





# Lecture 8: MPI Parallelization, part I (parallel::distributed::Triangulation)

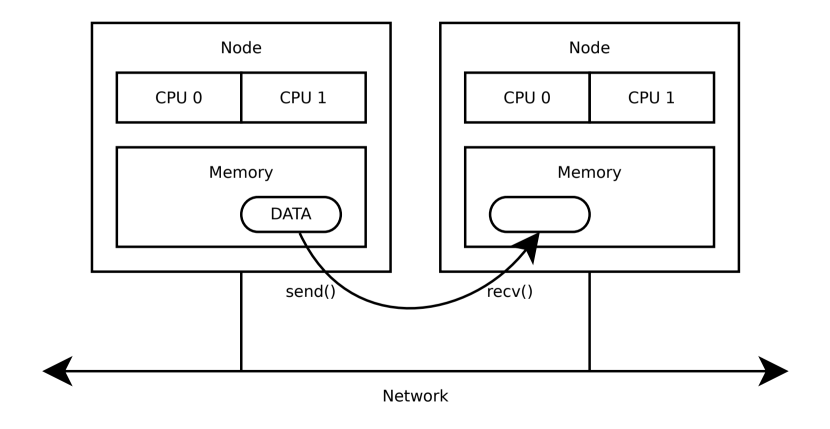
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# Parallel computing model: MPI









## General Considerations

- Goal: get the solution faster!
- If FEM with <500.000 dofs, and 2d, use direct solver!</li>
- If you need more, then you have to SPLIT the work
  - Distributed data storage everywhere
    - need special data structures
  - Efficient algorithms
    - not depending on total problem size
  - "Localize" and "hide" communication
    - point-to-point communication, nonblocking sends and receives

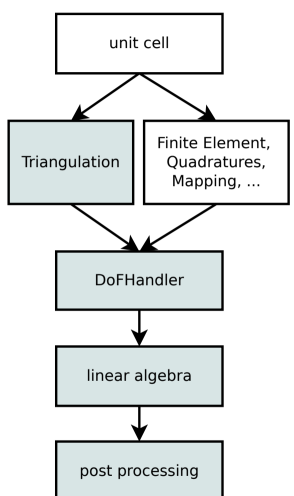




## Data Structures

#### Needs to be parallelized:

- 1. Triangulation (mesh with associated data)
  - hard: distributed storage, new algorithms
- 2. DoFHandler (manages degrees of freedom)
  - hard: find global numbering of DoFs
- 3. Linear Algebra (matrices, vectors, solvers)
  - use existing library
- 4. Postprocessing (error estimation, solution transfer, output, . . . )
  - do work on local mesh, communicate





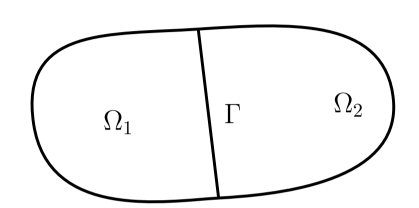




## How to Parallelize?

#### Option 1: Domain Decomposition

- Split up problem on PDE level
- Solve subproblems independently
- Converges against global solution
- Problems:
  - Boundary conditions are problem dependent:
    - → sometimes difficult!
    - → no black box approach!
  - Without coarse grid solver: condition number grows with # subdomains → no linear scaling with number of CPUs!





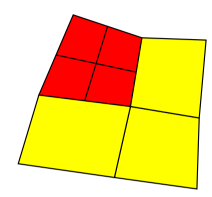




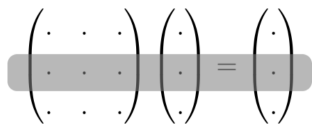
## How to Parallelize?

Option 2: Algebraic Splitting

Split up mesh between processors:



Assemble logically global linear system (distributed storage):



- Solve using iterative linear solvers in parallel
- Advantages:
  - Looks like serial program to the user
  - Linear scaling possible (with good preconditioner)



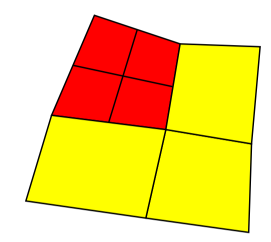




## Partitioning

Optimal partitioning (coloring of cells):

- same size per region
  - → even distribution of work
- minimize interface between region
  - → reduce communication



Optimal partitioning is an NP-hard graph partitioning problem.

