
GraphQNTK: Quantum Neural Tangent Kernel for Graph Data

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Abstract

1 Graph Neural Networks (GNNs) and Graph Kernels (GKs) are two fundamental
2 tools used to analyze graph-structured data. Efforts have been recently made in de-
3 veloping a composite graph learning architecture combining the expressive power
4 of GNNs and the transparent trainability of GKs. However, learning efficiency on
5 these models should be carefully considered as the huge computation overhead.
6 Besides, their convolutional methods are often straightforward and introduce severe
7 loss of graph structure information. In this paper, we design a novel quantum graph
8 learning model to characterize the structural information while using quantum
9 parallelism to improve computing efficiency. Specifically, a quantum algorithm is
10 proposed to approximately estimate the neural tangent kernel of the underlying
11 graph neural network where a multi-head quantum attention mechanism is intro-
12 duced to properly incorporate semantic similarity information of nodes into the
13 model. We empirically show that our method achieves competitive performance
14 on several graph classification benchmarks, and theoretical analysis is provided to
15 demonstrate the superiority of our quantum algorithm.

16 1 Introduction

17 Fusing quantum computing and classic machine learning has become a promising subject of research.
18 Quantum-based algorithms have been proposed in recent years, from naive quantum non-parametric
19 machine learning [51, 35, 42, 31] to classic-quantum hybrid deep learning [7, 10, 45, 36, 14]. Despite
20 that quantum machine learning (QML) has shown its potential in many machine learning tasks,
21 quantum computing for graph learning is still in its early stage [60]. Inspired by the two popular
22 classes of methods for learning on graph data, i.e., Graph Neural Networks (GNNs) [19, 38, 20, 66]
23 and Graph Kernels (GKs) [22], several works attempt to build quantum graph learning architecture
24 that captures the structural information of graph data, such as Quantum Graph Neural Networks
25 (QGNNs) [63, 7, 11, 16, 1] and Quantum Graph Kernel Methods (QGKs) [55, 3, 24, 4]. A brief
26 review about quantum graph learning is illustrated in Fig. 1.

27 Some quantum subroutines for attribute encoding [5, 69] and structural encoding [63, 45] have been
28 developed to dissolve the characteristics of the graph into the quantum model. However, most present
29 quantum graph learning models are hybrid such that the expressive capability depends more on the
30 complexity of the classic modules [69]. It is difficult to characterize the structure information and
31 attribute information of the graph by the quantum components without the participation of classic
32 modules. Even worse, the frequent interactions between classical systems and quantum environments
33 generally incur additional overhead [54]. It is unclear whether the introduced quantum module
34 can improve the performance of the model as well as the training efficiency. Besides, most of
35 existing proposals for quantum machine learning for graphs lack a clear demonstration of a quantum
36 superiority for tasks on classical datasets.

37 Using quantum computing power to boost the trainability and expressive behaviour of classic machine
 38 learning models provides one of the most promising direction for quantum machine learning. It
 39 is demonstrated that the power of quantum computing could be used to find atypical but useful
 40 patterns that classical systems are not considered to be able to generate effectively [14, 23, 27], and
 41 accelerate the training process of existing classic models [35, 42, 70]. Several quantum algorithms
 42 [51, 44, 43] based on the HHL algorithm [21] show the exponential speedup compared with their
 43 classical counterpart, with a assumption that a quantum random access memory (QRAM) [34] is
 44 accessible. Recent literature employ quantum algorithms to efficiently train deep neural networks
 45 [70], reconstruct unsupervised clustering [33] and supervised kernel classifier [42]. It is hopeful that
 46 quantum computing could provide a new learning paradigm. In addition, simulations and physical
 47 experiments have proved the potential of using quantum algorithms to encode and process regular
 48 classical data such as text and image [59, 6].

49 Beyond vanilla GNNs and GKs, composite graph learning studies have emerged that combine the
 50 advantages of both areas [17, 49, 10, 18]. However, the computation overheads is extremely large
 51 due to either the dense gram matrix [17], or the large number of substructures to be compared after
 52 graph decomposition [10]. Prospectively, the barrier that conventional model is difficult to train
 53 and scale up is expected to be circumvented with the help of the uniqueness of quantum computing.
 54 Early research involves altering the amplitude of quantum basis states to accomplish a quantum logic
 55 operations [8], which is profitable from the huge quantum Hilbert space to encode the normalized
 56 data. Recently, simultaneous transformation of basic states in quantum superposition using quantum
 57 parallelism is regarded as a remarkable manifestation of quantum superiority, which is successfully
 58 implemented in classic machine learning to reduce the computational overheads [35, 36, 70]. These
 59 strategies could be helpful in the regime of training graph models with either the non-convex nature
 60 of the training procedure, or the poor scalability w.r.t. training size.

61 In this paper, we focus on quantum machine learning of graph-structured data with attributed nodes
 62 and binary edges. Inspired by recent quantum neural network methods [36, 70] that efficiently
 63 reconstruct the dynamics of classic neural networks using quantum computing techniques, a new
 64 quantum graph learning model is proposed which is analogue to train an infinite-width GNN with
 65 attention mechanism, where the number of heads goes to infinity. Our contributions are:

- 66 • For the first time, we show that the infinite-width GNN, namely graph tangent neural network
 67 [17] can be trained in a quantum reconstructive paradigm, by introducing a quantum aggregation
 68 transformation and a quantum kernel estimation adaptive to the different sizes of graph. This
 69 composite quantum graph learning architecture preserves the expressive power of GNNs and the
 70 transparent trainability of GKs as the composite graph learning model does.
- 71 • A quantum-friendly self-attention mechanism ([transformer](#)) is employed to incorporate semantic
 72 correlation information of all node pairs of the graph into the model, which can be naturally adopted
 73 by quantum algorithms with additional relaxation. To our best acknowledge, this is the first for
 74 leveraging attention mechanism in quantum graph learning. In contrast, the present quantum graph
 75 learning algorithms can only undertake propagation between the adjoining neighbors [45, 24, 63],
 76 and constrain themselves to specify identical priority to distinct neighbours [1, 11].
- 77 • We evaluate our quantum model on graph classification benchmarks. The results show that our
 78 method achieves competitive performance compared with the classic baselines including both
 79 GNNs and GKs. Besides, we give a theoretical analysis showing that our GraphQNTK reduces the
 80 running time complexity of graph kernel neural networks from $O(N^2)$ to $O(N)$.

81 2 Methodology

82 2.1 Preliminaries

83 We first briefly review the most common setting
 84 for GNNs and the corresponding neural tangent
 85 kernel (NTK), and by the way the notation is given.
 86 A graph $G = (V, E)$ is denoted by a collection of
 87 nodes V and edges E . Each node has a d -
 88 dimensional feature vector $\mathbf{h}_v \in \mathbb{R}^d$, $v \in V$, and
 89 $\mathbf{H} \in \mathbb{R}^{n \times d}$ is the feature matrix stacking all nodes
 90 features. For graph classification, we consider the

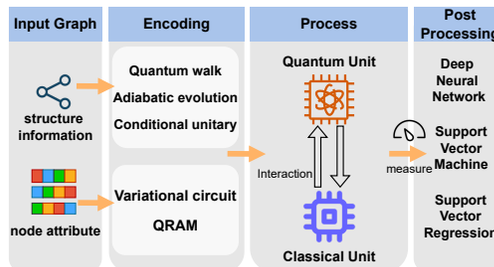


Figure 1: Overview of quantum graph learning.

91 dataset with a set of graphs $\{G_1, \dots, G_N\} \subseteq \mathcal{G}$ and their labels $\{y_1, \dots, y_N\} \subseteq \mathcal{Y}$. Our goal is to
 92 learn to predict labels of unseen graphs.

93 **The formulation of GNN** The differences of GNNs mainly depend on the different settings of
 94 message propagation process. Here we consider a simple message passing framework [19] and the
 95 propagation of the l -th ($l \in [L]$) layer is given as:

$$\hat{\mathbf{h}}_u^l := \sum_{v \in \mathcal{N}(u) \cup \{u\}} \mathbf{h}_v^{(l-1)}, \quad (1)$$

$$\mathbf{h}_u^l := \sqrt{\frac{c_\sigma}{d^L}} \sigma \left(\mathbf{W}_R^l \sqrt{\frac{c_\sigma}{d^{L-1}}} \sigma \left(\mathbf{W}_{R-1}^l \cdots \sqrt{\frac{c_\sigma}{d^1}} \cdot \sigma \left(\mathbf{W}_1^l \hat{\mathbf{h}}_u^l \right) \right) \right), \quad (2)$$

97 where $\mathcal{N}(u)$ denotes the neighbors of u , c_σ is the scaling factor, d^l is the output dimension of the l -th
 98 layer, σ is an element-wise activated function, and \mathbf{W}_R^l is learnable weights performing on the input
 99 for R times of the l -th layer (equivalent to R fully-connected layers without the bias term).

