

Parallelizing the CORAL LULESH microkernel

Goals:

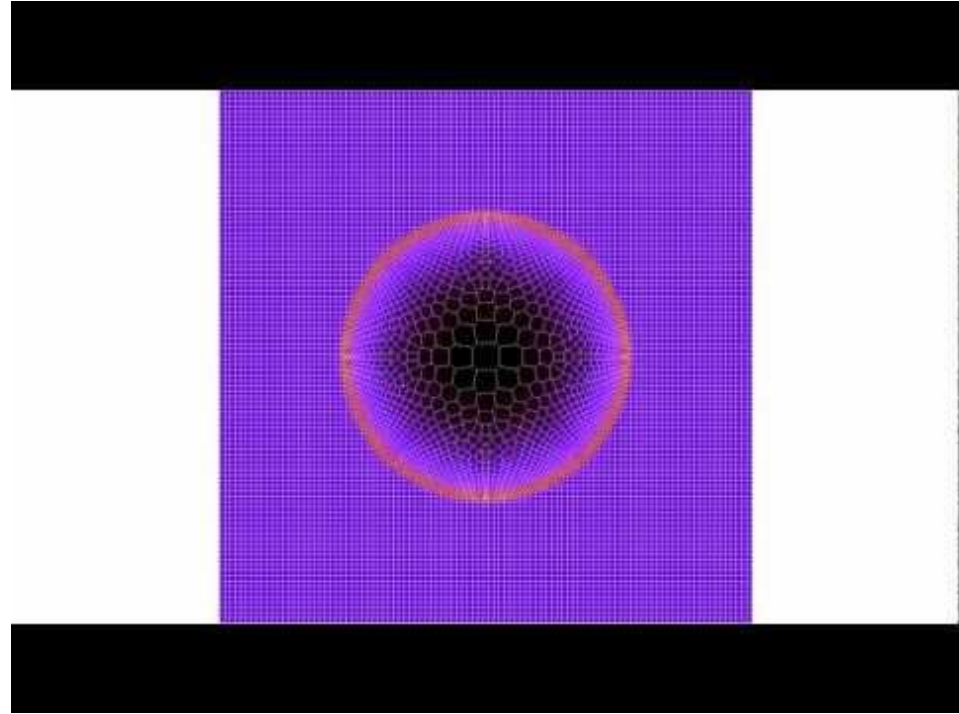
- Understand the code components
- Parallelize each component, including comparing performance of different implementations
- Combine parallel regions as required to improve performance

CORAL Benchmarks: LULESH

Livermore **U**nstructured **L**agrange
Explicit **S**hock **H**ydrodynamics

Part of a Physics Simulation
software (ALE3D)

Models the propagation of a Sedov blast
wave using Lagrangian hydrodynamics



Parallelware Trainer

Project Explorer

Code Editor

Version Manager

The screenshot displays the Parallelware Trainer IDE interface. On the left, the Project Explorer shows a project named 'pi' with files 'Makefile', 'pi.c', and 'README.md'. The main area contains two code editors. The left editor shows the source code for 'pi.c', which includes a main function and a 'getClock' helper function. The right editor shows the original source code for comparison. Below the code editors is the Output Console, which displays the results of a parallelization analysis. The console output indicates that a scalar reduction pattern was identified for the variable 'sum' and that the code was successfully parallelized using OpenMP directives. The analysis found 0 opportunities for further optimization.

```
File Edit Project Help
PI
  Makefile
  pi.c
  README.md
pi.c
35
36 out_result = 4.0 / N * sum;
37
38 // =====
39 double time_finish = getClock();
40
41 // Prints an execution report
42 printf("time (s)= %.6f\n", time_finish - time_start);
43 printf("result\t= %.8f\n", out_result);
44 const double realPiValue = 3.141592653589793238;
45 printf("error\t= %.1e\n", fabs(out_result - realPiValue));
46
47 return 0;
48 }
49
50 double getClock() {
51 #ifdef _OPENMP
52 #include <omp.h>
53 return omp_get_wtime();
54 #elif __linux__ || __APPLE__
55 #include <time.h>
56 struct timespec ts;
57 clock_gettime(CLOCK_MONOTONIC, &ts);
58 return ts.tv_sec + ts.tv_nsec / 1.0e9;
59 #else
60 #include <time.h>
61 return (double)clock() / CLOCKS_PER_SEC;
62 #endif
63 }
64
Original 1
32 out_result = 4.0 / N * sum;
33
34 // =====
35 double time_finish = getClock();
36
37 // Prints an execution report
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51 #include <time.h>
52 struct timespec ts;
53 clock_gettime(CLOCK_MONOTONIC, &ts);
54 return ts.tv_sec + ts.tv_nsec / 1.0e9;
55 #else
56 #include <time.h>
57 return (double)clock() / CLOCKS_PER_SEC;
58 #endif
59 }
60

[12:05:56] Parallelizing...
pi.c line 27: Parallel scalar_reduction pattern identified for variable 'sum' with associative, commutative operator '+'
pi.c line 27: Available parallelization strategies for variable 'sum'
pi.c line 27: #1 OpenMP scalar_reduction (* implemented)
pi.c line 27: #2 OpenMP atomic_access
pi.c line 27: #3 OpenMP explicit privatization
pi.c line 27: Loop parallelized with multithreading using OpenMP directive 'for'
pi.c line 27: Parallel_region defined by OpenMP directive 'parallel'
pi.c line 27: Make sure there is no aliasing among arguments in 'main': argc, argv

[12:05:56] Parallelization completed successfully

[12:05:57] Analysis completed: 0 opportunities found

Build output Execution output Parallelware output Analysis completed
```



