

Algorithm 813: SPG—Software for Convex-Constrained Optimization

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Fortran 77 software implementing the SPG method is introduced. SPG is a nonmonotone projected gradient algorithm for solving large-scale convex-constrained optimization problems. It combines the classical projected gradient method with the spectral gradient choice of steplength and a nonmonotone line-search strategy. The user provides objective function and gradient values, and projections onto the feasible set. Some recent numerical tests are reported on very large location problems, indicating that SPG is substantially more efficient than existing general-purpose software on problems for which projections can be computed efficiently.

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1. INTRODUCTION

In this article, we describe Fortran 77 software that implements the nonmonotone spectral projected gradient (SPG) algorithm. The SPG method applies to problems of the form

min
$$f(x)$$
 subject to $x \in \Omega$,

where Ω is a closed convex set in \mathbb{R}^n . It is assumed that f is defined and has continuous partial derivatives on an open set that contains Ω . Users of the software must supply subroutines to compute the function f(x), the gradient $\nabla f(x)$ and projections of an arbitrary point x onto Ω . Information about the Hessian matrix is not required and the storage requirements are minimal. Therefore, the algorithm is appropriate for large-scale convex-constrained optimization problems with affordable projections onto the feasible set. Notice that the algorithm is also suitable for unconstrained optimization problems simply by setting $\Omega = \mathbb{R}^n$.

The algorithm is fully described in Birgin et al. [2000] and combines the projected gradient method [Bertsekas 1976] with two new features in optimization. First, it extends the typical globalization strategies associated with these methods to the nonmonotone line search scheme developed by Grippo et al. [1986]. Second, it uses the spectral steplength, introduced by Barzilai and Borwein [1988] and analyzed by Raydan [1993]. This choice of steplength requires little computational work and greatly speeds up the convergence of gradient methods for unconstrained problems [Raydan 1997].

It is worth noting that two different versions of the projected gradient method were considered in Birgin et al. [2000]. The new features were applied to the classical curvilinear path (piecewise linear if Ω is a polyhedral set) to introduce Algorithm SPG1. They were also applied to the feasible continuous projected path to produce Algorithm SPG2. Based on numerical experimentation reported in Birgin et al. [2000], we concluded that there are no meaningful differences between the performances of SPG1 and SPG2. Therefore, since SPG1 tends to require more projections onto Ω per iteration, we only consider the SPG2 version of the method in the software presented and described in this article.

2. ALGORITHM

Given $\hat{x} \in \mathbb{R}^n$ we define $P_{\Omega}(\hat{x})$ to be the projection with respect to a given norm $\|\cdot\|$ onto Ω , that is, $P_{\Omega}(\hat{x}) = \arg\min_{x \in \Omega} \|x - \hat{x}\|$. We denote $g(x) = \nabla f(x)$. The algorithm starts with $x_0 \in \mathbb{R}^n$ and uses an integer $m \geq 1$; a small parameter $\lambda_{\min} > 0$; a large parameter $\lambda_{\max} > \lambda_{\min}$; a sufficient decrease parameter $\gamma \in (0,1)$; and safeguarding parameters $0 < \sigma_1 < \sigma_2 < 1$. Initially, $\lambda_0 \in [\lambda_{\min}, \lambda_{\max}]$ is arbitrary.

Algorithm SPG

```
Set k \leftarrow 0. If x_0 \notin \Omega, replace x_0 by P_\Omega(x_0). While (the stopping criterion is not satisfied) do Compute d_k = P_\Omega(x_k - \lambda_k \, g(x_k)) - x_k, \alpha_k using the line search algorithm described below and x_{k+1} = x_k + \alpha_k d_k. Compute s_k = x_{k+1} - x_k, y_k = g(x_{k+1}) - g(x_k) and \beta_k = \langle s_k, y_k \rangle.
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If (\beta_k \leq 0) set \lambda_{k+1} = \lambda_{\max} else compute \lambda_{k+1} = \min(\lambda_{\max}, \max(\lambda_{\min}, \langle s_k, s_k \rangle / \beta_k)). Set k \leftarrow k+1. end while Set x_* \leftarrow x_k.
```

