Steiner Minimal Trees: An Introduction, Parallel Computation, and Future Work

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1 Introduction

Minimizing a network’s length is one of the oldest optimization problems in mathematics and, consequently, it has been worked on by many of the leading mathematicians in history. In the mid-seventeenth century a simple problem was posed: Find the point $P$ that minimizes the sum of the distances from $P$ to each of three given points in the plane. Solutions to this problem were derived independently by Fermat, Torricelli, and Cavalieri. They all deduced that either $P$ is inside the triangle formed by the
given points and that the angles at $P$ formed by the lines joining $P$ to the
three points are all $120^\circ$, or $P$ is one of the three vertices and the angle at
$P$ formed by the lines joining $P$ to the other two points is greater than or
equal to $120^\circ$.

In the nineteenth century a mathematician at the University of Berlin,
named Jakob Steiner, studied this problem and generalized it to include an
arbitrarily large set of points in the plane. This generalization created a
star when $P$ was connected to all the given points in the plane, and is a
geometric approach to the 2-dimensional center of mass problem.

In 1934 Jarník and Kössler generalized the network minimization prob-
lem even further [38]: Given $n$ points in the plane find the shortest possible
connected network containing these points. This generalized problem, how-
ever, did not become popular until the book, *What is Mathematics*, by
Courant and Robbins [14], appeared in 1941. Courant and Robbins linked
the name Steiner with this form of the problem proposed by Jarník and
Kössler, and it became known as the Steiner Minimal Tree problem. The
general solution to this problem allows multiple points to be added, each of
which is called a Steiner Point, creating a tree instead of a star.

Much is known about the exact solution to the Steiner Minimal Tree
problem. Those who wish to learn about some of the spin-off problems
are invited to read the introductory article by Bern and Graham [3], the
excellent survey paper on this problem by Hwang and Richards [34], or the
recent volume in The Annals of Discrete Mathematics devoted completely
to Steiner Tree problems [35]. Some of the basic pieces of information about
the Steiner Minimal Tree problem that can be gleaned from these articles
are: (i) the fact that all of the original $n$ points will be of degree 1, 2, or
3, (ii) the Steiner Points are all of degree 3, (iii) any two edges meet at an
angle of at least $120^\circ$ in the Steiner Minimal Tree, and (iv) at most $n - 2$
Steiner Points will be added to the network.

This paper concentrates on the Steiner Minimal Tree problem, hence-
forth referred to as the SMT problem. We present several algorithms for
calculating Steiner Minimal Trees, including the first parallel algorithm for
doing so. Several implementation issues are discussed, some new results are
presented, and several ideas for future work are proposed.

In Section 2 we review the first fundamental algorithm for calculating
SMT’s. In Section 3 we present a proposed heuristic for SMT’s. In Section 4
problem decomposition for SMT’s is outlined. In Section 5 we present Win-
ter’s sequential algorithm which has been the basis for most computerized
calculation of SMT’s to the present day. In Section 6 we present a parallel
algorithm for SMT's. Extraction of the correct answer is discussed in Section 7. Computational Results are presented in Section 8 and Future Work and open problems are presented in Section 9.

2 The First Solution

A typical problem-solving approach is to begin with the simple cases and expand to a general solution. As we saw in Section 1, the trivial three point problem had already been solved in the 1600's, so all that remained was the work toward a general solution. As with many interesting problems this is harder than it appears on the surface.

The method proposed by the mathematicians of the mid-seventeenth century for the three point problem is illustrated in Figure 1. This method stated that in order to calculate the Steiner Point given points A, B, and C, you first construct an equilateral triangle (ACX) using the longest edge between two of the points (AC) such that the third (B) lies outside the triangle. A circle is circumscribed around the triangle, and a line is constructed from the third point (B) to the far vertex of the triangle (X). The location of the Steiner Point (P) is the intersection of this line (BX) with the circle.

![Diagram of Steiner Point](image)

Figure 1: AP + CP = PX.

The next logical extension of the problem, going to four points, is attributed to Gauss. His son, who was a railroad engineer, was apparently
designing the layout for tracks between four major cities in Germany and was trying to minimize the length of these tracks. It is interesting to note at this point that a general solution to the SMT problem has recently been uncovered in the archives of a school in Germany [25].

For the next thirty years after Kössler and Jarník presented the general form of the SMT problem, only heuristics were known to exist. The heuristics were typically based upon the Minimum-Length Spanning Tree (MST), which is a tree that spans or connects all vertices whose sum of the edge lengths is as small as possible, and tried in various ways to join three vertices with a Steiner Point. In 1968 Gilbert and Pollak [24] linked the length of the SMT to the length of a MST. It was already known that the length of an MST is an upper bound for the length of an SMT, but their conjecture stated that the length of an SMT would never be any shorter than \( \frac{3}{2} \) times the length of an MST. This conjecture, was recently proved [15], and has led to the MST being the starting point for most of the heuristics that have been proposed in the last 20 years including a recent one that achieves some very good results [28].

In 1961 Melzak developed the first known algorithm for calculating an SMT [41]. Melzak’s Algorithm was geometric in nature and was based upon some simple extensions to Figure 1. The insight that Melzak offered was the fact that you can reduce an \( n \) point problem to a set of \( n-1 \) point problems. This reduction in size is accomplished by taking every pair of points, A and C in our example, calculating where the two possible points, \( X_1 \) and \( X_2 \), would be that form an equilateral triangle with them, and creating two smaller problems, one where \( X_1 \) replaces A and C, and the other where \( X_2 \) replaces A and C. Both Melzak and Cockayne pointed out however that some of these sub-problems are invalid. Melzak’s algorithm can then be run on the two smaller problems. This recursion, based upon replacing two points with one point, finally terminates when you reduce the problem from three to two vertices. At this termination the length of the tree will be the length of the line segment connecting the final two points. This is due to the fact that \( BP + AP + CP = BP + PX \). This is straightforward to prove using the law of cosines, for when \( P \) is on the circle, \( \angle APX = \angle CPX = 60^\circ \). This allows the calculation of the last Steiner Point (P) and allows you to back up the recursive call stack to calculate where each Steiner Point in that particular tree is located.

This reduction is important in the calculation of an SMT, but the algorithm still has exponential order, since it requires looking at every possible
reduction of a pair of points to a single point. The recurrence relation for an \( n \)-point problem is stated quite simply in the following formula:

\[
T(n) = 2 \cdot \left( \frac{n}{2} \right) \cdot T(n - 1).
\]

This yields what is obviously a non-polynomial time algorithm. In fact Garey, Graham, and Johnson [16] have shown that the Steiner Minimal Tree problem is NP-Hard (NP-Complete if the distances are rounded up to discrete values).

In 1967, just a few years after Melzak’s paper, Cockayne [9] clarified some of the details from Melzak’s proof. This clarified algorithm proved to be the basis for the first computer program to calculate SMTs. The program was developed by Cockayne and Schiller [13] and could compute an SMT for any placement of up to 7 vertices.

3 A Proposed Heuristic

3.1 Background and Motivation

By exploring a structural similarity between stochastic Petri nets (see [42] and [43]) and Hopfield neural nets (see [26] and [32]), Geist was able to propose and take part in the development of a new computational approach for attacking large, graph-based optimization problems. Successful applications of this mechanism include I/O subsystem performance enhancement through disk cylinder remapping [21, 22], file assignment in a distributed network to reduce disk access conflict [20], and new computer graphics techniques for digital halftoning [19] and color quantization [18]. The mechanism is based on maximum-entropy Gibbs measures, which is described in Reynold’s dissertation [47], and provides a natural equivalence between Hopfield nets and the simulated annealing paradigm. This similarity allows you to select the method that best matches the problem at hand. For the SMT problem we will implement the Simulated Annealing approach.

Simulated Annealing [39] is a probabilistic algorithm that has been applied to many optimization problems in which the set of feasible solutions is so large that an exhaustive search for an optimum solution is out of the question. Although Simulated Annealing does not necessarily provide an optimum solution, it usually provides a good solution in a user-selected amount of time. Hwang and Richards [34] have shown that the optimal placement
of \( s \) Steiner Points to \( n \) original vertices yields a feasible solution space of the size

\[
2^{-n} \binom{n}{s+2} \frac{(n-s-2)!}{s!}
\]

provided that none of the original points have degree 3 in the SMT. If the degree restriction is removed they showed that the number is even larger. The SMT problem is therefore a good candidate for this approach.

