## Machine Learning of Many Body Localization

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The goal of this study was to find the quantum phase transition at intermediate local disorder strengths on a Heisenberg chain. Exact diagonalization was used to find the reduced density matrices for a different number of consecutive spins for the lowest energy eigenstate of the Heisenberg Model with an additional random field in z-direction at low and high disorder strengths. The resulting dataset representing extended and localized phases was used to train a neural network. Afterwards, the trained network was applied on intermediate disorder strengths to deduct the critical disorder strength for a phase transition. This phase transition was for all system sizes predicted to be around  $W_c = 1.5J$  for the system sizes  $L \in \{9, 10, 11, 12\}$  and block sizes  $n \in [1, 7]$ . Low block sizes suffered from a low accuracy in the machine learning model, whereas higher block size  $W_c$  values approached  $W_c = J$ .

#### I. INTRODUCTION

The physical model and the concept of exact diagonalization is presented first. As we use reduced density matrices as features for the neural network, we explain briefly their computation and meaning.

## A. Physical model

#### 1. Hamiltonian of the Heisenberg model

The Hamiltonian of the Heisenberg model is shown in equation 1. In the course of further analysis, we choose J=1 and sample h from a uniform distribution such that  $h_i \in [-W,W]$ .

$$H = J \sum_{i} \vec{S}_{i} \cdot \vec{S}_{i+1} - \sum_{i} h_{i} S_{i}^{z}$$
Exchange Energy Random Field (1)

## 2. Expectations for the ground state

The expectation for the ground state is dependent on the ratio of the coupling and the local random field.

For  $\frac{W}{J} \ll 1$ , we expect an delocalized, extended phase, since the exchange energy dominates over the small external field. Therefore, the system can relax to thermal equilibrium serving as its own heat bath in the limit of large system size  $L \to \infty$ . Here, the reduced density operator of a finite subsystem converges to the equilibrium thermal distribution for  $L \to \infty$ .[1]

For  $\frac{W}{J} \gg 1$ , we can expect a localized phase, since the  $h_i$  factors dominate over the exchange energy. The resulting states are expected to be product states of spins "up" or "down", as the external field points in z-direction. Also an infinite system cannot equilibrate itself. The local configurations are set by the initial conditions at all times and are adiabatically connected to the trivial state.[1]

#### B. Exact diagonalization

Exact diagonalization (ED) is a numerical technique we can use to solve the time independent Schrödinger Equation  $H | \psi \rangle = E | \psi \rangle$  for the eigenvalues E and eigenvectors  $| \psi \rangle$ . This only works of the Hamiltonian H represents a discrete and finite system. Most quantum many-particle problems lead to a sparse matrix representation of the Hamiltonian, where only a very small fraction of the matrix elements is non-zero.[2] An efficient method to find ground states is the Lanczos algorithm.[3] At first, the algorithm was numerically unstable. This issue was overcome in 1970 by Ojalvo and Newman.[4] In this study, we rely on the Lanczos algorithm for the eigensolver.

### C. Reduced Density Matrix

The usefulness of reduced density matrices has already been shown by White in 1992 with ground states of Heisenberg chains [5]. In our case we use areal density matrices as features for the neural network to predict the critical disorder strength of a phase change from delocalized to localized. The reduced density matrix is defined in equation 3. Physically, the reduced density matrix  $\rho_A$ , provides correct measurement statistics for subsystem A.

$$\rho_{AB} = |\psi_A\rangle \langle \psi_A| \otimes |\psi_B\rangle \langle \psi_B| \tag{2}$$

$$\rho_A = \operatorname{Tr}_B(\rho_{AB}) = |\psi_A\rangle \langle \psi_A| \operatorname{Tr}(|\psi_B\rangle \langle \psi_B|) \quad (3)$$

The reduced density matrix was also used by Zhang in 2019 to learn the localization transition in disordered quantum Ising spin chains. Here, the motivation was to reduce the dimension and filter out redundant information. However, it proved to be inferior in comparison to the full density matrix in the analysis. [6] However, due to RAM limitations, we will rely on reduced density matrices.

#### D. Artificial Neural Networks

Rosenblatt published in 1958 his concept of the probabilistic model for information storage and organization in the brain, which greatly inspired others to use those models for computation.[7] Over the course of years, they have evolved to a tool that can be used for a variety of applications including computer vision, speech recognition, medical diagnosis, playing games or even artistic painting.[8]

The reduced density matrices are essentially complex 2D arrays with length  $2^n \times 2^n$ . As we want to classify for an arbitrary W whether we have a localized or delocalized phase, it is convenient to use a machine learning classifier. The density matrices can then be thought of as a complex and real image that can be fed into it analogously to classical image classification.

#### II. COMPUTATIONAL METHODS

The strategy for implementation was as follows:

- 1. Generate Hamiltonian from random disorder strength and system size. Then calculate lowest eigenstate near Energy E=0.
- 2. Generate density matrix from the eigenstate and the respective reduced density matrices for defined block sizes n.
- 3. Set up machine learning model per n, L that takes density matrices of different W as an input and predicts whether the state represents an extended or a localized phase.
- 4. Make predictions for different system sizes L and block sizes n and plot the predictions over W. Then extract  $W_c$  from the data by using a fit function.

Critical decisions and specifications for each steps are listed below. Afterwards, a brief motivation for the parameter range and resolution is given.

## A. Eigenvalue solver

For the eigenvalue solution, we use SciPy's method eigsh through QuTiP's method groundstate [9, 10]. In comparison, a naive parameter choice for eigsh for N=10 lattice sites needed 70 s to calculate the ground state, whereas groundstate only took 0.7 s, by choosing an optimized parameter set for eigsh. Of course, eigsh supplies the user with k eigenvalues instead of only one, but this feature was not found to be critical for the further analysis. Therefore, groundstate is used throughout the program, to avoid making a non optimal parameter choice.

