PC40 Hands-on: UPC

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Abstract

For this hands-on assignement, I wanted to measure the performance of the different versions of the code and compare them to measure speedups. For this reason, my code diverges slightly from the template that was given to us, as I needed to insert a timing method and avoid refactoring the code.

You will find in this report listings of the different versions of the code, alongside experimental measurements and commentaries.

The source code itself, this report's source code and instructions on how to build and run the code yourself can be found on this project's git repository: https://github.com/adri326/pc40-upc/.

1 Simplified 1D Laplace solver

1.1 C implementation

The C implementation of the laplace solver has been slightly modified to time the main loop. It is single-threaded but the implementation allows for compiler SIMD optimizations. Settings common to every version (vector size, epsilon, max number of iterations) have been placed in a file called settings.h.

This code has been compiled with gcc v4.9.0 and run on the mesoshared server, yielding the following results:

Options	Time (avg)	$CI (\sigma = 0.01)$
-00	$450\mu s/{\rm iter}$	$\pm 10 \mu s/{\rm iter}$
-03	$140 \mu s/{ m iter}$	$\pm 10 \mu s/\text{iter}$

Table 1: Timing results for the C implementation of the 1D Laplace solver (Listing 1)

```
#include <stdio.h>
#include <stdlib.h>
#include <time.h>
#include "settings.h"

#define LEN DESIRED_LEN

void init();
```

```
9 void iteration();
  void copy_array();
10
11
  double x new [LEN];
12
  double x [LEN];
13
  double b[LEN];
15
  int main() {
16
       init();
17
18
19
       clock_t begin = clock();
       for (size t n = 0; n < ITERATIONS; n++) {
20
           iteration();
21
           if (n != ITERATIONS - 1) copy_array();
22
23
       clock t end = clock();
24
25
       double diff = (double)(end - begin) / CLOCKS PER SEC / ITERATIONS;
26
       double mean squared = 0.0;
27
       if (DISPLAY) {
28
           printf("
29
                     b
                                X
                                        x_new | diff
                                                            | \langle \mathbf{n}^{"} \rangle ;
           printf(" | :-
                            - | :--
                                        | :---- | :-
30
           for (size t i = 0; i < LEN; i++) {
31
               printf("| %1.41f | %1.41f | %1.41f | % 1.41f | \n", b[i], x[i], x new[i
32
      [ ], x_{new}[i] - x[i] );
33
           }
34
      35
       - x | i | );
       printf("\nIterations: %d\n", ITERATIONS);
36
       printf("Mean squared difference: %1.41f\n", mean squared / LEN);
37
       printf("Took \%.3lfms/iter!\n", diff * 1000);
38
39
40
  void init() {
41
      \operatorname{srand}(\operatorname{time}(0));
42
       for (size t i = 0; i < LEN; i++) {
43
           b[i] = (double) rand() / RAND MAX;
44
           x[i] = (double) rand() / RAND MAX;
45
           x_{new}[i] = 0;
46
       }
47
48
49
  void iteration() {
50
       for (size t i = 1; i < LEN - 1; i++) {
51
           x \text{ new}[i] = 0.5 * (x[i-1] + x[i+1] + b[i]);
52
53
      x \text{ new}[0] = x[0];
      x \text{ new}[LEN - 1] = x[LEN - 1];
55
56
  void copy_array() {
58
59
       for (size t i = 0; i < LEN; i++) {
           x[i] = x_new[i];
60
61
62
```

Listing 1: C implementation of the 1D Laplace solver

1.2 Porting the C code to UPC

The base C implementation was designed to be able to quickly port it to UPC. A new if statement had to be inserted in the for loop within iteration(). Additionally, several lines had to only be executed by the thread 0, so additional conditionals were added when needed.

Finally, the x, xnew and b arrays were made shared and upc_barrier statements were added at the end of init, iteration and copy_array: this is to prevent threads from beginning processing the next iteration when the x and xnew arrays aren't ready, and to prevent the thread 0 from stopping early and printing the wrong timing.

