Calculating Stack Distances Efficiently

George Almási, Călin Cascaval and David A. Padua
Department of Computer Science
University of Illinois at Urbana-Champaign
{galmasi, cascaval, padua}@cs.uiuc.edu

ABSTRACT

This paper describes our experience using the stack processing algorithm [6] for estimating the number of cache misses in scientific programs. By using a new data structure and various optimization techniques we obtain instrumented run-times within 50 to 100 times the original optimized run-times of our benchmarks.

Keywords: memory systems, measurement and monitoring, simulation, cache modeling

1. INTRODUCTION

Caches have become one of the most important components in computer systems. As processor speed increases much faster than memory speed, the memory subsystem becomes the bottleneck. To alleviate this behavior, computer architects have designed deeper memory hierarchies, including several levels of cache.

The stack algorithm [6] was originally designed for modeling virtual paging, i.e. to operate on a program trace consisting of virtual page references, but in the recent past has been used mainly to model cache behavior, by tracing cache line references [7, 8, 4, 10].

Different architectures exhibit different memory systems with caches organized in different configurations. In order to make the best use of the memory hierarchy, either the programmer or the compiler must be aware of the cache organization of the machine. We believe that the programmer should not be burdened with this responsibility, therefore we have developed a memory hierarchy model that helps the compiler to predict the number of cache misses, and thus reorganize the code to optimize the cache behavior. This memory model is based on the stack distances obtained from a stack processing algorithm [6].

The main advantage of the stack algorithm in simulating cache behavior is that it allows the estimation of the number of misses for caches of any size in a single pass through the trace. Variants of the algorithm have been used to simulate caches of multiple line sizes.

Unfortunately, since most programs reference about one hundred times more cache lines than virtual pages, the stack algorithm in its original form becomes very time-consuming. Several researchers [1, 4, 7] proposed changes to the original algorithm to improve its speed.

Seeking to further improve the performance of the stack algorithm, we introduce two new data structures and corresponding algorithms, each of which is more suitable for a particular kind of application. The interval tree approach works well for programs with long traces and relatively good locality, whereas the preallocated tree approach is more suited to shorter traces with bad locality.

The rest of this paper is organized as follows. First we give a short description of the LRU stack algorithm, and briefly describe the work done by others. In section 4 we describe the new data structures and algorithms we used. Section 5 details the experiments we ran using the Perfect Benchmarks [2] benchmark suite. We conclude by qualifying our results and identifying possible areas of future research.

2. STACK DISTANCES

Simulation of memory hierarchies dates back to 1970, when Mattson et al. [6] presented an evaluation of virtual memory page replacement strategies with a stack. His algorithm takes a trace of memory references – cache line references or virtual page references in a program – and builds a stack as follows: every memory location is pushed onto the stack when it is referenced. Locations that have been referenced a long time ago sink towards the bottom of the stack, but locations that are referenced again are extracted from the stack and pushed back on top.

The depth from which a certain reference had to be extracted in the stack is called the stack distance of that reference. By convention, the stack distance of first-time references is ∞.

The result of the stack processing algorithm is a trace of stack distances. These can be coalesced into a histogram of stack distances. Figure 1 shows the stack distance histogram for the QCD benchmark of the Perfect benchmark suite.

The stack distance histogram is a useful measure of locality, both visually and numerically. It can be used to calculate the number of out-of-core page references, or equivalently, the number of cache misses, for any memory or cache size. For physical memory size C, all the accesses at stack distances of less than C are in-core, and all the others are out-of-core. Splitting the stack depth histogram at C, the
area under the histogram curve at the left of $C$ is the number of in-core references, whereas the area to the right of $C$ represents the out-of-core accesses.

This is true because caches and virtual memory systems have the inclusion property: a cache of size $C+1$ will always include a cache of size $C$ for any replacement strategy.

The LRU replacement policy ensures that if a cache line $L$ is accessed several times and the number of distinct references to other cache lines between two consecutive accesses to $L$ is less than the total number of cache lines in the cache, the subsequent accesses to $L$ are hits.

