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MASSIVELY PARALLEL ALGORITHMS FOR TRACE-DRIVEN CACHE SIMULATIONS

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Massively Parallel Algorithms for Trace-Driven Cache Simulations

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Abstract

Trace-driven cache simulation is central to computer design. A trace is a very long sequence, $z_1, \ldots, z_N$, of references to lines (contiguous locations) from main memory. At the $t^{th}$ instant, reference $z_t$ is hashed into a set of cache locations, the contents of which are then compared with $z_t$. If at the $t^{th}$ instant $z_t$ is not present in the cache, then it is said to be a miss, and is loaded into the cache set, possibly forcing the replacement of some other memory line, and making $z_t$ present for the $(t+1)^{th}$ instant. The problem of parallel simulation of a subtrace of $N$ references directed to a $C$ line cache set is considered, with the aim of determining which references are misses and related statistics.

A simulation method is presented for the Least-Recently-Used (LRU) policy, which regardless of the set size $C$ runs in time $O(\log N)$ using $N$ processors on the exclusive read, exclusive write (EREW) parallel model. A simpler LRU simulation algorithm is given that runs in $O(C \log N)$ time using $N/\log N$ processors. We present timings of the second algorithm's implementation on the MasPar MP-1, a machine with 16384 processors. A broad class of reference-based line replacement policies are considered, which includes LRU as well as the Least-Frequently-Used and Random replacement policies. A simulation method is presented for any such policy that on any trace of length $N$ directed to a $C$ line set runs in time $O(C \log N)$ time with high probability using $N$ processors on the EREW model. The algorithms are simple, have very little space overhead, and are well-suited for SIMD implementation.

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1 Introduction

A cache is a high-speed memory on the access path to a larger, slower main memory. Cache performance is critical to the overall performance of computer systems [10], and consequently a tremendous amount of effort is put into the evaluation of cache designs. This is particularly true for RISC microprocessor designs, where the ratio of the time needed to access an off-chip cache to that needed to access the main memory can be as high as 10 [10], and the off-chip cache is typically at least 10 times smaller than the main memory. Trace-driven simulations, which evaluate cache performance on actual reference streams taken from characteristic programs, are the most reliable and widely used tools for cache design evaluation. These simulations require a great deal of computation, because of the many different design possibilities that are simulated, and because of the length of the reference traces that drive the simulation [19].

Data is moved between main memory and the cache in contiguous blocks called lines. Every memory line is hashed to some fixed cache set, but may be placed in any one of the $C$ physical cache lines in the set. In emerging computer designs, a microprocessor might be supported by a 1 Megabyte off-chip cache, with a line size of 128 bytes, and a set size $C = 4$. A miss occurs whenever a memory line is referenced, but is not found in its set. The cache hardware then fetches the desired line from main memory, overwriting another line in the same set if the set is full. The rule used to select which line to replace is called the replacement policy. An effective, widely used policy is Least-Recently-Used (LRU), which simply replaces the line accessed least recently. The objective of a trace-driven simulation is to determine which references in the trace are misses. Given the identities of the misses, statistics of chief interest in cache design are easily computed, such as the fraction of read misses, the fraction of write misses, and the number of write-backs (stores of modified lines) from cache to main memory.

Heidelberger and Stone [9] showed that it is valuable to simulate a long trace directed to a few sets, when cache miss statistics between sets are highly correlated. High correlation removes the need to simulate all sets, but also removes the easy parallelism that might be exploited by simulating a large number of sets in parallel on different processing elements (PEs). A massively parallel method to handle the simulation of a long trace targeted to a single set allows more powerful, flexible solutions.

We consider the problem of determining the misses in a given reference trace, $x_1, ..., x_N$, directed to a set of size $C$. An algorithm is presented (Section 3) that solves this problem in $O(\log N)$ time using $N/\log N$ PEs, on the exclusive-read, exclusive-write (EREW) model of a parallel machine. The algorithm and its complexity do not depend on $C$. The algorithm computes the stack distance $\Delta_t$ associated with each reference $x_t$ [16]. If $x_t$ is not a first reference to a line then $\Delta_t$ is the smallest set size for which $x_t$ would be a hit; otherwise $\Delta_t = \infty$.

In Section 4, we present an alternative LRU simulation, with running time $O(C \log N)$ time using $N/\log N$ PEs.

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1 Recent experiments (private communication from Harold Stone) have validated that high correlation exists between sets, but have also shown that special care must be taken when selecting the sets which are analyzed, as the measured miss ratio from an arbitrary set simulation may not be an accurate predictor of the overall miss ratio.
PEs on the EREW model. The algorithm computes the stack distance at level \( C \), \( \Delta(C) \), for each reference \( x \). If \( x \) is not a first reference to a line then \( \Delta(C) \) is the smallest set size \( \leq C \) for which \( x \) would be a hit; otherwise \( \Delta(C) = \infty \). The algorithm is simple and the implicit constant in the time bound is favorable. We report timings of this algorithm’s performance on a MasPar [4] SIMD computer having 16384 PEs.

In Section 5, a broad class of reference-based replacement policies is considered. Roughly, the class contains all stack replacement policies where priorities controlling line replacement are static and can be computed efficiently in parallel. This class includes LRU as well as:

- **OPT**: Replace the line referenced most remotely in the future. This unrealizable policy provably minimizes the number of misses. Its simulation gives a baseline against which realizable policies can be measured.

- **Least-Frequently-Used or LFU**: Replace the line accessed least often in the past. Ties can be broken by, for example, giving higher priority to the reference that has been in the cache the shortest length of time.

- **Random**: Replace one of the \( C \) lines, chosen independently and uniformly at random. Random replacement is easy to implement; furthermore, there is evidence that if the total number of lines in the cache (not just the lines in one set) is sufficiently large, the policy works nearly as well as any other implementable policy[10].

In Section 5, an algorithm is presented for reference-based policy simulation. Given any trace of \( N \) references targeted to a \( C \) line set, the algorithm runs in time in \( O(C \log N) \) with high probability using \( N \) PEs on the EREW model. (The algorithm is probabilistic, the choice of trace is not.) In Section 5.3, we extend the class of reference-based replacement policies to include an aging mechanism, whereby stale lines lose priority and tend to be flushed from the cache. Accommodating this mechanism increases the algorithm’s running time to \( O(C \log^2 N) \).

Our algorithms are simple, require at most \( O(\log N) \) space per PE, and break the computation down into calls to a few primitive parallel subroutines. As a result the algorithms are well-suited for SIMD architectures, such as the Connection Machine [11] or MasPar [4]. The \( O(C \log N) \) with high probability bound holds because we have assumed that a fast probabilistic parallel algorithm [18] is used to solve a certain trapezoidal decomposition problem (Sections 2, 5). Adopting the notation of [18], this algorithm runs in \( \tilde{O}(\log N) \) time using \( N \) PEs, meaning that there is a constant \( k \) such that the time exceeds \( km \log N \) with probability less than \( N^{-m} \) for any \( m > 1 \). In practice, simpler, deterministic methods may do better, while raising the asymptotic time bound to \( O(C \log^2 N) \).

