### **PQC** Function Evaluation

Carnegie Vacation Scholarship

David Amorim

Week 4 (22/07/2024 - 26/07/2024)

#### Aims for the Week

The following aims were set at the last meeting (22/07/2024):

### 1. Change Input Layer Structure

Improve the connectivity of input layers. Each input qubit should ideally control each target qubit at some point in the network.

#### 2. Fix Parameters

Add the option to keep parameters fixed for each type of network layer.

### 3. Improve Loss Function

Develop a distance measure taking into account digital encoding. Either incorporate this into weights for an existing loss function or define a new loss function on this basis.

David Amorim PQC Function Evaluation 29/07/2024 2 / 25

### Glossary

Acronym	Meaning
CL	convolutional layer
AA-CL	all-to-all convolutional layer
NN-CL	neighbour-to-neighbour convolutional layer
IL	input layer
SAM	sign-adjusted mismatch

Table 1: Acronyms and short-hands used in the following.

Variable	Meaning
$\overline{n}$	input register size
m	target register size
L	number of network layers

Table 2: Variables used in the following.

- 1 Changing Input Layer Structure
- Pixing Parameters
- Improving the Loss Function
- A Results
- 6 Next Steps

David Amorim PQC Function Evaluation 29/07/2024 4/25

- Previously, the jth input qubit controlled an operation on the jth target qubit (with wrap-around for n>m)
- An optional shift parameter, s, has now been added so that the jth input qubit controls an operation on the j+sth target qubit
- This shift parameter is incremented for each successive IL
- The QCNN is padded with additional ILs to ensure that the number of ILs is  $\geq m$
- Thus, each input qubit now controls an operation on each target qubit at some point in the QCNN
- Note that ILs still alternate between control states 0 and 1

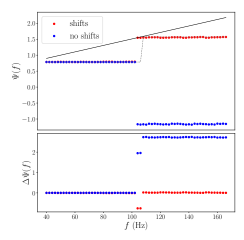


Figure 1: Effects of shifted ILs for  $\Psi(f) \sim f$  and m=3 (L=6, 600 epochs, SAM)

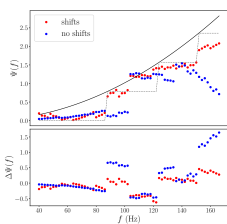


Figure 2: Effects of shifted ILs for  $\Psi(f)\sim f^2$  and m=3 (L=6, 600 epochs, SAM)

6/25

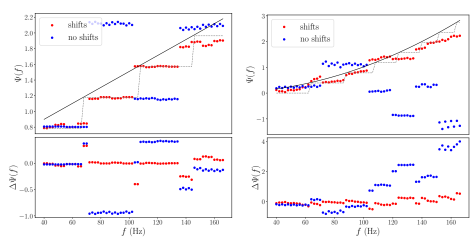


Figure 3: Effects of shifted ILs for  $\Psi(f)\sim f$  and m=4 (L=6, 600 epochs, SAM)

Figure 4: Effects of shifted ILs for  $\Psi(f)\sim f^2$  and m=4 (L=6, 600 epochs, SAM)

7/25

- The data for 'no shifts' was obtained by setting s=0 for all ILs instead of incrementing s
- This should be equivalent to last week's circuit structure
- However, the 'no shifts' results are significantly worse than the results shown last week (???)
- Thus, the improvements due to the new IL structure are somewhat exaggerated
- Nonetheless, increased IL connectivity clearly leads to improved performance

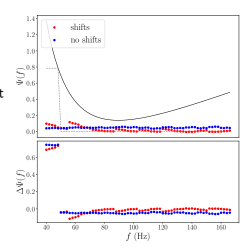


Figure 5: Effects of shifted ILs for  $\Psi_{\rm Hayes2023}$  and m=3 (L=6, 600 epochs, SAM)

8 / 25

- Changing Input Layer Structure
- Pixing Parameters
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- A Results
- 6 Next Steps