3.2 Adding One Junction

Georgakopoulos and Papadimitriou [23] have provided an \( \mathcal{O}(n^2) \) solution to the 1-Steiner problem, wherein exactly one Steiner Point is added to the original set of points. Since at most \( n - 2 \) Steiner Points are needed in an SMT solution, repeated application of the algorithm offers a “greedy” \( \mathcal{O}(n^3) \) approach. Using their method, the first Steiner Point is selected by partitioning the plane into Oriented Dirichlet Cells, which they describe in detail. Since these cells do not need to be discarded and recalculated for each addition, subsequent additions can be accomplished in linear time. Deletion of a candidate Steiner Point requires regeneration of the MST, which Shamos showed can be accomplished in \( \mathcal{O}(n \log n) \) time if the points are in the plane [44], followed by the cost for a first addition (\( \mathcal{O}(n^2) \)). This approach can be regarded as a natural starting point for Simulated Annealing by adding and deleting different Steiner Points.

3.3 The Heuristic

The Georgakopoulos and Papadimitriou 1-Steiner algorithm and the Shamos MST algorithm are both difficult to implement. As a result, we have chosen to investigate the potential effectiveness of this Annealing Algorithm using a more direct, but slightly more expensive \( \mathcal{O}(n^3) \) approach. As previously noted, all Steiner Points have degree 3 with edges meeting in angles of 120°. We consider all \( \binom{n}{3} \) triples where the largest angle is less than 120°, compute the Steiner Point for each (a simple geometric construction), select that Steiner Point giving greatest reduction, or least increase in the length of the modified tree (increases are allowed since the Annealing Algorithm may go uphill) and update the MST accordingly. Again, only the first addition requires this (now \( \mathcal{O}(n^3) \)) step. We use the straightforward, \( \mathcal{O}(n^2) \) Prim’s algorithm to generate the MST initially and after each deletion of a Steiner Point.
The Annealing Algorithm can be described as a non-deterministic walk on a surface. The points on the surface correspond to the lengths of all feasible solutions, where two solutions are adjacent if they can be reached through the addition or deletion of one Steiner Point. The probability of going uphill on this surface is higher when the temperature is higher but decreases as the temperature cools. The rate of this cooling typically will determine how good your solution will be. The major portion of this algorithm is presented in Figure 2. This non-deterministic walk, starting with the MST has led to some very exciting results.

3.4 Results

Before we discuss large problems, a simple introduction into the results from a simple six point problem is in order. The Annealing Algorithm is given the coordinates for six points: (0,0), (0,1), (2,0), (2,1), (4,0), (4,1). The first step is to calculate the MST, which has a length of 7, as shown in Figure 3. The output of the Annealing Algorithm for this simple problem is shown in Figure 4. In this case the Annealing Algorithm calculates the exact SMT solution which has a length of 6.616994.

We propose as a measure of accuracy the percentage of the difference between the length of the MST and the exact SMT solution that the Annealing Algorithm achieves. This is a new measure which has not been discussed (or used) because exact solutions have not been calculated for anything but the most simple layouts of points. For the six point problem discussed above this percentage is 100.0% (the exact solution is obtained).

After communicating with Cockayne, data sets were obtained for exact solutions to randomly generated 100 point problems that were developed for [12]. This allows us to use the measure of accuracy previously described. Results for some of these data sets provided by Cockayne are shown in Table 1.

An interesting aspect of the Annealing Algorithm that cannot be shown in the table is the comparison of execution times with Cockayne’s program. Whereas Cockayne mentioned that his results had an execution cut-off of 12 hours, these results were obtained in less than 1 hour. The graphical output for the first line of the table, which reaches over 96% of the optimal value, appears as follows: the data points and the MST are shown in Figure 5, the Simulated Annealing Result is in Figure 6, and the Exact SMT Solution is in Figure 7. The solution presented here is obtained in less than \(\frac{1}{16}\) of the time with less than 4% of the possible range not covered. This indicates that
Figure 2: Simulated Annealing algorithm.
we could hope to extend our Annealing Algorithm to much larger problems, perhaps as large as 1,000 points. If we were to extend this approach to larger problems we would definitely need to implement the Georgakopoulos-Papadimitriou 1-Steiner Algorithm and the Shamos MST Algorithm.

<table>
<thead>
<tr>
<th>Exact Solution</th>
<th>Spanning Tree</th>
<th>Simulated Annealing</th>
<th>Percent Covered</th>
</tr>
</thead>
<tbody>
<tr>
<td>6.255463</td>
<td>6.448690</td>
<td>6.261797</td>
<td>96.39%</td>
</tr>
<tr>
<td>6.759661</td>
<td>6.935189</td>
<td>6.763405</td>
<td>98.29%</td>
</tr>
<tr>
<td>6.667217</td>
<td>6.923836</td>
<td>6.675194</td>
<td>96.89%</td>
</tr>
<tr>
<td>6.719102</td>
<td>6.921413</td>
<td>6.721283</td>
<td>99.01%</td>
</tr>
<tr>
<td>6.759659</td>
<td>6.935187</td>
<td>6.763403</td>
<td>98.29%</td>
</tr>
<tr>
<td>6.285690</td>
<td>6.484320</td>
<td>6.289342</td>
<td>98.48%</td>
</tr>
</tbody>
</table>
Figure 5: Spanning tree.

Figure 6: Simulated Annealing solution.
4 Problem Decomposition

After the early work by Melzak [41], many people began to work on the Steiner Minimal Tree problem. The first major effort was to find some kind of geometric bound for the problem. In 1968 Gilbert and Pollak [24] showed that the SMT for a set of points, $\mathcal{S}$, must lie within the Convex Hull of $\mathcal{S}$. This bound has since served as the starting point of every bounds enhancement for SMT’s.

As a brief review, the Convex Hull is defined as follows: Given a set of points $\mathcal{S}$ in the plane, the Convex Hull is the convex polygon of smallest area containing all the points of $\mathcal{S}$. A polygon is defined to be convex if a line segment connecting any two points inside the polygon lies entirely within the polygon. An example of the Convex Hull for a set of 100 randomly generated points is shown in Figure 8.

Shamos in his PhD thesis [48] proposed a Divide and Conquer Algorithm which has served as the basis for many parallel algorithms calculating the Convex Hull. One of the first such approaches appeared in the PhD thesis by Chow [6]. This approach was refined and made to run in optimal $O(\log n)$ time by Aggarwal et.al. [1], and Attalah and Goodrich [2].
The next major work on the SMT problem was in the area of problem decomposition. As with any non-polynomial algorithm, the most important theorems are those that say “If property $P$ exists, then the problem may be split into the following sub-problems.” For the Steiner Minimal Tree Problem property $P$ will probably be geometric in nature. Unfortunately, decomposition theorems have been few and far between for the SMT problem. In fact, at this writing there have been only three such theorems.

### 4.1 The Double Wedge Theorem

The first decomposition theorem, known as the Double Wedge Theorem, was proposed by Gilbert and Pollak [24]. This is illustrated in Figure 9 and can be summarized quite simply as follows: If two lines intersect at point $x$ and meet at 120°, they split the plane into two 120° wedges and two 60° wedges. If $R_1$ and $R_2$ denote the two 60° wedges, and all the points of $S$ are contained in $R_1 \cup R_2$, then the problem can be decomposed. There are two cases to be considered. In case 1, $x$ is not a point in $S$; therefore, the Steiner Minimal Tree for $S$ is the SMT for $R_1$, the SMT for $R_2$, and the shortest edge connecting the two trees. In case 2, $x$ is a point in $S$; therefore, the Steiner Minimal Tree for $S$ is the SMT for $R_1$ and the SMT for $R_2$. Since
$\mathcal{X}$ is in both $R_1$ and $R_2$ the two trees are connected.

