# Iterative Solution of Sparse Linear Least Squares using LU Factorization

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### Abstract

In this paper, we are interested in computing the solution of an overdetermined sparse linear least squares problem  $Ax \simeq b$  via the normal equations method. Transforming the normal equations using the L factor from a rectangular LU decomposition of A usually leads to a better conditioned problem. Here we explore a further preconditioning by  $L_1^{-1}$  where  $L_1$  is the  $n \times n$  upper part of the lower trapezoidal  $m \times n$  factor L. Since the condition number of the iteration matrix can be easily bounded, we can determine whether the iteration will be effective, and whether further preconditioning is required. Numerical experiments are performed with the Julia programming language. When the upper triangular matrix U has no near zero diagonal elements, the algorithm is observed to be reliable. When A has only a few more rows than columns, convergence requires relatively few iterations and the algorithm usually requires less storage than the Cholesky factor of  $A^TA$  or the R factor of the QR factorization of A.

**Keywords**: Sparse linear least squares, iterative methods, preconditioning, conjugate gradient algorithm, lsqr algorithm, LU factorization.

### 1 Introduction

Linear least squares (LLS) problems arise in many high-performance computing (HPC) applications when the number of linear equations is not equal to the number of unknown parameters. We consider the overdetermined full rank LLS problem

$$\min_{x \in \mathbb{R}^n} \|Ax - b\|_2,\tag{1}$$

with  $A \in \mathbb{R}^{m \times n}$ ,  $m \ge n$  and  $b \in \mathbb{R}^m$ .

LLS solvers can be based on either direct methods (e.g., Cholesky factorization of the normal equations, sparse  $LDL^T$  factorization of the augmented system, or QR factorization of A), or iterative methods (e.g., preconditioned

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Krylov subspace methods). In this paper, we focus on the iterative conjugate gradient (CG) solution of the normal equations

$$A^T A x = A^T b. (2)$$

For the CG method, the linear rate of convergence is bounded by the quantity

$$\frac{\kappa-1}{\kappa+1}$$
,

where  $\kappa = cond_2(A) = \sqrt{cond_2(A^TA)}$  is the 2-norm condition number of A (see [3, p. 289] for more details). Then having a better conditioning of the normal equations matrix will influence positively the convergence of the solution of Equation (2) via the CG algorithm.

As with linear systems, a variety of preconditioners have been proposed to overcome the potential ill-conditioning of the normal equations (see [12] for a comprehensive overview of these preconditioners). One possibility is to precondition the normal equations by an incomplete Cholesky decomposition of  $A^T A$  (see, e.g., the RIF preconditioner [2] or its new left-looking variant in [23]).

There are also approaches based on LU factorization preconditioning [1, 4, 5, 21]. LU preconditioning has been also studied in [14] with the objective of exploiting the recent progress in sparse LU factorization. For example, both Matlab and Octave use fast sparse LU factorizations built on the UMFPACK package [6]. Other direct sparse solvers (e.g., [16, 17, 19, 22]) offer scalable sparse LU factorizations for large problems and are commonly used in HPC applications.

For the overdetermined case m > n, we can perform a rectangular LU factorization PAQ = LU of the  $m \times n$  matrix A, where L is a lower trapezoidal  $m \times n$  matrix, U is an upper triangular  $n \times n$  matrix, P is a permutation matrix obtained with threshold pivoting (swap of rows so that the lower trapezoidal L has 1's on the diagonal) and Q is a permutation matrix obtained from column pivoting to avoid fill-in in the L and U factors. As L tends to be better conditioned than A [24, p. 231], it was explored in [14] transforming Equation (2) to

$$L^T L y = L^T c, (3)$$

with  $y = UQ^Tx$ , and c = Pb, when U is nonsingular. As illustrated in [14], L is usually better conditioned than A and fewer iterations are required for the L iteration.