100 For graph classification, the output is a permutation invariance function acting on the collection of all
 101 node features in the last layer. The popular `sum_pooling` function is adopted: $\mathbf{h}_G = \sum_{u \in V} \mathbf{h}_u^L$.

102 **NTK of the infinite-width GNN.** Consider a training set $\{(\mathbf{x}_i, y_i)\}_{i=1}^N \subset \mathbb{R}^d \times \mathbb{R}$. When an over-
 103 parameterized fully connected network $f(\theta, \mathbf{x}) : \mathbb{R}^d \rightarrow \mathbb{R}$ whose width is allowed to go to infinity
 104 and parameters θ are randomly initialized and trained with gradient descent, the dynamics of the
 105 network is equivalent to the kernel regression [30]. This is the so called neural tangent kernel (NTK):

$$\mathbf{H}(t)_{ij} = \left\langle \frac{\partial f(\theta(t), x_i)}{\partial \theta}, \frac{\partial f(\theta(t), x_j)}{\partial \theta} \right\rangle, \quad (3)$$

106 which remains constant during training, i.e., $\mathbf{H}(t) = \mathbf{H}(0)$. And we replace $\mathbf{H}(t)$ with \mathbf{H} for
 107 convenience. The final prediction for a test datapoint \mathbf{x}_* is

$$f(\mathbf{x}_*) = \mathbf{k}_* \mathbf{H}^{-1} \mathbf{y}, \quad (4)$$

108 where $\mathbf{y}_i = y_i$ and $\mathbf{k}_* \in \mathbb{R}^N$ is the vector whose i -th element denotes the NTK value between \mathbf{x}_* and
 109 \mathbf{X}_* .

110 It is discovered that convolutional neural networks (CNNs) with infinite-width channels and infinite
 111 number of filters also have the same behaviour [2]. Inspired by this, Du et al. [17] adopts the designing
 112 strategy of NTK and leverages a GNN architecture to design new graph kernels, which is called
 113 graph neural tangent kernel (GNTK). The dynamics of training the GNTK is equivalent to train an
 114 infinitely-wide GNN initialized with random weights trained with gradient descent. Specifically,
 115 consider two input graph $G = (V, E)$ and $G' = (V', E')$ with $|V| = n$ and $|V'| = n'$, the GNTK
 116 $\Theta \in \mathbb{R}^{n \times n'}$ and the relative covariance matrix $\Sigma \in \mathbb{R}^{n \times n'}$ in the l -th layer of the feature aggregation
 117 phase as described in Eq. 1 after R fully-connected layers are given by

$$\begin{aligned} [\Sigma_0^l(G, G')]_{uu'} &= \sum_{v \in \mathcal{N}(u) \cup \{u\}} \sum_{v' \in \mathcal{N}(u') \cup \{u'\}} [\Sigma_R^{l-1}(G, G')]_{vv'}, \\ [\Theta_0^l(G, G')]_{uu'} &= \sum_{v \in \mathcal{N}(u) \cup \{u\}} \sum_{v' \in \mathcal{N}(u') \cup \{u'\}} [\Theta_R^{l-1}(G, G')]_{vv'}, \end{aligned} \quad (5)$$

118 which is an affine transformation of the input GNTK and covariance respectively where
 119 $[\Theta_R^0(G, G')]_{uu'}$ and $[\Sigma_R^0(G, G')]_{uu'}$ are both defined to be $\mathbf{h}_u^T \mathbf{h}_{u'}$. We replace them with
 120 $[\Theta^0(G, G')]_{uu'}$ and $[\Sigma^0(G, G')]_{uu'}$ respectively without ambiguity.

121 The successive fully-connected layers defined in Eq. 2 are used to update the node hidden feature
 122 after aggregation. Specifically, the GNTK of the fully-connected layer is recursively associated to
 123 that of the previous layer, and the transformation is given by

$$[\Sigma_r^l(G, G')]_{uu'} = \hat{\sigma}^{(r-1)}([\Sigma_{r-1}^l(G, G')]_{uu'}), r \in [R], \quad (6)$$

124 where $\hat{\sigma}^{(r)} : [-1, 1] \rightarrow \mathbb{R}$ denotes the the conjugate activation function corresponding to the activated
 125 function σ with centered Gaussian processes of covariance at the r -th fully-connected layer, as
 126 described in [15]. And the derivation of the covariance is

$$[\dot{\Sigma}_r^l(G, G')]_{uu'} = \hat{\sigma} \left(\hat{\sigma}^{(r-1)}([\Sigma_{r-1}^l(G, G')]_{uu'}) \right), \quad (7)$$

127 where $\hat{\sigma}$ denotes the derivative of σ . Given Eq. 6 and Eq. 7, the transformation of the GNTK for the
 128 feature update phase denoted by Eq. 2 is given by

$$[\Theta_R^l(G, G')]_{uu'} = \sum_{r=1}^R [\Sigma_0^l(G, G')]_{uu'} \left(\prod_{r'=r}^R [\dot{\Sigma}_0^l(G, G')]_{uu'} \right). \quad (8)$$

129 Therefore, computing each element of the GNTK (or covariance) matrix is only reliant on the element
 130 at the same place of the GNTK (or covariance) matrix in the previous fully-connected layer. The final
 131 GNTK corresponding the two input graphs G and G' determined by the sum_pooling function:

$$\Theta(G, G') = \sum_{u \in V, u' \in V'} [\Theta_R^L(G, G')]_{uu'}. \quad (9)$$

132 Intuitively, calculating each element of the GNTK of fully-connected layers could be accelerating
 133 by a proper quantum kernel estimation algorithm. However, it is indirect to realize an end-to-end
 134 speedup for GNTK since calculating the element of GNTK requires an affine transformation. To
 135 circumvent this barrier, we derive a unitary quantum aggregation transformation to bridge the gap
 136 between quantum kernel methods and estimation of GNTK.

137 2.2 QNTK with Attention Mechanism

138 Before giving the analytical quantum reconstruction
 139 of GNTK with multi-head attention mechanism, we first
 140 elaborate on how to integrate the transformer layer
 141 into the GNN as described in Sec. 2.1. The result-
 142 ing GraphQNTK can be efficiently reconstructed by quantum
 143 computing paradigm, which gives a quadratic speed-
 144 up over the classic estimation of GNTK. The mechanism
 145 to build the GNN and estimate the GNTK is shown in Fig. 2.

154 **GNN with multi-head attention.** The aggregation process
 155 of vanilla GCN [38] regards the contribution of each
 156 node's neighbor to the central
 157 node as equally important, which can be viewed as learning
 158 an averaged filter across the whole graph

159 [65], leading to a great loss of structure information. Besides, the aggregation only is performed
 160 within the adjoining neighbors under the assumption that the graph is homophilous. The method may
 161 fail to learn effective graph structures for message passing [12]. To capture the global node similarity
 162 semantics of the provided graph, numerous attempts that employ transformer for graph learning have
 163 been developed [26, 50, 52, 67]. Consider the input feature matrix $\mathbf{H}_{\text{in}}^l \in \mathbb{R}^{N \times s^l}$ where N denotes
 164 the number of samples and s^l is the dimension of feature at layer l before implementation of the
 165 transformer. The single transformer layer is to project the input $\mathbf{H}_{\text{in}}^l \in \mathbb{R}^{N \times s^l}$ by three matrices,
 166 i.e., $\mathbf{W}_Q^l \in \mathbb{R}^{s^l \times s_K^l}$, $\mathbf{W}_K^l \in \mathbb{R}^{s^l \times s_K^l}$ and $\mathbf{W}_V^l \in \mathbb{R}^{s^l \times s_V^l}$, to the corresponding representations
 167 $\mathbf{Q}^l, \mathbf{K}^l, \mathbf{V}^l$. The formulation is given as

$$\mathbf{Q}^l = \mathbf{H}_{\text{in}}^l \mathbf{W}_Q^l, \quad \mathbf{K}^l = \mathbf{H}_{\text{in}}^l \mathbf{W}_K^l, \quad \mathbf{V}^l = \mathbf{H}_{\text{in}}^l \mathbf{W}_V^l, \quad \hat{\mathbf{H}}^l = \zeta(\mathbf{G}^l) \mathbf{V}^l, \quad \mathbf{G}^l = \frac{\mathbf{Q}^l \mathbf{K}^{l \top}}{\sqrt{s_K^l}}, \quad (10)$$

