Redundant arrays of distributed disks (RADD) can be used in a distributed computing system or database system to provide recovery in the presence of disk crashes and temporary and permanent failures of single sites. In this paper, we look at the problem of partitioning the sites of a distributed storage system into redundant arrays in such a way that the communication costs for maintaining the parity information are minimized. We show that the partitioning problem is NP-hard. We then propose and evaluate several heuristic algorithms for finding approximate solutions. Simulation results show that significant reduction in remote parity update costs can be achieved by optimizing the site partitioning scheme.

1. INTRODUCTION

Redundant disk arrays are used for the purpose of providing reliable storage while increasing the I/O bandwidth in high performance systems [1, 2]. Redundant disk arrays can also be used in a distributed setting to increase availability in the presence of temporary site failures, disk failures, or major disasters. Stonebraker and Schloss have proposed the redundant arrays of distributed disks (RADD) scheme [3] as an alternative to multicopy schemes, which are much more costly in terms of storage requirements. Cabrera and Long [4] have proposed the use of redundant distributed disk striping in a high speed local area network to support such I/O-intensive applications as scientific visualization, image processing, and recording and playback of color video. The RADD concept can also be used in multicomputer I/O subsystems such as the one proposed by Reddy and Banerjee [5] for hypercubes.

The IDA approach proposed by Rabin [6] provides another way to tolerate failures in distributed storage systems with limited extra storage cost. However, in that approach, updates are more costly since all the fragments of the dispersed data are needed to reconstruct the encoding which involves multiple remote accesses. In the case of RADD, a local update will generate a single remote access for updating the parity.

When RADDs are used, sites are grouped together to form a redundant array containing data and parity and capable of recovering from a single site failure. The size of each array is fixed and is determined by the tradeoff between the availability requirements of the system and the cost of the storage overhead. Hence, a large distributed data storage system may have to be divided into several arrays of fixed size. In this paper we look at the problem of partitioning the distributed storage system into fixed-size arrays in such a way as to minimize the cost of remote accesses that have to be performed to update the parity information. This problem is somewhat related to the problem of file allocation and replica placement in a distributed system, which has been studied extensively in the literature [7, 8]. However, the two problems are different in nature because, in the RADD case, there is one redundant item for N data items while in the file allocation problem each file is replicated several times. More importantly, in the replica placement problem there is no stringent constraint on the number of sites “sharing” a replica because. When the replica becomes unavailable, those sites can access the second nearest replica while in the RADD case there is a hard constraint on the number of sites in an array. Note that the assignment of sites to redundant arrays (parity groups) can occur after all decisions on placing the data have been made. Data placement decisions are governed by a different set of criteria and are more influenced by the read access patterns since reads are usually more frequent than updates. Decisions on site assignment to redundant arrays are based on the update rate at each site and the cost of communication between sites and are independent of the read access rate. Changing the assignment of sites to redundant arrays does not change the placement of the data. The purpose of site assignment is to reduce the cost of the parity traffic and does not directly affect the data traffic.

In the following section, we describe the RADD organization. In Section 3, we present the model used to formulate the problem mathematically and we prove that the problem is NP-hard. In Section 4, heuristic algorithms for solving the problem are described and results from an
In terms of performance, both approaches require one remote access per update, while the RADD scheme may be mathematically formulated as follows:

\[ P_{\text{new}} = (D_{\text{old}} \oplus D_{\text{new}}) \oplus P_{\text{old}}. \]

Spare blocks are provided to make it possible to reconstruct data blocks that become inaccessible due to site failure. The failed data block is reconstructed by XORing all other data blocks and the parity block in its parity group. The bitwise parity of the data blocks in each parity group is computed and written at a different site. In Fig. 1, \( D_{ij} \) denotes a data block, \( P_i \) denotes a parity block, and \( S_j \) denotes a spare block, all at site \( i \). The number under block in the first column of the figure denotes the physical block number on disk. Each row in the figure represents a parity group. The position of the parity block is rotated among the sites in order to avoid creating a bottleneck at the site where parity is stored. For every update to one of the data blocks in the parity group, the parity block needs to be updated using the following formula:

\[ P_{\text{new}} = (D_{\text{old}} \oplus D_{\text{new}}) \oplus P_{\text{old}}. \]

The problem of partitioning the sites into arrays of size \( N \) in such a way that parity update costs are minimized can be mathematically formulated as follows:

\[ N = K + 2 \]

where \( K \) denotes the number of data blocks per parity group when \( N = K + 2 \) denotes the number of sites in a distributed disk array. The storage overhead for the parity and spare blocks required by RADDs is \((200/K)\%\) compared to a 100% overhead for the case of two copy schemes. In terms of performance, both approaches require one remote access per update, while the RADD scheme may require two additional local accesses per update to read the old data and old parity in order to compute the new parity. Under failure, RADD will perform much worse than the two-copy scheme because it requires \( K \) remote accesses for reconstructing a data block from a failed site. However, if failures are expected to be rare, the performance degradation associated with RADD may be justifi-

