#### A few words about gcov. This command tells us: Profiling on a higher level • how often each line of code executes Most unix systems have prof and gprof which can be used to find the most time consuming routines. gcov (Linux) (tcov Sun) • what lines of code are actually executed can find the loops (statements), in a routine, that are executed Compile without optimization. It works only with gcc. So it most frequently. should work with g95 and gfortran as well. There may, howman prof, man gprof, man gcov for details. ever, be problems with different versions of gcc and the gcclibraries. See the web-page for the assignment for the latest This is how you use gprof on the student system. details. The flags are not standardised, so you have to read the documentation, as usual. To use gcov on the student system (not Intel in this case) one ifort -03 -qp prog.f90 sub.f90 should be able to type: icc -03 -qp prog.c sub.f90 g95 -fprofile-arcs -ftest-coverage prog.f90 sub.f90 ./a.out gfortran -03 -pg prog.f90 sub.f90 -O3 -pg prog.f90 sub.f90 g95 gcov prog.f90 creates prog.f90.gcov gcc -03 -pg prog.c sub.c gcov sub.f90 creates sub.f90.gcov -03 -pg prog.cc sub.c a++ less prog.f90.gcov etc. ./a.out produces gmon.out gprof and for C One can use other options, of course, and have more than two gcc -fprofile-arcs -ftest-coverage prog.c sub.c files. One should link with the profiling options as well since it similarly for gfortran and g++. may include profiled libraries. Profiling disturbs the run; it takes more time. The Intel compilers have support for "Profile-guided Optimization", i.e. the information from the profiled run can be used by the compiler (the second time you compile) to generate more efficient code. 133 134 Example: Arpack, a package for solving large and sparse eigen-Each sample counts as 0.01 seconds. value problems, $Ax = \lambda x$ and $Ax = \lambda Bx$ . I fetched a comself % cumulative self total pressed tar-file, unpacked, read the README-file, edited the time seconds seconds calls s/call s/call name configuration file, and compiled using make. After having cor-79.10 8.10 8.10 322 0.03 0.03 dgemv\_ rected a few Makefiles everything worked. I then recompiled 8.50 8.97 0.87 60 0.01 0.01 dger\_ using the compiler options for gprof and tcov (on a Sun; I have 4.10 9.39 0.42 58 0.01 0.01 dgttrs\_ not run this one the AMD-system). 9.72 3.22 0.33 519 0.00 0.00 dcopy\_ 2.25 9.95 0.23 215 0.00 0.00 dnrm2 I used the f90-compiler even though Arpack is written in For-0.05 562 0.00 \_\_open 0.49 10.00 0.00 tran77. (There is also Arpack++, a collection of classes that ... lots of lines deleted ... offers C++ programmers an interface to Arpack.) 0.00 0.00 1 0.00 10.24 10.14 main ... lots of lines deleted ... First gprof: 0.00 0.00 1 0.00 0.00 strchr 10.24 % gprof | less (1662 lines, less is a pager) name is the name of the routine (not the source file). The Sunor compiler converts the routine name to lower case and adds . % gprof | more (or m with alias m more) **\_open** is a system (compiler?) routine. (I have alias m less) The columns are: or % gprof > file\_name (emacs file\_name, for example) % time the percentage of the total running time of the program etc. used by this function. Not the one it calls, look at main. The first part of the output is the flat profile, such a profile can cumulative seconds a running sum of the number of seconds be produced by prof as well. Part of it, in compressed form, accounted for by this function and those listed above it. comes on the next page. The flat profile may give a sufficient

**self seconds** the number of seconds accounted for by this function alone. This is the major sort for this listing.

calls the number of times this function was invoked, if this function is profiled, else blank.

**self ms/call** the average number of milliseconds spent in this function per call, if this function is profiled, else blank.

total ms/call the average number of milliseconds spent in this function and its descendents per call, if this function is profiled, else blank. Note main.

amount of information.

dgemv is a BLAS routine, double general matrix vector multiply:

dgemv - perform one of the matrix-vector operations
y := alpha\*A\*x + beta\*y or y := alpha\*A'\*x + beta\*y

I have compiled the Fortran code instead of using a faster performance library so we can look at the source code.

Let us run tcov on dgemv.

Part of the output (compressed):

		• • •	
168 ->		DO 60	, J = 1, N
4782 ->		IF	( X(JX).NE.ZERO )THEN
4740 ->			TEMP = ALPHA*X(JX)
			DO 50, $I = 1$ , M
77660160 ->			Y(I) = Y(I) + TEMP*A(I, J)
	50		CONTINUE
		EN	D IF
4782 ->		JX	= JX + INCX
	60	CONTI	NUE
		Top 10	Blocks
		Line	Count
		211	77660160
		238	50519992
		177	871645

Note that this code is very poor. Never use the simple Fortran BLAS- or Lapack routines supplied with some packages. One lab deals with this issue.

