CIRJE-F-478

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> > February 2007

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## On the estimation of a large sparse Bayesian system: the Snaer program<sup>\*</sup>

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February 26, 2007

<sup>\*</sup>An early version of this paper was presented at the 3rd IASC World Conference on Computational Statistics & Data Analysis, Cyprus, October, 2005. We thank the participants for their constructive comments. We are grateful to Jan van Tongeren for his long-term support in this project.

#### AMS Subject Classifications: 62F15, 65F50, 93A15, 62F30.

**Key words:** Bayesian linear estimation, sparse matrices, large-scale optimization, nonlinear constraints.

Abstract: The Snaer program calculates the posterior mean and variance of variables on some of which we have data (with precisions), on some we have prior information (with precisions), and on some prior indicator ratios (with precisions) are available. The variables must satisfy a number of exact restrictions. The system is both large and sparse. Two aspects of the statistical and computational development are a practical procedure to solve a linear integer system, and a stable linearization routine for ratios. We test our numerical method to solve large sparse linear least-squares estimation problems, and find that it performs well, even when the  $n \times k$  design matrix is large ( $nk \approx 2^{27.5}$ ).

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## 1 Introduction

Suppose we are given a system of latent variables to be estimated. The information consists of incomplete data (with precisions), priors on a subset of the variables or linear combinations thereof (with precisions), and exact linear restrictions. Some of the priors are so-called indicator ratios, hence nonlinear.

The system that we are going to be interested in is large and sparse. In this context, a system with  $2^6$  variables and  $2^{12}$  observations, thus giving  $2^{18} \approx 260,000$  entries in the design matrix is *not* considered to be large. On the other hand, a system with  $2^{11}$  variables and  $2^{13}$  observations, thus giving  $2^{24} \approx 16.8$  million entries in the design matrix is considered large.

Our system is sparse, because information is often available on one variable at a time, and restrictions are often definitions involving only a small number of variables. A matrix is sparse when it has many structural zeros. If a  $p \times n$  matrix possesses s structural zeros, then the matrix can be stored as a  $(pn-s) \times 3$  matrix, where the *i*-th row contains the row-index, column-index, and value of the *i*-th nonzero entry. This is often useful if storage space is more important than access speed. The number s/(pn) is called the sparsity of the matrix.

In this paper we analyze how we can estimate such a large and sparse system efficiently and accurately. The analysis contains both statistical and computational aspects. The statistical problem was first considered by Magnus, Van Tongeren, and De Vos (2000), but the sparse and large-dimension aspects were not considered in that paper. In addition, a number of their proposed solutions — especially for the treatment of indicator ratios — do not work well in large dimensions. Also, the Bayesian solution involves difficult inversions that can be avoided by rewriting the system in a different format. The purpose of this paper is to describe both the statistical and the computational aspects, and to perform tests for the stability, accuracy, and speed of the proposed routines.

The concept of sparsity is now widespread in modern numerical methods, and huge sparse matrices often appear in science and engineering when solving problems of mathematical physics and computational chemistry. Recently, large sparse matrices have become important in economics, for example in studying longitudinal samples of over one million workers from more than 500,000 employing firms; see Abowd, Kramarz, and Margolis (1999) and Abowd, Creecy, and Kramarz (2002), using an algorithm due to Dongarra, Duff, Sorenson, and Van der Vorst (1991). There exists a large literature on sparse matrices. Some recent references can be found in Björck (1996, Chapters 6 and 7) and Pauletto (1997). The search for accurate and fast software for large and sparse linear estimation problems was driven by the difficulties encountered in estimating national accounts data, especially in developing countries, where one may encounter 5000 or more variables and 20,000 observations. The relevant matrices in such applications are sparse but not structured, that is they do not have a Kronecker, Toeplitz, or other well-known structure. The literature on solving models with structured matrices utilizes the fact that iterative algorithms — which require only matrix-vector products — can provide storage or computational advantages, but since we only consider general (nonstructured) large and sparse matrices we cannot benefit from these advantages.

The Snaer (Software for National Accounts Estimation and Reconciliation) program, developed for this purpose, thus calculates the posterior mean and variance of variables, some of which are given as data (with precisions), on some we have prior information (with precisions) or prior indicator ratios (with precisions), and exact restrictions apply.

Our new theory for the estimation of national accounts has a number of characteristics:

- It specifies data and their precisions, in addition to priors and their precisions;
- It allows indicator ratios (with precisions) as input priors;
- It takes full account of all accounting identities;
- Solutions are continuous rather than discrete;
- It allows for multiple priors on variables or linear combinations of variables;
- The posterior estimates take all prior and data input into account and come with precisions;
- The system is transparent, flexible, and fast;
- Sensitivity analysis is easy to perform, and alternative scenarios may be analyzed;
- It can be applied without major alterations in existing compilation methods.

A first major application of the methodology and the software will be in St Vincent; see Magnus, Van Tongeren, Giesberts, and Danilov (2007). But also applications in developed countries are now underway. These applications

concern not only large economic data sets for national accounts, but also input-output analysis, flow-of-funds analysis, and the integration of quarterly and annual accounts.

There are many other potential applications of the proposed method, outside national accounting, for example in large employment matrices, material flow analysis in environmental accounting, socio-economic and other satellite accounts, integration of quarterly and annual accounts, business accounting, and demographic projections.

The plan of this paper is as follows. We formally describe the set-up in Section 2. In Sections 3 and 4 we present two solutions. First a Bayesian solution to a Bayesian problem. Then an equivalent least-squares solution, which involving less inversions and is much easier to estimate. The implementation of the theory together with some extensions and refinements is discussed in Section 5. An important aspect of our theory is the possibility to include indicator ratios as priors. These ratios must be linearized, but this can be done in many ways. Two solutions and their merits are presented in Section 6. In Sections 7 and 8 we perform Monte Carlo experiments to test the accuracy and speed of our proposed sparse least-squares routine (which also calculates variances). Section 9 concludes and discusses possible extensions.

## 2 Set-up: data, priors, and restrictions

We consider a vector x of n latent variables to be estimated. Data are available on p components (or linear combinations of components) of x. Let  $d_1$  denote the  $p \times 1$  data vector, generated by the measurement equation

$$d_1 | x \sim \mathcal{N}_p(D_1 x, \Sigma_1). \tag{1}$$

Typically, the  $p \times n$  matrix  $D_1$  is a selection matrix, say  $D_1 = (I_p, 0)$ , so that  $D_1 x$  is a subvector of x, but this is not required. Neither is it required that the matrix  $D_1$  has full row-rank; in fact, p may be larger than n. Measurements are unbiased in the sense that  $E(d_1|x) = D_1 x$ . The  $p \times p$  matrix  $\Sigma_1$  denotes a positive definite (hence nonsingular) variance matrix, typically (but not necessarily) diagonal.

In addition to the p data, we have access to two further pieces of information: prior views concerning the latent variables or linear combinations thereof, and deterministic (accounting) constraints. In particular, we have  $m_1$  random priors:

$$A_1 x \sim \mathcal{N}_{m_1}(h_1, H_1) \tag{2}$$

and  $m_2$  exact restrictions:

$$A_2 x = h_2 \text{ (a.s.)},\tag{3}$$

in total  $m := m_1 + m_2$  pieces of prior information.

We shall assume that the  $m_1 \times m_1$  matrix  $H_1$  is positive definite (hence nonsingular) and that the  $m_2 \times n$  matrix  $A_2$  has full row-rank  $m_2$ . This guarantees that the exact restrictions are linearly independent and thus form a consistent set of equations. Neither of these two restrictions constitutes any loss of generality, because if  $H_1$  is singular some of the priors can be interpreted as exact restrictions, and if  $A_2$  does not have full row-rank, we simply throw away the redundant restrictions, obviously without loss of information.

