can be extended for eliminating edges:

**Lemma 26** Let T be a Steiner minimal tree and assume that T contains an edge  $(v_i, v_j)$ . Then  $c(v_i, v_j) + d(v_i, z_{i,1}) + d(v_j, z_{j,1}) + \sum_{t=1}^{r-2} radius(z_t)$  is a lower bound for the weight of T.

Proof: Analogous to the proof of Lemma 25.

One can also define a test performing the same actions as  $NTD_k$  when it is successful, using the following lemma:

**Lemma 27** Let T be a Steiner minimal tree and assume that  $v_i$  is a Steiner node whose degree in T is at least three. Then  $d(v_i, z_{i,1}) + d(v_i, z_{i,2}) + d(v_i, z_{i,3}) + \sum_{t=1}^{r-3} radius(z_t)$  is a lower bound for the weight of T.

Proof: Analogous to the proof of Lemma 25.

Intuitively, one expects that an even better lower bound should be achievable through this line of argument, because the paths between the terminals in a Steiner tree not only leave the corresponding Voronoi regions, but also span all terminals. Indeed, one can use this idea:

**Lemma 28** Consider the auxiliary network G' = (R, E', d'), in which two terminals are adjacent if and only if they are neighbors in the original network, defining:

 $d'(z_i, z_j) := \min\{\min\{d(z_i, v_i), d(z_j, v_j)\} + c(v_i, v_j) \mid v_i \in N(z_i), v_j \in N(z_j)\}.$ 

The weight of a minimum spanning tree for G' is a lower bound for the weight of any Steiner tree for the original instance (G, R).

**Proof:** We will prove the lemma by transforming a Steiner minimal tree  $T_G(R)$  into a spanning tree T' in G' without increasing the cost. For guiding this transformation we construct an auxiliary tree T'' by contracting all edges of  $T_G(R)$  that are entirely in one Voronoi region. We consider T'' as a rooted tree with an arbitrary root  $z_r$ . Beginning with isolated terminals as T', each step of the transformation removes the path from one leaf of T'' to its parent and inserts an edge of G' into T'. Throughout the transformation the following invariant ( $\dagger$ ) holds: In each component of T', there is exactly one terminal that has not been removed from T''. In the beginning ( $\dagger$ ) holds trivially.

Each step of the transformation is performed as follows: Choose any leaf  $z_i$  of T'' such that all vertices  $v_i \in N(z_i) \setminus \{z_i\}$  have at most one successor in T''. Notice that there is always such a  $z_i$  in T'', because the number of leaves is greater than the number of non-terminals with more than one successor. From  $z_i$  we move in the direction of the root until we reach a terminal  $z_l$ . The path from  $z_i$  to  $z_l$  in T'' is denoted by  $P''_i$ . The corresponding path in  $T_G(R)$  is denoted by  $P_i$ . Now we look at the bases of the vertices on  $P_i$ . Let  $v_j$  be the last vertex on  $P_i$  whose base  $z_j$  is connected to  $z_i$  in T', and  $v_k$  the first whose base  $z_k$  is not connected to  $z_i$  in T'. The invariant  $(\dagger)$  guarantees that such vertices  $v_j$  and  $v_k$  exist, because not all bases  $z_i, \ldots, z_l$  of the vertices of  $P_i$  can belong to the same component of T', since  $z_i$  and  $z_l$  have not been removed from T''. We denote with  $P'_i$  the part of  $P_i$  between  $z_i$  and  $v_k$ . The length of  $P'_i$  is at least  $d'(z_j, z_k)$ , because  $d'(z_j, z_k) \leq d(z_j, v_j) + c(v_j, v_k) \leq d(z_i, v_j) + c(v_j, v_k)$ . Remove the subpath of  $P''_i$  beginning from  $z_i$  until a vertex in T'' with degree greater than 2 or  $z_l$  is reached. The edge  $(z_j, z_k)$  is inserted in T', so  $(\dagger)$  remains valid.

After all terminals except  $z_r$  have been removed from T'', r-1 edges have been inserted into T' without creating a cycle, so T' is a spanning tree at the end.

We show now that each two paths  $P'_a$  and  $P'_b$  corresponding to terminals  $z_a$  and  $z_b$  are edge-disjoint (in the following simply denoted as disjoint). There are two cases:

I)  $z_a$  is a successor of  $z_b$  in T'' (or vice versa): If there are common edges in  $P'_a$  and  $P'_b$ , there must be common edges or (if they are contracted) at least common vertices in  $P''_a$  and  $P''_b$ . But the paths  $P''_a$  and  $P''_b$  are disjoint and have at most one vertex in common, namely  $z_b$ . Hence, common edges of  $P'_a$  and  $P'_b$  must lie entirely inside the Voronoi region of  $z_b$  and have been contracted in T'' into  $z_b$ . When  $z_a$  is chosen, neither  $z_a$  nor  $z_b$  has been removed from T''. Because of (†), at this time  $z_a$  and  $z_b$  are not connected in T'. So, for  $P_a$  the vertex  $v_j$  (as defined above) must be outside  $N(z_b)$ , and there is no edge in  $P'_a$  that is entirely inside  $N(z_b)$ . Therefore, the paths are disjoint.

II)  $z_a$  and  $z_b$  are successors of a terminal  $z_c$  in T'': Assume that  $z_a$  is chosen before  $z_b$ . There are disjoint paths in  $T_G(R)$  from a vertex  $v_d$  to  $z_a, z_b$  and  $z_c$  ( $v_d = z_c$  is possible). Let  $z_d := base(v_d)$ . Suppose that the paths  $P'_a$  and  $P'_b$  are not disjoint. So, they must both contain edges of the path between  $v_d$  and  $z_c$ . Thus, when  $z_a$  is chosen,  $z_d$  must be connected to  $z_a$  in T'. Because  $z_a$  has not been removed from T'', it follows from ( $\dagger$ ) that  $z_d$  was removed from T'' before. This is only possible if  $v_d \neq z_d$ . When  $z_d$  was chosen,  $v_d$  had only one successor in T''. Thus, the path between  $z_d$  and  $z_b$  has been removed even sooner. This means that  $z_b$  had to be chosen before  $z_a$ , a contradiction.

Since each edge of T' corresponds to a path in  $T_G(R)$  with at least the same length and all these paths are disjoint, the spanning tree T' is not longer than  $T_G(R)$ , and the lemma follows.<sup>1</sup>

This lemma can be extended to a test condition; for example, for any non-terminal  $v_i$ , the weight of such a spanning tree minus the length of its longest edge plus  $d(v_i, z_{i,1}) + d(v_i, z_{i,2})$  is a lower bound for the weight of any Steiner minimal tree that contains  $v_i$ . The resulting test is very fast: The network G' can be determined without much extra work while computing the Voronoi regions, and a minimum spanning tree for it can be computed in time  $O(m + r \log r)$ .

For computing upper bounds in this context, we use a modified path heuristic with time  $O(m + n \log n)$ , which is described in Section 4.2. So, all these tests can be performed in time  $O(m+n \log n)$ ; we call this combined test **VR** (standing for Voronoi Regions). With a heuristic solution available, all these tests can be easily extended to the case of equality of lower and upper bound. As intuition suggests, the VR test is most effective for sparse networks with relatively few terminals; in this sense, it is a nice complement to the alternative-based tests, which are often especially successful if the proportion of terminals to all vertices is high. Additionally, this test was the basis for the development of the strong PRUNE-heuristics, which are presented in Section 4.3.

## 3.4.2 Using Dual Ascent

The information provided by the algorithm DUAL-ASCENT (section 2.9.2), namely the lower bound with value *lower* and the reduced costs can be used to design another bound-based reduction test. Here we use a simple, yet very helpful lemma, which we will exploit frequently later on:

**Lemma 29** Let G = (V, A, c) be a (directed) network (with a given set of terminals) and  $\tilde{c} \leq c$ . Let *lower'* be a lower bound for the value of any (directed) Steiner tree in G' = (V, A, c') with  $c' := c - \tilde{c}$ . For each  $\tilde{x}$  representing a feasible Steiner tree for G with  $\cot c \cdot \tilde{x}$ , it holds that  $lower' + \tilde{c} \cdot \tilde{x} \leq c \cdot \tilde{x}$ .

**Proof:**  $c \cdot \tilde{x} = c' \cdot \tilde{x} + \tilde{c} \cdot \tilde{x} \ge lower' + \tilde{c} \cdot \tilde{x}$ .

<sup>&</sup>lt;sup>1</sup>In [PV00], we gave an easier proof that used the equivalence between this lower bound and the lower bound produced by some primal dual algorithm for  $LP_C$ .

Now consider the reduced costs provided by DUAL-ASCENT as  $\tilde{c}$ : One can observe that the lower bound *lower'* provided by DUAL-ASCENT in G' is the same as *lower*. So for any  $\tilde{x}$  representing a feasible (directed) Steiner tree  $\vec{T}$ , *lower* +  $\tilde{c} \cdot \tilde{x}$  represents a lower bound on the weight of  $\vec{T}$ .

This lemma can be used to compute lower bounds for the value of an optimal Steiner tree with additional constraints, for example, that the tree contains a certain non-terminal. The resulting tests are basically identical to the tests IRA and IRAe, which are introduced in [Dui93], using a somewhat more tedious argumentation.

Let  $v_k$  be a non-terminal, and  $\vec{T}$  any optimal (directed) Steiner tree containing  $v_k$ , represented by  $\tilde{x}$ . The lower bound  $\tilde{c} \cdot \tilde{x}$  on the weight of  $\vec{T}$  minus *lower* can be further estimated from below by the length of a shortest path (with respect to the costs  $\tilde{c}$ ) from the root to  $v_k$  plus the length of an (arc-disjoint) shortest path from  $v_k$  to another terminal; and the last value can be again estimated from below by the distance of  $v_k$  to its nearest terminal, as described in Section 3.3.1. The non-terminal  $v_k$  can be eliminated if this lower bound exceeds a known upper bound. Similar tests can be developed for the elimination of edges and for the elimination of vertices after replacing incident edges (as in NTD<sub>k</sub>). All these tests can be performed in time  $O(m + n \log n)$  after a run of DUAL-ASCENT (and computation of an upper bound). With a heuristic solution available, these tests can be easily extended to the case of equality of lower and upper bound. We call this collection of tests **DA** (standing for Dual Ascent).

When dealing with the Steiner problem in undirected networks, it is a good idea to try different terminals as the root. Although the optimal value  $DLP_C$  is independent of this choice, the value of the lower bound provided by DUAL-ASCENT is not, and, much more important, different roots can lead to the elimination of different parts of the network, even if the value of the lower bound does not change. Trying a constant number (at most 10) of terminals as roots, we have substantially improved the effectiveness of this test. Notice also that each repetition profits from the reductions achieved by the previous ones.

The test DA is very effective, and usually it is fast empirically. But the time bound  $O(a \cdot \min\{a, rn\})$  (resulting from DUAL-ASCENT) is, in comparison to the time  $O(m + n \log n)$  of the other tests hitherto presented, somewhat unsatisfactory, especially because the other parts of the test can indeed be performed in time  $O(m + n \log n)$ .

One can try to achieve a better time bound by using a faster dual ascent algorithm, even if the provided lower bounds are worse: The tests described above use both the reduced costs and the lower bound, and a worse lower bound can be compensated to some degree by larger reduced costs.

One successful variant with running time  $O(m + n \log n)$  uses the observation that it is possible to increase many dual variables around a terminal simultaneously.

**Lemma 30** Choose a terminal  $z_t \in R^{z_1}$ . Define  $d'(v_i) := \min\{d(v_i, z_t), d(z_1, z_t)\}$ . For all Steiner cuts  $\{\bar{W}, W\}$  set the dual variable  $u_W := \max\{0, \min_{v_j \notin W}\{d'(v_j)\} - \max_{v_j \in W}\{d'(v_j)\}\}$ . Then  $\sum_{W, [v_a, v_b] \in \delta^-(W)} u_W = \max\{0, d'(v_a) - d'(v_b)\} \le c_{ab}$  for all edges  $[v_a, v_b] \in A$ .

**Proof:** Let  $v_1, v_2, \ldots, v_n$  be the vertices of V sorted by their distances to  $z_t$  in ascending order. Consider a Steiner cut  $\{\overline{W}, W\}$ . Obviously  $u_W = \max\{0, d'(v_h) - d'(v_i)\}$  for  $h = \min\{j \mid v_j \notin W\}$ ,  $i = \max\{j \mid v_j \in W\}$ . If there are two vertices  $v_h$  and  $v_i$  with  $h < i, v_h \notin W$ , and  $v_i \in W$ , then  $u_W = 0$ . So if  $u_W > 0$ , there must be a vertex  $v_k$  with  $v_l \in W$  for all  $l \le k$  and  $v_l \notin W$  for all l > k; so we can denote W by  $W_k$ :  $W_k = \{v_1, \ldots, v_k\}$ ;  $u_{W_k} = d'(v_{k+1}) - d'(v_k)$ . For any edge  $[v_a, v_b]$  we have:

$$\sum_{\substack{W, [v_a, v_b] \in \delta^-(W) \\ b \leq k < a}} u_W = \sum_{\substack{b \leq k < a \\ b < k < a}} u_{W_k} = \max\{0, d'(v_a) - d'(v_b)\} = \max\{0, d'(v_a) - d'(v_b)\} = 0$$

#### 3.4. BOUND-BASED REDUCTIONS

$$\max\{0, \min\{d(v_a, z_t), d(z_1, z_t)\} - \min\{d(v_b, z_t), d(z_1, z_t)\}\} \le \max\{0, \min\{c_{ab} + d(v_b, z_t), d(z_1, z_t) + c_{ab}\} - \min\{d(v_b, z_t), d(z_1, z_t)\}\} = c_{ab}.$$

It follows immediately that u is feasible for  $DLP_C$ . Since the dual variables u are not used explicitly in the reduction process, it is sufficient to work with the reduced costs and the calculated lower bound; so the updating process for one terminal can be performed very quickly, because we just need a shortest paths tree rooted at  $z_t$  that spans  $z_1$ . Then the reduced costs for an edge  $[v_a, v_b]$  are decreased by  $\max\{0, d'(v_a) - d'(v_b)\}$  and the lower bound is increased appropriately. After each such updating there may still be terminals that are not reachable from the root by edges of zero reduced costs, so the updating can be repeated with other terminals, but then with respect to the remaining reduced costs. We guide this calculation by the structure of a heuristic solution: The terminals are sorted according to non-decreasing distances from the root in this solution and considered one at a time.

Note that using this method, an edge can be visited by several terminals. To limit the effort, we simply abort the calculation of a shortest paths tree if it reaches a vertex that has already been visited by a constant number of terminals (e.g., 5). This leads to a worst-case running time for the calculation of a lower bound and reduced costs of  $O(m + n \log n)$ . The other operations of the test can be performed in the same time, as previously described. To construct a heuristic solution, we use a heuristic described in Section 4.2, which has the same running time. So the whole test can be performed in total time  $O(m + n \log n)$ . We call this test **LDA** (Limited Dual Ascent). Despite its low running time, it is fairly effective, especially if the proportion of terminals to all vertices is not very high.

#### 3.4.3 Using the Row Generation Strategy

The modification above aimed at making the reduction technique based on reduced costs faster. A legitimate question is if it is possible to make that technique stronger. For this, we use the row generation method described in Section 2.9.2.

Every iteration of the row generation method provides a dual feasible solution for  $LP_C$  (or  $LP_{C'}$ ) and appropriate reduced costs. Using this information, the same reduction techniques as described in Section 3.4.2 can be used. The only enhancement here is that edges are allowed to be deleted only in one direction during the row generation process (remember that the relaxations  $LP_C$  and  $LP_{C'}$  use directed networks). This can amplify the effect of subsequent reductions considerably. In the linear program itself, the deletion of edges is realized by fixing the corresponding variables to zero.

In many cases the mentioned reductions during the row generation make further alternative-based reductions possible. But it would be a bad idea to delay these reductions until the row generation terminates, because they could possibly accelerate the computation and raise the optimal value of the relaxation. On the other hand, it would be problematic to abort the row generation, do the alternative-based reductions and then start it again, because the constraints generated in the meantime could not be used anymore, at least not directly. Our approach for dealing with this problem is to perform alternative-based reductions in an undirected copy of the current directed instance (which is not necessarily bidirected). After that, the reduced undirected instance is translated back into a directed instance, with the performed reductions translated into fixing of variables. We call the whole reduction method **RG** (for Row Generation).

The row generation approach can be exploited for even stronger reductions in combination with the extended reduction techniques that will be described in the next sections.

Note that using a lower bound and reduced costs produced by an LP-solver in the context of reduction techniques is a delicate thing. As the LP-solver works with floating point arithmetic and gives hardly any guarantee for the quality of the returned solution, the output of the LP-solver cannot

be used directly. One reliable and very efficient solution is the use of fix point integer arithmetic. The floating point numbers are scaled by some factor and rounded to integers.

In this process, we distinguish between the primal and the dual variables. The primal variables are less critical, as they are only used for the computation of new constraints, e.g., by minimum cut computations. Note that the preflow-push algorithm used only works reliably using exact arithmetic. As we always round up while transforming the primal variables, we make it less probable, yet not impossible, to find an already satisfied cut constraint that has a capacity less then one according to the rounded capacity values. Also, it is possible that we miss some violated cut constraints with a capacity very close to one. Neither of the two situations is really harmful. As primal variables are always between zero and one, we can choose a very high factor for the scaling.

The dual side is more interesting. The basic idea is to round the dual variables to fixed point numbers and compute a valid lower bound and reduced costs in exact arithmetic. A minor issue is that we have to store the integer constraint matrix A of the linear program. The problem is that the dual variables may not be feasible any more. Here we can use a nice extension of Lemma 29.

Lemma 31 For any linear program

 $\min\{c \cdot x \mid Ax \ge b; \ 1 \ge x \ge 0\} = \max\{b \cdot \lambda - 1 \cdot \mu \mid A^T \lambda - \mu \le c; \ \lambda \ge 0; \ \mu \ge 0\},$ 

any (feasible or infeasible) dual values  $\bar{\lambda} \ge 0$ ,  $\bar{\mu} \ge 0$  and any feasible primal values  $\tilde{x}$  it holds that:

$$b \cdot \bar{\lambda} - 1 \cdot \bar{\mu} + (c + \bar{\mu} - A^T \bar{\lambda}) \cdot \tilde{x} \le c \cdot \tilde{x}.$$

**Proof:** Let  $x^+$  be an optimal solution for the linear program  $\min\{c^+ \cdot x \mid Ax \ge b; x \ge 0\}$  with  $c^+ = A^T \overline{\lambda}$ . Note that  $\tilde{x}$  is feasible for this linear program, and  $\overline{\lambda}$  is feasible for the dual of the program. It follows that  $b \cdot \overline{\lambda} \le c^+ \cdot x^+ \le c^+ \cdot \tilde{x}$ . As  $\tilde{x} \le 1$ , it follows that  $\overline{\mu} \cdot \tilde{x} \le 1 \cdot \overline{\mu}$ . Both inequalities together prove the claim.  $\Box$ 

**Corollary 31.1** If  $\bar{\mu} \ge A^T \bar{\lambda} - c$ , it follows that for any primal feasible  $\tilde{x}, b \cdot \bar{\lambda} - 1 \cdot \bar{\mu}$  is a lower bound for  $c \cdot \tilde{x}$ .

To use the inexact dual variables in the context of bound-based reductions, we do the following:

- 1. We round the dual variables to fixed point numbers  $\overline{\lambda}$ . We choose the scaling factor such that the current difference between upper and lower bound corresponds to a fixed point number *rest* that uses the nearly full bit length of the computer's integers. We can ensure that all numbers that are computed in the bound-based reductions are not greater than 2 *rest* + 1. On the one hand, this presents overflows. On the other, the accuracy is increased in the row generation iterations automatically, as the difference between upper and lower bound decreases.
- 2. We set any negative  $\bar{\lambda}_i$  to zero.
- 3. We try a "test and repair" strategy: If  $A^T \overline{\lambda} \leq c$  is violated for some arc  $[v_i, v_j]$ , we try to shift the dual variables associated to constraints containing  $x_{ij}$  a little so that the violation gets smaller. This done in a heuristical way.
- 4. Using exact arithmetic, we compute  $\bar{\mu} := \max\{0, A^T \bar{\lambda} c\}$ . From Corollary 31.1 it follows that *lower* :=  $b \cdot \bar{\lambda} 1 \cdot \bar{\mu}$  is an exact lower bound for the value of an optimal Steiner tree.
- 5. We compute reduced costs  $\tilde{c} := c + \bar{\mu} A^T \bar{\lambda}$ . Note that  $\tilde{c} \ge 0$ .
- 6. Now, we can perform bound-based reductions as described in the previous section with *lower* and  $\tilde{c}$ : *lower* +  $\tilde{c} \cdot \tilde{x}$  is a lower bound for the value of a Steiner tree represented by  $\tilde{x}$ .

# 3.5 Extended Reduction Techniques: Combining Alternative- and Bound-based Approaches

The classical reduction tests just consider single vertices or edges. Recent and more sophisticated tests extend the scope of inspection to more general patterns. In the next sections, we present such an extended reduction test, which generalizes various tests from the literature. We use the new approach of combining alternative- and bound-based methods, which substantially improves the impact of the tests. We also present several algorithmic contributions. The experimental results show a large improvement over previous methods using the idea of extension, leading to a drastic speed-up in the optimal solution process and the solution of several previously unsolved benchmark instances.

