Homework 1

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List of Algorithms

&& : 1 || : 0 | : 0 $^{\land}$: 0

2 Problem 2

Suppose OpenMP did not have the reduction clause. Show how to implement an efficient parallel reduction by adding a private variable and using the critical pragma.

```
/* File: problem2.cpp
 * Purpose: Alternates sign of integer added to sum
                sum = 0 + 1 + -2 + 3 + -4...
  Compile: g++ -Wall -fopenmp -o problem2 problem2.cpp -std=c++11
            g++ -Wall -fopenmp -o problem2 problem2.cpp -DDEBUG -std=c++11
 * Run:
            ./problem2
 * Input:
           none
 * Output: Times for each of the three runs
 * Notes:
            If ran with the -DDEBUG flag you can see what the sum should
      1.
            be based on n
#include <inttypes.h>
                       // Better integer functionality
#include <stdio.h>
                       // Printing to console
#include <omp.h>
                        // Multithreading
#include <chrono>
                       // High precision clock
using namespace std::chrono;
// Global
uint8_t
            thrds
                   = omp_get_num_procs();
int main(int argc, char* argv[]) {
    uint8_t times = 20;
    high_resolution_clock::time_point t1 = high_resolution_clock::now();
   high_resolution_clock::time_point t2 = high_resolution_clock::now();
    duration<double> no_omp_time = duration_cast<duration<double>>\
        (high_resolution_clock::now() - high_resolution_clock::now());
    duration<double> omp_time = duration_cast<duration<double>>\
        (high_resolution_clock::now() - high_resolution_clock::now());
    duration<double> no_reduc_time = duration_cast<duration<double>>\
        (high_resolution_clock::now() - high_resolution_clock::now());
    for(uint8_t j = 0; j < times; ++j)
```

```
{
        uint64_t
                          = 80000000,
                    n
                             = 0;
                    k
        int64_t
                             = 0;
                    \operatorname{\mathtt{sum}}
        // RESET for baseline
        t1 = high_resolution_clock::now();
        for (k = 0; k < n; ++k)
            sum += ((k \& 1) == 0 ? 1.0 : -1.0) * k;
        }
        t2 = high_resolution_clock::now();
        no_omp_time += duration_cast<duration<double>>(t2 - t1);
#ifdef DEBUG
        if (j == 0){
            printf("No OMP sum : %" PRIi64 "\n", sum);
        }
#endif
        // RESET for reduction + omp
        sum = 0;
        t1 = high_resolution_clock::now();
        #pragma omp parallel for num_threads(thrds) reduction(+: sum) private(k)
        for (k = 0; k < n; ++k)
        {
            sum += ((k \& 1) == 0 ? 1.0 : -1.0) * k;
        }
        t2 = high_resolution_clock::now();
        omp_time += duration_cast<duration<double>>(t2 - t1);
#ifdef DEBUG
        if (j == 0){
                                  : %" PRIi64 "\n", sum);
            printf("OMP sum
#endif
        // RESET for no reduction
        sum = 0;
        k = 0;
        t1 = high_resolution_clock::now();
        #pragma omp parallel num_threads(thrds)
            int64_t thread_sum = 0;
            #pragma omp for
            for(uint64_t i = k; i < n; ++i){</pre>
                // Locally (privately) runs this
                thread_sum += ((i \& 1) == 0 ? 1.0 : -1.0) * i;
            }
```

```
#pragma omp critical
            sum += thread_sum;
        }
        t2 = high_resolution_clock::now();
        no_reduc_time += duration_cast<duration<double>>(t2 - t1);
#ifdef DEBUG
        if (j == 0){
            printf("No Reduc sum : %" PRIi64 "\n", sum);
#endif
   printf("Averages over %" PRIu8 " runs:\n", times);
                      : %.14f\n", no_omp_time.count() / times);
                      : %.14f\n", omp_time.count() / times);
   printf("OMP
   printf("No Reduc
                     : %.14f\n", no_reduc_time.count() / times);
    return 0;
}
```

```
fopenmp -o problem2 problem2.cpp -DDEBUG -std=c++11
cvle@:HW1S
           ./problem2
No OMP sum
               -40000000
OMP sum
               -40000000
No Reduc sum : -40000000
Averages over 20 runs:
No OMP
          : 0.40371242310000
OMP
          : 0.06024930730000
No Reduc
          : 0.06116008355000
kyle@:HW1S
```

Figure 1: Example debug output.

<mark>/le@:HW1</mark>\$./problem2 Averages over 20 runs: No OMP : 0.40169680075000 OMP : 0.05187247365000 No Reduc : 0.05121839645000 kyle@:HW1\$./problem2 Averages over 20 runs: No OMP : 0.40076352375000 OMP : 0.05140341830000 0.05126510895000 kyle@:HW1\$./problem2 Averages over 20 runs: No OMP 0.40068608615000 OMP 0.05138620015000 0.05121338355000 No Reduc

Figure 2: Better performance without reduction.

As can be seen in the figures the sums are performing as expected. An interesting, and expected outcome is that in Figure 1 it takes 0.06 seconds to run OMP and $No\ Reduc$ but in Figure 2 it takes 0.05 seconds. The $No\ OMP$ takes 0.40 seconds regardless. The reason for this behavior is that OMP uses the cores you give it and at the time of recording the first figure the browser was open and running a video. When I recorded the second Figure I had closed my browser to maximize performance for multi-core processing. The $No\ OMP$ section of code was only running on one core, so it did not care that I had a video playing.

Please see my repository for the full code breakdown for this problem.

