QuCumber Documentation

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PIQuIL

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INSTALLATION

QuCumber only supports Python 3, not Python 2. If you are using Python 2, please update! You may also want to install PyTorch v1.0 (https://pytorch.org/), if you have not already.

If you're running a reasonably up-to-date Linux or macOS system, PyTorch should get installed automatically when you install QuCumber with *pip*.

1.1 Github

Navigate to the qucumber page on github (https://github.com/PIQuIL/QuCumber) and clone the repository by typing:

git clone https://github.com/PIQuIL/QuCumber.git

Navigate to the main directory and type:

```
python setup.py install
```

1.2 Windows

Navigate to the directory (through command prompt) where pip.exe is installed (usually C:\Python\Scripts\pip.exe) and type:

pip.exe install qucumber

1.3 Linux / macOS

Open up a terminal, then type:

pip install qucumber

TWO

THEORY

For a basic introduction to Restricted Boltzmann Machines, click here.

DOWNLOAD THE TUTORIALS

Once you have installed QuCumber, we recommend going through our tutorial that is divided into two parts.

- 1. Training a wave function to reconstruct a positive-real wave function (i.e. no phase) from a transverse-field Ising model (TFIM) and then generating new data.
- 2. Training an wave function to reconstruct a complex wave function (i.e. with a phase) from a simple two qubit random state and then generating new data.

We have made interactive python notebooks that can be downloaded (along with the data required) here. Note that the linked examples are from the most recent stable release (relative to the version of the docs you're currently viewing), and may not match the examples shown in the following pages. It is recommended that you refer to documentation for the latest stable release: https://qucumber.readthedocs.io/en/stable/.

If you wish to simply view the static, non-interactive notebooks, continue to the next page of the documentation.

Alternatively, you can view interactive notebooks online at: launch binder, though they may be slow.

RECONSTRUCTION OF A POSITIVE-REAL WAVEFUNCTION

This tutorial shows how to reconstruct a **positive-real** wavefunction via training a *Restricted Boltzmann Machine* (RBM), the neural network behind QuCumber. The data used for training are σ^z measurements from a onedimensional transverse-field Ising model (TFIM) with 10 sites at its critical point.

4.1 Transverse-field Ising model

The example dataset, located in tfimld_data.txt, comprises 10,000 σ^z measurements from a one-dimensional TFIM with 10 sites at its critical point. The Hamiltonian for the TFIM is given by

$$\mathcal{H} = -J\sum_{i}\sigma_{i}^{z}\sigma_{i+1}^{z} - h\sum_{i}\sigma_{i}^{x}$$

$$\tag{4.1}$$

where σ_i^z is the conventional spin-1/2 Pauli operator on site *i*. At the critical point, J = h = 1. By convention, spins are represented in binary notation with zero and one denoting the states spin-down and spin-up, respectively.

4.2 Using QuCumber to reconstruct the wavefunction

4.2.1 Imports

To begin the tutorial, first import the required Python packages.

```
[1]: import numpy as np
import matplotlib.pyplot as plt
from qucumber.nn_states import PositiveWaveFunction
from qucumber.callbacks import MetricEvaluator
import qucumber.utils.training_statistics as ts
import qucumber.utils.data as data
```

The Python class PositiveWaveFunction contains generic properties of a RBM meant to reconstruct a positivereal wavefunction, the most notable one being the gradient function required for stochastic gradient descent.

To instantiate a PositiveWaveFunction object, one needs to specify the number of visible and hidden units in the RBM. The number of visible units, num_visible, is given by the size of the physical system, i.e. the number of spins or qubits (10 in this case), while the number of hidden units, num_hidden, can be varied to change the expressiveness of the neural network.

Note: The optimal num_hidden: num_visible ratio will depend on the system. For the TFIM, having this ratio be equal to 1 leads to good results with reasonable computational effort.

4.2.2 Training

To evaluate the training in real time, we compute the fidelity between the true ground-state wavefunction of the system and the wavefunction that QuCumber reconstructs, $|\langle \psi | \psi_{RBM} \rangle|^2$, along with the Kullback-Leibler (KL) divergence (the RBM's cost function). As will be shown below, any custom function can be used to evaluate the training.

First, the training data and the true wavefunction of this system must be loaded using the data utility.

```
[2]: psi_path = "tfimld_psi.txt"
    train_path = "tfimld_data.txt"
    train_data, true_psi = data.load_data(train_path, psi_path)
```

As previously mentioned, to instantiate a PositiveWaveFunction object, one needs to specify the number of visible and hidden units in the RBM; we choose them to be equal.

```
[3]: nv = train_data.shape[-1]
nh = nv
nn_state = PositiveWaveFunction(num_visible=nv, num_hidden=nh)
# nn_state = PositiveWaveFunction(num_visible=nv, num_hidden=nh, gpu = False)
```

By default, QuCumber will attempt to run on a GPU, and default to CPU if GPU is not available. To run QuCumber on a CPU, add the flag gpu=False in the PositiveWaveFunction object instantiation (i.e. uncomment the line above).

Now we specify the hyperparameters of the training process:

- 1. epochs: the total number of training cycles that will be performed (default = 100)
- 2. pbs (pos_batch_size): the number of data points used in the positive phase of the gradient (default = 100)
- 3. nbs (neg_batch_size): the number of data points used in the negative phase of the gradient (default = 100)
- 4. k: the number of contrastive divergence steps (default = 1)
- 5. lr: the learning rate (default = 0.001)

Note: For more information on the hyperparameters above, it is strongly encouraged that the user to read through the brief, but thorough theory document on RBMs located in the QuCumber documentation. One does not have to specify these hyperparameters, as their default values will be used without the user overwriting them. It is recommended to keep with the default values until the user has a stronger grasp on what these hyperparameters mean. The quality and the computational efficiency of the training will highly depend on the choice of hyperparameters. As such, playing around with the hyperparameters is almost always necessary.

For the TFIM with 10 sites, the following hyperparameters give excellent results:

```
[4]: epochs = 500
  pbs = 100
  nbs = pbs
  lr = 0.01
  k = 10
```

For evaluating the training in real time, the MetricEvaluator is called every 100 epochs in order to calculate the training evaluators. The MetricEvaluator requires the following arguments:

- 1. period: the frequency of the training evaluators being calculated (e.g. period=100 means that the MetricEvaluator will do an evaluation every 100 epochs)
- 2. A dictionary of functions you would like to reference to evaluate the training (arguments required for these functions are keyword arguments placed after the dictionary)

The following additional arguments are needed to calculate the fidelity and KL divergence in the training_statistics utility:

- target_psi: the true wavefunction of the system
- space: the Hilbert space of the system

The training evaluators can be printed out via the verbose=True statement.

Although the fidelity and KL divergence are excellent training evaluators, they are not practical to calculate in most cases; the user may not have access to the target wavefunction of the system, nor may generating the Hilbert space of the system be computationally feasible. However, evaluating the training in real time is extremely convenient.

Any custom function that the user would like to use to evaluate the training can be given to the MetricEvaluator, thus avoiding having to calculate fidelity and/or KL divergence. Any custom function given to MetricEvaluator must take the neural-network state (in this case, the PositiveWaveFunction object) and keyword arguments. As an example, we define a custom function psi_coefficient, which is the fifth coefficient of the reconstructed wavefunction multiplied by a parameter A.

```
[5]: def psi_coefficient(nn_state, space, A, **kwargs):
    norm = nn_state.compute_normalization(space).sqrt_()
    return A * nn_state.psi(space)[0][4] / norm
```

Now the Hilbert space of the system can be generated for the fidelity and KL divergence.

```
[6]: period = 10
space = nn_state.generate_hilbert_space(nv)
```

Now the training can begin. The PositiveWaveFunction object has a property called fit which takes care of this. MetricEvaluator must be passed to the fit function in a list (callbacks).

```
[7]: callbacks = [
        MetricEvaluator(
            period,
            {"Fidelity": ts.fidelity, "KL": ts.KL, "A_rbm_5": psi_coefficient},
            target_psi=true_psi,
            verbose=True,
            space=space,
            A=3.0,
        )
    ]
    nn_state.fit(
        train_data,
        epochs=epochs,
        pos_batch_size=pbs,
        neg_batch_size=nbs,
        lr=lr,
        k=k,
        callbacks=callbacks,
    )
    Epoch: 10
                    Fidelity = 0.526148
                                            KL = 1.310731
                                                            A_rbm_5 = 0.125463
                    Fidelity = 0.631814
    Epoch: 20
                                            KL = 0.875887
                                                            A_{rbm_5} = 0.193193
    Epoch: 30
                    Fidelity = 0.736986
                                            KL = 0.577408
                                                            A_rbm_5 = 0.249697
                    Fidelity = 0.794626
                                                            A_rbm_5 = 0.267554
    Epoch: 40
                                            KL = 0.445550
                    Fidelity = 0.828487
    Epoch: 50
                                            KL = 0.363523
                                                            A_rbm_5 = 0.263156
                                            KL = 0.284768
    Epoch: 60
                    Fidelity = 0.861033
                                                            A_rbm_5 = 0.255909
    Epoch: 70
                    Fidelity = 0.888133
                                            KL = 0.226607
                                                            A_rbm_5 = 0.251317
```

				1 1 2 /
Epoch:	80	Fidelity = 0.904473	KL = 0.191903	$A_rbm_5 = 0.230342$
Epoch:	90	Fidelity = 0.916896	KL = 0.168523	$A_rbm_5 = 0.232834$
Epoch:	100	Fidelity = 0.925543	KL = 0.151414	$A_rbm_5 = 0.226578$
Epoch:	110	Fidelity = 0.933069	KL = 0.136249	$A_rbm_5 = 0.227657$
Epoch:	120	Fidelity = 0.939533	KL = 0.122066	$A_rbm_5 = 0.216086$
Epoch:	130	Fidelity = 0.945398	KL = 0.109634	$A_rbm_5 = 0.210336$
Epoch:	140	Fidelity = 0.950329	KL = 0.099964	$A_rbm_5 = 0.214536$
Epoch:	150	Fidelity = 0.954255	KL = 0.092397	$A_rbm_5 = 0.212398$
Epoch:	160	Fidelity = 0.957539	KL = 0.086165	$A_rbm_5 = 0.213869$
Epoch:	170	Fidelity = 0.959890	KL = 0.081415	$A_rbm_5 = 0.205124$
Epoch:	180	Fidelity = 0.961762	KL = 0.077955	$A_{rbm_5} = 0.207600$
Epoch:	190	Fidelity = 0.963395	KL = 0.075018	$A_rbm_5 = 0.203214$
Epoch:	200	Fidelity = 0.965103	KL = 0.071877	$A_{rbm_5} = 0.207948$
Epoch:	210	Fidelity = 0.966435	KL = 0.069428	$A_rbm_5 = 0.216086$
Epoch:	220	Fidelity = 0.967274	KL = 0.067780	$A_rbm_5 = 0.215082$
Epoch:	230	Fidelity = 0.968685	KL = 0.064706	$A_{rbm_5} = 0.211092$
Epoch:	240	Fidelity = 0.969841	KL = 0.062323	$A_{rbm_5} = 0.213523$
Epoch:	250	Fidelity = 0.971052	KL = 0.059850	$A_{rbm_5} = 0.212783$
Epoch:	260	Fidelity = 0.971965	KL = 0.057842	$A_rbm_5 = 0.208115$
Epoch:	270	Fidelity = 0.973736	KL = 0.054289	$A_{rbm_5} = 0.215748$
Epoch:	280	Fidelity = 0.974085	KL = 0.053346	$A_{rbm_5} = 0.212171$
Epoch:	290	Fidelity = 0.976066	KL = 0.049299	$A_{rbm_5} = 0.219986$
Epoch:	300	Fidelity = 0.977303	KL = 0.046733	$A_{rbm_5} = 0.225259$
Epoch:	310	Fidelity = 0.978261	KL = 0.044790	$A_rbm_5 = 0.228821$
Epoch:	320	Fidelity = 0.979351	KL = 0.042555	$A_{rbm_5} = 0.225733$
Epoch:	330	Fidelity = 0.980212	KL = 0.040565	$A_{rbm_5} = 0.223765$
Epoch:	340	Fidelity = 0.981664	KL = 0.037660	$A_{rbm_5} = 0.226980$
Epoch:	350	Fidelity = 0.982528	KL = 0.035918	$A_{rbm_5} = 0.230829$
Epoch:	360	Fidelity = 0.983351	KL = 0.034181	$A_{rbm_5} = 0.224962$
Epoch:	370	Fidelity = 0.984213	KL = 0.032504	$A_{rbm_5} = 0.225617$
Epoch:	380	Fidelity = 0.984872	KL = 0.031177	$A_{rbm_5} = 0.227120$
Epoch:	390	Fidelity = 0.985186	KL = 0.030594	$A_{rbm_5} = 0.222515$
Epoch:	400	Fidelity = 0.985662	KL = 0.029606	$A_{rbm_5} = 0.220782$
Epoch:	410	Fidelity = 0.986466	KL = 0.028079	$A_{rbm_5} = 0.227727$
Epoch:	420	Fidelity = 0.986970	KL = 0.027100	$A_{rbm_5} = 0.233300$
Epoch:	430	Fidelity = 0.987040	KL = 0.026978	$A_{rbm_5} = 0.232759$
Epoch:	440	Fidelity = 0.987675	KL = 0.025714	$A_{rbm_5} = 0.224514$
Epoch:	450	Fidelity = 0.988244	KL = 0.024636	A_rbm_5 = 0.229669
Epoch:	460	Fidelity = 0.988569	KL = 0.023975	$A_{rbm_5} = 0.230897$
Epoch:	470	Fidelity = 0.988666	KL = 0.023802	$A_rbm_5 = 0.229378$
Epoch:	480	Fidelity = 0.988781	KL = 0.023565	$A_rbm_5 = 0.236488$
Epoch:	490	Fidelity = 0.989243	KL = 0.022694	$A_rbm_5 = 0.228858$
Epoch:	500	Fidelity = 0.988991	KL = 0.023196	A_rbm_5 = 0.235301