This code was compiled with upcc v2.22.0 + gcc v4.2.4 and run on the mesoshared server, yielding the following results:

Threads	Time (avg)	$CI (\sigma = 0.01)$
2	$515.2 \mu s/{ m iter}$	$\pm 3.9 \mu s/{\rm iter}$
3	$536.2 \mu s/{ m iter}$	$\pm 2.4 \mu s/{\rm iter}$
4	$312.4 \mu s/{ m iter}$	$\pm 2.0 \mu s/{\rm iter}$
8	$204.0 \mu s/{ m iter}$	$\pm 2.0 \mu s/{\rm iter}$
16	$166.2 \mu s/{ m iter}$	$\pm 2.5 \mu s/{\rm iter}$
32	$150.4 \mu s/{ m iter}$	$\pm 3.2 \mu s/{\rm iter}$

Table 2: Timing results for the first UPC implementation of the 1D Laplace solver (Listing 2)

When compiled and run with 3 threads, the code runs noticeably slower. For curiosity, I ran the code with 24 and 31 threads and obtained a similar slowdown:

```
Threads = 24, Time = 252.6\mu s/\text{iter} \pm 3.7\mu s/\text{iter} (expected \approx 160 \ \mu s/\text{iter})
Threads = 31, Time = 317.6\mu s/\text{iter} \pm 4.4\mu s/\text{iter} (expected \approx 150 \ \mu s/\text{iter})
```

```
1 #include <upc relaxed.h>
2 #include <stdio.h>
3 #include <stdlib.h>
4 #include <time.h>
  #include "settings.h"
  #define LEN DESIRED LEN
  void init();
9
  void iteration();
  void copy array();
11
  shared double x new[LEN];
  shared double x[LEN];
  shared double b[LEN];
15
17
  int main() {
18
       init();
19
       clock t begin = clock();
20
       for (size t n = 0; n < ITERATIONS; n++) {
21
            iteration();
22
            if (n != ITERATIONS - 1) copy_array();
23
24
       \operatorname{clock\_t} = \operatorname{end} = \operatorname{clock}();
```

```
26
       if (MYTHREAD = 0) {
27
            double diff = (double)(end - begin) / CLOCKS PER SEC / ITERATIONS;
            double mean squared = 0.0;
29
           if (DISPLAY) {
30
                printf("
                                    X
                                              x new
                                                         diff
                                                                    | \langle n'' \rangle ;
                           b
31
                printf(" | :-
                                  - | :--
                                            - | :---
32
                for (size t i = 0; i < LEN; i++) {
                     printf("| \%1.41f | \%1.41f | \%1.41f | \% 1.41f | \n", b[i], x[i],
34
      x_{new[i]}, x_{new[i]} - x[i]);
35
36
            for (size t i = 0; i < LEN; i++) mean squared += (x new[i] - x[i]) * (
37
      x \text{ new}[i] - x[i];
            printf("\nIterations: %d\n", ITERATIONS);
38
            printf("Mean squared difference: %1.41f\n", mean squared / LEN);
39
            printf("Took \%.31fms/iter!\n", diff * 1000);
40
       }
41
42
43
44
  void init() {
       if (MYTHREAD = 0) {
45
           \operatorname{srand}(\operatorname{time}(0));
46
            for (size t i = 0; i < LEN; i++) {
47
                b[i] = (double) rand() / RAND_MAX;
                x[i] = (double) rand() / RAND MAX;
49
                x_new[i] = 0;
            }
       }
       upc barrier;
54
  void iteration() {
       for (size t i = 1; i < LEN - 1; i++) {
57
            if (i % THREADS == MYTHREAD) {
58
                x \text{ new}[i] = 0.5 * (x[i-1] + x[i+1] + b[i]);
59
            }
       }
61
       if (MYTHREAD == 0) x_new[0] = x[0];
63
       if (MYTHREAD = (LEN - 1) \% THREADS) x_new[LEN - 1] = x[LEN - 1];
65
       upc_barrier;
66
67
68
  void copy_array() {
69
       for (size t i = 0; i < LEN; i++) {
70
            if (i \% THREADS = MYTHREAD) x[i] = x new[i];
71
72
       upc barrier;
73
74
```

Listing 2: First UPC implementation of the 1D Laplace solver

1.3 Optimizing the inner for loop

The first step in optimizing our UPC implementation of the 1D Laplace equation solver is to replace the for (...) if (...) with a single, more efficient for loop.

