# Conjugate gradient methods and other optimization methods

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#### Motivation

Our aim with this part of the project is to be able to

- find an optimal value for the variational parameters using only some few Monte Carlo cycles
- use these optimal values for the variational parameters to perform a large-scale Monte Carlo calculation

To achieve this will look at methods like *Steepest descent* and the *conjugate gradient method*. Both these methods allow us to find the minima of a multivariable function like our energy (function of several variational parameters). Alternatively, you can always use Newton's method. In particular, since we will only have one variational parameter, Newton's method can be easily used in finding the minimum of the local energy.

#### Simple example and demonstration

Let us illustrate what is needed in our calculations using a simple example, the harmonic oscillator in one dimension. For the harmonic oscillator in one-dimension we have a trial wave function and probability

$$\psi_T(x) = e^{-\alpha^2 x^2}$$
  $P_T(x)dx = \frac{e^{-2\alpha^2 x^2} dx}{\int dx e^{-2\alpha^2 x^2}}$ 

with  $\alpha$  being the variational parameter. We obtain then the following local energy

$$E_L[\alpha] = \alpha^2 + x^2 \left(\frac{1}{2} - 2\alpha^2\right),\,$$

which results in the expectation value for the local energy

$$\langle E_L[\alpha] \rangle = \frac{1}{2}\alpha^2 + \frac{1}{8\alpha^2}$$

#### Simple example and demonstration

The derivative of the energy with respect to  $\alpha$  gives

$$\frac{d\langle E_L[\alpha]\rangle}{d\alpha} = \alpha - \frac{1}{4\alpha^3}$$

and a second derivative which is always positive (meaning that we find a minimum)

$$\frac{d^2\langle E_L[\alpha]\rangle}{d\alpha^2} = 1 + \frac{3}{4\alpha^4}$$

The condition

$$\frac{d\langle E_L[\alpha]\rangle}{d\alpha}=0,$$

gives the optimal  $\alpha = 1/\sqrt{2}$ , as expected.

a) Derive the local energy for the harmonic oscillator in one dimension and find its expectation value.

dimension and find its expectation value. b) Show also that the optimal value of optimal  $\alpha = 1/\sqrt{2}$ .

# Variance in the simple model

We can also minimize the variance. In our simple model the variance is

$$\sigma^{2}[\alpha] = \frac{1}{2}\alpha^{4} - \frac{1}{4} + \frac{1}{32\alpha^{4}},$$

with first derivative

$$\frac{d\sigma^2[\alpha]}{d\alpha} = 2\alpha^3 - \frac{1}{8\alpha^5}$$

and a second derivative which is always positive

$$\frac{d^2\sigma^2[\alpha]}{d\alpha^2} = 6\alpha^2 + \frac{5}{8\alpha^6}$$

#### Computing the derivatives

In general we end up computing the expectation value of the energy in terms of some parameters  $\alpha_0, \alpha_1, \ldots, \alpha_n$  and we search for a minimum in this multi-variable parameter space. This leads to an energy minimization problem where we need the derivative of the energy as a function of the variational parameters.

In the above example this was easy and we were able to find the expression for the derivative by simple derivations. However, in our actual calculations the energy is represented by a multi-dimensional integral with several variational parameters. How can we can then obtain the derivatives of the energy with respect to the variational parameters without having to resort to expensive numerical derivations?

## Expressions for finding the derivatives of the local energy

To find the derivatives of the local energy expectation value as function of the variational parameters, we can use the chain rule and the hermiticity of the Hamiltonian.

Let us define

$$\bar{\mathcal{E}}_{\alpha} = \frac{d\langle \mathcal{E}_{L}[\alpha] \rangle}{d\alpha}.$$

as the derivative of the energy with respect to the variational parameter  $\alpha$  (we limit ourselves to one parameter only). In the above example this was easy and we obtain a simple expression for the derivative. We define also the derivative of the trial function (skipping the subindex T) as

$$\bar{\psi}_{\alpha} = \frac{d\psi[\alpha]}{d\alpha}.$$

#### Derivatives of the local energy

The elements of the gradient of the local energy are then (using the chain rule and the hermiticity of the Hamiltonian)

$$ar{\mathcal{E}}_{lpha} = 2 \left( \langle rac{ar{\psi}_{lpha}}{\psi[lpha]} \mathcal{E}_{L}[lpha] 
angle - \langle rac{ar{\psi}_{lpha}}{\psi[lpha]} 
angle \langle \mathcal{E}_{L}[lpha] 
angle 
ight).$$