4.2 The Steiner Hull

The next decomposition theorem is due to Cockayne [10] and is based upon what he termed the *Steiner Hull*. The Steiner Hull is defined as follows: let $P_1$ be the convex Hull. $P_{i+1}$ is constructed from $P_i$ by finding an edge $(p, r)$ of $P_i$ that has a vertex $(q)$ near it such that $\angle pqr \geq 120^\circ$ and there is not a vertex inside the triangle $pqr$. The final polygon, $P_i$, that can be created in such a was is called the Steiner Hull. The Algorithm for this construction is shown in Figure 10. The Steiner Hull for the 100 points shown in Figure 8 is given in Figure 11.

After defining the Steiner Hull, Cockayne showed that the SMT for $S$ must lie within the Steiner Hull of $S$. This presents us with the following decomposition: The Steiner Hull can be thought of as an ordered sequence of points, $\{p_1, p_2, \ldots, p_n\}$, where the hull is defined by the sequence of line segments, $\{p_1p_2, p_2p_3, \ldots, p_np_1\}$. If there exists a point $p_i$ that occurs twice in the Steiner Hull, then the problem can be decomposed at point $p_i$. If a Steiner Hull contains such a point, then the Steiner Hull is referred to as *degenerate*. This decomposition is accomplished by showing that the Steiner Hull splits $S$ into two contained subsets, $R_1$ and $R_2$, where $R_1$ is the set of points contained in the Steiner Hull from the first time $p_i$ appears until the last time $p_i$ appears, and $R_2$ is the set of points contained in the Steiner Hull from the last time $p_i$ appears until the first time $p_i$ appears. With this decomposition it can be shown that $S = R_1 \cup R_2$, and the SMT for S is
The initial Steiner Polygon, $P_1$, is the Convex Hull. Repeat
Create Next Steiner Polygon $P_{i+1}$ from $P_i$ by
1) find a set of points $pqr \in S$ such that:
   $p$ and $r$ are adjacent on $P_i$
   $\angle pqr \geq 120^\circ$
   $\exists$ a point from $S$ in the triangle $pqr$
2) remove the edge $pr$.
3) add edges $pq$ and $qr$.
Until($P_i = P_{i+1}$)
Steiner Hull = $P_i$

Figure 10: The Steiner Hull algorithm.

Figure 11: The Steiner Hull for a random set of 100 points.
the union of the SMT for $R_1$ and the SMT for $R_2$. This decomposition is illustrated in Figure 12. Cockayne also proved that the Steiner Hull decomposition includes every decomposition possible with the Double Wedge Theorem.

In their work on 100 point problems, Cockayne and Hewgill [12] mention that approximately 15% of the randomly generated 100 point problems have degenerate Steiner Hull’s. The Steiner Hull shown in Figure 11 is not degenerate, while that in Figure 12 is.

![Figure 12: An illustration of the Steiner Hull decomposition.](image)

### 4.3 The Steiner Hull Extension

The final decomposition belongs to Hwang, Song, Ting, and Du [36]. They proposed an extension to the Steiner Hull as defined by Cockayne. Their extension is as follows:

If there exist four points $a, b, c, d$ on a Steiner Hull such that:

1. $a, b, c, d$ form a convex quadrilateral,
2. there does not exist a point from $S$ in the quadrilateral $(a, b, c, d),$
3. $\angle a \geq 120^\circ$ and $\angle b \geq 120^\circ$,
4. the two diagonals $(ac)$ and $(bd)$ meet at $O$, and $\angle bOa \geq \angle a + \angle b - 150^\circ$,
then the SMT for $S$ is the union of the SMT’s for $R_1$ and $R_2$ and the edge $ab$
where $R_1$ is the set of points contained in the Steiner Hull from $c$ to $b$ with
the edge $bc$, and $R_2$ is the set of points contained in the Steiner Polygon
from $a$ to $d$ with the edge $ad$. This decomposition is illustrated in Figure 13.
These three decomposition theorems were combined into a parallel algo-

![Figure 13: An illustration of the Steiner Hull extension.](image)

rithm for decomposition presented in [27].

5 Winter’s Sequential Algorithm

5.1 Overview and Significance

The development of the first working implementation of Melzak’s algorithm
sparked a move into the computerized arena for the calculation of SMT’s. As
we saw in Section 2, Cockayne and Schiller [13] had implemented Melzak’s
Algorithm and could calculate the SMT for all arrangements of 7 points.
This was followed almost immediately by Boyce and Seery’s program which
they called STEINER72 [4]. Their work, done at Bell Labs could calculate
the SMT for all 10 point problems. They continued to work on the problem
and in personal communication with Cockayne said they could solve 12 point
problems with STEINER73. Yet even with quite a few people working on
the problem, the number of points that any program could handle was still
very small.

As mentioned towards the end of Section 2, Melzak’s algorithm yields
invalid answers and invalid tree structures for quite a few combinations
of points. It was not until 1981 that anyone was able to characterize a
few of these invalid tree structures. These characterizations were accom-
plished by Pawel Winter and were based upon several geometric construc-
tions which enable one to eliminate many of the possible combinations previously generated. He implemented these improvements in a program called GeoSteiner [52]. In his work he was able to calculate in under 30 seconds SMT’s for problems having up to 15 vertices and stated that “with further improvements, it is reasonable to assert that point sets up to 30 V-points could be solved in less than an hour [52].”

5.2 Winter’s Algorithm

Winter’s breakthrough was based upon two things: the use of extended binary trees, and what he termed pushing. Winter proposed an extended binary tree as a means of constructing trees only once and easily identifying a Full Steiner Tree (FST: trees with $n$ vertices and $n - 2$ Steiner Points) on the same set of vertices readily.

Pushing came from the geometric nature of the problem and is illustrated in Figure 14. It was previously known that the Steiner Point for a pair of points, $a$ and $b$, would lie on the circle that circumscribed that pair and their equilateral third point. Winter set out to limit this region even further. This limitation was accomplished by placing a pair of points, $a'$ and $b'$, on the circle at $a$ and $b$ respectively, and attempting to push them closer and closer together. In his work Winter proposed and proved various geometric properties that would allow you to push $a'$ towards $b$ and $b'$ towards $a$. If the two points ever crossed then it was impossible for the current branch of the sample space tree to contain a valid answer.

Unfortunately, the description of Winter’s algorithm is not as clear as one would hope, since the presence of goto statements rapidly makes his program difficult to understand, and almost impossible to modify. Winter’s goal is to build a list of FST’s which are candidates for inclusion in the final answer. This list, called $T_{list}$, is primed with the edges of the MST, thereby guaranteeing that the length of the SMT does not exceed the length of the MST.

The rest of the algorithm sets about to expand what Winter termed as $Q_{list}$, which is a list of partial trees that the algorithm attempts to combine until no combinations are possible. $Q_{list}$ is primed with the original input points. The legality of a combination is determined in the Construct procedure, which uses pushing to eliminate cases. While this combination proceeds, the algorithm also attempts to take newly created members of $Q_{list}$ and create valid FST’s out of them. These FST’s are then placed onto $T_{list}$. 

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This algorithm was a turning point in the calculation of SMT's. It sparked renewed interest into the calculation of SMT's in general. This renewed interest has produced new algorithms such as the Negative Edge Algorithm [51] and the Luminary Algorithm [33]. Winter's algorithm has also served as the foundation for most computerized computation for calculating SMT's and is the foundation for the parallel algorithm we present next.

6 A Parallel Algorithm

6.1 A Introduction to Parallelism

Parallel computation is allowing us to look at problems that have previously been impossible to calculate, as well as allowing us to calculate faster than ever before problems we have looked at for a long time. It is with this in mind that we begin to look at a parallel algorithm for the Steiner Minimal Tree problem.

There have been volumes written on parallel computation and parallel algorithms; therefore, we will not rehash the material that has already been so excellently covered by many others more knowledgeable on the topic, but will refer the interested readers to various books currently available. For a thorough description of parallel algorithms, and the PRAM Model the
reader is referred to the book by Joseph JáJá [37], and for a more practical approach to implementation on a parallel machine the reader is referred to the book by Vipin Kumar et.al. [40], the book by Michael Quinn [45], or the book by Justin Smith [49].

6.2 Overview and Proper Structure

When attempting to construct a parallel algorithm for a problem the sequential code for that problem is often the starting point. In examining sequential code, major levels of parallelism may become self-evident. Therefore for this problem the first thing to do is to look at Winter’s algorithm and convert it into structured code without *gotos*. The Initialization (Step 1) does not change, and the translation of steps 2 through 7 appears in Figure 15.

