# Algorithm 778: L-BFGS-B: Fortran Subroutines for Large-Scale Bound-Constrained Optimization

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L-BFGS-B is a limited-memory algorithm for solving large nonlinear optimization problems subject to simple bounds on the variables. It is intended for problems in which information on the Hessian matrix is difficult to obtain, or for large dense problems. L-BFGS-B can also be used for unconstrained problems and in this case performs similarly to its predecessor, algorithm L-BFGS (Harwell routine VA15). The algorithm is implemented in Fortran 77.

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## 1. INTRODUCTION

The purpose of algorithm L-BFGS-B is to minimize a nonlinear function of n variables,

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 $\min f(x)$ 

subject to

$$l \leq x \leq u$$
,

where the vectors l and u represent lower and upper bounds on the variables. Not all the variables need to have bounds; in fact the algorithm is also appropriate and efficient for solving unconstrained problems. The user must supply the gradient g, but knowledge about the Hessian matrix of f is not required. For this reason the algorithm can be useful for solving large problems in which the Hessian is difficult to compute or is dense.

The algorithm is described in detail in Byrd et al. [1995] and proceeds roughly as follows. At each iteration a limited-memory BFGS approximation to the Hessian is updated. This limited-memory matrix is used to define a quadratic model of the objective function f. A search direction is then computed using a two-stage approach: first, the gradient projection method [Bertsekas 1982; Conn et al. 1988; Levitin and Polyak 1966; Moré and Toraldo 1989] is used to identify a set of active variables, i.e., variables that will be held at their bounds; then the quadratic model is approximately minimized with respect to the free variables. The search direction is defined to be the vector leading from the current iterate to this approximate minimizer. Finally a line search is performed along the search direction using the subroutine described in Moré and Thuente [1994]. A novel feature of the algorithm is that the limited-memory BFGS matrices are represented in a compact form [Byrd et al. 1994] that is efficient for bound-constrained problems.

The user can control the amount of storage required by L-BFGS-B by selecting a parameter m that determines the number of BFGS corrections saved. The algorithm requires roughly (12+2m)n storage locations, and since small values of m (say  $3 \le m \le 20$ ) are recommended, it can be used to solve very large problems. The computational cost of one iteration of the algorithm is modest, ranging from 4mn + n multiplications when no bounds are active, to approximately  $m^2n$  multiplications when all variables are at their bounds.

L-BFGS-B is an extension of the limited-memory algorithm (L-BFGS) for unconstrained optimization described in Liu and Nocedal [1989] and implemented as Harwell routine VA15. The main improvement is the ability of L-BFGS-B to deal with bounds on the variables. Even though this requirement makes the new algorithm far more complex than its predecessor, the two codes perform similarly on unconstrained problems. Therefore L-BFGS-B could be considered to supersede L-BFGS, except for one fact that

<sup>&</sup>lt;sup>1</sup>For information regarding how to obtain VA15, send email to Scott Roberts (Scott.Roberts@aeat.co.uk), Richard Lee (Richard.Lee@aeat.co.uk), or Maria Woodbridge (Maria.Woodbridge@aeat.co.uk).

can be important in some applications: L-BFGS-B requires 8 more n-vectors of storage.

L-BFGS-B is, at present, the only limited-memory quasi-Newton algorithm capable of handling bounds on the variables; other published codes [Buckley 1989; Buckley and LeNir 1985; Gilbert and Lemaréchal 1989]<sup>2</sup> are only able to solve unconstrained problems. We note also that the nonlinear conjugate gradient method [Gill et al. 1981], which is used for solving many large unconstrained problems, has not been adequately extended to handle bounds on the variables, and L-BFGS-B can be used in its place.

The advantages of L-BFGS-B are (1) the code is easy to use, and the user need not supply information about the Hessian matrix or the structure of the objective function, (2) the storage requirements are modest and can be controlled by the user, and (3) the cost of the iteration is low and is independent of the properties of the objective function. As a result, L-BFGS-B is recommended for large problems in which the Hessian is not sparse or is difficult to compute.

However, L-BFGS-B suffers from the following drawbacks: (1) it is not rapidly convergent, and on difficult problems it can take a large number of function evaluations to converge; (2) on highly ill-conditioned problems it may fail to obtain high accuracy in the solution; (3) it cannot make use of knowledge about the structure of the problem to accelerate convergence.

