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AN ANALYSIS OF SCATTER DECOMPOSITION

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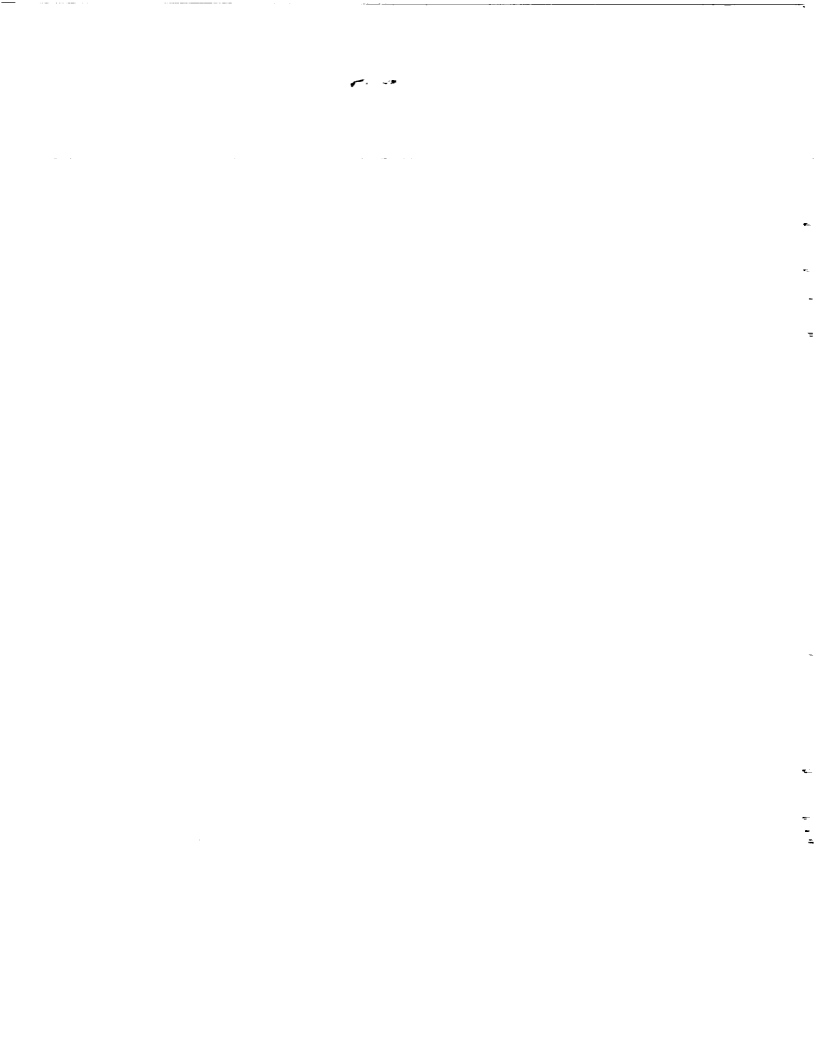
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An Analysis of Scatter Decomposition

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Abstract

This paper provides a formal analysis of a powerful mapping technique known as scatter decomposition. Scatter decomposition divides an irregular computational domain into a large number of equal sized pieces, and distributes them modularly among processors. We use a probabilistic model of workload in one dimension to formally explain why, and when scatter decomposition works. Our first result is that if correlation in workload is a convex function of distance, then scattering a more finely decomposed domain yields a lower average processor workload variance. Our second result shows that if the workload process is stationary Gaussian and the correlation function decreases linearly in distance until becoming zero and then remains zero, scattering a more finely decomposed domain yields a lower expected maximum processor workload. Finally we show that if the correlation function decreases linearly across the entire domain, then among all mappings that assign an equal number of domain pieces to each processor, scatter decomposition minimizes the average processor workload variance. The dependence of these results on the assumption of decreasing correlation is illustrated with situations where a coarser granularity actually achieves better load balance.

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1 Introduction

Scatter decomposition [1], (also described as modular mapping [4]) is an effective method for parallelizing a large class of irregular scientific programs that are tied to physical domains. Examples include a wide variety of techniques for numerically solving time dependent partial differential equations, and other, less numerical domain-oriented simulations. Scatter decomposition divides the domain into a set of rectangular regions with the same spatial size and geometry. The regions are labeled using Cartesian coordinates, and are mapped to processors by applying the mod function to the label in each coordinate. For example, Figure 1 shows how a two dimensional irregular grid for a PDE is decomposed into strips (marked by the heavy lines) and assigned to processors. The execution of all workload related to a subregion is a basic unit of schedulable work which we call a *cluster*. A cluster's granularity is controlled by the parameters defining the region size, in this case the strip width.

Scatter decomposition's success lies in its ability to balance workload without ever actually analyzing it. Any region of high workload tends to be subdivided and distributed among processors. Scatter decomposition is a technique applied to many problems in many contexts [1, 2, 4, 5, 9, 11, 14, 17]. Its success has been explained informally in [1] and [4], by appealing to the physics and numerics of many scientific computations. While these explanations suffice for most practitioners, the literature lacks a full formal analysis of why scatter decomposition balances workload. This paper provides some such analysis, identifying model assumptions under which scatter decomposition can be expected to effectively balance load. As such, our work is a necessary prerequisite for any future formal treatment of the very important problem of managing the inherent tensions between load imbalance and communication costs in a scatter decomposition.

The object of this paper is to construct and analyze a performance model to explain when and why scatter decomposition works. The model is based on a number of simplifying assumptions to promote tractability. As such, it should not be viewed as a model that accurately *predicts* performance quantitatively. Rather, it should be viewed as a model that *explains* performance qualitatively. Specifically, we model workload in a one dimensional domain as a continuous second-order stationary process. This means that we associate a random workload with every point in the domain, assume that the mean workload at every point is the same, assume that the workload variance at every point is the same, and assume that the covariance between the workloads at any two points is uniquely determined by their distance. The model takes the domain to be divided into some $n = 2^d$ clusters of equal size, mapped modularly onto $P = 2^p$ processors. Throughout this paper we take P to be fixed, and $d \ge p$. The *degree* of the decomposition is defined to be d. Given one scatter decomposition, another of higher degree can be constructed by splitting each cluster into two, then by modularly mapping the resulting set of clusters.

