Deep learning study on the Dirac eigenvalue spectrum of staggered quarks

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We study the chirality of staggered quarks on the Dirac eigenvalue spectrum using deep learning (DL) techniques. The Kluberg-Stern method to construct staggered bilinear operators conserves continuum property such as recursion relations, uniqueness of chirality, and Ward identities, which leads to a unique and characteristic pattern (we call it “leakage pattern (LP)”) in the matrix elements of the chirality operator sandwiched between two quark eigenstates of staggered Dirac operator. DL analysis gives 99.4(2)% accuracy on normal gauge configurations and 0.998 AUC (Area Under ROC Curve) for classifying non-zero mode octets in the Dirac eigenvalue spectrum. It confirms that the leakage pattern is universal on normal gauge configurations. The multi-layer perceptron (MLP) method turns out to be the best DL model for our study on the LP.

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1. Introduction

Here, we present technical details on how to do a deep learning analysis on low non-zero modes for staggered Dirac operators. This work is a follow-up paper on our previous papers: Ref. [1–4].

2. Staggered Dirac eigenstate

Here, we adopt the same notation as in Ref. [4]. Staggered Dirac operator, $D_s$, is anti-Hermitian ($D_s^\dagger = -D_s$) and its eigenvalues are pure imaginary.

$$D_s |f_i\rangle = i\lambda_i |f_i\rangle$$

(1)

where $\lambda_i$ is a real eigenvalue. Here, the eigenstates are normalized such that

$$h_{f_i f_i} = \delta_{i k}.$$

Under conserved $U(1)_A$ axial transformation $\Gamma_e$, Dirac operator transforms into its Hermitian conjugate,

$$\Gamma_e D_s \Gamma_e = D_s^\dagger = -D_s.$$

(2)

And an eigenstate $|f_i\rangle$ is related with its parity partner $|f_{-i}\rangle$ as follows,

$$D_s \Gamma_e |f_i\rangle = -i\lambda_i \Gamma_e |f_i\rangle,$$

$$D_s |f_{-i}\rangle = -i\lambda_i |f_{-i}\rangle.$$

(3)

(4)

Hence,

$$\Gamma_e |f_i\rangle = e^{i\theta} |f_{-i}\rangle \quad \leftrightarrow \quad \Gamma_e |f_{-i}\rangle = e^{-i\theta} |f_{+i}\rangle.$$

(5)

The eigenvectors can be classified into two categories: one is a set of zero modes and the other is a set of non-zero modes. At the finite lattice spacing, there is no exact zero mode ($\lambda = 0$) with staggered fermions [5]. Those eigenmodes which correspond to zero modes in the continuum, are named “would-be zero modes”. For more details, refer to Ref. [4].

3. Kluberg-Stern method

In general, there are two independent methods to transcribe a continuum operator to its lattice version using staggered fermions: one is the Golterman (Gol) method [6] and the other is the Kluberg-Stern (Klu) method [7]. The Klu method has a number of advantages in the study on staggered Dirac eigenvalue spectrum, compared with the Gol method, since it satisfies (a) recursion relations, (b) uniqueness of chirality, and (c) Ward identities.

In the Klu method, a quark bilinear operator with spin $S$ and taste $T$ is

$$O_{S\times T}(x) \equiv \tilde{\chi}(x_A) [y_S \otimes \xi_T]_{AB} \chi(x_B) = \tilde{\chi}(x_A) \overline{(y_S \otimes \xi_T)}_{AB} U_{AB} \chi(x_B)$$

(6)

where $\chi$ is a staggered quark field, $x_A = 2x + A$, $x$ is the coordinate of a hypercube, and $A_{\mu}, B_{\mu} \in \{0, 1\}$. Here, $\overline{(y_S \otimes \xi_T)}_{AB} = \frac{1}{4} \text{Tr} \left( Y_A y_S Y_B y_T^\dagger \right)$. The $U_{AB}$ is inserted to make the operator gauge-invariant,

$$U_{AB} \equiv \mathbb{P}_{SU(3)} \left[ \sum_{p \in C} V(x_A, x_{p_1}) V(x_{p_1}, x_{p_2}) \cdots V(x_{p_n}, x_B) \right]$$

(7)
where $P_{SU(3)}$ is the $SU(3)$ projection, $C$ denotes the complete set of shortest paths from $x_A$ to $x_B$, and $V(x, y)$ is HYP-smearred fat link [8]. The matrix elements for the chirality operator $[\gamma_5 \otimes 1]$ and the shift operator $[1 \otimes 5]$ are

$$
\Gamma^i_{5,k} = \Gamma_5(\lambda_i, \lambda_k) = \langle f_i | [\gamma_5 \otimes 1] | f_k \rangle \equiv \sum_{x,A,B} [f_i(x_A)]^\dagger (\gamma_5 \otimes 1)_{A,B} U_{AB} f_k(x_B),
$$

