



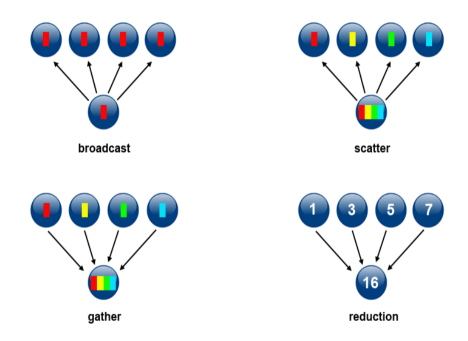
#### **Exercise sheet III**

Simon Scheidegger simon.scheidegger@gmail.com August 1<sup>st</sup>, 2017 Open Source Macroeconomics Laboratory – BFI/UChicago

Supplementary material for the exercises is provided in OSM\_Lab/HPC\_day3/supplementary\_material

Including adapted teaching material from of G. Hager & G. Wellein, B. Barney and the Swiss Supercomputing Centre (CSCS)

#### Reminder - collective communication



## 1. MPI exercise – Bcast

- Have a look at the code:
   OSM\_Lab/HPC\_day3/supplementary\_material/broadcast.f90 or broadcast.cpp
- write a makefile to compile the code.
- hard code a value (any double precision value), and broadcast the value of rank 0 to all Ranks.
- submit the job via slurm.

## 2. MPI exercise – Allreduce

- Have a look at the code:
   OSM\_Lab/HPC\_day3/supplementary\_material/allreduce.f90
   or allreduce.cpp
- write a makefile to compile the code.
- calculate the sum of all ranks.
- submit the job via slurm.

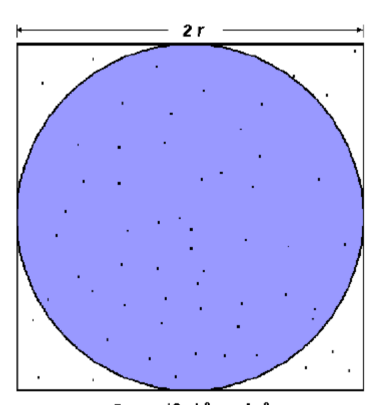
#### 3. MPI exercise – Scatter

- Have a look at the code:
   OSM\_Lab/HPC\_day3/supplementary\_material/scatter.f90
   or scatter.cpp
- write a makefile to compile the code.
- scatter the value of senddata of rank 0 to receivedata of all ranks.
- submit the job via slurm.

# 4. MPI Exercise (I)

Revisit the following method of approximating PI:

- 1. Inscribe a circle in a square.
- 2. Randomly generate points in the square.
- 3. Determine the number of points in the square that are also in the circle.
- 4. approximate PI.



$$A_S = (2r)^2 = 4r^2$$
  
 $A_C = \pi r^2$   
 $\pi = 4 \times \frac{A_C}{A_S}$ 

# MPI exercise (II)

Serial pseudo code for this procedure:

- → "embarrassingly parallel" solution:
- Break the loop iterations into chunks that can be executed by different tasks simultaneously.
- Each task executes its portion of the loop a number of times.
- Each task can do its work without requiring any information from the other tasks (there are no data dependencies).
- Master task receives results from other tasks using send/receive point-to-point operations.
- → red: highlights changes for parallelism.
- → Try to implement this example (probably using a reduction clause).

```
npoints = 10000
circle_count = 0

do j = 1,npoints
   generate 2 random numbers between 0 and 1
   xcoordinate = random1
   ycoordinate = random2
   if (xcoordinate, ycoordinate) inside circle then circle_count = circle_count + 1
end do

PI = 4.0*circle_count/npoints
```

```
npoints = 10000
circle count = 0
p = number of tasks
num = npoints/p
find out if I am MASTER or WORKER
do j = 1.num
  generate 2 random numbers between 0 and 1
  xcoordinate = random1
  vcoordinate = random2
  if (xcoordinate, ycoordinate) inside circle
  then circle count = circle count + 1
end do
if I am MASTER
  receive from WORKERS their circle counts
  compute PI (use MASTER and WORKER calculations)
else if I am WORKER
 send to MASTER circle count
endif
```

# 5. Work on Projects: American Options

- Parallelize the problem with MPI (for both Europen as well as the Asian option) (look at BS.cpp – take care of the random seeds!).
- Check the speed-up for combinations of threads and a variety of sample points on one node of MIDWAY (use slurm and the MPI\_Wtime()).
- Generate Speed-up graphs (normalize to the 1 CPU result).
- Ensure that the serial and parallel return the same results.

