On the Sublinear Processor Gap for Multi-Core Architectures
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Abstract. In the past, parallel algorithms were developed, for the most part, under the assumption that the number of processors is $\Theta(n)$ and that if in practice the actual number was smaller, this could be resolved using Brent’s Lemma to simulate the highly parallel solution on a lower-degree parallel architecture. In this paper, however, we argue that design and implementation issues of algorithms and architectures are significantly different—both in theory and in practice—between computational models with high and low degrees of parallelism.

We report an observed gap in the behavior of a CMP/parallel architecture depending on the number of processors. This gap appears repeatedly in both empirical cases, when studying practical aspects of architecture design and program implementation as well as in theoretical instances when studying the behaviour of various parallel algorithms. It separates the performance, design and analysis of systems with a sublinear number of processors and systems with linearly many processors. More specifically we observe that systems with either logarithmically many cores or with $O(n^\alpha)$ cores (with $\alpha < 1$) exhibit a qualitatively different behavior than a system with a linear number of cores on the size of the input, i.e. $\Theta(n)$. The evidence we present suggests the existence of a sharp theoretical gap between the classes of problems that can be efficiently parallelized with $o(n)$ processors and with $\Theta(n)$ processors unless $NC = P$.

1 Introduction

There is a vast experience in the study and development of algorithms for the PRAM architecture. In this case, the standard assumption (though often unstated) was that the number of processors $p$ was linear on the size of the input, i.e. $p = O(n)$ (see for example [15] for a thorough discussion). Indeed, the definition of the class NC which is often equated with the class of problems that can be efficiently parallelized in a PRAM allows for up to polynomially many processors. Hence algorithms were designed to handle the case when $p = \Theta(n)$ or $p = \Theta(n^k)$ for $k \geq 1$ and if the actual number of processors available was lower, this could readily be handled by Brent’s Lemma using a suitable scheduler [8, 5]. A fruitful theory was developed under this assumptions, and papers in which $p = o(n)$ were relatively rare.

2 Overview of arguments

Here we briefly list the arguments in favour of considering a limited degree of parallelism. We emphasize that we did not start from the outset with this goal, but rather we sought to develop algorithms and tools (both practical and theoretical) for current multi-core architectures. The observations within are derived from both theoretical investigations and practical experiences in which time and time again we found that there seems to be a qualitative difference between a model with $O(\log(n))$ processors and one with $O(n)$ processors, with, surprisingly, the advantage being for the weaker, i.e. $O(\log(n))$ model. There is strong evidence of a sublinear cliff, beyond which
development and implementation of PRAM algorithms is substantially harder if not completely impossible, unless $P = NC$. In several instances among the evidence observed the phenomenon had been observed earlier by others [15, 17, 12].

1. The number of cores is nearly a constant, but first, if it is truly a constant there is nothing we can say, and second, it seems to be steadily though slowly growing.

2. In analogous fashion to the word-PRAM the number of bits could be an arbitrary $w$ but really it is most likely $O(\log n)$ since it is also an index into memory and memory is usually polynomial on $n$.

3. The probability of collision on a memory access is only acceptably low for up to $O(\sqrt{n})$ processors.

4. The number of interconnects on a CPU network grows too fast for anything else.

5. Serialization at the network end is too costly, i.e. if more than two processors want to talk to you at the same time you have to listen to them serially.

6. There are natural $\log n$ and $n^\epsilon$ barriers in the complexity of designing algorithms.

7. Efficient cache performance requires bounded number of processors in terms of cache sizes which are always assumed to be below $n$ as well as the ratio of cache sizes which is well below 100.

8. We define the class of problems which can be sped up using a logarithmic number of processors and show that it contains $NC$ and furthermore, this containment is strict.

9. For Turing Machines we can automatically increase performance by a $\log n + \log \log n$ factor when simulating with a multi-core computer and this works for $\log n$ processors.

