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2.1 Gauss-Jordan Elimination

For inverting a matrix, *Gauss-Jordan elimination* is about as efficient as any other method. For solving sets of linear equations, Gauss-Jordan elimination produces *both* the solution of the equations for one or more right-hand side vectors **b**, and also the matrix inverse \mathbf{A}^{-1} . However, its principal weaknesses are (i) that it requires all the right-hand sides to be stored and manipulated at the same time, and (ii) that when the inverse matrix is *not* desired, Gauss-Jordan is three times slower than the best alternative technique for solving a single linear set (§2.3). The method's principal strength is that it is as stable as any other direct method, perhaps even a bit more stable when full pivoting is used (see below).

If you come along later with an additional right-hand side vector, you can multiply it by the inverse matrix, of course. This does give an answer, but one that is quite susceptible to roundoff error, not nearly as good as if the new vector had been included with the set of right-hand side vectors in the first instance.

For these reasons, Gauss-Jordan elimination should usually not be your method of first choice, either for solving linear equations or for matrix inversion. The decomposition methods in §2.3 are better. Why do we give you Gauss-Jordan at all? Because it is straightforward, understandable, solid as a rock, and an exceptionally good "psychological" backup for those times that something is going wrong and you think it *might* be your linear-equation solver.

Some people believe that the backup is more than psychological, that Gauss-Jordan elimination is an "independent" numerical method. This turns out to be mostly myth. Except for the relatively minor differences in pivoting, described below, the actual sequence of operations performed in Gauss-Jordan elimination is very closely related to that performed by the routines in the next two sections.

For clarity, and to avoid writing endless ellipses (\cdots) we will write out equations only for the case of four equations and four unknowns, and with three different right-hand side vectors that are known in advance. You can write bigger matrices and extend the equations to the case of $N \times N$ matrices, with M sets of right-hand side vectors, in completely analogous fashion. The routine implemented below is, of course, general.

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Elimination on Column-Augmented Matrices

Consider the linear matrix equation

$$\begin{bmatrix} a_{11} & a_{12} & a_{13} & a_{14} \\ a_{21} & a_{22} & a_{23} & a_{24} \\ a_{31} & a_{32} & a_{33} & a_{34} \\ a_{41} & a_{42} & a_{43} & a_{44} \end{bmatrix} \cdot \begin{bmatrix} \begin{pmatrix} x_{11} \\ x_{21} \\ x_{31} \\ x_{41} \end{pmatrix} \sqcup \begin{pmatrix} x_{12} \\ x_{22} \\ x_{32} \\ x_{42} \end{pmatrix} \sqcup \begin{pmatrix} x_{13} \\ x_{23} \\ x_{33} \\ x_{43} \end{pmatrix} \sqcup \begin{pmatrix} y_{11} & y_{12} & y_{13} & y_{14} \\ y_{21} & y_{22} & y_{23} & y_{24} \\ y_{31} & y_{32} & y_{33} & y_{34} \\ y_{41} & y_{42} & y_{43} & y_{44} \end{bmatrix}$$

$$= \left[\begin{pmatrix} b_{11} \\ b_{21} \\ b_{31} \\ b_{41} \end{pmatrix} \sqcup \begin{pmatrix} b_{12} \\ b_{22} \\ b_{32} \\ b_{42} \end{pmatrix} \sqcup \begin{pmatrix} b_{13} \\ b_{23} \\ b_{33} \\ b_{43} \end{pmatrix} \sqcup \begin{pmatrix} 1 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 \\ 0 & 0 & 1 & 0 \\ 0 & 0 & 0 & 1 \end{pmatrix} \right]$$
(2.1.1)

Here the raised dot (\cdot) signifies matrix multiplication, while the operator \sqcup just signifies column augmentation, that is, removing the abutting parentheses and making a wider matrix out of the operands of the \sqcup operator.

It should not take you long to write out equation (2.1.1) and to see that it simply states that x_{ij} is the ith component (i=1,2,3,4) of the vector solution of the jth right-hand side (j=1,2,3), the one whose coefficients are $b_{ij}, i=1,2,3,4$; and that the matrix of unknown coefficients y_{ij} is the inverse matrix of a_{ij} . In other words, the matrix solution of

$$[\mathbf{A}] \cdot [\mathbf{x}_1 \sqcup \mathbf{x}_2 \sqcup \mathbf{x}_3 \sqcup \mathbf{Y}] = [\mathbf{b}_1 \sqcup \mathbf{b}_2 \sqcup \mathbf{b}_3 \sqcup \mathbf{1}] \tag{2.1.2}$$

where A and Y are square matrices, the b_i 's and x_i 's are column vectors, and 1 is the identity matrix, simultaneously solves the linear sets

$$\mathbf{A} \cdot \mathbf{x}_1 = \mathbf{b}_1 \qquad \mathbf{A} \cdot \mathbf{x}_2 = \mathbf{b}_2 \qquad \mathbf{A} \cdot \mathbf{x}_3 = \mathbf{b}_3 \tag{2.1.3}$$

and

$$\mathbf{A} \cdot \mathbf{Y} = \mathbf{1} \tag{2.1.4}$$

Now it is also elementary to verify the following facts about (2.1.1):