This algorithm is based on the spectral projected gradient direction $P_{\Omega}(x_k - \lambda_k g(x_k)) - x_k$, where λ_k is the safeguarded "inverse Rayleigh quotient" $\langle s_{k-1}, s_{k-1} \rangle / \langle s_{k-1}, y_{k-1} \rangle$. (Observe that $\langle s_{k-1}, y_{k-1} \rangle / \langle s_{k-1}, s_{k-1} \rangle$ is a Rayleigh quotient corresponding to the average Hessian matrix $\int_0^1 \nabla^2 f(x_{k-1} + ts_{k-1}) dt$.)

The line search is based on a safeguarded quadratic interpolation. The safeguarding procedure acts when the minimum of the one-dimensional quadratic $q(\cdot)$, such that $q(0) = f(x_k)$, $q(\alpha) = f(x_k + \alpha d_k)$, and $\nabla q(0) = \nabla f(x_k)^T d_k$, lies outside $[\sigma_1, \sigma_2 \alpha]$, and not when it lies outside $[\sigma_1 \alpha, \sigma_2 \alpha]$ as usually implemented. This means that, when interpolation tends to reject 90% (for $\sigma_1 = 0.1$) of the original search interval ([0, 1]), we judge that its prediction is not reliable and we prefer the more conservative bisection. This procedure turned out to be more efficient than the classical one. The complete line search procedure is described below.

Line search

```
Compute f_{\max} = \max\{f(x_{k-j}) \mid 0 \leq j \leq \min\{k, m-1\}\}, x_+ \leftarrow x_k + d_k, \delta \leftarrow \langle g(x_k), d_k \rangle and set \alpha \leftarrow 1. While (f(x_+) > f_{\max} + \alpha \gamma \delta) do Compute \alpha_{temp} \leftarrow -\frac{1}{2}\alpha^2\delta/(f(x_+) - f(x_k) - \alpha \delta). If (\alpha_{temp} \geq \sigma_1 and \alpha_{temp} \leq \sigma_2\alpha) set \alpha \leftarrow \alpha_{temp} else set \alpha \leftarrow \alpha/2. Compute x_+ \leftarrow x_k + \alpha d_k. end while Set \alpha_k \leftarrow \alpha.
```

In the case of rejection of the first trial point, the next ones are computed along the same direction. As a consequence, the projection operation is performed only once per iteration.

We use the convergence criteria given by

$$||P_{\Omega}(x_k - g(x_k)) - x_k||_{\infty} \le \epsilon_1 \tag{1}$$

or

$$||P_{\Omega}(x_k - g(x_k)) - x_k||_2 \le \epsilon_2.$$
 (2)

In addition, the algorithm is also stopped when the number of iterations is larger than *maxit* or the number of function evaluations exceeds *maxfun*.

In the experiments presented in the next section, we chose m=10, $\lambda_{\min}=10^{-3}$, $\lambda_{\max}=10^{3}$, $\lambda_{0}=\min(\lambda_{\max},\max(\lambda_{\min},1/\|P_{\Omega}(x_{k}-g(x_{k}))-x_{k}\|_{\infty}))$, $\gamma=10^{-4}$, $\sigma_{1}=0.1$, $\sigma_{2}=0.9$, $\epsilon_{1}=0$, $\epsilon_{2}=10^{-6}$, maxit=1000, maxfun=2000. The stopping criterion (1) was inhibited in order to encourage a fair comparison with an alternative algorithm. For projections onto the feasible set the 2-norm was adopted. In this case, it can be ensured that d_{k} is a descent direction, which is

the essential property that guarantees convergence. Other norms can be used, depending on the structure of the feasible set.