#### B. Computation of reduced density matrix

To get the reduced density matrix of system A, one has to "trace out" all states outside of A. Luckily, the library QuTiP supplies a method ptrace, which does exactly that. It is important to note that the method takes those indices as an argument which should be kept.[10]

A demonstration of the functionality can be found in Figure 1.

```
density matrix:
 [[0. 0. 0. 0. 0. 0. 0. 0.]
 [0. 1. 0. 0. 0. 0. 0. 0.]
 [0. 0. 2. 0. 0. 0. 0. 0.]
 [0. 0. 0. 3. 0. 0. 0. 0.]
 [0. 0. 0. 0. 4. 0. 0. 0.]
 [0. 0. 0. 0. 0. 5. 0. 0.]
 [0. 0. 0. 0. 0. 6. 0.]
 [0. 0. 0. 0. 0. 0. 0. 7.]]
Summation over all but first lattice site:
 [[ 6.+0.j 0.+0.j]
 [ 0.+0.j 22.+0.j]]
Summation over first lattice site:
 [[ 4.+0.j 0.+0.j 0.+0.j 0.+0.j]
  0.+0.j 6.+0.j 0.+0.j 0.+0.j]
 [ 0.+0.j 0.+0.j 8.+0.j 0.+0.j]
 [ 0.+0.j 0.+0.j 0.+0.j 10.+0.j]]
```

Figure 1: Proof of concept for partial trace calculation similar to QuTiP-Guide/ptrace.

The algorithm of selecting the position vector of n consecutive sites was implemented as follows:

- 1. Find the center spin rounded to next lowest integer.
- 2. Determine left chain length  $n_{\text{left}}$  as n/2 rounded to the next lowest integer.
- 3. Determine right chain length  $n_{\text{right}}$  as  $n n_{\text{left}}$ .
- 4. Select spins from left chain end to right chain end around center spin.

This results in a behavior that picks left indices more favorably, but succeeds if equally spaced ends exist. Let the spins be numbered as  $\{1, 2, 3, 4, 5\}$  for N = 5, then n = 3 results in  $\{2, 3, 4\}$ , whereas n = 2 results in  $\{2, 3\}$ .

These lattice sites serve then as an input to the partial trace function, such that the density matrix represents the measurement statistics of the center system.

#### C. Machine learning models and error metrics

The decision for the machine learning framework keras was motivated by its flexibility and simplicity. [11]

When setting up the machine learning model, one can already specify the first and last layer: The first (input) layer has to match the sample size of the incoming data, this can be already computed in advance. The length len for block size n is  $2 \cdot (2^n \times 2^n)$ . The factor 2 comes from a preprocessing step, where the complex values are

mapped to a second real picture, since the fitting procedure usually does not expect complex numbers. The last layer is a one node sigmoid, as the target output is the one-dimensional classification in [0,1].

For small sample sizes, there exist various approaches to choose the right amount of layers and regularization methods [12, 13], which cannot be generalized, as they heavily depend on feature size and target dimension. As a rule of thumb the approximation was used that each weight should be influenced by at least seven samples. Using this we get from 500 samples roughly 70 weights.

The optimizer Adam was chosen, because it is computationally efficient, has little memory requirements. [14]

For a two label classification problem, it is useful to use cross-entropy as a loss metric, as the penalty increases exponentially the further one deviates from the correct prediction. [15] The definition for a two class cross-

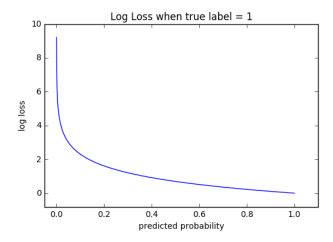


Figure 2: Cross-Entropy Loss

entropy loss can be found in equation 4, where  $y \in \{0, 1\}$  is the true class and  $\hat{y} \in [0, 1]$  the predicted probability. This loss is also plotted in Figure 2.

$$L(\hat{y}, y) = -(y \log(\hat{y}) + (1 - y) \log(1 - \hat{y})) \tag{4}$$

### D. Extraction of critical disorder strength $W_c$

To fit for the critical disorder strength  $W_c$ , two functions were compared. The logistic Fermi-Dirac like function:

$$L: \mathbb{R} \to [0,1] \tag{5}$$

$$W_{pred} \mapsto \frac{1}{\exp\left(-\alpha \left(W_{pred} - W_c\right)\right) + 1} \tag{6}$$

and the heaviside function:

$$H\colon \mathbb{R} \to \{0,1\} \tag{7}$$

$$W_{pred} \mapsto \begin{cases} 0: & W_{pred} < W_c \\ 1: & W_{pred} \ge W_c \end{cases} \tag{8}$$

The fully delocalized phase is defined as 0 and fully localized as 1. Whereas the heaviside function has an abrupt step and only maps to the extrema, the logistic function serves as a smoother option for a transition, depending on the parameter alpha. The motivation came also from an optimizers view: Differentiable functions are easier to fit for the computer.[16] Therefore, the logistic function was used to extract  $W_c$  with the empiric decision of  $\alpha = 50$ .

#### E. Limitations for parameter range and resolution

- 1. System size L: Limited by computing time of eigenvalue solver. For the system size L=12, one calculation lasted approximately one minute.
- 2. Block size n: 500 samples, L=9, n=8 required 4 GB of storage for the training set, exceeding the machines performance during model fitting. Therefore, n=7 was found to be sufficient for all system sizes.
- 3. Sample size: 500 samples can be generated for L = 12,  $n_{max} = 7$  in approximately 9 hours. This was found to be a sufficient sample size per system and block size.
- 4. Disorder strength W for the testing set: Since each point of a test set comes with a Hamiltonian with randomly drawn  $h_i \in [-W, W]$ , a decent amount of variance can be expected for the phase prediction. As we want to extract the phase change in general, and are not interested in the particular phase predictions of one specific Hamiltonian we choose to regularize the prediction by averaging over five predicted samples.