- Interchanging any two rows of A and the corresponding rows of the b's and of 1, does not change (or scramble in any way) the solution x's and Y. Rather, it just corresponds to writing the same set of linear equations in a different order.
- Likewise, the solution set is unchanged and in no way scrambled if we replace any row in **A** by a linear combination of itself and any other row, as long as we do the same linear combination of the rows of the **b**'s and **1** (which then is no longer the identity matrix, of course).
- Interchanging any two columns of A gives the same solution set only
 if we simultaneously interchange corresponding rows of the x's and of
 Y. In other words, this interchange scrambles the order of the rows in
 the solution. If we do this, we will need to unscramble the solution by
 restoring the rows to their original order.

Gauss-Jordan elimination uses one or more of the above operations to reduce the matrix $\bf A$ to the identity matrix. When this is accomplished, the right-hand side becomes the solution set, as one sees instantly from (2.1.2).

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Pivoting

In "Gauss-Jordan elimination with no pivoting," only the second operation in the above list is used. The first row is divided by the element a_{11} (this being a trivial linear combination of the first row with any other row — zero coefficient for the other row). Then the right amount of the first row is subtracted from each other row to make all the remaining a_{i1} 's zero. The first column of $\bf A$ now agrees with the identity matrix. We move to the second column and divide the second row by a_{22} , then subtract the right amount of the second row from rows 1, 3, and 4, so as to make their entries in the second column zero. The second column is now reduced to the identity form. And so on for the third and fourth columns. As we do these operations to $\bf A$, we of course also do the corresponding operations to the $\bf b$'s and to $\bf 1$ (which by now no longer resembles the identity matrix in any way!).

Obviously we will run into trouble if we ever encounter a zero element on the (then current) diagonal when we are going to divide by the diagonal element. (The element that we divide by, incidentally, is called the *pivot element* or *pivot*.) Not so obvious, but true, is the fact that Gauss-Jordan elimination with no pivoting (no use of the first or third procedures in the above list) is numerically unstable in the presence of any roundoff error, even when a zero pivot is not encountered. You must *never* do Gauss-Jordan elimination (or Gaussian elimination, see below) without pivoting!

So what *is* this magic pivoting? Nothing more than interchanging rows (*partial pivoting*) or rows and columns (*full pivoting*), so as to put a particularly desirable element in the diagonal position from which the pivot is about to be selected. Since we don't want to mess up the part of the identity matrix that we have already built up, we can choose among elements that are both (i) on rows below (or on) the one that is about to be normalized, and also (ii) on columns to the right (or on) the column we are about to eliminate. Partial pivoting is easier than full pivoting, because we don't have to keep track of the permutation of the solution vector. Partial pivoting makes available as pivots only the elements already in the correct column. It turns out that partial pivoting is "almost" as good as full pivoting, in a sense that can be made mathematically precise, but which need not concern us here (for discussion and references, see [1]). To show you both variants, we do full pivoting in the routine in this section, partial pivoting in §2.3.

We have to state how to recognize a particularly desirable pivot when we see one. The answer to this is not completely known theoretically. It is known, both theoretically and in practice, that simply picking the largest (in magnitude) available element as the pivot is a very good choice. A curiosity of this procedure, however, is that the choice of pivot will depend on the original scaling of the equations. If we take the third linear equation in our original set and multiply it by a factor of a million, it is almost guaranteed that it will contribute the first pivot; yet the underlying solution of the equations is not changed by this multiplication! One therefore sometimes sees routines which choose as pivot that element which *would* have been largest if the original equations had all been scaled to have their largest coefficient normalized to unity. This is called *implicit pivoting*. There is some extra bookkeeping to keep track of the scale factors by which the rows would have been multiplied. (The routines in §2.3 include implicit pivoting, but the routine in this section does not.)

