The Palm Vein Graph
Feature Extraction and Matching

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Abstract: We present a graphical representation for palm vein patterns for use as biometric identifiers. The palm vein image captured from an infra red camera is converted into a spatial graph. After image enhancement and binarisation, the palm vein features are extracted from the skeleton using a novel two stage spur removal technique. The location of the features and the connections between them are used to define a Palm Vein Graph. Palm vein graphs are compared using the Biometric Graph Matching (BGM) Algorithm. We propose a graph registration algorithm that improves over existing state of the art algorithms for graph registration. We introduce 10 graph topology-based measures for comparing palm vein graphs. Experiments are conducted on a public palm vein database. One of the introduced measures, an edge-based similarity, gave a definite improvement in matching accuracies over other published results on the same database, especially for samples with only a small common overlap area due to displacement. In addition, when the edge-based measure was combined with one of three other topological features, we demonstrate a further improvement in matching accuracy.

1 INTRODUCTION

Palm vein biometrics refers to the use of the unique vascular pattern in the human palm for authentication. Palm veins have the following advantages shared with most vascular biometrics:
1. Internal location, which makes it hard to to spoof or covertly acquire from an individual.
2. In-built liveness detection in the capture process. Palm-vein images are captured using infra-red (IR) cameras. The palm vein images are created by the property that infra red light is absorbed by the de-oxygenated haemoglobin creating dark regions corresponding to the veins. The rest of the human tissues and skin reflect the IR creating bright regions in the image. This property ensures that a live subject is available for biometric capture.
3. The capture is harmless and often contactless.
4. The capture process is non-intrusive with respect to an individual’s personal space and is therefore more socially acceptable.
5. A very low failure to enrol (FTE) rate.

Palm vein has been identified as the easiest (Watanabe et al., 2005) vascular biometric to work with as the palm area is free of hair and skin color variations thereby reducing capture noise. Also the palm vein pattern is very detailed and intricate and covers a larger area than other vascular biometrics like the dorsal vein, finger vein and wrist vein.

In this paper, we investigate the palm vein patterns captured in the PUT database (Kabacinski and Kowalski, 2011). The palm vein patterns are represented as biometric graphs. Biometric graphs have been used to represent fingerprints (Horadam et al., 2011), retina patterns (Lajevardi et al., 2013) and dorsal hand vein patterns (Lajevardi et al., 2014). The palm vein graphs are compared using Biometric Graph Matching (Lajevardi et al., 2013). The main contributions of this work are:
1. A novel image feature extraction process that creates a palm vein graph representing the vascular structure.
2. An improved registration algorithm that enhances the speed and registration accuracy compared to the algorithm in (Lajevardi et al., 2014).
3. Graph-based distance measures that effectively separate genuine and imposter comparisons of palm vein patterns.
2 IMAGE PREPROCESSING AND GRAPH EXTRACTION

2.1 The Database

This research uses palm vein images captured by researchers at the Poznan University of Technology (PUT) (Kabacinski and Kowalski, 2011). The database has vein pattern images from the left and right palms of 50 individuals. Each individual gave 12 samples of each palm, totalling 24 samples. Each set of 12 samples was acquired over 3 sessions with 4 images each, at least one week apart. The database was chosen for two main reasons - a large number of samples per individual palm (12 images) and 3 separate sessions, one week apart. Several other databases exist in the literature (Kumar and Prathyusha, 2009), (Shahin et al., 2007), (Wang et al., 2007), but they all take samples within one session. The across session comparison is valuable to understand the natural variation that will occur in a practical scenario where individuals leave after enrolment and return after a period to be verified without remembering the exact location of their hand at enrolment.

2.2 Image Processing and Feature Extraction

The infra red capture process often generates noisy images with low contrast between the vein pattern and background. The challenges of feature extraction in the PUT database include noise of wrinkles and other skin features on the hand, non-uniform background brightness and palm displacement within and across the sessions. In order to extract features from the palm vein image, specific image processing steps are required to enhance the vein pattern. We used Discrete Fourier Transform based method, gradient-based method and thresholding method (Kabacinski and Kowalski, 2010; Wenxiong and Qiuxia, 2014; Gaikwad and Narote, 2013) to extract the vein pattern and found that they failed to provide the continuous rich structure of palm vein in the PUT database. This paper has modified and combined several noise reduction and vein enhancement methods for the better performance of palm vein graph extraction.

