NEW YORK UNIVERSITY

Master's Thesis

An L-BFGS-B Optimizer for Non-Smooth Functions

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in the

Courant Institute of Mathematical Sciences

Department of Mathematics

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Declaration of Authorship

I, Wilmer Henao, declare that this thesis titled, 'An L-BFGS-B Optimizer for Non-Smooth Functions' and the work presented in it are my own. I confirm that:

- This work was done wholly or mainly while in candidature for a research degree at this University.
- Where any part of this thesis has previously been submitted for a degree or any other qualification at this University or any other institution, this has been clearly stated.
- Where I have consulted the published work of others, this is always clearly attributed.
- Where I have quoted from the work of others, the source is always given. With the exception of such quotations, this thesis is entirely my own work.
- I have acknowledged all main sources of help.
- Where the thesis is based on work done by myself jointly with others, I have made clear exactly what was done by others and what I have contributed myself.

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Abstract

Faculty Name

Department of Mathematics

Master of Science in Scientific Computing

An L-BFGS-B Optimizer for Non-Smooth Functions

by Wilmer Henao

The Thesis Abstract is written here \dots

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I also would like to thank Allan Kaku and Anders Skaaja for earlier contributions to other versions of the code and Ariana Promessi for contributing to the language in the final version.

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$\mathbf{L}\mathbf{A}\mathbf{H}$ List Abbreviations Here Dedicated to my mother, my brother and Dr. Ian Malcolm

Chapter 1

Introduction

The problem addressed is to find a local minimizer of the nonsmooth minimization problem

$$\min_{x \in \mathbb{R}^n} \quad f(x)
\text{s.t.} \quad l_i \le x_i \le u_i,
\qquad i = 1, \dots, n.$$
(1.1)

where $f: \mathbb{R}^n \to \mathbb{R}$, is continuous but not differentiable everywhere and n is a very large but finite number.

The L-BFGS-B algorithm [1] is a standard method for solving large instances of 1.1 when f is a smooth function. The original name of BFGS stands for Broyden, Fletcher, Goldfarb and Shanno, the authors of the original "BFGS" quasi-Newton algorithm for unconstrained optimization discovered and published independently by them in 1970 [2] [3] [4] [5]. This method requires storing and updating a matrix which approximates the inverse of the Hessian matrix $\nabla^2 f(x)$ and hence requires $\mathcal{O}(n^2)$ operations per iteration. The L-BFGS variant [6] where the L stands for "Large", is based on BFGS but requires only $\mathcal{O}(mn)$ operations per iteration. And not only it requires less FLOPS, but it also requires much less memory. Instead of storing the $n \times n$ hessian approximations, L-BFGS stores only m vectors of dimension n, where m is much smaller than n. Finally, the last letter B in L-BFGS-B stands for bounds, meaning the lower and upper bounds l_i and u_i in 1.1. The L-BFGS-B algorithm is implemented in a well known Fortransoftware package by the same name [7]

In this thesis, there is a brief description of the L-BFGS-B algorithm at a high level and then an explanation of how the modified algorithm is more suitable for functions f which may not be differentiable at their local or global optimizers. We call the new algorithm L-BFGS-B-NS where NS stands for Non-Smooth. These changes were implemented in a modified version of the Fortran code [8] which can be downloaded from a web repository. We report on some numerical experiments that strongly suggest that the new code should be useful for the nonsmooth bound-constrained optimization problem (1.1).

We are grateful to Jorge Nocedal and his coauthors for allowing us to modify the L-BFGS-B code and post the modified version.

Chapter 2

L-BFGS-B

This section is a description of the original L - BFGS - B code at a very high level [7]. The original software is intended to work well with smooth functions. This thesis discusses how to modify the algorithm for Non-Smooth functions. The original FORTRAN code contains as well some documentation [9] about how the software was built

