Gaussian process classifiers for convolutional neural network uncertainty

Sebastian Borgeaud dit Avocat

spb61@cam.ac.uk

LE49 - Probabilistic Machine Learning Project 28 February 2018

Abstract

In this report, I show how a Gaussian process can be used in conjunction with a convolutional neural network to obtain prediction uncertainty bounds without decreasing the performance of the model. Furthermore, I inspect how these uncertainty bounds increase when presented with input data coming from a different underlying distribution than the training data. Finally, I show how the uncertainty bounds can be incorporated into the model to give the model the option to reject classifying examples for which it is too uncertain and evaluate this model using a metric that weighs misclassification and classification rejection according to user-defined weights.

1 Introduction

Model uncertainty is of crucial importance in many regression and classification tasks such as medical diagnosis, autonomous vehicle steering or high frequency trading [1]. With model uncertainty it is possible to treat uncertain inputs and special cases explicitly. For example, if model uncertainty is high, a human expert could be asked to decide whether a particular MRI scan shows signs of a tumour or not. Another example showing how model uncertainty could be utilised is when the model is allowed to reject classifying certain examples. In this case the model could reject classification for those examples for which it is too uncertain.

Although deep learning models have become the state-of-the-art in many of those tasks, they do not capture model uncertainty. In classification tasks, the last layer is often passed through a softmax activation which returns a probability distribution over the classes [2]. However, a high probability doesn't mean that the model is certain of its prediction: the model could give a high probability with a high uncertainty in which case the prediction might be worthless [3].

Finding ways to integrate Bayesian uncertainty with deep learning models has become a hot research topic recently. For example, Yarin Gal shows how dropout in neural networks can be used to obtain uncertainty bounds [3]. In this project, I focus on one method to obtain uncertainty bounds for convolutional neural networks: using a Gaussian process on the last-layer features learned by the convolutional neural network. In particular, I focus on the task of classifying hand-written digits using the MNIST dataset [4].

2 Background

2.1 Convolutional neural networks

Convolutional neural networks (CNNs) have become standard in many deep learning applications, especially in image processing or vision tasks [2]. A convolutional neural network is a type of

feedforward neural network, typically consisting of convolutional layers, pooling layers and fully connected layers [5]:

• Convolutional layers are composed of several convolution kernels each computing a different feature map. The output feature maps are obtained by convolving the input with the convolution kernel and then applying an element-wise nonlinearity. Mathematically, the feature value $z_{i,j,k}^l$ at location (i,j) of the k^{th} feature map in the l^{th} layer is computed as:

$$z_{i,j,k}^l = \mathbf{w}_k^{l} \mathbf{x}_{i,j}^l + b_k^l$$

where \mathbf{w}_k^l and b_k^l are the weight and bias vectors for the k^{th} convolution kernel in the l^{th} layer and $\mathbf{x}_{i,j}^l$ is the input patch centered around (i,j) in the l^{th} layer. The ouput value is computed by apply a nonlinearity $a(\cdot)$ point-wise:

$$x_{i,j,k}^{(l+1)} = a(z_{i,j,k}^l)$$

Pooling layers aim to achieve shift-invariance and reduce the number of parameters in the
network by reducing the resolution of the feature maps. The pooling layer operates on each
feature map independently. Mathematically, the output of a pooling layer with pooling
operation pool(·) is given by

$$y_{i,j,k}^l = \operatorname{pool}(x_{m,n,k}^l), \forall (m,n) \in \mathcal{R}_{i,j}$$

where $\mathcal{R}_{i,j}$ is a local neighbourhood around (i,j). Typically, the pooling operation computes the average or the maximum.

• Fully connected layers connect every neuron in the previous layer to every neuron in the current layer. Mathematically, the output of a fully connected layer is given by:

$$x_i^{(l+1)} = a\Big(\big(\sum_j w_{i,j}^l x_j^l\big) + b_i^l\big)$$

where $a(\cdot)$ is a nonlinearity, $w_{i,j}^l$ is the weight connecting neuron j in the l^{th} layer to neuron i in layer $(l+1)^{\text{th}}$, and b_i^l is the bias weight for neuron i.

