

# Geometric Tree Classification by Markov Chain Modeling

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## Abstract

Geometric Tree Classification is an important problem in data mining and computer vision, with applications in object detection and medical imaging. In this paper, we study the problem of learning tree graph objects from images. We first show standard Convolutional Neural Networks (CNNs) do *not* learn the *structure* of graph images, rather they learn the background objects. Next, we propose a novel method for Geometric Graph Classification based on Markov Chain Modeling. We model the structure of a tree as a Markov Chain in two ways, (i) each bifurcation point in the tree is represented as a node and each branch is represented as an edge, (ii) each branch is represented as a node. We then use the Markov Chain to define a probability distribution over the possible tree structures, which we use to classify the input tree. Our method achieves superior results over several state of the art models in both synthetic and real datasets. Overall, this research provides a promising new direction for geometric tree classification.

## 1 Introduction

Geometric Trees are commonly used to represent hierarchical structures in many different applications, particularly in biomedical imaging and linguistics. In this work, we will explore geometric tree classification in images which contain tree structures. Our motivating example is a curated tree dataset we have produced. The state of the art methods for object classification in images are deep convolutional neural networks, [12]. Convolutional Networks are translationally invariant, but they are not invariant to rotations in the 3-dimensional space which is then projected back onto to the image plane. Our real and synthetic datasets consist of tree images undergoing various affine transformations and rotations in the 3-dimensional space which are then imaged and can be seen in Figures 1a, 1b, 1c, 1d.

To show Convolutional Neural Networks are unable to learn a good representation of the trees, even with a good amount of data, we run a network similar to AlexNet on 5 real trees, each with 120 training images and 30 testing images. The results can be seen in table 1. As we can see in the figures 1b and 1d, our synthetic



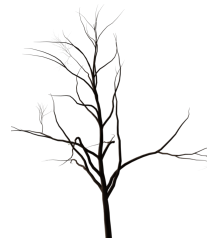
(a) Real



(b) Synthetic



(c) Real



(d) Synthetic

Figure 1: Figures 1a and 1c are the same tree from a different angle and figures 1b and 1d are the same tree from a different angle.

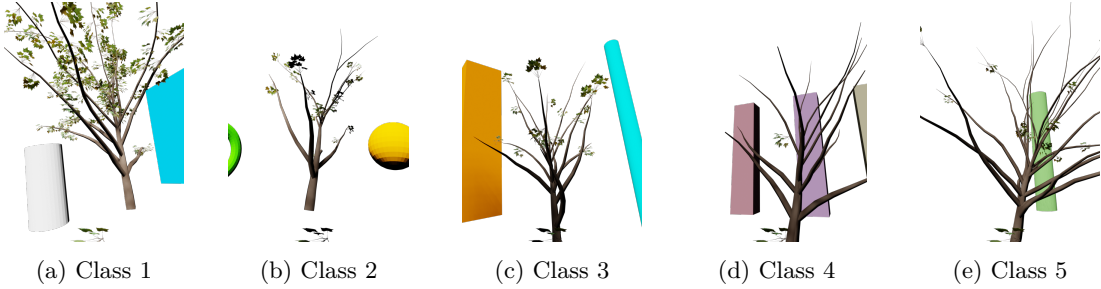


Figure 2: Tree Images with Random Objects in the Background that are unique to each class of trees. Each class represents the same tree with images taken from different angles and lighting conditions.

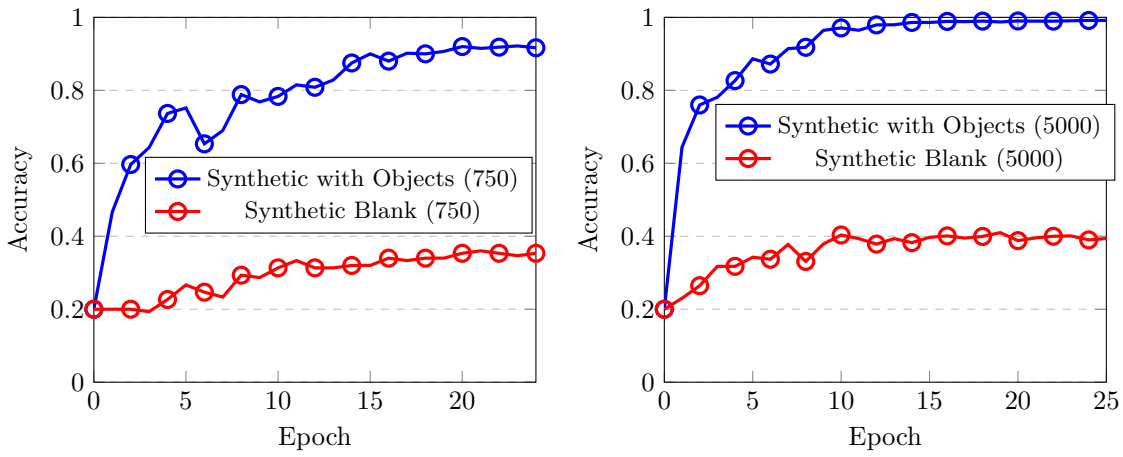


Figure 3: Train Accuracy is given in blue and Test Accuracy is given in red. We train on data with random objects in the background and test on images with no background. In (*Left*), we have 750 total images in the dataset. In (*Right*), we have 5000 total images in the dataset. This figure shows the increase in data does not overcome the distributional shift of removing background objects.

dataset trees have no background. The very poor results of the neural network in this synthetic case supports our informal hypothesis that the CNN is only able to learn an accurate prediction due to it learning features in the background of the real images. To further verify this, we create two datasets of synthetic trees, one where the background is blank, and one where the background has 4 – 5 randomly selected and shaped objects. We can see in table 1, the **AlexNet** model learns a significantly better representation when there exists the random objects in the background compared to when the background is blank. Our results can be seen in figure 3

This can be very dangerous in biomedical applications where it is essential to learn the structure of the graph underlying the image. Our research question is to find a more data-efficient method to learn the tree structure in image classification with higher accuracy. Now we will show some of the segmentations from the Segment-Anything-Model [11]. As we see in figures 7a and 7b, the Segment-Anything-Model is able to remove the background from the lower half of the tree near the trunk, where the branches are large. However, as we go up the tree, we see it is unable to segment the fine-structure of the tree and often leaves parts of the building and the sky. With the segmentation from the Segment-Anything Model, we will now do classification and see if there are better results in the small data case. We will use the underlying graph structure of the image as can be seen in figure 9.

Our approaches utilize the graph structure of the image, which we have annotated in Figures 9a, 9b, 9c, and 9d.

Many approaches have been made for Tree Graph Classification. Such include the use of Geometric Tree Distance ([6]), feature-matching approaches ([15]), geometric graph matching ([18, 1]).