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# Profiling in Matlab

Matlab has a built-in profiling tool. help profile for more details. Start Matlab (must use the GUI).

>> profile on >> run % The assignment Elapsed time is 1.337707 seconds. Elapsed time is 13.534952 seconds. >> profile report % in mozilla or netscape >> profile off

You can start the profiler using the GUI as well

(click in "Profiler" using "Desktop" under the main meny). The output comes in a new window and contains what looks like the flat profile from gprof.

One can see the details in individual routines by clicking on the routine under Function Name. This produces a gcov-type of listing. It contains the number of times a line was executed and the time it took.

# More about gprof

**gprof** produces a call graph as well. It shows, for each function, which functions called it, which other functions it called, and how many times. There is also an estimate of how much time was spent in the subroutines called by each function. This list is edited.

index	%time	self	children	n called	name
					rm lines
		0.01	10.13	1/1	main [1]
[3]	99.0	0.01	10.13	1	MAIN_ [3]
		0.00	7.19	59/59	dsaupd_ [5]
		0.00	2.45	1/1	dseupd_ [8]
		0.42	0.00	58/58	dgttrs_ [14]
• • •	lines (	delete	d		
		0.83	0.00	33/322	dsapps_ [11]
		1.48	0.00	59/322	dlarf_ [9]
		5.79	0.00	230/322	dsaitr_ [7]
[4]	79.1	8.10	0.00	322	dgemv_ [4]
		0.00	0.00	1120/3179	lsame_ [50]

Each routine has an index (see table at the end) and is presented between ---lines. 8.10s was spent in dgemv itself, 79.1% of total (including calls from dgemv). dsapps, dlarf, dsaitr (parents) called dgemv which in turn called lsame, a child. dsapps made 33 out of 322 calls and dgemv took 0.83s for the calls. dgemv called lsame 1120 of 3179 times, which took no measurable time (self).

children: For dgemv it is the total amount of time spent in all its children (lsame). For a parent it is the amount of that time that was propagated, from the function's children (lsame), into this parent. For a child it is the amount of time that was propagated from the child's children to dgemv.

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# Using Lapack from Fortran and C

Use Lapack to solve a problem like:

1	$^{-1}$	$^{-2}$	-3	-4		-9
1	1	-1	-2	-3		-4
<b>2</b>	1	1	$^{-1}$	$^{-2}$	x =	1
3	<b>2</b>	1	1	-1		6
4	3	<b>2</b>	1	1		11

The solution is the vector of ones. We use the Lapack-routine dgesv from Lapack. Here is a man-page:

### NAME

DGESV - compute the solution to a real system of linear equations A \* X = B,

#### SYNOPSIS

 SUBROUTINE DGESV( N, NRHS, A, LDA, IPIV, B, LDB, INFO )

 INTEGER
 INFO, LDA, LDB, N, NRHS

 INTEGER
 IPIV( \* )

 DOUBLE
 PRECISION A( LDA, \* ), B( LDB, \* )

#### PURPOSE

DGESV computes the solution to a real system of linear equations A \* X = B, where A is an N-by-N matrix and X and B are N-by-NRHS matrices. The LU decomposition with partial pivoting and row interchanges is used to factor A as A = P \* L \* U, where P is a permutation matrix, L is unit lower triangular, and U is upper triangular. The factored form of A is then used to solve the system of equations A \* X = B.