<table>
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<th>Site2</th>
<th>Site3</th>
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<td>D43</td>
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</tr>
</tbody>
</table>

FIG. 1. Organization of a distributed redundant disk array \((N = 6)\).

expensive evaluation are presented. In Section 5 we develop heuristics with guaranteed bounds on the deviation from the optimal cost. In Section 6 we address the issue of hot spots and non-uniform site capacity and discuss the use of RADD for disaster recovery in OLTP systems as well as the issue of when and how often site reassignment should be initiated.

2. DISTRIBUTED REDUNDANT DISK ARRAY ORGANIZATION

The RADD organization is shown in Fig. 1. The data at each site are partitioned into blocks. Data blocks from different sites are grouped into a block parity group. The bitwise parity of the data blocks in each parity group is computed and written at a different site. In Fig. 1, \( D_{ij} \) denotes a data block, \( P_i \) denotes a parity block, and \( S_j \) denotes a spare block, all at site \( i \). The number under block in the first column of the figure denotes the physical block number on disk. Each row in the figure represents a parity group. The position of the parity block is rotated among the sites in order to avoid creating a bottleneck at the site where parity is stored. For every update to one of the data blocks in the parity group, the parity block needs to be updated using the following formula:

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\[ P_{\text{new}} = (D_{\text{old}} \oplus D_{\text{new}}) \oplus P_{\text{old}}. \]

3. THE MODEL

We model the distributed computing system as an undirected connected graph \( G = (V, E) \), where \( V \) is the set of sites and each edge \( e \in E \) represents a bidirectional communication link between two sites. For each \( e \in E \), \( w_e \) denotes the cost of communication over link \( e \). We assume that if \( n \) is the number of sites in \( V \) then \( n = mN \) for some integer \( m \). We assume that the site capacity is uniform. In Section 6.2 we show how to deal with nonuniform site capacity. In the pattern shown in Fig. 1, the parity blocks of the \( N - 2 \) data blocks from site \( i \) reside on sites \((i + 1) \mod N \) through \((i + N - 2) \mod N \). If the same pattern is repeated throughout the range of blocks then there will be no parity update traffic from site \( i \) to site \((i - 1) \mod N \). In order to make the problem symmetrical and thus easier to tackle, we assume that for the next set of \( N \) blocks the pattern shown in Fig. 2 is used. In all, there are \( N - 1 \) such patterns obtained by changing the distance between the parity block and the spare block on a given row. These \( N - 1 \) patterns should alternate throughout the range of blocks so that update traffic from a given site is distributed over the remaining \( N - 1 \) sites. This will also provide more load balancing for the parity update traffic in the array.

Let \( \mu_v \) designate the rate of update accesses to data blocks at site \( v \). Each update will cause communication between the site where the update took place and the site holding the parity for the given data block. At each site the set of data blocks that have their corresponding parity blocks on the same remote site is called a data group. To simplify the model, assume that the \( N - 1 \) data groups share equally the update rate. This implies that the rate at which site \( v \) sends parity update information to each other site in its redundant array is \( \alpha_v = \mu_v/(N - 1) \). This assumption is supported by the fact that consecutive data blocks have their parity blocks on different sites, which implies that accesses to a heavily used file that is stored on consecutive disk blocks will be spread over different data groups. In Section 6, the above assumption will be removed.

The problem of partitioning the sites into arrays of size \( N \) in such a way that parity update costs are minimized can be mathematically formulated as follows:

\[ \alpha_v = \mu_v/(N - 1) \]

FIG. 2. Alternative placement pattern for parity and spare blocks.
Problem 1 (SP). Find a partition of $V$ into $m$ disjoint subsets $V_1, V_2, ..., V_m$ of size $N$ such that if $d(u, v)$ denotes the length of the shortest path between $u$ and $v$ then $\sum_{i=1}^{m} \sum_{u \in V_i} \lambda_u \sum_{v \in V_i} d(u, v)$ is minimum.

**Theorem 1.** Problem SP is NP-hard for any fixed $N \geq 3$.

**Proof.** We prove that problem SP is NP-hard by showing that there is a polynomial time transformation from the problem of partitioning a graph into cliques of size $N$ to problem SP. The partition into cliques of size $N$ (PC) problem can be stated as follows:

**Instance.** A graph $G = (V, E)$, where $|V| = Nm$ for some positive integer $m$.

**Problem.** Is there a partition of $V$ into $m$ disjoint subsets $V_1, V_2, ..., V_m$ such that the subgraph of $G$ induced by $V_i$ is a clique of size $N$ (complete graph with $N$ nodes)?