The automatic processing of graph feature extraction from palm vein images can be summarized in the following steps - Image pre-processing and vein enhancement; image binarisation; and skeleton and graph feature extraction. It is important to identify the Region of Interest (ROI): the spatial region where the vein patterns are most visible and useful. A well defined ROI will enable a good registration between compared palm vein graphs and will hence improve overall processing time. However, the PUT database has shown large displacements in captured images, especially between the sessions. In fact, the region in one image may only be partially present in another. In addition, if the ROI is too small, it limits the number of features that can be extracted from the image. This paper selects the entire captured image area as the ROI. Then, 40 pixels inwards from the border was masked to eliminate possible misidentification of the vein.

Image pre-processing and vein enhancement aim to remove the artifacts and increase the contrast between vein and background. The palm vein image was first converted to greyscale. To avoid amplifying the noise in the image, especially in homogeneous areas, contrast-limited adaptive histogram equalization (Zuiderveld, 1994) was applied on a small window (8x8 pixels) to make image contrast uniform at different regions. The noise in the image was reduced using an anisotropic diffusion filter (Perona and Malik, 1990). Linear anisotropic diffusion performs image smoothing and restoration that minimizes the noise without removing significant components of the image. An example of the original image and vein-enhanced image is shown in Figures 1(a) and 1(b).

The next step is image binarisation. Skeleton extraction requires image binarization to distinguish vessel and background. The process of binarization was described previously in (Lajevardi et al., 2013). Briefly, a family of 2D matched filters was used to search for vessel segments along all possible directions in frequency domain. The maximum responses over all possible orientations were calculated for each

Figure 1: This figure shows the stages that a palm vein image goes through before feature extraction. (a) (top-left) Original image; (b) (top-right) Palm vein enhanced image; (c) (bottom-left) Image after binarisation; (d) (bottom right) Skeleton extracted from binarised image.
pixel position. A threshold was chosen based on empirical observation that the veins occupy around 15% of the pixels in the image. The image was binarised using this optimum threshold. Then the binarised image was morphologically dilated to connect broken pixels. Isolated objects, in which the connected components were less than 200 pixels in area, were removed. A binarized image is shown in Figure 1(c).

The final step is skeleton and graph feature extraction. The skeleton was extracted from the binary image by morphologically removing pixels on the boundaries and keeping pixels connected, shown in Figure 1(d). Graph features of vertices and edges were then detected from the skeleton.

### 2.2.1 The Biometric Graph

The biometric graph is a spatial graph with features in a biometric represented as vertices and the connections between those features represented as edges. Often the spatial coordinates of the features become the spatial attributes of the corresponding vertices. In a vascular biometric, the vein bifurcations and crossovers represent the vertices and their locations in the image represent their attributes. A pair of vertices are connected by an edge if their features were linked by a vein in the original vein pattern. The biometric graph representation for retina (Lajevardi et al., 2013) and handvein (Lajevardi et al., 2014) have been shown to be extremely effective when compared to other existing templates. Mathematically, a biometric graph is represented as $g = (V,E,\mu,\nu)$, where $V$ represents a set of vertices, $E$ represents the set of edges between the vertices, $\mu$ is the vertex labeling function and $\nu$ is the edge labeling function. For the palm vein graphs in this research, $\mu$ associates each vertex with its unique two dimensional spatial coordinate while $\nu$ associates each edge with attributes of edge length and slope. The main advantages of using a graphical representation is that it is concise and modular. There are no limits to the vertex, edge or graph attributes that can be added to the graph. Thus additional features deemed important for representing the biometric in the future can easily be added on the existing graph representation as new attributes.

To obtain the palm vein graph after skeletonisation, a 3 by 3 window was applied on each pixel of the skeleton to test if it was an end or a junction (crossing point or branching point). All junction and end points were marked as vertices of the graph. An edge is the link between two vertices. Edges were detected by neighborhood tracking along the skeleton. All possible branches were tracked from the junction vertex.

