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SAND2010-1422  
Unlimited Release  
Printed March 2010

## **Poblano v1.0: A Matlab Toolbox for Gradient-Based Optimization**

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# Poblano v1.0: A Matlab Toolbox for Gradient-Based Optimization

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## Abstract

We present Poblano v1.0, a Matlab toolbox for solving gradient-based unconstrained optimization problems. Poblano implements three optimization methods (nonlinear conjugate gradients, limited-memory BFGS, and truncated Newton) that require only first order derivative information. In this paper, we describe the Poblano methods, provide numerous examples on how to use Poblano, and present results of Poblano used in solving problems from a standard test collection of unconstrained optimization problems.

## Acknowledgments

The development of Poblano was supported by Sandia's Laboratory Directed Research & Development (LDRD) program. We thank Dianne O'Leary for providing the Matlab translation of the Moré-Thuente line search from MINPACK; MINPACK was developed by the University of Chicago, as Operator of Argonne National Laboratory.

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# 1 Toolbox Overview

Poblano is a toolbox of large-scale algorithms for nonlinear optimization. The algorithms in Poblano require only first-order derivative information (e.g., gradients for scalar-valued objective functions).

## 1.1 Introduction

Poblano optimizers find local minimizers of scalar-valued objective functions taking vector inputs. Specifically, the problems solved by Poblano optimizers are of the following form:

$$\min_{x \in \mathbb{R}^N} f(x), \quad \text{where } f : \mathbb{R}^N \rightarrow \mathbb{R}. \quad (1)$$

The gradient of the objective function,  $\nabla f(x)$ , is required for all Poblano optimizers. The optimizers converge to a stationary point,  $x^*$ , where

$$\nabla f(x^*) \approx 0.$$

The Moré-Thuente cubic interpolation line search (`cvsrch`) [8], satisfying the strong Wolfe conditions is used to guarantee global convergence of the Poblano optimizers.

The following methods for solving (1) are available in Poblano; see §2 for detailed descriptions.

- Nonlinear conjugate gradient (`ncg`) [9]
  - Uses Fletcher-Reeves, Polak-Ribiere, and Hestenes-Stiefel conjugate direction updates
  - Includes restart strategies based on number of iterations or orthogonality of gradients across iterations
  - Can do steepest descent method as a special case
- Limited-memory BFGS (`lbfgs`) [9]
  - Uses a two-loop recursion for approximate Hessian-gradient products
- Truncated Newton (`tn`) [1]
  - Uses finite differencing for approximate Hessian-vector products

## 1.2 Creating an Objective Function

Functions are passed to Poblano using Matlab function handles. The Matlab function should take a vector as input ( $\mathbf{x} \in \mathbb{R}^N$ ) and return a scalar function value ( $f \in \mathbb{R}$ ) as its first return value and a vector gradient ( $\mathbf{g} \in \mathbb{R}^N$ ) as its second return value. Optimizing

$$f(x) = \sum_{i=1}^N \sin(x_i).$$

requires the following Matlab function (which is the same for any value of  $N$ ):

```
function [f,g] = example0(x)
f = sum(sin(x));
g = cos(x);
```

In the examples that follow, we use a more general function:

$$f(x) = \sum_{i=1}^N \sin(ax_i) \tag{2}$$

where  $a$  is a user-defined parameter. The following `example1` function (distributed with Poblano) evaluates the function and gradient of (2). The input parameter `a` is optional and defaults to  $a = 1$ . In the sections below, we show how to specify  $a$  when the function handle is passed to the optimization method.

```
function [f,g] = example1(x,a)
if nargin < 2
    a = 1;
end
f = sum(sin(a*x));
g = a*cos(a*x);
```

### 1.3 Calling a Poblano Optimizer

Poblano methods have two required inputs as well as optional parameters discussed in the next subsection. The required arguments are 1) a function handle for the objective and gradient calculations, and 2) an initial guess of the solution ( $x_0 \in \mathbb{R}^N$ ). For example, the call below optimizes (2) with  $a = 1$  and  $N = 1$ ; the initial guess is  $x_0 = \pi/4$ :

```
>> out = ncg(@example1, pi/4);
```

Iter	FuncEvals	F(X)	G(X)  /N
0	1	0.70710678	0.70710678
1	6	-0.99999998	0.00017407
2	7	-1.00000000	0.00000000

At each iteration, Poblano prints the iteration number (`Iter`), the total number of function evaluations (`FuncEvals`), the function value at the current iterate (`F(X)`), and the norm of the gradient at the current iterate scaled by the problem size (`||G(X)||/N`). The final iterate is passed back as `out.X`; see §1.5 for details.

Parameterized functions can be optimized using Poblano as well by using more advanced Matlab function handles. We once again consider (2), but this time with  $N = 2$ ,  $a = 3$ , and  $x_0 = [\pi/3 \ \pi/4]^T$ . The call below specifies that  $a = 3$  by passing that as the second argument to `example1`. The choice of  $N = 2$  is implicitly specified since `x0` is in  $\mathbb{R}^2$ .

```
>> out = ncg(@(x) example1(x,3), [pi/3 pi/4]')
```

Iter	FuncEvals	F(X)	G(X)  /N
0	1	0.70710678	1.83711731
1	6	-1.50761836	1.37972368
2	10	-1.83422986	0.82720647
3	13	-1.99999926	0.00182812
4	15	-2.00000000	0.00000001



## 1.4 Optimization Input Parameters

In this section, we discuss the input parameters that are common to all the Poblano methods and give examples of their use. Typing `help poblano_params` lists these parameters. Method-specific parameters are discussed in §2.

Table 1 lists the parameter controlling how much information displayed. For example, we can limit the display to just the final iteration as follows:

```
>> out = ncg(@(x) example1(x,3), [pi/3 pi/4]', 'Display', 'final');
```

Iter	FuncEvals	F(X)	G(X)  /N
4	15	-2.00000000	0.00000001

Table 2 lists the parameters controlling the stopping criteria of the optimization methods. The relative change in the function value is defined as

$$\frac{|f - f_{\text{old}}|}{d} \quad \text{where} \quad d = \begin{cases} f_{\text{old}} & \text{if } f_{\text{old}} \geq \epsilon_{\text{mach}} \\ 1 & \text{if } f_{\text{old}} < \epsilon_{\text{mach}} \end{cases}$$

with  $f$  representing the function value at the current iterate  $F(X)$ ,  $f_{\text{old}}$  the function value at the previous iterate, and  $\epsilon_{\text{mach}}$  machine epsilon (`eps` in Matlab). We can get a more accurate solution by tightening the tolerances, as the following example shows:

```
>> out = lbfgs(@(x) example1(x,2), ones(4,1), 'Display', 'final');
```

Iter	FuncEvals	F(X)	G(X)  /N
2	11	-4.00000000	0.00000001

```
>> out = lbfgs(@(x) example1(x,2), ones(4,1), ...  
'Display', 'final', 'StopTol', 1e-10, 'RelFuncTol', 0);
```

Iter	FuncEvals	F(X)	G(X)  /N
3	13	-4.00000000	0.00000000

Table 3 lists the parameters controlling the information that is saved for each iteration; see §1.5 for further details.