PC is NP-complete for any fixed $N \geq 3$ (see partition into isomorphic subgraphs [9]). To transform an instance of PC into an instance of SP, it is sufficient to set $\lambda_v = 1$ for all $v \in V$, and $w_e = 1$ for all $e \in E$. Then graph $G$ can be partitioned into cliques of size $N$ if and only if the cost of the optimal solution to the above instance of problem SP is $n(N - 1)$. ■

The cost function $\sum_{i=1}^{m} \sum_{u \in V_i} \lambda_u \sum_{v \in V_i} d(u, v)$ can be rewritten as $\sum_{i=1}^{m} \sum_{u \in V_i} \sum_{v \in V_i} \lambda_u \lambda_v d(u, v) = \sum_{i=1}^{m} \sum_{u \in V_i} \sum_{v \in V_i, u \neq v} D(u, v)$, where $D(u, v)$ is defined as $D(u, v) = (\lambda_u + \lambda_v) d(u, v)$. In this form the general problem is reduced to a uniform load problem with the pseudo-distance $D$ replacing $d$. However, $D$ is not a true distance since it does not necessarily satisfy the triangular inequality.

4. APPROXIMATION ALGORITHMS

4.1. Description of the Heuristics

The first heuristic is based on a greedy strategy that consists of satisfying first the sites with the largest update rate. Let $\Lambda$ be the list of update rates for all sites. When sites are grouped into clusters (redundant arrays) their update rates are removed from $\Lambda$ and replaced by a single update rate for the cluster. The cluster update rate is the average update rate of the sites in the cluster.

**Algorithm 1**

**Step 1.** Select the largest value in $\Lambda$ and let $a$ be the corresponding site (or cluster). Find the site (or cluster) $b$ such that merging $a$ and $b$ results in the smallest increase in the cost function. Merge the two sites (or clusters) if the resulting cluster has less than $N$ sites and the total number of clusters does not exceed $m$. If the clusters cannot be merged, find the next best choice for $b$ and repeat. Remove the update rates of the merged sites (or clusters) from $\Lambda$ and replace them with the cluster update rate.

**Step 2.** Repeat Step 1 until $m$ clusters having $N$ sites each have been formed.

The computational cost of Algorithm 1 is $O(Nn^2)$. But it requires that the all-pair shortest path algorithm be performed first, which requires $O(n^3)$ operations.

The second approach consists of two stages: in the first stage $m$ sites are identified to be used as cluster seeds and in the second stage the remaining sites are allocated to the clusters to form $m$ subsets of $N$ sites each.

**Algorithm 2**

**Step 1.** Select the two sites with the largest distance between them and include them in the set $S$ of cluster seeds.

**Step 2.** Select the site $v$ with the largest average distance to the sites already in $S$ and add it to $S$.

**Step 3.** Repeat Step 2 above until $|S| = m$. Each cluster initially contains one of the $m$ seeds in $S$.

**Step 4.** For each of the $m$ clusters, compute the average update rate of the sites in the cluster. In decreasing order of their average update rate, allocate to each cluster the site that is closest to it in terms of the pseudo-distance $D$.

**Step 5.** Repeat Step 4 above until all sites have been allocated to the $m$ clusters.

We use the pseudo-distance metric $D$ in Step 4 because it provides the actual increase in the cost function of a cluster when a node is added to it. The computational cost of the Algorithm 2 is $O(Nn^2)$. It also requires that the all-pair shortest path algorithm be performed first.

The third approach is based on the hierarchical clustering technique [10]. We use the distance matrix whose entries are $d(u, v)$ for all $u, v \in V$. Clusters are formed by merging together sites or smaller clusters that are close to each other. When two sites (or clusters) are grouped together, the distance matrix is modified by eliminating the columns and rows corresponding to the merged sites (or clusters) and replacing them with a single column and a single row reflecting the average distance between the merged sites and other sites (or clusters). The procedure is as follows:

**Algorithm 3**

**Step 1.** Find the smallest entry in the distance matrix and merge the two sites (or clusters) together if the resulting cluster has $N$ sites or less and if the total number of clusters does not exceed $m$. If any of the latter conditions is not satisfied, select the next smallest entry and repeat. Once two sites (or clusters) have been merged, update the distance matrix and the number of clusters accordingly.

**Step 2.** Repeat Step 1 above until $m$ clusters having $N$ sites each have been formed.

The complexity of Algorithm 3 is $O(n^3)$.

After an initial partition has been found, the following procedure may be used to improve it.
Each entry represents the average over 1000 randomly generated graphs. The costs of the approximate solutions are within 10% of the cost of the optimal solution. In the first column of the table, we have listed the cost of a random solution. For each number the half-width of the corresponding 95% confidence interval is shown between parenthesis.