Notice that the code in Figure 15 lies within a *for* loop. In a first attempt to parallelize anything you typically look at loops that can be split across multiple processors. Unfortunately, upon further inspection, the loop continues while p<q and, in the large if statement in the body of the loop, is the statement q++ (line 30). This means that the number of iterations is data dependent and is not fixed at the outset. This loop cannot be easily parallelized.

Since the sequential version of the code does not lend itself to easy parallelization, the next thing to do is back up and develop an understanding of how the algorithm works. The first thing that is obvious from the code is that you select a left subtree and then try to mate it with possible right subtrees. Upon further examination we come to the conclusion that a left tree will mate with all trees that are shorter than it, and all trees of the same height that appear after it on Q_list, but it will never mate with any tree that is taller.

6.3 First Approach

The description of this parallel algorithm is in a master–slave perspective. This perspective was taken due to the structure of most parallel architectures at the time of its development, as well as the fact that all nodes on the Q_list need a sequencing number assigned to them. The master will therefore be responsible for numbering the nodes and maintaining the main Q_list and T_list.

The description from the slave’s perspective is quite simple. A process is spawned off for each member of Q_list that is a proper left subtree (Winter’s
/* Step 2 */
1 for(p=0; p<q; p++) {
2     AP = A(p);
3     /* Step 3 */
4     for(r=0; (H(p) > H(r)) AND (r!=q); r++) {
5         if((H(p) == H(r)) AND (r<p))
6             r = p;
7         if(Subset(V(r), AP)) {
8             p3 = p;
9             r3 = r;
10           for(Label = PLUS; Label <= MINUS; Label++) {
11               /* Step 4 */
12               AQ = A(q);
13               if(Construct(p3, r3, &E(q))) {
14                   L(q) = p;
15                   R(q) = r;
16                   LBL(q) = Label;
17                   LF(q) = LF(p);
18                   H(q) = H(p) + 1;
19                   /* next line is different */
20                   Min(q) = max(Min(p)-1, H(r));
21                   if(Lsp(p) != 0)
22                       Lsp(q) = Lsp(p)
23                   else
24                       Lsp(q) = Lsp(r)
25                   if(Rsp(r) != 0)
26                       Rsp(q) = Rsp(r)
27                   else
28                       Rsp(q) = Rsp(p)
29                   q3 = q;
30                   q++;
31                   /* Step 5 */
32                   if(ProperAddTreeAddList(q3)) {
33                       for(j in AP with Lf(R(q3)) < j) {
34                           SRoot(t) = j;
35                           Root(t) = q3;
36                           t++;
37                     }
38                   }
39                   /* Step 6 */
40                   p3 = p;
41                   r3 = r;
42               }
43           }
44       }
45   }
46 }

Figure 15: The main loop properly structured.
algorithm allows members of \( Q \) list that are not proper left subtrees). Each new process is then given all the current nodes on \( Q \) list. With this information the slave then can determine with which nodes its left subtree could mate. This mating creates new nodes that are sent back to the master, assigned a number and added to the master’s \( Q \) list. The slave also attempts to create an FST out of the new \( Q \) list member, and if it is successful, this FST is sent to the master to be added to the \( T \) list. When a process runs out of \( Q \) list nodes to check it sends a request for more nodes to the master.

The master also has a simple job description. It has to start a process for each initial member of the \( Q \) list, sends them all the current members of the \( Q \) list and wait for their messages.

This structure worked quite well for smaller problems (up to about 15 points), but for larger problems it reached a grinding halt quite rapidly. This was due to various reasons such as the fact that for each slave started the entire \( Q \) list had to be sent. This excessive message passing quickly bogged down the network. Secondly in their work on 100 point problems Cockayne and Hewgill [12] made the comment that \( T \) list has an average length of 220, but made no comment about the size of \( Q \) list, which is the number of slaves that would be started. From our work on 100 point problems this number easily exceeds 1,000 which means that over 1,000 processes are starting, each being sent the current \( Q \) list. From these few problems, it is quite easy to see that some major changes needed to be made in order to facilitate the calculation of SMT’s for large problems.

### 6.4 Current Approach

The idea for a modification to this approach came from a paper by Quinn and Deo [46], on parallel algorithms for Branch-and-Bound problems. Their idea was to let the master have a list of work that needs to be done. Each slave is assigned to a processor. Each slave requests work, is given some, and during its processing creates more work to be done. This new work is placed in the master’s work list, which is sorted in some fashion. When a slave runs out of work to do, it requests more from the master. They noted that this leaves some processors idle at times (particularly when the problem was starting and stopping), but this approach provides the best utilization if all branches are independent.

This description almost perfectly matches the problem at hand. First, we will probably have a fixed number of processors which can be determined at runtime. Secondly we have a list of work that needs to be done. The hard
part is implementing a sorted work list in order to obtain a better utilization. This was implemented in what we term the Proc list, which is a list of the processes that either are currently running or have not yet started. This list is primed with the information about the initial members of Q list, and for every new node put on the Q list, a node which contains information about the Q list node, is placed on the Proc list in a sorted order.

The results for this approach are quite exciting, and the timings are discussed in Section 8.

7 Extraction of the Correct Answer

7.1 Introduction and Overview

Once the T list discussed in Section 5 is formed, the next step is to extract the proper answer from it. Winter described this in step 7 of his algorithm. His description stated that unions of FST’s saved in T list were to be formed subject to constraints described in his paper. The shortest union is the SMT for the original points.

The constraints he described were quite obvious considering the definition of an SMT. First, the answer had to cover all the original points. Second, the union of FST’s could not contain a cycle. Third, the answer is bounded in length by the length of the MST.

This led Winter to implement a simple exhaustive search algorithm over the FST’s in T list. This approach yields a sample space of size $O(2^m)$ (where m is the number of trees in T list) that has to be searched. This exponentiality is born out in his work where he stated that for problems with more than 15 points “the computation time needed to form the union of FST’s dominates the computation time needed for the construction of the FST’s [52].” An example of the input the last step of Winter’s algorithm receives (T list) is given in Figure 16. The answer it extracts (the SMT) is shown in Figure 17.

7.2 Incompatibility Matrix

Once Cockayne published the clarification of Melzak’s proof in 1967 [9] and Gilbert and Pollak published their paper giving an upper bound the the SMT length [24] many people were attracted to this problem. From this time until Winter’s work was published in 1985 [52] quite a few papers were published dealing with various aspects of the SMT Problem, but the attempt
Figure 16: $T_\text{list}$ for a random set of points.

Figure 17: SMT extracted from $T_\text{list}$ for a random set of points.
to computerize the solution of the SMT problem bogged down around 12 vertices. It wasn’t until Winter’s algorithm was published that the research community received the spark it needed to work on computerized computation of the SMT problem with renewed vigor. With the insight Winter provided into the problem, an attempt to computerize the solution of the SMT problem began anew.

Enhancement of this algorithm was first attempted by Cockayne and Hewgill at the University of Victoria. For this implementation Cockayne and Hewgill spent most of their work on the back end of the problem, or the extraction from T_list, and used Winter’s algorithm to generate T_list. This work on the extraction focused on what they termed an *incompatibility matrix*. This matrix had one row and one column for each member of T_list. The entries in this matrix were flags corresponding to one of three possibilities: *compatible*, *incompatible*, or *don’t know*. The rationale behind the construction of this matrix is the fact that it is faster to look up the value in a matrix than it is to check for the creation of cycles and improper angles during the union of FST’s.

The first value calculations for this matrix were straightforward. If two trees do not have any points in common then we *don’t know* if they are incompatible or not. If they have two or more points in common then they form a cycle and are *incompatible*. If they have only one point in common and the angle at the intersection point is less than 120° then they are also *incompatible*. In all other cases they are *compatible*.

This simple enhancement to the extraction process enabled Cockayne and Hewgill to solve all randomly generated problems of size up to 17 vertices in a little over three minutes [11].

### 7.3 Decomposition

The next focus of Cockayne and Hewgill’s work was in the area of the decomposition of the problem. As was discussed earlier in Section 4, the best theorems for any problem, especially non-polynomial problems, are those of the form “If property $\mathcal{P}$ exists then the problem can be decomposed.” Since the formation of unions of FST’s is exponential in nature any theorem of this type is important.

