# **Uncertainty Estimation Practical Part I**

There are many ways to estimate predictive uncertainty in deep learning. However, in this practical, we focus on simple yet effective methods. One such method that has achieved impressive performance is the Deep Ensembles method[1]. In this practical, we first study this method in Part I, then a recent extension in Part II.

## Simple and Scalable Predictive Uncertainty Estimation Using Deep Ensembles --Lakshminarayanan B, Pritzel A, Blundell C. [1]

## **Abstract:**

Deep neural networks (NNs) are powerful black box predictors that have recently achieved impressive performance on a wide spectrum of tasks. Quantifying predictive uncertainty in NNs is a challenging and yet unsolved problem. Bayesian NNs, which learn a distribution over weights, are currently the state-of-the-art for estimating predictive uncertainty; however these require significant modifications to the training procedure and are computationally expensive compared to standard (non-Bayesian) NNs. We propose an alternative to Bayesian NNs that is simple to implement, readily parallelizable, requires very little hyperparameter tuning, and yields high quality predictive uncertainty estimates. Through a series of experiments on classification and regression benchmarks, we demonstrate that our method produces well-calibrated uncertainty estimates which are as good or better than approximate Bayesian NNs. To assess robustness to dataset shift, we evaluate the predictive uncertainty on test examples from known and unknown distributions, and show that our method is able to express higher uncertainty on out-of-distribution examples. We demonstrate the scalability of our method by evaluating predictive uncertainty estimates on ImageNet.

## **Notations**

Assume we have i.i.d. training data points (x, y), where  $x \in \mathbb{R}^D$  is the D-dimensional input features. And for a classification problem with K classes,  $y \in \{1, \dots, K\}$ ; for a regression problem,  $y \in \mathbb{R}$ . We use a NN with parameters  $\theta$  to model the probablistic predictive distribution  $p_{\theta}(y|x)$  over the labels.

## **Proper scoring rules**

Scoring rules measure the quality of predictive uncertainty [2]. Suppose we have  $(y, \boldsymbol{x})$  following the true distribution  $q(y, \boldsymbol{x})$ , and we want to evaluate the quality of the predictive distribution,  $p_{\theta}(y|\boldsymbol{x})$ . The scoring rule, a function,  $S(p_{\theta}, (y, \boldsymbol{x}))$ , assigns a numerical score to the predictive distribution  $p_{\theta}(y|\boldsymbol{x})$ . Here we consider scoring rules where a high score means better quality. The expected scoring rule is then

 $S_{\mathbb{E}}(p_{\theta},q)=\int q(y,m{x})S(p_{\theta},(y,m{x}))dydm{x}$ . A proper scoring rule is, for all  $p_{\theta}$  and q,  $S_{\mathbb{E}}(p_{\theta},q)\leq S_{\mathbb{E}}(q,q)$  with equality if and only if  $p_{\theta}(y|m{x})=q(y|m{x})$ . Since a larger value  $S_{\mathbb{E}}(p_{\theta},q)$  means a better quality of the predictive uncertainty, we could train NNs by minimizing the loss  $\mathcal{L}(\theta)=-S_{\mathbb{E}}(p_{\theta},q)$ .

Log-likelihood,  $\log p_{\theta}(y|\boldsymbol{x})$ , turns out to be a proper scoring rule due to [Gibbs inequality]:

$$S_{\mathbb{E}}(p_{\theta}, q) = \mathbb{E}_{q(\boldsymbol{x})}q(y|\boldsymbol{x})\log p_{\theta}(y|\boldsymbol{x}) \leq \mathbb{E}_{q(\boldsymbol{x})}q(y|\boldsymbol{x})\log q(y|\boldsymbol{x}). \tag{1}$$

Thus, minimizing the negative log-likelihood (NLL), which is equivalent to the softmax cross entropy loss in classification, is a proper scoring rule. Interestingly minimizing the squared error between the predictive probability of a label and one-hot encoding of the correct label, is also a proper scoring rule known as the Brier score. In regression, if we assume  $p_{\theta}$  to be Gaussian  $\mathcal{N}(\mu_{\theta}(\boldsymbol{x}), \sigma_{\theta}^{2}(\boldsymbol{x}))$ :

## **TODO**: Derive the NLL loss for Gaussian assumption.

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$$-\log p_{ heta}(y|oldsymbol{x}) = rac{1}{2}\log(2\pi) + \log(\sigma_{ heta}(oldsymbol{x})) + rac{(y-\mu_{ heta}(oldsymbol{x}))^2}{2\sigma_{ heta}^2(oldsymbol{x})}$$
 (2)