169 where ζ denotes an element-wise activated function. The multi-head attention alternative is given by

$$\mathbf{H}_{\text{out}}^l = \left[\hat{\mathbf{H}}_{\text{head}_1}^l, \dots, \hat{\mathbf{H}}_{\text{head}_M}^l \right] \mathbf{W}_O^l, \quad (11)$$

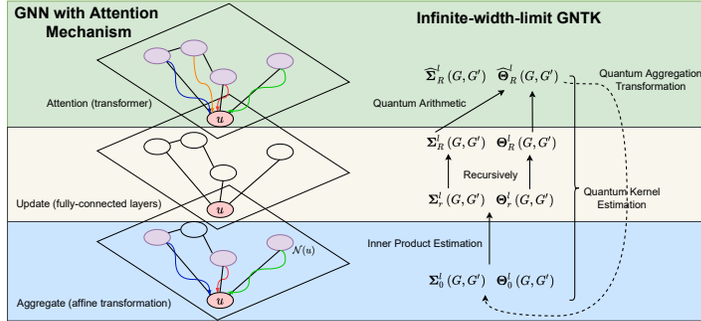


Figure 2: Framework for GNN with attention mechanism and its corresponding GNTK. The GNN comprises a message transmission process similar to the vanilla GCN but involves a transformer at the tail of the model (excluding the last layer), which characterizes the global semantic similarity between each pairs of nodes. The neighbor aggregation is kept since the two nodes connected by an edge often have stronger semantic relationship. The dynamics of the infinite-width-limit GNN is analogous to kernel methods and we reconstruct it by quantum algorithms to estimate the kernel.

170 where $\mathbf{W}_O^l \in \mathbb{R}^{(Ms_V^l) \times s^l}$ projects the $N \times Ms_V^l$ concatenated multi-head feature matrix back to
 171 $N \times s^l$ matrix.

172 Let \mathbf{Y} and Θ denote the neural network Gaussian Process Kernel (NNGP) [50] and NTK after
 173 the transformer layer, and let $\tilde{\mathbf{Y}}$ and $\tilde{\Theta}$ be the input NNGP and NTK before the transformer layer.
 174 Consider two input feature vector \mathbf{x} and \mathbf{x}' . When the output dimension of the transformer layer and
 175 the number of heads go to infinity, i.e., $s^l \rightarrow \infty, s_K^l \rightarrow \infty, s_V^l \rightarrow \infty, M \rightarrow \infty$, the output NTK is:

$$\begin{aligned}\Theta(\mathbf{x}, \mathbf{x}') &= 2\mathbf{Y}(\mathbf{x}, \mathbf{x}') + \zeta \left(\tilde{\mathbf{Y}}(\mathbf{x}, \mathbf{x}) \right) \tilde{\Theta}(\mathbf{x}, \mathbf{x}') \zeta \left(\tilde{\mathbf{Y}}(\mathbf{x}', \mathbf{x}') \right)^\top, \\ \mathbf{Y}(\mathbf{x}, \mathbf{x}') &= \zeta \left(\tilde{\mathbf{Y}}(\mathbf{x}, \mathbf{x}) \right) \tilde{\mathbf{Y}}(\mathbf{x}, \mathbf{x}') \zeta \left(\tilde{\mathbf{Y}}(\mathbf{x}', \mathbf{x}') \right)^\top,\end{aligned}\tag{12}$$

176 where the under the restriction that 1) \mathbf{W}_Q^l and \mathbf{W}_K^l share the same weights, and 2) scaling the dot
 177 products between \mathbf{Q}^l and \mathbf{K}^l by their dimension instead of the square root of the same quantity, i.e.,
 178 $\mathbf{G}^l = \frac{\mathbf{Q}^l \mathbf{K}^{l \top}}{s_K^l}$. The detailed proof can be found in [25].

179 To efficiently estimate the element of the NTK defined by the transformer layer using quantum
 180 parallel, we consider the identity activated function, i.e., $\zeta = I$, and slightly modify the Eq. 12 as

$$\begin{aligned}\Theta(\mathbf{x}, \mathbf{x}') &= 2\mathbf{Y}(\mathbf{x}, \mathbf{x}') + \tilde{\mathbf{T}}(\mathbf{x}, \mathbf{x}') \odot \Theta(\mathbf{x}, \mathbf{x}'), \\ \mathbf{Y}(\mathbf{x}, \mathbf{x}') &= \tilde{\mathbf{T}}_\zeta(\mathbf{x}, \mathbf{x}') \odot \tilde{\mathbf{Y}}(\mathbf{x}, \mathbf{x}'),\end{aligned}\tag{13}$$

181 where $\tilde{\mathbf{T}}(\mathbf{x}, \mathbf{x}')$ is the result of matrix multiplication between the column vector of the diagonal of
 182 $\tilde{\mathbf{Y}}(\mathbf{x}, \mathbf{x})$ and row vector of the diagonal of $\tilde{\mathbf{Y}}(\mathbf{x}', \mathbf{x}')$, and $\tilde{\mathbf{T}}_\zeta$ is the result of matrix multiplication
 183 between the diagonal of those two matrix after activated operation. It is reasonable to accept this
 184 modification since in the limit of infinite width neural network the output converges in distribution to
 185 a multivariate normal with a block diagonal covariance [50]. Notice that the difference between the
 186 definition of NNGP and the covariance of NTK is that the former denotes the expectation with respect
 187 to the output before the activated operation, while the later after the denotes the expectation with
 188 respect to the output after the activated operation [30]. Consequently, we consider that \mathbf{Y} is equal to
 189 the covariance of NTK within the transformer layer as the result of the identity activated function.

190 **GNTK with infinite-width-limit attention.** To appropriately incorporate semantic similarity in-
 191 formation of nodes into the model, a multi-head attention mechanism is implemented at the tail
 192 of the each GNN layer except the first and the last layer, and the calculation of the GNTK with
 193 infinite-width-limit attention is to insert an additional procedure after the fully connected layers. For
 194 the two input graphs G and G' , the formulation derived by Eq. 13 is given as

$$\begin{aligned}\left[\hat{\Theta}_R^l(G, G') \right]_{uu'} &= 2 \left[\Sigma_R^l(G, G') \right]_{uu'} + \left[\mathbf{T}^l(G, G') \right]_{uu'} \left[\Theta_R^l(G, G') \right]_{uu'}, \\ \left[\hat{\Sigma}_R^l(G, G') \right]_{uu'} &= \left[\mathbf{T}^l(G, G') \right]_{uu'} \left[\Sigma_R^l(G, G') \right]_{uu'},\end{aligned}\tag{14}$$

195 where $\mathbf{T}^l(G, G')$ is the result of matrix multiplication between the column vector of the diagonal of
 196 $\Sigma_R^l(G, G)$ and row vector of the diagonal of $\Sigma_R^l(G', G')$. The affine transformation of the input
 197 GNTK corresponding to the aggregation phase as described in Eq. 5 is changed to (similar to Σ_0^l):

$$\left[\Theta_0^l(G, G') \right]_{uu'} = \sum_{v \in \mathcal{N}(u) \cup \{u\}} \sum_{v' \in \mathcal{N}(u') \cup \{u'\}} \left[\hat{\Theta}_R^{l-1}(G, G') \right]_{vv'},\tag{15}$$

198 2.3 The Proposed GraphQNTK

199 We first show that estimating the single-layer GraphQNTK and its covariance with infinite-width-
 200 limit attention mechanism can be efficiently reconstructed in the regime of quantum computing, and
 201 generalize to the multi-layer model. The following statements only consider two input graphs $G =$
 202 (V, E) and $G' = (V', E')$ with $|V| = n$ and $|V'| = n'$, and the corresponding feature matrix $\mathbf{H} =$
 203 $[\mathbf{h}_1^\top, \dots, \mathbf{h}_u^\top, \dots, \mathbf{h}_n^\top] \in \mathbb{R}^{n \times d}$ and $\mathbf{H}' = [\mathbf{h}_1^\top, \dots, \mathbf{h}_{u'}^\top, \dots, \mathbf{h}_{n'}^\top] \in \mathbb{R}^{n' \times d}$. The approximate
 204 estimation of GNTK is denoted as $\Theta \in \mathbb{R}^{n \times n'}$ and its element is $\Theta_{uu'}$. The corresponding
 205 covariance is $\tilde{\Sigma} \in \mathbb{R}^{n \times n'}$ and $\tilde{\Sigma}_{uu'}$. We use $\Theta_{GG'} \in \mathbb{R}$ to represent GraphQNTK after readout. We
 206 omit the subscript R for clarity. The same setting can be easily generalized to the arbitrary pair of
 207 graphs $G, G' \in \mathcal{G}$ by introducing auxiliary index registers. First, we introduce the quantum data
 208 structure accessible to the classical data, as commonly used by QML algorithms [51, 35, 33, 70].