Palm vein graph features represent the vascular pattern with limited vertices and edges. The undesirable spurs caused by noise and tiny veins have an impact on the registration and matching process. There were two stages to remove undesirable spurs. Firstly, morphological operations were applied to remove undesirable short spurs on the skeleton. However, removing spurs also reduces the vertices on the major skeleton that may alter the shape of the skeleton. Therefore, the threshold for morphological operations involved in defining short spurs was set to 10 pixels to ensure correct representation of the vein shape. Secondly, there were a number of short to medium spurs in the skeleton due to palm principal lines and wrinkles. These lines and wrinkles vary in the captured images at different sessions. Palm principal lines and wrinkles can be considered as part of the biometric and are difficult to completely remove; our experiments have shown that removing the short-medium spurs improves the process of registration and matching. The short-medium spurs were removed after the detection of vertices and edges. We used the following algorithm: 1) calculate the lengths of all branches. 2) identify the short-medium branches that are not linked to the end vertices (these branches should not be removed) 3) the short-medium branches, that are linked to end vertices and whose lengths are less than 40 pixels are removed. The connecting vertex was kept to form the correct vein shape while the edge and
Figure 3: Average Degree Distribution of Palm Vein Graphs.

the vertex at the end of the edge was removed.

Examples of palm vein graphs from an individual captured at two different sessions are shown in the right panels of Figure 2(a) and Figure 2(b). The Palm vein graphs from the PUT database have an average of 101 vertices and 112 edges. The average edge to vertex ratio is 1.11 and the average size of the largest component is 91. The average size of the largest component is almost equal to the average size of the palm vein graph. This indicates that the image processing can retrieve most of the features and connecting veins in a given image. Smaller disconnected components exist in a palm vein graph because the quality of the image in certain parts is not good enough to retrieve the connecting veins. Figure 3 shows the degree distribution of the Palm Vein Graphs.

3 BIOMETRIC GRAPH MATCHING

Biometric Graph Matching (BGM) is a two-stage algorithm for the error-tolerant comparison of biometric graphs and is described in Lajevardi et al (Lajevardi et al., 2013). The first stage is registration and the second is error tolerant graph matching. Registration is the process of bringing a pair of compared graphs into the same reference frame. The Iterative Closest Point (ICP) algorithm (Chen et al., 2009) and the Modified Hausdorff Distance (MHD) (Dubuisson and Jain, 1994) are algorithms commonly used to register point clouds. It has been demonstrated by Lajevardi et al (Lajevardi et al., 2014) that the registration process in BGM is either as good as or does better than these standard algorithms, especially when the graphs are small. This paper uses a registration algorithm that is a slight modification of Algorithm1 in Lajevardi et al (Lajevardi et al., 2013). The modified algorithm is denoted here as Algorithm 2 (see comparison results in Table 1). The registration process in Algorithm 1 translates and rotates a pair of graphs based on a corresponding pair of edges that define the coordinate system. Every pair of edges with one edge from each compared graph, is given a score based on a function of their edge attributes of length and slope. The edge pairs are ranked on this score. 2% of the top ranked edges are then shortlisted for the registration process. For each of the shortlisted edge pairs, the two graphs are translated and rotated to make the start vertex of each edge the origin and the end vertex indicate the positive x-direction. The vertex labels of both the translated and rotated graphs are recalculated based on the new orientation. The Euclidean distance between the vertex labels, which in this case are the cartesian coordinates of the vertex positions, are compared to count the number of vertex pairs that lie within a tolerance $\epsilon$ of each other. The edge pair that gets the largest number of vertex pairs within $\epsilon$ is chosen as the edge pair to register the graphs and the corresponding transform (translation and rotation) is taken as the correct transform to achieve the best registration.

On implementing Algorithm 1 for registering the palm vein graphs, we found one drawback that caused several genuine palm vein samples to be misaligned. Most edge pairs in the palm vein graph were short and shorted pairs often scored a high rank compared to longer pairs. This caused longer pairs that gave a better registration to not appear on the top 2% shortlist. To overcome this, we modified Algorithm1 to split the set of edge pairs into long and short edge pairs. The mean of the medians of the edge lengths in the two graphs was selected as the threshold. If both edges of an edge pair had length greater than this threshold, the edge pair was categorised as long. All other edge pairs were labeled as short. The new shortlist comprised the top 2% of long edge pairs and the top 2% of short edge pairs. This modification ensured that long edge pairs that potentially gave better alignment could be included in the shortlist to get a better registration of the graphs. This is called as Algorithm 2 in this paper.

Once a pair of graphs is registered, the graph matching process gives a numerical measure of the distance between the pair of graphs by using the topological properties of a graph edit path-based induced subgraph. The error tolerant graph matching is adapted from that proposed by Riesen and Bunke (Riesen and Bunke, 2009). Their proposed algorithm computes a graph edit distance and graph edit path between a pair of compared graphs by treating the two graphs as two vertex sets of a bipartite graph.
Every primary vertex is mapped to one secondary vertex using the Hungarian optimisation algorithm with a cost function based on the vertex attributes. In this paper, the cost function between a pair of vertices is the Euclidean distance between the vertex labels and is described in (Lajevardi et al., 2013).

