

ARGONNE NATIONAL LABORATORY
9700 South Cass Avenue
Argonne, Illinois 60439

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SCALABILITY OF OPTIMIZATION ALGORITHMS**

Steven J. Benson, Lois Curfman McInnes, and Jorge J. Moré

Mathematics and Computer Science Division

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GPCG: A Case Study in the Performance and Scalability of Optimization Algorithms*

Steven J. Benson, Lois Curfman McInnes, and Jorge J. Moré

Mathematics and Computer Science Division
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Abstract

GPCG is an algorithm within the Toolkit for Advanced Optimization (TAO) for solving bound constrained, convex quadratic problems. Originally developed by Moré and Toraldo [19], this algorithm was designed for large-scale problems but had been implemented only for a single processor. The TAO implementation is available for a wide range of high-performance architecture, and has been tested on up to 64 processors to solve problems with over 2.5 million variables.

1 Introduction

The Toolkit for Advanced Optimization (TAO) focuses on the design and implementation of component-based optimization software for the solution of large-scale optimization applications. Our approach is motivated by the scattered support for parallel computations and lack of reuse of linear algebra software in currently available optimization software. We exploit numerical abstractions in the optimization software design so that we can leverage external parallel computing infrastructure (for example, communication libraries and visualization packages) and linear algebra tools in the development of optimization algorithms. The algorithms in the toolkit place strong emphasis on the reuse of external tools where appropriate. Our design enables connection to lower-level support (parallel sparse matrix data structures, preconditioners, solvers) provided in toolkits such as PETSc [2, 3], and thus we are able to build on top of these toolkits instead of having to redevelop code. The advantages in terms of development time are significant.

Initial work in the TAO project [4, 5] has centered on the development of a core library of components for various types of optimization problems, including unconstrained and bound-constrained minimization and nonlinear least squares. To explain the TAO design strategy and analyze parallel performance issues, we focus on the gradient projection conjugate gradient (GPCG) algorithm for the solution of the bound-constrained quadratic programming problem

$$\min \{q(x) : l \leq x \leq u\}, \quad (1.1)$$

where $q : \mathbb{R}^n \mapsto \mathbb{R}$ is a strictly convex quadratic function, and the vectors l and u define bounds on the variables. Although GPCG had been originally designed [19] for large-scale

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problems, implementation of GPCG on a parallel architecture presented significant obstacles that are typical of a large class of optimization algorithms. The most significant obstacle arises from the method used to compute the step between iterates. Specifically, in modern active set methods for solving (1.1), the step between iterates is usually defined via the approximate solution of a linear system of the form

$$A_k w_k = -r_k,$$

where the matrix A_k and the vector r_k are, respectively, the reduced Hessian matrix and the reduced gradient of q with respect to the free variables. In a parallel environment, the efficient implementation of the conjugate gradient method requires that A_k be evenly distributed over the processors, but since the set of free variables can change drastically between iterates, the reduced matrix is unlikely to be well distributed. Hence, a redistribution of the rows of A_k over the processors may be necessary at each iteration.

This observation implies that the scalability of the GPCG is limited not only by the efficiency of the redistribution algorithm but by the sizes of the matrices A_k . If the set of free variables is large, then performance is likely to improve because the communication overhead is small, while performance is likely to suffer when there are few free variables. Thus, the GPCG algorithm is prime candidate for a case study in the performance and scalability of optimization algorithms in parallel architectures.

Our implementation of GPCG uses object-oriented techniques to leverage the parallel computing and linear algebra infrastructure offered by PETSc [2, 3], which relies on MPI [14] for all interprocessor communication. As a result, our implementation runs on a wide variety of high-performance architectures. Biros and Ghattas [6, 7] have been using a similar approach for the solution of PDE-constrained optimization problems. They have also been concerned with efficiency and scalability issues, but for quadratic problems with linear equality constraints. As we have pointed out, inequality constrained optimization problems give rise to different performance issues. Hohmann [16], Deng, Gouveia and Scales [11], Meza [18], Bruhwiler et al. [8], and Gockenbach, Petro, and Symes [13] have employed object-oriented design for nonlinear optimization, but their work does not address the reuse of linear algebra toolkits and is restricted to uniprocessor environments. Our use of object-oriented techniques and linear algebra toolkits also distinguishes our implementation of GPCG from the data-parallel implementation of McKenna, Mesirov, and Zenios [17]. In particular, they can rely only on diagonal preconditioners, while our approach allows a wide range of preconditioners.

Sections 2 and 3 are dedicated to background material on the bound-constrained optimization problem (1.1) and to a brief overview of the GPCG algorithm, while Section 4 has a discussion of our design philosophy and its benefits in developing robust and scalable solutions strategies.