Output Consoles

Run Parallelware Trainer on CORI

Step 1: Log in to CORI enabling X11 forwarding

```
$ ssh -X <your_login>@cori.nersc.gov
```

Step 2: Copy the sample codes to be used during the course

```
$ cp -a /global/common/cori/software/pwtrainer-1.2.0_linux-x64/examples.zip ~
```

Step 3: Prepare the environment by loading the appropriate modules

```
$ module purge && module load esslurm cuda/10.0 gcc/8.1.1-openacc-gcc-8-branch-20190215 pgi/19.4 pwtrainer
```

Step 4: Open an interactive session in a GPU node

```
$ salloc -t 60 -N 1 -c 16 -C gpu --gres=gpu:2 --mem=32GB
```



Step 5: Run the Parallelware Trainer tool from the GPU node

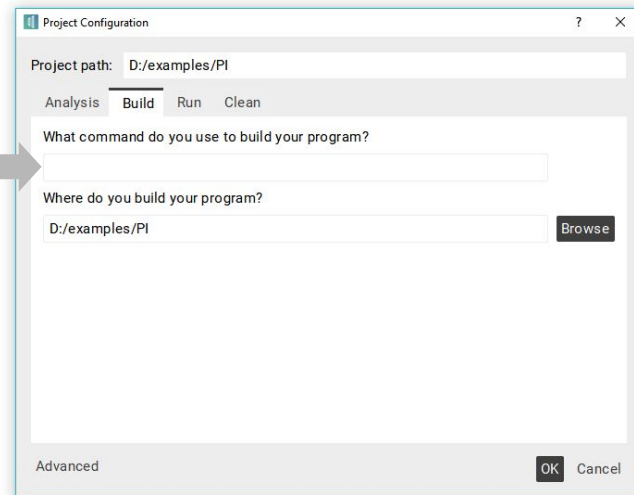
```
$ pwtrainer &
```

Step 6: Activate Parallelware Trainer

Click on 'Click here to install a license' and enter the following in the 'File name' text box:
`/global/common/cori/software/pwtrainer-1.2.0_linux-x64/pwtrainer-Jun19-NERSC.lic`



Build in Parallelware Trainer

		
GCC	<pre>gcc -fopenmp -foffload=nvptx-none="-Ofast -lm -misa=sm_35" -lm -o luleshmk luleshmk.c</pre>	<pre>gcc -fopenacc -foffload=nvptx-none="-Ofast -lm -misa=sm_35" -lm -D_OPENMP -o luleshmk luleshmk.c</pre>
PGI	<pre>pgcc -mp -ta=tesla:cc70,cuda10.0 -Minfo=accel -lm -o luleshmk luleshmk.c</pre>	<pre>pgcc -acc -ta=tesla:cc70,cuda10.0 -Minfo=accel -lm -D_OPENMP -o luleshmk luleshmk.c</pre>