We derive three main results, each of which has a different set of assumptions concerning the correlation function.

- 1. Assumption: The correlation function is convex. Result: Increasing the degree of a scatter decomposition does not increase the common processor workload variance.
- 2. Assumptions: The workload process is stationary and Gaussian. The correlation function decreases linearly until reaching zero, then remains zero (an elbow function). Result: There exists a degree d_0 , such that if $d_0 \leq d_1 < d_2$, then the expected maximum processor workload under a scatter decomposition of degree d_2 is no larger than the expected maximum processor workload under a scatter decomposition of degree d_1 .
- Assumption: The correlation function decreases linearly across the entire domain. Result: For any number of clusters 2ⁱ, among all mappings that assign 2^{i-p} clusters per processor the modular mapping minimizes the average processor workload variance.

Performance ultimately is measured in terms of finishing time, so that the expected load of the most heavily loaded processor is an appropriate metric. One of our results addresses this metric directly. Average processor workload variance is a secondary measure, although intuition does suggest that decreasing the variance while keeping the mean constant will decrease the expected maximum. Consequently, all these results confirm our intuition that modularly mapping increasingly finer grained workload leads to better load balance. It should be noted that increased communication overhead is the price paid for this balance, and is a cost we do not include in this model. One should not interpret these results as saying that better overall performance can always be achieved by increasing the degree. For a given domain, there will be an optimal degree that balances the conflicting goals of low communication costs and good load balance.

A brief analysis of scatter decomposition can be found in [15]. However, that analysis assumes statistical independence between all cluster workloads, and seems to consider the effects of scatter decomposition on a given architecture as the problem size is increased. As such it is an inappropriate model for studying the effects of changing the mapping of a single given problem. Treatments of other problems have used stochastic models of workload to estimate the expected finishing time; but invariably those models concern statistically independent workloads, e.g. the analyses in [3] and [6]. These results are inadequate for analyzing scatter decomposition. When all workload is independent, then aggregated workload is independent, and there is no performance benefit to be gained by scattering. Scatter decomposition is successful precisely because the workload is not independent. Our contribution is to propose and analyze a model that includes workload correlation, and explain why increasingly finer partitions mapped modularly tend to balance the load better.

2 Analysis

In this section we study a probabilistic model of workload, and the performance of different mappings. For the sake of simplicity we constrain our model to be one-dimensional. This assumption does not negate the utility of the model; any multi-dimensional problem partitioned into hyper-strips can be viewed as a one-dimensional problem. Such partitions greatly simplify the programming needed to exchange information between processors. In fact, our experience in mapping a land-battle simulation using scatter decomposition was that strip partitions minimized the execution time [10]. This was also our experience in mapping a regular scientific code onto the Intel iPSC/1 [16].

Our analysis concerns the effect of scatter decomposition on load balance, in the absence of communication or synchronization costs. By understanding how load balance in isolation is affected by the decomposition/mapping decisions, we are better able to understand the tension between load imbalance and communication/synchronization overheads. The model we use is intended to be descriptive, rather than predictive; the analysis is qualitative rather than quantitative. We doubt that the end benefits of fitting a model to performance data will justify the costs of doing so. Nevertheless we feel there is worth in formally affirming the intuition behind scatter decomposition.

2.1 When and Why Scatter Decomposition Works

Our model explains the success of scatter decomposition by showing that it induces correlation between processors' workloads. To see the performance benefits of correlated workloads, imagine that a random workload is generated and partitioned so that the same amount of work is assigned to every processor. A processor's workload is random, but all processors always finish at the same time, because their workloads are perfectly correlated. This situation is optimal, because all processors are busy all the time. Now imagine that the workload at every point is statistically independent of any other. No matter what the domain decomposition or mapping, processor workloads are statistically independent. In fact, the expected maximum processor workload is the same regardless of granularity, so long as the same volume of domain is assigned to each processor. The "ideal" of random but highly correlated processor workloads cannot be achieved in this artificial scenario.

Scatter decomposition works because irregular workloads are *not* statistically independent: high workload tends to appear in contiguous regions. A sufficiently fine-grained decomposition will split the region up, modular assignment will spread its workload around. The contribution of that region to one processor's workload is highly correlated with the contribution of a nearby region to a different processor's workload. If the underlying workload is highly correlated in nearby regions, then scatter decomposition induces correlation between processors' workloads. We have observed this phenomenon in our own experiments with a one-

dimensional fluid flow computation using adaptive gridding [11]. The fluids problem exhibits irregular grids similar to those in Figure 1.

For a given problem, the sample autocorrelation function [12](p. 437) is a statistical estimate of correlation between point workloads, as a function of the distance between them. Autocorrelations range between 1 and -1; the larger the autocorrelation, the more similar the workloads of two points at a given distance tend to be. Zero correlation implies statistical independence; increasingly negative correlations imply increasing dissimilarity between workloads. Figure 2 shows the sample autocorrelation function at one time-step in a fluid flow computation. Not only does correlation diminish as a function of distance, it can reasonably be modeled as a convex "elbow" function $d_{\alpha}(t) = \sigma^2 \max\{0, 1 - \alpha t\}$ over an appropriate range of t, and some $\alpha \ge 0$. This corresponds nicely with two of our results, one of which assumes elbow correlation, the other of which assumes a convex correlation function.

There are situations where scatter decomposition will not work well. Consider a one dimensional domain discretized into 1000 points, numbered between 0 and 999, to be mapped onto ten processors. Randomly choose some "base" number $b \in [0, 99]$, and imagine that every hundredth point beginning with b has a computational cost of 1000, while all other points have a computational cost of 1. If one evenly divides the domain into ten subregions and maps them modularly, every processor has 1099 units of computational cost of 2098, while the other half each have a cost of 100. Modularly assign each point individually, and processor (b mod 10) has a cost of 10090, while every other processor has a cost of 1000. In this situation mapping increasingly finer-grained workload leads to decreasing performance. Due to b's randomness this workload model is stochastic, and is second-order stationary. Two points at a distance 100m for $m = 1, \ldots, 9$ will always have the same workload. The correlation function at all distances 100m consequently has value one. It has some fixed smaller value for all other distances. The principle reason this problem defeats finegrained scatter decomposition is the periodicity. One should be extremely careful using scatter decomposition in the presence of strong periodic behavior, if there is any chance that the periodicity of the modular mapping can align with the periodicity of workload. The assumptions of the models we study do not admit periodicity.