The performance results in Section 5 are noteworthy in several ways. First, the number of faces visited by GPCG is remarkably small. Other strategies can lead to a large number of gradient projection iterates, but the GPCG algorithm is remarkably efficient. Another

interesting aspect is that because of the low memory requirements of iterative solvers, we are able to solve problems with over 2.5 million variables with only 8 processors. Strategies that rely on direct solvers are likely to need significantly more storage, and thus more processors. Finally, these results show that the GPCG implementation has excellent efficiency.

Section 6 examines the scalability of the GPCG component functions and the performance of GPCG as the number of variables and the number of active variables at the solution change. These results illustrate the complex performance behavior for constrained optimization problems as well as the observation that performance results that focus only on efficiency can be deceiving if the total computing time is not taken into account.

Section 7 considers the performance of GPCG as the preconditioners change. The ability to use various preconditioners is a result of our design, which allows the connection to external linear algebra toolkits. Our results in this section show that for our benchmark problem, a block Jacobi preconditioner with one block per processor, where each subproblem is solved with a standard, sparse ILU(2) factorization, is faster than a variant with ILU(0). We also show that both block Jacobi variants are faster than a simple point Jacobi method, although the point Jacobi preconditioner exhibits better scalability.

2 Bound-Constrained Quadratic Optimization Problem

A classical result shows that the bound-constrained quadratic optimization problem (1.1) has a unique solution on the feasible region

$$\Omega = \{x \in \mathbb{R}^n : l \leq x \leq u\} \quad (2.1)$$

when the quadratic $q : \mathbb{R}^n \mapsto \mathbb{R}$ is strictly convex, so that

$$q(x) = \frac{1}{2}x^T A x + b^T x + c, \quad (2.2)$$

where $A \in \mathbb{R}^{n \times n}$ is symmetric and positive definite, $b \in \mathbb{R}^n$, and $c \in \mathbb{R}$. This result holds for unbounded Ω , and we thus allow the components of l and u to be infinite. Solutions to problem (1.1) satisfy the Kuhn-Tucker conditions

$$\begin{aligned} \partial_i q(x) &= 0 & \text{if } x_i \in (l_i, u_i) \\ \partial_i q(x) &\geq 0 & \text{if } x_i = l_i \\ \partial_i q(x) &\leq 0 & \text{if } x_i = u_i, \end{aligned}$$

where $\partial_i q(x)$ is the partial derivative of q with respect to the i th variable. Approximate solutions can be defined in terms of the projected gradient, defined by

$$[\nabla_{\Omega} q(x)]_i = \begin{cases} \partial_i q(x) & \text{if } x_i \in (l_i, u_i) \\ \min\{\partial_i q(x), 0\} & \text{if } x_i = l_i \\ \max\{\partial_i q(x), 0\} & \text{if } x_i = u_i \end{cases} \quad (2.3)$$

This definition of a projected gradient is appropriate because x^* is a solution of (1.1) if and only if $\nabla_{\Omega} q(x^*) = 0$.

Given $x_0 \in \Omega$, and a tolerance τ , an approximate solution to the bound constrained problem (1.1) is any vector $x \in \Omega$ such that

$$\|\nabla_{\Omega} q(x)\| \leq \tau. \quad (2.4)$$

Note that (2.4) holds whenever x is sufficiently close to x^* and in the face of Ω that contains x^* . The concept of a face is standard in convex analysis; for the convex set (2.1), the face of Ω that contains x is

$$\left\{ y \in \Omega : y_i = x_i \text{ if } x_i \in \{l_i, u_i\} \right\}.$$

Thus, the face of the feasible set that contains x can be described in terms of the set of active constraints

$$\mathcal{A}(x) = \{i : x_i = l_i \text{ or } x_i = u_i\}.$$

Variables with indices in $\mathcal{A}(x)$ are the active variables, and those with indices outside $\mathcal{A}(x)$ are the free variables. Similarly, the binding variables are those with indices in

$$\mathcal{B}(x) = \{i : x_i = l_i \text{ and } \partial_i q(x) \geq 0, \text{ or } x_i = u_i \text{ and } \partial_i q(x) \leq 0\}.$$

The Kuhn-Tucker conditions show that $\mathcal{B}(x) = \mathcal{A}(x)$ at a solution, so that if all the active variables are not binding, then x is not on the face that contains the solution.

3 The GPCG Algorithm

The GPCG algorithm uses a gradient projection method to identify a face of the feasible region Ω that contains the solution, and the conjugate gradient method to search the face. This section provides an outline of the algorithm and notes any differences between our implementation and the implementation of Moré and Toraldo [19].