In order to identify all n variables from the information (data and priors) we need at least n pieces of information:  $m + p \ge n$ . But this is not sufficient for identification, because some of the information may be on the same variables. The condition

$$\operatorname{rk} \begin{pmatrix} D_1 \\ A_1 \\ A_2 \end{pmatrix} = n \tag{4}$$

is necessary and sufficient for identification.

## **3** Bayesian solution

In order to estimate the n latent variables x, we define

$$A := \begin{pmatrix} A_1 \\ A_2 \end{pmatrix}, \quad h := \begin{pmatrix} h_1 \\ h_2 \end{pmatrix}, \quad H := \begin{pmatrix} H_1 & 0 \\ 0 & 0 \end{pmatrix},$$

and we make the additional assumption that

$$\operatorname{rk}(A) = m,$$

which, of course, implies that both  $A_1$  and  $A_2$  have full row-rank. In fact, as shown in Danilov and Magnus (2006), the rank condition on A is not a binding restriction, because we can always move priors to data (and vice versa). This condition will be dropped in the next section.

Since  $\operatorname{rk}(A) = m$  we have  $m \leq n$ . If m = n, then all variables are identified. If m < n, we define a semi-orthogonal  $n \times (n - m)$  matrix L such that AL = 0 and  $L'L = I_{n-m}$ . The condition

$$\operatorname{rk}(D_1L) = n - m.$$

is then an alternative and equivalent identifiability condition to (4), because the definition of L implies that

$$\operatorname{rk}\begin{pmatrix}A\\D_1\end{pmatrix} = \operatorname{rk}(A) + \operatorname{rk}(D_1L).$$

From Theorem 1 of Magnus, Van Tongeren, and De Vos (2000), we know that the posterior distribution of x is given by

$$x|d_1 \sim \mathcal{N}_n(\mu, V),$$

where

$$\mu = A^{+}h - (A^{+}HA^{+'} + CK)D_{1}'\Sigma_{0}^{-1}(D_{1}A^{+}h - d_{1})$$
(5)

and

$$V = A^{+}HA^{+'} - A^{+}HA^{+'}D_{1}'\Sigma_{0}^{-1}D_{1}A^{+}HA^{+'} + CKC',$$
(6)

and the following definitions have been employed:

$$\Sigma_0 := \Sigma_1 + D_1 A^+ H A^{+'} D_1', \quad C := I_n - A^+ H A^{+'} D_1' \Sigma_0^{-1} D_1,$$

and

$$K := \begin{cases} L(L'D_1'\Sigma_0^{-1}D_1L)^{-1}L' & \text{if } m < n, \\ 0 & \text{if } m = n. \end{cases}$$

The matrix  $A^+ := (A'A)^{-1}A'$  denotes the Moore-Penrose inverse of A.

This is a useful theoretical result, but Equations (5) and (6) do not lend themselves well to applications where the dimensions are large. Hence we seek an alternative but equivalent formulation which allows us to tackle largedimensional problems.

## 4 Least-squares solution

Because of the normality assumption for both data and priors, there is no mathematical or statistical difference between them (although there is a conceptual difference); see Danilov and Magnus (2006). We may therefore move all random priors to the data and consider a new "data" vector

$$d := \begin{pmatrix} d_1 \\ h_1 \end{pmatrix}.$$

Defining

$$D := \begin{pmatrix} D_1 \\ A_1 \end{pmatrix}, \quad \Sigma := \begin{pmatrix} \Sigma_1 & 0 \\ 0 & H_1 \end{pmatrix},$$

we may write the new measurement equation as

$$d|x \sim N_{p+m_1}(Dx, \Sigma)$$

together with the priors (exact restrictions)

$$A_2 x = h_2$$
 (a.s.).

The Bayesian problem is then equivalent to the constrained least-squares estimation problem of estimating x from

$$d = Dx + \varepsilon, \qquad \varepsilon \sim \mathcal{N}_{p+m_1}(0, \Sigma),$$

under the restriction  $A_2 x = h_2$ .

In many cases, an unconstrained problem is easier to solve in large dimensions than a constrained problem. Thus, let

$$A_2 = (A_{21} : A_{22}),$$

where  $A_{21}$  is an  $m_2 \times (n - m_2)$  matrix and  $A_{22}$  is a nonsingular  $m_2 \times m_2$  matrix. Partitioning x accordingly, we write the restriction as

$$A_{21}x_1 + A_{22}x_2 = h_2,$$

so that

$$x = \begin{pmatrix} x_1 \\ x_2 \end{pmatrix} = \begin{pmatrix} I \\ -A_{22}^{-1}A_{21} \end{pmatrix} x_1 + \begin{pmatrix} 0 \\ A_{22}^{-1}h_2 \end{pmatrix} \equiv Qx_1 + q.$$

Let

$$B := \Sigma^{-1/2} D = \begin{pmatrix} \Sigma_1^{-1/2} D_1 \\ H_1^{-1/2} A_1 \end{pmatrix}, \quad b := \Sigma^{-1/2} d = \begin{pmatrix} \Sigma_1^{-1/2} d_1 \\ H_1^{-1/2} h_1 \end{pmatrix}.$$

Then the constrained problem of minimizing  $(d - Dx)'\Sigma^{-1}(d - Dx)$  subject to  $A_2x = h_2$  is equivalent to the unconstrained problem

minimize 
$$||(b - Bq) - BQx_1||$$

with respect to  $x_1$ . The solution, say  $\mu_1$ , is our estimator for  $x_1$  with associated variance  $\operatorname{var}(\mu_1) = (Q'B'BQ)^{-1}$ . Since  $x_2$  is a linear function of  $x_1$  we can estimate the complete x-vector and obtain the full variance matrix V.

## 5 Implementation

Our information thus consists of p data:

$$d_1|x \sim \mathcal{N}_p(D_1x, \mathcal{L}_1),$$

 $m_1$  random priors:

$$A_1 x \sim \mathcal{N}_{m_1}(h_1, H_1),$$

and  $m_2$  exact priors:

$$A_2 x = h_2$$
 (a.s.).

We need to estimate this system taking into account that the dimensions may be very large and that the matrices are sparse. We proceed as follows.

Step 1: Rank. Two rank-conditions need to be satisfied:

$$\operatorname{rk}(A_2) = m_2, \quad \operatorname{rk} \begin{pmatrix} D_1 \\ A_1 \\ A_2 \end{pmatrix} = n.$$

If the rank condition on  $A_2$  is not satisfied, then there are redundant rows in  $A_2$  which should be deleted. But if the second rank condition is not satisfied, then there is insufficient information in the data and priors to estimate the whole system, and more information needs to be acquired. The computation of the rank of a (sparse) matrix is no trivial exercise. Several good routines are available, for example in the NAG (2006) or Harwell (2004) libraries. Our routine Snaer-RANK has two special features. First, it performs a preliminary (iterative) dimension reduction by looking for rows and columns with a single nonzero entry. Second, it provides information about where rank deficiencies (if any) occur, which is of great practical use in repairing the inputs.

Step 2: Scaling. The matrices  $\Sigma_1$  and  $H_1$  must be nonsingular. We scale by defining

$$B := \begin{pmatrix} \Sigma_1^{-1/2} D_1 \\ H_1^{-1/2} A_1 \end{pmatrix}, \quad b := \begin{pmatrix} \Sigma_1^{-1/2} d_1 \\ H_1^{-1/2} h_1 \end{pmatrix}$$

of orders  $(p + m_1) \times n$  and  $p + m_1$ , respectively.