The graph matching process is simplified to a set of edit operations to convert the first graph (source graph) to the second graph (destination graph). The edit operations allowed are insertion, deletion and substitution of vertices. Each vertex in the source graph has a fixed cost associated with deleting it \((D(n) \times \alpha)\) where \(D(n)\) is the degree of the vertex \(n\) and \(\alpha\) is called the insertion and deletion cost and is a non zero integer. If \(n\) is a vertex in the destination graph, the cost of inserting it is also \(\alpha\). In this research we keep the insertion cost and deletion cost the same \((\alpha)\), however, it is possible to have different costs. Once the Hungarian algorithm computes the cheapest edit path to convert the source graph to the destination graph, this edit path is used to build the maximum common subgraph (mcs). The vertices of the destination path could be associated with two operations from the edit path - substitutions of vertices from the source graph and insertions. The mcs is a vertex induced subgraph of the destination graph. In particular, the mcs will comprise only those vertices of the destination graph that were associated with substitutions in the cheapest edit path. An edge will exist in the mcs if the edge existed in corresponding vertex pairs of both the source and destination graphs. Figure 4 shows the mcses from a genuine graph comparison and an imposter graph comparison. Note the difference in topology of the two types of mces. Mces from a genuine comparison are observed to have more vertices and more edges with larger connected components than the mces from an imposter comparison. It is this difference in topology that will be used to define similarity scores to distinguish genuine and imposter comparisons.

4 GRAPH FEATURES

The topological features of the mcs are used to differentiate the comparisons of samples from the same palm and samples from different palms. Let \(g_1 = (V_1, E_1, \mu, v)\) and \(g_2 = (V_2, E_2, \mu, v)\) be the two graphs compared using the BGM algorithm and \(g = (V, E, \mu, v)\) be the mcs of the graphs. Let \(C_1\) and \(C_2\) be the vertex counts of the largest and second largest components, \(c_1\) and \(c_2\) in \(g\).

Similarity scores based on the graph features are defined as follows:

1. Normalised vertex count, \(S_n = \frac{|V|}{\sqrt{|V_1| \times |V_2|}}\)
2. Normalised edge count, \(S_e = \frac{|E|}{\sqrt{|E_1| \times |E_2|}}\)
3. Edge to vertex ratio, \(S_p = \frac{|E|}{|V|}\)
4. Size of the largest component, \(S_{i1} = \frac{C_1}{\sqrt{|V_1| \times |V_2|}}\)
5. Size of the second largest component, \(S_{i2} = \frac{C_2}{\sqrt{|V_1| \times |V_2|}}\)
6. Ratio of the number of isolated vertices in the mcs to the number of connected vertices, \(S_i\)
7. Total length of the edges in the largest component in the mcs. Let \(L()\) be a function that takes a graph as input and returns the sum of the lengths of the edges in the graph. Then \(S_l = \frac{L(g_1)}{L(g_1) \times L(g_2)}\)
8. sum of the sizes of two largest components in the mcs, \(S_{i1i2} = \frac{C_1 + C_2}{\sqrt{|V_1| \times |V_2|}}\)
9. Edge to vertex ratio of the largest component, \(S_{p1}\)
10. Ratio of the number of vertices with degree greater than 3 in the mcs, to the total number of vertices in the mcs, \(S_d\)

We investigate if one or more of these features can be combined to achieve maximum separation between genuine and imposter scores of comparisons from the palm vein database.
Table 1: Comparison of Registration Algorithms Algorithm 1 (Lajevardi et al., 2013) and Algorithm 2, proposed in this paper.