Table 4 lists the parameters controlling the behavior of the Moré-Thuente line search method [8]. Type `help poblano_linesearch` and `help cvsrch` for more details.

An alternative to passing a long list of parameter-value pairs is to specify the parameters in a structure. The easiest way to do this is to extract the default parameters (by passing a single string argument of `'defaults'` as input to the Poblano optimizer to be used) and modify them, as shown in the example below.

```
>> params = ncg('defaults');  
>> params.Display = 'final';  
>> out = ncg(@(x) example1(x,3), [pi/3 pi/4]', params);
```

Iter	FuncEvals	F(X)	G(X)  /N
4	15	-2.00000000	0.00000001

Parameter	Description	Type	Default
Display	Controls amount of printed output 'iter': Display information every iteration 'final': Display information only after final iteration 'off': Display no information	string	'iter'

Table 1: Input parameters for controlling amount of information displayed.

Parameter	Description	Type	Default
MaxIters	Maximum number of iterations	int	100
MaxFuncEvals	Maximum number of function evaluations	int	100
StopTol	Gradient norm stopping tolerance, i.e., the method stops when $\ G(X)\ /N < \text{StopTol}$	double	$10^{-5}$
RelFuncTol	Tolerance on relative function value change	double	$10^{-6}$

Table 2: Input parameters for controlling termination.

Parameter	Description	Type	Default
TraceX	Save iterates ( $X$ )	boolean	false
TraceFunc	Save function values ( $F(X)$ )	boolean	false
TraceRelFunc	Save relative change in function value	boolean	false
TraceGrad	Save the gradients ( $G(X)$ )	boolean	false
TraceGradNorm	Save the norm of the gradients	boolean	false
TraceFuncEvals	Save the number of function evaluations	boolean	false

Table 3: Optimization method input parameters for controlling what information is saved for each iteration.

Parameter	Description	Type	Default
LineSearch_xtol	Stopping tolerance for minimum change in $X$	double	$10^{-15}$
LineSearch_ftol	Stopping tolerance for sufficient decrease condition	double	$10^{-4}$
LineSearch_gtol	Stopping tolerance for directional derivative condition	double	$10^{-2}$
LineSearch_stpmin	Minimum step size allowed	double	$10^{-15}$
LineSearch_stpmax	Maximum step size allowed	double	$10^{15}$
LineSearch_maxfev	Maximum number of iterations allowed	int	20
LineSearch_initialstep	Initial step to be taken in the line search	double	1

Table 4: Optimization method input parameters for controlling line search methods.

## 1.5 Optimization Output Parameters

The methods in Poblano outputs a structure containing the final iteration, function value, etc. The fields are described in Table 5. The `Trace` fields are optional, depending on the input parameters.

Parameter	Description	Type
<code>X</code>	Final iterate	vector, double
<code>F</code>	Function value at <code>X</code>	double
<code>G</code>	Gradient at <code>X</code>	vector, double
<code>Params</code>	Input parameters used for the minimization method (as parsed Matlab <code>inputParser</code> object)	struct
<code>FuncEvals</code>	Number of function evaluations performed	int
<code>Iters</code>	Number of iterations performed (see individual optimization routines for details on what each iteration consists of)	int
<code>ExitFlag</code>	Termination flag, with one of the following values 0 : scaled gradient norm < <code>StopTol</code> input parameter) 1 : maximum number of iterations exceeded 2 : maximum number of function values exceeded 3 : relative change in function value < <code>RelFuncTol</code> input parameter 4 : NaNs found in <code>F</code> , <code>G</code> , or <code>  G  </code>	int
<code>TraceX</code>	History of iterates	matrix, double
<code>TraceFunc</code>	History of function values	vector, double
<code>TraceRelFunc</code>	History of relative change in function value	vector, double
<code>TraceGrad</code>	History of gradients	matrix, double
<code>TraceGradNorm</code>	History of norm of the gradients	vector, double
<code>TraceFuncEvals</code>	History of the number of function evaluations	vector, int

Table 5: Output parameters returned by Poblano optimization methods. The `Trace` parameters are optional, depending on whether or not tracing has been enabled

An example is:

```
>> out = ncg(@(x) example1(x,3), pi/4, 'Display', 'final')
```

```

  Iter  FuncEvals      F(X)          ||G(X)||/N
-----
     2         16    -1.00000000    0.00000147
```

```

out =
  Params: [1x1 inputParser]
  ExitFlag: 0
         X: 70.6858
         F: -1.0000
         G: -1.4734e-06
  FuncEvals: 16
  Iters: 2
```

The exit flag (`out.ExitFlag`) is extremely important. A value other than zero indicates a failure to converge to the specified tolerance. In the following example, the method uses up all its function evaluations:

```
>> out = lbfgs(@(x) example1(x,2), ones(4,1), 'Display', 'final', 'MaxFuncEvals', 5)
```

Iter	FuncEvals	F(X)	G(X)  /N
1	9	-3.99999981	0.00030988

```
out =
```

```

    Params: [1x1 inputParser]
  ExitFlag: 2
        X: [4x1 double]
        F: -4.0000
        G: [4x1 double]
  FuncEvals: 9
    ITERS: 1

```

The number of function evaluations is only checked at the end of each iteration, so it is possible to exceed the allowed number by up to the value specified by the input parameter `LineSearch_maxfev` (see [Table 4](#)).

If tracing is turned on (see [Table 3](#)), then additional outputs will be returned. Here is an example:

```
>> out = ncg(@(x) example1(x,3), [1 2 3]', 'TraceX', true, 'TraceFunc', true, ...
    'TraceRelFunc', true, 'TraceGrad', true, 'TraceGradNorm', true, 'TraceFuncEvals', true)
```

Iter	FuncEvals	F(X)	G(X)  /N
0	1	0.27382300	1.65292785
1	5	-2.65134210	0.79522946
2	11	-2.93709563	0.35196906
3	14	-2.99999975	0.00070154
4	16	-3.00000000	0.00000000

```
out =
```

```

    Params: [1x1 inputParser]
  ExitFlag: 0
        X: [3x1 double]
        F: -3
        G: [3x1 double]
  FuncEvals: 16
    ITERS: 4
    TraceX: [3x5 double]
  TraceFunc: [0.2738 -2.6513 -2.9371 -3.0000 -3]
  TraceRelFunc: [10.6827 0.1078 0.0214 8.2026e-08]
    TraceGrad: [3x5 double]
  TraceGradNorm: [4.9588 2.3857 1.0559 0.0021 2.5726e-09]
  TraceFuncEvals: [1 4 6 3 2]

```

```
>> X = out.X    % Final function value
```

```

X =
    3.6652
   -0.5236
    5.7596