Step 3: Re-ordering. If necessary, we re-order the x-variables such that the matrix  $A_{22}$  in the restriction

$$A_2 x = A_{21} x_1 + A_{22} x_2 = h_2$$

is a nonsingular  $m_2 \times m_2$  matrix. We re-order the columns of the matrices  $A_2$  and B accordingly. There is no unique way to select  $m_2$  linearly independent columns of  $A_2$ . But some selections are more robust than others computationally. Also, some selections require less covariances than others in the computation of the diagonal elements of the variance matrix (Step 8). These considerations are taken into account at the selection stage.

Step 4: Solving. The inverse of  $A_{22}$  appears in the formulae, but in fact we only need the expressions  $A_{22}^{-1}A_{21}$  and  $A_{22}^{-1}h_2$ . Hence, we solve the sparse matrix equation

$$A_{22}X = (-A_{21}:h_2)$$

and we write the resulting  $m_2 \times (n - m_2 + 1)$  matrix X as  $X = (Q_1 : q_1)$ .

There exist a number of sparse routines to solve linear equations. Our sparse subroutine Snaer-SOLVE builds on the existing routines, but it contains two special features.

First, not only the left-hand side matrix  $A_{22}$  but also the right-hand side matrix  $(-A_{21}:h_2)$  is (or can be) sparse.

Second, there is an integer refinement possibility in the routine, based on the fact that the matrix  $A_{22}$  is often an integer matrix, that is, all nonzero elements of  $A_{22}$  are integers. The reason for this is that in many applications the restrictions are in fact definitions, such as  $x_1 = x_2 + x_3 + x_4$ . The nonzero entries in  $A_{22}$  are then  $\pm 1$ . In the case when  $A_{22}$  is an integer matrix, the determinant  $|A_{22}|$  will be an integer, and the determinant of every submatrix of  $A_{22}$  will also be an integer. Hence, the adjoint  $A_{22}^{\#}$  of  $A_{22}$  is an integer matrix.

Now let c be a column of  $(-A_{21} : h_2)$ , all of whose nonzero elements are integers, and let x be the corresponding column of the solution matrix X, so that  $A_{22}x = c$ . Then,

$$x = A_{22}^{-1}c = \frac{A_{22}^{\#}c}{|A_{22}|},$$

each element of which is the ratio of two integers. We know  $|A_{22}|$  (integer) and x (real) and we can improve on the solution for x using the integer feature, as follows:

(a) Compute  $\Delta_1 := |A_{22}|$  and  $z_1 := A_{22}^{\#} c = \Delta_1 x$ ;

(b) Use the fact that  $\Delta_1$  and the elements of  $z_1$  are all integers, and obtain "exact" integer solutions  $\Delta_2$  and  $z_2$ ;

(c) Now obtain an improved solution for  $x = z_2/\Delta_2$ , each element of which is a rational number.

The new solution is "exact". This "integer smoothing" can be done for all integer columns of the matrix  $(-A_{21} : h_2)$ .

Step 5: Multiplication. Compute the matrix and vector

$$Q := \begin{pmatrix} I_{n-m_2} \\ Q_1 \end{pmatrix}, \quad q := \begin{pmatrix} 0 \\ q_1 \end{pmatrix},$$

and the matrix and vector

$$R := BQ, \quad r := b - Bq.$$

Step 6: Minimization. We need to minimize the norm

$$||r - Rx_1||,$$

the solution of which we call  $\mu_1$ . Notice that the matrix R has full column-rank  $n - m_2$ . The least-squares problem

$$\min_{x_1} (r - Rx_1)' (r - Rx_1)$$

can be rewritten as

$$\min_{x_1} \left( r'r - 2r'Rx_1 + x_1'R'Rx_1 \right),\,$$

which in turn is equivalent to the quadratic minimization problem

$$\min_{x_1} \left( \frac{1}{2} x_1' H x_1 - c' x_1 \right),$$

where H := R'R and c := R'r. There are many libraries that offer routines dealing with sparse least-squares, in particular Matlab (2002), NAG (2006), and Harwell (2004), but they are all quite different. We offer a brief comparison of these routines in Sections 7 and 8. Our Snaer-OLS routine is based on the Gould-Nocedal (1998) algorithm, which also underlies the Harwell (2004) approach. This matrix H is decomposed as H = PLDL'P', where P is a permutation matrix, L is unit lower triangular, and D is block-diagonal with blocks of dimension one or two.

Least-squares routines typically do not compute (elements of) the variance matrix, but the Snaer-OLS routine does, in the following way. Let  $e^{(j)}$  denote the *j*-th unit vector, that is, the *j*-th column of  $I_{n-m_2}$ . The *j*-th column  $v^{(j)}$  of the matrix  $V_{11} := (R'R)^{-1}$  can be found by minimizing  $||e^{(j)} - Hv^{(j)}||^2$  for all *j*, that is by solving the quadratic minimization problem

$$\min_{v^{(j)}} \left( \frac{1}{2} v^{(j)'} H^2 v^{(j)} - e^{(j)'} H v^{(j)} \right), \qquad j = 1, \dots, n - m_2,$$

whose value at the minimum should be -1/2. It is, however, faster and more accurate to obtain the columns of  $V_{11}$  by minimizing  $||H^{-1/2}e^{(j)} - H^{1/2}v^{(j)}||^2$  for all j, that is by solving the quadratic minimization problem

$$\min_{v^{(j)}} \left( \frac{1}{2} v^{(j)'} H v^{(j)} - e^{(j)'} v^{(j)} \right), \qquad j = 1, \dots, n - m_2,$$

using the Snaer-OLS algorithm. The value at the minimum equals  $-v_i^{(j)}/2$ .

In practice we will not need to calculate (or store) all elements of  $v^{(j)}$ . To store all elements is a physical impossibility in large dimensions. We have to indicate in advance which elements need to be stored. These typically include all diagonal elements of  $V_{11}$  (the variances) and a selection of covariances. The selected covariances are either of direct interest or they are required to calculate the diagonal (or other) elements of  $V_{22} := \operatorname{var}(\mu_2)$ . For example, if  $x_{21} = x_{11} + x_{12}$ , where  $x_{21}$  is an element of  $x_2$  and  $x_{11}$  and  $x_{12}$ are elements of  $x_1$ , then we need  $\operatorname{cov}(\mu_{11}, \mu_{12})$  in order to calculate  $\operatorname{var}(\mu_{21})$ .

Step 7: Completion of  $\mu$ . We compute

$$\mu_2 = Q_1 \mu_1 + q_1.$$

Step 8: Completion of V. The variance of  $\mu$  is given by

$$V = \operatorname{var}(\mu) = \begin{pmatrix} I \\ Q_1 \end{pmatrix} (R'R)^{-1} (I, Q'_1).$$

But only selected elements of V are required and only selected elements of  $V_{11} = (R'R)^{-1}$  are stored in Step 6. The subroutine Snaer-OLS automatically selects which elements of  $V_{11}$  need to be stored in order to calculate the required elements of V.

#### 6 The treatment of ratios

One of the features of our program is that also ratios can be used as priors. This is of much practical use, for example in the estimation of national accounts, where many ratios are used as indicators and are assumed to be relatively stable, hence predictable.

Let x and y be two latent variables and assume information is available on the ratio R := y/x. If this ratio is part of an otherwise linear system, then we can either transform the linear system into a nonlinear one, or we can linearize the ratio. Especially if the system is large, the latter option is often more practical. Thus, we consider a linearization y - rx for some suitably chosen value of r. If information is available on the first two moments of R, the question then becomes how to transform these moments into moments of the linearization.

Consider first the following special case. Let r denote the expected value of R, if it exists, and suppose that R and x are independent. Then we write y - rx = (R - r)x, so that

$$\mathbf{E}(y - rx) = \mathbf{E}(R - r)\mathbf{E}(x) = 0$$

and

$$\operatorname{var}(y - rx) = \operatorname{E}(y - rx)^2 = \operatorname{E}(R - r)^2 \operatorname{E}(x^2) = \operatorname{var}(R) \cdot (\operatorname{var}(x) + (\operatorname{E} x)^2).$$

Hence, in this special case, we easily obtain the moments of the linearization. In general, however, this is less straightforward.

