A Unified Algorithm for Load-balancing Adaptive Scientific Simulations

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Abstract

Adaptive scientific simulations require that periodic repartitioning occur dynamically throughout the course of the computation. The repartitionings should be computed so as to minimize both the inter-processor communications incurred during the iterative mesh-based computation and the data redistribution costs required to balance the load. Recently developed schemes for computing repartitionings provide the user with only a limited control of the tradeoffs among these objectives. This paper describes a new Unified Repartitioning Algorithm that can tradeoff one objective for the other dependent upon a user-defined parameter describing the relative costs of these objectives. We show that the Unified Repartitioning Algorithm is able to reduce the precise overheads associated with repartitioning as well as or better than other repartitioning schemes for a variety of problems, regardless of the relative costs of performing inter-processor communication and data redistribution. Our experimental results show that this scheme is extremely fast and scalable to large problems.

Keywords: Unified Repartitioning Algorithm, Dynamic Graph Partitioning, Multilevel Diffusion, Scratch-remap, Adaptive Mesh Computations

1 Introduction

For large-scale scientific simulations, the computational requirements of techniques relying on globally refined meshes become very high, especially as the complexity and size of the problems increase. By locally refining and de-refining the mesh either to capture flow-field phenomena of interest [1] or to account for variations in errors [14], adaptive methods make standard computational methods more cost effective. The efficient execution of such adaptive scientific simulations on parallel computers requires a periodic repartitioning of the underlying computational mesh. These repartitionings should minimize both the inter-processor communications incurred in the iterative mesh-based computation and the data redistribution costs required to balance the load. Recently developed schemes for computing repartitionings provide the user with only a limited control of the tradeoffs among these two objectives. This paper describes a new Unified Repartitioning Algorithm that can tradeoff one objective for the other dependent upon a user-defined parameter describing the relative costs of these objectives.

Figure 1 illustrates the steps involved in the execution of adaptive mesh-based simulations on parallel computers. Initially, the mesh is distributed among the processors. A number of iterations of the simulation

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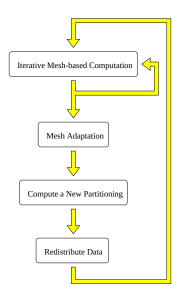


Figure 1: A diagram illustrating the execution of adaptive scientific simulations on high performance parallel computers.

are performed in parallel, after which mesh adaptation occurs. Here, each processor refines and de-refines its local regions of the mesh resulting in some amount of load imbalance. A new partitioning based on the adapted mesh is computed to re-balance the load, and then the mesh is redistributed among the processors, respectively. The simulation can then continue for another number of iterations until either more mesh adaptation is required or the simulation terminates.

If we consider each round of executing a number of iterations of the simulation, mesh adaptation, and load-balancing to be an *epoch*, then the run time of an epoch can be described by

$$(t_{comp} + f(|E_{cut}|))n + t_{repart} + g(|V_{move}|)$$
(1)

where n is the number of iterations executed, t_{comp} is the time to perform the computation for a single iteration of the simulation, $f(|E_{cut}|)$ is the time to perform the communications required for a single iteration of the simulation, and t_{repart} and $g(|V_{move}|)$ represent the times required to compute the new partitioning and to redistribute the data accordingly. Here, the inter-processor communication time is described as a function of the edge-cut of the partitioning and the data redistribution time is described as a function of the total amount of data that is required to be moved in order to realize the new partitioning.

Adaptive repartitioning affects all of the terms in Equation 1. How well the new partitioning is balanced influences t_{comp} . The inter-processor communications time is dependent on the edge-cut of the new partitioning. The data redistribution time is dependent on the total amount of data that is required to be moved in order to realize the new partitioning. Recently developed adaptive repartitioning schemes [4, 5, 11, 12, 13, 14, 15, 16, 17, 20, 21, 22, 23, 24, 25] tend to be very fast, especially compared to the time required to perform even a single iteration of a typical scientific simulation. They also tend to balance the new partitioning to within a few percent of optimal. Hence, we can ignore both t_{repart} and t_{comp}^{-1} . However, depending on the nature of the application, both $f(|E_{cut}|)$ and $g(|V_{move}|)$ can seriously affect parallel run times and drive down parallel efficiencies. Therefore, it is critical for adaptive partitioning schemes to attempt to minimize both the edge-cut and the data redistribution when computing the new partitioning. Viewed in this way, adaptive graph partitioning is a multi-objective optimization problem.