Regarding the storage issues, we plot in Figure 1 the storage required for an LU factorization of A (with partial pivoting) and for the Cholesky factorization of  $A^TA$  for matrices from the Davis collection (modified such that most of these matrices had ten percent more rows than columns). We observe that the storage of L and U for a nearly square rectangular matrix is usually less than that of the Cholesky factor of  $A^TA$  (or equivalently, in exact arithmetic, the R factor from the QR factorization of A). In fact, if every set of k columns of k

<sup>1</sup>http://www.cise.ufl.edu/research/sparse/matrices/

has nonzero entries in at least k+1 rows, for all  $1 \le k \le n-1$  (i.e. A is "strong Hall" [6, pp. 83-84]), the storage required by U is always bounded by that of R (see [8, 9, 10]). Moreover, if the "strong Hall" property holds, the storage required for L is bounded by that needed for an orthogonal Householder derived Q, QR = AP [9, 10].

## 

Figure 1: Storage for LU factorization of A vs Cholesky factorization of  $A^TA$ .

In this paper, we consider a partition of the lower trapezoidal matrix L as

$$L = \left[\frac{L_1}{L_2}\right]$$

where  $L_1$  is  $n \times n$  lower triangular and  $L_2$  is  $(m-n) \times n$ . We explore iterating with the normal equations in terms of

$$F = LL_1^{-1} = \left[\frac{I_n}{L_2L_1^{-1}}\right].$$

Note that a partition of  $A = \left[\frac{A_1}{A_2}\right]$  (instead of L here) is considered in [5] where an LU factorization of  $A_1$  is used to precondition the normal equations. In [1], a partition  $A = \left[\frac{A_1}{A_2}\right]$  is also used (where  $A_1$  is a set of basis obtained with an LU factorization), and the augmented system is transformed into an equivalent symmetric quasi-definite system.

We will show that iterating on  $LL_1^{-1}$  results in faster convergence than with L, due to a better conditioning of  $LL_1^{-1}$  and does not require additional storage.

Moreover, since a bound on the condition number of  $LL_1^{-1}$  can be explicitly computed, this can indicate if further preconditioning is required.

The plan of this paper is as follows. Section 2 motivates the use of  $LL_1^{-1}$  preconditioning. Section 3 describes the experimental framework including the test matrices and some implementation issues. Section 4 presents numerical results on the test matrices. Section 5 concludes the paper.

### 2 $LL_1^{-1}$ preconditioning

In this section we motivate our choice to precondition the conjugate gradient solution of the normal equations by using the matrix  $F = LL_1^{-1}$  obtained from the partition of the rectangular matrix L as

$$L = \left\lceil \frac{L_1}{L_2} \right\rceil$$

where  $L_1$  is square and lower triangular and  $L_2$  is  $(m-n) \times n$  so that

$$LL_1^{-1} = \left[\frac{I_n}{L_2L_1^{-1}}\right] = \left[\frac{I_n}{C}\right]$$

with  $C = L_2 L_1^{-1}$ . Then Equation (3) becomes

$$[I_n|C^T]$$
  $\left[\frac{I_n}{C}\right]z = [I_n|C^T]c,$ 

or equivalently,

$$F^T F z = F^T c, (4)$$

with  $z = L_1 U Q^T x$ , and c = Pb. Since  $L_1$  and U are respectively lower and upper triangular, x is computed from z using forward and backward substitutions. For matrices with only a few more rows than columns, explicit computation of  $C = L_2 L_1^{-1}$  may be worthwhile, being easily performed in parallel, and allowing easy parallelization of the iteration. If we do not explicitly compute C, there is no additional storage beyond that required for L.

In this paper, we solve Equation (4) using the 1sqr algorithm [20], with the iteration matrix  $F = LL_1^{-1}$ . The iteration requires multiplying both by F and  $F^T$ . We compute

$$v = Fu = \left[\frac{I_n}{L_2 L_1^{-1}}\right] u = \left[\frac{u}{L_2 L_1^{-1} u}\right]$$

entailing a forward triangular solve with  $L_1$ . Similarly, we compute

$$F^T v = [I_n | L_1^{-T} L_2^T] v = v + L_1^{-T} (L_2^T v),$$

which requires a triangular backsolve. The multiplications by F and  $F^T$  take almost the same number of flops as multiplications by L, but the backward and

forward solves are likely to be harder to parallelize than multiplication by L. In the remainder of this paper, the lsqr algorithm where we iterate with

$$F = LL_1^{-1} = \left\lceil \frac{I}{C} \right\rceil$$

instead of A is referred to as lsqrLinvL and is given in Algorithm 1. The adapted lsqr algorithm which is called in lsqrLinvL is detailed in Algorithm 2.