```

```

>> G = out.G      % Final gradient

G =
    1.0e-08 *
    0.0072
   -0.2052
   -0.1550

>> TraceX = out.TraceX      % All iterates (columnwise)

TraceX =
    1.0000    3.7424    3.7441    3.6652    3.6652
    2.0000   -0.6598   -0.6106   -0.5238   -0.5236
    3.0000    5.5239    5.7756    5.7594    5.7596

>> TraceGrad = out.TraceGrad      % All gradients (columnwise)

TraceGrad =
   -2.9700    0.6886    0.7031    0.0001    0.0000
    2.8805   -1.1918   -0.7745   -0.0017   -0.0000
   -2.7334   -1.9486    0.1437   -0.0013   -0.0000

```

One of the outputs returned by the Poblano optimization methods is the `inputParser` object of the input parameters used in that run. That object contains a field called `Results`, which can be passed as the input parameters to another run. For example, this is helpful when running comparisons of methods where only one parameter is changed. Shown below is such an example, where default parameters are used in one run, and the same parameters with just a single change are used in another run.

```

>> out = ncg(@(x) example1(x,3), pi./[4 5 6]');

  Iter  FuncEvals      F(X)          ||G(X)||/N
-----
    0         1      2.65816330      0.77168096
    1         7     -0.63998759      0.78869570
    2        11     -0.79991790      0.60693819
    3        14     -0.99926100      0.03843827
    4        16     -0.99999997      0.00023739
    5        18     -1.00000000      0.00000000

>> params = out.Params.Results;
>> params.Display = 'final';
>> ncg(@(x) example1(x,3), pi./[4 5 6]', params);

  Iter  FuncEvals      F(X)          ||G(X)||/N
-----
    5        18     -1.00000000      0.00000000

```

## 1.6 A More Complex Example Using Matrix Decomposition

The following example is a more complicated function involving matrix variables. Given an  $m \times n$  matrix  $A$ , the goal is to find a rank- $k$  approximation of the form  $UV^T$ . The variable  $x$  is just all the entries of  $U$

and  $V$ , i.e.,  $x = [U(:); V(:)]$ . Thus, the function we wish to optimize is

$$f(x) = \frac{1}{2} \|A - UV^T\|_F^2 \quad (3)$$

where  $A$  and  $k$  have been specified. The gradient of  $f$  is

$$\begin{aligned} \nabla_U f &= -(A - UV^T)V \\ \nabla_V f &= -(A - UV^T)^T U. \end{aligned}$$

Listed below are the contents of the `example2.m` file distributed with the Poblano code. The input `Data` contains  $A$  and the desired rank,  $k$ . This example illustrates how  $x$  is converted to  $U$  and  $V$  for the function and gradient computations.

```
function [f,g] = example2(x,Data)

% Data setup
[m,n] = size(Data.A);
k = Data.rank;
U = reshape(x(1:m*k), m, k);
V = reshape(x(m*k+1:m*k+n*k), n, k);

% Function value (residual)
AmUVt = Data.A - U * V';
f = 0.5 * norm(AmUVt, 'fro')^2;

% First derivatives computed in matrix form
g = zeros((m+n)*k,1);
g(1:m*k) = -reshape(AmUVt * V, m*k, 1);
g(m*k+1:end) = -reshape(AmUVt' * U, n*k, 1);
```

The function `example2_init.m` (included with Poblano) generates a test problem and starting point.

```
>> randn('state',0);
>> m = 4; n = 3; k = 2;
>> [x0,Data] = example2_init(m,n,k)

x0 =
    -0.588316543014189
         2.1831858181971
    -0.136395883086596
         0.11393131352081
         1.06676821135919
    0.0592814605236053
    -0.095648405483669
    -0.832349463650022
         0.29441081639264
        -1.3361818579378
         0.714324551818952
         1.62356206444627
    -0.691775701702287
         0.857996672828263
```

```
Data =
    rank: 2
      A: [4x3 double]

>> out = ncg(@(x) example2(x,Data), x0, 'RelFuncTol', 1e-16, 'StopTol', 1e-8, ...
    'MaxFuncEvals',1000,'Display','final')
```

Iter	FuncEvals	F(X)	G(X)  /N
29	67	0.28420491	0.00000001

```
out =
    Params: [1x1 inputParser]
    ExitFlag: 0
      X: [14x1 double]
      F: 0.284204907556308
      G: [14x1 double]
    FuncEvals: 67
      Iters: 29
```

The function `example2_extract.m` extracts the  $U$  and  $V$  matrices from  $x$ . We can verify that we have found the optimal solution since the Euclidean norm of the difference between the matrix and the approximate solution is equal to the  $k + 1$  singular value of  $A$  [4, Theorem 2.5.3].

```
>> [U,V] = example2_extract(m,n,k,out.X);
>> norm_diff = norm(Data.A-U*V')
```

```
norm_diff =
    0.753929582330216
```

```
>> sv = svd(Data.A);
>> sv_kp1 = sv(k+1)
```

```
sv_kp1 =
    0.753929582330215
```

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## 2 Optimization Methods

### 2.1 Nonlinear Conjugate Gradient

Nonlinear conjugate gradient (NCG) methods [9] are used to solve unconstrained nonlinear optimization problems. They are extensions of the conjugate gradient iterative method for solving linear systems adapted to solve unconstrained nonlinear optimization problems. The Poblano function for the nonlinear conjugate gradient methods is called `n cg`. The general steps of the NCG method are given below in pseudo-code:

1. Input:  $x_0$ , a starting point
2. Evaluate  $f_0 = f(x_0), g_0 = \nabla f(x_0)$
3. Set  $p_0 = -g_0, i = 0$
4. **while**  $\|g_i\| > 0$
5.     Compute a step length  $\alpha_i$  and set  $x_{i+1} = x_i + \alpha_i p_i$
6.     Set  $g_i = \nabla f(x_{i+1})$
7.     Compute  $\beta_{i+1}$
8.     Set  $p_{i+1} = -g_{i+1} + \beta_{i+1} p_i$
9.     Set  $i = i + 1$
10. **end while**
11. Output:  $x_i \approx x^*$

In cases where the update coefficient is negative, i.e.,  $\beta_{i+1} < 0$ , it is set to 0 to avoid directions that are not descent directions [9].

The choice of  $\beta_{i+1}$  in Step 7 leads to different NCG methods. The update methods for  $\beta_{i+1}$  are listed in Table 6. The special case of  $\beta_{i+1} = 0$  leads to the steepest descent method [9]; see Table 7.

Update Type	Update Formula
Fletcher-Reeves [3]	$\beta_{i+1} = \frac{g_{i+1}^T - g_{i+1}}{g_i^T g_i}$
Polak-Ribiere [12]	$\beta_{i+1} = \frac{g_{i+1}^T (g_{i+1} - g_i)}{g_i^T g_i}$
Hestenes-Stiefel [5]	$\beta_{i+1} = \frac{g_{i+1}^T (g_{i+1} - g_i)}{p_i^T (g_{i+1} - g_i)}$

Table 6: Conjugate direction updates available in Poblano.

The NCG iterations are restarted every  $k$  iterations, where  $k$  is specified via the `RestartIters` parameter. The default is 20.

Another restart modification that was suggested by Nocedal and Wright [9] is taking a step in the direction of steepest descent when two consecutive gradients are far from orthogonal. Specifically, a steepest descent step is taken whenever

$$\frac{|g_{i+1}^T g_i|}{\|g_{i+1}\|} \geq \nu$$

where  $\nu$  is specified via the `RestartNWTol` parameter. This modification is off by default, but can be used by setting `RestartNW` to `true`.