Since, in the first experiment, an exhaustive search was used to find the optimal solution, the number of nodes \( n \) could not be very large. In a second experiment, we compared the performance of the three heuristics for larger values of \( n \). Figure 3 shows the results for the second experiment. For clarity of the figure, we plotted the cost of the approximate solution divided by 1000. For each data point, the 95% confidence interval is shown. In the case \( N = 10 \), Algorithm 3 outperforms Algorithms 1 and 2 for all values of \( n \) except \( n = 20 \), in which case Algorithm 2 performs better. For the first and second environments, Algorithm 1 outperforms Algorithm 2 for large values of \( n \), but for the last environment Algorithm 2 outperforms Algorithm 1. For \( N = 5 \), Algorithm 2 does not do very well except in the last environment in which the range of site loads is much larger than the range of edge weights. Algorithm 1 performs best in the first two environments.

The main point that can be deduced from this experiment is that, in spite of the fact that Algorithm 3 does not use any information about site loads, it outperforms the other two algorithms when \( n \) and \( N \) are relatively large and, in the other cases, its performance is always close to that of the best algorithm. This means that, in a large system, it is more important to minimize the sum of the edge weights within each cluster than to use the greedy approach that attempts to assign to the sites with large loads their nearest neighbors. Since site loads vary with time, the solution found by Algorithm 3 will remain close to optimal as the site loads change while solutions based on estimates of site loads will degrade with time as the site loads deviate from the estimates. This is especially true for large \( N \). A large value for \( N \) means lower storage costs but also lower reliability and worse performance under failure.

### 5. Heuristics with Performance Guarantees

The heuristics described in Section 4 provide a good approximate solution. However, there is no guarantee that the approximate solution will not diverge significantly from the optimal one in certain cases. In this section, we seek to find a heuristic that has a bound on the error between the approximate solution and the optimal one. We develop

<table>
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<th>( K_\alpha, K_\nu )</th>
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<th>Algorithm 2</th>
<th>Algorithm 3</th>
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such a heuristic first for the case of a system with balanced load, \( \lambda_v = \lambda \), for all \( v \in V \), and uniform edge weights, then we look at the more general case of a balanced load system with arbitrary edge weights. Since a problem with arbitrary site loads can always be transformed into a problem with uniform site load as shown in Section 3, then the heuristic for the balanced load case with arbitrary edge weights will also provide performance guarantees for the arbitrary load case.

5.1. Balanced Load and Uniform Edge Weights

The heuristic requires the use of a spanning tree with many leaves. The problem of finding a spanning tree with a maximum number of leaves is NP-hard [9], however, there exist polynomial time algorithms for generating spanning trees with many leaves. Typically these methods guarantee that a certain fraction of the nodes will be leaves. The fraction of leaves is a function of the minimum degree \( k \) of the graph. Kleitman and West proved the following result [13]:

**Theorem 2 (Kleitman–West).** If \( k \) is sufficiently large, then there is an algorithm that constructs a spanning tree with at least \( (1 - b \ln k/k)n \) leaves in any graph with minimum degree \( k \), where \( b \) is any constant exceeding 2.5.

It was also conjectured that a spanning tree can be constructed with a larger fraction of leaves. More specifically, Linial conjectured that the number of leaves could be at
least \((k - 2/k + 1)n + c_k\). This stronger result was proved for \(k = 3\) with \(c_3 = 2\) and for \(k = 4\) with \(c_4 = 8/5\) [13].

**Algorithm**

**Step 1.** Find a spanning tree with many leaves.

**Step 2.** Partition the spanning tree into \(m\) clusters of \(N\) nodes each using procedure `Partition_Tree` described below.

The partition found for the tree will be used as the approximate solution for the partitioning problem in the original graph. We first describe a basic version of the procedure `Partition_Tree` which insures that every edge in the tree is used by at most two clusters. Then we describe an optimization that reduces the cost in the tree but that is not needed to establish the bound on the cost of the heuristic solution. In the description of the procedure `Partition_Tree`, we assume that the tree is levelized starting from the root. Figure 4 shows an example of a tree partitioned using this procedure.

**Procedure Partition_Tree**

The procedure partitions the tree from the bottom up and from left to right. As the clusters are built, whenever the size of a cluster reaches \(N\) nodes, that cluster is removed from the tree. Starting from the deepest leaf of the leftmost branch in the tree, the leaf is assigned to the first cluster. After a node has been assigned to a cluster, its sibling to the right is considered next. If no siblings are left to right of the node then the parent is assigned next. If the sibling to the right is a leaf, it is included in the cluster, otherwise the leftmost branch of the subtree rooted at the sibling is followed to its deepest leftmost leaf and that leaf is included in the cluster. Then the procedure continues from that point moving to the right sibling (if any) or to the parent in the same fashion. When a node is to be assigned, it is either assigned to the current cluster if that cluster has not reached \(N\) nodes or a new cluster is formed and the node assigned to it. The tree remains connected as newly completed clusters are removed.

