The Feed Forward Neural Network library

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FFNN (Feed Forward Neural Network) is a C++ library which allows the implementation of a feed forward neural network with few simple calls. The library includes a training method, which currently utilizes GSL nonlinear multifit routines (trust region based Levenberg-Marquardt) to achieve least square fitting against target data. A data split into training, validation and testing is supported and enables early stopping. The library features inherent data normalization, completely hidden from the user. In terms of derivatives of the NN value, the library currently supports its first and second derivatives with respect to the NN input, the first derivative with respect to the variational weights within the FFNN and first/second order cross derivatives, with respect to both input (first / second order) and variational parameters (only first order).

The code has been developed using the standard C++11.

In the following we will present some classes made available by the library. At the beginning we will report the necessary #include call and the prototype of the class (not necessarily equal to actual code).

1 FeedForwardNeuralNetwork

```
#include "FeedForwardNeuralNetwork.hpp"
  class FeedForwardNeuralNetwork
2
3 {
       // Constructor and destructor
      FeedForwardNeuralNetwork(const int &insize,
5
           const int &hidlaysize,
           const int &outsize);
      FeedForwardNeuralNetwork(const char * filename);
      ~FeedForwardNeuralNetwork();
9
10
      // Getters for the basic information
11
      int getNHiddenLayers();
12
      int getNLayers();
13
      int getLayerSize(const int &li);
14
       ActivationFunctionInterface *
15
            getLayerActivationFunction(const int &li);
16
```

```
// Getters for the variational parameters
18
      int getNBeta();
19
20
       double getBeta(const int &ib);
21
       void setBeta(const int &ib, const double &beta);
22
       // Set the structure of the FFNN
23
       void pushHiddenLayer(const int &size);
24
       void popHiddenLayer();
25
       void setLayerSize(const int &li , const int &size);
26
       void setGlobalActivationFunctions(
27
            ActivationFunctionInterface * actf);
28
      void setLayerActivationFunction(const int &li,
29
            ActivationFunctionInterface * actf);
30
31
       // Set the computations required
32
      void addFirstDerivativeSubstrate();
33
       void addSecondDerivativeSubstrate();
34
       void addVariationalFirstDerivativeSubstrate();
35
       void addCrossFirstDerivativeSubstrate();
36
       void addCrossSecondDerivativeSubstrate();
37
38
      // Connect the FFNN,
39
      void connectFFNN();
40
       void disconnectFFNN();
41
42
       // Set the input
43
       void setInput(const int &n, const double * in);
45
       // Feed Forward propagate the input signal
46
      void FFPropagate();
47
48
       // Get the ouput after the FFNN Propagation
49
       double getOutput(const int &i);
50
       double getFirstDerivative(const int &i,
51
                                 const int &ild);
52
       double getSecondDerivative(const int &i,
53
                                  const int &i2d);
      double getVariationalFirstDerivative(const int &i,
55
                                             const int &iv1d);
56
      double getCrossFirstDerivative(const int &i,
57
                                       const int &i1d
58
                                       const int &iv1d);
59
       double getCrossFirstDerivative(const int &i,
60
                                       const int &i2d
61
                                       const int &iv1d);
62
63
       // Store the FFNN structure on a file
       void storeOnFile(const char * filename);
66 };
```

Fig. 1 illustrates the structure of a Feed Forward Neural Network (FFNN). A FFNN has always at least one *hidden layer*, an *input layer*, and an *output layer*. Each layer is made of at least one *unit* (in the figure, the empty circles

containing a label) plus an offset unit (in the figure, the filled circles). The number of units in a layer (including the offset unit) is called layer size. Each unit contains a value, labeled with the symbol u_i^l , where l is the layer index and i the unit index. The units with index i=0 are not reported in the figure, but represent the offset units. The units are connected through a net of connections, parametrised by a value $\beta_n^{l,p}$, where l is the index of the layer where the connections are ending, p refers to the index of the origin-unit, and the index n to the index of the destination-unit.