The input parameters specific to the `n cg` method are presented in Table 7.

## 2.2 Limited-memory BFGS

Limited-memory quasi-Newton methods [9] are a class of methods that compute and/or maintain simple, compact approximations of the Hessian matrices of second derivatives, which are used determining search directions. Poblano includes the limited-memory BFGS (L-BFGS) method, a variant of these methods whose Hessian approximations are based on the BFGS method (see [2] for more details). The Poblano function for the L-BFGS method is called `lbfgs`. The general steps of L-BFGS methods are given below in pseudo-code [9]:

1. Input:  $x_0$ , a starting point;  $M > 0$ , an integer
2. Evaluate  $f_0 = f(x_0)$ ,  $g_0 = \nabla f(x_0)$
3. Set  $p_0 = -g_0$ ,  $\gamma_0 = 1$ ,  $i = 0$
4. **while**  $\|g_i\| > 0$
5.     Choose an initial Hessian approximation:  $H_i^0 = \gamma_i I$
6.     Compute a step direction  $p_i = -r$  using `TwoLoopRecursion` method
7.     Compute a step length  $\alpha_i$  and set  $x_{i+1} = x_i + \alpha_i p_i$
8.     Set  $g_i = \nabla f(x_{i+1})$
9.     **if**  $i > M$
10.         Discard vectors  $\{s_{i-m}, y_{i-m}\}$  from storage
11.     **end if**
12.     Set and store  $s_i = x_{i+1} - x_i$  and  $y_i = g_{i+1} - g_i$
13.     Set  $i = i + 1$
14. **end while**
15. Output:  $x_i \approx x^*$

In Step 6 in the above method, the computation of the step direction is performed using the following method (assume we are at iteration  $i$ ) [9]:

- TwoLoopRecursion**
1.  $q = g_i$
  2. **for**  $k = i - 1, i - 2, \dots, i - m$
  3.      $a_k = (s_k^T q) / (y_k^T s_k)$
  4.      $q = q - a_k y_k$
  5. **end for**
  6.  $r = H_i^0 q$
  7. **for**  $k = i - m, i - m + 1, \dots, i - 1$
  8.      $b = (y_k^T r) / (y_k^T s_k)$
  9.      $r = r + (a_k - b) s_k$
  10. **end for**
  11. Output:  $r = H_i g_i$

The input parameters specific to the `lbfgs` method are presented in [Table 8](#).

## 2.3 Truncated Newton

Truncated Newton (TN) methods for minimization are Newton methods in which the Newton direction is only approximated at each iteration (thus reducing computation). Furthermore, the Poblano implementation of the truncated Newton method does not require an explicit Hessian matrix in the computation of the approximate Newton direction (thus reducing storage requirements). The Poblano function for the truncated

Newton method is called `tn`. The general steps of TN methods are given below in pseudo-code [1]:

1. Input:  $x_0$ , a starting point
2. Evaluate  $f_0 = f(x_0), g_0 = \nabla f(x_0)$
3. Set  $i = 0$
4. **while**  $\|g_i\| > 0$
5.     Compute the conjugate gradient stopping tolerance,  $\eta_i$
6.     Compute  $p_i$  by solving  $\nabla^2 f(x_i)p = -g_i$  using a linear conjugate gradient (CG) method
7.     Compute a step length  $\alpha_i$  and set  $x_{i+1} = x_i + \alpha_i p_i$
8.     Set  $g_i = \nabla f(x_{i+1})$
9.     Set  $i = i + 1$
10. **end while**
11. Output:  $x_i \approx x^*$

In Step 5, the linear conjugate gradient (CG) method stopping tolerance is allowed to change at each iteration. The input parameter `CGTolType` determines how  $\eta_i$  is computed.

In Step 6, we note the following.

- One of Matlab's CG methods is used to solve for  $p_i$ : `symmlq` (designed for symmetric indefinite systems) or `pcg` (the classical CG method for symmetric positive definite systems). The input parameter `CGSolver` controls the choice of CG method to use.
- The maximum number of CG iterations is specified using the input parameter `CGIters`.
- The CG method stops when  $\| -g_i - \nabla^2 f(x_i)p_i \| \leq \eta_i \|g_i\|$  .
- In the CG method, matrix-vector products involving  $\nabla^2 f(x_i)$  times a vector  $v$  are approximated using the following finite difference approximation [1]:

$$\nabla^2 f(x_i)v \approx \frac{\nabla f(x_i + \sigma v) - \nabla f(x_i)}{\sigma}$$

The difference step,  $\sigma$ , is specified using the input parameter `HessVecFDStep`. The computation of the finite difference approximation is performed using the `hessvec_fd` provided with Poblano.

The input parameters specific to the `tn` method are presented in [Table 9](#).

Parameter	Description	Type	Default
Update	Conjugate direction update 'FR': Fletcher-Reeves 'PR': Polak-Ribiere 'HS': Hestenes-Stiefel 'SD': Steepest Descent	string	'PR'
RestartIters	Number of iterations to run before conjugate direction restart	int	20
RestartNW	Flag to use restart heuristic of Nocedal and Wright	boolean	false
RestartNWTol	Tolerance for Nocedal and Wright restart heuristic	double	0.1

Table 7: Method-specific parameters for Poblano's `ncg` optimization method.

Parameter	Description	Type	Default
M	Limited memory parameter (i.e., number of vectors $s$ and $y$ to store from previous iterations)	int	5

Table 8: Method-specific parameters for Poblano's `lbfgs` optimization method.

Parameter	Description	Type	Default
CGSolver	Matlab CG method used to solve for search direction 'symmlq' : Symmetric LQ method [11] 'pcg' : Classical CG method [5]	string	'symmlq'
CGIters	Maximum number of CG iterations allowed	int	5
CGTolType	CG stopping tolerance type used 'quadratic' : $\ R\ /\ G\  < \min(0.5, \ G\ )$ 'superlinear' : $\ R\ /\ G\  < \min(0.5, \sqrt{\ G\ })$ 'fixed' : $\ R\  < \text{CGTol}$ $R$ is the residual and $G$ is the gradient	string	'quadratic'
CGTol	CG stopping tolerance when <code>CGTolType</code> is 'fixed'	double	$10^{-6}$
HessVecFDStep	Hessian vector product finite difference step 0 : Use iterate-based step: $10^{-8}(1 + \ X\ _2)$ > 0 : Fixed value to use as the difference step	double	$10^{-10}$

Table 9: Method-specific parameters for Poblano's `tn` optimization method.

### 3 Checking Gradient Calculations

Analytic gradients can be checked using finite difference approximations. The Poblano function `gradientcheck` computes the gradient approximations and compares the results to the analytic gradient using a user-supplied objective function/gradient M-file. The user can choose one of several difference formulas as well as the difference step used in the computations.

#### 3.1 Difference Formulas

The difference formulas for approximating the gradients in Poblano are listed in [Table 10](#). For more details on the formulas, see [\[9\]](#).