2.1 BFGS

BFGS is a standard tool for optimization of smooth functions.[10] It is a line search method and its goal is to find a search direction starting from its current position x. The search direction is of type $d = -B_k \nabla f^{-1}$ where B_k is the k^{th} approximation to the inverse Hessian. ² This k^{th} step approximation is calculated via the BFGS formula

$$B_{k+1} = \left(I - \frac{s_k y_k^T}{y_k^T s_k}\right) B_k \left(I - \frac{y_k s_k^T}{y_k^T s_k}\right) + \frac{s_k s_k^T}{y_k^T s_k}$$
(2.1)

where $y_k = \nabla f(x_{k+1}) - \nabla f(x_k)$ and $s_k = x_{k+1} - x_k$ and where the first approximation of B_0 is assumed to be the identity matrix I in this thesis ³. BFGS exhibits superlinear convergence but it also requires $\mathcal{O}(n^2)$ operations per iteration. [10]

In the case of Non-Smooth functions. BFGS typically succeeds in finding a local minimizer. This however requires some modifications of the line search conditions. This line

 $^{^{1}}$ Notice that when B is the identity, this is the same direction as steepest descent. Another common line search method of optimization

²When it is exactly the inverse Hessian the method is known as Newton's method. Newton's method has quadratic convergence but requires the explicit calculation of the Hessian at every single step.

 $^{^3{\}rm a}$ different starting matrix is suggested in Nocedal's book [10]

search conditions are known as the Armijo and weak Wolfe line search conditions and they will be explained later in this thesis.

2.2 L-BFGS

L-BFGS stands for Limited-memory BFGS. This algorithm approximates BFGS using only a limited amout of computer memory to approximate the inverse of the Hessian B. Instead of storing a dense $n \times n$ matrix, L-BFGS keeps a record of the last m iterations where m is a small number that is chosen according to the problem at hand.⁴ It is for this reason that during the first m iterations, BFGS and L-BFGS produce exactly the same search directions.

Because of this construction, the L-BFGS algorithm is less computationally intensive and it only requires $\mathcal{O}(mn)$ operations per iteration. So it is much better suited for problems where the number of dimensions n is large. For this reason it is the algorithm of choice in this thesis.

2.3 L-BFGS-B

Finally L-BFGS-B comes naturally as an extension of L-BFGS. The B stands for the inclusion of Boundaries. L-BFGS-B requires two extra steps on top of L-BFGS. First, there is a step called gradient projection that reduces the dimensionality of the problem. Depending on the problem, the gradient projection could potentially save a lot of iterations by eliminating those variables that are at bound at the optimum thus reducing the initial dimensionality of the problem and the number of iterations and running time. After this gradient projection comes the second step or subspace minimization. During the subspace minimization phase, an approximate quadratic model of 1.1 is solved iteratively in a similar way that the original L-BFGS algorithm is solved. The only difference is that during the search step phase the step length is restricted as much as necessary in order to remain within the box defined by 1.1.

2.3.1 Gradient Projection

The original algorithm was created for the case when n is large and f is smooth. Its first step is a gradient projection similar to the one outlined in [11, 12] which is used to

 $^{^4}$ In this thesis m < 20, and in practice numbers between 5 and 10 are regularly used. There is no way of knowing a priori what choice of m will provide the best results

determine an active set corresponding to those variables that are on either their lower or upper bounds. The active set defined at point x^* is:

$$\mathcal{A}(x^*) = \{ i \in \{1 \dots n\} | x_i^* = l_i \lor x_i^* = u_i \}$$
(2.2)

Working with this active set is more efficient in large scale problems. A pure line search algorithm would have to choose a step length short enough to remain within the box defined by u_i and l_i . So if at the optimum, a large number \mathcal{B} of variables are either on the lower or the upper bound. At least a number \mathcal{B} of iterations might be needed. Gradient projection tries to reduce this number of iterations. In the best case, only 1 iteration is needed instead of \mathcal{B} .

Gradient projection works on the approximation model:

$$m_k(x) = f(x_k) + \nabla f(x_k)^T (x - x_k) + \frac{(x - x_k)^T H_k(x - x_k)}{2}$$
 (2.3)

where H_k is a L - BFGS - B approximation to the Hessian $\nabla^2 f$ stored in the implicit way defined by L - BFGS.