<table>
<thead>
<tr>
<th>Setting</th>
<th>Algorithm 1</th>
<th>Algorithm 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Best $\varepsilon$</td>
<td>6</td>
<td>6</td>
</tr>
<tr>
<td>Average time (sec)</td>
<td>15.99</td>
<td>25.94</td>
</tr>
<tr>
<td>EER (as %)</td>
<td>2</td>
<td>0</td>
</tr>
</tbody>
</table>

5 EXPERIMENT

The PUT database has for left and right hands respectively, 600 palm vein images from 50 individuals, giving 6600 genuine comparisons across all sessions, 1800 genuine comparisons within same sessions and 352800 imposter comparisons. To tune the parameters in the experiment, we chose a random sample of 50 genuine comparisons and 50 imposter comparisons from all possible genuine and imposter comparisons. This set of 100 comparisons is called the training set and will be consistently used throughout all the experiments described in this paper. The BGM algorithm has two parameters to be tuned, the tolerance $\varepsilon$ in the registration algorithm and the insertion and deletion cost $\alpha$ in the graph matching algorithm. The first experiment was to compare the registration algorithm by Lajevardi et al (Lajevardi et al., 2013) with the improved algorithm, Algorithm 2, presented in this paper. The graph pairs in the testing set were aligned first using Algorithm 1 and then Algorithm 2 and in each case, the number of vertex pairs that lay within a tolerance $\varepsilon$ of each other were counted. A distance measure based on the Similarity score $S_e$ (Section 4. Score 1), given by $d_{min} = 1 - S_e$ is computed from the number of common vertex pairs and the sizes of the two graphs compared. The $d_{min}$ values between genuine and imposter comparisons in the training set are used to define score distributions to compare the alignment performance of the two algorithms. This experiment is run over range of $\varepsilon$ values between 0 and 0.1 to find the $\varepsilon$ value that gave the lowest EER for each algorithm. The Equal Error Rates (EER) at the best $\varepsilon$ value for each algorithm and the average registration times are presented in Table 1.

Setting $\varepsilon = 6$ and using Algorithm 2 for registration, the BGM algorithm was run on all the graphs from the palm vein database, excluding the comparisons used in the training set. BGM was run using a range of $\alpha$ values to determine the parameters that best separated the genuine and imposter scores. It was found that $\alpha = 11$ best compensated for the variations within samples of the same hand in determining the graph edit path when comparing pairs of graphs.

The BGM algorithm was run on Left and Right hands were tested separately. There were two distinct types of experiments based on the type of genuine comparisons made. First, Across Session Genuines using all the genuine comparisons including those across sessions and the second Within Session Genuines, with genuine comparisons taken only from within the same session. In both types of experiments, distances between a pair of graphs was measured using each of the 10 topological measures listed in Section 4. Table 2 shows the EER based on the the 10 topological features for across session and within session experiments for left and right hands. The Table 2 shows that $S_e$ does the best job of separating the genuine comparisons from the imposters.

The next step was to determine if any combination of similarity scores could improve the performance compared to using a single similarity measure. To do this, first the pairwise Spearman’s correlation coefficient between the 10 features was calculated. We found that most of the topological features were strongly correlated, with correlation coefficients between 0.8 to 0.95. In fact only $S_{p1}$ showed moderate correlations ranging between 0.58 and 0.78 with the other features. Nevertheless, in the absence of perfect correlation between features, there is a potential that pairwise combination of $S_e$ with one of the other features could produce better matching results. To test this hypothesis, $S_e$ was combined with every other feature to give 9 different pairings of topological features. For every pair of features, the following experiment was done. A Support Vector Machine (SVM) was used to build a classifier with a radial basis function (RBF) kernel that was tuned on the score pairs from the training set to determine the best parameters for the RBF kernel. The remaining comparisons on the database were divided into 10 parts and a 10 fold test was conducted where the SVM classifier was trained on 9 parts of the data and tested on the one other part. The false match rate (FMR), false non match rate (FNMR) and total misclassification error (TE) were computed in every fold. The average over 10 folds was taken as the matching performance using the chosen pair of features. The results for the 9 pairings are shown in the Table 4.

6 RESULTS AND DISCUSSION

Table 1 shows that Algorithm 2 significantly improves the registration process over Algorithm 1 by Lajevardi et al (Lajevardi et al., 2013), evidenced by the 0%
Table 2: Equal Error Rate as %, when comparing genuine and imposter palm vein graphs using each of the 10 graph measures.