## 3.5.1 Additional Definitions for Extended Reduction Techniques

For every tree T in G, we denote by V(T) the vertices of T, by L(T) the leaves of T, and by c(T) the sum of the costs of edges in T. Let T' be a subtree of T. The **linking set** between T and T' is the set of vertices  $v_i \in V(T')$  such that there is a fundamental path from  $v_i$  to a leaf of T not containing any edge of T'. Note that the paths can have zero length and if a leaf of T' is also a leaf of T it will be in the linking set. If the linking set between T and T' is equal to L(T'), T' is said to be **peripherally contained** in T (Figure 3.5). This means that for every leave  $v_j$  of T the fundamental path connecting  $v_j$  to T' ends in a leaf of T'. A set  $L' \subseteq V(T)$ , |L'| > 1, induces a subtree  $T_{L'}$  of T containing for every two vertices  $v_i, v_j \in L'$  the fundamental path between  $v_i$  and  $v_j$  in T. We define L' to be a **pruning set** if L' contains the linking set between T and  $T_{L'}$ .



For the depicted tree T, let T' be the subtree of T after removing the edges  $(v_4, v_6)$ and  $(v_6, v_7)$ . The linking set between T and T' is  $\{v_1, v_2, v_4, v_5\}$ , and therefore T' is not peripherally contained in T. But if we add  $(v_4, v_6)$  to T', it is.

Figure 3.5: Depiction of some central notions for extended reductions.

#### 3.5.2 Extending Reduction Tests

The classical reduction tests for the Steiner problem inspect only simple patterns (a single vertex or a single edge). There have been some approaches in the literature for extending the scope of inspection [Dui00, UdAR99, Win95]. The following function EXTENDED-TEST describes in pseudocode a general framework for many of these approaches. The argument of EXTENDED-TEST is a tree T that is expanded recursively. For example, to eliminate an edge e, T is initialized with e. The function returns 1 if the test is successful, i.e., it is established that there is an optimal Steiner tree that does not peripherally contain T.

In the pseudocode, the function RULE-OUT(T, L) contains the specific test conditions (see Section 3.5.3): RULE-OUT(T, L) returns 1 if it is established that T is not contained with linking set L in at least one optimal Steiner tree. The function TRUNCATE checks some criterion to truncate the recursive expansion, and PROMISING tries to identify promising candidates for expansion.

```
EXTENDED-TEST(T):
    (returns 1 only if T is not contained peripherally in an optimal Steiner tree)
1
       if RULE-OUT(T, L(T)):
\mathcal{2}
            return 1
                                (test successful)
\mathcal{B}
       if TRUNCATE(T):
            return 0
                                (test truncated)
4
5
       forall leaves v_i of T:
            if v_i \notin R and PROMISING(v_i):
6
\tilde{7}
               success := 1
8
              forall nonempty extension \subseteq \{(v_i, v_j) :
                      not RULE-OUT(T \cup \{(v_i, v_j)\}, L(T) \cup \{v_j\})\}:
                   if not EXTENDED-TEST(T \cup extension):
g
10
                      success := 0
11
              if success :
12
                                        (no acceptable extension at v_i)
                   return 1
13
                        (in all inspected cases, there was an acceptable extension)
       return 0
```

Assuming that RULE-OUT is correct, the correctness of EXTENDED-TEST can be proven easily by induction, using the fact that if T' is a subtree of an optimal Steiner tree  $T^*$  and contains no inner terminals, all leaves of T' are connected to some terminal by paths in  $T^* \setminus T'$ .

Clearly the decisive factor for the performance of this algorithm is the realization of the functions *RULE-OUT*, *TRUNCATE* and *PROMISING*.

Using this framework, previous extension approaches can be outlined easily:

- In [Win95] the idea of expansion was introduced for the rectilinear Steiner problem.
- In [UdAR99] this idea was adopted to the Steiner problem in networks. This variant of the test tries to replace vertices with degree three; if this is successful, the newly introduced edges are tested again with an expansion test. The expansion is performed only if there is a single possible extension at a vertex, thus eliminating the need for backtracking.
- In [Dui00] backtracking was explicitly introduced, together with a number of new test conditions to rule out subnetworks, dominating those mentioned in [UdAR99].
- In Section 3.3.4, we used a different test that tries to eliminate edges. Expansion is performed only if there is at most one possible extension (thus inspecting a path) and only if the elimination of one edge implies the elimination of all edges of the path.

All previous approaches use only alternative-based methods. We present an expansion test that explicitly combines the alternative-based and bound-based methods. This combination is far more effective than previous tests, because the two approaches have complementary strengths. Intuitively speaking, the alternative-based method is especially effective if there are terminals in the vicinity of the currently inspected subgraph T, because it uses the bottleneck Steiner distances. On the other hand, the bound-based method is especially effective if there are no close terminals, because it uses the distances (with respect to reduced costs) to terminals. Furthermore, for the expansion test to be successful, usually many possible extensions must be considered and it is often the case that not all of them can be ruled out using exclusively the alternative- or the bound-based methods, whereas an explicit combination of both methods can do the job.

#### 3.5. EXTENDED REDUCTION TECHNIQUES

Although the pseudocode of *EXTENDED-TEST* is simple, designing an efficient and effective implementation requires many algorithmic ideas and has to be done carefully, taking the interaction between different actions into account, which is highly nontrivial. Since writing down many pages of pseudocode would be less instructive, we prefer to explain the main building blocks. In the following, we first describe the test conditions for ruling out trees (the function *RULE-OUT*), using the results of [Dui00] and introducing new ideas. Then we explain the criteria used for truncation and choice of the leaves for expansion (the functions *TRUNCATE* and *PROMISING*). Finally, we will address some implementation issues, particularly data structures for querying different types of distances.

#### 3.5.3 Test Conditions

For the following test conditions we always consider a tree T where terminals may appear only as leaves of the tree, i.e.,  $V(T) \cap R \subseteq L(T)$ . A very general formulation of the alternative-based test condition is the following:

**Lemma 32** Consider a pruning set L' for T. If  $c(T_{L'})$  is larger than the cost of a Steiner tree T' in  $G' = (V, V \times V, s)$  with L' as terminals, then there is an optimal Steiner tree that does not peripherally contain T. This test can be strengthened to the case of equality if there is a vertex v in  $T_{L'}$  that is not in any of the paths used for defining the *s*-values of the edges of T'.

**Proof:** Assume that T is peripherally contained in an (optimal) Steiner tree  $T^*$  in G. As L' is a pruning set for T and the leaves of T are a pruning set for  $T^*$ , L' is also a pruning set for  $T^*$ . It follows that after removing the edges of  $T_{L'}$  from  $T^*$ , each of the remaining subtrees contains one vertex of L'. The plan is to reconnect these subtrees to a new Steiner tree by replacing each necessary edge of T' with a path in G of no larger cost. Consider the forest F consisting of these subtrees together with the remaining nodes of T' (i.e., nodes that are not in any of these subtrees). Merge all vertices of T' that are in one component of F, breaking emerging cycles by deleting an arbitrary edge of each cycle. This operation does not increase the cost of T'. Now, each component  $C_i$  of F corresponds to one vertex  $t_i$  of T'. We will ensure this invariant during the whole process of updating T' and F.

Choose a shortest edge  $(t_i, t_j)$  of T'. Let  $P_{ij}$  be a path of bottleneck Steiner distance  $s_{ij}$  between  $v_i$  and  $v_j$ , vertices of V corresponding to  $t_i$  and  $t_j$  (before merging). Let  $P_{kl}$  be a subpath of  $P_{ij}$  in which only the endpoints  $v_k$  and  $v_l$  are in  $R \cup \{v_i, v_j\}$ , and  $v_k$  and  $v_l$  are in different components  $C_k$  and  $C_l$  of F. Remove an arbitrary edge on the fundamental path in T' between  $t_k$  and  $t_l$  and merge  $t_k$  and  $t_l$  in T'. Finally, connect  $C_k$  and  $C_l$  in F by adding the necessary edges from  $P_{kl}$ . The sum of the costs of these edges is not larger than  $s_{ij}$ . Because  $(t_i, t_j)$  was a shortest edge of T', the added cost in F is also not larger than the cost of the edge that was removed from T'.

Repeating this procedure leads to a new network that connects all terminals of G and has cost at most  $c(T^*) - c(T_{L'}) + c(T')$ . If  $c(T') < c(T_{L'})$  or  $c(T') = c(T_{L'})$  and there is at least one vertex in  $V(T_{L'})$  that is not in the new tree (because it was not in any of the paths that were used for defining the *s*-values of the edges in T'), we have a Steiner tree of cost not larger then  $c(T^*)$  that does not peripherally contain T.<sup>2</sup>

A typical choice for L' is L(T), often with some leaves replaced by vertices added to T in the first steps of the expansion. If computing an optimal Steiner tree T' is considered too expensive, the cost of a minimum spanning tree for L' with respect to s can be used as a valid upper bound.

A relaxed test condition compares bottleneck Steiner distances with tree bottlenecks:

<sup>&</sup>lt;sup>2</sup>There are proofs in [Dui00, HRW92] for similar (but weaker) conditions, but they are not complete.

**Lemma 33** If  $s_{ij} < t_{ij}$  for any  $v_i, v_j \in T$ , there is an optimal Steiner tree that does not peripherally contain T. Again, the test can be strengthened to the case of equality if a path corresponding to  $s_{ij}$  does not contain a tree bottleneck of T between  $v_i$  and  $v_j$ .

**Proof:** Consider  $v_i, v_j$  and all key nodes on the fundamental path  $P_{ij}$  between  $v_i$  and  $v_j$  in T as the pruning set L' in the previous lemma. The induced subtree  $T_{L'}$  is the path  $P_{ij}$  itself. Removing a tree bottleneck from  $P_{ij}$ , inserting an edge  $(v_i, v_j)$  of cost  $s_{ij}$  and substituting the *c*-values for the other edges with the (not larger) *s*-values leads to a Steiner tree for L' in G' with cost at most  $c(P_{ij}) + s_{ij} - t_{ij}$ .

The bound-based test condition uses a dual feasible solution for  $LP_C$  of value *lower'* and corresponding reduced costs c'' (with resulting distances d''):

**Lemma 34** Let  $\{l_1, l_2, \ldots, l_k\} = L(T)$  be the leaves of T. Then  $lower_{constrained} := lower' + min_i\{d''(z_1, l_i) + c''(\vec{T_i}) + \sum_{j \neq i} min_{z_p \in \mathbb{R}^{z_1}} d''(l_j, z_p)\}$  defines a lower bound for the cost of any Steiner tree with the additional constraint that it peripherally contains T, where  $\vec{T_i}$  denotes the directed version of T when rooted at  $l_i$ .

**Proof:** If T is peripherally contained in an optimal Steiner tree  $T^*$ , then there is a path in  $T^*$  from the root terminal  $z_1$  to a leaf  $l_i$  of T. After rooting T from  $l_i$ , each (possibly single-vertex) subtree of  $T^*$  corresponding to other leaves  $l_j$  contains a terminal. Now the lemma follows directly using Lemma 29.

In the context of replacement of edges, one can use the following lemma.

**Lemma 35** Let  $e_1$  and  $e_2$  be two edges of T in a reduced network. If both edges originate from a common edge  $e_3$  by a series of replacements, then no optimal Steiner tree for the reduced network that corresponds to an optimal Steiner tree in the unreduced network contains T.

**Proof:** Assume that there is an optimal Steiner tree  $T^*$  for the reduced network containing both  $e_1$  and  $e_2$ . Back-substituting the edges of  $T^*$  leads to a solution in the original network in which  $e_3$  is used twice. This means that the solution value in the unreduced and consequently in the reduced network can be decreased by  $c(e_3)$ , which contradicts the optimality of  $T^*$ .

The conditions above cover the calls RULE-OUT(T, L) with L = L(T). In case other vertices than the leaves need to be considered in the linking set (as in line 8 of the pseudocode), one can easily establish that all lemmas above remain valid if we treat all vertices of L as leaves.

## 3.5.4 Criteria for Expansion and Truncation

The basic truncation criterion is the number of backtracking steps, where there is an obvious tradeoff between the running time and the effectiveness of the test. A typical number of backtracking steps in our implementations is five.

Additionally, there are other criteria that guide and limit the expansion:

 If a leaf is a terminal, we cannot easily expand over this leaf, because we cannot assume anymore that an optimal Steiner tree must connect this leaf to a terminal by edges not in the current tree. However, if all leaves are terminals (a situation in which no expansion is possible for the original test), we know that at least one leaf is connected by an edge-disjoint path to another terminal (as long as not all terminals are spanned by the current tree). This can be built into the test by another level of backtracking and some modifications of the test conditions. But we do

#### 3.5. EXTENDED REDUCTION TECHNIQUES

not describe the modifications in detail, because the additional cost did not pay off in terms of significantly more reductions.

- 2. If the degree d of a leaf is large, considering all  $2^{(d-1)} 1$  possible extensions would be too costly and the desired outcome, namely that we can rule out all of these extended subtrees, is less likely. Therefore, we limit the degree of possible candidates for expansion by a small constant, e.g. 8.
- 3. It has turned out that a depth-first realization of backtracking is quite successful. In each step, we consider only those leaves for expansion that have maximum depth in T when rooted at the starting point. In this way, the bookkeeping of the inspected subtrees becomes much easier and the whole procedure can be implemented without recursion. A similar idea was already mentioned (but not explicitly used) in [Dui00].
- 4. In case we do not choose the depth-first strategy, a tree T could be inspected more than once. As an example, consider a tree T resulting from an expansion of T' at leaf  $v_i$  and then at  $v_j$ . If T cannot be ruled out, it is possible that we return to T', expand it at  $v_j$  and then at  $v_i$ , arriving at T again. This problem can be avoided by using a (hashing-based) dictionary.

## 3.5.5 Implementation Issues

**Precomputing (Steiner) Distances:** A crucial issue for the implementation of the test is the calculation of bottleneck Steiner distances as defined in Section 3.2. An exact calculation of all  $s_{ij}$  needs time  $O(n(m + n \log n))$  [Dui00] and space  $\Theta(n^2)$ , which would make the test impractical even for mid-size instances. So we need a good approximation of these distances and some appropriate data structures for retrieving them. Building upon a result of Mehlhorn [Meh88], Duin [Dui93] gave a nice suggestion for the approximation of the bottleneck Steiner distances, which needs preprocessing time  $O(m + n \log n)$  and a small running time for each query. As described in Section 3.3.1 the query time can even made constant if all necessary queries are known in advance. Although the resulting approximate values  $\hat{s}_{ij}$  produce quite satisfactory results for the original s-test (PT<sub>m</sub> in Section 3.3.1), for the extended test the results are unfortunately much worse than with the exact values. But we observed that  $\tilde{s} = \min\{\hat{s}, d\}$  is almost always equal to the exact s-values, and therefore can be used in the extended test as well. Still there remains the problem of computing the *d*-values: For each vertex  $v_i$  we compute and store in a neighbor list the distances to a constant number (e.g. 50) of nearest vertices. But we consider only vertices  $v_j$  with  $d_{ij} < \hat{s}_{ij}$ . This is justified by the observation that in case  $d_{ij} \geq \hat{s}_{ij}$ , for all descendants  $v_k$  of  $v_j$  in the shortest paths tree with the root  $v_i$  there is a path  $P_{ik}$  of bottleneck Steiner distance  $s_{ik}$  containing at least one terminal, and in such cases  $s_{ik}$  is usually quite well approximated by  $\hat{s}_{ik}$ .

Now, we use different methods for different variants of the test:

- 1. If we only replace vertices in an expansion test, the *d* and *s*-values do not increase and we can use the precomputed neighbor lists during the whole test using binary search.
- 2. If we limit the number of (backtracking) steps, then we can confine the set of all possible queries in advance. When a vertex is considered for replacement by the expansion test, we first compute the set of possibly visited neighbors (adjacent vertices, and vertices adjacent to them, and so on, up to the limited depth). Then we compute a distance matrix for this set according to the  $\tilde{s}$ -values. Using this matrix, each query can be answered in constant time.

3. If we also want to delete edges, we have to store for each vertex in which neighbor list computations it was used. When an edge is deleted, we can redo the computation of the  $\hat{s}$ -value (or at least those parts that may have changed due to the edge deletion) and restore the affected neighbor lists. This can even be improved by a lazy calculation of the neighbor lists.

**Tree Bottlenecks:** The tree bottleneck test of Lemma 33 can be very helpful, because every distance between tree vertices calculated for a minimum spanning tree or a Steiner minimal tree computation can be tested against the tree bottleneck; and in many of the cases where a tree can be ruled out, already an intermediate bottleneck test can rule out this tree, leading to a shortcut in the computation. This is especially the case if there are long chains of nodes with degree two in the tree. We promote the building of such chains while choosing a leaf for extension: We first check whether there is a leaf at which the tree can be expanded by only one edge. In this case we immediately perform this expansion, without creating a new key node and without the need of backtracking through all possible combinations of expansion edges.

The tree bottleneck test can be sped up by storing for each node of the tree the length of a tree bottleneck on the path to the starting vertex. For each two nodes  $v_i$  and  $v_j$  in the tree, the maximum of these values gives an upper bound for the actual tree bottleneck length  $t_{ij}$ . Only if this upper bound is greater than the (approximated) bottleneck Steiner distance, an exact tree bottleneck computation is performed.

**Computations for the Bound-Based Tests:** An efficient method for generating the dual feasible solution needed for the bound-based test of Lemma 34 is the DUAL-ASCENT algorithm described in Section 3.4.2. We improve the test by calculating a lower bound and reduced costs for different roots. Although the optimal value of the directed cut relaxation  $LP_C$  does not change with the choice of the root, this is not true concerning the value of the dual feasible solution generated by DUAL-ASCENT and, more importantly, the resulting reduced costs can have significantly different patterns, leading to a greater potential for reductions.

Even more reductions can be achieved by using stronger lower bounds, as computed with a row generating algorithm, see Section 3.4.3. Concerning the tests for the replacement of vertices, we use only the result of the final iteration, which provides an optimal dual solution of the underlying linear relaxation. The dual feasible solutions of the intermediate iterations are used only for the tests dealing with the deletion of edges, because the positive effect of the replacement of a vertex (see the NTD<sub>k</sub> test in Section 3.3.2) cannot be translated easily into linear programs.

**Replacement History:** Our program package can transform a tree in a reduced network back into a tree in the original instance. For this purpose, we assign a unique ID number to each edge. When a vertex is replaced, we store for each newly inserted edge a triple with the new ID and the two old IDs of the replaced edges. We use this information to implement the test described in Lemma 35. First we do some preprocessing, determining for each ID the edges it possibly originates from (here called ancestors); this can be done in time and space linear in the number of IDs. Later, a test for a conflict between two edges (i.e., they originate from the same edge) can be performed by marking the IDs of the ancestors of one edge and then checking the IDs of ancestors of the other edge; so each such test can be done in time linear in the number of ancestors. We perform this test each time the current tree T is to be extended over a leaf  $v_i$  (with  $(v_k, v_i)$  in T) by an edge  $(v_i, v_j)$ . Then we check for a conflict between  $(v_i, v_j)$  and  $(v_k, v_i)$ . This procedure implements an idea briefly mentioned in [Dui00], where a coloring scheme was suggested for a similar purpose. Our scheme has the advantage that it may even discover conflicts in situations where an edge is the result of a series of replacements.

## 3.5.6 Variants of the Test

A general principle for the application of reduction tests is to perform the faster tests first so that the stronger (and more expensive) tests are applied to (we hope) sufficiently reduced graphs. In the present context, different design decisions (e.g., trying to delete edges or replace vertices) lead to different consequences for an appropriate implementation and quite different versions of the test, some faster and some stronger.

We have implemented four versions of expansion tests and integrated them into the reduction process. Some details of the corresponding implementations were already given in Section 3.5.5.

- 1. For a fast preprocessing we use the linear time expansion test that eliminates paths, as described in Section 3.3.4.
- 2. A stronger variant tries to replace vertices, but only expands at leaves that are the most number of edges away from the starting vertex.
- 3. Even stronger but more time-consuming is a version that performs full backtracking.
- 4. The most time-consuming variant tries to eliminate edges.

# **3.6** Partitioning as a Reduction Technique

Partitioning is a basic principle in many algorithms. Although the basic ideas behind the approach presented in the following are relatively simple, the way we exploit them is new, and our approach could also be useful for other problems.

There are several reasons that motivate the use of partitioning in the present or similar contexts:

- **Efficiency:** For any algorithm with superlinear running time, a suitable partitioning of the instance leads to a superlinear speedup. Note that because exact algorithms in this context use very time-consuming components (like LP-solvers) and are even exponential in the worst case, the speedup for solving subproblems can be highly superlinear.
- **Effectiveness:** Sometimes, a method (a component of an exact algorithm) works well on some group of instances; but it fails on larger instances of the same type. Methods that are based on LP-relaxations of the problem are good examples, because any LP-relaxation of polynomial size is bound to have some (integrality) errors in this context. In larger instances, such errors can accumulate and become more and more relevant. Partitioning can help against this accumulation of errors, as we will show in Section 3.6.3.
- **Implementation:** A reasonable partitioning offers a direct path to a distributed implementation, because different (reasonably independent) subproblems can be processed on different processors.