Although the algorithm implemented in the Fortran code is essentially the same as the one described in Byrd et al. [1995], it differs in a few important details. As a result of these improvements, the operation counts given in Byrd et al. [1995] slightly overestimate the computational work of L-BFGS-B; we return to this in Section 4. Several new limited-memory algorithms have recently been proposed that reduce the amount of storage needed; see for example Siegel [1992]. We have not followed such an approach because it is not clear at present that their performance is as good as that of L-BFGS-B.

## 2. THE DRIVERS

L-BFGS-B is written in Fortran 77, in double precision. The user is required to calculate the function value f and its gradient g. In order to allow the user complete control over these computations, reverse communication is used. Thus, the routine setulb.f must be called repeatedly from the user's program.

The simplest way to use the code is to modify one of the sample drivers provided in the package. Most users will need to make only a few changes to one of the drivers to run their applications.

—**driver1.f** is the simplest driver. It demonstrates how to solve a problem using default parameters. We recommend that every user of L-BFGS-B read this driver. It gives a good idea of how the code works, and at the

<sup>&</sup>lt;sup>2</sup>Including VA15.

end of the program there is a detailed description of the parameters used in L-BFGS-B.

- —**driver2.f** is a more sophisticated driver. It illustrates various ways of terminating the run and alternative ways of generating output. This driver is designed for users who need specially formatted output or for users who wish to have more control over the execution of the run.
- —**driver3.f** is a time-controlled driver. It shows how to terminate a run after some prescribed CPU time has elapsed and how to print the desired output before exiting.

## 3. TERMINATION AND ERROR MESSAGES

The code may terminate for a variety of reasons described in this section. First of all, the user can force termination by including an appropriate instruction in the driver, e.g., see driver2. The code may also terminate if one of the two built-in stopping tests is activated. The first stopping test is

$$\frac{(f_k - f_{k+1})}{\max(|f_{k+1}|, |f_k|, 1)} \leq \mathbf{factr} * \mathbf{epsmch}, \tag{1}$$

where **epsmch** is the machine precision, which is automatically generated by the code, and **factr** is a parameter controlled by the user. This test is designed to terminate the run when the change in the objective function f is sufficiently small. Typical values for **factr** on a computer with 15 digits of accuracy in double precision are as follows: **factr** = 1.d + 12 for low accuracy; **factr** = 1.d + 7 for moderate accuracy; **factr** = 1.d + 1 for extremely high accuracy. If **factr** = 0, the test will stop the algorithm only if the objective function remains unchanged after one iteration.

The second built-in stopping test is based on the projected gradient. This is the projection of the gradient vector onto the space tangent to the active bounds, and it must equal zero at a local minimizer of the bound-constrained problem. The test is designed to terminate the run when the (infinity) norm of the projected gradient becomes sufficiently small:

$$\|\operatorname{proj} g\|_{\infty} \leq \mathbf{pgtol}.$$
 (2)

The parameter **pgtol** is controlled by the user, but the test will be hard to satisfy if **pgtol** is set smaller than the square root of machine precision.

Both tests can be almost disabled by setting factr = pgtol = 0.

The code may also terminate because an input error has been detected, or because no further progress can be made during the line search, as described in Section 4.

#### 4. IMPLEMENTATATION

The algorithm implemented in L-BFGS-B is described in detail in Byrd et al. [1995]. However, a few additions and modifications were made during the development of the code.

Section 5 of Byrd et al. [1995] describes three methods for performing the subspace minimization: direct primal, primal CG, and dual. Extensive numerical tests since the publication of Byrd et al. [1995] indicate that the CG approach is the least effective. Moreover we were able to show that the primal and dual approaches can be implemented in a unified framework in which they are very similar; they require essentially the same amount of computation and perform equally well in practice. Tests supporting these observations are reported in the technical report by Zhu et al. [1995], an earlier version of this article. Because of this, the L-BFGS-B code uses only the primal method for subspace minimization.