209 **Feature encoding.** Using the storage structure as stated in our proposed Theorem 1 in Appendix,
 210 the feature matrix can be prepared into the QRAM at the initialization of the algorithm. The data
 211 encoding only occurs a single time and readout operation only takes logarithmic complexity time
 212 with respect to the number of samples n and dimension of feature d . The quantum representations
 213 corresponding to the encoded feature vector and feature matrix are as follows

$$\begin{aligned} |u\rangle|0\rangle &\rightarrow |u\rangle|\mathbf{h}_u\rangle, & |0\rangle &\rightarrow \frac{1}{\|\mathbf{H}\|_F} \sum_u \|\mathbf{h}_u\| |u\rangle, \\ |u'\rangle|0\rangle &\rightarrow |u'\rangle|\mathbf{h}_{u'}\rangle, & |0\rangle &\rightarrow \frac{1}{\|\mathbf{H}'\|_F} \sum_{u'} \|\mathbf{h}_{u'}\| |u'\rangle. \end{aligned} \quad (16)$$

214 **Estimation of the initialized NTK.** The empirical uncentered covariance of inputs $[\Sigma^0(G, G')]_{uu'}$
 215 and the initialized GNTK $[\Theta^0(G, G')]_{uu'}$, is the inner product between \mathbf{h}_u and $\mathbf{h}_{u'}$. Fol-
 216 lowing a similar approach to [36], the inner product between two vectors with respect to
 217 their quantum representations can be estimate efficiently by introducing an auxiliary register.
 218 Specifically, estimation of the inner product $\mathbf{h}_u^\top \mathbf{h}_{u'}$ can be performed by constructing the state
 219 $\frac{1}{\sqrt{2}}(|u\rangle|u'\rangle|0\rangle|\mathbf{h}_u\rangle + |u\rangle|u'\rangle|1\rangle|\mathbf{h}_{u'}\rangle)$. Applying a Hadamard gate on the third register gives the
 220 state $|u\rangle|u'\rangle (\sqrt{P_{uu'}}|0, g_{uu'}\rangle + \sqrt{1 - P_{uu'}}|1, g'_{uu'}\rangle)$, where $P_{uu'} = \frac{1 + \mathbf{h}_u^\top \mathbf{h}_{u'}}{2}$ is the estimation of
 221 the inner product. This procedure takes $O(\log d)$ time and we denote this quantum operation by \mathcal{D}^0 ,
 222 and we add a subscript to denote the corresponding conditioned operator, i.e, $\mathcal{D}_{uu'}^0$ represents \mathcal{D}^0 is
 223 conditioned acting on the basis state coupled with state $|0\rangle \rightarrow |u\rangle|u'\rangle$. We can perform the $\mathcal{D}_{uu'}^0$ in su-
 224 perposition such that the state $\frac{1}{\sqrt{nn'}} \sum_{u \in V} \sum_{u' \in V'} |u\rangle|u'\rangle (\sqrt{P_{uu'}}|0, g_{uu'}\rangle + \sqrt{1 - P_{uu'}}|1, g'_{uu'}\rangle)$
 225 can be generated in time $O(\log(nd))$.

226 **Quantum aggregation transformation.** Recall that an affine transformation (refer to Eq. 5 and
 227 Eq. 15) acting on the GNTK and its covariance is relative to the neighborhood aggregation defined by
 228 Eq. 1. Therefore, it is indirect to realize an end-to-end speedup similar to the estimation of the inner
 229 product since the transformation of each element of NTK and the covariance is not independent. To
 230 circumvent this barrier, we derive a unitary quantum aggregation transformation to approximately
 231 reconstruct the affine transformation. Consider the quantum operation $\mathcal{D}_{uu'}^0 : |u\rangle|u'\rangle|0\rangle|0\rangle \rightarrow$
 232 $|u\rangle|u'\rangle (\sqrt{P_{uu'}}|0, g_{uu'}\rangle + \sqrt{1 - P_{uu'}}|1, g'_{uu'}\rangle)$ that is employed to estimate the inner product of
 233 two feature vectors. Define a unitary operator which is used to perform aggregation transformation

$$\mathcal{U} = \sum_{v \in \mathcal{N}(u) \cup \{u\}} \sum_{v' \in \mathcal{N}(u') \cup \{u'\}} |v\rangle|v'\rangle \langle v|\langle v'| \otimes \mathcal{D}_{vv'}^0, \quad (17)$$

234 which can be generated by introducing conditional quantum evolution [21]. The operation \otimes denotes
 235 the tensor product. We apply the \mathcal{U} with Hadamard gates to the given initial state, which is given as

$$\begin{aligned} H^{\otimes} \mathcal{U} H^{\otimes} |0\rangle^{\otimes} |0\rangle |0\rangle &\rightarrow H^{\otimes} \mathcal{U} \sum_{v, v'} |v, v'\rangle |0\rangle |0\rangle \\ &\rightarrow H^{\otimes} \sum_{v \in \mathcal{N}(u) \cup \{u\}} \sum_{v' \in \mathcal{N}(u') \cup \{u'\}} |v, v'\rangle (\sqrt{P_{vv'}}|0, g_{vv'}\rangle + \sqrt{1 - P_{vv'}}|1, g'_{vv'}\rangle) \\ &\rightarrow \sum_{v \in \mathcal{N}(u) \cup \{u\}} \sum_{v' \in \mathcal{N}(u') \cup \{u'\}} \sqrt{P_{vv'}} |0\rangle^{\otimes} + \sqrt{\cdot} |\text{other}\rangle + \dots \end{aligned} \quad (18)$$

236 where $\sqrt{\cdot} |\text{other}\rangle$ represents other computational basis states except for $|0\rangle^{\otimes}$ with amplitude $\sqrt{\cdot}$, and
 237 the detailed mathematical expression and the scalar for state normalization are omitted since the
 238 result of the affine transformation has been embedded into the amplitude of $|0\rangle^{\otimes}$. The $(\cdot)^{\otimes}$ denotes
 239 that there could be multiple unitary operations acting on multiple registers, depending on the number
 240 of qubits required to encode the classic data. Similar to the inner product estimation, the quantum
 241 aggregation transformation can be performed in superposition and the resulting superposition is

$$\begin{aligned} &\frac{1}{\sqrt{nn'}} \sum_{u \in V} \sum_{u' \in V'} |u\rangle|u'\rangle (\sqrt{A_{uu'}}|0, y_{uu'}\rangle + \sqrt{1 - A_{uu'}}|1, y'_{uu'}\rangle), \\ \sqrt{A_{uu'}} &= \frac{\sum_{v \in \mathcal{N}(u) \cup \{u\}} \sum_{v' \in \mathcal{N}(u') \cup \{u'\}} \sqrt{P_{vv'}}}{|v| \times |v'|}. \end{aligned} \quad (19)$$

242 The amplitude $\sqrt{A_{uu'}}$ can be encoded into an ancillary register by using Amplitude Es-
 243 timation (Theorem 3) and Median Evaluation (Theorem 4). The obtained quantum state

244 $\frac{1}{\sqrt{nn'}} \sum_{u \in V} \sum_{u' \in V'} |u\rangle|u'\rangle|\bar{A}_{uu'}\rangle|y_{uu'}\rangle$ whose third register carries the approximate result after
 245 aggregation transformation as described in Eq. 5 and Eq. 15, where $|A_{uu'} - \bar{A}_{uu'}| \leq \epsilon$ and $|y_{uu'}\rangle$ is a
 246 garbage state. The runtime is $O(\log(nd)\log(1/\Delta)/\epsilon)$ and Δ is the proximity defined by the Median
 247 Evaluation. Note that $\bar{A}_{uu'}$ is actually the polynomial combination of the element-wise square root
 248 of the NTK from the previous layer, thus it is an approximate aggregation transformation. In the
 249 experiment, we empirically show that this approximation has a restrictive effect on the performance.