In this first stage of the algorithm a piecewise linear segment starts on the current point x_k in the direction $-\nabla f(x_k)$. Whenever this direction encounters one of the constraints, the line segment turns corners in order to remain feasible. The path is nothing but the feasible piecewise projection of the negative gradient direction on the constraint box determined by the values \overrightarrow{l} and \overrightarrow{u} . At the end of this stage, the value of x that minimizes $m_k(x)$ restricted to this piecewise gradient projection path is known as the "Cauchy point" x^c .

2.3.2 Subspace Minimization

The problem with gradient projection is that its search direction does not take advantage of information provided implicitly by the Hessian H_k , and therefore the speed of convergence is at best linear. It is for this reason that a stage two is necessary. Stage 2 or subspace minimization uses an L - BFGS implicit approximation of the Inverse Hessian matrix restricted to the free variables that are not in the active set $\mathcal{A}(x^c)$.

The idea at a higher level is to solve the constrained problem 2.3, but only on those dimensions that are free. The starting point for this new stage will be the previously found Cauchy point x^c , the L-BFGS algorithm provides a new search direction \hat{d}^u that takes implicit advantage of second order approximations of the Hessian matrix.

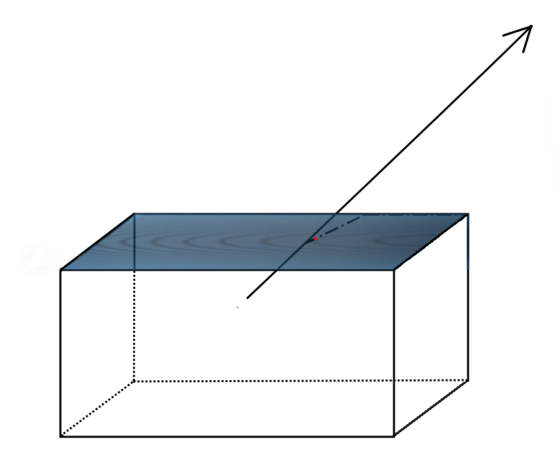


FIGURE 2.1: The arrow represents the direction of the gradient. The dotted path represents the projected gradient path from the gradient and onto the box. The region represents the level sets of the model. The optimal point (in red) is the Cauchy point x^c

The algorithm will move in the direction $\hat{d}^* = \alpha^* \hat{d}^u$ where α^* (the step length) is chosen so that the new point \bar{x}_i satisfies Armijo and weak wolfe descent and curvature conditions, A third restriction on the step length is added so that the next iteration stays feasible. Once this step is finished, the next and final step will be the termination condition. If the termination condition fails a new gradient projection and subspace minimization will be needed.

Chapter 3

Modifications to the L-BFGS-B algorithm

We made three main changes to the original L-BFGS-B algorithm. They concern the line search Wolfe conditions, the line search methodology, and the termination condition.

3.1 The Wolfe conditions

Probably the most important change made to the original code was the change in the curvature condition. It is accepted that the Wolfe conditions work very well whenever the function f is smooth [13]. Originally there are two Wolfe conditions: one of them is the Armijo condition, also known as the sufficient decrease requirement. The other one is the curvature condition, of which the most popular version is the "strong Wolfe" curvature condition:

$$|d_k^T \nabla f(x_k + \alpha_k d_k)| \le |d_k^T \nabla f(x_k)| \tag{3.1}$$

Where d_k represents the search direction. The strong Wolfe condition is natural for smooth optimization. Its goal is to find a step length long enough that the slope has been reduced "sufficiently", but the problem is that it does not work well for the Non-Smooth case. This is because near the minimal points, there may be abrupt changes in the gradient. For example the curvature condition can never be satisfied when f(x) = |x|, because the slope never becomes flat. The "weak wolfe" condition

$$d_k^T \nabla f(x_k + \alpha_k d_k) \ge d_k^T \nabla f(x_k)$$
(3.2)

is all that is needed to guarantee that the BFGS updated inverse Hessian approximation is positive definite [14]. This weak version is the one that will be used in this thesis. It was changed inside of the line search algorithm explained in the next section.