### III. RESULTS

# A. Generation of reduced density matrix training set

The parameter range for the computation of the reduced density matrices can be found in Table I. The total computation time was 16.5 h, where 12.5 h where solely needed to compute the ground states of the L=12 system.

Parameter	Range or Set
System size:	$L \in \{9, 10, 11, 12\}$
Block size:	$n \in \{1,2,3,4,5,6,7\}$
Repetitions:	r = 500

Table I: Parameter choice for training set generation

In order to give some visual intuition, Figure 3 shows realizations for different block sizes and phases.

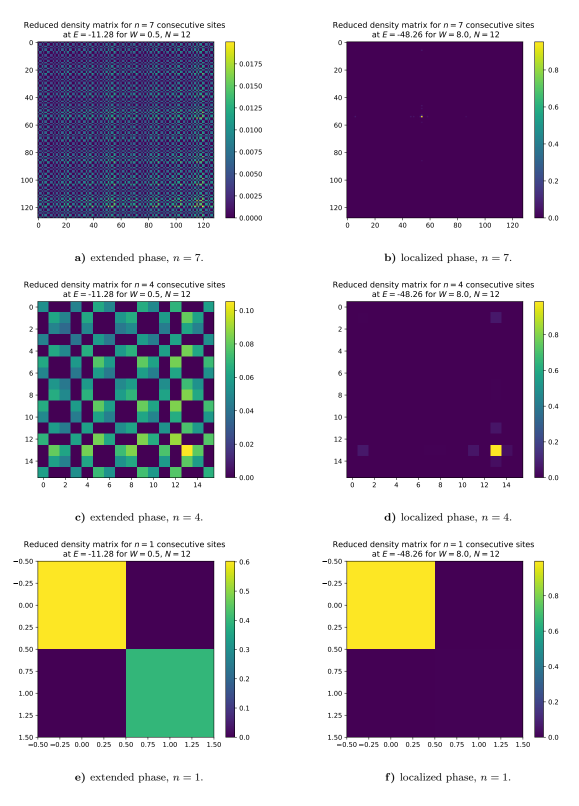


Figure 3: Real part of the density matrix of an ergodic and localized phases for block sizes  $n = \{4, 5\}$  and system size L = 12.

The visual inspection indicates that the density matrix of the localized phase has a sharp maximum at the

preferred state that is forced by the random disorder strength. The extended phase shows a checkerboard pattern structure, which reflects that some configurations are more preferred than others. These energetically expensive states are related to neighboring unaligned spins. Another observation is that the density matrix reductions of the full ground state conserved these properties for n > 2, when comparing n = 7 to  $n = \{6, 5, 4, 3, 2\}$ . The similarity between the two phases gets smaller the smaller the block size n. For n = 1, one could argue that the density matrices are very similar, as they only deviate for half the matrix elements.

#### B. Model training

Before we can predict the phase of a newly generated test set, we have to train the neural network with our available training data. For each system and block size a separate model was trained, as a different system size might influence the physical behavior due to open boundary conditions.

The neural networks are generated as a sequential keras model with the following configuration, as discussed in section II C:

```
model = models.Sequential()
model.add(layers.Flatten())
model.add(layers.Dense(64, activation='relu',
```

```
bias_regularizer='12'))

model.add(layers.Dense(64, activation='relu',
    bias_regularizer='12'))

model.add(layers.Dense(1, activation='sigmoid'))

model.compile(optimizer='adam', loss='
    binary_crossentropy', metrics=['accuracy'])
```

Two strategies are employed to prevent over-fitting:

- 30 % of the training set was used for validation. To avoid a biased split, we relied on sklearn's method train\_test\_split that samples randomly from the training set.
- 2. A bias regularizer was introduced to move the output functions closer to the origin. Even though some further regularization might still be possible, a kernel regularizer did not prove to be useful and impacted the resulting scores heavily.

The model training was executed by using a batch size of 70 and 40 epoch, where the batch size was limited by the CPU performance and no significant loss or accuracy improvements where noted after 30 epochs.

An example of the accuracy and loss dependency on the number of epochs for system size L=10, and block sizes  $n=\{1,7\}$  is presented below in Figure .

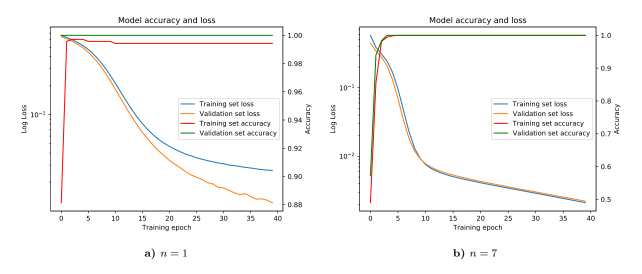


Figure 4: Loss and accuracy on training and validation set for system size L=10.

The overall losses for n>1 where always found to be  $\approx 0.01$  with accuracies of 1.0, whereas the models at n=1 showed larger losses of  $\approx 0.1$  and mostly acceptable accuracies near 1.

Figure 4 illustrates that not only the scores for small block sizes were lower, but also the convergence rates. In conclusion, the scores show that the learning of the phases was prone to severe over-fitting and resulted in

acceptable scores for the next step of prediction, where n=1 can be expected to have a worse performance during the prediction. This behavior was expected, when we took a closer look at the training set.