To achieve this, only the iteration() function had to be changed. This transformation is shown in Figure 1

This change greatly increases the speed of the program:

Threads	Time (avg)	$CI (\sigma = 0.01)$
2	$482.0 \mu s/{\rm iter}$	$\pm 3.2 \mu s/{\rm iter}$
4	$265.2 \mu s/{ m iter}$	$\pm 2.3 \mu s/{\rm iter}$
8	$128.8 \mu s/{ m iter}$	$\pm 1.6 \mu s/{\rm iter}$
16	$74.6 \mu s/{ m iter}$	$\pm 3.1 \mu s/{\rm iter}$
32	$46.0 \mu s/{ m iter}$	$\pm 2.5 \mu s/{\rm iter}$

Table 3: Timing results for the second UPC implementation of the 1D Laplace solver

```
// ex3.upc:
void iteration() {
  for (size_t i = 1; i < LEN - 1; i++) {
      if (i % THREADS == MYTHREAD) {
            x_new[i] = 0.5 * (x[i-1] + x[i +1] + b[i]);
      }
  }
  if (MYTHREAD == 0) x_new[0] = x[0];
  if (MYTHREAD == (LEN - 1) % THREADS)
      x_new[LEN - 1] = x[LEN - 1];
  upc_barrier;
}</pre>
```

```
// ex4.upc:
void iteration() {
    size_t i = MYTHREAD;
    if (MYTHREAD == 0) {
        i += THREADS;
        x_new[0] = x[0];
    }

    for (; i < LEN - 1; i += THREADS) {
        x_new[i] = 0.5 * (x[i-1] + x[i+1] + b[i]);
    }

    if (MYTHREAD == THREADS - 1) x_new[
    LEN - 1] = x[LEN - 1];

    upc_barrier;
}</pre>
```

(a) Previous iteration() function from Listing 2 (b) New iteration() function without the inner if

Figure 1: Modification in iteration() to remove the reduce the number of loop iterations.

1.4 Optimizing the array blocking factor

Our next optimization is to increase the blocking factor B for the x, x_{new} and b arrays. Each operation on the item of index j accesses b[j], x[j-1], x[j+1] and $x_{new}[j]$.

With the default block factor of 1, accesses to both x[j-1] and x[j+1] will always be outside of the current thread's affinity. With a greater block factor, these outside accesses can be reduced to only 2/B accesses on average.

Two parts of the code had to be changed: the array declarations (Figure 2) and the loop in iteration() and copy_array() (Figure 3).

The effects of this change is dependent on B: similar to the results found in Table 2, the code runs fastest when $B = 2^n$. Whichever value is chosen, however, the program did not run faster than ex4.upc (Table 3), but rather ran much slower (the source code of this version can be found here):

Threads	Time (avg)	$CI (\sigma = 0.01)$
2	$959.2\mu s/{\rm iter}$	$\pm 3.9 \mu s/{\rm iter}$
4	$863.9 \mu s/{ m iter}$	$\pm 16.1 \mu s/{\rm iter}$
8	$691.6 \mu s/{ m iter}$	$\pm 6.1 \mu s/{\rm iter}$
16	$581.5 \mu s/{\rm iter}$	$\pm 4.1 \mu s/{\rm iter}$
32	$536.9 \mu s/\text{iter}$	$\pm 4.0 \mu s/{\rm iter}$

Table 4: Timing results for the third UPC implementation of the 1D Laplace solver, B=32

When ran with B=1, THREADS=32, the code takes $318\mu s/\text{iter} \pm 7.8\mu s/\text{iter}$. I do not know what causes this slowdown, but it seems like upc_forall could contribute to it.

When limiting the modification of ex4.upc to only replace the for loop with a upc_forall loop, the example becomes 83% slower ($206\mu s/\text{iter} \rightarrow 376\mu s/\text{iter}$, 8 threads, AMD Ryzen 2700X, upcc v2.28.2 + gcc v9.3.0).