<table>
<thead>
<tr>
<th>Topological Measure</th>
<th>Across Session Genuine, Left</th>
<th>Within Session Genuine, Left</th>
<th>Across Session Genuine, Right</th>
<th>Within Session Genuine, Right</th>
</tr>
</thead>
<tbody>
<tr>
<td>( S_0 )</td>
<td>4.18</td>
<td>2.02</td>
<td>4.17</td>
<td>2.13</td>
</tr>
<tr>
<td>( S_e )</td>
<td><strong>2.71</strong></td>
<td><strong>1.11</strong></td>
<td><strong>2.55</strong></td>
<td><strong>1.23</strong></td>
</tr>
<tr>
<td>( S_0 )</td>
<td>5.32</td>
<td>2.9</td>
<td>5.27</td>
<td>3.42</td>
</tr>
<tr>
<td>( S_{c1} )</td>
<td>6.06</td>
<td>3.91</td>
<td>6.63</td>
<td>4.39</td>
</tr>
<tr>
<td>( S_{c2} )</td>
<td>7.33</td>
<td>5.49</td>
<td>7.76</td>
<td>5.58</td>
</tr>
<tr>
<td>( S_l )</td>
<td>8.43</td>
<td>5.66</td>
<td>8.25</td>
<td>5.93</td>
</tr>
<tr>
<td>( S_l )</td>
<td>5.64</td>
<td>3.5</td>
<td>6.59</td>
<td>4.19</td>
</tr>
<tr>
<td>( S_{c12} )</td>
<td>4.45</td>
<td>2.51</td>
<td>4.79</td>
<td>2.91</td>
</tr>
<tr>
<td>( S_{2l1} )</td>
<td>19.22</td>
<td>19.96</td>
<td>13.04</td>
<td>12.67</td>
</tr>
<tr>
<td>( S_d )</td>
<td>13.33</td>
<td>9.9</td>
<td>11.5</td>
<td>8.57</td>
</tr>
</tbody>
</table>

EER for Algorithm 2. This demonstrates that for palm vein graphs, a better registration could be achieved by considering longer edge pairs which represent longer vein segments that were often reliably captured across samples. The algorithm takes longer on average due to the additional step of separation of long and short edges.

Table 2 shows that for \( S_e \), average EER for left and right palms is 2.63% for across session comparisons and this markedly improves on the published benchmark of 3.86% (Kabacinski and Kowalski, 2011) on the same database. BGM performs on par with the published result 1.11% for within session comparisons using \( S_e \). Note the large difference between EERs of across session and within session data. Genuine samples of palm veins captured from different sessions tend to have large displacement in captured area. This reduces the common overlap area between genuine samples and hence causes a lower score. Comparing our results to the published benchmark illustrates that BGM performs better than the benchmark when palm vein samples have only a partial overlap area. Five out of the 10 features give competitive EERs of under 5% for within session comparisons while three of these features \( (S_e, S_c, S_{c12}) \) also give competitive EERs for across session comparisons.

Table 4 shows that there are three topological features, that when combined with \( S_e \) give a competitive matching performance with total misclassification rate under 1.9%. To understand how much better feature combination does over using the single measure \( S_e \), we adopt the following procedure. From the ROC curve when using single feature \( S_e \) (see Figure 5) we locate the FNMR when using \( S_e \) at a FMR value, close to that obtained using two features. If the FNMR value using two features is actually lower than the corresponding one feature value, that particular pair of features gives a better result than using a single feature for comparison. The performance comparison between two features and one feature \( S_e \) is shown in Table 3.

Table 3 shows that the topological feature pairs \( (S_e, S_{c1}) \), \( (S_e, S_l) \) and \( (S_e, S_{c12}) \) do better than using \( S_e \) alone, with \( (S_e, S_l) \) showing the maximum gain in performance.

7 CONCLUSION

This paper proposes a graph-based approach to represent a palm vein template and proposes the Biometric Graph Matching algorithm (BGM) to match the templates. The method is tested on the PUT public palm vein database where each palm vein has 12 samples captures across 3 sessions. This gave us a good number of genuine comparisons that were within the same
The paper shows that using the graph representation and BGM matching approach for palm vein graphs is practical and efficient. The graph-based measure based on edge count outperforms the measure based on vertex count. This reinforces the advantage of using the edge information in representing the palm vein graph. We further demonstrate that by combining graph features, the matching performance can be improved over using just a single feature. The ability to explore and combine different graph features based on the spatial characteristics of the palm vein graph is one of the biggest advantages of using the graph-based approach.

Table 3: Comparison of matching performance when using a pair of features and a single feature $S_i$. The rows where the FNMRs are in boldface show the feature pairs whose performance is better than using $S_i$ alone.