Optimising a convolutional neural network is done in the same way as optimising standard neural networks. A differentiable loss function is computed for the training examples (often done in batches) and the gradients w.r.t. the weights of the network are computed. Using these gradients, the weights are updated in a gradient descent step. Typically, more complex update rule that take into account change momentum (e.g. Adam optimisation[6]) are used as they converge faster to a local minimum.

2.2 Gaussian processes

Formally, a Gaussian Process is defined as a collection of random variables, any finite number of which have (consistent) joint Gaussian distributions. A Gaussian process therefore defines a distribution over functions and is fully specified by a mean function m(x) and a covariance function k(x,x'). Write $f \sim \mathcal{GP}(m,k)$ meaning f is distributed as a GP with mean m and covariance k.

Using the GP we can draw samples from the function for any finite number n of locations. Given locations $\mathbf{x} = [x_1, \dots, x_n]$, first compute $\mu_i = m(x_i)$, $\Sigma_{i,j} = k(x_i, x_j)$. We can then sample a vector from this distribution: $\mathbf{f} \sim \mathcal{N}(\mu, \mathbf{\Sigma})$.

2.2.1 Regression

We can now use this GP as a prior for Bayesian inference. Let f be the known function values for the training examples an let f_* be the set of function values corresponding to the set of test inputs X_* . The joint distribution is given by

$$\begin{bmatrix} \mathbf{f} \\ \mathbf{f}_* \end{bmatrix} = \mathcal{N} \Big(\begin{bmatrix} \boldsymbol{\mu} \\ \boldsymbol{\mu}_* \end{bmatrix}, \begin{bmatrix} \boldsymbol{\Sigma} & \boldsymbol{\Sigma}_* \\ \boldsymbol{\Sigma}_*^T & \boldsymbol{\Sigma}_{**} \end{bmatrix} \Big)$$

where μ_* are the test means, Σ_* are the training-test covariances, and Σ_{**} are the test-test covariances. Since we know the training values f, we are interested in the conditional distribution of f_* given f:

$$\mathbf{f} \big| \mathbf{f}_* \sim \mathcal{N} \big(\mu_* + \boldsymbol{\Sigma}_*^T \boldsymbol{\Sigma}^{-1} (\mathbf{f} - \boldsymbol{\mu}), \boldsymbol{\Sigma}_{**} - \boldsymbol{\Sigma}_*^T \boldsymbol{\Sigma}^{-1} \boldsymbol{\Sigma}_* \big)$$

This corresponds to a posterior Gaussian process, given by

$$f | \mathcal{D} \sim \mathcal{GP}(m_{\mathcal{D}}, k_{\mathcal{D}}),$$

$$m_{\mathcal{D}}(x) = m(x) + \Sigma(X, x)^T \Sigma^{-1}(\mathbf{f} - \mathbf{m})$$

$$k_{\mathcal{D}}(x, x') = k(x, x') - \Sigma(X, x)^T \Sigma^{-1} \Sigma(X, x')$$

where $\Sigma(X,x)$ is a vector of covariances between every training case in X and x. Furthermore, it is easy to incorporate noise in the observations. Assuming i.i.d. additive Gaussian noise, every f(x) now has extra covariance with itself with a magnitude equal to the noise variance σ_n^2 :

$$f|\mathcal{D} \sim \mathcal{GP}(m_{\mathcal{D}}, k_{\mathcal{D}} + \delta_{ii}\sigma_n^2)$$

where $\delta_{ii'} = 1$ iff i = i' is the Kronecker's delta.