250 **Quantum kernel estimation** For fully-connected neural network, the calculation of each element of the NTK and its covariance is only reliant on the element at the same position of the
 251 covariance matrix in the previous fully-connected layer. Besides, the affine transformation of the
 252 GNTK and its covariance can be efficiently approximated by quantum aggregation transformation and the result has been embedded into the basis states of a superposition. In general, there
 253 exists a unitary $V : \sum_x |x, 0\rangle \rightarrow \sum_x |x, f(x)\rangle$ for any classical function f with the same time
 254 complexity to evaluate each element of the NTK and each element of the covariance [47, 70].
 255 Specifically, an oracle which operates as the same as classical function defined by Eq. 8 is im-
 256 plemented on the third register of $\frac{1}{\sqrt{nn'}} \sum_{u \in V} \sum_{u' \in V'} |u\rangle|u'\rangle|\bar{A}_{uu'}\rangle|y_{uu'}\rangle$. The resulting NTK is
 257 $\frac{1}{\sqrt{nn'}} \sum_{u \in V} \sum_{u' \in V'} |u\rangle|u'\rangle|\bar{\Theta}_{uu'}\rangle|y_{uu'}\rangle$, where $\bar{\Theta}_{uu'}$ is the approximate estimation of its classical
 258 counterpart after R fully-connected layers. The oracle is expected to be with the same complexity
 259 of its classical counterpart, which is associative to the number of fully-connected layers and is
 260 independent on the number of training samples n . For estimation of the GNTK after a transformer
 261 layer (Eq. 14), the covariance $\Sigma_R^l(G, G)$ for any $G \in \mathcal{G}$ requires to be estimated in advance. It
 262 means that the state $\frac{1}{n} \sum_{u \in V} \sum_{u' \in V'} |u\rangle|u'\rangle|\bar{\Sigma}_{uu'}\rangle|y_{uu'}\rangle$ must be estimated for any $G(V, E) \in \mathcal{G}$
 263 before input the different graphs, and we only consider the element when $u = u'$. By taking the
 264 partial trace on the second register, we obtain the state $\frac{1}{\sqrt{n}} \sum_{u \in V} |u\rangle|\bar{\Sigma}_{uu}\rangle|y_{uu}\rangle$ for graph G and
 265 $\frac{1}{\sqrt{n'}} \sum_{u' \in V'} |u'\rangle|\bar{\Sigma}_{u'u'}\rangle|y_{u'u'}\rangle$ for graph G' . Thus, estimation of the GNTK and its covariance
 266 corresponding to the multiplication part in Eq. 14 is given as

$$\begin{aligned} \frac{1}{\sqrt{nn'}} \sum_{u \in V} \sum_{u' \in V'} |u\rangle|u'\rangle|\bar{\Theta}_{uu'}\rangle|y_{uu'}\rangle &\rightarrow \frac{1}{\sqrt{nn'}} \sum_{u \in V} \sum_{u' \in V'} |u\rangle|u'\rangle|\bar{\Theta}_{uu'} \times \bar{\Sigma}_{uu} \times \bar{\Sigma}_{u'u'}\rangle|y_{uu'}\rangle, \\ \frac{1}{\sqrt{nn'}} \sum_{u \in V} \sum_{u' \in V'} |u\rangle|u'\rangle|\bar{\Sigma}_{uu'}\rangle|y_{uu'}\rangle &\rightarrow \frac{1}{\sqrt{nn'}} \sum_{u \in V} \sum_{u' \in V'} |u\rangle|u'\rangle|\bar{\Sigma}_{uu'} \times \bar{\Sigma}_{uu} \times \bar{\Sigma}_{u'u'}\rangle|y_{uu'}\rangle. \end{aligned} \quad (20)$$

269 This is performed by using the conditional quantum adder and the multiplier conditioned on the
 270 index register, i.e. $|u\rangle$ and $|u'\rangle$, which are designed by [62, 53, 40]. The final GNTK after the
 271 transformer layer can be directly generated by additional quantum arithmetic operations that perform
 272 an element-wise addition between the covariance to the GNTK.

273 **Estimation the GNTK for multiple layers** The quantum aggregation transformation requires that the
 274 approximate NTK and its covariance are embedded into the amplitudes of a superposition. However,
 275 after the quantum kernel estimation, these matrix are embedded into the quantum basis states of a
 276 superposition. To extract them back to the amplitudes, we apply Conditional Rotation [36] on the
 277 register containing the approximate GNTK (and the covariance), which is given by

$$\begin{aligned} \frac{1}{\sqrt{nn'}} \sum_{u \in V} \sum_{u' \in V'} |u\rangle|u'\rangle|\bar{\Theta}_{uu'}\rangle &\rightarrow \frac{1}{\sqrt{nn'}} \sum_{u \in V} \sum_{u' \in V'} |u\rangle|u'\rangle(a_{uu'}|0\rangle + \sqrt{1 - a_{uu'}^2}|1\rangle), \\ \frac{1}{\sqrt{nn'}} \sum_{u \in V} \sum_{u' \in V'} |u\rangle|u'\rangle|\bar{\Sigma}_{uu'}\rangle &\rightarrow \frac{1}{\sqrt{nn'}} \sum_{u \in V} \sum_{u' \in V'} |u\rangle|u'\rangle(b_{uu'}|0\rangle + \sqrt{1 - b_{uu'}^2}|1\rangle), \end{aligned} \quad (21)$$

278 where $a_{uu'} = \sqrt{\frac{\bar{\Theta}_{uu'}}{\max_{u,u'}(\bar{\Theta}_{uu'})}}$ and $b_{uu'} = \sqrt{\frac{\bar{\Sigma}_{uu'}}{\max_{u,u'}(\bar{\Sigma}_{uu'})}}$. We denote this quantum operation as
 279 $\mathcal{D}^l, l \in \{1, \dots, L\}$, where \mathcal{D}^L is used for the quantum readout operation. Similar to the operation
 280 \mathcal{D}^0 , the quantum aggregation transformation can be performed by generating a unitary operator by
 281 introducing conditional quantum evolution. Notice that $a_{uu'}$ and $b_{uu'}$ can be viewed as $\sqrt{P_{uu'}}$ in the
 282 setting of the single-layer GraphQNTK.

283 **Quantum readout** The resulting NTK is embedded into the basis states of a superposition since
 284 the algorithm ends up in the fully-connected layers. Similar to the classic readout operation, the
 285 summation of all the elements of the NTK matrix at the L -th layer is required. We use Conditional
 286 Rotation to extract the NTK back to the amplitude, and define a unitary \mathcal{O} which is a generalization

287 of the unitary \mathcal{U} , where

$$\mathcal{O} = \sum_{v \in V} \sum_{v' \in V} |v\rangle |v'\rangle \langle v| \langle v'| \otimes \mathcal{D}_{vv'}^L. \quad (22)$$

288 The unitary \mathcal{O} sums the square root of all the elements of the GraphQNTK matrix. And the
 289 resulting GraphQNTK between two input graphs is $\bar{\Theta}_{GG'} = \frac{(\sum_{u \in V, u' \in V'} \sqrt{\bar{\Theta}_{uu'}})^2}{n \times n'}$, where $\bar{\Theta}_{uu'}$ is
 290 the GraphQNTK of the last layer.

291 **Quantum inference to unseen data** We assume that the test data and the label of the training set
 292 are already encoded into the QRAM such that $|\mathbf{k}_*\rangle \in \mathbb{R}^N$, the GraphQNTK between the test graph
 293 G^* , can be evaluated as the same way to the evaluation between the training data. Let $\bar{\Theta} \in \mathbb{R}^{N \times N}$
 294 denote the GraphQNTK. The final prediction for a test datapoint G_* is

$$f^*(G^*) = \langle \mathbf{k}_* | \bar{\Theta}^{-1} | \mathbf{y} \rangle, \quad (23)$$

295 which requires solving the linear equation $|\mathbf{E}\rangle = \bar{\Theta}^{-1} |\mathbf{y}\rangle$ and performing inner product es-
 296 timation on $\langle \mathbf{k}_* | \mathbf{E} \rangle$. A popular quantum algorithm which is designed to solve the quantum
 297 linear systems problem (QLSP) is developed by [13], and its runtime is $O(\log(N)\kappa s \text{ poly-}$
 298 $\log(\kappa s/\epsilon))$ where s is the sparsity of matrix $\bar{\Theta}$ and κ is the condition number. To realize
 299 the quantum speedup, we assume a specific sparsity pattern is created in the quantum stor-
 300 age that only keeps $O(\log N)$ number of non-zero elements of the $N \times N$ GraphQNTK ma-
 301 trix and the well-conditioning is achieved by using Gershgorin circle theorem similar to [70].
 302