The stack processing algorithm can therefore be used exactly compute cache misses in fully associative caches, and to approximate them in set associative caches.

3. RELATED WORK

Following Mattson’s work, other researchers extended the usefulness of the stack algorithm. Traiger and Slutz [9] described a one-pass algorithm to simulate multiple page sizes in virtual memory. Their technique can be used to simulate multiple cache line sizes in one pass. Thompson and Smith [8] extended the stack algorithm to simulate write-back caches, which do not satisfy the inclusion property. By adding a dirty bit to each block in the stack they were able to count the number of writes that can be avoided when the dirty block is still resident in the cache.

The stack algorithm is however very expensive to run, especially if the stack becomes large enough. It was soon recognized that more efficient data structures were needed to do the job of the stack search. Bennett and Kruskal [1] presented an algorithm which replaces the stack with a pre-allocated hierarchy of partial sums. Hill and Smith [4] used a forest of trees to simulate multiple cache associativities; Sugumar and Abraham [7] used a generalized binomial tree for the same purpose.

4. IMPLEMENTATIONS OF THE LRU STACK ALGORITHM

Figure 2 shows a formal three-step description of the LRU stack algorithm, as first described by Mattson in [6]. We use this description as the basis for the algorithms we present.

It is assumed henceforth that the algorithm is operating on a memory trace of length $N$ that contains $M$ distinct memory references (obviously $M \leq N$). The trace is indexed with the letter $\tau$, where $1 \leq \tau \leq N$. For the notations used in this paper refer to Section 7.

4.1 Naive Implementation

This implementation directly follows the algorithm presented above. The stack is represented as a doubly linked list. For each reference in the trace, the first two operations (search and count) are executed simultaneously by traversing the stack top to bottom. If the element exists in the stack, its distance from the top of the stack is recorded. Finally the element is moved from its current position to the top of the stack — the update stage. If the element is not found, $\infty$ is recorded as its stack distance and the element is pushed on top of the stack.

4.1.1 Analysis

For each reference in the trace the work done is, in the worst case, $M$ (due to the traversal of the linked list). The total complexity is thus $O(NM)$.

The worst case doesn’t happen very often. In fact, many programs exhibit excellent locality, causing many references to lie close to the top of the stack. Unfortunately the few references that are near the bottom of the stack cause huge slow-downs, resulting in terrible overall performance.

4.2 Markers Algorithm

The slowest part of the naive algorithm is the linear traversal of the linked list that makes up the stack. The markers algorithm, presented next, replaces linear search with a combination of hashtable lookup and two-level linked lists.

The search phase of the markers algorithm is done using a hash table that associates a cache line/memory/page reference with its current place in the linked list. Given enough hash buckets, hashtable access and update are $O(1)$ operations. The number of necessary hash buckets can be approximated with $M$, the number of distinct references in the trace.
Unfortunately finding an element in the middle of the stack by using the hashtable is not enough. The stack depth of the element needs to be counted. To avoid traversing the stack from top to bottom, a set of markers are interspersed in the linear list implementing the stack, one about every D elements. The markers form another doubly linked list, and each marker records its distance from the top. To find out the depth of a memory reference in the stack, one needs to find the nearest marker by traversing the stack (a marker would be at most D steps away) and then look up the marker’s distance from the top.

When an element is removed from the stack and inserted at the top, the markers between the top and the element need to be updated. This involves at most M/D steps.

4.2.1 Analysis

The cost per memory reference of this algorithm is at most D + M/D (the cost of finding a marker, plus the cost of updating all markers up to the beginning of the stack). D can be varied at runtime by adding or removing markers, in order to minimize the cost: assuming can be varied at runtime by adding or removing markers, in order to minimize the cost: assuming \( D = \lceil \sqrt{M} \rceil \), the cost evaluates to \( O(\sqrt{M}) \) per element, or \( O(N + \sqrt{M}) \) total.