2.2 Model Preliminaries

We consider the behavior of a computation over a real line interval, divided into n clusters, and mapped onto P processors. Both n and P are taken to be powers of two, and $n \ge P$. We are interested in the average processor workload variance, and in the expected workload of the processor that takes the longest time to complete. Without loss of generality we take the real interval to be [0, 1]. Assume that every point $p \in [0, 1]$ has a certain work intensity W(t). The time required to process [a, b] is the integral of W(t) from t = a to t = b. We assume that the intensities W(t) are unknown, but we are willing to model our uncertainity by

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assuming that W(t) is a random variable, and that W(t) can be viewed as a second-order stationary process [13] over $t \in [0, 1]$. Thus we suppose that $E[W(t)] = \mu$ for all $t \in [0, 1]$, that $Var[W(t)] = \sigma^2$ for all $t \in [0, 1]$, and that Cov[W(t), W(s)] depends only on |t - s|. To emphasize this point we will denote the covariance function as Cov(|t - s|). These assumptions are reasonable if we are unwilling or unable to differentiate between the likely behavior of the computation at t and at s. We do not assume that W(t) = W(s), we simply assume that we have the same degree of uncertainity about W(t) and W(s).

The execution time for [a, b] is

$$T(a,b) = \int_a^b W(t) \ dt.$$

T(a, b) has mean value $(b - a)\mu$. The variance of T(a, b) is

$$Var[T(a,b)] = E[T(a,b)^{2}] - (b-a)^{2}\mu^{2}$$

= $E\left[\left(\int_{a}^{b} W(t) dt\right)\left(\int_{a}^{b} W(s) ds\right)\right] - (b-a)^{2}\mu^{2}$
= $\int_{a}^{b} \int_{a}^{b} E[W(t)W(s)] dt ds - (b-a)^{2}\mu^{2}$
= $\int_{a}^{b} \int_{a}^{b} Cov(|s-t|) dt ds - (b-a)^{2}\mu^{2}.$ (1)

Following a decomposition into n clusters, the *ith* cluster's workload is T(i/n, (i+1)/n), and is denoted as $c_i(n)$. The random vector of cluster workloads is denoted $C(n) = \langle c_0(n), \ldots, c_{n-1}(n) \rangle$.

We are interested in the covariance matrix σ_C^2 for the cluster workloads. For $i \neq j$ we have

$$Cov[c_i(n), c_j(n)] = (\sigma_C^2)_{ij} = \int_{i/n}^{(i+1)/n} \int_{j/n}^{(j+1)/n} Cov(t-s) dt ds.$$
(2)

 $Var[c_i(n)]$ is simply Var[T(i/n, (i+1)/n)], given above. The sequence $c_0(n), c_1(n), \ldots, c_{n-1}(n)$ is second-order stationary, a fact easily deduced from equations (1) and (2). To emphasize this we define the function ϕ :

$$\phi(|j-i|,n) = Cov[c_i(n),c_j(n)].$$

Note that $\phi(0, n)$ is a cluster's variance.

An assignment of clusters to processors is described by a $P \times n$ assignment matrix whose *ij*-th entry is 1 if $c_j(n)$ is assigned to processor *i*, and is 0 otherwise. Given assignment matrix \mathcal{A} , the multiplication $\mathcal{A}C$ yields a $P \times 1$ random vector whose *j*th component is the sum of the execution times of all clusters assigned to processor *j*. The vector of mean processor loads is the matrix-vector product $\mathcal{A}\bar{\mu}_n$, where $\bar{\mu}_n$ is the *n* element vector with μ/n in each coordinate. The covariance matrix of $\mathcal{A}C$ is the product $\mathcal{A}\sigma_C^2 \mathcal{A}^T$, where \mathcal{A}^T is the transpose of \mathcal{A} . The overall execution time is the maximum processor execution time, or $\max\{(\mathcal{A}C)^T\}$. This quantity is random. For any processor P_i , let $\mathcal{A}(i)$ denote the set of clusters assigned to it under \mathcal{A} , and let $L_i(\mathcal{A}, n)$ be P_i 's random workload. By definition the variance of $\overline{L_i}(\mathcal{A}, n)$ is given by

$$Var[L_{i}(\mathcal{A}, n)] = \left(\mathcal{A}\sigma_{C}^{2}\mathcal{A}^{T}\right)_{ii}$$

=
$$\sum_{c_{j}(n)\in\mathcal{A}(i)} \phi(0, n) + \sum_{\langle c_{j}(n), c_{k}(n)\rangle\in\mathcal{A}(i)\times\mathcal{A}(i)} \phi(|j-k|, n).$$
(3)

The first component of this expression is the sum of variances of all clusters assigned to P_i . The second component is a sum of *cluster covariance terms* (we will call these *cc* terms), that depends on the assignment. Similarly, the covariance between processors $L_i(\mathcal{A}, n)$ and $L_j(\mathcal{A}, n)$ is given by a sum of *cc* terms:

$$Cov[L_i(\mathcal{A}, n), L_j(\mathcal{A}, n)] = \sum_{\langle c_k(n), c_m(n) \rangle \in \mathcal{A}(i) \times \mathcal{A}(j)} \phi(|k - m|, n)$$
(4)

The sum of all cluster covariance matrix terms always equals the sum of all processor workload variances and covariances¹

$$\sum_{i=0}^{n-1} \sum_{j=0}^{n-1} (\sigma_C^2)_{ij} = \sum_{i=0}^{P-1} \sum_{j=0}^{P-1} (\mathcal{A} \sigma_C^2 \mathcal{A}^T)_{ij}.$$

This implies a balance between processor workload variances and covariances (and hence correlations); if by changing \mathcal{A} we reduce the average processor workload variance, then we are increasing the average interprocessor workload correlation.