10. Amdahl’s law suggests that programs can only noticeably benefit from parallelism if the number of processors is proportional to the relative difference between the execution time of the serial and parallel portions of a program.

### 3 Exposition

In this section we briefly expand on each of the points above. We aim to keep each argument as short as possible, since the entirety of the case is more important than any individual point.

#### 3.1 Limited Parallelism

In principle it is possible to build a computer with an arbitrary degree of parallelism. In practice PRAMs algorithms and architectures focused on $\Theta(n)$-processor architectures, while relying on
Brent’s Lemma for cases when the number of processors was below that. In contrast CMP processors have aimed for a much smaller number of cores. In principle this number could be modeled as a constant. However this is unrealistic as the number of cores continues to grow—albeit slowly—with desktop computers having transitioned over the last decade from single core to dual core to quad core and presently eight cores with sixteen cores already shipping at the higher end of the spectrum. Additionally, it has been observed that generally speaking larger inputs justify larger investments in RAM and CPU capacity, so a function of \( n \) is much more reflective of real life constraints. This suggests that the number of cores is a function which grows slowly on the input size \( n \) since there is a high processor cost. Let \( \mathcal{P}(n) \) denote this function. Natural candidates for \( \mathcal{P}(n) \) are \( \Theta(\log n) \) and \( \Theta(n^{\alpha}) \) for \( \alpha < 1 \), though there are other possibilities. Over the next subsections we shall consider various candidates for \( \mathcal{P}(n) \).

### 3.2 Natural Constraints

In the case of word-RAMs the ability to index using a word as an address suggest that a natural value for the word size \( w \) is \( w = O(\log M) \) where \( M \) is the size memory, though this does not necessarily need to be the case.\(^1\) Memory itself is usually a polynomial function of the input size, i.e. \( M = \Theta(n^k) \) for some \( k \geq 1 \), with \( k = 1 \) being a common value. Substituting \( M = \Theta(n^k) \) in \( w = O(\log M) \) gives \( w = O(\log n) \)\(^2\) which is the usual assumption in word-RAM papers.

Hence, the word size which in the early days of computing was treated as a constant, namely 4 or 8 bits, became better understood as in fact proportional to the logarithm of the input size, that is \( O(\log n) \). Similarly, in modern multi-core computers, the number of processors has remained relatively bounded (in contrast to commercial PRAMs or GPUs which support anywhere from thousands to hundreds of thousands of processors and still growing). This relatively slow growth (at least as compared to most other usually exponential growing performance hardware indices) on the number of processors can thus be best modeled as \( \log n \) in similar fashion to the word size.

### 3.3 Write Conflicts

Consider a program running in time \( T(n) = O(n^c) \) for \( c \geq 1 \). There are two natural assumptions for the size of memory: (1) linear memory, i.e., \( M = O(n) \), which is the minimum amount to hold the input in an offline computation, and (2) \( M = O(n^c) \), which is the maximum amount of memory cells that can be accessed in the given amount of time.

Modern CMP architectures use memory as the main means of interprocessor communication. The memory used by each core can roughly be classified in two parts, private and shared. The private part contains control variables, counters and other data that are exclusive to the computation being executed in this core. The shared data part consists of portions of the input as well as computation that is shared across threads. Since multi-core threads are not synchronous, even if they are executing the same basic functions we expect the execution to become somewhat out of kilter since execution of branching statements and other such might vary from thread to thread. The end effect is that write access to shared memory can be modeled as a random process with a

\(^1\) In practice there have been architectures in which the memory size was strictly greater than \( 2^w \). Currently in the Intel architecture the size \( w \) places a limit on the largest addressable space but this has not always been the case (e.g. the 8088 processor).

\(^2\) Observe that we use big-Oh in this case advisedly as it allows for values strictly smaller than \( \log n \).

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certain probability of collision. A reasonable first order approximation is to consider memory access to shared data as uniformly random with \( p \) processors contending for access to memory.