#### 6.1 Bayesian solution

Suppose we have data:  $d|x \sim (x, \sigma^2)$  and a prior on R = y/x,

$$\pi: \quad R \sim (r, \tau^2).$$

We wish to replace the prior on the ratio by a prior on its linearization, so we consider

$$\pi': \quad y - rx \sim (0, \omega^2).$$

The question is how to choose  $\omega^2$ . Let us calculate the posterior moments of y|d based on  $\pi$  and  $\pi'$  respectively. Based on the linearization  $\pi'$  we obtain the posterior moments of y given d as

$$y|d \sim (rd, r^2\sigma^2 + \omega^2). \tag{7}$$

Notice that the two pieces of information  $x \sim N(d, \sigma^2)$  and  $y - rx \sim N(0, \omega^2)$ yield  $y \sim N(rd, r^2\sigma^2 + \omega^2)$  if and only if x and y - rx are uncorrelated, that is, if and only if r = cov(x, y) / var(x).

Next we calculate the posterior moments of y|d based on  $\pi$ . Following the arguments in Magnus, Van Tongeren, and De Vos (2000), we obtain

$$\mathbf{E}(y|x) = rx, \quad \operatorname{var}(y|x) = \tau^2 x^2.$$

Let z := y | x. Then the posterior moments of y | d are

$$E(y|d) = E_z E(d|d, z) = E_z E(y|d, x)$$
$$= E_z E(y|x) = E_z(rx) = rd$$
(8)

and

$$\operatorname{var}(y|d) = \operatorname{E}_{z} \operatorname{var}(y|d, z) + \operatorname{var}_{z}(\operatorname{E}(y|d, z))$$
$$= \operatorname{E}_{z} \operatorname{var}(y|x) + \operatorname{var}_{z}(\operatorname{E}(y|x))$$
$$= \operatorname{E}_{z}(\tau^{2}x^{2}) + \operatorname{var}_{z}(rx)$$
$$= \tau^{2}(d^{2} + \sigma^{2}) + r^{2}\sigma^{2}.$$
(9)

In order to reconcile (7) with (8) and (9) we must have

$$\omega^2 = \tau^2 (d^2 + \sigma^2). \tag{10}$$

In a practical situation, we know r and  $\tau^2$ , but we do not necessarily know d and  $\sigma^2$  (the moments of x). If we would know the mean and variance of x, then we could replace the prior on R by a prior on y - rx. Since we don't know the moments of x, we may use a simple iterative procedure, based on the assumption that some reasonable starting values for E(x) and var(x) are available. We use these as proxies for d and  $\sigma^2$  and obtain posterior moments of all latent variables, in particular E(x) and var(x). Next, we set d = E(x) and  $\sigma^2 = var(x)$ , recalculate the prior variance of y-rx from (10), and obtain posterior moments again. We may continue this process until convergence. In practice we will have not one but many indicator ratios. The iteration procedure is then applied to all of them simultaneously.

The iterative procedure is justified by the fact that it leads to the correct first two posterior moments. The process does not, however, always converge, and it turns out to be rather unstable. Therefore we outline an alternative approach, based on a different philosophy, below.

#### 6.2 Invariance

Suppose again that the ratio y/x has a prior mean or median r, so that

$$\Pr\left(\frac{r}{a} < \frac{y}{x} < br\right) = 1 - \alpha,$$

where a and b are determined by the assumed prior standard deviation  $\tau_1$  of the ratio y/x. The associated confidence region is given by

$$C_1 = k\left(br - \frac{r}{a}\right) = kr\left(b - \frac{1}{a}\right)$$

for some constant k. Let us assume that y and x are positive (almost surely) and that a > 1, b > 1, and r > 0. This gives

$$\Pr\left(-(1-1/a)rx < y - rx < (b-1)rx\right) = 1 - \alpha$$

with associated confidence region

$$C_2 = k \left( (b-1)rx + (1-1/a)rx \right) = krx \left( b - 1/a \right) = C_1 x.$$

We wish to "translate" the prior information on the ratio into prior information on its linearization. Motivated by the comparison of the two confidence bands, we assume that y - rx is distributed around zero with some standard deviation  $\omega_1 = \tau_1 x$ .

Although we have written the prior information in terms of the ratio y/x, we could equally well have written it in terms of x/y. Two different linearizations do not, in general, yield the same result. The invariance approach demands that we obtain the same answer irrespective whether we start from y/x or x/y.

Thus motivated, we rewrite the ratio as

$$\Pr\left(\frac{1}{br} < \frac{x}{y} < \frac{a}{r}\right) = 1 - \alpha.$$

with confidence region

$$C_3 = k\left(\frac{a}{r} - \frac{1}{br}\right) = \frac{k}{r}\left(a - \frac{1}{b}\right).$$

We obtain the linearization

$$\Pr\left(-(1-1/b)\frac{y}{r} < x - (1/r)y < (a-1)\frac{y}{r}\right) = 1 - \alpha,$$

with associated confidence interval

$$C_4 = k\left((a-1)\frac{y}{r} + (1-1/b)\frac{y}{r}\right) = \frac{ky}{r}\left(a-1/b\right) = C_3y.$$

Let  $\tau_2$  be the prior standard deviation of x/y. Using the same argument as above, we assume that the linearization x - (1/r)y is distributed around zero with some standard deviation  $\omega_2 = \tau_2 y$ .

Since  $\omega_1 = r\omega_2$ , we obtain  $\tau_1 x = r\tau_2 y$ . If the prior values for y and x satisfy y/x = r exactly, then this gives

$$\frac{\tau_1}{r} = \frac{\tau_2}{1/r},$$

in other words: we must choose the *relative* precisions of the prior ratios to be the same.

However, the prior values for y and x will almost certainly not satisfy y/x = r exactly. Hence, we must massage the priors y and x in such a way that y/x = r.

Let  $y_0$  and  $x_0$  be the given prior values, and assume that at least one of them is available. Now define

$$A := \begin{cases} \frac{1}{1+r^2} x_0 + \frac{r^2}{1+r^2} (y_0/r) & \text{if both } x_0 \text{ and } y_0 \text{ are available;} \\ x_0 & \text{if only } x_0 \text{ is available;} \\ y_0/r & \text{if only } y_0 \text{ is available.} \end{cases}$$

Instead of  $x_0$  and  $y_0$ , we use  $\bar{x} := A$  and  $\bar{y} := rA$ , which obviously satisfy  $\bar{y} = r\bar{x}$ . Then, letting  $\omega_1 := \tau_1 \bar{x}$  and  $\omega_2 := \tau_2 \bar{y}$ , the procedure will be invariant to whether we choose y/x or x/y as our starting point. The weights  $1/(1+r^2)$  and  $r^2/(1+r^2)$  are "optimal" in the sense that if  $x_0$  and  $y_0$  are unbiased estimates of x and y, if  $E(y_0 - rx_0) = 0$ , and if  $x_0$  and  $y_0$  are uncorrelated with equal variances, then A is also an unbiased estimate of x and its variance is minimized.

One can look at the problem in a different way. If we think of the problem as minimizing a weighted sum of squares, then the relevant term in the objective function is

$$\phi_1 := \frac{(y/x - r)^2}{\tau_1^2} = \left(\frac{y - rx}{\tau_1 x}\right)^2$$

or

$$\phi_2 := \frac{(x/y - 1/r)^2}{\tau_2^2} = \left(\frac{y - rx}{\tau_2 ry}\right)^2.$$

Instead of  $\phi_1$  or  $\phi_2$ , we use the term

$$\phi := \left(\frac{y - rx}{\tau_1 \bar{x}}\right)^2 = \left(\frac{y - rx}{\tau_2 r \bar{y}}\right)^2,$$

where, as above,  $\tau_1 = r^2 \tau_2$ , and  $\bar{y}$  and  $\bar{x}$  are chosen from the "priors" such that  $\bar{y} = r\bar{x}$ . This provides further justification of the above method.