David Amorim PQC Function Evaluation 29/07/2024 9/25

### Fixing Parameters

- Implemented the option to fix parameters for each layer type
- This means that each instance of a layer type (IL, AA-CL, NN-CL) uses the same set of parameters
- $\bullet$  This significantly reduces the number of trainable parameters at large L
- Surprisingly, reducing the parameter space produces no noticeable speed-up (so-called qiskit primitives, i.e. the sampler, take up roughly 95% of the computational time)

David Amorim PQC Function Evaluation 29/07/2024 10/25

# Fixing Parameters

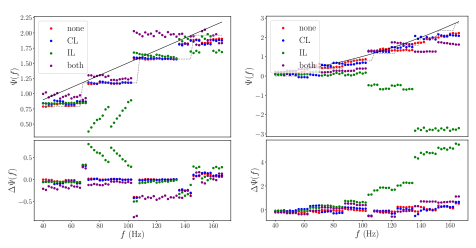


Figure 6: Effects of fixing parameters ILs for  $\Psi(f) \sim f$  and m=4 (L=6, 600 epochs, SAM)

Figure 7: Effects of fixing parameters for  $\Psi(f)\sim f^2$  and m=4 (L=6, 600 epochs, SAM)

### Fixing Parameters

- Legend for the plots on the previous slide:
  - 'none' : no parameters fixed
  - 'CL': only CL parameters fixed
  - 'IL': only IL parameters fixed
  - 'both': all parameters fixed
- Evidently, keeping parameters fixed leads to (slightly, for 'CL', or drastically, for 'IL' and 'both') worse performance
- This is likely due to a reduction of the search space
- Note that 'IL' as well as 'both' lead to incomplete phase extraction (formalise this concept!) and hence somewhat meaningless results
- Thus, especially taking into account the equivalent computational times, not keeping parameters fixed yields better results
- These results suggest a particular importance of ILs compared to CLs

- Changing Input Layer Structure
- Pixing Parameters
- 3 Improving the Loss Function
- 4 Results
- 6 Next Steps

David Amorim PQC Function Evaluation 29/07/2024 13/25

### **Preliminary Definitions**

- Consider a computational basis state,  $|k\rangle$ , of the combined input-register-target-register system
- The state  $|k\rangle$  is associated with a bit string  $k = \{0,1\}^{n+m}$
- Denote by  $[k]_n$  and  $[k]_m$  the bit strings of length n and m, respectively, associated with each of the registers and write their concatenation as

$$k \equiv [k]_n \diamond [k]_m \tag{1}$$

A general state of the two-register system is then written as

$$|z\rangle = \sum_{k=0}^{2^{n+m}-1} z_k |k\rangle \tag{2}$$

and referred to via its coefficients  $z_k$ 



David Amorim PQC Function Evaluation 29/07/2024 14 / 25

### **Preliminary Definitions**

When training in superposition, the desired state of the system is

$$|y\rangle = \sum_{j=0}^{2^{n}-1} \frac{1}{\sqrt{2^{n}}} |j\rangle_{i} |\Psi(j)\rangle_{t},$$
 (3)

target registers, respectively

ullet This state |y
angle can be written in terms of the combined basis  $\{|k
angle\}$  via

where the subscripts i and t indicate basis states of the input and

$$y_k = \begin{cases} \frac{1}{\sqrt{2^n}} & \text{if } k = [k]_n \diamond \Psi([k]_n) \\ 0 & \text{else} \end{cases}$$
 (4)

• Further denote the output state produced by the QCNN by  $|x\rangle$ , with associated coefficients  $x_k$ 

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### SAM and Beyond

Recall the definition of SAM:

$$\mathsf{SAM}(\ket{x},\ket{y}) = \left|1 - \sum_{k} x_{k} y_{k}\right| \tag{5}$$

By construction, this is closely related to the mismatch

$$M(|x\rangle, |y\rangle) = 1 - |\langle x|y\rangle| \tag{6}$$

• While effective, SAM's fundamental flaw is that it does not directly take into account the amplitudes  $x_k$  for k where  $y_k=0$  (i.e. where  $[k]_m \neq \Psi([k]_n)$ 