The indices of the sums (3) and (4) have special structure when \mathcal{A} describes a modular mapping. We know that if $c_j(n)$ and $c_k(n)$ are assigned to the same processor, then |j-k| is a multiple of P. Under a modular mapping each processor will have n/P clusters. Among these there are n/P - 1 pairs of clusters whose indices are exactly P apart, n/P - 2 pairs whose indices are exactly 2P apart, and so on. Since n and P determine the specifics of the mapping we may drop the notational dependence of $L_i(\mathcal{A}, n)$ on \mathcal{A} . Under the modular mapping we may write the common processor workload variance as

$$Var[L(n)] = (n/P)\phi(0,n) + 2\sum_{k=1}^{(n/P)-1} ((n/P) - k)\phi(kP,n).$$
(5)

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To consider processor workload covariance under a modular assignment take i < j, and consider a cluster $c_a(n)$ assigned to processor P_i . It has cc terms with all processor P_j clusters $c_m(n)$ such that $|a - m| \mod P = j - i$ or $|a - m| \mod P = P - j + i$. There are ((n/P) - k) cc terms arising from clusters whose indices are kP + j - i apart (for k = 0, ..., (n/P) - 1); there are ((n/P) - k) cc terms arising from clusters whose indices are kP - j + i apart (for k = 1, ..., (n/P) - 1). We may therefore write

$$Cov[L_i(n), L_j(n)] = \sum_{k=0}^{(n/P)-1} ((n/P) - k)\phi(kP + j - i, n) + \sum_{k=1}^{(n/P)-1} ((n/P) - k)\phi(kP - j + i, n)$$

¹ this conservation law proved to be invaluable when debugging detailed expressions for the processor workload variance and covariances, e.g. (12) and (13).

$$= (n/P)\phi(j-i,n) + \sum_{k=1}^{(n/P)-1} ((n/P) - k)\phi(kP + j - i,n) + \sum_{k=1}^{(n/P)-1} ((n/P) - k)\phi(kP - j + i,n).$$
(6)

2.3 Decreasing Workload Variance

Under very general assumptions one can show that increasing the degree of a scatter decomposition reduces the common processor workload variance. The necessary assumptions are that the workload process be second-order stationary, and that its covariance function be convex.

The first step is to show that $\phi(|j-i|, n)$ is a convex function of |j-i| over the range 1, 2, ..., n-1. Towards this end assume that $x \ge 1/n$ and define

$$I(n,x) = E\left[\int_{0}^{1/n} \int_{x}^{x+1/n} W(s)W(t) dt ds\right]$$

= $\int_{0}^{1/n} \int_{x}^{x+1/n} Cov(t-s) dt ds$
= $\int_{0}^{1/n} \left[\int_{x}^{\infty} Cov(t-s) dt - \int_{x+1/n}^{\infty} Cov(t-s) dt\right] ds$

Taking the derivative with respect to x we find that

$$\frac{\partial}{\partial x}I(n,x) = \int_0^{1/n} \left(Cov(x+1/n-s) - Cov(x-s) \right) \, ds.$$

The difference being integrated increases in x due to Cov(t) convexity, implying that the derivative of I(n, x)with respect to x increases in x—one characterization of a convex function. By stationarity $Cov[c_i(n), c_j(n)] = Cov[c_0(n), c_{|j-i|}(n)]$; furthermore $Cov[c_0(n), c_{|j-i|}(n)] = I(n, |j-i|/n)$. Consequently $Cov[c_i(n), c_j(n)]$ is a convex function of |j-i| once $|j-i| \ge 1$ (it may indeed be convex over the entire range, but that fact has not been shown, and is not needed).

We are interested in the effects of moving from a scatter decomposition with degree d - 1 to one with degree d. To analyze these effects we make the following observation. Consider a domain partitioned into $n = 2^d$ clusters, which is mapped by modularly assigning pairs of clusters: $c_0(n)$ and $c_1(n)$ are assigned to processor 0, $c_2(n)$ and $c_3(n)$ are assigned to processor 1, and so on. This mapping is identical to the scatter decomposition of degree d - 1; the pair of clusters $c_0(n), c_1(n)$ viewed from the d degree mapping is the same as the single cluster $c_0(n/2)$ viewed from the d-1 degree mapping. We will show that the modular mapping with degree d - 1 produces processor variances that are no smaller than those of the modular mapping with degree d.

Split each cluster $c_i(n/2)$ into two equal sized clusters. The sum of the two split cluster variances plus

twice their covariance must equal the variance of $c_i(n/2)$. That is,

$$\phi(0, n/2) = 2\phi(0, n) + 2\phi(1, n). \tag{7}$$

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Similarly, take two clusters $c_i(n/2)$ and $c_j(n/2)$, and split each into two equal sized clusters. The total covariance between the four split clusters must equal the covariance between the two unsplit clusters. Thus

$$\phi(|j-i|, n/2) = 2\phi(2|j-i|, n) + \phi(2|j-i|+1, n) + \phi(2|j-i|-1, n).$$
(8)

Note that the index values must double when taken with respect to n rather than n/2 clusters.

Substituting the right-hand-sides of equations (7) and (8) into equation (5) and working through the algebra, we find that

$$Var[L(n/2)] = (n/P)\phi(0,n) + (n/P)\phi(1,n) + 2\sum_{k=1}^{(n/(2P))-1} ((n/P) - 2k)\phi(2kP,n) + \sum_{k=1}^{(n/(2P))-1} ((n/P) - 2k) [\phi(2kP + 1, n) + \phi(2kP - 1, n)].$$

Using this expression and (5), we compute the difference Var[L(n/2)] - Var[L(n)]. All terms involving $\phi(2kP,n)$ cancel, for k = 0, ..., n/(2P) - 1. Each remaining term from Var[L(n)] has the form $2((n/P) - 2k - 1)\phi((2k + 1)P, n)$, for k = 0, ..., n/(2P) - 1. We split each such term into the sum $(n/P - 2k)\phi((2k + 1)P, n) + (n/P - 2k - 2)\phi((2k + 1)P, n)$, and pair these with Var[L(n/2)] terms as follows:

$$Var[L(n/2)] - Var[L(n)] = \sum_{k=0}^{(n/(2P))-1} ((n/P - 2k)(\phi(2kP + 1, n) - \phi((2k + 1)P, n)) - (n/P - 2k - 2)(\phi((2k + 1)P, n) - \phi((2k + 2)P - 1, n))).$$
(9)

One characteristic of a convex function g is that for fixed y the difference g(x) - g(x + y) is a decreasing function of x. Every two terms we have paired differ in their index arguments by exactly P - 1, e.g., $\phi(2kP + 1, n)$ and $\phi((2k + 1)P, n)$. Since ϕ is a convex function of the index argument once the index is at least 1, we have for every k

$$\phi(2kP+1,n) - \phi((2k+1)P,n) \ge \phi((2k+1)P,n) - \phi((2k+2)P-1,n).$$

The left-hand-side expression in this inequality is weighted more heavily in equation (9) than is the right-hand-side expression. It follows that $Var[L(n/2)] - Var[L(n)] \ge 0$, proving our first result.