Summarizing, we start from a prior  $y/x \sim (r, \tau^2)$  and reference data  $x_0$ and  $y_0$  (only one of them needs to be available). The invariant linearization is  $y - rx \sim (0, \omega^2)$ , where  $\omega = \tau A$ . Notice that no iterations are required in this approach. This procedure works well in practice, because it forces the linearization to be accurate in precisely the region where it matters most.

At the end of the estimation process we obtain estimates  $\mu_x$  for x and  $\mu_y$  for y, and also estimates for  $var(\mu_x)$ ,  $var(\mu_y)$ , and  $cov(\mu_x, \mu_y)$ . The posterior moments of the ratio R = y/x are then estimated by

$$E(R) = \frac{\mu_y}{\mu_x}, \quad var(R) = \frac{var(\mu_y - r\mu_x)}{A^2} \le \frac{\omega^2}{A^2} = \tau^2,$$

where we notice that the posterior variance of R is never larger than the prior variance, in accordance to standard Bayesian theory.

## 7 Simulation set-up for the Snaer-OLS routine

In the next two sections we describe some Monte Carlo experiments concerning the Snaer-OLS routine, which we consider the most critical of the program. Our framework is the standard linear regression model

$$y = X\beta + \varepsilon, \quad \varepsilon \sim \mathcal{N}(0, \sigma^2 I_n),$$

from which we obtain the ordinary least-squares (OLS) estimates and their variance matrix as

$$\hat{\beta} = (X'X)^{-1}X'y, \quad \operatorname{var}(\hat{\beta}) = \sigma^2 (X'X)^{-1}.$$

We shall be concerned with the situation when the  $n \times k$  design matrix X is of "large" dimension and is "sparse". (Notice that in Sections 7 and 8 we write the dimensions of the  $(p + m_1) \times (n - m_2)$  design matrix as  $n \times k$ , which is more common and provides and easier link to least squares.) The two questions are whether our method, based on the Gould-Nocedal (1998) algorithm, performs the task which it is supposed to perform, and also whether it compares favorably to other available methods.

We wish to test our method in terms of precision and speed as a function of the dimensions (n and k), the sparsity, and the ill-conditioning of the problem. To conduct the simulations within the framework of the linear regression model, we need values for the parameters  $\beta_1, \ldots, \beta_k$ , and  $\sigma^2$ , and for the  $n \times k$  design matrix X. Throughout the simulation experiment we set  $\sigma^2 = 1$  and  $\beta = 0$ . Given  $\beta$  and  $\sigma^2$ , we generate  $\varepsilon_1, \ldots, \varepsilon_n$  as independent draws from a N(0, 1) distribution, and obtain  $y = X\beta + \varepsilon = \varepsilon$ . The only difficulty in the set-up is the choice of X, because we wish to assess the precision of the method, and hence require an X-matrix such that  $(X'X)^{-1}$ is known analytically, and where the sparsity and the ill-conditioning can be controlled.

The sparsity  $d_s$  of a matrix X is defined as

$$d_s(X) := \frac{\text{number of structural zeros in } X}{\text{number of elements in } X},$$

which is a number between 0 (full matrix, no sparsity) and 1 (null matrix, complete sparsity). For example, the sparsity of a diagonal  $k \times k$  matrix is 1 - 1/k. The sparsity  $d_s$  of our problem is defined as the sparsity of X. The ill-conditioning of the problem is defined by the *condition number c*:

$$c := \sqrt{\frac{\text{largest eigenvalue of } X'X}{\text{smallest eigenvalue of } X'X}}.$$

Since X has full column-rank k, we can write

$$X = (X(X'X)^{-1/2}) (X'X)^{1/2} \equiv ST,$$

where S is a semi-orthogonal  $n \times k$  matrix such that  $S'S = I_k$  and T is a symmetric nonsingular  $k \times k$  matrix. We now construct S and T in such a way that T (and hence X'X) has an analytically known inverse.

#### 7.1 Construction of T

Let  $T_0$  be the tridiagonal symmetric  $k \times k$  matrix

$$T_0 = \frac{1}{4} \begin{pmatrix} 2 & -1 & 0 & \dots & 0 & 0 \\ -1 & 2 & -1 & \dots & 0 & 0 \\ 0 & -1 & 2 & \dots & 0 & 0 \\ \vdots & \vdots & \vdots & & \vdots & \vdots \\ 0 & 0 & 0 & \dots & 2 & -1 \\ 0 & 0 & 0 & \dots & -1 & 2 \end{pmatrix}$$

From Grenander and Szegö (1958, p. 67) and Samarsky (1971, p. 53), we know that the eigenvalues  $\lambda_{01} > \lambda_{02} > \cdots > \lambda_{0k}$  of  $T_0$  are given by

$$\lambda_{0j} = \sin^2\left(\frac{(k+1-j)\pi}{2(k+1)}\right), \quad j = 1, \dots, k,$$

and the corresponding eigenvectors  $u_1, u_2, \ldots, u_k$  by

$$u_{ij} = \sqrt{\frac{2}{k+1}} \sin\left(\frac{i(k+1-j)\pi}{k+1}\right),\,$$

where  $u_{ij}$  denotes the *i*-th component of  $u_j$ . Let  $\Lambda_0$  denote the diagonal matrix containing  $\lambda_{01}, \lambda_{02}, \ldots, \lambda_{0k}$ , and let U denote the orthogonal matrix  $U = (u_1, u_2, \ldots, u_k)$ . Then,

$$T_0 = U\Lambda_0 U', \quad T_0^{-1} = U\Lambda_0^{-1}U'.$$

Now consider two positive numbers  $0 < c_{min} < c_{max} < 1$ , and define the  $k \times k$  matrix T as

$$T := \alpha T_0 + \beta I_k,$$

where

$$\alpha := \frac{c_{max} - c_{min}}{\lambda_{01} - \lambda_{0k}}, \quad \beta := \frac{\lambda_{01}c_{min} - \lambda_{0k}c_{max}}{\lambda_{01} - \lambda_{0k}}$$

The eigenvalues  $\lambda_1 > \lambda_2 > \cdots > \lambda_k$  of T are then given by

$$\lambda_j = \alpha \lambda_{0j} + \beta, \quad j = 1, \dots, k,$$

so that, by construction,  $\lambda_1 = c_{max}$  and  $\lambda_k = c_{min}$ . Moreover, T possesses the same set of eigenvectors as  $T_0$ , so that

$$T = U\Lambda U', \quad T^{-1} = U\Lambda^{-1}U',$$

where

$$\Lambda = \operatorname{diag} \left( \lambda_1, \lambda_2, \ldots, \lambda_k \right).$$

Note that T is a sparse matrix (tridiagonal),  $T^2$  is also sparse (five-diagonal), but that  $T^{-1}$  is a full matrix for any dimension.

