Solving Large-Scale Linear Programs by Interior-Point Methods Under the MATLAB Environment *

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Abstract

In this paper, we describe our implementation of a primal-dual infeasible-interior-point algorithm for large-scale linear programming under the MATLAB¹ environment. The resulting software is called LIPSOL – Linear-programming Interior-Point SOLvers. LIPSOL is designed to take the advantages of MATLAB's sparse-matrix functions and external interface facilities, and of existing Fortran sparse Cholesky codes. Under the MATLAB environment, LIPSOL inherits a high degree of simplicity and versatility in comparison to its counterparts in Fortran or C language. More importantly, our extensive computational results demonstrate that LIPSOL also attains an impressive performance comparable with that of efficient Fortran or C codes in solving large-scale problems. In addition, we discuss in detail a technique for overcoming numerical instability in Cholesky factorization at the end-stage of iterations in interior-point algorithms.

Keywords: Linear programming, Primal-Dual infeasible-interior-point algorithms, MATLAB, LIPSOL.

Abbreviated Title: Solving Large Linear Programs under MATLAB

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1 Introduction

After over a decade of extraordinarily active research triggered by the seminal work of Karmarkar [5], the field of interior-point methods has finally come to maturity as far as linear programming is concerned. Not only do we have a solid theoretical foundation for interior-point methods for linear programming, but also a rather comprehensive understanding on their practical efficiency. Among many general algorithmic approaches, the most effective one in practice has proven to be the primal-dual infeasible-interior-point approach, including a number of variants and enhancements such as Mehrotra's predictor-corrector technique [13].

Recent experiments indicate that as the problem size increases, so does the frequency of interior-point methods outperforming the classic simplex method [12]. This trend has been observed for a while but is becoming more pronounced recently. In our opinion, this marks the beginning of a new era in which interior-point methods coexist with the simplex method but gradually assume the role of dominant computational engine for general large-scale linear programming

Beside computational efficiency, another advantage of interior-point algorithms is that their implementations for large-scale linear programming, through still quite involved, are considerably simpler than those for the simplex algorithms. This is particularly true under a high-level programming environment such as MATLAB that supports high-level sparse-matrix operations.

The main purpose of the paper is to describe our implementation of a large-scale primal-dual infeasible-interior-point algorithm under the MATLAB environment and to present computational results. Hence we will concentrate on the implementation aspects of interior-point methods. There is a vast volume of literature on the theoretic aspects of interior-point methods. We refer interested readers to the *Interior Methods Online* web page at the Argonne National Laboratory ² for more informations on interior-point methods, including a comprehensive bibliography containing works up to 1993. For survey papers concerning computation and implementation issues, we refer readers to [11] and [1].

In Section 2, we describe the general framework for primal-dual infeasible-interior-point algorithms. We present the framework as a modified form of the well-known Newton's method and give intuitions to justify the modifications to the Newton method. These motivations are easily understandable to readers with a basic knowledge on Newton's method. In doing so, we hope that this section can also serve a pedagogical purpose. Readers who are familiar with interior-point algorithms may choose to skip parts of Section 2.

Section 3 is a brief description of the MATLAB technical computing environment under which

 $^{^2\}mathrm{URL}$: http://www.mcs.anl.gov/home/otc/InteriorPoint/index.html

we developed our software LIPSOL. We discuss a number of major issues in implementation in Section 4. Section 5 is devoted to discussing a technique for overcoming numerical instability in Cholesky factorization. Numerical results are presented in Section 6 and finally concluding remarks in Section 7.

2 Infeasible-Interior-Point Methods

In this section, we will motivate the formulation of primal-dual infeasible-interior-point methods from a practical algorithmic point of view, exploring the close tie between the classic Newton's method and the primal-dual infeasible-interior-point methods. We start from a description of linear programming.

2.1 Linear Programming

We consider the following primal linear program:

(1)
$$\min \quad c^{T}x$$
s.t.
$$Ax = b$$

$$0 \le x_{i} \le u_{i} \quad i \in \mathcal{I}$$

$$0 \le x_{i} \quad i \in \mathcal{J}$$

where $A \in \mathbf{R}^{m \times n}$, which determines the sizes of other vectors involved, and \mathcal{I} and \mathcal{J} are disjoint index sets such that $\mathcal{I} \cup \mathcal{J} = \{1, 2, \dots, n\}$ and $\mathcal{I} \cap \mathcal{J} = \emptyset$.

Without loss of generality, let us assume that for some positive integer $n_u \leq n$

(2)
$$\mathcal{I} = \{1, 2, \dots, n_u\}, \text{ and } \mathcal{J} = \{n_u + 1, n_u + 2, \dots, n\}.$$

Given a vector x in \mathbf{R}^n , we use the notation x_u for the vector in \mathbf{R}^{n_u} whose elements are the first n_u elements of x, i.e.,

$$[x_u]_i = x_i, i = 1, 2, \cdots, n_u.$$

Moreover, we define the appending operator "app" from \mathbf{R}^{n_u} to \mathbf{R}^n that appends $n - n_u$ zeros to vectors in \mathbf{R}^{n_u} , i.e., for $w \in \mathbf{R}^{n_u}$

$$[\operatorname{app}(w)]_i = \begin{cases} w_i, & 1 \le i \le n_u \\ 0, & n_u < i \le n. \end{cases}$$

By adding the slack variables $s \in \mathbf{R}^{n_u}$, we can rewrite the above linear program into the standard

form:

(4)
$$\min c^{T} x$$
s.t.
$$Ax = b$$

$$x_{u} + s = u$$

$$x \ge 0, \ s \ge 0$$

The dual of the above standard primal linear program is:

(5)
$$\max \qquad b^{T}y - u^{T}w$$
$$\text{s.t.} \quad A^{T}y + z - \text{app}(w) = c$$
$$z > 0, \ w > 0$$

where y, z and w are the dual variables and slacks.

It is well-known that the solutions of the primal and the dual linear programs, if they exist, satisfy the following KKT conditions which is a system of linear-quadratic equations with nonnegativity constraints on some variables:

(6)
$$F(x, z, s, w, y) = \begin{pmatrix} Ax - b \\ x_u + s - u \\ A^T y + z - \operatorname{app}(w) - c \\ xz \\ sw \end{pmatrix} = 0, (x, z, s, w) \ge 0,$$

where xz and sw denote component-wise multiplications and the equations xz = 0 and sw = 0 are called the complementarity conditions for the linear program.

For nonnegative variables x, s, z, w, we will call the quantity $x^Tz + s^Tw$ the duality gap (regardless of feasibility or infeasibility of these variables). The duality gap measures the residual of the complementarity portion of F in ℓ_1 -norm when $(x, z, s, w) \geq 0$.

A straightforward calculation shows that the Jacobian matrix of F(x, z, s, w, y) is

(7)
$$F'(x, z, s, w, y) = \begin{bmatrix} A & 0 & 0 & 0 & 0 \\ E^T & 0 & I_u & 0 & 0 \\ 0 & I & 0 & -E & A^T \\ Z & X & 0 & 0 & 0 \\ 0 & 0 & W & S & 0 \end{bmatrix}.$$

where $E^T = \begin{bmatrix} I_u & 0 \end{bmatrix}$, I_u is the identity matrix of dimension n_u , $X = \operatorname{diag}(x)$, $Z = \operatorname{diag}(z)$, $S = \operatorname{diag}(s)$ and $W = \operatorname{diag}(w)$.

Unlike a primal (dual) method which concentrates on solving the primal (dual) program, a primal-dual interior-point method for linear programming solves the KKT system (6), which includes all the primal and dual variables and slacks.