4.3 Alternative Data Structures

The major stumbling block in implementing more efficient versions of the LRU stack algorithm is the implementation of the stack as a linear list. We will present a formulation of the LRU stack algorithm that does not use a stack. We will closely follow Bennett and Kruskal’s [1] notation.

4.3.1 Definition 1.

To aid us in the search phase of the algorithm, we formalize the concept of the hashtable \( \mathbf{P} \), which we already used informally in the markers algorithm. The hash table contains, for each memory reference in the trace, the time/index of its last use. We define \( \mathcal{J} \) as the set of indices of references to \( z \) that occurred before an index \( \tau \) in the trace:

\[
\mathcal{J}_z = \{ i \mid 0 \leq i < \tau \land x_i = z \}
\]

Using \( \mathcal{J} \) we define the hashtable \( \mathbf{P}_\tau \) as follows:

\[
\mathbf{P}_\tau(z) = \begin{cases}
\max \{ i \mid i \in \mathcal{J} \} & \text{if } \mathcal{J} \neq \emptyset \\
\text{undefined} & \text{otherwise}
\end{cases}
\]  

Note 1. \( \mathbf{P}_\tau \) is time dependent; hence the \( \tau \) subscript.

Note 2. \( \mathbf{P}_\tau(z) \) is undefined when a cold miss occurs, i.e. when there is no previous reference to \( z \).

4.3.2 Definition 2.

Next we define \( \mathbf{B} \), a mapping from the trace indices \( 0 \ldots N-1 \) to \( \{0, 1\} \). Intuitively, \( \mathbf{B} \) flags all the references that are on the stack at time \( \tau \). Like \( \mathbf{B} \), \( \mathbf{B} \) changes with time and therefore is subscripted with \( \tau \). \( \mathbf{B}_\tau(i) \) is defined as follows:

\[
\mathbf{B}_\tau(i) = \begin{cases}
1 & \text{if } \mathbf{P}_\tau(x_i) = i \\
0 & \text{otherwise}
\end{cases}
\]  

\( \mathbf{B} \) is an alternative representation of the stack: \( \mathbf{B}_\tau(i) = 1 \) precisely for the elements \( i \) in the trace that are part of the stack at any given moment.

4.3.3 Definition 3.

Given \( \mathbf{P} \) and \( \mathbf{B} \) we can define \( \text{dist}(\tau) \), the stack distance of the element \( x_\tau \) in the program trace: it is the number of 1’s in \( \mathbf{B} \) between the last reference to \( x_\tau \) and \( \tau \).

\[
\text{dist}(\tau) = \begin{cases}
|\mathcal{H}| & \text{if } \mathbf{P}_\tau(x_\tau) \text{ is defined} \\
\infty & \text{otherwise}
\end{cases}
\]  

where \( \mathcal{H} \) is the subset of trace indices between \( \mathbf{P}_\tau(x_\tau) \) and \( \tau \) whose values are 1:

\[
\mathcal{H} = \{ i \mid \mathbf{P}_\tau(x_\tau) < i < \tau \land \mathbf{B}_\tau(i) = 1 \}
\]

We can now reformulate the stack algorithm by using \( \mathbf{P} \) and \( \mathbf{B} \) instead of the stack.

Repeat for each reference \( x_\tau, 0 \leq \tau < N \):

- **search**: compute \( \mathbf{P}_\tau(x_\tau) \);
- **count**: evaluate \( \text{dist}(\tau) \). If \( \mathbf{P}_\tau(x) \) is undefined then \( \text{dist}(\tau) \) is defined as \( \infty \);
- **update**: change \( \mathbf{B} \) and \( \mathbf{P} \) as follows:

\[
\mathbf{B}_{\tau+1}(i) = \begin{cases}
1 & \text{if } i = \tau \\
0 & \text{if } i = \mathbf{P}_\tau(x_\tau) \\
\mathbf{B}_\tau(i) & \text{otherwise}
\end{cases}
\]

\[
\mathbf{P}_{\tau+1}(z) = \begin{cases}
\tau & \text{if } z = x_\tau \\
\mathbf{P}_\tau(z) & \text{otherwise}
\end{cases}
\]

Figure 3: Modified Stack Algorithm

4.4 Bennett and Kruskal’s Algorithm

Given the alternative data structures we just presented, the stack algorithm boils down to maintaining \( \mathbf{B} \) and \( \mathbf{P} \) and computing \( \text{dist}(\tau) \) from \( \mathbf{B} \) and \( \mathbf{P} \). All algorithms presented in the remainder of this paper attempt to compute \( \text{dist}(\tau) \) in a more efficient way.