# C. Prediction of extended vs localized phase and analysis of critical disorder strength

First, the testing set was generated. Following the parameter discussion in section II E, we generate five samples for each  $W \in [0, 4]$ , with step  $\Delta W = 0.05$ , resulting

in 400 samples per system and block size. Afterwards, the predictions were fitted with a logistic function to obtain  $W_c$  as described in section II D.

Five predicted phases are averaged at each point and plotted to a heat map. The fitted  $W_c$  is plotted along in Figure 5.

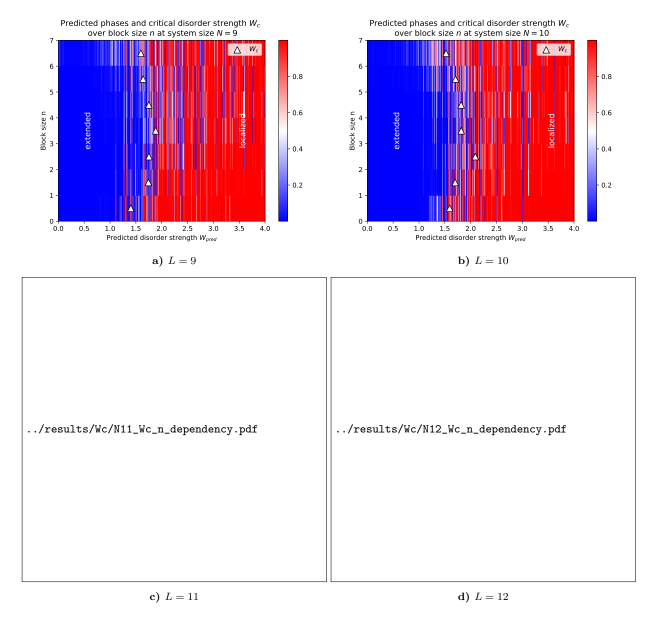


Figure 5: Dependency of the phase transition on block size n for different system sizes.

In conclusion, the predicted critical disorder strength  $W_c$  decayed, when models with larger block sizes n were used for prediction. The low  $W_c$  values for  $n=\{1,2\}$  might just as well be attributed to the poor loss and accuracy values shown in section III B. An explanation for

this decay might be that a bigger block size can more accurately reflect the level of disorder forced on the system. For smaller block sizes, for some spins the information is lost whether the configuration was the result of interacting lattice sites or the random disorder strength.

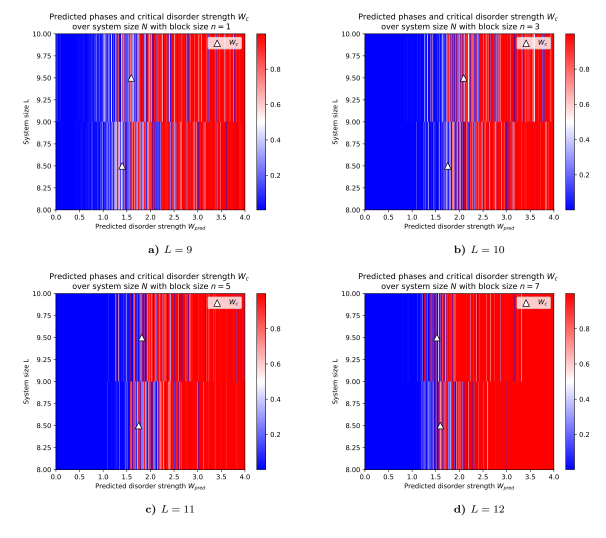


Figure 6: Dependency of the phase transition over system sizes L for different block sizes n.

The plots are indicating that a bigger system size requires a larger disorder strength to perform the phase transition.

These are our  $W_c$  depending on n, L.

## IV. CONCLUSION

 $W_c$  depends on n, L (yes/no).

 $W_c$  prediction coincides with the expectation (yes/no)

 $W_c$  is dependent on these and that effects = $\xi$  scaling analysis? (yes/no)

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## Appendix A: Code listing

The code consists essentially of four different files, which are callable through a main function, but can also be run separately. Every file serves a number of different purposes as listed below.

- 1. **generate\_training\_set.py**: Here, the training set is generated and some example plots of ground states are saved to the results folder. The training sets are saved in the **training\_sets** folder, where they are numbered with their system and block size.
- 2. **model\_save\_train.py**: First, models are generated that automatically match the input data of different block sizes n, afterwards, they are trained with a certain amount of epochs and batch sizes. The history of the validation and accuracy are plotted individually into the results folder.
- 3. **generate\_test\_set.py**: A set of reduced density matrices for ground states in the intermediate regime is generated.
- 4. **load\_model\_get\_wc.py**: The models for each system and block size make phase predictions to the respective test sets, extract  $W_c$  and plot everything together as a heat map.