# ARGUMENTS

N (input) INTEGER

The number of linear equations, i.e., the order of the matrix A.  $N \ge 0$ .

```
NRHS
        (input) INTEGER
                                                               In Fortran90, but using the F77 interface, and F77-type
        The number of right hand sides, i.e., the number
                                                               declarations (to get shorter lines) this may look like:
        of columns of the matrix B. NRHS >= 0.
                                                               program main
                                                                 integer, parameter :: n = 10, lda = n, &
        (input/output) DOUBLE PRECISION array, dimension
Α
                                                                                         ldb = n, nrhs = 1
        (LDA,N) On entry, the N-by-N coefficient matrix
                                                                                     :: info, row, col, ipiv(n)
                                                                 integer
        A. On exit, the factors L and U from the
                                                                 double precision
                                                                                    :: A(lda, n), b(ldb)
        factorization A = P*L*U; the unit diagonal
        elements of L are not stored.
                                                                 do col = 1, n
                                                                   do row = 1, n
LDA
        (input) INTEGER
                                                                     A(row, col) = row - col
        The leading dimension of the array A.
                                                                   end do
        LDA >= max(1,N).
                                                                   A(col, col) = 1.0d0
                                                                   b(col)
                                                                               = 1 + (n * (2 * col - n - 1)) / 2
        (output) INTEGER array, dimension (N)
IPIV
                                                                 end do
        The pivot indices that define the permutation
        matrix P; row i of the matrix was interchanged
                                                                 call dgesv ( n, nrhs, A, lda, ipiv, b, ldb, info )
        with row IPIV(i).
                                                                 if (info == 0) then
        (input/output) DOUBLE PRECISION array, dimension
в
                                                                   print*, "Maximum error = ", maxval(abs(b - 1.0d0))
        (LDB,NRHS) On entry, the N-by-NRHS matrix of
                                                                 else
        right hand side matrix B. On exit, if INFO = 0,
                                                                   print*, "Error in dgesv: info = ", info
        the N-by-NRHS solution matrix X.
                                                                 end if
LDB
        (input) INTEGER
                                                               end program main
        The leading dimension of the array B.
        LDB >= max(1,N).
                                                               % Compile and link, somehow, to Lapack
                                                               % a.out
INFO
        (output) INTEGER
                                                                Maximum error = 4.218847493575595E-15
        = 0: successful exit
        < 0: if INFO = -i, the i-th argument had an
                                                               Where can we find dgesv? There are several options. Fetching
                                                               the Fortran-code from Netlib, using a compiled (optimized)
               illegal value
                                                               library etc. One of the assignments, Lapack (Uniprocessor
        > 0: if INFO = i, U(i,i) is exactly zero. The
               factorization has been completed, but the
                                                               optimization), deals with these questions.
               factor U is exactly singular, so the
               solution could not be computed.
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                                                                                          142
The following optimized libraries contain Lapack and BLAS (and
                                                               #include <math.h>
perhaps routines for fft, sparse linear algebra, etc. as well).
                                                               #include <stdio.h>
                                                               #define N 10
 • AMD: ACML (AMD Core Math Library).
 • Intel: MKL (Intel Math Kernel library).
                                                               #ifdef __cplusplus
                                                                                                 /* For C++ */
                                                               extern "C" void
 • SGI: complib.sgimath (Scientific and Mathematical Library).
                                                               #else
 • IBM: ESSL (Engineering and Scientific Subroutine Library).
                                                                                                 /* For C */
                                                               extern void
                                                               #endif
 • Sun: Sunperf (Sun Performance Library).
                                                                  dgesv_(int *, int *, double *, int *, int[],
There may be parallel versions.
                                                                         double[], int *, int *);
                                                               /*
Now for C and C++
                                                                * int [] or int *. double [][] is NOT OK but
                                                                * double [][10] is, provided we
Fairly portable (do not use local extensions of the compiler).
                                                                * call dgesv_ with A and not &A[0][0].
Think about: In C/C++
                                                                */
 • matrices are stored by row (not by column as in Fortran)
                                                               int main()

    matrices are indexed from zero

                                                               {
                                                                 int
                                                                          n = N, lda = N, ldb = N, nrhs = 1,
 • call by reference for arrays, call by value for scalars
                                                                           info, row, col, ipiv[_N];
 • the Fortran compiler MAY add an underline to the name
                                                                 double
                                                                          A[_N][_N], b[_N], s, max_err;
 • you may have to link with Fortran libraries
                                                                 /* Make sure you have the correct mix of types. */
   (mixing C and Fortran I/O may cause problems, for example)
                                                                 printf("sizeof(int) = %d\n", sizeof(int));
 • C++ requires an extern-declaration, in C you do not have
   to supply it (but do)
                                                                 /* Indexing from zero. */
 • make sure that C and Fortran types are compatible (number
                                                                 for (col = 0; col < n; col++) {
   of bytes)
                                                                   for (row = 0; row < n; row++)
                                                                     A[col][row] = row - col; /* Note TRANSPOSE */
 • some systems have C-versions of Lapack
                                                                   b[col] = 1 + (n * (1 + 2 * col - n)) / 2;
In the example below I have linked with the Fortran-version since
                                                                   A[col][col] = 1;
not all systems have C-interfaces. Make sure not to call dgesv
                                                                 }
from C on the Sun, if you want the Fortran-version (dgesv gives
you the C-version).
```