In this subsection we investigate the expected number of collisions for \( p \) cores accessing \( m = O(n^k) \) memory cells, with \( k \geq 1 \), uniformly at random for a duration of time \( T_p(n) = T(n)/p = O(n^c/p) \) for some \( c \geq k \).

Clearly, the smaller the number of processors the lower the probability of collision. The question is for what value of \( p \) as a function of \( n \) does this probability become negligible.

This reduces to a balls-and-bins scenario (see, e.g. [14]). Let us first consider the total number of overall collisions in one step. Let \( C \) be a random variable denoting the number of collisions in one step. The probability that two memory accesses are to the same cell is \( 1/m \). Since there are \( p^2 \) pairs of memory accesses, the expected number of collisions in one step is

\[
E[C] = \frac{p(p-1)}{2m}.
\]

As \( m \) grows this expression tends to 0 if \( p < \sqrt{m} \), tends to infinity if \( p > \sqrt{m} \), and to \( 1/2 \) for \( p = \sqrt{m} \).

Since \( m = O(n^k) \) and \( T(n) = O(n^c) \) for \( c \geq k \), the number of collisions per step becomes negligible when \( p = O(n^{k/2}) = O(\sqrt{T}) \).

Now we consider an alternative expression for memory access conflicts, namely the number of cells involved in collisions at each step. Thus, if three or more accesses are to the same cell, the event counts as one collision. Let \( X \) be a random variable denoting the number of memory cells which suffer a collision when there are \( p \) simultaneous memory accesses. The probability of a memory cell not being accessed is \( 1 - 1/m \), and thus the expected number of accessed cells is

\[
m - m(1 - 1/m)^p.
\]

Then, the expected value of \( X \) is

\[
E[X] = p - m + m(1 - 1/m)^p.
\]

Assume that \( p = m^\alpha \) with \( \alpha \leq 1 \). The expression above is then

\[
E[X] \approx m^\alpha - m + me^{-m^{\alpha-1}}.
\]

Using the Taylor expansion of \( e^{-m^{\alpha-1}} \) we obtain

\[
E[X] \approx \frac{m^{2(\alpha-1)}}{2}.
\]

Again, when \( m \) tends to infinity, the above tends to 0, 1/2, or diverges if \( \alpha \) is less, equal, or greater than 1/2, and thus the threshold again is for \( p = \sqrt{m} \). Clearly the smaller \( p \) is, the fewer the expected the collisions.

Case 1. If \( p = m \), then \( E[X] = m/e \), and \( E[C] = (m - 1)/2 \). Thus in each step about 37\% of memory cells have more than one processor trying to access them and about half of the accesses result in collisions.

Case 2. If \( p = \sqrt{m} \) then on average there is a collision every two steps of an execution.

Case 3. If \( p < \sqrt{m} \) the number of collisions goes to zero as \( m \) grows.

We consider now the impact of collisions in the parallel time of a program. Suppose that all \( p \) processors are active for the duration of the program and execute during \( T(n)/p \) instructions each. Suppose that every instruction of the program takes unit time if there is no collision and a \( s \geq 1 \) units of time otherwise. The total number of operations that resulted in a collision during the execution of the program is \( Col = T(n) \cdot \frac{p(p-1)}{2m} = O\left(\frac{n^p^2}{m}\right) \). Since the parallel time of the
program equals the maximum of the time among all processors, this time is minimized when collisions are distributed equally among processors. Thus, assume that the number of collisions per thread is \( u = \frac{Col}{p} = O \left( \frac{n^c m}{p} \right) \). Then the total execution time including collisions is at least \( T'_p(n) = us + T_p(n) - u = O \left( \frac{n^c p(s-1)}{m} + \frac{n^c}{p} \right) \), and hence the slowdown factor due to collisions is \( f = \frac{p^2(s-1)}{m} + 1 \). Then, if \( p = \sqrt{m} \) there is a constant slowdown of approximately \( s/2 \) as \( m \) grows. The slowdown becomes negligible for smaller \( p \) and it grows with the input size for \( p \geq \sqrt{m} \).

