

# Introduction to High-Performance Computing

Session 08

Matlab Distributed Compute Server  
(MDCS)

## Questions

- Are you already using Matlab?
  - a) Yes, on my own/work computer
  - b) Yes, and I am already using the HPC cluster
  - c) No, not yet

## Questions

- Did you install Matlab on the computer you are using now?
  - a) Yes, Matlab R2019b
  - b) Yes, Matlab R2018b
  - c) Yes, another version
  - d) No

# Introduction to MDCS

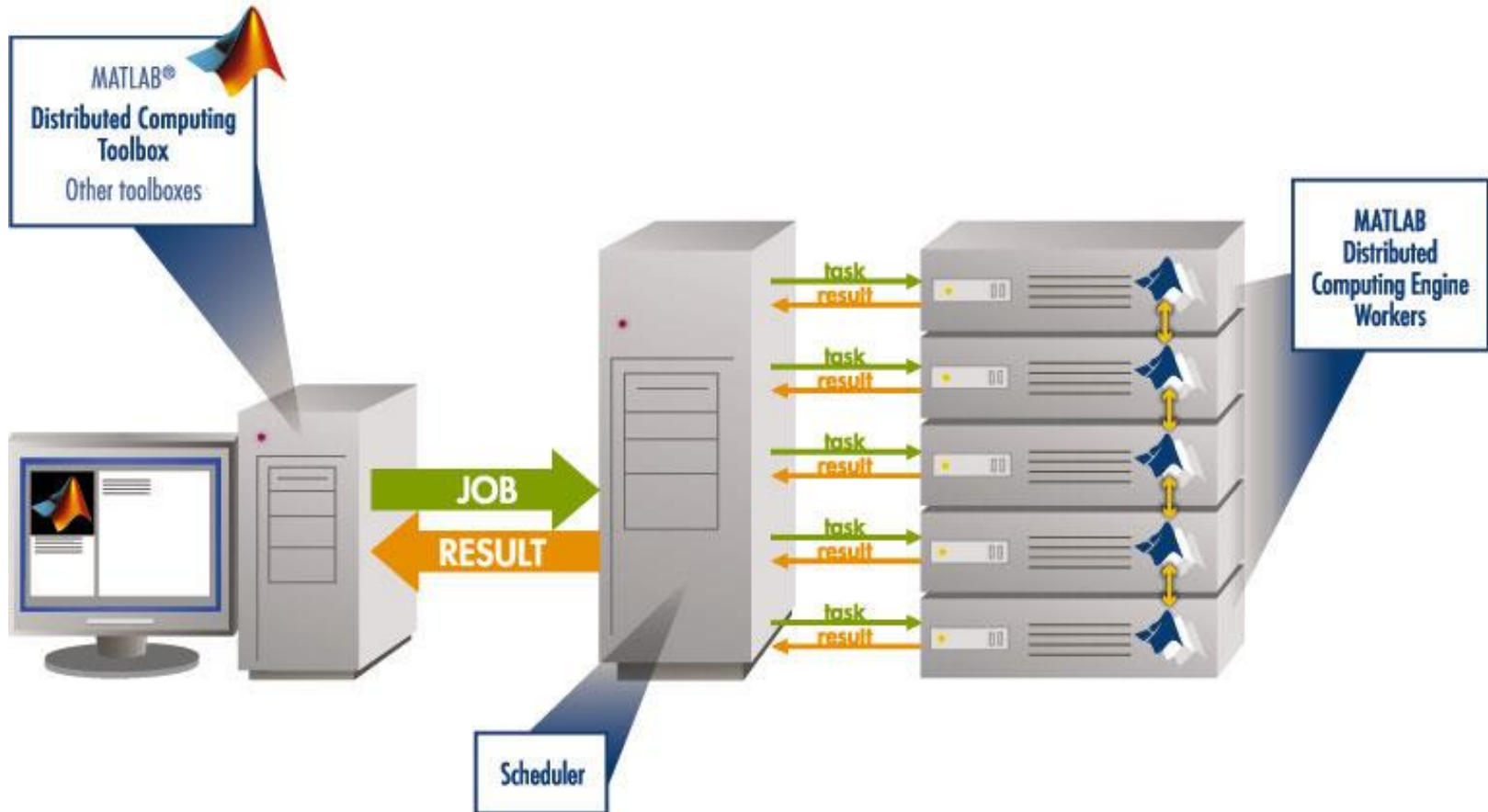
# What is MDCS

Matlab on your desktop computer:

- you are limited by the compute power of your local machine
  - memory
  - CPU speed
- you can only run one job at a time
- your machine may become unusable while your Matlab job is running