From a computational point of view it means that you need to compute the expectation values of

$$\langle \frac{\bar{\psi}_{\alpha}}{\psi[\alpha]} E_{L}[\alpha] \rangle$$
,

and

$$\langle \frac{\bar{\psi}_{\alpha}}{\psi[\alpha]} \rangle \langle \mathcal{E}_{L}[\alpha] \rangle$$

a) Show that

$$ar{E}_lpha=2\left(\langlerac{ar{\psi}_lpha}{\psi[lpha]} E_L[lpha]
angle-\langlerac{ar{\psi}_lpha}{\psi[lpha]}
angle\langle E_L[lpha]
angle
ight).$$

b) Find the corresponding expression for the variance.

a) The expectation value for the local energy of the Helium atom with a simple Slater determinant is given by (project 1)

$$\langle E_L \rangle = \alpha^2 - 2\alpha \left( Z - \frac{5}{16} \right)$$

Show that its derivative is

$$\frac{d\langle E_L[\alpha]\rangle}{d\alpha} = 2\alpha - 2\left(Z - \frac{5}{16}\right).$$

**b)** Use result when you test your numerical derivative, user either the steepest descent method, Newton's method or the conjugate gradient method.

The success of the CG method for finding solutions of non-linear problems is based on the theory of conjugate gradients for linear systems of equations. It belongs to the class of iterative methods for solving problems from linear algebra of the type

$$\hat{A}\hat{x}=\hat{b}.$$

In the iterative process we end up with a problem like

$$\hat{r} = \hat{b} - \hat{A}\hat{x},$$

where  $\hat{r}$  is the so-called residual or error in the iterative process. When we have found the exact solution,  $\hat{r}=0$ .

The residual is zero when we reach the minimum of the quadratic equation

$$P(\hat{x}) = \frac{1}{2}\hat{x}^T\hat{A}\hat{x} - \hat{x}^T\hat{b},$$

with the constraint that the matrix  $\hat{A}$  is positive definite and symmetric. If we search for a minimum of the quantum mechanical variance, then the matrix  $\hat{A}$ , which is called the Hessian, is given by the second-derivative of the variance. This quantity is always positive definite.

## Conjugate gradient method, Newton's method first

We seek the minimum of the energy or the variance as function of various variational parameters. In our case we have thus a function f whose minimum we are seeking. In Newton's method we set  $\nabla f = 0$  and we can thus compute the next iteration point

$$\hat{x} - \hat{x}_i = \hat{A}^{-1} \nabla f(\hat{x}_i).$$

Subtracting this equation from that of  $\hat{x}_{i+1}$  we have

$$\hat{x}_{i+1} - \hat{x}_i = \hat{A}^{-1}(\nabla f(\hat{x}_{i+1}) - \nabla f(\hat{x}_i)).$$

#### Simple example and demonstration

The function f can be either the energy or the variance. If we choose the energy then we have

$$\hat{\alpha}_{i+1} - \hat{\alpha}_i = \hat{A}^{-1}(\nabla E(\hat{\alpha}_{i+1}) - \nabla E(\hat{\alpha}_i)).$$

In the simple model gradient and the Hessian  $\hat{A}$  are

$$\frac{d\langle E_L[\alpha]\rangle}{d\alpha} = \alpha - \frac{1}{4\alpha^3}$$

and a second derivative which is always positive (meaning that we find a minimum)

$$\hat{A} = \frac{d^2 \langle E_L[\alpha] \rangle}{d\alpha^2} = 1 + \frac{3}{4\alpha^4}$$

## Simple example and demonstration

We get then

$$\alpha_{i+1} = \frac{4}{3}\alpha_i - \frac{\alpha_i^4}{3\alpha_{i+1}^3},$$

which can be rewritten as

$$\alpha_{i+1}^4 - \frac{4}{3}\alpha_i\alpha_{i+1}^4 + \frac{1}{3}\alpha_i^4$$
.