The choice of λ_{\min} , λ_{\max} , λ_0 , ϵ_1 and ϵ_2 is sensitive because these are dimensional parameters. Roughly speaking, the adopted choice is adequate if the problem is scaled in such a way that the Hessian eigenvalues are O(1).

The user must supply the external subroutines EvalF, EvalG, and Proj to evaluate the objective function and its gradient and to project an arbitrary point onto the feasible region. Each gradient evaluation is necessarily preceded by a function evaluation at the same point. For this reason, the user can take advantage of possible common expressions or tests in his/her gradient subroutine implementation. Subroutine Proj depends on the description of the feasible region.

3. TEST PROBLEMS

In Birgin et al. [2000] we showed the performance of SPG on a set of large-scale box-constrained problems from the CUTE collection [Bongartz et al. 1995] and compared it against LANCELOT [Conn et al. 1988, 1991]. The main conclusion of those tests is that SPG is competitive for box-constrained problems. However, the SPG theory allows one to deal with convex constraints, so it is natural to test the method in more general situations. The only restriction is that the projection onto the feasible region must be easy (affordable) to compute. In our software, we leave this task to the user-given subroutine Proj. Since most general nonlinear programming algorithms do not take explicit advantage of the possible simplicity of projections, our feeling is that SPG could outperform general NLP solvers in that case.

Here, we consider a family of *location* problems. Given a set of *npol* disjoint polygons $P_1, P_2, \ldots, P_{npol}$ in \mathbb{R}^2 , we wish to find the point $z^1 \in P_1$ that minimizes the sum of the distances to the other polygons. Therefore, the problem is

$$\min_{z^i, i=1,...,npol} \sum_{i=2}^{npol} \|z^i - z^1\|_2$$

subject to
$$z^i \in P_i$$
, $i = 1, ..., npol$.

Observe that the problem has $2 \times npol$ variables. The number of (linear inequality) constraints is $\sum_{i=1}^{npol} v_i$, where v_i is the number of vertices of polygon P_i . Each constraint defines a half-plane in \mathbb{R}^2 .

constraint defines a half-plane in \mathbb{R}^2 . Let us write $x=(z_1^1,z_2^1,\dots,z_1^{npol},z_2^{npol})$. For projecting x onto the feasible set, observe that we only need to project each z^i separately onto the corresponding polygon P_i . In the projection subroutine, we consider the half-planes that define the polygon. If z^i belongs to all these half-planes, then z^i is the projection onto P_i . Otherwise, we consider the set of half-planes to which z^i does not belong. We project z^i onto these half-planes and discard the projected points that do not belong to P_i . Let A_i be the (finite) set of nondiscarded half-plane projections and let V_i be the set of vertices of P_i . Then, the projection of z^i onto P_i is the point of $A_i \cup V_i$ that is closest to z^i . Both the subroutine for generating the problems and the projection subroutine are included in the test driver for SPG method.

Table I. Problems Set

Problem	# Polygons	# Constraints n			
1	#101/g6115 86	343	174		
$\frac{1}{2}$	86	700	$174 \\ 174$		
3	85	1018	$\frac{174}{172}$		
4	174	701	350		
5	181	1450	364		
6	176	$\frac{1430}{2104}$	354		
7	257	1033	516		
8	256	2103	514		
9	262	3216	526		
10	359	1437	$\frac{320}{720}$		
11	349	2854	700		
12	349	4328	700		
13	435	1743	872		
14	432	3522	866		
15	430	5257	862		
16	935	3729	1872		
17	935 928	7399	1858		
18	940	11200	1882		
19	1841	7362	3684		
20	1825	14574	3652		
$\begin{array}{c c} 20 \\ 21 \end{array}$	1841	22049	3684		
21 22	$\frac{1641}{2766}$	$\frac{22049}{11102}$	5534		
23	$\frac{2700}{2795}$	$\frac{11102}{22511}$	5592		
$\begin{array}{c c} 25 \\ 24 \end{array}$	2831	34445	5664		
25	3779	15104	7560		
26	3815	30699	7632		
27	$\frac{3815}{3852}$	46704	7706		
28	4735	18937	9472		
29	4767	38247	9536		
30	4836	58622	9674		
31	9680	38790	19362		
32	9644	77413	19290		
33	9639	116195	19290		
34	19117	76550	38236		
35	19117	153156	38188		
36	19093	230190	38230		
37	$\frac{19114}{28770}$	230190 115301	57542		
38	28770 28799	230878	57600		
39	28799 28767	250676 346048	57536		
40	38409	153784	76820		
40	38568	309252	77138		
41 42	38506	309252 463718	77014		
42	48004	$\frac{463718}{192152}$	96010		
43	48004 48209	386121	96420		
44	48209 48126	578648	96254		
40	40120	010040	30254		