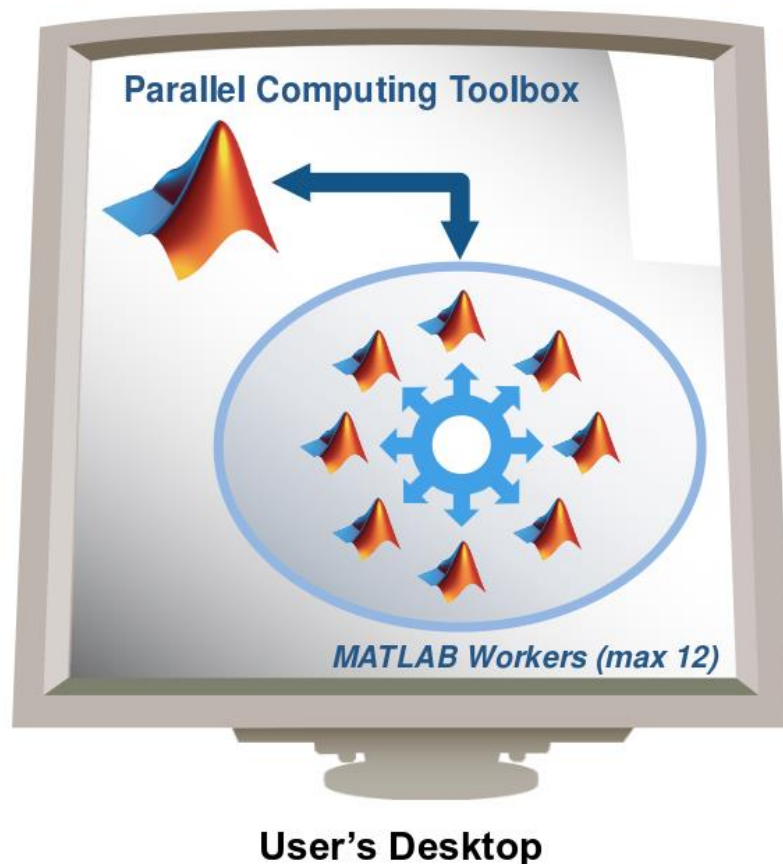


# What is MDCS



# Parallel Computing with Matlab

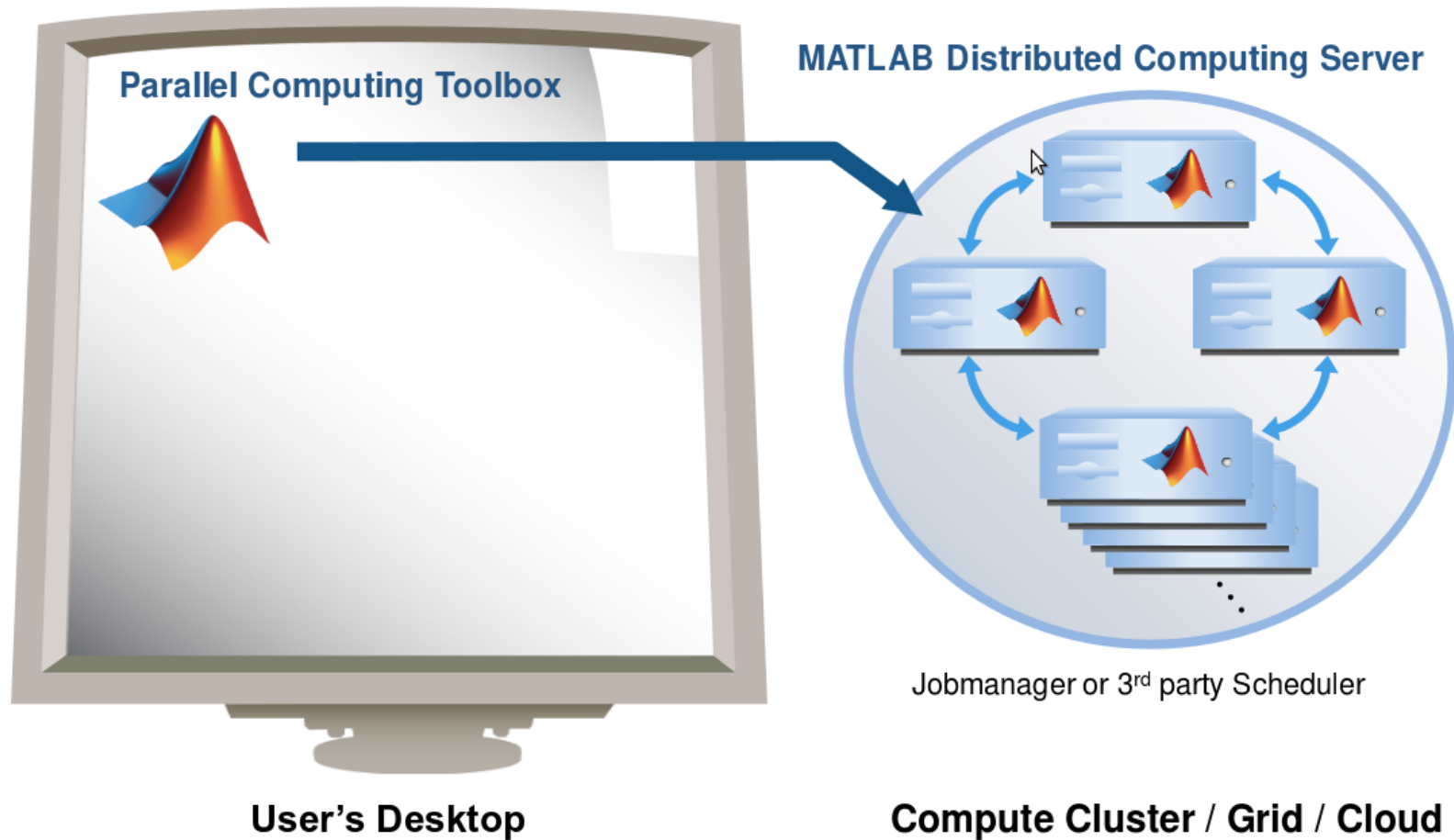
*(taken from MathWorks marketing)*



- easily experiment with explicit parallelism on multicore machines
- rapidly develop parallel applications on local computer
- take full advantage of desktop power, incl. GPUs
- separate compute cluster not required