However, for applying the idea of partitioning to problems like the Steiner problem, classical approaches are not very helpful. Divide-and-conquer techniques are not generally applicable, because one usually cannot find independent subproblems. Dynamic programming techniques can indeed be applied, but these techniques (at least in their classical form) do not lead to empirically efficient algorithms.

In this section, we introduce the new approach of using partitioning to design reduction methods. We partition instances by finding (small) terminal separators, i.e., after removing these terminals from the network it is no longer connected. This allows us to keep the dependence between the resulting subinstances manageable.

#### 3.6.1 Partitioning on the Basis of Terminal Separators

Although one cannot assume that a typical instance of the Steiner problem has small terminal separators, the situation often changes in the process of solving an instance, as described in the following.

#### **Reduced Instances**

There are several reduction methods which, when successful, tend to transform instances without useful terminal separators into instances with them.

Figure 3.6 shows a VLSI-instance from the library [SteinLib]. The terminals (black squares) are to be connected on the grid. Note that there are holes in the grid (corresponding to obstacles on the chip), so such instances are not geometric (rectilinear). In this figure, one does not detect any useful terminal separator. But the situation changes after applying some reduction methods: Figure 3.7 shows the reduced instance, which is produced after a couple of seconds: the black edges are chosen (and contracted); the grey edges remain as the reduced instance; this reduced instance is redrawn more compactly in Figure 3.8. Note that the visualization used is not geometric; edges which appear relatively long may actually have relatively small cost. (Our algorithm is a pure graph algorithm, which does not use the coordinates of the points anyway.) In this reduced instance, one easily detects many small terminal separators and corresponding components.



Figure 3.6: VLSI-instance taq0377 (|V| = 6836, |E| = 11715, |R| = 136).



Figure 3.7: taq0377, reduced (|V| = 193, |E| = 312, |R| = 67).



Figure 3.8: taq0377, reduced, redrawn (|V| = 193, |E| = 312, |R| = 67).

#### **Geometric Steiner Problems**

For geometric Steiner problems, an approach based on full Steiner trees has been empirically successful [WWZ00]: In geometric Steiner problems, a set of points (in the plane) is to be connected at minimum cost according to some geometric distance metric. The resulting interconnection, a Steiner minimal tree (SMT), can be decomposed into FSTs (full Steiner trees) by splitting its inner terminals. The FST-approach consists of two phases. In the first phase, the FST generation phase, a set of FSTs is generated that is guaranteed to contain an SMT. In the second phase, the FST concatenation phase, one chooses a subset of the generated FSTs whose concatenation yields an SMT. Although there are point sets that give rise to an exponential number of FSTs in the first phase, usually only a linear number (in |R|) of FSTs are generated, and empirically the bottleneck of this approach has usually been used. A breakthrough occured when Warme [War98] used the fact that FST concatenation can be reduced to finding a minimum spanning tree in a hypergraph whose vertices are the terminals and whose (hyper-)edges correspond to the generated FSTs. Although the minimum spanning tree in hypergraph (MSTH) problem is  $\mathcal{NP}$ -hard, a branch-and-cut approach based on the linear relaxation of an integer programming formulation of this problem has been empirically successful.

Note that by building the union of (the edge sets of) the FSTs generated in the first phase, we get a normal graph and the FST-concatenation problem is reduced to solving the classical Steiner problem in this graph. In Section 5.4.1, we report that our general graph algorithm often performs the second phase significantly faster than the MSTH-based method, especially for the more time-consuming instances. This is mainly due to the use of reduction techniques, where the methods presented here also play an important role, because usually the instances have many useful terminal separators with corresponding small components already at the beginning of the second phase, and this is even amplified after application of other reduction methods.

#### **Combination of Steiner Trees**

During the solution process, our program generates many heuristical solutions (Steiner trees). It would not be the best idea to just keep the best of them and throw the others away, because one could possibly combine parts of them to construct a new, even better solution. A simple but effective method is to build the instance corresponding to the union of the edges of different Steiner trees and find a (heuristical or even exact) solution in that instance (see Section 4.5). Such instances often have small terminal separators and the methods described here can be applied.

#### 3.6.2 Finding Terminal Separators

It is well known that the vertex connectivity problem can be solved by network flow techniques in the so-called split graph, which is generated by splitting each vertex into two vertices and connecting them by edges of low capacity; original edges have high (infinite) capacity. In this way, k-connectedness (finding a vertex separator of size less than k or verifying that no such separator exists) can be decided in time  $O(\min\{kmn, (k^3 + n)m\})$  [HRG00] (this bound comes from a combination of augmenting path and preflow-push methods). In case of undirected graphs (as in the present application), the job can be done in a sparse graph with O(kn) edges, which can be constructed in time O(m) [NI92], so m can be replaced in the above bound by kn.

However, the application here is less general: we search for vertex separators consisting of terminals only, so only terminals need to be split. Besides, we are interested in only small separators, where k is a very small constant (usually less than 5, say at most 10), so we can concentrate on the
(easier) augmenting flow methods. More importantly, we are not searching for a single separator of minimum size, but for many separators of small (not necessarily minimum) size. These observations have lead to the following implementation: we build the (modified) split graph (as described above), fix a random terminal as source, and try different terminals as sinks, each time solving a minimum cut problem using augmenting path methods. In this way, up to  $\Theta(r)$  terminal separators can be found in time O(rm). We accelerate the process by using some heuristics. A simple observation is that vertices that are reachable from the source by paths of non-terminals need not be considered as sinks. Similar arguments can be used to heuristically discard vertices that are reachable from already considered sinks by paths of non-terminals.

Empirically, this method is quite effective (it finds enough terminal separators if they do exist) and reasonably fast, so a more stringent method (e.g., trying to find all separators of at most a given size) would not pay off. Note that the running time is within the bound given above for the k-connectedness problem, which is mainly the time for finding a single vertex separator.

### **3.6.3 Reduction Methods**

In this section, we describe two methods that exploit a terminal separator  $S \subset R$  and the corresponding partitioning to reduce a given instance.

#### **Case Differentiation**

- warm-up (|S| = 1): The case |S| = 1 corresponds to biconnected components and articulation points (i.e., if one of the articulation points is removed from the network, it is no longer connected) and biconnected components, which can be found in linear time O(m). It is well known [HRW92] that the subinstances corresponding to the biconnected components can be solved independently. An example can be found in Figure 3.18: There is an articulation point in the middle; the upper and the lower components can be solved independently and the small component in the middle can be discarded (see Lemma 42 in Section 5.3.2).
- **base case** (|S| = 2): The case |S| = 2 corresponds to separation pairs (and triconnected components). All (non-trivial) triconnected components can be found in linear time O(m) [HT73]. Consider Figure 3.9, where a separator S of size 2 and the corresponding components  $G_1$  and  $G_2$  are shown (black edges for  $G_2$ ). Note that the two subinstances are no longer independent. Now, for any Steiner minimal tree T, two cases are possible :
  - I) The terminals in S are connected by T inside  $G_2$ . A corresponding Steiner tree can be found by solving the subinstance corresponding to  $G_2$ .
  - II) The terminals in S are connected by T inside  $G_1$ . Now there are two subtrees of T inside S, and we do not know in advance how the terminals of  $G_2$  are divided between them. But one can observe that the problem can be solved by merging the terminals in S and solving the resulting subinstance.

Since we do not know T in advance, for a direct solution we must also consider both cases for its complement  $G_1$ . But if  $G_2$  is relatively small, the solution of the complementary subinstance can be almost as time-consuming as the solution of the original instance, meaning that not much is gained (or time may even be lost, because now we have to solve it twice). A classical approach would search for components of almost equal size, but we choose a different approach. The idea is to solve only the small component twice, and then take edges that are common to both solutions and discard edges that are included in neither. The result is shown in Figure 3.10, where only one edge from  $G_2$  is left undecided (the other edges can be either contracted (black) or deleted (light grey)). In Figure 3.11, the reduced instance is redrawn; as one sees, the processed component has almost disappeared.

general case (|S| = k): As described in Section 3.6.2, we can find up to  $\Theta(r)$  separators of size at most k in time O(krm). The basic approach is the same as for the case |S| = 2; but a larger number of cases must be considered now. We put each subset of terminals in S that are connected by one subtree of T in  $G_1$  into one group. There can be  $i = 1, \ldots, k$  such groups. For each i, we must count the number of ways of partitioning a set of k elements into i non-empty subsets, which is a Stirling number of the second kind  ${k \atop i}$ . So there are  $\sum_{i=1}^{k} {k \atop i}$  cases. Table 1 contains the concrete numbers for small k.

k	2	3	4	5	6	7	8
cases	2	5	15	52	203	877	4140

Table 1: Possible cases for a terminal separator of size k

As the numbers in Table 1 suggest, this method can be used profitably usually only for  $k \le 4$  (and for  $k \ge 3$  only if the processed component is reasonably small). Actually, not all these cases must always be considered explicitly, because many of them can be ruled out at little extra cost using some heuristics. A basic idea for such heuristics uses the following lemma:

**Lemma 36** Let  $z_i$  and  $z_j$  be two terminals in the separator S and let  $b_{ij}^1$  and  $s_{ij}^2$  be the bottleneck distance in  $G_1$  and bottleneck Steiner distance in  $G_2$  between  $z_i$  and  $z_j$ , respectively. Then the cases in which  $z_i$  and  $z_j$  are connected in  $G_1$  can be discarded if  $b_{ij}^1 \ge s_{ij}^2$ .

**Proof:** Consider a Steiner tree T connecting  $z_i$  and  $z_j$  in  $G_1$ . A bottleneck on the fundamental path between  $z_i$  and  $z_j$  has at least cost  $b_{ij}^1$ . Removing such a bottleneck and reconnecting the two resulting subtrees of T with the subpath corresponding to  $s_{ij}^2$ , we get again a feasible solution of no larger cost in which  $z_i$  and  $z_j$  are connected in  $G_2$ .

Note also that for the cases in which we assume that  $z_i$  and  $z_j$  are connected in  $G_1$ , we do not merge  $z_i$  and  $z_j$  while solving the subinstance corresponding to  $G_2$ , but connect them with an edge of weight  $b_{ij}^1$ . In case this edge is not used in the solution of the subinstance, this can lead to more reductions.

This and similar observations can be used to rule out many cases in advance. Nevertheless, a question naturally arises: Can we find an alternative method that does not need explicit case differentiation? We will introduce such an alternative in the following section.

#### **Local Bounds**

Recall that the general principle of bound-based reduction methods is to compute an upper bound *upper* and a lower bound under some constraint *lower* constrained. The constraint cannot be satisfied by any optimal solution if *lower* constrained > upper. The constraint is usually that the solution must contain some pattern (e.g., a vertex or an edge, or even more complex patterns like paths and trees, as in Section 3.5 on extended reduction techniques). But it is usually too costly to recompute a (strong) lower bound from scratch for each constraint. Here one can use an approach based on Linear Programming presented in Section 3.4.2: Any linear relaxation can provide a dual feasible solution of value *lower* and reduced costs c'. We can use a fast method to compute a constrained lower bound



Figure 3.9: |S| = 2.



Figure 3.10: (|V| = 133, |E| = 214, |R| = 45).



Figure 3.11: (|V| = 133, |E| = 214, |R| = 45).

*lower'* with respect to c'. From Lemma 29 follows that *lower* constrained := *lower* + *lower'* is a lower bound for the value of any solution satisfying the constraint.

As an example for such a relaxation, consider the (directed) cut formulation  $P_C$  [Won84]. It has been described in Section 2.3.1. Here, we give an informal description: The Steiner problem is formulated as an integer program by introducing a binary x-variable for each arc (in case of undirected graphs, the bidirected counterpart is used, fixing a  $z_1 \in R$  as the root). For each cut separating the root from another terminal, there is a constraint requiring that the sum of x-values of the cut arcs is at least 1. In this way, every feasible binary solution represents a feasible solution for the Steiner problem and each minimum solution (with value  $v(P_C)$ ) a Steiner minimal tree (Figure 3.12). But in the optimal solution of the LP-relaxation  $LP_C$  variables can have fractional values. In Figure 3.13, the grey arcs have value 1/2 (the black arcs have value 1, the other arcs have value 0). One observes that a flow of one unit can still be sent from the root to each terminal, so all cut constraints are satisfied; and if the costs are adverse, an integrality gap can occur. This is in fact the case in this example, where the linear relaxation has optimum solution value  $v(LP_C) = 6392.5$ . As such gaps accumulate (e.g., in larger instances), the difference between the bounds grows, eventually causing the bound-based reductions to fail.

In the following, we show how to use locally computed bounds for bound-based reduction. This approach has two main advantages: The bounds can be computed faster; and there is less chance of accumulation of errors. The main difficulty is that the bounds must somehow take the dependence on the rest of the graph into account.

Let S be a terminal separator in G and  $G_1$  and  $G_2$  the corresponding subgraphs (Figure 3.14). The bounds will be computed locally in supplemented versions of  $G_2$ . Let C be a clique over S. We denote with (C, b) the weighted version of C with weights equal to bottleneck distances in  $G_1$ ; similarly for (C, s) with weights equal to bottleneck Steiner distances in G. Let  $G'_2$  and  $G''_2$  be the instances of the Steiner problem created by supplementing  $G_2$  with (C, s) and (C, b), respectively. We compute a lower bound  $lower_{constrained}(G''_2)$  for any Steiner tree satisfying a given constraint in  $G''_2$  and an upper bound  $upper(G'_2)$  corresponding to an (unrestricted) Steiner tree in  $G'_2$ . The test condition is:  $upper(G'_2) < lower_{constrained}(G''_2)$ .



Figure 3.12:  $v(P_C) = 6393$ .



Figure 3.13:  $v(LP_C) = 6392.5$ .



Figure 3.14: A terminal separator S and corresponding subgraphs  $G_1$  and  $G_2$ .

**Lemma 37** The test condition is valid, i.e., no Steiner minimal tree in G satisfies the constraint if  $upper(G'_2) < lower_{constrained}(G''_2)$ .

**Proof:** Consider  $T_{con}^{opt}(G)$ , an (unknown) optimum Steiner tree satisfying the constraint. The subtrees of this tree restricted to subgraphs  $G_1$  and  $G_2$  build two forests  $F_1$  (with connected components  $T_i$ ) and  $F_2$  (Figure 3.15, left). Removing  $F_2$  and reconnecting  $F_1$  with  $T^{upper}(G'_2)$  we get a feasible solution again, which is not necessarily a tree (Figure 3.15, middle). Let  $S_i$  be the subset of S in  $T_i$ . Consider two terminals of  $S_i$ : Removing a bottleneck on the corresponding fundamental path disconnects  $T_i$ into two connected components. Repeating this step until all terminals in  $S_i$  are disconnected in  $T_i$ , we have removed  $|S_i| - 1$  bottlenecks, which together build a spanning tree  $spanT_i$  for  $S_i$  (Figure 3.15, right). Repeating this for all  $T_i$ , we get again a feasible Steiner tree  $T^{upper}(G')$  for the graph G', which is created by adding the edges of (C, s) to G. Note that the optimum solution values in G'and G are the same (see Section 1.3). Let upper(G') be the weight of  $T^{upper}(G')$ . By construction of  $T^{upper}(G')$ , we have:

$$upper(G') = opt_{con}(G) + upper(G'_2) - c(F_2) - \sum_i c(spanT_i)$$

The edge weights of the trees  $spanT_i$  correspond to bottlenecks in  $F_1$ , so by definition they cannot be smaller than the corresponding bottleneck distances in  $G_1$ . By construction of  $G''_2$ , all these edges (with the latter weights) are available in  $G''_2$ . Since the trees  $spanT_i$  reconnect the forest  $F_2$ , together with  $F_2$  they build a feasible solution for  $G''_2$ , which even satisfies the constraint (because  $F_2$  did), so it has at least the cost  $opt_{con}(G''_2)$ . This means:

$$upper(G') \leq opt_{con}(G) + upper(G'_2) - opt_{con}(G''_2) \\ < opt_{con}(G) + lower_{constrained}(G''_2) - opt_{con}(G''_2)$$
 (because of the test condition)   
 
$$\leq opt_{con}(G),$$

thus  $opt_{con}(G) > upper(G') \ge opt(G') = opt(G)$ , meaning that the constraint cannot be satisfied without deteriorating the optimal solution value.  $\Box$ 



Figure 3.15: Construction of  $T^{upper}(G')$  from  $T^{opt}_{con}(G)$ .

Studying the test condition, one detects a weakness of the lower bound used relative to the upper bound: bottleneck distances used in the lower bound correspond to single edges, whereas the bottleneck Steiner distances used in the upper bound correspond to whole paths and can be much larger. The attempt to use some paths in  $F_1$  instead of bottlenecks fails because the tree  $T_{con}^{opt}(G)$  and consequently the forest  $F_1$  are unknown. But going through the proof, one observes that we can use the fact that the tree  $T^{upper}(G'_2)$  is available: Instead of removing a single edge in  $F_1$ , we can remove a (longest) key path on the corresponding fundamental path in  $T^{upper}(G'_2)$ . This leads to the following improvement of the test: While choosing the edge weights for constructing  $G''_2$ , we can use the length of a key path in  $T^{upper}(G'_2)$  whenever it is larger than the corresponding bottleneck distance in  $G_1$ . However, care must be taken to keep the key paths used disjoint.

An application example for this test is given in Figure 3.16, where a separator of size 4 and the corresponding component  $G_2$  are shown (black edges). Figure 3.17 shows the result of application of the reduction method presented (black edges contracted, grey edges remaining); in Figure 3.18 the reduced instance is redrawn.

This is also an example of how reduction methods based on partitioning can reduce the errors in an LP-relaxation: As shown in Figure 3.19, the relaxation  $LP_C$  has now an integer optimal solution. It is perhaps interesting that this improvement is mainly achieved by applying the same relaxation locally.

## **3.7 Integration and Implementation of Tests**

To study the effect of different combinations and orderings of the tests, we designed an interpreter for command-lines, where each test is encoded by a character. We also implemented a direct control of loops (through parentheses), their termination criteria, switching of parameters, etc. The main observation is that the (alternative-based) tests are not very sensitive to the order in which they are executed. On the other hand, the ordering often has an impact on the total time for reductions; in this sense the ordering cited in [HRW92] is a suitable one (although not necessarily the only one, as long as a fast version of  $PT_m$  is performed first).

For the implementation, we have chosen a kind of adjacency-list representation of networks (with all edges in a single array), but we sometimes switch to other auxiliary representations (all linear in



Figure 3.16: |S| = 4.



Figure 3.17: (|V| = 121, |E| = 200, |R| = 41).



Figure 3.18: (|V| = 121, |E| = 200, |R| = 41).



Figure 3.19:  $v(LP_C) = 6393$ .

the number of edges) for certain operations. For each test, we perform all actions in a single pass (and do not, for example, delete an edge and start the test from scratch). The details of the realization of the various actions are very technical and are omitted here; we merely mention that all actions following each test can be realized in a time dominated by the worst-case time  $O(m + n \log n)$  of the fast tests.

With the additional requirement that in each loop of the selected tests a constant proportion (say 5%) of vertices and edges must be eliminated and that instances of trivially small size are solved directly (by enumeration), one gets the same asymptotic time bound for the whole reduction process as for the first iteration  $(O(m + n \log n))$ , if one confines oneself to the fast tests).

Another technical aspect is the efficient reconstruction of a solution for the original instance out of a solution for the reduced instance (which often consists of a single terminal). Saving appropriate information during the reduction process, this can be done in time O(m). We always perform such a transformation after each run of the program, checking the feasibility and value of the solution in the original instance.

## **3.8 Some Experimental Results**

In this section, we present some summarized results of the reduction package on instance groups of SteinLib (see Section 1.2 for a description of the problem instances). To give some impression of the effect of the different approaches, we present results for different subsets of the reduction techniques:

- $O(m + n \log n)$ : In these runs, we have only included the fast versions of the reduction tests  $PT_m$ , Triangle,  $NTD_k$ , NV, SL, PS, VR, and LDA, which all can be implemented with a worst-case running time of  $O(m + n \log n)$ . For computing an upper bound, we used a fast path heuristic with the same running time, which will be described in Section 4.2.
- + **DUAL-ASCENT:** These runs additionally use the DUAL-ASCENT algorithm, for generating a lower bound, the calculation of an auxiliary graph for ASCEND-AND-PRUNE (see Section 4.4), and for bound-based reductions.
- + Extended Reduction Techniques: In this column, the extended reduction tests are also used. Note that using extended reductions, there is an obvious tradeoff between reduction results and running time, which can be configured by increasing or decreasing the parameter for the maximum backtracking depth. A more detailed study of the results using extended reduction techniques is given in [PV01b].
- + **Partitioning:** Here, we additionally use the partitioning based reductions, presented in Section 3.6. As they require an exact solution of small subinstances, we also had to include the RG-test and branching, but only for the solution of the subinstances. Results including the RG-test are presented in the context of the exact algorithm, in Section 5.4.