There is another significant difference between L-BFGS-B and the algorithm in Byrd et al. [1995], but it occurs at a fairly low level and is of interest only to those readers wishing to understand the code in detail. The definition of the reduced Hessian given in Eq. (5.10) of Byrd et al. [1995] makes use of the matrix

$$\left(I - rac{1}{ heta}MW^TZZ^TW
ight)^{-1}M.$$

This matrix can be written as the inverse of

$$\left[ egin{array}{ccc} -D - rac{1}{ heta} Y^T Z Z^T Y & L_A^T - R_Z^T \ L_A - R_z & heta S^T A A^T S \end{array} 
ight],$$

where  $L_A$  is the strict lower triangle of  $S^TAA^TS$  and  $R_Z$  is the upper triangle of  $Y^TZZ^TY$ . Although this matrix is not positive definite, it can be factorized symmetrically by using Cholesky factorizations of the submatrices, and we do so in the L-BFGS-B code.

Next we describe several devices for dealing with failures of the code and for trying to improve performance in the region where rounding errors begin to dominate the computation. The steplength parameter is computed by a procedure using the line search program of Moré and Thuente [1994]. If the line search is unable to find a point with a sufficiently lower value of the objective after 20 evaluations of the objective function, we conclude that the current direction is not useful.

In this case all correction vectors forming the limited-memory matrix are discarded and the iteration is restarted along the steepest descent direction. If the line search fails along this steepest descent direction, the algorithm terminates with an error message. This type of failure will usually occur only if the user has specified high accuracy in the solution and L-BFGS-B is having difficulties meeting this accuracy. Our restarting strategy sometimes leads to successful termination in these difficult cases, but not always.

Similarly, if during the course of the iteration the limited-memory BFGS matrix, or a related submatrix, becomes singular or indefinite, all correc-

tion vectors are discarded and the iteration is restarted along the steepest descent direction. This device is also used if the search direction d is not a descent direction (i.e., if  $g^T d \ge 0$ ).

We emphasize that all the difficulties just described occur only when rounding errors begin to dominate the computation.

## 4.1 Machine and Scale Dependencies

L-BFGS-B computes the machine precision **epsmch** by means of the routine **dpmeps** from MINPACK-2.<sup>3</sup> The machine precision **epsmch** is used twice in the algorithm: in the stopping test (1) and in the skipping criterion for BFGS updating described now.

The line search of Moré and Thuente [1994] enforces the Wolfe conditions whenever no bound is encountered in the line search. In particular, if no bounds are hit we always have  $y_k^T s_k \ge 0.9(-g_k^T s_k)$ , where  $y_k = g_{k+1} - g_k$ , and  $s_k = x_{k+1} - x_k$ . If a bound is hit, it may not be possible to satisfy this condition (see Dennis and Schnabel [1983]) and to ensure that the Hessian approximation is sufficiently positive definite. We therefore skip the BFGS update if

$$\frac{y_k^T s_k}{-g_k^T s_k} \le \mathbf{epsmch}. \tag{3}$$

Our numerical experience indicates that skipping occurs rarely. The test (3) is rather weak, and it is possible that in finite precision a small value of  $y_k^T s_k$  could result in the BFGS update being undefined or not positive definite. In such cases the restarting mechanism described above would allow the algorithm to continue. We have never observed any numerical indefiniteness that gives rise to a nondescent direction.

Effort was taken to ensure that L-BFGS-B is as scale-invariant as possible. However, complete scale-invariance was not possible to achieve; indeed the limited-memory algorithm itself is not invariant to linear transformations in the variables. However, the algorithm is invariant with respect to scalar multiples of the variables and the objective function, and we have been able to maintain that invariance in the code with only a few exceptions. One of them occurs in the first iteration, where the step is quite dependent on scaling of the variables.

Note that (2) and, when |f| is small, (1) are scale-dependent: if f is multiplied by a constant and the code is rerun, then termination may occur at a different solution point for positive **factr** and **pgtol**.

## 5. NUMERICAL RESULTS

We now present results of L-BFGS-B on a set of test problems from the CUTE collection [Bongartz et al. 1995]. We tested only bound-constrained

<sup>&</sup>lt;sup>3</sup>Currently being developed by Averick and Moré.