Of course we can model the  $p_{\theta}$  to be other distributions, e.g. [Laplacian distribution], which corresponds to L1 loss, MAE, rather than Gaussian distribution, which corresponds to L2 loss, MSE. If we assume  $p_{\theta}$  to be Laplacian  $\mathcal{L}(\mu_{\theta}(\boldsymbol{x}), \sigma_{\theta}(\boldsymbol{x}))$ , with mean  $\mu_{\theta}(\boldsymbol{x})$  and variance  $\sigma_{\theta}(\boldsymbol{x})$ :

## **TODO**: Derive the NLL loss for Laplacian assumption.

$$-\log p_{ heta}(y|oldsymbol{x}) = \log(2\sigma_{ heta}(oldsymbol{x})) + rac{|y - \mu_{ heta}(oldsymbol{x})|}{\sigma_{ heta}(oldsymbol{x})}$$
 (3)

## Adversarial training

Adversarial examples (see [3] and [4]) are those which are very close to the original training examples but are misclassified by the NN. In [4], the authors proposed the *fast gradient sign method* as a fast solution to generate adversarial examples. Given  $(\boldsymbol{x},y)$  and the loss  $\ell(\theta,\boldsymbol{x},y)$ , we can use the fast gradient sign method to generate an adversarial example as  $\boldsymbol{x}' = \boldsymbol{x} + \epsilon \mathrm{sign} \big( \nabla_{\boldsymbol{x}} \ell(\theta,\boldsymbol{x},y) \big)$ , where  $\epsilon$  is a small value to bound the perturbation. The procedure to use this adversarial example as an additional training

sample to augument the training set, referred to as *adversarial training* was found to improve the classifier's robustness [4].

There are more than one way to generate adversarial examples, for example, *virtual* adversarial training (VAT)[5], which does not require to know the true target y, could be used in *semi-supervised learning* (SSL).

## **Ensembles**

In general there are two classes of ensembles: randomization-based approaches and boosting-based approaches. Here we only focus on randomization-based approaches since it is more suitable for parallel computation. In practice, random initialization of the NN parameters  $\theta$  and with random shuffling of the data points during training were found to be sufficient to obtain good performance. During inference, the ensemble is treated as a uniformly-weighted mixture model and the predictions are combined as  $p(y|\boldsymbol{x}) = M^{-1} \sum_{m=1}^{M} p_{\theta_m}(y|\boldsymbol{x},\theta_m)$ . For classification, this corresponds to averaging the predicted probabilities. For regression, if we assume a Gaussian distribution over label y given point  $\boldsymbol{x}$ , i.e.  $p(y|\boldsymbol{x}) = \mathcal{N}(\mu,\sigma)$  for some  $\mu$  and  $\sigma$ , then the prediction's distribution is a mixture of Gaussian distributions (as we have one Gaussian output for each model in the ensemble). We then approximate the ensemble prediction as a Gaussian, whose mean and variance are respectively the mean and variance of the mixture:

$$\mu_*(oldsymbol{x}) = M^{-1} \sum_{m=1}^M \mu_{ heta_m}(oldsymbol{x})$$
 (4)

$$\sigma_*^2(\boldsymbol{x}) = M^{-1} \sum_{m=1}^M (\sigma_{\theta_m}^2(\boldsymbol{x}) + \mu_{\theta_m}^2(\boldsymbol{x})) - \mu_*^2(\boldsymbol{x}).$$
 (5)

And if we assume a Laplacian distribution over label y given input x, then the mean and variance of the mixture is:

$$\mu_*(\boldsymbol{x}) = M^{-1} \sum_{m=1}^{M} \mu_{\theta_m}(\boldsymbol{x})$$
 (6)

$$\sigma_*^2(\boldsymbol{x}) = M^{-1} \sum_{m=1}^M (2\sigma_{\theta_m}^2(\boldsymbol{x}) + \mu_{\theta_m}^2(\boldsymbol{x})) - \mu_*^2(\boldsymbol{x}).$$
 (7)

# Regression on toy dataset

```
In [21]: #!pip install -U matplotlib
In [35]: import jax
import jax.numpy as jnp
from jax import grad, jit, vmap, value_and_grad
from jax import random
import matplotlib.pyplot as plt
from jax.scipy.special import logsumexp
import numpy as np
from jax.example_libraries import optimizers
```

```
import time
import sys
# Add this at the top of your cell
```