2.2 Newton's Method and Variants

It is clear from (6) that solving a linear program is equivalent to solving a system of linearquadratic equations with nonnegativity constraints on a subset of variables. In order to simplify our coming discussion, we rewrite the KKT system (6) into a constrained algebraic system of ℓ equations and ℓ variables with nonnegativity constraints on ℓ_+ variables:

(8)
$$F(v) = 0, v_i \ge 0, 1 \le i \le \ell_+,$$

where $v = (x, z, s, w, y), \ell = 2n + 2n_u + m \text{ and } \ell_+ = 2n + 2n_u.$

Let us first drop the nonnegativity constraints from (8) and consider Newton's method for solving an unconstrained system of nonlinear equations

$$F(v) = 0,$$

which can be written as

(9)
$$v^{k+1} = v^k - F'(v^k)^{-1}F(v^k).$$

It is well-known that Newton's method has excellent local convergence properties. More specifically, if the Jacobian matrix F'(v) is nonsingular and Lipcshitz continuous at a solution v^* , and the initial point v^0 is sufficiently close to v^* , then the iterate sequence $\{v^k\}$ converges to v^* Q-quadratically. On the other hand, Newton's method generally does not have very good global convergence properties.

A variant of Newton's method is called the damped Newton:

(10)
$$v^{k+1} = v^k - \alpha^k F'(v^k)^{-1} F(v^k),$$

which introduces a damping factor, or step length, α^k , usually chosen from the interval (0, 1], to enhance global convergence.

Another variation of Newton's method is the so-called composite Newton's method. At each iteration, it calculates an intermediate point \hat{v}^k and uses the same Jacobian matrix twice:

$$\hat{v}^k = v^k - F'(v^k)^{-1}F(v^k), \quad v^{k+1} = \hat{v}^k - F'(v^k)^{-1}F(\hat{v}^k).$$

Equivalently, we can shorten the expression as

(11)
$$v^{k+1} = v^k - F'(v^k)^{-1}(F(v^k) + F(\hat{v}^k)).$$

In terms of linear algebra work, the composite Newton requires one matrix factorization per iteration, same as for Newton's method, but two back-solves instead of one. Since matrix factorizations are usually much more expensive than back-solves $(O(n^3) \text{ vs. } O(n^2) \text{ for dense matrices})$,

the required work per iteration for the composite Newton is comparable with that for Newton's method. However, under similar conditions, the composite Newton has a Q-cubic asymptotic convergence rate, one order faster than that of Newton's method.

Similarly, one can introduce a damping factor into the composite Newton's method:

(12)
$$v^{k+1} = v^k - \alpha^k F'(v^k)^{-1} (F(v^k) + F(\hat{v}^k)).$$

For more detailed information on Newton's method and its variants, we refer to the classic book by Ortega and Rheinboldt [16].

2.3 Infeasible-Interior-Point Methods

Let us now go back to the nonnegativity constrained system of equations (8) that represents a linear programming in a simplified notation. We recall that for linear programming, F(v) contains only linear and quadratic equations and the quadratic ones are complementarity pairs.

Obviously, in the presence of the nonnegativity constraints for the first ℓ_+ variables we cannot expect to obtain a nonnegative solution by just simply applying the Newton's method. However, a crucial observation is that the damped Newton (10) will generate a sequence of iterates $\{v^k\}$ that satisfies the nonnegativity constraints if (i) we start from a initial point v^0 whose first ℓ_+ components are positive, i.e., $v_i^0 > 0$, $1 \le i \le \ell_+$, and (ii) we choose the step-length α at each iteration so that these first ℓ_+ components remain positive in all the subsequent iterations.

It is important to note the necessity of keeping the nonnegative variables (the first ℓ_+ component) strictly positive at every iteration instead of just nonnegative. Consider a complementarity condition sw = 0. The Newton equation (i.e., the linearized equation) for sw = 0 at a given point (s, w) is $s\Delta w + w\Delta s = -sw$. If one variable, say w_i , is zero, then the Newton equation becomes $s_i\Delta w_i = 0$, leading to a zero update $\Delta w_i = 0$. Consequently, w_i will remain zero all the time once it becomes zero. This is fatal because an algorithm will never be able to recover once it sets a variable to zero by mistake. Therefore, it is critical to keep the nonnegative variables strictly positive at every iteration, not just nonnegative.

Even keeping nonnegative variables strictly positive, one would still expect difficulty in recovering from a situation where a variable is adversely set to too small a value. To decrease the chances of such mistakes at early stages, it would be desirable that all the complementarity pairs converge to zero at the same pace, say $s_i^k w_i^k = \mu^k \to 0$ as $k \to \infty$ for every index i. Towards this goal (and for other theoretic considerations as well), one can perturb each complementarity condition $s_i w_i = 0$ into $s_i w_i - \mu = 0$ and let μ go to zero in a controlled fashion. This idea, incorporated into the damped Newton (10), leads to the following framework: choose v^0 such that

all the linear equations in F(v)=0 (i.e., feasibility equations) are satisfied and $v_i^0>0,\ i\leq \ell_+;$ for $k=0,1,\cdots,$ do

$$v^{k+1} = v^k - \alpha^k F'(v^k)^{-1} (F(v^k) - \mu^k \hat{e}),$$

where $\mu^k > 0$, $\alpha^k > 0$ is chosen such that $v_i^{k+1} > 0$, $i \leq \ell_+$, and the vector $\hat{e} \in \mathbf{R}^n$ is defined as

$$\hat{e}_i = \begin{cases} 0, & \text{if } F_i(v) \text{ is linear,} \\ 1, & \text{if } F_i(v) \text{ is quadratic.} \end{cases}$$

Clearly, this definition indicates that perturbations are only applied to the complementarity conditions, which are all quadratic, but not to the feasibility conditions, which are all linear.

The above primal-dual interior-point approach was first proposed by Kojima, Mizuno and Yorshisi [6], in which a feasible initial point is required and the linear equations are always satisfied. According to the classification introduced in [19], this type of methods is called feasible-interior-point methods.

Lustig, Marsten and Shanno [9] implemented a modified version of the Kojima-Mizuno-Yorshisi interior-point approach. A fundamental modification is the removal of the requirement for a feasible initial point which is practically undesirable. The Lustig-Marsten-Shanno algorithm can be schematically cast into the following infeasible-interior-point (IIP) algorithmic framework:

Algorithm IIP-1:

Choose an initial point v^0 such that $v_i^0 > 0$, $1 \le i \le \ell_+$. Set k = 0.

Do until $||F(v^k)||$ is "small"

- (1) Choose $\mu^k > 0$ and compute $\Delta v^k = -F'(v^k)^{-1}(F(v^k) \mu^k \hat{e})$. (2) Choose $\alpha^k > 0$ and set $v^{k+1} = v^k + \alpha^k \Delta v^k$
- (2) Choose $\alpha^k > 0$ and set $v^{k+1} = v^k + \alpha^k \Delta v^k$ such that $v_i^{k+1} > 0$, $1 \le i \le \ell_+$. Increment k by 1.

End

Mehrotra [13] introduced an enhanced version of the infeasible-interior-point algorithm that he called a predictor-corrector method. It was observed in [17] that Mehrotra's algorithm can also be considered as a modification to the damped composite Newton's method (12), which is schematically very close to the following algorithmic framework:

Algorithm IIP-2:

Choose an initial point v^0 such that $v_i^0 > 0$, $1 \le i \le \ell_+$. Set k = 0.

Do until $||F(v^k)||$ is "small"

- (1) Compute $\Delta^p v^k = -F'(v^k)^{-1}F(v^k)$ and determine $\mu^k > 0$.
- (2) Compute $\Delta^c v^k = -F'(v^k)^{-1}(F(v^k + \Delta^p v^k) \mu^k \hat{e}).$
- (3) Choose $\alpha^k > 0$ and set $v^{k+1} = v^k + \alpha^k (\Delta^p v^k + \Delta^c v^k)$ such that $v_i^{k+1} > 0$, $1 \le i \le \ell_+$. Increment k by 1.

End

In this framework, two steps are computed at each iteration: the predictor step $\Delta^p v^k$ and the corrector step $\Delta^c v^k$, both using the same matrix factorization. Extensive computational experiments [10, 13] have indicated that Mohrotra's infeasible-interior-point framework is by far the most efficient interior-point framework in practice. It usually requires considerably less iterations than algorithms in the framework of Algorithm IIP-1. Furthermore, since it requires only one matrix factorization per iteration, it is more efficient than algorithms based on the Mizuno-Todd-Ye [14] predictor-corrector approach which requires two matrix factorizations per iteration.