### Algorithm 1 lsqrLinvL

```
function [x, its] = lsqrLinvL(A, b, tol, maxit)
% A is an input matrix of m rows and n columns
% b is an input m-vector
% tol (convergence tolerance) is a positive input scalar
% maxit is the maximal number of iterations
% output x is an n vector to minimize ||Ax - b||_2
% its is the actual number of iterations performed
[L,U,prow,qcol] \leftarrow lu(A,'vector');
r \leftarrow permute(b,prow);
[x,its] \leftarrow lsqr(L,r,tol,maxit);
if (its \geq maxit)," maxits exceeded", end
x \leftarrow U \setminus x;
x \leftarrow ipermute(qcol,x);
End
```

Figure 2 compares the condition numbers of A, L, and C for some rectangular matrices small enough that we can explicitly compute the SVD in Matlab and thus the condition number as the ratio of largest and smallest singular values. These matrices were created in a similar fashion to those discussed in the next section, with the number of columns at most 5,000. Most had ten per cent more rows than columns.

We have observed that  $F=LL_1^{-1}$  is usually better conditioned than L. Moreover we can easily estimate  $cond_2(F)$  even for large matrices. Consider that

$$F^T F = [I_n | C^T] \left[ \frac{I_n}{C} \right] = I_n + C^T C,$$

then we have

$$\sigma_i^2(F) = \lambda_i(F^T F) = 1 + \lambda_i(C^T C) = 1 + \sigma_i^2(C),$$

where  $\sigma_i(\cdot)$  and  $\lambda_i(\cdot)$  are the singular values and eigenvalues of the considered matrices. If  $\lambda_{max}$ ,  $\lambda_{min}$  denote the largest and smallest eigenvalues of  $C^TC$ , and since  $C^TC$  is real and symmetric, then we have  $0 \leq \lambda_{min} \leq \lambda_{max}$ . The condition number of  $F = LL_1^{-1}$  can be expressed as

$$cond_2(F) = \sqrt{\frac{\lambda_{max} + 1}{\lambda_{min} + 1}}.$$

```
Algorithm 2 1sqr: adapted from Paige and Saunders, 1982
```

```
function [x, its] = lsqr(L, b, tol, maxit)
     % output x is an n vector to minimize ||Ax - b||_2
     \% its is the actual number of iterations performed
     % Initialization
          \beta \leftarrow ||b||_2; \phi_b \leftarrow \beta; b \leftarrow b/\beta;
          x \leftarrow 0_n; \delta \leftarrow 1; its \leftarrow 0;
          [m, n] \leftarrow \operatorname{size}(L);
          L_1 \leftarrow L[1:n,1:n];
          L_2 \leftarrow L[n+1:m,1:n];
          v_1 \leftarrow L_2'u[n+1:m];
          v_1 \leftarrow L_1^7 \backslash u_2;
          v \leftarrow u[1:n] + v_1;
          \nu \leftarrow \|v\|_2;
          v \leftarrow v/\nu;
          w \leftarrow v;
          \phi_a \leftarrow 1;
      % Lanczos iteration
      While ((|\delta| > tol) \text{ and } (its < maxit))
          \% Continue the bidiagonalization
          u_1 \leftarrow L_1 \backslash v;
          u \leftarrow [v; L_2 u_1] - \nu u;
          \beta \leftarrow ||u||_2;
          u \leftarrow u/\beta;
          v_1 \leftarrow L_2' u[n+1:m];
          v_1 \leftarrow L_1' \backslash v_1;
          v_1 \leftarrow u[1:n] + v_1;
          v \leftarrow v_1 - \beta v;
          \nu \leftarrow ||v||_2;
          v \leftarrow v/\nu;
          % Construct and apply next orthog. transformation
          \rho \leftarrow \sqrt{\rho_b^2 + \beta^2}
          c \leftarrow \rho_b/\rho;
          s \leftarrow \beta/\rho;
          \theta \leftarrow s\nu;
          \rho_b \leftarrow -c\nu;
          \phi \leftarrow c\phi_b;
          \phi_b \leftarrow s\phi_b;
          \% update x, w
          \delta \leftarrow \phi/\rho;
          \phi_a = \sqrt{|\phi\delta|};
          x \leftarrow x + \delta w;
          w \leftarrow v - (\theta/\rho)w;
          its \leftarrow its + 1;
     End
     x \leftarrow L_1 \backslash x;
End
                                                               6
```