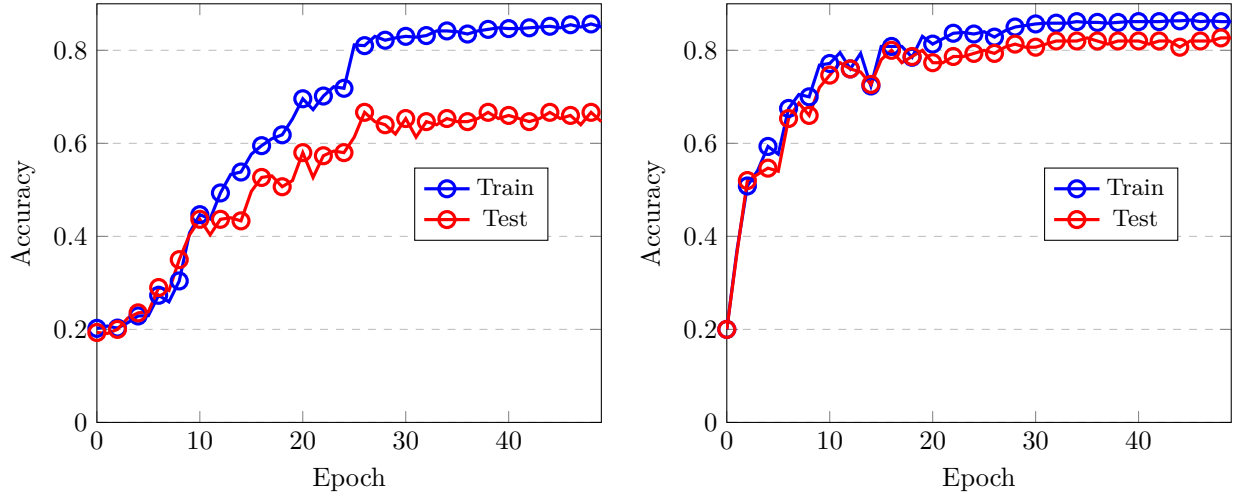


Figure 4: Training and testing results on 750 synthetic images (5 classes, 120 train, 30 test) from synthetically developed trees without background are displayed after training on AlexNet in (*Left*). Training and testing results on 750 synthetic images (5 classes, 120 train, 30 test) from synthetically developed trees with random items in the background unique to each class are displayed after training on AlexNet in (*Right*).

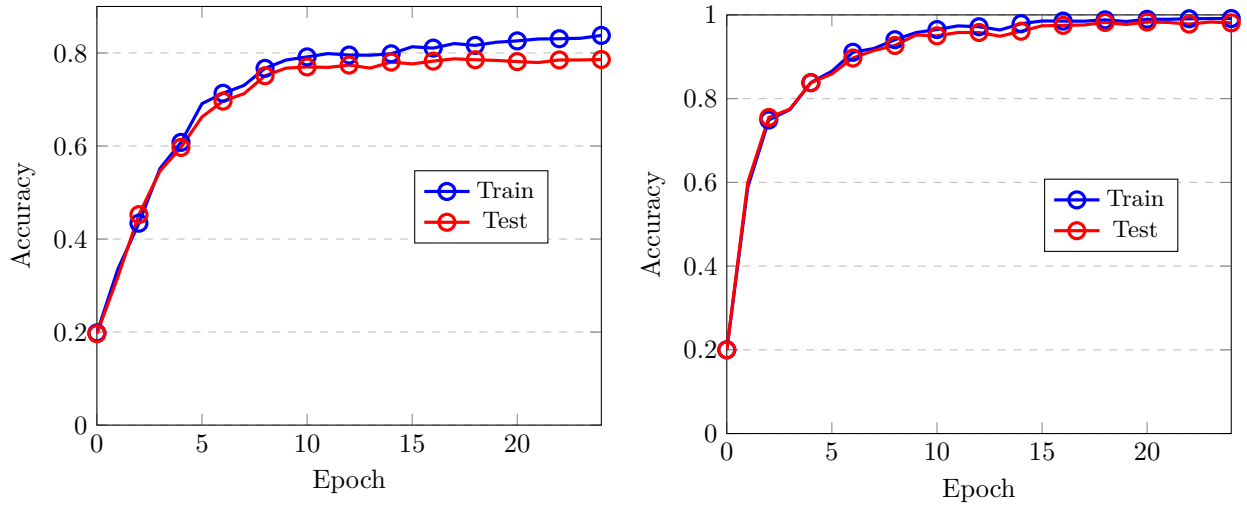


Figure 5: Training and testing results on 5000 synthetic images (5 classes, 800 train, 200 test) from synthetically developed trees with random items in the background unique to each class are displayed after training on AlexNet in (*Right*). As we see in the Background to Background case, more data helps with the training. Furthermore, we see better generalization with more data in the blank case, but we do not see a high increase in train accuracy. This leads us to believe learning the blank trees is in general difficult.

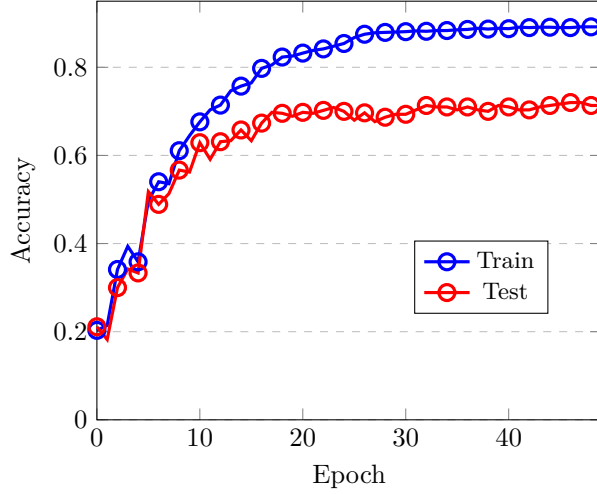


Figure 6: Training and Testing on 750 real images (5 classes, 600 train, 150 test) pictured in nature. The results are from training on AlexNet. We see the network overfits as it is likely learning local features such as the background as we have seen in our previous synthetic experiments.



Figure 7: Segmentation Results from Segment-Anything-Model

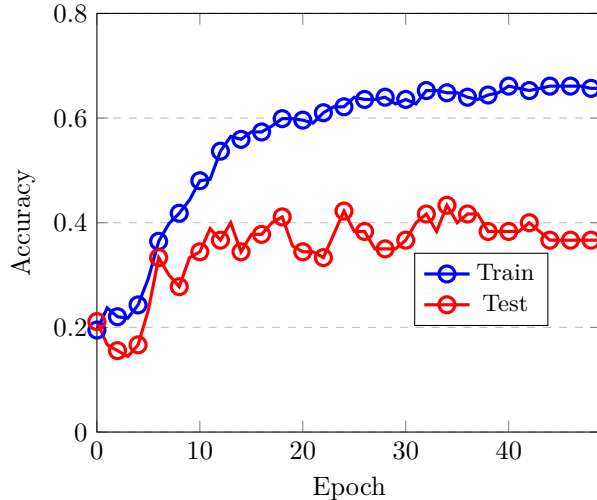


Figure 8: Training and Testing on 150 real images (5 classes, 120 train, 30 test) pictured in nature with segmentations by Segment Anything Model to remove the background to and focus as much as possible on the tree structure. The results are from training on AlexNet. We find the results in the small data case to be quite low.