Similarly, if instead of charging \( s \) for a collision we only charge \( s \) once to all processors involved in a collision, then the expected slowdown due to collisions is derived in terms of the expected number of cells involved in collisions. The total number of cells involved in collisions during the execution is \( T(n)E[X] \). Thus the average number of cells with collisions per processor is \( T_p(n)E[X] \). Again, the running time due to collisions is at least \( T'_p(n) = sT_p(n)E[X] + T_p(n) - T_p(n)E[X] = T_p(n)(E[X](s-1) + 1) \). Thus the slowdown factor is \( f = (E[X](s-1) + 1) \). Since the conditions for divergence of \( E[X] \) equal the ones for \( E[C] \) above, the same considerations apply for the significance of the slowdown in this case.

### 3.4 Processor Communication Network

Traditionally, parallel computers use either shared memory or a processor communication network (or both) to exchange information between the various processing units. The advantage of shared memory is that no additional hardware is required for it; the disadvantages are issues of synchronization and memory contention. Hence a widely explored alternative is the use of an ad-hoc processor communication networks connecting the processors. In general, from the perspective of performance a full communication network is the preferable network architecture. However when the number of processors is assumed to be very large this is unfeasible. For example for the case of \( \Theta(n) \)-processors of many commercial PRAM implementation the number of interconnects required would have been \( \Theta(n^2) \) which is prohibitive. Thus there was extensive study of alternative network topologies which reduced the complexity of the network while attempting to minimize the penalty in performance derived from the smaller network. Among the most successful such architectures we have the hypercube, the butterfly and the tori (see e.g. [18]).

We observe now that full processor communication network becomes a realistic possibility if the number of processors is \( O(\log n) \) or even possibly \( O(n^\alpha) \) for some \( \alpha \ll 1/2 \). For example for a modest (by present standards) input size of 100,000,000 even \( n^{1/2} \) processors would require and impossible number of interconnects on the full graph. A network of \( O(\log n) \) processors on the other hand would require 300 interconnects which are well within the realm of current architectures.

### 3.5 Buffer overflow

Aside from issues of network topology, in practice it is natural to assume that each processor can handle at most a small constant number of messages at once. If more than a constant number of processors send messages to a single processor, said messages would queue up at the receiving end for further processing. In this section we consider a natural communication model in which in each instruction cycle a processor may send a message to at most one other processor. In practice depending on the specific application the probability of collision may range anywhere from zero for the execution of independent threads to one for, say, a master processor serializing requests to some shared lock. As a compromise we model again this process as if the processors chose their
destination uniformly at random. Let \( p \) be the number of processors; then the maximum number of collisions observed at the most loaded buffer is \( O(\log p) \) with high probability \([14]\). If \( p = \Theta(n) \) then buffer handling can introduce delays of 20–100 instruction cycles. In this case even \( p = \Theta(n^\alpha) \) for \( 0 \leq \alpha < 1 \) might prove too costly. In contrast if we assume \( p = \Theta(\log n) \) the most congested buffer would contain \( O(\log \log n) \) elements which for all practical purposes is at most 6.

### 3.6 Divide-and-Conquer Algorithms

Divide-and-Conquer algorithms are naturally suited for parallelization. Instances at the same level of the recursion tree are independent and can be scheduled to be executed in parallel. This is especially well suited for multi-threaded systems, as each recursive call can simply be handled to a separate thread. This strategy requires no parallelization of the divide and combine phases of the recursion, which can be executed by each thread just as in the sequential algorithm. It has been shown that this easy parallelization yields optimal speedups for a large class of divide-and-conquer algorithms \([12]\), but only for a bounded number of processors. Thus, in a system with a logarithmic or sublinear number of processors, obtaining the maximum possible speedup for this class of algorithms is simple and can be realized with a general strategy that is independent of the algorithm itself.