We know of only a few other works where the reported reduction results reach a quality that is at least comparable to ours. One is the PhD thesis of Cees Duin [Dui93] who developed together with Ton Volgenant several of the most important classical tests. He modified these tests so that they are applicable at least to medium sized instances. Our results on these instances are similar, but in general we have better results: On the D-instances of SteinLib our fastest reductions already leave less than 2% of the edges, while his leave 24%. (Duin did not give results on larger benchmark instances.) The best other results on these instances that were published in a journal are from Koch and Martin [KM98]. For the D-instances they achieve reduction to 38% of the edges of the original network.

## 3.8. SOME EXPERIMENTAL RESULTS

For VLSI instances, Uchoa, Poggi de Aragão, and Ribeiro [UdAR02] presented the best other results: With their reduction techniques only 6.2% of the edges remain, and the reductions took 367 seconds on a Sun Ultra Sparc with 167 MHz. With our reductions, on the average only 0.05% of the edges remain after 4 seconds on a Sparc-III CPU with 900 MHz. A more detailed comparison of the two reduction packages is given in [PV01b, PV02a].

instance	O(m	$+ n \log n$ )	+ DUA	L-ASCENT	+ Exte	ended Red.	+ Par	titioning
group	time	remaining	time	remaining	time	remaining	time	remaining
	in sec.	edges in %	in sec.	edges in %	in sec.	edges in %	in sec.	edges in %
1R	0.08	57.41	0.12	45.60	0.32	0.00	0.20	0.00
2R	0.19	69.99	0.22	64.03	23.46	6.95	20.64	6.24
D	0.08	1.46	0.07	0.52	0.64	0.35	0.31	0.14
E	0.30	3.03	0.27	0.93	4.27	0.55	3.65	0.50
ES10000FST	9.42	62.35	92.68	62.39	874.23	46.05	1048.04	17.41
ES1000FST	0.58	62.67	0.80	62.79	4.82	46.83	9.63	20.32
I080	0.02	43.62	0.02	26.77	0.06	20.63	0.09	15.95
I160	0.05	63.98	0.06	33.61	0.31	28.77	0.43	25.60
I320	0.19	71.82	0.31	47.51	2.48	40.07	3.42	35.72
I640	0.96	79.88	1.89	53.17	15.32	45.12	25.33	43.29
LIN	2.31	31.84	1.72	27.12	207.55	9.29	194.75	6.08
MC	0.03	9.68	0.03	7.05	0.10	6.34	0.17	6.34
PUC	0.34	99.33	0.89	99.33	7.67	99.31	12.77	99.31
SP	0.17	37.50	1.39	37.50	5.75	37.50	7.81	37.50
TSPFST	0.24	30.71	0.80	29.93	7.53	20.37	15.48	7.84
VLSI	0.36	15.70	0.64	10.78	11.00	0.60	4.37	0.05
WRP3	0.15	92.68	0.24	91.41	42.24	58.01	44.26	52.28
WRP4	0.09	96.16	0.20	87.65	14.61	57.63	12.46	44.15
Х	0.34	0.39	0.31	0.00	0.28	0.00	0.32	0.00

Table 3.1: Some experimental results for reductions.

**Chapter 4** 

**Fast Computation of Short Steiner Trees** 

## 4.1 Introduction

In this chapter, we focus on methods for computing upper bounds, in particular computing short (but not necessary optimal) Steiner trees in short running times.

The number of papers on this topic is enormous, there are many heuristics and every popular meta-heuristic has been tried (typically in many variants). A very comprehensive overview of articles published before 1992 can be found in [HRW92]. But the best results have been published after that (or have not yet even been published in a journal). In [PV97] we gave a comprehensive overview of the most important publications before 1997. Important recent developments can be found in [RUW02, PW02].

We have developed a variety of heuristics for obtaining upper bounds. Especially in the context of exact algorithms very sharp upper bounds are highly desired. So, our main concern was achieving very strong bounds, reaching the optimum as often as possible. On the other hand, the goal of obtaining short total empirical running times prohibited us from using heuristics that achieve good solution values only after long runs. In the following, we describe some of the methods we used in our attempt to achieve both goals simultaneously.

## 4.2 Path Heuristics

The repetitive shortest paths heuristic (SPH) is one of the empirically most successful classical heuristics for the Steiner problem in networks [TM80, HRW92, WS92, Voß92, PV01c, PW02]. The basic idea is to build a Steiner tree similar to Prim's algorithm for the minimum spanning tree problem: Start with a subtree that initially consists of one vertex. In each step, connect the current subtree with a shortest path to the next terminal. Repeat this until all terminals are connected. A final pruning step computes a minimum spanning tree of the vertices of the heuristic tree and removes nonterminals that have degree one in this tree. The worst-case running time for such a tree construction is  $O(r(m + n \log n))$ . To improve the quality of the solution, one usually repeats this process with different start vertices (repetitive SPH).

For the computation of shortest paths in each iteration the algorithm of Dijkstra is used, which uses tentative distances  $\pi_i$  for every vertex  $v_i \in V$ . A tentative distance  $\pi_i$  represents an upper bound to the distance between the current subtree and  $v_i$ . In [PW02] Poggi de Aragão and Werneck made a simple, but very effective observation: One does not have to reset tentative distances when a terminal is connected to the current subtree. The reason for this is that the old tentative distances are still valid upper bounds for the distances from the extended subtree. All one has to do is reinsert every vertex  $v_i$  into the priority queue of Dijkstra's algorithm (with  $\pi_i = 0$ ) as soon as it becomes part of the current subtree. Although the empirical running time is improved drastically, it is unsatisfying that the worst-case running time is still  $O(r(m + n \log n))$ . Furthermore, as SPH is usually used in the repetitive variant, one can observe that very similar information is computed over and over without taking advantage of the previous computations.

Studying the repetitive shortest paths heuristic one observes that the actions can be divided into two phases (see [Dui93, DV97]): In the first phase, one can compute shortest paths from each terminal to all vertices; this can be done in time  $O(r(m+n \log n))$ . Using the information from the first phase, each run of the SPH in the second phase (constructing a Steiner tree by successively connecting the current tree (a single vertex at the beginning) to the closest terminal not in the tree by a shortest path) can easily be realized in time O(rn).

In [PV97, PV01c], we presented an efficient implementation of this approach and also a version

with a worst-case running time of  $O(m + n \log n)$ . We will describe these heuristics in the next two sections. Note that we use these variants only as components of our other algorithms, not as standalone heuristics.

## 4.2.1 Faster Preprocessing for the Repetitive Shortest Path Heuristic

Our concern here is achieving empirical acceleration of the two-phase version of repetitive SPH. Regarding the first phase, we observe that the shortest paths need not be always computed completely:

**Lemma 38** Let P be a shortest path between a terminal z and a vertex v, such that there is a vertex v'on P with  $z' := base(v') \neq z$  and  $d(z, z') \leq d(z, v)$ . If v, but not z, belongs to the current tree T in the second phase, there exists at least one other path connecting T to a terminal not in T which is not longer than P. So, when computing shortest paths from z, we need consider neither v nor any vertex that would become a successor of v in the shortest paths tree.

#### **Proof:** There are two cases:

I)  $z' \in T$ : Since  $d(z, z') \leq d(z, v)$ , we can choose the path between z' and z. II)  $z' \notin T$ : Since  $d(z', v) \leq d(z', v') + d(v', v) \leq d(z, v') + d(v', v) = d(z, v)$ , we can choose the path between v and z'.

As a consequence, one can stop computing the shortest paths tree from a terminal z in the first phase as soon as the Voronoi region of z and the neighboring terminals (as defined in 1.3) have been spanned, because the shortest path between z and every vertex v visited afterwards contains a vertex  $v' \in N(z')$ with z' a neighbor of z and  $d(z, z') \leq d(z, v)$ , since z' has already been spanned by the shortest paths tree. Furthermore, no shortest path via an intermediary terminal needs to be considered. These observations often lead to a considerable reduction in the empirical times, especially if the graph has many terminals and is not dense (the latter is almost always the case after reductions). Note that for graphs with few terminals, repetitive SPH is fast anyway.

For building the Steiner trees in the second phase, we prefer a realization that uses the concept of neighborhoods: Using the information from the first phase, we manage for each vertex v a list of neighborhoods: Using the information from the first phase, we manage for each vertex v a list of neighboring terminals, sorted by (increasing) distances to v. A priority queue manages candidates for expansion of the tree, using the distance to the nearest terminal not in the tree as the key for insertion. Each time a vertex v is extracted from the queue, two cases can arise: Either the terminal corresponding to the key is not yet in the tree, in which case the tree is expanded by the corresponding shortest path (and the queue is updated); or it is already in the tree, in which case the neighbor list of v is scanned further until either a terminal not in the tree is visited (which delivers the key for reinsertion of v into the queue) or the end of the list is reached (meaning that v can be ignored). Although the worst-case time of this implementation ( $O(rn \log n)$ ) is slightly worse than O(rn) of the straightforward implementations, it is usually much faster, and the worst-case time is dominated by the first phase anyway.

## 4.2.2 A Path Heuristic with Good Worst Case Running Time

In situations where the worst-case time is the primary concern, we used a strengthening of the ideas above to design a heuristic with time  $O(m + n \log n)$ . Motivated by the fact that the vicinity of each terminal relevant for SPH often gets smaller with a growing number of terminals, one can simply force the first phase not to perform more than  $O(m+n \log n)$  operations. But then it is no longer guaranteed that the relevant neighborhood of each terminal is captured. To remedy this defect, we simultaneously use the graph G' of Mehlhorn's fast implementation of  $T'_D(R)$  [HRW92, Meh88], which we also

#### 4.2. PATH HEURISTICS

compute in the first phase, see Section 1.3. In addition to the priority queue described above, a second priority queue, offering expansion of the current tree through edges of G', is managed in the second phase. For each expansion, the better offer is accepted and both queues are updated appropriately.

The information gained in the first phase can be used more economically if not only one, but a (constant) number of Steiner trees are computed in the second phase, using different terminals or vertices as starting points.

This heuristic can be implemented with time  $O(m + n \log n)$  and guarantees a performance ratio of 2.

#### 4.2.3 Some Experimental Results Of Path Heuristics

Although the path heuristics mentioned before were designed only to be used as a component of other algorithms (especially in combination with reductions), they yield reasonable results even on their own. In Table 4.1, we give average running time and average gap from the optimal solution for instances from SteinLib (see Section 1.2 for a description of the instance groups). The four algorithms compared are the following:

- **one-phase SPH:** One iteration of the simple shortest path heuristic, using the improvement of [PW02].
- **one-phase R-SPH:** Repetitive variant of the above: Start from up to 100 different starting points (terminals are preferred as starting points).
- **two-phase R-SPH:** Repetitive shortest path heuristic (up to 100 different starting points, terminals are preferred), using the restricted preprocessing of distances and  $T'_D(R)$  to achieve a worst-case running time of  $O(m + n \log n)$ .
- $T'_D(R)$ : The distance network heuristic, for comparison with the two phase R-SPH.

All variants use the final minimum spanning tree pruning step, described above in the beginning of Section 4.2.

One observes that due to the improvement of [PW02], the one phase SPH variant is very fast, but the quality of the results is relatively weak. For this reason, the repetitive variant is usually used. Here, we see that using the fast two-phase variant can sometimes improve the running time. Furthermore, we can give an  $O(m + n \log n)$  running time guarantee. The solutions quality produced by this twophase variant is often the same as for the one-phase variant. This is due to the fact that in many cases the computation is focussed very effectively by the application of Lemma 38, and in spite of the rigid running time guarantee, all necessary distances can be computed in the first phase. In these cases, differences in the quality of the solutions of the one-phase and the two-phase variant reflect only different decisions in situations when ties of paths of equal length are broken. A different situation happens for example for the WRP3 and WRP4 instances, where many vertices are similarly close to many terminals. In the one-phase variant this leads to comparably high running times, as distance values have to be rechecked again and again. In the two-phase  $O(m + n \log n)$  variant, many computations in the first phase have to be aborted. As a consequence, for many paths, a precomputed shortest path is not available and one has two use paths from  $T'_D(R)$  and the solution quality deteriorates.

As a comparison we included the minimum spanning tree in the distance network  $T'_D(R)$ . Here one sees that quality of the solutions is drastically improved with the two-phase variant, although the same asymptotic running time guarantee can be given.

instance	one pl	nase SPH	one pha	ase R-SPH	two pha	ase R-SPH	T	$_{D}^{\prime}(R)$
group	$O(r(m \cdot$	$+ n \log n))$	$O(r(m \cdot$	$+ n \log n))$	O(m -	$+ n \log n$ )	O(m -	$+ n \log n$ )
	time	gap in %	time	gap in %	time	gap in %	time	gap in %
1R	0.001	4.59	0.054	1.40	0.017	3.19	0.001	8.13
2R	0.002	5.82	0.114	2.36	0.032	4.42	0.002	8.98
D	0.002	3.25	0.176	1.27	0.128	1.43	0.004	5.16
E	0.007	4.58	0.521	1.60	0.382	1.58	0.009	7.77
ES10000FST	0.065	2.32	5.682	2.27	6.489	2.23	0.071	3.18
ES1000FST	0.005	2.41	0.446	2.22	0.452	2.13	0.005	3.26
I080	0.001	14.66	0.011	1.35	0.004	1.41	0.001	18.56
I160	0.001	17.52	0.047	1.83	0.013	1.92	0.001	22.36
I320	0.004	19.11	0.193	2.51	0.038	3.80	0.004	25.54
LIN	0.015	3.65	1.176	2.27	0.235	2.14	0.015	5.74
MC	0.001	3.75	0.072	2.39	0.069	2.44	0.002	5.94
TSPFST	0.003	1.39	0.267	1.05	0.323	1.08	0.003	1.95
VLSI	0.004	2.63	0.314	1.12	0.118	1.59	0.004	5.45
WRP3	0.007	0.0007	0.572	0.0003	0.059	0.08	0.001	49.08
WRP4	0.003	0.002	0.282	0.001	0.040	0.12	0.001	42.09
Х	0.017	1.05	0.778	0.34	0.285	0.34	0.053	1.05

Table 4.1: Experimental results of path heuristics.

Our implementation compares quite well with the implementation given in [PW02]. Although they use a faster computer (1.7 GHz Pentium 4), the running times for one iteration of the one-phase SPH are comparable (e.g., on the VLSI instances they need 8 milliseconds on average while we need only 4). The quality of the solutions is also similar. Note that one cannot expect exactly the same values, as there is some freedom in the choice of the starting vertex and in choosing a shortest path in the case of multiple shortest paths. For the two-phase variant they only implemented a simpler version that is much slower and runs into memory problems on larger instances (although they cite our paper where we also presented the fast variant).

## 4.3 Reduction-based Heuristics

Working with reductions, one often gets the impression that some of the tests are too cautious. Sometimes nice ideas for strengthening a test turn out to be not universally valid. Of course even the strangest exception is enough to make a reduction test completely useless for (direct) integration into an exact algorithm. But with respect to heuristics, the situation is fundamentally different: Here a much stronger orientation towards the frequent case can be adopted.

The idea used here is to support the normal (exact) reduction tests through some heuristic ones. It must be emphasized that the goal is not reducing the graphs by brute force, but only giving an impulse in situations where the exact reduction process is blocked, in order to activate it again. In this context, it is particularly advantageous if it can be assumed that the actions performed could have been carried out by a more powerful, but unknown exact test anyway.

A natural basis for such an approach is given by the test VR (see Section 3.4.1). This test is kept very cautious to make a comprehensible proof possible. Furthermore, one observes readily that in case the upper bound used is not optimal, the test could potentially perform more (exact) reductions if a

better upper bound was available. The idea is now to perform the usual actions of this test without an upper bound each time the other tests are blocked. At each application, a certain proportion of vertices is eliminated (directly or after replacing of incident edges) according to the same criteria as in the exact version of the test (sum of distances to the next two or three terminals). Motivated by the fact that for a large ratio r/n the alternative-based reductions are very successful anyway and the test VR is usually effective only for small r/n, the proportion of the vertices being eliminated is a function of n and r, getting smaller with growing r/n. With the additional postulation that during each application of the tests a constant percentage (say 5%) of vertices and edges is eliminated, the asymptotic time for all iterations together is the same as for the first one, namely  $O(m + n \log n)$ . To make sure that the instance is not made infeasible by the heuristic reductions, we further forbid direct elimination of vertices in the current tree  $T'_D(R)$ . The computation of  $T'_D(R)$  also yields as a side effect a guaranteed performance ratio of 2. We call this whole procedure **PRUNE**.

The idea of not eliminating the nodes of a Steiner tree can be further utilized by using a (good) heuristic solution instead of  $T'_D(R)$  for guiding the heuristic reductions. We use the implementation of repetitive SPH described in Section 4.2 (with a constant upper bound for the number of repetitions) for this purpose, but any other good solution would do, too. On the other hand, we make the actions of the heuristic reductions somewhat bolder, eliminating vertices only directly (without replacing of incident edges). Note also that even Steiner nodes of the guiding heuristic solution may be eliminated, but only by the exact tests; these tests are guaranteed not to deteriorate the optimum. We call this variant of the PRUNE heuristic **GUIDED-PRUNE**.

## 4.4 Relaxations and Upper Bounds

Computing lower bounds is not the only motivation for dealing with relaxations; the information obtained can also be used (among other things) to obtain upper bounds.

Consider the (directed) cut formulation  $P_C$  of the Steiner problem: Given an optimal solution  $\hat{x}$ of its linear relaxation  $LP_C$ , the complementary slackness conditions state that each edge  $[v_i, v_j]$  with  $\hat{x}_{ij} > 0$  has zero reduced cost. Assuming that there is some similarity between some optimal solutions of the integer program  $P_C$  and its linear relaxation  $LP_C$ , it is well motivated to search an (optimal) solution in a subgraph containing the edges with reduced cost zero.

The algorithm DUAL-ASCENT, attempting to construct an optimal solution for  $DLP_C$ , adjusts the reduced costs favorably. So it is very natural to search for a solution in the set of edges whose reduced costs are set to zero by this algorithm, an idea already used in [Won84, Voß92]. The auxiliary graph to be searched for a good solution need not contain all these edges; we have experimented with several schemes and gained the best overall results with a subgraph containing the (undirected edges corresponding to) edges on zero-cost ways (with respect to reduced costs) from the root to another terminal, although other variants are not inappropriate either.

Having chosen such an auxiliary graph, the key question is how to obtain an (optimal) solution for the corresponding instance. The structure of such instances is very suitable for the application of our PRUNE heuristics; in particular, there are often long chains of vertices that are replaced by long edges through the NTD<sub>2</sub> test, making other alternative-based reductions very effective; and the heuristic reductions do the rest of the job. We call the whole procedure of doing fast reductions, calling DUAL-ASCENT, determining a subgraph and performing a PRUNE heuristic in the subgraph **ASCEND-AND-PRUNE**.

Since we are working in a subgraph of G, the time bounds for the PRUNE heuristics (which are dominated by the worst case time of DUAL-ASCENT) are guaranteed in any case. Empirically, since

the PRUNE heuristics run extremely fast on the auxiliary graphs, this kind of computation of upper bounds should be performed after each call to DUAL-ASCENT.

Although the experimental solution quality of this heuristic is impressive, it still sometimes misses the optimum. We found out that in almost all such cases the reason is simply that the auxiliary graph does not contain an optimal solution (and not that the PRUNE heuristics do not find it). This observation suggests a supplementation of this heuristic: The Steiner tree found in the subgraph can be used as the guiding solution for a call to GUIDED-PRUNE in the original graph. In the cases mentioned, this approach often improves the solution value, frequently leading to the optimum.

By applying the idea of the PRUNE heuristics directly to the original graph, one can do without the auxiliary graphs altogether. Let *lower* and  $\tilde{c}$  be the lower bound and the reduced cost vector provided by DUAL-ASCENT and  $\hat{x}$  an optimal solution of  $P_C$  with value *optimum*. The inequality  $\tilde{c} \cdot \hat{x} \leq optimum - lower$  (see Lemma 29) strongly suggests that normally there cannot be many edges with large reduced costs in an optimal solution. This motivates another heuristic, **SLACK-PRUNE**, that basically follows the same scheme as GUIDED-PRUNE, but uses the criterion of the test DA for eliminating vertices. The guiding solution is computed by a call of PRUNE in the auxiliary graph described above, since the necessary information is available after performing DUAL-ASCENT anyway. The running time is dominated by the worst-case time of DUAL-ASCENT. Using the same arguments as in the case of PRUNE, one gets the time bound  $O(m \cdot \min\{m, nr\})$ . But in combination with reductions, the empirical times are much smaller than the above term suggests.

As in DUAL-ASCENT, dual feasible solutions and corresponding reduced costs for  $LP_C$  are calculated during the row generating algorithm (Section 2.9.2). This information can be used to generate auxiliary graphs similar to those in ASCEND-AND-PRUNE. But in this case there are not necessarily paths with reduced cost zero from the root to all terminals. The auxiliary graph in this context contains all vertices with the property that there is a path from the root over this vertex to another terminal not longer (with respect to reduced costs) than the longest shortest path from the root to another terminal. This auxiliary graph can be used as in ASCEND-AND-PRUNE.