(8)

$$
\Xi^i_{5,k} = \Xi_5(\lambda_i, \lambda_k) = \langle f_i | [1 \otimes 5] | f_k \rangle \equiv \sum_{x,A,B} [f_i(x_A)]^\dagger (1 \otimes 5)_{A,B} U_{AB} f_k(x_B).
$$

(9)

Unlike the Gol operators, the Klu operators satisfy the following:

(a) recursion relations

$$
[\gamma_5 \otimes 1]^{2n+1} = [\gamma_5 \otimes 1], \quad [\gamma_5 \otimes 1]^{2n} = [1 \otimes 1],
$$

(10)

(b) uniqueness of chirality, which is a natural consequence of the above recursion relations,

(c) Ward identity,

$$
\Gamma_e = [\gamma_5 \otimes 5] = [\gamma_5 \otimes 1] [1 \otimes 5] = [1 \otimes 5] [\gamma_5 \otimes 1],
$$

(11)

$$
\Gamma_e [\gamma_5 \otimes 1] = [\gamma_5 \otimes 1] \Gamma_e = [1 \otimes 5],
$$

(12)

$$
\Gamma_e [1 \otimes 5] = [1 \otimes 5] \Gamma_e = [\gamma_5 \otimes 1].
$$

(13)

Using the Ward identity in Eqs. (12) and (13), we can derive the following results:

$$
|\Gamma^i_{5,k}| = |\Xi^{-i,-k}_{5} | = |\Xi^{-i,-k}_{5} | = |\Gamma^{i,-k}_{5} | = |\Xi^{-k,i}_{5} | = |\Xi^{k,-i}_{5} | = |\Gamma^{k,-i}_{5} |.
$$

(14)

For more details on this, refer to Ref. [4]. Here, we choose $|\Gamma^i_{5,k}|$ to study leakage patterns, because $|\Xi^{i,-k}_{5} |$ is just a mirror image of $|\Gamma^{i,-k}_{5} |$.

4. Quartet structure

Let us describe the serial notation for Dirac eigenmodes. So far we have used a serial index $i$ for an eigenvalue $\lambda_i$ which is assigned in ascending order such that $i < j$ if $\lambda_i < \lambda_j$. It is convenient since we can use a compact notation such as $\Gamma^i_{5,k}$. Since the $SU(4)$ taste symmetry breaking effect is so small at $a \neq 0$, we can also introduce the quartet notation which reflects on the taste symmetry as in Figure 1. Here, we adopt the same quartet notation as in Ref. [4]:

$$
D_s |f_{j,m}\rangle = i\lambda_{j,m} |f_{j,m}\rangle \equiv i\lambda_{j,m} |j,m\rangle
$$

(15)

where $j$ is a quartet index and $m$ is the index of a quartet component ($m = 1, 2, 3, 4$). In the continuum ($a = 0$), $\lambda_{j,m} = \lambda_{j,n}$ for $m \neq n$ thanks to the $SU(4)$ taste symmetry. But on the lattice ($a > 0$), $\lambda_{j,m} \neq \lambda_{j,n}$ for $m \neq n$ as one can see in Fig. 1. The ono-to-one mapping between a serial index $i$ and a quartet index ($j, m$) is given in Table 1 for topological charge $Q = \pm 1$. Here, note that $\lambda_{2n-1} = -\lambda_{2n}$ for $n > 0$, and $\lambda_{j,m} = -\lambda_{-j,m}$. We find a minute deviation between eigenvalues within a quartet, $\lambda_{j,m} \neq \lambda_{j,n}$ for $m \neq n$ since the taste symmetry breaking is so tiny. We find that the size of taste symmetry breaking is only 7% even for the smallest non-zero quartet as one can see in Fig. 2.

\footnote{Here, we assume that we use the HYP staggered action [8] for valence quarks.}
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Figure 1: Staggered Dirac eigenvalue spectrum.

Table 1: One-to-one correspondence between serial index $i$ and quartet index $(j, m)$ for $Q = \pm 1$. Here, Z (NZ) represents zero (non-zero) modes.