#### Using the conjugate gradient method

- ► Start your program with calling a function which implements for example the CGM method.
- This function needs the function for the expectation value of the local energy and the derivative of the local energy.
- ➤ Your function **func** is now the Metropolis part with a call to the local energy function. For every call to the function **func** many practitionser use 1000-10000 Monte Carlo cycles for the trial wave function.
- ► This gives an expectation value for the energy which is returned by the function **func**.
- When one calls the local energy one also computes the first derivative of the expectaction value of the local energy

$$\frac{d\langle E_L[\alpha]\rangle}{d\alpha} = 2\langle \frac{\bar{\psi}_{T\alpha}}{\psi_T[\alpha]} \left( E_L[\alpha] - \langle E_L[\alpha]\rangle \right) \rangle.$$

In the CG method we define so-called conjugate directions and two vectors  $\hat{s}$  and  $\hat{t}$  are said to be conjugate if

$$\hat{s}^T \hat{A} \hat{t} = 0.$$

The philosophy of the CG method is to perform searches in various conjugate directions of our vectors  $\hat{x}_i$  obeying the above criterion, namely

$$\hat{x}_i^T \hat{A} \hat{x}_j = 0.$$

Two vectors are conjugate if they are orthogonal with respect to this inner product. Being conjugate is a symmetric relation: if  $\hat{s}$  is conjugate to  $\hat{t}$ , then  $\hat{t}$  is conjugate to  $\hat{s}$ .

An example is given by the eigenvectors of the matrix

$$\hat{\mathbf{v}}_i^T \hat{\mathbf{A}} \hat{\mathbf{v}}_j = \lambda \hat{\mathbf{v}}_i^T \hat{\mathbf{v}}_j,$$

which is zero unless i = j.

Assume now that we have a symmetric positive-definite matrix  $\hat{A}$  of size  $n \times n$ . At each iteration i+1 we obtain the conjugate direction of a vector

$$\hat{x}_{i+1} = \hat{x}_i + \alpha_i \hat{p}_i.$$

We assume that  $\hat{p}_i$  is a sequence of n mutually conjugate directions. Then the  $\hat{p}_i$  form a basis of  $R^n$  and we can expand the solution  $\hat{A}\hat{x}=\hat{b}$  in this basis, namely

$$\hat{\mathbf{x}} = \sum_{i=1}^{n} \alpha_i \hat{\mathbf{p}}_i.$$

The coefficients are given by

$$\mathbf{A}\mathbf{x} = \sum_{i=1}^{n} \alpha_i \mathbf{A} \mathbf{p}_i = \mathbf{b}.$$

Multiplying with  $\hat{p}_{k}^{T}$  from the left gives

$$\hat{\rho}_k^T \hat{A} \hat{x} = \sum_{i=1}^n \alpha_i \hat{\rho}_k^T \hat{A} \hat{\rho}_i = \hat{\rho}_k^T \hat{b},$$

and we can define the coefficients  $\alpha_k$  as

$$\alpha_k = \frac{\hat{\rho}_k^T \hat{b}}{\hat{\rho}_k^T \hat{A} \hat{\rho}_k}$$

#### Conjugate gradient method and iterations

If we choose the conjugate vectors  $\hat{p}_k$  carefully, then we may not need all of them to obtain a good approximation to the solution  $\hat{x}$ . So, we want to regard the conjugate gradient method as an iterative method. This also allows us to solve systems where n is so large that the direct method would take too much time. We denote the initial guess for  $\hat{x}$  as  $\hat{x}_0$ . We can assume without loss of generality that

$$\hat{x}_0 = 0$$
,

or consider the system

$$\hat{A}\hat{z}=\hat{b}-\hat{A}\hat{x}_0,$$

instead.

Important, one can show that the solution  $\hat{x}$  is also the unique minimizer of the quadratic form

$$f(\hat{x}) = \frac{1}{2}\hat{x}^T\hat{A}\hat{x} - \hat{x}^T\hat{x}, \quad \hat{x} \in \mathbb{R}^n.$$

This suggests taking the first basis vector  $\hat{p}_1$  to be the gradient of f at  $\hat{x} = \hat{x}_0$ , which equals

$$\hat{A}\hat{x}_0-\hat{b},$$

and  $\hat{x}_0 = 0$  it is equal  $-\hat{b}$ . The other vectors in the basis will be conjugate to the gradient, hence the name conjugate gradient method.