#### 7.2 Construction of S

All dimensions are integer powers of 2:

$$k := 2^j, \quad n := 2^{i+j}, \quad n_k := n/k = 2^i,$$

Let s be an integer satisfying  $0 \le s \le n_k$ . We construct the  $n \times k$  matrix S as

where the orthogonal  $k \times k$  matrix  $S_0$  is constructed as

$$S_0 := A_1 A_2 \cdots A_{k^*}, \quad 1 \le k^* \le k - 1,$$

with

$$A_j := \begin{pmatrix} I_{j-1} & 0 & 0\\ 0 & Q(\theta_j) & 0\\ 0 & 0 & I_{k-j-1} \end{pmatrix}, \quad j = 1, \dots, k^*,$$

and

$$Q(\theta) = \begin{pmatrix} \cos \theta & -\sin \theta \\ \sin \theta & \cos \theta \end{pmatrix}, \quad \theta_j = \frac{\pi}{2(k^* - j + 1)}$$

The sparsity of S is a weighted average of the sparsity of  $S_0$  and the sparsity of  $I_k$ . Since the number of nonzeros in  $S_0$  is  $k + k^*(k^* + 3)/2$ , the sparsity of S is given by

$$d_s(S) = 1 - \frac{1}{k} - \frac{k^*(k^* + 3)s}{2nk},$$

which depends on the parameters s and  $k^*$ . In most simulations we choose  $k^* = k - 1$ . For s = 0, we have maximal sparsity  $d_s(S) = 1 - 1/k$ ; for  $s = n_k$ , we have minimal sparsity  $d_s(S) = d_s(S_0) = 1 - 1/k - k^*(k^* + 3)/(2k^2)$ .

#### 7.3 Analytical solution

Given S and T, and the definition X = ST, we find

$$X'X = T'S'ST = T^{2}, \quad (X'X)^{-1} = T^{-2} = U\Lambda^{-2}U' = \sum_{j=1}^{k} \frac{1}{\lambda_{j}^{2}} u_{j}u'_{j},$$

and

$$\hat{\beta} = (X'X)^{-1}X'y = T^{-2}T'S'y = T^{-1}(S'y) = \sum_{j=1}^{k} \frac{1}{\lambda_j} u_j u'_j z,$$

where z := S'y. The estimator for  $\sigma^2$  can be found as

$$\hat{\sigma}^2 = \frac{1}{n-k} y' (I_n - X(X'X)^{-1}X') y = \frac{1}{n-k} (y'y - z'z)$$

and the condition number is

$$c := \sqrt{\frac{\lambda_1(X'X)}{\lambda_k(X'X)}} = \sqrt{\frac{\lambda_1(T^2)}{\lambda_k(T^2)}} = \frac{\lambda_1(T)}{\lambda_k(T)} = \frac{c_{max}}{c_{min}}$$

Throughout the simulation experiment we set  $c_{max} = 0.9$ . The simulations are then controlled by five parameters: i, j, c, s, and  $k^*$ , thus providing the dimensions  $k = 2^j$  and  $n = 2^{i+j}$ , the bounds  $c_{max} = 0.9$  and  $c_{min} = c_{max}/c$ , and the sparsity  $d_s$  (as a function of s and  $k^*$ ). The condition number c can be fully controlled, the sparsity  $d_s$  can be approximately controlled.

## 8 Simulation results

#### 8.1 Methods considered

We are concerned with accuracy and speed. We consider six sparse methods. The first four methods use various sparse Matlab functions. The next method is also a sparse method, but not used in Matlab. The final method is our Snaer-OLS routine. Method 1: Exact numerical inversion (Matlab). The standard way to calculate the OLS solution is to apply the explicit formulae  $\hat{\beta} = (X'X)^{-1}X'y$  and  $\widehat{\operatorname{var}}(\hat{\beta}) = \hat{\sigma}^2 (X'X)^{-1}$  with  $\hat{\sigma}^2 = (y - X\hat{\beta})'(y - X\hat{\beta})/(n-k)$ , where the matrix  $(X'X)^{-1}$  is obtained by applying the Matlab matrix inversion function.

Method 2: QR decomposition (Matlab). Matlab provides another way to calculate the OLS solution for linear regression through the "economy-size" QR decomposition available as part of the regress algorithm. We write X = QR, where Q is semi-orthogonal ( $Q'Q = I_k$ ) and R is an upper triangular  $k \times k$ matrix with positive diagonal elements; see Chatterjee and Hadi (1986), and Draper and Smith (1998, p. 94). The formulae now depend on the inversion of the triangular matrix R and are given by

$$\hat{\beta} = R^{-1}Q'y, \quad \widehat{\operatorname{var}}(\hat{\beta}) = \hat{\sigma}^2 R^{-1} (R^{-1})', \quad \hat{\sigma}^2 = \frac{y'(I_n - QQ')y}{n-k}.$$

Method 3: Least-squares minimization (Matlab). The next method considers the numerical minimization problem

$$\min_{\beta} \frac{1}{2} \|y - X\beta\|^2.$$

and uses the Matlab function *mldivide* (\) for the optimization. More specifically, the Matlab command  $X \setminus y$  uses Householder reflections to compute an orthogonal-triangular factorization XP = QR, where P is permutation matrix, Q is semi-orthogonal, and R is upper triangular. Then,

$$\hat{\beta} = X \setminus y = (X'X)^{-1}X'y = (PR'Q'QRP')^{-1}PR'Q'y$$
$$= P(R'R)^{-1}R'Q'y = P(R \setminus (Q'y)).$$

This method is thus closely related to the QR decomposition.

In order to compute the variance matrix, we require elements of the inverse of the matrix X'X. Using the same Matlab algorithm *mldivide*, the *j*-th column  $v^{(j)}$  of  $V := (X'X)^{-1}$  can be found by considering the matrix equation  $(X'X)V = I_k$  and solving the minimization problem

$$\min_{v^{(j)}} \frac{1}{2} \|e^{(j)} - X' X v^{(j)}\|^2, \quad j = 1, \dots, k,$$

where  $e^{(j)}$  denotes the *j*-th unit vector, that is the *j*-th column of  $I_k$ . For the calculation of  $\hat{\beta}$ , methods 2 and 3 give the same numerical results, but for the calculation of the elements of the variance matrix, the method of computation and hence the numerical results are different. Method 4: Conjugate gradient method (Matlab). Here we consider the normal equations

$$X'X\beta = X'y.$$

The Matlab procedure *pcg* finds a nonsingular matrix P such that  $P'X'XP = \Lambda$  (diagonal). This implies that  $(X'X)^{-1} = P\Lambda^{-1}P'$  and hence

$$\hat{\beta} = P\Lambda^{-1}P'X'y = Pa,$$

where the j-th element of a is given by

$$a_j = \frac{p'_j X' y}{p'_j X' X p_j}, \quad j = 1, \dots, k.$$

This is the so-called direct method. In practice, if we choose the conjugate vectors  $p_j$  carefully, we may not need to compute all of them to obtain a good approximation of  $\hat{\beta}$ . Another Matlab function, the generalized minimum residual method (*gmres*), is also based on solving the normal equations, and its performance will be briefly discussed when we present the results for the *pcg* method.

The variance matrix is obtained by considering the equations

$$X'Xv^{(j)} = e^{(j)}, \quad j = 1, \dots, k,$$

and applying the *pcg* routine.

Method 5: The iterative Lanczos algorithm. The Numerical Algorithms Group (NAG, 2006) library provides the subroutine f04qaf for solving large sparse OLS problems. This is an iterative method, based on the bidiagonalization procedure of Golub and Kahan (1965) and the Lanczos (1950) iteration. In essence it is the *lsqr* method of Paige and Saunders (1982a, 1982b) belonging to the family of Krylov methods; see Broyden and Vespucci (2004).

The Lanczos scheme requires a symmetric matrix, and hence we write our system as

$$\begin{pmatrix} I & X \\ X' & 0 \end{pmatrix} \begin{pmatrix} r \\ \beta \end{pmatrix} = \begin{pmatrix} y \\ 0 \end{pmatrix},$$

where r denotes the residual vector. Similarly, the variance matrix is obtained by considering the symmetric system

$$X'Xv^{(j)} = e^{(j)}, \quad j = 1, \dots, k.$$

The time taken by the routine is likely to be principally determined by the time taken in APROD, which is called twice on each iteration. APROD

performs a matrix-vector product, where the sparsity is explicitly taken into account. Although the Lanczos process will usually converge more quickly if some form of preconditioning is employed (Broyden and Vespucci, 2004, Chapter 11), we do not use preconditioners because we are investigating the performance of methods for general sparsity patterns, while preconditioners are structure-dependent.