A classical method for utilizing the information provided by linear relaxations is to use an ordinary heuristic in the original network with modified edge costs  $c'_{ij} = c_{ij}(1 - x_{ij})$  (where x is the primal solution of the current linear program). But this is not generally a good idea, because the structure of the primal solutions does not provide a good guide for a primal heuristic until the most advanced stages of the row generating algorithm.

These latter approaches only work in combination with explicit solution of linear programs and are therefore not suitable for fast, stand alone heuristics. But as a complement to the row generating strategy, they are frequently effective, especially in the advanced stages of the algorithm.

## 4.5 Combination of Steiner Trees

During the reduction process and especially while solving instances exactly, one usually gets several distinct heuristic solutions. In general, it is not the best idea to simply keep the best solution and forget the others. It is possible that solutions with a worse value are better locally, and one can try to keep the best part of each solution.

We have developed several techniques for realizing the idea above. One simple and effective way is to consider the graph consisting of the union of the edge sets of two (or more) Steiner trees. In this graph, one can call a (powerful) heuristic again or even try an exact solution. Such graphs have frequently several (nontrivial) biconnected components, which makes the (exact) solution considerably faster. Using such schemes, we frequently get improvements in solution values (in case they

#### 4.6. EXPERIMENTAL RESULTS AND EVALUATION

were not optimal anyway). Since the instances generated through such combinations (in the following called combination-instances) are almost always solved to optimality through (fast) reductions, these improvements are gained at no significant extra cost.

For the results reported in this section, we simply call a PRUNE heuristic in such combinationinstances; in particular, in the context of the heuristic SLACK-PRUNE we call the same heuristic (only without combinations) again in each combination-instance.

## 4.6 Experimental Results and Evaluation

In this section, we present some experimental results for reductions-based heuristics on instances from [SteinLib] (see Section 1.2 for a description of the instance groups). All of them are heuristics with a worst-case time describable by a polynomial of low order, as explained in the previous sections. Other good heuristics in the literature can be found in [Dui93, DV97, Esb95, Ver96, RUW02].

For comparison, we include the results of Ribeiro, Uchoa, and Werneck [RUW02], which are the best other results we are aware of. They use a combination of several techniques: A greedy randomized adaptive search procedure (GRASP), a weight perturbation strategy (which uses the knowledge of which edges are used frequently in many solutions), a local search procedure, a heuristic for the combination of good solutions, and many reduction techniques. Unfortunately, they include a preprocessing for the D, E, and VLSI-instances in their runs, but did not include the preprocessing time in the reported running times as we did. They report results only for those instances that could not be solved with their preprocessing routine, but we rescaled the values (assuming that instances solved by the preprocessing produced an optimal solution in zero time), so that at least the values for the average gaps are comparable. A stroke in the table means that no results were reported for these instances. The running times for [RUW02] where measured on a 300 MHz AMD and a Pentium II 400 MHz.

instance	P	RUNE	ASCE	ND-&-PRUNE	SLACK	-PRUNE	[RUW02]	
group	time	gap in %	time	gap in %	time	gap in %	time gap in %	
1R	0.13	1.36	0.07	1.03	0.22	0.00		_
2R	0.27	1.42	0.16	1.59	10.91	0.00		
D	0.10	0.07	0.07	0.02	0.14	0.00	> 11 0.04	
E	0.39	0.31	0.25	0.13	1.64	0.00	> 86 0.05	
ES10000FST	7.56	1.11	30.88	0.67	2101.89	0.38		
ES1000FST	0.57	1.01	0.38	0.53	18.57	0.19		
I080	0.07	1.15	0.02	1.65	0.43	0.06	> 2 0.01	
I160	0.31	1.97	0.07	1.69	1.68	0.10	> 16 0.13	
I320	1.63	2.84	0.30	1.81	7.46	0.14	> 108 0.15	
LIN	1.51	1.44	1.09	0.76	153.99	0.04		
MC	0.05	1.70	0.04	1.01	0.95	0.42		
TSPFST	0.21	0.42	0.38	0.31	32.01	0.04		
VLSI	0.33	0.39	0.40	0.35	7.17	0.004	> 830 0.01	
WRP3	0.15	0.0006	0.14	0.0003	17.79	0.00003		
WRP4	0.10	0.07	0.10	0.0006	6.19	0.00006		
Х	0.44	0.17	0.29	0.00	0.23	0.00		

Table 4.2: Some experimental results on upper bound calculation.

From Table 4.2, one can see that SLACK-PRUNE is clearly superior to the approach of Ribeiro, Uchoa, and Werneck. Even taking into account the different speed of the machines, our approach is faster in most cases. Remember also that in [RUW02] the preprocessing time was not recorded. Additionally, the quality of the solutions produced by SLACK-PRUNE is better, even on the larger incidence instances (I160, I320), which were constructed to be difficult for the known reduction techniques. Note that the percentage of pruned vertices is a simple parameter for the tradeoff between running time and solution quality, i.e., better solutions can be achieved simply by decreasing this parameter at the cost of longer running times.

The two heuristics PRUNE and ASCEND-AND-PRUNE are very fast while producing fairly good results. Note that PRUNE, although it has the better running time guarantee of  $O(m+n \log n)$ , in many cases takes longer than ASCEND-AND-PRUNE. The reason is that the techniques used in ASCEND-AND-PRUNE, in particular DUAL-ASCENT, are on the one hand much faster than the worst-case time bound suggests, and on the other hand reduce a problem instance much more efficiently.

# Chapter 5

# **Solving to Optimality**

## 5.1 Introduction

In this chapter, we present our results concerning the computation of optimal Steiner trees.

As with the upper bound calculations, the number of competitors is huge: For every classical approach (enumeration algorithms, dynamic programming, branch-and-bound, branch-and-cut) there are many published results and the presented algorithms are sometimes quite involved. Again, we point to [HRW92] for a survey and only mention the most important updates:

- Duin [Dui93] improved the preprocessing techniques and invented some heuristics that were integrated into a branch-and-bound algorithm, which was clearly the best at its time.
- Koch and Martin [KM98] used some of the improved preprocessing techniques and developed a very powerful and robust branch-and-cut algorithm.
- Uchoa et al. [UdAR02] contributed some new extended preprocessing techniques (which were further generalized in Section 3.5). Their branch-and-cut algorithm is currently, to the best of our knowledge, the best exact algorithm produced by other authors.

In the next section, we describe a dynamic programming approach that is useful for solving certain small subgraphs. After that we will describe the integration of the previously mentioned methods for reductions and upper and lower bound into an exact algorithm.

## 5.2 A Dynamic Programming Approach for Subgraphs

In this section, we present a practical algorithm for solving the Steiner problem in graphs with a small width-parameter (a formal definition of width is given in the next section). The running time of the algorithm is linear in the number of vertices when the width is constant, thus it belongs to the category of algorithms exploiting the fixed-parameter (FP) tractability of NP-hard problems. But the applicability of this algorithm is much broader in the context of our work. Due to the use of reduction techniques based on partitioning (Section 3.6) we can already profit from small width in subgraphs of the given instance. These techniques are in turn applicable to a very wide range of instances of the Steiner problem in networks after applying extended reduction methods (Section 3.5) and instances of geometric Steiner problems after FST-generation [PV03, WWZ00].

The width-concept here is closely related to pathwidth. (For an overview of subjects concerning pathwidth and the more general notion of treewidth see [Bod93].) There are already FP-polynomial algorithms for the Steiner problem in graphs; specifically, in [KS90] a linear-time algorithm for graphs with bounded treewidth is described. But this algorithm is more complicated than the one we present here, and its running time grows faster with the (tree-) width (it is given in [KS90] as O(nf(d)) with  $f(d) = \Omega(d^{4d})$ , where d is the treewidth of the graph); so it seems to be not as practical as our algorithm, and no experimental results are reported in [KS90]. In a different context (network reliability), a similar approach using pathwidth is described in [PT01], which is practical for a range of pathwidths similar to the one considered here. We also adapted this approach to the Steiner problem, but the experimental results were not as good as with the our approach.

A theoretically more powerful concept is the branch-width decomposition. It was formalized by Robertson and Seymour [RS91] and applied successfully to the TSP [CS02] and to the Steiner problem by Cook et. al. [Coo02]. We will not describe this approach, because experimental results showed that no gain can be expected from it: Cook was not able to solve many instances of the TSPFST group from [SteinLib], while we were able to solve them using a combination of our reduction techniques,

in particular the partitioning-based reductions (Section 3.6), and the Dynamic Programming approach for subgraph presented in this section.

## 5.2.1 Additional Definitions

A path-decomposition of a graph G = (V, E) is a sequence of subsets of vertices  $(X_1, X_2, \ldots, X_r)$ , such that

- 1.  $\bigcup_{1 \le i \le r} X_i = V,$
- 2.  $\forall (v, w) \in E \quad \exists i \in \{1, \dots, r\}: \quad v \in X_i \land w \in X_i,$
- 3.  $\forall i, j, k \in \{1, \dots, r\}: i \leq j \leq k \Rightarrow X_i \cap X_k \subseteq X_j.$

The **pathwidth** of a path-decomposition  $(X_1, X_2, ..., X_r)$  is  $\max\{|X_i| \mid 1 \le i \le r\} - 1$ . The **pathwidth** of a graph G is the minimum pathwidth over all possible path-decompositions of G.

## 5.2.2 The Algorithm

The outline of the algorithm is as follows: We maintain a set of already visited vertices and a subset of them (the **border**) that are adjacent to some non-visited vertex. In each step, the set of visited vertices is extended by one non-visited vertex adjacent to the border. For all possible partitions in each border, we calculate (the cost of ) a forest of minimum cost that contains all visited terminals with the property that each tree in the forest spans just one of the partition sets. We are finished when all vertices have been visited.

The motivation behind this approach is as follows: For any optimal Steiner tree T, the subgraph of T when restricted to the visited vertices is a forest, which also defines a partition in the border. The plan is to calculate these forests in a bottom-up manner, in each step using the values calculated in the previous step. If the size of all borders can be bounded by a constant, the total time can be bounded by the number of steps times another constant.

For an arbitrary fixed ordering  $v_1, \ldots, v_n$  of the vertices and any  $s \in \{1, \ldots, n\}$ , we define  $V_s := \{v_1, \ldots, v_s\}$  and denote with  $G_s$  the subgraph of G with vertex set  $V_s$ . In the following, we assume an ordering of the vertices with the property that all  $G_s$  are connected. (For example, a depth-first-search traversal of G delivers such an ordering.)

We denote with  $B_s$  the border of  $V_s$ , i.e.,  $B_s := \{v_i \in V_s \mid \exists (v_i, v_j) \in E : v_j \in V \setminus V_s\}$ . With  $L_s$  we denote the set of vertices that leave the border after step s, i.e.,  $L_s := (B_{s-1} \cup \{v_s\}) \setminus B_s$ . The inclusion of  $v_s$  in this definition should cover the case that  $v_s$  has no adjacent vertices in  $V \setminus V_s$ ; this simplifies some other definitions.

Consider a set  $Q, B_s \cap R \subseteq Q \subseteq B_s$ , and a partitioning  $\mathcal{P} = \{P_1, \ldots, P_t\}$  of Q into nonempty subsets, i.e.,  $\bigcup_{1 \leq i \leq t} P_i = Q$  and  $\emptyset \notin \mathcal{P}$ . For a partition  $\mathcal{P}$  and a set  $L \subseteq V$  we define  $\mathcal{P} - L :=$  $\{P'_i \mid P_i \in \mathcal{P}, P'_i = P_i \setminus L\}$ . Let  $F(s, \mathcal{P})$  be a forest of minimum cost in  $G_s$  containing all terminals in  $V_s$  and consisting of t (vertex-disjoint) trees  $T_1, \ldots, T_t$  such that  $T_i$  spans  $P_i$  for all  $i \in \{1, \ldots, t\}$ . With  $c(s, \mathcal{P})$  we denote the cost of  $F(s, \mathcal{P})$ .

Let  $V_0 = B_0 = \emptyset$  and set  $c(0, \emptyset) = 0$ . The value  $c(s, \mathcal{P})$  can be calculated recursively using a case distinction:

I)  $v_s \in Q$ :  $c(s, \mathcal{P}) = \min\{c(s-1, \mathcal{P}') + C \mid \mathcal{P}' = \{P_1, \dots, P_r\}, j \in \{0, \dots, r\},$   $\mathcal{P} = (\{\{v_s\} \cup \bigcup_{1 \le l \le j} P_l\} \cup \{P_{j+1}, \dots, P_r\}) - L_s,$   $\forall 1 \le l \le j : v_l \in P_l,$  $C = \sum_{1 \le l \le j} c(v_l, v_s)\},$ 

II)  $v_s \not\in Q$ :

$$c(s, \mathcal{P}) = \min \left\{ c(s-1, \mathcal{P}') \mid \mathcal{P} = \mathcal{P}' - L_s \right\}$$

The cost of an optimal Steiner tree in G is

$$\min \{ c(s, \mathcal{P}) \mid R \subseteq V_s, |\mathcal{P}| = 1 \}$$

Obviously the forests  $F(s, \mathcal{P})$  (and an optimal Steiner tree) can be calculated following the same pattern.

## 5.2.3 Dynamic Programming Implementation

By using the recursive formula above, the necessary values can be calculated in a bottom-up manner by memorizing, for each step s, the values  $c(s, \mathcal{P})$ . We assume  $c(s, \mathcal{P}) = \infty$  if no partition  $\mathcal{P}$  is calculated at step s. This leads to the following algorithm, written in pseudocode:

```
"r : number of visited terminals"
     s = 0; r = 0; opt = \infty;
1
\mathcal{2}
     c(s,\emptyset) = 0;
3
     while s < n:
            s = s + 1; determine v_s, B_s and L_s;
4
5
            if v_s \in R : r = r + 1;
            forall \mathcal{P} with c(s-1,\mathcal{P}) \neq \infty:
6
                  oldCost = c(s - 1, \mathcal{P});
\tilde{7}
                  if v_s \notin R and \emptyset \notin \mathcal{P} - L_s:
8
                        c(s, \mathcal{P} - L_s) = oldCost;
g
10
                   Pcandidates = \{ P_i \in \mathcal{P} \mid \exists v_i \in P_i : (v_i, v_s) \in E \};
                  forall Pconnect \subset Pcandidates:
11
                         connectionCost = \sum_{P_i \in Pconnect} \min_{v_i \in P_i, (v_i, v_s) \in E} c(v_i, v_s);
12
                         Pstay = \mathcal{P} \setminus Pconnect; \ Pnew = (\{\{v_s\} \cup \bigcup_{P_i \in Pconnect} P_i\} \cup Pstay) - L_s;
13
                        if \emptyset \notin Pnew and c(s, Pnew) > oldCost + connectionCost:
14
15
                               c(s, Pnew) = oldCost + connectionCost;
                               if r = |R| and |Pnew| = 1:
                                                                            "feasible Steiner tree"
16
17
                                      opt = \min(opt, c(s, Pnew));
```

## 5.2.4 Running Time

Let  $p_s$  denote the number of partitions at step s. We have

$$p_s = \sum_{R \cap B_s \subseteq Q \subseteq B_s} \sum_{i=1}^{|Q|} \left\{ \begin{matrix} |Q| \\ i \end{matrix} \right\} = \sum_{R \cap B_s \subseteq Q \subseteq B_s} B(|Q|),$$

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where B(b) is the *b*-th Bell number; so  $p_s = O(2^{b_s}B(b_s))$  with  $b_s := |B_s|$ . We only maintain one global list of partitions, which is updated after each step, keeping for each valid partition a solution of minimum cost. Because of the loop in line 11, this list can grow to at most  $l_s := 2^{b_s}p_s = O(2^{2b_s}B(b_s))$  partitions. Eliminating the duplicates can be done by sorting the list: Each partition can be represented as a (lexicographically) sorted string (of length at most  $2b_s$ ) of sorted substrings (of length at most  $b_s$ ) separated by some extra symbol. Using radix sort, all the individual sortings of  $l_s$  strings can be done in total time  $O(n + l_s b_s)$ . Sorting the resulting list of  $l_s$ strings takes again  $O(n+l_s b_s)$ . We set aside for now a total extra time of O(|E|) for the operations on edges; and assume that an ordering of vertices is given (these points are explained below). The (rest of the) operations in lines 12 - 17 can be carried out in time  $O(b_s)$ . This gives the total running time  $O(\sum_{s=1}^n b_s 2^{2b_s} B(b_s))$ . Note that this bound implicitly contains the extra amortized time O(|E|) by the following observation: After a vertex is visited for the first time, it remains in the border as long as it has some non-visited adjacent vertex; so each edge is accounted for by its first-visited endpoint.

Now if we can guarantee an upper bound b for the size of all borders, we have an upper bound of  $O(nb2^{2b}B(b))$  for the running time. Using the very rough upper bound  $(2b)^b$  for B(b) we get the running time:  $O(n2^{b\log b+3b+\log b})$ . This means that the algorithm runs in linear time for constant b and, for example, in time  $O(n^2)$  for  $b = O(\log n/\log \log n)$ .

For the actual implementation, some modifications are used. For example, avoiding duplicate partitions is done using hashing techniques, which reduces the amount of necessary memory.

## 5.2.5 Ordering the Vertices

In Section 5.2.6 we will show that finding an ordering of vertices such that the maximum border size equals b is (up to some easy transformations) equivalent to finding a path-decomposition of pathwidth b. The problem of deciding whether the pathwidth of a given graph is at most b, and if so, finding a path-decomposition of width at most b is  $\mathcal{NP}$ -hard for general b [ACP87], but for constant b, this problem can be solved in linear time [Bod96]. However, already for b > 4 the corresponding algorithm is no longer practical [R oh98], and it seems that no practical exact algorithm is known for more general cases [Bod02]. Furthermore, we have a more specific scenario; for example we differentiate between terminals and nonterminals. So for the actual implementation we use a heuristic, which has produced quite satisfactory results for our applications. The heuristic chooses in each step a vertex  $v_s$  adjacent to the border using a (ad hoc) priority function of the following parameters:

- size of resulting set  $L_s$ ,
- number of visited vertices in the adjacency list of  $v_s$ ,
- is  $v_s$  a terminal (1/0),
- number of edges connecting  $V_s$  and  $V \setminus V_s$ .

We select the starting vertex by trying a small number of terminals and performing a sweep through the graph without actually computing the partitions, but by estimating their number using another (ad hoc) function and selecting the one with the smallest overall value.

A straightforward implementation of this heuristic needs time  $O(n^2)$  for all choices. This bound could be improved using advanced data structures for priority queues and additional tricks, but the ordering has not been the bottleneck in our applications; and theoretically a better (linear for constant b as in our applications) time bound for path-decomposition is available anyway.

## 5.2.6 Relation to Pathwidth

In this section, we show that every path-decomposition with pathwidth k delivers a sequence of borders  $B = (B_1, \ldots, B_s, \ldots, B_n)$  such that  $\max\{|B_s| \mid 1 \le s < n\} \le k$  and vice versa.

First, note that the 3rd condition in the definition of path-decomposition can be rewritten as follows: There are functions *start*, *end*:  $|V| \rightarrow \{1, ..., r\}$  with  $v \in X_j \Leftrightarrow start(v) \leq j \leq end(v)$ . We call a path decomposition bijective if the manning start is a bijection

We call a path-decomposition **bijective** if the mapping *start* is a bijection.

**Lemma 39** Every path-decomposition  $X = (X_1, \ldots, X_r)$  can be transformed to a bijective pathdecomposition  $(X'_1, \ldots, X'_n)$  of no larger pathwidth.

**Proof:** We modify the sequence *X* as follows:

As long as there is some i with  $start^{-1}(\{i\}) = \emptyset$ , remove  $X_i$  from X; adapting the functions start and end.

As long as there are  $v \neq v'$  with start(v) = i = start(v'), define  $X_{j+1} = X_j$  for all  $j \geq i$  and remove v' from  $X_i$ ; adapting the functions start and end.

One easily observes that the final resulting sequence X' satisfies the properties for a pathdecomposition, is bijective, and has no greater pathwidth.

We call a path-decomposition with functions start, end minimal if it holds:

$$end(v) \ge i \Rightarrow start(v) = i \lor \exists (v, w) \in E : start(w) \ge i.$$

(Note that the other direction is satisfied for every path-decomposition.)

**Lemma 40** Every (bijective) path-decomposition X can be transformed to a minimal (bijective) path-decomposition of no larger pathwidth.

**Proof:** For every v, set  $end(v) := \max\{start(w) \mid (v, w) \in E \lor v = w\}$ . Delete v from all  $X_i$  with i > end(v). One easily observes that the resulting sequence satisfies the properties for a path-decomposition, has no greater pathwidth, is minimal, and remains bijective if the original decomposition was bijective.

**Lemma 41** Let  $(X_1, \ldots, X_n)$  be a minimal, bijective path-decomposition of G with the functions start and end. Assume that the vertices are ordered according to their start values, i.e.,  $start(v) = s \Leftrightarrow v \in V_s \setminus V_{s-1}$ . For each  $s \in \{1, \ldots, n\}$  it holds:  $X_s = \{v_s\} \cup B_{s-1}$ .