Then we get

$$cond_2(F) \le \sqrt{\lambda_{max} + 1}.$$

In the case where m - n < n, we have  $\lambda_{min} = 0$ , so we get an equality

$$cond_2(F) = \sqrt{\lambda_{max} + 1}.$$

### Condition number for LSQR iterate

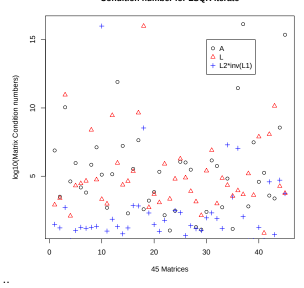


Figure 2: Comparison of the conditioning of various lsqr iteration matrices.

We can estimate  $\lambda_{max}$  with a few iterations of the symmetric power method [11, p. 406] applied to  $C^TC$ . A few iterations suffice for a  $cond_2(F)$  estimate. So having computed L, we can easily decide whether further preconditioning is needed. For the  $LL_1^{-1}$  preconditioning, the above analysis shows that we can estimate  $cond_2(F)$  for the partial pivoting case  $|l_{ij}| \leq 1, i > j$ , and also for the threshold pivoting case where  $|l_{ij}|$  is not bounded by 1. Inexpensive estimation of  $cond_2(F)$  does need  $L_1$  to be lower triangular and nonsingular (so that the symmetric power method can use forward and backward solves on each iteration).

### 3 Experimental framework

In this section, we define the test set for our experiments and the implementation choices based on the Julia programming language [15].

The numerical experiments are carried out with the lsqr algorithm where we iterate with

$$F = LL_1^{-1} = \left[\frac{I}{C}\right],\,$$

using Algorithms 1 and 2. In general, neither C nor  $L_1^{-1}$  are explicitly computed (except in some cases described in Section 4). The LU = PAQ factorization is performed in full precision and with both column and row permutations, using the UMFPACK routine [7]. UMFPACK chooses the Q column permutations to reduce the matrix fill-in and the P row permutations to reduce the Gaussian multipliers size (threshold pivoting). For our numerical experiments, we get L such that the  $l_{ij}$  entries satisfy

$$l_{ii} = 1 \text{ for } i = j, \ l_{ij} = 0 \text{ for } i < j, \ |l_{ij}| < 10 \text{ for } i > j$$

To construct our test problems, we took 235 matrices from the Davis collection, with most of the matrices having less than a maximum of twenty thousand rows or columns. For matrices with more columns than rows, we transposed. Since most matrices were square, we augmented the matrices with one hundred additional rows (or, for square matrices with fewer than 1,000 columns, we increased the number of rows by ten per cent). The additional rows were perturbed copies of randomly selected rows of the original square matrix. The randomly perturbed entries were the original entries times a factor in the range [0.9, 1.1], i.e.

$$a_{ij_{new}} = (1 + 0.1\tau_{ij})a_{ij},$$

where  $\tau_{ij}$  was randomly selected from a uniform distribution on [-1,1]. The minimum number of columns n was 25, the minimum of m + n was 182, the maximum of m + n was 39532. The average number of columns was around 5600. The average number of rows was around 7230.

We performed numerical experiments with the Julia language which is a high-level language for numerical computing and provides good performance (close to that of a C code). Julia codes require fairly minimal changes from Matlab or Octave codes, while being much faster, making Julia very suitable for prototyping HPC algorithms. Wherever there are loops, Julia runs much faster<sup>2</sup>. For example, for sparse matrix multiplications [18], a sparse matrix-vector multiplication Ax in C (gcc in Linux) and Julia (compiled with the same gcc), the C version was less than ten per cent faster, with Octave much slower. Moreover converting code from Octave to Julia is much easier than converting to C or Fortran.