We generated 45 problems of this class, varying npol and choosing randomly the location of the polygons and the number of vertices of each one. The details of the generation, including the way in which we guarantee empty intersections, are rather tedious but, of course, are available for interested readers. In Table I, we display the main characteristics of each problem (number of polygons,

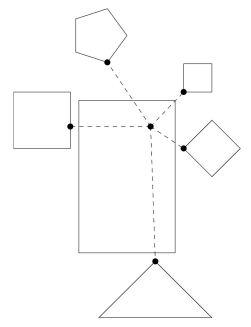


Fig. 1. Six-polygons problem.

number of constraints and dimension of the problem). Figure 1 shows the solution of a small six-polygons problem that has 12 variables and 24 constraints.

The test problems were solved both with SPG and with FSQP, the general nonlinear programming solver described in Zhou and Tits [1997]. (See also Bonnans et al. [1992], Lawrence and Tits [1996], Panier and Tits [1993], and Zhou and Tits [1993].) This is a sequential quadratic programming algorithm that uses feasible iterates and has good global and local convergence properties. For running FSQP, we used the default parameters indicated in the code documentation [Zhou and Tits 1993].

For both methods, we used the origin as initial approximation. Both algorithms found the solutions of the problems and stopped with the diagnostic of convergence. The quality of the solutions was always the same.

Tables II and III display the performance of SPG and FSQP, respectively. The columns mean: *Problem*, problem number; *iter*, iterations needed to reach the solution; *fe*, function evaluations; *Time*, CPU time (seconds); *f*, function value at the solution.

All the experiments were run on a Sun Ultra 60 Workstation with 2 UltraSPARC-II processors at 296-Mhz, $512\,\mathrm{Mb}$ of main memory, and SunOS 5.7 operating system. The compiler was Sun WorkShop Compiler Fortran $77\,4.0$ with flag-O to optimize the code.

We ran only 15 problems using FSQP because, for larger problems, the computation time for this algorithm becomes unaffordable. One should be careful when looking at the time for this algorithm because the implementation FSQP of the feasible sequential quadratic programming algorithm does not take