# 5. Pricing an Asian Option

The following algorithm illustrates the steps in simulating n paths of m transitions each. To be explicit, we use  $Z_{ij}$  to denote the j-th draw from the normal distribution along the i-th path:

```
for i = 1, ..., n

for j = 1, ..., m

generate Z_{ij}

set S_i(t_j) = S_i(t_{j-1}) \exp\left([r - \frac{1}{2}\sigma^2](t_j - t_{j-1}) + \sigma\sqrt{(t_j - t_{j-1})}Z_{ij}\right)

set \bar{S} = (S_i(t_1) + \cdots + S_i(t_m))/m

set C_i = e^{-rT}(\bar{S} - K)^+

set \hat{C}_n = (C_1 + \cdots + C_n)/n
```

#### 6. Project: Discrete State DP

- Parallelize the problem with MPI (look at the routine solver.cpp).
- Check the speed-up for combinations of threads and a variety of the discretization level on one node of MIDWAY (use slurm).
- Generate Speed up graphs (normalize to the 1 CPU result).
- Ensure that the serial and parallel return the same results.
- BONUS\* task Split the communicator (1 group per shock)

#### 6. Recall DSDP

$$V_{new}(k,\Theta) = \max_{c} \left( u(c) + \beta \mathbb{E} \{ V_{old}(k_{next}, \Theta_{next}) \} \right)$$

s.t. 
$$k_{next} = f(k, \Theta_{next}) - c$$

$$\Theta_{next} = g(\Theta)$$

#### States of the model:

- k : today's capital stock → **There are many independent k's**
- $\Theta$ : today's productivity state  $\rightarrow$  **The \Theta's are independent**

#### Choices of the model:

 $\rightarrow$  k,  $k_{next}$ ,  $\Theta$  and  $\Theta_{next}$  are limited to a finite number of values

# 6. solver.cpp

```
2). *split MPI communicator
for (int itheta=0; itheta<ntheta; itheta++) {
    Given the theta state, we now determine the new values and optimal policies corresponding to each
    capital state.
    */
                                                                                   1). distribute k's via MPI
    for (int ik=0; ik<nk; ik++) {
       // Compute the consumption quantities implied by each policy choice
                                                                                          loops to worry about
       c=f(kgrid(ik), thetagrid(itheta))-kgrid;
       // Compute the list of values implied implied by each policy choice
       temp=util(c) + beta*ValOld*p(thetagrid(itheta));
       /* Take the max of temp and store its location.
        The max is the new value corresponding to (ik, itheta).
        The location corresponds to the index of the optimal policy choice in kgrid.
       ValNew(ik, itheta)=temp.maxCoeff(&maxIndex);
       Policy(ik, itheta)=kgrid(maxIndex);
```

# 7. Project: Continuous State DP

- Parallelize the non-stochastic Growth model, implemented with TASMANIAN using MPI (distribute the sparse grid points equally among MPI processes).
- Check the speed-up for combinations of threads and a variety of the discretization level on one node of MIDWAY (use slurm).
- Generate Speed up graphs (normalize to the 1 CPU result).
- Ensure that the serial and parallel return the same results.
- BONUS\* task: parallelize the stochastic growth model.
   In particular, split the communicator (1 group per shock)

# 7. Recall VFI: there are many k's

$$V(\mathbf{k}) = \max_{\mathbf{I},\mathbf{c},\mathbf{l}} \left( u(c,l) + \beta \left\{ \underbrace{V_{next}(k^+)} \right\} \right),$$

$$s.t.$$

$$k_j^+ = (1 - \delta) \cdot k_j + I_j , \quad j = 1, ..., D$$

$$\Gamma_j = \frac{\zeta}{2} k_j \left( \frac{I_j}{k_j} - \delta \right)^2, \quad j = 1, ..., D$$

$$\sum_{j=1}^{D} (c_j + I_j - \delta \cdot k_j) = \sum_{j=1}^{D} (f(k_j, l_j) - \Gamma_j)$$

**State k:** sparse grid coordinates → **independent optimization problems** 

 $V_{next}$ : sparse grid interpolator from the previous iteration step

Solve this optimization problem at every point in the sparse grid!

Attention: Take care of the econ domain/ sparse grid domain

# 7. Growth model – Homework (II)

→ Add stochastic production to the model

$$f(k_i, l_i, \theta_i) = \theta_i A k_i^{\psi} l_i^{1-\psi}$$

- → Here we assume 5 possible values of  $\Theta_{i} = \{0.9, 0.95, 1.00, 1.05, 1.10\}$
- $\rightarrow$  for simplicity, we assume  $\Pi(*,*) = 1/5$  (no Markov chain)
- → solve

$$V_t(k,\theta) = \max_{c,l,I} u(c,l) + \beta \mathbb{E} \left\{ V_{t+1}(k^+,\theta^+) \mid \theta \right\}$$

 $\rightarrow$  Try to parallelize over the  $\Theta$ 's as well, they are independent