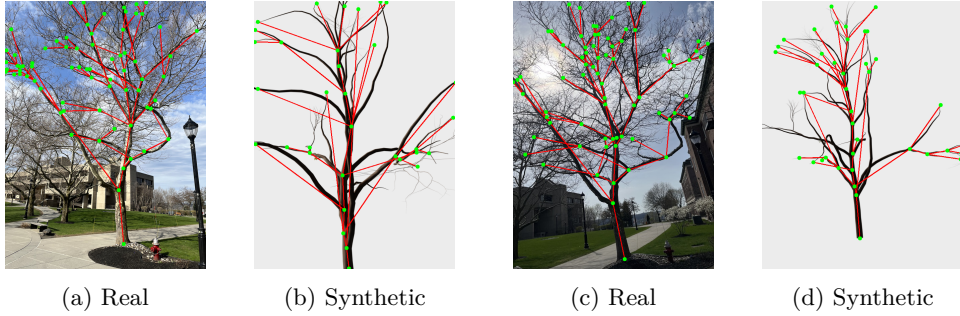


Figure 9: Geometric Tree Graphs overlaying the Tree Images

Model	Parameters	Dataset	Classes	Total Samples	Test Accuracy
AlexNet	46,767,493	Synthetic Blank	5	750	66.6 <sub>(4.80)</sub>
AlexNet	46,767,493	Synthetic with Objects	5	750	82.6 <sub>(2.63)</sub>
AlexNet	46,767,493	Synthetic Blank	5	5000	78.0 <sub>(8.07)</sub>
AlexNet	46,767,493	Synthetic with Objects	5	5000	98.9 <sub>(1.01)</sub>

Table 1: Statistics and Training Results for our Synthetic Datasets. We observe with more data we have higher accuracy. Furthermore, it is evident with objects in the background, the neural network obtains a higher accuracy.

### Contributions

In this paper, we assume there is a method to obtain the tree structure underlying an image and we propose multiple novel approaches for geometric tree classification by modelling the tree as a Markov Chain. We give two methods that use only the topological features of the graph for classification, and we propose one method that uses both the topological and geometric features of the graph for classification. Our methods perform better than the State of the Art in our geometric tree classification datasets.

## 2 Related Work

In this section we will describe relevant works with regards to learning geometric graphs and trees. A family of Geometric Tree Kernels is presented in [8] for the classification of Geometric Trees. The best performing kernels were root-path kernels, which are formulated as the following

$$K_r(T_1, T_2) = \sum_{v_i \in V_1, v_j \in V_2} k_p(p_{ir}, p_{jr}) \quad (1)$$

$$\text{where } k_p(x_{ij}, x'_{kl}) = \begin{cases} \langle x_{ij}, x'_{kl} \rangle & (\text{linear}) \\ e^{-\lambda \|x_{ij} - x'_{kl}\|_2^2} & (\text{Gaussian}) \end{cases}$$

Here the researchers model the difference between root-paths by landmarking a set of points on each path and comparing the difference, either with a linear dot product or with the Gaussian. They then use kernel methods to solve the classification task.

The spaces of Tree-Like shapes has been analyzed in [7]. Using the space of tree-like shapes in Principal Component Analysis (PCA) has been studied in [17].

Graph Neural Networks have gained considerable attention in the past few years. However, only a subset of such Graph Neural Networks are capable of Graph Classification in tasks where the Graphs can have different number of nodes and edges.

The Graph Convolutional Network [10] is a semi-supervised approach for Graph Classification. It is based on the principles of the popular Convolutional Neural Networks used typically in images. The layer wise

propagation rule is defined as:

$$H^{(l+1)} = \sigma \left( \tilde{D}^{-\frac{1}{2}} \tilde{A} \tilde{D}^{-\frac{1}{2}} H^{(l)} W^{(l)} \right) \quad (2)$$

The convolutional structure is similar to convolutional neural networks for images where in lower layers local features are learned and in later layers more holistic features about the graph are learned.

Graph Isomorphism Networks,[20] are a powerful GNN which has provably maximal discriminative power among GNNs. In each iteration the GIN updates node representations with the following equation:

$$h_v^{(k)} = \text{MLP}^{(k)} \left( \left( 1 + \epsilon^{(k)} \right) \cdot h_v^{(k-1)} + \sum_{u \in \mathcal{N}(v)} h_u^{(k-1)} \right) \quad (3)$$

### 3 Markov Chain Modeling

In this section we will discuss the background for Markov Chain Modeling and then present our algorithms.

#### 3.1 Markov Chain Preliminaries

A Markov Chain can be represented by a 3-tuple  $(\mathcal{S}, \mathcal{P}, s_0)$ , [16].  $\mathcal{S}$  represents the state space of the Markov Chain,  $\{s_0, s_1, \dots, s_n\}$ .  $\mathcal{P}$  represents the probabilistic transition matrix. Each entry  $\mathcal{P}_{ij}$  represents the probability of moving to  $s_j$  from  $s_i$ .  $s_0$  represents the start state of the Markov Chain. The Markov Property states  $\mathbb{P}[s_n | s_{n-1}, s_{n-2}, \dots, s_0] = \mathbb{P}[s_{n-1}]$ . All states in the Markov Chain must follow this property.

**Definition 1.** Let a Markov Chain be defined by  $(\mathcal{S}, \mathcal{P}, s_0)$ , then a Markov Chain is **irreducible** if for all  $s_1 \in \mathcal{S}$ , there exists a path to any  $s_2 \in \mathcal{S}$ .

**Definition 2.** Let a Markov Chain be defined by  $(\mathcal{S}, \mathcal{P}, s_0)$ . Let  $\mathbb{P}[s_i \rightarrow s_i | t]$  be the return probability after  $t$  steps. If there exists  $t$  s.t.  $\mathbb{P}[s_i \rightarrow s_i | nt] = 0$  for all  $n \in \mathbb{N}$ , then the Markov Chain is Periodic. If no such  $t$  exists, then the Markov Chain is **aperiodic**.

#### 3.2 Topological Features

Throughout this section, we will represent  $\phi(X) : \mathcal{T} \rightarrow \mathbb{R}^d$  represent the feature mapping from Tree to vector,  $\psi(i) : V \rightarrow \mathbb{Z}$  represent the distance in edges from node  $i$  to the root,  $\varphi : V \rightarrow \mathbb{N}_0$  represent the number of outgoing edges from node  $i$ . The state space  $\mathcal{S}$  represents all two-tuple points of the form  $(a, b)$ . Given a vertex,  $v$ , in a tree  $T$ , we determine the state of  $v$  as  $(\psi(v), \phi(v))$ . In practice, we set a maximum on  $a$  and  $b$ , in our datasets  $a \leq 10$  and  $b \leq 10$ , so our feature vector is not too sparse and to limit the effect of unseen vertices in the testing set.

##### 3.2.1 Markov Chain Likelihood Approach

In our first proposed method, we use only topological features of the tree for the graph. This means we do not utilize the node locations in  $\mathbb{R}^2$  for our training or prediction.

**Theorem 3.** Let the Markov Chain be defined as  $(\mathcal{S}, \mathcal{P}, s_0)$  and the classification problem be defined as follows:

$$c = \arg \max_{c \in C} \mathbb{P}[C = c | T] \quad (4)$$

under the condition

$$\sum_{c=1}^m \mathbb{P}[C = c | T] = 1 \quad \forall T \in \mathcal{T} \quad (5)$$

Let  $\mathcal{B}(T)$  represent the set of branches in tree  $T$ , then the negative log-likelihood of a tree  $T$  belonging to class  $c$  is given as

$$\arg \max_{c \in C} \mathbb{P}[C = c | T] = \arg \min_{c \in C} \frac{\left( \sum_{b \in \mathcal{B}(T)} \sum_{s_i \in b} - \log \left( \mathcal{P}_{s_{i-1}, s_i}^{(c)} \right) \right) \mathbb{P}[C = c]}{\sum_{j=1}^m \left( \left( \sum_{b \in \mathcal{B}(T)} \sum_{s_i \in b} - \log \left( \mathcal{P}_{s_{i-1}, s_i}^{(c)} \right) \right) \mathbb{P}[C = j] \right)} \quad (6)$$

This represents the conditional probability a geometric tree  $T$ , is in class  $c$  of the  $m$  total classes. Similar to [8], we identify a tree by its root-paths. This means a tree  $T$  is made by a set of root-paths, and on each path is a set of nodes from the root to a leaf. Thus, the number of root paths in a tree equals the number of leaves.