The mean function m(x) and the covariance function k(x,x') are typically parametrised in terms of hyper-parameters θ . During training we find the values of the hyper-parameters which optimise the marginal likelihood:

$$L = \log p(\boldsymbol{y}|\boldsymbol{x}, \boldsymbol{\theta}) = -\frac{1}{2}|\boldsymbol{\Sigma}| - \frac{1}{2}(\boldsymbol{y} - \boldsymbol{\mu})^T \boldsymbol{\Sigma}^{-1}(\boldsymbol{y} - \boldsymbol{\mu}) - \frac{n}{2}\log(2\pi)$$

This optimisation can be done using standard gradient methods.

2.2.2 Classification

Binary classification using Gaussian processes can be done by setting a GP prior over a latent function $f(\mathbf{x})$ and then using squashing function such as the sigmoid to obtain a probability:

$$\pi(\mathbf{x}) = p(y = +1|\mathbf{x}) = \sigma(f(\mathbf{x}))$$

Inference is done in two steps. First, the distribution of the latent variable corresponding to a new test input \mathbf{x}_* is computed:

$$p(f_*|\mathbf{X}, \mathbf{y}, \mathbf{x}_*) = \int p(f_*|\mathbf{X}, \mathbf{x}_*, \mathbf{f}) p(\mathbf{f}|\mathbf{X}, \mathbf{y}) d\mathbf{f}.$$

Second, a probabilistic prediction is computed using the distribution computed in the first step:

$$\bar{\pi}_* = p(y_* = +1|\mathbf{X}, \mathbf{y}, \mathbf{x}_*) = \int \sigma(f_*) p(f_*|\mathbf{X}, \mathbf{y}, \mathbf{x}_*) df_*$$

As the likelihood is non longer Gaussian, the first integral becomes analytically intractable. Similarly, depending on the sigmoid function, the second integral can also be intractable. Hence, we need to use approximations, either analytical or numerical, for example using Monte Carlo sampling, to solve the integrals.

Multi-class classification is typically [7] approached by assuming the following labelling rule for y_* given \mathbf{x}_* :

$$y_* = \operatorname*{arg\,max}_{k=1} f^k(\mathbf{x}_*)$$

where each $f^k(\cdot)$ is a nonlinear latent function with a GP prior and C is the number of classes. The likelihood is again non-Gaussian meaning that approximation techniques have to be used to perform inference and to optimise the hyper-parameters.

2.3 Uncertainty in Deep Learning

Current deep learning methods only give point estimates of parameters and predictions [2]. Despite the fact the output of a softmax layer in a classification task can be interpreted as a probability distribution $p(y_i = k) = f(x_i)_k$, the probabilities say nothing about how certain the model is: The model could assign a high probability to a class but can still be highly uncertain about it [3].

One way to obtain uncertainty bounds from deep learning models was discovered by Yarin Gal. In fact, Gal [1] showed that dropout training in neural networks can be casted as approximate Bayesian inference in deep Gaussian processes. In particular, he showed that using Dropout not only at training

time but also at test time gives a principled way of computing uncertainty bounds by sampling multiple estimates from the model and computing their sample variance [1].

In this project, I focus on a different way to obtain those uncertainty bounds: Using a Gaussian process trained on the features extracted by a CNN that was trained in a previous step on the same test training data. It should be noted that this is a less theoretically grounded approach than the one taking by Gal, but nonetheless provides a practical way of getting uncertainty bounds without losing the power of deep learning models.

3 Method

3.1 Convolutional Neural Network architecture

The first step consists of training the convolutional neural network, which can be done using one of the many deep learning libraries. For example, I use Keras [8] which defines an extra abstraction layer above TensorFlow. The network architecture is provided in the Keras tutorial for image classification on MNIST. The first two layers are convolutional layers with 3×3 kernels and ReLU activations, where ReLU(x) = $\max(x,0)$. These layers have respectively 32 and 64 feature maps. A maxpooling layer with kernel size 2×2 is then applied to the output of the convolutional layer. The final 2 layers are fully connected layers consisting of 128 and 10 neurons respectively. The first fully connected layer has a ReLU activation. The last fully connected layer uses a softmax activation, which outputs a probability distribution over the 10 classes representing the 10 digits. Furthermore, Dropout [9] is applied after the max-pooling layer with p=0.25 and after the first fully connected layer with p=0.5, where p is the probability of dropping a neuron. The network is trained using an Adadelta optimiser over 10 epochs with batches of size 128.