3.2 The line search methodology

The original FORTRAN code [7] contains a line search subroutine. This subroutine first establishes a maximum step. This maximum step is such that it guarantees that the step length stays within the bounding box delimited by l and u. This part of the line search was not changed in this thesis. The next part of the line search is finding a step length that satisfies a decrease and a curvature condition (aka. Armijo and Wolfe conditions). The next procedure was completely changed for the purpose of this thesis. and is only explained here with the purpose of showing why it needed to be modified. It was called on line 2662 A.1 of lbfgsnomessages.f90 and it has been left commented out in order to show how this call was before the modifications.

From then on, the line search required a lot of modifications. This new code is available online [8]

The original line search was a two stage process. In the first stage the interval was chosen so that it contains a minimizer of the modified function, which is nothing but a modification of the Armijo condition.

$$\Psi(\alpha_k) = f(x_k + \alpha_k d_k) - f(x_k) - c1\alpha_k d_k^T \nabla f(x_k)$$
(3.3)

If $\Psi(\alpha_k) < 0$ and $\alpha_k d_k^T \nabla f(x_k) > 0$. The interval is chosen so that it contains a minimizer of f. This first stage took place in the subroutine dcsrch; specifically on lines 3687 and 3709 A.2

The second stage was called on line 3713 and its mission was to find the step that satisfied Armijo and "Strong" Wolfe conditions on the original function (as opposed to the modified from stage 1) f. Both stage 1 and stage 2 called the subroutine destep. The subroutine still exists on file lbfgsbnomessages.f90 for illustration. Although it is never called in this thesis.

But there is a problem with the function *dcstep*. It turns out that function *dcstep* was designed to work only with smooth functions in mind. The code takes advantage of quadratic and cubic approximations to the function in order to calculate the next points

that satisfy armijo and wolfe conditions. Unfortunately, these second and third order approximations do not work in the Non-Smooth case, and the optimizer crashes under the line search as it was formulated. Function destep starts on line 3779 and a sample of the approximations is shown in between lines 3881 and 3902 A.3 of lbfgsbnomessages.f90.

The approach to this issue is to use a line search similar to the one in hanso [15]. The HANSO approach is to double the step length while the Armijo condition is violated (while the decresing conditions has not been satisfied). And once the interval has been bracketed, to do a bisection until both Armijo and Wolfe conditions are satisfied. The only difference with the HANSO approach in this thesis is that the line search in HANSO can double its step length up to 30 times ¹. Whereas in this thesis, the step length can double only as long as the step length is less than the maximum value that guarantees feasibility of the solution. This version of the bisection and expansion is found in between lines 4425 and 4456 A.4 of lbfqsbnomessages.f90

3.3 The termination condition

One important requirement of a practical algorithm is that it ends in a finite time. For the case of smooth functions, the usual way to check whether the algorithm has converged, is by using the projected gradient which is nothing but the projection of the negative gradient onto the bounding box defined by l and u. If this projected gradient has a small norm the algorithm has converged. In the case of nonsmooth functions however, this is not necessarily true and the function at the minimum, may have a wedge. In this wedge the projected gradient may not vanish. Furthermore, if there is a sequence of points that approaches the optimum x from the right, the projected gradients corresponding to this sequence of points might be completely different from the projected gradients associated to a sequence of points that approach the optimum x from the left.

Given this set of conditions, there is the need for a special set of rules to establish the finalization of each optimization.

3.3.1 Description of the Solution

Overton and Lewis formulate an algorithm that gives a practical solution to this problem on section 6.3 [14] The best practical methodology should be to calculate wheter zero 0 or a very small number is the norm of a vector that is part of the convex hull of

 $^{^{1}\}mathrm{for}$ all practical purposes this is considered as a good limit of iterations

gradients evaluated at points near the optimum candidate x. In order to make sure that the gradient zero $\vec{0}$ or a vector with a very small norm smaller than a very tiny tolerance $\tau_d > 0$ is part of the convex hull calculated near a neighbourhood of the optimum. The algorithm needs to keep a record of the latest gradient vectors in this small neighbourhood of the point where it is suspected that the optimum is located. The neighborhood is defined as those points with a distance to x smaller than a small tolerance $\tau_x > 0$ and no more than $J \in \mathbb{N}$ iterations back. This list of gradients is referred to as the set \mathcal{G} [14]

With this list \mathcal{G} of gradients at hand. The next step is to solve a quadratic program that locates the vector with the minimal norm that lives in the convex hull of these gradients. If the norm of this vector has a norm smaller than τ_d , the algorithm ends with a message of convergence success. If the minimum such norm is larger than the tolerance, the algorithm must continue to the following iteration and not terminate.