# Parallel Computing with Matlab

*(taken from MathWorks marketing)*





## What is MDCS

- MDCS allows you to off-load Matlab programs to a compute server
- simplified workflow
  - you can develop and test your application locally before submitting jobs, also in parallel
  - results are automatically returned to your local machine for post-processing
- the Parallel Computing Toolbox provides utilities for parallelization
  - task-parallel
  - data-parallel

## Why to use MDCS on the Cluster?

- easy to use
  - work on your local computer within known Matlab environment
  - files (scripts, data, results) are automatically transferred
  - no need to learn about job scripts (but it helps to know a little)
- parallelization across multiple nodes
  - make use of distributed memory
  - use parallel threads (CPU cores) for each worker

```
>> maxNumCompThreads(1);      % set the number of threads to 1
>> a = rand(4096); b = rand(4096); % create two matrices
>> tic;c=a*b;toc              % compute and time matrix multiplication
Elapsed time is 3.633846 seconds.
>> maxNumCompThreads(4);      % set the number of threads to 4
>> tic;c=a*b;toc              % compute and time matrix multiplication
Elapsed time is 1.019613 seconds.
>> maxNumCompThreads('automatic'); % set the number of threads to automatic
>> tic;c=a*b;toc              % compute and time matrix multiplication
Elapsed time is 0.257363 seconds.
```

## MDCS Licenses

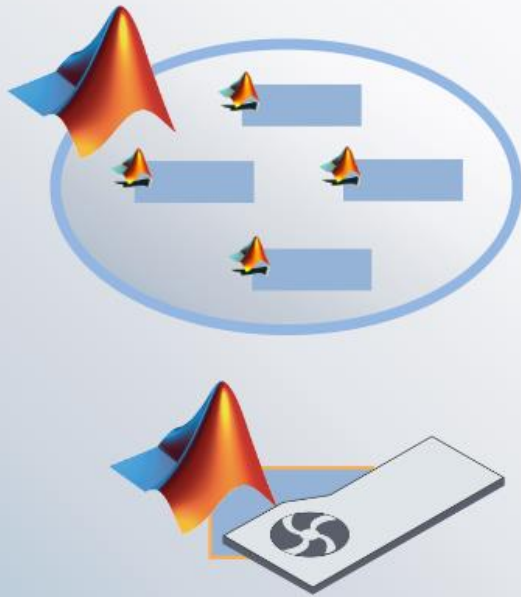
- MDCS on the HPC cluster includes 272 worker licenses
  - Matlab used to be limited to 200 licenses, now Campus license
  - for fair sharing not more than 36 MDCS licenses should be used per job and at most two jobs per user (hard limit)
  - check license use on the cluster:

```
[abcd1234@carl]$ scontrol show license mdcs  
LicenseName=mdcs  
Total=272 Used=47 Free=225 Remote=no
```