Cockayne and Hewgill’s theorem states: “Let $A_1$ and $A_2$ be subsets of $A$ satisfying (i) $A_1 \cup A_2 = A$ (ii) $|A_1 \cap A_2| = 1$ and (iii) the leaf set of each FST in $T_{\text{list}}$ is entirely contained in $A_1$ or $A_2$. Then any SMT on $A$ is the union of separate SMT’s on $A_1$ and $A_2$ [11].” This means that if you
break $T_{\text{list}}$ into biconnected components, the SMT will be the union of the
SMT's on those components.

Their next decomposition theorem allowed further improvements in the
calculation of SMT's. This theorem stated that if you had a component
of $T_{\text{list}}$ left from the previous theorem and if the $T_{\text{list}}$ members of that
component form a cycle, then it might be possible to break that cycle and
apply the previous algorithm again. The cycle could be broken if there
existed a vertex $v$ whose removal would change that component from one
biconnected component to more than one.

With these two decomposition theorems, Cockayne and Hewgill were able
to calculate the SMT for 79 of 100 randomly generated 30 point problems.
The remaining 21 would not decompose into blocks of size 17 or smaller, and
thus would have taken too much computation time [11]. This calculation
was implemented in the program they called EDSTEINER86.

7.4 Forest Management

Cockayne and Hewgill's next work focused on improvements to the *incompatibility matrix* previously described and was implemented in a program
called EDSTEINER89. Their goal was to reduce the number of *don't know's*
in the matrix and possibly remove some FST's from $T_{\text{list}}$ altogether.

They proposed two refinements for calculating the entry into the *incompatibility matrix* and one Tree Deletion Theorem. The Tree Deletion
Theorem stated that if there exists an FST in $T_{\text{list}}$ that is incompatible
with all FST's containing a certain point $a$ then the original FST can be
deleted since at least one FST containing $a$ will be in the SMT.

This simple change allowed Cockayne and Hewgill to calculate the SMT
for 77 of 100 randomly generated 100 point problems [12]. The other 23
problems could not be calculated in less than 12 hours and were therefore
terminated. For those that did complete, the computation time to generate
$T_{\text{list}}$ had become the dominate factor in the overall computation time.

So the pendulum had swung back from the extraction of the correct
answer from $T_{\text{list}}$ to the generation of $T_{\text{list}}$ dominating the computation
time. In Section 8 we will look at the results of the parallel algorithm
presented in Section 9 to see if the pendulum can be pushed back the other
way one more time.
8 Computational Results

8.1 Previous Computation Times

Before presenting the results for the parallel algorithm presented in Section 6, it is worthwhile to review the computation times that have preceded this algorithm in the literature. The first algorithm for calculating FST's was discussed in a paper by Cockayne [10] where he mentioned that preliminary results indicated his code could solve any problem up to 30 points that could be decomposed with the Steiner Hull into regions of 6 points or less.

As we saw in Section 2, the next computational results were presented by Cockayne and Schiller [13]. Their program, called STEINER, was written in FORTRAN on an IBM 360/50 at the University of Victoria. STEINER could calculate the SMT for any 7 point problem in less than 5 minutes of cpu time. When the problem size was increased to 8 it could solve them if 7 of the vertices were on the Steiner Hull. When this condition held it could calculate the SMT in under 10 minutes, but if this condition did not hold it would take an unreasonable amount of time.

Cockayne called STEINER a prototype for calculating SMT's and allowed Boyce and Serry of Bell Labs to obtain a copy of his code to improve the work. They improved the code, renamed itSTEINER72, were able to calculate the FST for all 9 point problems and most 10 point problems in a reasonable amount of time [4]. Boyce and Serry continued their work and developed another version of the code that they thought could solve problems of size up to 12 points, but no computation times were given.

The breakthrough we saw in Section 5 was by Pawel Winter. His program called GEOSTEINER [52] was written in SIMULA 67 on a UNIVAC-1100. GEOSTEINER could calculate SMT's for all randomly generated sets with 15 points in under 30 seconds. This improvement was put into focus when he mentioned that all previous implementations took more than an hour for non-degenerate problems of size 10 or more. In his work, Winter tried randomly generated 20 point problems but did not give results since some of them did not finish in his cpu time limit of 30 seconds. The only comment he made for problems bigger than size 15 was that the extraction discussed in Section 7 was dominating the overall computation time.

The next major program, EDSTEINER86, was developed in FORTRAN on an IBM 4381 by Cockayne and Hewgill [11]. This implementation was based upon Winter's results, but had enhancements in the extraction process. EDSTEINER86 was able to calculate the FST for 79 out of 100 ran-
domly generated 32 point problems. For these problems the cpu time for T\_list varied from 1 to 5 minutes, while for the 79 problems that finished the extraction time never exceeded 70 seconds.

Cockayne and Hewgill subsequently improved their SMT program and renamed it EDSTEINER89 [12]. This improvement was completely focused on the extraction process. EDSTEINER89 was still written in FORTRAN, but was run on a SUN 3/60 workstation. They randomly generated 200 32-point problems to solve and found that the generation of T\_list dominated the computation time for problems of this size. The average time for T\_list generation was 438 seconds while the average time for forest management and extraction averaged only 43 seconds. They then focused on 100 point problems and set a cpu limit of 12 hours. The average cpu time to generate T\_list was 209 minutes for these problems, but only 77 finished the extraction in the cpu time limit. These programs and their results are summarized in Table 2.

<table>
<thead>
<tr>
<th>Program</th>
<th>Author(s)</th>
<th>Points</th>
</tr>
</thead>
<tbody>
<tr>
<td>STEINER</td>
<td>Cockayne &amp; Schiller</td>
<td>7</td>
</tr>
<tr>
<td></td>
<td>Univ of Victoria</td>
<td></td>
</tr>
<tr>
<td>STEINER72</td>
<td>Boyce &amp; Serry</td>
<td>10</td>
</tr>
<tr>
<td></td>
<td>ATT Bell Labs</td>
<td></td>
</tr>
<tr>
<td>STEINER73</td>
<td>Boyce &amp; Serry</td>
<td>12</td>
</tr>
<tr>
<td></td>
<td>ATT Bell Labs</td>
<td></td>
</tr>
<tr>
<td>GEOSTEINER</td>
<td>Winter</td>
<td>15</td>
</tr>
<tr>
<td></td>
<td>Univ of Copenhagen</td>
<td></td>
</tr>
<tr>
<td>EDSTEINER86</td>
<td>Cockayne &amp; Hewgill</td>
<td>30</td>
</tr>
<tr>
<td></td>
<td>Univ of Victoria</td>
<td></td>
</tr>
<tr>
<td>EDSTEINER89</td>
<td>Cockayne &amp; Hewgill</td>
<td>100</td>
</tr>
<tr>
<td></td>
<td>Univ of Victoria</td>
<td></td>
</tr>
<tr>
<td>PARSTEINER94</td>
<td>Harris</td>
<td>100</td>
</tr>
<tr>
<td></td>
<td>Univ of Nevada</td>
<td></td>
</tr>
</tbody>
</table>
8.2 The Implementation

8.2.1 The Significance of the Implementation

The parallel algorithm we presented has been implemented in a program called PARSTEINER94 [27, 30]. This implementation is only the second SMT program since Winter’s GEOSTEINER in 1981 and is the first parallel code. The major reason that the number of SMT programs is so small is due to the fact that any implementation is necessarily complex.

PARSTEINER94 currently has over 13,000 lines of C code. While there is a bit of code dealing with the parallel implementation, certain sections of Winter’s Algorithm have a great deal of code buried beneath the simplest statements. For example line 13 of Figure 15 is the following:

\[
\text{if(Construct(p\_star, r\_star, \&(E(q))))}.
\]

To implement the function \text{Construct}() over 4,000 lines of code were necessary, and this does not include the geometry library with functions such as \text{equilateral\_third\_point()}, \text{center\_of\_equilateral\_triangle()}, \text{line\_circle\_intersect()}, and a host more.