Formula Type	Formula
Forward Differences	$\frac{\partial f}{\partial x_i}(x) \approx \frac{f(x + he_i) - f(x)}{h}$
Backward Differences	$\frac{\partial f}{\partial x_i}(x) \approx \frac{f(x) - f(x - he_i)}{h}$
Centered Differences	$\frac{\partial f}{\partial x_i}(x) \approx \frac{f(x + he_i) - f(x - he_i)}{2h}$

Note:  $e_i$  is a vector the same size as  $x$  with a 1 in element  $i$  and zeros elsewhere, and  $h$  is a user-defined parameter.

Table 10: Difference formulas available in Poblano for checking user-defined gradients.

The type of finite differences to use is specified using the `DifferenceType` input parameter, and the value of  $h$  is specified using the `DifferenceStep` input parameter. For a detailed discussion on the impact of the choice of  $h$  on the quality of the approximation, see [\[10\]](#).

#### 3.2 Gradient Check Input Parameters

The input parameters available for the `gradientcheck` function are presented in [Table 11](#).

Parameter	Description	Type	Default
<code>DifferenceType</code>	Difference formula to use 'forward': $g_i = (f(x + he_i) - f(x))/h$ 'backward': $g_i = (f(x) - f(x - he_i))/h$ 'centered': $g_i = (f(x + he_i) - f(x - he_i))/(2h)$	string	'forward'
<code>DifferenceStep</code>	Value of $h$ in difference formulae	double	$10^{-8}$

Table 11: Input parameters for Poblano's `gradientcheck` function.

### 3.3 Gradient Check Output Parameters

The fields in the output structure generated by the `gradientcheck` function are presented in [Table 12](#).

Parameter	Description	Type
G	Analytic gradient	vector, double
GFD	Finite difference approximation of gradient	vector, double
MaxDiff	Maximum difference between G and GFD	double
MaxDiffInd	Index of maximum difference between G and GFD	int
NormGradientDiffs	Norm of the difference between gradients: $\ G - GFD\ _2$	double
GradientDiffs	G - GFD	vector, double
Params	Input parameters used to compute approximations	struct

Table 12: Output parameters generated by Poblano's `gradientcheck` function.

### 3.4 Examples

We use `example1` (distributed with Poblano) to illustrate how to use the `gradientcheck` function to check user-supplied gradients. The user provides a function handle to the M-file containing their function and gradient computations, a point at which to check the gradients, and the type of difference formula to use. Below are examples of running the gradient check using each of the difference formulas.

```
>> outFD = gradientcheck(@(x) example1(x,3), pi./[4 5 6]','DifferenceType','forward')

outFD =
         G: [3x1 double]
        GFD: [3x1 double]
    MaxDiff: 6.4662e-08
  MaxDiffInd: 1
NormGradientDiffs: 8.4203e-08
  GradientDiffs: [3x1 double]
        Params: [1x1 struct]

>> outBD = gradientcheck(@(x) example1(x,3), pi./[4 5 6]','DifferenceType','backward')

outBD =
         G: [3x1 double]
        GFD: [3x1 double]
    MaxDiff: -4.4409e-08
  MaxDiffInd: 3
NormGradientDiffs: 5.2404e-08
  GradientDiffs: [3x1 double]
        Params: [1x1 struct]
```

```
>> outCD = gradientcheck(@(x) example1(x,3), pi./[4 5 6]','DifferenceType','centered')

outCD =
           G: [3x1 double]
          GFD: [3x1 double]
        MaxDiff: 2.0253e-08
        MaxDiffInd: 1
    NormGradientDiffs: 2.1927e-08
        GradientDiffs: [3x1 double]
           Params: [1x1 struct]
```

Note the different gradients produced using the various differencing formulas:

```
>> [outFD.G outFD.GFD outBD.GFD outCD.GFD]

ans =
-2.121320343559642 -2.121320408221550 -2.121320319403708 -2.121320363812629
-0.927050983124842 -0.927051013732694 -0.927050969323773 -0.927050991528233
 0.000000000000000 -0.000000044408921  0.000000044408921  0
```

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## 4 Numerical Experiments

To demonstrate the performance of the Poblano methods, we present results of runs of the different methods in this section. All experiments were performed using Matlab 7.9 on a Linux Workstation (RedHat 5.2) with 2 Quad-Core Intel Xeon 3.0GHz processors and 32GB RAM.

### 4.1 Description of Test Problems

The test problems used in the experiments presented here are from the Moré, Garbow, and Hillstom collection, which is described in detail in [7]. The Matlab code used for these problems, along with the known optima, is provided as part of the SolvOpt optimization software [6], available at <http://www.kfunigraz.ac.at/imawww/kuntsevich/solvopt/>. (SolvOpt is an implementation of Shor’s  $r$ -algorithm for non-smooth optimization [13] with additional heuristics for computing search directions and stopping criteria, but has been tested using this collection, which contains continuously differentiable problems.) There are 34 test problems in this collection.

### 4.2 Results

Results in this section are for optimization runs using the default parameters, with the exception of the stopping criteria parameters, which were changed to allow more computation and find more accurate solutions. The parameters changed from their default values are as follows:

```
params.Display = 'off'  
params.MaxIters = 20000;  
params.MaxFuncEvals = 50000;  
params.RelFuncTol = 1e-16;  
params.StopTol = 1e-12;
```

For the results presented in this section, the function value computed by the Poblano methods is denoted by  $\hat{F}^*$ , the solution (i.e., the best known function value reported in the literature) is denoted by  $F^*$ , and the (relative) error is  $|F^* - \hat{F}^*| / \max\{1, |F^*|\}$ .

For this test collection, we say the problem is solved if the relative error is less than  $10^{-8}$ . We see that the Poblano methods solve most of the problems using the default input parameters, but have difficulty with a few that require particular parameter settings. Specifically, the initial step size in the line search method appears to be the most sensitive parameter across the methods. More investigation into the effective use of this parameter for other problem classes is planned for future work. Below are more details of the results for the different Poblano methods.

#### 4.2.1 Nonlinear Conjugate Gradient

The results of the tests for the `ncg` method using Polak-Ribiere (PR) conjugate direction updates are presented in Table 13. Note that several problems (3, 6, 10, 11, 17, 20, 27 and 31) were not solved using the parameters listed above. With more testing, specifically with different values of the initial step size used in the line search (specified using the `LineSearch_initialstep` parameter) and the number of iterations to perform before restarting the conjugate directions using a step in the steepest direction (specified using the `RestartIters` parameter), solutions to all problems except #10 were found. Table 14 presents parameter choices leading to successful runs of the `ncg` method using PR conjugate direction updates.

The results of the tests for the `ncg` method using Hestenes-Stiefel (HS) conjugate direction updates are presented in [Table 15](#). Again, several problems (3, 6, 10, 11 and 31) were not solved using the parameters listed above. [Table 16](#) presents parameter choices leading to successful runs of the `ncg` method using HS conjugate direction updates. Note that we did not find parameters to solve problem #10 with this method either.