When we run inference, we choose the  $c \in C$  that minimizes the negative log likelihood in equation 6.

### 3.2.2 Calculating $\mathcal{P}$

Let  $\mathcal{T}^{(k)}$  represent the set of trees in the training data set from class  $k$ , and let  $N^{(k)}(\cdot)$  represent the number of occurrences of reaching  $\cdot$  within  $\mathcal{T}^{(k)}$ .

$$\mathcal{P}_{s_{i-1}, s_i}^{(k)} = \frac{N^{(k)}(s_{i-1} \rightarrow s_i)}{N^{(k)}(s_{i-1})} \quad (7)$$

### 3.2.3 Stationary Distribution Approach

We will now discuss our second approach. This approach also utilizes only the topological features of the Geometric Tree. The stationary distribution  $\pi$ , is a vector that satisfies the following conditions

$$\sum_{i=1}^{|S|} \pi_i = 1 \quad (8)$$

$$\pi \mathcal{P} = \pi \quad (9)$$

Let us note the probabilistic transition matrix of a geometric tree is always irreducible, however, this is a periodic Markov Chain. For example, if we are to start at the root, we can only come back to the root on steps 2, 4, 6, ... . This is because for all states we have self-transition probability equal to 0. To justify this mathematically, let  $\mathcal{L}$  represent the eigenvalues such that  $|\lambda_1| \geq |\lambda_2| \geq \dots \geq |\lambda_n|$ . Since  $\mathcal{P}$  has periodicity of 2, we end up with  $|\lambda_1| \approx |\lambda_2|$ . It thus follows in the power method:

$$\mathbf{x} \mathcal{P}^k = \alpha_1 \lambda_1^k \left( \mathbf{v}_1 + \frac{\alpha_2}{\alpha_1} \left( \frac{\lambda_2}{\lambda_1} \right)^k \mathbf{v}_2 + \dots + \frac{\alpha_m}{\alpha_1} \left( \frac{\lambda_m}{\lambda_1} \right)^k \mathbf{v}_m \right) \quad (10)$$

However since, we have  $|\lambda_1| \approx |\lambda_2|$  this simplifies to:

$$\mathbf{x} \mathcal{P}^k = \alpha_1 \lambda_1 \mathbf{v}_1 + \alpha_2 \lambda_2 \mathbf{v}_2 \quad (11)$$

We thus need to create an aperiodic Probabilistic Matrix. To create an aperiodic Probabilistic Matrix, we do the following adjustment:

$$\hat{\mathcal{P}} = (1 - \varepsilon) \mathcal{P} + \varepsilon \mathbf{R} \text{ where } \mathbf{R} = \begin{pmatrix} 1 & 0 & \dots & 0 \\ 1 & 0 & \dots & 0 \\ \vdots & \vdots & \ddots & \vdots \\ 1 & 0 & \dots & 0 \end{pmatrix} \text{ and } \varepsilon \in [0, 1] \quad (12)$$

We call  $\varepsilon$  the return probability. From any node in the tree, with probability  $\varepsilon$ , we can return back to the root of the tree. With higher  $\varepsilon$ , we will be less likely to explore areas of the tree from the root.

**Lemma 4.** *The probabilistic matrix  $\hat{\mathbf{P}} \triangleq (1 - \varepsilon) \mathcal{P} + \varepsilon \mathbf{R}$  where  $\mathcal{P}$  is a probabilistic matrix with columns summing to 1 s.t.  $P_{i,j} \geq 0$  for all  $i \in [n]$  and  $j \in [n]$ , then  $\hat{\mathbf{P}}$  is a aperiodic matrix.*

**Proof Sketch.** To be aperiodic means from any node, there exists a path of odd and even length to any other node in the Markov chain. Consider the path from the root to node adjacent to the root, call it  $v$ , let the length of this path be  $m$ , note since the tree is directed, the path to  $v$  from the root is unique and is of length  $m$ . Now, with the return probability, we can create a path where the first step is from the root to itself, now we will have a path of length  $m + 1$ , this can be generalized for all  $v \in \mathcal{S}$ . We provide a more

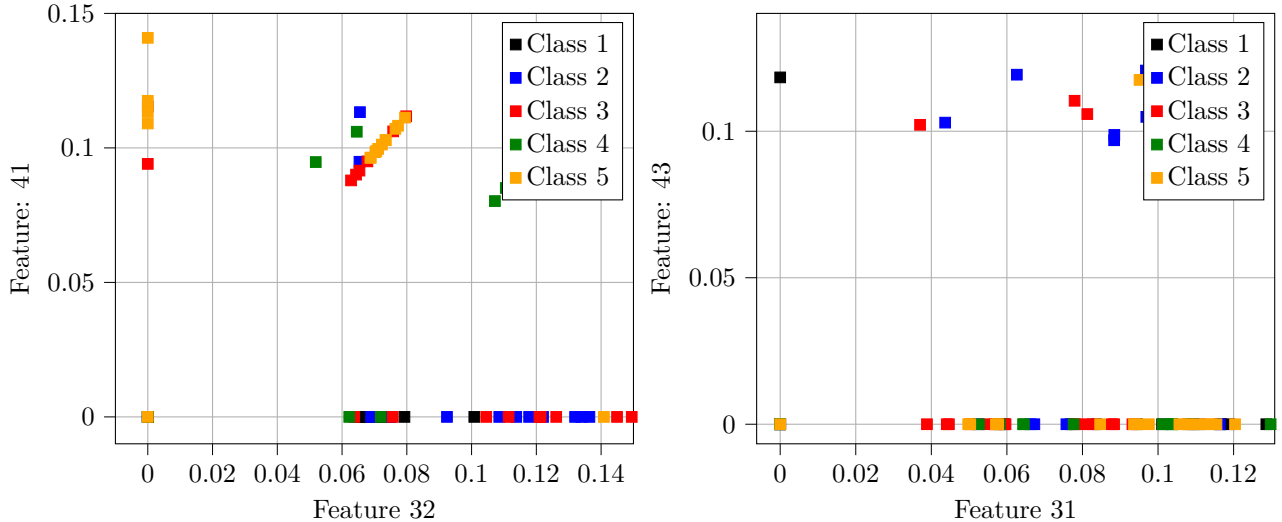


Figure 10: Graphs of 4 Highest Variance Features in Stationary Distribution Method

rigorous proof in the appendix. ■

We can now find the stationary distribution of  $\hat{\mathcal{P}}$ . We do this by calculating the eigenvector that is associated with the 1 eigenvalue. Let this be  $\hat{\mathbf{v}}$ , we then map this feature to the state space described in the beginning of § 3.2. This is done by a sum-aggregation. In other words, for each vertex,  $v$ , in the tree we obtain its state representation  $(\psi(v), \phi(v))$ , and the value assigned to the state  $(\psi(v), \phi(v))$  is equal to the summation of all probabilities of vertices with the same state representation. We then use this feature as an input to an SVM. Our results can be seen in Table 3 and ??.

To impose a better inductive bias on the task of classification of the geometric tree, we introduce the idea of a return probability. Let  $\varepsilon$  determine the return probability. At each state in the graph, we have a probability of  $\varepsilon$  chance of going back to the root. We want to see if this has any effect on the accuracy and if it is useful for the classification. In figure 11, we see that at low return probabilities, the accuracy is maximized. As  $\varepsilon$  is increased, the nodes with greatest distance from the node are explored less and less, thus leading the stationary vector to be less descriptive of the tree graph.