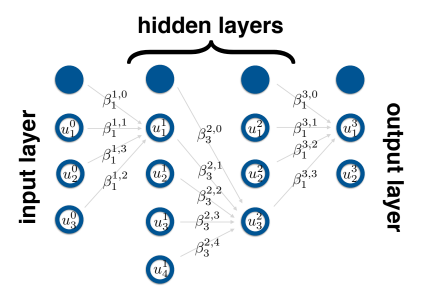


Figure 1: Graphical representation of a Feed Forward Neural Network (FFNN). For a sake of clearness, only some connections are displayed.

Despite the complex tasks that a FFNN can be used for, the math of a FFNN is extremely simple. A FFNN can be interpreted as a function $f_H: \mathbb{R}^N \to \mathbb{R}^M$, where N is the size of the input layer (i.e. its number of units, without counting the offset unit), and M the size of the output layer. The label H represents the number of hidden layers of the FFNN. Then, once that all the parameters β are set, the FFNN output can be computed with the recursive formula:

$$u_i^l = \mathbf{a}_f(u_k^{l-1}\beta_i^{l,k}) \tag{1}$$

where $a_f : \mathbb{R} \to \mathbb{R}$ is a so-called *activation function* and we have made use of the Einstein notation, for which repeated symbols on different levels imply a summation, e.g. $x_i^j y^i = \sum_i x_i^j y^i$. The activation function is required to account for non-linear effects. The FFNN library provides several well-known

activation functions by default, but the user has the possibility to define and use its own, even though its storage on a file it is not supported (explained later). These available activation functions are:

- 1. the logistic function: $\sigma_{lgs}(x) \equiv \frac{1}{1+e^{-x}}$ The current default for hidden and output units.
- 2. the tansig function: $\sigma_{\text{tans}}(x) \equiv \frac{2}{1+e^{-2x}} 1$ Faster approximation to the regular tanh. Similar to logistic function, but with output in [-1,1].
- 3. the gaussian: $\sigma_{\rm gss}(x) \equiv e^{-x^2}$ Unlike normal activation functions it is symmetric around origin and not monotonic. Still there are use cases for it.
- 4. the sine function: $\sigma_{\sin}(x) \equiv \sin(x)$ Periodic activation function for special use cases.
- 5. the *identity*: $\sigma_{id}(x) \equiv x$ This function is linear and therefore should not be used for (all) hidden units.
- 6. the $PReLU^1$ function: $\sigma_{ReLU}(x) \equiv \begin{cases} x & \text{if } x > 0 \\ \alpha x & \text{if } x <= 0 \end{cases}$

The fixed hyper-parameter α defaults to 0. Due to their efficiency and because their gradient doesn't vanish, (Parametric) Rectified Linear Units are widely used in deep neural networks. However, the function is not differentiable at x=0 and has flat derivatives elsewhere.

7. the $SELU^2$ function: $\sigma_{SELU}(x) \equiv \lambda \begin{cases} x & \text{if } x > 0 \\ \alpha e^x - \alpha & \text{if } x <= 0 \end{cases}$

The Scaled Exponential Units (with two fixed hyper-parameters α and λ) solve some problems of ReLUs and can provide a self-normalizing network. But still they are not differentiable at x=0.

The use of the FFNN library should be organised in four major steps:

- 1. generate the geometry of the FFNN;
- 2. connect the FFNN. This step will generate all the variational parameters β ;
- 3. add substrates, required for computing the derivatives, if necessary;

¹arXiv:1502.01852

²arXiv:1706.02515

4. set the input and propagate it through the FFNN in order to get the output.

Now that the notation of the FFNN has been introduced, and we have briefly seen its structure and math, let us see the methods of this class:

- FeedForwardNeuralNetwork: There are two possible constructors. The first one takes 3 int, which represents the size of the input, hidden (only one), and output layers. The user can change its shape later on. The values of β will be set to some random values. The second constructor takes the path to a file where a FFNN has been stored;
- getNHiddenLayers: Return the number of hidden layers H;
- getNLayers: Return the total number of layers, i.e. H + 2;
- getLayerSize: Return the size of the layer 1i. Remember that 1i = 0 is reserved for the input layer and 1i = H + 1 for the output layer;
- getLayerActivationFunction: Return a pointer to the activation function used to obtain the layer li. In the next section we will present the class ActivationFunctionInterface;
- getNBeta: Return the total number of β ;
- getBeta: Return the value of the $\beta_n^{l,p}$ corresponding to the ib:

$$\mathtt{ib} = \mathtt{p} + (\mathtt{n} - 1)\,\mathtt{getLayerSize}(\mathtt{l} - 1) + \sum_{i=1}^{\mathsf{l} - 1}\,\mathtt{getLayerSize}(i)\,\mathtt{getLayerSize}(i + 1)$$

- setBeta: Set the value of a beta;
- pushHiddenLayer: Add an hidden layer of size size between the last hidden layer and the output layer. Notice that all the beta related to this layer will be set randomly;
- popHiddenLayer: Remove the last hidden layer (the one connected to the output layer);
- setLayerSize: Set the size of a Layer;
- setGlobalActivationFunctions: Set a global activation function for all the connections in the FFNN;
- setLayerActivationFunction: Set the activation function used to generate the layer li;

- addFirstDerivativeSubstrate: Add a substrate that allows for the computation of the first derivatives of the FFNN in respect to the input, i.e. $\frac{\partial u_i^{H+1}}{\partial u_i^0}$;
- addSecondDerivativeSubstrate: Add a substrate that allows for the computation of the first derivatives of the FFNN in respect to the input, i.e. $\frac{\partial^2 u_i^{H+1}}{\partial u_i^{0.2}}$;
- addVariationalFirstDerivativeSubstrate: Add a substrate that allows for the computation of the first derivative of the FFNN in respect to the variational parameters β , i.e. $\frac{\partial u_i^{H+1}}{\partial \beta_i}$;
- addCrossFirstDerivativeSubstrate: Add a substrate that allows for the computation of the cross derivatives of the FFNN in respect to the input and the variational parameters β , i.e. $\frac{\partial^2 u_i^{H+1}}{\partial \beta_i \partial u_i^0}$;
- addCrossSecondDerivativeSubstrate: Add a substrate that allows for the computation of the cross second derivatives of the FFNN in respect to the input and the variational parameters β , i.e. $\frac{\partial^3 u_i^{H+1}}{\partial \beta_j \, \partial u_i^{0}}$;
- connectFFNN: Connect all the units in the FFNN. After this call it is possible to use the FFNN for computing quantities;
- disconnectFFNN: Disconnect all the units;
- setInput: Set the values of the units in the input layer. n is the size of the array in, and if it does not suite the size of the input layer an error message will be thrown;
- FFPropagate: Compute the output values, including the derivatives if the substrates have been set accordingly;
- getOutput: Get the value of the unit i in the output layer;
- getFirstDerivative: Get the first derivative of the unit value u_{i}^{H+1} in respect to u_{i1d}^{0} ;
- getSecondDerivative: Get the second derivative of the unit value u_{i}^{H+1} in respect to u_{i2d}^{0} ;
- getVariationalFirstDerivative: Get the first derivative of the unit value $u_{\mathbf{i}}^{H+1}$ in respect to β with $\mathbf{ib} = \mathbf{iv1d}$. See getBeta for a definition of \mathbf{ib} ;
- getCrossFirstDerivative: Get the cross derivative of the unit value $u_{\mathbf{i}}^{H+1}$ in respect to $u_{\mathbf{i}1d}^{0}$ and β with $\mathbf{ib} = \mathbf{iv1d}$;

- getCrossSecondDerivative: Get the second cross derivative of the unit value $u_{\mathbf{i}}^{H+1}$ in respect to $u_{\mathbf{i}+1}^{0}$ and β with $\mathbf{ib} = \mathbf{iv1d}$;
- storeOnFile: Store the FFNNon a file, making possible to retrieve it later on. If the activation functions have been customised, this procedure will not succeed.