3.2 Gaussian process

From the trained CNN model, the activations of the last hidden layer are extracted. These correspond to a vector of size 128 for each input image, and can be thought of as the features extracted by the CNN for classification. Using these features I then train a Gaussian Process. This has the advantage of being a lower dimensional classification task compared to classifying the entire input image directly, whilst still being fully automatic, i.e. there are no user-defined features. However, the MNIST dataset, with 60,000 training images, is considered a large dataset for GPs because inference takes $\mathcal{O}(n^3)$ time where n is the number of training instances [10]. To make the training possible, I use GPFlow [11], which implements various approximation algorithms for Gaussian Processes. Furthermore, the library is built on top of TensorFlow which has the further advantage of being usable on a GPU out-of-the-box, which provides a further speed-up. More precisely, I use the Sparse Variational Gaussian Process Classifier presented by Hensman et al. [10]. The model learns a set of m inducing points which are used instead of the training points in the Gaussian process. Typically, m < n, which makes the task tractable as its complexity is $\mathcal{O}(nm^2)$. For the experiments done in this project I chose to use 600 inducing points, which allowed the model to be trained in about 15 minutes on a NVidia Titan Xp GPU.

I train the Gaussian process with a simple kernel as it is sufficient to obtain good accuracy. The kernel consists of a Matern kernel [12] with $\nu=\frac{3}{2}$ and a white noise function to account for the noise in the input data.

3.3 Classifying with a reject option

One way to incorporate the model uncertainty is by allowing the model to say "I don't know". For example, if the model was presented with an image of the letter 'a', it would still classify the image into one of the 10 digits, which might not be a desired outcome. Instead, the model should be able to reject classification for the inputs in which the uncertainty is too high. This is known as **classification** with a reject option [13].

Given uncertainty bounds with each class probability, we can design the model to reject classification according to certain rejection rules. For example, we could decided to reject if the standard deviation of the highest probability is above a pre-defined threshold. Another rejection rule could be to reject if the second highest class probability lies within a certain factor ϵ of the standard deviations of the

highest probability, i.e. the model would reject if

$$|f_p(\mathbf{x}^{(i)})_1 - f_p(\mathbf{x}^{(i)})_2| < \epsilon f_{\text{std}}(\mathbf{x}^{(i)})_1$$

where $f_p(\mathbf{x}^{(i)})$ is the classification probability vector for input $\mathbf{x}^{(i)}$ with $f_p(\mathbf{x}^{(i)})_j$ being defined as the probability of the j^{th} most likely class, and $f_{\text{std}}(\mathbf{x}^{(i)})_1$ is the standard deviation of the most likely class returned by the model for input $\mathbf{x}^{(i)}$. Note that the ϵ parameter defines a trade-off between the number of misclassified examples at the number of examples that would have been correctly classified but were rejected instead.

3.4 Evaluation

3.4.1 Accuracy

The standard metric for evaluating a model f on a classification task is the accuracy:

$$accuracy = \frac{1}{N} \sum_{i=1}^{N} \mathbb{1}(f(\mathbf{x}^{(i)}) = y^{(i)})$$

where N is the number of test data points and $(\mathbf{x}^{(i)}, y^{(i)})$ is the i^{th} test point. Note that the accuracy does not take into account the uncertainty with which a model makes the prediction and only rewards correctly classified examples.

