Dynamic Adaptively Refined Mesh Simulations on 1M+ Cores

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ABSTRACT
This poster presents a clustering algorithm and a partitioning algorithm for patch-based dynamically adaptive mesh refinement (AMR) and shows scaling results to more than 1M cores.

Keywords
scalability, parallel algorithm, adaptive mesh refinement, dynamic adaptivity, mesh partitioning, clustering

1. INTRODUCTION
Patch-based structured adaptive mesh refinement (SAMR) is a meshing approach that assembles meshes from simple unpartitioned structured grids. The SAMR mesh is a hierarchy of box-shaped grids of increasing resolutions (Figure 1a & b). An advantage of SAMR is that the mesh is succinctly described because each box can be specified with very little metadata: just the indices of two corners and a few ancillary attributes, such as the MPI process that owns the box.

We use the term mesh management (MM) for the generation, acquisition (via communication), storage of and operations on metadata. Ultimately, MM must provide the boxes and their proximity data, such as that represented by the graph in Figure 1c. Traditional MM approaches duplicate global metadata on every process and use sequential algorithms. This has proved unfeasible on more than $O(10^5)$ cores due to the linear growth of global metadata. Distributing metadata is not trivial, because traditional MM all required global metadata.

To address this problem, we previously developed a distributed MM approach [2]. By saving and using key information typically discarded by traditional algorithms, MM operations on global metadata could be re-cast in terms of distributed metadata. Proximity data was generated and evolved with the boxes instead of by global searches. Although this proved to be scalable, two key MM algorithms bogged down at $O(10^5)$ cores: the clustering algorithm [3] and the partitioning algorithm [1]. These were plug-in components, which the SAMR framework accessed by abstract interfaces. This poster highlights the new clustering and partitioning algorithms that allowed scaling to $O(10^6)$ cores.

2. TILE CLUSTERING
Clustering generates the initial set of boxes when creating or replacing a level. The user tags cells for the new level to cover, and the algorithm finds a set of boxes that cover those cells. We allow the boxes to contain some non-tagged cells to avoid requiring too many boxes, but the over-refinement extreme of too many non-tagged cells should also be avoided.

The new clustering algorithm was an extension of the highly scalable tile clustering approach by Luijtjens [4]. The index space was divided into fixed-size tiles. We generated a box for each tile that overlapped at least one tag. We used tile sizes on the small side to reduce over-refinement. To reduce the number of boxes, we coalesced them. Thus, we achieved both the low over-refinement of small tiles and the low box-counts of large tiles. In addition, we extended Luijtjens’s algorithms to generate the edges necessary to integrate scalably in the MM framework. For flexibility, we allowed tiles to cross patch and processor boundaries.

3. CASCADE PARTITIONING
Cascade partitioning was a new approach. We used P2P communication to build on each process a $O(\log N)$-sized picture of the global load distribution, including the load of each half of the machine. Transferring work between the halves balanced their loads. Applying the algorithm recursively balanced within each half.
We grouped the tasks by recursive splitting, forming a binary tree wherein each node was a grouping (Figure 2). Data exchanges among intergroup communication pairs gave every process the load of each group it belonged to and of their sibling groups. After data exchanges using groupings 3, 2, then 1, all processes knew which half of the machine had more load and how much to transfer to the other half. Load transfer from the overloaded half to the underloaded used communication pairs in Grouping 1. Only metadata were exchanged, so all message lengths were short.

4. RESULTS

Figure 3 shows two scaling results obtained on a 1.5M-core Sequoia IBM BG/Q machine. The weak-scaling benchmark solved a linear advection equation, with 1 unknown/cell, propagating sinusoidal waves in the domain. The physics and numerics of the problem were simple. Regridding interval was once every 9 time steps. This benchmark averaged $1.64 \times 10^6$ cells/core. Regridding remained less than 25% of the total cost. The strong scaling benchmark solved the Euler equation, with 5 unknowns/cell, for an expanding spherical shock wave. Its mesh started with $4 \times 10^9$ cells and nearly doubled during the run. The regrid interval was 8 time steps. Both benchmarks used a simple forward Euler time-stepping scheme. Note that both were designed to be challenging for regridding, having little computation and frequent regridding. Changing computational work and/or regrid intervals would give different results.

Figure 4 shows scaling characteristics of mesh management for the scalar advection benchmark at 4 levels and $195 \times 10^3$ cells/core. All components, including the new clustering and partitioning components, scaled well and showed promise of continuing well past the largest runs performed. Note that if any one component failed to scale, regridding itself would also fail to scale. The strong scaling benchmark showed comparable performance but was harder to evaluate because in strong scaling, everything eventually degraded.

5. REFERENCES


Figure 2: Cascade groups for ranks 0-7. Underscores indicate process groupings. Each group is a binary tree node. Each process belongs to 4 groups. Arcs connecting processes indicate inter-group communication pairs.

Figure 3: Scaling results. a) Weak scaling to 1M cores. Advection equation with 1 unknown/cell. $1.640 \times 10^6$ cells/core. Regrid every 9 time steps. b) Strong scaling to 1.57M cores. Euler equations with 5 unknowns/cell. $4.03-8.41 \times 10^9$ cells globally. Regrid every 8 time steps.

Figure 4: Regrid scaling results