## 1. Training set generation

```
1 from ed import *
2 import time
3 import pickle
  from scipy.sparse.linalg import ArpackNoConvergence
  import qutip
8
  def generate_training_set(Ns, Ws, n_max, repetitions):
      start_time = time.time()
9
       for N in Ns:
           training_set_generator = TrainingSetGenerator(N, Ws, n_max, repetitions)
           print("Training Set N="+str(N)+" completed after %s seconds." % (time.time() - start_time))
12
13
           for n in range(1, n_max+1):
               save\_groundstate\_figures (\texttt{N}, training\_set\_generator.training\_set[\texttt{n}], \ \texttt{n})
14
               save_pickle("lanczos/training_sets/N" + str(N) + "n" + str(n) + "_Trainset",
      training_set_generator.training_set[n])
      print("--- Training set generation lasted %s seconds ---" % (time.time() - start_time))
17
```

```
18
19
    def save_groundstate_figures(N, training_set, n): # reduced_rho, W, self.N, n, E
20
            ergodic = [item for item in training_set if item[1] == 0.5 and item[-1] == 0][0] # len:
21
           repetitions
           localized = [item for item in training_set if item[1] == 8 and item[-1] == 0][0] # len:
           repetitions
23
           fig, ax1 = plt.subplots()
24
25
           pos = ax1.imshow(np.real(ergodic[0]), cmap='bwr')
26
           fig.colorbar(pos, ax=ax1)
           plt.title("Reduced density matrix for $n=$" + str(n) + " consecutive sites \n at $E=$" extraction of the consecutive sites of the consecutive si
27
                              + str(round(ergodic[4], 2)) + " for $W=$" + str(ergodic[1]) + ", $N = $" + str(N))
28
           plt.savefig(
29
                   "results/groundstates/N" + str(N) + "n" + str(n) + "_trainingset_groundstate_Wmax" + str(
30
           ergodic[1]) + ".pdf")
           plt.close()
31
32
           fig, ax1 = plt.subplots()
33
34
           pos = ax1.imshow(np.real(localized[0]), cmap='bwr')
           fig.colorbar(pos, ax=ax1)
35
           plt.title("Reduced density matrix for $n=$" + str(localized[3]) + " consecutive sites \n at $E=
36
                              + str(round(localized[4], 2)) + " for $W=$" + str(localized[1]) + ", $N = $" + str(N)
37
           plt.savefig(
38
39
                   "results/groundstates/N" + str(N) + "n" + str(localized[3]) + "
             trainingset_groundstate_Wmax" + str(localized[1]) + ".pdf")
40
           plt.close()
           pass
41
42
43
    def save_pickle(filename, data):
44
            with open(filename, 'wb') as f:
46
                   pickle.dump(data, f)
47
    class TrainingSetGenerator:
49
50
           def __init__(self, N, Ws, n_max, repetitions):
5.1
                   self.N = int(N) # Lattice sites
52
53
                   self.n_max = n_max
                   self.repetitions = repetitions
                   self.Ws = Ws
                   self.training_set = self.generate_training_set_m_lanczos_list() # self.
56
           generate_training_set_list()
57
           def generate_training_set_m_lanczos_list(self):
58
59
                   Returns training set with shape samples x [density matrix, W, lattice sites, block size,
60
           ground state energy]
61
                   :return: training set
62
                   training_set = {consecutive_spins: [] for consecutive_spins in range(1,self.n_max+1)}
63
                   for W in self.Ws:
64
                          for rep in range(self.repetitions):
65
                                 H = gen_hamiltonian_random_h(self.N, W=W, J=1.)
66
67
                                 E, v = qutip.Qobj(H).groundstate() # fixme might not be sparse, make sparse=True!!!
68
                                 rho = np.outer(v, v)
                                 for n in range(1, self.n_max+1):
69
                                         reduced_rho = self.get_partial_trace(rho, n) # must trace out something
70
                                         training_set[n].append([reduced_rho, W, self.N, n, E, rep])
71
                                 # training_set[self.N].append([rho, W, self.N, self.N, E, rep])
72
                   return training_set
73
74
           def get_partial_trace(self, rho, n):
75
76
                   calculates partial trace by reshaping the density matrix and adding along the axis
77
78
                   :param rho: full density matrix
79
                   :param n: block size
```

```
:return: reduced density matrix
80
81
           kept_sites = self.get_keep_indices(n)
82
           \label{eq:qutip_dm} qutip\_dm = qutip.Qobj(rho, dims=[[2]*self.N]*2)
83
           reduced_dm_via_qutip = qutip_dm.ptrace(kept_sites).full()
85
           return reduced_dm_via_qutip
86
87
       def diff(self, first, second):
           second = set(second)
88
           return [item for item in first if item not in second]
89
90
91
       def get_keep_indices(self, n):
92
           Determines the middle indices for lattice sites numbered from 0 to N-1. Picks left indices
93
       more favourably.
           :return: List of complement of n consecutive indices
94
95
           left_center = n // 2
96
           right_center = n - left_center
97
98
           middle = self.N // 2
           sites = np.arange(self.N)
99
           return sites[middle - left_center:middle + right_center].tolist()
100
   if __name__ == "__main__":
103
       Ns = [10]
104
105
       n_max = 7
       Ws = [0.5, 8.0] \# 0.5 \Rightarrow ergodic/delocalized phase, 8.0 localized phase
106
107
       repetitions = 500
108
       generate_training_set(Ns, Ws, n_max, repetitions)
       # N=09, n=7, rep=10 7s=> rep=500: 6 min
       # N=10, n=7, rep=10 31s => rep=500: 25 min
       # N=11, n=7, rep=10 182s=> rep=500: 2,5 h
     # N=12, n=7, rep=10 00s=> rep=500
113
```