For proper choices of parameters and starting points, the global convergence and computational complexity of Algorithm IIP-1 were first established in [7] and [19], respectively; and the computational complexity of Algorithm IIP-2 was first established in [21].

There are a number of effective enhancements to Algorithm IIP-2. Xu et al [18] proposed and implemented a homogeneous and self-dual algorithm which applies the basic framework of Algorithm IIP-2 to a homogeneous and self-dual reformulation of a linear program. At the cost of some extra work at each iteration, this algorithm can more reliably detect infeasibility. Hence in the case of solving untested models with unknown feasibility, the homogeneous and self-dual algorithm may be preferable. Recently, Gondzio [4] proposed a technique called multiple centrality corrections that can further reduce the number of iterations required for convergence by adaptively adding one or more corrector steps to the framework of Algorithm IIP-2.

3 The MATLAB Environment

Our implementation was done under the MATLAB environment. MATLAB, standing for matrix laboratory, is a high-level technical computing environment for numeric computation and visualization. It integrates numerical analysis, matrix computation and graphics in an easy-to-use environment where matrix-computational formulas can be expressed in MATLAB scripts,

called M-files, that are very close to what they would be written mathematically - thus avoiding low-level programming as is needed in Fortran or C programming. Because of this advantage, development-time in MATLAB is much shorter than in other languages such as Fortran and C. However, for computation-intensive tasks overhead incurred in MATLAB generally makes it slower, which is a price one has to pay in exchange for programming simplicity.

MATLAB provides external interface facilities to enable interaction with programs written in Fortran or C languages in the form of the MEX-files, where MEX stands for MATLAB Executable. MEX-files are dynamically linked subprograms compiled from C or Fortran source code that can be called and run from within MATLAB in the same way as MATLAB M-files or built-in functions. The external interface functions provide functionality to transfer data between MEX-files and MATLAB.

Our implementation includes both M-files and MEX-files. The routine tasks, mostly matrix and vector operations, are done by M-files, while the most computation-intensive tasks are done by MEX-files generated from Fortran programs. This combination enables us to take advantage of both worlds: programming simplicity in MATLAB and computational speed in Fortran.

4 Implementation Details

Our implementation is based on the framework of Algorithm IIP-2. We still need to specify the following: (1) the initial point and the stopping criteria; (2) how to compute the steps; and (3) how to choose the parameters μ^k and α^k . As in most implementations, we use different step lengths in the primal and dual spaces to achieve better efficiency. In addition, there are a number of implementation issues that usually do not appear in algorithms but are extremely important to the quality of software. These include the exploitation of sparsity and the treatment of numerical instability as well as dense-columns. We will discuss these issues in some detail.

The software package resulted from our implementation, currently for UNIX systems only, is called LIPSOL (Linear-programming Interior Point SOLvers). LIPSOL runs under MATLAB version 4.0 and above.

4.1 Preprocessing, Starting and Stopping

The purpose of preprocessing is to transform a given linear program into the standard form (see (1)) and get rid of some redundancy from the formulation. Currently, the de facto industrial standard input format for linear programs is the MPS format (see [15] for an introduction to the MPS format). LIPSOL includes an MPS reader program mps2mat to read MPS files and

translate the data into MATLAB data files. In the MPS reader, a free variable is split into the difference of two nonnegative variables.

The preprocessing function preprocessing.m performs a number of tasks such as checking obvious infeasibility, deleting fixed variables and zero rows and columns from the matrix A, solving equations with only one variables and shifting nonzero lower bounds to zero, whenever any of these events occurs.

The scaling function scaling.m performs a row and a column scaling to the matrix A. We believe that those scalings help the solution process in some way, though we do not have hard evidence on how much benefits these scalings actually bring.

The M-file initpoint.m chooses starting point for iterations. The starting-point selection in LIPSOL basically follows the one used in [10] with some minor modifications.

The function stopping.m tests stopping criteria and, if any met, terminates the iterations. The main stopping criterion is the standard one:

(13)
$$\frac{\|r_b\|}{\max(1,\|b\|)} + \frac{\|r_c\|}{\max(1,\|c\|)} + \frac{\|Ru\|}{\max(1,\|u\|)} + \frac{|c^Tx - b^Ty + u^Tw|}{\max(1,|c^Tx|,|b^Ty - u^Tw|)} \le tol$$

where $r_b = Ax - b$, $r_c = A^Ty + z - \text{app}(w) - c$ and $r_u = x_u + s - u$ are, respectively, the residuals in primal, dual and upper-bound feasibility, $c^Tx - b^Ty + u^Tw$ is the difference between the primal and dual objective values, and tol is the tolerance which is set to 10^{-8} by default. The sum on the left measures the total relative errors in the KKT system (6). A number of heuristics are also included to detect infeasibility, though they have not been well tested on infeasible problems at this point and may change over time.

4.2 Centering and Step-Length Parameters

The centering parameter μ^k is computed by the function centering.m. It is chosen largely as suggested by Mehrotra [13] but with some minor modifications.

The value of μ^k is a fraction of the normalized duality gap

$$g(x, z, s, w) = \frac{x^T z + s^T w}{n + n_u}$$

evaluated at the current iterate (x^k, z^k, s^k, w^k) , i.e.,

$$\mu^k = \sigma^k g^k,$$

for some $\sigma^k \in (0,1)$. After computing the predictor step $\Delta^p v^k$. A ratio test is performed to determine $\hat{\alpha}_p$ such that

$$(\hat{x}, \hat{s}) = (x^k, s^k) + \min(1, \hat{\alpha}_p)(\Delta^p x^k, \Delta^p s^k)$$

is either positive in the case $\hat{\alpha}_p > 1$ or on the boundary of the nonnegative orthant $(x, s) \ge 0$ in the case $\hat{\alpha}_p \le 1$. Similarly, we compute

$$(\hat{z}, \hat{w}) = (z^k, w^k) + \min(1, \hat{\alpha}_d)(\Delta^p z^k, \Delta^p s^k)$$

such that it is either positive or on the boundary of the nonnegative orthant $(z, w) \geq 0$. The parameter σ^k is then set to

$$\sigma^k = \left(\frac{\hat{x}^T \hat{z} + \hat{s}^T \hat{w}}{(x^k)^T z^k + (s^k)^T w^k}\right)^2.$$

The above choice was first suggested by Mehrotra [13]. We also impose an upper bound on σ^k at each iteration to limit σ^k to a value smaller than one. At the early stage of the iterations, we set the upper bound to a value around 0.2. Near the end we force it to be in the same order as the total relative error. This strategy is designed to achieve fact local convergence at the end stage of the iterations.

Let us recall the notation v = (x, z, s, w, y) and define

$$(\Delta x^k, \Delta z^k, \Delta s^k, \Delta w^k, \Delta y^k) \equiv \Delta v^k = \Delta^p v^k + \Delta^c v^k,$$

i.e., the combined predictor-corrector step. The primal and dual step-length parameters α_p^k and α_d^k at each iteration are chosen so that the new iterate

$$v^{k+1} \equiv \begin{pmatrix} x^{k+1} \\ z^{k+1} \\ s^{k+1} \\ w^{k+1} \\ y^{k+1} \end{pmatrix} = \begin{pmatrix} x^k \\ z^k \\ s^k \\ w^k \\ y^k \end{pmatrix} + \begin{pmatrix} \alpha_p \, \Delta x^k \\ \alpha_d \, \Delta z^k \\ \alpha_p \, \Delta s^k \\ \alpha_d \, \Delta w^k \\ \alpha_d \, \Delta y^k \end{pmatrix}$$

lies on or just inside the boundary of the set

$$\mathcal{N} = \{ v : (x, z, s, w) > 0, \min(xz, sw) \ge \phi_0 g(x, z, s, w) \},\$$

where $\phi_0 = 10^{-5}$ by default. It is known that the membership in \mathcal{N} guarantees the positivity of (x, z, s, w). The set \mathcal{N} is often referred as a neighborhood of the (generalized) central path defined as the set

$$C = \{v : (x, z, s, w) \ge 0, \min(xz, sw) = \max(xz, sw)\},\$$

i.e., the components of the vectors xz and sw are all the same and, of course, equal to their average g(x, z, s, w). Staying in \mathcal{N} prevents the elements of (x, z, s, w) from becoming too close to zero at an early stage.