**Theorem 3.** The cost (HEU) of the approximate solution found using a spanning tree with many leaves and the cost (OPT) of the optimal solution satisfy the following relationship:

\[
\frac{\text{HEU}}{\text{OPT}} \leq 2\alpha + (1 - \alpha) \frac{N^2}{N - 1},
\]

where \(\alpha\) is the fraction of leaves in the spanning tree.

**Proof.** We need to establish an upper bound on the cost of the approximate solution and a lower bound on that of the optimal one. The cost in the graph of the approximate solution is at most the cost of that solution in the tree. We evaluate the cost in the tree by adding up the contributions of each edge in the spanning tree to the overall cost. If an edge connects a leaf node to the tree it will be referred to as a leaf edge, otherwise it will be called an internal edge. A leaf edge will be used in only one cluster and it will be used only for communication between the leaf node and the other \((N - 1)\) nodes in the cluster. Therefore the contribution of a leaf edge to the overall cost is 2\((N - 1)\). An internal edge will be used in at most two clusters and in each cluster it will be used by \(i\) nodes to communicate with the other \(N - i\) nodes in the cluster. If \(\alpha\) designates the fraction of leaf nodes in the tree, we have

\[
\text{HEU} \leq \alpha n \times 2(N - 1) + (n - 1 - \alpha n)
\]

\[
\times 2 \times \max_{1 \leq i < N - 1} 2i(N - i)
\]

\[
\leq n(N - 1)(2\alpha + (1 - \alpha)N^2/(N - 1)).
\]

For the cost of the optimal solution, an obvious lower bound is the cost in a complete graph, which is \(n(N - 1)\). Hence, \(\text{HEU}/\text{OPT} \leq 2\alpha + (1 - \alpha)N^2/(N - 1)\).

As stated in Theorem 2, for large \(k\), \(\alpha\) converges to 1 and the upper bound approaches 2. Note that it is reasonable to assume that the minimum degree will be large in practice because the underlying network has to have sufficient connectivity to enable communication under node and link failures, and hence, has to have a reasonably large minimum degree.

The complexity of the algorithms for generating trees with many leaves [13] is \(O(|E|)\). The complexity of the `Partition_Tree` procedure is \(O(n)\).

There is an optimization to procedure `Partition_Tree` that reduces the cost of the solution in the tree (not necessarily the cost in the original graph) by reducing the number of tree edges that are used by two clusters. It can be described as follows: Consider the case where one subtree has been processed and there remains an incomplete cluster (less than \(N\) nodes) and assume that there is a subtree rooted at the sibling to the right. Procedure `Partition_Tree` would complete the cluster using the lowest leftmost nodes of the right subtree. However, if the right subtree is deep, a number of intermediate edges will contribute to the cost of communicating (in the tree) between the two components of the newly completed cluster. This can be avoided.
by successively removing from the right subtree complete clusters formed by connected branches of exactly N nodes whose removal does not disconnect the tree and then completing the cluster from the left subtree. After those clusters are removed Partition_Tree proceeds as described above to complete the cluster with what remains of the right subtree. The requirement that the removed clusters be connected ensures that edges remaining in the right subtree are not used by any of the removed clusters. Those remaining edges may then be used both by the cluster subtree and by a cluster formed with the remaining nodes in the right subtree.

5.2. Balanced Load and Arbitrary Edge Weights

For arbitrary edge weights the problem of finding a heuristic with guaranteed performance bounds is much harder. In the following we describe a heuristic for which a worst case performance bound can be established. The bound is more significant for systems where link communication costs (edge weights) do not vary widely. The heuristic consists of finding a minimum spanning tree, partitioning the tree into clusters using procedure Partition_Tree and using that partition as an approximate solution. The following result will be used to establish a lower bound on the cost of the optimal solution.

**Lemma 1.** In a complete graph, the average weight of the edges in a minimum spanning tree is at most the average weight of all edges.