1.4.1 Note on the update/copy loop

So far, the program has been running many iterations of the "main" loop, which calls iteration() and copy_array(). A upc_barrier; call is necessary at the end of each of those two operations, as copy_array() modifies x and depends on the processed value of x_{new} , while iteration() modifies x_{new} and depends on the copied value in x.

```
// settings.h:
#define BLOCKSIZE 32

// ex4.upc:
shared double x_new[LEN];
shared double x[LEN];
shared double x[LEN];
shared double b[LEN];
shared [BLOCKSIZE] double x[LEN];
shared [BLOCKSIZE] double b[LEN];
```

- (a) Previous array declarations
- (b) New array declarations with blocking factor

Figure 2: Modification of the array declarations to increase the blocking factor B.

```
// ex5.upc:
void iteration() {
    upc_forall (size_t i = 1; i < LEN - 1; i++; &x_new[i]) {
        x_new[i] = 0.5 * (x[i-1] + x[i+1] + b[i]);
    }

    if (MYTHREAD == 0) x_new[0] = x[0];
    if (MYTHREAD == THREADS - 1) x_new[LEN - 1] = x[LEN - 1];

    upc_barrier;
}

void copy_array() {
    upc_forall (size_t i = 0; i < LEN; i++; &x_new[i]) {
        x[i] = x_new[i];
    }
    upc_barrier;
}</pre>
```

Listing 3: New iteration() and copy_array() implementations

1.5 Detecting convergence

One of our last modifications is to measure $\delta_{max} = \max_{i \in [0; n-1]} |x[i] - x_{new}[i]|$ (named diffmax in the source code) and to stop once $\delta_{max} \leq \varepsilon$.

Because of the shared nature of x and x_{new} , we need to compute a partial δ_{max}^t and compute $\delta_{max} = \max_{t \in [0;T[}(\delta_{max}^t), \text{ with:}$

$$\delta_{max}^{t} = \max_{i \in [0; n-1], \text{ affinity}(x_{new}[i]) = t} \left| x[i] - x_{new}[i] \right|$$

In the implementation, we use one of the collective vector functions provided by UPC. This function contains implicit upc_barrier; instructions, so we do not need to add them ourselves.

The new timings are:

Threads	Time (avg)	$CI (\sigma = 0.01)$
2	$1400.4 \mu s/{ m iter}$	$\pm 6.0 \mu s/{\rm iter}$
4	$1092.8 \mu s/{ m iter}$	$\pm 17.0 \mu s/{\rm iter}$
8	$803.4 \mu s/{ m iter}$	$\pm 7.7 \mu s/\text{iter}$
16	$647.8 \mu s/{\rm iter}$	$\pm 18.9 \mu s/{\rm iter}$
32	$572.0 \mu s/{ m iter}$	$\pm 4.6 \mu s/{\rm iter}$

Table 5: Timing results for the fourth UPC implementation of the 1D Laplace solver, B=32, MAX_ITER = 1000

```
1 // ex6.upc (partial):
2 shared double diff[THREADS];
3 shared double diffmax;
5 void handle_diff(double d) {
       if (d < 0.0) d = -d;
6
       if (diff[MYTHREAD] < d) diff[MYTHREAD] = d;
  int main() {
10
       while (true) {
12
           iteration();
14
           upc barrier;
16
           if (MYTHREAD = 0) {
17
                for (size t n = 0; n < THREADS; n++) {
18
                    if (diffmax < diff[n]) diffmax = diff[n];</pre>
19
20
           }
21
22
           upc_barrier;
24
           if (diffmax <= EPSILON) break;</pre>
25
           if (++iter > MAX ITER) break;
26
           copy_array(); // upc_barrier; at the end of copy_array();
28
       }
29
30
31
```

```
32
33
void iteration() {
    upc_forall (size_t i = 1; i < LEN - 1; i++; &x_new[i]) {
        x_new[i] = 0.5 * (x[i-1] + x[i+1] + b[i]);
        handle_diff(x_new[i] - x[i]);
    }
}</pre>
```

Listing 4: Excerpt from ex6.upc: new handle_diff function and calculation of diffmax

1.5.1 Using reduction operations

We can further optimize this code by using reduction operations from upc_collective.h. The optimized variant is available at laplace/optimized.upc.

By removing the calls to upc_forall, we can reduce the time taken by a further $35\%(\pm 5\% pt)$.