Bennett and Kruskal’s algorithm [1] maintains a modified form of \( \mathbf{B} \): it uses a hierarchy of partial sums \( \mathbf{B}^1, \mathbf{B}^2, \ldots, \mathbf{B}^L \), where \( L = \lceil \log(N) \rceil \). Renaming \( \mathbf{B} \) to \( \mathbf{B}^0 \), the partial sum hierarchy is set up such that for some chosen interval \( m \), at any time \( \tau \),

\[
\mathbf{B}^\tau_s(j) = \sum_{i=j}^{i=m-1} \mathbf{B}^{\tau-1}_s(i)
\]

This formula describes an \( m \)-ary tree of nodes having the value of each node being equal to the sum of the values of its children.

Calculating the number of 1’s between the indices \( \mathbf{P}(x_\tau) \) and \( \tau \) is now a matter of traversing the partial sum hierarchy, as shown in Figure 4. The figure presents the first 31 elements of a trace. We trace the 31’s last access, and \( x_{31} \) last occurred in position 4.

The third step, updating \( \mathbf{B} \), also becomes a matter of tree traversal, since all partial sums on the path from the root of the hierarchy to the leaf node are affected.

The algorithm needs two traversals of the tree. The first traversal, from the root to index \( \mathbf{P}(x_\tau) \), deletes \( \mathbf{P}(x_\tau) \) as the last reference to \( x \) by setting \( B(\mathbf{P}(x_\tau)) \) value to 0 and adjusting all partial sums along the path. The second traversal is from the root to index \( \tau \) and sets \( B(\tau) \) to 1, again adjusting partial sums on the path. For reasons of brevity we are not going to fully explain the algorithm, except to mention
that our major improvement, to be presented in the next
sections, replaces the two traversals with a single traversal
of the tree.

4.4.1 Analysis
Since the tree traversal is an \(O(\log(N))\) operation and the
location finder works in constant time (hashable lookup is
\(O(1)\)), the total execution time is \(O(N\log(N))\). The memory
requirements of the algorithms are very large because \(B\) and
its partial sums are represented explicitly in memory.

4.5 Hole-based Algorithms
We define a hole as a memory reference in the program
trace that is not the latest reference to a particular location
at time \(\tau\). Holes are the zero elements in \(B_{\tau}\).

Whereas values of 1 in \(B\) (and corresponding latest ref-
ences) are newly created and then destroyed all the time,
holes have the property of being created and never destroyed.

Using the concept of holes, the stack distance at index \(\tau\)
can be expressed as

\[
dist(\tau) = \tau - P(x_{\tau}) - holes(\text{holes}(x_{\tau}))
\]

where \(\text{holes}(i)\) is the number of holes in the program trace
between indices \(i\) and \(\tau\). Here we are in effect counting the
\(0\)'s in \(B\) instead of counting the \(1\)'s, and adjusting Equation
(3) to reflect this.

Holes can be represented more efficiently than latest refer-
ences. We will present two kinds of algorithms based
on holes, a variant in which holes are held by an interval
tree and another which is a faster version of Bennett and
Kruskal's algorithm.

4.5.1 Interval Tree of Holes
An interval tree can be used to efficiently represent an
ordered set of mutually disjunct intervals \(I = \{[i_{11}, i_{12}],
[i_{21}, i_{22}], \ldots, [i_{n1}, i_{n2}]\}\). In our case the intervals in \(I\)
are all bounded by natural numbers (indices in the program
trace). The intervals represent contiguous sets of indices
that are holes in the trace.

