```
#include "string.h"
        #include "spparks.h"
 SPPARKS
         #include "memory.h"
         #include "error.h"
         #include "universe.h"
        #include "input.h"
         #in ]
Getting
         #in
                   SPPARKS for microstructural science
         #in
         #include "potential.h"
         #include "random mars.h"
         #include "timer.h"
                                     Brian DeCost
         #include "output.h"
         using namespace SPPARKS NS;
                                     -11 July 2014
            allocate fundamental classes (memory, error, universe, input)
            parse input switches
            initialize communicators, screen & logfile output
            input is allocated at end after MPI info is setup
         SPPARKS::SPPARKS(int narg, char **arg, MPI_Comm communicator)
```

error = new Error(this):

1/21

Stochastic Parallel PARticle Kinetic Simulator¹

SPPARKS

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Running SPPARKS in parallel

SPPARKS is an efficient parallel Kinetic Monte Carlo framework

- ▷ Mesoscale materials phenomena as stochastic processes
- ▷ Scalable to thousands of processors
- ▶ Modular C++ facilitates user-defined models/statistics
- ▷ Open Source (GPLv2)
- ▷ Related to the LAMMPS Molecular Dynamics framework

¹http://spparks.sandia.gov/

How SPPARKS is used

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Running SPPARKS in parallel SPPARKS is a command-line application. Simulations are defined by input scripts composed of SPPARKS commands

The most important commands are:

- ▷ app_style
- ▷ solve_style / sweep_style
- ▷ diag_style

What applications is SPPARKS useful for?

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▷ lattice-based models

- ▷ off-lattice models

In this session, we'll focus on the Potts model

- ▶ Modeling grain growth, recrystallization, and related phenomena
- Annealing synthetic microstructures to achieve local equilibrium along triple lines

The SPPARKS distribution

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Download SPPARKS from

http://www.sandia.gov/~sjplimp/download.html

Unpack the SPPARKS source archive

mkdir -p ~/Software && cd ~/Software
tar xvzf ~/Downloads/spparks.tar.gz
mv spparks-19Nov13 spparks
cd spparks/src
ls ~/Software/spparks/

SPPARKS uses the Unix build system make

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Kunning SPPARKS in parallel SPPARKS is distributed with several makefiles, found in src/MAKE/.

```
bdecost@unix6 > ls MAKE :~/Software/spparks-19Nov13/src
Makefile.blackrose Makefile.mac Makefile.serial
Makefile.cygwin Makefile.mac_debug
Makefile.g++ Makefile.mac mpi Makefile.serial_debug
Makefile.linux Makefile.redsky Makefile.xt5
```

You will likely have to edit Makefile.linux, Makefile.g++, or Makefile.mac_mpi to reflect your system to get SPPARKS to compile.²

 $^{^2\}mbox{You'll}$ likely need to supply the location of the MPI library files on your system.

Building SPPARKS

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Running SPPARKS in parallel Try to compile SPPARKS by running make with the appropriate build target for your system as an argument.³

Building SPPARKS using Makefile.demo

make demo

The result is an executable named spk_demo in src/.

³To use the demo makefiles, first copy Makefile.demo and Makefile.demo_serial are in the SPPARKS src/MAKE directory.

Building serial SPPARKS

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Running SPPARKS in parallel SPPARKS comes with a placeholder MPI library for building a serial version.

First build the MPI stubs:

cd STUBS

make

cd ..

Build the demo_serial version of SPPARKS

make demo_serial

"Installing" SPPARKS⁴

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Kunning SPPARKS in parallel To avoid copying the SPPARKS executable or typing the full path to it repeatedly, define a *shell alias*:

alias spk=~/Software/spparks/src/spk_demo_serial

Now typing spk into the shell prompt from any working directory runs SPPARKS. Try this now.

⁴To make this persistent, append this command to ~/.bashrc (~/.bash_profile on a mac)

Running SPPARKS with no input

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Running SPPARKS in

```
SPPARKS reads commands from stdin

brian@rsfern > spk

SPPARKS (19 Nov 2013)

Press Ctrl-C to exit.
```

Demo: SPPARKS input scripts

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Running SPPARKS in parallel SPPARKS is controlled by a script of commands which are executed line-by-line.

Run the basic demo

cd ~/Downloads/spparks-demo-master/demos/basic
spk < basic.spkin</pre>

SPPARKS will print some diagnostic information to the screen, and to a log file called log.spparks.

SPPARKS output

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```
Created box = (0 \ 0 \ 0) to (32 \ 32 \ 32)
 32768 settings made for site
```

Anatomy of an input script: basic.spkin

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Running SPPARKS in parallel # Minimal SPPARKS Potts script

seed 56789

app_style potts 100

dimension 3

lattice sc/26n 1.0

region box block 0 32 0 32 0 32

create_box box
create_sites box

set site range 1 100

solve_style tree temperature 0.0 diag_style energy

stats 10.0 run 100.0

Demo: Grain growth kinetics

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Running SPPARKS in parallel

Run the grain-growth-kinetics demo

cd ../grain-growth-kinetics
spk < potts2d.spkin</pre>

This script uses the cluster diagnostic, which computes grain size information.

Grain growth kinetics

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Running SPPARKS ir parallel This demo includes a python script which extracts grain growth kinetics from log.spparks and attempts to plot them.

Run the python script

python parse_log.py log.spparks

The script produces a file cluster-stats.csv that you can open in Excel.

Grain growth kinetics

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Introduction

Getting started with SPPARKS

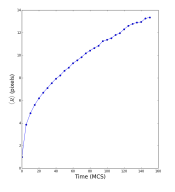
Demo: SPPARKS input scripts

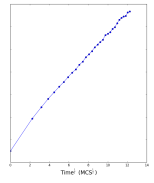
Demo: Grain growth kinetics

Demo: Visualization with

Running SPPARKS in parallel

Grain size is linear in sqrt-time!





The dump command

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Running SPPARKS in parallel $Post-processing.\ {\tt pizza.py}\ or\ custom\ scripts.$

Conversion to vtk with pizza.py

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Running SPPARKS ir parallel The python script pizza_dump2vtk.py uses tools from the open source pizza.py toolkit to create vtk files from the SPPARKS many-timestep dump file.

Run pizza_dump2vtk.py

python pizza_dump2vtk.py potts.dump

Visualizing with ParaView

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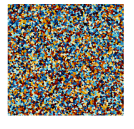
Demo: SPPARKS input scripts

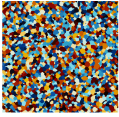
Demo: Grain growth kinetics

Demo: Visualization with ParaView

Running SPPARKS in

Open the vtk files in ParaView to create a movie.







mpirun

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Processor partitions

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