Assuming you know basic MPI

- This is a rare group that can discuss this topic meaningfully.

- I have mentioned MPI 3.0’s “improvements” to its hybrid capabilities. These are primarily tying up loose ends and formally specifying that things work as you would expect, and as they largely do. Your MPI 1/2 knowledge will be more than sufficient here.
#pragma acc data copy(Temperature_last), create(Temperature)
while (dt_global > MAX_TEMP_ERROR && iteration <= max_iterations) {
    #pragma acc kernels
    for(i = 1; i <= ROWS; i++) {
        for(j = 1; j <= COLUMNS; j++) {
            Temperature[i][j] = 0.25 * (Temperature_last[i+1][j] + Temperature_last[i-1][j] + Temperature_last[i][j+1] + Temperature_last[i][j-1]);
        }
    }
    if(my_PE_num != npes-1) {
        MPI_Send(&Temperature[ROWS][1], COLUMNS, MPI_DOUBLE, my_PE_num+1, DOWN, MPI_COMM_WORLD);
    }
    if(my_PE_num != 0) {
        MPI_Recv(&Temperature_last[0][1], COLUMNS, MPI_DOUBLE, my_PE_num-1, DOWN, MPI_COMM_WORLD, &status);
    }
    if(my_PE_num != 0) {
        MPI_Send(&Temperature[1][1], COLUMNS, MPI_DOUBLE, my_PE_num-1, UP, MPI_COMM_WORLD);
    }
    if(my_PE_num != npes-1) {
        MPI_Recv(&Temperature_last[ROWS+1][1], COLUMNS, MPI_DOUBLE, my_PE_num+1, UP, MPI_COMM_WORLD, &status);
    }
    dt = 0.0;
    #pragma acc kernels
    for(i = 1; i <= ROWS; i++) {
        for(j = 1; j <= COLUMNS; j++) {
            dt = fmax(fabs(Temperature[i][j]-Temperature_last[i][j]), dt);
        }
    }
    MPI_Reduce(&dt, &dt_global, 1, MPI_DOUBLE, MPI_MAX, 0, MPI_COMM_WORLD);
    MPI_Bcast(&dt_global, 1, MPI_DOUBLE, 0, MPI_COMM_WORLD);
    if((iteration % 100) == 0) {
        if (my_PE_num == npes-1) {
            #pragma acc update host(Temperature)
            track_progress(iteration);
        }
        iteration++;
    }
}
Hybrid OpenACC Programming (Fast & Wrong)

```c
#pragma acc data copy(Temperature_last), create(Temperature)
while (dt_global > MAX_TEMP_ERROR && iteration <= max_iterations) {

#pragma acc kernels
for(i = 1; i <= ROWS; i++) {
    for(j = 1; j <= COLUMNS; j++) {
        Temperature[i][j] = 0.25 * (Temperature_last[i+1][j] + Temperature_last[i-1][j] + Temperature_last[i][j+1] + Temperature_last[i][j-1]);
    }
}

if(my_PE_num != npes-1){
    MPI_Send(&Temperature[ROWS][1], COLUMNS, MPI_DOUBLE, my_PE_num+1, DOWN, MPI_COMM_WORLD);
}

if(my_PE_num != 0){
    MPI_Recv(&Temperature_last[0][1], COLUMNS, MPI_DOUBLE, my_PE_num-1, DOWN, MPI_COMM_WORLD, &status);
}

if(my_PE_num != 0){
    MPI_Send(&Temperature[1][1], COLUMNS, MPI_DOUBLE, my_PE_num-1, UP, MPI_COMM_WORLD);
}

if(my_PE_num != npes-1){
    MPI_Recv(&Temperature_last[ROWS+1][1], COLUMNS, MPI_DOUBLE, my_PE_num+1, UP, MPI_COMM_WORLD, &status);
}

dt = 0.0;

#pragma acc kernels
for(i = 1; i <= ROWS; i++) {
    for(j = 1; j <= COLUMNS; j++){
        dt = fmax(fabs(Temperature[i][j]-Temperature_last[i][j]), dt);
        Temperature_last[i][j] = Temperature[i][j];
    }
}

MPI_Reduce(&dt, &dt_global, 1, MPI_DOUBLE, MPI_MAX, 0, MPI_COMM_WORLD);
MPI_Bcast(&dt_global, 1, MPI_DOUBLE, 0, MPI_COMM_WORLD);

if((iteration % 100) == 0) {
    if (my_PE_num == npes-1){
        #pragma acc update host(Temperature)
        track_progress(iteration);
    }
}

iteration++;
}
```