2 ActivationFunctionInterface

```
// #include "ActivationFunctionInterface.hpp"
1
2 class ActivationFunctionInterface
3 {
4 public:
        // allocate a new copy of this to returned ptr
6
         virtual ActivationFunctionInterface * getCopy() = 0;
7
8
        // return an unique identifier code
9
         virtual std::string getIdCode() = 0;
10
11
        // return the ideal input mean value (mu) and standard
12
            deviation (sigma)
         virtual double getIdealInputMu() = 0;
13
        virtual double getIdealInputSigma() = 0;
14
15
        // return the output mean value (mu) and standard
16
            deviation (sigma)
        // (pretending a flat distribution)
17
         virtual double getOutputMu() = 0;
18
        virtual double getOutputSigma() = 0;
19
20
        // compute the activation function value
21
22
         virtual double f(const double \&) = 0;
23
        // first derivative of the activation function
24
         virtual double fld(const double \&) = 0;
25
26
         // second derivative of the activation function
27
         virtual double f2d (const double &) = 0;
28
29
        // third derivative of the activation function
30
        virtual double f3d (const double &) = 0;
31
32
        // calculate all derivatives together to save redundant
33
            calculations
34
         virtual void fad (const double &, const double &, const
            double &, const double &, const double &, const bool &,
             const bool &, const bool &);
35 };
```

The ActivationFunctionInterface is a pure virtual class, which can be used to generate customised activation functions to feed a FFNN. It is necessary to provide implementations of all virtual methods except fad and highly recommended to provide fad (otherwise fad will use the other methods via virtual function calls -> slow). As example we report the class for the logistic function:

```
class LogisticActivationFunction: public
      ActivationFunctionInterface
2
3
     public:
4
         ActivationFunctionInterface * getCopy(){return new
5
             LogisticActivationFunction();}
6
         std::string getIdCode(){return "LGS";}
7
8
         // input should be in the rage [-5:5] -> mu=0
                                                                sigma=
9
             sqrt(3)
         double getIdealInputMu(){return 0.;}
10
         double getIdealInputSigma(){return 2.886751345948129;}
11
12
         // output is in the range [0:1] \rightarrow mu=0.5
                                                           sigma = 1/(2*
13
             sqrt(3)
         double getOutputMu(){return 0.5;}
14
         double getOutputSigma() {return 0.288675134594813;}
15
16
         double f(const double &in)
17
18
            return (1./(1.+\exp(-in)));
19
20
21
         double fld (const double &in)
22
23
            double f=this->f(in);
24
            return (f*(1.-f));
25
26
27
         double f2d (const double &in)
28
29
            double f = this -> f(in);
30
            return (f*(1.-f)*(1.-2.*f));
31
32
33
         double f3d (const double &in)
34
35
            const double f = this \rightarrow f(in);
36
            return f * (1. - f) * (1. - 6.*f + 6*f*f);
37
38
39
         void fad (const double &in, double &v, double &v1d, double
40
             &v2d, double &v3d, const bool flag d1 = false, const
             bool flag d2 = false, const bool flag d3 = false)
```

```
41
             v = 1./(1.+\exp(-in));
42
43
44
             if (flag d1) {
45
                 \mathbf{v1d} = \mathbf{v} * (1. - \mathbf{v});
                 v2d = flag_d2 ? v1d * (1. - 2.*v) : 0.;
46
                 v3d = flag d3 ? v1d * (1. - 6.*v + 6.*v*v) : 0.;
47
48
             else {
49
                 v1d = 0.;
50
                 v2d = flag d2 ? v * (1. - v) * (1. - 2.*v) : 0.;
51
                 v3d = flag d3 ? v * (1. - v) * (1. - 6.*v + 6.*v*v)
52
             }
53
54
  };
```

3 Training

The library contains trainer classes with virtual base trainer class NNTrainer, which already provides the bestFit method to find the best NN fit among several fits to target data. Currently the only trainer implementation is the one using GSL nlinear multifit, namely NNTrainerGSL. This class provides the actual routine for an individual fit, findFit.