For simplicity, we have assumed the problem size \( N \) is comparable to the number of PEs, so that it is as if each PE handles a few references (up to \( \log N \)). However, a “supersaturated” setup [8] may be effective in practice, where a large block of consecutive references would be loaded in the local memory of each PE. Our algorithms generalize to that setup, by using efficient supersaturated implementations of the underlying
parallel primitives (cf. [12, 17]). Indeed, our implementation of the LRU algorithm is a supersaturated one, with complexity $O(C(N/P + \log P))$ for a reference trace with $N$ elements on an architecture with $P$ processors.

Collecting the cache miss statistics mentioned above adds just $O(\log N)$ time. Moreover, by the nature of the replacement policies and the simulation methods, statistics for each set size up to $C$ can be computed at this cost. All of our algorithms can be adapted for efficient simultaneous simulation of many sets, by the simple device of initially sorting the references on the basis of their set identifiers.

Heidelberger and Stone [9] had the original insight that trace-driven simulation of an LRU cache set could be parallelized. Their algorithm is intended for a network of $P$ MIMD processors, and requires $P \ll N$ for good speedup. Our work was motivated by theirs; our algorithms are different, apply to a larger class of replacement policies, and to a different class of architectures. Lin, Baer, and Lazowska have considered parallelizing cache simulations, in the context of multiprocessor cache protocols[15]. Their method assumes that each individual processor's cache is simulated on a different PE, so that the degree of parallelism is limited to the number of caches in the simulated system. An important and beautiful paper on cache simulation was published in 1970 by Mattson, Gecsei, Slutz, and Traiger[16]. Most of our notation is taken from that paper.

The practical utility of implementing trace-driven cache simulations on today's SIMD computers has yet to be shown, although our implementation proves the great promise of the approach. It seems likely that a very long reference trace will have to be partitioned into blocks, where one block is processed at a time. The I/O problem is to move the blocks to the processors fast enough to keep them busy. An attractive alternative is to use a synthetic trace; for example Thiebault, Stone, and Wolf [20] recently proposed a simple method for random generation of realistic traces.

## 2 Preliminaries

### 2.1 Cache Notation

Henceforth, we focus on a single set cache, and treat its size $C$ as a parameter. Let $B_t(C)$ denote the set of lines stored just after reference $x_t$. Each reference must be cached, so $x_t \in B_t(C)$ for all $C \geq 1$ and $t \geq 1$. (By convention, $B_t(0)$ is the empty set.) If the cache is full ($|B_t(C)| = C$) and $x_t$ is a miss ($x_t \notin B_{t-1}(C)$) then $x_t$ replaces a line in $B_{t-1}(C)$. We refer to this replaced line as $y_t$. All of the replacement policies we consider are stack policies [16], meaning if a reference is a hit given that the cache size is $C$ then the reference will remain a hit if the cache size is increased to $C + 1$. That is,

$$B_t(C) \subseteq B_t(C + 1) \quad \text{for all } C \geq 0.$$

This inclusion allows us to order the lines of the cache by the least size needed for their appearance.
Figure 1 gives an example, for the LRU replacement policy. The trace length $N = 18$, and the lines are labeled $a - d$. The first level, $s_t(1)$ coincides with the trace itself. Consider the first two levels; i.e., the cache contents given $C = 2$. The cache is initially empty. The first two references, $x_1 = a$, $x_2 = c$, miss, and as a result $B_1(2) = \{a\}$, $B_2(2) = \{a, c\}$. The third reference, $x_3 = b$, also misses, forcing the replacement of $y_3 = a$, yielding $B_3(2) = \{b, c\}$. The fourth reference, $x_4 = c$ hits, so $B_4(2) = B_3(2)$, and so forth.

Hit and miss statistics are easily extracted from stack distances, defined as follows. Let the level $i$ stack distance $\Delta_t(i)$ denote the smallest cache size $\leq i$ such that $x_t$ is a hit, or $\infty$ if $x_t$ is a miss for cache size $i$. Thus, given $\Delta_t(C)$, for $t = 1, \ldots, N$, we can extract the hits for any cache size $c < C$. More generally, define the stack distance $\Delta_t = \lim_{C \to \infty} \Delta_t(C)$ to be the smallest cache size such that $x_t$ is a hit, or $\infty$ if there is no prior reference to the same line ($x_s = x_t$ for some $s < t$). Stack distances are shown in Figure 1.

2.2 Parallel Processing Model

Our algorithms are well-suited for a wide variety of parallel architectures, because the algorithms transform data in simple ways using a small number of basic, highly parallelizable operations. However, to state precise time and processor requirements, we must choose a precise model of parallel computation. The EREW (exclusive read, exclusive write) model (cf. [14]) provides a nice blend of simplicity and realism. In this model, the PEs operate in lockstep. There is a global shared memory, supporting at unit cost any pattern of accesses except those where two PEs simultaneously access the same location.

We state the complexity of our algorithms with respect to the EREW model. We now list the parallel subroutines used in our algorithms, and their complexities on the EREW model.

- Merging: Two sorted lists each of length $N$ can be merged in time $O(\log N)$ using $N/\log N$ PEs, via Batcher's odd-even merge algorithm [2].
• Sorting: A list of length \( N \) can be sorted in time \( O(\log N) \) time using \( N \) PEs [5].

• 2d Ranking: Given points \((x_i, y_i), i = 1, \ldots, N\), compute for each point \((x_i, y_i)\) its rank, the number of other points \((x_j, y_j)\) strictly above and to the right: \( x_i < x_j \) and \( y_i < y_j \). In slightly different form, this is the problem of computing the empirical cumulative distribution function (ECDF), considered in [3]. The multidimensional divide-and-conquer serial algorithm given in [3] parallelizes easily to solve the problem in \( O(\log^2 N) \) time using \( N \) PEs. Atallah et al. improved on this, lowering the time to \( O(\log N) \) using \( N \) PEs.

• Closest Larger Right Neighbor (CLRN) Problem: Given input numbers \( a_1, \ldots, a_N \), find, for each \( a_i \), the index of the first larger number to the right; i.e., for each \( i = 1, \ldots, N-1 \), compute \( b_i = \min\{j > i : x_j > x_i\} \) if there is some \( j > i \) with \( x_j > x_i \), \( b_i = N + 1 \) otherwise. The CLRN problem can be reduced to trapezoidal decomposition [18]: given a set of line segments and points, from each point, report the line segment first hit (if any) by a ray shot horizontally to the right. To make the reduction, consider the polygonal path connecting consecutive points \((i, a_i), i = 1, \ldots, N\). If \( a_{i+1} > a_i \) then we know \( b_i = a_{i+1} \). Otherwise, \( b_i \) is the height of the right end point \( a_j \) of the segment from \((j-1, a_{j-1})\) to \((j, a_j)\) first hit by the ray shot horizontally to the right from \((i, a_i)\), if there is an \( a_j > a_i, j > i \). If not then \( b_i = N + 1 \). Reif and Sen give a probabilistic algorithm for trapezoidal decomposition. Applying that algorithm to the CLRN problems yields its solution in \( \tilde{O}(\log N) \) time using \( N \) PES. Alternatively, the CLRN problem can be solved by a binary search-like algorithm, given in Section 5, in \( O(\log^2 N) \) time using \( N \) PEs.