Interval trees (Figure 5) are represented as a quasi-balanced
binary trees \(BT\) (such as red-black trees [3] or AVL trees [5])
in which each node \(n\) represents the closed interval \([k_1(n), k_2(n)]\).
The tree ordering corresponds to the order of the intervals
in \(I\); thus \(k_1(n) > k_2(\text{left}(n))\) and \(k_2(n) < k_1(\text{right}(n))\),
where \(\text{left}(n)\) and \(\text{right}(n)\) are respectively the left and
right children of \(n\).

4.5.2 The Partial Sum Hierarchy
We use the interval tree to evaluate the number of holes
between \(P(x_{\tau})\) and \(\tau\). There are no holes beyond the current
index \(\tau\) (a logical impossibility considering the definition of
a hole). Thus we are left with counting the number of holes
at indices larger than \(P(x_{\tau})\). To do this, we follow Bennett
and Kruskal's method and associate a value \(\text{sum}(n)\) with
each interval node \(n\), to hold the sum of holes contained in
the children of \(n\). Our hole tree now becomes equivalent to
Bennett and Kruskal's partial sum hierarchy.

A slight optimization is to make \(\text{sum}(n)\) hold the sum of
holes in the right subtree of \(n\) instead of \(n\) itself. In Figure
6 the shaded boxes contain partial sums of the right
subtree of their nodes, as indicated by the dashed arrows.
This optimization reduces the number of right subtree derefer-
enciations when the next target of the tree traversal is the
left subtree (in Figure 6 the nodes marked with (*) will not
need to be dereferenced).

The counting algorithm works like this: we traverse the
interval tree from the root towards the leaf node closest to
index \(i = P(x_{\tau})\). We carry a partial sum along the path,
and add to it the sum of holes in all subtrees encountered
to the right of the path (i.e. having indices larger than \(i\)).

4.5.3 Updating the Interval Tree
We now extend the counting algorithm to include the third
component of the LRU stack algorithm: \texttt{update}. We
need to update the tree structure as well as the partial sum
hierarchy residing in it.

With regard to inserting a new hole \(p\) into the interval
tree, there are several cases to consider:
• $p$ may be adjacent to a single existing interval $[k_1, k_2]$ in the tree, i.e. $p = k_1 - 1$ or $p = k_2 + 1$. In this case the interval is adjusted to include $p$.

• $p$ may be adjacent to two intervals $[k_1, p - 1]$ and $[p + 1, k_3]$. In this case the intervals are fused into a single interval $[k_1, k_3]$ prompting the deletion of one of the nodes and the potential re-balancing of the whole tree.

• $p$ may not be adjacent to any intervals in $BT$. In this case a new node is created, to hold the interval $[p, p]$. Again, the tree may need to be rebalanced.

The partial sum hierarchy is updated by changing the sum values of the nodes on the path from the root to the affected interval. Figure 6 illustrates the operation of counting holes and inserting a new hole at location 101 in an example tree. The Appendix lists the algorithm that performs this operation.

### 4.5.4 Analysis

The algorithm presented in Figure 6 is based on a quasi-balanced binary tree. $dist(p, n)$ is a variant of the insertion operation for quasi-balanced binary trees, which makes it an $O(log(M))$ operation (the number of disjunct hole intervals, and thus nodes in the tree, is always less than $M+1$). Thus the total execution time of the Binary Tree Hole Algorithm is $O(Nlog(M))$.

### 4.6 Preallocated Tree of Holes

A tree of holes can also be implemented as a preallocated fixed tree $\{B^0, B^1, ... B^{b^{10}(N)}\}$, like the one of Bennett and Kruskal. Unfortunately the memory requirements for the whole tree get quickly out of hand: for a program trace of length $2^{31}$ (a realistic number for today’s programs) we need to allocate $1 + 2 + 2^2 + ... + 2^{30} = 2^{31} - 1$ locations.