3.4.2 R-accuracy

An appropriate metric for classification with a reject option assigns a cost α to misclassification and a cost β to the reject option \mathbb{R} :

$$\text{R-accuracy}_{\alpha,\beta} = \frac{1}{N} \sum_{i=1}^{N} \mathbb{1}(f(\mathbf{x}^{(i)}) = y^{(i)}) - \alpha \mathbb{1}(f(\mathbf{x}^{(i)}) \neq y^{(i)}) - \beta \mathbb{1}(f(\mathbf{x}^{(i)}) = \mathbb{R})$$

Here, α and β are parameters that are set by the user. A large value for α corresponds to cases where misclassification is expensive, for example in medical diagnosis or when a self-driving car has to make a steering decision. The β parameter intuitively indicates how much value the user places on obtaining a prediction, as a higher β will increase the cost of the reject option.

4 Results

4.1 MNIST dataset

The MNIST dataset is considered a toy dataset in deep learning. In fact, using the model described above, I obtain an accuracy of 99.12% on the held-out test images. This means that only 88 of the 10,000 test images are misclassified.

Using the Gaussian process with the CNN features, the accuracy of the model on the test images improves to 99.28%, reducing the number of misclassified images to 72. Although this increase in performance might seem surprising at first, it can be explained by the fact that the Gaussian process is more complex and more powerful than the last layer of the CNN, which just computes a linear combination of the features and then applies the softmax activation. The results obtained on the MNIST dataset are summarised in the table 1.

Model	Accuracy	Misclassified images
CNN	99.12%	88
GP	99.28%	72

Table 1: MNIST test images accuracy for the CNN and the GP.

Furthermore, it is interesting to inspect how uncertain the Gaussian process is for the different test images. In order to do this, I plot in figure 1 the cumulative distribution of the standard deviations

for the most likely class for both correctly and incorrectly classified test images. For the correctly classified examples, the model's uncertainty is low: the model gives the correct class probability with a standard deviation of less than 0.1 for 97.3% of the correctly classified images. However, for the incorrectly classified images, the standard deviation of the most likely class seems to be higher: The model predicts a standard deviation below 0.1 for only 19.4% of the incorrectly classified images. This suggests that the model is generally less certain of its prediction in the cases where it predicted the wrong class.

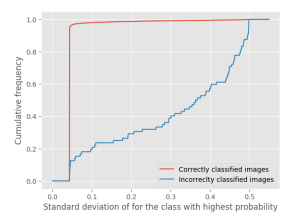


Figure 1: Cumulative distribution of the standard deviations for the most likely class for both correctly and incorrectly classified test images.

4.2 Noisy MNIST (n-MNIST)

Next, I evaluate both the CNN and the Gaussian process on the n-MNIST dataset [14]. The dataset consists of the MNIST images, but modified in 3 different ways so as to make the classification task harder. The first set of images is created by adding white Gaussian noise with a signal-to-noise ratio of 9.5 to the images. The second set of images is obtained by emulating a linear motion blur. The last set of images is the hardest to classify as it is created by reducing the contrast and also adding white Gaussian noise to the original images. Figure 2 shows an example image taken from the 3 datasets.

The task of classifying examples whose underlying distribution is different from the underlying distribution of the training images is called **out-of-distribution** classification [1]. The performance of the two models is shown in table 2. Although the GP performs better on the NMIST test images, the CNN performs better on all 3 noisy datasets. This could again be explained by the fact that the GP is a more complex model, therefore is overfitting more on the NMIST data and hence is not able to generalise as well on these different datasets. Especially noticeable is the difference in performance for the reduced contrast dataset, where the CNN obtains an accuracy of 77.71% compared to 70.13% for the GP. As the 3rd dataset is the least similar the original one, this higher discrepancy supports the claim that the GP is overfitting more on the MNIST dataset.