All of these training evaluators can be accessed after the training has completed. The code below shows this, along with plots of each training evaluator as a function of epoch (training cycle number).

```
[8]: # Note that the key given to the *MetricEvaluator* must be
# what comes after callbacks[0].
fidelities = callbacks[0].Fidelity
# Alternatively, we can use the usual dictionary/list subsripting
# syntax. This is useful in cases where the name of the
# metric contains special characters or spaces.
KLs = callbacks[0]["KL"]
coeffs = callbacks[0]["A_rbm_5"]
```

```
epoch = np.arange(period, epochs + 1, period)
 [9]: # Some parameters to make the plots look nice
      params = {
           "text.usetex": True,
           "font.family": "serif",
           "legend.fontsize": 14,
           "figure.figsize": (10, 3),
           "axes.labelsize": 16,
          "xtick.labelsize": 14,
          "ytick.labelsize": 14,
          "lines.linewidth": 2,
          "lines.markeredgewidth": 0.8,
          "lines.markersize": 5,
           "lines.marker": "o",
           "patch.edgecolor": "black",
      }
      plt.rcParams.update(params)
      plt.style.use("seaborn-deep")
[10]: # Plotting
      fig, axs = plt.subplots(nrows=1, ncols=3, figsize=(14, 3))
      ax = axs[0]
      ax.plot(epoch, fidelities, "o", color="CO", markeredgecolor="black")
      ax.set_ylabel(r"Fidelity")
      ax.set_xlabel(r"Epoch")
      ax = axs[1]
      ax.plot(epoch, KLs, "o", color="C1", markeredgecolor="black")
      ax.set_ylabel(r"KL Divergence")
      ax.set_xlabel(r"Epoch")
      ax = axs[2]
      ax.plot(epoch, coeffs, "o", color="C2", markeredgecolor="black")
      ax.set_ylabel(r"$A\psi_{RBM}[5]$")
      ax.set_xlabel(r"Epoch")
      plt.tight_layout()
      plt.savefig("fid_KL.pdf")
      plt.show()
         1.0
                                       Divergence
                                                                         0.25
                                                                       A\psi_{RBM}[5]
       Fidelity
8'0
                                                                         0.20
                                       КL
                                                                         0.15
         0.6
                                         0.0
                                                100
                100
                     200
                         300
                              400
                                   500
                                                          300
                                                               400
                                                                    500
                                                                                 100
                                                                                      200
                                                                                           300
                                                                                                400
                                                                                                     500
           0
                                            0
                                                     200
                                                                             0
                      \operatorname{Epoch}
                                                       Epoch
                                                                                       Epoch
```

It should be noted that one could have just ran nn_state.fit(train_samples), which uses the default hyperparameters and no training evaluators.

To demonstrate how important it is to find the optimal hyperparameters for a certain system, restart this notebook and comment out the original fit statement, then uncomment and run the cell below.

[11]: # nn_state.fit(train_samples)

Using the non-default hyperparameters yielded a fidelity of approximately 0.989, while the default hyperparameters yield approximately 0.523!

The trained RBM's parameters are saved to a pickle file with the name <code>saved_params.pt</code> for future use in other tutorials:

[12]: nn_state.save("saved_params.pt")

This saves the weights, visible biases and hidden biases as torch tensors with the following keys: "weights", "visible_bias", "hidden_bias".

RECONSTRUCTION OF A COMPLEX WAVEFUNCTION

In this tutorial, a walkthrough of how to reconstruct a **complex** wavefunction via training a *Restricted Boltzmann Machine* (RBM), the neural network behind QuCumber, will be presented.

5.1 The wavefunction to be reconstructed

The simple wavefunction below describing two qubits (coefficients stored in qubits_psi.txt) will be reconstructed.

$$|\psi\rangle = \alpha|00\rangle + \beta|01\rangle + \gamma|10\rangle + \delta|11\rangle \tag{5.1}$$

where the exact values of α,β,γ and δ used for this tutorial are

$$\alpha = 0.2861 + 0.0539i \tag{5.2}$$

$$\beta = 0.3687 - 0.3023i \tag{5.3}$$

$$\gamma = -0.1672 - 0.3529i \tag{5.4}$$

$$\delta = -0.5659 - 0.4639i. \tag{5.5}$$

The example dataset, qubits_train.txt, comprises of 500σ measurements made in various bases (X, Y and Z). A corresponding file containing the bases for each data point in qubits_train.txt, qubits_train_bases.txt, is also required. As per convention, spins are represented in binary notation with zero and one denoting spin-down and spin-up, respectively.

5.2 Using qucumber to reconstruct the wavefunction

5.2.1 Imports

To begin the tutorial, first import the required Python packages.

```
[1]: import numpy as np
import torch
import matplotlib.pyplot as plt
from qucumber.nn_states import ComplexWaveFunction
from qucumber.callbacks import MetricEvaluator
import qucumber.utils.unitaries as unitaries
import qucumber.utils.cplx as cplx
```

import qucumber.utils.training_statistics as ts
import qucumber.utils.data as data

The Python class ComplexWaveFunction contains generic properties of a RBM meant to reconstruct a complex wavefunction, the most notable one being the gradient function required for stochastic gradient descent.

To instantiate a ComplexWaveFunction object, one needs to specify the number of visible and hidden units in the RBM. The number of visible units, num_visible, is given by the size of the physical system, i.e. the number of spins or qubits (2 in this case), while the number of hidden units, num_hidden, can be varied to change the expressiveness of the neural network.

Note: The optimal num_hidden : num_visible ratio will depend on the system. For the two-qubit wavefunction described above, good results are yielded when this ratio is 1.

On top of needing the number of visible and hidden units, a ComplexWaveFunction object requires the user to input a dictionary containing the unitary operators (2x2) that will be used to rotate the qubits in and out of the computational basis, Z, during the training process. The unitaries utility will take care of creating this dictionary.

The MetricEvaluator class and training_statistics utility are built-in amenities that will allow the user to evaluate the training in real time.

Lastly, the cplx utility allows QuCumber to be able to handle complex numbers. Currently, PyTorch does not support complex numbers.

5.2.2 Training

To evaluate the training in real time, the fidelity between the true wavefunction of the system and the wavefunction that QuCumber reconstructs, $|\langle \psi | \psi_{RBM} \rangle|^2$, will be calculated along with the Kullback-Leibler (KL) divergence (the RBM's cost function). First, the training data and the true wavefunction of this system need to be loaded using the data utility.

```
[2]: train_path = "qubits_train.txt"
    train_bases_path = "qubits_train_bases.txt"
    psi_path = "qubits_psi.txt"
    bases_path = "qubits_bases.txt"
    train_samples, true_psi, train_bases, bases = data.load_data(
        train_path, psi_path, train_bases_path, bases_path
)
```

The file qubits_bases.txt contains every unique basis in the qubits_train_bases.txt file. Calculation of the full KL divergence in every basis requires the user to specify each unique basis.

As previously mentioned, a ComplexWaveFunction object requires a dictionary that contains the unitary operators that will be used to rotate the qubits in and out of the computational basis, Z, during the training process. In the case of the provided dataset, the unitaries required are the well-known H, and K gates. The dictionary needed can be created with the following command.

If the user wishes to add their own unitary operators from their experiment to $unitary_dict$, uncomment the block above. When $unitaries.create_dict()$ is called, it will contain the identity and the H and K gates by default under the keys "Z", "X" and "Y", respectively.

The number of visible units in the RBM is equal to the number of qubits. The number of hidden units will also be taken to be the number of visible units.

By default, QuCumber will attempt to run on a GPU if one is available (if one is not available, QuCumber will fall back to CPU). If one wishes to guarantee that QuCumber runs on the CPU, add the flag gpu=False in the ComplexWaveFunction object instantiation. Set gpu=True in the line above to run this tutorial on a GPU.

Now the hyperparameters of the training process can be specified.

- 1. epochs: the total number of training cycles that will be performed (default = 100)
- 2. pos_batch_size: the number of data points used in the positive phase of the gradient (default = 100)
- 3. neg_batch_size: the number of data points used in the negative phase of the gradient (default = pos_batch_size)
- 4. k: the number of contrastive divergence steps (default = 1)
- 5. lr: the learning rate (default = 0.001)

Note: For more information on the hyperparameters above, it is strongly encouraged that the user to read through the brief, but thorough theory document on RBMs. One does not have to specify these hyperparameters, as their default values will be used without the user overwriting them. It is recommended to keep with the default values until the user has a stronger grasp on what these hyperparameters mean. The quality and the computational efficiency of the training will highly depend on the choice of hyperparameters. As such, playing around with the hyperparameters is almost always necessary.

The two-qubit example in this tutorial should be extremely easy to train, regardless of the choice of hyperparameters. However, the hyperparameters below will be used.

```
[5]: epochs = 100
    pbs = 50  # pos_batch_size
    nbs = 50  # neg_batch_size
    lr = 0.1
    k = 5
```

For evaluating the training in real time, the MetricEvaluator will be called to calculate the training evaluators every 10 epochs. The MetricEvaluator requires the following arguments.

- 1. period: the frequency of the training evaluators being calculated (e.g. period=200 means that the MetricEvaluator will compute the desired metrics every 200 epochs)
- 2. A dictionary of functions you would like to reference to evaluate the training (arguments required for these functions are keyword arguments placed after the dictionary)

The following additional arguments are needed to calculate the fidelity and KL divergence in the training_statistics utility.

- target_psi (the true wavefunction of the system)
- space (the entire Hilbert space of the system)

The training evaluators can be printed out via the verbose=True statement.

Although the fidelity and KL divergence are excellent training evaluators, they are not practical to calculate in most cases; the user may not have access to the target wavefunction of the system, nor may generating the Hilbert space of the system be computationally feasible. However, evaluating the training in real time is extremely convenient.

Any custom function that the user would like to use to evaluate the training can be given to the MetricEvaluator, thus avoiding having to calculate fidelity and/or KL divergence. As an example, functions that calculate the the norm of each of the reconstructed wavefunction's coefficients are presented. Any custom function given to MetricEvaluator must take the neural-network state (in this case, the ComplexWaveFunction object) and keyword arguments. Although the given example requires the Hilbert space to be computed, the scope of the MetricEvaluator's ability to be able to handle any function should still be evident.

```
[6]: def alpha(nn_state, space, **kwargs):
        rbm_psi = nn_state.psi(space)
        normalization = nn_state.compute_normalization(space).sqrt_()
        alpha_ = cplx.norm(
            torch.tensor([rbm_psi[0][0], rbm_psi[1][0]], device=nn_state.device)
            / normalization
        )
        return alpha_
    def beta(nn_state, space, **kwargs):
        rbm_psi = nn_state.psi(space)
        normalization = nn_state.compute_normalization(space).sqrt_()
        beta_ = cplx.norm(
            torch.tensor([rbm_psi[0][1], rbm_psi[1][1]], device=nn_state.device)
            / normalization
        )
        return beta_
    def gamma(nn_state, space, **kwargs):
        rbm_psi = nn_state.psi(space)
        normalization = nn_state.compute_normalization(space).sqrt_()
        gamma_ = cplx.norm(
            torch.tensor([rbm_psi[0][2], rbm_psi[1][2]], device=nn_state.device)
            / normalization
        )
        return gamma_
    def delta(nn_state, space, **kwargs):
        rbm_psi = nn_state.psi(space)
        normalization = nn_state.compute_normalization(space).sqrt_()
        delta_ = cplx.norm(
            torch.tensor([rbm_psi[0][3], rbm_psi[1][3]], device=nn_state.device)
             / normalization
        )
        return delta_
```

Now the Hilbert space of the system must be generated for the fidelity and KL divergence and the dictionary of functions the user would like to compute every period epochs must be given to the MetricEvaluator. Note that some of the coefficients aren't being evaluated as they are commented out. This is simply to avoid cluttering the output, and may be uncommented by the user.

```
[7]: period = 2
    space = nn_state.generate_hilbert_space(nv)
    callbacks = [
        MetricEvaluator(
           period,
            {
                 "Fidelity": ts.fidelity,
                "KL": ts.KL,
                "norm": alpha,
                 # "norm": beta,
                 # "norm": gamma,
                 # "norm": delta,
            },
            target_psi=true_psi,
            bases=bases,
            verbose=True,
            space=space,
        )
    ]
```

Now the training can begin. The ComplexWaveFunction object has a function called fit which takes care of this.

