

# The VMC++ Library

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VMC++ (Variational Monte Carlo Plus Plus) is a C++ library which allows with few simple commands for the implementation of a Variational Monte Carlo calculation. It provides the tools for describing a trial wave function and an Hamiltonian, and it uses this information to find the optimal values of the variational parameters embedded in the trial wave function, and therefore find the best approximation of the ground state.

The code has been developed using the standard C++11, and requires the MCI++ (<https://github.com/francesco086/MCIntegratorPlusPlus>) and NoisyFunMin (<https://github.com/francesco086/NoisyFunMin>) libraries.

In the following we will present the classes made available by the library. At the beginning we will report the necessary `#include` call and the prototype of the class. The comment `TO DO` indicates that the method needs to be implemented (as in the case of a pure virtual class).

First, we will present the virtual classes that should be instantiated for describing the trial wave function and the Hamiltonian. Then, we will introduce the class that performs the whole Variational Monte Carlo calculation.

## 1 WaveFunction

```
1 // #import "WaveFunction.hpp"
2 class WaveFunction: public MCISamplingFunctionInterface
3 {
4     public:
5         // Constructor and destructor
6         WaveFunction(const int &nspacedim, const int &npart,
7                     const int &ncomp, const int &nvp);
8         virtual ~WaveFunction();
9
10        // Getters
11        int getNSpaceDim();
12        int getNPart();
13        int getNVP();
14
15        // Manage the variational parameters
16        virtual void setVP(const double *vp) = 0; // TO DO
17        virtual void getVP(double *vp) = 0; // TO DO
```

```

18
19 // // Heritage from MCISamplingFunctionInterface,
20 // // which must be implemented
21 // void samplingFunction(const double *in,
22 //     double *out) = 0; // TO DO
23 // double getAcceptance() = 0; // TO DO
24
25 // Derivatives of the trial wave function
26 virtual double d1(const int &i,
27     const double *in) = 0; // TO DO
28 virtual double d2(const int &i, const int &j,
29     const double *in) = 0; // TO DO
30 virtual double vd1(const int &i,
31     const double *in) = 0; // TO DO
32
33 }

```

The class **WaveFunction** is a pure virtual class introduced for representing the trial wave function. The first thing that should be noticed, is that it is a child class of the MCI++ class **MCISamplingFunctionInterface**. Please refer to the user manual of MCI++ to see the details of this class. The user must be able to correctly implement its methods **samplingFunction** and **getAcceptance**.

Let us now see in details the methods of the class **WaveFunction**:

- **WaveFunction**: The constructor. It is necessary to specify the number of dimensions of the space in which it is used (**nspacedim**, typically 3 dimensions), the number of particles that the wave function describes (**npart**), the number of components of the wave function (**ncomp**, which corresponds to **nproto** of the class **MCISamplingFunctionInterface**), and the number of variational parameters used (**nvp**);
- **getNSpaceDim**: Returns **nspacedim**;
- **getNPart**: Returns **npart**;
- **getNVP**: Returns **nvp**
- **setVP**: Set the variational parameters according to the provided **nvp**-dimensional array **vp**. Must be implemented by the user;
- **getVP**: Store in the **nvp**-dimensional array **vp** the values of the variational parameters; Must be implemented by the user;
- **d1**: Returns the value of  $\frac{\partial \psi(x)}{\partial x_i}$ , computed in  $x$  equal to the array given in **in**. Here  $\psi$  is the wave function, and **i** the coordinate's index. For example, in a system with **npart=2** and **nspacedim=3**,  $x_0, x_1, x_2$  are the  $x, y, z$  coordinates of the first particle, and  $x_3, x_4, x_5$  the coordinates of the second particle. Must be implemented by the user;

- d2: Returns the value of  $\frac{\partial^2 \psi(x)}{\partial x_i \partial x_j}$ . Must be implemented by the user;
- vd1: Returns the value of  $\frac{\partial \psi(x)}{\partial \alpha_i}$ , where  $\alpha_i$  is the  $i$ -th variational parameter of  $\psi$ . Must be implemented by the user.