Another important aspect of this implementation is the fact that there can now be comparisons made between the two current SMT programs. This would allow verification checks to be made between EDSTEINER89 and PARSTEINER94. This verification is important since with any complex program it is quite probable that there are a few errors hiding in the code. This implementation would also allow other SMT problems, such as those we will discuss in Section 9, to be explored independently, thereby broadening the knowledge base for SMT’s even faster.

8.2.2 The Platform

In the design and implementation of parallel algorithms you are faced with many decisions. One such decision is what will your target architecture be? There are times when this decision is quite easy due to the machines at hand or the size of the problem. In our case we decided not to target a specific machine, but an architectural platform called PVM [17].

PVM, which stands for Parallel Virtual Machine, is a software package available from Oak Ridge National Laboratory. This package allows a collection of parallel or serial machines to appear as a large distributed memory computational machine (MIMD model). This is implemented via two major
pieces of software, a library of PVM interface routines, and a PVM demon that runs on every machine that you wish to use.

The library interface comes in two languages, C and Fortran. The functions in this library are the same no matter which architectural platform you are running on. This library has functions to spawn off (start) many copies of a particular program on the parallel machine, as well as functions to allow message passing to transfer data from one process to another. Application programs must be linked with this library to use PVM.

The demon process, called *pvm*d in the user’s guide, can be considered the back end of PVM. As with any back end, such as the back end of a compiler, when it is ported to a new machine the front end can interface to it without change. The back end of PVM has been ported to a variety of machines, such as a few versions of Crays, various Unix machines such as Sun workstations, HP machines, Data General workstations, and DEC Alpha machines. It has also been ported to a variety of true parallel machines such as the iPSC/2, iPSC/860, CM2, CM5, BBN Butterfly and the Intel Paragon.

With this information it is easy to see why PVM was picked as the target platform. Once a piece of code is implemented under PVM it can be re-compiled on the goal machine, linked with the PVM interface library on that machine, and run without modification. In our case we designed PARSTEINER94 on a network of SUN workstations, but, as just discussed, moving to a large parallel machine should be trivial.

### 8.2.3 Errors Encountered

When attempting to implement any large program from another person’s description you often reach a point where you don’t understand something. At first you always question yourself, but as you gain an understanding of the problem you learn that there are times when the description you were given is wrong. Such was the case with the SMT problem. Therefore, to help some of those that may come along and attempt to implement this problem after us we recommend that you look at the list of errors we found while implementing Winter’s Algorithm [27].
8.3 Random Problems

8.3.1 100 Point Random Problems

From the literature it is obvious that the current standard for calculating SMT's has been established by Cockayne and Hewgill. Their work on SMT's has pushed the boundary of computation out from the 15 point problems of Winter to being able to calculate SMT's for a large percentage of 100 point problems.

Cockayne and Hewgill, in their investigation of the effectiveness of EDSTEINER89, randomly generated 100 problems with 100 points inside the unit square. They set up a CPU limit of 12 hours, and 77 of 100 problems finished within that limit. They described the average execution times as follows: $T_{\text{list}}$ construction averaged 209 minutes, Forest Management averaged 27 minutes, and Extraction averaged 10.8 minutes.

While preparing the code for this project, Cockayne and Hewgill were kind enough to supply us with 40 of the problems generated for [12] along with their execution times. These data sets were given as input to the parallel code PARSTEINER94, and the calculation timed. The Wall Clock time necessary to generate $T_{\text{list}}$ for the two programs appear in Table 3. For all 40 cases, the average time to generate $T_{\text{list}}$ was less than 20 minutes. This is exciting because we have been able to generate $T_{\text{list}}$ properly, while cutting an order of magnitude off the time.

These results are quite promising for various reasons. First, the parallel implementation presented in this work is quite scalable, and therefore could be run with many more processors, thereby enhancing the speedup provided. Secondly, with the PVM platform used, we can in the future port this work to a real parallel MIMD machine, which will have much less communication overhead, or to a shared memory machine, where the communication could all but be eliminated, and expect the speedup to improve much more.

It is also worth noting that proper implementation of the Cycle Breaking which Cockayne and Hewgill presented in [11], is important if extraction of the proper answer is to be accomplished. In their work, Cockayne and Hewgill mentioned that 58% of the problems they generated were solvable without the Cycle Breaking being implemented, which is approximately what we have found with the data sets they provided. An example of such a $T_{\text{list}}$ that would need cycles broken (possibly multiple times) is provided in Figure 18.
Table 3: Comparison of $T_{\text{list}}$ times.

<table>
<thead>
<tr>
<th>Test Case</th>
<th>PARSTEINER94</th>
<th>EDSTEINER89</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>650</td>
<td>8597</td>
</tr>
<tr>
<td>2</td>
<td>1031</td>
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</tr>
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<td>Avg.</td>
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<td>13748</td>
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</table>
8.3.2 Larger Random Problems

Once the 100 point problems supplied by Cockayne and Hewgill had been successfully completed, the next step was to try a few larger problems. This was done with the hope of gaining an insight into the changes that would be brought about from the addition of more data points.

For this attempt we generated several random sets of 110 points each. The length of T-list increased by approximately 38%, from an average of 210 trees to an average of 292 trees. The time to compute T-list also increased, but the growth more than doubled, going from an average of 15 minutes to an average of more than 40 minutes.

The interesting thing that jumped out the most was the increase in the number of large bi-connected components. Since the extraction process must do a complete search of all possibilities, the larger the component the longer it will take. This is a classic example of an exponential problem, where when the problem size increases by 1 the time doubles. With this increased component size, none of the random problems generated finished inside a 12 hour cut off time.

This rapid growth puts into perspective the importance of the work previously done by Cockayne and Hewgill. Continuation of their work with
incompatibility matrices as well as decomposition of $T_{j\text{list}}$ components appears at this point to be very important for the future of SMT calculations.

8.4 Grids

The problem of determining SMTs for grids was mentioned to the author by Ron Graham. In this context we are thinking of a grid as a regular lattice of unit squares. The literature has little of information regarding SMTs on grids, and most of the information that is given is conjectured and not proven. In Section 8.4.1 we will look at what is known about SMTs on grids. In the following sub-sections we will introduce new results for grids up through $7 \times m$ in size. These results presented are computational results from PARSTEINER94 [27, 29, 30] which was discussed previously.

8.4.1 $2 \times m$ and Square Grids

The first proof for anything besides a $2 \times 2$ grid came in a paper by Chung and Graham [8] in which they proved the optimality of their characterization of SMTs for $2 \times m$ grids. The only other major work was presented in a paper by Chung, Gardner, and Graham [7]. They argued the optimality of the SMT on $2 \times 2$, $3 \times 3$, and $4 \times 4$ grids and gave conjectures and constructions for those conjectures for SMTs on all other square lattices.

In their work Chung, Gardner, and Graham specified three building blocks from which all SMTs on square $(n \times n)$ lattices were constructed. The first, labeled $I$, is just a $K_2$, or a path on two vertices. This building block is given in Figure 19-A. The second, labeled $\mathcal{Y}$, is a Full Steiner Tree (FST) ($n$ vertices and $n - 2$ steiner points) on 3 vertices of the unit square. This building block is given in Figure 19-B. The third, labeled $\mathcal{X}$, is an FST on all 4 vertices of the unit square. This building block is given in Figure 19-C. For the generalizations we are going to make here, we need to introduce one more building block, which we will label $\mathcal{S}$. This building block is an FST on a $3 \times 2$ grid and appears in Figure 19-D.

SMTs for grids of size $2 \times m$ have two basic structures. The first is an FST on all the vertices in the $2 \times m$ grid. An example of this for a $2 \times 3$ grid is given in Figure 19-D. The other structure is constructed from the building blocks previously described. We hope that these building blocks, when put in conjunction with the generalizations for $3 \times m$, $4 \times m$, $5 \times m$, $6 \times m$, and $7 \times m$ will provide the foundation for a generalization of $m \times n$ grids in the future.
In their work on ladders \((2 \times m)\) grids, Chung and Graham established and proved the optimality of their characterization for \(2 \times m\) grids. Before giving their characterization, a brief review of the first few \(2 \times m\) SMTs is in order. The SMT for a \(2 \times 2\) grid is shown in Figure 19-C, the SMT for a \(2 \times 3\) grid is shown in Figure 19-D, and the SMT for a \(2 \times 4\) grid is given in Figure 20.