• Parallel Prefix (scan, segmented scan): Given inputs \( a_1, \ldots, a_N \) and an associative operator \( o \), compute the partial products \( p_1, \ldots, p_N \) where \( p_i = a_1 \circ a_2 \circ \ldots \circ a_i \). Solutions to this parallel prefix problem [13] are commonly called scan computations. The problem can be solved in \( O(\log N) \) time using \( N/\log N \) PEs [12].

A variation breaks the products over the indices \([1, N]\) into segments over these indices, with the segment boundaries also given as inputs. For example, an additional vector \( b_1, \ldots, b_N \), is given where \( b_1 = 0 \), for \( i > 1 \), \( b_i \) is either 0 or 1, and the 0's mark the segments' left boundaries. Specifically, if \( b_i = 0 \) then \( p_i = a_i \); otherwise, \( p_i = a_j \circ a_{j+1} \circ \ldots \circ a_i \) where \( j \) is the largest index \( k, 1 < k < i \), such that \( b_k = 0 \). The segmented problem has the same complexity as the original. In the algorithms below, we use copy-scans defined by \( \alpha \circ \beta = \alpha \), and add-scans where \( \circ \) is addition.

None of the algorithms listed above requires more than \( O(\log N) \) space per PE.

3 Fast Parallel LRU Simulation

In this section we present a fast parallel algorithm for computing stack distances under the LRU replacement policy.
LRU may be characterized as follows. Reference to \( \alpha = x_t \) places \( \alpha \) at the first level of the stack. Until \( \alpha \) is referenced again, it can only move down in the stack. Specifically, after \( \alpha \) has been pushed to level \( i \) it remains there until a reference is made either to \( \alpha \) (moving \( \alpha \) to level 1) or to a line not stored in levels 1 through \( i - 1 \) (moving \( \alpha \) to level \( i + 1 \)). As a result, the stack distance \( \Delta_i \) is one greater than the number of distinct lines in the subtrace between \( t \) and the closest prior reference to \( \alpha \) (or \( \infty \) if there is no prior reference to \( \alpha \)). For example, in Figure 1, consider the consecutive references to line \( b \) at \( t = 3 \) and \( t = 10 \). The stack distance \( \Delta_{10} = 4 \) because 3 distinct symbols belong to the subtrace \( x_4, \ldots, x_9 \). More generally, letting

\[
\text{prev}(t) = \begin{cases} 
\max\{s < t : \alpha = x_t\} & \text{if } \alpha = x_t \text{ for some } s < t \\
0 & \text{otherwise}
\end{cases}
\]

we obtain

\[
\Delta_t = \begin{cases} 
1 + \text{number of distinct symbols in } x_{\text{prev}(t)+1}, \ldots, x_{t-1} & \text{if } \text{prev}(t) > 0 \\
\infty & \text{otherwise}
\end{cases}
\]

Let us take a geometric view of this new problem of counting distinct symbols within subtraces. As illustrated in Figure 2, identify each reference \( x_t \) with the point \((t, \text{next}(t))\), where

\[
\text{next}(t) = \begin{cases} 
\max\{s > t : \alpha = x_t\} & \text{if } \alpha = x_t \text{ for some } s > t \\
N + 1 & \text{otherwise}
\end{cases}
\]

Note that the last references to symbols within the subtrace \( x_{\text{prev}(t)+1}, \ldots, x_{t-1} \) are identified by those points \((s, \text{next}(s))\) satisfying

\[
\text{prev}(t) < s < t < \text{next}(s).
\]

These are the points that lie strictly within the rectangle with lower left hand corner \((\text{prev}(t), t)\), lower right hand corner \((t, t)\) and sides extending upwards to \((\text{prev}(t), N + 1)\) and \((t, N + 1)\). Again, see Figure 2. Counting these points reduces to 2d-ranking. Specifically, suppose we know the 2d-rank, \( \text{rank}(u, v) \), of each point \((u, v)\) in the union of sets \( \{(t, \text{next}(t)) : 1 \leq t \leq N\} \) and \( \{(t, t) : 1 \leq t \leq N\} \). Then, the stack distance

\[
\Delta_t = \begin{cases} 
\text{rank}(\text{prev}(t), t) - \text{rank}(t, t) & \text{if } \text{prev}(t) > 0 \\
\infty & \text{otherwise}
\end{cases}
\]

We see from Figure 2 that \( \Delta_{10} = 4 \) because \( \text{rank}(3, 10) = 20 \) and \( \text{rank}(10, 10) = 16 \).

Now, let us present the detailed simulation method. Suppose that the trace is initially stored in the \( N \)-vector \( x \). We use the additional \( N \)-vectors \( p, \text{next}, \text{prev}, \) and \( \Delta \). Initially, let \( p_t = t \), so \( x_t \) identifies the line and \( p_t \) the trace index of reference \( x_t \). Vectors \( \text{next}, \text{prev}, \) and \( \Delta \) will hold permuted copies of the vectors \( \text{next}, \text{prev}, \) and \( \Delta \), respectively. The algorithm is as follows.

1. [Compute next and prev.] Sort the tuples \((x_t, p_t)\) using \( x_t \) as the primary key and \( p_t \) as the secondary key: \((x_t, p_t) < (x_s, p_s)\) if either \( x_t < x_s \) or \( x_t = x_s \) and \( p_t < p_s \). Thus, the data now in location \( t \) of \( x \) and \( p \) was in location \( p_{t-1} \) before the sort. For all \( t = 1, \ldots, N \), set

\[
\text{prev}_t = \begin{cases} 
p_{t-1} & \text{if } t > 1 \text{ and } x_{t-1} = x_t \\
0 & \text{otherwise}
\end{cases}
\]
Figure 2: The trace of Figure 1 is repeated, along with corresponding next and prev values. The values \((t, \text{next}(t))\) are plotted as +’s, and the values \((t, t)\) as o’s. The number of points strictly within the rectangle indicated by dashed lines is one less than the stack distance of \(x_{10} = b\).

\[
\text{next}_t = \begin{cases} 
  p_{t+1} & \text{if } t < N \text{ and } x_{t+1} = x_t \\
  N + 1 & \text{otherwise}
\end{cases}
\]

At this point, the prev and next vectors hold permuted copies of the prev and next vectors discussed above.

2. [2d-rank.] Compute the 2d-ranks of the set of points

\[\{(p_t, \text{next}_t) : 1 \leq t \leq N\} \cup \{(t, t) : 1 \leq t \leq N\}\]
and set

\[ \Delta_t = \text{rank}(\text{prev}_t, p_t) - \text{rank}(p_t, p_t). \]

As a result, \( \Delta_t = \Delta_{p_t} \), which completes the computation.

Sorting within the first step costs \( O(\log N) \) time on \( N \) PEs, using the EREW model [5]. The \textit{next} and \textit{prev} computations may be done within the same time and processor bounds using segmented copy-scans, with changes in the \( x \) vector marking the segment boundaries. The 2d-ranking within the second step costs \( O(\log N) \) time on \( N \) PEs [1]. Thus:

**Theorem 1** On the EREW model, given the trace \( x_1, \ldots, x_N \), the associated stack distances \( \Delta_1, \ldots, \Delta_N \) induced under the LRU replacement policy can be computed in \( O(\log N) \) time using \( N \) PEs.