Table I. Test Results of L-BFGS-B and Results of LANCELOT with SR1 and Exact Hessian Options, on Bound-Constrained Problems from the CUTE Collection

			L-BFGS-B m=5			L-BFGS-B m=17		LANCELOT SR1		LANCELOT Hessian	
Problem	n	nbnd	nfg	Time	nfg	Time	nf	Time	nf	Time	
BDEXP	1000	0	15	2.31	16	3.50	27	13.74	11	6.54	
BIGGS5	6	1	121	0.88	69	1.51	41	0.62	19	0.30	
BQPGASIM	50	7	25	0.28	23	0.43	8	0.58	4	0.34	
BQPGAUSS	2003	27	*F1	(7E-3)	*C1	(4E-4)	20	1957.59	9	1751.29	
HATFLDC	25	0	23	0.19	23	0.41	5	0.11	5	0.19	
HS110	50	50	2	0.02	2	0.02	2	0.17	2	0.16	
HS45	5	5	11	0.03	11	0.01	3	0.05	3	0.03	
JNLBRNGA	15625	5657	332	740.33	296	1133.88	24	1263.77	22	1502.96	
JNLBRNGB	1024	516	424	62.73	426	125.17	6	7.21	6	5.56	
LINVERSE	999	338	291	56.85	369	159.31	27	194.08	28	149.99	
MAXLIKA	8	1	1665	88.38	158	10.27	98	24.33	9	2.24	
MCCORMCK	1000	0	15	1.85	15	2.05	7	5.25	5	3.97	
NONSCOMP	1000	2	45	6.79	60	17.24	9	4.70	9	4.43	
OBSTCLAE	5625	2724	258	207.20	308	455.60	7	1442.00	6	1422.62	
OBSTCLAL	1024	508	40	5.84	40	10.45	11	9.45	9	7.69	
OBSTCLBL	1024	475	50	7.83	55	16.62	8	15.42	8	18.45	
OBSTCLBM	15625	4309	146	353.04	138	573.84	7	1106.37	6	2017.70	
OBSTCLBU	1024	475	44	6.57	41	11.48	9	16.10	8	8.45	
PALMER1A	6	0	799	4.95	262	4.50	113	2.29	68	1.37	
PALMER1E	8	0	*F1	(7E-2)	290	5.06	190	6.95	204	7.38	
PALMER2A	6	0	518	3.67	182	4.12	180	3.05	157	2.60	
PALMER2E	8	0	*F1	(2E-3)	291	6.98	268	8.01	113	3.89	
PALMER3A	6	0	716	5.13	140	3.31	176	3.02	147	2.48	
PALMER3E	8	0	*F1	(4E-4)	221	3.59	141	3.81	68	1.76	
PALMER4A	6	0	483	3.30	128	2.82	98	1.54	48	0.80	
PALMER4E	8	0	*F1	(3E-3)	172	2.89	206	4.78	67	1.95	
PROBPENL	500	0	3	0.10	3	0.11	3	1.72	2	1.67	
S368	100	29	21	16.84	21	16.93	37	91.24	8	21.14	
TORSION1	1024	436	43	6.35	32	8.16	13	11.04	11	10.18	
TORSION2	1024	436	61	10.08	55	18.33	10	12.31	5	12.69	
TORSION3	1024	748	23	2.76	22	3.61	7	8.43	6	4.05	
TORSION4	1024	748	49	5.87	43	8.32	7	6.73	6	4.85	
TORSION6	14884	12316	362	707.22	360	1157.74	10	130.73	9	130.31	
*Termination	because	e the nun	nber of t	function e	valuat	ions reac	hed 9	999 (the	valu	e in	
parentheses is	parentheses is the norm of the projected gradient at the final iterate); C1: gradient										
atonning toat	stanning test (2) was not mot but the final function value was at least as good as that										

\*Termination because the number of function evaluations reached 9999 (the value in parentheses is the norm of the projected gradient at the final iterate); C1: gradient stopping test (2) was not met, but the final function value was at least as good as that obtained by LANCELOT SR1; F1: gradient stopping test (2) was not met, and the final function value was greater than that obtained by LANCELOT SR1.

problems with  $n \ge 5$  and unconstrained problems with  $n \ge 100$ . As a benchmark we also present the results obtained by the SR1 and Exact Hessian methods of the LANCELOT package [Conn et al. 1992]. LANCELOT was run using all its default options. All runs were performed on a SPARCstation-2 with 32MB of main memory; the stopping test used **pgtol** =  $10^{-5}$  in (2).