Two approaches have primarily been taken when designing adaptive graph partitioners. The first approach is to focus on minimizing the edge-cut and to minimize the data redistribution only as a secondary objective [11, 12, 16, 17, 21, 22, 23, 24, 25]. A good example of such schemes are scratch-remap repartitioners

¹This is because, in the absence of load imbalance, t_{comp} will be primarily determined by the domain-specific computation and cannot be reduced further.

[11, 17, 21]. These use some type of state-of-the-art graph partitioner to compute a new partitioning from scratch and then attempt to intelligently remap the subdomain labels to those of the original partitioning in order to reduce the data redistribution costs. Since a state-of-the-art graph partitioner is used to compute the partitioning, the resulting edge-cut tends to be extremely good. However, since there is no guarantee as to how similar the new partitioning will be to the original partitioning, data redistribution costs can be high, even after remapping [2, 17, 19]. A second approach is to focus on minimizing the data redistribution cost and to minimize the edge-cut as a secondary objective [4, 13, 14, 15, 20]. Diffusion-based repartitioners [12, 16, 17, 23, 24, 25] are good examples of this approach. These schemes attempt to perturb the original partitioning just enough so as to balance it. This strategy usually leads to low data redistribution costs, especially when the partitioning is only slightly imbalanced. However, it can result in higher edge-cuts than scratch-remap methods because perturbing a partitioning in this way also tends to adversely affect its quality.

Both of these approaches to adaptive partitioning have two drawbacks. The first is that the two types of repartitioners allow the user to compute partitionings that focus on minimizing either the edge-cut or the data redistribution costs, but give the user only a limited ability to control the tradeoffs among these objectives. This control is sufficient if the number of iterations that a simulation performs during every epoch (i. e. the value of n in Equation 1) is either very high or very low. However, when n is neither very high nor very low, neither type of scheme precisely minimizes the combined costs of $f(|E_{cut}|)n$ and $g(|V_{move}|)$. The second disadvantage exists for applications in which n is difficult to predict or those in which n can change dynamically throughout the course of the computation. As an example, one of the key issues concerning the elastic-plastic soil-structure interaction computations required for earthquake simulation is that the number of iterations performed during each epoch is both unpredictable and dynamic. Here, zones in the 3D solid may load (i. e., become plastic) and then unload (become elastic again) so that the extent of the plastic zone is changing. The change can be both slow and rapid. Slow change usually occurs during initial loading phases, while the later deformation tends to localize in narrow zones rapidly and the rest of the solid unloads rapidly [6].

Recently, Castanos and Savage [2] presented an adaptive repartitioning algorithm that attempts to directly minimize the communication overheads of adaptive multigrid-based finite-element computations. During each load-balancing epoch, their algorithm computes a repartitioning of the coarsest mesh of the hierarchy so as to optimize an objective that is similar to Equation 2 (given below). The coarse-mesh repartitioning is then used to partition the entire hierarchy of nested meshes. While this approach addresses the multi-objective nature of the adaptive repartitioning problem, Castanos and Savage's repartitioning algorithm is serial. Therefore, the scheme is suited only for problems in which: (i) the entire mesh refinement history is retained (eg., multigrid solvers), and (ii) a nested partitioning of the successively finer meshes is employed (i. e., for each coarse element, the entire hierarchy of corresponding finer elements belongs to the same processor).

Our Contributions In this paper, we present a parallel adaptive repartitioning scheme (called the Unified Repartitioning Algorithm) for the dynamic load-balancing of scientific simulations that attempts to solve the precise multi-objective optimization problem. By directly minimizing the combined costs of $f(|E_{cut}|)n$ and $g(|V_{move}|)$, our scheme is able to tradeoff one objective for the other as required by the specific application. Our experimental results show that when inter-processor communication costs are much greater in scale than data redistribution costs, our scheme obtains results that are similar to those obtained by an optimized scratch-remap repartitioner and better than those obtained by an optimized diffusion-based repartitioner. When these two costs are of similar scale, our scheme obtains results that are similar to the diffusive repartitioner and better than the scratch-remap repartitioner. When the cost to perform data redistribution is much greater than the cost to perform inter-processor communication, our scheme obtains better results than the diffusive scheme and much better results than the scratch-remap scheme. Finally, our experimental results show that our Unified Repartitioning Algorithm is fast and scalable to large problems.