Aiming for a simpler implementation and smaller implicit constants, we may sacrifice a \( \log N \) factor in the running time. The natural parallelization of Bentley's multidimensional divide and conquer method [3] gives a 2d-ranking algorithm that runs in \( O(\log^2 N) \) time using \( N \) PEs. Using, for example, Batcher's sorting method [2] requires time \( O(\log^2 N) \) on \( N \) PEs.

### 4 Parallel Simulation of LRU Level by Level

An alternative approach is to simulate LRU level by level, at the \( i^{th} \) iteration computing the level \( i \) cache contents \( s_1(i), \ldots, s_N(i) \) and stack distances \( \Delta_1(i), \ldots, \Delta_N(i) \). Assuming a set size of \( C \), the final results are the stack distances \( \Delta_1(C), \ldots, \Delta_N(C) \).

Define reference \( x_t \) to be a prior hit (prior miss) at level \( i \) if \( x_t \) is a hit (miss) given that the cache size is \( i - 1 \). That is, \( x_t \) is a prior miss at level \( i \) if \( x_t \notin B_{i-1}(i - 1) \). If \( x_t \) is a prior hit at level \( i \) then \( \Delta_t(i - 1) < i \); otherwise \( \Delta_t(i) = \infty \). In Figure 3, we have marked the prior hits \( x_t \) at level 3 by underscoring the symbol at level 2 in column \( t \). In studying this figure one should remember that an underscore on symbol \( s_t(i) \) means that symbol \( x_t \) was a hit in a \((i - 1)\)-line cache, not that \( s_t(i) \) was. The placement of underscores was chosen to highlight the propagation of a symbol across a sequence of prior hit positions, to be described below. For example, of the first ten references four are prior hits at level 3—\( x_4, x_5, x_7, \) and \( x_8 \)—because \( c = x_4, x_5, x_7 \) is found in \( B_3(2), B_4(2) \) and \( B_6(2) \), and symbol \( a(x_8) \) is found in \( B_7(2) \).

Any prior hit at level \( i - 1 \) is also a prior hit at level \( i \) (for example, \( x_5 \) in Figure 3). Under LRU, the other prior hits at level \( i \) are the references \( x_t \) satisfying \( x_t = s_{i-1}(i - 1) \); i.e., the references that hit at the last level of the size \( i - 1 \) cache (for example, \( x_4 \) in Figure 3).

The key to the simulation method is that under LRU, for all \( t \geq 1 \) and \( i > 1 \),

\[
 s_t(i) = \begin{cases} 
 s_{i-1}(i - 1) & \text{if } x_t \text{ is a prior miss at level } i \\
 s_{i-1}(i) & \text{otherwise} 
\end{cases}
\]

(2)

where

\[
 s_t(1) \equiv x_t, s_0(i) \equiv \emptyset.
\]
Table 1

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<td>(\infty)</td>
<td>(\infty)</td>
<td>(\infty)</td>
<td>2</td>
<td>1</td>
<td>3</td>
<td>2</td>
<td>2</td>
<td>(\infty)</td>
<td>(\infty)</td>
<td>1</td>
<td>2</td>
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<td>2</td>
<td>(\infty)</td>
<td>1</td>
<td>(\infty)</td>
<td>2</td>
</tr>
</tbody>
</table>

Figure 3: LRU acting on an 18 line trace, assuming a cache size \(C = 3\). Each prior hit \(x_t\) at level 3 is identified by underscoring \(s_t(2)\).

To see this, suppose \(x_t \notin B_{t-1}(i-1)\). The LRU rule puts \(x_t\) into level one, and shifts lines 1, 2, ..., \(i-1\) down one level, which pushes \(s_{t-1}(i-1)\) to level \(i\). On the other hand, if \(x_t\) is a prior hit at level \(i\), the cache update leaves level \(i\) unchanged. In Figure 3, we see that the 3\(^{rd}\) and the 6\(^{th}\) references are prior misses at level 3, and that the intervening references are prior hits. As a result, \(s_2(2) = a\) enters level 3 at \(t = 3\) and propagates over prior hits at level 3 until \(t = 6\), where it is replaced with \(s_6(2) = b\), which in turn propagates up through \(t = 8\).

We now describe the simulation algorithm, taking special care with the details because similar methods are needed in Section 5. At the \((i-1)^{st}\) iteration we will overwrite vectors \(s = (s_0, s_1, \ldots, s_N)\) and \(d = (d_1, d_2, \ldots, d_N)\) with the level \(i\) cache contents and stack distances, \((s_0(i), s_1(i), \ldots, s_N(i))\) and \((\Delta_1(i), \Delta_2(i), \ldots, \Delta_N(i))\), respectively. A vector \(x = (x_1, x_2, \ldots, x_N)\) holds the trace \((x_1, x_2, \ldots, x_N)\), and another vector \(u = (u_1, \ldots, u_N)\) will hold a copy of \((s_0(i-1), s_1(i-1), \ldots, s_{N-1}(i-1))\). To initialize the computation, for \(t = 1, \ldots, N\), set \(d_t = \infty\), and \(s_t = x_t\), \(s_0 = \emptyset\). For \(i = 1, \ldots, C\), do as follows.

1. [Update the Level \(i\) Cache Contents via equation (2).] For \(t = 1, \ldots, N\), set \(u_t = s_{t-1}\). For \(t = 1, \ldots, N\), if \(d_t \neq \infty\) then set \(s_t = s_{t-1}\); otherwise, \(s_t = u_t\). This is to be understood, but not implemented, as a serial update: first \(s_1\) is updated, then \(s_2\), and so forth.

2. [Update the Stack Distances.] For \(t = 1, \ldots, N\), set \(d_t = i\) if \(d_t = \infty\) and \(u_t = x_t\); otherwise leave \(d_t\) unchanged.

The right shift of \(s\) into \(u\) in step 1 and the update to the stack distances in step 2 are naturally parallel operations. The update of \(s\) is a segmented copy-scan, with the coordinates \(t\) with \(d_t = \infty\) marking the segment boundaries. Hence, the cost of both steps is just \(O(\log N)\) time using \(N/\log N\) PEs. As there are a total of \(C\) iterations to perform, we obtain:

\textbf{Theorem 2} On the EREW model, given the trace \(x_1, \ldots, x_N\), the associated level \(C\) stack distances \(\Delta_1(C), \ldots, \Delta_N(C)\) under the LRU replacement policy can be computed in \(O(C \log N)\) time using \(N/\log N\) processors.

We implemented this algorithm on a MasPar MP-1 computer [4], with 16384 PEs. Each PE is a 4-bit processor with a clock cycle of 80 nanoseconds. A typical integer operation such as those common in our
algorithm requires a few ten's of clocks.

Our implementation supports "super-saturation" of the PE's, as described earlier. The PE memory size permit us to assign as many as 2048 references to each PE, thereby permitting the simultaneous simulation of a trace with over 33 million references. The performance data we present includes only the time spent in the solution phase of the algorithm. The traces were generated randomly. For a given trace length and cache set size we observed a 10-15% increase in running time between caches with a very low hit ratio, and caches with a high hit ratio. This is likely due to the fact that long segments accompany high hit ratios, requiring greater inter-PE communication to implement the copy-scan. The timings presented are from traces with nearly perfect hit ratios, and so represent an upper bound on the timing one might expect from an actual trace.