The stationary vectors are high dimensional vectors, so it is not possible to visualise all dimensions. To see the effectiveness of this feature, we graph the entries in the vector with the highest variance. The graphs can be seen in Figure 10. We can see the data is not separable with just the few features, it requires many more features.

### 3.3 Geometric Features

In our third proposed method, we utilize the locations of the nodes in  $\mathbb{R}^2$  in conjunction with the Markov Chains for classification.

We will define a new Markov Chain  $(\mathcal{S}, \mathcal{P}, s_0)$ , where  $s \in \mathcal{S}$  represents the branches of the tree.

**Definition 5.** The **Hitting Distance** of a Geometric Markov Chain is defined as:

$$D_A = \inf\{n \in \mathbb{R}^+ : \sum_{i=1}^m d(s_{i-1}, s_i) = n\} \quad (13)$$

where  $d$  represents a metric in the  $\mathbb{R}^2$  space.

**Definition 6.** The **Expected Hitting Distance** of a Geometric Markov Chain defined by the tuple  $(\mathcal{S}, \mathcal{P}, s_0)$  is defined as follows:

$$\mathbb{E}[D_{iA}] = \mathbb{E}[D_A | s_0 = i] \quad (14)$$

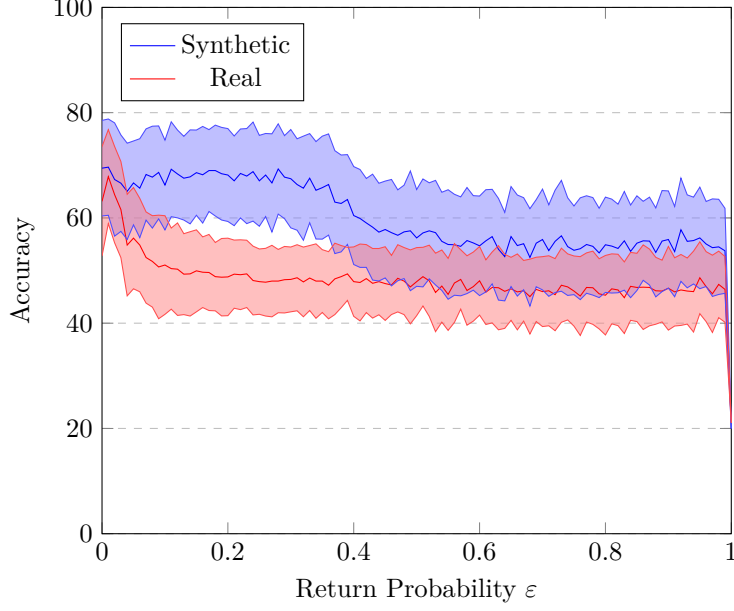


Figure 11: Effect of Return Probability on the accuracy

The Hitting Distance defined in equation 13 can be solved with the following set of equations which are based on the law of total probability.

$$D_{iA} = \begin{cases} \sum_{j \in S} P_{ij} D_{jA} & \text{if } i \neq A \\ 0 & \text{if } i = A \end{cases} \quad (15)$$

Similarly the Expected Hitting Distance defined in equation 14 can be solved with the following set of equations:

$$\mathbb{E}[D_{iA}] = \begin{cases} \sum_{j \in S} P_{ij} (d(i, j) + \mathbb{E}[D_{jA}]) & \text{if } i \neq A \\ 0 & \text{if } i = A \end{cases} \quad (16)$$

However, this system of equations grows exponentially with respect to the number of nodes. Thus we propose a novel approach to find the expected hitting distance of each node from the root.

To solve for the expected distance from the root to vertex  $A$ . Let  $Q$  denote all branches that can be reached from the root without having to go through  $A$ , and let  $R$  denote all branches that can only be reached from the root by going through  $A$ . Let us reorganize the probabilistic transition matrix into the following.

$$\mathcal{P}_B = \begin{pmatrix} Q & R \\ 0 & I_r \end{pmatrix} \quad (17)$$

To calculate the Expected Hitting Distance, we want to know the expected distance before we reach an absorptive state. Note by the formulation of the transient and absorptive states, the first absorptive state we reach is  $A$ . Thus we solve for the fundamental matrix:

$$N = \sum_{k=0}^{\infty} Q^k = (I - Q)^{-1} \quad (18)$$

**Theorem 7.** *The expected hitting distance can be solved by*

$$\mathbf{D}_A = (I - Q)^{-1} \mathbf{D} \quad \text{where} \quad \mathbf{D} = (0 \quad \ell(b_1) \quad \ell(b_2) \quad \dots \quad \ell(b_n)) \quad (19)$$

where  $\ell(b)$  represents the length of branch  $b$ .

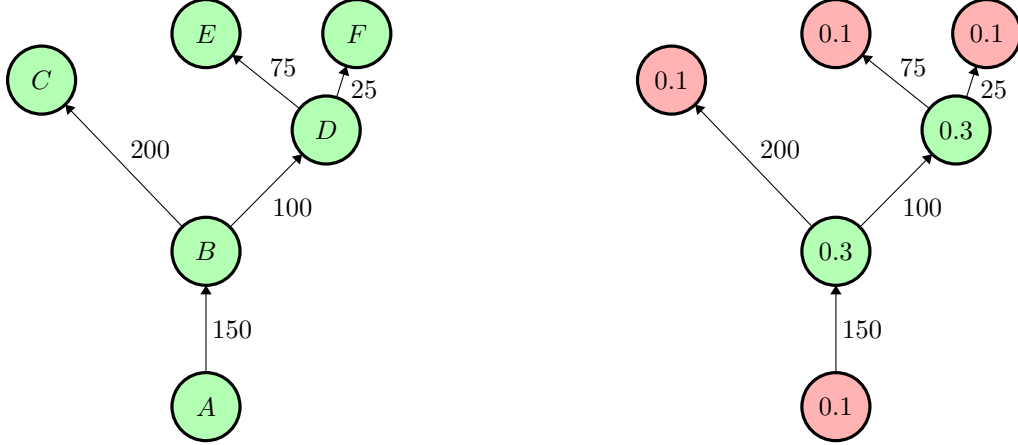


Figure 12: Stationary Distribution of Toy Example

### 3.3.1 Calculating Expected Hitting Distance to all nodes

Our proposed method which uses Geometric Features calculates the Expected Hitting Distance from the Root to all other Nodes in the Geometric Tree Graph. Since we want a normalized way to compare Geometric Trees, for each  $s \in \mathcal{S}$  we average over the expected distances in the original tree.

We will now define the new Markov Chain  $(\mathcal{S}_B, \mathcal{P}_B, s_0)$ . The  $s_0$  remains the same as in the original. All  $b = (s_i, s_{i+1}) \in \mathcal{S}_B$  such that  $(s_i, s_{i+1}) \in E$ , where  $E$  represents the edges of the orginal graph  $T$ .