# Parallel Computing with Matlab

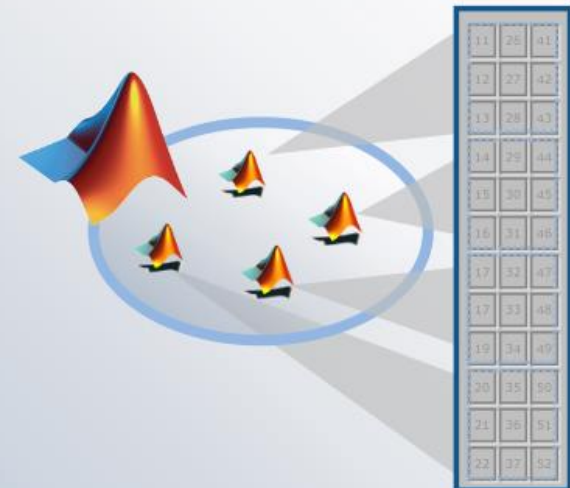
## Larger Compute Pool

Speed up Computations



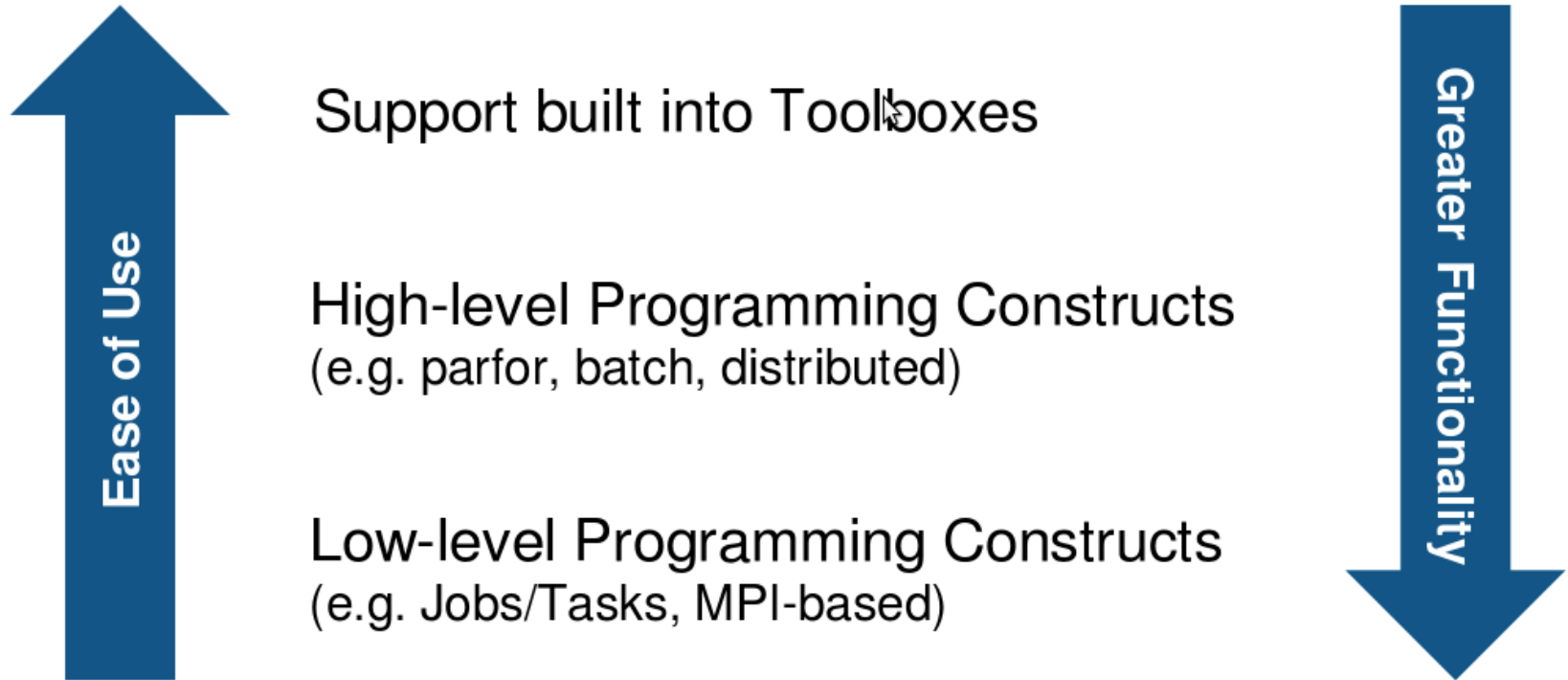
## Larger Memory Pool

Work with Large Data



# Parallel Computing with Matlab

## Three levels of Integration:



## Parallel Computing Support in Toolboxes

- Optimization Toolbox
- Global Optimization Toolbox
- Statistics Toolbox
- Simulink Design Optimization
- Bioinformatics Toolbox
- Communications Toolbox
- Model-Based Calibration Toolbox
- ... and more

see

<http://www.mathworks.com/products/parallel-computing/builtin-parallel-support.html>

# Configuration of MDCS

## Using MDCS on CARL/EDDY

- before you can use MDCS a few preparations are needed (**only needed to be done once**)
  - Matlab needs to be installed (see local web page) on your local machine, version must match to version on cluster (e.g. R2016b)
  - your local machine must be able to login to CARL/EDDY via ssh
    - Linux/Mac have ssh per default, for Windows you can use PuTTY
    - if you are not in the university network you also need to connect to a VPN (see HPC-Wiki for details)
  - a number of files (from a zipped archive from the HPC-Wiki) have to be copied to your local Matlab directory (depending on the setup of your local machine, your system admin has to help you)
  - a parallel configuration has to be setup with Matlab

see [https://wiki.hpcuser.uni-oldenburg.de/index.php?title=Configuration\\_MDCS\\_2016](https://wiki.hpcuser.uni-oldenburg.de/index.php?title=Configuration_MDCS_2016)



# Configuration of MDCS Cluster Profile

- the remote system is described in the cluster profile

**JobStorageLocation:** local directory for job data, e.g.  
`C:\Users\name\Documents\MATLAB\2019b\JobData`

**RemoteJobStorageLocation:** local directory for job data, e.g. on **\$WORK**  
`/gss/work/abcd1234/MATLAB/2019b/JobData`

- directories are sync'd at job submission and after the job has completed
- existing workspace is copied at job submission (can affect submission time)
- workspace of main process is copied back (can affect job load time), use e.g. `clear bigvar1 bigvar2;` (and save in separate files if needed)

## Configuration of MDCS Cluster Profile

- the remote system is described in the cluster profile

**JobStorageLocation:** local directory for job data, e.g.  
`C:\Users\name\Documents\MATLAB\2019b\JobData`

**RemoteJobStorageLocation:** local directory for job data, e.g. on `$WORK`  
`/gss/work/abcd1234/MATLAB/2019b/JobData`

**NumWorkers:** set to 36 for fair sharing

**NumThreads:** set to 1 (default), can be changed when useful

- change with e.g.: `sched.NumThreads=4;`
- maximum number of threads is the number of CPU cores in a node
- total number of cores allocated is `(worker+1)*NumThreads`
- benchmark your code to determine a good number of threads per worker.