Every vector in the convex hull can be expressed as a nonnegative linear combination Gz of those vectors in \mathcal{G} . Where G is the matrix with columns made up of gradients in \mathcal{G} ; and z is such that $\sum_{i=1}^{n} z_i = 1$ and $z_i \geq 0$.

In other words the objective is to find the right combination of z that minimizes the norm $||Gz||_2$. This is equivalent to solving the optimization problem

min
$$q(z) = ||Gz||_2^2 = z^T G^T G z$$

s.t. $\sum_{i=1}^J z_i = 1$ (3.4)
 $z_i >= 0$.

The solution to this problem z^* has the associated vector Gz^* . And if $||Gz^*|| < \tau_d$ the algorithm converges.

3.3.2 The Solution of the Quadratic Program

The solution of 3.4 was implemented with a practical primal-dual methodology. This methodology is the same methodology implemented by Skajaa [16] in his thesis. His code apprecial was implemented in FORTRAN and is part of lbfgsbnomessages.f90. The solution is a typical Mehrotra's Predictor-Corrector algorithm applied to quadratic programming.

This algorithm initially tries to solve the Karush Kuhn Tucker KKT conditions. The KKT is a system of equations whose solution characterizes the solution of the original optimization problem. In 3.4. The Karush Kuhn Tucker equations are:

$$G^{T}Gz - e^{T}s = 0$$

$$\sum_{i=1}^{J} x - 1 = 0$$

$$z_{i}s_{i} = 0; \quad i = 1, 2, ..., J$$

$$(s, z) \geq 0$$
(3.5)

Where s is the variable of the dual problem.

In order to solve this system of equations a variant of Newton's method is used. This Newton's method as usual, provides a search direction which requires a corresponding step length. Since this is an interior point method, it approaches the solution following a path inside the convex interior of the feasible set. In the best of cases, it would be nice to approach the solution through a central path where the third equation in 3.5 is allowed to take more relaxed values $z_i s_i = \tau$, where $\tau \geq 0$. If the solution is approached through this central path. The convergence will require fewer iterations.

The problem is that the pure Newton's method solution is not close to this central path. The step length is usually very small because of the nature of the third and fourth equations. If the step length is too large, the condition of z > 0 or s > 0 will be violated. The method by itself approaches the solution very close to the boundaries of the feasible set. Not close to the central path where the step lengths could be larger and the convergence therefore would be faster. It is for this reason that a centering of the path is performed. This first step is known as the "predictor" step.

Besides the centering; Mehrotra proposed a "correction" stage which pushes the solution closer to the central path by taking into account the curvature of the approaching *central path*. This second stage is also used to calibrate some of the parameters used during the centering in the "predictor" stage. The second stage is similar to the first stage in that it requires the solution of a new system of equations.

Mehrotra's algorithm requires only one Cholesky factorization for the first stage. This factorization is reused during the second stage, by reusing the factorization, the performance is improved. This algorithm is widely used in optimizers.

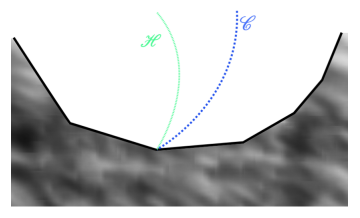


Figure 3.1: The central path ${\cal C}$ is approached from a current and noncentral point on the tangent path ${\cal H}$

Chapter 4

the solution test functions

There were some tests run on the high performance cluster machines at NYU. In order to run these tests it was necessary to create a series of PBS files using a PBS generator script B.1. This script generator created PBS files which in turn run bash scripts B.2. The main reason to run scripts this way is that it achieves parallelism and the system sends confirmation e-mails about the different stages of the processes.