The matrices for the toy example in figure 12 are as follows:

$$A \begin{pmatrix} A & B & C & D & E & F \\ 0 & 1 & 0 & 0 & 0 & 0 \\ 1 & 0 & 1 & 1 & 0 & 0 \\ 0 & 1 & 0 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 & 1 & 1 \\ 0 & 0 & 0 & 1 & 0 & 0 \\ 0 & 0 & 0 & 1 & 0 & 0 \end{pmatrix} \rightarrow B \begin{pmatrix} A & B & C & D & E & F \\ 0 & 1 & 0 & 0 & 0 & 0 \\ 1/3 & 0 & 1/3 & 1/3 & 0 & 0 \\ 0 & 1/3 & 0 & 0 & 0 & 0 \\ 0 & 1/3 & 0 & 0 & 1/3 & 1/3 \\ 0 & 0 & 0 & 1 & 0 & 0 \\ 0 & 0 & 0 & 1 & 0 & 0 \end{pmatrix} \rightarrow C \begin{pmatrix} A & B & C & D & E & F \\ 0.1 & 0.3 & 0.1 & 0.3 & 0.1 & 0.1 \end{pmatrix} \quad (20)$$

If we then consider the state space with  $\mathcal{S}$  with maximum outgoing degree 2 and maximum distance from root 3. The feature vector for the toy example in figure 12 is then:

$$\phi(X) = \left( \underbrace{0 \quad 0.1 \quad 0}_{\psi(i)=0} \quad \underbrace{0 \quad 0 \quad 0.3}_{\psi(i)=1} \quad \underbrace{0.1 \quad 0 \quad 0.3}_{\psi(i)=2} \quad \underbrace{0.2 \quad 0 \quad 0}_{\psi(i)=3} \right) \quad (21)$$

$$B = BD \begin{pmatrix} A & AB & BA & BC & CB & BD & DB & DE & ED & DF & FD \\ 0 & 1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\ 0 & 0 & 1 & 1 & 0 & 1 & 0 & 0 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 & 1 & 0 & 0 & 0 & 0 & 0 & 0 \\ 0 & 0 & 1 & 1 & 0 & 1 & 0 & 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 & 0 & 0 & 1 & 1 & 0 & 1 & 0 \\ 0 & 0 & 1 & 1 & 0 & 1 & 0 & 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 1 & 0 & 0 \\ 0 & 0 & 0 & 0 & 0 & 0 & 1 & 1 & 0 & 1 & 0 \\ 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 1 \end{pmatrix} \quad (22)$$

If we then consider the state space with  $\mathcal{S}$  with maximum outgoing degree 2 and maximum distance from root 3. The feature vector for the toy example in figure 12 is then:

$$\phi(X) = \left( \underbrace{0 \quad 0 \quad 0}_{\psi(i)=0} \quad \underbrace{0 \quad 0 \quad 150}_{\psi(i)=1} \quad \underbrace{1050 \quad 0 \quad 950}_{\psi(i)=2} \quad \underbrace{4000 \quad 0 \quad 0}_{\psi(i)=3} \right) \quad (23)$$

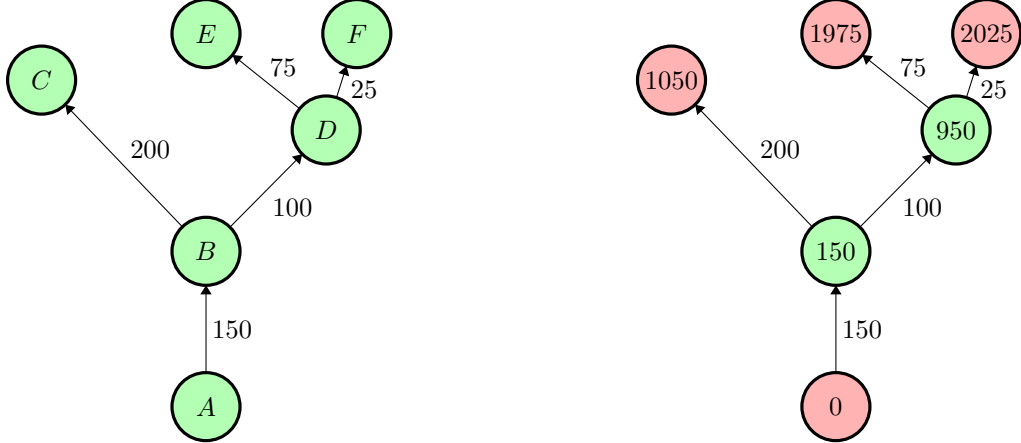


Figure 13: Expected Distance Representation of Toy Example

We then use  $\phi(X)$  as input to a RBF kernel SVM.

## 4 Empirical Evaluation

Here we test our methods against two different datasets. In table 2, we list different statistics of each dataset.

Dataset	Number of Graphs	Classes	Avg. Nodes	Avg. Edges
Synthetic	100	5	38.29	39.29
Real	100	5	55.84	56.84

Table 2: Statistics of the Synthetic and Real Dataset

### 4.1 Synthetic Data and Real Data

In this section we test our methods against the current state of the art methods in Geometric Tree Classification with our synthetically created dataset.

Method	Synthetic Dataset ( $\uparrow$ )	Real Dataset ( $\uparrow$ )
Markov Chain Maximum Likelihood ( <i>ours</i> )	<b>70.33</b> <sub>(9.304)</sub>	58.77 <sub>(1.026)</sub>
Stationary ('rbf', $p = 0.01$ ) ( <i>ours</i> )	<u>69.65</u> <sub>(9.145)</sub>	<b>68.84</b> <sub>(9.667)</sub>
Geometric Expected Hitting Distance ('rbf')( <i>ours</i> )	56.00 <sub>(7.461)</sub>	<u>62.81</u> <sub>(8.26)</sub>
Roothpath, Gaussian [8]	15.5 <sub>(0.212)</sub>	13.68 <sub>(5.367)</sub>
Graph Convolutional Network [10]	29.00 <sub>(8.699)</sub>	34.39 <sub>(10.677)</sub>
Graph Isomorphism Network [20]	24.67 <sub>(8.055)</sub>	26.49 <sub>(9.064)</sub>
Vertex Histogram Kernel	42.67 <sub>(10.71)</sub>	35.96 <sub>(10.141)</sub>
Spectral Baseline Method [5]	20.07 <sub>(3.392)</sub>	21.05 <sub>(4.611)</sub>

Table 3: Accuracy of our tree learning method and various state of the art algorithms for our Synthetic Dataset and Real Dataset

## 4.2 Experimental Details

Here we will describe the details of the baseline methods and current state of the art for Graph Classification. With the limited data, the neural networks, despite having a large amount of parameters, are unable to find good representations of the data that generalize well to the testing set.

## 5 Discussion and Further Research

As part of further work in tree classification from images, an algorithm to determine the graph structure of a tree is still an open problem when working with fine structure. For pixel-by-pixel segmentation, we estimate manual labeling to take close to three hours for each image. Semi-manual approaches will likely reduce the time significantly, [14]. Starting from a base segmentation with a foundation model, [11], would also likely save significant time, although the accuracy might suffer. Approaches to tree segmentation by 3d modeling in a software such as **Blender**, [4], will give good segmentation masks for synthetic trees, however the distributional shift from synthetic trees to real trees tends to be a difficult problem. One way to combat this issue could be to use a Style-GAN, [21]. The idea would be to i.) generate training data and masks with **Blender** for trees, then ii.) train a Cycle-GAN on the synthetic trees from **Blender** and the real trees from the original dataset and change the style of the synthetic trees to that of the real trees, and lastly iii.) train a segmentation model such as U-net [19] on the synthetic trees to learn the tree segmentation. After using one of these approaches, one might be able to use a deep reinforcement learning [13] model to create the center line of a tree after manually labeling graph data. Another approach would be to use standard centerline techniques on a binary mask of the tree to form the tree structure (points where the centerline meets would be vertices) [3].

We find our three Markov Chain Modeling Techniques as superior to other state of the art methods in graph classification. Typically, in images the size of the tree will not be very large, so we believe our methods can be used for a wide variety of applications in biomedical imaging. In general, it the stationary distribution method is very fast to run as for each tree it only needs to calculate the dominant eigenvector. The Expected Distance approach, although a strong approach, requires a calculation of the fundamental matrix for each subtree in the tree, so in the case of large trees it will not scale as well as the Stationary Distribution Approach.