**Proof.** We use induction on the number of nodes n. The lemma is obviously true for n = 2 or n = 3. Suppose it is true for graphs with n - 1 nodes and consider an n-node graph. Select node v such that the average weight of edges incident on v is at least the average weight of all edges in the graph. Remove v from the graph and find a minimum spanning tree in the remaining (n - 1)-node graph. Then add to this spanning tree the lightest edge e* connecting v to the other nodes to form an n-node spanning tree. Let MST_{n-1} and MST_n be the total weights of the (n - 1)-node and the n-node spanning trees, respectively. Let e'(v) be the set of edges incident on v. Using the induction hypothesis, we have

\[
\text{MST}_{n-1} \leq \frac{\sum_{e \in E \setminus \{v\}} w_e}{(n - 1)(n - 2)/2}.
\]

Therefore

\[
\text{MST}_n \leq \text{MST}_{n-1} + w_{e'} \leq \frac{\sum_{e \in E \setminus \{v\}} w_e}{(n - 1)/2} + \frac{\sum_{e \in e'(v)} w_e}{n - 1} \leq \frac{\sum_{e \in E \setminus \{v\}} w_e}{(n - 1)/2} + \frac{\sum_{e \in e'(v)} w_e}{n - 1} + \sum_{e \in E \setminus \{v\}} w_e \leq \frac{\sum_{e \in E} w_e}{n - 1} \leq \frac{\sum_{e \in E} w_e}{n/2}.
\]

Hence, the average weight of the edges in the minimum spanning tree is MST_{n/2}(n - 1) \leq \sum_{e \in E} w_e/(n(n - 1)/2).

To obtain a lower bound on the cost of the optimal solution, we consider the optimal partition and we build a spanning tree by first finding a minimum spanning tree in each cluster and then replacing each cluster by a single node and connecting each pair of these nodes by the lightest edge linking the initial clusters. An intercluster minimum spanning tree is then found. The intracluster spanning trees along with the intercluster spanning tree form a spanning tree for the entire graph.

**Lemma 2.** The list of edge weights of the intercluster minimum spanning tree (ICMST) is included in the list of edge weights of the global minimum spanning tree (GMST).

**Proof.** Let e be an edge in the ICMST that does not appear in the GMST. Let u and v be its endpoints in the original graph and let w be its weight. The path in the GMST from u to v induces a path in the intercluster graph from the cluster of u to that of v. If the path is a single edge then this edge must have weight w and could replace the edge e in the ICMST. If the induced path has more than one edge then, since the ICMST cannot contain a cycle, some of the edges on the induced path must not appear in the ICMST. At least one of these induced edges that do not appear in the ICMST forms a cycle containing e when added to the ICMST. Let e' be such an edge: e' must have weight at most w otherwise it could be replaced in the GMST by (u, v) to obtain a spanning tree with a smaller cost. In addition e' cannot have weight less than w because it would then be possible to replace e by e' in the ICMST and obtain a smaller intercluster spanning tree. Hence the weight of e' is w and we could remove e and replace it with e' in the ICMST. This process can be repeated until all edges in the ICMST also appear in the GMST.

The following theorem establishes a bound on the cost of the heuristic based on finding a minimum spanning tree in the graph and then using Partition_Tree to find a partitioning into clusters.

**Theorem 4.** The cost (HEU) of the approximate solution found using a minimum spanning tree and the cost (OPT) of the optimal solution satisfy the following relationship:

\[
\frac{\text{HEU}}{\text{OPT}} \leq \frac{N}{\text{MST}} \frac{\text{MST} - (m - 1)\bar{w}}{(n(n - 1)/2)}.
\]

where MST is the total weight of the edges in the minimum spanning tree and \(\bar{w}\) is the average weight of the m - 1 heaviest edges in the minimum spanning tree.

**Proof.** In evaluating an upper bound on the cost of the approximate solution, we follow the same procedure as in the proof of Theorem 3 but we will not distinguish between leaf edges and internal edges. Each edge e in the tree will
be used by at most two clusters and the contribution of e to the overall cost is bounded by $2 \times w_e \times \max_{1 \leq i \leq N-1} 2i(N - i)$. Hence, we have $\text{HEU} \leq N^2 \text{MST}$.

Let $\text{MST}_i$ be the weight of the minimum spanning tree of cluster $i$ for $1 \leq i \leq m$ and $\text{MST}_c$ be the weight of the intercluster tree. The intracluster minimum spanning trees and the ICMST form a spanning tree in the original graph. The total weight of the edges in that spanning tree is at least $\text{MST}_c + \text{MST}_i \geq \text{MST}$. By Lemma 2, every edge in the ICMST is also in the GMST. Hence $\text{MST}_c \leq (m - 1)\overline{w}$. This yields $\sum_{i=1}^{m} \text{MST}_i + (m - 1)\overline{w} \geq \text{MST}$. Let $\text{OPT}_i$ be the contribution to the optimal cost by cluster $i$. The average cost of the edges in cluster $i$ is $\text{OPT}_i / (N(N - 1))$ and the average cost of the edges in the corresponding spanning tree is $\text{MST}_i / (N - 1)$. Applying Lemma 1, we have $\text{OPT}_i / N \geq \text{MST}_i$, therefore $\text{OPT} \geq N(\text{MST} - (m - 1)\overline{w})$.