As example, we report the implementation of a trial wave function for 1 particle in a one-dimensional space, that uses 2 variational parameters, and has only one component. The functional form is:

$$\psi(x) = e^{-b(x-a)^2} \quad (1)$$

where  $a$  and  $b$  are the two variational parameters.

```

1 class QuadrExponential1D1POrbital: public WaveFunction
2 {
3     protected:
4         double _a, _b;
5
6     public:
7         QuadrExponential1D1POrbital(const double a, const double b
8             ): WaveFunction(1,1,1,2) {_a=a; _b=b;}
9
10        void setVP(const double *in)
11        {
12            _a=in[0];
13            // if (_a<0.01) _a=0.01;
14            _b=in[1];
15            // if (_b<0.01) _b=0.01;
16            using namespace std;
17            //cout << "change a and b! " << _a << " " << _b <<
18                endl;
19        }
20        void getVP(double *out)
21        {
22            out[0]=_a; out[1]=_b;
23        }
24
25        void samplingFunction(const double *in, double *out)
26        {
27            *out = -2.*(_b*(in[0]-_a)*(in[0]-_a));
28        }
29
30        double getAcceptance()
31        {
32            return exp(getProtoNew(0)-getProtoOld(0));
33        }
34
35        double d1(const int &i, const double *in)
36        {
37            return (-2.*_b*(in[0]-_a));
38        }
39
40        double d2(const int &i, const int &j, const double *in)

```

```

39     {
40         return ( -2.*_b + (-2.*_b*(in[0]-_a))*(-2.*_b*(in[0]-_a
41             )) ) ;
42     }
43
44     double vd1(const int &i, const double *in)
45     {
46         if (i==0)
47         {
48             return (2.*_b*(in[0]-_a));
49         } else if (i==1)
50         {
51             return (-(in[0]-_a)*(in[0]-_a));
52         } else
53         {
54             using namespace std;
55             cout << "ERROR vd1 QuadrExponential! " << endl;
56             return 0.;
57         }
58     };

```

## 2 Hamiltonian

```

1 // import "Hamiltonian.hpp"
2 class Hamiltonian: public MCIObservableFunctionInterface
3 {
4     public:
5         // Constructor and destructor
6         Hamiltonian(const int &nspacedim, const int &npart,
7             WaveFunction * wf):
8             MCIObservableFunctionInterface(nspacedim*npart,4)
9             virtual ~Hamiltonian() {}
10
11         // Getters
12         int getNSpaceDim();
13         int getNPart();
14
15         // Kinetic energy
16         double localPBKineticEnergy(const double *in);
17         double localJFKineticEnergy(const double *in);
18
19         // Potential energy
20         virtual double localPotentialEnergy(const double *in) = 0;
21         // TO DO
22
23         // Heritage from MCIObservableFunctionInterface
24         // already implemented
25         void observableFunction(const double * in, double *out)
26 };

```

This pure virtual class is used to define the Hamiltonian of the system. The user must take care of implementing only the `localPotentialEnergy` method, while all the rest is already provided.

- **Hamiltonian:** The constructor. It requires the spacial number of dimensions (`nspacedim`), the number of particles (`npart`), and a pointer to an implementation of a `WaveFunction`. Notice that the constructor inform the constructor of `MCIObservableFunctionInterface` that the Monte Carlo integral will be performed in a `nspacedim × npart` space, and that there will be 4 observables, corresponding to the total, potential, kinetic, and Jackson-Feenberg kinetic energies, respectively. All the energies are computed as *energy unit per particle*;
- `getNSpaceDim`: Returns `nspacedim`;
- `getNPart`: Returns `getNPart`;
- `localPBKineticEnergy`: Returns the standard (Pandharipande-Bethe) local kinetic energy, in units of energy per particle. It requires the particle positions, provided through the array `in`;
- `localJFKineticEnergy`: Returns the Jackson-Feenberg local kinetic energy;
- `localPotentialEnergy`: Returns the local potential energy. Must be implemented by the user;
- `observableFunction`: Returns the 4 observables for the particle coordinates contained in the array `in`. The observables will be contained in the array `out`, and are the local total, potential, kinetic, and Jackson-Feenberg kinetic energies.