In this paper, we introduce Markov Chain Modeling of Geometric Tree Graphs. We test our method against various baselines and state of the art methods. Thus, we believe modeling the geometric tree as a Markov Chain imposes a good inductive bias in the Geometric Tree Classification Problem and hope this work encourages researchers to see Markov Chain Modeling as a promising new direction in the problem of Geometric Tree Classification.

Our two datasets are small scale, since they have on average only 20 images per class it makes this a very difficult dataset for traditional learning algorithms. As with the limited amount of data, the Markov Chain Model is able to find a statistically significant good representation of the graph. However, it would be interesting to see how well our Markov-based methods fair on large scale datasets.

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## A Deferred Theory

In this section we will theoretically prove all claims made in the main text.

### A.1 Notation

Define  $\mathcal{S}$  as the state space,  $\mathcal{P}$  be all the one step transition probabilities. Let  $\mathbb{P}[s_i \rightarrow s_j] = \mathcal{P}_{s_i, s_j}$ , where  $s_i, s_j \in \mathcal{S}$ , be the transition probability between states  $s_i$  and  $s_j$ . Let  $\pi(s_i, s_j)$  be a path from  $s_i \rightarrow s_j$ . The number of the states in a path between  $s_i, s_j \in \mathcal{S}$  is given as  $|\pi(s_i, s_j)|$ . Using the above definitions, the notation  $\mathbb{P}[s_0 \rightarrow s_j || \pi(s_0, s_j) = k]$  represents the probability of a random walker going from state  $s_0 \in \mathcal{S}$  to another state  $s_j \in \mathcal{S}$  in a path with  $k$  vertices. Branches are can be considered as adjacent state pairs and denoted by the set  $\mathcal{B}$ . A branch  $b \in \mathcal{B}$  exists iff there exists  $s_1, s_2 \in \mathcal{S}$  s.t.  $\mathcal{P}_{s_1, s_2} > 0$  or  $\mathcal{P}_{s_2, s_1} > 0$ .

### A.2 Proof of Theorem 3

**Proof.** We want to calculate the probability a Tree  $T$  is from class  $c$ . We first start with Bayes' Theorem [2].

$$\arg \max_{c \in \mathcal{C}} \mathbb{P}[C = c | T] = \frac{\mathbb{P}[T | C = c] \mathbb{P}[C = c]}{\mathbb{P}[T]} \quad (24)$$

$$= \arg \max_{c \in \mathcal{C}} \frac{\mathbb{P}[T | C = c] \mathbb{P}[C = c]}{\mathbb{P}[T | C = c] \mathbb{P}[C = c] + \mathbb{P}[T | C \neq c] \mathbb{P}[C \neq c]} \quad (25)$$

$$= \arg \max_{c \in \mathcal{C}} \frac{\mathbb{P}[T | C = c] \mathbb{P}[C = c]}{\mathbb{P}[T | C = c] \mathbb{P}[C = c] + \sum_{j=1}^m \mathbb{I}\{j \neq c\} \mathbb{P}[T | C = j] \mathbb{P}[C = j]} \quad (26)$$

$$= \arg \max_{c \in \mathcal{C}} \frac{\mathbb{P}[T | C = c] \mathbb{P}[C = c]}{\sum_{j=1}^m \mathbb{P}[T | C = j] \mathbb{P}[C = j]} \quad (27)$$

let  $\mathcal{B}(T)$  represent set of branches in tree  $T$

$$= \arg \max_{c \in \mathcal{C}} \frac{\left( \prod_{b \in \mathcal{B}(T)} \mathbb{P}[b | C = c] \right) \mathbb{P}[C = c]}{\sum_{j=1}^m \left( \left( \prod_{b \in \mathcal{B}(T)} \mathbb{P}[b | C = j] \right) \mathbb{P}[C = j] \right)} \quad (28)$$

Let us here note that a branch is a sequence of states from the root to the branch of the markov chain

$$= \arg \max_{c \in \mathcal{C}} \frac{\left( \prod_{b \in \mathcal{B}(T)} \prod_{s_i \in b} \mathcal{P}_{s_{i-1}, s_i}^{(c)} \right) \mathbb{P}[C = c]}{\sum_{j=1}^m \left( \left( \prod_{b \in \mathcal{B}(T)} \prod_{s_i \in b} \mathcal{P}_{s_{i-1}, s_i}^{(c)} \right) \mathbb{P}[C = j] \right)} \quad (29)$$

$\mathcal{P}_{s_{i-1}, s_i}^{(c)}$  represents the transition probability from state  $s_{i-1}$  to  $s_i$  in the Markov Chain for the trees in class  $(c)$ . Then from the Principle of Maximum Likelihood.

$$= \arg \min_{c \in \mathcal{C}} \frac{\left( \sum_{b \in \mathcal{B}(T)} \sum_{s_i \in b} - \log \left( \mathcal{P}_{s_{i-1}, s_i}^{(c)} \right) \right) \mathbb{P}[C = c]}{\sum_{j=1}^m \left( \left( \sum_{b \in \mathcal{B}(T)} \sum_{s_i \in b} - \log \left( \mathcal{P}_{s_{i-1}, s_i}^{(c)} \right) \right) \mathbb{P}[C = j] \right)} \quad (30)$$

This completes the proof. ■

### A.3 Proof of Lemma 4

**Proof.** Recall  $\hat{\mathbf{P}} \triangleq (1 - \varepsilon)\mathbf{P} + \varepsilon\mathbf{R}$ . We will first prove  $\hat{\mathbf{P}}$  is an irreducible matrix, i.e. we will prove for all  $i \in [n], j \in [n]$ , there exists  $k \in \mathbb{N}$ , s.t.  $\hat{\mathbf{P}}_{i,j}^k > 0$ . After adding  $\mathbf{R}$ , we know have  $\mathbb{P}[s_i \rightarrow s_0 | 1] = \varepsilon$ . Thus we have,

$$\mathbb{P}[s_i \rightarrow s_j || \pi(s_i, s_j) = n] \geq \mathbb{P}[s_i \rightarrow s_0 || \pi(s_i, s_0) = 1] \cdot \mathbb{P}[s_0 \rightarrow s_j || \pi(s_0, s_{j-1}) = n-1] = \varepsilon \mathcal{P}_{0,j} > 0 \quad (31)$$

when we choose  $n$  s.t. the distance from the root to  $s_j$  is equal to  $n - 1$ . We have therefore proved  $\hat{\mathbf{P}}$  is irreducible. We will now prove aperiodicity. It suffices to prove  $\mathbb{P}[s_i \rightarrow s_j | n+1] > 0$ .

$$\mathbb{P}[s_i \rightarrow s_j || \pi(s_i, s_j)] = n + 1 \quad (32)$$

$$\geq \mathbb{P}[s_i \rightarrow s_0 || \pi(s_i, s_0)] = 1 \cdot \mathbb{P}[s_0 \rightarrow s_i | 1] \cdot \mathbb{P}[s_0 \rightarrow s_0 || \pi(s_0, s_0)] = 1 = \varepsilon^2 \mathcal{P}_{0,j} > 0 \quad (33)$$