Table II. SPG Performance

Problem	iter	fe	Time	f
1	14	15	0.01	1.853D+02
$\frac{1}{2}$	14	15	0.01	1.959D+02
3	14	15	0.02	1.914D+02
4	$\overline{24}$	27	0.03	1.945D+02
5	23	26	0.04	2.021D+02
6	20	21	0.05	1.948D+02
7	19	20	0.04	1.952D + 02
8	24	25	0.07	1.989D + 02
9	34	36	0.12	1.948D + 02
10	19	20	0.05	1.969D + 02
11	29	30	0.11	1.969D + 02
12	23	24	0.11	1.943D+02
13	14	15	0.05	1.968D + 02
14	25	26	0.11	1.983D + 02
15	26	29	0.15	1.974D + 02
16	20	21	0.14	1.286D+03
17	30	33	0.30	1.324D + 03
18	28	30	0.36	1.304D+03
19	65	78	0.86	1.311D+03
20	51	63	0.98	1.315D+03
21	42	47	1.07	1.293D+03
22	33	35	0.67	1.308D+03
23	55	67	1.62	1.266D+03
24	80	117	3.22	1.312D+03
25	69	103	1.91	1.311D + 03
26	50	60	2.06	1.301D+03
27	37	42	2.02	1.321D+03
28	144	219	5.02	1.305D+03
29	50	60	2.56	1.306D+03
30	44	47	3.15	1.309D+03
31	18	20	1.39	1.968D+03
32	24	30	2.94	1.966D+03
33	20	21	3.30	1.959D + 03
34	14	15	2.36	1.968D+03
35	15	17	3.59	1.964D + 03
36	19	21	6.09	1.973D+03
37	14	15	3.54	1.973D+03
38	18	20	6.52	1.974D+03
39	13	14	6.19	1.974D+03
40	14	15	4.81	1.975D+03
41	25	28	12.11	1.980D+03
42	$\frac{26}{24}$	26	14.97	1.980D+03
43	18	20	7.54	1.973D+03
44	13	14	8.06	1.978D+03
45	17	19	12.96	1.980D+03
10	<u> </u>	10	12.00	1.000D 00

advantage of sparsity of the matrix of constraints or the Hessian of the objective function. An implementation using sparse factorizations would be much more efficient in terms of computer time. However, we decided to maintain the comparison against FSQP because the results in terms of number of iterations and function evaluations would not change (at least, certainly, could not be better) in a sparse implementation of the method.

Problem iter fe Time 1.853D+021.959D + 021.914D + 021.945D+022.021D+021.948D + 021.952D+021.989D + 021.948D+021.969D + 021.969D + 021.943D+021.968D + 021.983D + 021.974D + 02

Table III. FSQP Performance

Clearly, independently of the linear-algebra savings of sequential quadratic programming (SQP) implementations, any SQP iteration is (much) more expensive than a single iteration of the algorithm described in this paper. So, it is remarkable that the number of iterations used by our algorithm is from 8 to 20 times smaller than the number of iterations used by FSQP. Approximately, the same relation exists between the number of function and gradient evaluations used by both methods. Therefore, the advantage of solving the problems using SPG in place of a general nonlinear programming algorithm is quite impressive.

4. CONCLUSIONS

We have presented a computational algorithm for minimizing functions of many variables restricted to a convex set. The algorithm tends to work well when projections onto the feasible set are easy to compute. The user is required to provide a subroutine to compute these projections. A previous paper [Birgin et al. 2000] shows that the new algorithm is effective for solving many large-scale box-constrained problems. Besides the tests of Birgin et al. [2000], SPG has been shown to be efficient in several applied box-constrained problems; see Birgin et al. [1999a, 1999b], Birgin and Evtushenko [1998], and Mulato et al. [2000]. Here we show that, perhaps, this effectiveness is even more evident when the constraints are given in some other form, provided that projections are not complicated. The key point is that most general nonlinear programming algorithms do not take advantage of the easy-projection property at all.

An interesting family of problems to which SPG can be applied is the norm-constrained regularization problem [Heinkenschloss 1993; Martínez and Santos 1995, 1997; Vogel 1990], defined by

$$\min f(x) \text{ subject to } x^T A x \le r, \tag{3}$$

where A is symmetric positive definite. This problem can be reduced to ball-constrained minimization by a change of variables and, in this case, projections can be trivially computed. A particular case of (3) is the classical trust-region

problem where f is quadratic. Recently (see Lucidi et al. [1998] and Pham Dinh and Hoai An [1998]) procedures for escaping from nonglobal stationary points of this problem have been found and, so, it becomes increasingly important to obtain fast algorithms for finding critical points, especially in the large-scale case. (See Rendl and Wolkowicz [1997], Rojas et al. [2000], and Sorensen [1997].)

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