Some of the issues we encountered are as follows: In some instances, we had to be aware that in the statement A=B for Julia arrays, the copy of B is "shallow" i.e., no new copy of A is produced, so changes in A also change B. Though Matlab, Octave and Julia all use sparse LU and QR factorizations based on SuiteSparse [7], the Julia lufact and qrfact functions do not offer user options. The lufact wrapper that we used for UMFPACK is the default in Julia, using a threshold pivoting of 10, and both row and column pivoting. To disable row scaling, we had to find what line to change in the lufact wrapper for UMFPACK, else we would have been solving a weighted least squares problem. Also, using Octave, we could obtain a factorization of the form  $QR = \begin{bmatrix} I_n \\ C_{drop} \\ P \end{bmatrix}$ 

<sup>&</sup>lt;sup>2</sup>See https://julialang.org/benchmarks/ for Julia benchmarks

 $(C_{drop})$  is the matrix C on which we apply a drop tolerance, see Section 4), where we could explicitly obtain the orthogonal matrix Q, upper triangular matrix R, and column permutation P. Alas the qrfact function in Julia currently returns a least squares solution, but does not offer options to return R, Q, or P. In exact arithmetic,  $R^TR = R^TQ^TQR$ , so in this sense the storage for R from the Cholesky factorization (cholmod routine from SuiteSparse, as wrapped in Julia) is the same as the storage for R from the QR factorization. We used this equivalence as a work-around for the Julia qrfact, so long as  $\left[\frac{I_n}{C_{drop}}\right]$  was not so poorly conditioned that the Cholesky factorization failed.

### 4 Numerical experiments

We performed julia lufact LU = PAQ decompositions for the 235 matrices of the test set defined in Section 3.

We plot in Figure 3 the condition number of  $LL_1^{-1}$  for these 235 matrices. We observe that  $LL_1^{-1}$  tends to be acceptably well conditioned (192 of 235 matrices had  $cond_2(LL_1^{-1}) < 400$ ). As noted in Section 2, the condition number is easily computed, using the symmetric power method. Of the 43 more poorly conditioned matrices, 29 were among the 47 matrices with more than one hundred rows in C (i.e. correspond to "tall" A matrices), indicating that  $LL_1^{-1}$  preconditioning is appropriate for near square matrices.

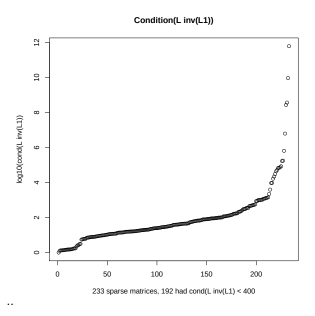


Figure 3: Condition number of  $LL_1^{-1}$  for the 235 test matrices.

We plot in Figure 4 the iteration count obtained using test matrices for which

lsqrLU converged in n iterations (iteration matrix L). For these matrices, we observe that the convergence is significantly accelerated by iterating on  $LL_1^{-1}$ .

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Figure 4: Convergence for 79 matrices for which lsqr iteration with L converged in n iterations.

After the lsqrLinvL algorithm has converged, a final solve with U is needed. However we can observe in Figure 5 that for some matrices U may have near zero diagonal elements. If we consider the normalized (by column norm) size of the diagonal elements, 30 of the 235 matrices had diagonal values  $u_{ii}$  such that  $|u_{ii}|/max_i|u_{ij}| < 10^{-10}$ , so for these matrices neither L nor  $LL_1^{-1}$  preconditioning was judged to be feasible.