- Typically done: heuristics (existing tools: METIS)
- Problem: worse than linear runtime
- Large graphs: several minutes, memory restrictions
- → Alternative: avoid graph partitioning

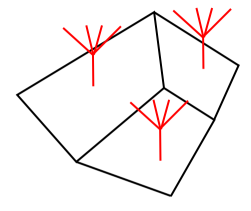


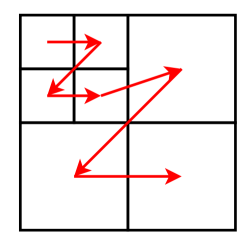


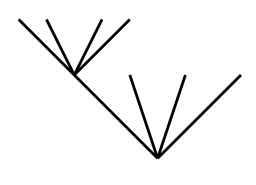


#### Partitioning with "Space filling curves"

- \* p4est library: parallel quad-/octrees
- Store refinement flags from a base mesh
- Based on space-filling curves
- Very good scalability









Burstedde, Wilcox, and Ghattas.

p4est: Scalable algorithms for parallel adaptive mesh refinement on forests of octrees.

SIAM J. Sci. Comput., 33 no. 3 (2011), pages 1103-1133.

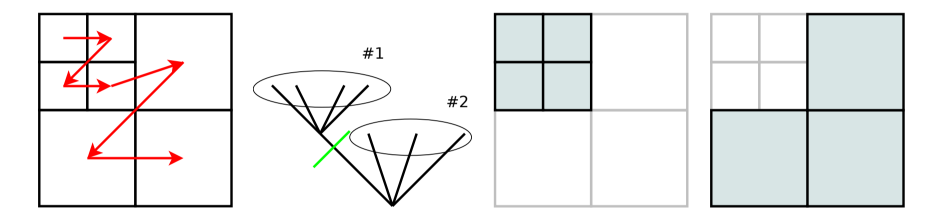






# Triangulation

Partitioning is cheap and simple:



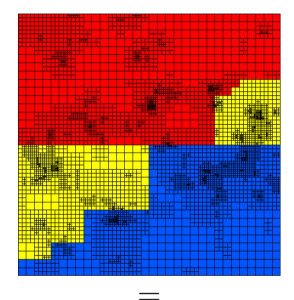
- Then: take *p4est* refinement information
- Recreate rich *deal.II* Triangulation only for local cells (stores coordinates, connectivity, faces, materials, . . . )
- \* How? recursive queries to *p4est*
- Also create ghost layer (one layer of cells around own ones)