Note that for the sake of numerical stability, we assume that the prediction variance is modeled as follows:

$$\sigma = \log(1 + \exp(\rho)) + \epsilon 
= \operatorname{softplus}(\rho) + \epsilon$$
(8)

where  $\rho$  is the model output in  $\mathbb{R}$ , and  $\sigma$  is in  $(\epsilon, +\infty)$ . Usually,  $\epsilon$ , a very small number, is used to avoid division by zero.

```
In [99]: # A helper function to randomly initialize weights and biases
         # for a dense neural network layer
         def random_layer_params(m, n, key, scale=1e-2):
             w_key, b_key = random.split(key)
             return scale * random.normal(w_key, (n, m)), scale * random.normal(b_key, (n)
         # Initialize all layers for a fully-connected neural network with sizes "sizes"
         def init_network_params(sizes, key, scale):
             keys = random.split(key, len(sizes))
             return [random_layer_params(m, n, k, scale) for m, n, k in zip(sizes[:-1], s
         def predict(params, inputs):
             # per-example predictions
             activations = inputs
             for w, b in params[:-1]:
                 outputs = jnp.dot(w, activations) + b
                 activations = jax.nn.relu(outputs)
             final_w, final_b = params[-1]
             scores = jnp.dot(final_w, activations) + final_b
             mean, rho = jnp.split(scores, 2)
             variance = jax.nn.softplus(rho) + 1e-6
             return mean, variance # for NLL loss
             # return scores # for MSE loss
         batched_predict = vmap(predict, in_axes=(None, 0))
         def loss(params, inputs, targets):
             preds = batched predict(params, inputs)
             return jnp.mean(jnp.square(preds - targets))
         def loss_l1(params, inputs, targets):
             preds = batched predict(params, inputs)
             return jnp.mean(jnp.absolute(preds - targets))
         def NLLloss_gaussian(params, inputs, targets):
             Negative log-likelihood loss function.
             _____
             TODO: Implementation required.
             The ``inputs`` argument and ``targets`` of this function are both of shape (
             1. Feed forward the data to NN to obtain mean and variance (use batched_pred
```

```
2. Calculate the negative log-likelihood using the outputs of the NN, based
   mean, variance = batched_predict(params, inputs)
   nll_vector = 0.5 * jnp.log(2*jnp.pi) + 0.5 * jnp.log(variance) + ((targets
   nll_loss = jnp.mean(nll_vector)
   #raise NotImplementedError("Task: Implement!")
   #nll loss = NotImplemented
   return nll_loss
def NLLloss_laplacian(params, inputs, targets):
   Negative log-likelihood loss function.
   TODO: Implementation required.
   _____
   The ``inputs`` argument and ``targets`` of this function are both of shape (
   1. Feed forward the data to NN to obtain mean and variance (use batched_pred
   2. Calculate the negative log-likelihood using the outputs of the NN, based
   mean, variance = batched_predict(params, inputs)
   nll_vector = jnp.log(2 * variance) + jnp.abs(targets - mean) / variance
   nll_loss = jnp.mean(nll_vector)
   #raise NotImplementedError("Task: Implement!")
   #nll_loss = NotImplemented
   return nll_loss
def NLLlossAT_gaussian(params, inputs, targets, eps=0.08):
   Negative log-likelihood loss function with adversarial training