3.1 Problem 3a

```
#pragma omp for
for(i = 0; i < (long) sqrt(x); ++i) {
   a[i] = 2.3 * i;
   if (i < 10)
       b[i] = a[i];
}</pre>
```

3.2 Problem 3b

This code section is not suitable for OpenMP because of the && operator in comparison. Per the OpenMP documentation §2.6, p53:

```
\begin{array}{ccc} test\text{-}expr & \text{One of the following:} \\ & var\ relational\text{-}op\ b \\ & b\ relational\text{-}op\ var \\ \end{array} \begin{array}{cccc} relational\text{-}op & \text{One of the following:} \\ & < \\ & < \\ > \\ > = \\ \end{array}
```

3.3 Problem 3c

This code can be ran with OMP but it is dependent on whether or not foo() is threadsafe.

```
#pragma omp for
for(i = 0; i < n; ++i) {
    a[i] = foo(i);
}</pre>
```

3.4 Problem 3d

This problem is similar to 3c in that it is dependent on foo(). Nothing else is preventing this from being made parallel. Only comparisons and assignments are used. Access of b[i] is fine because nothing is writing to it so there is no opportunity for trouble.

```
#pragma omp for
for(i = 0; i < n; ++i) {
    a[i] = foo(i);
    if(a[i] < b[i])
        a[i] = b[i];
}</pre>
```

3.5 Problem 3e

Cannot be ran in parallel because the break.

3.6 Problem 3f

Standard use of OMP, though I feel that it didn't acutally use OMP. I tried to throw a reduction at it but I kept getting an error about dotp being private. I threw a private local sum into the for loop and added $pragma\ omp\ critical$ followed by dotp+=sum but it gave no performance increase.

```
dotp = 0;
#pragma omp for
for(i = 0; i < n; i++){
    dotp += a[i] * b[i];
}</pre>
```

3.7 Problem 3g

I thought this one was going to have trouble because of the $i \leftarrow k$ portion but testing showed the values matched for the sequential sanity test and the OMP portion. Of note though, OMP was signficantly slower than sequential until I ramped up k.

```
#pragma omp for
for(i = k; i < 2 * k; ++i){
    a[i] = a[i] + a[i-k];
}</pre>
```

3.8 Problem 3h

This problem is the same as 3g except we are using what is essentially a shared constant, b and we are running through the whole list, depending on what k is. Other than that it behaves the same. Passed sanity check and ran without issue.

```
#pragma omp for
for(i = k; i < n; ++i){
    a[i] = b * a[i-k];
}</pre>
```

Given a task that can be divided into m subtasks, each requiring one unit of time, how much time is needed for an m-stage pipeline to process n tasks?

How many of the subtasks are dependent on other subtasks? If all m subtasks require the previous subtask then m*n time is needed to process n tasks. If they can all be ran independently and if we have enough threads we can run it in n time. It is more likely that we will have some combination of subtasks that require other subtask completion, therefore the only blanket assumption that can be safely made is that it will take m*n time.

If the address of the nodes in a hypercube has n bits. How many nodes can it be at the most and how many edges does each node have? Give an algorithm that routes a message from node u to node v in this k-node hypercube in no more than log(k) steps.

Hypercubes are generalized by dimensionality. A hypercube address containing n bits will exist in n dimensions (d). The maximum number of nodes is determined by d, and is defined as 2^d . A trait of hypercubes is that each node will have d edges. The total amount of edges would be defined as $d*2^{d-1}$. The number of nodes you have to travel between u and v is equal to the number of differing bits.

For example:

```
\begin{array}{cccc} 000 & \Longrightarrow & 111 & : 3 \text{ edges} \\ 001 & \Longrightarrow & 100 & : 2 \text{ edges} \\ 110 & \Longrightarrow & 011 & : 2 \text{ edges} \end{array}
```

A traversal algorithm would be along the lines of Algorithm1

Algorithm 1 Hypercube Traversal

```
● Move from node u to v
i \leftarrow 0
current\_bit \leftarrow u[i]
left\_bit \leftarrow u[i]
while i < dimension(hypercube) do
current\_bit \leftarrow u[i]
if current\_bit = v[i] then
i \leftarrow i + 1
else
route(current\_bit)
route(current\_bit)
end if
end while
```

If built properly there will only be one node the algorithm can route to during the $route(current_bit)$ section. Essentially we look at the left-most bit and if it is the same we move to the next bit, if it is different we must move to a new node. There will only be one node available because if they are on the same dimension they would have the same bit in that slot. For example on a 3-dimensional hypercube $000 \implies 100$ looks at the left bit and sees it is on a different dimension. There is only one node connected to 000 that follows the format 1xx, so that is the only place it can go. If we wanted to route from $000 \implies 010$ we would identify we are on the correct 3rd dimension, then look at the middle bit. From there we would see a need to move in the second dimension. There is only one connected node that matches in the second dimension, so that is where it would go.

6 Graduate Assignment

Do a search on Shuffle-exchange network topology. Draw the network with 16 processor nodes (carefully numbering each node binary and showing what is a shuffle link, what is an exchange link). If k is the number of digits in the binary address, how many nodes (n) are there? With n nodes what is the diameter (d) of the networks, the bisection width (b) and how many edges/node?

When you first drew a shuffle-exchange network, also sometimes called a butterfly network, I immediately thought that looked like a sorter I came up with in CSC300. I later learned that sort wasn't something I had uniquely developed and was in fact a bitonic sort.

If there are k digits in the address then there will be a maximum of $(k + 1) * 2^k$ nodes. With n nodes the diameter will be 2 * k. If the network is setup to wrap around then the diameter could be k. Bisection on butterfly networks is easy to see, visually, and is defined as k + 1. Most nodes would contain 4 edges, where two come from "below" and two come from "above". The boundary nodes would contain 2 edges, ethier connecting "up" or "down".