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```
/* Note underline and & for the scalar types.
                                                                                        Java
   * &A[0][0] not to get a
                                                               It is possible to mix Java with other languages using JNI, the
   * conflict with the prototype.
                                                               Java Native Interface. Wikipedia is a good starting point (look
                                                               for ini).
  dgesv_(&n, &nrhs, &A[0][0], &lda, ipiv, b,
         &ldb, &info);
                                                               Here are a few words on Java.
  if (info) {
                                                               % cat test.java
    printf("Error in dgesv: info = %d\n", info);
                                                               public class test {
    return 1;
                                                                 public static void main (String[] args) {
  } else {
                                                                   int n = 10;
    \max_{err} = 0.0;
                                                                   double[] a = new double[n];
    for (row = 0; row < n; row++) {
      s = fabs(b[row] - 1.0);
                                                                   for(int j = 0; j < n; j++)</pre>
      if (s > max_err)
                                                                     a[j] = j;
        max err = s:
                                                                   System.out.println("a[n-1] = " + a[n-1]);
    printf("Maximum error = %e\n", max err);
                                                                 }
    return 0;
                                                               }
  }
}
                                                               % javac test.java
                                                                                         Produces test.class
                                                               % java test
                                                                                         Execute (can optimize, later...)
On a Sun. See the lab for AMD.
                                                               a[n-1] = 9.0
                                                               javac produces the file test.class containing the bytecode.
% cc -fast extern.c -xlic lib=sunperf
% a.out
                                                               java is the Java interpreter that reads and executes test.class.
sizeof(int) = 4
                                                               We can study the bytecode (instructions) using javap, the Java
Maximum error = 4.218847e-15
                                                               class file disassembler. The interpreter uses a stack and has local
                                                               variables; I have called them var_1 etc. To make the bytecode
% CC -fast extern.c -xlic_lib=sunperf
                                                               easier to read I have used our variable names. Half of the listing
% a.out
                                                               (mostly dealing with the print, has been deleted). I have not
sizeof(int) = 4
                                                               printed all the pops from the stack.
Maximum error = 4.218847e-15
% a.out
           If you call dgesv and not dgesv_
                                                               See Wikipedia (java bytecode) for more details.
sizeof(int) = 4
** On entry to DGESV , parameter number 1 has an
   illegal value. Error in_{145} dgesv: info = -1
                                                                                          146
% javap -verbose test
                                                               One can profile Java programs. from man java:
                                                               -Xprof
public static void main(java.lang.String[]);
                                                                 Profiles the running program, and sends profiling
Code:
                                                                 data to standard output. This option is provided
  0: bipush 10
                       10
                             -> stack
                                                                 as a utility that is useful in program development
  2: istore 1
                       stack -> var 1
                                                                 and is not intended to be be used in production
  3: iload 1
                       var_1 -> stack
                                                                 systems.
  4: newarray double create double a[10], &a[0]->stack
  6: astore 2
                       &a[0] -> var 2
                                                               -Xrunhprof[:help][:suboption=value,...]
  7: iconst 0
                       0 -> stack
                                                                 Enables cpu, heap, or monitor profiling. This
                       0 -> var_3 (corresponds to j)
  8: istore 3
                                                                 option is typically followed by a list of
                                                                 comma-separated suboption=value pairs. Run the
  9: iload_3
                       j -> stack
                                                                 command java -Xrunhprof:help to obtain a
 10: iload 1
                       n -> stack
                                                                 list of suboptions and their default values.
 11: if_icmpge 25
                       if ( j \ge n ) goto line 11+25
 14: aload 2
                       &a[0] -> stack
 15: iload 3
                       j -> stack
 16: iload_3
                       j -> stack (used as index)
 17: i2d
                       double(j) -> stack
 18: dastore
                       a[j] = double(j), "index reg"
 19: iinc3, 1
                       j++
 22: goto9
                       goto line 9:
 54:return
}
To speed things up the bytecode interpreter (java) often uses
a JIT (Just In Time) technique. A JIT compiler converts all of
the bytecode into native machine code just as a Java program
is run. This results in run-time speed improvements over code
that is interpreted by a Java virtual machine.
java -client test or
java -server test (usually much faster; default).
```

# Interfacing Matlab with C

It is not uncommon that we have a program written in C (or Fortran) and need to communicate between the program and Matlab.

The simplest (but not the most efficient) way the fix the communication is to use ordinary text files. This is portable and cannot go wrong (in any major way). The drawback is that it may be a bit slow and that we have to convert between the internal binary format and text format. We can execute programs by using the unix-command (or ! or system).

One can do more, however:

- Reading and writing binary MAT-files from C
- Calling Matlab as a function (Matlab engine)
- Calling a C- or Fortran-function from Matlab (using MEXfiles, compiled and dynamically linked C- or Fortran-routines)

In the next few pages comes a short example on how to use MEX-files.

#### MEX-files

Let us write a C-program that can be called as a Matlab-function. The MEX-routine will call a band solver, written in Fortran, from Lapack for solving an Ax=b-problem. The routine uses a Cholesky decomposition, where A is a banded, symmetric and positive definite matrix.

b contains the right hand side(s) and x the solution(s). I fetched the routines from www.netlib.org.

Matlab has support for solving unsymmetric banded systems, but has no special routines for the positive definite case.

We would call the function by typing:

>> [x, info] = bandsolve(A, b);

where A stores the matrix in compact form. info returns some status information (A not positive definite, for example).

bandsolve can be an m-file, calling a MEX-file. Another alternative is to let **bandsolve** be the MEX-file. The first alternative is suitable when we need to prepare the call to the MEX-file or clean up after the call.

The first alternative may look like this:

function [x, info] = bandsolve(A, b) A\_tmp = A; % copy A b\_tmp = b; % copy b % Call the MEX-routine [x, info] = bandsolve\_mex(A\_tmp, b\_tmp);

I have chosen to make copies of A and b. The reason is that the Lapack-routine replaces A with the Cholesky factorization and b by the solution. This is not what we expect when we program in Matlab. If we have really big matrices, and if we do not need A and b afterwards we can skip the copy (although the Matlab-documentation says that it "may produce undesired side effects").

I will show the code for the second case where we call the MEXfile directly. Note that we use the file name, bandsolve, when invoking the function. There should always be a  ${\tt mexFunction}$  in the file, which is the entry point. This is similar to a C-program, there is always a main-routine.

It is possible to write MEX-files in Fortran, but is more natural to use C.

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<pre>First some details about how to store the matrix (for the band solver). Here an example where we store the lower triangle. The dimension is six and the number of sub- (and super-) diagonals is two. all a22 a33 a44 a55 a66 a21 a32 a43 a54 a65 * a31 a42 a53 a64 * * Array elements marked * are not used by the routine. The Fortran-routine, dpbsv, is called the following way: call dpbsv( uplo, n, kd, nB, A, lda, B, ldb, info ) where uplo = 'U': Upper triangle of A is stored 'L': Lower triangle of A is stored 'L': Lower triangle of A is stored 'L': Lower triangle of A is stored 'L': anumber of sub-diagonals nB = number of right hand sides (in B) A = packed form of A lda = leading dimension of B info = 0, successful exit &lt; 0, if info = -i, the i-th argument had an illegal value &gt; 0, if info = i, the leading minor of order i of A is not positive definite, so the factorization could not be completed, and the solution has not been computed. Here comes bandsolve.c (I am using C++-style comments):</pre>	<pre>#include <math.h> // For Matlab #include "mex.h" void dpbsv_(char *, int *, int *, int *, double *,</math.h></pre>