Threads	Time (avg)	CI $(\sigma = 0.01)$
2	$1216.0 \mu s/{\rm iter}$	$\pm 27.1 \mu s/\text{iter}$
4	$778.0 \mu s/{ m iter}$	$\pm 15.1 \mu s/{\rm iter}$
8	$415.8 \mu s/{ m iter}$	$\pm 10 \mu s/\text{iter}$
16	$225.8 \mu s/{ m iter}$	$\pm 4.8 \mu s/\text{iter}$
32	$130.0 \mu s/{ m iter}$	$\pm 3.1 \mu s/\text{iter}$
64	$98.2 \mu s/{ m iter}$	$\pm 2.7 \mu s/\text{iter}$

Table 6: Timing results for the fully optimized UPC implementation of the 1D Laplace solver, B=32, MAX_ITER = 1000

1.6 Conclusion (1D Laplace Solver)

The first implementation in UPC of the 1D Laplace solver algorithm and its subsequent first optimization (flattening the for (...) if (...) gave very promising results for the improvement of the speed of the program, bringing the timings down from $140\mu s/\text{iteration}$ to only $46\mu s/\text{iteration}$.

Unfortunately, the next optimization attempt, which introduces a blocking factor to the primary arrays and used upc_forall, brought the speeds down and made the UPC implementation much slower than the C implementation.

The addition of δ_{max} , as a way to stop the algorithm once a sufficiently accurate solution is found, inevitably added another overhead to the program. With my best efforts, I could only bring its performance to somewhere between those of the first implementation and its first optimization.

Nonetheless, the resulting program runs slightly faster than the original C implementation, given enough threads).

2 2D Heat conduction

For this second algorithm, we will try to improve on the knowledge gathered from the first algorithm and implement a simple, 2D heat conduction simulation. As with Section 1, we begin with a simple C implementation of the algorithm, which will work as our performance base value. The provided template came bundled with a performance measurement method, but I swapped it out with clock() in the subsequent UPC implementations for consistency with the last section's code.

2.1 First C implementation

Following are the performance results for the first C implementation, ran on mesoshared with gcc v4.9.0 and on an AMD Ryzen 2700X with clang v12.0.1:

Options	Time (avg)	$CI (\sigma = 0.01)$
-00	$89.1 \mu s/{\rm iter}$	$\pm 0.3 \mu s/{\rm iter}$
-03	$15.0 \mu s/{ m iter}$	$\pm 0.1 \mu s/{\rm iter}$

Table 7: Timing results for heat_c.c (mesoshared, Listing 5)

Options	Time (avg)	$CI (\sigma = 0.01)$
-00	$80.9 \mu s/{ m iter}$	$\pm 2.6 \mu s/{\rm iter}$
-03	$17.9 \mu s/{\rm iter}$	$\pm 2.2 \mu s/{\rm iter}$

Table 8: Timing results for heat_c.c (Ryzen 2700X, Listing 5)

```
1 #include <stdio.h>
2 #include <math.h>
3 #include <sys/time.h>
4 #include <stdbool.h>
5 #include "settings.h"
  double grid [N+2][N+2], new grid [N+2][N+2];
  void initialize(void) {
9
      int j;
11
       /* Heat one side of the solid */
12
       for (j=1; j<N+1; j++) {
13
           grid[0][j] = 1.0;
14
           new_grid[0][j] = 1.0;
15
       }
17
18
  int main(void) {
19
       struct timeval ts_start, ts_end;
20
21
       double dTmax, dT, time;
22
       int i, j, k, l;
       bool finished = false;
23
       double T;
24
       int n iter = 0;
25
26
       initialize();
27
2.8
       /* Set the precision wanted */
```

```
finished = 0;
30
31
      /* and start the timed section */
32
      gettimeofday(&ts start, NULL);
33
34
      do {
35
           dTmax = 0.0;
36
           for (i=1; i< N+1; i++) {
37
               for (j=1; j<N+1; j++) {
38
                   T = 0.25 \ \ast
39
                        (grid[i+1][j] + grid[i-1][j] +
40
                         grid[i][j-1] + grid[i][j+1]); /* stencil */
41
                   dT = T - grid[i][j]; /* local variation */
42
                   new grid [i][j] = T;
43
                   if (dTmax < fabs(dT)) dTmax = fabs(dT); /* max variation in this
44
      iteration */
45
46
           if (dTmax < EPSILON) { // is the precision reached good enough ?
47
               finished = true;
48
           } else { // It isn't: preparing for a new iteration
49
               for (k=0; k<N+2; k++) {
                   for (1=0; 1<N+2; 1++) {
51
                        grid[k][1] = new_grid[k][1];
52
               }
54
           n_iter++;
56
      } while (!finished);
57
58
      gettimeofday(&ts end, NULL); /* end the timed section */
59
60
      /* compute the execution time */
      time = ts end.tv sec + (ts end.tv usec / 1000000.0);
62
      time = ts_start.tv_sec + (ts_start.tv_usec / 1000000.0);
63
64
      printf("%d iterations in %.31f sec \n", n_iter, time);
      printf("Took %.31f ms/iter\n", time * 1000.0 / n iter);
66
67
      return 0;
68
69
```