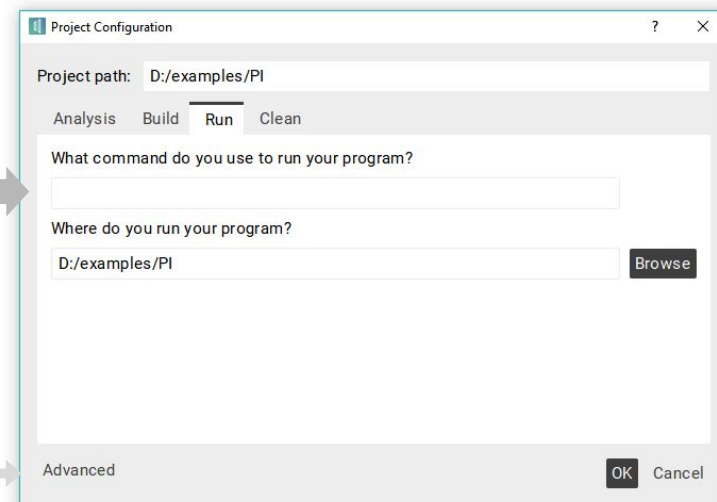


N.B. For Macs, check that your compiler supports OpenMP/OpenACC, and update with the appropriate compiler (e.g. gcc-mp-7)

Run in Parallelware Trainer

	<code>srun env OMP_NUM_THREADS=4 ./luleshmk</code>
	<code>srun env ACC_DEVICE_TYPE=nvidia ./luleshmk</code>

N.B. Instead of using the `env` command, you can add this variable by clicking the `Advanced` button.



Profiling of LULESH microkernel

```
$ gcc -pg -o luleshmk luleshmk.c -lm
$ ./luleshmk
$ gprof ./luleshmk
```

Flat profile:

Each sample counts as 0.01 seconds.

% time	cumulative seconds	self seconds	calls	self s/call	total s/call	name
56.23	24.62	24.62	223680000	0.00	0.00	CalcElemFBHourglassForce_workload
22.76	34.59	9.97	932	0.01	0.01	ApplyMaterialPropertiesForElems_workload
15.26	41.27	6.68	27960000	0.00	0.00	CalcElemVelocityGradient_workload
2.26	42.26	0.99	932	0.00	0.03	CalcFBHourglassForceForElems
2.24	43.24	0.98	27960000	0.00	0.00	CalcElemFBHourglassForce
0.75	43.57	0.33	27960000	0.00	0.00	CalcElemVelocityGradient
0.34	43.72	0.15	1	0.15	43.76	luleshmk
0.09	43.76	0.04	932	0.00	0.01	CalcKinematicsForElems
0.09	43.80	0.04				frame_dummy
0.00	43.80	0.00	932	0.00	0.01	ApplyMaterialPropertiesForElems
0.00	43.80	0.00	2	0.00	0.00	calculate_checksum
0.00	43.80	0.00	2	0.00	0.00	getClock
0.00	43.80	0.00	1	0.00	0.00	Parameters_create
0.00	43.80	0.00	1	0.00	0.00	Parameters_free
0.00	43.80	0.00	1	0.00	0.00	VerifyAndWriteFinalOutput

How to verify correctness of LULESH

```
$ ./luleshmk
```

```
- Configuring the test...
```

```
- Executing the test...
```

```
gprof ./luleshmk
```

```
- Verifying the test...
```

```
Run completed:
```

```
  Problem size          = 30
```

```
  MPI tasks             = 1
```

```
  Iteration count       = 932
```

```
  Final Origin Energy   = 1.000000e+00
```

```
  Number of nodes       = 27000
```

```
  Number of elements    = 30000
```

```
  Number of regions     = 1
```

```
    Region 1 of size 30000
```

```
  Testing Plane 0 of Energy Array on rank 0:
```

```
    MaxAbsDiff          = 8.410000e+02
```

```
    TotalAbsDiff        = 1.303550e+05
```

```
    MaxRelDiff          = 9.655568e-01
```

```
Elapsed time           = 71.00 (s)
```

```
Grind time (us/z/c)    = 2.821491 (per dom) ( 2.821491  
overall)
```

```
FOM                     = 354.42254 (z/s)
```


Decomposition of LULESH into patterns

Code file "luleshmk.c"		Pattern				
Function	Line	Forall		Scalar reduction	Sparse reduction	Convergence loop
CalcElemFBHourglassForce_workload()	60			sum		
CalcElemFBHourglassForce()	73	hgfx				
CalcFBHourglassForceForElems()	131				domain_m_fx	
ApplyMaterialPropertiesForElems_workload()	187					
ApplyMaterialPropertiesForElems()	210 217	-----				
CalcElemVelocityGradient_workload()	231					
CalcKinematicsForElems()	258					
luleshmk()	292					iter (loop index)
main()	349					
	358					
	365					
VerifyAndWriteFinalOutput()	485					