Consider a divide-and-conquer algorithm whose time complexity can be written as \( T(n) = aT(n/b) + f(n) \). The master theorem \([11]\) yields the time bounds for a sequential execution of such an algorithm. A parallel version of this theorem can be obtained by analyzing the parallel time \( T_p(n) \) of an execution in which recursive calls are executed in parallel and scheduled with work-stealing \([9]\) with a bounded number of processors \([12]\):

\[
T_p(n) = \begin{cases} 
O(T(n)/p), & \text{if } f(n) = O(n^{\log_b(a) - \epsilon}) \text{ and } p = O(n^{\epsilon}) \quad \text{(Case 1)} \\
O(T(n)/p), & \text{if } f(n) = \Theta(n^{\log_b(a)}) \text{ and } p = O(\log n) \quad \text{(Case 2)} \\
\Theta(f(n)), & \text{if } f(n) = \Omega(n^{\log_b(a) + \epsilon}) \text{ and } af(n/b) \leq cf(n), \text{ for some } c < 1 \quad \text{(Case 3)}
\end{cases}
\]

Optimal speedups are achieved in Cases 1 and 2 only for \( p = O(n^{\epsilon}) \) for \( \epsilon > 0 \), and \( p = O(\log n) \), respectively. In Case 3, the time is dominated by the sequential divide and conquer time \( f(n) \) at the top of the recursion \([12]\).

### 3.7 Cache imposed bounds

Cache contention is a key factor in the efficiency of multi-core systems. Various multi-core cache models have been studied, with a focus on algorithms and schedulers with provable cache performance. Many of the results involving shared and private caches performance require bounds on the number of processors related to the relative size of the private and shared caches.

The parallel external memory model \([3]\) models \( p \) processors, each with a private cache of size \( M \), partitioned in blocks of size \( B \). A sorting algorithm given in this model is asymptotically optimal for the I/O bounds for at most \( p \geq n/B^2 \) processors, and it is actually proven that \( p \geq n/(B \log B) \) is a lower bound for optimal processor utilization. This algorithm is used in further results in the model for graph and geometry problems \([4, 1, 2]\). Thus the assumption that \( p \geq n/B^2 \) is carried on to these results as well, some of which actually require \( p \leq n/(B \log n) \).

Shared cache performance is studied in \([7]\), which compares the number of cache misses of a multi-threaded computation running on a system with \( p \) processors and shared cache of size \( C_2 \)
to those of a sequential computation with a private cache of size $C_1$. They show that the parallel number of misses is at most the sequential one if $C_p \geq C_1 + pd$, where $d$ is the critical path of the computation. This implies $p \leq (C_p - C_1)/d$, is less than $n$ (as otherwise all the input would fit in the cache) and is usually sublinear, as $d$ is rarely constant and is $\Omega(\log n)$ for many algorithms.

Gibbons et al. [6] extend the analysis of multi-core divide-and-conquer algorithms in a multi-core cache model of $p$ processors with private L1 caches of size $C_1$ and a shared L2 cache of size $C_2$. An assumption of the model is that $p \leq \frac{C_2}{C_1} \ll n$, since the input size is assumed not to fit in L2. This model is used to show that an online scheduler achieves optimal speedup and cache complexity within constant factors of the sequential cache complexity for a class of hierarchical divide-and-conquer algorithms (whose divide and conquer phases are in turn divide-and-conquer algorithms as well). Optimality for some algorithms, such as Strassen’s matrix multiplication and associative matrix inversion even require $p \leq \left( \frac{C_2}{C_1} \right)^{\frac{1}{1+\alpha}}$ [6].