Table 2: Details on gauge configuration [12].
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5. Leakage pattern

First, let us consider the continuum case ($a = 0$). A shift operator, which is also a generator for the $SU(4)$ taste symmetry, is

$$\Xi_F = [1 \otimes \xi_F]$$

(16)

where $\xi_F \in \{\xi_5, \xi_\mu, \xi_\mu^5, \xi_\mu^5\}$ (for $\mu \neq \nu$). Here, $\xi_\mu$ satisfies the Clifford algebra $\{\xi_\mu, \xi_\nu\} = 2\delta_{\mu\nu}$. Since the taste symmetry is conserved, $\Xi_F$ commutes with staggered Dirac operator,

$$0 = \langle f_i | [D_s, \Xi_F] | f_k \rangle = i(\lambda_i - \lambda_k) |\Xi_F^{i,k} | .$$

(17)

If $\lambda_i \neq \lambda_k$, $|\Xi_F^{i,k} | = 0$. If $\lambda_i = \lambda_k$, $|\Xi_F^{i,k} | \neq 0$ is possible. $|\Xi_F^{i,k} |^2$ is a probability that one eigenstate transforms into another by $\Xi_F$. Hence, we call $|\Xi_F^{i,k} |$ the leakage parameter for the $\Xi_F$ operator. Here, note that the leakage by $\Xi_F$ can occur only within a quartet. And among the set of $\{\Xi_F \}$, $\Xi_5$ is especially important because it is related to $\Gamma_5$ by the Ward identity in Eqs. (12) and (13). Hence, $|\Gamma_5^{jm,-jn} |$ is just a mirror image of $|\Xi_5^{jm,jn} |$ for the non-zero modes ($j \neq 0$). Therefore, in the continuum ($a = 0$),

$$\Gamma_5^{jm,\ell n} = \langle j, m | \gamma_5 \otimes 1 | \ell, n \rangle = \begin{cases} \delta_{j\ell} \delta_{mn} & \text{for zero modes,} \\ \delta_{j,-\ell} \delta_{mn} & \text{for non-zero modes.} \end{cases}$$

(18)

On the lattice ($a \neq 0$),

$$\Gamma_5^{jm,\ell n} = \langle j, m | \gamma_5 \otimes 1 | \ell, n \rangle \simeq \begin{cases} \delta_{j\ell} \delta_{mn} & \text{for zero modes,} \\ \delta_{j,-\ell} c_{mn} & \text{for non-zero modes,} \end{cases}$$

(19)

where $c_{mn} \neq 0$ due to the taste symmetry breaking. Hence, $|\Gamma_5^{i,k} |$ has a specific pattern which we call “leakage pattern”. For zero modes it is diagonal within a quartet. For non-zero modes it is off-diagonal and non-trivial only between a quartet and its parity partner (we call them “octet” collectively because the parity partner can be determined from a quartet completely). Therefore, zero modes and non-zero modes can be distinguished by the leakage pattern. For example, we show

![Figure 3: An example of leakage patterns.](image)
6. Deep learning study

Here, we want to address a question: Is the leakage pattern valid and universal over the entire gauge configurations? Since there are too many zero modes and non-zero modes with various $Q$ values, it is practically impossible to check the leakage pattern over the entire gauge configurations by analytic methods or by visual examination. Here, we introduce the deep learning (DL) techniques to solve the problem, which recognize a specific pattern embedded in the data if it exists, even though the data look completely random in the eye sight. The DL method allows us to check the leakage pattern for non-zero modes through the whole gauge configurations.

For the numerical study on the DL method, we use the lowest $200 \times 200$ matrix elements of $|\Gamma_{5}^{i,k}|$ ($0 \leq i, k < 200$) on the lattice specified in Table 2.

6.1 Data sampling

Here, our task belongs to a classification problem in DL. We select $15 \times 15$ sub-matrices randomly along the diagonal ($i = k$) line of the $200 \times 200$ matrix of $|\Gamma_{5}^{i,k}|$. There is no overlap
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| DL model structure | our specific choice |
|--------------------|---------------------|
| Loss function      | categorical cross-entropy |
| Hidden layer’s activation function | ReLU |
| Output layer’s activation function | Softmax |
| Optimization algorithm for loss function | Adam |
| Neural network layer | MLP, CNN, CNN + MLP |
| Auto-hyperparameter tuner | Keras tuner (random search) |

Table 3: DL model structure

among the sub-matrices so that the probability distribution should be independent and identical. The sub-matrices are classified by the location of the border line between two adjacent octets as in Fig. 4, where an octet is composed of a quartet and its parity partner of non-zero modes. We choose the size of the sub-matrices as $15 \times 15$ from the following guidelines.