The meaning of some of the variables used in the tables is as follows.

—**nbnd**: the number of active bounds at the solution of LANCELOT-SR1.

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Table II.	Test Results	of L-BFGS-B	and Results	of LANCEL	OT with	SR1 and	Exact	Hessian
	Options	s, on Unconstr	ained Proble	ems from the	CUTE (	Collection		

-											
				FGS-B		GS-B		CELOT	LANCELOT		
			n	n=5		=17		SR1		ssian	
	Problem	n	nfg	Time	nfg	Time	nf	Time	nf	Time	
	ARWHEAD	1000	13	1.09	**C1	(2E-5)	5	4.66	6	4.79	
	BDQRTIC	100	101	1.28	47	1.29	11	1.06	12	1.07	
	BROYDN7D	1000	373	66.30	398	104.51	112	62.72	125	66.52	
	CRAGGLVY	1000	95	13.33	89	19.08	15	9.81	15	9.89	
	DIXMAANA	1500	12	1.34	13	1.66	8	8.71	6	7.96	
	DIXMAANB	1500	12	1.36	12	1.43	9	10.18	8	8.96	
	DIXMAANC	1500	14	1.61	14	1.85	10	8.91	12	11.28	
	DIXMAAND	1500	15	1.70	15	2.08	13	13.07	20	15.73	
	DIXMAANE	1500	188	24.28	169	41.07	14	13.01	7	8.74	
	DIXMAANF	1500	163	21.04	126	30.71	26	21.17	33	22.11	
	DIXMAANG	1500	158	20.38	127	30.94	32	24.78	25	18.05	
	DIXMAANH	1500	156	20.30	124	30.01	37	28.07	36	24.44	
	DIXMAANI	1500	1237	166.37	1066	273.08	11	11.56	8	9.30	
	DIXMAANK	1500	130	16.59	146	35.36	34	25.98	51	32.56	
	DIXMAANL	1500	134	16.93	120	28.04	105	67.67	50	35.88	
	DQDRTIC	1000	19	1.47	19	1.73	3	2.47	3	2.55	
	DQRTIC	500	43	1.46	43	2.96	34	6.13	34	6.11	
	EIGENALS	110	574	17.21	302	15.77	21	4.72	22	4.36	
	EIGENBLS	110	1116	33.36	1041	55.73	186	98.47	193	95.55	
	EIGENCLS	462	2900	563.81	2507	599.32	456	2010.40	543	2299.42	
	ENGVAL1	1000	23	2.02	20	2.38	8	6.28	8	6.03	
	FREUROTH	1000	**C1	(2E-5)	**C1	(1E-3)	11	7.53	11	7.27	
	GENROSE	500	1244	60.86	1315	116.82	590	103.92	586	99.79	
	MOREBV	1000	79	6.85	77	12.22	2	3.89	2	3.85	
	NONDIA	1000	23	1.79	23	2.56	C2	C2	30	12.54	
	NONDQUAF	R 100	1001	10.09	828	25.82	16	0.86	16	0.86	
	PENALTY1	1000	60	3.91	60	7.58	64	118.89	64	117.61	
	PENALTY3	100	**C1	(3E-3)	**C1	(3E-3)	100	436.12	**C1	(2E-4)	
	QUARTC	1000	47	3.10	47	5.86	36	12.74	36	12.68	
	SINQUAD	1000	183	17.17	210	32.76	132	81.51	132	79.20	
	SROSENBR	1000	20	1.18	19	1.77	14	6.85	11	5.92	
	TQUARTIC	1000	27	1.77	27	2.80	13	7.76	13	5.93	
	TRIDIA	1000	763	48.90	534	78.98	3	3.96	3	3.91	

\*\*Termination because the code could make no further progress in reducing f; the value in parentheses is the norm of the projected gradient at the final iterate; C1: gradient stopping test (2) was not met, but the final function value was at least as good as that obtained by LANCELOT SR1; C2: the SR1 option of LANCELOT converged to a different solution point than the other methods.

Tables I and II indicate that L-BFGS-B is a competitive code in terms of CPU time. This came as a surprise since L-BFGS-B does not use any specific knowledge of the objective function, as is the case in both versions

<sup>-</sup>nfg: the total number of function or gradient evaluations.