A few test functions were evaluated. The most important function to test this non-smooth optimizer is a modified version of the rosenbrock function:

$$f(x) = (x_1 - 1)^2 + \sum_{i=2}^{n} |x_i - x_{i-1}^2|^p$$
(4.1)

Where the value of p changes the behaviour of the optimizer. This function can be proven to be lipschitz continuous whenever p > 1 if restricted to the domain defined by

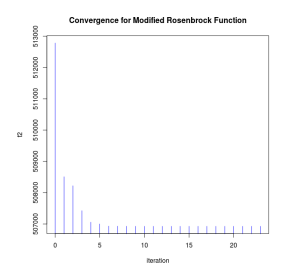
$$x_i = \begin{cases} [-100, 100] & \text{if } i \in \text{ even numbers} \\ [10, 100] & \text{if } i \in \text{ odd numbers} \end{cases}$$

$$(4.2)$$

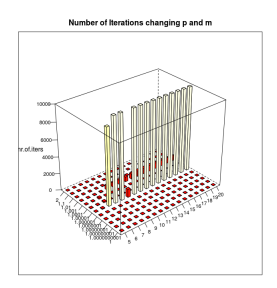
and in fact, whenever the function is restricted to a finite domain this function will be lipschitz continuous for p > 1. Whenever p > 1 the function $f(z) = |z|^p$ is zero 0 around zero because the derivative $p|z|^{p-1}$ is zero whenever z tends to zero from the right. (this is also the case from the left because it is an even function). However the second derivative will not be as nice.

For the case when $p \leq 1$ the second derivative tends to infinity. $\lim_{x \to 0^+} f' = \infty$. Which is already well known given the "heavyside" look of f(z) = |z|.

The convergence of the algorithm smoothly descends to the objective



The converge is adversely affected by the selection of p as one would expect. Values of p descending to 1 make the function less "smooth" and have the adverse effect of making the convergence much more difficult. In this exercise it is noticeable how slow the convergence becomes for a few specific values of p. In particular for 1.0001



$$f(x_k + \alpha_k p_k) \le f(x_k) + c_1 \alpha_k p_k^T \nabla f(x_k)$$
(4.3)

Appendix A

Samples of Code

A.1 Ignoring old code

All the code is available on [8] . This is also true for the code that is not being used anymore. Here the old function dcsrch is commented out. lineww is replacing it instead

```
2607
2608 ! call dcsrch(f,gd,stp,ftol,gtol,xtol,zero,stpmx,csave,isave,dsave)
2609 call lineww(f,gd,stp,ftol,gtol,xtol,zero,stpmx,csave,isave,dsave)
```

A.2 Old Stage 1 of the line search

Stage 1 of the line search[7] . This stage is not being run in lbfgsbNS. But is kept there for comparisson. For the new line search that replaces it A.4

```
if (stage .eq. 1 .and. f .le. fx .and. f .gt. ftest) then
3687
3688
       Here we define the modified function and derivative values.
3689 !
       This line search will only aim to satisfy the condition in (3.3) modified
3691
              fm = f - stp*gtest
3692
              fxm = fx - stx*gtest
3693
             fym = fy - sty*gtest
3694
              gm = g - gtest
3695
              gxm = gx - gtest
3696
              gym = gy - gtest
3697
       Call dcstep to update stx, sty, and to compute the new step.
3698 !
3699 !
3700
              call dcstep(stx,fxm,gxm,sty,fym,gym,stp,fm,gm,brackt,stmin,stmax)
3701
             Reset the function and derivative values for f
3702 !
3703
```

A.3 Cubic and Quadratic line search sample

This part of the code does not work in the case when the function is Non-Smooth . For this reason it was eliminated from execution on lbfgsbNS and replaced with A.4. stx in this case is a variable that represents the step with the minimum value so far.

```
First case: A higher function value. The minimum is bracketed.
3881
3882
           If the cubic step is closer to stx than the quadratic step, the
3883
           cubic step is taken, otherwise the average of the cubic and
3884
           quadratic steps is taken.
      ! theta, three, gamma are parameters
3886
          if (fp .gt. fx) then
3887
              theta = three*(fx - fp)/(stp - stx) + dx + dp
3888
              s = max(abs(theta),abs(dx),abs(dp))
              gamma = s*sqrt((theta/s)**2 - (dx/s)*(dp/s))
3889
3890
              if (stp .lt. stx) gamma = -gamma
3891
              p = (gamma - dx) + theta
              q = ((gamma - dx) + gamma) + dp
3892
3893
              r = p/q
              stpc = stx + r*(stp - stx) !quadratic step
3894
              stpq = stx + ((dx/((fx - fp)/(stp - stx) + dx))/two)* &
3895
3896
                                                                    (stp - stx) !The
        cubic step
              if (abs(stpc-stx) .lt. abs(stpq-stx)) then
3897
3898
                 stpf = stpc
3899
                 stpf = stpc + (stpq - stpc)/two
3900
3901
              endif
3902
              brackt = .true.
```