We can again inspect how uncertain the model is when classifying the noisy dataset by plotting the cumulative distributions of the standard deviations of the most likely digit. Figure 3 shows the resulting plots for each dataset, which indeed suggest that the uncertainty increases as the model is asked to classify images that are increasingly different from those it was trained on. The uncertainty



Figure 2: Example digits from the noisy-MNIST dataset. From left to right: Additive white Gaussian noise, Motion blur, and Reduced contrast with AWGN

Model	Accuracy		
	AWGN	Motion blur	Reduced contrast
CNN	94.35%	93.27%	77.71%
GP	93.34%	92.09%	70.13%

Table 2: MNIST test images accuracy for the CNN and the GP.

for the additive white Gaussian noise dataset is higher than the uncertainty for the MNIST test dataset. This is shown by the fact that the cumulative distribution for the correctly classified images in the AWGN images is lower. Similarly, the cumulative distribution further decreases for the motion blur and reduced contrast images, with a lower classification accuracy corresponding to a higher uncertainty.

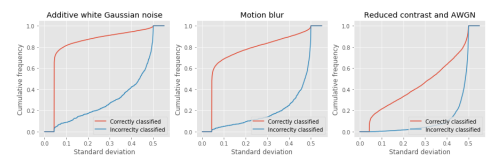


Figure 3: Cumulative distribution of standard deviations for correctly and incorrectly classified test images on the 3 n-MNIST datasets.

4.3 Classifying with a reject option

Finally, I also evaluate the performance of the GP augmented with a reject option according to the rejection rule described in 3.3 on the MNIST test dataset. Figure 4 (left) shows the R-accuracy with varying α and fixed $\beta=0$ for different values of ϵ . In the cases where $\alpha>1$, that is when the cost of misclassification is higher than the reward obtained by a correct classification, there seem to be an optimal value of ϵ lying between 20 and 40.

Furthermore, there is indeed a trade-off between the number of misclassified but accepted images and the number of correctly classified but rejected images. Figure 4 (right) illustrates this trade-off.

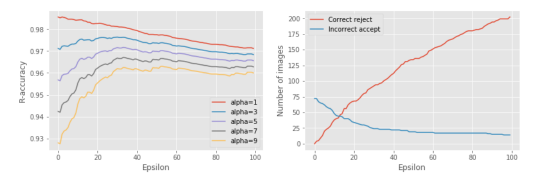


Figure 4: Left: R-accuracy obtained using varying ϵ and α but setting $\beta=0$. For $\alpha>0$ there seems to be an optimal value of ϵ around 30. Right: Number of examples misclassified or rejected. This illustrates the trade-off between the number of misclassified but accepted and correctly classified but rejected examples.

5 Discussion

Only 72 of the 10,000 MNIST test images are misclassified by the GP. To gain some insights into why only those were misclassified, the 72 misclassified input images are shown in figure 8 in appendix A. The images reveal that many of those 72 images are indeed very ambiguous, even for a human annotator. For example, it is hard to say whether the first image should be a '1' or a '2'. This also explains why the uncertainty given by the model is higher for the misclassified images than it is for the correctly classified images. In fact, one could argue that for those ambiguous images the model should indeed be less certain of its prediction.

5.1 Uncertainty with additive white noise

A further property the model uncertainty should obey is that as an image becomes more and more distorted, the uncertainty should increase. To verify that this holds I add increasingly more white Gaussian noise to a test digit and then ask the GP to make predictions for those images. The results are shown in figure 5 for an image of the digit '6'. Note that for the last few images, it becomes

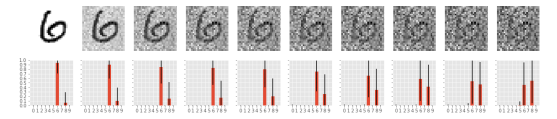


Figure 5: Top: Test images with increasingly more additive white Gaussian noise. Bottom: Gaussian process predictions for the test images. The error bars show the standard error for each class.

almost impossible to discern the original digit from the noise (especially when looked at individually, rather than next to the original image). Indeed, the model wrongly classifies the last image as a '5', although it does so with a high uncertainty. Figure 6 shows the probability and uncertainty assigned to the digit '6' for the 10 images. As hoped for, the uncertainty increases as more noise is added to the image. Especially noteworthy is the increase of the standard deviation relative to the class probability.