Listing 5: C implementation of the 2D Heat simulation

2.2 Porting the C code to UPC

Porting the template C version to UPC is straightforward. The performance, however, takes a hit, as the default work sharing causes a lot of remote accesses to memory, despite the blocking factor that was put in place. We obtain the following measurements on mesoshared:

Threads	Time (avg)	$CI (\sigma = 0.01)$
2	$259.9 \mu s/{ m iter}$	$\pm 1.9 \mu s/\text{iter}$
4	$193.6 \mu s/{ m iter}$	$\pm 1.5 \mu s/{\rm iter}$
8	$164.1 \mu s/{ m iter}$	$\pm 1.2 \mu s/{\rm iter}$
16	$150.6 \mu s/{ m iter}$	$\pm 1.4 \mu s/{\rm iter}$
32	$151.7 \mu s/{ m iter}$	$\pm 1.7 \mu s/{\rm iter}$

Table 9: Timing results for heat_1.upc

```
// heat_c.c:
for (i=1; i< N+1; i++) {
  for (j=1; j< N+1; j++) {
    T = 0.25 *
                                                 // heat 1.upc:
                                                 for (size_t i = 1; i \le N; i++) {
       ( grid [i+1][j] + grid [i-1][j] +
         grid[i][j-1] + grid[i][j+1];
                                                   upc\_forall (size_t j = 1; j <= N; j++;
                                                     &grid[i][j]) { double T = 0.25 * (grid[i+1][j] +
     stencil */
    dT = T - grid[i][j]; /* local
   variation */
                                                     \operatorname{grid}[i-1][j] + \operatorname{grid}[i][j-1] + \operatorname{grid}[i]
    new grid [i][j] = T;
                                                     [[j+1]];
    if (dTmax < fabs(dT)) dTmax = fabs(
                                                      double dT = fabs(T - grid[i][j]);
   dT); /* max variation in this
                                                     new grid [i][j] = T;
   iteration */
                                                      if (dTmax < dT) dTmax = dT;
                                                 }
```

(a) Main loop in heat_c.c

(b) Main loop in heat_1.upc

Figure 3: Porting the main loop from C to UPC

2.3 Optimizing the array accesses

One simple optimization is to avoid copying the destination array (new_grid) into the source array (grid).

To do this, we store two shared pointers that will point on either array, and we swap them at the end of each iteration, simulating a copy of grid into new_grid, with a $\mathcal{O}(1)$ time complexity instead of $\mathcal{O}(n)$. Doing so effectively halves the amount of synchronization that needs to happen for each loop, which can be observed in the timing results:

Threads	Time (avg)	$CI (\sigma = 0.01)$
2	$134.0 \mu s/{ m iter}$	$\pm 1.4 \mu s/{\rm iter}$
4	$103.9 \mu s/{ m iter}$	$\pm 4.1 \mu s/{\rm iter}$
8	$87.4 \mu s/{ m iter}$	$\pm 1.0 \mu s/{\rm iter}$
16	$81.4 \mu s/{ m iter}$	$\pm 1.2 \mu s/{\rm iter}$
32	$82.9 \mu s/{ m iter}$	$\pm 1.5 \mu s/{\rm iter}$

Table 10: Timing results for heat_3.upc

This optimization was also done on the C version for comparison, and we obtain the following speeds:

Options	Time (avg)	$CI (\sigma = 0.01)$
-00	$64.0 \mu s/{\rm iter}$	$\pm 0.5 \mu s/{\rm iter}$
-03	$11.3\mu s/\text{iter}$	$\pm 0.4 \mu s/\text{iter}$