```
// Create a matrix for the return argument
                                                                Some comments:
  // and for A. dpbsv destroys A and b).
                                                                nrhs is the number of input arguments to the MEX-routine.
  // Should check the return status.
                                                                prhs is an array of pointers to input arguments. prhs[0] points
  plhs[0]=mxCreateDoubleMatrix(b_rows, b_cols, mxREAL);
                                                                to a so-called, mxArray, a C-struct containing size-information
  plhs[1]=mxCreateDoubleMatrix(1, 1, mxREAL);
                                                                and pointers to the matrix-elements.
  A_tmp =mxCreateDoubleMatrix(A_rows, A_cols, mxREAL);
                                                                prhs[0] corresponds to the first input variable, A etc.
  px = mxGetPr(plhs[0]);
                               // Solution x
                                                                Since one should not access the member-variables in the struct
 pA = mxGetPr(prhs[0]);
                               // A
                                                                directly, there are routines to extract size and elements.
  pA_tmp = mxGetPr(A_tmp);
                              // temp for A
                                                                A_rows = mxGetM(prhs[0]); extracts the number of rows and
  pb = mxGetPr(prhs[1]);
                               // b
                                                                A_cols = mxGetN(prhs[0]); extracts the number of columns.
  for (k = 0; k < b_rows * b_cols; k++) // b -> x
                                                                The lines
    *(px + k) = *(pb + k);
                                                                  plhs[0]=mxCreateDoubleMatrix(b_rows, b_cols, mxREAL);
                                                                  plhs[1]=mxCreateDoubleMatrix(1, 1, mxREAL);
  for (k = 0; k < A_rows * A_cols; k++) // A -> A_tmp
                                                                allocate storage for the results (of type mxREAL, i.e. ordinary
    *(pA_tmp + k) = *(pA + k);
                                                                double).
  dpbsv_(&uplo, &A_cols, &kd, &b_cols, pA_tmp,
         &A_rows, px, &b_rows, &info);
                                                                A_tmp = mxCreateDoubleMatrix(A_rows, A_cols, mxREAL);
                                                                allocates storage for a copy of A, since the Lapack-routine de-
                                                                strovs the matrix.
  *mxGetPr(plhs[1]) = info; // () higher prec. than *
                                                                px = mxGetPr(plhs[0]); extracts a pointer to the (real-part)
  if (info)
    mexWarnMsgTxt("Non zero info from dpbsv.");
                                                                of the matrix elements and stores it in the pointer variable, px.
  // Should NOT destroy plhs[0] or plhs[1]
                                                                The first for-loop copies b to x (which will be overwritten by the
                                                                solution). The second loop copies the matrix to the temporary
 mxDestroyArray(A_tmp);
                                                                storage, pointed to by A_tmp. This storage is later deallocated
}
                                                                using mxDestroyArray.
                                                                Note that neither the input- nor the output-arguments should
                                                                be deallocated.
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                                                                                           154
It is now time to compile and link. This is done using the
                                                                Now it is time to compile:
Bourne-shell script mex. We must also make a symbolic link.
                                                                % mex -f ./mexopts.sh bandsolve.c lapack/*.f
Since we would like to change some parameters when compiling,
                                                                which creates bandsolve.mexglx.
we will copy and edit an options file, mexopts.sh.
% which matlab
                                                                Now we can test a simple example in Matlab:
/chalmers/sw/sup/matlab-2007b/bin/matlab
                                                                >> A = [2 * ones(1, 5); ones(1, 5)]
(ls -ld /chalmers/sw/sup/matlab* to see the versions)
                                                                A =
Make the link:
                                                                     2
                                                                            2
                                                                                  2
                                                                                         2
                                                                                                2
% ln -s /usr/lib/libstdc++.so.6.0.3 libstdc++.so
                                                                                  1
                                                                                         1
                                                                                                1
