1. [20 points]
   
   (a) Find a 2-by-2 matrix $A$ that is symmetric and nonsingular, but for which neither $A$ nor $-A$ is positive definite. What are the eigenvalues of $A$? Find a 2-vector $y$ such that $y^T Ay < 0$.

   (b) For $A$ as above, find a 2-vector $b$ such that the conjugate gradient algorithm, when started with the zero vector as an initial guess, does not converge to the solution of $Ax = b$. Show what happens on the first two iterations of CG, as described on Slide 3 of the September 26 class. How do you know it won’t converge to the right answer?

2. [40 points] In this problem you’ll actually prove that CG works in at most $n$ steps, assuming that real numbers are represented exactly. (This is not a realistic assumption in floating-point arithmetic, or on any computer with a finite amount of hardware, but it gives a solid theoretical underpinning to CG.) Let $A$ be an $n$-by-$n$ symmetric, positive definite matrix, and let $b$ be an $n$-vector.

   We start with the idea of searching through $n$-dimensional space for the value of $x$ that minimizes $f(x) = \frac{1}{2} x^T Ax - b^T x$, which is the $x$ that satisfies $Ax = b$. We begin by picking a set of $n$ linearly independent search directions, called $d_0, d_1, \ldots, d_{n-1}$. (Actually we don’t know them in advance, but that’s a detail.) At each iteration we proceed along the next direction until we are “lined up” with the final answer, the value of $x$ at which $Ax = b$. In $n$-space, once we are lined up with the answer from $n$ independent directions, we will be exactly on the answer.

   The first magic of CG is that for the right kind of search directions, there is a way to define “lined up” for which we can actually compute how far to go along each search direction. The key definition uses $A$-conjugate vectors. Then “lined up” means that the error $e_i = x_i - x$ is exactly crossways to the search direction $d_{i-1}$, not in the sense of being perpendicular (which would mean $e_i^T d_i = 0$), but in the sense of being $A$-conjugate: $e_i^T Ad_i = 0$.

   An informal way to say that is, we proceed along the search direction until we are lined up with the solution as seen through $A$-glasses. The reason for lining up through $A$-glasses rather than bare eyes is that we can compute where to stop without knowing where the final answer is. We can’t see and compute with $x$-space directly, but we can see the space where $Ax$ and $b$ live. And after lining up each of $n$ independent directions in an $n$-dimensional space we are guaranteed to be sitting on top of the right answer, whether the independent directions are the conventional coordinate axes or the $A$-conjugate axes we see through our $A$-glasses.

   To go along with this, we need to choose the search directions themselves to be mutually $A$-conjugate: we will require each $d_i$ to be $A$-conjugate to all the earlier $d_j$’s, so $d_i^T Ad_j = 0$ if $i \neq j$.

   (a) Suppose we are given $i$ mutually $A$-conjugate vectors $d_0, \ldots, d_{i-1}$. Suppose $x_0 = 0$, and for each $j < i$ we have $x_j = x_{j-1} + \alpha_j d_{j-1}$. Write down and prove correct an expression for a scalar $\alpha_i$ such that, if we take $x_i = x_{i-1} + \alpha_i d_{i-1}$, then the error $e_i = x_i - x$ is $A$-conjugate to $d_{i-1}$.
Now, how do we get a sequence of $A$-conjugate directions to search along? In fact, we can start with any sequence of linearly independent directions, and convert them to $A$-conjugate directions by projecting out all the earlier search directions from each one, using Gram-Schmidt orthogonalization, as follows.

(b) Suppose we are given $i$ mutually $A$-conjugate vectors $d_0, \ldots, d_{i-1}$, and one more vector $u_i$ that does not lie in their span. Write down and prove correct an expression for scalars $\beta_{i,j}$ such that, if we take

$$d_i = u_i + \sum_{j=0}^{i-1} \beta_{i,j} d_j,$$

then $d_i$ is $A$-conjugate to all the earlier $d_j$.

Finally, the second magic of CG is that there is a way to choose a particular sequence of directions for which the Gram-Schmidt orthogonalization is really easy. If we choose the right directions to start with, we only need to project out one earlier direction, not all $i$ of them. This is why the cost of one CG iteration is only $O(n)$, not $O(n^2)$.

(c) Suppose the vectors $d_0, \ldots, d_{i-1}$, the vectors $x_0, \ldots, x_{i-1}$, and the scalars $\alpha_j$ and $\beta_{i,j}$ are as above. Suppose in addition that at each stage we take $u_i = b - Ax_i$ (which is also known as $r_i$, the residual). First, prove that if this choice of $u_i$ lies in the span of $d_0, \ldots, d_{i-1}$, the CG iteration can stop with $x_i = x$. Second, show that this direction $u_i$ is already $A$-conjugate to all of the $d_j$ except $d_{i-1}$, and therefore we can take $\beta_{i,j} = 0$ for $j < i - 1$.

(d) One last detail: Prove that the CG code on the course slide does in fact compute the residual $r_i$ correctly; that is, prove that $r_{i-1} - \alpha_i A d_{i-1}$ is in fact equal to $b - Ax_i$.

3. [40 points] You may do the programming assignments for this course in C or Fortran; I recommend C. In each case, you will set your code up with an interface so that it can be called from Matlab as a “mexFunction”. This will let you use Matlab to test and debug your code, and to plot results. This warmup assignment is just for you to learn how to write a Matlab interface using sparse matrices.

Write a C or Fortran mexFunction that can be called from Matlab as $y = \text{matvec}(A,x)$, which takes as input a sparse matrix $A$ and a full column vector $x$, and returns a full column vector $y$ whose value is the matrix-vector product $Ax$. Your routine can assume the matrix is real, but should not assume that it’s square. (You might want to check to make sure the sizes of $A$ and $x$ are compatible.)

Test your routine from Matlab with several sparse matrices you make up, verifying that it gives the same answer as Matlab’s $y = A\times x$. (The norm of the difference, $\text{norm}(A\times x - \text{matvec}(A,x))$, should be tiny. It may not be exactly zero because floating-point addition is not associative, and your routine may be doing arithmetic in a different order than Matlab’s.) See the Matlab functions sprand or sprandn to generate random sparse test matrices. Or download the Matlab interface to the UF Sparse Matrix Collection, at http://www.cise.ufl.edu/research/sparse.

For documentation on mexFunctions, open up a help window in Matlab, and look in the External Interfaces/API section, under the Matlab section. The C mexFunction syntax is quite a bit nicer than the Fortran. All mexFunctions have the routine name mexFunction.

You’ll need to use the following Matlab “mx” and “mex” routines:
mxGetM returns the number of rows of a matrix
mxGetN returns the number of columns of a matrix
mxGetJc returns a pointer to the column pointer array (Ap)
mxGetIr returns a pointer to the row indices (Ai)
mxGetPr returns a pointer to the numerical values (Ax)

There is a more complex but very well-written example of mexFunctions for sparse matrices in Tim Davis’s LDL code, which is at http://www.cise.ufl.edu/research/sparse/ldl and also on the CS290H reference page on the web. LDL computes a sparse Cholesky factorization; you don’t need to understand how that works, but you can see from that code how a C program can get access to a Matlab sparse matrix.

Turn in all your code, and also a Matlab transcript of a session that tests your code and verifies that the output agrees with Matlab’s. (Say “help diary” to Matlab to see how to record a transcript.)