Given $y_0 = x_k$, the gradient projection method generates a sequence of vectors $\{y_j\}$ in the feasible region Ω such that

$$y_{j+1} = P[y_j - \alpha_j \nabla q(y_j)], \quad (3.1)$$

where P is the projection onto (2.1), and the step size α_j is chosen such that

$$q(y_{j+1}) \leq q(y_j) + \mu \langle \nabla q(y_j), P[y_j - \alpha_j \nabla q(y_j)] - y_j \rangle \quad (3.2)$$

for some $\mu \in (0, 1/2)$. The projection P can be computed in n operations by

$$P[x] = \text{mid}(l, u, x),$$

where $\text{mid}(l, u, x)$ is the vector whose i th component is the median of the set $\{l_i, u_i, x_i\}$. The step size is computed by a projected search [19] by setting α_j to the first member of the sequence $\alpha_0 (\frac{1}{2})^j$ for $j = 0, 1, \dots$ such that y_{j+1} satisfies the sufficient decrease condition (3.2). In our implementation, we use

$$\alpha_0 = \arg \min \{q(y_k - \alpha \nabla_{\Omega} q(y_k)) : \alpha > 0\}. \quad (3.3)$$

Computation of α_0 is straightforward, since the mapping $\alpha \mapsto q(y_k - \alpha \nabla_{\Omega} q(y_k))$ is a quadratic.

We generate gradient projection iterates until sufficient progress is not made or the active set settles down. Thus, we generate iterates until either

$$\mathcal{A}(y_j) = \mathcal{A}(y_{j-1}) \quad (3.4)$$

or

$$q(y_{j-1}) - q(y_j) \leq \eta_1 \max\{q(y_{l-1}) - q(y_l) : 1 \leq l < j\}. \quad (3.5)$$

If either test is satisfied, we proceed to the conjugate gradient part of the algorithm.

The first test (3.4) measures when the active set settles down. For nondegenerate problems, (3.4) holds in a neighborhood of the solution. The gradient projection could be followed until the optimal face is found, but experience has shown that a large number of iterates may be required. The second test (3.5) measures when the gradient projection method is not making sufficient progress.

Given an iterate x_k and the active set $\mathcal{A}(x_k)$, the conjugate gradient method computes an approximate minimizer to the subproblem

$$\min\{q(x_k + d) : d_i = 0, i \in \mathcal{A}(x_k)\}. \quad (3.6)$$

This problem is unconstrained in the free variables. Note that if x_k lies in the same face as the solution and d_k solves (3.6), then $x_k + d_k$ is the solution of (1.1).

The conjugate gradient algorithm for solving (3.6) is implemented by expressing this subproblem in terms of an equivalent subproblem in the free variables. If i_1, \dots, i_{m_k} are the indices of the free variables, and the matrix Z_k is defined as the matrix in $\mathbb{R}^{n \times m_k}$ whose j th column is the i_j th column of the identity matrix in $\mathbb{R}^{n \times n}$, then subproblem (3.6) is equivalent to the unconstrained subproblem

$$\min\{q_k(w) : w \in \mathbb{R}^{m_k}\}, \quad (3.7)$$

where

$$q_k(w) \equiv q(x_k + Z_k w) - q(x_k) = \frac{1}{2} \langle w, A_k w \rangle + \langle r_k, w \rangle.$$

The matrix A_k and the vector r_k are, respectively, the reduced Hessian matrix of q and reduced gradient of q at x_k with respect to the free variables. If A is the Hessian matrix of the quadratic q , then

$$A_k = Z_k^T A Z_k, \quad r_k = Z_k^T \nabla q(x_k).$$

Also note that A_k is the matrix obtained from A by taking those rows and columns whose indices correspond to free variables; similarly, r_k is obtained from $\nabla q(x_k)$ by taking the components whose indices correspond to free variables.

Given a starting point $w_0 \in \mathbb{R}^{m_k}$, the conjugate gradient algorithm generates a sequence of iterates w_0, w_1, \dots that terminates at a solution of subproblem (3.7) in at most m_k iterations. We use the conjugate gradient algorithm until it generates w_j such that

$$q_k(w_{j-1}) - q_k(w_j) \leq \eta_2 \max\{q_k(w_{l-1}) - q_k(w_l) : 1 \leq l < j\} \quad (3.8)$$

for some tolerance $\eta_2 > 0$. The approximate solution of (3.6) is then $d_k = Z_k w_{j_k}$, where j_k is the first index j that satisfies (3.8).

The termination test (3.8) is not standard. Iterative solvers usually terminate when

$$\|r_j + A_j w_j\| \leq \eta_2 \|r_j\|$$

for some tolerance $\eta_2 \in (0, 1)$. This test suffers from the erratic behavior of the residual $\|r_j + A_j w_j\|$. On the other hand, the termination test (3.8) depends on whether the conjugate gradient method is making sufficient progress.