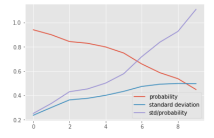
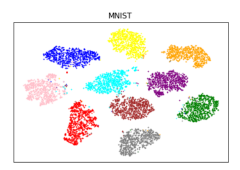


Figure 6: Probability and standard deviation assigned to class '6' for the 10 images shown in figure 5. Also plotted is the standard deviation relative to the class proability.

5.2 CNN features visualisation

Finally, I show the features learned by the CNN by reducing the number of dimensions of the features to 2 using the t-SNE [15] dimensionality reduction technique. The learned features are shown for the MNIST test dataset and for the additive white Gaussian noise images of the n-MNIST dataset.



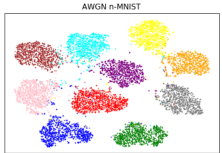


Figure 7: Learned features for the MNIST test dataset and the AWGN n-MNIST test dataset. The dimensionality of the features were reduced using t-SNE.

References

- [1] Yarin Gal. Uncertainty in Deep Learning. PhD thesis, University of Cambridge, 2016.
- [2] Ian Goodfellow, Yoshua Bengio, and Aaron Courville. *Deep Learning*. MIT Press, 2016. http://www.deeplearningbook.org.
- [3] Yarin Gal and Zoubin Ghahramani. Dropout as a bayesian approximation: Representing model uncertainty in deep learning. In *international conference on machine learning*, pages 1050–1059, 2016.
- [4] Yann LeCun, Léon Bottou, Yoshua Bengio, and Patrick Haffner. Gradient-based learning applied to document recognition. *Proceedings of the IEEE*, 86(11):2278–2324, 1998.
- [5] Jiuxiang Gu, Zhenhua Wang, Jason Kuen, Lianyang Ma, Amir Shahroudy, Bing Shuai, Ting Liu, Xingxing Wang, and Gang Wang. Recent advances in convolutional neural networks. CoRR, abs/1512.07108, 2015.
- [6] Diederik P Kingma and Jimmy Ba. Adam: A method for stochastic optimization. arXiv preprint arXiv:1412.6980, 2014.
- [7] Carlos Villacampa-Calvo and Daniel Hernández-Lobato. Scalable multi-class gaussian process classification using expectation propagation. *arXiv* preprint arXiv:1706.07258, 2017.
- [8] François Chollet et al. Keras. https://github.com/keras-team/keras, 2015.
- [9] Nitish Srivastava, Geoffrey Hinton, Alex Krizhevsky, Ilya Sutskever, and Ruslan Salakhutdinov. Dropout: A simple way to prevent neural networks from overfitting. *The Journal of Machine Learning Research*, 15(1):1929–1958, 2014.
- [10] James Hensman, Alexander G de G Matthews, and Zoubin Ghahramani. Scalable variational gaussian process classification. 2015.
- [11] Alexander G. de G. Matthews, Mark van der Wilk, Tom Nickson, Keisuke. Fujii, Alexis Boukouvalas, Pablo Le'on-Villagr'a, Zoubin Ghahramani, and James Hensman.
- [12] Carl Edward Rasmussen and Christopher KI Williams. Gaussian processes for machine learning. 2006. The MIT Press, Cambridge, MA, USA, 38:715–719, 2006.
- [13] C Chow. On optimum recognition error and reject tradeoff. *IEEE Transactions on information theory*, 16(1):41–46, 1970.
- [14] Saikat Basu, Manohar Karki, Sangram Ganguly, Robert DiBiano, Supratik Mukhopadhyay, and Ramakrishna R. Nemani. Learning sparse feature representations using probabilistic quadtrees and deep belief nets. CoRR, abs/1509.03413, 2015.
- [15] Laurens van der Maaten and Geoffrey Hinton. Visualizing data using t-sne. Journal of machine learning research, 9(Nov):2579–2605, 2008.

A Misclassified MNIST images by the Gaussian process model



Figure 8: Test images misclassified images by the GP model.