<table>
<thead>
<tr>
<th>Features</th>
<th>Left Palm</th>
<th></th>
<th>Right Palm</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>FMR</td>
<td>FNMR</td>
<td>TE</td>
<td>FMR</td>
</tr>
<tr>
<td>$S_r, S_n$</td>
<td>0.37 (±0.07)</td>
<td>4.72 (±0.38)</td>
<td>2.34 (±0.17)</td>
<td>0.38 (±0.06)</td>
</tr>
<tr>
<td>$S_r, S_p$</td>
<td>0.52 (±0.09)</td>
<td>4.02 (±0.35)</td>
<td>2.1 (±0.13)</td>
<td>0.36 (±0.08)</td>
</tr>
<tr>
<td>$S_r, S_{c1}$</td>
<td>0.38 (±0.09)</td>
<td>3.90 (±0.32)</td>
<td>1.97 (±0.11)</td>
<td>0.29 (±0.04)</td>
</tr>
<tr>
<td>$S_r, S_{c2}$</td>
<td>0.41 (±0.06)</td>
<td>4.23 (±0.3)</td>
<td>2.12 (±0.12)</td>
<td>0.44 (±0.07)</td>
</tr>
<tr>
<td>$S_r, S_i$</td>
<td>0.31 (±0.08)</td>
<td>4.37 (±0.35)</td>
<td>2.15 (±0.12)</td>
<td>0.4 (±0.08)</td>
</tr>
<tr>
<td>$S_r, S_{c1c2}$</td>
<td>0.38 (±0.1)</td>
<td>3.83 (±0.29)</td>
<td>1.94 (±0.09)</td>
<td>0.27 (±0.03)</td>
</tr>
<tr>
<td>$S_r, S_{p1}$</td>
<td>0.45 (±0.08)</td>
<td>3.82 (±0.3)</td>
<td>1.97 (±0.12)</td>
<td>0.36 (±0.03)</td>
</tr>
<tr>
<td>$S_r, S_d$</td>
<td>0.51 (±0.11)</td>
<td>4.02 (±0.33)</td>
<td>2.1 (±0.12)</td>
<td>0.46 (±0.09)</td>
</tr>
</tbody>
</table>

Table 4: Matching performance when combining $S_r$ with each of the other 9 features. The table represents the mean over 10 folds with the standard errors in brackets. All results are expressed as percentages. The rows corresponding to the combinations with lowest TE are highlighted.

<table>
<thead>
<tr>
<th>Feature Pairs</th>
<th>FMR</th>
<th>FNMR</th>
<th>FMR ($S_r$)</th>
<th>FNMR ($S_r$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$S_r, S_n$</td>
<td>Left 0.37</td>
<td>4.72</td>
<td>0.37</td>
<td>4.28</td>
</tr>
<tr>
<td>$S_r, S_p$</td>
<td>Right 0.38</td>
<td>4.29</td>
<td>0.38</td>
<td>4.06</td>
</tr>
<tr>
<td>$S_r, S_{c1}$</td>
<td>Left 0.38</td>
<td>3.90</td>
<td>0.38</td>
<td>4.22</td>
</tr>
<tr>
<td>$S_r, S_{c2}$</td>
<td>Right 0.29</td>
<td>3.93</td>
<td>0.29</td>
<td>4.46</td>
</tr>
<tr>
<td>$S_r, S_i$</td>
<td>Left 0.41</td>
<td>4.23</td>
<td>0.41</td>
<td>4.16</td>
</tr>
<tr>
<td>$S_r, S_{c1c2}$</td>
<td>Right 0.44</td>
<td>4.26</td>
<td>0.44</td>
<td>4.03</td>
</tr>
<tr>
<td>$S_r, S_{p1}$</td>
<td>Left 0.31</td>
<td>4.37</td>
<td>0.31</td>
<td>4.39</td>
</tr>
<tr>
<td>$S_r, S_d$</td>
<td>Right 0.38</td>
<td>3.83</td>
<td>0.38</td>
<td>4.22</td>
</tr>
</tbody>
</table>

We remark that as the graph edit algorithm we used gave significant results we did not need to try other cost functions. However, for future work these could be applied to help evaluate the efficiency trade-off of graph matching.

session and across different sessions, to estimate the difference in performance when biometrics are captured in different sessions. Note that multi session comparison is typical in a real-world scenario. The proposed method beats the existing state of art results on the PUT public palm vein database (Kabacinski and Kowalski, 2011) for genuine samples captured across sessions. This establishes BGM as an effective algorithm when genuine palm vein samples have partial overlap.
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REFERENCES


