ALPSCore Tutorial

DRAFT Version 2

Alexander Gaenko 2/24/2017

ALPSCore tutorial: links

- These slides:
- https://git.io/alpscore-tut-2017-02-24

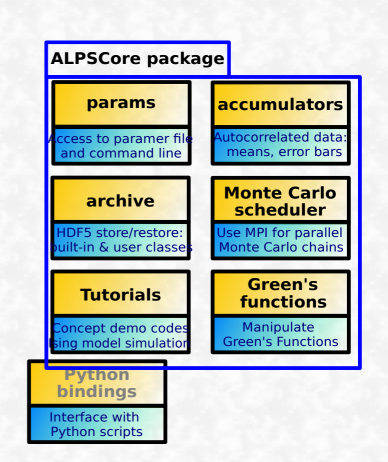
- Accompanying code:
- https://github.com/galexv/ALPSCore_Tutorial2

What is ALPSCore?

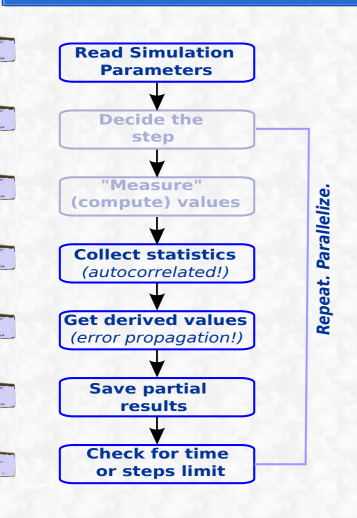
- Software library for writing physics simulation codes
- > Primarily for Monte Carlo methods (as of now).

However...

- > Has support for Green's Functions;
- > Support for lattices is planned.



Why use ALPSCore?



Look at a typical simulation:

- > A lot of "boilerplate" code;
- > Often non-trivial;
- > Performance is important!
- > Python (or other scripting): hard to combine with C++

ALPSCore helps to focus on science!

Brief history of ALPSCore

- > ALPSCore is the "core ALPS library"
- > ALPS is "Algorithms and Libraries for Physics Simulations" (see http://alps.comp-phys.org)
- > ALPSCore goals:
 - →Be more compact
 - → Faster development cycle
 - More extensive documentation.

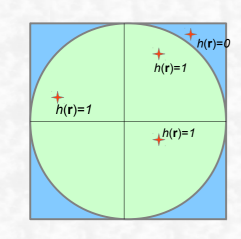
Focuses on the most often used components.

What is the plan for this tutorial?

- 1. Assume that ALPSCore is installed!
- 2.Make a very simple Monte Carlo simulation demo.
- 3.Parallelize the demo.
- 4.Make a 2D Ising model simulaton code.

Exercise 1: Preface

Task: Compute π by Monte Carlo method.



$$\frac{S_{cir}}{S_{sqr}} = \frac{\pi}{4}$$

"hit function" $h(r) = \begin{cases} 1 \text{ for } r \in \text{Cir} \\ 0 \text{ for } r \notin \text{Cir} \end{cases}$

$$\frac{S_{cir}}{S_{sqr}} = \frac{N_{cir}}{N_{sqr}} = \frac{1}{N} \sum_{i=1}^{N} h(\mathbf{r}_i) = \langle h \rangle$$

It is not hard to write the program to compute the mean <h>.
But...

What about error bars?

Do you really remember those formulas?

Exercise 1: Implementation specifics

```
Directory: step1 pi/
               specifies application structure
CMakeLists.txt
simulation.hpp
               describes simulation class (the model)
simulation.cpp
                implements the model
 constructor:
             constructs the model
 fraction_completed(): are we done yet?
                                             Let's look at the code!
   main.cpp
               sets up and runs the simulation
```

Exercise 1: Implementation specifics

```
simulation.cpp
```

main.cpp

Code uses parameters: to allow user to pass input data

Each parameter has:

- ✓ Name (e.g. "trials")
- ▼ Type (e.g. int)
- ✓ Description (e.g. "Number of trials")
- ✓ Optionally: a default value

define_parameters() is responsible for defining
 simulation-specific parameters

Let's look at the code!

Exercise 1: Hands-on

Task: Computing π by Monte Carlo method.

- 1. Build the program:
- \$ mkdir 000build
- \$ cd 000build
- \$ cmake ...
- \$ make
- 2. Run without arguments:
- \$./alpsdemo
- 3. Request help:
- \$./alpsdemo --help

- 4. Timed runs, varying trial count:
- \$ time -p ./alpsdemo --trials=100

- 5. Timed runs, varying time limits:
- \$ time -p ./alpsdemo \
 --trials=999999 \
 --timelimit=10
- 6. Start a long run; then press **Ctrl-C**; observe graceful termination.