#### **Proof:**

$$\begin{array}{ll} v \in X_s & \Leftrightarrow \quad start(v) \leq s \wedge end(v) \geq s \\ & \Leftrightarrow \quad (start(v) = s \vee start(v) < s) \wedge (start(v) = s \vee \exists (v,w) \in E : \ start(w) \geq s) \\ & \Leftrightarrow \quad start(v) = s \vee (start(v) < s \wedge \exists (v,w) \in E : \ start(w) \geq s) \\ & \Leftrightarrow \quad v = v_s \vee (v \in V_{s-1} \wedge \exists (v,w) \in E : \ w \in V \setminus V_{s-1}) \\ & \Leftrightarrow \quad v = v_s \vee v \in B_{s-1} \end{array}$$

It follows that every path-decomposition of G can be transformed to a path-decomposition  $X = (X_1, \ldots, X_n)$  of no larger pathwidth such that for an ordering of vertices according to the *start* function of X it holds:  $X_s = \{v_s\} \cup B_{s-1}$ . On the other hand, it is easy to verify that each ordering of vertices and the corresponding sequence of borders  $(B_1, \ldots, B_n)$  deliver a (minimal, bijective) path-decomposition X by setting  $X_s = \{v_s\} \cup B_{s-1}$ . In each case, we have:  $\max\{|X_s| \mid 1 \le s \le n\} - 1 = \max\{|B_{s-1}| \mid 1 \le s \le n\}$ .

## **5.3** Putting the Pieces Together: The Exact Algorithm

In this section we describe the synthesis of an exact algorithm from the components described in the previous sections.

## 5.3.1 Interaction of the Components

A central feature of our exact algorithm is that the various components (reduction tests, lower bounds and upper bounds) do not act independently of each other, as described in detail in previous sections:



Figure 5.1: Interaction of the different components.

The bound-based reductions depend on upper and lower bounds; and the computation of upper and lower bounds profits from reductions, both in terms of running time and quality of results. The idea behind reduction tests is also the central part of the reduction-based heuristics for computing upper bounds. Further we use the structure of heuristic solutions (corresponding to good upper bounds) to guide the computation of lower bounds; and the information gained during the computation of lower bounds is used to guide the computation of upper bounds. All in all, there is a mutual dependence between the three major components: reductions, upper bounds, and lower bounds. This is not a drawback, but an advantage: The scenario is that performing (alternative-based) reductions accelerates the computation of upper and lower bounds and enhances their qualities; the information gained during the computation of bounds is used to reduce the instance further (using bound-based reductions), and then the whole pattern repeats. We call this whole process the reduction process, beginning with fast reductions and switching to more and more powerful ones as the process advances. This strategy is not only a major reason for the short solution times our algorithm very often achieves, but also enables instances to be solved that we could not solve in a reasonable time otherwise. Note especially that the value of the lower bound corresponding to a certain relaxation can be enhanced through reductions; this helps to solve instances that otherwise could not be solved (without branching) using the same techniques for computing upper and lower bounds.

For the experimental results given in this chapter, we use the following components: For computing lower bounds, we use the relaxation  $LP_C$  through the algorithm DUAL-ASCENT and, in advanced stages, row generation with  $LP_{C'}$  (Section 2.9.2) or, if the proportion of terminals to all vertices is high, the Lagrangian relaxation  $LaP_{T_0}$  (Section 2.9.1). To reduce the instances, we use all described alternative-based techniques (Section 3.3). In addition, we use the bound-based techniques DA (Section 3.4.2) and, in combination with row-generation, the test RG (Section 3.4.3). Furthermore, we use the extended reduction techniques (Section 3.5) and the partitioning-based reductions (Section 3.6). For computing upper bounds we use our PRUNE heuristics (Sections 4.3, 4.4), including the combination of Steiner trees (Section 4.5). Before starting the time-consuming RG reduction test, we check with the heuristic mentioned in Section 5.2 whether we can solve the (sub-)instance with the pathwidth approach.

As described above, the fast methods are applied first, with switching to more time-consuming ones only if an instance has not already been solved to optimality. Apart from this general principle, the exact ordering of the components has usually not been critical.

## 5.3.2 Branch-and-Bound

The reduction process described in the previous subsection is an extremely powerful device, but it is not guaranteed to solve every instance of the problem. To get an exact algorithm, we integrate it into a branch-and-bound framework. But one should not be misled by the name *branch*-and-bound: Branching is something we generally (and often successfully) try to avoid, it is only a safety net in case the reduction process is blocked. This also means that we invest a lot of work in each node of the branch-and-bound tree to keep the tree small, and do not try to gain speed by limiting the work in each node.

We use binary, vertex-oriented forward branching [HRW92]. Both depth-first and best-first search strategies are available in our implementation, with depth-first as default. There are usually not many nodes in our branch-and-bound trees anyway; moreover, only the currently processed node needs to be kept in the main memory.

As the branching variable, we choose the non-terminal with the largest degree in the best available Steiner tree. The intuitive motivation for this choice is an intensification of the search in an area where a good solution has been found (in case of inclusion) and a diversification of the search to other areas (in case of exclusion). This strategy also supports the building of several blocks (biconnected components). It is known [HRW92] that in case several blocks exist, the problem can be solved by solving the instances corresponding to each intermediate block separately, which generally reduces the total running time substantially. Although it usually cannot be assumed that the original instance is not biconnected, this often changes later during the reduction process and after branching. We use this fact frequently in our algorithms: Whenever a more time-consuming part is to be performed, we check whether the graph is biconnected. If this is not the case, we solve the corresponding subinstances separately and transform the information gained back to information for the original instance. Here one can use the following observation to identify the blocks that must be considered:

**Lemma 42** Let T be a Steiner tree with all leaves being terminals in a network G. A block of G is intermediate if and only if it contains an edge of T.

**Proof:** For a block B to be intermediate, there must be two terminals  $z_k$  and  $z_l$  such that every path between  $z_k$  and  $z_l$  contains an edge in B. Hence, every Steiner tree must contain an edge in B. Conversely, consider a Steiner tree T with all leaves being terminals that contains an edge in a block B. So there are two terminals  $z_k$  and  $z_l$  such that at least one path between  $z_k$  and  $z_l$  contains an edge in B. If  $z_k$  (or  $z_l$ ) is in B and it is not an articulation point, B is obviously intermediate. Otherwise there must be two articulation points  $v_i$  and  $v_j$  of B such that a path between  $z_k$  and  $v_i$  and a path between  $v_j$  and  $z_l$  contain no edge in B. Now suppose B is not intermediate. Then there is a path between  $z_k$  and  $z_l$  that does not contain an edge in B. Hence, there is also a path between  $v_i$  and  $v_j$ that has no edge in B, which contradicts the definition of B as a biconnected component.

## **5.4 Summary on Experimental Results**

Here we report on what the components presented achieve together, acting as an "orchestra". As stated in Section 1.2, results for different types of instances from the benchmark SteinLib are presented, including the instances of the OR-Library. Here we give only averages for some groups of instances. In the appendix, we report on the results for each instance separately.

We leave it mainly to the reader to compare the given running times to those of other exact algorithms in the literature (see for example [Bea89, BL98, CGR92, Dui93, KM98, Uch01]). As an orientation, we provide in Table 5.1 the average times (in seconds) for the exact solution of some instances groups, which have frequently been used by other authors. It is true that we used the fastest machine (it is approximately half as fast as the PC we used in Section 5.4.1); however improved machine speed explaines our improved times only to a small extend.

instance-	[Bea89]	[BL98]	[CGR92]	[Dui93]	[KM98]	[Uch01]*	here
group	Cray X-MP	SG Indigo	VAX 8700	i486	Sun Sparc 20	UltraSparc II	Sunfi re
D	556	3545	14260	176	117	> 4	0.2
E		_		_	4415	> 181	1.7
$TAQ^{\dagger}$		_		_	197	4	0.08
TAQ		—		—		162	1.1

Table 5.1: Average running times of different exact algorithms.

We solved all instances that (to our knowledge) have been solved before. Furthermore, we solved 32 instances from different instance classes from SteinLib (geometric (TSPFST), VLSI (LIN), wire routing problems (WRP3), constructed difficult instances (I640, PUC, SP) that have not been solved by other authors. See the appendix for details.

## 5.4.1 Experimental Results on Geometric Instances

As mentioned already in Sections 1.4.2 and 3.6.1 for geometric Steiner problems, an approach based on full Steiner trees (FSTs) has been empirically successful [WWZ00]: In the first phase, the FST generation phase, a set of FSTs is generated that is guaranteed to contain an SMT. In the second phase, the FST concatenation phase, one chooses a subset of the generated FSTs whose concatenation yields an SMT. Empirically, the second phase was usually the bottleneck, where originally methods like backtracking or dynamic programming have been used. A breakthrough occured when Warme [War98] used the fact that FST concatenation can be reduced to finding a minimum spanning tree in a hypergraph whose vertices are the terminals and whose (hyper-)edges correspond to the generated FSTs. We use a different approach: By building the union of (the edge sets of) the FSTs generated in the first phase, we get a normal graph and the FST-concatenation problem is reduced to solving the classical Steiner problem in this graph. Note that in this way the useful information about the individual FSTs is lost. Still, our approach is much faster. In Table 5.2, we compare for the second phase the running times of our program and of the program package GeoSteiner [WWZ01]. For more detailed results on this, see [PV03, PV01d].

<sup>\*</sup>In [Uch01] only running times for 5 of the 40 D/E instances are reported, we estimated the averages by assuming that all other instances were solved in zero time.

<sup>&</sup>lt;sup>†</sup>Excluding instances not solved by [KM98].

For both programs all computation was performed on a PC with an AMD Athlon XP 1800+ (1.53 GHz) processor and 1 GB of main memory, using the operating system Linux 2.4.9. We used the gcc 2.96 compiler and CPLEX 7.0 as LP-solver.

instance	GeoSteiner	our program
group	time (s)	time (s)
ES1000FST	150.6	10.3
TSPFST <sup>‡</sup>	261.8	3.0

Table 5.2: Comparison of GeoSteiner (2nd phase) and our program (average running time for exact solution).

Additionally, we are able to solve some instances (fl1400, fl3795, fnl4461) that have not been solved before. In Table 5.3, we show results for instances not solved by GeoSteiner in one day.

instance		size		optimum	our program
	R	V	E		time (s)
es10000	10000	27019	39407	716174280	758
fl1400	1400	2694	4546	17980523	118
fl3795	3795	4859	6539	25529856	139
fnl4461	4461	17127	27352	182361	6148
pcb3038	3038	5829	7552	131895	2.4
pla7397	7397	8790	9815	22481625	0.1

Table 5.3: Instances not solved by GeoSteiner in one day.

## 5.5 Concluding Remarks

We have presented several algorithmic contributions for solving the Steiner problem in networks. The experimental results strongly recommend the chosen approach based on reductions and underline the utility of the techniques presented here. In particular, the reduction-based heuristics have proven to be extremely strong and robust. Also, the running times of the exact algorithm are often surprisingly small; and for many instances, there is not much room left for improvements. But this is not always the case:

For some instances, fast reductions come to a halt at a time when the relaxations used are still not strong enough; this is for example the case for some of the MC- and I-instances, which were insightfully constructed to fool the currently available techniques. Here, the algorithm has gone into branching to solve the instances exactly. Until new techniques are developed, instances of this type with thousands of vertices will not be solvable in reasonable time. But the results for the other groups of instances seem to indicate that such cases rarely arise naturally.

With respect to instances that were not artificially constructed, even instances with tens of thousands of vertices can usually be solved in reasonable time. Of course, very large instances are still a challenge. There are two main problems:

First, even though the running time and memory consumption for many components can be described by polynomials of low degree, they can grow to a critical value for very large instances. Especially, the  $\Theta(rn)$  memory consumption of the DUAL-ASCENT procedure, which is used relatively

<sup>&</sup>lt;sup>‡</sup>Excluding instances not solved by GeoSteiner.

#### 5.5. CONCLUDING REMARKS

early in the reduction process, can be a limiting factor. To overcome this problem, the development of new fast reduction techniques could be a solution. If they combine low memory consumption, fast (empirical) execution times and relevant reduction results, they could prepare the ground for currently available reduction techniques.

A second reason is even more decisive. As already discussed in Section 3.6.3, the LP-based approaches rely on the quality of the Linear Programming relaxation used, but any LP-relaxation of polynomial size is bound to have some (integrality) gap in this context. In larger instances, such gaps can accumulate and become more and more relevant. In particular, the important bound-based reductions lose their impact if the available bounds are too weak. There are several approaches to address this:

- **Branch-and-cut:** Currently, we do not use the branch-and-cut approach to improve the quality of the lower bound calculation. The reason is that we found it more advantageous to use the calculated lower bound to reinforce the reduction process, i.e., we use the lower bound and reduced costs for bound-based reductions and then try all reduction tests on the reduced graph. This procedure is often sufficient to solve problem instances without branching. Even if branching is necessary, the whole reduction processes can try to make use of the vertex branching, which is not the case in a branch-and-cut setting, because for many operations performed by reduction tests (e.g., replacement of a vertex by a clique over its neighbors, see Section 3.3.2) there are no methods available to translate them profitably into the linear program maintained by the branch-and-cut algorithm. On the other hand, our approach has the disadvantage that the generated constraints will be discarded after the row generation test (RG) is finished and for the next RG test, they have to be computed from scratch. For medium sized instances this disadvantage is not important, as the row generation is not too time consuming. For larger instances where most of the time is spent solving linear programs, a better integration of the reductions into a row generation or branch-and-cut approach could be an improvement.
- **Using tighter relaxations:** In Section 2.9.3, we showed how to use a tighter linear relaxation efficiently. Other, even tighter relaxations are available in the hierarchy of relaxations, presented in Section 2.8, but currently no technique is at hand to use them in such an efficient way that they could be applied to larger problem instances.
- **Improving linear relaxations outside the template paradigm:** We have already discussed two advanced techniques, the local cuts approach and the graph expansion approach, in the concluding remarks on lower bounds, Section 2.11. A further development of these techniques could enlarge the range of solvable instances.

For geometric instances, we have shown that a combination of full Steiner tree (FST) approach (see the results in the last section) that exploits the geometric properties of the problem and our algorithm for the Steiner problem in general networks is currently the most successful approach. However, it should be possible to improve on this, as a lot of useful information is discarded when the set of FSTs is converted into an instance for the network Steiner tree problem. Exploiting this information in our approach, or integrating our reduction techniques into the FST approach could be fruitful.

Another algorithmic challenge for Steiner tree algorithms is the development of algorithms for reliable computation [Meh02], i.e., algorithms that come with some kind of evidence that they produce correct results. As a formal verification of the whole program is unlikely, certifying algorithms as components would already be an advantage. Such algorithms return additional output (frequently

called a witness) that enables a (fast) verification of the results. A weak version of a witness is mentioned in Section 3.7: After all reductions are performed, the algorithm can transform a tree in the reduced instance into one for the unreduced instance, thus certifying that a tree with the same value exists in the original instance. Of course, this says nothing about the optimality of this tree. Other paradigms of reliable computation, namely "exact arithmetic" and "test and repair" have been mentioned in the context of using linear programming packages for reductions in Section 3.4.3. We list some of the major problems for certifying algorithms in the context of reduction techniques:

• A difficulty arises from a pattern that is frequently used in the efficient implementation of reduction tests: First, in a preprocessing step, some data structures are set up (e.g., representing a Steiner tree produced by some heuristic, a minimum spanning tree T'<sub>D</sub>(R) in the distance network, approximations of distances and bottleneck Steiner distances, a lower bound and reduced costs). Then, reduction tests use these data and modify the network. These modifications are done in such a way that the precomputed information is still valid (e.g., the extension of the PT<sub>m</sub>-test to the case of equality in Section 3.3.1) or that invalid parts are marked such they do not cause harm (e.g., the neighbor lists used for tagging precomputations as invalid if an edge is deleted in Section 3.5.5).

Even if it was possible to verify that the data structures were correct for the original network, it could be very difficult to verify that they are used correctly after the network is modified. A simple example is the following: When the first edge is deleted by the  $PT_m$ -test (Section 3.3.1), it is easy to verify that a path exists with Steiner distances not longer than the length of the edge by following the parent pointers in the shortest path tree and the  $T'_D(R)$  edges. For edges that are deleted later, this is no longer possible directly, as edges on this path may have been deleted.

• For many certifying algorithms the running times for producing the witness and verification of the result is (at least asymptotically) dominated by the running time for the algorithm. For the presented reduction tests with running time  $O(m + n \log n)$  this is not an easy task. Note that one of the major contributions of this work are the techniques for performing reduction tests so fast that they can be applied extensively. As a consequence, if a certifying reduction test is much slower than the original version, it cannot be applied in the same way as before.

In the example above, checking the edges of the alternative path can take time O(n), while the test condition of the  $PT_m$  test can be checked in nearly constant time.

Finally, it is an interesting open question whether the concepts presented in this work can be adapted to other combinatorial optimization problems. Promising candidates for such a recycling of ideas are the reduction-based heuristics (Section 4.3) and extended reduction techniques (Section 3.5) for problems where reduction techniques are already known, and partitioning as a reduction technique (Section 3.6) and graph expansion for improving lower bounds (sketched in Section 2.11) for general optimization problems in networks.

## **Appendix A**

# **Experimental Results of the Program Package**

Here we report on what the components presented achieve together, acting as an "orchestra". In Tables A.2-A.11 we report on each instance of SteinLib, except for the instance groups I080, B, C, P, and ES10FST–ES500FST, which are too easy. Overall we report on 831 instances.

All results were produced by a single run of the same program with the same parameter values, with the exception of the instance groups I and PUC. These instances were constructed with the aim of being difficult for known techniques, thus currently many of them can only be solved using many branching nodes in a branch-and-bound framework. Therefore, we made the solution faster by omitting the time consuming row generation method (Section 2.9.2) in the reduction process. Note that instances that arise naturally from some application can often be solved without branching by using the sophisticated reduction techniques.

For each instance, we give the value of the optimal solution and the total time until the exact solution of the instance (in seconds). We set a time limit of five hours on each run. Within this time, we have solved most of the considered instances. Only 64 instances have not been solved within the time limit. For these instances we give the upper and the lower bound after five hours (in italics). However, 9 of these instances could be solved using stronger extended reductions (see Section 3.5), and 12 could be solved by longer runs. For them we give the time for the exact solution in Tables A.14 and A.15. In total, there are only 43 unsolved instances left. These instances are from the instance groups SP, PUC, and I640, which where constructed with the aim of being difficult for known techniques.

We solved all instances that (to our knowledge) have been solved before. Furthermore, we solved 32 instances from different instance classes from SteinLib that have not been solved by other authors (see Table A.1).

In many cases our program performs several orders of magnitude faster than programs of other authors, see Section 5.4 for a short comparison with others' results. It is acknowledged to be the strongest program for solving general Steiner problem instances at the time being [CD01, SteinLib].

All results in this work were produced single-threaded on a Sunfire 15000 with 900 MHz SPARC III+ CPUs, using the operating system SunOS 5.9. We always used the GNU g++ 2.95.3 compiler with the -O4 flag. As it is a multi-processor machine with shared memory, it is slower than a single processor system with the same processor. A comparison with the running times reported in Section 5.4.1 shows that the machine is approximately half as fast as a PC with an AMD Athlon XP 1800+ (1.53 GHz) processor.

Instance Group	Table	Instances
LIN	A.5	lin31, lin32, lin33, lin34, lin35, lin36, lin37
WRP	A.4	wrp3-83
TSPFST	A.6	fl1400fst, fl3795fst
PUC	A.12	bipe2p, bipe2u, cc5-3p, cc5-3u, hc8p, hc8u
SP	A.13	w3c571
I640	A.11	i640-211, i640-212, i640-213, i640-214, i640-215,
		i640-211, i640-212, i640-213, i640-214, i640-215, i640-321, i640-322, i640-323, i640-324, i640-325,
		i640-341, i640-342, i640-343, i640-344, i640-345

Table A.1: Instances solved by us and not solved by others.

	i i	Size		Opt	Time	Instance	I	Size		Opt	
stance	V		R	Opt	Time	mstanee	V		R	Opt	
101	1000	1250	5	106	0.1	e01	2500	3125	5	111	
102	1000	1250	10	220	0.1	e02	2500	3125	10	214	
103	1000	1250	167	1565	0.1	e03	2500	3125	417	4013	
104	1000	1250	250	1935	0.1	e04	2500	3125	625	5101	
105	1000	1250	500	3250	0.1	e05	2500	3125	1250	8128	
106	1000	2000	5	67	0.3	e06	2500	5000	5	73	
d07	1000	2000	10	103	0.3	e07	2500	5000	10	145	
d08	1000	2000	167	1072	0.1	e08	2500	5000	417	2640	
d09	1000	2000	250	1448	0.1	e09	2500	5000	625	3604	
d10	1000	2000	500	2110	0.1	e10	2500	5000	1250	5600	
d11	1000	5000	5	29	0.2	e11	2500	12500	5	34	
d12	1000	5000	10	42	0.1	e12	2500	12500	10	67	
d13	1000	5000	167	500	0.1	e13	2500	12500	417	1280	
d14	1000	5000	250	667	0.1	e14	2500	12500	625	1732	
d15	1000	5000	500	1116	0.1	e15	2500	12500	1250	2784	
d16	1000	25000	5	13	0.2	e16	2500	62500	5	15	
d17	1000	25000	10	23	0.2	e17	2500	62500	10	25	
d18	1000	25000	167	223	0.7	e18	2500	62500	417	564	
d19	1000	25000	250	310	0.5	e19	2500	62500	625	758	
d20	1000	25000	500	537	0.1	e20	2500	62500	1250	1342	

Table A.2: Results on the D and E-instances. Type: Sparse random with varying graph parameters, OR-Library.