Table 11: Timing results for heat_c_ptr.c

```
// heat_3.upc:
shared[BLOCKSIZE] double (*ptr)[N+2] = grid;
shared[BLOCKSIZE] double (*new_ptr)[N+2] = new_grid;

// ...

if (n_iter % 2 == 0) {
    ptr = grid;
    new_ptr = new_grid;
} else {
    ptr = new_grid;
    new_ptr = grid;
}

// ...

double T = 0.25 * (ptr[i+1][j] + ptr[i-1][j] + ptr[i][j-1] + ptr[i][j+1]);
double dT = fabs(T - ptr[i][j]);
new ptr[i][j] = T;
```

Listing 6: Excerpt from heat_3.upc, where pointer-swapping was implemented

2.4 Array privatization

As observed in the later examples of the 1D Laplace equation solver (Table 4), upc_forall has a significant performance cost over a regular for loop. We also know that local shared

memory accesses have a considerable cost when the shared pointer math is done without dedicated hardware support [1].

To make use of less shared accesses, we can copy a chunk of the *grid* array into private memory using upc_memget (into ptr_priv and new_ptr_priv). We specifically only retrieve the memory that has the affinity to the current thread, so that we can write to it and broadcast it in one go. The memory is copied also duplicated to make use of pointer-swapping as in Section 2.3.

We then operate on the private array, only using remote accesses when necessary, and store the results in the new private array. Once the work is finished, we put the new private array into new_grid with upc_memput and synchronize δ_{max} .

Threads	Time (avg)	$CI (\sigma = 0.01)$
2	$13.5 \mu s/{ m iter}$	$\pm 1.4 \mu s/{\rm iter}$
4	$12.6 \mu s/{ m iter}$	$\pm 1.2 \mu s/{\rm iter}$
8	$13.3 \mu s/{ m iter}$	$\pm 1.0 \mu s/{\rm iter}$
16	$15.0 \mu s/{ m iter}$	$\pm 1.2 \mu s/{\rm iter}$
32	$21.5 \mu s/{\rm iter}$	$\pm 1.1 \mu s/\text{iter}$