Finally, the results of the tests for the `ncg` method using Fletcher-Reeves (FR) conjugate direction updates are presented in [Table 17](#). We see that even more problems (3, 6, 10, 11, 17, 18, 20 and 31) were not solved using the parameters listed above. [Table 18](#) presents parameter choices leading to successful runs of the `ncg` method using FR conjugate direction updates. Note that we did not find parameters to solve problem #10 with this method either.

#### 4.2.2 Limited-memory BFGS

The results of the tests for the `lbfgs` method are presented in [Table 19](#). Compared to the `ncg` methods, fewer problems (6, 11, 18 and 31) were not solved using the parameters listed above with the `lbfgs` method. Most notably, `lbfgs` was able to solve problem #10, illustrating that some problems are better suited to the different Poblano methods. With more testing, specifically with different values of the initial step size used in the line search (specified using the `LineSearch_initialstep` parameter), solutions to all problems were found. [Table 20](#) presents parameter choices leading to successful runs of the `lbfgs` method.

#### 4.2.3 Truncated Newton

The results of the tests for the `tn` method are presented in [Table 21](#). Note that several problems (10, 11, 20 and 31) were not solved using this method. However, problem #6 was solved (which was not solved using any of the other Poblano methods), again illustrating that some problems are better suited to `tn` than the other Poblano methods. With more testing, specifically with different values of the initial step size used in the line search (specified using the `LineSearch_initialstep` parameter), solutions to all problems were found, with the exception of problem #10. [Table 22](#) presents parameter choices leading to successful runs of the `tn` method.

#	Exit	Iters	FuncEvals	$\hat{F}^*$	$F^*$	Error
1	3	20	90	1.3972e-18	0.0000e+00	1.3972e-18
2	3	11	71	4.8984e+01	4.8984e+01	5.8022e-16
3	3	3343	39373	6.3684e-07	0.0000e+00	<b>6.3684e-07</b>
4	3	14	126	1.4092e-21	0.0000e+00	1.4092e-21
5	3	12	39	1.7216e-17	0.0000e+00	1.7216e-17
6	0	1	8	2.0200e+03	1.2436e+02	<b>1.5243e+01</b>
7	3	53	158	5.2506e-18	0.0000e+00	5.2506e-18
8	3	28	108	8.2149e-03	8.2149e-03	1.2143e-17
9	0	5	14	1.1279e-08	1.1279e-08	2.7696e-14
10	3	262	1232	9.5458e+04	8.7946e+01	<b>1.0844e+03</b>
11	0	1	2	3.8500e-02	0.0000e+00	<b>3.8500e-02</b>
12	0	15	62	3.9189e-32	0.0000e+00	3.9189e-32
13	3	129	352	1.5737e-16	0.0000e+00	1.5737e-16
14	3	47	155	2.7220e-18	0.0000e+00	2.7220e-18
15	3	58	165	3.0751e-04	3.0751e-04	3.9615e-10
16	3	34	168	8.5822e+04	8.5822e+04	4.3537e-09
17	3	3122	7970	5.5227e-05	5.4649e-05	<b>5.7842e-07</b>
18	3	1181	2938	2.1947e-16	0.0000e+00	2.1947e-16
19	3	381	876	4.0138e-02	4.0138e-02	2.9355e-10
20	3	14842	30157	1.4233e-06	1.4017e-06	<b>2.1546e-08</b>
21	3	20	90	6.9775e-18	0.0000e+00	6.9775e-18
22	3	129	352	1.5737e-16	0.0000e+00	1.5737e-16
23	0	90	415	2.2500e-05	2.2500e-05	2.4991e-11
24	3	142	520	9.3763e-06	9.3763e-06	7.4196e-15
25	3	5	27	7.4647e-25	0.0000e+00	7.4647e-25
26	3	47	144	2.7951e-05	2.7951e-05	2.1878e-13
27	3	2	8	8.2202e-03	0.0000e+00	<b>8.2202e-03</b>
28	3	79	160	1.4117e-17	0.0000e+00	1.4117e-17
29	3	7	16	9.1648e-19	0.0000e+00	9.1648e-19
30	3	30	75	1.0410e-17	0.0000e+00	1.0410e-17
31	3	41	97	3.0573e+00	0.0000e+00	<b>3.0573e+00</b>
32	0	2	5	1.0000e+01	1.0000e+01	0.0000e+00
33	3	2	5	4.6341e+00	4.6341e+00	0.0000e+00
34	3	2	5	6.1351e+00	6.1351e+00	0.0000e+00

Table 13: Results of `ncg` using PR updates on the Moré, Garbow, Hillstrom test collection. Errors greater than  $10^{-8}$  are highlighted in bold, indicating that a solution was not found within the specified tolerance.

#	Input Parameters	Iters	FuncEvals	$\hat{F}^*$	Error
3	LineSearch.initialstep = 1e-5 RestartIters = 40	19	152	9.9821e-09	9.9821e-09
6	LineSearch.initialstep = 1e-5 RestartIters = 40	19	125	1.2436e+02	3.5262e-11
11	LineSearch.initialstep = 1e-2 RestartIters = 40	407	2620	2.7313e-14	2.7313e-14
17	LineSearch.initialstep = 1e-3	714	1836	5.4649e-05	1.1032e-11
20	LineSearch.initialstep = 1e-1 RestartIters = 50	3825	8066	1.4001e-06	1.6303e-09
27	LineSearch.initialstep = 0.5	11	38	1.3336e-25	1.3336e-25
31	LineSearch.initialstep = 1e-7	17	172	1.9237e-18	1.9237e-18

Table 14: Parameter changes that lead to solutions using `ncg` with PR updates on the Moré, Garbow, Hillstrom test collection.