   _____
   TODO: Implementation required.
   ______
   The ``inputs`` argument and ``targets`` of this function are both of shape (
   1. Use the ``value and grad`` function together with the
       ``NLLloss_gaussian`` function you implemented to get
      the loss value and the gradients with respect to the ``inputs``.
      Note that you also need to pass ``argnums`` to ``value_and_grad``.
   2. Generate adversarial examples. Note that you need to call jnp.array
      on gradients to be able to compute the sign.
      So use jnp.sign(jnp.array(grads)) instead of jnp.sign(grads).
   3. Calculate the NLL loss based on the adversarial examples and average the
   value, grads = value_and_grad(NLLloss_gaussian, argnums=1)(params, inputs, t
   nll = NLLloss_gaussian(params, inputs, targets)
   sign_gradients = jnp.sign(jnp.array(grads))
   adversarial_inputs = inputs + sign_gradients * eps
   nll_ad = NLLloss_gaussian(params, adversarial_inputs, targets)
   return (nll + nll_ad)/2
def NLLlossAT laplacian(params, inputs, targets, eps=0.08):
   Negative log-likelihood loss function with adversarial training
   _____
   TODO: Implementation required.
   _____
   The ``inputs`` argument and ``targets`` of this function are both of shape (
   1. Use the ``value_and_grad`` function together with the
```

```
the loss value and the gradients with respect to the ``inputs``
                 Note that you also need to pass ``argnums`` to ``value_and_grad``.
              2. Generate adversarial examples. Note that you need to call jnp.array
                 on gradients to be able to compute the sign.
                 So use jnp.sign(jnp.array(grads)) instead of jnp.sign(grads).
              3. Calculate the NLL loss based on the adversarial examples and average the
              value, grads = value_and_grad(NLLloss_laplacian, argnums=1)(params, inputs,
              nll = NLLloss_laplacian(params, inputs, targets)
              sign_gradients = jnp.sign(jnp.array(grads))
              adversarial_inputs = inputs + sign_gradients * eps
              nll_ad = NLLloss_laplacian(params, adversarial_inputs, targets)
              return (nll + nll_ad)/2
          @jit
          def update(params, x, y, opt_state):
              """ Compute the gradient for a batch and update the parameters """
              # value, grads = value_and_grad(loss)(params, x, y)
              # value, grads = value_and_grad(NLLloss_gaussian)(params, x, y)
              #value, grads = value_and_grad(NLLlossAT_gaussian)(params, x, y)
              value, grads = value_and_grad(NLLloss_laplacian)(params, x, y)
              #value, grads = value_and_grad(NLLlossAT_laplacian)(params, x, y)
              opt_state = opt_update(0, grads, opt_state)
              return get_params(opt_state), opt_state, value
          def data set(points=20, xrange=(-4, 4), std=3.):
              xx = jnp.array([[np.random.uniform(*xrange)] for i in range(points)])
              yy = jnp.array([[x**3 + np.random.normal(0, std)] for x in xx])
              return xx.reshape(-1,1), yy.reshape(-1,1)
In [101...
         # network parameters
          layer sizes = [1, 10, 6, 30, 2]
          # param scale = 0.02
          num epochs = 600
          step size = 8e-2
          # to ensure reproducibility
```