A full implementation would have to spend time loading the trace; the I/O time required depends on the available I/O hardware and the organization of the trace on the I/O devices. In light of our timings, it is clear that moving the trace onto the machine may well be the most serious bottleneck an actual implementation would face.

Our experiments vary the length of the trace from $2^{14}$ to $2^{25}$, and the set size $C$ from 2 to 32. The presented timings are averages, given in milliseconds, taken by executing the solution loop many times in succession.

Observe that about five seconds of execution time were required to analyze the behavior of a 32-line set on a trace with $2^{25} = 33,554,432$ references. This is 710 times faster than the solution time (with trace generation costs subtracted off) of an optimized serial algorithm we implemented on a Sparc-1+ workstation. These timings demonstrate the remarkable promise of massive parallelism for trace-driven cache simulation.

### 5 Reference Based Replacement Policies

We now broaden the scope of our methods, to handle a large class of line replacement policies, which we term reference-based. In Section 5.3, we extend the class to handle policies that allow priorities associated with cached lines to "age" so that stale lines tend to be flushed.

```
<table>
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<th>2^{16}</th>
<th>2^{17}</th>
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<th>2^{22}</th>
<th>2^{23}</th>
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<td>5.0</td>
<td>7.3</td>
<td>11.4</td>
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<td>139</td>
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<td>8.4</td>
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<td>12</td>
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<td>660</td>
<td>1286</td>
<td>2539</td>
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</tr>
</tbody>
</table>
```

Table 1: Execution time of the LRU algorithm on a MasPar MP-1 with $2^{14}$ PEs, in milliseconds, as a function of trace length and set size.
Mattson et al. [16] show that a stack policy is obtained if a numerical priority $P(s_t(i))$ is assigned to each line $s_t(i)$ at reference $x_t$, and the line $y_t(C)$ chosen for replacement on loading $x_t$ is the one with least priority among the members of $B_{t-1}(C)$. In the class of policies we now consider, a line's priority is established at the point it appears in the reference stream, after which it remains constant until the line is referenced again.

Of course we must be able to calculate the priorities from the reference trace. Thus we limit attention to policies that support efficient parallel priority calculations. As practical policies seem to use very simple priority assignments, this limitation is mild. Here, "efficient" means within the resource bounds needed for the rest of our simulation method: $O(\log N)$ time using $N$ processors. Recall (Section 2.2) that $O(\log N)$ means $O(\log N)$ with high probability.

Let us define the class of reference-based replacement policies as those stack policies induced by priorities satisfying the following conditions.

**R1:** All $P(x_t)$ values can be computed quickly in parallel: in $O(\log N)$ time using $N$ PE’s. For example, the priorities for LFU (Least Frequently Used) can be established with a sort on the reference tags, followed by a segmented sum-scan.

**R2:** A line’s replacement priority does not change except when the line appears in the reference stream.

Several important replacement policies are reference-based, including

- **LRU:** $P(x_t) = t$.
- **LFU:** $P(x_t) = \text{Count}(x_t, t)$, the number of references $x_u = x_t$ for $u \leq t$. Ties can be broken, for example, by lexicographic ordering of the lines, or by giving higher priority to the line that has been in the cache the shortest length of time. ($P(x_t) = \text{Count}(x_t, t) - 1/(t + 1)$ would serve the latter purpose.)
- **OPT:** $P(x_t)$ is the negation of the smallest index $u > t$ such that $x_u = x_t$.

In addition, the Random replacement (RR) policy shares most of the properties we need to quickly simulate reference-based policies, and we include it in this class as a special case. Under RR, priorities are chosen that determine a uniform random ranking of the cache contents; details are given below.

Figure 4 gives an example of the operation of LFU with ties broken by lexicographic ordering, $a < b < c < d$. A line's subscript equals the number of earlier references to the line. First, note that the stack order and the priority order may differ. A line with low priority can be buried in the middle of the stack order; for example, line $d$ at $t = 13$. There are important departures from the behavior of the LRU policy. Under LRU, the replaced line is the one at the lowest stack level. Here, we see that if the cache size is 2 then at $t = 9$, line $d$ misses and replaces line $a$ at the first stack level, leaving line $c$ in place at the second stack level. To illustrate the entry and propagation of lines across a given level, each prior miss $x_t$ at level 3 is marked by underscoring $s_t(2)$. As in LRU, a line propagates across all prior hits. Unlike LRU, a line may propagate across some prior misses. For example, line $a$ enters level 3 at $t = 9$ and propagates until $t = 15$, across the prior misses at $t = 10$ and $t = 12$. 

11
### 5.1 Parallel Simulation Level by Level

In this section, we present a rapid parallel simulation algorithm for any reference-based replacement policy. For any given $C > 0$, the objective is to compute the level $C$ stack distances $\Delta_1(C), \ldots, \Delta_N(C)$. The algorithm works level by level, like the LRU simulation algorithm presented in Section 4. Specifically, we compute the cache contents $s_t(i)$, the replacees $y_t(i)$, and the stack distances $\Delta_t(i)$ at level $i$, given these same quantities at level $i - 1$. As in the LRU simulation, just $O(1)$ space per reference is needed, with the results of the level $i$ computation overwriting those for level $i - 1$.

Level 1 is easy: $s_t(1) \equiv x_t$, $y_t(1) \equiv s_{t-1}(1)$, $\Delta_t(1) = 1$ if $s_t(1) = s_{t-1}(1)$, and $\Delta_t(1) = \infty$ otherwise. Two facts, which follow from equation (3), are crucial to our approach for computing the desired results for level $i > 1$:

Notice that the only lines that ever enter level $i$ are the least priority lines $y_{t-1}(i)$ from lower levels.

#### Table 4: LFU; ties are broken by $a < b < c < d$. The subscripts count the number of earlier references, and underscores mark prior hits at level 3. Also shown are the stack distances $\Delta_t(2)$ at level 2 and the lowest priority lines $y_t(2) \in B_{t-1}(2)$ at level 2.