Theorem 1 Suppose the workload process W(t) is second-order stationary with a convex covariance function. Then increasing the degree of a scatter decomposition does not increase the processor workload variance.

2.4 Decreasing Expected Maximum Workload

Next we demonstrate circumstances where increasing the degree of a scatter decomposition reduces the expected workload of the most heavily loaded processor. The argument is to show that under appropriate assumptions the correlation between any two processors' workloads increases as the degree increases. We then site a result from the literature proving that the expected maximum decreases in this situation.

We assume that the workload process $\{W(t)\}$ is a stationary Gaussian process² [7]. Additivity properties of the Gaussian then ensure that the vector of *n* clusters has a jointly normal distribution [7](Chapter 6) and that under any assignment, the processors' workloads are jointly normal. We also assume that the correlation function is $Cov(t) = \sigma^2 \max\{0, 1 - \alpha t\}$, where $\alpha = 2^v/m_\alpha \ge 1$ for some integers $v, m_\alpha \ge 0$. The restriction on α is used to simplify certain calculations. $\delta = 1/\alpha$ is the smallest distance *t* at which Cov(t) = 0. Our results apply when the degree is large enough so that subinterval $[0, \delta]$ is partitioned into at least $P = 2^p$ clusters. If the degree is *d*, then the number of clusters in $[0, \delta]$ is $\delta 2^d$. Now let d_0 be the least *d* such that $\delta 2^d \mod 2^p = 0$. Equivalently, d_0 is the least integer *d* such that $m_\alpha 2^{d-p-\nu}$ is an integer. Clearly $d_0 \le p+v$. Our results apply when the degree is at least d_0 .

We can compute functional forms for $\phi(|j-i|, n)$ given this explicit definition of Cov(t). Performing the integration given by (2) one determines that

$$\phi(|j-i|,n) = \begin{cases} \frac{\sigma^2}{n^3}(n-\alpha|j-i|) & \text{if } |j-i| < \delta n\\ \frac{\sigma^2\alpha}{\delta n^3} & \text{if } |j-i| = \delta n\\ 0 & \text{if } |j-i| > \delta n \end{cases}$$
(10)

These calculations take advantage of the fact that δ is a multiple of 1/n. The variance of a cluster is determined by evaluating (1), yielding

$$\phi(0,n) = \frac{\sigma^2}{n^3} (n - \alpha/3).$$
(11)

Given equations (10) and (11) we can compute processor workload variance and covariance under scatter decomposition. General expressions for these quantities are given by (5) and (6). For large values of k, some terms in those sums vanish, being zero. Our assumption that the scatter decomposition has degree d_0 or larger ensures that terms which vanish are easily characterized,³ and that those clusters whose indices are exactly δn apart are assigned to the same processor. All $\phi(kP, n)$ terms in (5) vanish for $k > \delta n/P$; we have $\phi(kP, n) = \sigma^2 \alpha/(6n^3)$ for $k = \delta n/P$. We may rewrite the variance as

$$Var[L(n)] = (n/P)\frac{\sigma^{2}(n-\alpha/3)}{n^{3}} + \sum_{k=1}^{(\delta n/P)-1} \left((n/P-k)\frac{\sigma^{2}(n-\alpha kP)}{n^{3}} \right) + (n/P-\delta n/P) \left(\frac{\sigma^{2}\alpha}{6n^{3}} \right)$$

 $^{^{2}}$ note that this assumption is stronger than we have used so far, due both to stationarity rather than second-order stationarity, and due to the assumption of a specific workload distribution

³This is not the case for smaller degrees. A large number of special cases must be constructed and analyzed. This task seemed to us to be more tedious than is warranted by the anticipated correspondingly stronger result.

$$= \sigma^{2} \left(\frac{(\delta - \delta^{2}/3)}{P^{2}} + \frac{1 - \alpha/P}{3n^{2}} - \frac{1 - \alpha}{3n^{2}P} \right).$$
(12)

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Calculation of this equality is much simplified with the use of a symbolic mathematics package.

The processor workload covariance is similarly handled. Assume that i < j. $k = \delta n/P$ again delineates where ϕ terms vanish: $\phi(kP + j - i, n) = 0$ for all $k \ge \delta n/P$, and $\phi(kP - i + j, 0) = 0$ for all $k > \delta n/P$. We may rewrite (6) as

$$Cov[L_{i}(n), L_{j}(n)] = (n/P)\frac{\sigma^{2}}{n^{3}}(n - \alpha(j - i)) + \sum_{k=1}^{(\delta n/P)-1} ((n/P) - k)\frac{\sigma^{2}(n - \alpha(kP + j - i))}{n^{3}} + \sum_{k=1}^{\delta n/P} ((n/P) - k)\frac{\sigma^{2}(n - \alpha(kP - j + i))}{n^{3}} = \sigma^{2}\left(\frac{\delta - \delta^{2}/3}{P^{2}} + \frac{1}{3n^{2}} - \frac{j - i}{Pn^{2}}\right).$$
(13)

The correlation between $L_i(n)$ and $L_j(n)$ is the ratio $Cov[L_i(n), L_j(n)]/Var[L(n)]$. For all $d \ge d_0$ we obtain the correlation using (13) and (12), and can treat the ratio as a continuous function of n. It is interesting to note that as n increases the correlation approaches unity. This supports our intuition that partitioning the domain into increasingly finer clusters and mapping them modularly induces correlation between processor workloads. In fact, the tendency towards unity is monotonic. Taking the derivative with respect to n we find that the derivative is positive if

$$(4/3 - 2\delta/3)(j - i) + 2\delta/9 - 2/3 > 0.$$

This inequality holds, since $(4/3 - 2\delta/3) \ge 2/3$. Consequently, for all $n = 2^d \ge 2^{d_0}$ we must have

$$Cov[L_i(2n), L_j(2n)]/Var[L(2n)] > Cov[L_i(n), L_j(n)]/Var[L(n)].$$

Next we use this relationship to analyze the expected maximum processor workload.