#	Exit	Iters	FuncEvals	$\hat{F}^*$	$F^*$	Error
1	3	20	89	1.0497e-16	0.0000e+00	1.0497e-16
2	3	10	69	4.8984e+01	4.8984e+01	2.9011e-16
3	2	4329	50001	4.4263e-08	0.0000e+00	<b>4.4263e-08</b>
4	3	9	83	1.3553e-18	0.0000e+00	1.3553e-18
5	3	12	39	4.3127e-19	0.0000e+00	4.3127e-19
6	0	1	8	2.0200e+03	1.2436e+02	<b>1.5243e+01</b>
7	3	27	98	1.8656e-20	0.0000e+00	1.8656e-20
8	3	25	92	8.2149e-03	8.2149e-03	1.2143e-17
9	3	5	31	1.1279e-08	1.1279e-08	2.7696e-14
10	3	49	332	1.0856e+05	8.7946e+01	<b>1.2334e+03</b>
11	0	1	2	3.8500e-02	0.0000e+00	<b>3.8500e-02</b>
12	0	15	63	5.8794e-30	0.0000e+00	5.8794e-30
13	3	84	258	2.7276e-17	0.0000e+00	2.7276e-17
14	3	44	152	3.4926e-24	0.0000e+00	3.4926e-24
15	3	50	165	3.0751e-04	3.0751e-04	3.9615e-10
16	3	39	176	8.5822e+04	8.5822e+04	4.3537e-09
17	3	942	2754	5.4649e-05	5.4649e-05	4.3964e-12
18	3	748	1977	2.7077e-18	0.0000e+00	2.7077e-18
19	3	237	607	4.0138e-02	4.0138e-02	2.9356e-10
20	3	4685	9977	1.3999e-06	1.4017e-06	1.7946e-09
21	0	22	93	6.1630e-32	0.0000e+00	6.1630e-32
22	3	84	258	2.7276e-17	0.0000e+00	2.7276e-17
23	0	83	381	2.2500e-05	2.2500e-05	2.4991e-11
24	3	170	691	9.3763e-06	9.3763e-06	7.3568e-15
25	3	5	47	7.4570e-25	0.0000e+00	7.4570e-25
26	3	43	144	2.7951e-05	2.7951e-05	2.1878e-13
27	3	9	40	2.5738e-20	0.0000e+00	2.5738e-20
28	3	33	70	8.3475e-18	0.0000e+00	8.3475e-18
29	3	7	16	9.1699e-19	0.0000e+00	9.1699e-19
30	3	29	73	3.4886e-18	0.0000e+00	3.4886e-18
31	3	33	117	3.0573e+00	0.0000e+00	<b>3.0573e+00</b>
32	0	2	5	1.0000e+01	1.0000e+01	0.0000e+00
33	3	2	5	4.6341e+00	4.6341e+00	0.0000e+00
34	3	2	5	6.1351e+00	6.1351e+00	0.0000e+00

Table 15: Results of `ncg` using HS updates on the Moré, Garbow, Hillstrom test collection. Errors greater than  $10^{-8}$  are highlighted in bold, indicating that a solution was not found within the specified tolerance.

#	Input Parameters	Iters	FuncEvals	$\hat{F}^*$	Error
3	LineSearch.initialstep = 1e-5 RestartIters = 40	57	395	3.3512e-09	3.3512e-09
6	LineSearch.initialstep = 1e-5	18	105	1.2436e+02	3.5262e-11
11	LineSearch.initialstep = 1e-2 RestartIters = 40	548	3288	8.1415e-15	8.1415e-15
31	LineSearch.initialstep = 1e-7	17	172	1.6532e-18	1.6532e-18

Table 16: Parameter changes that lead to solutions using `ncg` with HS updates on the Moré, Garbow, Hillstrom test collection.

#	Exit	Iters	FuncEvals	$\hat{F}^*$	$F^*$	Error
1	3	41	199	2.4563e-17	0.0000e+00	2.4563e-17
2	3	25	104	4.8984e+01	4.8984e+01	4.3517e-16
3	3	74	538	1.2259e-03	0.0000e+00	<b>1.2259e-03</b>
4	3	40	304	2.9899e-13	0.0000e+00	2.9899e-13
5	3	27	76	4.7847e-17	0.0000e+00	4.7847e-17
6	0	1	8	2.0200e+03	1.2436e+02	<b>1.5243e+01</b>
7	3	48	153	3.5829e-19	0.0000e+00	3.5829e-19
8	3	46	144	8.2149e-03	8.2149e-03	3.4694e-18
9	3	7	72	1.1279e-08	1.1279e-08	2.7696e-14
10	3	388	2098	3.0166e+04	8.7946e+01	<b>3.4201e+02</b>
11	0	1	2	3.8500e-02	0.0000e+00	<b>3.8500e-02</b>
12	3	19	65	2.6163e-18	0.0000e+00	2.6163e-18
13	3	308	720	1.8955e-16	0.0000e+00	1.8955e-16
14	3	53	219	3.0620e-17	0.0000e+00	3.0620e-17
15	3	83	232	3.0751e-04	3.0751e-04	3.9615e-10
16	3	42	206	8.5822e+04	8.5822e+04	4.3537e-09
17	3	522	2020	5.5389e-05	5.4649e-05	<b>7.4046e-07</b>
18	3	278	644	5.6432e-03	0.0000e+00	<b>5.6432e-03</b>
19	3	385	1003	4.0138e-02	4.0138e-02	2.9355e-10
20	3	14391	29061	2.7316e-06	1.4017e-06	<b>1.3298e-06</b>
21	3	41	199	1.2282e-16	0.0000e+00	1.2282e-16
22	3	308	720	1.8955e-16	0.0000e+00	1.8955e-16
23	3	192	783	2.2500e-05	2.2500e-05	2.4991e-11
24	3	1239	4256	9.3763e-06	9.3763e-06	7.3782e-15
25	0	5	29	1.7256e-31	0.0000e+00	1.7256e-31
26	3	60	187	2.7951e-05	2.7951e-05	2.1877e-13
27	3	10	34	1.0518e-16	0.0000e+00	1.0518e-16
28	3	112	226	1.7248e-16	0.0000e+00	1.7248e-16
29	3	7	16	1.9593e-18	0.0000e+00	1.9593e-18
30	3	30	74	1.1877e-17	0.0000e+00	1.1877e-17
31	3	58	166	3.0573e+00	0.0000e+00	<b>3.0573e+00</b>
32	0	2	5	1.0000e+01	1.0000e+01	3.5527e-16
33	3	2	5	4.6341e+00	4.6341e+00	0.0000e+00
34	3	2	5	6.1351e+00	6.1351e+00	0.0000e+00

Table 17: Results of `ncg` using FR updates on the Moré, Garbow, Hillstrom test collection. Errors greater than  $10^{-8}$  are highlighted in bold, indicating that a solution was not found within the specified tolerance.

#	Input Parameters	Iters	FuncEvals	$\hat{F}^*$	Error
3	LineSearch_initialstep = 1e-5 RestartIters = 50	128	422	2.9147e-09	2.9147e-09
6	LineSearch_initialstep = 1e-5	52	257	1.2436e+02	3.5262e-11
11	LineSearch_initialstep = 1e-2 RestartIters = 40	206	905	4.3236e-13	4.3236e-13
17	LineSearch_initialstep = 1e-3 RestartIters = 40	421	1012	5.4649e-05	1.7520e-11
18	LineSearch_initialstep = 1e-4	1898	12836	3.2136e-17	3.2136e-17
20	LineSearch_initialstep = 1e-1 RestartIters = 50	3503	7262	1.4001e-06	1.6392e-09
31	LineSearch_initialstep = 1e-7	16	162	8.2352e-18	8.2352e-18

Table 18: Parameter changes that lead to solutions using `ncg` with FR updates on the Moré, Garbow, Hillstrom test collection.

