I compiled LAPACK with lines 19-24 changed to following in make.inc because running tests caused segfault with -frecursive:

FORTRAN = gfortran

OPTS = -02

DRVOPTS = \$(OPTS)

NOOPT = -00

LOADER = gfortran

LOADOPTS =

Then I compiled fortran version of the given program with "gfortran -o ex2_p3 ex2_p3.f90 -llapack -lblas" (flags to link lapack) and ran it first with "./ex2_p3 < matrix6" which yielded following x:

-22.413899 12.990382 29.555568 -9.3848441 -21.079137 10.385575

After that I gave it matrix100, vector x for which was following:

7.2144883	11.163655	3.0358445	1.0799059	1.8689894
4.7119332	-8.2343102	5.5900076	-4.1903471	-3.5101576
10.645714	-4.0814602	-8.6241006	-0.63710851	2.7890658
8.7764311	-4.1028957	-4.7785748	-4.4614943	-11.498610
-11.963623	11.075400	5.6697070	3.5490224	
0.58301116E-0	1 -5.0599416	-1.9754759	-1.2538670	-3.8985944
8.2744529	-5.6839025	-5.6691980	0.37960791	-4.1624764
10.597685	-0.34778509	-1.6239494	-3.8423147	-4.8322450
0.62900360				
-2.6086740	6.8657889	-8.9785345	4.6703364	0.54891920
-8.3940974	-5.2656037	5.1105622	2.7500810	7.5109883
5.4796523	9.1413935	2.0645985	-3.7819297	-3.8121316
-1.5216914	10.472063	10.756644	-12.808315	-3.2649371
4.2395374	-8.4943304	-3.9817579	-5.1799615	2.5102437
-1.9911264	0.23276902E-01	4.8850823	-0.69939738	3.1103510
0.94131700	0.10284713	4.3865474	5.2776985	4.2953560
-2.3233743	15.556085	-4.8151092	6.9185871	-8.0440978
-0.75193463	2.3261727	-1.4131884	-1.7420666	3.6335352
-13.360496	6.8480197	-7.3610078	-2.3170964	3.3144431
-1.6072032	-3.5258292	4.2756271	-6.1851535	-5.3759474
-8.5609332	-3.9181126	-0.66337326	8.2351565	8.6265645

Ex. 4

Residuals:

m	matrix6	matrix100
0	1.952974e-07	1.003843e-06
1	5.412062e-07	2.496027e-05
2	2.742264e-07	3.031577e-06

I used dgemm for matrix multiplication. Makefile for compiling is attached. I did not manage to use LAPACK with C-standard newer than C89, is that how it just is or did I fail at googling?

Program expexts inputs in order N, M, A, b, x.