303 2.4 Complexity Study

304 In Sec. 2.3, we discuss how to approximately estimate
 305 GNTK using quantum computing paradigm between
 306 two input graphs. The time complexity is dominated by
 307 the quantum aggregation transformation procedure
 308 as it requires encoding the amplitude into an addi-
 309 tional register, which takes $O(\log(nd)\log(1/\Delta)/\epsilon)$
 310 time. Other quantum operations including esti-
 311 mation of the inner product, estimation of the GNTK
 312 within the neighborhood

313 aggregation and the fully-connected feature updating and quantum readout are totally unitary
 314 operations which can be efficiently performed under the regime of quantum computing. For
 315 estimating GNTK of each pairs of the graphs (G, G') where $G, G' \in \mathcal{G}$, each element of GraphQNTK
 316 $\bar{\Theta}$ can be generated simultaneously by introducing auxiliary index registers. The quantum runtime is
 317 $O(\log(Nnd))$. However, evaluating GNTK of the infinite-width-limit attention requires computing
 318 the kernel where the input is two same graphs, which can be implemented in time $O(N)$. The result
 319 should be stored in QRAM in advance which will be used to update GNTK corresponding the
 320 multi-head attention as described in Eq. 14. Therefore, it takes $O(N \log(Nnd))$ time to train the
 321 proposed quantum graph learning model, which achieves quadratic speedup compared to the existing
 322 GKs and completed approaches with $O(N^2)$ time.

331 3 Experiments

332 We evaluate our method for both GNTK and GraphQNTK with attention mechanism on several
 333 graph classification datasets involving either discrete or continuous attributes. All the experiments
 334 are performed on a workstation with a single machine with 1TB memory, one physical CPU with 28
 335 cores Intel(R) Xeon(R) W-3175X CPU @ 3.10GHz, and a single GPU (Nvidia Quadro RTX 8000).
 336 For our method and all the compared models, We follow the same setting as [17, 66], and report the
 337 average test accuracy and its standard deviation over a 10-fold cross validation on each dataset.

Table 1: Classification accuracies on graphs with discrete node attributes. The AttentionGNTK denotes the GNTK with attention mechanism without both sparsity and well conditioning, while the GraphQNTK is the kernel after performing these two transformations to meet the conditions for the use of quantum matrix inversion. The results of other models are taken from [17] except QS-CNN, which we evaluate on our dataset separation.

Dataset	MUTAG	PROTEINS	PTC	NCI1	IMDB-B	IMDB-M
WL subtree [56]	90.4 ± 5.7	75.0 ± 3.1	59.9 ± 4.3	86.0 ± 1.8	73.8 ± 3.9	50.9 ± 3.8
AWL [29]	87.9 ± 9.8	-	-	-	74.5 ± 5.9	51.5 ± 3.6
RetGK [68]	90.3 ± 1.1	75.8 ± 0.6	62.5 ± 1.6	84.5 ± 0.2	71.9 ± 1.0	47.7 ± 0.3
GNTK [17]	90.0 ± 8.5	75.6 ± 4.2	67.9 ± 6.9	84.2 ± 1.5	76.9 ± 3.6	52.8 ± 4.6
GCN [38]	85.6 ± 5.8	76.0 ± 3.2	64.2 ± 4.3	80.2 ± 2.0	74.0 ± 3.4	51.9 ± 3.8
GraphSAGE [20]	85.1 ± 7.6	75.9 ± 3.2	63.9 ± 7.7	77.7 ± 1.5	72.3 ± 5.3	50.9 ± 2.2
PatchySAN [48]	92.6 ± 4.2	75.9 ± 2.8	60.0 ± 4.8	78.6 ± 1.9	71.0 ± 2.2	45.2 ± 2.8
GIN [66]	89.4 ± 5.6	76.2 ± 2.8	64.6 ± 7.0	82.7 ± 1.7	75.1 ± 5.1	52.3 ± 2.8
QS-CNN [69]	93.1 ± 4.7	78.2 ± 4.6	66.0 ± 4.4	81.4 ± 2.6	72.1 ± 3.7	46.2 ± 4.2
AttentionGNTK	90.0 ± 8.5	76.2 ± 3.8	66.2 ± 5.1	84.1 ± 1.2	76.9 ± 3.2	52.9 ± 3.5
GraphQNTK	88.4 ± 6.5	71.1 ± 3.2	62.9 ± 5.0	77.2 ± 2.7	73.3 ± 3.6	48.1 ± 4.3

338 **3.1 Experiments setup**

339 **Datasets.** For graph with discrete attributes, the benchmark datasets include four bioinformatics
 340 datasets MUTAG, PTC, NC11, PROTEINS and three social network datasets IMDB-BINARY, IMDB-
 341 MULTI. For each graph, the input attributes is category of the node and they are transformed to
 342 one-hot encoding representations. For datasets where the graphs have no node features, i.e. only
 343 graph structure matters, we use degrees as input node features. For graph with continuous attributes,
 344 we select four benchmark datasets including ENZYMES, PROTEINS full, BZR, COX2. All the
 345 datasets can be found in [37]. The statistic information of the datasets are given in Tab. 3 in Appendix.

346 **Compared baselines.** We compare our method with state-of-the-art GKs such as WL kernel [56],
 347 AWL [29], RetGK [68], GNTK [17], WWL [61], and GNNs including GCN [38], PatchySAN [48],
 348 GCKN [10], GraphSAGE [20] and GIN [66]. For quantum graph learning, there are very few baseline
 349 available. We report the performance of the quantum walk based subgraph convolutional neural
 350 network (QS-CNN) developed by [69]. The data separation we use is the same as [66] for graph
 351 datasets with discrete attributes. For graph dataset with continuous attributes, we follow the same
 352 protocol as used in [61] to normalize the input feature vectors for a fair comparison.

353 **Results.** We apply dif-
 354 ferent hyper-parameter set-
 355 tings to $L \in \{2, 4, 6, 8\}$ and
 356 $R \in \{1, 2, 3\}$ and select
 357 the model with the best av-
 358 eraged accuracy. We test
 359 the kernel regression using
 360 SVM classifier and the reg-
 361 ularization parameter is de-
 362 termined using the search
 363 protocol which is the same
 364 as the [17]. We report the
 365 performance of the quan-
 366 tum approximate GNTK be-
 367 fore and after the matrix
 368 sparsity and conditioning
 369 operations. The numerical
 370 results are listed in Tab. 2.3

Table 2: Classification accuracies on graphs with continuous attributes. The accuracies of other models are taken from [10]. We only take the results of GCKN under the supervised learning for a fair comparison. We utilize the similar settings that preprocess the continuous node features to a normalized feature vector as in [61] for fair comparison (Note that the data encoded into the QRAM requires normalization, thus it is reasonable to use this data-preprocessing operation).

Dataset	ENZYMES	PROTEINS	BZR	COX2
RBF-WL [61]	68.4 ± 1.5	75.4 ± 0.3	81.0 ± 1.7	75.5 ± 1.5
HGK-WL [46]	63.0 ± 0.7	75.9 ± 0.2	78.6 ± 0.6	78.1 ± 0.5
HGK-SP [46]	66.4 ± 0.4	75.8 ± 0.2	76.4 ± 0.7	72.6 ± 1.2
WWL [61]	73.3 ± 0.9	77.9 ± 0.8	84.4 ± 2.0	78.3 ± 0.5
GNTK [17]	69.6 ± 0.9	75.7 ± 0.2	85.5 ± 0.8	79.6 ± 0.4
GCKN [10]	72.8 ± 1.0	77.6 ± 0.4	86.4 ± 0.5	81.7 ± 0.7
AttentionGNTK	69.2 ± 1.1	76.8 ± 1.2	86.7 ± 1.3	82.1 ± 0.4
GraphQNTK	64.8 ± 0.7	72.5 ± 0.3	80.1 ± 1.7	74.3 ± 1.9

371 for datasets with discrete attributes and Tab. 3.1 for datasets with continuous attributes. The attention
 372 method we integrate to the infinite-width GNNs brings to an improvement in the performance of
 373 the model. The results show that the GNTK with attention mechanism achieves better classification
 374 accuracy for graph data with medium number of nodes and edges. It is demonstrated that the infinite-
 375 width-limit attention captures global node similarity semantics and learns effective structure of the
 376 provided graph, which brings an remarkable accuracy improvement of the model compared with
 377 the vanilla GNTK [17]. Moreover, our model performs better than QS-CNN on more than 60% of
 378 the datasets with discrete attributes, given the caveat that QS-CNN is a hybrid graph learning model
 379 where the contribution of the classic components (CNNs, spatial message passing) in their model
 380 cannot be ignored. While the matrix sparsity and conditioning operations have a great influence on
 381 the model’s performance, it can be found that the classification performance of GNTK evaluated by
 382 quantum algorithms is still comparable with that of GKs and vanilla GNNs, where a tradeoff exists
 383 between the performance of the model and the quantum computational efficiency.