The silver lining is that not all locations need to be of the same size. Elements of $B^0$, for instance, need to hold only one bit; elements of $B^1$ need to be two bits each, and so on; the total memory requirement is $\frac{1}{2} \times (2^{30} + 2 \times 2^{29} + 3 \times 2^{28} + \ldots + 31 \times 2^0) \approx 536.87MB$, which fits into the virtual memory of most modern workstations. Also, the algorithm does not use all of this memory at once, but rather progresses slowly through it as the trace is analyzed. This allows for huge portions of the preallocated tree to reside in virtual storage.

#### 4.6.1 Analysis

Since the tree is preallocated, and has $N$ leaf nodes, tree traversal is now an $O(log(N))$ operation rather than $O(log(M))$, which would seem to make this algorithm impractical. Also, the tree needs to be allocated before the program is run, which means that the user has to guess $N$.

However, once $N$ is calibrated, the algorithm becomes the fastest we tried so far, outperforming Bennett and Kruskal’s by a factor of up to 2:1. The reason for this is that only one tree traversal is needed per element, as opposed to two for Bennett and Kruskal’s algorithm.

## 5. EXPERIMENTAL RESULTS

We selected the Perfect Benchmarks [2] as our experimental base and instrumented them with a source-to-source translator to generate a program trace. Rather than storing the program trace we hooked up the analyzer to the instrumented benchmark directly, and generated the trace and the histogram on the fly.

At first we used the naive implementation of the LRU stack algorithm, and experienced a drastic slowdown. In an effort to find better implementations of the LRU algorithm we experimented with all algorithms described in this paper.

We ran our experiments on a 270MHz Ultrasparc Solaris machine. Table 1 summarizes the results we obtained. Benchmarks are listed by name; the total number of references and the maximum stack are included.

The algorithms we measured are the following:

- **orig** is the runtime of the original non-instrumented benchmark.
- **null** measures the trace generation overhead, but the stack processing part is not implemented. We measured “null” to find out how much the benchmarks are affected by just generating the trace.
- **kru and pre** are preallocated implementations of Bennett and Kruskal’s, and the preallocated tree hole based algorithm’s, respectively.
- **avl and rb** are interval tree implementations using AVL and red-black trees respectively.
- **mrk** is the markers algorithm. Many of the numbers are missing because we had to abort runs that were taking too long.

We also measured the relative overhead of the stack computation. Figure 7 breaks down the total runtime of each benchmark into the time spent in the original benchmark, instrumentation overhead (i.e. time spent generating the program trace), hash table lookup overhead and stack computation overhead.

The interval tree based algorithms have better theoretical bounds than the preallocated tree algorithms, $O(Nlog(M))$ versus $O(Nlog(N))$. There are several reasons why the preallocated algorithms tend to yield better execution times in practice:

![Figure 7: Execution Time Breakdown for the Preallocated Hole Tree Algorithm](image)
### Table 1: Perfect Benchmark run times (seconds)

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- The interval tree implementation severely stresses the memory bandwidth of the host processor. For each element in the program trace the interval tree algorithm generates about $3 \cdot \log(M)$ additional references while traversing the interval tree: in each tree node at least one node key is accessed; in addition the node’s partial sum is accessed and one of the leaves is dereferenced.

The value of $M$ can be approximated with the measured maximum stack depth, which for most of our algorithms yields an AVL tree height of around 20 to 25, resulting in up to 75 extra memory accesses per element in the memory trace. In the case of red-black trees the number of references is even higher.

By comparison the preallocated tree implementation generates only $\log(N)$ (or $2 \cdot \log(N)$, in the case of Kruskal’s algorithm) references. In practice we limited $N$ to $2^{31}$, which means 31 memory references for each tree. In addition the preallocated tree is built such that adjacent nodes at lower levels tend to be clustered into the same cache line, resulting in good spatial locality.

- The interval tree implementation relies on dynamic memory allocation as the interval tree shrinks and expands in the course of the process. We were able to gain up to 33% in execution speed by writing our own memory allocators (this gain is included in the performance figures).