| $t$ | 0 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | 11 | 12 | 13 | 14 | 15 | 16 | 17 | 18 |
|-----|---|---|---|---|---|---|---|---|---|---|----|----|----|----|----|----|----|----|
| $s_t(1)$ | \emptyset | a | c | b | c | c | a | c | a | d | b | b | d | c | d | a | a | b | a |
| $s_t(2)$ | \emptyset | \emptyset | a_0 | c_0 | b_0 | c_0 | a_1 | c_0 | b_1 | d_0 | b_1 | d_0 | c_1 | d_0 | c_2 | d_0 | c_3 | d_2 | a_3 | c_4 | d_2 | a_4 | b_3 | c_5 |
| $s_t(3)$ | \emptyset | \emptyset | a_0 | a_0 | a_0 | b_0 | a_0 | b_0 | a_0 | b_0 | b_0 | a_2 | a_2 | a_2 | a_2 | a_2 | a_2 | a_2 | a_2 | a_2 | a_2 | a_2 | a_2 | a_2 | a_2 | a_2 | a_2 | a_2 |
| $s_t(4)$ | \emptyset | \emptyset | \emptyset | \emptyset | \emptyset | \emptyset | \emptyset | \emptyset | \emptyset | \emptyset | \emptyset | \emptyset | \emptyset | \emptyset | \emptyset | \emptyset | \emptyset | \emptyset | \emptyset | \emptyset | \emptyset | \emptyset | \emptyset | \emptyset | \emptyset | \emptyset | \emptyset | \emptyset |
| $\Delta_t(2)$ | $\infty$ | $\infty$ | $\infty$ | $\infty$ | 2 | 1 | $\infty$ | 2 | 2 | $\infty$ | $\infty$ | 1 | $\infty$ | 2 | 2 | $\infty$ | 1 | $\infty$ | $\infty$ | $\infty$ | $\infty$ | 2 | 1 | $\infty$ | 2 | 2 | $\infty$ | 1 | $\infty$ | $\infty$ |
| $y_t(2)$ | \emptyset | \emptyset | a | b | b | a | a | a | d | b | b | d | b | d | d | d | d | d | d | d | d | d | d | d | d | d | d | d | d | d | d | d | d | d | d | d |
If a new line, say $\alpha$, enters level $i > 1$ at time $t$ (meaning $s_t(i) = \alpha$, $s_{t-1}(i) \neq \alpha$) then $x_t$ must be a prior miss at level $i$ and $\alpha$ must be the replacee $y_t(i-1)$.

Assuming $\alpha = y_t(i-1)$ does enter level $i > 1$, it propagates until coming to the first reference $x_u$, $u > t$, where either $x_u = \alpha$ or $x_u$ is a prior miss and $P(y_t(i-1)) < P(y_u(i-1))$. That is, $s_t(i) = \ldots = s_{u-1}(i) = \alpha$. If $u < N + 1$ then replacee $y_u(i-1)$ enters level $i$ at time $u$. If there is no such $u < N + 1$ then the replacee $y_t(i-1)$ propagates on through time $N$. For every reference $x_t$ let $u(t)$ denote the index so identified.

Consider the graph where the vertices are the indices $t$ of all prior misses $x_t$ and there is an edge from $t$ to $u(t)$. This edge records the fact that if $y_t(i-1)$ enters level $i$ then $s_t(i) = \ldots = s_{u(t)-1}(i) = y_t(i-1)$, whereupon it is replaced by $y_{u(t)}(i-1)$ (if $u(t) < N + 1$). Observe that not all such $y_t(i-1)$ actually do enter level $i$—for instance, in Figure 4, $y_{10}(2) = d$ does not enter level 3, because $P(y_{10}(2)) < P(s_0(3))$. However, the set of references that do actually enter level $i$ can be determined by following the maximal path through the graph, starting at vertex 1. Replacee $y_t(i) = \emptyset$ enters at time 1 and propagates until time $v = u(1) - 1$. If $v < N + 1$ then replacee $y_{v+1}(i-1)$ enters level $i$, and propagates until time $w = u(v) - 1$. If $w < N + 1$ then replacee $y_{w+1}(i-1)$ enters level $i$, and so forth.

Converting this serial process for simulating level $i$ into a parallel one, we simulate level $i > 1$ as follows:

1. [Compute tentative propagation intervals.] For each prior miss $x_t$, compute

$$
\begin{align*}
\text{stop}(t) &= \text{least } s > t \text{ such that } P(x_s) > P(x_t), \text{ and } x_s \text{ is a prior miss at level } i \\
\text{next}(t) &= \text{least } s > t \text{ such that } x_s = x_t,
\end{align*}
$$

where, by convention, $\text{stop}(t)$ and $\text{next}(t)$ equal $N + 1$ if the index $s$ above does not exist. Set $p(t) = \min\{\text{next}(t), \text{stop}(t)\}$. By earlier remarks, if $y_t(i-1)$ enters level $i$ then $s_t(i) = \ldots = s_{p(t)-1}(i) = y_t(i-1)$.

2. [Follow propagation chain.] The pointers $u(t)$ determine the replacees that enter level $i$, namely, those with indices: $0$, $u(0)$, $u(u(0))$, ..., with the sequence stopping at $v = u(\ldots u(0)\ldots) \neq N + 1$, $u(v) = N + 1$. Mark these replacees.

3. [Compute level $i$ results.] For each marked replacee $y_t(i-1)$ and each $v$ in the interval $[t, u(t) - 1]$, set $s_v(i) = y_t(i-1)$. For every $x_t$ set $y_{t+1}(i)$ to $\max\{s_t(i), y_{t+1}(i-1)\}$. Following this, if $x_t$ is a prior miss at level $i-1$ and if $s_t(i) = x_t$ set $\Delta_t(i) = i$. Set $\Delta_t(i) = \Delta_t(i-1)$ for prior hits.

Computing the $\text{stop}(t)$ values is an instance of the closest larger right neighbor (CLRN) problem, discussed in Section 2.2. It can be solved in $\tilde{O}(\log N)$ time using $N$ PEs. A simpler $O(\log^2 N)$ time solution is described below. Computing the $\text{next}(t)$ values via sorting is described in Section 3; the time needed is $O(\log N)$ using $N$ PEs. Marking the replacees on the chain of pointers from 0 to $N + 1$ is a pointer jumping problem [14], which can be solved in $O(\log N)$ time using $N/\log N$ PEs. The final step of updating the level $i$ cache...
contents and stack distances is essentially the same as was done for LRU simulation in Section 4. We do not repeat the details. The time needed is $O(\log N)$ time using $N/\log N$ PEs. Summing up, the total time is $\tilde{O}(\log N)$ using $N$ PEs. To produce the level $C$ results, the computation must be repeated for each $i = 2, \ldots, C$. Thus,

**Theorem 3** On the EREW model, given the trace $x_1, \ldots, x_N$, the associated level $C$ stack distances $\Delta_1(C), \ldots, \Delta_N(C)$ induced by any reference-based replacement policy can be computed in time $\tilde{O}(C \log N)$ using $N$ PEs.

We close this section with a simple method for solving the CLRN problem. This method plays the key role in generalizing the simulation method to accommodate priority aging (Section 5.3).

Let $a_1, \ldots, a_N$ be a sequence of $N$ numbers, and, for simplicity, assume that $N$ is a power of two and $a_N = +\infty$. For each $a_i$, $i = 1, \ldots, N - 1$, we wish to find the closest right larger neighbor $a_j$; i.e., $j > i$ is as small as possible and $a_i < a_j$. Construct a binary tree over the inputs as illustrated in Figure 5. Each node is labeled with the maximum value of the inputs in its subtree and with the corresponding subrange of indices. To find the closest larger right neighbor $a_j$ of $a_i$ a two phase search is initiated. Phase one starts at the leaf node $a_i$, and progresses in steps up the tree. At each step we move from the present node to the nearest internal node at the next higher level whose span includes a node to the right. This phase ends upon visiting (i) an internal node which is rightmost at its level, or (ii) a node whose value is (strictly) greater than $a_i$. In the example of Figure 5, the first phase for $a_3$ visits nodes representing ranges $[3, 4]$ and $[5, 8]$. In general, the first phase stops at a node spanning an interval $[m + 2^k, m + 2^{k+1}]$, with $i \leq m + 2^k$, which must contain the sought $a_j$. In phase two the search descends down to $a_j$, at each step moving to the left child if the left child's value exceeds $a_i$, or to the right child otherwise. On a concurrent read model, carrying out all $N$ searches in parallel gives an $O(\log N)$ time solution. On an exclusive read model, standard methods [14] for resolving the read conflicts add an $O(\log N)$ factor, bringing the total time to $O(\log^2 N)$. 