In our implementation, we use a back-tracking strategy to ensure that the new iterate v^{k+1} is on the boundary or just inside of \mathcal{N} . We start with

$$(\alpha_p, \alpha_d) = \tau_0(\bar{\alpha}_p, \bar{\alpha}_d)$$

where $(\bar{\alpha}_p, \bar{\alpha}_d)$ are computed by ratio tests and τ_0 is set to 0.9995 by default, and repeatedly decrease (α_p, α_d) by a certain factor, if needed at all, until the defining inequality of the set \mathcal{N} is satisfied. This process is implemented in the function update.m.

4.3 Computing the Steps

The major computation at each iteration is to solve the a system with the coefficient matrix F'(v) (see (7)) and two different right-hand sides, one for the predictor direction and the other for the corrector one. Both right-hand sides can be written in terms of vectors r_b, r_u, r_c, r_{xz} and r_{sw} . More specifically, the right-hand sides for the predictor and the corrector directions are, respectively, the negative of the following two vectors:

$$\begin{pmatrix} r_b \\ r_u \\ r_c \\ r_{xz} \\ r_{sw} \end{pmatrix} = \begin{pmatrix} Ax - b \\ x_u + s - u \\ A^Ty + z - \operatorname{app}(w) - c \\ xz \\ sw \end{pmatrix} \text{ and } \begin{pmatrix} r_b \\ r_u \\ r_c \\ r_{xz} \\ r_{sw} \end{pmatrix} = \begin{pmatrix} 0 \\ 0 \\ (\Delta^p x^k)(\Delta^p z^k) - \mu^k e_n \\ (\Delta^p s^k)(\Delta^p w^k) - \mu^k e_{n_u} \end{pmatrix}$$

where e_n and e_{n_u} are vector of all ones of dimension n and n_u , respectively. It is worth noting that the first vector is just $F(v^k)$, and the second $F(v^k + \Delta^p v^k)$ subtract the perturbation vector involving μ^k .

Since F'(v) is a highly structured sparse matrix, one can solve it by block Gaussian elimination which leads to the following procedure for both the predictor and the corrector steps corresponding to different right-hand sides:

Procedure Steps:

- (1) Form the matrix $D = [X^{-1}Z + \text{diag}(\text{app}(S^{-1}w))]^{-1}$.
- (2) Overwrite: $r_c \Leftarrow r_c X^{-1}r_{xz} + \operatorname{app}(S^{-1}(r_{sw} Wr_u))$.
- (3) Solve for Δy by Cholesky factorization from the linear system: $(ADA^T)\Delta y = -(r_b + ADr_c)$.

(4) Compute
$$\Delta x = DA^{T}(\Delta y + r_{c}),$$

$$\Delta z = -X^{-1}(Z\Delta x + r_{xz}),$$

$$\Delta s = -(\Delta x_{u} + r_{u}),$$

$$\Delta w = -S^{-1}(W\Delta s + r_{sw}).$$

This procedure is implemented in the M-file direction.m. Similar procedures were used in earlier implementations, see [9], [10] and [13].

4.4 Sparse Cholesky Factorization

For our large-scale interior-point code, the major computation at each iteration is in step (3) of Procedure Steps where the coefficient matrix ADA^T is formed and factored. Since D is a positive diagonal matrix, if A has full rank ADA^T is positive definite with the same sparsity pattern as AA^T . The efficiency of a large-scale interior-point code lies predominantly on the efficiency of the sparse Cholesky factorization code used.

Cholesky factorization of a sparse matrix, say $P = LL^T$ where L is lower triangular, generally involves three stages. First one need to reorder the rows and columns of the matrix P in an attempt to minimize the fillings – the new nonzero elements created during factorization. Then one performs a symbolic factorization to figure out the sparsity pattern of L. Finally, one carries out the actual factorization. In the context of interior-point algorithms, since the sparsity pattern of $P = ADA^T$, disregarding cancelation, remains the same for any positive diagonal matrix D, one only need to perform an ordering and a symbolic factorization once at the very beginning.

MATLAB does provide functions for ordering and factoring a sparse symmetric positive definite matrices. However, for the sake of numerical stability, we must be able to modify the standard Cholesky factorization procedure (see Section 5). This can not be done under MATLAB since source code access is unavailable for MATLAB's build-in functions. Therefore, we decided to use MATLAB's external interface facilities and existing Fortran sparse Cholesky code in our LIPSOL implementation. We constructed MEX-files from two Fortran packages: a sparse Cholesky factorization package (version 0.3) developed by Esmond Ng and Barry Peyton at the Oak Ridge National Laboratory (ORNL), and a multiple minimum-degree ordering package by Joseph Liu at University of Waterloo which is included in the ORNL package. The ORNL package uses relatively new techniques such as supernode block factorization to take advantage of modern computer architectures [8].

4.5 Dense Columns

Let us denote the *i*-th column of A by a_i . It is clear from

$$ADA^T = \sum_{i=1}^m d_{ii} a_i a_i^T$$

that ADA^T will be generally dense as long as there is one dense column in A. Although for most large-scale applications the occurrence of dense columns is rare, it does occur from time to time

and must be dealt with.

Our strategy for dense columns, like in many other codes, is to split the relatively dense columns, if any, from the sparse ones to form

$$ADA^T = P + UU^T$$

where P is the product of the sparse columns and U is the collection of dense columns multiplied by corresponding diagonal elements of $D^{1/2}$. Matrix U usually has a very low rank. If P is nonsingular, then by the Sherman-Morrison formula

$$(P + UU^{T})^{-1} = P^{-1} - P^{-1}U(I + U^{T}P^{-1}U)^{-1}U^{T}P^{-1},$$

the solution to the linear system $(P + UU^T)\Delta y = h$ is

$$\Delta y = \hat{h} - P^{-1}U(I + U^T P^{-1}U)^{-1}U^T \hat{h},$$

where $\hat{h} = P^{-1}h$. Of course, we never compute any inverse matrices, but just solve the relevant linear systems. Since U has very low rank, the matrix $I + U^T P^{-1}U$ is very small and requires a small number of back-solves to form once P is factored.

This approach is not always stable because P, the sparse portion of ADA^T , can be severely ill-conditioned or actually singular. To stabilize the process, we utilize a preconditioned conjugate gradient method whenever we find the residual for the solution from the Sherman-Morrison formula is unsatisfactory. The pre-conditioner is constructed from the Cholesky factor of $P = LL^T$ (possibly modified, see Section 5). Recall that the system we want to solve is

$$(LL^T + UU^T)\Delta y = h.$$

Our preconditioned conjugate gradient method mathematically amounts to applying the conjugate gradient method to the following preconditioned, equivalent system

$$[I + (L^{-1}U)(L^{-1}U)^T]\Delta \bar{y} = \bar{h},$$

where $\Delta \bar{y} = L^T \Delta y$ and $\bar{h} = L^{-1}h$. The new coefficient matrix generally has a better eigenvalue distribution.

This combination of the Sherman-Morrison formula and preconditioned conjugate gradient method was proposed in Choi, Monma and Shanno [2]. Throughout the computation, we only factor the sparse portion of the matrix ADA^T . Computational results have shown that the above dense column strategy performs adequately for the Netlib set of test problems (see Section 6 for a description of the Netlib set). The implementations of the Sherman-Morrison formula and the preconditioned conjugate gradient method are, respectively, sherman.m and pcg.m.