Given the direction d_k , we use a projected search [19] to define $x_{k+1} = P[x_k + \alpha_k d_k]$, where α_k is the first element in the sequence $(\frac{1}{2})^k$ for $k = 0, 1, \dots$ such that

$$q(x_{k+1}) \leq q(x_k) + \mu \langle \nabla q(x_k), x_{k+1} - x_k \rangle. \quad (3.9)$$

More sophisticated projected searches are possible [19], but this simple search has proved to be sufficient in all cases tried. If

$$\mathcal{B}(x_{k+1}) = \mathcal{A}(x_{k+1}), \quad (3.10)$$

then we find a more accurate solution to subproblem (3.7) by reducing η_2 and continuing with the conjugate gradient method. Otherwise, we terminate this iteration.

Algorithm GPCG

Choose $x_0 \in \Omega$.

For $k = 0, \dots$,

Set $y_0 = x_k$, and generate gradient projection iterates y_1, \dots, y_{j_k} , where j_k is the first index to satisfy (3.4) or (3.5). Set $x_k = y_{j_k}$.

Set $w_0 = 0$, and generate conjugate gradient iterates w_1, \dots, w_{j_k} for the reduced system (3.6). Set $d_k = Z_k w_{j_k}$, where j_k is the first index that satisfies (3.8).

Use a projected search to generate x_{k+1} . If (3.10) holds, reduce η_2 , and continue with the conjugate gradient method.

Our outline of algorithm GPCG does not include the termination test. An advantage of the termination test (2.4) is that this test is satisfied [9] in a finite number of iterations. On nondegenerate problems GPCG terminates [19] at the solution in a finite number of iterations.

Algorithm GPCG is suitable for large problems. As opposed to some other active set methods, each iteration is capable of adding or removing multiple constraints from the active set. Moreover, as we shall see, GPCG tends to require few iterations for convergence. Another advantage of the GPCG algorithm is that convergence can be achieved while requiring only approximate solutions to the linear systems.

4 Software Design

The TAO design philosophy uses object-oriented techniques of data and state encapsulation, abstract classes, and limited inheritance to create a flexible optimization toolkit. This section provides a short introduction to our design philosophy by describing the objects needed to create GPCG.

Our current implementation leverages the parallel computing and linear algebra infrastructure offered by PETSc [2, 3], which employs MPI [14] for all interprocessor communication. TAO optimization algorithms use high-level abstract data objects that are provided by PETSc, including vectors, matrices, and index sets. In this context, a vector (**Vec**) is an abstraction of an array of values that represent a discrete field, and a matrix (**Mat**) represents a discrete linear operator that maps between vector spaces. An index set (**IS**) is a generalization of a set of integer indices, which can be used for selecting, gathering, and scattering subsets of vector and matrix elements. TAO also interfaces to the linear solvers (**SLES**) within PETSc. Because each of these abstractions has several underlying representations, TAO has easy access to a variety of parallel vector and sparse matrix implementations as well as preconditioners and Krylov subspace methods.

Solving an optimization problem with TAO requires first creating a context data type called **TAO_SOLVER**, which encapsulates information about the solution process, including the algorithm, convergence tolerances, options, and parameters. All of the computations and communications related to a particular solution process are managed in the solver context variable. After defining the optimization problem, the user then calls **TaoSolve** to determine the solution. Finally, the user destroys the TAO solver via **TaoDestroy**. The code fragment in Figure 4.1 shows the main functions needed to solve bound-constrained quadratic programming problems with TAO.

```
TaoCreate(MPI_Comm comm,TaoMethod method,TAO_SOLVER *tao);
TaoSetQuadraticFunction(TAO_SOLVER tao,Vec X,Vec G,Mat A,Vec B,double c);
TaoSetVariableBounds(TAO_SOLVER tao,Vec XL,Vec XU);
TaoSolve(TAO_SOLVER tao);
TaoDestroy(TAO_SOLVER tao);
```

Figure 4.1: TAO interface for GPCG

The function **TaoCreate** creates the **TAO_SOLVER** context for one of several possible methods (denoted by **TaoMethod**) for solving the problem. This interface serves several algorithms for bound-constrained quadratic problems in addition to GPCG, including limited memory variable metric, trust region Newton, and interior point techniques. Moreover, this single interface serves other types of optimization problems as well. The function **TaoSetQuadraticFunction** in Figure 4.1 defines the objective function (2.2) in terms of the **Mat** object **A**, **Vec** object **B**, and scalar **c** and provides the **Vec** objects **X** and **G** that are used for the solution and gradient.