David Amorim PQC Function Evaluation 29/07/2024 16/25

# SAM and Beyond

• Consider k=a and k=b with  $[k]_m \neq \Psi([k]_n)$  for both and

$$\left| [a]_m - \Psi([a]_n) \right| \ge \left| [b]_m - \Psi([b]_n) \right| \tag{7}$$

- To improve performance, the loss function should punish a non-zero  $x_a$  more than a non-zero  $x_b$  which is not the case for SAM
- This could be achieved via a weighted mismatch,

$$WIM(|x\rangle, |y\rangle) = \left|1 - \sum_{k} \tilde{w}_{k} x_{k} y_{k}\right|, \tag{8}$$

where  $ilde{w}_k \in \mathbb{R}_+$  are appropriate weights

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David Amorim PQC Function Evaluation 29/07/2024 17 / 25

### SAM and Beyond

It was discussed at the last meeting to base the weights on

$$w_k = \sum_{\substack{l=0, \\ l \neq [k]_m}}^{2^m - 1} \left| x_{[k]_n \diamond l} \right| \left| l - \Psi([k]_n) \right|, \tag{9}$$

with the  $\tilde{w}_k$  obtained from the  $w_k$  via normalisation and smoothing  $(\tilde{w}_k \sim e^{\tau w_k})$ 

- Implementing this proved to be uneffective, with no improvement on SAM
- This raises broader questions about the feasibility of WIM

David Amorim PQC Function Evaluation 29/07/2024 18/25

#### The Limits of WIM

- SAM is effective as it learns to maximise the  $x_k$  for k with  $y_k \neq 0$  without regard for the remaining  $x_k$
- It effectively optimises over a reduced set of states
- This is, presumably, why it outperforms more global loss functions, which directly take into account all coefficients, like L<sub>1</sub> loss
- Thus, it seems that SAM's 'fundamental flaw', of disregarding most  $x_k$ , is really the basis of its success
- Adding weights to SAM (WIM) likely is insufficient to alter its fundamental dynamic (or will alter it detrimentally)
- Therefore, WIM will be abandoned for the foreseeable future

David Amorim PQC Function Evaluation 29/07/2024 19/25

# Introducing WILL

- To improve on SAM, it could be beneficial to return to a loss function, which more directly takes into account all  $x_k$  (e.g.  $L_1$  loss)
- More generally, define L<sub>p</sub> loss as

$$\mathsf{LL}_{\mathsf{p}}(|x\rangle, |y\rangle) = \left(\sum_{k} |x_{k} - y_{k}|^{p}\right)^{1/p},\tag{10}$$

for some  $p \ge 1$ 

- Note that for phase encoding, computational basis states are not equidistant: their distance depends on the value they encode on the input and target registers
- A weighted L<sub>p</sub> loss (WILL) can factor in an appropriate distance measure for the state space:

$$\mathsf{WILL}_{\mathsf{p},\mathsf{q}} = \left(\sum_{k} \left| x_k - y_k \right|^p \left| [k]_m - \Psi([k]_n) \right|^q \right)^{1/p} \tag{11}$$

David Amorim PQC Function Evaluation 29/07/2024 20 / 25

### Testing WILL

• SPEND SOME TIME PLAYING AROUND WITH p AND q!

David Amorim PQC Function Evaluation 29/07/2024 21/25

- Changing Input Layer Structure
- Pixing Parameters
- Improving the Loss Function
- 4 Results
- 6 Next Steps

David Amorim PQC Function Evaluation 29/07/2024 22 / 25

#### Results

Show phase encoding with improved methods show the full waveform [new frame]

David Amorim PQC Function Evaluation 29/07/2024 23/25

- Changing Input Layer Structure
- Pixing Parameters
- 3 Improving the Loss Function
- 4 Results
- 6 Next Steps

David Amorim PQC Function Evaluation 29/07/2024 24/25

### Next Steps

- formalise and investigate 'completeness of phase extraction'
- look into adding more ILs compared to CLs?



David Amorim PQC Function Evaluation 29/07/2024 25 / 25