*Method 6: Snaer-OLS.* This is our method, based on the Gould-Nocedal algorithm, and described before.

All computations were performed on a desktop personal computer, Intel Pentium 4 processor, 2.66 GHz speed, 2 GB RAM, and endowed with Microsoft Windows XP Professional (SP2). The Matlab results were obtained using Matlab version 6.5, release 13. The Fortran programs were compiled using the Compaq Visual Fortran 6.6 compiler.

#### 8.2 Accuracy

In the first two experiments, we set

$$n = 2k, \quad k = 2^{j}, \quad c = 100,000,$$

and we let j = 5, ..., 12. The total number of elements in X thus equals  $nk = 2^{2j+1}$ , which ranges from  $2^{11}$  (about two thousand) to  $2^{25}$  (over 33 million). Since  $n_k = n/k = 2$ , we have  $0 \le s \le 2$ . We consider the two extremes s = 0 and s = 2, both for  $k^* = k - 1$ .

The accuracy of the various solutions is controlled by the average relative deviation

ARD = 
$$\sqrt{\frac{1}{k} \sum_{j=1}^{k} (\hat{v}_{jj}/v_{jj} - 1)^2},$$

where  $\hat{v}_{jj}$  denotes the *j*-th diagonal element of  $\hat{V} := \hat{\sigma}^2 (X'X)^{-1}$  in one of the six algorithms, and  $v_{jj}$  denotes its true value. The ARD measure thus takes into account both the accuracy of  $\hat{\beta}$  (through  $\hat{\sigma}^2$ ) and the accuracy of  $(X'X)^{-1}$ . We have chosen the diagonal elements of the variance matrix for simplicity, but also because of their obvious practical importance. In the national accounts applications we always calculate the diagonal elements of  $(X'X)^{-1}$ , but only occasionally some of the offdiagonal elements. Controlling the accuracy of the variances is therefore our primary concern.

The accuracies are reported in Table 1 for very sparse systems (s = 0) in the top panel and less sparse systems (s = 2) in the bottom panel. In

	Matlab				NAG	Snaer-OLS	
j	1	2	3	4	5	6	sparsity
5	2.24	0.000015	2.24	2.38	4.58	2.32	0.9082
6	2.49	0.000004	2.49	2.51	22.4	2.51	0.9536
7	1.57	0.000070	1.57	1.56	174	1.76	0.9767
8	1.42	0.000006	1.42	1.38	$9,\!984,\!000$	1.40	0.9883
9	4.05	0.000031	4.05	4.01		3.97	0.9941
10	2.42	0.000015	2.42	2.43		2.39	0.9971
11	0.81	0.000056	0.81	0.80		0.63	0.9985
12						1.45	0.9993
5	0.56	0.000015	0.56	0.45	859,670	0.14	0.4248
6	0.28	0.000031	0.28	0.17	$7,\!622,\!500$	0.61	0.4617
7	0.07	0.000025	0.07	0.05	$9,\!670,\!600$	0.14	0.4807
8	0.03	0.000006	0.03	0.02		0.26	0.5419
9	0.06	0.000029	0.06	0.06		0.21	0.7435
10	0.08	0.000016	0.08	0.07		0.16	0.8744
11	0.07	0.000008	0.08	0.08		0.36	0.9412

Table 1 — Relative accuracy ARD (×10<sup>-7</sup>), n = 2k,  $k = 2^{j}$ , c = 100,000, and s = 0 (top panel) and s = 2 (bottom panel).

the top panel the sparsity increases from 90.8% to 99.9% as the dimension increases. Methods 2 (QR) is the most accurate, while the Lanczos algorithm (method 5) is very inaccurate and has not been calculated for j > 8. (The related Matlab *lsqr* function behaves similarly and does not reach acceptable accuracy for any reasonable choice of tolerances and number of iterations.) An analysis of the breakdown causes of the Lanczos method can be found in Broyden and Vespucci (2004, pp. 24–26). The other methods (1, 3, 4, and 6) are all of the same degree of accuracy, indicating that on average  $\hat{v}_{jj}/v_{jj} \approx 1.0000001$ , which seems sufficiently accurate. If the system becomes large, all methods except Snaer-OLS break down due to memory allocation restrictions. The accuracy of the *gmres* function, related to *pcg* (method 4), is comparable with *pcg*.

In the bottom panel (s = 2) there is much less sparsity, ranging only from 42.5% for j = 5 to 94.1% for j = 11. The conclusions are the same as for the sparse system. Clearly, a sparse procedure will be very inefficient (in terms of computing time) when employed in a nonsparse situation, but the accuracy seems unaffected.

The chosen accuracy measure ARD is quite robust. If instead we calculate the average squared deviation taking *all* elements of the variance matrix, this produces only a marginal increase. For example for j = 8 and s = 0 (line 4 from the top in Table 1), the alternative measure gives  $1.44 \cdot 10^{-7}$  instead of  $1.42 \cdot 10^{-7}$  for method 1 and has essentially no effect on method 3 and other methods.

#### 8.3 Tolerances

In the first three Matlab routines (exact, QR, and *mldivide*) no tolerances can be set, so that the defaults must be used.

The *pcg* algorithm was used with tolerance 1.0D-15 and the maximum number of iterations: 5000 for j < 9, 10,000 for j = 9, 20,000 for j = 10, and 120,000 for j = 11.

For NAG f04qaf the tolerances  $tol_1$  and  $tol_2$  were set to 1.0D-14 and 1.0D-15, respectively, and the routine accepts  $\hat{\beta}$  as a solution of  $X'X\beta = X'y$  if it is estimated that one of the following two conditions is satisfied:

$$||r|| \le tol_1 ||X|| \cdot ||\hat{\beta}|| + tol_2 ||y||, \qquad ||X'r|| \le tol_1 ||X|| \cdot ||r||,$$

where  $r := y - X\hat{\beta}$  is the residual vector. The parameter *conlim* is the upper limit on condition number of X, and was set to *conlim* = 1.0D+12. The parameter *itnlim* is the upper limit on the number of iterations and was set to *itnlim* = 1,000,000. Further reduction in tolerances and increase in iteration and condition numbers did not lead to higher accuracy.

For the Snaer-OLS functions the default tolerances have been used.

#### 8.4 Computing time

The computing time (in seconds) that each method requires is measured as follows. For the Matlab procedures 1–4, we employ the tic and toc functions: tic saves the current time that toc uses later to measure the elapsed time that Matlab takes to complete the required operations. For the NAG-Lanczos (5) and Snaer-OLS (6) routines we use the RTC function.

Naturally, computing time grows with the dimension. In Figure 1 we present the computing time T as a function of the dimension k in the sparse case (s = 0). The figures are plotted on a log-log scale, where  $j := \log k$ and  $\tau := \log T$  are logarithms to base 2, so that  $k = 2^j$  and  $T = 2^{\tau}$ . The relationship on this scale appears to be close to linear. This suggests that  $T \approx \text{constant} \times ((1 - d_s)nk)^{\delta}$ . We expect  $\delta > 1$ , because all elements of the matrix must be evaluated at least once. A value of  $\delta$  close to one thus indicates a near-optimal slope; the lower is  $\delta$ , the faster is the method. The Gould-Nocedal method (Snaer-OLS) is the best in this sense with  $\delta = 1.9$ .



Figure 1 — Computing time,  $k = 2^{j}$ , n = 2k, c = 100,000, and s = 0.