Chung and Graham [8] proved that SMTs for ladders fell into one of two categories. If the length of the ladder was odd, then the SMT was the FST on the vertices of the ladder. The SMT for the \(2 \times 3\) grid in Figure 19-D is an example of this. If the length of the ladder was even, the SMT was made up of a series of \(\left(\frac{m}{2} - 1\right)X\)'s followed by one last \(X\). The SMT for the \(2 \times 4\) grid in Figure 20 is an example of this.

8.4.2 3 \(\times\) \(m\) Grids

The SMT for \(3 \times m\) grids has a very easy characterization which can be seen once the initial cases have been presented. The SMT for the \(3 \times 2\) grid is presented in Figure 19-D. The SMT for the \(3 \times 3\) grid is presented in Figure 21.

From here we can characterize all \(3 \times m\) grids. Except for in the \(3 \times 2\) grid, which is an \(S\) building block, there will be only two basic building
blocks present, $\mathcal{X}$’s and $\mathcal{I}$’s. There will be exactly two $\mathcal{I}$’s and $(m - 1)\mathcal{X}$’s. The two $\mathcal{I}$’s will appear on each end of the grid. The $\mathcal{X}$’s will appear in a staggered checkerboard pattern, one on each column of the grid the same way that the two $\mathcal{X}$’s are staggered in the $3 \times 3$ grid. The $3 \times 5$ grid is a good example of this and is shown in Figure 22.

Figure 21: SMT for a $3 \times 3$ Grid

Figure 22: SMT for a $3 \times 5$ Grid

8.4.3 $4 \times m$ Grids

The foundation for the $4 \times m$ grids has already been laid. In their most recent work, Cockayne and Hewgill presented some results on Square Lattice Problems [12]. They looked at $4 \times m$ grids for $m = 2$ to $m = 6$. They also looked at the SMTs for these problems when various lattice points in that grid were missing. What they did not do, however, was characterize the structure of the SMT’s for all $4 \times m$ grids.

The $4 \times 2$ grid is given in Figure 20. From the work of Chung, Gardner, and Graham [7], we know that the SMT for a $4 \times 4$ grid is a checkerboard pattern of $5 \mathcal{X}$’s. This layout gives us the first two patterns we will need to describe the $4 \times m$ generalization. The first pattern, which we will call
pattern \( \mathcal{A} \), is the same as the \( 3 \times 4 \) grid without the two \( \mathcal{I} \)'s on the ends. This pattern is given in Figure 23. The second pattern, denoted as pattern \( \mathcal{B} \), is the \( 2 \times 4 \) grid in Figure 20 without the connecting \( \mathcal{I} \). This is shown in Figure 24.

![Figure 23: 4 \times m Pattern \( \mathcal{A} \)](image)

![Figure 24: 4 \times m Pattern \( \mathcal{B} \)](image)

Before the final characterization can be made, two more patterns are needed. The first one, called pattern \( \mathcal{C} \), is a \( 4 \times 3 \) grid where the pattern is made up of two non-connected \( 2 \times 3 \) SMTs, shown in Figure 25. The next pattern, denoted pattern \( \mathcal{D} \), is quite simply a \( \mathcal{Y} \) centered in a \( 2 \times 4 \) grid. This is shown in Figure 26. The final pattern, denoted \( \mathcal{E} \), is just an \( \mathcal{I} \) on the right side of a \( 2 \times 4 \) grid. This is shown in Figure 27.

Now we can begin the characterization. The easiest way to present the characterization is with some simple string rewriting rules. Since the \( 4 \times 2 \), \( 4 \times 3 \), and \( 4 \times 4 \) patterns have already been given, the rules will begin with a \( 4 \times 5 \) grid. This grid has the string \( \mathcal{AC} \). The first rule is that whenever there is a \( \mathcal{C} \) on the right end of your string replace it with \( \mathcal{BDB} \). Therefore a \( 4 \times 6 \) grid is \( \mathcal{CABDB} \). The next rule is that whenever there is a \( \mathcal{B} \) on the right end of your string replace it with a \( \mathcal{C} \). The final rule is whenever there is a \( \mathcal{DC} \) on the right end of your string replace it with an \( \mathcal{EAB} \). These rules are summarized in Table 4. A listing of the strings for \( m \) from 5 to 11 is given in Table 5.
Figure 25: $4 \times m$ Pattern $C$

Figure 26: $4 \times m$ Pattern $D$

Figure 27: $4 \times m$ Pattern $E$

<p>| | | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>$B \rightarrow C$</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>$C \rightarrow BDB$</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>$DC \rightarrow EAB$</td>
<td></td>
</tr>
</tbody>
</table>

Table 4: Rewrite rules for $4 \times m$ Grids.
<table>
<thead>
<tr>
<th>m =</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
</tr>
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<tr>
<td>String</td>
<td>AC</td>
<td>ABDB</td>
<td>ABDC</td>
<td>ABEB</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>m =</th>
<th>9</th>
<th>10</th>
<th>11</th>
</tr>
</thead>
<tbody>
<tr>
<td>String</td>
<td>ABEC</td>
<td>ABEBED</td>
<td>ABEBEDC</td>
</tr>
</tbody>
</table>

Table 5: String Representations for $4 \times m$ Grids

### 8.4.4 $5 \times m$ Grids

For the $5 \times m$ grids there are 5 building blocks (and their mirror images which are donated with an ') that are used to generate any $5 \times m$ grid. These building blocks appear in Figure 28, Figure 29, Figure 30, Figure 31, and Figure 32.

![Figure 28: $5 \times m$ Pattern A](image)

![Figure 29: $5 \times m$ Pattern B](image)

With the building blocks in place, the characterization of $5 \times m$ grids is quite easy using grammar rewrite rules. The rules used for rewriting strings representing a $5 \times m$ grid are given in Table 6. The SMTs for $5 \times 2$, $5 \times 3$, and $5 \times 4$ have already been given. For a $5 \times 5$ grid the SMT is made up of the following string: $\mathcal{E}A'BD$. As a reminder, the $A'$ signifies the mirror
Figure 30: $5 \times m$ Pattern $\mathcal{C}$

Figure 31: $5 \times m$ Pattern $\mathcal{D}$

Figure 32: $5 \times m$ Pattern $\mathcal{E}$
of building block $A$. A listing of the strings for $m$ from 5 to 11 is given in Table 7.

<table>
<thead>
<tr>
<th>$m$</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
</tr>
</thead>
<tbody>
<tr>
<td>String</td>
<td>$A'BA'D'$</td>
<td>$A'BA'E'$</td>
<td>$A'BA'AC$</td>
<td>$A'BA'AB'D'$</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>$m$</th>
<th>9</th>
<th>10</th>
<th>11</th>
</tr>
</thead>
<tbody>
<tr>
<td>String</td>
<td>$A'BA'AB'AE'$</td>
<td>$A'BA'AB'AA'C'$</td>
<td>$A'BA'AB'AA'BD$</td>
</tr>
</tbody>
</table>

Table 7: String Representations for $5 \times m$ Grids

### 8.4.5 $6 \times m$ Grids

For the $6 \times m$ grids there are 5 building blocks that are used to generate any $6 \times m$ grid. These building blocks appear in Figure 33, Figure 34, Figure 35, Figure 36, and Figure 37.

![Figure 33: 6 × m Pattern $A$](image-url)
Figure 34: $6 \times m$ Pattern $\mathcal{B}$

Figure 35: $6 \times m$ Pattern $\mathcal{C}$

Figure 36: $6 \times m$ Pattern $\mathcal{D}$

Figure 37: $6 \times m$ Pattern $\mathcal{E}$
The solution for $6 \times m$ grids can now be characterized by using grammar rewrite rules. The rules used for rewriting strings representing a $6 \times m$ grid are given in Table 8. The basis for this rewrite system is the SMT for the $6 \times 3$ grid which is $\mathcal{AC}$. It is also nice to see that for the $6 \times m$ grids there is a simple regular expression which can characterize what the string will be. That regular expression has the form: $A(BE)^*(C|BD)$. A listing of the strings for $m$ from 6 to 11 is given in Table 9.

\begin{tabular}{|c|c|}
\hline
1 & $C \rightarrow BD$ \\
\hline
2 & $D \rightarrow EC$ \\
\hline
\end{tabular}

Table 8: Rewrite rules for $6 \times m$ Grids

\begin{tabular}{|c|c|c|c|}
\hline
m = & 6 & 7 & 8 \\
\hline
String & $ABEBD$ & $ABEBEC$ & $ABEBEBD$ \\
\hline
\end{tabular}

\begin{tabular}{|c|c|c|c|}
\hline
m = & 9 & 10 & 11 \\
\hline
String & $ABEBEBEC$ & $ABEBEBEBD$ & $ABEBEBEBEC$ \\
\hline
\end{tabular}

Table 9: String Representations for $6 \times m$ Grids

### 8.4.6 $7 \times m$ Grids

For the $7 \times m$ grids there are 6 building blocks that are used to generate any $7 \times m$ grid. These building blocks appear in Figure 38, Figure 39, Figure 40, Figure 41, Figure 42, and Figure 43.