Figure 2: The three phases of multilevel k-way graph partitioning. During the coarsening phase, the size of the graph is successively decreased. During the initial partitioning phase, a k-way partitioning is computed. During the multilevel refinement (or uncoarsening) phase, the partitioning is successively refined as it is projected to the larger graphs. G_0 is the input graph, which is the finest graph. G_{i+1} is the next level coarser graph of G_i . G_4 is the coarsest graph.

Initial Partitioning Phas

2 Unified Repartitioning Algorithm

We have developed a new parallel Unified Repartitioning Algorithm (URA) for dynamic load-balancing of scientific simulations that improves upon the best characteristics of scratch-remap and diffusion-based repartitioning schemes. A key parameter used in URA is the *Relative Cost Factor* (RCF). This parameter describes the relative times required for performing the inter-processor communications incurred during parallel processing and to perform the data redistribution associated with balancing the load. Therefore, it depends not only on n from Equation 1, but also on the specific machine and algorithm used. Using the RCF, it is possible to unify the two minimization objectives of the adaptive graph partitioning problem into the precise cost function

$$|E_{cut}| + \alpha |V_{move}| \tag{2}$$

where α is the Relative Cost Factor, $|E_{cut}|$ is the edge-cut of the partitioning, and $|V_{move}|$ is the total amount of data redistribution. The Unified Repartitioning Algorithm computes a partitioning while attempting to directly minimize this cost function.

The Unified Repartitioning Algorithm is based upon the multilevel paradigm that is illustrated in Figure 2. We next describe its three phases: graph coarsening, initial partitioning, and uncoarsening/refinement. In the graph coarsening phase, coarsening is performed using a purely local variant of heavy-edge matching [7, 16, 17, 25]. That is, vertices may be matched together only if they are in the same subdomain on the original partitioning. This matching scheme has been shown to be effective at reducing both edge-cuts and data redistribution costs compared to global matching when the original partitioning is of high quality, although possibly imbalanced [17, 25]. (Note that for the adaptive partitioning problem, we can assume a high quality original partitioning.) Local matching is also inheritly more scalable than global matching [16, 17, 25].

Selecting an initial partitioning scheme for an adaptive repartitioner is complicated for a number of reasons. Experimental results [19] have shown that for some types of repartitioning problem instances, scratch-remap repartitioners tend to obtain better results compared to diffusive repartitioners, while for other types of problem instances, diffusive repartitioners tend to do better than scratch-remap repartitioners. Furthermore, the effectiveness of each type of scheme is highly dependent on the value of the Relative Cost Factor. For

| Graph | Num of Verts | Num of Edges | Description |
|--------|--------------|--------------|----------------------|
| auto | 448,695 | 3,314,611 | 3D mesh of GM Saturn |
| mdual2 | $988,\!605$ | 1,947,069 | dual of a 3D mesh |
| mrng3 | 4,039,160 | 8,016,848 | dual of a 3D mesh |

Table 1: Characteristics of the graphs used in some of the experiments.

these reasons, in the initial partitioning phase of URA, repartitioning is performed on the coarsest graph twice by alternative methods. That is, optimized variants of scratch-remap and global diffusion [17] are both used to compute new partitionings. The costs from Equation 2 are then computed for each of these and the one with the lowest cost is selected. This technique tends to give a very good point from which to start multilevel refinement, regardless of the type of repartitioning problem or the value of the Relative Cost Factor. Note that the fact that URA computes two initial partitionings does not impact the scalability of the algorithm as long as the size of the coarsest graph is suitably small [8].

Most adaptive graph partitioning algorithms perform partition refinement in order to reduce the edge-cut of the new partitioning. Some of these [16, 17] also attempt to reduce the data redistribution cost as a tie breaking scheme. (That is, this objective is considered when the gain to the edge-cut that will result from moving a vertex is the same for two or more different subdomains.) However, even with such a tie-breaking scheme, these do not directly reduce the precise cost function described in Equation 2. A refinement algorithm that does so, especially one that is applied in the multilevel context, can potentially reduce the cost function better than current refinement schemes. The refinement algorithm employed in the uncoarsening phase of URA attempts to minimize the precise cost function from Equation 2. Except for this important modification, the refinement algorithm used in URA is similar to the parallel adaptive refinement algorithm described in [17]. Note that since each time a vertex is moved, the cost function can be updated in constant time using information local to the processor, these two algorithms have the same asymptotic run times.