For large systems, say when  $k = 2^{12}$  and  $n = 2^{13}$ , exact inversion (method 1) and QR (method 2) break down due to memory allocation problems. The Lanczos algorithm (method 5) breaks down earlier, but for another reason: the accuracy becomes unacceptably low, as we have seen. For large sparse systems with j = 12 or larger (giving an X-matrix with 33.6 million or more entries), only Snaer-OLS works. For j = 11, all methods except the NAG routine work with reasonable and comparable accuracies except the QR method which is much more accurate (see Table 1). However, the computing times of the methods differ greatly. Method 4 takes 158 hours, to complete its calculations. (For the gmres function, related to pcg of method 4, the number of iterations required and thus the computation time is so large that the method becomes unusable for j > 8.) In contrast, method 3 takes 78 seconds, while Snaer-OLS's Gould-Nocedal method 6 only requires 2.1 seconds, and thus performs best for large sparse systems.

Figure 2 illustrates the performance of the methods in a less sparse situation, where s = 2. Computations have only been performed for  $j \leq 11$ . The exact (1) and QR (2) routines work best here, while the Lanczos method, again, was not able to reach sufficient accuracy. Snaer-OLS's Gould-Nocedal method is not the favorite choice now.

Let us consider the largest system, where j = 11, so that  $k = 2^{11}$  and  $n = 2^{12}$ , and compare the sparse situation (Figure 1, sparsity 94.1%) with



Figure 2 — Computing time,  $k = 2^j$ , n = 2k, c = 100,000, and s = 2.

the less sparse situation (Figure 2, sparsity 99.8%). All methods take longer in the less sparse situation. In particular, Snaer-OLS's method requires 2 seconds in the sparse situation and 26 minutes in the less sparse situation. Apparently, a sparsity of 94% is not sufficient for a sparse procedure to work well. What degree of sparsity, then, is sufficient? Our experiments suggest that a matrix should be called "sparse" when at least 98% or 99% of its elements are structural zeros. With sparsity thus defined, a sparse method will work well on a sparse matrix.

#### 8.5 Dimension limitations

All methods are constrained by computer memory limitations. When a nonsparse method is used, Matlab is unable to store an  $n \times k$  matrix when  $nk > 2^{25}$ . For example, a matrix with  $k = 2^{13}$  (about 8000) and n = 2k or with  $k = 2^{12}$  (about 4000) and n = 8k can not be stored.

This upper bound also provides a limit to the number of elements in the coordinate storage representation of a sparse matrix. Since three numbers are required to store a sparse element (row index, column index, value), the maximum number of elements in a sparse matrix in Matlab can not be more than approximately  $2^{25}/3$ . Snaer-OLS's Gould-Nocedal routine, however, works for dimensions even higher than this.

## **9** Conclusions and extensions

The Snaer program discussed in this paper calculates the posterior mean and variance of variables on some of which we have data (with precisions), on some we have prior information (with precisions), and on some prior indicator ratios (with precisions) are available. The variables must satisfy a number of exact restrictions. The system is both large and sparse.

Two aspects of the statistical and computational development are a practical procedure to solve a linear integer system, and a stable linearization routine for ratios.

We tested our numerical method to solve large sparse linear least-squares estimation problems, and found that it performs well, even when the  $n \times k$ design matrix is large  $(nk > 2^{23})$ . In fact, Snaer-OLS can handle design matrices of dimension 16,000 × 12,000, where  $nk \approx 2^{27.5}$ .

Our procedures work well if the system is sparse, and in this context we call a matrix "sparse" when at least 98% or 99% of its elements are structural zeros.

We now discuss three practical difficulties in applying our methods: how to obtain prior precisions, how to deal with skewness, and how to include dynamics in the system.

*Prior precisions.* A major advantage but also a major challenge is that we allow priors and prior precisions. The priors themselves (that is, the means of the priors) are based on expert opinion, historical data, neighboring information, or otherwise. Their precisions are more difficult to obtain.

If exact precisions are available, then we can use these. But in practice we will only have indications. For example, banking data are more precise than survey data. For practical applications we adopt a small number (say four) levels of precision, defined by the coefficient of variation (CV), that is, the ratio of the standard deviation to the mean:  $CV = \sigma/\mu$ . The coefficient of variation is a dimensionless number that allows comparison of the variation of populations that have significantly different mean values, and it is often used in the context of the normal distribution when the means are positive (as in our applications).

A problem arises when the mean is near zero, because the coefficient of variation then becomes sensitive to changes in the standard deviation; in that case we have to assign precisions on an *ad hoc* basis.

We also have to decide which values are appropriate for the four levels of precision. This depends on the situation and requires expert knowledge of the data. Recall that the variance of  $\mu$  is given by

$$\operatorname{var}(\mu) = \sigma^2 \begin{pmatrix} I \\ Q_1 \end{pmatrix} (R'R)^{-1} (I, Q_1'),$$

where, up to now, we have set  $\sigma^2 = 1$ . In fact we may estimate  $\sigma^2$  by the usual unbiased estimator as

$$\hat{\sigma}^2 = \frac{||r - R\mu_1||}{p + m_1 + m_2 - n}.$$

Including  $\sigma^2$  implies that the prior precisions only need to be assigned in *relative* terms rather than in absolute terms, thus greatly improving their credibility.

In practical applications one will usually only define prior precisions and not also correlations, which seems even more demanding. But there is no theoretical constraint to include correlations if they appear to be available.

Skewness. Our data and priors are assumed to be normal, hence symmetric. This assumption allows us to rewrite the problem in a manageable format so that we can estimate very large systems. But there is also a cost, because a number of data and priors are known to be skewed. Although we can find *ad hoc* solutions by introducing indicator ratios which provide a one-directional force, the general problem remains unresolved.

Dynamics. There is no problem in combining data from various years and define the interactions. We simply think of  $x_t$  and  $x_{t-1}$  as two different variables, linked by some ratio or linear relationship. This allows a simple form of dynamics in the system.

Finally we briefly discuss four topics that are currently under investigation as part of the Snaer program.

Sensitivity analysis. The program make it easy to assess how small changes in one assumption will work through the system and affect one or more key variables and their precisions. For example, if more precise information were available on a set of variables (say by financing a survey), then we can calculate the impact of these improved precisions *before* the survey has in fact taken place. The improved precisions of a subset of the data may or may not matter for the key variables in our system — it is hard to decide in such a large system with so many constraints. But in our program such assessments are relatively easy to perform. Aggregation and hierarchy. On the whole there is a tendency to believe that the more data the better. Certainly this is the view in national accountants circles. But is it true? How useful is it to work with large (micro) data sets when we are primarily interested in macro estimates? How much do the (often not very precise) micro data help in improving the macro estimates? Of course, it depends. But precisely on what does it depend in practical nonlinear situations? This we don't know yet.

Forecasting. The set-up and focus of this paper is estimation, not forecasting. We want to estimate last year's national accounts. In essence, however, our procedure is a forecasting (more accurately, prediction) procedure. Given all available information, we provide the best possible forecast at a particular point in time. If more information becomes available a new estimate (forecast) can be produced quickly. The same set-up and software can then also be used to produce next year's national accounts, of course primarily based on priors because data will not yet be available. This is unusual, but it will be useful in policy debates based on the national accounts.

Interaction between data collection and estimation. Data collection and data analysis are typically performed by different groups of experts. Data collection is the expertise of national accountants working in national statistics offices or in international organizations like the United Nations. Data analysis and economic modeling is the expertise of academics at universities and other research environments. How do the data collection process and the subsequent econometric analysis interact? The econometrician usually takes the data as given, but in reality the data are also based on models, and these could be closely correlated with the research question of the econometrician. This can have serious effects on the reliability and interpretation of the analysis.

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