Instance		Size		Opt	Time	Instance	1	Size		Opt	Time
	V	E	R				V	E	R		
1r111	625	2352	6	28000	0.2	2r111	1000	5800	9	28000	1.0
1r112	625	2352	6	28000	0.1	2r112	1000	5800	9	32000	1.0
1r113	625	2352	6	26000	0.1	2r113	1000	5800	9	28000	0.5
1r121	625	2340	6	36000	0.1	2r121	1000	5766	9	28000	0.2
1r122	625	2342	6	45000	0.7	2r122	1000	5772	9	29000	0.5
1r123	625	2343	6	40000	0.3	2r123	1000	5754	9	25000	0.7
1r131	625	2336	6	43000	0.4	2r131	1000	5726	9	27000	0.4
1r132	625	2340	6	37000	0.1	2r132	1000	5725	9	33000	6.6
1r133	625	2326	6	36000	0.1	2r133	1000	5729	9	29000	0.6
1r211	625	2352	31	77000	0.5	2r211	1000	5800	50	89000	384.5
1r212	625	2352	30	81000	0.5	2r212	1000	5800	49	80000	3.7
1r213	625	2352	29	70000	0.2	2r213	1000	5800	48	76000	45.9
1r221	625	2341	31	79000	0.3	2r221	1000	5764	50	83000	4.5
1r222	625	2343	31	68000	0.1	2r222	1000	5765	50	84000	39.7
1r223	625	2340	31	77000	0.2	2r223	1000	5770	49	84000	74.0
1r231	625	2331	30	80000	0.2	2r231	1000	5737	50	86000	51.0
1r232	625	2335	29	86000	0.2	2r232	1000	5733	49	87000	71.4
1r233	625	2327	31	71000	0.1	2r233	1000	5730	47	83000	18.3
1r311	625	2352	56	108000	0.2	2r311	1000	5800	95	129000	70.6
1r312	625	2352	60	113000	0.2	2r312	1000	5800	92	126000	78.9
1r313	625	2352	58	106000	0.2	2r313	1000	5800	97	128000	41.7
1r321	625	2338	59	118000	0.2	2r321	1000	5771	92	125000	2.1
1r322	625	2336	60	113000	0.2	2r322	1000	5753	92	130000	34.0
1r323	625	2341	60	117000	0.3	2r323	1000	5764	96	142000	92.9
1r331	625	2319	58	103000	0.1	2r331	1000	5736	93	134000	26.2
1r332	625	2333	58	109000	0.1	2r332	1000	5745	95	136000	130.7
1r333	625	2331	58	113000	0.1	2r333	1000	5741	98	143000	100.3

Table A.3: Results on the 1R and 2R-instances. Type: 2D (respectively 3D) cross grid graph [Fre97].

nstance		Size		Opt	Time	Instance C'	0
	V	E	R	*		Instance Size $ V   E $	Opt R
vrp3-11	128	227	11	1100361	0.1		1 11001
vrp3-12	84	149	12	1200237	0.1		3 13007
vrp3-13	311	613	13	1300497	0.5		4 14002
vrp3-14	128	247	14	1400250	0.1	wrp4-15 193 369	5 15004
vrp3-15	138	257	15	1500422	0.1		6 16011
vrp3-16	204	374	16	1600208	0.2		7 17005
vrp3-17	177 189	354 353	17 19	1700442 1900439	0.1 0.1	wrp4-18 211 380	8 18014
vrp3-19		353 454	20				9 190144
vrp3-20 vrp3-21	245 237	454 444	20 21	2000271 2100522	0.3 0.2		21 210328
vrp3-21 vrp3-22	237	444	21	2100322 2200557	0.2	wrp4-22 294 568	220039
vrp3-22 vrp3-23	132	230	22	2200337 2300245	0.5	wrp4-23 257 515	23 23003
vrp3-23 vrp3-24	262	487	23 24	2300243	0.1	wrp4-24 493 963	24033
vrp3-24 vrp3-25	262	487 468	24 25	2400623 2500540	0.7		25008
vrp3-25 vrp3-26	402	468 780	25 26	2500540 2600484	0.2		26 26004
vrp3-26 vrp3-27	370	721	20	2700502	1.2		27 27004
vrp3-27 vrp3-28	307	559	27	2700302 2800379	0.3		28 28004
vrp3-28 vrp3-29	245	436	28 29	2800379 2900479	0.3		29 29004
vrp3-29	467	896	30	3000569	3.6		30005
vrp3-30	323	592	31	3100635	0.8		31005
vrp3-33	437	838	33	3300513	0.6		32005
vrp3-33	1244	2474	34	3400646	397.0		33 33006
vrp3-34	435	818	36	3600610	4.2		34005
vrp3-37	1011	2010	37	3700485	181.1		350060
vrp3-38	603	1207	38	3800656	17.7		36005
vrp3-39	703	1616	39	3900450	725.6		37006
vrp3-41	178	307	41	4100466	0.8		38 38006
vrp3-42	705	1373	42	4200598	46.4		39037
vrp3-43	173	298	43	4300457	0.8		40007
vrp3-45	1414	2813	45	4500860	592.5		41006
vrp3-48	925	1738	48	4800552	24.7		420070
rp3-49	886	1800	49	4900882	131.3		430150 440150
vrp3-50	1119	2251	50	5000673	2769.9		440150
vrp3-52	701	1352	52	5200825	207.7		450072 460075
vrp3-53	775	1471	53	5300847	7.3		470131
vrp3-55	1645	3186	55	[5500887-	-5500890]		470131
vrp3-56	853	1590	56	5600872	53.3	wrp4-49 557 1080	480222
vrp3-60	838	1763	60	6001164	262.2		50 500162
vrp3-62	670	1316	62	6201016	68.5		51 510161
vrp3-64	1822	3610	64	6400931	1549.1		52 520108
vrp3-66	2521	4858	66	6600922	4483.6		52 520108
vrp3-67	987	1923	67	6700776	46.4		54 540153
vrp3-69	856	1621	69	6900841	21.3		55 550195
vrp3-70	1468	2931	70	7000890	208.1		56 56022
rp3-71	1221	2414	71	7101028	248.3		58 58014
rp3-73	1890	3613	73	7301207	7104.5		59 590159
vrp3-74	1019	1941	74	7400759	263.0		60017
vrp3-75	729	1395	75	7501020	11.5		61022
vrp3-76	1761	3370	76	7601028	865.7		62021
vrp3-78	2346	4656	78	7801094	3306.0		63014
/rp3-79	833	1595	79	7900444	28.0		640199
vrp3-80	1491	2831	80	8000849	212.2		66029
vrp3-83	3168	6220	83	[8300888-			670280
/rp3-84	2356	4547	84	8401094	1243.2		68017
vrp3-85	528	1017	85	8500739	564.8		690232
vrp3-86	1360	2607	86	86000746	677.7		70 700302
vrp3-88	743	1409	88	88001175	26.4		71 710232
vrp3-91	1343	2594	91 92	91000866	265.1	wrp4-72 1151 2274	720280
vrp3-92 vrp3-94	1765 1976	3613 3836	92 94	92000764 94001181	518.2 851.2		73 730264
	2518	3830 4985	94 96	94001181 96001172	851.2 3421.5		740204
vrp3-96 vrp3-98	2518	4985	96 98	98001172	3421.5 3812.8		75 750171
							76 760204

Table A.4: Results on the WRP-instances. Type: Wire routing problems from industry [ZR00]. Instances not solved here could be solved using stronger reductions, see Table A.14.

alue2087 alue2105 alue3146 alue5067 alue5345 alue5623 alue65901 alue6457 alue6457 alue6457 alue6951 alue7065 alue7066 alue7080 alue7229 alut0787	V  1244 1220 3620 3524	4	E  1971	R		
alue2105 alue3146 alue5067 alue5345 alue5623 alue5623 alue6179 alue6457 alue6735 alue6951 alue7065 alue7066 alue7080 alue7229	1220 3620			34	1049	0.1
alue3146 alue5067 alue5345 alue5623 alue6503 alue6179 alue6457 alue6735 alue6951 alue7065 alue7066 alue7080 alue7229	3626		1858	34	1032	0.1
alue5345 alue5623 alue5901 alue6179 alue6457 alue6735 alue6951 alue7065 alue7066 alue7080 alue7229	3524		5869	64	2240	0.4
alue5623 alue5901 alue6179 alue6457 alue6735 alue6735 alue7065 alue7066 alue7080 alue7229			5560	68	2586	0.9
alue5901 alue6179 alue6457 alue6735 alue6951 alue7065 alue7066 alue7080 alue7229	5179	9	8165	68	3507	3.9
alue6179 alue6457 alue6735 alue6951 alue7065 alue7066 alue7080 alue7229	4472		6938	68	3413	1.9
alue6457 alue6735 alue6951 alue7065 alue7066 alue7080 alue7229	1154		18429	68	3912	3.3
alue6735 alue6951 alue7065 alue7066 alue7080 alue7229	3372		5213	67	2452	0.8
alue6951 alue7065 alue7066 alue7080 alue7229	3932	2	6137	68	3057	1.1
alue7065 alue7066 alue7080 alue7229	4119		6696	68	2696	0.9
alue7066 alue7080 alue7229	2818	8	4419	67	2386	0.8
alue7080 alue7229	3404	46	54841	544	23881	94.5
alue7229	6405	15	10454	16	2256	7.3
	3447	79	55494	2344	62449	68.3
abut0787	940	0	1474	34	824	0.1
alutoror	1160	60	2089	34	982	0.1
alut0805	966	6	1666	34	958	0.1
alut1181	3041	1	5693	64	2353	0.5
alut2010	6104	14	11011	68	3307	1.4
alut2288	9070		16595	68	3843	3.2
alut2566	5021		9055	68	3073	2.5
alut2610	3390		62816	204	12239	95.6
alut2625	3671		68117	879	35459	305.9
alut2025 alut2764	387		626	34	640	0.1
gap1307	342		552	17	549	0.1
gap1307 gap1413	542		906	17	457	0.1
gap1415 gap1500	220		908 374	10	254	0.1
						0.1
gap1810	429 735		702 1256	17 21	482 763	0.1
gap1904						
gap2007	2039		3548	17	1104	0.2
gap2119	1724		2975	29	1244	0.2
gap2740	1190		2084	14	745	0.1
gap2800	386		653	12	386	0.1
gap2975	179		293	10	245	0.1
gap3036	346		583	13	457	0.1
gap3100	921		1558	11	640	0.1
gap3128	1039		18043	104	4292	4.3
msm0580	338		541	11	467	0.1
msm0654	1290		2270	10	823	0.1
msm0709	1442		2403	16	884	0.1
msm0920	752		1264	26	806	0.1
msm1008	402		695	11	494	0.1
msm1234	933		1632	13	550	0.1
msm1477	1199		2078	31	1068	0.1
msm1707	278		478	11	564	0.1
msm1844	90		135	10	188	0.1
msm1931	875		1522	10	604	0.1
msm2000	898	8	1562	10	594	0.1
msm2152	2132	2	3702	37	1590	0.3
msm2326	418	8	723	14	399	0.1
msm2492	4045	5	7094	12	1459	0.4
msm2525	3031	1	5239	12	1290	0.3
msm2601	2961	51	5100	16	1440	0.5
msm2705	1359		2458	13	714	0.1
msm2802	1709		2963	18	926	0.1
msm2846	3263		5783	89	3135	0.8
msm3277	1704		2991	12	869	0.1
msm3676	957		1554	10	607	0.1
msm3727	4640		8255	21	1376	0.7
msm3829	4221		7255	12	1570	1.8
msm4038	237		390	11	353	0.1
msm4038 msm4114	402		690	16	393	0.1
msm4114 msm4190	391		666	16	393	0.1
msm4224	191		302	11	311	0.1
msm4312	5181		8893	10	2016	3.7
msm4414	317		476	11	408	0.1
msm4515	777		1358	13	630 5226	0.1
taq0014	6466		11046	128	5326	2.9
taq0023	572		963	11	621	0.1
taq0365	4186		7074	22	1914	0.9
taq0377	6836		11715	136	6393	6.2
taq0431	1128		1905	13	897	0.2
taq0631	609		932	10	581	0.1
taq0739	837	7	1438	16	848	0.1
taq0741	712	2	1217	16	847	0.1
taq0751	105		1791	16	939	0.2
taq0891	331		560	10	319	0.1
taq0903	6163		10490	130	5099	5.5
taq0900	310		514	17	370	0.1
taq0910	122		194	17	210	0.1
taq0920 taq0978	777		1239	17	566	0.1

Table A.5: Results on the VLSI and LIN-instances. Type: Grid graph with holes (not metric) from VLSI design. Instances not solved here could be solved using stronger reductions, see Table A.14.
Instance	V	Size  E	R	Opt	Time
es10000fst	27019	39407	10000	716174280	1398.9
				•	
Instance		Size		Opt	Time
	V	E	R		
es1000fst01	2865	4267	1000	230535806	19.7
es1000fst02	2629	3793	1000	227886471	33.4
es1000fst03	2762	4047	1000	227807756	9.3
es1000fst04	2778	4083	1000	230200846	15.0
es1000fst05	2676	3894	1000	228330602	11.1
es1000fst06	2815	4162	1000	231028456	24.3
es1000fst07	2604	3756	1000	230945623	5.7
es1000fst08	2834	4207	1000	230639115	17.6
es1000fst09	2846	4187	1000	227745838	14.1
es1000fst10	2546	3620	1000	229267101	5.5
es1000fst11	2763	4038	1000	231605619	18.8
es1000fst12	2984	4484	1000	230904712	19.2
es1000fst13	2532	3615	1000	228031092	6.7
es1000fst14	2840	4200	1000	234318491	17.1
es1000fst15	2733	3997	1000	229965775	13.6
	•				
Instance		Size		Opt	Time
	V	E	R		
a280fst	314	329	279	2502	0.1
att48fst	139	202	48	30236	0.3
att532fst	1468	2152	532	84009	4.5
berlin52fst	89	104	52	6760	0.1
bier127fst	258	357	127	104284	0.1
d1291fst	1365	1456	1291	481421	0.1
d1655fst	1906	2083	1655	584948	0.1
d198fst	232	256	198	129175	0.1
d2103fst	2206	2272	2103	769797	0.1
d493fst	1055	1473	493	320137	0.7
d657fst	1416	1978	657	471589	2.8
dsj1000fst	2562	3655	1000	17564659	1.9
eil101fst	330	538	101	605	1.5
eil51fst	181	289	51	409	3.4
eil76fst	237	378	76	513	1.1
fl1400fst	2694	4546	1400	17980523	263.2
fl1577fst	2413	3412	1577	19825626	1.4
fB795fst	4859	6539	3795	25529856	279.7
f#17fst	732	1084	417	10883190	1.3
fnl4461fst	17127	27352	4461	182361	12967.0
gil262fst	537	723	262	2306	0.1
kroA100fst	197	250	100	20401	0.1
kroA150fst	389	562	150	25700	0.9
kroA200fst	500	714	200	28652	0.4
kroB100fst	230	313	100	21211	0.1
kroB150fst	420	619	150	25217	0.6
kroB200fst	480	670	200	28803	0.7
kroC100fst	244	337	100	20492	0.1
kroD100fst	216	288	100	20437	0.1
kroE100fst	226	306	100	21245	0.1
	•				

Instance	l	Size		Opt	Time
	V	E	R	-	
lin105fst	216	323	105	13429	0.1
lin318fst	678	1030	318	39335	0.6
linhp318fst	678	1030	318	39335	0.6
nrw1379fst	5096	8105	1379	56207	207.6
p654fst	777	867	654	314925	0.1
pcb1173fst	1912	2223	1173	53301	0.1
pcb3038fst	5829	7552	3038	131895	2.9
pcb442fst	503	531	442	47675	0.1
pla7397fst	8790	9815	7397	22481625	0.2
pr1002fst	1473	1715	1002	243176	0.1
pr107fst	111	110	107	34850	0.1
pr124fst	154	165	124	52759	0.1
pr136fst	196	250	136	86811	0.1
pr144fst	221	285	144	52925	0.1
pr152fst	308	431	152	64323	0.1
pr226fst	255	269	226	70700	0.1
pr2392fst	3398	3966	2392	358989	0.1
pr264fst	280	287	264	41400	0.1
pr299fst	420	500	299	44671	0.1
pr439fst	572	662	439	97400	0.1
pr76fst	168	247	76	95908	0.1
rat195fst	560	870	195	2386	1.3
rat575fst	1986	3176	575	6808	23.6
rat783fst	2397	3715	783	8883	18.1
rat99fst	269	399	99	1225	0.2
rd100fst	201	253	100	764269099	0.1
rd400fst	1001	1419	400	1490972006	1.9
rl11849fst	13963	15315	11849	8779590	0.8
rl1304fst	1562	1694	1304	236649	0.1
rl1323fst	1598	1750	1323	253620	0.1
rl1889fst	2382	2674	1889	295208	0.1
r15915fst	6569	6980	5915	533226	0.2
rl5934fst	6827	7365	5934	529890	0.1
st70fst	133	169	70	626	0.2
ts225fst	225	224	225	1120	0.1
tsp225fst	242	252	225	356850	0.1
u1060fst	1835	232	1060	21265372	1.5
u10001st u1432fst	1432	1431	1432	1465	0.1
u14521st u159fst	1432	1451	1452	390	0.1
u1817fst u2152fst	1831 2167	1846 2184	1817 2152	5513053	0.1 0.1
				6253305	
u2319fst	2319	2318	2319	2322	0.1
u574fst	990	1258	574	3509275	0.2
u724fst	1180	1537	724	4069628	0.3
vm1084fst	1679	2058	1084	2248390	0.6
vm1748fst	2856	3641	1748	3194670	7.2

Table A.6: Results on the ES10000, ES1000 and TSP-instances. Type: Rectilinear, derived with geosteiner from 10000 (respectively 1000) random points in the plane, respectively instances from TSPLIB.

Instance		Size		Opt	Time
	V	E	R		
berlin52	52	1326	16	1044	0.1
brasil58	58	1653	25	13655	0.1
world666	666	221445	174	122467	0.8

Table A.7: Results on the X-instances. Type: Complete with euclidean weights.

	Size		Opt	Time
V	E	R	-	
400	760	213	11689	0.1
150	11175	80	92	2.6
120	7140	60	71	1.7
97	4656	45	47	5.4
400	760	170	3417	0.1
400	760	188	1566	0.1
	400 150 120 97 400	$\begin{array}{                                    $	$\begin{array}{ c c c c c c c c c c c c c c c c c c c$	$\begin{array}{ c c c c c c c c c c c c c c c c c c c$

Table A.8: Results on the MC-instances. Type: Constructed difficult instances.

$\begin{array}{c c c c c c c c c c c c c c c c c c c $	nstance	V	Size  E	R	Opt	Time	Instance	V	Size  E	R	Opt	1
i160-002    160    240    7    2158    0.1    i160-203    160    240    24    7630      i160-004    160    240    7    2370    0.1    i160-205    160    240    24    7243      i160-011    160    240    7    2495    0.1    i160-211    160    240    24    7123      i160-012    160    812    7    1677    0.1    i160-213    160    812    24    5643      i160-013    160    812    7    1778    0.3    i160-213    160    812    24    5547      i160-012    160    12720    7    1355    0.2    i160-221    160    12720    24    4779      i160-022    160    12720    7    1351    0.2    i160-223    160    12720    24    4730      i160-023    160    1270    7    1351    0.2    i160-231    160    12720    24    4728      i160-023    160    1270    7    1370    0.1	160.001				2400	0.1	:160 201				6022	
160-003    160    240    7    2297    0.1    160-024    160    240    24    7243      1160-005    160    240    7    2395    0.1    160-024    160    240    24    7083      1160-012    160    812    7    177    0.1    1160-212    160    812    24    5833      1160-013    160    812    7    1778    0.1    1160-213    160    812    24    5543      1160-014    160    812    7    1778    0.1    1160-215    160    812    24    5518      1160-022    160    12720    7    1355    0.2    1160-222    160    12720    24    4797      1160-024    160    12720    7    1356    0.2    1160-224    160    12720    24    4721      1160-024    160    12720    7    1376    0.2    1160-231    160    12720    24    4721      1160-023    160    1270    7    1370    0.1												
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$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$		160										
	160-021	160	12720	7	1352	0.2	i160-221	160	12720	24	4729	
	160-022	160	12720	7	1365	0.2	i160-222	160	12720	24	4697	
$ \begin{array}{c c c c c c c c c c c c c c c c c c c $	160-023	160	12720	7	1351	0.2	i160-223	160	12720	24	4730	
	160-024	160	12720	7	1371	0.2	i160-224	160	12720	24	4721	
	160-025	160	12720	7	1366	0.2	i160-225	160	12720	24	4728	
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	160-031	160	320	7	2170	0.1	i160-231	160	320	24	6662	
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$\begin{array}{c ccccccccccccccccccccccccccccccccccc$		160		12	2363			160		40		
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$		160								40		
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	160-123	160		12	2355			160	12720	40		
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$		160		12				160		40		
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	160-125	160	12720	12	2351	0.2	i160-325	160	12720	40	7862	
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	160-131	160	320	12	3356	0.1	i160-331	160	320	40	10414	
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	160-132	160	320	12	3450	0.1	i160-332	160	320	40	10806	
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$		160								40		
i160-135    160    320    12    3716    0.1    i160-335    160    320    40    10589      i160-141    160    2544    12    2549    0.3    i160-341    160    2544    40    8331      i160-142    160    2544    12    2562    1.5    i160-342    160    2544    40    8348      i160-142    160    2544    12    2557    0.6    i160-343    160    2544    40    8375      i160-144    160    2544    12    2607    1.2    i160-344    160    2544    40    8307												
$\begin{array}{cccccccccccccccccccccccccccccccccccc$												
i160-142    160    2544    12    2562    1.5    i160-342    160    2544    40    8348      i160-143    160    2544    12    2557    0.6    i160-343    160    2544    40    83275      i160-144    160    2544    12    2607    1.2    i160-344    160    2544    40    8307												
i160-143      160      2544      12      2557      0.6      i160-343      160      2544      40      8275        i160-144      160      2544      12      2607      1.2      i160-344      160      2544      40      8275												
i160-144 160 2544 12 2607 1.2 i160-344 160 2544 40 8307												
1160-145 1 160 2544 17 1 2578 0.8 1160-345 1 160 2544 40 1 9227	160-144 160-145	160	2544	12	2578	0.8	i160-344	160	2544	40	8307	

Table A.9: Results on the I160-instances. Type: Incidence networks, constructed with the aim of being difficult for known techniques.