![Figure 5: Search tree identifying maximum values over subintervals, which is used to solve the nearest right neighbor problem.](image)
5.2 Random Replacement

The Random Replacement (RR) rule selects the line to replace on a cache miss independently and uniformly at random from the set of cached lines. Mattson et al. [16] observed that the selections can be made so as preserve the stack property \((B_t(i - 1) \subseteq B_t(i))\) for all \(t, i \geq 1\), by coupling the random decisions as follows. Suppose the members of \(B_t(i - 1)\) have been ranked randomly from 1 to \(i - 1\), with the understanding that the higher a line’s rank the lower its priority. To build \(B_t(i)\), insert \(s_t(i)\) into the priority structure by randomly choosing an integer rank for it from \([1, i]\), say \(k\). Members of \(B_t(i - 1)\) with ranks \(\geq k\) have their ranks incremented by one in \(B_t(i - 1)\). Other members of \(B_t(i - 1)\) retain their rank. Thus, for each prior miss \(x_t\) at level \(i\), we may decide the line \(y_t(i)\) of least rank in \(B_t(i)\) by a coin toss: with probability \(1/i\), line \(y_t(i) = s_{t-1}(i)\), and with the complementary probability \(y_t(i) = y_t(i - 1)\). Moreover, the outcome is completely independent of \(s_{t-1}(i)\).

This independence can be exploited to considerably simplify the simulation of level \(i\) over that described above. Step 1 becomes: For each prior miss \(x_t\), compute \(next(t)\) as before, and use a coin toss to decide whether to label index \(t\) as a “stopper” (probability \(1/i\)) or leave the index unlabeled (probability \(1 - 1/i\)). Let \(stop(t)\) be the least stopper \(u > t\), or \(N + 1\) if no such \(u\) exists. Let \(u(t) = \min\{stop(t), next(t)\}\) as before. Steps 2 and 3 remain the same.

Computing the \(stop(t)\) values entails \(N\) independent coin tosses and a segmented copy-scan (cf. Section 2.2), operations that net \(O(\log N)\) time using \(N\) PEs. As a result, we obtain

**Theorem 4** On the EREW model, given the trace \(x_1, \ldots, x_N\), the associated level \(C\) stack distances \(\Delta_1(C), \ldots, \Delta_N(C)\) induced by the RR policy can be computed in time \(O(C \log N)\) using \(N\) PEs.

Comparing with Theorem 3 we see that dropping the CLRN problem and the probabilistic algorithm used to solve it strengthens the running time bound from one that holds with high probability to one that holds deterministically.

5.3 Priority Aging

Under any reference-based replacement rule other than RR, a line’s priority is fixed when it enters the cache. If the policy is a practical one, then it is likely that the priority is a simple function of the previous references. For example, under LFU a line’s priority is the number of earlier references to the line. Past cache activity gives an imperfect indication of future cache activity. Under LFU a flurry of references to a small set of lines might lead to their long retention during a subsequent period when the lines are not needed. To counter this, it is natural to consider policies that allow a line’s priority to age; i.e., to decrease monotonically while the line remains unreferenced. In this Section, we extend our reference-based simulation method to accommodate aging.

Let \(\phi : \mathbb{R} \rightarrow \mathbb{R}\) be a monotonically decreasing operator, and let \(\phi^d\) represent the \(d\)-fold application of \(\phi\). Let \(P_t(\alpha)\) denote the priority of a line \(\alpha\) held in the cache at the time of \(x_t\). We consider replacement
policies where the initial priority of line \( x_t \) is reference-based \((P_t(x_t) \text{ satisfies } R1 \text{ of Section 5})\), but the line’s priority “ages” to \( P_{t+d}(x_t) = \phi^d(P_t(x_t)) \) in the cache \( B_{t+d}(C) \) if it remains unreferenced throughout time \( t + 1, \ldots, t + d \). As before, the replacement policy always selects the line with least priority. Some natural \textit{aging operators} \( \phi \) are \( \phi(x) = x - \alpha \) for some fixed \( \alpha > 0 \) or \( \phi(x) = \alpha x \) for some fixed \( \alpha \in (0, 1) \). We assume that for any \( d = 1, \ldots, N - 1 \), \( \phi^d(x) \) can be computed in \( O(1) \) time.

Equation (3) describing the evolution of the stack levels continues to hold. A little thought shows that to adapt the simulation method to accommodate aging, we need only change the definition of \( \text{stop}(t) \) in equation (4) to

\[
\text{stop}(t) = \text{least } s > t \text{ such that } P_s(x_s) > \phi^{s-t}(P_t(x_t)), \text{ and } x_s \text{ is a prior miss at level } i.
\]

Computing these new values \( \text{stop}(t) \) can be posed as the following variant of the CLRN problem. Let \( a_1, \ldots, a_N \) be a sequence of \( N \) numbers, and, for simplicity, assume that \( N \) is a power of two and \( a_N = +\infty \). For each \( i = 1, \ldots, N - 1 \), we wish to find the smallest \( j > i \) such that

\[
\phi^{j-i}(a_i) < a_j.
\]

We now sketch how to extend the binary search solution given in Section 5 to solve the new problem. Let \( \hat{\phi} \) denote the inverse of \( \phi \). Since \( \hat{\phi} \) is monotone increasing, the inequality above implies to

\[
\phi^u(a_i) < \hat{\phi}^v(a_j) \quad \text{for all nonnegative integers } u, v \text{ with } u + v = j - i.
\]

Letting \([u, v]\) be any range of indices with \( u \geq i \),

\[
\phi^{j-i}(a_i) < a_j \quad \text{for all } j \in [u, v] \iff \phi^{u-i}(a_i) < \max\{a_u, \hat{\phi}(a_{u+1}), \ldots, \hat{\phi}^{v-u}(a_v)\}.
\]

This equivalence reveals a way to determine whether \( a_i \) ages below some \( a_j \), \( j \in [u, v] \), and \( i < u \); i.e., whether \( \phi^{j-i}(a_i) < a_j \). For \( j \geq u \) define \( \hat{\phi}^{j-u}(a_j) \) to be the “rejuvenated” value of \( a_j \) with respect to index \( u \) (i.e., \( a_j \)'s priority if “de-aged” back to position \( u \)). Let \( r_{u,v} = \max\{a_u, \hat{\phi}(a_{u+1}), \ldots, \hat{\phi}^{v-u}(a_v)\} \) denote the maximum (over \( j \in [u, v] \)) rejuvenated value of any \( a_j \) with respect to \( u \). To find out if \( a_i \) ages below some \( a_j \) with \( j \in [u, v] \), we may simply compare \( \phi^{u-i}(a_i) \) and \( r_{u,v} \). Since \( a_i \) is arbitrary in this discussion, it is possible to use \( r_{u,v} \) concurrently in many searches.