We discard 30 matrices because  $|u_{ii}|/max_i|u_{ij}| < 10^{-10}$ . For the remaining 205 of the matrices U, we obtain, as explained in Section 2, an upper bound  $K = \sqrt{\lambda_{max} + 1}$  of  $cond_2(LL_1^{-1})$ , where  $\lambda_{max}$  is the largest eigenvalue of  $C^TC$  (computed using the symmetric power method.) Then we select the 1sqr iteration matrix by the size of K, as follows:

- If K < 400, we use lsqrLinvL (lsqr algorithm iterating with  $LL_1^{-1}$ ). This situation concerns 173 of the 205 matrices.
- If  $400 < K < 10^8$ , we explicitly compute C and we perform a drop tolerance on C by zeroing elements  $c_{ij}$  such that  $|c_{ij}| < |c_i|/K^{.25}$ , where  $c_i$  denotes the maximal entry of the ith column of C, leading to the matrix  $C_{drop}^{3}$ .

<sup>&</sup>lt;sup>3</sup> Taking  $K^{.25}$  means for example that if  $K=10^4$ , then all entries less than 1/10 of the column max in absolute value are discarded. Taking a constant less than .25 would reduce fill, but might retard convergence.

### minimal values of diagonals of U from PAQ = LU

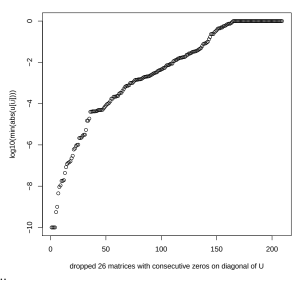


Figure 5: Size of the diagonal elements of U.

We then compute the Cholesky factor R satisfying

$$R^T R = \left[I_n | C_{drop}^T\right] \left[\frac{I_n}{C_{drop}}\right],$$

and we use 1sqr iteration with  $LL_1^{-1}R^{-1}$  (28 of 205 matrices).

- If  $K > 10^8$ , we have  $cond_2(F^TF) > 10^{16}$ . Then the Cholesky decomposition in double precision arithmetic gives a floating point error. In this case, we try lsqr iteration with L (4 of 205 matrices).

The algorithm where we choose, as mentioned above, the iteration matrix according to the size of K will be referred to as the "hybrid"  $\mathtt{lsqrLinvL}$  algorithm in the remainder of this paper. In our experiments, we iterated with a stopping criterion of  $|\delta| < 10^{-10}$  in Algorithm 2. The results<sup>4</sup> are summarized in Table 1. In this table, the number of iterations to convergence is expressed as a multiple of n (number of columns of A) and the storage required is expressed as a multiple of the nonzero entries of A. As expected, the case  $400 < K < 10^8$  requires more storage (×13 in average) due to the use of R but decreases the number of iterations by a factor 2.3 in average. The decreased storage for the largest condition numbers is the average storage for A + L + U, which is the same storage required for K < 400, but with the average computed over a much smaller set.

 $<sup>^4{</sup>m The}$  results are stored in a comma separated file with 235 rows and 19 numbers per line, loadable as an R language data frame.

As we can see in Figure 6, when using a drop tolerance on C and then computing  $R^TR = F_{drop}^T F_{drop}$  (where  $F_{drop} = \left[\frac{I_n}{C_{drop}}\right]$ ), the resulting R factor is relatively dense matrix (R has relatively few entries when A had only 25 columns and many rows).

		Cases	Iterations	Nonzeros
Condition	Iteration	of	Multiple	Multiple
Number	Matrix	205	of $n$	nnz(A)
K < 400	$LL_1^{-1}$	173	.32 n	12.6
$400 < K < 10^8$	$LL_1^{-1}R^{-1}$	28	.138 n	165
$10^8 < K$	L	4	.6 n	6.52

Table 1: Storage and iterations for 205 matrices using the hybrid lsqrLinvL algorithm. The last two columns give averages over 173, 28, and 4 test matrices, respectively.

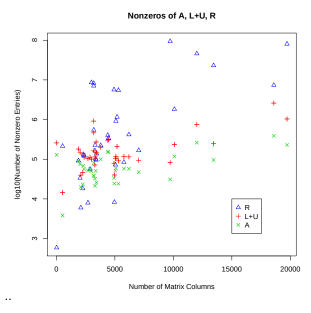


Figure 6: Fill-in for the Cholesky factorization of  $F_{drop}^T F_{drop}$ .