A.4 Line Search Enforcing Weak Wolfe Conditions

This is the implementation of the line search that enforces the weak Wolfe conditions. Bisections and expansions (as long as it doesn't mean leaving the bounding box) of the step length. This version tries to be as similar as possible to the interior of the while loop in qpspecial.m at hanso [15]

```
4425 if (fp .ge. ftest) then
4426 sty = stp
```

```
4427
              fy = fp
4428
             dy = dp
4429
              brackt = .true.
4430
4431 !
          if second condition is violated not gone far enough (Wolfe)
              if (-dp .ge. gtol*(-ginit)) then
4432
4433
                 stx = stp
4434
                 fx = fp
4435
                 dx = dp
4436
              else
4437
                 {\tt return}
4438
              endif
4439
           endif
4440
4441
          !Bisection and expansion
4442
          if (brackt) then
4443
              stp = (sty + stx)/two
4444
          else
4445
              if (two * stp .le. stpmax) then !Remain within boundaries
4446
                 ! Still in expansion mode
4447
                 stp = two * stp
4448
              else
4449
                 brackt = .true.
4450
                 sty = stp
4451
                 fy = fp
                 dy = dp
4452
4453
              endif
4454
           endif
4455
           return
4456
           end
```

Appendix B

Automation

This appendix includes some of the files that were used to run examples in parallel at the High Performance Clusters at NYU. All of these files can be found in the repository [8]

B.1 Script Generator

This is a sample of the script generator that creates pbs files to be sent to the High Performance Machines. The value "i" creates a different set of files, in this case one file for each value of p that we want to test. To see a result of running this script look at B.2.

```
1 #!/bin/bash
3 \text{ for i in } \{0..20\}
    cat > pbs.script.rosenbrock.$i <<EOF</pre>
6 #!/bin/bash
8 #PBS -1 nodes=1:ppn=8, walltime=48:00:00
9 #PBS -m abe
10 #PBS -M weh227@nyu.edu
11 #PBS -N rosenbrockHD$((i))
13 \text{ module load gcc/}4.7.3
14
15~\mathrm{cd}~/\mathrm{scratch/weh227/rosenbrock/}
16 p=$(bc -1 <<< "1+10 ^(-$((i)))")
18 do
19
     echo 10 \$n \$p
     ~/Documents/thesis/lbfgsfortran/./rosenbrockp 10 \$n \$p >> outputrosenbrock$
      ((i)).txt
```

```
21 done
22
23 exit 0;
24
25 EOF
26 done
```

After they are created I could fire all of the runs at once by using the command for i in 0..20; do qsub pbs.script.rosenbrock.\$i\$; done

B.2 PBS File Sample

This is a sample file general with B.1. It runs a special value p. These are the ones that get started with the qsub command.

Lines 4 to 5 provide the user with emails and critical information while running the process. Line 13 defines a run for different sizes of n. Line 16 is the actual run that will be sent to an output file. In this case "outputrosenbrock9.txt"

```
1
2 #!/bin/bash
4\ \mbox{\#PBS}\ \mbox{-l}\ \mbox{nodes=1:ppn=8,walltime=}48:00:00
5 #PBS -m abe
6 #PBS -M weh227@nyu.edu
7 #PBS -N rosenbrockHD9
9 module load gcc/4.7.3
11 cd /scratch/weh227/rosenbrock/
12 p=1.00000000100000000000
14 do
15
     echo 10 $n $p
     "/Documents/thesis/lbfgsfortran/./rosenbrockp 10 %n %p >> outputrosenbrock9.
17 done
18
19 exit 0;
```

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