3 Experimental Results

In this section, we present experimental results comparing the cost function and run time results of the Unified Repartitioning Algorithm with optimized versions of scratch-remap (LMSR) [17] and multilevel diffusion (Wavefront Diffusion) [17] repartitioners.

Experimental Setup The experiments presented in this section were conducted on graphs derived from finite-element computations. These graphs are described in Table 1. For each graph, we modified the vertex and edge weights in order to simulate various types of repartitioning problems. Specifically, we constructed four repartitioning problems for each graph that simulate adaptive computations in which the work imbalance is distributed globally throughout the mesh. An example of an application in which this might occur is a particle-in-mesh computation. Here, particles may be located anywhere within the mesh and may be free to move to any other regions of the mesh. The result is that both the densely and sparsely populated regions are likely to be distributed globally throughout the mesh. Typically, this type of repartitioning problem is easier for diffusion-based schemes compared to scratch-remap schemes [17, 19]. We also constructed four problems that simulate adaptive mesh computations in which adaptation occurs in localized regions of the mesh. An example of this type of problem is a simulation of a helicopter blade. Here, the finite-element mesh must be extremely fine around both the helicopter blade and in the vicinity of the sound vortex that is created by the blade in order to accurately capture flow-field phenomena of interest. It should be coarser in other regions of the mesh for maximum efficiency. As the simulation progresses, neither the blade nor the sound vortex remain stationary. Therefore, the new regions of the mesh that these enter need to be refined, while those regions that are no longer of key interest should be de-refined. In this case, mesh refinement and

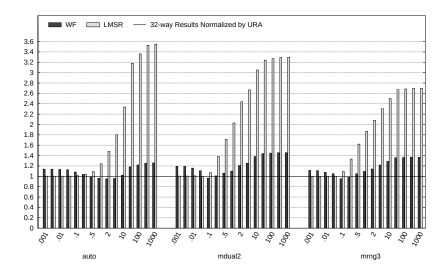


Figure 3: The cost function results obtained from the Unified Repartitioning Algorithm (URA) compared to the results obtained from optimized multilevel diffusion (WF) and scratch-remap (LMSR) algorithms on 32 processors of a Cray T3E.

de-refinement are often performed in very localized regions of the mesh. This type of repartitioning problem tends to be easier for scratch-remap schemes than diffusion-based schemes [17, 19].

Further experiments were performed on a real problem set from the simulation of a diesel internal combustion engine². This is a particles-in-cells computation. The mesh consists of 175-thousand mesh elements. At first, no fuel particles are present in the combustion chamber. As the computation progresses, fuel particles are injected into the chamber at a single point and begin to spread out. Thus, they may enter regions of the mesh belonging to different processors. Load imbalance occurs as processors are required to track different numbers of particles.

Results on Synthetic Data Sets Figures 3 through 5 show the costs from Equation 2 obtained by the Unified Repartitioning Algorithm compared to those obtained by the optimized scratch-remap and multilevel diffusion algorithms, LMSR and Wavefront Diffusion [17], as implemented in PARMEIIS, Version 2.0 [9] on 32, 64, and 128 processors of a Cray T3E. Specifically, these figures show three sets of results, one for each of the graphs described in Table 1. Each set is composed of fifteen pairs of bars. These pairs represent the averaged results from the eight experiments (simulating global and localized imbalances) that are described above. For each pair of bars, the Relative Cost Factor was set to a different value. These values are .001, .002, .01, .02, .1, .25, .5, 1, 2, 4, 10, 50, 100, 500, and 1000. Therefore, for each set of results, minimizing the edge-cut is the dominate objective for the results on the left, while minimizing the data redistribution cost is the dominate objective for the results on the right. The results in the middle represent varying tradeoffs between the two objectives. The bars in Figures 3 through 5 give the average costs obtained by the optimized scratch-remap and multilevel diffusion repartitioners normalized by the average costs obtained by URA. Therefore, a result above the 1.0 index line indicates that the corresponding scheme obtained worse results on average for Equation 2 than URA.

Figures 3 through 5 show that the Unified Repartitioning Algorithm is able to compute partitionings with comparable or lower costs compared with either of the other two schemes. Specifically, when the Relative Cost Factor is set low (i. e., minimizing the edge-cut is the key objective), URA minimizes the cost function as well as the scratch-remap scheme and better than the multilevel diffusion scheme. Note that URA does quite well here, because when the RCF is set low, it means that the edge-cuts of the partitionings are

²These test sets were provide by Boris Kaludercic, HPC Product Coordinator, Computational Dynamics Ltd, London, England.