5 Numerical Stability in Cholesky Factorization

It is well known that in a primal-dual interior-point method the Jacobian matrices at any primal degenerate solution is necessarily singular. In Procedure Steps, this singularity is reflected in the matrix $P = ADA^T$ where D is a diagonal matrix varying from iteration to iteration. It is known that for linear programming all solutions in the relative interior of a non-singleton solution set satisfy the strict complementarity

(14)
$$xz = 0, x + z > 0; sw = 0, s + w > 0.$$

This implies that the diagonal elements of D either approach zero or infinity, as can be seen from the construction of D. Whenever D has too many small diagonal elements, near-singularity could occur and the standard Cholesky factorization may become unstable in that very small diagonal pivots are encountered. For degenerate problems, this near-singularity can cause ill-conditioning so severe that it often prevents the standard Cholesky factorization method from getting an acceptable approximate solution.

In this section, we discuss in detail a technique for overcoming the numerical instability encountered in Cholesky factorization. Similar techniques have been used in earlier implementations such as in OB1 [9]. However, we have not seen any discussion in the literature on the theoretical foundation and justification for these techniques. In the sequel, we will motivate and construct a mathematically well-founded procedure for treating numerical instability in Cholesky factorization in the context of interior-point algorithms.

Let us define for a strict complementary solution $v^* = (x^*, z^*, s^*, w^*, y^*)$ the index sets

(15)
$$\mathcal{B} = \{i : x_i^* > 0 \text{ and } x_i^* < u_i \text{ for } i \le n_u\},\$$

(16)
$$\mathcal{N} = \{i : x_i^* = 0 \text{ or } x_i^* = u_i \text{ for } i \le n_u\}.$$

It is well-known that these index sets are invariant with respect to the choices of strict complementary solution v^* . Without loss of generality, let us assume that for some positive integer $p \leq n$

(17)
$$\mathcal{B} = \{1, 2, \cdots, p\}, \quad \mathcal{N} = \{p+1, p+2, \cdots, n\}.$$

Now consider the linear system in step (3) of the Procedure Steps

$$ADA^{T}\Delta y = -(r_b + ADr_c),$$

where $r_b = Ax - b$. As the iterates approach the interior of the solution set, from the definition

of D we see that

(19)
$$D_{ii} \to \begin{cases} +\infty, & i \in \mathcal{B}, \\ 0, & i \in \mathcal{N}. \end{cases}$$

We introduce the following partitioning in terms of the index sets \mathcal{B} and \mathcal{N} specified in (15), (16) and (17) for the matrices D, A and r_c :

$$D = egin{bmatrix} D_B & 0 \ 0 & D_N \end{bmatrix}, \quad A = [A_B \ A_N], \quad r_c = egin{pmatrix} (r_c)_B \ (r_c)_N \end{pmatrix}.$$

Using this partitioning, we rewrite the linear system (18) as

(20)
$$(A_B D_B A_B^T + A_N D_N A_N^T) \Delta y = -(r_b + A_B D_B(r_c)_B + A_N D_N(r_c)_N).$$

As the iterates converge to the solution set, we know that $D_B \to \infty$, $D_N \to 0$ and $r_b \to 0$. Moreover, the overwritten r_c (see (2) in Procedure Steps) is bounded under normal conditions. Hence we have the following asymptotic behavior for terms in (20):

$$A_B D_B A_B^T \to \infty,$$

 $A_N D_N A_N^T \to 0,$
 $A_B D_B (r_c)_B \to \infty,$
 $A_N D_N (r_c)_N \to 0.$

As a result, in a small neighborhood of the solution set linear systems of the form (20) are all small perturbations to the the following consistent linear system

$$(21) (A_B D_B A_B^T) \Delta y = -A_B D_B(r_c)_B.$$

If the rank of A_B is less than m, such as in the case of degeneracy where p < m, then $A_B D_B A_B^T$ is singular. Consequently, as the iterates progresses towards the solution set, we encounter increasingly ill-conditioned or even numerically singular systems. In this case, the standard solution procedure based on Cholesky factorization will break down and fail to return an acceptable approximate solution. Unfortunately, in practical applications degeneracy occurs surprisingly often. Therefore, in order to have a robust and accurate interior-point code one must be able to solve to an acceptable accuracy the linear system (21) as well as arbitrarily small perturbations to it.

Let us now consider solving (21). Without loss of generality, we assume that (i) A_B has full rank p but p < m, and (ii) the first p rows of A_B is linearly independent. The second assumption implies that the first p rows, and columns by symmetry, of $A_B D_B A_B^T$ are linearly independent.

However, we would like to avoid to be directly involved in the partitioning of \mathcal{B} and \mathcal{N} and prefer to have a single algorithm that can stably solve (18) in all situations, be it far away from or close to the solution set. So later when we construct the algorithm, we will assume no knowledge on the specific structure of the coefficient matrix P except that it is symmetric positive semidefinite but not necessarily definite.

Now suppose that we are given the coefficient matrix P and right-hand side h, where

$$P = A_B D_B A_B^T$$
, $h = -A_B D_B (r_c)_B$,

and want to solve

$$(22) P\Delta y = h.$$

This linear system has multiple solutions. Fortunately, it is known that the convergence theory for primal-dual interior-point algorithms only require a solution to (18), so any solution will suffice.

We make a further partitioning by grouping the first p rows and the last (m-p) rows of A, y and h:

$$A_B = \begin{bmatrix} A_B^1 \\ A_B^2 \end{bmatrix}, \quad \Delta y = \begin{pmatrix} \Delta y_1 \\ \Delta y_2 \end{pmatrix}, \quad h = \begin{pmatrix} h_1 \\ h_2 \end{pmatrix}.$$

By our assumption, A_B^1 is a $p \times p$ nonsingular matrix. From the transpose of QR decomposition, there exists a $p \times p$ orthogonal matrix Q such that

$$A_B D_B^{1/2} = \begin{bmatrix} L \\ M \end{bmatrix} Q$$

where L is $p \times p$, lower triangular and nonsingular. It is not difficult to see that

(23)
$$P = A_B D_B A_B^T = \begin{bmatrix} L \\ M \end{bmatrix} \begin{bmatrix} L^T & M^T \end{bmatrix} = \begin{bmatrix} L & 0 \\ M & 0 \end{bmatrix} \begin{bmatrix} L^T & M^T \\ 0 & 0 \end{bmatrix}$$

and the right-hand side is nothing but the Cholesky factorization of P. Moreover, all the solutions of (22) are given by the first p linearly independent equations

$$(LL^T)\Delta y_1 + (LM^T)\Delta y_2 = h_1.$$

In particular, we are interested in the so-called basic solution

(24)
$$\Delta y^* = \begin{pmatrix} L^{-T}L^{-1}h_1\\0 \end{pmatrix},$$

corresponding to $\Delta y_2 = 0$.

For convenience of discussion, in the sequel we will use the extended arithmetic; namely, we will include ∞ (only positive infinity will suffice for our purpose) as a real number such that $1/\infty = 0$. Let us define D_{∞} as the $(m-p) \times (m-p)$ diagonal matrix with infinity on the diagonal and as a result $D_{\infty}^{-1} = 0$.

Replacing the zero diagonal blocks by D_{∞} in the Cholesky factors of P in (23), we obtain the following matrix product associated with P,

(25)
$$P \Rightarrow \begin{bmatrix} L & 0 \\ M & D_{\infty} \end{bmatrix} \begin{bmatrix} L^T & M^T \\ 0 & D_{\infty} \end{bmatrix}.$$

We will call (25) the Cholesky-Infinity factorization of the positive semidefinite matrix P.