The following result is based on the Normal Comparison Lemma [8](p.81) and is the key to our observations concerning the expected maximum processor workload.

Theorem 2 (Leadbetter et al.) Let ξ_0, \ldots, ξ_k be standardized jointly normal random variables, and let η_0, \ldots, η_k be standardized jointly normal random variables, such that $Cov(\xi_i, \xi_j) \leq Cov(\eta_i, \eta_j)$ for each $i, j, i \neq j$. Then for every u,

$$Pr\{\max\{\xi_0,\ldots,\xi_k\} \le u\}\} \le Pr\{\max\{\eta_0,\ldots,\eta_k\} \le u\}\},\$$

and hence

$$E[\max\{\xi_0,\ldots,\xi_k\}] \ge E[\max\{\eta_0,\ldots,\eta_k\}].$$

The standardization of a random variable X is the scaled random variable Z = (X - m)/s, where m and s are X's mean and standard deviation, respectively. The mean of a standardized random variable is zero and its variance is one; the covariance between two standardized random variables is the correlation between their corresponding unstandardized forms. Let $Z_i(n)$ be the standardized workload of processor P_i given a domain of n clusters. $Cov[Z_i(n), Z_j(n)] = Cov[L_i(n), L_j(n)]/Var[L(n)]$, which we have shown to be increasing in n. If $\hat{n} > n$ (equivalently, if one scatter decomposition has higher degree than another), then

$$E[\max\{Z_0(n),\ldots,Z_{P-1}(n)\}] \ge E[\max\{Z_0(\hat{n}),\ldots,Z_{P-1}(\hat{n})\}].$$
(14)

The expected maximum workload is

$$E[\max\{L_0(n), \dots, L_{P-1}(n)\}] = E[\max_{0 \le i \le P-1} \{L_i(n) + Var[L(n)]^{1/2} Z_i(n)\}]$$

= $\mu/P + Var[L(n)]^{1/2} E[\max_{0 \le i \le P-1} \{Z_i(n)\}].$

Theorem 1 shows that $Var[L(n)] \ge Var[L(2n)]$; this along with inequality (14) proves our second result.

Theorem 3 Let $\{W(t)\}$ be a stationary Gaussian process, with a covariance function $Cov(t) = \sigma^2 \max\{0, 1 - \alpha t\}$, where $\alpha = 2^{\nu}/m_{\alpha} \ge 1$ for some positive integers m_{α} , ν . Let there be 2^p processors, and let d_0 be the least integer d such that $m_{\alpha}2^{d-p-\nu}$ is an integer. If $d_2 > d_1 \ge d_0$, then the expected maximum processor workload of a scatter decomposition with degree d_2 is no greater than that of a scatter decomposition with degree d_1 .

2.5 Minimization of Average Workload Variance

Our final result gives conditions where for a given n, among all "balanced" assignments—those placing n/P clusters per processor—the modular mapping minimizes the average processor workload variance. To prove this result we assume that the covariance function decreases linearly across the entire domain: $Cov(s) = \sigma^2(1 - \alpha s)$, for some α satisfying $0 \le \alpha \le 2$. The result is based on a procedure that takes any assignment and constructs another whose sum of processor workload variances is no larger. The repeated application of this procedure produces a modular assignment. Consequently, modular assignments minimize the average processor workload variance.

The arguments to follow specify individual covariance terms. These arguments are clearer using the $Cov[c_i(n), c_j(n)]$ notation rather than $\phi(|j-i|, n)$. It is straightforward to determine the form of $Cov[c_i(n), c_j(n)]$ under the present assumptions:

$$Cov[c_i(n), c_j(n)] = \begin{cases} \frac{\sigma^2}{n^3}(n - \alpha|j - i|) & \text{if } |j - i| > 0\\ \frac{\sigma^2}{n^3}(n - \alpha/3) & \text{if } |j - i| = 0 \end{cases}$$
(15)

Let \mathcal{A}_1 be any assignment matrix describing a balanced assignment. Without loss of generality, we assume that under \mathcal{A}_1 the processors are numbered so that P_0 is assigned $c_0(n)$, P_1 is assigned the smallest indexed $c_i(n)$ that is not assigned to P_0 , and in general P_j is assigned the smallest indexed cluster that is not assigned to any of $P_0, P_1, \ldots, P_{j-1}$.

We will say that $c_j(n)$ is in place if it is assigned to processor $P_{j \mod P}$. Note that all clusters are in place under a modular assignment. We construct another balanced assignment \mathcal{A}_2 by finding the smallest indexed $c_i(n)$ that is not in place, and by putting it in place. Let c_f denote this cluster, let P_S denote the source processor that has c_f under \mathcal{A}_1 , and let P_T denote the target processor $P_{f \mod P}$. Let c_g be the smallest indexed cluster assigned to P_T such that g > f. \mathcal{A}_2 is constructed from \mathcal{A}_1 by giving c_f to P_T , and c_g to P_S . Figure 3 illustrates these definitions. We will prove that the sum of processor variances under \mathcal{A}_2 bounds that sum under \mathcal{A}_1 from below; consequently the average workload variance under \mathcal{A}_2 is no greater than that under \mathcal{A}_1 .