Cache efficient dynamic programming algorithms have been designed in this multi-core model with the same $p \leq \frac{C_2}{C_1}$ assumption [10], as well as in a shared cache model with $p \leq C_2/B$, where $B$ is the block size. Thus although the time complexity of parallel dynamic programming allows a large number of processors for optimal speedups (e.g., $T_p = O(n^3/p + n)$) for Gaussian elimination paradigm problems, which is optimal for $p \leq n^2$, the efficiency in cache performance restricts the level of parallelism.

Observe that presently the ratio between L2 shared cache and private L1 cache is in the order of 4 to 100 depending on the specific processor architecture.

### 3.8 The class $E(p(n))$

The class $NC$ can be defined as the class of problems which can be solved in polylog time using polynomially many processors. It is believed that $NC \neq P$ and hence that there are known problems which do not admit a solution in time $O(\log^kn)$. In our case we are interested in the study of problems which can be sped up using $O(\log n)$ or $O(n^{\alpha})$ processors for $\alpha < 1$. Kruskal et al. [17] introduced the classes $ENC$ and $EP$ which encode the classes of problems that allow optimal speed up (up to constant factors) using polynomially many processors. The class $ENC$ has polylogarithmic running time, while the class $EP$ has polynomial running time. Following their notation we introduce the class of problems $E(p(n))$ which is the class of problems that can be solved using $O(p(n))$ processors in time $O(T(n)/p(n))$ where $T(n)$ is the running time of any sequential solution to the problem. In this work we are particularly interested in the classes $E(\log n)$ and $E(n^{\alpha})$ for $\alpha < 1$.

The class $ENC$ is sharpening of the well known class $NC$. Recall that the class $NC$ requires maximal speedup down to polylogarithmic time even at the cost of a polynomial amount of inefficiency (i.e. the ratio between sequential and parallel work). In contrast $ENC$ requires the same speedup but bounds the inefficiency to a constant factor.

The class $E(\log n)$ bounds the inefficiency to a constant which implies a speed up of $O(\log n)$ on the sequential solution to the problem. Observe that by Brent’s Lemma $ENC \subset EP \subset E(\log n)$. The reverse is not the case, i.e. $E(\log n) \neq EP, ENC$ unless $NC = P$ since there are known $P$-complete problems which allow optimal speedup using $O(\sqrt{n})$.

Similarly $E(n^{\alpha})$ bounds the inefficiency to a constant which implies a speed up of $O(n^{\alpha})$ on the sequential solution to the problem. Again we have $ENC \subset EP \subset E(n^{\alpha})$ while $E(n^{\alpha}) \neq EP, ENC$ for $\alpha \leq 1/2$ as the $P$-complete speed up referred to above uses $O(\sqrt{n})$ processors.
This gives a theoretical separation between the problems that can be speed up optimally using polynomially many processors and those that can be speed up using a sublinear number of processors $O(n^\alpha)$ for $\alpha \leq 1/2$.

### 3.9 Optimal Time Simulation of Turing Machines by $O(\log n)$ Processors

We show that any computation on a Turing machine that takes time $T(n) \geq n \log n$ can be carried out in parallel by a multi-core system with $p = \lg n$ processors in time $T_p(n) = O(T(n)/(\lg n + \lg \lg n))$. There are known simulation results for Turing Machines by a sequential RAM [16] as well as by a PRAM [13]. In the latter it is shown that a deterministic machine running in $T(n)$ time can be simulated by a PRAM in time $O(\sqrt{T(n)})$ using an exponential number of processors and memory addressing on words of size $O(\sqrt{T(n)})$. We adapt this simulation to a more realistic logarithmic number of processors and word size.