                                                                     1
                                                                            1
Copy mexopts.sh:
                                                                >> [x, info] = bandsolve(A, [3 4 4 4 3]')
% cp /chalmers/sw/sup/matlab-2007b/bin/mexopts.sh .
                                                                x =
                                                                    1.0000
and edit the file (after glnx86):
                                                                    1,0000
  change CC='gcc' to CC='gcc4'
                                                                    1.0000
                                                                    1.0000
if you are using the latest Matlab-version. In the CFLAGS-line,
                                                                    1.0000
change -ansi to -Wall, to use C++-style comments and to get
                                                                info =
more warnings.
                                                                     0
Add -L. to CLIBS, and add linker-info, to get Goto-blas:
   CLIBS="SRPATH SMLIBS -lm -L. -lstdc++
                                                                Here a case when A is not positive definite:
           -L/chalmers/sw/unsup/libgoto/lib
           -lgoto_opt32-r0.96"
                                   NOTE: in one long line
                                                                >> A(1, 1) = -2; % Not positive definite
  change -O to -O3 in FOPTIMFLAGS
                                                                >> [x, info] = bandsolve(A, [3 4 4 4 3]')
                                                                Warning: Non zero info from dpbsv.
Make sure your LD_LIBRARY_PATH contains the name of the
                                                                                   % Since b is copied to x
                                                                x =
directory where Goto-blas resides.
                                                                     3
                                                                     4
I have fetched the lapack-routines from Netlib:
                                                                     4
% ls lapack
                                                                     4
          dpbtf2.f dpbtrf.f dpbtrs.f dpotf2.f
dpbsv.f
                                                                     3
ieeeck.f ilaenv.f lsame.f xerbla.f
                                                                info =
                                                                     1
                           155
                                                                                            156
```

<pre>Note that the first call of bandsolve may take much more time, since the mex-file has to be loaded. Here a small test when n=10000, kd=10: &gt;&gt; tic; [x, info] = bandsolve(A, b); toc Elapsed time is 0.147128 seconds. &gt;&gt; tic; [x, info] = bandsolve(A, b); toc Elapsed time is 0.034625 seconds. &gt;&gt; tic; [x, info] = bandsolve(A, b); toc Elapsed time is 0.034950 seconds. Now to some larger problems: With n=100000 and kd=10, dpbsv takes 0.25 s and sparse backslash 0.41 s on a student AMD-computer. kd=20 gives the times 0.48 s and 0.77 s respectively. On an Opteron with more memory: with n=1000000, kd=10 the times are 2.9 s, 4.7 s. Increasing kd to 50, the times are 15.4 s and 27.6 s.</pre>	Libraries, ar, ld Numerical (and other software) is often available in libraries. To use a subroutine from a library one has to use the linker to include the routine. Advantages: • Fewer routines to keep track of. • There is no need to have source code for the library routines that a program calls. • Only the needed modules are loaded. These pages deal with how one can make libraries and use the linker, link-editor, 1d. * cat subl.f90 subroutine sub1 print*, 'in sub1' end * cat sub2.f90 subroutine sub2 print*, 'in sub2' end * cat sub3.f90 subroutine sub3 print*, 'in sub3' call sub2 end * cat main.f90 program main call sub3 end
157	156
<pre>% ls sub*.f90 sub1.f90 sub2.f90 sub3.f90 % g95 -c sub*.f90 sub1.f90: sub2.f90: sub3.f90: % ls sub* sub1.f90 sub1.o sub2.f90 sub2.o sub3.f90 sub3.o % ar -r libsubs.a sub*.o % ar -t libsubs.a sub*.o % ar -t libsubs.a sub1.o sub2.o sub3.o % g95 main.f90 -Llsubs % a.out in sub3 in sub2 g95 calls the link-editor, 1d, to combine main.o and the object files in the library to produce the executable a.out-file. Note that the library routines become part of the executable. If you write -lname the link-editor looks for a library file with name libname.a (or libname.so). On some systems you may have to give the location of the library using the flag -L (1d does not look everywhere) means current working directory, but you could have a longer path, of course. You can have several -L flags.</pre>	<pre>From man ar: ar creates an index to the symbols defined in relocatable object modules in the archive when you specify the modifier s.  An archive with such an index speeds up linking to the library, and allows routines in the library to call each other without regard to their placement in the archive. ar seems to do this even with ar -r as well. If your library does not have this index: % g95 main.f90 -Llsubs ./libsubs.a: could not read symbols: Archive has no index; run ranlib to add one % ranlib libsubs.a % g95 main.f90 -Llsubs The order of libraries is important: % g95 -c sub4.f90 sub5.f90 sub4.f90: sub5.f90: % ar -r libsub45.a sub[45].o % ar -t libsub45.a sub4.o sub5.o</pre>