Recall that under any assignment matrix \mathcal{A} the variance of P_i 's work load is given by

$$Var[L_{i}(\mathcal{A}, n)] = \left(\mathcal{A}\sigma_{C}^{2}\mathcal{A}^{T}\right)_{ii}$$

$$= \sum_{c_{j}(n)\in\mathcal{A}(i)} Var[c_{j}(n)] + \sum_{\langle c_{j}(n), c_{k}(n)\rangle\in\mathcal{A}(i)\times\mathcal{A}(i)} Cov[c_{j}(n), c_{k}(n)], \quad (16)$$

and that

$$Cov[L_i(\mathcal{A}, n), L_j(\mathcal{A}, n)] = \sum_{\langle c_k, c_m \rangle \in \mathcal{A}(i) \times \mathcal{A}(j)} Cov[c_k(n), c_m(n)]$$

It is clear from (16) that the variance of any processor other than P_S or P_T is by unaffected by swapping c_f and c_g . To prove the desired result we need only show that the swap does not increase the sum of P_T and P_S variances. The change in processor variances caused by the swap is entirely due to changes in the sum of cluster covariance (cc) terms in each processor. After swapping c_f and c_g , each cluster $c_i(n)$ assigned to P_S loses the cc term $Cov[c_f(n), c_i(n)]$ and gains the term $Cov[c_g(n), c_i(n)]$. We let Δ_{L_S} denote the sum of all such changes among clusters in P_S to the left of c_f , and let L_S denote the number of such clusters. Similarly Δ_{R_S} denotes the sum of changes among clusters in P_S to the right of c_g and R_S denotes the number of such clusters; Δ_{M_S} denotes the sum of changes among clusters in P_S with indices between f and g. Expressions for these quantities are derived using equation (15):

$$\Delta_{L_{S}} = \sum_{\substack{c_{i}(n) \in \mathcal{A}_{2}(S) \\ i < f}} (Cov[c_{g}(n), c_{i}(n)] - Cov[c_{f}(n), c_{i}(n)]) = -\frac{\sigma^{2}}{n^{3}}(g - f)L_{S}\alpha;$$

$$\Delta_{R_{S}} = \sum_{\substack{c_{j}(n) \in \mathcal{A}_{2}(S) \\ j > g}} (Cov[c_{g}(n), c_{j}(n)] - Cov[c_{f}(n), c_{j}(n)]) = \frac{\sigma^{2}}{n^{3}}(g - f)R_{S}\alpha;$$

$$\Delta_{M_{S}} = \sum_{\substack{c_{j}(n) \in \mathcal{A}_{2}(S) \\ f < k < g}} (Cov[c_{g}(n), c_{k}(n)] - Cov[c_{f}(n), c_{k}(n)]) = \frac{\sigma^{2}}{n^{3}} \sum_{\substack{c_{k}(n) \in \mathcal{A}_{2}(S) \\ f < k < g}} (2k - f - g)\alpha.$$

The change in P_S 's variance after the swap is the sum $\Delta_{L_S} + \Delta_{M_S} + \Delta_{R_S}$.

We can similarly describe the change in P_T 's variance with the definitions

$$\Delta_{L_T} = \sum_{\substack{c_i(n) \in \mathcal{A}_2(T) \\ i < f}} (Cov[c_f(n), c_i(n)] - Cov[c_g(n), c_i(n)]) = \frac{\sigma^2}{n^3} (g - f) L_T \alpha;$$

$$\Delta_{R_T} = \sum_{\substack{c_j \in \mathcal{A}_2(T) \\ i > g}} (Cov[c_f(n), c_j(n)] - Cov[c_g(n), c_j(n)]) = -\frac{\sigma^2}{n^3} (g - f) R_T \alpha.$$

No term analogous to Δ_{M_S} is necessary since there are no clusters in P_T with indices between f and g.

The change in the sum of P_S 's variance with P_T 's variance is given by the sum of all the Δ terms defined above. We will show that the sum of Δ terms is bounded from above by 0. At this point a number of observations are useful. Since all $c_i(n)$ with i < f are in order, it follows that $L_T \leq L_S$. Thus $\Delta_{L_S} + \Delta_{L_T} \leq 0$. It remains to show that $\Delta_{R_S} + \Delta_{R_T} + \Delta_{M_S} \leq 0$. We know that

$$\Delta_{R_S} + \Delta_{R_T} = -\frac{\sigma^2}{n^3} (R_T - R_S)(g - f)\alpha; \qquad (17)$$

furthermore, since $n/P = L_T + R_T + 1$, we must also have $R_S \leq R_T$. We proceed to show that the magnitude of Δ_{M_S} is no greater than the magnitude of (17) and consequently prove the larger result.

 $m = n/P - L_S - R_S - 1$ is the number of clusters in P_S whose indices lie strictly between f and g. Δ_{M_S} is maximized when the indices of these clusters are as large as possible; when $k = g - 1, g - 2, \dots, g - m$. With such indices, the sum of c_g 's cc terms in P_S is

$$\frac{\sigma^2}{n^3}\sum_{i=1}^m (n-i\cdot\alpha)$$

Likewise, the sum of c_f 's cc terms in P_S is

$$\frac{\sigma^2}{n^3}\sum_{i=1}^m (n-(g-f-i)\alpha).$$

From this, we see that Δ_{M_S} when maximized can be written as

$$\Delta_{M_S} = \frac{\sigma^2}{n^3} \sum_{i=1}^m (n-i\cdot\alpha) - \frac{\sigma^2}{n^3} \sum_{i=1}^m (n-(g-f-i)\alpha) = \frac{\sigma^2}{n^3} m(g-f)\alpha.$$

But note that

$$m = n/P - L_S - R_S - 1$$

$$\leq n/P - L_T - R_S - 1$$

$$= (n/P - L_T - R_T - 1) + (R_T - R_S)$$

$$= (R_T - R_S),$$

so that

$$\Delta_{R_S} + \Delta_{R_T} + \Delta_{M_S} = \frac{\sigma^2}{n^3} (-(R_T - R_S)(g - f)\alpha + m \cdot (g - f)\alpha) \le 0$$

Consequently, swapping c_f and c_g does not increase the sum of P_S and P_T 's variance. Furthermore, the swap does not affect the sum of other processors' variances. Repeatedly applying this procedure puts every cluster in place, which is the modular assignment. This discussion has proved the following theorem.