Finally, let us consider the storage requirements of the method. With a little reflection you will see that at every stage of the algorithm, *either* an element of **A** is

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Here is the routine for Gauss-Jordan elimination with full pivoting:

₹

```
#include <math.h>
#include "nrutil.h"
#define SWAP(a,b) {temp=(a);(a)=(b);(b)=temp;}
void gaussj(float **a, int n, float **b, int m)
Linear equation solution by Gauss-Jordan elimination, equation (2.1.1) above. a [1..n] [1..n]
is the input matrix. b[1..n][1..m] is input containing the m right-hand side vectors. On
output, a is replaced by its matrix inverse, and b is replaced by the corresponding set of solution
vectors.
    int *indxc,*indxr,*ipiv;
    int i,icol,irow,j,k,l,ll;
    float big,dum,pivinv,temp;
    indxc=ivector(1,n);
                                           The integer arrays ipiv, indxr, and indxc are
    indxr=ivector(1,n);
                                               used for bookkeeping on the pivoting.
    ipiv=ivector(1,n);
    for (j=1;j<=n;j++) ipiv[j]=0;</pre>
    for (i=1;i<=n;i++) {
                                           This is the main loop over the columns to be
        big=0.0;
                                               reduced.
        for (j=1;j<=n;j++)
                                           This is the outer loop of the search for a pivot
            if (ipiv[j] != 1)
                for (k=1;k\leq n;k++) {
                     if (ipiv[k] == 0) {
                         if (fabs(a[j][k]) >= big) {
                             big=fabs(a[j][k]);
                             irow=j;
                             icol=k:
                         }
                     } else if (ipiv[k] > 1) nrerror("gaussj: Singular Matrix-1");
                }
        We now have the pivot element, so we interchange rows, if needed, to put the pivot
        element on the diagonal. The columns are not physically interchanged, only relabeled:
        indxc[i], the column of the ith pivot element, is the ith column that is reduced, while
        indxr[i] is the row in which that pivot element was originally located. If indxr[i] \neq
        indxc[i] there is an implied column interchange. With this form of bookkeeping, the
        solution b's will end up in the correct order, and the inverse matrix will be scrambled
        by columns.
        if (irow != icol) {
            for (l=1;l<=n;l++) SWAP(a[irow][1],a[icol][1])</pre>
            for (l=1;l<=m;l++) SWAP(b[irow][1],b[icol][1])
                                           We are now ready to divide the pivot row by the
        indxr[i]=irow;
        indxc[i]=icol;
                                               pivot element, located at irow and icol.
        if (a[icol][icol] == 0.0) nrerror("gaussj: Singular Matrix-2");
        pivinv=1.0/a[icol][icol];
        a[icol][icol]=1.0:
        for (l=1;l<=n;l++) a[icol][l] *= pivinv;
        for (l=1;l<=m;l++) b[icol][l] *= pivinv;</pre>
```