```
% cat sub4.f90
                                                                % cat main.c
subroutine sub4
                                                                #include <math.h>
 print*, 'in sub4'
                                                                #include <stdio.h>
  call sub2
end
                                                                int main()
                                                                {
                                                                  double x = 1.0e - 15;
% cat main.f90
program main
                  ! A NEW main
 call sub4
                                                                  printf("expml(x) = %e\n", expml(x));
                                                                  printf("exp(x) - 1 = %e n", exp(x) - 1.0);
end
% g95 main.f90 -L. -lsubs -lsub45
                                                                  return 0;
./libsub45.a(sub4.o)(.text+0x6f): In function `sub4_':
                                                                }
: undefined reference to `sub2 '
                                                                % gcc main.c
                                                                /tmp/cc40PH10.o(.text+0x2b): In function `main':
1d \operatorname{does} \underline{\operatorname{not}} go back in the list of libraries.
                                                                : undefined reference to `expml'
                                                                /tmp/cc40PH10.o(.text+0x53): In function `main':
% g95 main.f90 -L. -lsub45 -lsubs
                                                                : undefined reference to 'exp'
% a.out
in sub4
                                                                % gcc main.c -lm
in sub2
                                                                % a.out
The compiler uses several system libraries, try g95 -v ....
                                                                expm1(x)
                                                                          = 1.000000e - 15
One such library is the C math-library, /usr/lib/libm.a.
                                                                exp(x) - 1 = 1.110223e-15
% ar -t /usr/lib/libm.a | grep expm1 | head -1
s_expm1.o
% man expml
NAME expm1, expm1f, expm11 - exponential minus 1
       #include <math.h>
       double expm1(double x);
. . .
                           159
                                                                                            160
                  Shared libraries
                                                                It is easier to handle new versions, applications do not have to
                                                                be relinked.
More about libm. The following output has been shortened.
                                                                If you link with -lname, the first choice is libname.so and the
% ls -l /usr/lib/libm.*
                                                                second libname.a.
  /usr/lib/libm.a
  /usr/lib/libm.so -> ../../lib/libm.so.6
                                                                /usr/lib/libm.so -> ../../lib/libm.so.6 is a soft link
                                                                (an "alias").
% ls -l /lib/libm.*
  /lib/libm.so.6 -> libm-2.3.4.so
                                                                % ln -s full_path alias
                                                                The order is not important when using shared libraries (the
% ls -1 /lib/libm-2.3.4.so
                                                                linker has access to all the symbols at the same time).
-rwxr-xr-x 1 root root 176195 Aug 20 03:21
    /lib/libm-2.3.4.so
                                                                A shared library is created using 1d (not ar) or the compiler,
What is this last file?
                                                                the ld-flags are passed on to the linker.
                                                                % g95 -o libsubs.so -shared -fpic sub*.f90
% ar -t /lib/libm-2.3.4.so
                                                                % g95 main.f90 -L. -lsubs
ar: /lib/libm-2.3.4.so: File format not recognized
                                                                % ./a.out
                                                                 in sub4
Look for symbols (names of functions etc.):
                                                                 in sub2
% objdump -t /lib/libm-2.3.4.so | grep expm1
                                                                From man gcc (edited):
00009420 w
              F.text 0000005c expml
                                                                -shared
 . . .
                                                                  Produce a shared object which can then be linked with
so means shared object. It is a library where routines are loaded
                                                                  other objects to form an executable. Not all systems
to memory during runtime. This is done by the dynamic link-
                                                                  support this option. For predictable results, you must
er/loader ld.so. The a.out-file is not complete in this case, so
                                                                  also specify the same set of options that were used
it will be smaller.