Let $r$ be the ratio of the largest edge weight to the smallest edge weight. A looser but simpler bound than the one established in Theorem 4 can be derived using the parameter $r$. To do so, we first rewrite the bound as follows:

$$\frac{\text{HEU}}{\text{OPT}} \leq N \frac{\text{MST}}{\text{MST} - (m - 1)\overline{w}}$$

$$= N \left(1 + \frac{(m - 1)\overline{w}}{\text{MST} - (m - 1)\overline{w}}\right)$$

$$= N \left(1 + \frac{(m - 1)\overline{w}}{(n - m)\overline{w}}\right).$$

FIG. 5. Evaluation of the heuristics for the refined model.
where \( \bar{w} \) is the average weight of the \( n-m \) lightest edges in the GMST. Since \( \bar{w}/\bar{w} \leq r \) and \( n = mN \), we have

\[
\text{HEU/OPT} \leq N \left( 1 + \frac{m-1}{n-m} r \right) < N(1 + r/(N-1)).
\]

6. GENERALIZATION AND APPLICATION OF THE MODEL

6.1. Non-Uniform Load within Site

In our model, we assumed that each site sends parity updates to each other site in its partition at the same rate. This implies a uniform update rate to each of the \( N-1 \) data groups of a given site that have parity information on each of the \( N-1 \) other sites. If the update rate information for each data group at each site is available then the model can be refined to account for the difference in the rate of parity update requests issued by a given site and destined to the other sites in the array. The refined model should yield better results in the presence of static hot spots. The update rate \( \lambda_u \) of site \( u \) is replaced by \( N - 1 \) update rates \( \lambda_{u1}, ..., \lambda_{u(N-1)} \) corresponding to each of its data groups. In this case, an obvious optimization would be to have the parity of the \( i \)th most frequently accessed data group of a given site placed on the \( i \)th nearest site in its partition. We call this optimization \texttt{LocalOpt}. Note that \texttt{LocalOpt} can be implemented without having to reshuffle the data on disk by saving the permutation describing the remapping of the \( N-1 \) data groups for each site and using it to route parity update requests to the proper site. Given the above optimization, the algorithms of Section 4 with some minor modifications can still be used to partition the sites. The site update rate used in Algorithm 1 and 2 is set to the sum of all \( N-1 \) data group update rates at that site. We have evaluated the three algorithms of Section 4 in the case of the refined model, along with a new greedy strategy that looks at data groups instead of sites and tries to place the parity of the data groups with the largest update rates on the closest sites. Details of the greedy algorithm are provided in the Appendix.

Figure 5 shows the results of the comparison between the four algorithms. The results shown assume that \texttt{LocalOpt} is performed. The individual data group update rates are chosen randomly from the interval \([1, K_s]\) while the edge weights are chosen from \([1, K_w]\). We found that Algorithms 2 and 3 perform best for \( N = 10 \) with Algorithm 2 being the winner for lower values of \( n \) while Algorithm 3 is better for the high values of \( n \). For \( N = 5 \) Algorithm 3 performs best in almost all situations. The reason that Algorithm 3 performs better for \( N = 5 \) in this case compared with the uniform load case (Fig. 3) can be explained by the fact that the site loads have smaller variance because they are the sum of \( N-1 \) rates drawn from the uniform distribution over \([1, K_s]\). The performance of the greedy algorithm indicates that basing assignment decisions on individual data group loads produces poorer results than using total site loads.

We also found that the parity assignment within a cluster is as important as the problem of partitioning the sites into clusters. Using \texttt{LocalOpt} reduces the cost of the solution by 15 to 20%. This is shown in Fig. 6 for the case \( N = 10, K_w = 100, \) and \( K_s = 100 \). Similar results were obtained for the other environments.

6.2. Non-Uniform Site Capacity

The case of nonuniform site capacity can be handled in the same fashion as proposed by Stonebraker and Schloss [3]. We assume that the total number of disks is \( np \) for some integer\(^5\) \( p \) and that the number of disks at any given site is at most \( p \). The system could then be partitioned using the following procedure.

\begin{enumerate}
  \item Select the \( N|V/N| \) sites with the largest number of disks and apply one of the partitioning algorithms described in the previous sections to assign one disk from each of the selected sites to an array.
  \item Remove the assigned disks and remove sites with no disks left.
  \item Repeat the above steps until all disks have been assigned.
\end{enumerate}

Nonuniform disk capacity can be dealt with by using logical disks of size \( B \) blocks such that the site capacities are multiples of \( B \) [3].