- We want to maximize statistics or the number of samples, which requires minimizing the size of the sub-matrices.
- We want to make the whole samples contain at least one complete set of an octet.

The total number of different classes or labels (in terms of DL) is eight, which is equal to the number of components in an octet. We define the class ID, $C$ as follows,

$$C = (s - i_b) \mod 8$$ (20)

where $s$ is a serial index of the left-upper corner in a sub-matrix (e.g. $i = k = 34$, the red blob in Fig. 4b), and $i_b$ is a serial index for the border line (e.g. $i = 40$ and $k = 40$, black lines in Fig. 4b). Examples for the eight classes are presented in Fig. 10 of Ref. [4].

6.2 Deep learning analysis

Details on the structure of our DL model are summarized in Table 3. Here, ReLU, MLP, and CNN represents rectified linear unit, multi layer Perceptron, and convolutional neural network, respectively. In order to perform the DL study, we need three types of data sets by construction: one for training, another for validation, and the other for test. Details on the DL data sets are given in Table 4a.

When we perform the auto hyperparameter tuning, we find out that our best performance model is MLP. The hyperparameters for MLP are given in Table 4b. There are about ten samples ($15 \times 15$ sub-matrices) per gauge configuration as shown in Table 4b. We adopt the accuracy metric to measure the performance of DL model. The accuracy is defined as a ratio $TC/TND$, where $TC$ represents the number of truly classified samples (DL prediction is equal to true class), and $TND$ represents the total number of the data samples.

In the test data set, we find that there are two independent sets of gauge configurations which are distinct by the accuracy: one set of five gauge configurations has accuracy less than 0.5, and the other set of 137 gauge configurations has accuracy of about 0.9. The former are called “abnormal set” and the latter “normal set”. The abnormal gauge configurations are entirely caused by the ghost
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Table 4: DL data sets and the best performance model. (a) The ngc (nds) represents the number of gauge configurations (the number of data samples). (b) The NN denotes neural network and the act.f. activation function.

| data set     | ngc | nds |
|--------------|-----|-----|
| training     | 120 | 1223|
| validation   | 30  | 308 |
| test         | 142 | 1441|

| layer         | NN type | # of units | act.f. |
|---------------|----------|------------|--------|
| input         | -        | 225        | -      |
| 1st hidden    | MLP      | 160        | ReLU   |
| 2nd hidden    | MLP      | 1210       | ReLU   |
| 3rd hidden    | MLP      | 1490       | ReLU   |
| output        | MLP      | 8          | softmax|

The hyperparameters of the best performance model.

(a) DL data sets

The eigenvalue problem of Lanczos algorithm [14]. Before we begin our numerical study on the DL method, we have excluded by hand those abnormal gauge configurations on which the topological charge $Q$ measured by the index theorem is different from that of the direct measurement. In fact, the DL method is the best to identify the abnormal gauge configurations such that we can filter them out.

Let us consider the accuracy $A_i$ measured on a gauge configuration with an index $i$. Then we can consider a set of the accuracies $S_a = \{A_i\}$ for $1 \leq i \leq N$. We also have a set of the number of samples $M_i$ for each gauge configuration $S_n = \{M_i\}$. Then we can consider an weighted average of the accuracy $\bar{A}$ over the gauge configurations:

$$\bar{A} = \frac{\sum_i M_i A_i}{\sum_i M_i}$$

$$\sigma^2_{\bar{A}} = \frac{\sum_i M_i (A_i - \bar{A})^2}{(N - 1) \sum_i M_i}$$

(21)

where $\sigma_{\bar{A}}$ is the statistical error for $\bar{A}$. In Table 5, we present results for $\bar{A}$ measured over the entire set (mixed = normal + abnormal), a subset of the abnormal gauge configurations (abnormal), and a subset of the normal gauge configurations (normal). The accuracy for the abnormal subset is so small that the DL method can identify it easily. We find that the large error for the mixed set comes from that for the abnormal subset. We also find that the extremely high accuracy for the normal subset reflects on the fact that the DL method succeeds in classifying the samples of the normal subset, which leads to the key point that if we select the majority of the DL prediction on each gauge configuration, we can distinguish all the non-zero mode octets perfectly over the entire normal subset.

We apply the DL method on the non-zero modes, because they are more appropriate for the DL analysis due to the following reasons: 1) abundant data, 2) complex structure, and 3) randomness in leakage pattern. The DL analysis gives a border line of the non-zero modes octets robustly. The leakage pattern for zero modes combined with the DL method is as robust as the SF method in determining the topological charge $Q$. However the key point is that the computational cost for the DL method is much cheaper than the SF method by at least a factor of 1000.