The function `TaoSetVariableBounds` defines upper and lower bounds for the variables \mathbf{X} with the `Vec` objects `XL` and `XU`. Additional routines may be used to specify the starting point and various options for the optimization solver, but the structure in Figure 4.1 is needed in all cases. Detailed information can be found in the TAO User Guide [4, 5].

TAO implements the GPCG algorithm as a sequence of well-defined routines. The evaluation of the function and gradient of the quadratic q , for instance, can be implemented through the standard numerical operations of matrix-vector multiplication, vector inner product, and vector `saxpy`. TAO passes `Mat` and `Vec` objects, whose representation is independent of our implementation of GPCG, to external tools that perform the numerical computations. Additional work vectors required by the algorithm are created by calling a routine that clones the variable vector \mathbf{X} in Figure 4.1.

Users working in a parallel environment must provide TAO with data structures `A`, `B`, `X`, `G`, `XL`, and `XU` that are properly distributed over the processors. Appropriate distribution allows efficient executions of the matrix-vector multiplication, vector inner product, and vector `saxpy` operations. Numerical toolkits such as PETSc facilitate the creation of these objects and provide the functionality for most of the required numerical operations.

The operations required to implement the GPCG algorithm as outlined in Section 3 include the vector and matrix operations listed in the preceding paragraph, functions to compute the pointwise minimum and maximum of two vectors, and a function that creates an index set that defines the indices where the elements of two vectors are equal.

At each iteration of the GPCG algorithm, we also need to apply the conjugate gradient method to the matrix A_k corresponding to the free variables. This is an important phase of the computation because, as we shall see in Section 5, at least 70% of the GPCG computing time is due to the conjugate gradient method. An efficient parallel implementation of the conjugate gradient method requires that the reduced matrix A_k be evenly distributed over the processors, but since the set of free variables may not be well distributed over the processors, the reduced matrix may not well distributed—regardless of how the matrix A is distributed. Since an unbalanced load can result in tremendous losses in performance, a redistribution of the rows of A_k over the processors may be necessary. We end this section by discussing the implementation of the conjugate gradient method for solving the reduced problem in the free variables.

At least two techniques exist for applying the conjugate gradient method to the reduced system of equations. One technique creates a second matrix A_k that contains the rows and columns of A corresponding to the free variables, and then applies the conjugate gradient method to the reduced system. An alternative technique applies the conjugate gradient method to the rows and columns of the full matrix A specified by the index set of the free variables. In our implementation, we chose the first method. Despite the additional memory requirements and cost of copying data, this method is simpler, facilitates the preconditioning and load-balancing of the reduced matrix, and was easily implemented with the utilities provided by PETSc.

Our implementation of GPCG calls `MatExtractSubmatrix(Mat,IS,IS,Mat *)`, which

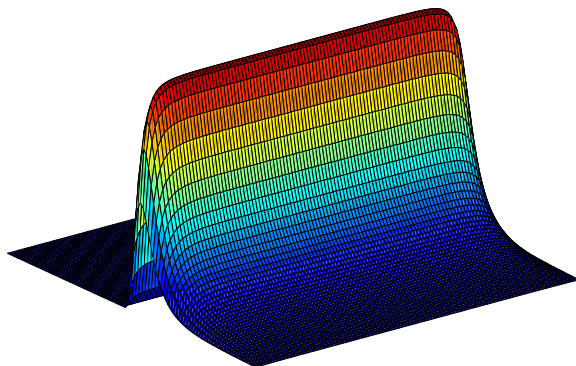


Figure 5.1: The journal bearing problem with $\varepsilon = 0.9$.

accepts the matrix A and the index set that identifies the set of free variables, and creates the reduced matrix A_k . A call to `VecCreateSubVec(Vec,IS,Vec*)` accepts the gradient vector and index set identifying the free variables to create a new, reduced vector. In a parallel environment, the index sets also define the distribution of the reduced matrix over the processors. These operations require a careful implementation when load balancing issues are taken into consideration.

We interface to the preconditioned conjugate gradient method provided by the **SLES** component of PETSc. We use the **SLES** object to define this iterative method, its preconditioner, the solution tolerance, and an initial point. The routine `LinearSolve(SLES,Mat,Vec,Vec)` computes an approximate solution to the linear system using the **SLES** object. At each iterate we create the conjugate gradient solver, apply it to the reduced linear system, and then destroy it.

In the entire implementation of GPCG no assumptions are made about the representations of data in the vectors and matrices. This approach eliminates some of the barriers in using independently developed software components by accepting data that is independent of representation and interfacing to numerical routines with the appropriate data formats.

5 Performance

We have evaluated the performance of the GPCG implementation on a variety of architectures. The data presented in this section was generated on the IBM SP (each processor has 256 MB RAM, 128 KB cache for data, and a 32 KB cache for instructions) at Argonne National Laboratory; performance trends were similar on other machines.