Theorem 4 Let $\{W(t)\}$ be a second-order stationary process, with a covariance function $Cov(s) = \sigma^2(1 - \alpha s)$, where $0 \le \alpha \le 2$. Let P and n be given such that P divides n evenly, and let \mathcal{A}_M be the $P \times n$ assignment matrix describing the modular mapping. Then for any $P \times n$ assignment matrix \mathcal{A} describing a balanced assignment,

$$(1/P)\sum_{i=0}^{P-1}(\mathcal{A}_M\sigma_C^2\mathcal{A}_M^T)_{ii}\leq (1/P)\sum_{i=0}^{P-1}(\mathcal{A}\sigma_C^2\mathcal{A}^T)_{ii}.$$

In the event that the workload process is Gaussian and stationary, we can show that increasing the degree reduces the expected maximum processor workload. We determine the processor variance and covariance under scatter decomposition by substituting the values given by (15) into (5) and (6). Assume that i < j. Working through the algebra one determines that

$$Var[L(n)] = \sigma^2 \left(\frac{1-\alpha/3}{P^2} + \frac{\alpha(1-1/P)}{3n^2} \right),$$

and that

$$Cov[L_i(n), L_j(n)] = \sigma^2 \left(\frac{1 - \alpha/3}{P^2} + \frac{\alpha}{3n^2} - \frac{(j - i)\alpha}{Pn^2} \right).$$

The derivative with respect to n of $C[L_i(n), L_j(n)]/Var[L(n)]$ is positive if

$$(4/3 - 2\alpha/3)(j-i) + 2\alpha/9 - 2/3 > 0.$$

This is always true over the range $\alpha \in [0, 2]$. Consequently the same arguments used to prove Theorem 3 can be applied here.

3 Summary

Scatter decomposition is an attractive method for mapping domain-oriented computations with irregular workloads to parallel architectures. Scatter decomposition partitions the domain into n equal-size pieces, and maps them modularly onto P processors. This paper uses a formal probabilistic model of correlated workload in a one-dimensional domain to explain why and when scatter decomposition works. First, we show that periodicity in workload correlation can lead to load imbalance under scatter decomposition if the

correlation period aligns with the period of the modular mapping. Consequently we consider nonperiodic workload correlation functions.

Our first result shows that if workload correlation is a convex function of distance, then scattering with increasingly finer grained clusters decreases a processor's workload variance, thereby increasing the average inter-processor workload correlation. Since the processor workload mean is unaffected by this change, one anticipates that the expected maximum workload will correspondingly decrease.

Our second result affirms this intuition under a stronger set of assumptions: the workload process is Gaussian, and the correlation function decreases linearly in distance until it reaches zero and then stays at zero. We then show that once a scatter decomposition is sufficiently fine-grained, making the grain-size finer reduces the expected maximum processor workload.

Our third result shows that under slightly different assumptions still, among all possible "balanced" mappings scatter decomposition minimizes the average processor workload variance. This result depends on the correlation function decreasing linearly across the entire domain. In this case it is also true that if the workload process is Gaussian, then scattering a finer-grained decomposition reduces the expected maximum processor workload.

These analytic results serve to formally verify the intuition behind scatter decomposition. However, the results only concern load balance. The additional communication cost of decreasing granularity is not built into this model. Extensions to this work might find the optimal granularity by determining a quantitative estimator of the expected maximum workload and the expected communication cost as a function of granularity. An overall execution time model would be constructed depending on the influence of architecture on the communication costs, and then optimized.

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Processor Assignment

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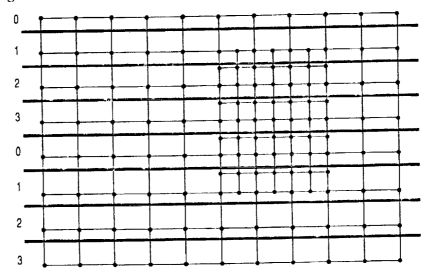


Figure 1: Scatter decomposition of an irregular grid

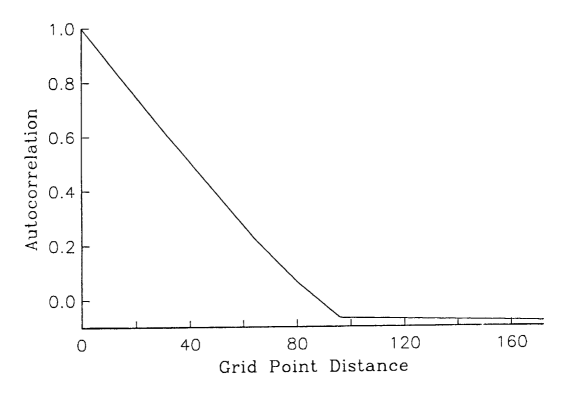
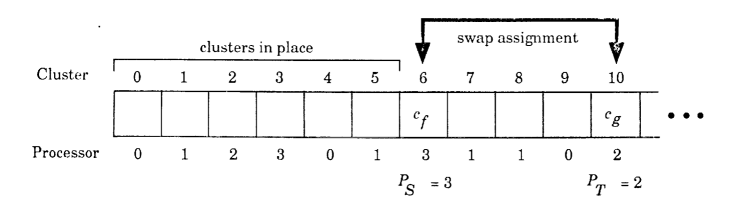


Figure 2: Correlation as function of distance in 1D fluids problem



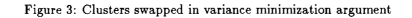
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16. Abstract This paper provides a formal analysis of a powerful mapping technique known as scatter decomposition. Scatter decomposition divides an irregular computational domain into a large number of equal sized pieces, and distributes them modularly among processors. We use a probabilistic model of workload in one dimension to for- mally explain why, and when scatter decomposition works. Our first result is that if correlation in workload is a convex function of distance, then scattering a more finely decomposed domain yields a lower average processor workload variance. Our second result shows that if the workload process is stationary Gaussian and the cor- relation function decreases linearly in distance until becoming zero and then re- mains zero, scattering a more finely decomposed domain yields a lower expected maxi- mum processor workload. Finally we show that if the correlation function decreases linearly across the entire domain, then among all mappings that assign an equal num- ber of domain pieces to each processor, scatter decomposition minimizes the average processor workload variance. The dependence of these results on the assumption of decreasing correlation is illustrated with situations where a coarser granularity actually achieves better load balance. 17. Key Words (Suggested by Author(s))					
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