

## HMMs: Biometric Signature Verification

In this exercise, we aim to (partially) reproduce a work on biometric signature verification. During training, the task is to fit a Hidden Markov Model to handwritten signatures of a person. During testing, the task is to determine for a new signature whether it belongs to that person or to someone else who imitates that signature.

This exercise loosely builds on a paper by Fierrez *et al.*, “HMM-based on-line signature verification: Feature extraction and signature modeling”, Pattern Recognition Letters, vol. 28, Aug. 2007, pp. 2325–2334. The paper is also available in studOn.

### Exercise 1 Using hmmlearn

You will likely need to locally install `hmmlearn`. To do so, please run

```
pip3 install --user setuptools  
pip3 install --upgrade --user hmmlearn
```

for integration into python 3. Note that the package is about 190MB large.

### Exercise 2 Dataset

We will use some writer data from the “mobisig” dataset. The data is uploaded StudOn in the *Exercise 8* folder as a zip file. Please download and unpack the data before using it. Let us use signatures from up to 10 writers, denoted as `user_0` to `user_9`. Each signature is stored in an individual CSV file. “Original” signatures are from the writer herself or himself. “Imitated” signatures are from an imposter who only pretends to be that writer, i.e., they should not be considered for training.

### Exercise 3 Feature Extraction and Processing

The purpose of this task is to prepare the signature data for classification in a HMM. Each signature file provides several feature information: the time stamp of a stroke,  $x$ - and  $y$ -position, pen pressure  $p$ , velocity in  $x$ - and  $y$ -direction, and acceleration in  $x$ - and  $y$ -direction, and other information that we will not consider.

The task is to use the available features to compute the seven-dimensional feature vector from Eqn. (3) of the paper. To this end, you will notice that  $x$ ,  $y$ ,  $p$  are identical to the features in the paper. The remaining four features are relatively straightforward to calculate with the equations in the bullet list above Eqn. (3).

There is one notable caveat for the feature computation: if during your calculation a division by 0, or a log of 0 occurs, the result in that dimension will be `nan` instead of a numeric value — this can really screw up your result. One standard approach in pattern recognition to deal with such issues is to add a small epsilon to “dangerous” calculations.

The final feature vector should be mean-free and with unit variance (confer Sec. 2.5 in the paper).

We found in our own experiments that you can omit the feature rotation (Eqn. (2)) and a resampling of the timestamps to equidistance. These two steps would only be required if our goal would be to achieve a performance that is competitive to what the authors reported. Don't worry about this for now, let us just aim to be better than guessing.

## Exercise 4 HMM Preliminary Remarks

Have a look at the `hmmlearn` API documentation and tutorial.

Considering the HMM that we discussed in the lecture, and considering our feature set, you will notice a difference: in the lecture, we assumed that an observation  $o_i$  and the associated output probability  $b_{s_j}(o_i)$  of state  $s_j$  is a categorical value. In our dataset, we do not have such clearly defined categorical values. To accomodate this, `hmmlearn` offers that each state internally holds a parametric probability density estimate of “its” features. In other words, a state fits during training a Gaussian density or a Gaussian Mixture model to a subsequence of the features. During testing, it evaluates the probability that a sample at time step  $t$  is drawn from that distribution.

Therefore, you should use `hmm.GMMHMM` from the `hmmlearn` library to achieve this internal representation. We made good experiences with a commonly used constraint, namely to set the covariance matrices to be diagonal matrices with the (preset) parameter `covariance.type='diag'`.

We also made good experiences with constraining the model to a left-right HMM (this is also what the paper does). This can be done by pre-setting the transition matrix to a left-right model, as shown in the `hmmlearn` tutorial. Note that if you do this, then you have to prevent that the transition matrix is freshly re-initialized by the library, i.e., you have to omit `'t'` from the parameter `init_params`.

The same applies to the vector of starting probabilities: the paper sets these to  $(1, 0, \dots, 0)$ . Do this analogously (again, please refer to the tutorial), and drop its initialization from `init_params`.