As a benchmark application we have used a journal bearing model, a variational problem over a two-dimensional region. This problem arises in the determination of the pressure distribution in a thin film of lubricant between two circular cylinders. The infinite-dimensional

version of this problem is of the form

$$\min\{q(v) : v \geq 0, v = 0 \text{ on } \partial D\},$$

where $v : \mathcal{D} \mapsto \mathbb{R}$ is piecewise continuously differentiable, $q : H^1 \rightarrow \mathbb{R}$ is the quadratic

$$q(v) = \int_{\mathcal{D}} \left\{ \frac{1}{2} w_q(x) \|\nabla v(x)\|^2 - w_l(x) v(x) \right\} dx,$$

$\mathcal{D} = (0, 2\pi) \times (0, 2b)$ for some constant $b > 0$, and

$$w_q(\xi_1, \xi_2) = (1 + \varepsilon \cos \xi_1)^3, \quad w_l(\xi_1, \xi_2) = \varepsilon \sin \xi_1,$$

where ε in $(0, 1)$ is the eccentricity parameter. The eccentricity parameter influences, in particular, the difficulty of the problem. Figure 5.1 shows the solution of the journal bearing problem for $\varepsilon = 0.9$. The steep gradient in the solution makes this problem a difficult benchmark.

Discretization of the journal bearing problem with either finite differences or finite elements leads to a problem of the form (1.1) with $l \equiv 0$ and $u \equiv +\infty$. The number of variables is $n = n_x n_y$, where n_x and n_y are, respectively, the number of grid points in each coordinate direction of the domain \mathcal{D} . See [19] for a description of the finite element discretization.

We now analyze the performance of GPCG on large problems, that is, problems that will not fit into the memory of a single processor. Specifically, we used a grid with 1600 points in each direction, leading to a problem with $n = 2.56 \cdot 10^6$ variables.

The initial point x_0 was set to the lower bound l . We used $\eta_1 = 0.1$ in the test (3.5) to terminate the gradient projection algorithm and $\eta_2 = 0.05$ in the test (3.8) to terminate the conjugate gradient algorithm. We stopped GPCG when the convergence test (2.4) was satisfied with $\tau = 10^{-4}$.

Table 5.1 presents performance data for GPCG. We show the number of processors p , the number of GPCG iterates (iters), the number of conjugate gradient iterations n_{GP} , the wall clock solution time (in seconds), the percentage of time ($t_{CG}\%$) used by the conjugate gradient algorithm, and the efficiency (\mathcal{E}) of GPCG in going from 8 to 64 processors. The time in the conjugate gradient algorithm includes the time spent computing the preconditioner. Our design allows the use of several preconditioners, but for the results in this section we used a block Jacobi preconditioner with one block per processor, where each subproblem was solved with ILU(2).

The results in Table 5.1 are noteworthy in several ways. First, the number of iterations of GPCG is remarkably small. This is surprising because the feasible set (2.1) has 3^n faces, and the GPCG visits only one face on each iteration. Other strategies can lead to a large number of iterates, but the GPCG algorithm is remarkably efficient.

Another interesting aspect of the results in Table 5.1 is that due to the low memory requirements of iterative solvers, we were able to solve these problems with only $p = 8$ processors. Strategies that rely on direct solvers are likely to need significantly more storage,

Table 5.1: Performance of GPCG on the journal bearing problem with $n = 2.56 \cdot 10^6$.

ε	p	iters	n_{GP}	time	$t_{CG}\%$	\mathcal{E}
0.1	8	46	431	7419	86	100
0.1	16	45	423	3706	83	100
0.1	32	45	427	2045	82	91
0.1	64	45	427	1279	82	73
0.9	8	37	105	2134	70	100
0.9	16	37	103	1124	71	95
0.9	32	38	100	618	69	86
0.9	64	38	99	397	68	67

and thus more processors. Finally, these results show that the GPCG implementation has excellent efficiency with respect to $p = 8$ processors, ranging between 67% and 100%. This sustained efficiency is remarkable because the GPCG algorithm is solving a sequence of linear problems with a coefficient matrix set to the submatrix of the Hessian of q with respect to the free variables for the current iterate. Thus, our implementation’s repartitioning of submatrices deals effectively with the load-balancing problem that is inherent in the GPCG algorithm.