We now show that the basic solution Δy^* given in (24) is the uniquely solution to the extended system associated with the Cholesky-Infinity factorization of P:

(26)
$$\begin{bmatrix} L & 0 \\ M & D_{\infty} \end{bmatrix} \begin{bmatrix} L^T & M^T \\ 0 & D_{\infty} \end{bmatrix} \begin{pmatrix} \Delta y_1 \\ \Delta y_2 \end{pmatrix} = \begin{pmatrix} h_1 \\ h_2 \end{pmatrix}.$$

This system can be solved by the forward and backward substitution procedure

$$\begin{bmatrix} L & 0 \\ M & D_{\infty} \end{bmatrix} \begin{pmatrix} t_1 \\ t_2 \end{pmatrix} = \begin{pmatrix} h_1 \\ h_2 \end{pmatrix} \quad \text{and} \quad \begin{bmatrix} L^T & M^T \\ 0 & D_{\infty} \end{bmatrix} \begin{pmatrix} \Delta y_1 \\ \Delta y_2 \end{pmatrix} = \begin{pmatrix} t_1 \\ t_2 \end{pmatrix},$$

which leads to

$$t_1 = L^{-1}h_1,$$

$$t_2 = D_{\infty}^{-1}(h_2 - Mt_1) = 0,$$

$$\Delta y_2 = D_{\infty}^{-1}t_2 = 0,$$

$$\Delta y_1 = L^{-T}(t_1 - M^T \Delta y_2) = L^{-T}L^{-1}h_1,$$

namely, $\Delta y = \Delta y^*$. Therefore, solving (22) for the basic solution reduces to finding the Cholesky-Infinity factorization of P. We provide a simple modification to the Cholesky factorization procedure to obtain an algorithm that will return the Cholesky factorization for a positive definite matrix, but the Cholesky-Infinity factorization for a positive semidefinite, but not definite, matrix.

A (column version) Cholesky factorization procedure for $P = LL^T$, where $P = [p_{ij}]$ is symmetric and $L = [\ell_{ij}]$ is lower triangular, can be written as follows.

```
Procedure Cholesky:

For j = 1: n
d = p_{jj} - \sum_{k=1}^{j-1} \ell_{jk}^2 \text{ (diagonal pivot)}.
If d <= 0, then exit (not positive-definite).
\ell_{jj} = \sqrt{d}.
For i = (j+1): n
\ell_{ij} = \left(p_{ij} - \sum_{k=1}^{j-1} \ell_{ik} \ell_{jk}\right) / \ell_{jj}.
End
End
```

The algorithm stops whenever a diagonal pivot is non-positive, indicating that P is not positive definite.

We give a simple, two-line modification to the Cholesky factorization procedure. For any positive definite matrix P, the procedure still returns its Cholesky factor L. Moreover, for any positive semidefinite, but not definite, matrix P, it returns its Cholesky-Infinity factorization.

```
Procedure Cholesky-Infinity:

For j = 1: n
d = p_{jj} - \sum_{k=1}^{j-1} \ell_{jk}^2 \text{ (diagonal pivot)}.
If d < 0, then exit (not positive-semidefinite).

If d = 0, then set d = \infty.
\ell_{jj} = \sqrt{d}.
For i = (j+1): n
\ell_{ij} = \left(p_{ij} - \sum_{k=1}^{j-1} \ell_{ik} \ell_{jk}\right) / \ell_{jj}.
End
End
```

In this procedure, once a diagonal pivot is found to be zero, the corresponding diagonal element of L is set to infinity. Consequently, the rest of the elements in that column will be zero due to division by infinity. Under the assumption that the rank of P is p and the first p rows (columns) of P are linearly independent, the procedure will return the Cholesky-Infinity factor of P in (25). In general, the columns containing ∞ and zeros may be scattered around, instead of clustered at the end.

In an actual implementation of Procedure Cholesky-Infinity, we replace 0 by a tiny positive number and ∞ by a huge positive number. More specifically, we make the following change to

the procedure for some tiny $d_{tiny} > 0$ and huge $d_{huge} > 0$:

If
$$d = 0$$
, then set $d = \infty$. \Longrightarrow If $d < d_{tiny}$, then set $d = d_{huge}$.

This modification enables the procedure to solve not only (21) but also its small perturbations in a stable manner.

Extensive computational experiments have proved that the above strategy works surprisingly well in general. In practice, it effectively circumvents numerical difficulty encountered in the standard Cholesky factorization procedure and can attain an accuracy up to the square root of the machine epsilon and beyond.

6 Numerical Results

In this section, we describe our numerical experiments and present computational results. Our numerical experiments were performed on a Silicon Graphics Challenge XL computer at the University of Maryland Baltimore County. It is a shared-memory, symmetric multi-processor machine with 20 MIPS R4400/150MHz CPU's, 1 gigabytes of memory, and 16k data cache per processor (plus 16k instruction cache and 1 Mbyte secondary cache). Since we always used only a single processor at a time, our numerical results can be reproduced on a single processor workstation with the same CPU and ample memory (say, 128 Mbytes).

The test problems used in our experiments were the Netlib set of linear programs. Netlib set is an electronically distributed collection of linear programs originated and maintained by David Gay [3] at the Bell Laboratory. As of January of 1996, the collection consists of 95 problems, including a number of fairly large problems from real-world applications. The sizes of the problems range from a few dozen variables to over fifteen thousands. Over the years, the Netlib set has become the standard set of test problems for testing and comparing algorithms and software for linear programming.

6.1 Test Results for LIPSOL

We tested LIPSOL v0.3 on all the 95 linear programming problems in the Netlib test set. We tabulate the test results in Tables 2-4 in the Appendix. In each table, the seven columns give, respectively, the problem name, the number of constraints in the unprocessed model, the number of variables in the unprocessed model, the number of iterations required for convergence, the total relative residual as given by (13), the primal objective value and the CPU seconds.

6.2 Fortran Package OB1

OB1 (Optimization with Barrier 1) was developed by Irv Lustig, Roy Marsten and David Shanno. Their work [9], largely based on the OB1 package, won the Orchard-Hays Prize from the Mathematical Programming Society in 1991 as the best computational research of the year. Until its retirement in 1993, OB1 was widely recognized as the best interior-point code for linear programming. The version of OB1 available to us and used in our experiments was version 5.0 released in 1991. We understand that the final version of OB1 was version 6.0, which contained enhancements and improvements to version 5.0.

Our purpose is to compare qualitatively the performance of LIPSOL v0.3 with an efficient interior-point code written in Fortran or C. In addition to its availability to us, OB1 version 5.0 is well suited for this purpose of comparison since, being the best interior-point code during the 1991-1992 period, it is still an adequately efficient code even by today's standard.

We compiled OB1 version 5.0 under SGI IRIX 5.3 F77 compiler using the flags -O -mips2. These same flags were also used to compile the MEX-files in LIPSOL. We used all the default options to run OB1 version 5.0 except specifying the cache size as 16k.

6.3 Comparison with OB1

We also solved all 95 Netlib problems using OB1 version 5.0 on the same SGI computer. Since the purpose of the comparison was to qualitative rather than quantitative to save space we will not list all the computational results obtained for OB1. Instead, we will summarize the comparison by a graph in Figure 1.

Our general observation is that for most problems, in particular for the smaller problems, LIPSOL v0.3 is slower than OB1 5.0, but for the a few largest problems LIPSOL v0.3 performs as well as, or even slightly better than, OB1 5.0. To capture the trend, we rearranged the problems in an ascending order according to the average time per iteration required by LIPSOL and plotted the total solution times required by both LIPSOL v0.3 and OB1 5.0 in Figure 1, where the left-most on the horizontal axis is the least expensive problem and the right-most the most expensive one. The plot is in logarithmic scales the the unit is in CPU second. To filter out random fluctuation on the small problems, we took average of ten (for the smallest problems) or five (for others) problems in groups. Since there are fewer very large problems, no average was taken for the four most expensive problems.