**Outline** Let $M$ be a single-tape deterministic Turing Machine. The idea of the simulation is to treat contiguous blocks of $b = b(n)$ bits of $M$’s tape as a word in RAM. By precomputing $M$’s resulting configuration after $b$ steps when starting with each possible block, we can then simulate $b$ steps of $M$ at a time by successively looking up the next configuration of $M$ from the precomputed table. Let $g(n)$ denote the precomputation time. If each access to the precomputed table takes constant time, then the total time of the simulation is

$$T_p(n) = \frac{T(n)}{b(n)} + g(n)$$

**Precomputation phase.** Since in $b$ steps $M$ can only alter the contents of $b$ cells, for a given position within the tape we need only to consider the content of the $b$ cells to the left and $b$ cells to the right of the current positions in order to compute the resulting configuration after $b$ steps. A block configuration of $M$ is a tuple $(s, B)$, where $s$ is a state, $B$ is a $(2b - 1)$-bit string representing the contents of a segment of $M$’s tape around some position of the head. For each possible block configuration $c$, we store in $A[c]$ the resulting configuration when running $M$ starting from $c$ (i.e., the new state and block contents), plus information about how many positions the head moved, and in which direction. The latter is necessary to know where the new block should be centered in $M$’s tape. A block configuration $c$ uses $|c| = 2b - 1 + d = O(b)$ bits, where $d$ is the constant number of bits required to indicate a state of $M$. Let $k$ be a constant such that $|c| \leq kb$. Then there are at most $2^{kb}$ starting block configurations. Since the resulting configurations starting from all possible configurations can be computed independently in parallel and each computation takes $O(b)$ time by direct simulation of $M$, the total precomputation time is $g(n) = O\left(\frac{2^{kb}}{p}\right)$ using $p$ processors. Note that the precomputation requires only $M$’s specification but is independent of a particular input.

**Simulation phase.** Suppose the configuration table $A$ has already been computed and it is stored in the RAM of the multi-core machine. If the length of each configuration is smaller than the machine word’s length, then $A$ can be indexed by configuration and each entry can be accessed in constant time. For this sake we set $b = \frac{\lg n + \lg \lg n}{k}$, and thus $|c| \leq \lg n + \lg \lg n = \Theta(\log n)$. Therefore $A$ can be stored as an array of configurations, indexed by configurations. Given $M$ and an input $x$,
and starting with the initial configuration $c_0$, the multi-core simulates $M$ (using one processor) by applying $c_{i+1} = A[c_i]$, and updating the contents of $M$’s tape at each step, until $c_{i+1}$ contains a final state. Since at each step $A$ can be accessed in constant time and the relevant part of $M$’s tape can be updated in constant time, the simulation takes $O(T(n)/(\lg n + \lg \lg n))$ time and the precomputation takes $O\left(\frac{2^{\lg n + \lg \lg n} (\lg n + \lg \lg n)}{p}\right)$ time, which is $O(n \lg n)$ for $p = \lg n$. Thus the total time is

$$T_p(n) = O\left(\frac{T(n)}{\lg n + \lg \lg n} + n \lg n\right)$$

**Faster recursive precomputation.** The approach described above requires $T(n) = \Omega(n \lg^2 n)$ to be optimal. However, we can relax this requirement by speeding up the precomputation phase. The idea is to recursively apply the simulation on the computation of each entry of $A$. Let $g_i$ and $b_i$ denote the precomputation time and block length of the $i$-th level simulation, respectively. Thus $g_m = g(n)$ is the total precomputation time and $b_m = b(n)$ is the block length of the final simulation as described above. Since the computation of each entry of $A$ can now be sped up by $b_m - 1$, we have

$$g_m = 2^{kb_m/p} \frac{b_m}{b_{m-1}} + g_{m-1},$$

where $k$ is a constant such that for all $i$, a configuration in level $i$ has size at most $kb_i$. We then set $b_{m-i} = \frac{\lg n + \lg \lg n}{k^2}$ for all $0 \leq i \leq m = \lg \left(\frac{\lg n + \lg \lg n}{k}\right)$. Then, $b_{m-i}/b_{m-i-1} = 2$, which is the length of the critical path at each recursive level. Then,

$$g_{m-i} = \max\left\{\frac{2^{kb_i+1}}{p}, 2\right\} + g_{m-i-1} = \max\left\{\frac{2(n \lg n)^{\frac{i}{p}}}{p}, 2\right\} + g_{m-i-1}$$