384 **4 Conclusion and Broader Impact**

385 This paper has presented a quantum graph learning model to characterize the structural information
 386 while using quantum parallelism to improve computing efficiency. We propose quantum algorithm to
 387 approximately estimate the neural tangent kernel of the underlying graph neural network where a
 388 multi-head quantum attention mechanism is introduced to incorporate semantic similarity of nodes.
 389 Empirical results on graph classification tasks as well as theoretical analysis show the superiority of
 390 our method. The limitation of the paper is that currently it only addresses graph-level embedding and
 391 we leave node-level quantum learning for future work. Our work may raise concerns for encryption,
 392 privacy protection etc. when the quantum hardware become more feasible.

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550 classical neural networks. *arXiv:2107.09200*, 2021.

551 Checklist

552 The checklist follows the references. Please read the checklist guidelines carefully for information on
553 how to answer these questions. For each question, change the default **[TODO]** to **[Yes]**, **[No]**, or
554 **[N/A]**. You are strongly encouraged to include a **justification to your answer**, either by referencing
555 the appropriate section of your paper or providing a brief inline description. For example:

- 556 • Did you include the license to the code and datasets? **[Yes]** See Section ??.
- 557 • Did you include the license to the code and datasets? **[No]** The code and the data are
558 proprietary.
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560 Please do not modify the questions and only use the provided macros for your answers. Note that the
561 Checklist section does not count towards the page limit. In your paper, please delete this instructions
562 block and only keep the Checklist section heading above along with the questions/answers below.

- 563 1. For all authors...
- 564 (a) Do the main claims made in the abstract and introduction accurately reflect the paper's
565 contributions and scope? **[Yes]**
- 566 (b) Did you describe the limitations of your work? **[Yes]**
- 567 (c) Did you discuss any potential negative societal impacts of your work? **[N/A]**
- 568 (d) Have you read the ethics review guidelines and ensured that your paper conforms to
569 them? **[Yes]**
- 570 2. If you are including theoretical results...
- 571 (a) Did you state the full set of assumptions of all theoretical results? **[N/A]**
- 572 (b) Did you include complete proofs of all theoretical results? **[N/A]**
- 573 3. If you ran experiments...
- 574 (a) Did you include the code, data, and instructions needed to reproduce the main experi-
575 mental results (either in the supplemental material or as a URL)? **[Yes]**
- 576 (b) Did you specify all the training details (e.g., data splits, hyperparameters, how they
577 were chosen)? **[Yes]**
- 578 (c) Did you report error bars (e.g., with respect to the random seed after running experi-
579 ments multiple times)? **[Yes]**
- 580 (d) Did you include the total amount of compute and the type of resources used (e.g., type
581 of GPUs, internal cluster, or cloud provider)? **[Yes]**
- 582 4. If you are using existing assets (e.g., code, data, models) or curating/releasing new assets...
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- 586 (d) Did you discuss whether and how consent was obtained from people whose data you're
587 using/curating? [N/A]
- 588 (e) Did you discuss whether the data you are using/curating contains personally identifiable
589 information or offensive content? [N/A]
- 590 5. If you used crowdsourcing or conducted research with human subjects...
- 591 (a) Did you include the full text of instructions given to participants and screenshots, if
592 applicable? [N/A]
- 593 (b) Did you describe any potential participant risks, with links to Institutional Review
594 Board (IRB) approvals, if applicable? [N/A]
- 595 (c) Did you include the estimated hourly wage paid to participants and the total amount
596 spent on participant compensation? [N/A]

597 A Related Works

598 In this section, we provide background on quantum graph learning and graph neural networks that
599 have the potential to be trained using quantum computing.

600 A.1 Graph Kernel Neural Network

601 Graph kernel neural networks [17, 10, 18] is a class of graph learning method combining the properties
602 of both GNNs and GKs. Forward process of the model tends to transmit node information like GNNs
603 [65], layer by layer, whereas the node (or graph) features live in the implicit reproducing kernel
604 Hilbert space (RKHS) of a specific kernel [39]. The mainstream of graph kernel neural networks
605 can be divided into completed and approximated approaches. For the former, the output is a kernel
606 matrix where each entry denotes the similarity of graph pairs, afterwards support vector machin is
607 used to perform classification or regression task. While the later generates the approximate feature of
608 the finite projected RKHS at the expense of information loss.

609 We consider the completed approaches for the basis of our proposed quantum graph learning model,
610 since the access to the explicit feature information requires measuring the relative quantum repre-
611 sentations, which incurs quantum collapse [32]. In the next section, we demonstrate that the graph
612 tangent kernel neural networks coincides with the condition of quantum parallel implementation by
613 introducing the quantum aggregation transformation and the quantum kernel estimation techniques.

614 A.2 Quantum Graph Learning

615 Quantum graph learning aims at leveraging quantum physics to extract graph structural information,
616 bringing up new possibilities for quantum computing applications. It is generally nontrivial to
617 analyze classical data under the regime of quantum computing, since the encoding and decoding
618 between classical vectors (or matrices) and their corresponding quantum states should be carefully
619 designed. In addition, encoding the irregular graph data and diverse structure topology may incur
620 different configurations of quantum models. Advanced contributions has developed some techniques
621 to overcome these issues. A hybrid graph learning method developed by [69, 16] encode the structure
622 information and generate a new adjacent matrix evaluating by the using quantum walk. The resulting
623 adjacent information captures the global topological arrangement information for graph substructures.
624 Adiabatic evolution [63] and conditional unitary [45] are applied to evolve the quantum systems
625 dependent on the underlying graph structure. In addition, the node attribute is encoded using
626 variational circuit [1] or a quantum random access memory [57]. Processing the encoded quantum
627 representation of the original graph can be realized via either a naive quantum algorithm [63] or a
628 hybrid method [11]. Then a post-processing operation is performed to further analyze the quantum
629 output. A brief review about quantum graph learning is illustrated in Fig. 1. **Generally speaking, the
630 researchers exploit to encode the graph structure and node features in the quantum system through
631 various schemes, and then process the information through quantum layers and auxiliary classical
632 layers. Finally, the (quantum) results are decoded through post-processing.** However, most quantum
633 graph learning models requires that adjustable parameters in the quantum algorithm need to be
634 updated frequently, where takes great computational overheads. Moreover, the classical components
635 in post-processing may dominate the performance of the model, thus weakening the role of the
636 quantum part. In this paper, We seek to establish a parameter-free quantum graph learning model to
637 maximize the efficacy of quantum computing.

Table 3: Statistic information of the used datasets.

Dataset	MUTAG	PROTEINS	PTC	NCI1	IMDB-B	IMDB-M	ENZYMES	BZR	COX2
size	188	1113	344	4110	1000	1500	600	405	467
classes	2	2	2	2	2	3	6	2	2
attr. dim.	-	-	-	-	-	-	18	3	3
avg. nodes	18	39	26	30	20	13	32.6	35.8	41.2
avg. edges	20	73	51	32	97	66	62.1	38.3	43.5

638 We notice that there are researches which are abbreviated as QNTK [58, 41], similar to ours nominally.
639 But their definition is quite different from ours. The motivation of these two papers is to analyze the
640 trainability and expressive power of variational quantum circuits through NTK. In contrast, in our
641 work, QNTK is a metric measuring the similarity of two input graphs. In this context, NTK is the
642 kernel that captures the dynamics of infinite-width GNNs, as well as the multi-head attention where
643 the number of heads and the dimension of output go to infinity.

644 B More Analysis

645 B.1 Quantum Access Memory

646 **Theorem 1** Let $|\mathbf{X}_p\rangle = \frac{1}{\|\mathbf{X}_p\|} \sum_{q=0}^{d-1} \mathbf{X}_{pq}|j\rangle$ denotes the amplitude encoding of the p -th row of data
647 $\mathbf{X} \in \mathbb{R}^{n \times d}$. There exists a data structure to store the entries of \mathbf{X} into the QRAM which is stated as

648 $i) |p\rangle |0\rangle \rightarrow |p\rangle |\mathbf{X}_p\rangle$

649 $ii) |0\rangle \rightarrow \frac{1}{\|\mathbf{X}\|_F} \sum_p \|\mathbf{X}_p\| |p\rangle$

650 in time T for $p \in [n]$. Using the binary tree QRAM architecture proposed by [34], the time T to store
651 and readout a new element scale logarithmically with respect to both n and d .