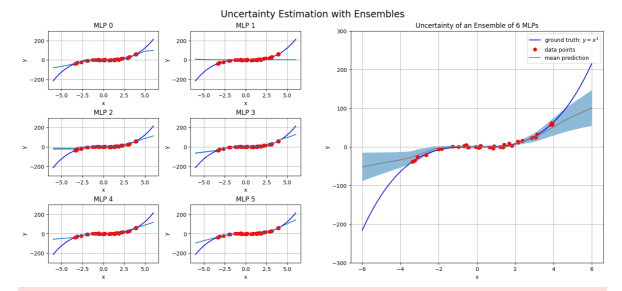
``NLLloss\_laplacian`` function you implemented to get

```
np.random.seed(173)
xx, yy = data_set(points=40, xrange=(-4, 4), std=3.) # generate data set of 20 s
x = np.linspace(-6, 6, 100).reshape(-1, 1)
y = x^{**}3
x = jnp.array(x)
y = jnp.array(y)
# number of ensembles
M = 6
params all = [None]*M
opt_state_all = [None]*M
```

```
mean_all, var_all = [], []
opt_init, opt_update, get_params = optimizers.adam(step_size)
for i in range(M):
   # use random seed to randomly initialize the NN
   SEED = 100 + abs(int(np.random.normal(0,1)*10000))
   np.random.seed(SEED)
   param_scale = abs(np.random.normal(0,0.1))
   print("-----param_scale "+str(pa
   params_all[i] = init_network_params(layer_sizes, random.PRNGKey(i*10), param
   opt_state_all[i] = opt_init(params_all[i])
   log_acc_train, log_train_loss = [], []
   for epoch in range(num_epochs):
       start_time = time.time()
       params_all[i] = get_params(opt_state_all[i])
       params_all[i], opt_state_all[i], train_loss = update(params_all[i], xx,
       if epoch == 0:
           print("initial loss: ",train_loss)
       epoch_time = time.time() - start_time
       log_train_loss.append(train_loss)
   print("final loss: ",train_loss)
   mean, var = batched_predict(params_all[i], x)
   mean_all.append(mean)
   var_all.append(var)
mean all = np.array(mean all)
var all = np.array(var all)
mean = mean all.mean(axis=0)
# TODO: Implementation required.
# Calculate the variance for the mixture model, according to your assumption, Ga
#Gaussian assumption
mean_gaussian = jnp.sum(mean_all, axis=0) / M
var_gaussian = jnp.sum(var_all + mean_all**2, axis=0) / M - mean_gaussian**2
#laplacian assumption
mean laplacian = jnp.sum(mean all, axis=0) / M
var_laplacian = jnp.sum(2 * var_all + mean_all**2, axis=0) / M - mean_laplacian*
#raise NotImplementedError("Task: Implement!")
var = var laplacian
std_ = np.sqrt(var_)
```

```
-----training network 0-----param_scale 0.05341477341995916
      initial loss: 21.935171
      final loss: 2.6328175
      -----param_scale 0.12899350046504218
      initial loss: 25.6118
      final loss: 4.399467
      -----param_scale 0.06450740460784611
      initial loss: 20.565968
      final loss: 2.847689
      -----training network 3-----param_scale 0.1859958185502153
      initial loss: 20.35938
      final loss: 2.4713166
      -----training network 4-----param_scale 0.021813948079358225
      initial loss: 21.813131
      final loss: 2.4738007
      -----training network 5-----param_scale 0.17893712367557665
      initial loss: 22.986002
      final loss: 2.1825185
In [ ]: # visualize ensemble output
        fig = plt.figure(constrained_layout=True,
                       figsize=(15, 7),
                       dpi=100)
        fig.suptitle('Uncertainty Estimation with Ensembles',
                    fontsize='xx-large')
        subfigs = fig.subfigures(1, 2, wspace=0.07)
        axs_left = subfigs[0].subplots(3, 2)
        for i,ax in zip(range(M),axs_left.flat):
           ax.plot(x, y, "b-")
           ax.plot(xx,yy,"or")
           ax.plot(x, mean_all[i])
           ax.fill_between(x.reshape(-1,), (mean_all[i]-np.sqrt(var_all[i])).reshape(-1
           ax.grid()
           ax.set_title("MLP " + str(i))
           ax.set_xlabel("x")
           ax.set ylabel("y")
           ax.set_ylim((-300,300))
        axs_right = subfigs[1].subplots(1, 1)
        axs_right.plot(x, y, "b-", label="ground truth: $y=x^3$")
        axs_right.plot(xx,yy,"or", label="data points")
        axs_right.plot(x, mean_, label="mean prediction", color="grey")
        axs right.fill between(x.reshape(-1,), (mean -std ).reshape(-1,), (mean +std ).r
        axs_right.grid()
        axs right.set title("Uncertainty of an Ensemble of "+str(M)+" MLPs")
        axs_right.set_xlabel("x")
        axs_right.set_ylabel("y")
        axs_right.set_ylim((-300,300))
        axs_right.legend()
```