[8]:	nn_state.fit(
------	---------------

<pre>train_samples, epochs=epochs, pos_batch_size=pbs, neg_batch_size=nbs, lr=lr, k=k, input_bases=train_bases, callbacks=callbacks,</pre>					
Epoch: 2	Fidelity = 0.623747	KL = 0.226386	norm = 0.272518		
Epoch: 4	Fidelity = 0.744691	KL = 0.142639	norm = 0.248872		
Epoch: 6	Fidelity = 0.818254	KL = 0.094584	norm = 0.263589		
Epoch: 8	Fidelity = 0.867098	KL = 0.067506	norm = 0.278453		
Epoch: 10	Fidelity = 0.900217	KL = 0.051592	norm = 0.281094		
Epoch: 12	Fidelity = 0.922993	KL = 0.041311	norm = 0.276052		
Epoch: 14	Fidelity = 0.937807	KL = 0.034972	norm = 0.274676		
Epoch: 16	Fidelity = 0.947232	KL = 0.030543	norm = 0.283873		
Epoch: 18	Fidelity = 0.955277	KL = 0.027313	norm = 0.278906		
Epoch: 20	Fidelity = 0.959930	KL = 0.025034	norm = 0.290271		
Epoch: 22	Fidelity = 0.963333	KL = 0.023719	norm = 0.296183		
Epoch: 24	Fidelity = 0.969419	KL = 0.021086	norm = 0.276108		
Epoch: 26	Fidelity = 0.972300	KL = 0.020200	norm = 0.290305		
Epoch: 28	Fidelity = 0.974777	KL = 0.018635	norm = 0.284231		
Epoch: 30	Fidelity = 0.976208	KL = 0.017865	norm = 0.282036		
Epoch: 32	Fidelity = 0.978382	KL = 0.016862	norm = 0.282498		
Epoch: 34	Fidelity = 0.980578	KL = 0.015977	norm = 0.279435		
Epoch: 36	Fidelity = 0.980983	KL = 0.015545	norm = 0.277835		
Epoch: 38	Fidelity = 0.982651	KL = 0.014751	norm = 0.280070		
Epoch: 40	Fidelity = 0.983155	KL = 0.014353	norm = 0.276912		
Epoch: 42	Fidelity = 0.983996	KL = 0.013827	norm = 0.278844		
Epoch: 44	Fidelity = 0.982731	KL = 0.015100	norm = 0.305219		
Epoch: 46	Fidelity = 0.984791	KL = 0.013417	norm = 0.293674		

				· · · · · · · · · · · · · · · · · · ·	1 107
Epoch:	48	Fidelity = 0.98539	5 KL = 0.012845	norm = 0.280658	
Epoch:	50	Fidelity = 0.98676	7 KL = 0.012093	norm = 0.277599	
Epoch:	52	Fidelity = 0.98779	5 KL = 0.011650	norm = 0.278886	
Epoch:	54	Fidelity = 0.98705	7 KL = 0.011843	norm = 0.271735	
Epoch:	56	Fidelity = 0.98712	5 KL = 0.011552	norm = 0.280304	
Epoch:	58	Fidelity = 0.98729	5 KL = 0.011382	norm = 0.288229	
Epoch:	60	Fidelity = 0.98820	1 KL = 0.011201	norm = 0.266736	
Epoch:	62	Fidelity = 0.98918	1 KL = 0.010504	norm = 0.288520	
Epoch:	64	Fidelity = 0.98930	8 KL = 0.010293	norm = 0.292218	
Epoch:	66	Fidelity = 0.98932	1 KL = 0.009901	norm = 0.282069	
Epoch:	68	Fidelity = 0.98934	7 KL = 0.009836	norm = 0.275723	
Epoch:	70	Fidelity = 0.98949	4 KL = 0.009838	norm = 0.293840	
Epoch:	72	Fidelity = 0.99011	5 KL = 0.009225	norm = 0.282556	
Epoch:	74	Fidelity = 0.99019	9 $KL = 0.009095$	norm = 0.278911	
Epoch:	76	Fidelity = 0.98997	9 $KL = 0.009214$	norm = 0.273241	
Epoch:	78	Fidelity = 0.98963	3 KL = 0.009275	norm = 0.274384	
Epoch:	80	Fidelity = 0.98997	2 KL = 0.008976	norm = 0.275430	
Epoch:	82	Fidelity = 0.98992	0 KL = 0.008871	norm = 0.285605	
Epoch:	84	Fidelity = 0.99117	7 KL = 0.008183	norm = 0.282607	
Epoch:	86	Fidelity = 0.99124	9 $KL = 0.008095$	norm = 0.276934	
Epoch:	88	Fidelity = 0.99085	7 KL = 0.008273	norm = 0.272151	
Epoch:	90	Fidelity = 0.99080	2 KL = 0.008071	norm = 0.280823	
Epoch:	92	Fidelity = 0.99109	0 KL = 0.007838	norm = 0.279963	
Epoch:	94	Fidelity = 0.99099	5 KL = 0.007861	norm = 0.275772	
Epoch:	96	Fidelity = 0.99032	6 KL = 0.008202	norm = 0.289882	
Epoch:	98	Fidelity = 0.99101	2 KL = 0.007690	norm = 0.277037	
Epoch:	100	Fidelity = 0.99173	6 KL = 0.007292	norm = 0.275516	

All of these training evaluators can be accessed after the training has completed, as well. The code below shows this, along with plots of each training evaluator versus the training cycle number (epoch).

```
[9]: # Note that the key given to the *MetricEvaluator* must be
  # what comes after callbacks[0].
  fidelities = callbacks[0].Fidelity
  # Alternatively, we may use the usual dictionary/list subscripting
  # syntax. This is useful in cases where the name of the metric
  # may contain special characters or spaces.
  KLs = callbacks[0]["KL"]
  coeffs = callbacks[0]["norm"]
  epoch = np.arange(period, epochs + 1, period)
```

```
[10]: # Some parameters to make the plots look nice
params = {
    "text.usetex": True,
    "font.family": "serif",
    "legend.fontsize": 14,
    "figure.figsize": 14,
    "figure.figsize": 16,
    "xtick.labelsize": 14,
    "ytick.labelsize": 14,
    "ytick.labelsize": 14,
    "lines.linewidth": 2,
    "lines.markeredgewidth": 0.8,
    "lines.markersize": 5,
    "lines.markersize": 5,
    "lines.marker": "o",
    "patch.edgecolor": "black",
```



It should be noted that one could have just ran nn_state.fit(train_samples) and just used the default hyperparameters and no training evaluators.

At the end of the training process, the network parameters (the weights, visible biases and hidden biases) are stored in the ComplexWaveFunction object. One can save them to a pickle file, which will be called saved_params.pt, with the following command.

[12]: nn_state.save("saved_params.pt")

This saves the weights, visible biases and hidden biases as torch tensors with the following keys: "weights", "visible_bias", "hidden_bias".

SIX

SAMPLING AND CALCULATING OBSERVABLES

6.1 Generate new samples

Firstly, to generate meaningful data, an RBM needs to be trained. Please refer to the tutorials 1 and 2 on training an RBM if how to train an RBM using QuCumber is unclear. An RBM with a positive-real wavefunction describing a transverse-field Ising model (TFIM) with 10 sites has already been trained in the first tutorial, with the parameters of the machine saved here as saved_params.pt. The autoload function can be employed here to instantiate the corresponding PositiveWaveFunction object from the saved RBM parameters.

```
[1]: import numpy as np
import matplotlib.pyplot as plt
from qucumber.nn_states import PositiveWaveFunction
from qucumber.observables import ObservableBase
from quantum_ising_chain import TFIMChainEnergy, Convergence
nn_state = PositiveWaveFunction.autoload("saved_params.pt")
```

A PositiveWaveFunction object has a property called sample that allows us to sample the learned distribution of TFIM chains. The it takes the following arguments (along with a few others which are not relevant for our purposes):

- 1. k: the number of Gibbs steps to perform to generate the new samples. Increasing this number will produce samples closer to the learned distribution, but will require more computation.
- 2. num_samples: the number of new data points to be generated

```
[2]: new_samples = nn_state.sample(k=100, num_samples=10000)
print(new_samples)
tensor([[1., 0., 1., ..., 1., 1., 1.],
       [1., 1., 1., ..., 1., 0., 0.],
       [0., 1., 1., ..., 1., 1.],
       [1., 0., 0., ..., 0., 1., 1.],
       [0., 0., 1., ..., 1., 1.],
       [1., 1., 1., ..., 1., 0., 0.]], dtype=torch.float64)
```

6.1.1 Magnetization

With the newly generated samples, the user can now easily calculate observables that do not require any details associated with the RBM. A great example of this is the magnetization. To calculate the magnetization, the newly-

generated samples must be converted to ± 1 from 1 and 0, respectively. The function below does the trick.

```
[3]: def to_pml(samples):
    return samples.mul(2.0).sub(1.0)
```

Now, the (absolute) magnetization in the Z-direction is calculated as follows.

```
[4]: def Magnetization (samples):
    return to_pml(samples).mean(1).abs().mean()
magnetization = Magnetization(new_samples).item()
print("Magnetization = %.5f" % magnetization)
Magnetization = 0.55246
```

The exact value for the magnetization is 0.5610.

The magnetization and the newly-generated samples can also be saved to a pickle file along with the RBM parameters in the *PositiveWaveFunction* object.

```
[5]: nn_state.save(
    "saved_params_and_new_data.pt",
    metadata={"samples": new_samples, "magnetization": magnetization},
)
```

The metadata argument in the save function takes in a dictionary of data that you would like to save alongside the RBM parameters.

6.2 Calculate an observable using the Observable module

6.2.1 Magnetization (again)

QuCumber provides the Observable module to simplify estimation of expectations and variances of observables in memory efficient ways. To start off, we'll repeat the above example using the SigmaZ Observable module provided with QuCumber.

```
[6]: from qucumber.observables import SigmaZ
```

We'll compute the absolute magnetization again, for the sake of comparison with the previous example. We want to use the samples drawn earlier to perform this estimate, so we use the statistics_from_samples function:

```
[7]: sz = SigmaZ(absolute=True)
sz.statistics_from_samples(nn_state, new_samples)
[7]: {'mean': 0.5524600000000002,
'variance': 0.09724167256725606,
'std_error': 0.0031183597061156376}
```

With this function we get the variance and standard error for free. Now you may be asking: "That isn't too difficult, I could have computed those myself!". The power of the Observable module comes from the fact that it simplifies estimation of these values over a large number of samples. The statistics function computes these statistics by generating the samples internally. Let's see it in action:

```
[8]: %time sz.statistics(nn_state, num_samples=10000, burn_in=100)
# just think of burn_in as being equivalent to k for now
CPU times: user 1.77 s, sys: 5.24 ms, total: 1.78 s
Wall time: 582 ms
```

```
[8]: {'mean': 0.548680000000001,
    'variance': 0.10007226482647404,
    'std_error': 0.0031634200610490227}
```

Let's consider what is taking place under the hood at the moment. The statistics function is drawing 10000 samples from the given RBM state, and cycling it through the visible and hidden layers for 100 Block Gibbs steps before computing the statistics. This means that, at any given time it has to hold a matrix with 10000 rows and 10 (the number of lattice sites) columns in memory, which becomes infeasible for large lattices or if we want to use more samples to bring our standard error down. To bypass this issue, the statistics function allows us to specify the number of Markov Chains to evolve using the RBM, and will sample from these chains multiple times to produce enough samples. It takes the following arguments:

- num_samples: the number of samples to generate internally
- num_chains: the number of Markov chains to run in parallel (default = 0, meaning num_chains = num_samples)
- burn_in: the number of Gibbs steps to perform before recording any samples (default = 1000)
- steps: the number of Gibbs steps to perform between each sample; increase this to reduce the autocorrelation between samples (default = 1)

The statistics function will also return a dictionary containing the mean, standard error (of the mean) and the variance with the keys "mean", "std_error" and "variance", respectively.

```
[9]: %time sz.statistics(nn_state, num_samples=10000, num_chains=1000, burn_in=100, _

→ steps=2)

CPU times: user 292 ms, sys: 0 ns, total: 292 ms
```

```
Wall time: 75.9 ms
```

```
[9]: {'mean': 0.548,
    'variance': 0.09837783778377833,
    'std_error': 0.0031365241555546537}
```

In addition to using less memory (since the matrix held in memory is now of size num_chains x num_sites = 1000 x 10), we've also achieved a decent speed boost! Next, we'll try increasing the total number of drawn samples:

```
[10]: sz.statistics(nn_state, num_samples=int(le7), num_chains=1000, burn_in=100, steps=2)
```

Note how much we've decreased our standard error just by increasing the number of drawn samples. Finally, we can also draw samples of measurements of the observable using the sample function:

```
[11]: sz.sample(nn_state, k=100, num_samples=50)
```

Note that this function does not perform any fancy sampling tricks like statistics and is therefore susceptible to "Out of Memory" errors.

6.2.2 TFIM Energy

Some observables cannot be computed directly from samples, but instead depend on the RBM as previously mentioned. For example, the magnetization of the TFIM simply depends on the samples the user gives as input. While we did provide the nn_state as an argument when calling statistics_from_samples, SigmaZ ignores it. The TFIM energy, on the other hand, is much more complicated. Consider the TFIM Hamiltonian:

$$H = -J\sum_i \sigma_i^z \sigma_{i+1}^z - h\sum_i \sigma_i^x$$

As our RBM was trained in the Z-basis, the off-diagonal transverse-field term is impossible to compute just from the samples; we need to know the value of the wavefunction for each sample as well. An example for the computation of the energy is provided in the python file quantum_ising_chain.py, which takes advantage of QuCumber's Observable module.

quantum_ising_chain.py comprises of a class that computes the energy of a TFIM (TFIMChainEnergy) that inherits properties from the Observable module. To instantiate a TFIMChainEnergy object, the $\frac{h}{J}$ value must be specified. The trained RBM parameters are from the first tutorial, where the example data was from the TFIM with 10 sites at its critical point ($\frac{h}{T} = 1$).

$$[12]: h = 1$$

```
tfim_energy = TFIMChainEnergy(h)
```

To go ahead and calculate the mean energy and its standard error from the previously generated samples from this tutorial (new_samples), the statistics_from_samples function in the Observable module is called upon.

```
[13]: energy_stats = tfim_energy.statistics_from_samples(nn_state, new_samples)
print("Mean: %.4f" % energy_stats["mean"], "+/- %.4f" % energy_stats["std_error"])
print("Variance: %.4f" % energy_stats["variance"])
Mean: -1.2347 +/- 0.0005
Variance: 0.0022
```

The exact value for the energy is -1.2381.

To illustrate how quickly the energy converges as a function of the sampling step (i.e. the number of Gibbs steps to perform to generate a new batch of samples), steps, the Convergence function in quantum_ising_chain. py will do the trick. Convergence creates a batch of random samples initially, which is then used to generate a new batch of samples from the RBM. The TFIM energy will be calculated at every Gibbs step. Note that this function is not available in the QuCumber API; it is only used here as an illustrative example.

```
[14]: steps = 200
num_samples = 10000
dict_observables = Convergence(nn_state, tfim_energy, num_samples, steps)
energy = dict_observables["energies"]
err_energy = dict_observables["error"]
step = np.arange(steps + 1)
E0 = -1.2381
```

```
ax = plt.axes()
ax.plot(step, abs((E0 - energy) / E0) * 100, color="red")
ax.hlines(abs((E0 - energy_stats["mean"]) / E0) * 100, 0, 200, color="black")
ax.set_xlim(0, steps)
ax.set_ylim(0, 0.6)
ax.set_xlabel("Gibbs Step")
ax.set_ylabel("% Error in Energy")
```

[14]: Text(0, 0.5, '% Error in Energy')



One can see a brief transient period in the magnetization observable, before the state of the machine "warms up" to equilibrium (this explains the burn_in argument we saw earlier). After that, the values fluctuate around the estimated mean (the horizontal black line).