652 B.2 Inner Product Estimation

653 **Theorem 2** There exists a quantum operation \mathcal{A} that evaluates the inner product of two quantum
654 representations with respect to their d -dimensional classical vectors in time $O(\log d)$.

655 *Proof.* By introducing an auxiliary register, with the initial state $|p\rangle|q\rangle \frac{1}{\sqrt{2}}(|0\rangle + |1\rangle)|0\rangle$, the map
656 $\frac{1}{\sqrt{2}}(|p\rangle|q\rangle|0\rangle|0\rangle + |p\rangle|q\rangle|1\rangle|0\rangle) \rightarrow \frac{1}{\sqrt{2}}(|p\rangle|q\rangle|0\rangle|\mathbf{X}_p\rangle + |p\rangle|q\rangle|1\rangle|\mathbf{X}_q\rangle)$ can be performed in
657 $O(\log d)$ for two quantum representations $|\mathbf{X}_p\rangle$ and $|\mathbf{X}_q\rangle$ with respect to their classical vectors
658 $\mathbf{X}_p \in \mathbb{R}^d$ and $\mathbf{X}_q \in \mathbb{R}^d$. Applying a Hadamard gate on the third register, the state becomes

$$\frac{1}{2}|p\rangle|q\rangle (|0\rangle (|\mathbf{X}_p\rangle + |\mathbf{X}_q\rangle) + |1\rangle (|\mathbf{X}_p\rangle - |\mathbf{X}_q\rangle)). \quad (24)$$

659 The probability of measuring 0 on the third register is given by $P_{pq} = \frac{1 + \langle \mathbf{X}_p | \mathbf{X}_q \rangle}{2}$. Thus the state
660 defined by Eq. 24 can be reformulated as $|p\rangle|q\rangle (\sqrt{P_{pq}}|0, g_{pq}\rangle + \sqrt{1 - P_{pq}}|1, g'_{pq}\rangle)$ where $|g_{pq}\rangle$
661 and $|g'_{pq}\rangle$ are garbage states.

662 B.3 Amplitude Estimation

663 **Theorem 3** Given a unitary operator U such that $U : |0\rangle \mapsto \sqrt{p}|y\rangle|0\rangle + \sqrt{1-p}|y'\rangle|1\rangle$ in time T ,
664 where $p > 0$ is the probability of measuring 0, it is possible to obtain the state $|y\rangle|0\rangle$ using $O(\frac{T}{\sqrt{p}})$
665 queries to U , or to estimate p with relative error δ using $O(\frac{T}{\delta\sqrt{p}})$ queries to U . The detailed proof
666 can be found in [9].

667 B.4 Median Evaluation

668 **Theorem 4** Consider a unitary $U : |0^{\otimes m}\rangle \mapsto \sqrt{\alpha}|v, 1\rangle + \sqrt{1-\alpha}|g, 0\rangle$ for some $1/2 \leq \alpha \leq 1$ in
669 time T . There exists a quantum algorithm that, for any $\Delta > 0$ and for any $1/2 < \alpha_0 \leq \alpha$, produce a

Table 4: Running time comparison between different models.

	MUTAG	NCI1	IMDB-B	IMDB-M
GIN	22 sec	67 min	19 min	24 min
GNTK	9 sec	18 min	4 min	7 min
Ours	14 sec	21 min	4 min	9 min

670 state $|\psi\rangle$ such that $\| |\psi\rangle - |0^{\otimes mL}\rangle |x\rangle \| \leq \sqrt{2\Delta}$ for some integer L in time

$$2T \left[\frac{\log(1/\Delta)}{2(|\alpha_0| - 1/2)^2} \right]. \quad (25)$$

671 Refer to [64] for a detailed proof.

672 C Additional Experiments and Discussion

673 C.1 Running time comparison

674 We supplement the training time of our model on four selected datasets and compare it with two other
 675 models. To make a fair comparison, we set the layers of all the models to 2. All the experiments are
 676 performed on a workstation with a single machine with 1TB memory, one physical CPU with 28
 677 cores Intel® Xeon® W-3175X CPU @ 3.10GHz, and a single GPU (Nvidia Quadro RTX 8000). The
 678 results are shown in Tab. 4. Although the speedup introduced by the quantum algorithm depends on
 679 the quantum devices, it shows that our proposed model still has a computational overhead reduction
 680 when training on classic computers. The running time is slightly higher than that of GNTK which is
 681 a lack of attention mechanism. It is noticed that our model. The runtime of our model is apparently
 682 faster than that of GIN.

683 It is worth mentioning that the quadratic quantum speedup will be realized when the quantum
 684 hardware becomes more feasible.

685 C.2 Model Sensitivity to the Number of Layers

686 In the main body of the paper, we report the best classification accuracy of the model when the
 687 the number of layers L is selected from $\{2, 4, 6, 8\}$. We compare the graph classification accuracy
 688 bewteen GIN and our model at the same number of layers and the results are given in Tab. 5 and
 689 Tab. 6. The number in parentheses in the table indicates the number of layers.

690 From Tab. 5, it is shown that our model (AttentionGNTK) is more robust compared with GIN when
 691 the number of layers becomes larger. The main reason is that an additional feature aggregation, e.g.,
 692 the transformer, can slow down the convergence rate, which is consistent with the observations in
 693 [28] that connectivity enhancement can help wide and deep GNNs to avoid a discrepancy between
 694 prediction and the ground truth.

695 While in Tab. 6, the results empirically demonstrate that our model (attentionGNTK) reaches the
 696 peak of classification accuracy when the number of layers is small, while GNTK needs more layers
 697 to reach, indicating that our model is easier to capture the global structure information of the graph.
 698 This could be interpreted from the theoretical perspective. The transformer (Eq. 12) captured the
 699 semantic information between each pair of (connected and disconnected) nodes with similar features.

700 Consider $\mathbf{G} = \frac{\mathbf{Q}\mathbf{K}^T}{\sqrt{s}}$ in Eq. 10, where \mathbf{Q} and \mathbf{K} are linear transformation of the node feature matrix
 701 and we ignore the superscript and the subscript for simplicity. The \mathbf{G} can be viewed as a matrix
 702 whose element corresponds to the similarity of each pair of nodes. Then the operation $\hat{\mathbf{H}} = \zeta(\mathbf{G})\mathbf{V}$
 703 transforms the node features of the last layer to the next layer depending on the node similarity. This
 704 enables the model to make better use of the graph structure to transmit information and perceive
 705 topology information over long distances.

Table 5: Classification accuracy between GIN [66] and ours with respect to different layers.

	GIN(4)	Ours(4)	GIN(6)	Ours(6)	GIN(8)	Ours(8)
MUTAG	87.6 ± 6.2	89.1 ± 7.8	88.5 ± 5.6	90.0 ± 8.5	86.2 ± 6.4	88.4 ± 7.4
PROTEINS	75.5 ± 3.0	75.0 ± 4.1	74.3 ± 3.0	76.1 ± 3.8	72.8 ± 3.5	74.2 ± 4.4
PTC	62.8 ± 5.0	64.9 ± 5.3	62.0 ± 6.2	66.2 ± 5.1	61.2 ± 7.1	63.4 ± 6.6
NCII	82.3 ± 3.6	84.1 ± 1.2	80.1 ± 2.4	83.8 ± 1.2	77.2 ± 3.3	82.3 ± 2.2
IMDB-B	73.2 ± 4.1	75.7 ± 2.8	74.4 ± 6.0	76.9 ± 4.3	72.1 ± 5.2	75.1 ± 4.0
IMDB-M	51.7 ± 3.7	52.0 ± 4.1	52.0 ± 2.6	51.9 ± 3.7	48.2 ± 4.3	50.3 ± 4.5

Table 6: Classification accuracy between GNTK [17] and ours with respect to different layers.

	GNTK(4)	Ours(4)	GNTK(6)	Ours(6)	GNTK(8)	Ours(8)
PTC	62.9 ± 7.2	64.9 ± 5.3	63.5 ± 6.8	66.2 ± 5.1	65.2 ± 7.9	63.4 ± 6.6
NCII	83.6 ± 2.1	84.1 ± 1.2	84.0 ± 0.9	83.8 ± 1.2	82.9 ± 1.8	82.3 ± 2.2