For these results we have noted that as ε increases, both $t_{CG}\%$ and the overall efficiency decrease. This observation follows from the empirical result that the number of free constraints at the solution is inversely proportional to the eccentricity parameter ε . In particular, roughly 68% of the constraints are free at the solution when $\varepsilon = 0.1$, and 54% are free for $\varepsilon = 0.9$. Since the size of the linear system that the conjugate gradient algorithm needs to solve increases as ε decreases, the time required by the conjugate gradient algorithm increases. Since the parallel efficiency of larger problems is greater than the parallel efficiency for smaller problems, the overall efficiency of GPCG increases.

6 Performance Analysis

GPCG is typical of optimization algorithms that must deal with constrained problems in the sense that these algorithms have dynamically changing active sets. In this section we analyze the performance of GPCG.

Table 6.1 presents performance results for the journal bearing problem with dimension 640,000. In comparing these results with those of the larger problem in Table 5.1, note that while the number of variables increases by a factor of four, the number of iterations, the number of gradient projection iterates, and the time for solving the problem, increase by about a factor of two. This seems to be fairly typical of GPCG but may not hold for other optimization algorithms. Some algorithms for unconstrained problems exhibit mesh invariance in the sense that the number of iterations is independent of the number of variables, but this does not generally hold for constrained problems.

When analyzing the parallel performance of an algorithm, we must bear in mind that

Table 6.1: Performance of GPCG on the journal bearing problem with $n = 640,000$.

ε	p	iters	n_{GP}	time	$t_{CG}\%$	\mathcal{E}
0.1	2	27	227	2057	79	100
0.1	4	26	227	1173	79	89
0.1	8	27	232	639	78	80
0.1	16	26	231	365	75	70
0.1	32	27	230	220	74	58
0.1	64	27	228	152	75	42
0.9	2	21	58	645	65	100
0.9	4	20	54	368	63	88
0.9	8	20	52	199	64	81
0.9	16	21	54	128	64	63
0.9	32	20	52	74	61	54
0.9	64	23	54	58	62	35

a problem can scale well only when the ratio of computation to communication time is sufficiently large. Thus, for a particular problem size, scalability tapers off when more processors are added than can be used effectively. For GPCG, this effect can be seen clearly by comparing the results in Table 6.1 with those in Table 5.1.

An important aspect of the results in Table 6.1 is that for this particular problem of dimension 640,000, the efficiency of GPCG is acceptable for $p \leq 8$ processors but drops rapidly with more processors. To explain the drop in efficiency, we list in Table 6.2 the percentage of time spent in the main operations of GPCG. Note that some of these operations overlap, so the sum of the percentages always exceed 100%. In this table *Vec Red* refers to vector reductions, such as dot products and norms, while *Vec Local* refers to vector operations such as $y \leftarrow \alpha x + y$.

Table 6.2: Scalability of GPCG functions ($n = 640,000, \varepsilon = 0.1$)

Number Proc.	Percentage of time					Total MFlops	
	Mat-Vec Multiply	Vec Local	Vec Red	Linear Solve	Extract Submatrix	Linear Solve	TAO Solve
1	27	15	7	81	1	26	23
2	30	15	8	83	2	47	42
4	30	12	8	82	2	94	82
8	29	11	10	81	2	179	156
16	26	10	14	78	2	333	279
32	24	9	22	78	2	563	473
64	20	5	36	78	2	790	665

The percentage of time spent in the various functions of GPCG generally decreases slightly as the number of processors increases, with the exception of the vector reductions. Since vector reductions require communication among all processors, they have a significant effect on the efficiency of the algorithm. Note that the time for vector reductions remains

fairly constant at about 8% of the total computation time for 1–8 processors but that the efficiency of the algorithm declines quickly as the percentage of time doing vector reductions increases to 36% on 64 processors. This analysis shows that the ratio of computation to communication for this problem is too small for large number of processors and is responsible for the loss in scalability of GPCG for $p > 8$.

In this discussion of efficiency bear in mind that the Hessian matrix of the journal bearing problem is relatively sparse with 5 nonzeros per row on average. The efficiency is likely to improve if we deal with matrices with more nonzeros per row, since then the amount of computation per conjugate gradient iteration increases. These problems arise, for example, in three-dimensional simulations or in variational problems with vector functions, that is, variational problems that require determining a vector-valued $v : \mathcal{D} \mapsto \mathbb{R}^m$ for $m > 1$ that minimizes the quadratic q .

A surprising aspect of the results in Table 6.1 is that the percentage of time required to extract the submatrix remains nearly constant at 2% of the total computation time, demonstrating the relative efficiency of this phase of the computation. These results are surprising because at first sight the need to extract an arbitrary submatrix and to rebalance the distribution of rows across the processors would destroy the efficiency of the algorithm. On the other hand, the creation of a second matrix to hold the submatrix requires additional storage. For large problems the additional storage may exceed the memory capacity of a small number of processors.