![Figure 38: 7 × m Pattern A](image)

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Figure 39: $7 \times m$ Pattern $\mathcal{B}$

Figure 40: $7 \times m$ Pattern $\mathcal{C}$

Figure 41: $7 \times m$ Pattern $\mathcal{D}$

Figure 42: $7 \times m$ Pattern $\mathcal{E}$

Figure 43: $7 \times m$ Pattern $\mathcal{F}$
The grammar rewrite rules for strings representing a $7 \times m$ grid are given in Table 10. The basis for this rewrite system is the SMT for the $7 \times 5$ grid which is $F'AF'$. A listing of the strings for $m$ from 6 to 11 is given in Table 11.

$$
\begin{align*}
1 & : E'F' \rightarrow BA'F \\
2 & : F \rightarrow CD \\
3 & : CD \rightarrow AEF \\
4 & : EF \rightarrow B'AF' \\
5 & : F' \rightarrow C'D' \\
6 & : C'D' \rightarrow A'E'F'
\end{align*}
$$

Table 10: Rewrite rules for $7 \times m$ Grids

<table>
<thead>
<tr>
<th>$m =$</th>
<th>6</th>
<th>7</th>
<th>8</th>
<th>9</th>
</tr>
</thead>
<tbody>
<tr>
<td>String</td>
<td>$FABA'F$</td>
<td>$FABA'CD$</td>
<td>$FABA'AEF$</td>
<td>$FABA'AB'AEF'$</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>$m =$</th>
<th>10</th>
<th>11</th>
<th>12</th>
</tr>
</thead>
<tbody>
<tr>
<td>String</td>
<td>$FABA'AB'AC'D'$</td>
<td>$FABA'AB'AA'E'F'$</td>
<td>$FABA'AB'AA'E'F'$</td>
</tr>
</tbody>
</table>

Table 11: String Representations for $7 \times m$ Grids

9 Future Work

9.1 Grids

In this work we reviewed what is known about SMTs on grids and then presented results from PARSTEINER94 [27, 30] which characterize SMTs for $3 \times m$ to $7 \times m$ grids. The next obvious question is what is the characterization for an $8 \times m$ grid, or an $n \times m$ grid? Well, this is where things start getting nasty. Even though PARSTEINER94 cuts the computation time of the previous best program for SMTs by an order of magnitude, the computation time for an NP-Hard problem blows up sooner or later, and $8 \times m$ is where we run into the computation wall.

We have been able to make small chips into this wall though, and have some results for $8 \times m$ grids. The pattern for this seems to be based upon
repeated use of the $8 \times 8$ grid which is shown in Figure 44. This grid solution seems to be combined with smaller $8 \times$ solutions in order to build larger solutions. However, until better computational approaches are developed further characterizations of SMTs on grids will be very hard, and tedious.

9.2 Further Parallelization

There remains a great deal of work that can be done on the Steiner Minimal Tree problem in the parallel arena. The first thing to consider is whether there are other ways to approach the parallel generation of $T_j$ that would be more efficient. Improvement in this area would push the computation pendulum even further away from $T_j$ generation and towards SMT extraction.

The next thing to consider is the entire extraction process. The initial generation of the incompatibility matrix has the appearance of easy parallelization. The forest management technique introduced by Cockayne and Hewgill could also be put into a parallel framework, thereby speeding up the preparation for extraction quite a bit.

With this initialization out of the way, decomposition could then be considered. The best possible enhancement here might be the addition of thresholds. As with most parallel algorithms, for any problem smaller than a particular size it is usually faster to solve it sequentially. These thresholds could come into play in determining whether to call a further decomposition, such as the cycle decomposition introduced by Cockayne and Hewgill that was discussed in Section 7.
The final option for parallelization is one that may yield the best results, and that is in the extraction itself. Extraction is basically a branch and bound process, using the incompatibility matrix. This branch and bound is primed with the length of the MST as the initial bound, and continues until all possible combinations have been considered. The easiest implementation here would probably be the idea presented in the paper by Quinn and Deo [46] that served as the basis for the parallel algorithm in Section 6.

9.3 Additional Problems

9.3.1 1-Reliable Steiner Tree Problem

If we would like to be able to sustain a single failure of any vertex, without interrupting communication among remaining vertices, the minimum length network problem takes on a decidedly different structure. For example, in any FST all of the original vertices are of degree 1, and hence any one can be disconnected from the network by a single failure of the adjacent Steiner Point.

We would clearly like a minimum length 2-connected network. The answer can be the minimum length Hamiltonian cycle (consider the vertices of the unit square), but it need not be, as shown in the \( \Theta \) graph given in Figure 45.

Here we can add Steiner points near the vertices of degree 3, and reduce the network length without sacrificing 2-connectivity. This is not just a single graph, but is a member of a family of graphs that look like ladders, where the \( \Theta \) graph has only one internal rung. We hope to extend earlier work providing constructions on 2-connected graphs [31] to allow effective application of an Annealing Algorithm that could walk through graphs within the 2-connected class.

9.3.2 Augmenting Existing Plane Networks

In practical applications, it frequently happens that new points must be joined to an existing Steiner Minimal Tree. Although a new and larger SMT can, in principle, be constructed which connects both the new and the existing points, this is typically impractical. e.g. in cases where a fiber optic network has already been constructed. Thus the only acceptable approach is to add the new points to the network as cheaply as possible. Cockayne has presented this problem which we can state as follows:
Augmented Steiner Network: Given a connected plane graph $G = (V, E)$ (i.e. an embedding of a connected planar graph in $E^2$) and a set $V'$ of points in the plane which are not on edges of $G$, construct a connected plane supergraph $G'' = (V'', E'')$, such that $V''$ contains $V \cup V'$, $E''$ contains $E$, and the sum of the Euclidean lengths of the set of edges in $E'' - E$ is a minimum. In constructing the plane graph $G''$ it is permitted to add an edge connecting a point in $V'$ to an interior point of an edge in $G$. It is also permitted to add Steiner points. Thus, strictly speaking, $G''$ need not be a supergraph of $G$.

The Augmented Steiner Network Problem clearly has applications in such diverse areas as canal systems, rail systems, housing subdivisions, irrigation networks and computer networks. For example, given a (plane) fiber optic computer network $G = (V, E)$ and a new set $V'$ of nodes to be added to the network, the problem is to construct a set $F'$ of fiber optic links with minimum total length that connects $V'$ to $G$. The set $F'$ of new links is easily seen to form a forest in the plane, because the minimum total length requirement ensures that there cannot be cycles in $F'$.

As an example, consider the situation in Figure 46 where $G$ consists of a single, long edge and $V' = v_1, \ldots, v_8$. The optimal forest $F'$ consists of three trees joining $G$ at $f_1$, $f_2$ and $f_3$. It is necessary that extra Steiner points $s_1$, $s_2$ and $s_3$ be added so that $F$ has minimum length.

While we are aware of several algorithms for solving special cases of the Augmented Existing Plane Network Problem, such as those by Chen [5] and Trietsch [50] or the special case where the graph $G$ consists of a single vertex, in which case the problem is equivalent to the classical Steiner Minimal Tree

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Problem, we are not aware of any algorithms or computer programs available for exact solutions to the general form of this problem. Here, “exact” means provably optimal except for round-off error and machine representation of real numbers. Non-exact (i.e. heuristic) solutions are sub-optimal although they may often be found considerably faster.

**References**


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