0.9s

MPI routines using host data
#pragma acc data copy(Temperature_last), create(Temperature)
while ( dt_global > MAX_TEMP_ERROR && iteration <= max_iterations ) {
    #pragma acc kernels
    for(i = 1; i <= ROWS; i++) {
        for(j = 1; j <= COLUMNS; j++) {
            Temperature[i][j] = 0.25 * (Temperature_last[i+1][j] + Temperature_last[i-1][j] + Temperature_last[i][j+1] + Temperature_last[i][j-1]);
        }
    }
    #pragma acc update host(Temperature, Temperature_last)
    if(my_PE_num != npes-1){
        MPI_Send(&Temperature[ROWS][1], COLUMNS, MPI_DOUBLE, my_PE_num+1, DOWN, MPI_COMM_WORLD);
    }
    if(my_PE_num != 0){
        MPI_Recv(&Temperature_last[0][1], COLUMNS, MPI_DOUBLE, my_PE_num-1, DOWN, MPI_COMM_WORLD, &status);
    }
    if(my_PE_num != 0){
        MPI_Send(&Temperature[1][1], COLUMNS, MPI_DOUBLE, my_PE_num-1, UP, MPI_COMM_WORLD);
    }
    if(my_PE_num != npes-1){
        MPI_Recv(&Temperature_last[ROWS+1][1], COLUMNS, MPI_DOUBLE, my_PE_num+1, UP, MPI_COMM_WORLD, &status);
    }
    #pragma acc update device(Temperature, Temperature_last)
    dt = 0.0;
    #pragma acc kernels
    for(i = 1; i <= ROWS; i++){
        for(j = 1; j <= COLUMNS; j++){
            dt = fmax(fabs(Temperature[i][j]-Temperature_last[i][j]), dt);
            Temperature_last[i][j] = Temperature[i][j];
        }
    }
    MPI_Reduce(&dt, &dt_global, 1, MPI_DOUBLE, MPI_MAX, 0, MPI_COMM_WORLD);
    MPI_Bcast(&dt_global, 1, MPI_DOUBLE, 0, MPI_COMM_WORLD);
    if((iteration % 100) == 0) {
        if (my_PE_num == npes-1){
            #pragma acc update host(Temperature)
            track_progress(iteration);
        }
    }
    iteration++;
}
Hybrid OpenACC Programming (Slow and Right)

```c
#pragma acc data copy(Temperature_last), create(Temperature)
while ( dt_global > MAX_TEMP_ERROR && iteration <= max_iterations ) {
    #pragma acc kernels
    for(i = 1; i <= ROWS; i++) {
        for(j = 1; j <= COLUMNS; j++) {
            Temperature[i][j] = 0.25 * (Temperature_last[i+1][j] + Temperature_last[i-1][j] + Temperature_last[i][j+1] + Temperature_last[i][j-1]);
        }
    }
    #pragma acc update host(Temperature, Temperature_last)
    if(my_PE_num != npes-1){
        MPI_Send(&Temperature[ROWS][1], COLUMNS, MPI_DOUBLE, my_PE_num+1, DOWN, MPI_COMM_WORLD);
    }
    if(my_PE_num != 0){
        MPI_Recv(&Temperature_last[0][1], COLUMNS, MPI_DOUBLE, my_PE_num-1, DOWN, MPI_COMM_WORLD, &status);
    }
    if(my_PE_num != 0){
        MPI_Send(&Temperature[1][1], COLUMNS, MPI_DOUBLE, my_PE_num-1, UP, MPI_COMM_WORLD);
    }
    if(my_PE_num != npes-1){
        MPI_Recv(&Temperature_last[ROWS+1][1], COLUMNS, MPI_DOUBLE, my_PE_num+1, UP, MPI_COMM_WORLD, &status);
    }
    #pragma acc update device(Temperature, Temperature_last)
    dt = 0.0;
    #pragma acc kernels
    for(i = 1; i <= ROWS; i++) {
        for(j = 1; j <= COLUMNS; j++) {
            dt = fmax(fabs(Temperature[i][j]-Temperature_last[i][j]), dt);
            Temperature_last[i][j] = Temperature[i][j];
        }
    }
    MPI_Reduce(&dt, &dt_global, 1, MPI_DOUBLE, MPI_MAX, 0, MPI_COMM_WORLD);
    MPI_Bcast(&dt_global, 1, MPI_DOUBLE, 0, MPI_COMM_WORLD);
    if((iteration % 100) == 0) {
        if (my_PE_num == npes-1){
            #pragma acc update host(Temperature)
            track_progress(iteration);
        }
    }
    iteration++;
}
```