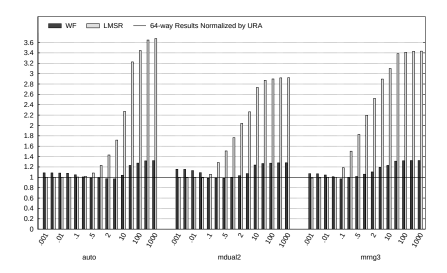


Figure 4: The cost function results obtained from the Unified Repartitioning Algorithm (URA) compared to the results obtained from optimized multilevel diffusion (WF) and scratch-remap (LMSR) algorithms on 64 processors of a Cray T3E.

primarily being compared. Therefore, in order to obtain costs that are similar to the scratch-remap scheme, URA must compute partitionings of similar edge-cut to a multilevel graph partitioner.

For the experiments in which the Relative Cost Factor is set high (i. e., minimizing the data redistribution cost is the key objective), the Unified Repartitioning Algorithm minimizes the cost function better than the multilevel diffusion scheme and much better than the scratch-remap scheme. URA beat the diffusion scheme here because it attempts to minimize the true cost function during multilevel refinement. The multilevel diffusion scheme, on the other hand, minimizes the edge-cut as the primary objective and the data redistribution cost as the secondary objective during refinement.

For the experiments in which the Relative Cost Factor was set near one, URA tended to do about as well as the diffusion-based scheme and somewhat better than the scratch-remap scheme in minimizing the cost function. It is interesting to note that the multilevel diffusion algorithm performs well in this region. This is because in the initial partitioning phase of this algorithm, the partitioning is balanced while aggressively minimizing the data redistribution cost. During the uncoarsening phase, the multilevel refinement algorithm focuses on aggressively minimizing the edge-cut. The result is that this scheme presents an almost even tradeoff between the edge-cut and the data redistribution cost.

The results presented in Figures 3 through 5 indicate that the Unified Repartitioning Algorithm is able to meet or beat the results obtained by either the scratch-remap or the multilevel diffusion repartitioner for a variety of experiments regardless of the value of the Relative Cost Factor. The other two schemes perform well only for limited ranges of values of the RCF.

Results from a Simulation of a Diesel Combustion Engine Table 2 gives the edge-cut, the total amount of data redistribution, and the cost function results for the optimized multilevel diffusion algorithm (WF), the optimized scratch-remap algorithm (LMSR), and URA on the diesel combustion engine test set with 16 processors of a Cray T3E. The numbers at the top of each column indicate the Relative Cost Factor of the experiments in that column. Table 2 shows that across the board, URA computes partitionings with lower costs than either of the other two schemes. These results confirm the trends shown in Figures 3 through 5. Table 2 also shows that URA is able to tradeoff one objective for another as the input value for RCF is changed. WF and LMSR compute the same partitioning regardless of the value for RCF.

| Metric | 0.001 | 0.01 | 0.1 | 1 | 10 | 100 | 1000 |
|-------------|--------|--------|------------|------------|-------------|-----------|------------|
| WF | | | | | | | |
| edge-cut | 19,059 | 19,059 | 19,059 | 19,059 | 19,059 | 19,059 | 19,059 |
| data redist | 45,491 | 45,491 | $45,\!491$ | 45,491 | 45,491 | 45,491 | $45,\!491$ |
| cost | 19,104 | 19,514 | 23,608 | $64,\!550$ | 473,969 | 4,568,159 | 45,510,059 |
| LMSR | | | | | | | |
| edge-cut | 12,022 | 12,022 | 12,022 | 12,022 | 12,022 | 12,022 | 12,022 |
| data redist | 73,312 | 73,312 | 73,312 | 73,312 | 73,312 | 73,312 | 73,312 |
| cost | 12,095 | 12,755 | 19,353 | 85,334 | 745,142 | 7,343,222 | 73,324,022 |
| URA | | | | | | | |
| edge-cut | 11,812 | 11,812 | 11,625 | 21,217 | 26,202 | 27,463 | 27,463 |
| data redist | 72,905 | 72,905 | $72,\!507$ | 41,006 | $37,\!330$ | 36,293 | 36,293 |
| cost | 11,885 | 12,541 | 18,876 | 62,223 | $399,\!502$ | 3,656,763 | 36,320,463 |

