

The Hidden Uncertainty in a Neural Network’s Activations

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Abstract

The distribution of a neural network’s latent representations has been successfully used to detect out-of-distribution (OOD) data. This work investigates whether this distribution moreover correlates with a model’s epistemic uncertainty, thus indicates its ability to generalise to novel inputs. We first empirically verify that epistemic uncertainty can be identified with the surprise, thus the negative log-likelihood, of observing a particular latent representation. Moreover, we demonstrate that the output-conditional distribution of hidden representations also allows quantifying aleatoric uncertainty via the entropy of the predictive distribution. We analyse epistemic and aleatoric uncertainty inferred from the representations of different layers and conclude that deeper layers lead to uncertainty with similar behaviour as established - but computationally more expensive - methods (e.g. deep ensembles). While our approach does not require modifying the training process, we follow prior work and experiment with an additional regularising loss that increases the information in the latent representations. We find that this leads to improved OOD detection of epistemic uncertainty at the cost of ambiguous calibration close to the data distribution. We verify our findings on both classification and regression models.

1. Introduction

The recent success of deep neural networks in a variety of applications (Hinton et al., 2012; Redmon et al., 2016) has been primarily driven by improvements in predictive performance, while progress on safety-related issues remained relatively slow (Amodei et al., 2016; Bozhinowski et al., 2019; Janai et al., 2017). Thereby, quantifying the uncertainty associated with the predictions of a neural network is a central

challenge and a prerequisite for their broader deployment. One commonly distinguishes between two types of uncertainty (Kiureghian & Ditlevsen, 2009). Uncertainties arising from noise in the data regardless of the model are called *aleatoric*. Uncertainties originating from the model choice and parameter fitting are referred to as *epistemic*. This differentiation also has practical reasons, since *epistemic uncertainty is reducible* - e.g. by using more data - while aleatoric uncertainty, as a property of the data, is not.

The predominant *holistic* framework for both aleatoric and epistemic uncertainty is Bayesian Deep Learning (BDL) (MacKay, 1992; Hinton & Van Camp, 1993). However, scaling BDL to neural networks of practical size remains an open research question. Although recent research proposed various *scalable* approaches (Kingma et al., 2015; Gal & Ghahramani, 2016; Zhang et al., 2017; Postels et al., 2019), they still fall short of delivering a widely adopted practical solution. Moreover, the quality of the resulting posterior distribution, in particular for scalable approximations, is open to debate (Osband, 2016; Wenzel et al., 2020).

Prior work used the hidden representations for detecting OOD samples (Papernot & McDaniel, 2018; Lee et al., 2018). Moreover, density estimates of latent distributions have been implied to yield epistemic uncertainty when regularised appropriately (Alemi et al., 2018; van Amersfoort et al., 2020). The prospect that the distribution of latent representations holds information about the uncertainty associated with the predictions of a neural network promises efficient and scalable uncertainty estimates. However, prior work enforced additional constraints on the architecture, focused on classification and only evaluated on OOD detection. Nevertheless, epistemic uncertainty should not only identify far-away OOD samples, but also indicate how well a model generalises to its input, thus correlate with model performance *in proximity of* the training data distribution. Furthermore, hidden representations have, thus far, not been shown to also quantify aleatoric uncertainty. We demonstrate the latter by estimating the output-conditional density.

We show that the latent distribution in fact gives rise to a holistic framework for estimating both epistemic and aleatoric uncertainty for classification and regression - even without additional regularisation of the latent space en-

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abling widespread application. To this end, we examine uncertainty estimates extracted from different architectures and layers of various depth. Aleatoric uncertainty is assessed by its correlation with the prediction error on the training data distribution. Epistemic uncertainty is evaluated using OOD data - far away from and close to the training data distribution. We find that density estimates of shallow layers yield more conservative epistemic uncertainty, i.e. are more likely to label new data as OOD. The density of deeper layers behaves less conservatively and more similar to established - but computationally more expensive - epistemic uncertainty estimates (section 4.1). Moreover, we discuss practical challenges that arise from uncertainty estimation based on hidden representations (section 3.3). We further investigate how additional regularisation at training time in our framework impacts epistemic uncertainty (section 4.3).

2. Related Work

2.1. Density Estimation for Out-of-Distribution Detection

Density estimation is an established technique for OOD detection. While research also developed alternative methods (Pimentel et al., 2014; Chalapathy & Chawla, 2019), generative models are particularly appealing, since they denote a principled, unsupervised approach to the problem by learning the data distribution (Schlegl et al., 2017; Li et al., 2018). However, concerns have been voiced regarding their effectiveness (Škvára et al., 2018) and, in particular for explicit generative models, vulnerability to OOD samples (Nalisnick et al., 2019a). Ensembles of generative models (Choi et al., 2018) have been proposed to solve the latter, as well as considering the distribution of log-likelihoods (Nalisnick et al., 2019b; Morningstar et al., 2020).

Since learning high-dimensional densities is challenging, another line of research focuses on applying machine learning approaches to the latent space of neural networks (Ruff et al., 2018; Papernot & McDaniel, 2018) or directly estimating its density, which is often simpler than for the data itself. (Lee et al., 2018) fits a Gaussian mixture model (GMM) to the representations of a neural network. (Blum et al., 2019) trains a normalizing flow on the latent space. Moreover, (Alemi et al., 2018) utilizes the rate term in the Information Bottleneck (VIB) (Alemi et al., 2016) for OOD detection.