9.3 s

Update data entering and leaving MPI section
#pragma acc data copy(Temperature_last), create(Temperature)
while ( dt_global > MAX_TEMP_ERROR && iteration <= max_iterations ) {
    #pragma acc kernels
    for(i = 1; i <= ROWS; i++) {
        for(j = 1; j <= COLUMNS; j++) {
            Temperature[i][j] = 0.25 * (Temperature_last[i+1][j] + Temperature_last[i-1][j] + Temperature_last[i][j+1] + Temperature_last[i][j-1]);
        }
    }
    #pragma acc update host(Temperature[1:1][1:COLUMNS],Temperature[ROWS:1][1:COLUMNS])

    if(my_PE_num != npes-1){
        MPI_Send(&Temperature[ROWS][1], COLUMNS, MPI_DOUBLE, my_PE_num+1, DOWN, MPI_COMM_WORLD);
    }
    if(my_PE_num != 0){
        MPI_Recv(&Temperature_last[0][1], COLUMNS, MPI_DOUBLE, my_PE_num-1, DOWN, MPI_COMM_WORLD, &status);
    }
    if(my_PE_num != 0){
        MPI_Send(&Temperature[1][1], COLUMNS, MPI_DOUBLE, my_PE_num-1, UP, MPI_COMM_WORLD);
    }
    if(my_PE_num != npes-1){
        MPI_Recv(&Temperature_last[ROWS+1][1], COLUMNS, MPI_DOUBLE, my_PE_num+1, UP, MPI_COMM_WORLD, &status);
    }
    #pragma acc update device(Temperature_last[0:1][1:COLUMNS], Temperature_last[ROWS+1:1][1:COLUMNS])
    dt = 0.0;
    #pragma acc kernels
    for(i = 1; i <= ROWS; i++) {
        for(j = 1; j <= COLUMNS; j++) {
            dt = fmax( fabs(Temperature[i][j]-Temperature_last[i][j]), dt);
            Temperature_last[i][j] = Temperature[i][j];
        }
    }
    MPI_Reduce(&dt, &dt_global, 1, MPI_DOUBLE, MPI_MAX, 0, MPI_COMM_WORLD);
    MPI_Barrier(&dt_global, 1, MPI_DOUBLE, 0, MPI_COMM_WORLD);
    if((iteration % 100) == 0) {
        if (my_PE_num == npes-1) {
            #pragma acc update host(Temperature)
            track_progress(iteration);
        }
    }
    iteration++;
}
#pragma acc data copy(Temperature_last), create(Temperature)
while ( dt_global > MAX_TEMP_ERROR && iteration <= max_iterations ) {
    #pragma acc kernels
    for(i = 1; i <= ROWS; i++) {
        for(j = 1; j <= COLUMNS; j++) {
            Temperature[i][j] = 0.25 * (Temperature_last[i+1][j] + Temperature_last[i-1][j] + Temperature_last[i][j+1] + Temperature_last[i][j-1]);
        }
    }
    #pragma acc update host(Temperature[1:1][1:COLUMNS], Temperature[ROWS:1][1:COLUMNS])
    if(my_PE_num != npes-1){
        MPI_Send(&Temperature[ROWS][1], COLUMNS, MPI_DOUBLE, my_PE_num+1, DOWN, MPI_COMM_WORLD);
    }
    if(my_PE_num != 0){
        MPI_Recv(&Temperature_last[0][1], COLUMNS, MPI_DOUBLE, my_PE_num-1, DOWN, MPI_COMM_WORLD, &status);
    }
    if(my_PE_num != 0){
        MPI_Send(&Temperature[1][1], COLUMNS, MPI_DOUBLE, my_PE_num-1, UP, MPI_COMM_WORLD);
    }
    if(my_PE_num != npes-1){
        MPI_Recv(&Temperature_last[ROWS+1][1], COLUMNS, MPI_DOUBLE, my_PE_num+1, UP, MPI_COMM_WORLD, &status);
    }
    #pragma acc update device(Temperature_last[0:1][1:COLUMNS], Temperature_last[ROWS+1:1][1:COLUMNS])
    dt = 0.0;
    #pragma acc kernels
    for(i = 1; i <= ROWS; i++){
        for(j = 1; j <= COLUMNS; j++){
            dt = fmax(fabs(Temperature[i][j]-Temperature_last[i][j]), dt);
            Temperature_last[i][j] = Temperature[i][j];
        }
    }
    MPI_Reduce(&dt, &dt_global, 1, MPI_DOUBLE, MPI_MAX, 0, MPI_COMM_WORLD);
    MPI_Bcast(&dt_global, 1, MPI_DOUBLE, 0, MPI_COMM_WORLD);
    if((iteration % 100) == 0) {
        if (my_PE_num == npes-1){
            #pragma acc update host(Temperature)
            track_progress(iteration);
        }
    }
    iteration++;
}
Hybrid OpenMP Programming
(Most “complex” version: MPI_THREAD_MULTIPLE)