Another important component of our scalability analysis is the flop rate per processor. As noted in Table 6.1, the flop rate for the linear solve component of GPCG is 26 MFlops for one processor and decreases to about 12.3 for 64 processors. For comparison purposes, the flop rate of a Newton algorithm in PETSc is about 42 MFlops for one processor on a system of nonlinear equations with the same sparsity as the journal bearing problem. This rate is higher than the rate achieved by the GPCG algorithm, but this is to be expected because, as previously mentioned, the GPCG algorithm spends a significant amount of time on tasks with no arithmetic operations. The extraction of the submatrix, creating the reduced linear system and determining the free variables, typically requires more than 10% of the time. Hence, it is unlikely that the GPCG algorithm, or any active set algorithm for constrained problems, can achieve a computation rate as high as a Newton algorithm.

While these computations employed a standard compressed, sparse row format for matrix data, higher flop rates could be obtained on some problems, changing the matrix format. Alternative storage schemes that exploit the structured sparsity of these problems would achieve higher flop rates for matrix operations by alleviating unnecessary memory references. Likewise, block sparse storage variants for problems with multiple unknowns per grid point would achieve higher flop rates [15]. Since our optimization algorithms use a data-structure-neutral interface to matrix and vector operations, we can easily experiment with such alternatives without altering any of the optimization code.

7 Preconditioners

The ability to experiment with various preconditioners is a direct result of our design philosophy, which enables connection to the linear algebra infrastructure provided in toolkits such as PETSc. In particular, we compared the diagonal Jacobi preconditioner with a block Jacobi preconditioner that used one block per processor. We employed sparse matrix based ILU as a subdomain solver for the block Jacobi method, where we considered both $\text{ILU}(0)$, which produced a factored matrix that maintained the same sparsity pattern as the subdomain matrix, and $\text{ILU}(2)$, which allowed two levels of fill.

The statistics summarized in Table 7.1 are the eccentricity parameter ε , the number of processors p , the number of iterations of GPCG, the time required to solve the problem, and the number of conjugate gradient iterations. We present results only for $n = 640,000$, since similar results were obtained for $n = 2,560,000$.

Table 7.1: Performance of preconditioners in GPCG ($n = 640,000$)

ε	p	Diagonal			Block Jacobi - $\text{ILU}(0)$			Block Jacobi - $\text{ILU}(2)$		
		iters	time	CG iters	iters	time	CG iters	iters	time	CG iters
0.1	4	26	2928	37045	27	1324	8679	26	1173	6312
0.1	16	26	851	37045	27	409	9105	26	364	6712
0.9	4	21	1216	18118	20	416	2654	20	368	1864
0.9	16	22	390	18118	23	150	3390	21	128	2303

The number of GPCG iterations in Table 7.1 is independent of the number of processors and of the preconditioner. In general we expect small variations in the number of iterations because different preconditioners create different approximate solutions to linear systems and different paths to the solution.

In these experiments we were interested in the impact of the preconditioner on the total time to solution. The Jacobi method is scalable, so the main issue is whether the higher computational cost of the block Jacobi is justified. As expected, the block Jacobi preconditioner with subdomain solver $\text{ILU}(2)$ required fewer conjugate gradient iterations than subdomain solver $\text{ILU}(0)$, and both block Jacobi preconditioners required fewer iterations than the point Jacobi method. In addition, the block Jacobi methods also required less time. In general, better preconditioners require more time to compute, and this additional cost sometimes negates the savings achieved from fewer iterations of the linear solver. In this problem, the block Jacobi preconditioners used about half of the time required by the diagonal preconditioner, and the additional cost of computing better preconditioners is justified. The most expensive preconditioner to compute of the three under consideration in this work, namely, the block Jacobi method with subdomain solver $\text{ILU}(2)$, produced the fewest iterations by the conjugate gradient method and the smallest overall solution time.

The ability to experiment easily with a variety of preconditioners is an advantage because we can then choose a technique that is most suitable to the problem. In this spirit, we plan to

experiment with the evolving interfaces under development by the Equation Solver Interface (ESI) [12] and Common Component Architecture (CCA) [1, 10] working groups, with a goal of enabling dynamic use within TAO of any ESI-compliant preconditioning components.

8 Concluding Remarks

We have shown that the TAO design leverages external parallel computing infrastructure and linear algebra toolkits to solve large-scale optimization problems on high-performance architectures. With the exception of the work of Biros and Ghattas [6, 7], other codes for large-scale optimization problems are either custom-written or restricted to uni-processor environments.

TAO [4, 5] extends to general nonlinearly bound-constrained optimization, but the performance issues are more subtle due to the impact of user-supplied function, gradient and Hessian code. Extensions of TAO to large linearly-constrained and nonlinearly-constrained optimization problems is currently an active research area.

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