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```
for (ll=1;ll<=n;ll++)
                                      Next, we reduce the rows...
        if (ll != icol) {
                                      ...except for the pivot one, of course.
            dum=a[11][icol];
            a[11][icol]=0.0;
            for (l=1;l<=n;l++) a[l1][l] -= a[icol][l]*dum;
            for (l=1;l<=m;l++) b[l1][l] -= b[icol][l]*dum;
This is the end of the main loop over columns of the reduction. It only remains to unscram-
ble the solution in view of the column interchanges. We do this by interchanging pairs of
columns in the reverse order that the permutation was built up.
for (l=n;l>=1;l--) {
    if (indxr[1] != indxc[1])
        for (k=1;k<=n;k++)
            SWAP(a[k][indxr[1]],a[k][indxc[1]]);
                                      And we are done.
free_ivector(ipiv,1,n);
free_ivector(indxr,1,n);
free_ivector(indxc,1,n);
```

Row versus Column Elimination Strategies

The above discussion can be amplified by a modest amount of formalism. Row operations on a matrix A correspond to pre- (that is, left-) multiplication by some simple matrix R. For example, the matrix R with components

$$R_{ij} = \begin{cases} 1 & \text{if } i = j \text{ and } i \neq 2, 4\\ 1 & \text{if } i = 2, j = 4\\ 1 & \text{if } i = 4, j = 2\\ 0 & \text{otherwise} \end{cases}$$
 (2.1.5)

effects the interchange of rows 2 and 4. Gauss-Jordan elimination by row operations alone (including the possibility of *partial* pivoting) consists of a series of such left-multiplications, yielding successively

$$\mathbf{A} \cdot \mathbf{x} = \mathbf{b}$$

$$(\cdots \mathbf{R}_3 \cdot \mathbf{R}_2 \cdot \mathbf{R}_1 \cdot \mathbf{A}) \cdot \mathbf{x} = \cdots \mathbf{R}_3 \cdot \mathbf{R}_2 \cdot \mathbf{R}_1 \cdot \mathbf{b}$$

$$(1) \cdot \mathbf{x} = \cdots \mathbf{R}_3 \cdot \mathbf{R}_2 \cdot \mathbf{R}_1 \cdot \mathbf{b}$$

$$\mathbf{x} = \cdots \mathbf{R}_3 \cdot \mathbf{R}_2 \cdot \mathbf{R}_1 \cdot \mathbf{b}$$

$$(2.1.6)$$

The key point is that since the \mathbf{R} 's build from right to left, the right-hand side is simply transformed at each stage from one vector to another.

Column operations, on the other hand, correspond to post-, or right-, multiplications by simple matrices, call them C. The matrix in equation (2.1.5), if right-multiplied onto a matrix A, will interchange A's second and fourth *columns*. Elimination by column operations involves (conceptually) inserting a column operator, *and also its inverse*, between the matrix A and the unknown vector \mathbf{x} :

$$\mathbf{A} \cdot \mathbf{x} = \mathbf{b}$$

$$\mathbf{A} \cdot \mathbf{C}_1 \cdot \mathbf{C}_1^{-1} \cdot \mathbf{x} = \mathbf{b}$$

$$\mathbf{A} \cdot \mathbf{C}_1 \cdot \mathbf{C}_2 \cdot \mathbf{C}_2^{-1} \cdot \mathbf{C}_1^{-1} \cdot \mathbf{x} = \mathbf{b}$$

$$(\mathbf{A} \cdot \mathbf{C}_1 \cdot \mathbf{C}_2 \cdot \mathbf{C}_3 \cdot \dots) \cdots \mathbf{C}_3^{-1} \cdot \mathbf{C}_2^{-1} \cdot \mathbf{C}_1^{-1} \cdot \mathbf{x} = \mathbf{b}$$

$$(1) \cdots \mathbf{C}_3^{-1} \cdot \mathbf{C}_2^{-1} \cdot \mathbf{C}_1^{-1} \cdot \mathbf{x} = \mathbf{b}$$

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which (peeling of the C^{-1} 's one at a time) implies a solution

$$\mathbf{x} = \mathbf{C}_1 \cdot \mathbf{C}_2 \cdot \mathbf{C}_3 \cdots \mathbf{b} \tag{2.1.8}$$

Notice the essential difference between equation (2.1.8) and equation (2.1.6). In the latter case, the \mathbf{C} 's must be applied to \mathbf{b} in the *reverse order* from that in which they become known. That is, they must all be stored along the way. This requirement greatly reduces the usefulness of column operations, generally restricting them to simple permutations, for example in support of full pivoting.

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2.2 Gaussian Elimination with Backsubstitution

The usefulness of Gaussian elimination with backsubstitution is primarily pedagogical. It stands between full elimination schemes such as Gauss-Jordan, and triangular decomposition schemes such as will be discussed in the next section. Gaussian elimination reduces a matrix not all the way to the identity matrix, but only halfway, to a matrix whose components on the diagonal and above (say) remain nontrivial. Let us now see what advantages accrue.

Suppose that in doing Gauss-Jordan elimination, as described in $\S 2.1$, we at each stage subtract away rows only *below* the then-current pivot element. When a_{22} is the pivot element, for example, we divide the second row by its value (as before), but now use the pivot row to zero only a_{32} and a_{42} , not a_{12} (see equation 2.1.1). Suppose, also, that we do only partial pivoting, never interchanging columns, so that the order of the unknowns never needs to be modified.

Then, when we have done this for all the pivots, we will be left with a reduced equation that looks like this (in the case of a single right-hand side vector):

$$\begin{bmatrix} a'_{11} & a'_{12} & a'_{13} & a'_{14} \\ 0 & a'_{22} & a'_{23} & a'_{24} \\ 0 & 0 & a'_{33} & a'_{34} \\ 0 & 0 & 0 & a'_{44} \end{bmatrix} \cdot \begin{bmatrix} x_1 \\ x_2 \\ x_3 \\ x_4 \end{bmatrix} = \begin{bmatrix} b'_1 \\ b'_2 \\ b'_3 \\ b'_4 \end{bmatrix}$$
 (2.2.1)

Here the primes signify that the a's and b's do not have their original numerical values, but have been modified by all the row operations in the elimination to this point. The procedure up to this point is termed *Gaussian elimination*.

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