                                                                  to generate code (-fpic, -fPIC, or model suboptions)
                                                                  when you specify this option.[1]
One problem with these libraries is that they are needed at
runtime which may be years after the executable was created.
                                                                -fpic
Libraries may be deleted, moved, renamed etc.
                                                                  Generate position-independent code (PIC) suitable for
                                                                  use in a shared library, if supported for the target
One advantage is shared libraries can be shared by every process
                                                                  machine. Such code accesses all constant addresses
that uses the library (provided the library is constructed in that
                                                                  through a global offset table (GOT). The dynamic
way).
                                                                  loader resolves the GOT entries when the program
                           161
```

starts (the dynamic loader is not part of GCC; it is It is possible to store the location of the library when creating part of the operating system). ... a.out. % unsetenv LD\_LIBRARY\_PATH Since the subroutines in the library are loaded when we run the % g95 -o libsubs.so -shared -fpic sub\*.f90 program (they are not available in **a.out**) the dynamic linker % g95 main.f90 -L. -lsubs must know where it can find the library. % a.out a.out: error while loading shared libraries: % cd .. libsubs.so: cannot open shared object file: % Examples/a.out No such file or directory Examples/a.out: error while loading shared libraries: libsubs.so: cannot open shared object file: No such Add the directory in to the runtime library search path (stored file or directory in a.out): -W1, means pass -rpath 'pwd' to ld % setenv LD\_LIBRARY\_PATH \$LD\_LIBRARY\_PATH\:Examples % Examples/a.out % g95 -Wl,-rpath 'pwd' main.f90 -L. -lsubs in sub4 in sub2 % cd .. or cd to any directory  $\verb"LD_LIBRARY_PATH" contains a colon separated list of paths where$ % Examples/a.out ld.so will look for libraries. You would probably use a full path in sub4 and not Examples. in sub2 A useful command is 1dd (print shared library dependencies): \$LD\_LIBRARY\_PATH is the old value (you do not want to do setenv LD\_LIBRARY\_PATH Examples unless LD\_LIBRARY\_PATH % ldd a.out libsubs.so => ./libsubs.so (0x00800000) is empty to begin with. libm.so.6 => /lib/tls/libm.so.6 (0x009e2000) The backslash is needed in [t]csh (since colon has a special libc.so.6 => /lib/tls/libc.so.6 (0x008b6000) meaning in the shell). In sh (Bourbe shell) you may do some-/lib/ld-linux.so.2 (0x00899000) thing like: Used on our a.out-file it will, in the first case, give: \$ LD\_LIBRARY\_PATH=\$LD\_LIBRARY\_PATH:Example % ldd Examples/a.out \$ export LD\_LIBRARY\_PATH (or on one line) libsubs.so => not found Some form of LD\_LIBRARY\_PATH is usually available (but the In the second case, using **rpath**, **ldd** will print the full path. name may be different). The SGI uses the same name for the path but the linker is called rld. Under HPUX 10.20, for example, the dynamic loader is called dld.sl and the path SHLIB\_PATH. 163 164 And now to something related: Let us now create a tar-file for our package. % tar cvf My\_package.tar My\_package Large software packages are often spread over many directories. My\_package/ When distributing software it is customary to pack all the di-My package/src/ rectories into one file. This can be done with the tar-command My\_package/src/main.f (tape archive). Some examples: My package/src/sub.f My\_package/doc/ % ls -FR My\_package hin/ doc/ install\* lib/ README . . . My\_package/Makefile configure\* include/ INSTALL Makefile src/ One would usually compress it: My\_package/bin: binaries % gzip My\_package.tar (or using bzip2) This command produces the file  ${\tt My_package.tar.gz}$  . My package/doc: documentation .tgz is a common suffix as well (tar.bz2 or .tbz2 for bzip2). userguide.ps or in pdf, html etc. To unpack such a file we can do (using gnu tar) (z for gunzip, My package/include: header files or zcat, x for extract, v for verbose and f for file): params.h sparse.h % tar zxvf My\_package.tar.gz My\_package/lib: libraries My package My\_package/src/ My\_package/src: source . . . main.f sub.f Using tar-commands that do not understand z: Other common directories are man (for manual pages), examples, % zcat My\_package.tar.gz | tar vxf - $\mathbf{or}$ util (for utilities). % gunzip -c My\_package.tar.gz | tar vxf or % gunzip < My\_package.tar.gz | tar vxf -</pre> or  ${\tt README}$  usually contains general information,  ${\tt INSTALL}$  contains % gunzip My\_package.tar.gz followed by details about compiling, installation etc. There may be an install-% tar xvf My\_package.tar script and there is usually a Makefile (probably several). I recommend that you first try: If the package is using X11 graphics there may be an **Imakefile**. % tar ztf My\_package.tar.gz The tool xmkmf (using imake) can generate a Makefile using lo-My package/ ... cal definitions and the Imakefile. To see that files are placed in a new directory (and that are no name conflicts). In a Linux environment binary packages (such as the Intel com-Under GNOME there is an Archive Manager (File Roller) with pilers) may come in RPM-format. See http://www.rpm.org/ a GUI. Look under Applications/System Tools or type man rpm, for details. 165 166