Let  $\hat{r}_k$  be the residual at the k-th step:

$$\hat{r}_k = \hat{b} - \hat{A}\hat{x}_k.$$

Note that  $\hat{r}_k$  is the negative gradient of f at  $\hat{x}=\hat{x}_k$ , so the gradient descent method would be to move in the direction  $\hat{r}_k$ . Here, we insist that the directions  $\hat{p}_k$  are conjugate to each other, so we take the direction closest to the gradient  $\hat{r}_k$  under the conjugacy constraint. This gives the following expression

$$\hat{
ho}_{k+1} = \hat{r}_k - rac{\hat{
ho}_k^T \hat{A} \hat{r}_k}{\hat{
ho}_k^T \hat{A} \hat{
ho}_k} \hat{
ho}_k.$$

We can also compute the residual iteratively as

$$\hat{r}_{k+1} = \hat{b} - \hat{A}\hat{x}_{k+1},$$

which equals

$$\hat{b} - \hat{A}(\hat{x}_k + \alpha_k \hat{p}_k),$$

or

$$(\hat{b} - \hat{A}\hat{x}_k) - \alpha_k \hat{A}\hat{p}_k,$$

which gives

$$\hat{r}_{k+1} = \hat{r}_k - \hat{A}\hat{p}_k,$$

## Conjugate gradient method, our case

If we consider finding the minimum of a function f using Newton's method, that is search for a zero of the gradient of a function. Near a point  $x_i$  we have to second order

$$f(\hat{x}) = f(\hat{x}_i) + (\hat{x} - \hat{x}_i)\nabla f(\hat{x}_i) \frac{1}{2}(\hat{x} - \hat{x}_i)\hat{A}(\hat{x} - \hat{x}_i)$$

giving

$$\nabla f(\hat{x}) = \nabla f(\hat{x}_i) + \hat{A}(\hat{x} - \hat{x}_i).$$

In Newton's method we set  $\nabla f = 0$  and we can thus compute the next iteration point (here the exact result)

$$\hat{x} - \hat{x}_i = \hat{A}^{-1} \nabla f(\hat{x}_i).$$

Subtracting this equation from that of  $\hat{x}_{i+1}$  we have

$$\hat{x}_{i+1} - \hat{x}_i = \hat{A}^{-1}(\nabla f(\hat{x}_{i+1}) - \nabla f(\hat{x}_i)).$$

```
#include <cmath>
#include <iostream>
#include <fstream>
#include <iomanip>
#include "vectormatrixclass.h"
using namespace std;
// Main function begins here
int main(int argc, char * argv[]){
  int dim = 2;
  Vector x(dim), xsd(dim), b(dim), x0(dim);
  Matrix A(dim,dim);
  // Set our initial quess
  x0(0) = x0(1) = 0;
  // Set the matrix
  A(0,0) = 3; A(1,0) = 2; A(0,1) = 2; A(1,1) = 6;
  b(0) = 2; b(1) = -8;
```

```
cout << "The Matrix A that we are using: " << endl;
A.Print();
cout << endl;
x = ConjugateGradient(A,b,x0);
xsd = SteepestDescent(A,b,x0);
cout << "The approximate solution using Conjugate Gradient is: " << x.Print();
cout << endl;
cout << "The approximate solution using Steepest Descent is: " << en xsd.Print();
cout << endl;
cout << endl;</pre>
```

```
Vector SteepestDescent(Matrix A, Vector b, Vector x0){
  int IterMax, i;
  int dim = x0.Dimension();
  const double tolerance = 1.0e-14;
  Vector x(dim),f(dim),z(dim);
  double c,alpha,d;
  IterMax = 30;
  x = x0;
  f = A*x-b;
  i = 0;
```

```
while (i <= IterMax){
  z = A*f;
  c = dot(f,f);
  alpha = c/dot(f,z);
  x = x - alpha*f;
  f = A*x-b;
  if(sqrt(dot(f,f)) < tolerance) break;
  i++;
}
return x;</pre>
```

```
Vector ConjugateGradient(Matrix A, Vector b, Vector x0){
int dim = x0.Dimension();
const double tolerance = 1.0e-14;
Vector x(dim),r(dim),v(dim),z(dim);
double c,t,d;

x = x0;
r = b - A*x;
v = r;
c = dot(r,r);
```

```
int i = 0; IterMax = dim;
while(i <= IterMax){
    z = A*v;
    t = c/dot(v,z);
    x = x + t*v;
    r = r - t*z;
    d = dot(r,r);
    if(sqrt(d) < tolerance)
        break;
    v = r + (d/c)*v;
    c = d; i++;
}
return x;
}</pre>
```