#include <mpi.h>
#include <omp.h>

//Last thread of PE 0 sends its number to PE 1

main(int argc, char* argv[]){
    int provided, myPE, thread, last_thread, data=0, tag=0;
    MPI_Status status;

    MPI_Init_thread(&argc, &argv, MPI_THREAD_MULTIPLE, &provided);
    MPI_Comm_rank(MPI_COMM_WORLD, &myPE);

    #pragma omp parallel firstprivate(thread, data, tag, status)
    {
        thread = omp_get_thread_num();
        last_thread = omp_get_num_threads()-1;

        if ( thread==last_thread && myPE==0 )
            MPI_Send(&thread, 1, MPI_INT, 1, tag, MPI_COMM_WORLD);
        else if ( thread==last_thread && myPE==1 )
            MPI_Recv(&data, 1, MPI_INT, 0, tag, MPI_COMM_WORLD, &status);

        printf("PE %d, Thread %d, Data %d\n", myPE, thread, data);
    }

    MPI_Finalize();
}
Mix and Match

- **PGI Compile:**
  ```
  mpicc -acc laplace_hybrid.c
  mpf90 -acc laplace_hybrid.f90
  mpicc -mp -acc laplace_hybrid.c
  etc...
  ```

- **Running:**
  ```
  interact ?
  -n 4
  -N1 -n4
  -p GPU -N1 -n4
  -p GPU -N4 -n4
  -N1 -n28
  -N4 -n112
  etc...
  ```
Bottom Line...

• Each one of these approaches occupies its own space.

• If you understand this, you will not be confused as to how they fit together.

• Once again...
In Conclusion...
In Conclusion...

OpenMP
In Conclusion...

OpenMP
In Conclusion...

OpenACC

OpenMP
In Conclusion...

OpenMP

OpenACC
In Conclusion...

OpenMP

OpenACC

MPI
Hybrid Challenge
Head Start

John Urbanic
Parallel Computing Scientist
Pittsburgh Supercomputing Center
In the parallel case, we will break this up into 4 processors. There is only one set of boundary values. But when we distribute the data, each processor needs to have an extra row for data distribution, these are commonly called the “ghost cells”.

The program has a local view of data. The programmer has to have a global view of data. The ghost cells don’t exist in the global dataset. They are only copies from the “real” data in the adjacent PE.
Simplest Decomposition for Fortran Code

Then we send strips to ghost zones like this:

Same ghost cell structure as the C code, we have just swapped rows and columns.
Both C and Fortran will need to set proper boundary conditions based upon the PE number.
How do you know you are correct?

Your solution converges at 3372 timesteps!
How do you know you are correct?

Your solution converges at 3372 timesteps!
How do you know you are correct?

Both converge at 3372 steps!
How do you know you are correct?

Both converge at 3372 steps!
All the action is here.
void output(int my_pe, int iteration) {
    FILE* fp;
    char filename[50];
    sprintf(filename,"output%d.txt",iteration);
    for (int pe = 0; pe<4; pe++){
        if (my_pe==pe){
            fp = fopen(filename, "a");
            for(int y = 1; y <= ROWS; y++){
                for(int x = 1; x <= COLUMNS; x ++){
                    fprintf(fp, "%5.2f ",Temperature[y][x]);
                }
                fprintf(fp,"\n");
            }
            fflush(fp);
            fclose(fp);
        }
        MPI_Barrier(MPI_COMM_WORLD);
    }
}

• Human Readable
• 1M entries
• Visualize. I used Excel (terrible idea).
void output(int my_pe, int iteration) {
    FILE* fp;
    char filename[50];
    sprintf(filename,"output%d.txt",iteration);
    for (int pe = 0; pe<4; pe++){
        if (my_pe==pe){
            fp = fopen(filename, "a");
            for(int y = 1; y <= ROWS; y++){
                for(int x = 1; x <= COLUMNS; x ++){
                    fprintf(fp, "%5.2f ",Temperature[y][x]);
                }
                fprintf(fp,"\n");
            }
            fflush(fp);
            fclose(fp);
        }
    }
    MPI_Barrier(MPI_COMM_WORLD);
}

• Human Readable
• 1M entries
• Visualize. I used Excel (terrible idea).