As example, we report the Hamiltonian for a one-dimensional, one-particle, harmonic oscillator:

$$H = -\frac{\partial^2 \psi}{\partial x^2} + \frac{1}{2} \omega^2 x^2 \quad (2)$$

```

1 class HarmonicOscialltor1D1P: public Hamiltonian
2 {
3     protected:
4         double _w;
5
6     public:
7         HarmonicOscialltor1D1P(const double w, WaveFunction * wf):
8             Hamiltonian(1, 1, wf) {_w=w;}
9
10        double localPotentialEnergy(const double *r)
11        {
12            return (0.5*_w*_w*(*r)*(*r));
13        }
14 };

```

### 3 VMC

```

1 // #include "VMC.hpp"
2 class VMC{
3
4 public:
5     VMC(WaveFunction * wf, Hamiltonian * H);
6     ~VMC();
7
8
9     // Monte Carlo Integral within VMC should be performed using
10    the MCI object provided by VMC
11    MCI * getMCI(){return _mci;}
12
13    // Computation of the variational energy
14    void computeVariationalEnergy(const long & Nmc, double * E,
15                                  double * dE);
16
17    // Wave Function Optimization Methods
18    void conjugateGradientOptimization(const long &E_Nmc, const
19                                       long &grad_E_Nmc);
20
21    void stochasticReconfigurationOptimization(const long &Nmc);
22
23    void simulatedAnnealingOptimization(const long &Nmc, const
24                                         double &iota, const double &kappa, const double &lambda,
25                                         gsl_siman_params_t &params);
26
27 };

```

The Variational Monte Carlo (VMC) method is a method that uses a minimisation method in order to find the variational parameters  $\alpha$  of a trial wave function  $\psi_\alpha$  such that the variational energy

$$E(\psi_\alpha) = \frac{\int dR \psi_\alpha(R) H \psi_\alpha(R)}{\int dR \psi_\alpha(R) \psi_\alpha(R)} \quad (3)$$

is minimised. In Eq. (3)  $H$  is the Hamiltonian, and we have assumed that both the Hamiltonian and the wave functions are real, as it is done by this library. The minimisation can be achieved by using several optimisation algorithms. At the end of the optimization process, the wave function will be set to have the optimal variational parameters

- **VMC:** The constructor. It requires a wave function, a Hamiltonian, and the number of sampling points to use for computing the energy (**ENmc**) and the energy gradient (**GNmc**) during the optimisation process. Notice that the constructor inform the constructor of **NoisyFunctionWithGradient** about the number of the number of variational parameters used by the wave function;

- `getMCI`: Returns a pointer to the MCI object used to compute the variational energy and the wave function derivatives;
- `computeVariationalEnergy`: Compute the total, kinetic, and potential energies for the given wave function. In fact `E` is expected to be an array of size 4, where `E[0]` is the total energy, `E[1]` is the potential energy, `E[2]` is the Pandharipande-Bethe kinetic energy (i.e. the "real" kinetic energy), and `E[3]` is the Jackson-Feenberg kinetic energy;
- `conjugateGradientOptimization`: Optimise the trial wave function using the conjugate gradient method of the [NoisyFunMin](#) library. It requires the number of sampling points for computing the energy during the linear search, and the number of samplings to use for computing the energy gradient;
- `stochasticReconfigurationOptimization`: Optimise the trial wave function using the Dynamic Descent method of the [NoisyFunMin](#) library using as direction the one obtained using the Stochastic Reconfiguration (SR) approach. It requires the number of sampling points for computing the SR direction;
- `simulatedAnnealingOptimization`: Optimise the trial wave function using the Simulated Annealing algorithm contained in the GSL library (read [here](#)). The simulated annealing will try to minimise the target function

$$\iota E + \kappa \sigma_E + \lambda \frac{\sqrt{\sum_{i=1}^{N_\alpha} \alpha_i^2}}{N_\alpha}$$

where  $E$  is the energy,  $\sigma_E$  is the energy's standard deviation, and the last term represents a normalization factor, often used in Machine Learning. The weight of each term is controlled by the three parameters  $\iota$  (`iota`),  $\kappa$  (`kappa`), and  $\lambda$  (`lambda`), which must be provided to the method. Furthermore the method requires `Nmc`, the number of sampling points for computing the target function integrals ( $E$  and  $\sigma_E$ ), and the GSL `siman` library parameters.

Examples can be found in the folder `examples`.