6.2.3 Adding observables

One may also add / subtract and multiply observables with each other or with real numbers. To illustrate this, we will build an alternative implementation of the TFIM energy observable. First, we will introduce the built-in NeighbourInteraction observable:

```
[15]: from qucumber.observables import NeighbourInteraction
```

The TFIM chain we trained the RBM on did not have periodic boundary conditions, so periodic_bcs=False. Meanwhile, c specifies the between interacting spins, that is, a given site will only interact with a site c places away from itself; we set this to 1 as the TFIM chain has nearest-neighbour interactions.

```
[16]: nn_inter = NeighbourInteraction(periodic_bcs=False, c=1)
```

Next, we need the SigmaX observable, which computes the magnetization in the X-direction:

```
[17]: from qucumber.observables import SigmaX
```

Next, we build the Hamiltonian, setting h = J = 1:

```
[18]: h = J = 1
sx = SigmaX()
tfim = -J * nn_inter - h * sx
```

The same statistics of this new TFIM observable can also be calculated.

```
[19]: new_tfim_stats = tfim.statistics_from_samples(nn_state, new_samples)
print("Mean: %.4f" % new_tfim_stats["mean"], "+/- %.4f" % new_tfim_stats["std_error"])
print("Variance: %.4f" % new_tfim_stats["variance"])
Mean: -1.2347 +/- 0.0005
Variance: 0.0022
```

The statistics above match with those computed earlier.

6.2.4 Renyi Entropy and the Swap operator

We can estimate the second Renyi Entropy using the Swap operator as shown by Hastings et al. (2010). The 2nd Renyi Entropy, in terms of the expectation of the Swap operator is given by:

$$S_2(A) = -\ln\langle \operatorname{Swap}_A \rangle$$

where A is the subset of the lattice for which we wish to compute the Renyi entropy.

```
[20]: from qucumber.observables import SWAP
```

As an example, we will take the region A consist of sites 0 through 4 (inclusive).

```
[21]: A = [0, 1, 2, 3, 4]
swap = SWAP(A)
swap_stats = swap.statistics_from_samples(nn_state, new_samples)
print("Mean: %.4f" % swap_stats["mean"], "+/- %.4f" % swap_stats["std_error"])
print("Variance: %.4f" % swap_stats["variance"])
Mean: 0.7937 +/- 0.0083
Variance: 0.3484
```

The 2nd Renyi Entropy can be computed directly from the sample mean. The standard error of the entropy, from firstorder error analysis, is given by the standard error of the Swap operator divided by the mean of the Swap operator.

```
[22]: S_2 = -np.log(swap_stats["mean"])
S_2_error = abs(swap_stats["std_error"] / swap_stats["mean"])
print("S_2: %.4f" % S_2, "+/- %.4f" % S_2_error)
S_2: 0.2310 +/- 0.0105
```

6.2.5 Custom observable

QuCumber has a built-in module called Observable which makes it easy for the user to compute any arbitrary observable from the RBM. To see the the Observable module in action, an example observable called PIQuIL, which inherits properties from the Observable module, is shown below.

The PIQuIL observable takes a σ^z measurement at a site and multiplies it by the measurement two sites away from it. There is also a parameter, P, that determines the strength of each of these interactions. For example, for the dataset (-1, 1, 1, -1), (1, 1, 1, 1) and (1, 1, -1, 1) with P = 2, the PIQuIL for each data point would be $(2(-1 \times 1) + 2(1 \times -1) = -4), (2(1 \times 1) + 2(1 \times 1) = 4)$ and $(2(1 \times -1) + 2(1 \times 1) = 0)$, respectively.

```
[23]: class PIQuIL (ObservableBase):
         def init (self, P):
              self.name = "PIOuIL"
              self.symbol = "Q"
              self.P = P
          # Required : function that calculates the PIQuIL. Must be named "apply"
          def apply(self, nn_state, samples):
              samples = to_pm1(samples)
              interaction_ = 0.0
              for i in range(samples.shape[-1]):
                  if (i + 3) > \text{samples.shape}[-1]:
                      continue
                  else:
                      interaction_ += self.P * samples[:, i] * samples[:, i + 2]
              return interaction_
     P = 0.05
     piquil = PIQuIL(P)
```

The apply function is contained in the Observable module, but is overwritten here. The apply function in Observable will compute the observable itself and must take in the RBM (nn_state) and a batch of samples as arguments. Thus, any new class inheriting from Observable that the user would like to define must contain a function called apply that calculates this new observable. For more details on apply, we refer to the documentation:

```
[24]: help(ObservableBase.apply)
```

```
Help on function apply in module qucumber.observables.observable:
apply(self, nn_state, samples)
   Computes the value of the observable, row-wise, on a batch of
   samples.
   If we think of the samples given as a set of projective measurements
   in a given computational basis, this method must return the expectation
   of the operator with respect to each basis state in `samples`.
   It must not perform any averaging for statistical purposes, as the
   proper analysis is delegated to the specialized
    `statistics` and `statistics_from_samples` methods.
   Must be implemented by any subclasses.
    :param nn_state: The WaveFunction that drew the samples.
    :type nn_state: qucumber.nn_states.WaveFunctionBase
    :param samples: A batch of sample states to calculate the observable on.
    :type samples: torch.Tensor
    :returns: The value of the observable of each given basis state.
    :rtype: torch.Tensor
```

Although the PIQuIL observable could technically be computed without the first argument of apply since it does not ever use the nn_state, we still include it in the list of arguments in order to conform to the interface provided in the ObservableBase class.

Since we have already generated new samples of data, the PIQuIL observable's mean, standard error and variance on the new data can be calculated with the statistics_from_samples function in the Observable module.

The user must simply provide the RBM and the samples as arguments.

```
[25]: piquil_stats1 = piquil.statistics_from_samples(nn_state, new_samples)
```

The statistics_from_samples function returns a dictionary containing the mean, standard error and the variance with the keys "mean", "std_error" and "variance", respectively.

Exercise: We notice that the PIQuIL observable is essentially a scaled next-nearest-neighbours interaction. (a) Construct an equivalent Observable object algebraically in a similar manner to the TFIM observable constructed above. (b) Compute the statistics of this observable on new_samples, and compare to those computed using the PIQuIL observable.

[27]: # solve the above exercise here

6.3 Estimating Statistics of Many Observables Simultaneously

One may often be concerned with estimating the statistics of many observables simultaneously. In order to avoid excess memory usage, it makes sense to reuse the same set of samples to estimate each observable. When we need a large number of samples however, we run into the same issue mentioned earlier: we may run out of memory storing the samples. QuCumber provides a System object to keep track of multiple observables and estimate their statistics efficiently.

```
[28]: from qucumber.observables import System
from pprint import pprint
```

At this point we must make a quick aside: internally, System keeps track of multiple observables through their name field (which we saw in the definition of the PIQuIL observable). This name is returned by Python's built-in repr function, which is automatically called when we try to display an Observable object in Jupyter:

```
[29]: piquil
[29]: PIQuIL
```

```
[30]: tfim
```

[30]: ((-1 * NeighbourInteraction(periodic_bcs=False, c=1)) + -(1 * SigmaX))

Note how the TFIM energy observable's name is quite complicated, due to the fact that we constructed it algebraically as opposed to the PIQuIL observable which was built from scratch and manually assigned a name. In order to assign a name to tfim, we do the following:

```
[31]: tfim.name = "TFIM"
tfim
```

```
[31]: TFIM
```

Now, back to System. We'd like to create a System object which keeps track of the absolute magnetization, the energy of the chain, the Swap observable (of region A, as defined earlier), and finally, the PIQuIL observable.

```
[32]: tfim_system = System(sz, tfim, swap, piquil)
```

[33]: pprint(tfim_system.statistics_from_samples(nn_state, new_samples))

These all match with the values computed earlier. Next, we will compute these statistics from fresh samples drawn from the RBM:

```
[34]: %%time
     pprint(
         tfim_system.statistics(
             nn_state, num_samples=10000, num_chains=1000, burn_in=100, steps=2
          )
      )
      {'PIQuIL': {'mean': 0.17418,
                  'std_error': 0.0015624906072324945,
                  'variance': 0.02441376897689769},
      'SWAP': {'mean': 0.7977556285445141,
                'std_error': 0.006037075970673986,
                'variance': 0.3644628627568925},
       'SigmaZ': {'mean': 0.55228,
                 'std_error': 0.0031312075316178864,
                 'variance': 0.09804460606060576},
       'TFIM': {'mean': -1.235210919302773,
                'std_error': 0.000462796110346464,
                'variance': 0.0021418023975181646}}
     CPU times: user 913 ms, sys: 0 ns, total: 913 ms
     Wall time: 232 ms
```

Compare this to computing these statistics on each observable individually:

```
'std_error': 0.005957763454588121,
    'variance': 0.3549494538082578},
'SigmaZ': {'mean': 0.557,
    'std_error': 0.003132424174360939,
    'variance': 0.098120812081208088},
'TFIM': {'mean': -1.2349736426229931,
    'std_error': 0.00047315568939076296,
    'variance': 0.0022387630640284817}}
CPU times: user 1.96 s, sys: 3.97 ms, total: 1.96 s
Wall time: 493 ms
```

Note the slowdown. This is, as mentioned before, due to the fact that the System object uses *the same samples* to estimate statistics for *all* of the observables it is keeping track of.

6.3.1 Template for your custom observable

Here is a generic template for you to try using the Observable module yourself.

```
[36]: import torch
     from qucumber.observables import ObservableBase
     class YourObservable(ObservableBase):
         def __init__(self, your_constants):
             self.your_constants = your_constants
             self.name = "Observable_Name"
              # The algebraic symbol representing this Observable.
              # Returned by Python's built-in str() function
             self.symbol = "O"
         def apply(self, nn_state, samples):
              # arguments of "apply" must be in this order
              # calculate your observable for each data point
             obs = torch.tensor([42] * len(samples))
              # make sure the observables are on the same device and have the
              # same dtype as the samples
             obs = obs.to(samples)
              # return a torch tensor containing the observable values
              return obs
```

SEVEN

TRAINING WHILE MONITORING OBSERVABLES

As seen in the first tutorial that went through reconstructing the wavefunction describing the TFIM with 10 sites at its critical point, the user can evaluate the training in real time with the MetricEvaluator and custom functions. What is most likely more impactful in many cases is to calculate an observable, like the energy, during the training process. This is slightly more computationally involved than using the MetricEvaluator to evaluate functions because observables require that samples be drawn from the RBM.

Luckily, QuCumber also has a module very similar to the MetricEvaluator, but for observables. This is called the ObservableEvaluator. This tutorial uses the ObservableEvaluator to calculate the energy during the training on the TFIM data in the first tutorial. We will use the same training hyperparameters as before.

It is assumed that the user has worked through Tutorial 3 beforehand. Recall that $quantum_ising_chain.py$ contains the TFIMChainEnergy class that inherits from the Observable module. The exact ground-state energy is -1.2381.

```
[1]: import os.path
    import numpy as np
    import matplotlib.pyplot as plt
    from qucumber.nn_states import PositiveWaveFunction
    from qucumber.callbacks import ObservableEvaluator
    import qucumber.utils.data as data
    from quantum_ising_chain import TFIMChainEnergy
[2]: train_data = data.load_data(
        os.path.join("..", "Tutoriall_TrainPosRealWaveFunction", "tfimld_data.txt")
    )[0](
    nv = train_data.shape[-1]
    nh = nv
    nn_state = PositiveWaveFunction(num_visible=nv, num_hidden=nh)
    epochs = 1000
    pbs = 100 # pos_batch_size
    nbs = 200 # neg_batch_size
    lr = 0.01
    k = 10
    period = 100
```

```
h = 1
num_samples = 10000
burn_in = 100
steps = 100
tfim_energy = TFIMChainEnergy(h)
```

Now, the ObservableEvaluator can be called. The ObservableEvaluator requires the following arguments.

- 1. period: the frequency of the training evaluators being calculated (e.g. period=200 means that the MetricEvaluator will compute the desired metrics every 200 epochs)
- 2. A list of Observable objects you would like to reference to evaluate the training (arguments required for generating samples to calculate the observables are keyword arguments placed after the list). The ObservableEvaluator uses a System object (discussed in the previous tutorial) under the hood in order to estimate statistics efficiently.

The following additional arguments are needed to calculate the statistics on the generated samples during training (these are the arguments of the statistics function in the Observable module, minus the nn_state argument; this gets passed in as an argument to fit). For more detail on these arguments, refer to either the previous tutorial or the documentation for Observable.statistics.