Table 12: Timing results for heat_4.upc (Listing 7)

```
1 #include <upc relaxed.h>
2 #include <bupc collectivev.h>
з #include <stdio.h>
4 #include <math.h>
5 #include <time.h>
6 #include <stdbool.h>
7 #include "settings.h"
9 #if ((N+2) % THREADS) != 0
10 #error N+2 must be divisible by THREADS
  #endif
12
  // Change blocksize to (N+2)^2 / THREADS?
13
  \#define BLOCKSIZE ((N+2) * (N+2) / THREADS)
  // \# define BLOCKSIZE ((N+2) * THREADS)
16 #define LOCALWIDTH ((N+2) / THREADS)
  \#define LOCALSIZE (LOCALWIDTH * sizeof(double) * (N+2))
17
  shared [BLOCKSIZE] double grid [N+2][N+2];
  shared [BLOCKSIZE] double new_grid [N+2][N+2];
20
  shared double dTmax[THREADS];
21
22
  void init() {
23
      for (size t j = 1; j < N+1; j++) {
24
           grid[0][j] = 1.0;
           new_grid[0][j] = 1.0;
26
27
28
29
30
  int main() {
31
      shared [BLOCKSIZE] double (*ptr)[N+2] = grid;
      shared [BLOCKSIZE] double (*new_ptr) [N+2] = new_grid;
32
      double (*ptr_priv)[N+2] = malloc(LOCALSIZE);
33
      double (*new_ptr_priv) [N+2] = malloc(LOCALSIZE);
34
35
       if (MYTHREAD = 0) {
36
           init();
37
```

```
39
         upc_barrier;
40
        upc_memget(ptr_priv, &grid[LOCALWIDTH * MYTHREAD], LOCALSIZE);
42
        upc_memget(new_ptr_priv, &new_grid[LOCALWIDTH * MYTHREAD], LOCALSIZE);
43
44
         bool finished = false;
45
         int n iter = 0;
46
         clock_t begin = clock();
        do {
49
              double dTmax = 0.0;
50
              size t o = LOCALWIDTH * MYTHREAD;
              size t i = 0;
53
54
              // Local block start
              if (i + o > 0) {
56
                         (\hspace{.05cm} \mathtt{size\_t} \hspace{.15cm} \mathtt{j} \hspace{.15cm} = \hspace{.15cm} \mathtt{1}; \hspace{.15cm} \mathtt{j} \hspace{.15cm} < \hspace{.15cm} \mathtt{N}; \hspace{.15cm} \mathtt{j} + \hspace{.15cm} +) \hspace{.15cm} \{
57
                         \begin{array}{lll} \textbf{double} \ T = \ 0.25 \ * \ (\ ptr\_priv [\ i+1][\ j\ ] \ + \ ptr[\ o+i-1][\ j\ ] \ + \ ptr\_priv [\ i\ ][ \end{array}
58
       j-1] + ptr_priv[i][j+1];
                          // printf("%zu+%zu %zu: %lf\n", o, i, j, T);
                         double dT = fabs(T - ptr_priv[i][j]);
60
                         new_ptr_priv[i][j] = T;
61
                          if (dTmax < dT) dTmax = dT;
                    }
63
              }
64
65
              // Local block middle
66
              for (i += 1; i < LOCALWIDTH - 1; i++) {
67
                    for (size_t j = 1; j \le N; j++) {
68
                         double T = 0.25 * (ptr_priv[i+1][j] + ptr_priv[i-1][j] + ptr_priv[i+1][j]
69
       i \mid \mid j-1 \mid + ptr_priv[i] \mid j+1 \mid j;
                          // printf("%zu+%zu %zu: %lf\n", o, i, j, T);
70
                         double dT = fabs(T - ptr_priv[i][j]);
71
                         new_ptr_priv[i][j] = T;
72
                          if (dTmax < dT) dTmax = dT;
                    }
              }
76
              // Local block end
              if (i + o < N + 1) {
78
                    for (size_t j = 1; j \le N; j++) {
79
                         \begin{array}{lll} \textbf{double} \ T = \ 0.25 \ * \ (\ ptr \ [o+i+1][j] \ + \ ptr \ \_priv \ [i-1][j] \ + \ ptr \ \_priv \ [i][ \end{array}]
80
       j-1 + ptr priv[i][j+1];
                           / printf("%zu+%zu %zu: %lf\n", o, i, j, T);
81
                         double dT = fabs(T - ptr_priv[i][j]);
82
                         new\_ptr\_priv[i][j] = T;
                          if (dTmax < dT) dTmax = dT;
                    }
85
              }
86
87
                  printf("%d: %lf\n", MYTHREAD, dTmax);
88
89
              // upc barrier;
90
              // Update ptr priv and new ptr priv
92
              \label{localwidth} \mbox{upc} \ \mbox{memput}(\&\mbox{new\_ptr}[\mbox{LOCALWIDTH} * \mbox{MYTHREAD}] \;, \; \mbox{new\_ptr\_priv} \;, \; \mbox{LOCALSIZE}) \;;
93
94
              // printf("\%lf \%lf \n", new_ptr_priv[1][1], new_ptr[o+1][1]);
95
96
```

```
// Implicit barrier here:
97
           double dTmax_g = bupc_allv_reduce_all(double, dTmax, UPC_MAX);
98
99
            if (dTmax g < EPSILON) {
100
                finished = true;
101
            } else {
102
                // Swap ptr and new ptr
103
                shared[BLOCKSIZE] double (*ptr_tmp)[N+2] = ptr;
104
                ptr = new_ptr;
105
106
                new_ptr = ptr_tmp;
107
                double (*ptr_tmp_priv)[N+2] = ptr_priv;
108
                ptr_priv = new_ptr_priv;
109
110
                new_ptr_priv = ptr_tmp_priv;
            }
111
           n iter++;
           // upc_barrier;
113
       } while (!finished);
114
       clock_t = end = clock();
115
116
       if (MYTHREAD = 0) {
117
118
           double seconds = (double)(end - begin) / CLOCKS_PER_SEC;
            printf("%d iterations in %.31f sec\n", n_iter, seconds);
119
            printf("Took %.31f ms/iter\n", seconds * 1000.0 / n_iter);
120
       }
121
122
```

Listing 7: Optimized UPC implementation of the 2D Heat algorithm (heat_4.upc)

References

[1] Olivier Serres. Hardware Support for Productive Partitioned Global Address Space (PGAS) Programming. PhD thesis, George Washington University, 2015.