Thus we have proved  $\mathbb{P}[s_i \rightarrow s_j | n] > 0$  and  $\mathbb{P}[s_i \rightarrow s_j | n+1] > 0$ , we also have the greatest common divisor for  $n$  and  $n+1$  is equal to 1. Therefore there exists no  $t$  s.t.  $\mathbb{P}[s_i \rightarrow s_j | tn] = 0$ . Thus  $\hat{\mathbf{P}}$  is aperiodic. This completes the proof.  $\blacksquare$

## A.4 Proof of Theorem 7

**Proof.** From the definition of the Expected Hitting Distance given in Definition 6, we have

$$\mathbb{E}[D_{iA}] = \mathbb{E}[D_A | \text{initial state is } s_0] \quad (34)$$

$$= \sum_{j=1}^{|\mathcal{S}|} \mathcal{P}_{s_i, s_j} (d(s_i, s_j) + \mathbb{E}[D_{s_j, A}]) \quad (35)$$

$$= \sum_{j=1}^{|\mathcal{S}|} \mathcal{P}_{s_i, s_j} \left( d(s_i, s_j) + \left( \sum_{k=1}^{|\mathcal{S}|} \mathcal{P}_{s_j, s_k} (d(s_j, s_k) + \dots) \right) \right) \quad (36)$$

$$= \sum_{j=1}^{|\mathcal{S}|} d(s_i, s_j) \mathcal{P}_{s_i, s_j} + \sum_{j=1}^{|\mathcal{S}|} \sum_{k=1}^{|\mathcal{S}|} \mathcal{P}_{s_i, s_j, s_k} d(s_j, s_k) + \dots \quad (37)$$

$$\stackrel{(a)}{=} \sum_{i=1}^{|\mathcal{S}|} \sum_{j=1}^{|\mathcal{S}|} d(s_i, s_j) \sum_{k=1}^{\infty} \mathbb{P}[s_i \rightarrow s_j] \mathbb{P}[s_0 \rightarrow s_i || \pi(s_0, s_i)] = k - 1 \quad (38)$$

$$\stackrel{(b)}{=} \sum_{i=1}^{|\mathcal{B}|} \ell(b_i) \sum_{k=1}^{\infty} \mathbb{P}[b_0 \rightarrow b_i || \pi(b_0, b_i)] = k \quad (39)$$

$$\stackrel{(c)}{=} \sum_{i=1}^{|\mathcal{B}|} \ell(b_i) \sum_{k=1}^{\infty} (\mathbf{Q}^k)_{1,i} \stackrel{(d)}{=} \sum_{i=1}^{|\mathcal{B}|} \ell(b_i) \left( (\mathbf{I} - \mathbf{Q})^{-1} \right)_{1,i} \stackrel{(e)}{=} (\mathbf{I} - \mathbf{Q})_{(1,:)}^{-1} \mathbf{D} \quad (40)$$

In (a), we expand out all the inner summations we always have the distance multiplies by the probability of some path, we make the observation this path starts at  $s_i$  and ends at  $s_j$ . Furthermore, we also make the note this covers all possible paths as we are summing over all possible states. In (b), we convert everything to our branch notation to simplify the analysis. In (c), if we let  $\mathbf{Q}$  be defined as in the original theorem statement, then we have  $(\mathbf{Q}^k)_{i,j}$  is the probability after  $k$  steps, a random walker in branch  $i$  will be in branch  $j$ . In (d), since  $\mathbf{Q}$  is a probabilistic matrix, therefore  $\|\mathbf{Q}\| = 1$ , thus we have

$$\sum_{k=1}^{\infty} \mathbf{Q}^k = (\mathbf{I} - \mathbf{Q})^{-1} \quad (41)$$

In (e), we define  $\mathbf{D}$  as the vector of branch lengths given in the Theorem 7 statement. This completes the proof.  $\blacksquare$

## B Reproducability

In this section we give the neural network and graph neural network architectures used in the experiments in § 4.1.

### B.1 Convolutional Neural Network Architectures

Here we provide the architecture used for image classification for the 5 classes in section 1. We also resize all images from  $600 \times 800 \times 3$  to  $224 \times 224 \times 3$ . Trained with ADAM optimizer [9],  $\alpha = 0.0001$ , momentum is set to 0.1 or 0.9, batch size is 16 or 64, and trained for 25 – 50 epochs.

Table 4: Convolutional Neural Network

<b>Layer</b>	<b>Filter</b>	<b>Depth</b>	<b>Stride</b>	<b>Padding</b>	<b>Activation</b>
Convolution	$11 \times 11$	96	(4, 4)	(1, 1)	ReLU
Max Pooling	$3 \times 3$	-	2	0	-
Convolution	$5 \times 5$	256	(1, 1)	(2, 2)	ReLU
Max Pooling	$3 \times 3$	-	2	0	-
Convolution	$3 \times 3$	384	(1, 1)	(1, 1)	ReLU
Convolution	$3 \times 3$	384	(1, 1)	(1, 1)	ReLU
Convolution	$3 \times 3$	256	(1, 1)	(1, 1)	ReLU
Dropout	$p = 0.5$	-	-	-	-
Linear	-	2048	-	-	ReLU
Dropout	$p = 0.5$	-	-	-	-
Linear	-	4096	-	-	ReLU
Dropout	$p = 0.5$	-	-	-	-
Linear	-	4096	-	-	ReLU
Softmax	-	5	-	-	-

## B.2 Graph Neural Network Architectures

Here we provide the architectures used for the Graph Convolutional Network and the Graph Isomorphism Network for the experiments in section 4.1.

Table 5: Graph Neural Network Architectures

(a) GCN Network			(b) GIN Network		
Layer	Dim	Act	Layer	Dim	Act
GCNConv	$2 \times 250$	ReLU	Linear	$2 \times 250$	ReLU
GCNConv	$250 \times 250$	ReLU	Batch Normalization	-	-
GCNConv	$250 \times 250$	ReLU	Linear	$250 \times 250$	ReLU
Global Mean Pool	-	-	GINConv	-	-
Dropout	$p = 0.5$	-	Global Add Pool	-	-
Linear	$250 \times 5$	-	Linear	$250 \times 250$	-
Log Softmax	$5 \times 1$	-	Batch Normalization	-	ReLU
			Linear	$250 \times 250$	ReLU
			GINConv	-	-
			Global Add Pool	-	-
			Linear	$250 \times 250$	-
			Batch Normalization	-	ReLU
			Linear	$250 \times 250$	ReLU
			GINConv	-	-
			Global Add Pool	-	-
			Linear	$750 \times 750$	ReLU
			Dropout	$p = 0.5$	-
			Linear	$750 \times 5$	-
			Log Softmax	$5 \times 1$	-

## C Geometric Tree Generation Interface

We also provide details on our software implementation of a Geometric Tree Generation Interface. We provide details on the general user cycle.

1. Upload folder of tree images to the software.
2. Select a folder to save graph annotations to.
3. Annotate the Graph

There are two modes the user can be in while annotating the graph. Either in *parent* mode or not in *parent* mode. If the user is in *parent* mode, then either the user can place a new node which will be a *parent*, or if the user clicks within 5 pixels of another node, then that node will become a parent. After you click on parent mode, you will be taken out of *parent* mode. When you are not in parent mode, any node you click will have an edge from the *parent* node.

If you want to delete any nodes, click middle-button on the mouse and it will delete the whole subtree of the node within 5 pixels of your click point.

4. Save Graph

The graph is saved in a pickle file to the save directory the user already chose as an adjacency list. The keys in the adjacency list are the pixel locations of the nodes, and the list is the pixel locations of the neighbor nodes.

The software is free to download at [https://github.com/caped-doshi/tree\\_identification](https://github.com/caped-doshi/tree_identification)