## Configuration of MDCS Cluster Profile

- the remote system is described in the cluster profile

<b>JobStorageLocation:</b>	local directory for job data, e.g. C:\Users\name\Documents\MATLAB\2019b\JobData
<b>RemoteJobStorageLocation:</b>	remote directory for job data, e.g. on \$WORK /gss/work/abcd1234/MATLAB/2019b/JobData
<b>NumWorkers:</b>	set to 36 for fair sharing
<b>NumThreads:</b>	set to 1 (default), can be changed when useful
<b>AdditionalProperties:</b>	set at least <b>ClusterHost</b> and <b>RemoteJobStorageLocation</b> (see above), additional options for password-free login are described in HPC wiki

# Validation of MDCS Cluster Profile

**CARL** Type: Generic ([How to configure](#))

Properties Validation

	Stage	Status	Description
<input checked="" type="checkbox"/>	Cluster connection test (parcluster)	✓ Passed	
<input checked="" type="checkbox"/>	Job test (createJob)	✓ Passed	
<input checked="" type="checkbox"/>	SPMD job test (createCommunicatingJob)	✓ Passed	Job ran with 4 workers.
<input checked="" type="checkbox"/>	Pool job test (createCommunicatingJob)	✓ Passed	Job ran with 4 workers.
<input type="checkbox"/>	Parallel pool test (parpool)	⊘ Skipped	Not included in validation.

Number of workers to use:

STAGE DETAILS

Stage started at 15:38:53. Completed in 0 min 0 sec.

Validate Show Report

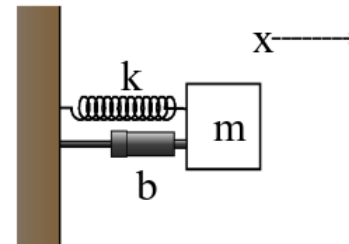
last test fails but that is not a problem (can be skipped)

recommended number of workers 4

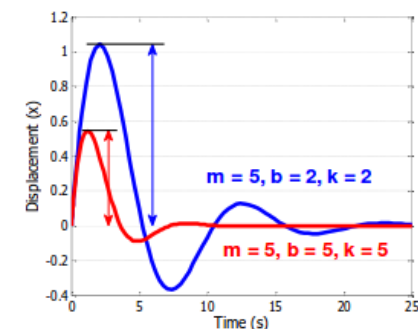
## Using MDCS on CARL/EDDY

- once you have completed the setup you can submit jobs to the cluster
  - example parameter sweep for 2<sup>nd</sup>-order ODE (taken from the [HPC-Wiki](#))
  - dampened oscillator

$$\underbrace{m}_{5} \ddot{x} + \underbrace{b}_{1,2,\dots} \dot{x} + \underbrace{k}_{1,2,\dots} x = 0$$



- simulate with different values for  $b$  and  $k$
- record peak value for each run



## 2<sup>nd</sup>-order ODE for example

odesystem.m

```
function dy = odesystem(t, y, m, b, k)
% 2nd-order ODE
%
%  $m \cdot X'' + b \cdot X' + k \cdot X = 0$ 
%
% --> system of 1st-order ODEs
%
%  $y = X'$ 
%  $y' = -1/m * (k \cdot y + b \cdot y')$ 
% Copyright 2009 The MathWorks, Inc.

dy(1) = y(2);
dy(2) = -1/m * (k * y(1) + b * y(2));

dy = dy(:); % convert to column vector
```

# Parameter Sweep: serial Matlab code

paramSweep\_batch.m

```
%% Initialize Problem
m      =      5; % mass
bVals  = 0.1:.1:15; % damping values (step .1)
kVals  = 1.5:.1:15; % stiffness values (step .1) damping
[kGrid, bGrid] = meshgrid(bVals, kVals);
peakVals = nan(size(kGrid));

%% Parameter Sweep
tic;

for idx = 1:numel(kGrid)
    % Solve ODE
    [T,Y] = ode45(@(t,y) odesystem(t, y, m, bGrid(idx), kGrid(idx)), ...
        [0, 25], ... % simulate for 25 seconds
        [0, 1]); % initial conditions

    % Determine peak value
    peakVals(idx) = max(Y(:,1));
end

t1 = toc;
```

# Parameter Sweep: parallel Matlab code

paramSweep\_batch.m

```
%% Initialize Problem
m      =      5; % mass
bVals = 0.1:.1:15; % damping values (step .1)
kVals = 1.5:.1:15; % stiffness values (step .1) damping
[kGrid, bGrid] = meshgrid(bVals, kVals);
peakVals = nan(size(kGrid));

%% Parameter Sweep
tic;

parfor idx = 1:numel(kGrid)
    % Solve ODE
    [T,Y] = ode45(@(t,y) odesystem(t, y, m, bGrid(idx), kGrid(idx)), ...
        [0, 25], ... % simulate for 25 seconds
        [0, 1]); % initial conditions

    % Determine peak value
    peakVals(idx) = max(Y(:,1));
end

t1 = toc;
```



## Using MDCS on CARL/EDDY

- submitting jobs to the cluster

```
 sched = parcluster('CARL');  
 job = batch(sched, 'paramSweep_batch', 'Pool', 7, ...  
            'AttachedFiles', {'odesystem.m'});
```

- first command creates a handle for the cluster using the available configuration
- second command creates a job and sends it to the cluster
  - Matlab script is executed on the cluster
  - requests a pool of workers (number of processes is +1 for master)
  - uses default resources unless modified
  - files can be attached but Matlab also automatically attaches needed files (if it can find them and if not disabled)

## Using MDCS on CARL/EDDY

- changing resource allocation

```
 sched.AdditionalProperties.runtime='0:30:00' ;  
 sched.AdditionalProperties.memory='4G' ;  
 remove(sched.AdditionalProperties, 'memory') ;
```