<sup>—</sup>**nf**: the total number of function evaluations. (In LANCELOT, the number of function evaluations may differ from the number of gradient evaluations.)

Table III. Test Results of L-BFGS-B with Various Values for m, on Bound-Constrained Problems from the CUTE Collection

		IRI	FGS-B	IR	FGS-B	I. B	FGS-B	IR1	FGS-B	
			m=3		1=5		m=17		m=29	
Problem	n	nfg	Time	nfg			Time	nfg	Time	
BDEXP	1000	15	1.91	15	2.31	16	3.50	16	3.61	
BIGGS5	6	109	0.57	121	0.88	69	1.51	71	3.23	
BQPGASIM		28	0.25	25	0.28	23	0.43	23	0.43	
BQPGAUSS		*F1	(3E-2)	*F1	(7E-3)	*C1	(4E-4)	**C1	(5E-5)	
HATFLDC	25	25	0.14	23	0.19	23	0.41	23	0.36	
HS110	50	2	0.01	2	0.02	2	0.02	2	0.02	
HS45	5	11	0.02	11	0.03	11	0.01	11	0.01	
JNLBRNGA	15625	389	763.79	332	740.33	296	1133.88	323	1758.70	
JNLBRNGB	1024	569	65.13	424	62.73	426	125.17	447	228.05	
LINVERSE	999	564	91.89	291	56.85	369	159.31	416	315.31	
MAXLIKA	8	*F1	(5E-3)	1665	88.38	158	10.27	118	10.67	
MCCORMCI	K 1000	15	2.00	15	1.85	15	2.05	15	2.04	
NONSCOMI	P 1000	46	5.38	45	6.79	60	17.24	61	20.14	
OBSTCLAE	5625	261	182.05	258	207.20	308	455.60	282	578.10	
OBSTCLAL		39	4.74	40	5.84	40	10.45	39	11.71	
OBSTCLBL	1024	55	7.07	50	7.83	55	16.62	53	22.27	
OBSTCLBM	15625	161	338.97	146	353.04	138	573.84	146	828.85	
OBSTCLBU	1024	46	5.62	44	6.57	41	11.48	41	15.12	
PALMER1A	6	*F1	(2E-1)	799	4.95	262	4.50	197	8.01	
PALMER1E	8	*F1	(2E-1)	*F1	(7E-2)	290	5.06	254	10.81	
PALMER2A	6	2888	16.26	518	3.67	182	4.12	170	9.69	
PALMER2E	8	*F1	(1E-3)	*F1	(2E-3)	291	6.98	221	13.29	
PALMER3A	6	2460	14.12	716	5.13	140	3.31	134	7.45	
PALMER3E	8	*F1	(2E-3)	*F1	(4E-4)	221	3.59	182	7.50	
PALMER4A	6	1985	11.38	483	3.30	128	2.82	90	4.32	
PALMER4E	8	*F1	(5E-2)	*F1	(3E-3)	172	2.89	142	5.42	
PROBPENL	500	3	0.11	3	0.10	3	0.11	3	0.10	
S368	100	19	15.23	21	16.84	21	16.93	21	16.86	
TORSION1	1024	60	7.38	43	6.35	32	8.16	33	9.51	
TORSION2	1024	59	7.87	61	10.08	55	18.33	63	30.45	
TORSION3	1024	27	2.86	23	2.76	22	3.61	22	3.65	
TORSION4	1024	50	5.96	49	5.87	43	8.32	42	10.17	
TORSION6	14884	309	565.25	362	707.22	360	1157.74	422	1994.78	
*Torminatio	n hogour	o the nu	mbon of f	ination	ovoluotio	ma mood	.0000 bad	**+0****	action	

\*Termination because the number of function evaluations reached 9999; \*\*termination because the code could make no further progress in reducing f (in cases \*\* and \* the value in parentheses is the norm of the projected gradient at the final iterate); C1: gradient stopping test (2) was not met, but the final function value was at least as good as that obtained by LANCELOT SR1; F1: gradient stopping test (2) was not met, and the final function value was greater than that obtained by LANCELOT SR1.

of LANCELOT. On the other hand, LANCELOT used much fewer function evaluations. It is an interesting fact that L-BFGS-B is sometimes unable to reduce the projected gradient sufficiently to satisfy the stopping condition even though the function value obtained is very good. More specifically, in the runs marked by C1 in the tables, L-BFGS-B obtained at least as good function value (to five digits) as LANCELOT but the gradient did not meet the stopping condition (2). We do not interpret these as failures of the algorithm, and feel that this property of L-BFGS-B deserves further study.