Out[]: <matplotlib.legend.Legend at 0x15694e26d20>



The Kernel crashed while executing code in the current cell or a previous cell.

Please review the code in the cell(s) to identify a possible cause of the failur e.

Click <a href='https://aka.ms/vscodeJupyterKernelCrash'>here</a> for more info.

View Jupyter <a href='command:jupyter.viewOutput'>log</a> for further details.

## **Discussion points:**

- In terms of regression, compared to only use NLL loss with a single network, what is the effect of adding Adversarial Training? How about adding ensembles of M networks?
- How is the number of networks, M, affecting the prediction and uncertainty estimation?

# Classification on toy dataset:

The NLL loss for categorical assumption is:

$$-\log p_{\theta}(y|\boldsymbol{x}) = -\log(\prod_{i=1}^{k} p^{y_i})$$

$$= -\sum_{i=1}^{k} \log(p_i^{y_i})$$
(10)

$$= -\sum_{i=1}^k \log(p_i^{y_i}) \tag{11}$$

$$= -\sum_{i=1}^{k} y_i \log(p_i) \tag{12}$$

$$= -\log(p_i) \tag{13}$$

where, for notational simplicity we assume  $y = [y_1, y_2, \dots, y_k]$  is the one-hot encoded representation of the random variable y.

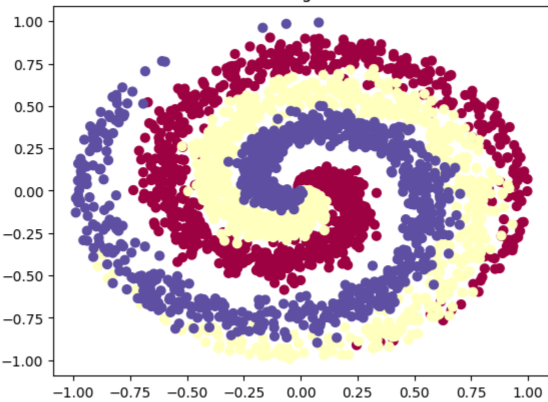
```
In [87]:
         def one_hot(x, k, dtype=jnp.float32):
             """Create a one-hot encoding of x of size k."""
             return jnp.array(x[:, None] == jnp.arange(k), dtype)
```

```
def generate_datasets(N,K,noise):
    X = np.zeros((N*K,D)) # data matrix (each row = single example)
   y = np.zeros(N*K, dtype='uint8') # class labels
   for j in range(K):
        ix = range(N*j,N*(j+1))
        r = np.linspace(0.0,1.,N) # radius
        t = np.linspace(j*8,(j+1)*8,N) + np.random.randn(N)*noise * (r+1.0) # th
        print(j, np.amin(t), np.amax(t))
        X[ix] = np.c_[r*np.sin(t), r*np.cos(t)]
        y[ix] = j
   X = jnp.array(X)
   y = jnp.array(y)
   y_onehot = one_hot(y, K)
   return X, y, y_onehot
def shuffle_dataset(X, y, y_onehot, seed):
   indices = jax.random.permutation(seed, jnp.arange(X.shape[0]))
   X_shuffled = jnp.take(X, indices, axis=0)
   y_oh_shuffled = jnp.take(y_onehot, indices, axis=0)
   y_shuffled = y[indices]
   return X_shuffled, y_shuffled, y_oh_shuffled
```

```
In [88]: N = 1000 # number of points per class
         D = 2 # dimensionality
         K = 3 # number of classes
         np.random.seed(0)
         X, y, y_onehot = generate_datasets(N, K, noise=0.3)
         # lets visualize the data:
         plt.figure()
         plt.scatter(X[:, 0], X[:, 1], c=y, s=40, cmap=plt.cm.Spectral)
         plt.title("Training Data")
         print("Training X:", X.shape)
         print("Training y:", y.shape)
```

```
0 -0.6210700567572351 9.160761050479454
1 7.506810982132371 16.88345105834359
2 15.192239293196302 25.21297645733773
Training X: (3000, 2)
Training y: (3000,)
```





```
In [95]: # A helper function to randomly initialize weights and biases
         # for a dense neural network layer
         def random_layer_params(m, n, key, scale=1e-1):
             w_key, b_key = random.split(key)
             return scale * random.normal(w_key, (n, m)), scale * random.normal(b_key, (n)
         # Initialize all layers for a fully-connected neural network with sizes "sizes"
         def init_network_params(sizes, key):
             keys = random.split(key, len(sizes))
             return [random layer params(m, n, k) for m, n, k in zip(sizes[:-1], sizes[1:
         @jit
         def predict(params, image):
             # per-example predictions
             activations = image
             for w, b in params[:-1]:
                 outputs = jnp.dot(w, activations) + b
                 activations = jax.nn.relu(outputs)
             final_w, final_b = params[-1]
             scores = jnp.dot(final_w, activations) + final_b
             probs = jax.nn.softmax(scores)
             return scores, probs
         batched_predict = vmap(predict, in_axes=(None, 0))
         def accuracy(params, inputs, targets):
             target_class = jnp.argmax(targets, axis=1)
             scores, _ = batched_predict(params, inputs)
             predicted_class = jnp.argmax(scores, axis=1)
             return jnp.mean(predicted_class == target_class)
         def NLLloss_categorical(params, inputs, targets):
```