Table 2: The edge-cut, total amount of data redistribution, and cost function results of the adaptive graph partitioners WF, LMSR, and URA for various RCF values on problems derived from a particles-in-cells simulation on a Cray T3E. The numbers at the top of each column indicate the RCF of the experiments in that column.

| Graph | Scheme | 8-processors | 16-processors | 32-processors | 64-processors | 128-processors |
|--------|--------|--------------|---------------|---------------|---------------|----------------|
| auto | WF | 2.35 | 1.25 | 0.62 | 0.37 | 0.35 |
| auto | LMSR | 2.31 | 1.16 | 0.62 | 0.58 | 0.45 |
| auto | URA | 2.42 | 1.20 | 0.73 | 0.55 | 0.49 |
| mdual2 | WF | 3.29 | 1.61 | 0.80 | 0.48 | 0.41 |
| mdual2 | LMSR | 3.25 | 1.56 | 0.80 | 0.50 | 0.42 |
| mdual2 | URA | 3.21 | 2.54 | 0.87 | 0.53 | 0.56 |
| mrng3 | WF | 11.11 | 5.55 | 2.88 | 1.47 | 0.93 |
| mrng3 | LMSR | 11.13 | 5.54 | 2.86 | 1.47 | 0.96 |
| mrng3 | URA | 11.32 | 5.71 | 3.06 | 1.64 | 1.11 |

Table 3: Parallel run times of selected experiments for the adaptive graph partitioners WF, LMSR, and URA on a Cray T3E.

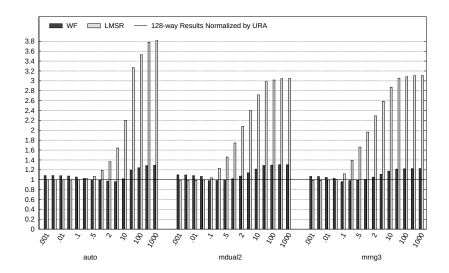


Figure 5: The cost function results obtained from the Unified Repartitioning Algorithm (URA) compared to the results obtained from optimized multilevel diffusion (WF) and scratch-remap (LMSR) algorithms on 128 processors of a Cray T3E.

| Scheme | 2-processors | 4-processors | 8-processors |
|--------|--------------|--------------|--------------|
| WF | 12.88 | 6.98 | 4.08 |
| LMSR | 12.90 | 6.86 | 4.04 |
| URA | 13.08 | 7.55 | 4.44 |

Table 4: Parallel run times of experiments performed on the graph *auto* for the adaptive graph partitioners WF, LMSR, and URA on a cluster of Pentium Pro workstations connected by a Myrinet switch.

Parallel Run Time Results Tables 3 and 4 give the run time results of the optimized multilevel diffusion algorithm (WF), the optimized scratch-remap algorithm (LMSR), and URA for selected experiments from Figures 3 through 5 on a Cray T3E and for similar experiments performed on up to 8 processors of a cluster of Pentium Pro workstations connected by a Myrinet switch. Tables 3 and 4 show that the repartitioning algorithms studied in this paper are very fast. For example, they are all able to compute a 128-way repartitioning of a four million vertex graph in about a second on 128 processors of a Cray T3E. Tables 3 and 4 show that URA tends to be slightly slower than the other two schemes. Note that all of the reported run times were obtained on non-dedicated machines. Therefore, these results contain a certain amount of noise. This reason, along with cache effects, explains the few super-linear speedups observed. The run time results of the diesel combustion engine follow these same trends, and so, were not reported.

4 Conclusions

We have presented a Unified Repartitioning Algorithm for dynamic load-balancing of scientific simulations and shown that this scheme is fast and effective. URA is significant because it is able to tradeoff the objectives of minimizing the edge-cut and the amount of data redistribution required to balance the load. The URA is also significant because it is a key component in developing tools for automatically performing dynamic load-balancing. Load-balancing tools such as DRAMA [10] and Zoltan [3] can measure the times required to perform inter-processor communications and data redistribution for an application, use this information to automatically compute an accurate Relative Cost Factor, and then call URA with the correct value as the input. Therefore, the user need not determine a good value each time load balancing is required.

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Related papers are available at: http://www-users.cs.umn.edu/~karypis.

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