- num_samples: the number of samples to generate internally
- num_chains: the number of Markov chains to run in parallel (default = 0)
- burn_in: the number of Gibbs steps to perform before recording any samples (default = 1000)
- steps: the number of Gibbs steps to perform between each sample (default = 1)

The training evaluators can be printed out by setting the verbose keyword argument to True.

```
[3]: callbacks = [
        ObservableEvaluator(
             period,
             [tfim_energy],
             verbose=True,
             num samples=num samples,
             burn_in=burn_in,
             steps=steps,
         )
    1
    nn_state.fit(
        train_data,
        epochs=epochs,
        pos_batch_size=pbs,
        neg_batch_size=nbs,
        lr=lr,
         k=k,
         callbacks=callbacks,
    )
    Epoch: 100
      TFIMChainEnergy:
        mean: -1.193770
                             variance: 0.024622
                                                       std_error: 0.001569
    Epoch: 200
      TFIMChainEnergy:
```
(continued from previous page)

			1 10,
mean: -1.215802 Epoch: 300	variance: 0.013568	std_error: 0.001165	
TFIMChainEnergy: mean: -1.221930	variance: 0.009081	std_error: 0.000953	
Epoch: 400			
TFIMChainEnergy:			
mean: -1.227180	variance: 0.006347	std_error: 0.000797	
Epoch: 500			
TFIMChainEnergy:			
mean: -1.2300/4	variance: 0.004502	std_error: 0.0006/1	
Epoch: 600			
moon: -1 222001	warriance = 0.002641	atd arrar. 0 000602	
Epoch: 700	Vallance: 0.005041	sta_e1101. 0.000803	
TEIMChainEnergy			
mean: -1 233434	variance. 0 002839	std error, 0 000533	
Epoch: 800	Variance. 0.002033	<u>Sta_error</u> . 0.0000000	
TFIMChainEnergy:			
mean: -1.235324	variance: 0.002306	std error: 0.000480	
Epoch: 900			
TFIMChainEnergy:			
mean: -1.235313	variance: 0.001936	std_error: 0.000440	
Epoch: 1000			
TFIMChainEnergy:			
mean: -1.235257	variance: 0.001590	std_error: 0.000399	

The callbacks list returns a list of dictionaries. The mean, standard error and the variance at each epoch can be accessed as follows:

```
[4]: # Note that the name of the observable class that the user makes
    # must be what comes after callbacks[0].
    energies = callbacks[0].TFIMChainEnergy.mean
    # Alternatively, we can use the usual dictionary/list subscripting
    # syntax, which is useful in the case where the observable's name
    # contains special characters
    errors = callbacks[0]["TFIMChainEnergy"].std_error
    variance = callbacks[0]["TFIMChainEnergy"]["variance"]
```

A plot of the energy as a function of the training cycle is presented below.

```
[5]: epoch = np.arange(period, epochs + 1, period)
E0 = -1.2381
plt.figure(figsize=(10, 5))
ax = plt.axes()
ax.plot(epoch, energies, color="red")
ax.set_xlim(period, epochs)
ax.axhline(E0, color="black")
ax.fill_between(epoch, energies - errors, energies + errors, alpha=0.2, color="black")
ax.set_xlabel("Epoch")
ax.set_ylabel("Energy")
ax.grid()
```



EIGHT

RBM

class qucumber.rbm.BinaryRBM(num_visible, num_hidden, zero_weights=False, gpu=True)
Bases: torch.nn.Module

effective_energy(v)

The effective energies of the given visible states.

$$\mathcal{E}(v) = -\sum_{j} b_{j} v_{j} - \sum_{i} \log \left[1 + \exp\left(c_{i} + \sum_{j} W_{ij} v_{j}\right) \right]$$

Parameters v (*torch*. *Tensor*) – The visible states.

Returns The effective energies of the given visible states.

Return type torch.Tensor

effective_energy_gradient(v, reduce=True)

The gradients of the effective energies for the given visible states.

Parameters

- **v** (torch. Tensor) The visible states.
- **reduce** If *True*, will sum over the gradients resulting from each visible state. Otherwise will return a batch of gradient vectors.
- **Returns** Will return a vector (or matrix if *reduce=False* and multiple visible states were given as a matrix) containing the gradients for all parameters (computed on the given visible states v).

Return type torch.Tensor

gibbs_steps (k, initial_state, overwrite=False)

Performs k steps of Block Gibbs sampling. One step consists of sampling the hidden state h from the conditional distribution p(h | v), and sampling the visible state v from the conditional distribution p(v | h).

Parameters

- k (int) Number of Block Gibbs steps.
- initial_state (torch.Tensor) The initial state of the Markov Chains.
- **overwrite** (bool) Whether to overwrite the initial_state tensor, if it is provided.

initialize_parameters (zero_weights=False)

Randomize the parameters of the RBM

partition (space)

Compute the partition function of the RBM.

Parameters space (torch.Tensor) – A rank 2 tensor of the visible space.

Returns The value of the partition function evaluated at the current state of the RBM.

Return type torch.Tensor

prob_h_given_v(v, out=None)

Given a visible unit configuration, compute the probability vector of the hidden units being on.

Parameters

- h (torch. Tensor) The hidden unit.
- **out** (*torch*. *Tensor*) The output tensor to write to.

Returns The probability of hidden units being active given the visible state.

Return type torch.Tensor

prob_v_given_h(h, out=None)

Given a hidden unit configuration, compute the probability vector of the visible units being on.

Parameters

- h (torch. Tensor) The hidden unit
- out (torch. Tensor) The output tensor to write to.

Returns The probability of visible units being active given the hidden state.

Return type torch.Tensor

sample_h_given_v(v, out=None)

Sample/generate a hidden state given a visible state.

Parameters

- **h** (torch. Tensor) The visible state.
- out (torch. Tensor) The output tensor to write to.

Returns The sampled hidden state.

Return type torch.Tensor

sample_v_given_h(h, out=None)

Sample/generate a visible state given a hidden state.

Parameters

- **h** (torch. Tensor) The hidden state.
- **out** (*torch*. *Tensor*) The output tensor to write to.

Returns The sampled visible state.

CHAPTER

NINE

QUANTUM STATES

9.1 Positive WaveFunction

Bases: qucumber.nn_states.WaveFunctionBase

Class capable of learning wavefunctions with no phase.

Parameters

- **num_visible** (*int*) The number of visible units, ie. the size of the system being learned.
- num_hidden (*int*) The number of hidden units in the internal RBM. Defaults to the number of visible units.
- gpu (bool) Whether to perform computations on the default GPU.
- **module** (qucumber.rbm.BinaryRBM) An instance of a BinaryRBM module to use for density estimation. Will be copied to the default GPU if *gpu=True* (if it isn't already there). If *None*, will initialize a BinaryRBM from scratch.

amplitude(v)

Compute the (unnormalized) amplitude of a given vector/matrix of visible states.

amplitude(σ) = $|\psi_{\lambda}(\sigma)| = e^{-\mathcal{E}_{\lambda}(\sigma)/2}$

Parameters \mathbf{v} (torch. Tensor) – visible states σ

Returns Matrix/vector containing the amplitudes of v

Return type torch.Tensor

static autoload(location, gpu=False)

Initializes a WaveFunction from the parameters in the given location.

Parameters

- location (*str or file*) The location to load the model parameters from.
- **gpu** (*bool*) Whether the returned model should be on the GPU.

Returns A new WaveFunction initialized from the given parameters. The returned WaveFunction will be of whichever type this function was called on.

compute_batch_gradients(k, samples_batch, neg_batch)

Compute the gradients of a batch of the training data (samples_batch).

- **k** (*int*) Number of contrastive divergence steps in training.
- **samples_batch** (*torch*. *Tensor*) Batch of the input samples.
- **neg_batch** (*torch.Tensor*) Batch of the input samples for computing the negative phase.

Returns List containing the gradients of the parameters.

Return type list

compute_normalization(space)

Compute the normalization constant of the wavefunction.

$$Z_{\lambda} = \sqrt{\sum_{\sigma} |\psi_{\lambda}|^2} = \sqrt{\sum_{\sigma} p_{\lambda}(\sigma)}$$

Parameters space (torch. Tensor) - A rank 2 tensor of the entire visible space.

property device

The device that the model is on.

fit (data, epochs=100, pos_batch_size=100, neg_batch_size=None, k=1, lr=0.001, progbar=False, starting_epoch=1, time=False, callbacks=None, optimizer=torch.optim.SGD, **kwargs) Train the WaveFunction.

Parameters

- data (np.array) The training samples
- **epochs** (*int*) The number of full training passes through the dataset. Technically, this specifies the index of the *last* training epoch, which is relevant if *starting_epoch* is being set.
- **pos_batch_size** (*int*) The size of batches for the positive phase taken from the data.
- **neg_batch_size** (*int*) The size of batches for the negative phase taken from the data. Defaults to *pos_batch_size*.
- **k** (*int*) The number of contrastive divergence steps.
- **lr** (*float*) Learning rate
- **progbar** (*bool or str*) Whether or not to display a progress bar. If "notebook" is passed, will use a Jupyter notebook compatible progress bar.
- **starting_epoch** (*int*) The epoch to start from. Useful if continuing training from a previous state.
- **callbacks** (*list* [qucumber.callbacks.CallbackBase]) Callbacks to run while training.
- optimizer (torch.optim.Optimizer) The constructor of a torch optimizer.
- kwargs Keyword arguments to pass to the optimizer

generate_hilbert_space (size=None, device=None)

Generates Hilbert space of dimension 2^{size} .

Parameters

• **size** (*int*) – The size of each element of the Hilbert space. Defaults to the number of visible units.

• **device** – The device to create the Hilbert space matrix on. Defaults to the device this model is on.

Returns A tensor with all the basis states of the Hilbert space.

Return type torch.Tensor

gradient (v)

Compute the gradient of the effective energy for a batch of states.

 $\nabla_{\lambda} \mathcal{E}_{\lambda}(\sigma)$

Parameters \mathbf{v} (torch.Tensor) – visible states σ

Returns A single tensor containing all of the parameter gradients.

Return type torch.Tensor

load(location)

Loads the WaveFunction parameters from the given location ignoring any metadata stored in the file. Overwrites the WaveFunction's parameters.

Note: The WaveFunction object on which this function is called must have the same parameter shapes as the one who's parameters are being loaded.

Parameters location (*str or file*) – The location to load the WaveFunction parameters from.

property max_size

Maximum size of the Hilbert space for full enumeration

property networks

A list of the names of the internal RBMs.

phase(v)

Compute the phase of a given vector/matrix of visible states.

In the case of a PositiveWaveFunction, the phase is just zero.

Parameters \mathbf{v} (torch.Tensor) – visible states σ

Returns Matrix/vector containing the phases of v

Return type torch.Tensor

probability (v, Z)

Evaluates the probability of the given vector(s) of visible states.

Parameters

- **v** (torch. Tensor) The visible states.
- **Z** (*float*) The partition function.

Returns The probability of the given vector(s) of visible units.

Return type torch.Tensor

psi(v)

Compute the (unnormalized) wavefunction of a given vector/matrix of visible states.

$$\psi_{\lambda}(\sigma) = e^{-\mathcal{E}_{\lambda}(\sigma)/2}$$

Parameters \mathbf{v} (torch. Tensor) – visible states σ

Returns Complex object containing the value of the wavefunction for each visible state

Return type torch.Tensor

property rbm_am

The RBM to be used to learn the wavefunction amplitude.

reinitialize_parameters()

Randomize the parameters of the internal RBMs.

sample (k, num_samples=1, initial_state=None, overwrite=False)

Performs k steps of Block Gibbs sampling. One step consists of sampling the hidden state h from the conditional distribution $p_{\lambda}(h|v)$, and sampling the visible state v from the conditional distribution $p_{\lambda}(v|h)$.

Parameters

- **k** (*int*) Number of Block Gibbs steps.
- num_samples (*int*) The number of samples to generate.
- initial_state (torch.Tensor) The initial state of the Markov Chains. If given, num_samples will be ignored.
- **overwrite** (*bool*) Whether to overwrite the initial_state tensor, if it is provided.

save (location, metadata=None)

Saves the WaveFunction parameters to the given location along with any given metadata.

Parameters

- **location** (*str or file*) The location to save the data.
- **metadata** (*dict*) Any extra metadata to store alongside the WaveFunction parameters.

property stop_training

If True, will not train.

If this property is set to *True* during the training cycle, training will terminate once the current batch or epoch ends (depending on when *stop_training* was set).

subspace_vector (num, size=None, device=None)

Generates a single vector from the Hilbert space of dimension 2^{size}.

Parameters

- **size** (*int*) The size of each element of the Hilbert space.
- **num** (*int*) The specific vector to return from the Hilbert space. Since the Hilbert space can be represented by the set of binary strings of length *size*, *num* is equivalent to the decimal representation of the returned vector.
- device The device to create the vector on. Defaults to the device this model is on.

Returns A state from the Hilbert space.

9.2 Complex WaveFunction

```
class qucumber.nn_states.ComplexWaveFunction (num_visible, num_hidden=None, uni-
tary_dict=None, gpu=True, mod-
ule=None)
```

Bases: qucumber.nn_states.WaveFunctionBase

Class capable of learning wavefunctions with a non-zero phase.

Parameters

- **num_visible** (*int*) The number of visible units, ie. the size of the system being learned.
- **num_hidden** (*int*) The number of hidden units in both internal RBMs. Defaults to the number of visible units.
- unitary_dict (dict[str, torch.Tensor]) A dictionary mapping unitary names to their matrix representations.
- gpu (bool) Whether to perform computations on the default GPU.
- **module** (qucumber.rbm.BinaryRBM) An instance of a BinaryRBM module to use for density estimation; The given RBM object will be used to estimate the amplitude of the wavefunction, while a copy will be used to estimate the phase of the wavefunction. Will be copied to the default GPU if *gpu=True* (if it isn't already there). If *None*, will initialize the BinaryRBMs from scratch.

amplitude(v)

Compute the (unnormalized) amplitude of a given vector/matrix of visible states.

amplitude(σ) = $|\psi_{\lambda\mu}(\sigma)| = e^{-\mathcal{E}_{\lambda}(\sigma)/2}$

Parameters \mathbf{v} (torch. Tensor) – visible states σ .

Returns Vector containing the amplitudes of the given states.

Return type torch.Tensor

static autoload(location, gpu=False)

Initializes a WaveFunction from the parameters in the given location.

Parameters

- location (*str or file*) The location to load the model parameters from.
- **gpu** (bool) Whether the returned model should be on the GPU.

Returns A new WaveFunction initialized from the given parameters. The returned WaveFunction will be of whichever type this function was called on.

compute_batch_gradients (k, samples_batch, neg_batch, bases_batch=None)

Compute the gradients of a batch of the training data (*samples_batch*).

If measurements are taken in bases other than the reference basis, a list of bases (*bases_batch*) must also be provided.

- **k** (*int*) Number of contrastive divergence steps in training.
- **samples_batch** (*torch.Tensor*) Batch of the input samples.
- **neg_batch** (*torch.Tensor*) Batch of the input samples for computing the negative phase.

• **bases_batch** (*np.array*) – Batch of the input bases corresponding to the samples in *samples_batch*.

Returns List containing the gradients of the parameters.