Note that $2(n \lg n)^{\frac{1}{p}}/p \leq 2$ when $i \geq \lg \left(\frac{\lg n + \lg \lg n}{\lg p}\right)$. Let $i^* = \lg \left(\frac{\lg n + \lg \lg n}{\lg p}\right)$. Since $g_0 = 0$,

$$g_m = \sum_{i=0}^{m-1} g_{m-i} - g_{m-i-1}$$

$$= \frac{2}{p} \sum_{i=0}^{i^*} (n \lg n)^{\frac{i}{p}} + \sum_{i=i^*+1}^{m-1} 2$$

$$\leq \frac{2}{p} (n \lg n + i^* \sqrt{n \lg n}) + 2(\lg \lg p - \lg k - 1)$$

$$\leq \frac{4n \lg n}{p} + 2 \lg \lg p$$

Therefore, the total simulation time is now

$$T_p(n) = O\left(\frac{T(n)}{\lg n + \lg \lg n} + n \frac{\lg n}{p} + \lg \lg p\right),$$

which for $p = \lg n$ and $T(n) \geq n(\lg n + \lg \lg n)$ is $O\left(\frac{T(n)}{\lg n + \lg \lg n}\right)$. 

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Larger number of processors. A simulation like the one described above would not be optimal if $p = w(\lg n)$. If the length of the blocks is kept at $b = O(\lg n + \lg \lg n)$, then the total time is $T_p(n) = \Omega \left( \frac{T(n)}{\lg n + \lg \lg n} + \frac{n}{p} + \lg \lg p \right)$, which for $p = w(\lg n)$ can never be $O(T(n)/p)$. A longer block length $b = \omega(\lg n)$ could reduce the simulation phase time, but would require an infeasible superpolynomial-size precomputed table. This, of course, does not preclude the existence of other approaches that could result in optimal simulation time with a larger number of processors.

3.10 Amdahl’s Law

Consider a program whose execution has a serial part that cannot be parallelized (unless $P = NC$) represented by $S(n)$ and a fully parallelizable part denoted by $P(n)$ then the parallel time with $p$ processors is:

$$T_p(n) = S(n) + P(n)/p$$

and the speedup is represented by

$$\frac{T_1(n)}{T_p(n)} = \frac{S(n) + P(n)}{S(n) + P(n)/p}.$$ 

Observe now that for $p = \Theta(n)$ we get that the parallel program is noticeably faster only if $S(n) = O(P(n)/n)$.

For $p = \Theta(n^\alpha)$ we get that the parallel program is noticeably faster only if $S(n) = O(P(n)/n^\alpha)$. Lastly, for $p = \Theta(\log n)$ we get that the parallel program is noticeable faster if $S(n) = P(n)/\log n$.

Observe that most practical algorithms on large data sets run in time $O(n \log n)$ or less, with the sequential part often corresponding to I/O operations, i.e. reading the input. This means that the likeliest value for which one can obtain optimal speedup corresponds to $P(n)/S(n)$ which is often (though not always) $\log n$.

4 Conclusions

We presented a list of theoretical arguments and practical evidence as to the existence of a qualitative difference between the classes of problems that can be sped up with a sublinear number of processors and those that can be sped up with polynomially many processors.

We also show that in various specific instances even though there are optimal algorithms for either case, it is conceptually and practically much simpler to design an algorithm for a sublinear number of processors. The benefits of a low processor count extend to issues of processor communication, buffering, memory access and cache bounds.

We introduced classes that describe the problems that allow for optimal speed up, up to a constant factors, for sublinear number of processors and show that they contain a strictly larger class of problems that the PRAM equivalents introduced by Kruskal, Rudolph, and Snir in 1990 [17], unless $NC = P$.

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References