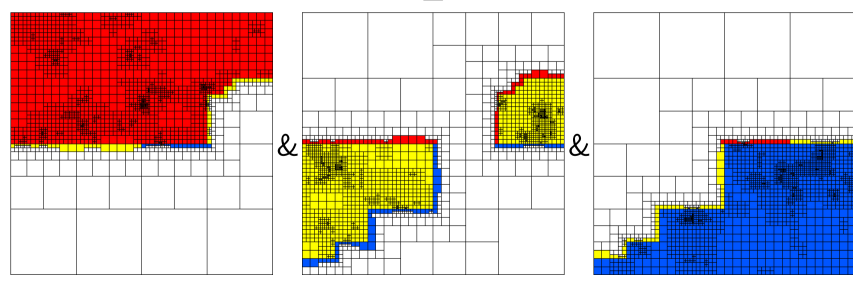






# Example (color by CPU ID)





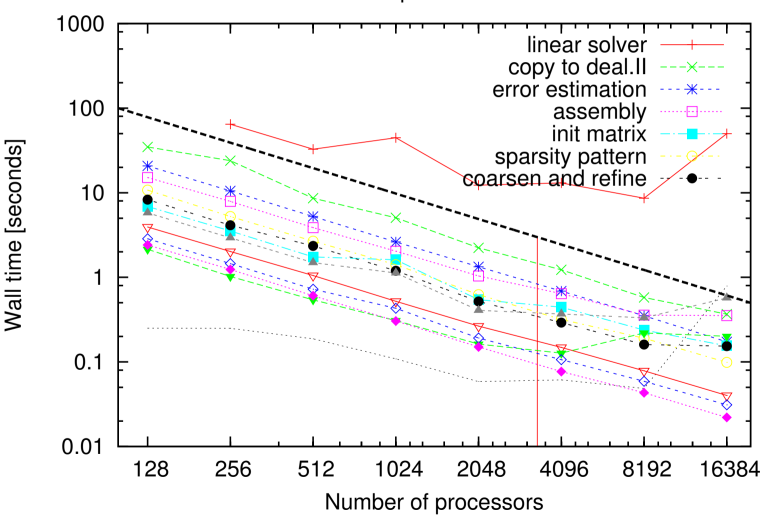






# Strong Scaling: 2d Poisson

Wall clock times for problem of fixed size 335M

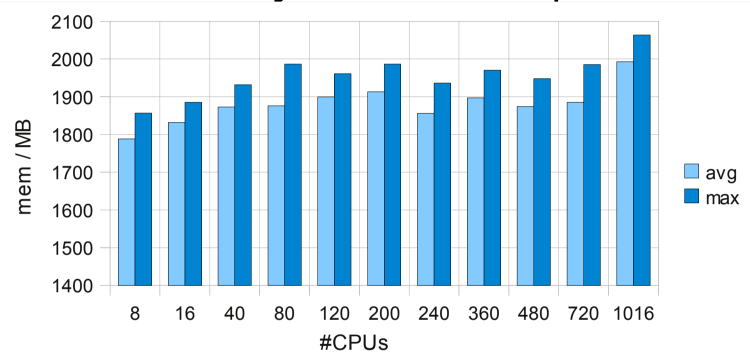








## Memory Consumption

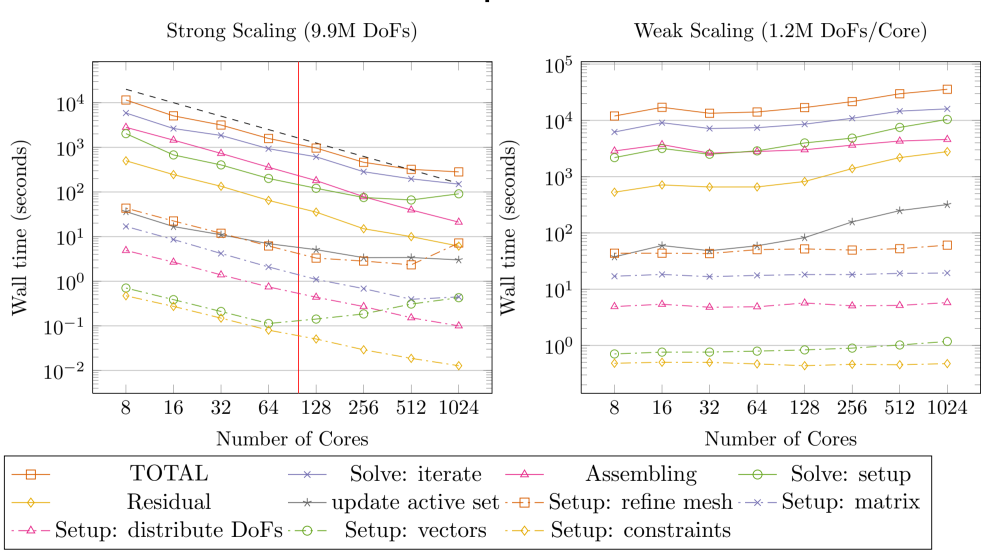


average and maximum memory consumption (VmPeak) 3D, weak scalability from 8 to 1000 processors with about 500.000 DoFs per processor (4 million up to 500 million total)





# Step 40









## Trilinos VS PETSc

#### What should I use?

- Similar features and performance
- Pro Trilinos: more development, some more features (automatic differentation, . . . ), cooperation with deal.II
- \* Pro PETSc: stable, easier to compile on older clusters
- But: being flexible would be better! "why not both?"
  - you can! Example: new step-40
  - can switch at compile time
  - need #ifdef in a few places (different solver parameters TrilinosML vs BoomerAMG)
  - some limitations, somewhat work in progress







## Trilinos VS PETSc

```
#define USE_PETSC_LA // uncomment this to run with Trilinos
namespace LA
#ifdef USE_PETSC_LA
  using namespace dealii::LinearAlgebraPETSc;
#else
  using namespace dealii::LinearAlgebraTrilinos;
#endif
   LA::MPI::SparseMatrix system_matrix;
   LA::MPI::Vector solution;
   LA::SolverCG solver(solver_control, mpi_communicator);
   LA::MPI::PreconditionAMG preconditioner;
   LA::MPI::PreconditionAMG::AdditionalData data:
#ifdef USE_PETSC_LA
    data.symmetric_operator = true;
#else
    //trilinos defaults are good
#endif
    preconditioner.initialize(system_matrix, data);
```