The better speed of the preallocated strategy comes, however, at the cost of extremely high memory usage (about 600 MBytes of virtual memory for the preallocated tree) and a hard limit of $2^{31}$ references in the memory trace. For a few of the benchmarks this limit is exceeded.

The interval tree implementation, if slower, has no inherent limitation with respect to the trace size and delivers reasonable performance. We see it as a more useful tool on the whole. The AVL tree is preferable to the red-black tree, since the higher reordering cost is clearly amortized by the lower average tree height.

In conclusion, the preallocated implementation works better for programs with short traces, bad locality and large cores (that is, large $M$ and relatively small $N$ values), whereas the interval tree implementation works better on long traces with good locality and small cores (larger $N$, smaller $M$ values).

### 6. CONCLUSIONS, FUTURE WORK

In this paper we have presented our experience using stack algorithms [6] to simulate the cache behavior of scientific programs. We have implemented several versions of an algorithm that was devised to simulate virtual paging, and although accurate for caches, is not performing very well due to the large number of memory references.

The main contribution of this paper is a novel way of computing stack distances based on holes. This allows the algorithm to perform fast enough to be convenient to use.

In its present form, the algorithm is easily modifiable for simulating multiple cache line sizes, as described in [9]. However we have simulated fully associative caches. Finding a way to extract information on arbitrary associativity [4] using a single stack, or interval tree, is a matter of future research. Further improvements in simulation speed, and reductions in memory footprint, would also be welcome.

### 7. NOTATION

This section enumerates and explains some of the symbols we used throughout this paper.

- $N$: number of references in the program trace under consideration.
- $M$: number of distinct memory references in the program trace. In effect $M$ is equal to the size of the memory used by the program we are analyzing.
- $\tau$: used as the current index in the trace. As such, $0 \leq \tau < N$. The stack algorithm processes the program trace sequentially; $\tau$ always denotes the current position processed by the algorithm.
- $x_{\tau}$: denotes the memory reference at index $\tau$ in the program trace.
- $P$: a mapping from memory references to trace indices. Since $P$ changes in time, it is normally indexed with $\tau$, the current index in the program trace.
- $B$: a mapping from trace indices to booleans. $B_{\tau}(i)$ is set if at moment $\tau$ the location referenced at position $i$ in the trace is the latest reference to its location.
- $\text{dist}(\tau)$: the stack distance corresponding to the location referenced in position $\tau$ in the trace. This is the number we compute for each element in the trace.
holes, (i): the number of holes in the program trace between indices i and \( \tau \) at moment \( \tau \). \( i < \tau \) by definition.

8. REFERENCES


APPENDIX

The Interval Tree Algorithm

function dist (n, p) begin
    if (p < key1(n) - 1) /* continue search left */
        if (left(n) ≠ nil)
            return sum(n) + dist(left(n))
        else /* can’t continue left - no nodes left */
            left(n) := new interval (p, p)
            return 0
    else if (p > key2(n) + 1) /* continue search right */
        if (right(n) ≠ nil)
            sum(n) := sum(n) + 1
            return dist(right(n))
        else /* can’t continue right - no nodes left */
            right(n) := new interval (p, p)
            sum(right(n)) := 0
            return 0
    else if (p = key1(n) - 1 AND p = key2(left(n)) + 1)
        /* merge left node */
        key1(n) := key2(left(n)) + 1
        remove_node(left(n))
        rebalance(n)
        return key2(n) - p + sum(n)
    else if (p = key1(n) - 1) /* add to node */
        key1(n) := p
        return key2(n) - p + sum(n)
    else if (p = key2(n) + 1 AND p = key1(right(n)) - 1)
        /* merge right node */
        key2(n) := key2(right(n))
        remove_node(right(n))
        rebalance(n)
        return sum(n)
    else if (p = key2(n) + 1) /* add to node */
        key2(n) := p
        return sum(n)
    else /* internal error */
    end if
end