#	Exit	Iters	FuncEvals	$\hat{F}^*$	$F^*$	Error
1	3	43	145	1.5549e-20	0.0000e+00	1.5549e-20
2	3	14	73	4.8984e+01	4.8984e+01	4.3517e-16
3	3	206	730	5.8432e-20	0.0000e+00	5.8432e-20
4	3	14	29	7.8886e-31	0.0000e+00	7.8886e-31
5	3	20	55	1.2898e-24	0.0000e+00	1.2898e-24
6	0	1	8	2.0200e+03	1.2436e+02	<b>1.5243e+01</b>
7	0	40	103	8.6454e-28	0.0000e+00	8.6454e-28
8	0	27	63	8.2149e-03	8.2149e-03	1.7347e-18
9	0	4	9	1.1279e-08	1.1279e-08	2.7696e-14
10	3	705	2406	8.7946e+01	8.7946e+01	4.2594e-13
11	0	1	2	3.8500e-02	0.0000e+00	<b>3.8500e-02</b>
12	3	24	76	3.2973e-20	0.0000e+00	3.2973e-20
13	3	77	165	1.7747e-17	0.0000e+00	1.7747e-17
14	3	66	187	1.2955e-19	0.0000e+00	1.2955e-19
15	0	29	72	3.0751e-04	3.0751e-04	3.9615e-10
16	3	22	96	8.5822e+04	8.5822e+04	4.3537e-09
17	3	221	602	5.4649e-05	5.4649e-05	9.7483e-13
18	3	56	165	5.6556e-03	0.0000e+00	<b>5.6556e-03</b>
19	3	267	713	4.0138e-02	4.0138e-02	2.9355e-10
20	3	5118	10752	1.3998e-06	1.4017e-06	1.9194e-09
21	3	44	146	2.8703e-24	0.0000e+00	2.8703e-24
22	3	77	165	1.7747e-17	0.0000e+00	1.7747e-17
23	0	218	786	2.2500e-05	2.2500e-05	2.4991e-11
24	3	458	1492	9.3763e-06	9.3763e-06	7.3554e-15
25	0	4	31	1.4730e-29	0.0000e+00	1.4730e-29
26	3	41	135	2.7951e-05	2.7951e-05	2.1879e-13
27	3	15	37	9.7350e-24	0.0000e+00	9.7350e-24
28	3	67	140	1.2313e-16	0.0000e+00	1.2313e-16
29	3	7	15	2.5466e-21	0.0000e+00	2.5466e-21
30	3	25	54	1.5036e-17	0.0000e+00	1.5036e-17
31	3	27	100	3.0573e+00	0.0000e+00	<b>3.0573e+00</b>
32	0	2	4	1.0000e+01	1.0000e+01	7.1054e-16
33	3	2	4	4.6341e+00	4.6341e+00	0.0000e+00
34	3	2	4	6.1351e+00	6.1351e+00	0.0000e+00

Table 19: Results of `lbfgs` on the Moré, Garbow, Hillstrom test collection. Errors greater than  $10^{-8}$  are highlighted in bold, indicating that a solution was not found within the specified tolerance.

#	Input Parameters	Iters	FuncEvals	$\hat{F}^*$	Error
6	<code>LineSearch_initialstep = 1e-5</code>	29	293	1.2436e+02	3.5262e-11
11	<code>LineSearch_initialstep = 1e-2</code>	220	1102	1.5634e-20	1.5634e-20
18	<code>LineSearch_initialstep = 0</code> <code>M = 1</code>	220	1102	9.3766e-17	9.3766e-17
31	<code>LineSearch_initialstep = 1e-7</code>	16	209	8.1617e-19	8.1617e-19

Table 20: Parameter changes that lead to solutions using `lbfgs` on the Moré, Garbow, Hillstrom test collection.

#	Exit	Iters	FuncEvals	$\hat{F}^*$	$F^*$	Error
1	0	105	694	4.9427e-30	0.0000e+00	4.9427e-30
2	3	9	107	4.8984e+01	4.8984e+01	2.9011e-16
3	3	457	3607	3.1474e-09	0.0000e+00	3.1474e-09
4	0	6	36	0.0000e+00	0.0000e+00	0.0000e+00
5	0	13	96	4.4373e-31	0.0000e+00	4.4373e-31
6	3	13	133	1.2436e+02	1.2436e+02	3.5262e-11
7	0	39	245	6.9237e-46	0.0000e+00	6.9237e-46
8	0	9	80	8.2149e-03	8.2149e-03	8.6736e-18
9	0	3	28	1.1279e-08	1.1279e-08	2.7696e-14
10	2	6531	50003	1.1104e+05	8.7946e+01	<b>1.2616e+03</b>
11	0	1	5	3.8500e-02	0.0000e+00	<b>3.8500e-02</b>
12	0	9	74	1.8615e-28	0.0000e+00	1.8615e-28
13	0	25	241	2.3427e-29	0.0000e+00	2.3427e-29
14	0	36	263	1.0481e-26	0.0000e+00	1.0481e-26
15	0	449	4967	3.0751e-04	3.0751e-04	3.9615e-10
16	3	28	237	8.5822e+04	8.5822e+04	4.3537e-09
17	3	2904	30801	5.4649e-05	5.4649e-05	3.3347e-10
18	3	2622	28977	6.3011e-17	0.0000e+00	6.3011e-17
19	3	526	5671	4.0138e-02	4.0138e-02	2.9355e-10
20	2	4654	50006	6.0164e-06	1.4017e-06	<b>4.6147e-06</b>
21	0	96	638	3.9542e-28	0.0000e+00	3.9542e-28
22	0	25	241	2.3427e-29	0.0000e+00	2.3427e-29
23	0	17	207	2.2500e-05	2.2500e-05	2.4991e-11
24	0	63	827	9.3763e-06	9.3763e-06	7.3554e-15
25	0	3	19	2.3419e-31	0.0000e+00	2.3419e-31
26	3	18	249	2.7951e-05	2.7951e-05	2.1878e-13
27	0	6	53	2.7820e-29	0.0000e+00	2.7820e-29
28	3	82	882	1.9402e-17	0.0000e+00	1.9402e-17
29	0	4	33	9.6537e-32	0.0000e+00	9.6537e-32
30	3	16	104	3.4389e-21	0.0000e+00	3.4389e-21
31	3	13	145	3.0573e+00	0.0000e+00	<b>3.0573e+00</b>
32	3	4	63	1.0000e+01	1.0000e+01	5.3291e-16
33	3	4	48	4.6341e+00	4.6341e+00	1.9166e-16
34	3	3	19	6.1351e+00	6.1351e+00	0.0000e+00

Table 21: Results of `tn` on the Moré, Garbow, Hillstrom test collection. Errors greater than  $10^{-8}$  are highlighted in bold, indicating that a solution was not found within the specified tolerance.

#	Input Parameters	Iters	FuncEvals	$\hat{F}^*$	Error
11	LineSearch.initialstep = 1e-3 HessVecFDStep = 1e-12	1018	17744	4.0245e-09	4.0245e-09
20	CGIters = 50	34	1336	1.3998e-06	1.9715e-09
31	LineSearch.initialstep = 1e-7	7	133	4.4048e-25	4.4048e-25

Table 22: Parameter changes that lead to solutions using `tn` on the Moré, Garbow, Hillstrom test collection.

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## 5 Conclusions

We have presented Poblano v1.0, a Matlab Toolbox for unconstrained optimization requiring only first order derivatives. Details of the methods available in Poblano as well as how to use the toolbox to solve unconstrained optimization problems were provided. Demonstration of the Poblano solvers on the Moré, Garbow and Hillstom collection of test problems indicates good performance in general of the Poblano optimizer methods across a wide range of problems.

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