6.3 ROC curve

In the previous subsection, we use the accuracy metric for the DL analysis. In general, for a skewed probability distribution of classes, the accuracy is not such a good metric that it might give a
misleading information on the DL model performance [15]. Since the class distribution of our data samples is not skewed but uniform, the accuracy is a good metric for our DL analysis. There are tens of alternative metrics in the DL market. Among them, the AUC (area under ROC curve) [16] is one of the most popular metrics, and can be used instead of accuracy even if the class distribution is skewed. Here, the ROC stands for receiver operating characteristic. In order to crosscheck results from the accuracy metric, we adopt the AUC metric, even though we expect that both metrics will give the same answer for our data sets in the end of day.

A ROC curve is defined in binary class (positive and negative classes). As in Fig. 5, y-axis of ROC curve is true positive rate (TPR) and x-axis is false positive rate (FPR).

\[
\text{TPR} = \frac{TP}{P} \quad \text{FPR} = \frac{FP}{N}
\]  \hspace{1cm} (22)

Here, \(TP\) (FP) represents the number of the samples for which the DL prediction is positive and their actual class is positive (negative). The \(P\) (N) is the total number of samples in the actual positive (negative) class. Here, the DL binary model gives a probability that each sample belongs to the positive class. The DL model prediction for each sample is determined by a threshold applied to the probability to accept it (positive prediction) or reject it (negative prediction). The number of samples in the positive prediction changes according to the threshold value, and so do the \(TP\) and \(FP\), which lead to a corresponding change in TPR and FPR. Hence, we may obtain the ROC curve for the DL binary model by running the threshold from 0 to 1.

If we set the threshold to zero, all the predictions by the DL binary model are positive, which results in a trivial case that \(TP = P\) and \(FP = N\), which corresponds to \((1, 1)\) in the ROC curve. If the threshold is set to 1, all the predictions are negative, which makes \(TP = 0\) and \(FP = 0\), which corresponds to \((0, 0)\) in the ROC curve. Hence, the ROC curves, in general, share the two points \((1, 1)\) and \((0, 0)\) in common. If the DL predictions are perfect (i.e. \(TP = P\) and \(FP = 0\)), then the ROC curve passes through \((1, 0)\). AUC is the area under the ROC curve, and so AUC for a perfect DL model is 1. If the DL predictions are random (i.e. \(TP = 0.5P\) and \(FP = 0.5N\)), the ROC curve corresponds to the blue dashed line in Fig. 5, which makes AUC=0.5. Hence, AUC is an alternative good metric for the DL model performance. If the DL model prediction gets better, its AUC becomes closer to 1. If the DL model prediction gets worse, its AUC becomes closer to 0.5.

Since our DL model belongs to the multiclass classification, we need to extend the binary-class picture of the ROC curve to the multiclass picture. We use the one-versus-rest method to extend the ROC curve to the multiclass classification [16, 17]. The one-versus-rest method converts multiclass classification into binary class classification regarding \(c_i\) as a positive class and the rest \((\{c_j| j \neq i\})\) as a negative class. In Fig. 5, we present the ROC curve and its AUC(\(c_i\)) for each class \(c_i\) in our

| configuration | # conf. | \(\hat{A}\) [%] |
|---------------|---------|---------------|
| mixed         | 142     | 96.5(13)      |
| abnormal      | 5       | 17.3(78)      |
| normal        | 137     | 99.4(2)       |

Table 5: Results for the accuracy \(\hat{A}\) for mixed, abnormal, and normal set of gauge configurations in the test data set.
our best performance model (MLP) gives $AUC_{tot} = 0.998 \approx 1$ for the normal set. The $AUC_{tot}$ value implies that our best DL model works almost perfectly. Results for the AUC metric are highly consistent with those for the accuracy metric in the previous subsection. Its physical meaning is that the leakage pattern for zero and non-zero modes is universal over the normal gauge configurations.

7. Conclusions

Thanks to the $U(1)_{A}$ symmetry and the $SU(4)$ taste symmetry, the leakage pattern (LP) for zero modes is quite different from that for non-zero modes. We use the deep learning (DL) method to verify that the LPs for the chirality operator are universal over the normal gauge configurations, which we cannot prove analytically nor visually. We find that, using the LP method combined with the DL method, we can determine topological charge as robustly as the spectral flow (SF) method. Since the computational cost for the LP/DL method is much cheaper than that for SF at least by a factor of 1000, the LP/DL method is highly promising.

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