- changes maximum runtime and memory per worker
  - remove previous setting to get default
  - older Matlab versions use a different format (see HPC wiki)
- path-dependency as alternative to attaching files
    - use `addpath` within script (.m-files)
    - use `AdditionalPath` property of scheduler object
    - use absolute path names
    - copy files to the cluster before submitting job

## Using MDCS on CARL/EDDY

- recovering jobs
  - it is possible to terminate the local Matlab session while jobs are running (or waiting on the cluster)
  - to reconnect

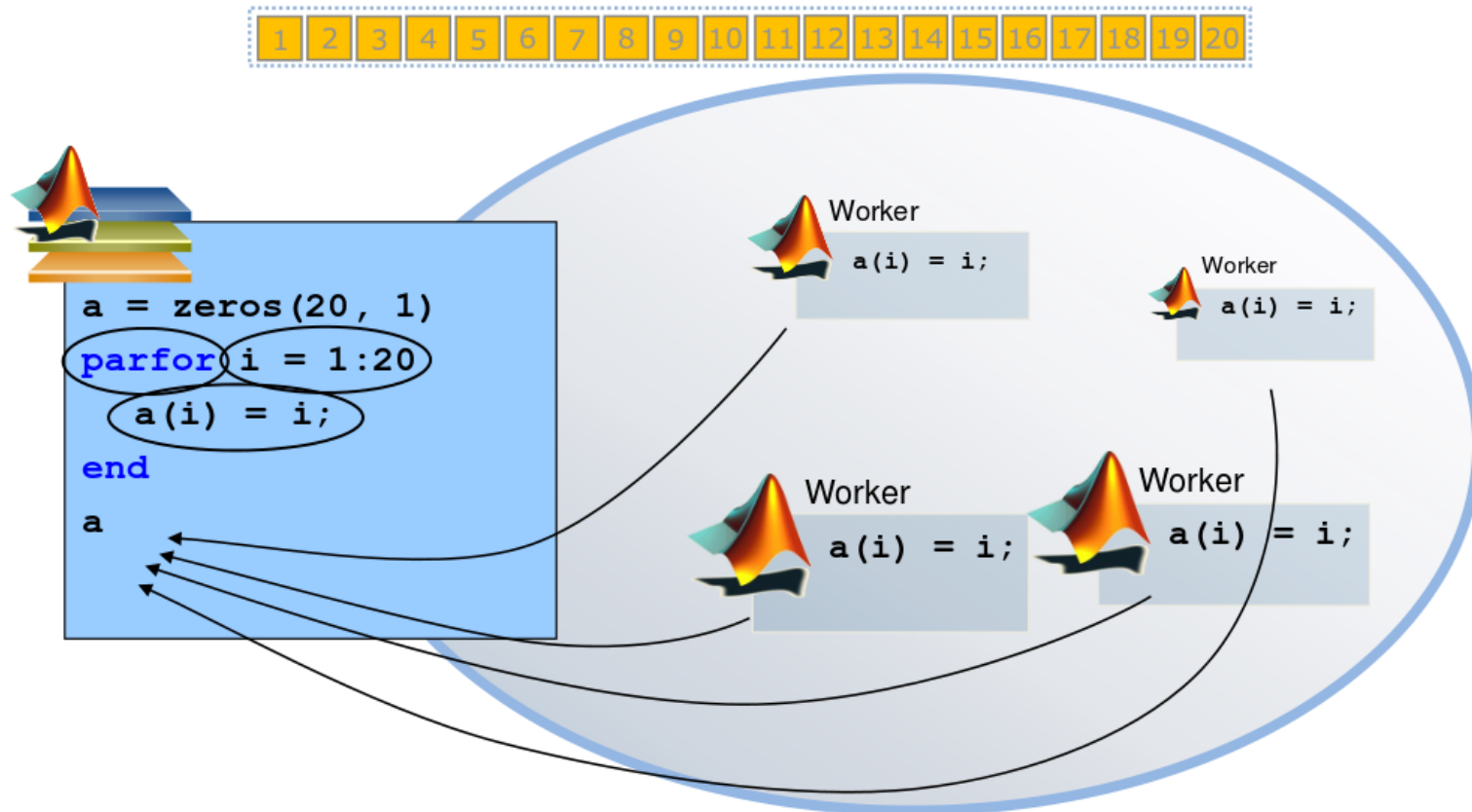
```
sched = parcluster(,CARL');  
sched.Jobs % to list available jobs  
job = sched.Jobs(1) % to get job information  
jobData = load(job);
```

## Monitoring Jobs and Error Tracking

- Matlab Job Monitor for basic information
- use `squeue` and `sacct` for additional information from SLURM
- job handle can be used to get information about errors
- Matlab diary for additional log output
- files in the job directory

# MDCS with parfor

# Mechanics of `parfor` Loops



Pool of MATLAB Workers

## Converting `for` to `parfor`

- requirements for `parfor` loops
  - task independent
  - order independent
- constraints on the loop body
  - cannot introduce variables (e.g. `eval`, `load`, `global`)
  - cannot contain `break` or `return` statements
  - cannot contain another `parfor` loop

<https://de.mathworks.com/help/parallel-computing/troubleshoot-variables-in-parfor-loops.html>

## Variable Classification

- all variables referenced at the top level of the **parfor** must be resolved and classified