Return type list

compute_normalization(space)

Compute the normalization constant of the wavefunction.

$$Z_{\lambda} = \sqrt{\sum_{\sigma} |\psi_{\lambda\mu}|^2} = \sqrt{\sum_{\sigma} p_{\lambda}(\sigma)}$$

Parameters space (torch. Tensor) – A rank 2 tensor of the entire visible space.

property device

The device that the model is on.

fit (data, epochs=100, pos_batch_size=100, neg_batch_size=None, k=1, lr=0.001, input_bases=None, progbar=False, starting_epoch=1, time=False, callbacks=None, optimizer=torch.optim.SGD, **kwargs)

Train the WaveFunction.

Parameters

- data (np.array) The training samples
- **epochs** (*int*) The number of full training passes through the dataset. Technically, this specifies the index of the *last* training epoch, which is relevant if *starting_epoch* is being set.
- **pos_batch_size** (*int*) The size of batches for the positive phase taken from the data.
- **neg_batch_size** (*int*) The size of batches for the negative phase taken from the data. Defaults to *pos_batch_size*.
- **k** (*int*) The number of contrastive divergence steps.
- lr (float) Learning rate
- **input_bases** (*np.array*) The measurement bases for each sample. Must be provided if training a ComplexWaveFunction.
- **progbar** (*bool* or *str*) Whether or not to display a progress bar. If "notebook" is passed, will use a Jupyter notebook compatible progress bar.
- **starting_epoch** (*int*) The epoch to start from. Useful if continuing training from a previous state.
- **callbacks** (*list* [qucumber.callbacks.CallbackBase]) Callbacks to run while training.
- **optimizer** (torch.optim.Optimizer) The constructor of a torch optimizer.
- kwargs Keyword arguments to pass to the optimizer

generate_hilbert_space (size=None, device=None)

Generates Hilbert space of dimension 2^{size} .

Parameters

• **size** (*int*) – The size of each element of the Hilbert space. Defaults to the number of visible units.

• **device** – The device to create the Hilbert space matrix on. Defaults to the device this model is on.

Returns A tensor with all the basis states of the Hilbert space.

Return type torch.Tensor

gradient (basis, sample)

Compute the gradient of a sample, measured in different bases.

Parameters

- **basis** (*np.array*) A set of bases.
- **sample** (*np.array*) A sample to compute the gradient of.

Returns A list of 2 tensors containing the parameters of each of the internal RBMs.

Return type list[torch.Tensor]

load(location)

Loads the WaveFunction parameters from the given location ignoring any metadata stored in the file. Overwrites the WaveFunction's parameters.

Note: The WaveFunction object on which this function is called must have the same parameter shapes as the one who's parameters are being loaded.

Parameters location (*str or file*) – The location to load the WaveFunction parameters from.

property max_size

Maximum size of the Hilbert space for full enumeration

property networks

A list of the names of the internal RBMs.

phase(v)

Compute the phase of a given vector/matrix of visible states.

phase(
$$\sigma$$
) = $-\mathcal{E}_{\mu}(\sigma)/2$

Parameters \mathbf{v} (torch. Tensor) – visible states σ .

Returns Vector containing the phases of the given states.

Return type torch.Tensor

probability (v, Z)

Evaluates the probability of the given vector(s) of visible states.

Parameters

- **v** (torch. Tensor) The visible states.
- **Z** (*float*) The partition function.

Returns The probability of the given vector(s) of visible units.

psi(v)

Compute the (unnormalized) wavefunction of a given vector/matrix of visible states.

 $\psi_{\lambda\mu}(\sigma) = e^{-[\mathcal{E}_{\lambda}(\sigma) + i\mathcal{E}_{\mu}(\sigma)]/2}$

Parameters \mathbf{v} (torch. Tensor) – visible states σ

Returns Complex object containing the value of the wavefunction for each visible state

Return type torch.Tensor

property rbm_am

The RBM to be used to learn the wavefunction amplitude.

property rbm_ph

RBM used to learn the wavefunction phase.

reinitialize_parameters()

Randomize the parameters of the internal RBMs.

sample (k, num_samples=1, initial_state=None, overwrite=False)

Performs k steps of Block Gibbs sampling. One step consists of sampling the hidden state h from the conditional distribution $p_{\lambda}(h|v)$, and sampling the visible state v from the conditional distribution $p_{\lambda}(v|h)$.

Parameters

- **k** (*int*) Number of Block Gibbs steps.
- **num_samples** (*int*) The number of samples to generate.
- initial_state (torch.Tensor) The initial state of the Markov Chains. If given, *num_samples* will be ignored.
- **overwrite** (bool) Whether to overwrite the initial_state tensor, if it is provided.

save (location, metadata=None)

Saves the WaveFunction parameters to the given location along with any given metadata.

Parameters

- **location** (*str or file*) The location to save the data.
- **metadata** (*dict*) Any extra metadata to store alongside the WaveFunction parameters.

property stop_training

If True, will not train.

If this property is set to *True* during the training cycle, training will terminate once the current batch or epoch ends (depending on when *stop_training* was set).

subspace_vector (num, size=None, device=None)

Generates a single vector from the Hilbert space of dimension 2^{size} .

Parameters

- **size** (*int*) The size of each element of the Hilbert space.
- **num** (*int*) The specific vector to return from the Hilbert space. Since the Hilbert space can be represented by the set of binary strings of length *size*, *num* is equivalent to the decimal representation of the returned vector.
- device The device to create the vector on. Defaults to the device this model is on.

Returns A state from the Hilbert space.

9.3 Abstract WaveFunction

Note: This is an Abstract Base Class, it is not meant to be used directly. The following API reference is mostly for developers.

class qucumber.nn_states.WaveFunctionBase

Bases: abc.ABC

Abstract Base Class for WaveFunctions.

amplitude(v)

Compute the (unnormalized) amplitude of a given vector/matrix of visible states.

amplitude(σ) = $|\psi(\sigma)|$

Parameters \mathbf{v} (torch.Tensor) – visible states σ

Returns Matrix/vector containing the amplitudes of v

Return type torch.Tensor

abstract static autoload(*location*, *gpu=False*)

Initializes a WaveFunction from the parameters in the given location.

Parameters

- location (*str or file*) The location to load the model parameters from.
- **gpu** (*bool*) Whether the returned model should be on the GPU.
- **Returns** A new WaveFunction initialized from the given parameters. The returned WaveFunction will be of whichever type this function was called on.

compute_batch_gradients (k, samples_batch, neg_batch, bases_batch=None)

Compute the gradients of a batch of the training data (*samples_batch*).

If measurements are taken in bases other than the reference basis, a list of bases (*bases_batch*) must also be provided.

Parameters

- **k** (*int*) Number of contrastive divergence steps in training.
- **samples_batch** (*torch.Tensor*) Batch of the input samples.
- **neg_batch** (*torch.Tensor*) Batch of the input samples for computing the negative phase.
- **bases_batch** (*np.array*) Batch of the input bases corresponding to the samples in *samples_batch*.

Returns List containing the gradients of the parameters.

Return type list

compute_normalization(space)

Compute the normalization constant of the wavefunction.

$$Z_{\lambda} = \sqrt{\sum_{\sigma} |\psi_{\lambda\mu}|^2} = \sqrt{\sum_{\sigma} p_{\lambda}(\sigma)}$$

Parameters space (torch. Tensor) – A rank 2 tensor of the entire visible space.

abstract property device

The device that the model is on.

fit (data, epochs=100, pos_batch_size=100, neg_batch_size=None, k=1, lr=0.001, input_bases=None, progbar=False, starting_epoch=1, time=False, callbacks=None, optimizer=torch.optim.SGD, **kwargs) Train the WaveFunction.

Parameters

- data (np.array) The training samples
- **epochs** (*int*) The number of full training passes through the dataset. Technically, this specifies the index of the *last* training epoch, which is relevant if *starting_epoch* is being set.
- **pos_batch_size** (*int*) The size of batches for the positive phase taken from the data.
- **neg_batch_size** (*int*) The size of batches for the negative phase taken from the data. Defaults to *pos_batch_size*.
- **k** (*int*) The number of contrastive divergence steps.
- **lr** (*float*) Learning rate
- **input_bases** (*np.array*) The measurement bases for each sample. Must be provided if training a ComplexWaveFunction.
- **progbar** (*bool or str*) Whether or not to display a progress bar. If "notebook" is passed, will use a Jupyter notebook compatible progress bar.
- **starting_epoch** (*int*) The epoch to start from. Useful if continuing training from a previous state.
- **callbacks** (*list* [qucumber.callbacks.CallbackBase]) Callbacks to run while training.
- **optimizer** (torch.optim.Optimizer) The constructor of a torch optimizer.
- **kwargs** Keyword arguments to pass to the optimizer

generate_hilbert_space (size=None, device=None)

Generates Hilbert space of dimension 2^{size}.

Parameters

- **size** (*int*) The size of each element of the Hilbert space. Defaults to the number of visible units.
- **device** The device to create the Hilbert space matrix on. Defaults to the device this model is on.

Returns A tensor with all the basis states of the Hilbert space.

Return type torch.Tensor

abstract gradient()

Compute the gradient of a set of samples.

load (location)

Loads the WaveFunction parameters from the given location ignoring any metadata stored in the file. Overwrites the WaveFunction's parameters.

Note: The WaveFunction object on which this function is called must have the same parameter shapes as the one who's parameters are being loaded.

Parameters location (*str or file*) – The location to load the WaveFunction parameters from.

property max_size

Maximum size of the Hilbert space for full enumeration

abstract property networks

A list of the names of the internal RBMs.

abstract phase(v)

Compute the phase of a given vector/matrix of visible states.

phase(σ)

Parameters \mathbf{v} (torch. Tensor) – visible states σ

Returns Matrix/vector containing the phases of v

Return type torch.Tensor

probability (v, Z)

Evaluates the probability of the given vector(s) of visible states.

Parameters

- **v** (torch. Tensor) The visible states.
- **Z** (*float*) The partition function.

Returns The probability of the given vector(s) of visible units.

Return type torch.Tensor

abstract psi(v)

Compute the (unnormalized) wavefunction of a given vector/matrix of visible states.

 $\psi(\sigma)$

Parameters \mathbf{v} (torch. Tensor) – visible states σ

Returns Complex object containing the value of the wavefunction for each visible state

Return type torch.Tensor

abstract property rbm_am

The RBM to be used to learn the wavefunction amplitude.

reinitialize_parameters()

Randomize the parameters of the internal RBMs.

sample (k, num_samples=1, initial_state=None, overwrite=False)

Performs k steps of Block Gibbs sampling. One step consists of sampling the hidden state h from the conditional distribution $p_{\lambda}(h|v)$, and sampling the visible state v from the conditional distribution $p_{\lambda}(v|h)$.

- k (int) Number of Block Gibbs steps.
- num_samples (*int*) The number of samples to generate.

- initial_state (torch.Tensor) The initial state of the Markov Chains. If given, *num_samples* will be ignored.
- **overwrite** (bool) Whether to overwrite the initial_state tensor, if it is provided.

save (location, metadata=None)

Saves the WaveFunction parameters to the given location along with any given metadata.

Parameters

- **location** (*str or file*) The location to save the data.
- **metadata** (*dict*) Any extra metadata to store alongside the WaveFunction parameters.

property stop_training

If *True*, will not train.

If this property is set to *True* during the training cycle, training will terminate once the current batch or epoch ends (depending on when *stop_training* was set).

subspace_vector (num, size=None, device=None)

Generates a single vector from the Hilbert space of dimension 2^{size} .

Parameters

- **size** (*int*) The size of each element of the Hilbert space.
- **num** (*int*) The specific vector to return from the Hilbert space. Since the Hilbert space can be represented by the set of binary strings of length *size*, *num* is equivalent to the decimal representation of the returned vector.
- device The device to create the vector on. Defaults to the device this model is on.

Returns A state from the Hilbert space.

CHAPTER

TEN

CALLBACKS

class qucumber.callbacks.**CallbackBase** Base class for callbacks.

on_batch_end (*nn_state*, *epoch*, *batch*) Called at the end of each batch.

Parameters

- nn_state (qucumber.nn_states.WaveFunctionBase) The WaveFunction being trained.
- **epoch** (*int*) The current epoch.
- **batch** (*int*) The current batch index.

on_batch_start (*nn_state*, *epoch*, *batch*) Called at the start of each batch.

Parameters

- nn_state (qucumber.nn_states.WaveFunctionBase) The WaveFunction being trained.
- **epoch** (*int*) The current epoch.
- **batch** (*int*) The current batch index.

on_epoch_end (nn_state, epoch)

Called at the end of each epoch.

Parameters

- nn_state (qucumber.nn_states.WaveFunctionBase) The WaveFunction being trained.
- **epoch** (*int*) The current epoch.

on_epoch_start (*nn_state*, *epoch*) Called at the start of each epoch.

Parameters

- nn_state (qucumber.nn_states.WaveFunctionBase) The WaveFunction being trained.
- **epoch** (*int*) The current epoch.

on_train_end(nn_state)

Called at the end of the training cycle.

- Parameters nn_state (qucumber.nn_states.WaveFunctionBase) The Wave-Function being trained.
- on_train_start (nn_state)

Called at the start of the training cycle.

Parameters nn_state (qucumber.nn_states.WaveFunctionBase) - The Wave-Function being trained.

Class for creating simple callbacks.

This callback is constructed using the passed functions that will be called at the appropriate time.

Parameters

- on_train_start (callable or None) A function to be called at the start of the training cycle. Must follow the same signature as CallbackBase.on_train_start.
- **on_train_end**(*callable or None*) A function to be called at the end of the training cycle. Must follow the same signature as *CallbackBase.on_train_end*.
- on_epoch_start (callable or None) A function to be called at the start of every epoch. Must follow the same signature as CallbackBase.on_epoch_start.
- on_epoch_end (callable or None) A function to be called at the end of every epoch. Must follow the same signature as CallbackBase.on_epoch_end.
- on_batch_start (callable or None) A function to be called at the start of every batch. Must follow the same signature as CallbackBase.on_batch_start.
- on_batch_end (callable or None) A function to be called at the end of every batch. Must follow the same signature as CallbackBase.on_batch_end.

class qucumber.callbacks.**ModelSaver** (*period*, *folder_path*, *file_name*, *save_initial=True*, *meta-data=None*, *metadata* only=False)

Callback which allows model parameters (along with some metadata) to be saved to disk at regular intervals.