C:
if (my_PE_num==2)
    printf("Global coord [750,900] is %f \n", Temperature[250][900]);

Fortran:
if (mype==2) then
    print*, 'magic point', temperature(900,250)
endif

• If about 1.0, probably good
• Otherwise (like 0.02 here) probably not
Running to convergence

3372 iterations to converge to 0.01 max delta.

<table>
<thead>
<tr>
<th></th>
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<th>4 Nodes</th>
<th>Speedup</th>
</tr>
</thead>
<tbody>
<tr>
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<td>16.0</td>
<td>4.9</td>
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</table>

Note that all versions converge on the same iteration. This kind of repeatability should be expected. However, exact binary repeatability is usually not possible due simply to floating point operation reordering.

Scaling off the node will typically be much better than scaling on the node for a well written problem of this type run at normal scale.

To run on 4 nodes you need to request 4 nodes from the queue: `interact -N 4 -n 4`
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To run on 4 nodes you need to request 4 nodes from the queue:

- interact
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Note that all versions converge on the same iteration. This kind of repeatability should be expected. However, exact binary repeatability is usually not possible due simply to floating point operation reordering.

Scaling off the node will typically be much better than scaling on the node for a well written problem of this type run at normal scale.

To run on 4 nodes you need to request 4 nodes from the queue:  

`interact -N 4 -n 4`
call MPI_Send_Init(temperature(1,columns), rows, MPI_DOUBLE_PRECISION, right, lr, MPI_COMM_WORLD, request(1), ierr)
call MPI_Recv_Init(temperature_last(1,0), rows, MPI_DOUBLE_PRECISION, left, lr, MPI_COMM_WORLD, request(2), ierr)

// 8 of these as winning solution did a 2D (left, right, up, down) decomposition on 10,000 x 10,000 size problem

do while ( dt_global > max_temp_error .and. iteration <= max_iterations)

do j=1,columns
    do i=1,rows
        temperature(i,j)=0.25*(temperature_last(i+1,j)+temperature_last(i-1,j)+ &
        temperature_last(i,j+1)+temperature_last(i,j-1))
    enddo
enddo

call MPI_StartAll(8,request,statuses)

dt=0.0

do j=1,columns
    do i=1,rows
        dt = max( abs(temperature(i,j) - temperature_last(i,j)), dt )
        temperature_last(i,j) = temperature(i,j)
    enddo
enddo

call MPI_WaitAll(8,request,statuses,ierr)
enddo
Rules and Regulations of the 2\textsuperscript{nd} Annual IHPCSS Challenge

Trophy bears no relationship to reality.
General Rules

- Due Thursday midnight (!)
- 4 Nodes of Bridges
- Use any combination of MPI, OpenACC, Python and OpenMP
- How fast can you run a 10K x 10K Laplace code to convergence?
Some Specifics

- Can’t change kernel (Must retain two core loops source)
- Can change number of MPI processes (Does not have to be 112 or 4)
- 1 Source File
- 1 Combined Environment/Compile/Submit/Execute script to make it easy for us to run your solutions!
- Mail to d.henty@epcc.ed.ac.uk by deadline
  - Mail a ping earlier if you want to be informed of any developments
Rules For Lawyers

- No libraries
- Don’t mess with timer placement
- ?
Reality Checks

- Serial code converges at 3578 time steps. Yours should too.

- As we know, this is not enough to verify correctness. You should find point [7500][9950] in C and (9950,7500) in Fortran converges to 17 (±1) degrees.

- As discussed, the 10K result differs from the 1K result.*
  - Plugging in Gauss-Seidel or Successive Over Relaxation (SOR) would be easy and interesting. But, not for our contest.

http://www.cs.berkeley.edu/~demmel/cs267/lecture24/lecture24.html is a brief analysis of these issues.
Suggested Things to Explore

- Compiler flags
  - fast

- Compiler
  - see Bridges documentation for how to use different modules

- MPI Environment Variables
  - man mpi

- Thread placement
  - google for KMP_AFFINITY

MPI

OpenMP

OpenACC
Suggested Things to Explore

- **Compiler flags**
  - `-fast`

- **Compiler**
  - see Bridges documentation for how to use different modules

- **MPI Environment Variables**
  - `man mpi`

- **Thread placement**
  - google for `KMP_AFFINITY`

---

**Blue Waters User Guide is your friend!**
https://bluewaters.ncsa.illinois.edu/user-guide
On Thursday evening we will take the top self-reported speeds and run them in an interactive session.

Timings not within 10% of self-reported time will be disqualified.

Codes should print out “test point” at [7500][9950] for C, (9950,7500) for Fortran at conclusion of run.

Best of two runs for each finalist will determine winner.