Exercise 1: Hands-on, continued

Task: Use parameter file; play with the code.

```
7. Use parameter file:
$ ./alpsdemo ../demo.ini
```

- 8. Override params from file:
- \$./alpsdemo ../demo.ini \
 --trials=9999
- 9. Change the code: make --trials=0 mean "till timeout" (solution)

```
File "demo.ini":

# how long to run?
timelimit = 100

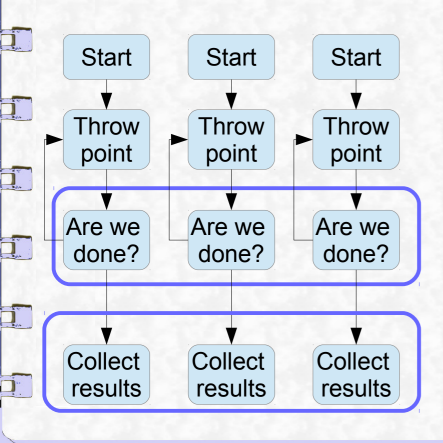
# how many trials?
trials = 1000
```

Exercise 1: Take-home message

- 1. Linking ALPSCore to your application is very simple.
- 2. Most of the framework is ready-made for you; you only code "science";
- 3. You get parameters, scheduling, graceful termination, and error bars all provided by the library.

Exercise 2: Preface

Point throwing can be done independently!



- Completion check requires occasional communications;
- Results are collected from all processes;
- > The rest is completely parallel!

Exercise 2: Hands-on

Task: Compute π in parallel.

- 1. Examine the differences (main.diff):
 - 1. MPI is initialized.
 - 2. A few operations are parallel-aware: params ctor simulation ctor
 - 3. An adapter class is used.
 - 4. Only master process should print.

Let's look at the code!

Directory:
step2_parallel_pi/

- 2. Build and run the program:
 - \$ mkdir 000build
 - \$ cd 000build
 - \$ cmake ...
 - \$ make
 - \$./alpsdemo --help

- 3. Do timed runs:
- \$ time -p mpiexec -n 1 \
 ./alpsdemo --trials=999999
- \$ time -p mpiexec -n 2 \
 ./alpsdemo --trials=999999

Exercise 2: Take-home message

- 1. Once you implemented serial MC, it's easy to make it parallel.
 - 2. The requested MC steps are distributed among the parallel processes.
 - 3. The processes occasionally communicate, to check if it's time to stop.

Exercise 3: Preface

Task: "Physics" problem: 2D Ising model.

Energy:
$$E = -\frac{1}{N} \sum_{(i,j)} s_i s_j$$

Magnetization:
$$M = \frac{1}{N} \sum_{i} s_{i}$$

Temperature
$$T$$
; $\beta = 1/T$
Boltzmann distribution: $p_i \sim e^-$

Mean values:
$$\langle E \rangle$$
, $\langle M \rangle$, $\langle |M| \rangle$, $\langle M^2 \rangle$, $\langle M^4 \rangle$

Binder Cumulant:
$$U=1-\frac{\langle M^4 \rangle}{3\langle M^2 \rangle^2}$$

Exercise 3: Approach

Task: "Physics" problem: 2D Ising model.

Directory: step3_ising/

Monte Carlo step:

- 1. Flip a random spin;
- 2. Update energy;
- 3. Update magnetization;
- 4. Accept or reject update.

Let's look at the code!

Problem: Measurements are correlated.

- 1. Naive approach: too narrow error bars;
- 2. Use LogBinningAccumulator for the correlated data;
- 3. Use FullBinningAccumulator if you need arithmetic.

Exercise 3: Hands-on

Task: Try different simulation parameters and accumulators.

- 1. Build and run the code
- 2. Use parameter file with different accumulators:

```
$ ./ising2_mc ../ising.ini --acc=fullbin
```

- \$./ising2 mc ../ising.ini --acc=nobin
- \$./ising2_mc ../ising.ini --acc=logbin

Look at energy and Binder Cumulant. Note different error bars!

- 3. Extra points: parallelize!
 - 1) Use mcmpiadapter 3) Pass communicator to:
 - 2) Initialize MPI params simulation constructors

4) Make sure only master prints.

Exercise 3: Take-home message

- 1. ALPSCore provides means to collect statistics properly;
 - 2. Autocorrelation results in too optimistic error bars;
- 3. Use binning accumulators (preferably FullBinning) for your Markov chain simulations.