How a complete code for NN training looks, you can see in the example 10. Leaving out the FFNN and data setup however, it comes down to the following:

```
#include "NNTrainerGSL.hpp"
1
2
    // create data and config structs
3
       (for reference see respective header files)
4
    NNTrainingData tdata = {ndata, ntraining, nvalidation, xndim,
5
        yndim, xdata, ydata, d1data, d2data, weights};
    NNTrainingConfig tconfig = {flag r, flag d1, flag d2, lambda r
6
        , lambda d1, lambda d2, maxn steps, maxn novali};
    // create trainer and find best fit
8
    NNTrainerGSL * trainer = new NNTrainerGSL(tdata, tconfig);
9
10
    // (optional) setup ffnn's internal normalization before
11
        fitting
    trainer -> setNormalization(ffnn);
12
13
    // find a fit out of nfits with minimal testing residual
14
    trainer -> bestFit (ffnn, nfits);
15
16
    // (optional) print output files along all axis
```

Afterwards, the ffnn will be in best fit configuration.

Make sure that your ffnn has the necessary derivatives set before training (to be safe, always set d1,d2,vd1 and d1vd1 (+ d2vd1) depending on flag_d1/flag_d2).

Also, it is important to understand the meaning of training, validation and testing here:

- training: The data that will be used by the least square solver itself. Especially the residual gradients will only be calculated from training data.
- validation: Every time the least square solver finds a new training residual minimum, the residual of the validation data set will be calculated. If it didn't decrease (for too long, controlled by maxn_novali), the fitting stops early.
- testing: When the individual fitting routine terminates (because maxn_steps reached or early stop), for the resulting fit the residual with respect to the testing data is calculated. Out of all fits, the fit with best testing residual is returned.

Note that validation can be disabled by setting nvalidation in the NNTrainingData struct to 0 (this disables early stopping). testing however is disabled, if ndata is equal to ntraining plus nvalidation (in this case the training data set is used as replacement).

It is also possible to fine tune the GSL nlinear multifit algorithm by passing a gsl_multifit_nlinear_parameters (see GSL docs) struct at trainer creation:

Finally, you can pass more arguments to bestFit to tweak the behavior:

This will terminate the whole bestFit routine, if a testing residual below resi_target is found. Depending on int verbose (0, 1 or 2), it will print different levels of information. And if flag_smart_beta is true, it will use smart beta generation to randomize betas before fitting. More about that in the next section.

4 Smart Beta

To understand this section, first read the pdf smart_initial_beta.pdf in this folder. It should be noted that this feature is a bit experimental, so double check that it actually works well for you.

It is possible to initialise the beta of a FFNN by doing the following:

For each unit we computed the desired μ_{β} and σ_{β} using the formula reported in smart_initial_beta.pdf, and sampled the beta accordingly.

To enforce the orthogonality we proceeded in the following way. We sampled the first unit beta, a vector that we will now label as v. Then we sampled the second unit one, u, and applied the following modification:

$$u \rightarrow u - \frac{\langle u, v \rangle}{\langle v, v \rangle} v$$
 (2)

where $\langle u,v\rangle$ denotes the scalar product between u and v. This modification enforces the orthogonality. We then multiply it by a scalar to make u have the same norm as before the modification in Eq. 2. In case we sampled u such that after the modification its norm is very small, we then sample it again. We proceed in this way as long as possible. When we have enough beta to form a complete basis set, we stop applying the orthogonality procedure and we sample them randomly again.

5 Serializable Components

All classes deriving from SerializableComponent (i.e. all components of our network) support serialization of their restorable configuration (i.e. no pointers) into simple strings, by a code system. It is explained in detail in the StringCodeUtilities.hpp header file and the examples, but will quickly be illustrated here.

Consider an output unit (type id OUT) of our neural network. It has two simple parameters shift and scale and itself two SerializableComponent members, the feeder RAY and the activation function LGS (default). If we assume the ray has no beta parameters set yet, the full serialized object string (treeCode) looks like:

```
string treeCode = "OUT ( shift 0.0 , scale 1.0 ) { RAY , LGS }"
```

If RAY and LGS had own parameters or members, they would be appended after their identifier, enclosed in () or {} brackets, respectively.

NOTE: Because the system relies on the spacing between every token, the spaces must not be neglected. Also a parameter value of string type may not contain any spaces, commas or brackets of kind () or $\{\}$.