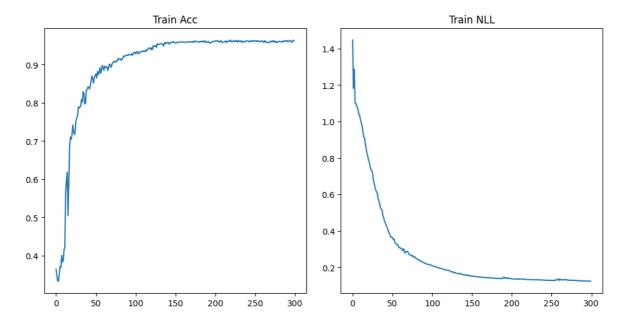
```
scores, preds = batched_predict(params, inputs)
    log_prob = scores - logsumexp(scores, axis=1)[:,None]
    return -jnp.mean(log_prob * targets)
def weight_decay_regularization(params):
    return jax.example libraries.optimizers.12 norm(params)
def loss(params, inputs, targets,wd):
    return NLLloss_categorical(params,
                              inputs,
                              targets) +\
           wd * weight_decay_regularization(params)
def NLLlossAT_categorical(params, inputs, targets, eps=0.02):
   Negative log-likelihood loss function with adversarial training
    _____
   TODO: Implementation required.
    The ``inputs`` argument and ``targets`` of this function are both of shape (

    Use the ``value_and_grad`` function together with the

       ``NLLloss_categorical`` function you implemented to get
      the loss value and the gradients with respect to the ``inputs``.
      Note that you also need to pass ``argnums`` to ``value_and_grad``
    2. Generate adversarial examples. Note that you need to call jnp.array
      on gradients to be able to compute the sign.
      So use jnp.sign(jnp.array(grads)) instead of jnp.sign(grads).
    3. Calculate the NLL loss based on the adversarial examples and average the
    value, grads = value_and_grad(NLLloss_categorical, argnums=1)(params, inputs
   loss_ori = NLLloss_categorical(params, inputs, targets)
   sign_gradients = jnp.sign(jnp.array(grads))
   ad_inputs = inputs + sign_gradients * eps
   loss ad = NLLloss categorical(params, ad inputs, targets)
   #raise NotImplementedError("Task: Implement!")
   #loss ori = NotImplemented
   #loss ad = NotImplemented
    return (loss ori + loss ad)/2
def lossAT(params, inputs, targets,wd):
    return NLLlossAT categorical(params,
                               inputs,
                               targets) +\
           wd * weight_decay_regularization(params)
@jit
def update(params, x, y, opt state, wd):
    """ Compute the gradient for a batch and update the parameters """
    \#value, qrads = value and qrad(loss)(params, x, y, wd)
   value, grads = value_and_grad(lossAT)(params, x, y, wd)
    opt state = opt update(0, grads, opt state)
    return get params(opt state), opt state, value
def logging(params_all, X, y, K, M, log_acc_train, log_nll_train):
    probs = jnp.mean( jnp.array([batched_predict(params_all[i], X)[1] for i in r
   y_pred = jnp.argmax(probs,axis=1)
```

```
train_acc = jnp.mean(y_pred == y)
             y_conf = probs[np.arange(y.shape[0]),y]
             train_nll = -jnp.mean(jnp.log(y_conf))
             log_acc_train.append(train_acc)
             log_nll_train.append(train_nll)
In [96]: layer_sizes = [2, 150, 150, 3]
         num_epochs = 300
         step_size = 0.1
         wd=1e-3
         M = 4
         K = layer_sizes[-1]
         params_all = [None]*M
         opt_state_all = [None]*M
         log_acc, log_nll = [], []
         opt_init, opt_update, get_params = optimizers.adam(step_size)
         for i in range(M):
             # use random seed to randomly initialize the NN
             params_all[i] = init_network_params(layer_sizes, random.PRNGKey(i))
             opt_state_all[i] = opt_init(params_all[i])
         key = random.PRNGKey(0)
         for epoch in range(num_epochs):
             for i in range(M):
                 key, seed = random.split(key)
                 X_shuffled, y_shuffled, y_oh_shuffled = shuffle_dataset(X, y, y_onehot,
                 params_all[i] = get_params(opt_state_all[i])
                 params_all[i], opt_state_all[i], train_loss = update(params_all[i], X_sh
             logging(params_all, X, y, K, M, log_acc, log_nll)
             print('\r', f'[Epoch {epoch+1}]: Train Acc: {log acc[-1]:.3f} | Train NLL: {
         [Epoch 300]: Train Acc: 0.962 | Train NLL: 0.124
In [97]: fig, axes = plt.subplots(1,2, figsize=(10,5))
         fig.tight_layout()
         axes[0].plot(log_acc)
         axes[1].plot(log_nll)
         axes[0].title.set_text('Train Acc')
```

axes[1].title.set text('Train NLL')



# Measuring Uncertainty - Total, Knowledge and Data Uncertainty

The Shannon entropy measures the information/uncertainty in a categorical distribution  $m{p}=[p_1,p_2,\ldots,p_K]$ 

$$H(p) = -\sum_{i=1}^K p_i \log p_i$$

The entropy is high when the probability is spread out over the classes and low when most of the probability is in a single class.

We can use entropy to measure the uncertainty in a prediction. Using the same notation as Malinin et al. [1], we call the average entropy of each prediction of the ensemble the **data uncertainty**. Let  $\boldsymbol{p}^{(1)},\ldots,\boldsymbol{p}^{(M)}$  be the M predictive distributions of the ensemble, then the data uncertainty is

$$rac{1}{M}\sum_{i=1}^{M}H\Big(oldsymbol{p}^{(i)}\Big)$$

The entropy of the mean prediction is called the **total uncertainty**, which is high if the data uncertainty is high or if the predictions are diverse.