### Codes from numerical recipes

The codes are taken from chapter 10.7 of Numerical recipes. We use the functions *dfpmin* and *Insrch*. You can load down the package of programs from the webpage of the course, see under project 1. The package is called *NRcgm*107.*tar.gz* and contains the files *dfmin.c*, *Insrch.c*, *nrutil.c* and *nrutil.h*. These codes are written in C

```
void dfpmin(double p[], int n, double gtol, int *iter, double *fret,
double(*func)(double []), void (*dfunc)(double [], double []))
```

### What you have to provide

#### The input to **dfpmin**

```
void dfpmin(double p[], int n, double gtol, int *iter, double *fret,
double(*func)(double []), void (*dfunc)(double [], double []))
```

is

- ► The starting vector *p* of length *n*
- ▶ The function *func* on which minimization is done
- ▶ The function *dfunc* where the gradient i calculated
- ▶ The convergence requirement for zeroing the gradient *gtol*.

It returns in *p* the location of the minimum, the number of iterations and the minimum value of the function under study *fret*.

```
#include "nrutil.h"
using namespace std;
       Here we define various functions called by the main program
double E_function(double *x);
void dE_function(double *x, double *g);
void dfpmin(double p[], int n, double gtol, int *iter, double *fret,
    double(*func)(double []), void (*dfunc)(double [], double []));
// Main function begins here
int main()
     int n, iter;
     double gtol, fret;
     double alpha;
     n = 1;
     cout << "Read in guess for alpha" << endl;</pre>
     cin >> alpha;
```

```
// reserve space in memory for vectors containing the variational
parameters
double *p = new double [2];
gtol = 1.0e-5;
now call dfmin and compute the minimum
p[1] = alpha;
dfpmin(p, n, gtol, &iter, &fret,&E_function,&dE_function);
cout << "Value of energy minimum = " << fret << endl;
cout << "Number of iterations = " << iter << endl;
cout << "Value of alpha at minimu = " << p[1] << endl;
delete [] p;</pre>
```

```
// this function defines the Energy function
double E_function(double x[])
{
   double value = x[1]*x[1]*0.5+1.0/(8*x[1]*x[1]);
   return value;
} // end of function to evaluate
```

```
// this function defines the derivative of the energy
void dE_function(double x[], double g[])
{
   g[1] = x[1]-1.0/(4*x[1]*x[1]*x[1]);
} // end of function to evaluate
```

```
using namespace std;
// Here we define various functions called by the main program
double E_function(double *x);
void dE_function(double *x, double *g);
void dfpmin(double p[], int n, double gtol, int *iter, double *fret,
    double(*func)(double []), void (*dfunc)(double [], double []));
// Main function begins here
int main()
     int n, iter;
     double gtol, fret;
     double alpha;
     n = 1:
     cout << "Read in guess for alpha" << endl;</pre>
     cin >> alpha;
```

```
// reserve space in memory for vectors containing the variational
parameters
double *p = new double [2];
gtol = 1.0e-5;
now call dfmin and compute the minimum
p[1] = alpha;
dfpmin(p, n, gtol, &iter, &fret,&E_function,&dE_function);
cout << "Value of energy minimum = " << fret << endl;
cout << "Number of iterations = " << iter << endl;
cout << "Value of alpha at minimu = " << p[1] << endl;
delete [] p;</pre>
```

```
// this function defines the Energy function
double E_function(double x[])
{

// Change here by calling your Metropolis function which
// returns the local energy

double value = x[1]*x[1]*0.5+1.0/(8*x[1]*x[1]);
  return value;
} // end of function to evaluate
```

You need to change this function so that you call the local energy for your system. I used 1000 cycles per call to get a new value of  $\langle E_L[\alpha] \rangle$ .

```
// this function defines the derivative of the energy
void dE_function(double x[], double g[])
{
// Change here by calling your Metropolis function.
// I compute both the local energy and its derivative for every call
    g[1] = x[1]-1.0/(4*x[1]*x[1]*x[1]);
} // end of function to evaluate
```

You need to change this function so that you call the local energy for your system. I used 1000 cycles per call to get a new value of  $\langle E_L[\alpha] \rangle$ . When I compute the local energy I also compute its derivative. After roughly 10-20 iterations I got a converged result in terms of  $\alpha$ .