This callback is called at the end of each epoch. If *save_initial* is *True*, will also be called at the start of the training cycle.

Parameters

- **period** (*int*) Frequency of model saving (in epochs).
- **folder_path** (*str*) The directory in which to save the files
- **file_name** (*str*) The name of the output files. Should be a format string with one blank, which will be filled with either the epoch number or the word "initial".
- **save_initial** (bool) Whether to save the initial parameters (and metadata).
- **metadata** (*callable or dict or None*) The metadata to save to disk with the model parameters Can be either a function or a dictionary. In the case of a function, it must take 2 arguments the RBM being trained, and the current epoch number, and then return a dictionary containing the metadata to be saved.
- **metadata_only** (bool) Whether to save only the metadata to disk.

Callback which logs output at regular intervals.

This callback is called at the end of each epoch.

Parameters

- **period** (*int*) Logging frequency (in epochs).
- **logger_fn** (*callable*) The function used for logging. Must take 1 string as an argument. Defaults to the standard *print* function.
- msg_gen (callable) A callable which generates the string to be logged. Must take 2 positional arguments: the RBM being trained and the current epoch. It must also be able to take some keyword arguments.
- ****kwargs** Keyword arguments which will be passed to *msg_gen*.

Stop training once the model stops improving.

There are three different stopping criteria available:

relative, which computes the relative change between the two model evaluation steps:

$$\left|\frac{M_{t-p} - M_t}{M_{t-p}}\right| < \epsilon$$

absolute computes the absolute change:

$$|M_{t-p} - M_t| < \epsilon$$

variance computes the absolute change, but scales the change by the standard deviation of the quantity of interest, such that the tolerance, *epsilon* can now be interpreted as the "number of standard deviations":

$$\left|\frac{M_{t-p} - M_t}{\sigma_{t-p}}\right| < \epsilon$$

where M_t is the metric value at the current evaluation (time t), p is the "patience" parameter, σ_t is the standard deviation of the metric, and ϵ is the tolerance.

This callback is called at the end of each epoch.

Parameters

- **period** (*int*) Frequency with which the callback checks whether training has converged (in epochs).
- **tolerance** (*float*) The maximum relative change required to consider training as having converged.
- **patience** (*int*) How many intervals to wait before claiming the training has converged.
- **evaluator_callback** (*MetricEvaluator* or *ObservableEvaluator*) An instance of *MetricEvaluator* or *ObservableEvaluator* which computes the metric that we want to check for convergence.
- **quantity_name** (*str*) The name of the metric/observable stored in *evalua-tor_callback*.
- **criterion** (*str*) The stopping criterion to use. Must be one of the following: *relative*, *absolute*, *variance*.

class qucumber.callbacks.VarianceBasedEarlyStopping(period, tolerance, patience, evaluator_callback, quantity_name, variance name=None)

Deprecated since version 1.2: Use *EarlyStopping* instead.

Stop training once the model stops improving. This is a variation on the *EarlyStopping* class which takes the variance of the metric into account.

The specific criterion for stopping is:

$$\left|\frac{M_{t-p} - M_t}{\sigma_{t-p}}\right| < \kappa$$

where M_t is the metric value at the current evaluation (time t), p is the "patience" parameter, σ_t is the standard deviation of the metric, and κ is the tolerance.

This callback is called at the end of each epoch.

Parameters

- **period** (*int*) Frequency with which the callback checks whether training has converged (in epochs).
- **tolerance** (*float*) The maximum (standardized) change required to consider training as having converged.
- **patience** (*int*) How many intervals to wait before claiming the training has converged.
- **evaluator_callback** (*MetricEvaluator* or *ObservableEvaluator*) An instance of *MetricEvaluator* or *ObservableEvaluator* which computes the metric/observable that we want to check for convergence.
- quantity_name (str) The name of the metric/obserable stored in evaluator_callback.
- **variance_name** (*str*) The name of the variance stored in *evaluator_callback*. Ignored, exists for backward compatibility.

Evaluate and hold on to the results of the given metric(s).

This callback is called at the end of each epoch.

Note: Since callbacks are given to *fit* as a list, they will be called in a deterministic order. It is therefore recommended that instances of *MetricEvaluator* be among the first callbacks in the list passed to *fit*, as one would often use it in conjunction with other callbacks like *EarlyStopping* which may depend on *MetricEvaluator* having been called.

Parameters

- **period** (*int*) Frequency with which the callback evaluates the given metric(s).
- **metrics** (*dict* (*str*, *callable*)) A dictionary of callables where the keys are the names of the metrics and the callables take the WaveFunction being trained as their positional argument, along with some keyword arguments. The metrics are evaluated and put into an internal dictionary structure resembling the structure of *metrics*.
- **verbose** (*bool*) Whether to print metrics to stdout.
- log (*str*) A filepath to log metric values to in CSV format.
- ****metric_kwargs** Keyword arguments to be passed to *metrics*.

```
___getattr__(metric)
```

Return an array of all recorded values of the given metric.

Parameters metric (*str*) – The metric to retrieve.

Returns The past values of the metric.

Return type np.array

___getitem___(metric)

Alias for <u>______</u> getattr___ to enable subscripting.

len()

Return the number of timesteps that metrics have been evaluated for.

Return type int

clear_history()

Delete all metric values the instance is currently storing.

property epochs

Return a list of all epochs that have been recorded.

Return type np.array

get_value (name, index=None)

Retrieve the value of the desired metric from the given timestep.

Parameters

- **name** (*str*) The name of the metric to retrieve.
- **index** (*int* or *None*) The index/timestep from which to retrieve the metric. Negative indices are supported. If None, will just get the most recent value.

property names

The names of the tracked metrics.

Return type list[str]

class qucumber.callbacks.ObservableEvaluator(period, observables, verbose=False,

log=None, **sampling_kwargs)

Evaluate and hold on to the results of the given observable(s).

This callback is called at the end of each epoch.

Note: Since callback are given to *fit* as a list, they will be called in a deterministic order. It is therefore recommended that instances of *ObservableEvaluator* be among the first callbacks in the list passed to *fit*, as one would often use it in conjunction with other callbacks like *EarlyStopping* which may depend on *ObservableEvaluator* having been called.

- **period** (*int*) Frequency with which the callback evaluates the given observables(s).
- **observables** (*list* (qucumber.observables.ObservableBase)) A list of Observables. Observable statistics are evaluated by sampling the WaveFunction. Note that observables that have the same name will conflict, and precedence will be given to the one which appears later in the list.
- **verbose** (*bool*) Whether to print metrics to stdout.
- log (*str*) A filepath to log metric values to in CSV format.
- ****sampling_kwargs** Keyword arguments to be passed to *Observable.statistics*. Ex. *num_samples, num_chains, burn_in, steps*.

_getattr__(observable)

Return an ObservableStatistics containing recorded statistics of the given observable.

Parameters observable (*str*) – The observable to retrieve.

Returns The past values of the observable.

Return type ObservableStatistics

___getitem___(observable)

Alias for <u>______</u> getattr___ to enable subscripting.

__len_()

Return the number of timesteps that observables have been evaluated for.

Return type int

clear_history()

Delete all statistics the instance is currently storing.

property epochs

Return a list of all epochs that have been recorded.

Return type np.array

get_value (name, index=None)

Retrieve the statistics of the desired observable from the given timestep.

Parameters

- **name** (*str*) The name of the observable to retrieve.
- **index** (*int or None*) The index/timestep from which to retrieve the observable. Negative indices are supported. If None, will just get the most recent value.

Return type dict(str, float)

property names

The names of the tracked observables.

Return type list[str]

class qucumber.callbacks.LivePlotting(period, evaluator_callback, quantity_name, er-

ror_name=None, total_epochs=None, smooth=True)

Plots metrics/observables.

This callback is called at the end of each epoch.

Parameters

- **period** (*int*) Frequency with which the callback updates the plots (in epochs).
- **evaluator_callback** (*MetricEvaluator* or *ObservableEvaluator*) An instance of *MetricEvaluator* or *ObservableEvaluator* which computes the metric/observable that we want to plot.
- **quantity_name** (*str*) The name of the metric/observable stored in *evalua-tor_callback*.
- **error_name** (*str*) The name of the error stored in *evaluator_callback*.

class qucumber.callbacks.Timer(verbose=True)

Callback which records the training time.

This callback is always called at the start and end of training. It will run at the end of an epoch or batch if the given model's *stop_training* property is set to True.

Parameters verbose (bool) – Whether to print the elapsed time at the end of training.

CHAPTER

ELEVEN

OBSERVABLES

11.1 Pauli Operators

class qucumber.observables.**SigmaZ** (*absolute=False*) Bases: qucumber.observables.ObservableBase

The σ_z observable.

Computes the magnetization in the Z direction of a spin chain.

Parameters absolute (bool) – Specifies whether to estimate the absolute magnetization.

apply (*nn_state*, *samples*)

Computes the magnetization along Z of each sample given a batch of samples.

Assumes that the computational basis that the WaveFunction was trained on was the Z basis.

Parameters

- **nn_state** (qucumber.nn_states.WaveFunctionBase) The WaveFunction that drew the samples.
- samples (torch.Tensor) A batch of samples to calculate the observable on. Must be using the $\sigma_i = 0, 1$ convention.

property name

The name of the Observable.

sample (nn_state, k, num_samples=1, initial_state=None, overwrite=False)
Draws samples of the observable using the given WaveFunction.

Parameters

- nn_state (qucumber.nn_states.WaveFunctionBase) The WaveFunction to draw samples from.
- k (int) The number of Gibbs Steps to perform before drawing a sample.
- num_samples (*int*) The number of samples to draw.
- initial_state (torch.Tensor) The initial state of the Markov Chain. If given, *num_samples* will be ignored.
- **overwrite** (*bool*) Whether to overwrite the initial_state tensor, if provided, with the updated state of the Markov chain.

Returns The samples drawn through this observable.

statistics (nn_state, num_samples, num_chains=0, burn_in=1000, steps=1)

Estimates the expected value, variance, and the standard error of the observable over the distribution defined by the WaveFunction.

Parameters

- nn_state (qucumber.nn_states.WaveFunctionBase) The WaveFunction to draw samples from.
- **num_samples** (*int*) The number of samples to draw. The actual number of samples drawn may be slightly higher if *num_samples* % *num_chains* != 0.
- **num_chains** (*int*) The number of Markov chains to run in parallel; if 0 or greater than *num_samples*, will use a number of chains equal to *num_samples*. This is not recommended in the case where a *num_samples* is large, as this may use up all the available memory.
- burn_in (*int*) The number of Gibbs Steps to perform before recording any samples.
- **steps** (*int*) The number of Gibbs Steps to take between each sample.

Returns A dictionary containing the (estimated) expected value (key: "mean"), variance (key: "variance"), and standard error (key: "std_error") of the observable.

Return type dict(str, float)

statistics_from_samples (nn_state, samples)

Estimates the expected value, variance, and the standard error of the observable using the given samples.

Parameters

- **nn_state** (qucumber.nn_states.WaveFunctionBase) The WaveFunction that drew the samples.
- samples (torch.Tensor) A batch of sample states to calculate the observable on.
- **Returns** A dictionary containing the (estimated) expected value (key: "mean"), variance (key: "variance"), and standard error (key: "std_error") of the observable.

Return type dict(str, float)

property symbol

The algebraic symbol representing the Observable.

class qucumber.observables.**SigmaX**(*absolute=False*) Bases: qucumber.observables.ObservableBase

The σ_x observable

Computes the magnetization in the X direction of a spin chain.

Parameters absolute (bool) – Specifies whether to estimate the absolute magnetization.

apply (*nn_state*, *samples*)

Computes the magnetization along X of each sample in the given batch of samples.

Assumes that the computational basis that the WaveFunction was trained on was the Z basis.

- nn_state (qucumber.nn_states.WaveFunctionBase) The WaveFunction that drew the samples.
- samples (torch.Tensor) A batch of samples to calculate the observable on. Must be using the $\sigma_i = 0, 1$ convention.

property name

The name of the Observable.

```
sample (nn_state, k, num_samples=1, initial_state=None, overwrite=False)
```

Draws samples of the observable using the given WaveFunction.

Parameters

- **nn_state** (qucumber.nn_states.WaveFunctionBase) The WaveFunction to draw samples from.
- **k** (*int*) The number of Gibbs Steps to perform before drawing a sample.
- num_samples (*int*) The number of samples to draw.
- initial_state (torch.Tensor) The initial state of the Markov Chain. If given, num_samples will be ignored.
- **overwrite** (*bool*) Whether to overwrite the initial_state tensor, if provided, with the updated state of the Markov chain.

Returns The samples drawn through this observable.

Return type torch.Tensor

statistics (nn_state, num_samples, num_chains=0, burn_in=1000, steps=1)

Estimates the expected value, variance, and the standard error of the observable over the distribution defined by the WaveFunction.

Parameters

- nn_state (qucumber.nn_states.WaveFunctionBase) The WaveFunction to draw samples from.
- **num_samples** (*int*) The number of samples to draw. The actual number of samples drawn may be slightly higher if *num_samples* % *num_chains* != 0.
- **num_chains** (*int*) The number of Markov chains to run in parallel; if 0 or greater than *num_samples*, will use a number of chains equal to *num_samples*. This is not recommended in the case where a *num_samples* is large, as this may use up all the available memory.
- burn_in (*int*) The number of Gibbs Steps to perform before recording any samples.
- **steps** (*int*) The number of Gibbs Steps to take between each sample.
- **Returns** A dictionary containing the (estimated) expected value (key: "mean"), variance (key: "variance"), and standard error (key: "std_error") of the observable.

Return type dict(str, float)

statistics_from_samples(nn_state, samples)

Estimates the expected value, variance, and the standard error of the observable using the given samples.

Parameters

- nn_state (qucumber.nn_states.WaveFunctionBase) The WaveFunction that drew the samples.
- **samples** (*torch.Tensor*) A batch of sample states to calculate the observable on.

Returns A dictionary containing the (estimated) expected value (key: "mean"), variance (key: "variance"), and standard error (key: "std_error") of the observable.