#### 2. Model Training

```
1 from sklearn.model_selection import train_test_split
2 import pickle
3 from tensorflow.keras import layers, models, regularizers
4 import numpy as np
5 import matplotlib.pyplot as plt
6 import tensorflow.keras.backend as k
7 import time
9
  def load_pickle(filename, to_numeric=1):
10
      with open(filename, 'rb') as f:
11
          data = pickle.load(f)
      return data
13
14
15
16 def preprocess_training_data(path): # reduced_rho, W, self.N, n, E
      data = load_pickle(path)
17
18
      X = data
      X = [item[0] for item in X]
19
      X = np.reshape(X, (np.shape(X)[0], np.shape(X)[1], np.shape(X)[2], 1))
20
      X = np.asarray(np.concatenate((np.real(X), np.imag(X)), axis=3))
21
22
      y = data
      y = np.reshape(np.asarray([map_target(item[1]) for item in data]), (np.shape(y)[0], 1))
23
24
      return X, y
2.5
def map_target(item):
27
      if item == 0.5:
          return 0 # ergodic/delocalized phase
28
29
      elif item == 8.0:
         return 1 # localized phase
30
31
    print("Invalid training data.")
32
```

```
33
def mean_pred(y_true, y_pred):
      return k.mean(y_pred)
35
36
37
38
  class ModelTrainer:
39
40
      def __init__(self, x, y, N, n_max):
          self.N = N
41
          self.n_max = n_max
42
          self.X_train, self.X_test, self.y_train, self.y_test = train_test_split(x, y, test_size
43
      =0.3, random_state=42)
44
          self.model = self.generate_model_sparse()
45
      def generate_model(self):
46
          model = models.Sequential()
47
          model.add(layers.Flatten())
48
          model.add(layers.Dense(64, activation='relu'))
49
          model.add(layers.Dense(128, activation='relu'))
50
51
          model.add(layers.Dense(64, activation='relu'))
          model.add(layers.Dense(32, activation='relu'))
52
53
          model.add(layers.Dense(1, activation='sigmoid'))
          model.compile(optimizer='rmsprop', loss='mae', metrics=['accuracy'])#loss used to be mae
54
      loss # metrics: 'mean_absolute_error', 'mean_squared_error',
          return model
56
57
      def generate_model_sparse(self):
          model = models.Sequential()
58
          # if self.N != 12:
59
          # model.add(layers.Conv2D(32, (6, 6), activation='relu', input_shape=(np.shape(self.X_train
60
      )[1], np.shape(self.X_train)[1], 2)))
          # model.add(layers.MaxPooling2D((4, 4)))
          62
       2)))
          model.add(layers.Dense(64, activation='relu', bias_regularizer='12')), # #
63
          model.add(layers.Dense(64, activation='relu', bias_regularizer='12'))) # fixme use kernel
64
      regularizer!! 11 loss as squared error is dangerous below 1
          model.add(layers.Dense(1, activation='sigmoid'))
65
          model.compile(optimizer='adam', loss='binary_crossentropy', metrics=['accuracy'])#loss used
66
       to be mae loss # metrics: 'mean_absolute_error', 'mean_squared_error',
          return model
67
      def score(self):
69
          score = self.model.evaluate(self.X_test, self.y_test, verbose=0)
70
          print("test loss: %.3E, test acc: %.3E" % (score[0], score[1]))
71
72
          pass
      def fit_model(self, batch_size, epochs):
74
          history = self.model.fit(self.X_train, self.y_train,
75
                         batch_size=batch_size,
76
                         epochs=epochs,
                         verbose=0.#2
78
                         validation_data=(self.X_test, self.y_test)
79
80
          return history
81
82
      def save_model(self, filepath):
83
          self.model.save(filepath)
84
85
      def training_history(self, history, n, N):
86
          fig, ax1 = plt.subplots()
88
          plt.title('Model accuracy and loss for n='+str(n)+', N='+str(N))
89
          plt.xlabel('Training epoch')
90
91
          # "Loss"
92
          ax1.set_ylabel('Log Loss') # we already handled the x-label with ax1
93
          ax1.set_yscale('log')
94
95
          ax1.tick_params(axis='y')
96
          ln1 = ax1.plot(history.history['loss'], label='Training set loss')
```

```
ln2 = ax1.plot(history.history['val_loss'], label='Validation set loss')
97
98
            # "Accuracy"
99
            ax2 = ax1.twinx() # instantiate a second axes that shares the same x-axis
100
            ax2.set_ylabel('Accuracy') # we already handled the x-label with ax1
            ax2.tick_params(axis='y')
           ln3 = ax2.plot(history.history['acc'], 'r', label='Training set accuracy')
ln4 = ax2.plot(history.history['val_acc'], 'g', label='Validation set accuracy')
104
            # Joined Legend
106
           lns = ln1 + ln2 + ln3 + ln4
            labs = [l.get_label() for l in lns]
108
            ax1.legend(lns, labs, loc="center right")
            plt.tight_layout()
            \verb|plt.savefig("results/accuracy_loss_epochs/N"+str(self.N)+"n"+str(n)+"_accuracy_loss_epochs.||
       pdf")
            print("Scores for N=" + str(N) + ", n=" + str(n))
113
            self.score()
114
            pass
116
   def train_save_model(Ns, n_max, batch_size, epochs):
117
118
       start_time = time.time()
       for N in Ns:
119
120
            start_model_time = time.time()
            for n in range(1, n_max+1):
                X, y = preprocess_training_data("lanczos/training_sets/N"+str(N)+"n"+str(n)+"_Trainset"
       )
                model_trainer = ModelTrainer(X, y, N, n_max)
                history = model_trainer.fit_model(batch_size=batch_size,
124
                                                      epochs = epochs)
                model_trainer.training_history(history, n, N)
126
                model_trainer.save_model("lanczos/models/N"+str(N)+"n"+str(n)+"_Model")
            print("--- Model trainings for N=" + str(N) + " lasted %s seconds ---" % (
128
                             time.time() - start_model_time))
       print("--- Model training lasted %s seconds ---" % (time.time() - start_time))
130
       pass
133
   if __name__ == "__main__":
134
       # Ns = [10, 11, 12]
135
       Ns = [10]
136
       n_max = 7
       train_save_model(Ns, n_max,
138
                          batch_size=70,
                          epochs=40)
140
141
       # N = 12 Model training lasted 537.23 seconds
142
```