Classification	Description
loop	serves as a loop index for arrays
sliced	an array whose segments are operated on by different iterations
broadcast	a variable defined before the loop whose value is used inside the loop, but never assigned in the loop
reduction	accumulates a value across iterations of the loop, regardless of iteration order
temporary	variable created inside the loop but unlike sliced or reduction variables, not available outside the loop



# Variable Classification Example

- matrix-vector multiplication

```
N=2048 ;                                % N is broadcast  
b=rand(N,1) ;                            % b is broadcast  
A=rand(N,N) ;                            % A is slices input  
  
parfor i=1:N                              % i is loop index  
    c(i)=A(i,:) * b(:) ;                  % c is sliced output  
end
```

## parfor Examples

- this example cannot be parallized in **parfor**

```
j=zeros(100);      %pre-allocate vector  
j(1)=5;  
for i=2:100;  
    j(i)=j(i-1)+5;  
end;
```

- order of iterations is important

## parfor Examples

- functions with multiple output may confuse Matlab

```
for i=1:10
    [x{i}(:,1), x{i}(:,2)]=functionName(z,w);
end;
```

- use this instead

```
for i=1:10
    [x1, x2]=functionName(z,w);
    x{i}=[x1 x2];
end;
```

## parfor Examples

- be careful not to broadcast unnecessary data

```
data.raw = ...  
data.processed = ...  
  
% Inefficient variant:  
parfor idx = 1 : N  
    % do something with data.processed  
end  
  
% This is better:  
processedData = data.processed;  
parfor idx = 1 : N  
    % do something with processedData  
end
```

<https://undocumentedmatlab.com/blog/a-few-parfor-tips>

## parfor Considerations

- **parfor** often only involves minimal code changes
- if a for loop cannot be converted to **parfor**, consider wrapping a subset of loop body in a function
  - e.g. `load` works not in **parfor**, however it does work in function that is called inside a **parfor** loop
- more information  
<http://blogs.mathworks.com/loren/2009/10/02/using-parfor-loops-getting-up-and-running/>
- there is a Code-Analyzer to diagnose **parfor** issues

# MDCS with spmd (single program multiple data)

# SPMD

	Client			Worker 1			Worker 2				
	a	b	e		c	d	f		c	d	f
a = 3;	3	-	-		-	-	-		-	-	-
b = 4;	3	4	-		-	-	-		-	-	-
spmd											
c = labindex();	3	4	-		1	-	-		2	-	-
d = c + a;	3	4	-		1	4	-		2	5	-
end											
e = a + d{1};	3	4	7		1	4	-		2	5	-
c{2} = 5;	3	4	7		1	4	-		5	6	-
spmd											
f = c * b;	3	4	7		1	4	4		5	6	20
end											

## SPMD

- when a SPMD block ends the workspace is saved, the worker is paused
- data is preserved from one block to the next
- does not apply to SPMD block in a function after the function is completed (as regular variables local to a function)



## SPMD Example

```
x = imread ( 'balloons.tif' );  
y = imnoise ( x, 'salt & pepper', 0.30 );  
yd = distributed ( y );  
  
spmd  
    yl = getLocalPart ( yd );  
    yl = medfilt2 ( yl, [ 3, 3 ] );  
end  
  
z(1:480,1:640,1) = yl{1};  
z(1:480,1:640,2) = yl{2};  
z(1:480,1:640,3) = yl{3};
```

- read image
- add noise to image
- distribute data
- parallel working on image data (filter)
- on master process put together filtered image



## Distributed Data

- Matlab provides different functions to manage distributed data
  - with `distributed(X)` you can distribute data among workers
  - with `distributed.METHOD` you can create data distributed among workers
  - workers can create codistributed data structures which become distributed data outside of the SPMD block
  - a datastore can be distributed to read manage large data files with multiple workers
  - see ‘help distributed’ for more information

## Distributed Data

distributing data  
from client

```
p = parpool('local', 4); % create a local pool of workers
A = zeros(4);           % create a 4x4 matrix with zeros
A                        % print A on client
B = distributed(A);     % distribute A to the workers
spmd                    % begin parallel spmd region
    B = B + labindex;   % modify distributed data in B
end                      % end parallel spmd region
B                        % print B on client
delete(p);
```

vs.

codistributed data  
created on workers

```
p = parpool('local', 4); % create a pool of workers
spmd                      % begin parallel spmd region
    codist = codistributor(2, [1,1,1,1]); % define distribution
    B = zeros(4, codist); % created codistributed array
    B = B + labindex;     % modify distributed data in B
end                        % end parallel spmd region
B                          % print B on client
delete(p);
```

## Example: Image Contrast

- a Matlab script that uses a simple function to change the contrast of an gray-scale image

```
% read an image (gray-scale)
y = imread('low_contrast.jpg');

% setup function for contrast manipulation
c = 1.7;
adjustContrast = @(x) c*x(2,2)+(1.0-c)*(mean(x(:)-x(2,2)/9.0));

% apply filter
z = nlfilter(y, [3,3], adjustContrast);

% save image side-by side
imwrite(cat(1,y,z), 'contrast_serial.jpg');
```