Return type dict(str, float)

property symbol

The algebraic symbol representing the Observable.

class qucumber.observables.**SigmaY**(*absolute=False*) Bases: qucumber.observables.ObservableBase

The σ_u observable

Computes the magnetization in the Y direction of a spin chain.

Parameters absolute (bool) – Specifies whether to estimate the absolute magnetization.

apply (*nn_state*, *samples*)

Computes the magnetization along Y of each sample in the given batch of samples.

Assumes that the computational basis that the WaveFunction was trained on was the Z basis.

Parameters

- **nn_state** (qucumber.nn_states.WaveFunctionBase) The WaveFunction that drew the samples.
- samples (torch.Tensor) A batch of samples to calculate the observable on. Must be using the $\sigma_i = 0, 1$ convention.

property name

The name of the Observable.

sample (nn_state, k, num_samples=1, initial_state=None, overwrite=False)
Draws samples of the observable using the given WaveFunction.

Parameters

- nn_state (qucumber.nn_states.WaveFunctionBase) The WaveFunction to draw samples from.
- **k** (*int*) The number of Gibbs Steps to perform before drawing a sample.
- **num_samples** (*int*) The number of samples to draw.
- initial_state (torch.Tensor) The initial state of the Markov Chain. If given, num_samples will be ignored.
- **overwrite** (*bool*) Whether to overwrite the initial_state tensor, if provided, with the updated state of the Markov chain.

Returns The samples drawn through this observable.

Return type torch.Tensor

statistics (nn_state, num_samples, num_chains=0, burn_in=1000, steps=1)

Estimates the expected value, variance, and the standard error of the observable over the distribution defined by the WaveFunction.

- nn_state (qucumber.nn_states.WaveFunctionBase) The WaveFunction to draw samples from.
- **num_samples** (*int*) The number of samples to draw. The actual number of samples drawn may be slightly higher if *num_samples* % *num_chains* != 0.
- **num_chains** (*int*) The number of Markov chains to run in parallel; if 0 or greater than *num_samples*, will use a number of chains equal to *num_samples*. This is not recommended in the case where a *num_samples* is large, as this may use up all the available memory.

- burn_in (*int*) The number of Gibbs Steps to perform before recording any samples.
- **steps** (*int*) The number of Gibbs Steps to take between each sample.
- **Returns** A dictionary containing the (estimated) expected value (key: "mean"), variance (key: "variance"), and standard error (key: "std_error") of the observable.

Return type dict(str, float)

statistics_from_samples (nn_state, samples)

Estimates the expected value, variance, and the standard error of the observable using the given samples.

Parameters

- **nn_state** (qucumber.nn_states.WaveFunctionBase) The WaveFunction that drew the samples.
- samples (torch.Tensor) A batch of sample states to calculate the observable on.

Returns A dictionary containing the (estimated) expected value (key: "mean"), variance (key: "variance"), and standard error (key: "std_error") of the observable.

Return type dict(str, float)

property symbol

The algebraic symbol representing the Observable.

11.2 Neighbour Interactions

class qucumber.observables.NeighbourInteraction(periodic_bcs=False, c=1)
Bases: qucumber.observables.ObservableBase

The $\sigma_i^z \sigma_{i+c}^z$ observable

Computes the *c*-th nearest neighbour interaction for a spin chain with either open or periodic boundary conditions.

Parameters

- **periodic_bcs** (bool) Specifies whether the system has periodic boundary conditions.
- **c** (*int*) Interaction distance.

apply (nn_state, samples)

Computes the energy of this neighbour interaction for each sample given a batch of samples.

Parameters

- **nn_state** (qucumber.nn_states.WaveFunctionBase) The WaveFunction that drew the samples.
- samples (torch.Tensor) A batch of samples to calculate the observable on. Must be using the $\sigma_i = 0, 1$ convention.

property name

The name of the Observable.

sample (nn_state, k, num_samples=1, initial_state=None, overwrite=False)
Draws samples of the observable using the given WaveFunction.

Parameters

• nn_state (qucumber.nn_states.WaveFunctionBase) - The WaveFunction to draw samples from.

- **k** (*int*) The number of Gibbs Steps to perform before drawing a sample.
- **num_samples** (*int*) The number of samples to draw.
- initial_state (torch.Tensor) The initial state of the Markov Chain. If given, num_samples will be ignored.
- **overwrite** (bool) Whether to overwrite the initial_state tensor, if provided, with the updated state of the Markov chain.

Returns The samples drawn through this observable.

Return type torch.Tensor

statistics (nn_state, num_samples, num_chains=0, burn_in=1000, steps=1)

Estimates the expected value, variance, and the standard error of the observable over the distribution defined by the WaveFunction.

Parameters

- nn_state (qucumber.nn_states.WaveFunctionBase) The WaveFunction to draw samples from.
- **num_samples** (*int*) The number of samples to draw. The actual number of samples drawn may be slightly higher if *num_samples* % *num_chains* != 0.
- num_chains (*int*) The number of Markov chains to run in parallel; if 0 or greater than *num_samples*, will use a number of chains equal to *num_samples*. This is not recommended in the case where a *num_samples* is large, as this may use up all the available memory.
- **burn_in** (*int*) The number of Gibbs Steps to perform before recording any samples.
- **steps** (*int*) The number of Gibbs Steps to take between each sample.

Returns A dictionary containing the (estimated) expected value (key: "mean"), variance (key: "variance"), and standard error (key: "std_error") of the observable.

Return type dict(str, float)

statistics_from_samples(nn_state, samples)

Estimates the expected value, variance, and the standard error of the observable using the given samples.

Parameters

- **nn_state** (qucumber.nn_states.WaveFunctionBase) The WaveFunction that drew the samples.
- **samples** (*torch.Tensor*) A batch of sample states to calculate the observable on.
- **Returns** A dictionary containing the (estimated) expected value (key: "mean"), variance (key: "variance"), and standard error (key: "std_error") of the observable.

Return type dict(str, float)

property symbol

The algebraic symbol representing the Observable.

11.3 Abstract Observable

Note: This is an Abstract Base Class, it is not meant to be used directly. The following API reference is mostly for developers.

class qucumber.observables.ObservableBase

Bases: abc.ABC

Base class for observables.

abstract apply (nn_state, samples)

Computes the value of the observable, row-wise, on a batch of samples.

If we think of the samples given as a set of projective measurements in a given computational basis, this method must return the expectation of the operator with respect to each basis state in *samples*. It must not perform any averaging for statistical purposes, as the proper analysis is delegated to the specialized *statistics* and *statistics_from_samples* methods.

Must be implemented by any subclasses.

Parameters

- nn_state (qucumber.nn_states.WaveFunctionBase) The WaveFunction that drew the samples.
- **samples** (*torch.Tensor*) A batch of sample states to calculate the observable on.

Returns The value of the observable of each given basis state.

Return type torch.Tensor

property name

The name of the Observable.

sample (nn_state, k, num_samples=1, initial_state=None, overwrite=False)

Draws samples of the *observable* using the given WaveFunction.

Parameters

- nn_state (qucumber.nn_states.WaveFunctionBase) The WaveFunction to draw samples from.
- **k** (*int*) The number of Gibbs Steps to perform before drawing a sample.
- **num_samples** (*int*) The number of samples to draw.
- initial_state (torch.Tensor) The initial state of the Markov Chain. If given, *num_samples* will be ignored.
- **overwrite** (*bool*) Whether to overwrite the initial_state tensor, if provided, with the updated state of the Markov chain.

Returns The samples drawn through this observable.

Return type torch.Tensor

statistics (nn_state, num_samples, num_chains=0, burn_in=1000, steps=1)

Estimates the expected value, variance, and the standard error of the observable over the distribution defined by the WaveFunction.

- nn_state (qucumber.nn_states.WaveFunctionBase) The WaveFunction to draw samples from.
- **num_samples** (*int*) The number of samples to draw. The actual number of samples drawn may be slightly higher if *num_samples* % *num_chains* != 0.
- num_chains (*int*) The number of Markov chains to run in parallel; if 0 or greater than *num_samples*, will use a number of chains equal to *num_samples*. This is not recommended in the case where a *num_samples* is large, as this may use up all the available memory.
- **burn_in** (*int*) The number of Gibbs Steps to perform before recording any samples.
- **steps** (*int*) The number of Gibbs Steps to take between each sample.
- **Returns** A dictionary containing the (estimated) expected value (key: "mean"), variance (key: "variance"), and standard error (key: "std_error") of the observable.

Return type dict(str, float)

statistics_from_samples (nn_state, samples)

Estimates the expected value, variance, and the standard error of the observable using the given samples.

Parameters

- nn_state (qucumber.nn_states.WaveFunctionBase) The WaveFunction that drew the samples.
- samples (torch. Tensor) A batch of sample states to calculate the observable on.
- **Returns** A dictionary containing the (estimated) expected value (key: "mean"), variance (key: "variance"), and standard error (key: "std_error") of the observable.

Return type dict(str, float)

property symbol

The algebraic symbol representing the Observable.

CHAPTER

TWELVE

COMPLEX ALGEBRA

qucumber.utils.cplx.absolute_value(x)

Computes the complex absolute value elementwise.

Parameters x (torch. Tensor) – A complex tensor.

Returns A real tensor.

Return type torch.Tensor

qucumber.utils.cplx.conjugate(x)

A function that takes the conjugate transpose of the argument.

Parameters x (*torch.Tensor*) – A complex vector or matrix.

Returns The conjugate of x.

Return type torch.Tensor

qucumber.utils.cplx.elementwise_division(x, y)

Elementwise division of x by y.

Parameters

- **x** (torch. Tensor) A complex tensor.
- y (torch. Tensor) A complex tensor.

Return type torch.Tensor

qucumber.utils.cplx.elementwise_mult(x, y)
Alias for scalar mult().

qucumber.utils.cplx.imag(x)

Returns the imaginary part of a complex tensor.

Parameters x (torch. Tensor) – The complex tensor

Returns The imaginary part of *x*; will have one less dimension than *x*.

Return type torch.Tensor

qucumber.utils.cplx.inner_prod(x, y)

A function that returns the inner product of two complex vectors, x and y (<xly>).

Parameters

- **x** (torch. Tensor) A complex vector.
- **y** (torch. Tensor) A complex vector.

Raises ValueError – If x and y are not complex vectors with their first dimensions being 2, then the function will not execute.

Returns The inner product, $\langle x|y\rangle$.

Return type torch.Tensor

qucumber.utils.cplx.kronecker_prod(x, y)

A function that returns the tensor / kronecker product of 2 complex tensors, x and y.

Parameters

- **x** (torch. Tensor) A complex matrix.
- **y** (torch. Tensor) A complex matrix.
- **Raises ValueError** If x and y do not have 3 dimensions or their first dimension is not 2, the function cannot execute.

Returns The tensorproduct of x and y, $x \otimes y$.

Return type torch.Tensor

qucumber.utils.cplx.make_complex(x, y=None)

A function that combines the real (x) and imaginary (y) parts of a vector or a matrix.

Note: x and y must have the same shape. Also, this will not work for rank zero tensors.

Parameters

- **x** (torch. Tensor) The real part
- **y** (*torch.Tensor*) The imaginary part. Can be None, in which case, the resulting complex tensor will have imaginary part equal to zero.

Returns The tensor [x,y] = x + yi.

Return type torch.Tensor

qucumber.utils.cplx.matmul(x, y)

A function that computes complex matrix-matrix and matrix-vector products.

Note: If one wishes to do matrix-vector products, the vector must be the second argument (y).

Parameters

- **x** (torch. Tensor) A complex matrix.
- **y** (torch. Tensor) A complex vector or matrix.

Returns The product between x and y.

Return type torch.Tensor

qucumber.utils.cplx.norm(x)

A function that returns the norm of the argument.

Parameters x (torch. Tensor) – A complex scalar.

Returns |x|.

qucumber.utils.cplx.norm_sqr(x)

A function that returns the squared norm of the argument.

Parameters x (torch. Tensor) – A complex scalar.

Returns $|x|^2$.

Return type torch.Tensor

qucumber.utils.cplx.outer_prod(x, y)

A function that returns the outer product of two complex vectors, x and y.

Parameters

- **x** (torch.Tensor) A complex vector.
- **y** (torch. Tensor) A complex vector.
- **Raises ValueError** If x and y are not complex vectors with their first dimensions being 2, then an error will be raised.

Returns The outer product between x and y, $|x\rangle\langle y|$.

Return type torch.Tensor

qucumber.utils.cplx.real(x)

Returns the real part of a complex tensor.

Parameters x (torch. Tensor) – The complex tensor

Returns The real part of *x*; will have one less dimension than *x*.

Return type torch.Tensor

qucumber.utils.cplx.scalar_divide(x, y)
 A function that computes the division of x by y.

Parameters

- **x** (torch.Tensor) The numerator (a complex scalar, vector or matrix).
- **y** (*torch*. *Tensor*) The denominator (a complex scalar).

Returns x / y

Return type torch.Tensor

qucumber.utils.cplx.scalar_mult(x, y, out=None)

A function that computes the product between complex matrices and scalars, complex vectors and scalars or two complex scalars.

Parameters

- **x** (*torch*. *Tensor*) A complex scalar, vector or matrix.
- **y** (*torch*. *Tensor*) A complex scalar, vector or matrix.

Returns The product between x and y. Either overwrites *out*, or returns a new tensor.
CHAPTER

THIRTEEN

DATA HANDLING

qucumber.utils.data.extract_refbasis_samples(train_samples, train_bases)
Extract the reference basis samples from the data.

Parameters

- train_samples (torch.Tensor) The training samples.
- train_bases (np.array (dtype=str)) The bases of the training samples.

Returns The samples in the data that are only in the reference basis.

Return type torch.Tensor

Load the data required for training.

Parameters

- **tr_samples_path** (*str*) The path to the training data.
- **tr_psi_path** (*str*) The path to the target/true wavefunction.
- **tr_bases_path** (*str*) The path to the basis data.
- **bases_path** (*str*) The path to a file containing all possible bases used in the tr_bases_path file.

Returns A list of all input parameters.

Return type list

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