For each of the 205 matrices, the hybrid  ${\tt lsqrLinvL}$  algorithm converged to a solution. For 152 of 205 matrices, we have

$$\frac{\|x_{LinvL} - x_{qr}\|_{2}}{\|x_{LinvL}\|_{2}} < 10^{-8},$$

where  $x_{LinvL}$  and  $x_{qr}$  are the solution computed using the hybrid lsqrLinvL

and SparseQR algorithms, respectively. When the solutions differ, which of SparseQR and lsqrLinvL is better in terms of finding x with a lower LLS residual? Figure 7 plots the log of

$$\frac{\|b - Ax_{qr}\|_2}{\|b - Ax_{LinvL}\|_2}.$$

This graph shows that in most cases, the x values do not differ significantly, so the ratio is 1 and its log is zero. But when x differs (i.e. the log of the ratio is larger than zero), then residuals are smaller for the hybrid lsqrLinvL algorithm.

### 

Figure 7: Comparison between LLS residuals for  $x_{qr}$  and  $x_{LinvL}$ .

Before claiming victory of lsqrLinvL over SparseQR, we should note that for the 53 solutions that differed significantly,  $||x||_2$  tended to be larger for the hybrid lsqrLinvL method than for SparseQR (42 of 53 cases). Then users concerned with  $||x||_2$  might well prefer SparseQR.

Balancing concerns for size of the residual vs. size of the solution can be thought of as "regularization". For example, the Tikhonov regularization (see, e.g., [13, p. 193]) chooses x to minimize the function

$$T(x) = \sqrt{\|Ax - b\|_2^2 + \lambda^2 \|x\|_2^2}.$$

Figure 8 plots the log of the ratio

$$\sqrt{\frac{\|b - Ax_{qr}\|_2^2 + \lambda^2 \|x_{qr}\|_2^2}{\|b - Ax_{LinvL}\|_2^2 + \lambda^2 \|x_{LinvL}\|_2^2}},$$

with the Tikhonov regularization parameters  $\lambda=10^{-8}$  and  $\lambda=10^{-4}$ . We recall that for most matrices,  $\|x_{qr}-x_{LinvL}\|$  is quite small. In these cases, residuals and solution norms match and the ratio is close to one. In some cases, residuals and norms do not match, then the ratios  $T(x_{qr})/T(x_{LinvL})$  are larger than one (points greater than zero on the plot) indicating that T(x) is larger for SparseQR than for lsqrLinvL solutions. For  $\lambda=10^{-8}$ , lsqrLinvL usually gives a solution with a smaller T(x). For  $\lambda=10^{-4}$ , the  $T(x_{qr})$  quantity is usually smaller.

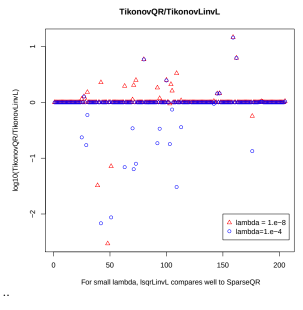


Figure 8: Comparison between Tikhonov regularizations of  $x_{qr}$  and  $x_{LinvL}$ .

### 5 Conclusion and future work

When the number of equations is not much larger than the number of variables, LU factorization usually allows iterative least squares solves with less storage than QR factorization or Cholesky factorization of the normal equations. Moreover, when the number of equations is not much more than the number of unknowns, iterating in the 1sqr algorithm using  $LL_1^{-1}$  usually requires relatively few iterations for the solution of a least squares problem. A main limitation is that the method fails when U has near zero diagonal elements, i.e., when the problem is numerically not overdetermined. The near singularity of U is typically not discovered until the relatively expensive LU factorization has been attempted.

As future work, a possible way to further reduce the required storage is to perform the original LU factorization in a lower precision (or to use an incomplete factorization of A). The iteration is then with  $AU^{-1}L_1^{-1}$ , possibly with a

preliminary cheaper iteration with  $\left[\frac{I}{L_2L_1^{-1}}\right]$ . That iteration can be compared to the RIF preconditioner [2], which is a sparse Cholesky-like factorization of  $A^TA$ . Other future tasks will be to address larger problems using the distributed computation possibilities provided by Julia, and to analyze the resulting execution time and speedup.

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