From Figure 1, we can see that for less expensive problems, LIPSOL v0.3 was clearly slower than OB1 5.0. However, as the problems become more and more expensive, the difference between the two shrinks gradually. For the a few most expensive (not necessarily the largest) problems,

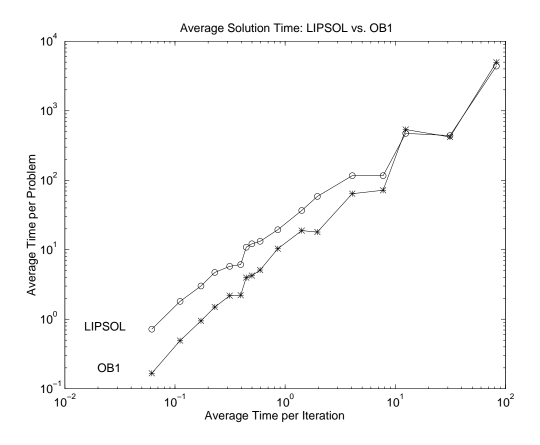


Figure 1: Solution Time: LIPSOL v0.3 vs. OB1 5.0

the amount of solution time required by LIPSOL v0.3 is similar to or even less than that required by OB1 5.0, as is shown in Table 1.

Table 1: LIPSOL v0.3 and OB1 5.0 on Large problems

Problem	Rows	Cols	LIP-Iter	OB1-Iter	LIP-Time	OB1-Time
dfl001	6072	12230	53	48	4384.76	4990.18
Maros-r7	3137	9408	14	13	439.51	418.47
pilot87	2031	4883	38	41	472.62	537.54
stocfor3	16676	15695	34	33	176.49	79.76

As one can see, the largest problem in terms of sizes is Stocfor3. However, this problem does not require a long time to solve because it is very sparse.

Our interpretation of the comparative results is as follows. The overhead induced by MAT-LAB consumes a significant portion of the total computation time when the problem size is small, which is the reason why LIPSOL is generally slower on small problems under the MATLAB environment. As the problem size increase, the time spent on Cholesky factorization and back-solves, the most computation-intensive tasks at each iteration, grows quickly to become the dominant portion of the total time, while the MATLAB overhead assuming a lesser role. Our results appear to indicate that for larger problems the ORNL sparse Cholesky code used in LIPSOL is faster than its counterpart in OB1 version 5.0, which was written earlier and did not seem to include some of the newer techniques.

6.4 Asymptotic Convergence Rates

We have observed cubic-like convergence in many of the tested problems. In Figure 2, we give edited printout from LIPSOL on the last several iterations for three Netlib problems: adlittle, beaconfd and stocfor3. We call the reader's attention to the column named Gap, representing the duality gap x^Tz+s^Tw . Since the equations xz=0 and sw=0 are the only nonlinear portion of the KKT system while the primal, dual and upper-bound feasibility conditions are all linear, the convergence rate of the duality gap usually dictates the asymptotic convergence rate of the whole residual vector. The last column is the total relative error and has no apparent correlation to the convergence rate of the residual vector.

Residuals:	Primal	Dual	U-bounds	Gap	TR-error
Iter 10:	9.22e-09	2.33e-11	0.00e+00	4.35e+00	1.93e-05
Iter 11:	4.53e-11	9.31e-11	0.00e+00	5.91e-03	2.61e-08
Iter 12:	2.11e-11	9.31e-11	0.00e+00	6.44e-10	3.12e-14
Name:	ADLITTLE				
Residuals:	Primal	Dual	U-bounds	Gap	TR-error
Iter 11:	1.34e-06	8.06e-08	0.00e+00	6.78e-01	1.97e-05
Iter 12:	4.45e-09	4.80e-11	0.00e+00	1.34e-03	3.84e-08
Iter 13:	1.57e-10	1.30e-11	0.00e+00	1.55e-10	7.69e-14
Name:	BEACONFD				
Residuals:	Primal	Dual	U-bounds	Gap	TR-error
Iter 32:	1.27e-05	3.23e-08	0.00e+00	4.09e-01	1.02e-05
Iter 33:	2.32e-07	1.26e-10	0.00e+00	2.45e-03	6.13e-08
Iter 34:	6.33e-08	2.97e-13	0.00e+00	2.09e-08	9.59e-10
Name:	STOCHFOR				

Figure 2: Examples of Cubic-like Convergence

As can be seen from these examples, that cubic-like convergence not only happens for smaller problems such as addittle and beaconfd, but also to large problems such as stocfor3, which

is the largest in the 95 Netlib problems in terms of problem sizes.

However, on other problems cubic convergence is not observed up to the time when the algorithm terminates. In Figure 3, we present printout from LIPSOL for three problems: stair, pilot87 and ship08s.

Residuals:	Primal	Dual	U-bounds	Gap	TR-error
Iter 12:	1.34e-04	5.24e-07	0.00e+00	4.29e-01	1.69e-03
Iter 13:	5.93e-05	6.76e-08	0.00e+00	6.38e-02	2.51e-04
Iter 14:	1.64e-06	1.78e-10	8.88e-16	7.64e-04	3.02e-06
Iter 15:	1.09e-05	4.42e-15	1.26e-15	1.36e-08	4.47e-08
Iter 16:	1.59e-10	2.98e-15	1.26e-15	1.36e-15	6.55e-13
Name:	STAIR				
Residuals:	Primal	Dual	U-bounds	Gap	TR-error
Iter 32:	1.86e-07	3.16e-11	5.07e-08	2.05e-02	6.75e-05
Iter 32:	5.26e-08	9.49e-12	1.43e-08	5.86e-03	1.93e-05
Iter 34:	1.94e-08	3.79e-12	5.28e-09	2.04e-03	6.69e-06
Iter 35:	3.72e-09	3.07e-12	1.00e-09	4.29e-04	1.41e-06
Iter 36:	9.29e-10	3.81e-12	2.03e-10	1.32e-04	4.35e-07
Iter 37:	9.27e-10	4.82e-12	4.59e-11	2.07e-05	6.79e-08
Iter 38:	2.21e-09	3.01e-12	1.65e-12	1.82e-06	5.99e-09
Name:	PILOT87				
				_	
Residuals:	Primal	Dual	U-bounds	Gap	TR-error
Iter 12:	7.31e-05	8.61e-11	0.00e+00	9.25e+01	4.72e-05
Iter 13:	2.12e-07	9.17e-11	0.00e+00	3.05e-01	1.56e-07
Iter 14:	1.56e-11	9.85e-11	0.00e+00	3.88e-07	2.13e-13
Name:	SHIP08S				

Figure 3: Examples of Quadratic, Linear, and "Big bang" Convergence

The convergence rate appears to be quadratic for stair and linear for pilot87, which is a well-known, ill-conditioned and difficult problem. On the other hand, for ship08s, the convergence rate appears, surprisingly but pleasantly, to be even faster than cubic. It looks like that after a "big bang" iteration, the last iterate just drops into the optimal facet so the algorithm terminates right away.

7 Concluding Remarks

In this paper, we provided intuitive motivations to some infeasible-primal-dual interior-point methods as perturbed and damped Newton's method. We described major steps in implementing an interior-point code for large-scale linear programming. We also discussed in detail a technique for stabilizing Cholesky factorization in the context of interior-point algorithms with mathematical justification.

Our implementation was carried out under the MATLAB environment. Yet it is still suffi-

ciently fast for large-scale problems and comparable in performance with a good Fortran code. The software package resulted from our implementation is called LIPSOL. LIPSOL is free software and available from the internet sites:

```
http://pc5.math.umbc.edu/~yzhang/lipsol/ftp://ftp.math.umbc.edu/pub/zhang/lipsol/v0.3/
```

where the first is the LIPSOL Web page and the second an anonymous FTP site. In addition to providing an economic means for solving relatively large-scale linear programs, LIPSOL can be a convenient tool for both research and teaching. For more details on LISPOL, we refer readers to the User's Guide for LIPSOL [20] available in the aforementioned sites for LIPSOL distribution.