A “rejuvenation-max” tree can be built in parallel using the following observation: for any \( M \) (assumed to be a power of 2)

\[
\max_{1 \leq i \leq M} \{\phi^{i-1}(a_i)\} = \max\{\max_{1 \leq i \leq M/2} \{\hat{\phi}^{i-1}(a_i)\}, \max_{M/2+1 \leq i \leq M} \{\phi^{i-1}(a_i)\}\}
\]

\[
= \max\{\max_{1 \leq i \leq M/2} \{\phi^{i-1}(a_i)\}, \hat{\phi}^{M/2} \max_{1 \leq i \leq M/2} \{\phi^{i-1}(a_{M/2+i})\}\}
\]

This recursion shows we can build a rejuvenation-max tree over \( a_1, \ldots, a_N \) in \( \log N \) steps. At every step, all nodes at a given level of the tree are constructed. Leaves are understood to be at level \( \log N \), the root is at level 0. A node spanning an interval \([u, v]\) is labeled with \( r_{u,v} \). The recursion shows that to compute
the label of a node at level \(k\), one computes the maximum of (i) the label on the node’s left-child, and (ii) the label on the node’s right-child promoted by an operator \(\hat{\phi}\), with \(c = 2^{\log N - k}\). Thus, the parent of a left node with value \(a\) and right node with value \(b\) has label \(\max\{a, \hat{\phi}(b)\}\). Figure 6 illustrates how the tree of Figure 5 is modified to accommodate the aging operator \(\phi(x) = x/2\).

Given \(a_i\), we wish to find the closest \(a_j\) to the right such that \(\phi^{-1}(a_i) < a_j\). The same two phase strategy described in Section 5 works, replacing the comparison of the value \(a_i\) with the label of a node spanning an interval \([u, v]\) with the comparison of \(\phi^{-1}(a_i)\) with \(r_{u,v}\), or an equivalent comparison. For example, consider \(a_3\)’s search for the setup of Figure 5. In the first phase, the search moves up and to the right in the tree, looking over successively larger intervals for a value that \(a_3\) ages below. First, we compare \(a_3 = 6\) with \(r_{3,4} = 6\), and since \(a_3\) is not smaller continue the first phase. The next node visited represents \([5, 8]\). We compare \(\hat{\phi}(a_3) = 1.5\) with \(r_{5,8} = 32\), and as \(a_3\) is smaller, phase one stops at this node. In the second phase, the search moves down from \([5, 8]\) to locate the leftmost \(a_j\) in \([5, 8]\) that \(a_3\) ages below. First, we branch left to \([5, 6]\), because \(r_{5,6} = 14\) is larger than \(\hat{\phi}(a_3) = 1.5\). Second, we branch left again to \([5, 5]\) because \(r_{5,5} = 3 > \hat{\phi}(a_3) = 1.5\). Since \([5, 5]\) is a leaf, the search stops, having located the right match, \(a_5\), for \(a_3\).

Building the rejuvenation-max tree costs \(O(\log N)\) time using \(N\) PEs. On the CREW model, we may assign one processor to the search for each input, and so obtain an \(O(\log N)\) time solution using \(N\) PEs. This in turn implies an \(O(\log^2 N)\) time solution using \(N\) PEs on the EREW model. The final result is:

**Theorem 5** On the EREW model, given the trace \(x_1, \ldots, x_N\), the level \(C\) stack distances \(\Delta_1(C), \ldots, \Delta_N(C)\) induced by any reference-based policy with aging can be computed in time \(O(C \log^2 N)\) using \(N\) PEs.

### 6 Summary

Trace driven cache simulation is an important tool used in the design of computer systems. Parallel processing offers the promise of reducing the time required to execute a cache simulation, and hence reduce the
overall cache design time. We have shown how massively parallel SIMD architectures can be applied to this important problem area.

We note that the bottleneck problem in our simulation of reference-based replacement policies is the closest larger right neighbor problem. An improvement in the solution of this problem to $O(\log N)$ time (without using probabilistic methods) and $N$ PEs on the EREW model appears possible [7]. This would improve the reference-based simulation method to deterministic $O(C \log N)$ time using $N$ PEs.

A number of important issues remain. There is a class of "clock-based" stack algorithms which do not appear to fit within our framework. The classical clock algorithm [6] associates one bit with each physical line in the cache. The bit is set whenever a new line is written into the physical location. A clock counter determines replacement lines. On a hit the counter is untouched, but on a miss, the counter scans the set for a clear bit; the first clear bit found identifies the replacement line. The scan begins where the counter was last left, and any set bit encountered in the scan is cleared. Thus, at most one scan of the set is needed to find a clear bit. The clock algorithm is induced if we assign priority $d$ to line $r$ if $d$ lines must be scanned by the clock before choosing $r$ as the replacement. A line's priority ages as it sits in the cache, but in a highly state-dependent way. One object of our future research is to determine whether clock-based stack algorithms can be simulated in parallel.

We believe that the geometric methods used to obtain the fast, set size independent LRU simulation method of Section 3 might yield similar simulation methods for OPT and for general reference-based policies. Another important issue is whether these techniques can be extended to the simulation of multiprocessor caches. Yet another issue is the use of SIMD processors to generate synthetic cache traces. The method discussed in [20] is basically LRU "in reverse": given the stack distances, compute the reference string. We believe we can implement this method in poly-log time using ideas similar to those developed here. Given the promise of SIMD trace-driven simulation, a more comprehensive study of parallelized synthetic trace generation will be useful.

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References


Trace-driven cache simulation is central to computer design. A trace is a very long sequence, \( z_1, \ldots, z_N \), of references to lines (contiguous locations) from main memory. At the \( t \)th instant, reference \( z_t \) is hashed into a set of cache locations, the contents of which are then compared with \( z_t \). If at the \( t \)th instant \( z_t \) is not present in the cache, then it is said to be a miss, and is loaded into the cache set, possibly forcing the replacement of some other memory line, and making \( z_t \) present for the \((t+1)\)th instant. The problem of parallel simulation of a subtrace of \( N \) references directed to a \( C \) line cache set is considered, with the aim of determining which references are misses and related statistics.

A simulation method is presented for the Least-Recently-Used (LRU) policy, which regardless of the set size \( C \) runs in time \( O(\log N) \) using \( N \) processors on the exclusive read, exclusive write (EREW) parallel model. A simpler LRU simulation algorithm is given that runs in \( O(C \log N) \) time using \( N/\log N \) processors. We present timings of the second algorithm’s implementation on the MasPar MP-1, a machine with 16384 processors. A broad class of reference-based line replacement policies are considered, which includes LRU as well as the Least-Frequently-Used and Random replacement policies. A simulation method is presented for any such policy that on any trace of length \( N \) directed to a \( C \) line set runs in time \( O(C \log N) \) time with high probability using \( N \) processors on the EREW model. The algorithms are simple, have very little space overhead, and are well-suited for SIMD implementation.