#### 3. Test set generation

```
1 from generate_training_set import TrainingSetGenerator, save_pickle
2 from model_save_train import *
3 import time
  def generate_test_set(Ns, Ws, n_max, repetitions):
5
      start_time = time.time()
      for N in Ns:
          training_set_generator = TrainingSetGenerator(N, Ws, n_max, repetitions)
9
          print("Testing Set N=" + str(N) + " completed after %s seconds." % (time.time() -
      start_time))
          for n in range(1, n_max+1):
              save\_pickle("lanczos/test\_sets/N"+str(N)+"n"+str(n)+"\_Testset", training\_set\_generator.
      training_set[n])
      print("--- Testing set generation lasted %s seconds ---" % (time.time() - start_time))
      pass
14
15
16 if __name__ == "__main__":
Ns = [9]
```

#### 4. Prediction and evaluation of $W_c$

```
1 from generate_training_set import TrainingSetGenerator, save_pickle
 from model save train import *
     from scipy.optimize import curve_fit
 6
     def preprocess_test_data(path):
              :param path: Path to pickled test_set
              :return: X: reduced density matrices, W: Disorder strength that was used for generating the
 9
              sample
              print("Accessing ",path)
              data = load_pickle(path)
              X = [item[0] for item in data]
              # print("Input shape (Ws, Imagedim1, Imagedim2): ", np.shape(X))
14
              X = np.reshape(X, (np.shape(X)[0], np.shape(X)[1], np.shape(X)[2], 1))
1.5
              X = np.asarray(np.concatenate((np.real(X), np.imag(X)), axis=3))
              W = np.reshape(np.asarray([item[1] for item in data]), (np.shape(data)[0], 1))
17
              return X, W
18
19
20
     def logistic(x, a):
21
              return 1 / (1 + np.exp(-50 * (x - a)))
23
24
def heaviside(x, a):
              return 0.5*np.sign(x-a)+0.5
27
28
29 def load_model(path):
              return models.load_model(path)
30
31
32
33
     def get_wc(N, n, Ws, repetitions):
34
              Calculates Wc
35
36
              :param N: system size for Model and Testset
37
              :param n: block size for Model and Testset
38
              :param Ws: chosen interval for fitting
39
              :param repetitions: Number of datapoints per W_pred
40
41
              :return: Wc, n
42
              model = load_model('lanczos/models/N' + str(N) + 'n' + str(n) + '_Model')
43
             X, W = preprocess_test_data('lanczos/test_sets/N' + str(N) + 'n' + str(n) +'_Testset')
44
              state_prediction = model.predict(X)
46
              state\_prediction = np.reshape(state\_prediction, (int(len(state\_prediction)/repetitions), length of the context of the contex
47
              repetitions))
              state_prediction = np.mean(state_prediction, axis=1)
48
              popt, pcov = curve_fit(logistic, Ws, np.reshape(state_prediction, (len(state_prediction)))) #
50
              state_prediction.astype(np.float))
5.1
              # plot_wc_fit(N,popt,state_prediction)
              return popt[0]# , n #, N #, np.shape(X[0])[0]
52
53
def get_wc_N(N, n, Ws, repetitions):
             Calculates Wc
56
```

```
57
58
       :param N: system size for Model and Testset
       :param n: block size for Model and Testset
59
       :param Ws: chosen interval for fitting
60
       :param repetitions: Number of datapoints per W_pred
61
62
       :return: Wc, n
63
       model = load_model('lanczos/models/N' + str(N) + 'n' + str(n) + '_Model')
64
       X, W = preprocess_test_data('lanczos/test_sets/N' + str(N) + 'n' + str(n) +'_Testset')
65
66
       state_prediction = model.predict(X)
67
       state_prediction = np.reshape(state_prediction, (int(len(state_prediction)/repetitions),
68
       repetitions))
       state_prediction = np.mean(state_prediction, axis=1)
69
70
       popt, pcov = curve_fit(logistic, Ws, np.reshape(state_prediction, (len(state_prediction)))) #
71
       state_prediction.astype(np.float))
72
       # plot_wc_fit(N,popt,state_prediction)
       return popt[0]# , N #, N #, np.shape(X[0])[0]
74
75
   class HeatMapPlotter:
76
       def __init__(self, Ns, Ws, n_max, repetitions):
78
79
           self.Ns = Ns
           self.Ws = Ws
80
81
            self.n_max = n_max
           self.repetitions = repetitions
82
83
84
       def predict_w_n(self):
85
            W_preds = {system_size : [] <mark>for</mark> system_size <mark>in</mark> self.Ns}
86
           for N in self.Ns:
87
88
                for n in range(1, self.n_max + 1):
                    model = load_model('lanczos/models/N' + str(N) + 'n' + str(n) + '_Model')
89
                    X, W = preprocess_test_data('lanczos/test_sets/N' + str(N) + 'n' + str(n) + '
90
       _Testset')
                    W_preds[N].append(model.predict(X))
91
           return W_preds
92
93
       def fit_wc_n(self):
94
           W_c_fit = {system_size : [] for system_size in self.Ns}
95
           for N in self.Ns:
96
                for n in range(1, self.n_max + 1):
97
                    \label{lem:wc_fit_N_append} $$ W_c_fit[N].append((get_wc(N, n, self.Ws, self.repetitions), n)) $$
98
           return W_c_fit
99
100
       def predict_w_N(self):
            W_preds = {block_size : [] for block_size in range(1, self.n_max+1)}
           for n in range(1, self.n_max + 1):
                for N in self.Ns:
104
                    model = load_model('lanczos/models/N' + str(N) + 'n' + str(n) + '_Model')
                    X, W = preprocess_test_data('lanczos/test_sets/N' + str(N) + 'n' + str(n) + '
106
       _Testset')
                    W_preds[n].append(model.predict(X))
           return W_preds
108
109
       def fit_wc_N(self):
           W_c_fit = {block_size : [] for block_size in range(1, self.n_max+1)}