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Appendix: Test Results for LIPSOL v0.3

Table 2: Test Results for LIPSOL v0.3 (I)

Problem	Rows	Cols	Iter	Residual	Primal Objective	CPU sec.
25fv47	822	1571	25	2.60e-13	5.5018458883e+03	15.27
80bau3b	2263	9799	37	6.39e-09	9.8722419330e+05	76.11
adlittle	57	97	12	3.12e-14	2.2549496316e+05	0.71
afiro	28	32	7	3.68e-09	-4.6475314219e+02	0.35
agg	489	163	18	1.01e-09	-3.5991767273e+07	5.85
agg2	517	302	16	2.57e-10	-2.0239252353e+07	6.84
agg3	517	302	16	6.69e-12	1.0312115935e+07	6.91
bandm	306	472	17	8.58e-11	-1.5862801844e+02	2.86
beaconfd	174	262	13	7.69e-14	3.3592485807e+04	2.41
blend	75	83	12	8.25e-13	-3.0812149846e+01	0.80
bnl1	644	1175	26	8.90e-09	1.9776295714e + 03	7.68
bnl2	2325	3489	33	3.84e-13	1.8112365404e+03	39.87
boeing1	351	384	21	3.04e-09	-3.3521356672e+02	6.21
boeing2	167	143	18	1.67e-11	-3.1501872801e+02	2.52
bore3d	234	315	18	8.08e-09	1.3730803987e+03	2.58
brandy	221	249	16	4.04e-09	1.5185098958e+03	2.34
capri	272	353	20	5.39e-11	2.6900129139e+03	3.81
cycle	1904	2857	27	2.98e-10	-5.2263930249e+00	40.16
czprob	930	3523	31	5.96e-12	2.1851966989e+06	14.80
d2q06c	2172	5167	31	8.50e-12	1.2278421081e+05	86.28
d6cube	416	6184	25	1.70e-11	3.1549166666e + 02	38.54
degen2	445	534	14	1.29e - 11	-1.4351780000e+03	4.61
degen3	1504	1818	20	2.65e-10	-9.8729399974e+02	60.33
dfl001	6072	12230	53	2.90e-09	1.1266396043e+07	4384.76
e226	224	282	21	1.17e-09	-1.8751929046e+01	3.20
etamacro	401	688	28	4.23e-09	-7.5571523297e+02	6.13
fffff800	525	854	25	5.62e-10	5.5567956497e + 05	10.97
finnis	498	614	27	2.33e-09	1.7279106571e+05	5.88
fit1d	25	1026	18	6.94e - 10	-9.1463780919e+03	8.34
fit1p	628	1677	15	3.38e-10	9.1463780925e+03	53.10
fit2d	26	10500	22	1.07e-10	-6.8464293294e+04	106.72
${ m fit}2{ m p}$	3001	13525	20	3.17e-10	6.8464293295e+04	197.93

Table 3: Test Results for LIPSOL v0.3 (II)

Problem	Rows	Cols	Iter	Residual	Primal Objective	CPU sec.
forplan	162	421	20	6.17e-10	-6.6421896090e+02	4.29
ganges	1310	1681	18	1.09e - 10	-1.0958573613e+05	9.62
giffpin	617	1092	20	1.46e-11	6.9022359996e+06	4.32
greenbea	2393	5405	44	3.24e-10	-7.2555248106e+07	83.28
greenbeb	2393	5405	37	5.02 e-09	-4.3022602396e+06	68.05
grow15	301	645	17	1.02e-09	-1.0687094129e+08	6.22
grow22	441	946	19	1.03e-09	-1.6083433648e+08	9.64
grow7	141	301	16	4.88e - 10	-4.7787811815e+07	3.06
israel	175	142	22	8.64e-13	-8.9664482186e+05	6.83
kb2	44	41	15	3.44e-09	-1.7499001299e+03	1.07
lotfi	154	308	18	9.83e-10	-2.5264706056e+01	1.65
maros	847	1443	32	8.10e-13	-5.8063743701e+04	17.58
maros-r7	3137	9408	14	1.02e-09	1.4971851676e+06	439.51
modszk1	688	1620	24	2.71e-12	3.2061972906e+02	6.95
nesm	663	2923	30	4.17e-09	1.4076036546e+07	25.64
perold	626	1376	33	5.18e-12	-9.3807552782e+03	15.10
pilot	1442	3652	34	8.63e-09	-5.5748972915e+02	152.53
pilot4	411	1000	28	4.69e-09	-2.5811392559e+03	12.66
pilot87	2031	4883	38	5.99e-09	3.0171034740e+02	472.62
pilotja	941	1988	30	1.20e-09	-6.1131364638e+03	29.16
pilotnov	723	2172	20	1.60e-13	-4.4972761882e+03	18.15
pilotwe	976	2789	36	1.54e-09	-2.7201075328e+06	18.65
recipe	92	180	9	8.63e-11	-2.6661600000e+02	0.91
sc105	106	103	10	1.46e - 13	-5.2202061212e+01	0.61
sc205	206	203	11	6.38e-09	-5.2202061119e+01	0.87
sc50a	51	48	10	2.52e-14	-6.4575077059e+01	0.49
sc50b	51	48	7	9.44e-10	-6.9999999976e+01	0.38
scagr25	472	500	16	3.05 e-12	-1.4753433061e+07	2.19
scagr7	130	140	12	5.62 e-11	-2.3313898243e+06	0.81
$\operatorname{scfxm1}$	331	457	18	5.20 e-12	1.8416759028e+04	2.93
$\operatorname{scfxm2}$	661	914	19	4.10e - 11	3.6660261565e+04	5.76
scfxm3	991	1371	20	7.65 e-11	5.4901254551e+04	8.72

Table 4: Test Results for LIPSOL v0.3 (III)

Problem	Rows	Cols	Iter	Residual	Primal Objective	CPU sec.
scorpion	389	358	14	4.16e-11	1.8781248228e+03	1.65
scrs8	491	1169	24	1.12e-11	9.0429695380e+02	4.58
$\operatorname{scsd} 1$	78	760	10	2.94e-11	8.6666666746e+00	1.23
$\operatorname{scsd}6$	148	1350	12	6.43e-11	5.0500000078e+01	2.30
scsd8	398	2750	11	4.00e-11	9.0499999995e+02	4.05
sctap1	301	480	17	5.75e-13	1.4122500000e+03	2.03
$\operatorname{sctap} 2$	1091	1880	18	3.45 e-09	1.7248071473e+03	7.21
$\operatorname{sctap}3$	1481	2480	18	6.61e-14	1.4240000000e+03	9.56
seba	516	1028	20	1.62e-10	1.5711600003e+04	15.26
share1b	118	225	20	1.05e-10	-7.6589318575e+04	1.65
${ m share 2b}$	97	79	12	2.09e-09	-4.1573224024e+02	0.87
shell	537	1775	18	3.07e-09	1.2088253485e+09	4.41
ship04l	403	2118	12	7.48e-11	1.7933245380e+06	3.83
m ship 04s	403	1458	13	1.27e-09	1.7987146998e+06	2.96
ship08l	779	4283	15	2.25e-09	1.9090552133e+06	9.18
ship08s	779	2387	14	2.13e-13	1.9200982105e+06	4.96
ship12l	1152	5427	15	8.58e-09	1.4701879275e+06	11.63
ship12s	1152	2763	15	6.73e-09	1.4892361411e+06	6.21
sierra	1228	2036	16	1.31e-12	1.5394362184e+07	12.55
stair	357	467	16	6.55e-13	-2.5126695119e+02	5.06
standata	360	1075	16	3.60e-11	1.2576995000e+03	3.49
$\operatorname{standgub}$	362	1184	16	7.83e-10	1.2576995007e+03	3.61
standmps	468	1075	24	3.98e-13	1.4060175000e+03	5.81
stocfor1	118	111	17	1.38e-11	-4.1131976219e+04	1.10
stocfor2	2158	2031	22	3.24e-10	-3.9024408538e+04	14.44
stocfor3old	16676	15695	34	9.59e-10	-3.9976783944e+04	176.49
truss	1001	8806	18	3.73e-10	4.5881584721e+05	22.85
tuff	334	587	23	2.39e-11	2.9214776512e-01	6.24
vtpbase	199	203	28	4.36e-13	1.2983146246e+05	3.37
wood1p	245	2594	20	1.12e-11	1.4429024116e+00	43.97
woodw	1099	8405	27	3.98e-11	1.3044763331e+00	42.31