**Knowledge uncertainty** (think: uncertainty in parameters) is related to the diversity of the predictions, which we measure with the multi-distribution Jensen-Shannon divergence

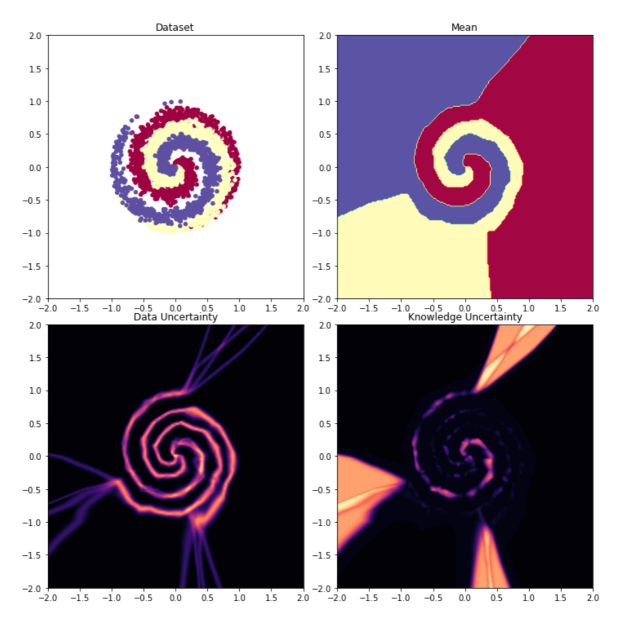
$$H\Big(rac{1}{M}\sum_{i=1}^{M}oldsymbol{p}^{(i)}\Big)-rac{1}{M}\sum_{i=1}^{M}H\Big(oldsymbol{p}^{(i)}\Big)$$

Indeed, the divergence measures the difference between total and data uncertainty. It is low when the predictions are similar and high when the predictions are different but confident.

Below, we visualize the data and knowledge uncertainty for our deep ensemble.

```
In [ ]: def entropy(preds):
            preds = np.clip(preds, 1e-7, 1.0)
            return -np.sum( preds * np.log(preds), axis=1)
        def js_terms(distributions):
            return entropy(np.mean(distributions, axis=0)), np.mean([entropy(p) for p in
        def visualize_predictions(X, y, params_list, min=-2.0, max=2.0, res=200, num_net
            xs = np.linspace(min, max, res)
            ys = np.linspace(min, max, res)
            N, M = len(xs), len(ys)
            xy = np.asarray([(_x,_y) for _x in xs for _y in ys])
            num_samples = xy.shape[0]
            predictions = [batched_predict(params, xy)[1] for params in params_list]
            predictions_ensemble = np.mean(predictions, axis=0)
            total, data = js_terms(predictions)
            Z, Z3 = np.zeros((N,M)), np.zeros((N,M)), np.zeros((N,M))
            indices = np.unravel_index(np.arange(num_samples), (N,M))
            Z[indices] = jnp.argmax(predictions_ensemble, axis=1)
            Z2[indices] = total - data
            Z3[indices] = data
            fig, axes = plt.subplots(2,2, figsize=(10,10))
            axes = axes.flatten()
            fig.tight_layout()
            axes[0].scatter(X[:, 0], X[:, 1], c=y, s=20, cmap=plt.cm.Spectral)
            axes[1].contourf(xs, ys, Z.T, cmap=plt.cm.Spectral, levels=50)
            axes[2].contourf(xs, ys, Z3.T, cmap='magma',levels=50)
            axes[3].contourf(xs, ys, Z2.T, cmap='magma', levels=50)
            axes[0].set_xlim([min, max]); axes[0].set_ylim([min, max]);
            axes[0].title.set_text('Dataset')
            axes[1].title.set text('Mean')
            axes[2].title.set_text('Data Uncertainty')
            axes[3].title.set_text('Knowledge Uncertainty')
```

In [ ]: visualize\_predictions(X, y, params\_all, num\_nets=M)



where dark to light increases in value.

#### **Discussion points:**

- Are there any regions where the network makes overconfident predictions?
- Where is the ensemble most uncertain (high entropy)? Why?
- Where are the members of the ensemble the most diverse? Why?
- How does adversarial training affect the results? Why?
- Finally, it is valuable for us to know how long did it take you to complete this practical?