## Example: Image Contrast

- parallelize with SPMD

```

% read an image (gray-scale)
y = imread('low_contrast.jpg');

% setup function for contrast manipulation
c = 1.7;
adjustContrast = @(x) c*x(2,2)+(1.0-c)*(mean(x(:))-x(2,2)/9.0);

% distribute image by columns
yd = distributed(y);

% now work in parallel
spmd
    y1 = getLocalPart(yd);

    % apply filter
    y1 = nlfilter(y1, [3,3], adjustContrast);
end

% combine local images
z = [ y1{:}];

% save image side-by-side
imwrite(cat(1,y,z), 'contrast_spmd.jpg');

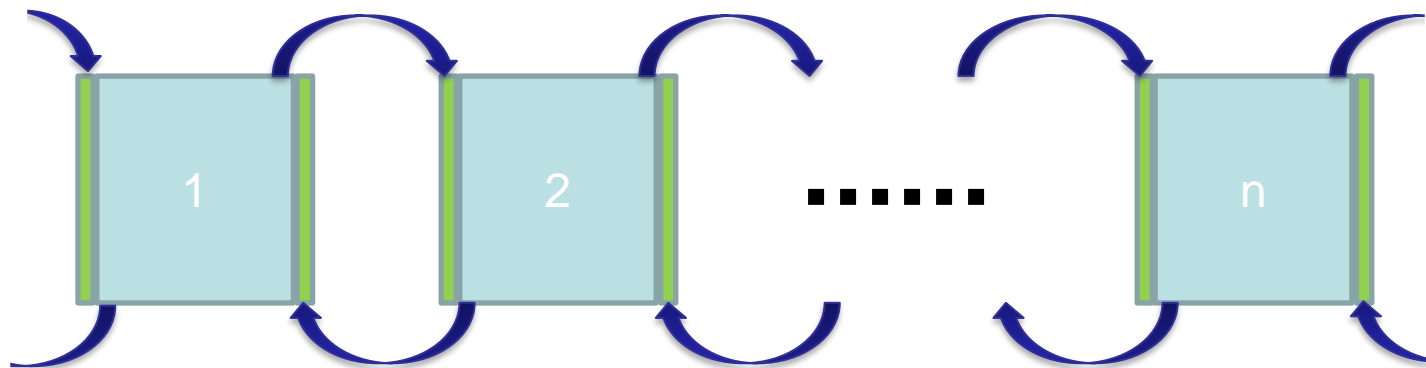
```

- algorithm produces artifacts when parallelized on multiple workers
  - problem is that increasing contrast requires information from neighbouring pixel
  - distributing the data adds additional boundaries



## labSendReceive

- solution is communication between workers
  - each worker has to send one boundary left and one right
  - each worker has to receive one boundary from left and one from right
  - extra columns are added before filter is applied, and need to be removed again afterwards



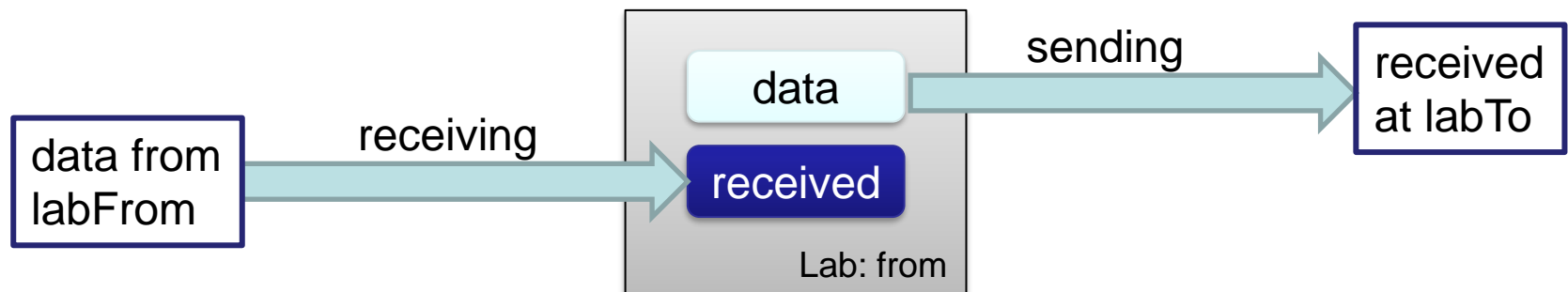


## labSendReceive

- the function `labSendReceive` simultaneously sends and receives data

```
received = labSendReceive(labTo, labFrom, data)
```

- sends data to `labTo`
- receives data from `labFrom` and stores it in `received`



## labSendReceive

```
column = labSendReceive ( previous, next, x1(:,1) );  
  
if ( labindex() < numlabs() )  
    x1 = [ x1, column ];  
end  
  
column = labSendReceive ( next, previous, x1(:,end - 1) );  
  
if ( 1 < labindex() )  
    x1 = [ column, x1 ];  
end
```



## Exercise: Heat Example in Matlab

```
% 2d-heat example in Matlab
% initial setup
NXPROB = 20;           % number of grid rows
NYPROB = 20;           % number of grid columns
STEPS   = 100;         % number of iterations
TIME    = 0;           % initial and current time

uvals = zeros(2, NXPROB, NYPROB); % allocate grid
uvals = inidat(uvals);           % initialize grid

plotdat(uvals, 1, TIME);         % make plot

it = 1;
for TIME=1:STEPS                 % time iteration
    uvals = updateu(uvals, it);   % update thermal energy
    it     = 3 - it;
end

plotdat(uvals, 1, TIME);         % make plot
```