6.3. Disaster Recovery in OLTP Systems

Disaster recovery is an important issue in on-line transaction processing (OLTP) systems [14–16]. However, in such systems, updating the remote parity after each disk

\(^{5}\)This replaces the assumption that \(|V| = mN\).
Typically, disaster recovery in OLTP systems is implemented by duplicating the data of a given site at a remote backup site and shipping redo log information to the backup site where the updates are applied to the backup database. There are two approaches used in shipping the log [17]. In the first approach, the log records are sent to the backup at commit time and the transaction waits for an acknowledgment before it is allowed to commit. No transactions are lost in this case. This configuration is called 1-safe. In the second approach, log records are sent to the backup site and shipped to that site while the transaction waits for an acknowledgment. If the updates performed by the transaction involve only one of the \( N-1 \) data groups then only one remote message has to be sent by the committing transaction and the delay will be the same as in the traditional remote backup scheme. The advantage of RADD over the traditional schemes is that it uses much less storage space than full duplication.

Our model can still be used to solve the site assignment problem in both of the above implementations. However, instead of using the update rate at each site, the frequency of the periodic updates should be used in the 1-safe case and the update transaction rate should be used in the 2-safe case.

Another optimization that might be useful in OLTP environments consists of using the scheme proposed by Bhide and Dias in [18] to reduce the number of random I/O's performed in updating the parity at the remote site. The scheme consists of storing the parity updates in non-volatile memory or sequentially on a dedicated disk and then periodically propagating them to their permanent locations. The scheme was originally proposed for use with a RAID level 4 organization [1] to reduce the load on the parity disk. When the parity updates are stored sequentially on a dedicated disk, disk sorting is used to apply the parity updates to their permanent location.

6.4. Applying the Algorithms

Another important question is when and how often to apply the algorithm in order to obtain a lower cost site assignment. Clearly the algorithms can be used when the RADD scheme is first implemented as long as information on site loads is available. As these loads change, the performance of the system degrades and the site assignment may need to be modified. Changing the site assignment is a costly operation. It involves reading large amounts of data to recompute the new parity and then updating the parity. This operation should be performed when the following two conditions are met: (1) the difference between the cost of the current assignment and the cost of the best solution found by the algorithms should be large enough, and (2) the parameters of the system (site loads) should be relatively stable so that the benefits of the new site assignment last long enough to offset the cost of performing the reassignment.

The cost of reassignment can be reduced if some clusters are kept unchanged. Hence one might be better off choosing a solution that is not the best possible but that preserves most of the current clustering. Procedure Improve described in Section 4 can be used to perform a limited number of swaps that decrease the cost of updating the parity without a full scale reassignment.

7. SUMMARY

We looked at the problem of partitioning the sites of a distributed storage system into redundant disk arrays while minimizing the communication costs for updating the parity information. The problem was shown to be NP-hard in its general form. Several heuristic methods were investigated to obtain approximate solutions to the site partitioning problem. It was found that the heuristic that minimizes the sum of distances between sites within each cluster (Algorithm 3) performs consistently well in all environments, especially in large systems with a relatively large array size. In such systems, the above approach outperforms greedy methods that attempt to satisfy first the sites with the largest loads by placing their nearest neighbors in their partition. The solutions produced by Algorithm 3 are also more robust because they provide good performance under different site loads. Guaranteed upper bounds were established on the deviation from the optimal cost for some of the heuristics. It was also found that modifying the parity assignment within each cluster to place the parity of the heavily accessed data groups on the nearest sites within the cluster can significantly decrease the parity update cost. Finally, we discussed implementations of the RADD scheme for disaster recovery in OLTP systems and described various optimizations that can be helpful in those environments.

APPENDIX

Algorithm Greedy

Let \( \Lambda \) be the list of update rates for all data groups at all sites.

Let \( p_v \) be the number of site \( v \)'s partition. Initially \( p_v = -1 \) for all \( v \in V \).

Let \( n_i \) be the number of sites in partition \( i \). Initially, \( n_i = 0 \). Assume \( n_{-1} = 1 \) throughout.

Let \( k \) be the current number of partitions. Initially \( k = 0 \).
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Let \( V \) be the set of vertices in the graph, and let \( l = 0 \).

Step 1. Select the largest value \( \lambda \) in \( \Lambda \) and let \( a \) be the corresponding site. If \( n_{p_a} = N \) go to Step 4.

Step 2. Find the site \( v \) in \( \mathcal{V} \) that is nearest to \( u \) and satisfies \( p_u = p_v \) for all \( v \in V \) and \( k < m \). If none exist go to Step 4.

Step 3. Remove \( v \) from \( \mathcal{V} \).

Step 4. Remove \( \lambda \) from \( \Lambda \).

Step 5. If \( \sum_i n_i < n \), go to Step 1; otherwise stop.

The algorithm is similar to Algorithm 1 in that it tries to satisfy first the nodes with the highest data group update rates. The complexity of the algorithm is \( O(Nn^2) \), but as in the case of Algorithm 1, it requires the all-pair shortest path algorithm.

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REFERENCES


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