When experimenting with the `hmm.GMMHMM` model, we found that it helps convergence if transition probabilities and starting probabilities are completely excluded from training, i.e., also not refined later. To achieve this, drop `t` and `s` from the not-so-excellently-named parameter `params`.

(as a sidenote, you probably see from these caveats that training can be a bit brittle with increasing model complexity. This is a general rule in statistical model fitting — which is why so many Ph.D. students are spending so much time on deep networks, our most complex models to date.)

## Exercise 5 HMM Results per Signature

The task is to fit a `GMMHMM` to the data. To do this, split the 45 original signatures from a writer directory into training and testing. The training should contain at least 25 signatures to prevent overfitting. Start with a modest number of states, e.g., 3 or 4, to get some first results.

The paper proposes to statically segment the input sequence into  $H$  segments (Eqn. (9)), where  $H$  is the number of states in the model. Additionally, they pre-initialize their `GMM` model with the  $k$ -means algorithm. Ignore this! You can

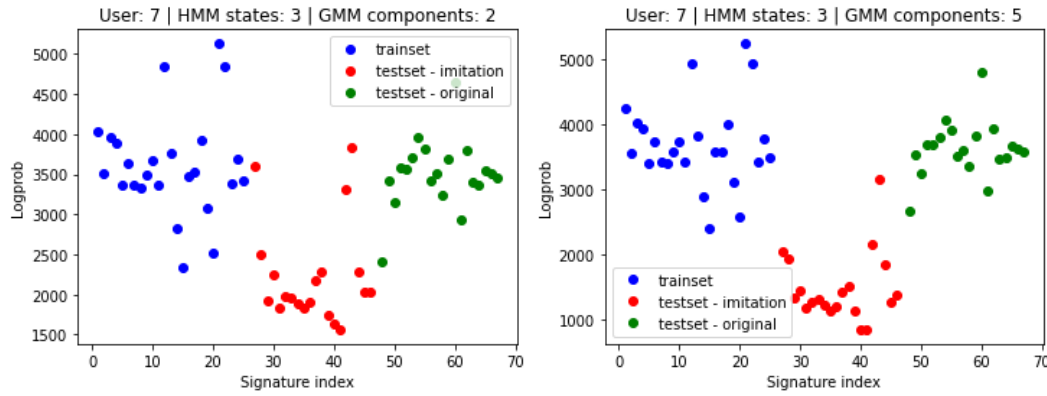


Figure 1: Log-probabilities for the trainset (blue), testset using the original signatures (green) and testset using the imitated signatures (red).

achieve some reasonable results by just feeding whole signatures to the training process. To this end, consult the tutorial: you will need a 2-D list where one row contains the seven features per time step. All time steps and multiple feature sequences are just concatenated.

Also ignore the threshold in Eqn. (11). Instead, compute the log-probabilities individually for each signature in one user directory using the method `score` from `hmmlearn`. Create a plot with the matching score per training signature in blue, each original testing signature in green, and each imitated signature in red. It should look like one of our example results in Fig. 1, two models were trained on 25 training samples. For the testing, 40 signatures were used, 20 original and 20 imitated.

“Success” is defined as having a set of imitated signatures with lower scores than the original (green or blue) signatures. Thus, if we had a good threshold, we could reject the imitations! For example, in Fig. 1, the model using 5 components for the GMM (right plot) would allow us to separate better the imitated signatures than the one with 2 components (left plot).

**Exercise 6** Our goal is to obtain a list of results that are a simplified version of Tab. 1 in the paper. As a simplification, we will not report error rates. Instead, calculate the average matching scores on the test set for original and imitated signatures. If the average score for the originals is larger than for imitated, we count it as success. Which combinations of  $H$  and  $M$  work better, which work worse for your implementation?

Please post a figure of one result with a  $H$  and  $M$  of your choice to the forum and add a short text why you chose it, and how it performed, compared to other combinations of  $H$  and  $M$ .

*Comments:*

*We ask for only one figure per group. Please also state your group number. Bring your code to the joint meeting on July 5 or July 6 for a potential little extra experiment.*