Instance	V	Size  E	R	Opt	Time
	320	480	8	2672	0.1
002	320	480	8	2847	0.1
320-003	320	480	8	2972	0.1
i320-004	320	480	8	2905	0.1
320-005	320	480	8	2991	0.1
i320-011	320	1845	8	2053	0.5
i320-012	320	1845	8	1997	0.1
i320-013	320	1845	8	2072	1.3
i320-014	320	1845	8	2061	0.3
i320-015	320	1845	8	2059	0.7
i320-021	320	51040	8	1553	1.4
i320-022	320	51040	8	1565	1.4
i320-023	320	51040	8	1549	1.2
i320-024	320	51040	8	1553	1.1
i320-025	320	51040	8	1550	1.1
i320-031	320	640	8	2673	0.1
i320-032	320	640	8	2770	0.1
i320-032	320	640	8	2769	0.1
i320-033	320	640	8	2521	0.1
i320-034	320	640	8	2385	0.1
i320-041	320	10208	8	1707	0.7
i320-042	320	10208	8	1682	0.2
i320-042	320	10208	8	1723	0.2
i320-043	320	10208	8	1681	0.3
i320-044	320	10208	8	1686	0.2
i320-045	320	480	17	5548	0.2
i320-101	320	480	17	5556	0.1
			17		
i320-103 i320-104	320 320	480 480	17	6239 5703	0.1 0.1
		480	17		0.1
i320-105 i320-111	320 320	480 1845	17	5928 4273	0.2
i320-112	320	1845	17	4213	3.6
i320-113	320	1845	17	4205	2.9
i320-114	320	1845 1845	17 17	4104 4238	2.4 2.9
i320-115	320				
i320-121	320	51040	17	3321	1.7
i320-122	320	51040	17	3314	1.4
i320-123	320	51040	17	3332	1.8
i320-124	320	51040	17	3323	1.8
i320-125	320	51040	17	3340	1.8
i320-131	320	640	17	5255	0.6
i320-132	320	640	17	5052	0.1
i320-133	320	640	17	5125	0.1
i320-134	320	640	17	5272	0.1
i320-135	320	640	17	5342	0.1
i320-141	320	10208	17	3606	4.8
i320-142	320	10208	17	3567	3.6
i320-143	320	10208	17	3561	2.1
i320-144	320	10208	17	3512	0.2
i320-144	320	10208	17	3601	3.2

Table A.10: Results on the I320-instances. Type: Incidence networks, constructed with the aim of being difficult for known techniques. Instances not solved here could be solved using longer runs, see Table A.15.

Instance	I	Size		Opt	Time
	V	E	R		
-001	640	960	9	4033	0.1
i640-002	640	960	9	3588	0.1
i640-003	640	960	9	3438	0.1
i640-004	640	960	9	4000	0.1
i640-005	640	960	9	4006	0.1
i640-011	640	4135	9	2392	0.1
i640-012	640	4135	9	2465	1.2
i640-013	640	4135	9	2399	0.8
i640-014	640	4135	9	2171	0.1
i640-015	640	4135	9	2347	0.1
i640-021	640	204480	9	1749	10.1
i640-022	640	204480	9	1756	10.1
i640-023	640	204480	9	1754	10.2
i640-024	640	204480	9	1751	8.3
i640-025	640	204480	9	1745	10.2
i640-031	640	1280	9	3278	0.1
i640-032	640	1280	9	3187	0.1
i640-033	640	1280	9	3260	0.1
i640-034	640	1280	9	2953	0.1
i640-035	640	1280	9	3292	0.1
i640-041	640	40896	9	1897	5.0
i640-042	640	40896	9	1934	2.3
i640-043	640	40896	9	1931	1.3
i640-044	640	40896	9	1938	2.5
i640-045	640	40896	9	1866	0.9
i640-101	640	960	25	8764	0.2
i640-102	640	960	25	9109	0.1
i640-103	640	960	25	8819	0.1
i640-104	640	960	25	9040	0.2
i640-105	640	960	25	9623	1.0
i640-111	640	4135	25	6167	20.0
i640-112	640	4135	25	6304	21.6
i640-113	640	4135	25	6249	32.9
i640-114	640	4135	25	6308	17.2
i640-115	640	4135	25	6217	21.6
i640-121	640	204480	25	4906	12.1
i640-122	640	204480	25	4911	12.2
i640-123	640	204480	25	4913	12.2
i640-124	640	204480	25	4906	10.7
i640-125	640	204480	25	4920	12.3
i640-131	640	1280	25	8097	3.4
i640-132	640	1280	25	8154	1.6
i640-133	640	1280	25	8021	0.3
i640-134	640	1280	25	7754	0.1
i640-135	640	1280	25	7696	0.6
i640-141	640	40896	25	5199	32.3
i640-142	640	40896	25	5193	34.0
i640-142	640	40896	25	5194	20.5
i640-143	640	40896	25	5205	18.6
i640-144	640 640	40896	25	5205	39.7
1040-145	040	40890	23	3210	39.7

Table A.11: Results on the I640-instances. Type: Incidence networks, constructed with the aim of being difficult for known techniques. Instances i640-211, i640-34[1-5] could be solved using longer runs, see Table A.15.

Testeres		0'		0.4	T
Instance	V	Size  E	R	Opt	Time
cc10-2p	1024	5120	135	[34]33-	-356871
cc10-2u	1024	5120	135	[331-	
cc11-2p	2048	11263	244	[61773-	
cc11-2u	2048	11263	244	[600-	-620]
cc12-2p	4096	24574	473	[117941-	-122925]
cc12-2u	4096	24574	473	[1144-	_1197]
cc3-10p	1000	13500	50	[12173-	-12964]
cc3-10u	1000	13500	50	[115-	-127]
cc3-11p	1331	19965	61	[14883-	-15816]
cc3-11u	1331	19965	61	[140-	
cc3-12p	1728	28512	74	[17947-	
cc3-12u	1728	28512	74	[171-	
cc3-4p	64	288	8	2338	10.6
cc3-4u	64	288	8	23	7.8
cc3-5p	125	750	13	3661	447.6
cc3-5u	125	750	13	36	636.3
cc5-3p	243	1215	27	[6773-	,
cc5-3u	243	1215	27	[66–	
cc6-2p	64	192	12	3271	2.2
cc6-2u	64	192	12	32	5.0
cc6-3p	729	4368	76	[19847-	
cc6-3u	729	4368	76	[194-	
cc7-3p	2187	15308	222		-57459]
cc7-3u	2187	15308	222	[531-	,
cc9-2p	512	2304	64	[16520-	,
cc9-2u	512	2304	64	[161-	-172]

Table A.12: Results on the PUC-instances. Type: Constructed difficult instances: hypercubes, from code covering, and bipartite graphs [RdAR<sup>+</sup>01].

Instance		Size		Opt	Time
	V	E	R	Ŷ	
antiwheel5	10	15	5	7	0.1
design432	8	20	4	9	0.1
oddcycle3	6	9	3	4	0.1
oddwheel3	7	9	4	5	0.1
se03	13	21	4	12	0.1
w13c29	783	2262	406	[500	-508]
w23c23	1081	3174	552	[684	-694]
w3c571	3997	10278	2284	2854	15910.5

Table A.13: Results on the SP-instances. Type: Constructed difficult instances, combination of odd wheels and odd circles, difficult for Linear Programming approaches

Instance	I	Size		Opt	Time
	V	E	R		
lin31	19100	35653	40	31696	1002
lin32	19112	35665	53	39832	3559
lin33	19177	35730	117	56061	1416
lin34	38282	71521	34	45018	9144
lin35	38294	71533	45	50559	8194
lin36	38307	71546	58	55608	763693
lin37	38418	71657	172	99560	297795
wrp3-55	1645	3186	55	5500888	15569
wrp3-83	3168	6220	83	8300906	115224

Table A.14: Instances solved using stronger extended reductions.

Instance		Size		Opt	Time
	V	E	R		
i320-312	320	1845	80	18122	33542
i320-314	320	1845	80	18088	45856
i320-315	320	1845	80	17987	24918
i640-211	640	4135	50	11984	67237
i640-341	640	40896	160	32042	402451
i640-342	640	40896	160	31978	18682
i640-343	640	40896	160	32015	99206
i640-344	640	40896	160	31991	77280
i640-345	640	40896	160	31994	68199
cc5-3p	243	1215	27	7299	41856
cc5-3u	243	1215	27	71	220668
hc8p	256	1024	128	15322	400071
hc8u	256	1024	128	148	511646

Table A.15: Instances solved in longer runs.

# Summary

The Steiner problem in networks is the problem of connecting a set of required vertices in a weighted graph at minimum cost. This is a classical  $\mathcal{NP}$ -hard problem with many important applications in network design in general and VLSI layout in particular. The primary goal of our research has been the development of empirically successful algorithms. This means we designed and implemented algorithms that

- 1. generate Steiner trees of low cost in reasonable running times (upper bounds),
- 2. prove the quality of a Steiner tree by providing a lower bound on the optimal value (lower bounds),
- 3. or find an optimal Steiner tree (exact algorithms).
- 4. As an important prerequisite for the first three tasks, we used preprocessing techniques to reduce the size of the original problem without changing the optimal solution (reduction tests).

The value of our algorithms is measured by comparing our results to those of other research groups on the huge and well-established benchmark library for Steiner tree problems, SteinLib. In the case of exact algorithms, this measure is the best one can get because of the  $\mathcal{NP}$ -hardness result. For upper and lower bounds, the extensive experimental evaluation gives much sharper and much more relevant information than a worst-case analysis.

In the following, we will give a list of our main contributions:

## Lower Bounds:

- There are many (mixed) integer programming formulations of the Steiner problem. The corresponding linear programming relaxations are of great interest particularly, but not exclusively, for computing lower bounds, but not much was known about the relative quality of these relaxations. We compare the linear relaxations of all classical, frequently cited integer programming formulations of this problem from a theoretical point of view with respect to their optimal values. We present several new results, establishing very clear relations between relaxations, which have often been treated as unrelated or incomparable, forming a **hierarchy of relaxations**.
- We introduce a **collection of new relaxations** that are stronger than any known relaxation that can be solved in polynomial time, and place the new relaxations into our hierarchy. Further, we show how such a relaxation can be used in practical algorithms. Except for the flow-balance constraints introduced by Koch and Martin, this is the first successful attempt to use a relaxation that is stronger than Wong's directed cut relaxation from 1984.

#### **Preprocessing/Reduction techniques:**

• For some of the classical reduction tests, which would have been too time-consuming for large instances in their original form, we design **efficient realizations**, improving the worst-case running time to  $O(m + n \log n)$  in many cases. Furthermore, we design new tests, filling some of the gaps left by the classical tests.

- Previous reduction tests were either alternative based or bound based. That means to simplify the problem they either argued with the existence of alternative solutions, or they used some constrained lower bound and upper bound. We develop a framework for **extended reduction tests**, which extends the scope of inspection of reduction tests to larger patterns and combines for the first time alternative-based and bound-based approaches effectively.
- We introduce the new concept of **partitioning-based reduction techniques**, which has a significant impact on the reduction results in some cases.
- We integrate all tests into a **reduction packet**, which performs stronger reductions than any other package we are aware of. Additionally, the reduction results of other packages can be achieved typically in a fraction of the running time.

### **Upper Bounds:**

- We present variants of known path heuristics, including an empirically fast variant with a fast worst-case running time of  $O(m + n \log n)$ . The previous running time for this kind of path heuristic was  $O(rm \log n)$ .
- We introduce a new meta-heuristic, **reduction-based heuristics**. On the basis of this concept, we develop heuristics that achieve typically sharper upper bounds than the strongest known heuristics for this problem despite running times that are smaller by orders of magnitude.

## **Exact Algorithm:**

We integrate the previously mentioned building blocks into an exact algorithm that achieves very good running times. Additionally, we present a procedure that uses the fixed-parameter tractability of the Steiner problem for subgraphs of small width.

- For most benchmark instances the program computes the exact solution in running times that are shorter than the running times of other authors by orders of magnitude.
- There are 73 instances in SteinLib that have not been solved by any other research group. We have been able to solve 32 of them.
- For geometric Steiner problems, our algorithm for general networks is (together with a preprocessing algorithm from Warme, Winter, and Zachariasen that exploits some of the geometric properties) the fastest algorithm and beats the specially tailored MSTH approach, which has received much attention.

From the current experiences one can expect that instances of the Steiner problem with thousands of vertices and edges can be solved in minutes or hours if the instances were not intentionally constructed with the aim of being difficult for the known techniques.

# Zusammenfassung

Das Steiner Problem in Netzwerken ist das Problem, eine Menge von Basisknoten in einem gewichteten Graphen kostenminimal zu verbinden. Es ist ein klassisches  $\mathcal{NP}$ -schweres Problem mit vielen wichtigen Anwendungen in der Netzwerkoptimierung im Allgemeinen und im VLSI-Entwurf im Besonderen. Der Schwerpunkt unserer Arbeit hat in der Entwicklung empirisch erfolgreicher Algorithmen gelegen. Das heißt, wir haben Algorithmen entworfen, implementiert und getestet,

- 1. die schnell kosteng ünstige Steinerb aume berechnen (obere Schranken),
- 2. die die Qualit at eines Steinerbaums durch die Berechnung einer unteren Schranke f ur den Wert einer optimalen L osung nachweisen (untere Schranken),
- 3. oder die einen optimalen Steinerbaum finden (exakte Algorithmen).
- 4. Als wichtige Voraussetzung f`ur die ersten drei Aufgaben benutzen wir Vorverarbeitungs (*preprocessing*) Techniken, die die Schwierigkeit einer Probleminstanz reduzieren sollen, ohne die optimale L`osung zu ver`andern (Reduktionstests).

F'ur die Bewertung unserer Algorithmen vergleichen wir unsere Ergebnisse auf der umfangreichen und in der Forschung weithin akzeptierten Benchmark-Bibliothek SteinLib mit denen anderer Forschergruppen. Bez'uglich exakter Algorithmen ist diese Bewertung wegen der  $\mathcal{NP}$ -schwere Resultate die bestm'ogliche. Bez'uglich oberer und unterer Schranken liefert die ausf'uhrliche, experimentelle Bewertung sch'arfere und relevantere Informationen als eine worst-case Analyse.

Im folgenden stellen wir eine Liste der wesentlichen Ergebnisse der Arbeit vor:

# **Untere Schranken:**

- Es gibt viele (gemischt-) ganzzahlige Formulierungen f'ür das Steiner Problem. Die dazugeh örigen Linearen Relaxationen sind von großem Interesse insbesondere (aber nicht ausschließlich) f'ür die Berechnung von unteren Schranken. Allerdings war nur wenig "über die relative Qualit at dieser Relaxationen bekannt. Wir vergleichen die Linearen Relaxationen von allen klassischen, h "aufig zitierten ganzzahligen Formulierungen dieses Problems von einem theoretischen Standpunkt aus bez "uglich ihrer optimalen Werte. Dabei zeigen wir einige neue Resultate und stellen sehr klare Beziehungen zwischen Relaxationen auf, die bisher oft als unvergleichbar behandelt wurden. Die Beziehungen werden in einer **Hierarchie von Relaxationen** dargestellt.
- Wir stellen eine Klasse von neuen Relaxationen vor, die st ärker ist als alle anderen bekannten Relaxationen, die in Polynomialzeit gel ost werden k onnen. Wir plazieren die neuen Relaxationen in unserer Hierarchie. Weiter zeigen wir, wie eine Relaxation der Klasse in einem praktischen Algorithmus eingesetzt werden kann. Abgesehen von den Flussgleichgewichts-Nebenbedingungen, die von Koch und Martin eingef ührt wurden, ist dies der erste erfolgreiche Versuch, eine Relaxation algorithmisch zu benutzen, die st ärker ist als Wongs gerichtete Schnitt-Relaxation von 1984.

#### **Reduktionstechniken:**

• F'ur einige der klassischen Reduktionstests, die in ihrer Originalform f'ur große Instanzen zu zeitaufw'andig sind, entwerfen wir **effiziente Realisierungen**. In vielen F'allen k'onnen

wir die worst-case Laufzeit auf  $O(m + n \log n)$  verbessern. Weiterhin haben wir einige neue Tests entworfen, die einige der L'ucken schließen, die die bisherigen Tests gelassen hatten.

- Alle bisherigen Reduktionstests waren entweder alternativenbasiert oder schrankenbasiert. Das heißt, um das Problem zu vereinfachen argumentierten sie entweder mit der Existenz (oder Nichtexistenz) von alternativen Lösungen, oder sie benutzten obere Schranken und (eingeschränkte) untere Schranken. Wir haben ein Gerütst für **erweiterte Reduktionstests** entwickelt, bei dem der Beobachtungsbereich von Reduktionstests auf größere Muster erweitert wird und zum ersten Mal alternativenbasierte und schrankenbasierte Ansätze effektiv miteinander kombiniert werden.
- Wir entwickeln das neue Konzept der **partitionierungsbasierten Reduktionstechniken**, das in einigen F<sup>-</sup>allen deutliche Auswirkung auf die Reduktionsresultate hat.
- Wir verbinden alle Tests zu einem **Reduktionspaket**, das st ärkere Reduktionen durchf ühren kann als alle anderen uns bekannten Pakete. Dar überhinaus lassen sich die Reduktionsergebnisse anderer Pakete typischerweise in einem Bruchteil der Laufzeit erreichen.

# **Obere Schranken:**

- Wir entwickeln Varianten einer bekannten Pfadheuristik, unter anderem eine empirisch schnelle Variante mit einer schnellen worst-case Laufzeit von  $O(m + n \log n)$ . Die bisherige Laufzeitgarantie füur diese Art von Pfadheuristik war  $O(rm \log n)$ .
- Wir stellen eine neue Meta-Heuristik vor, **reduktionsbasierte Heuristiken**. Auf der Basis dieses Konzepts entwickeln wir Heuristiken, die typischerweise bessere obere Schranken erzielen als die st arksten bekannten Heuristiken und dabei um Gr oßenordnungen kleinere Laufzeiten haben.

#### **Exakter Algorithmus:**

Wir verbinden die zuvor erwähnten Bausteine zu einem exakten Algorithmus, der sehr gute Laufzeiten erzielt. Dar überhinaus zeigen wir, wie Lösbarkeit des Steinerproblems bez üglich bestimmter Problemparameter (*fixed-parameter tractability*) f ür Teilgraphen mit kleiner Breite benutzt werden kann.

- F<sup>°</sup>ur die meisten Benchmarkinstanzen berechnet unser Programm die exakten L<sup>°</sup>osungen in Laufzeiten, die um Gr<sup>°</sup>oßenordnungen k<sup>°</sup>urzer sind als die anderer Forschergruppen.
- Von den 73 Instanzen, die bisher von keiner anderen Forschergruppe gel"ost wurde, konnten wir 32 l"osen.
- Für geometrische Steinerbaum Probleme ist unser Algorithmus für Steinerprobleme in allgemeinen Netzwerken (zusammen mit einer Vorverarbeitung von Warme, Winter, und Zachariasen, die die geometrischen Eigenschaften ausnutzt) der schnellste Algorithmus und schlägt den speziell für dieses Problem entwickelten MSTH-Ansatz.

Nach den derzeitigen Erfahrungen ist zu erwarten, dass Instanzen des Steinerproblems mit tausenden von Knoten und Kanten in einem Zeitraum von Minuten oder Stunden gel öst werden können, wenn die Instanzen nicht speziell mit der Absicht konstruiert wurden, schwer für die bekannten Methoden zu sein.

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