           for n in range(1, self.n_max + 1):
                for N in self.Ns:
113
                    W_c_fit[n].append((get_wc_N(N, n, self.Ws, self.repetitions),N))
114
           return W_c_fit
117
       def plot_wc_heatmap_n(self):
118
119
           Plots Heatmap with blocksize and W_pred
120
           W_pred: W x n array
           W_c_fit: W_c(n) x 1 array
```

```
124
           self.W_preds = self.predict_w_n()
           self.W_c_fit = self.fit_wc_n()
126
           for N in self.Ns:
127
                W_pred = np.asarray(self.W_preds[N])
128
                W_pred = np.reshape(W_pred, (np.shape(W_pred)[0],np.shape(W_pred)[1]))
130
                W_c_fit = np.array(self.W_c_fit[N])
                 \begin{tabular}{ll} \# \ \ W_c_fit = np.reshape(W_c_fit), (np.shape(W_c_fit)[0], np.shape(W_c_fit)[1])) \end{tabular} 
                fig, ax = plt.subplots()
134
                plt.title("Predicted phases and critical disorder strength $W_c$ \n over block size $n$
        at system size $N=$" + str(N))
                plt.text(0.5, 3.5, 'extended', {'color': 'w', 'fontsize': 12},
136
                         horizontalalignment='left',
                         verticalalignment='center',
138
                         rotation=90,
140
141
                plt.text(3.5, 3.5, 'localized', {'color': 'w', 'fontsize': 12},
                         horizontalalignment='left',
142
                          verticalalignment='center',
                         rotation=90,
144
145
146
                pos = ax.imshow(W_pred, extent=(0, 4, 0, 7), aspect=0.5, cmap='bwr')
                fig.colorbar(pos, ax=ax)
147
148
                ax.scatter(W_c_fit[:,0], W_c_fit[:,1]-0.5, s=100, c="w", marker='^', label='$W_c$',
       edgecolors="k")
               plt.ylabel("Block size n")
149
                plt.xlabel("Predicted disorder strength $W_{pred}$")
                ax.legend()
                plt.tight_layout()
                plt.savefig('results/Wc/N'+str(N)+'_Wc_n_dependency.pdf')
154
                plt.close()
           pass
       def plot_wc_heatmap_N(self):
157
158
           Plots Heatmap with blocksize and W_pred
159
160
           W_pred: W x n array
161
           W_c_fit: W_c(n) x 1 array
           self.W_preds = self.predict_w_N()
164
           self.W_c_fit = self.fit_wc_N()
165
166
           for n in range(1, self.n_max+1):
167
                W_pred = np.asarray(self.W_preds[n])
168
                W_pred = np.reshape(W_pred, (np.shape(W_pred)[0],np.shape(W_pred)[1]))
                W_c_fit = np.array(self.W_c_fit[n])
                # W_c_fit = np.reshape(W_c_fit, (np.shape(W_c_fit)[0], np.shape(W_c_fit)[1]))
173
                fig, ax = plt.subplots()
174
                \textbf{plt.title("Predicted phases and critical disorder strength $W_c$ \n over system size}
       N with block size n=" + str(n))
                plt.text(0.5, 3.5, 'extended', {'color': 'w', 'fontsize': 12},
                         horizontalalignment='left',
177
178
                         verticalalignment='center',
                         rotation=90,
179
                plt.text(3.5, 3.5, 'localized', {'color': 'w', 'fontsize': 12},
181
                         horizontalalignment='left',
182
                         verticalalignment='center',
183
                         rotation=90,
184
185
186
                pos = ax.imshow(W_pred, extent=(0, 4, self.Ns[0]-1, self.Ns[-1]), aspect='auto', cmap='
       bwr')
187
                # Shift ticks to be at 0.5, 1.5, etc
                # ax.yaxis.set(ticks=np.arange(0.5, len(self.Ns)), ticklabels=map(str, input(self.Ns)))
188
```

```
189
                                           fig.colorbar(pos, ax=ax)
190
 191
                                           # print(W_c_fit)
                                          ax.scatter(W_c_fit[:,0], W_c_fit[:,1]-0.5, s=100, c="w", marker='^', label='$W_c$', label='$W_
192
                    edgecolors="k")
193
                                          plt.ylabel("System size L")
                                           plt.xlabel("Predicted disorder strength $W_{pred}$")
194
 195
                                           ax.legend()
                                          plt.tight_layout()
196
197
                                          plt.savefig('results/Wc/n'+str(n)+'_Wc_N_dependency.pdf')
                                          plt.close()
198
199
200
                    def plot_wc_fit(self, N, popt, state_prediction):
201
                               fig, ax1 = plt.subplots()
202
                               ax1 = plt.scatter(self.Ws, state_prediction)
203
                               ax1 = plt.plot(self.Ws, logistic(self.Ws, *popt), 'k')
204
205
                               plt.title('Phase prediction N = ' + str(N) + , W_c = ' + (0:.3g)'.format(popt[0]))
206
207
                               plt.ylabel('Probability of localized phase')
                               plt.xlabel('$W_{max}$')
208
                               plt.legend(['Logistic fit', 'Predicted phase'], loc='upper left')
209
                               plt.savefig('results/N' + str(N) + '_predict_wc.pdf')
210
211
                               pass
212
213
         if __name__ == "__main__":
214
                    Ns = [9, 10]
215
                    Ws = np.arange(0., 4.0, 0.05)
216
217
                    n_max = 7
                    repetitions = 5
218
                    heat_map_plotter = HeatMapPlotter(Ns, Ws, n_max, repetitions)
                    heat_map_plotter.plot_wc_heatmap_n()
220
221
                    print("done")
222
                    heat_map_plotter.plot_wc_heatmap_N()
                  print("done")
223
```