Table IV. Test Results of L-BFGS-B with Various Values for m, on Unconstrained Problems from the CUTE Collection

-		I DEGG D				100 P		IGG P			
				L-BFGS-B		FGS-B		L-BFGS-B m=17		L-BFGS-B m=29	
				n=3		=5					
_	Problem	n	nfg	Time	nfg	Time	nfg	Time	nfg	Time	
		1000	12	0.95	13	1.09	**C1	(2E-5)	**C1	(2E-5)	
	BDQRTIC	100	124	1.34	101	1.28	47	1.29	39	1.59	
	BROYDN7D	1000	393	64.47	373	66.30	398	104.51	384	146.08	
	CRAGGLVY	1000	99	12.79	95	13.33	89	19.08	85	24.45	
	DIXMAANA	1500	11	1.09	12	1.34	13	1.66	13	1.61	
	DIXMAANB	1500	12	1.24	12	1.36	12	1.43	12	1.44	
	DIXMAANC	1500	14	1.47	14	1.61	14	1.85	14	1.81	
	DIXMAAND	1500	15	1.56	15	1.70	15	2.08	15	2.04	
	DIXMAANE	1500	214	23.53	188	24.28	169	41.07	166	60.31	
	DIXMAANF	1500	164	18.14	163	21.04	126	30.71	124	45.14	
	DIXMAANG	1500	191	20.93	158	20.38	127	30.94	132	47.22	
	DIXMAANH	1500	157	17.37	156	20.30	124	30.01	127	46.04	
	DIXMAANI	1500	828	97.87	1237	166.37	1066	273.08	922	364.27	
	DIXMAANK	1500	146	16.10	130	16.59	146	35.36	133	47.63	
	DIXMAANL	1500	164	17.93	134	16.93	120	28.04	125	44.08	
	DQDRTIC	1000	23	1.64	19	1.47	19	1.73	19	1.76	
	DQRTIC	500	43	1.28	43	1.46	43	2.96	43	4.06	
	EIGENALS	110	769	21.30	574	17.21	302	15.77	145	13.02	
	EIGENBLS	110	1445	39.91	1116	33.36	1041	55.73	870	86.35	
	EIGENCLS	462	2613	493.70	2900	563.81	2507	599.32	1969	593.89	
	ENGVAL1	1000	23	1.78	23	2.0	20	2.38	20	2.39	
	FREUROTH	1000	82	7.73	**C1	(2E-5)	**C1	(1E-3)	38	6.40	
	GENROSE	500	1323	57.99	1244	60.86	1315	116.82	1306	198.67	
	MOREBV	1000	73	5.50	79	6.85	77	12.22	76	18.04	
	NONDIA	1000	21	1.48	23	1.79	23	2.56	23	2.67	
	NONDQUAR	100	866	6.96	1001	10.09	828	25.82	588	43.50	
	PENALTY1	1000	60	3.26	60	3.91	60	7.58	60	10.96	
	PENALTY3	100	**C1	(9E-3)	**C1	(3E-3)	**C	(3E-3)	**C1	(1E-3)	
	QUARTC	1000	47	2.68	47	3.10	47	5.86	47	7.62	
	SINQUAD	1000	211	17.45	183	17.17	210	32.76	231	52.93	
	SROSENBR	1000	18	0.90	20	1.18	19	1.77	19	1.81	
	TQUARTIC	1000	23	1.34	27	1.77	27	2.80	27	2.97	
	TRIDIA	1000	882	44.16	763	48.90	534	78.98	474	120.90	
		-									

\*\*Termination because the code could make no further progress in reducing f (the value in parentheses is the norm of the projected gradient at the final iterate); C1: gradient stopping test (2) was not met, but the final function value was at least as good as that obtained by LANCELOT SR1.

Tables III and IV show the effect of varying the number m of updates saved. Increasing m definitely improves the reliability of the algorithm. Although increasing m often reduces the number of function evaluations, this effect is not consistent, and it does cause an increase in CPU time in most cases.

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