2.2. Uncertainty Estimation

The idea that uncertainty should increase outside of the training data distribution relates uncertainty estimation to OOD detection. Thus, uncertainty estimation has been applied to the latter (Hendrycks & Gimpel, 2016; Li & Gal, 2017; Snoek et al., 2019). We discuss this relation in section 3.1. BDL (Neal, 2012; Neal et al., 2011; Hinton & Van Camp,

1993) constitutes the predominant way to quantify uncertainty, since it handles both aleatoric and epistemic uncertainty in a principled manner. Due to well known difficulties of applying BDL to large neural networks, numerous attempts have been proposed (Kingma et al., 2015; Gal & Ghahramani, 2016; Zhang et al., 2017; Teye et al., 2018; Postels et al., 2019). However, current approaches still lack scalability or quality of the predicted uncertainty.

A parallel line of work uses the predictive conditional distribution of neural networks to quantify uncertainty (Bishop, 1994; Hendrycks & Gimpel, 2016; Lee et al., 2017). However, these approaches meet fundamental difficulties outside of the training data distribution. Furthermore, neural networks have been found to be poorly calibrated (Guo et al., 2017), which poses a challenge to all uncertainty estimates.

Recently, it has been suggested that the distribution of a neural network’s latent representations quantifies epistemic uncertainty. (Mandelbaum & Weinshall, 2017) fits it with a kernel density. They outperform the softmax entropy and Monte-Carlo (MC) dropout (Gal & Ghahramani, 2016) when imposing an additional regularising loss on the latent space. (van Amersfoort et al., 2020) replaces the softmax activation with a target-conditional Gaussian distribution with fixed variance for classification. Their approach paired with a gradient penalty on the input, boosts OOD performance by enforcing distance awareness of the representations. They refer to the distance to the closest target-conditional mean as epistemic uncertainty and evaluate it on OOD detection. Similarly, (Liu et al., 2020) enforces the representations of a neural network to be distance-aware by constraining its Lipschitz constant. In (Alemi et al., 2018) the authors consider the per-sample rate loss in the VIB objective (Alemi et al., 2016) and the softmax entropy as proxies for uncertainty. They show that the softmax entropy excels at quantifying aleatoric uncertainty, and the rate-loss at OOD detection. Above works are united by the fact that they introduce additional regularisation which forces the hidden representations to store more information about the input. Additional hyperparameters are tuned to maximise OOD performance. We discuss this further in section 3.3. On the contrary, our work reveals that similar conclusions can be drawn for deterministic neural networks without altering the training procedure while additionally incorporating aleatoric uncertainty.

3. Uncertainty from Latent Representations

3.1. Types of Uncertainty

Uncertainty estimation in machine learning (ML) tries to assign a level of confidence to a model’s output. Thus, the true uncertainty correlates with a model’s performance. While a correct prediction can have high uncertainty, average model performance degrades when uncertainty increases.

A model’s performance fluctuates within the training data distribution. One reason for this is inherent noise in the data, which causes irreducible *aleatoric* uncertainty (Kiureghian & Ditlevsen, 2009). Additionally, the performance depends on the choice of training data, model, and parameters. Uncertainties and performance drops resulting from these choices are reducible and should be distinguished from their irreducible counterparts. (Kiureghian & Ditlevsen, 2009) call these uncertainties ‘*epistemic*’. The same terminology was introduced to the ML community (Senge et al., 2014; Gal, 2016). It is worth noting that some works aim at separating *distributional* from *epistemic* uncertainty (Malinin & Gales, 2018; Harang & Rudd, 2018). We follow the definitions from (Kiureghian & Ditlevsen, 2009), where *epistemic* uncertainty encompasses all reducible uncertainty.

The Two Faces of Epistemic Uncertainty. Many related works solely evaluate epistemic uncertainty using OOD detection (Mandelbaum & Weinshall, 2017; Alemi et al., 2018; van Amersfoort et al., 2020; Liu et al., 2020). This wrongly equates epistemic uncertainty estimation with OOD detection. While OOD detection, as an important failure source, is related to epistemic uncertainty estimation and, thus, is a necessary evaluation method, we stress that OOD detection performance alone is not sufficient. This becomes apparent when considering that OOD detection represents a model agnostic task and epistemic uncertainty should be aligned with a model’s generalisation. Consequently, high epistemic uncertainty is only expected for those OOD samples that lead to degradation of model performance. In this work we therefore evaluate epistemic uncertainty along two dimensions - its behaviour close to and far away from the training distribution. We expect that a good epistemic uncertainty estimate successfully detects samples that reside far away from the data distribution as well as indicates how well the model generalises to data close to it. We particularly investigate the latter in section 4.1 by applying perturbations of increasing magnitude to the input while tracking estimated uncertainty and model performance.

3.2. Uncertainty of Deterministic Neural Networks

We now establish a theoretical connection between the density of hidden representations and the information-theoretic surprise of observing a particular sample or prediction under the assumption of a deterministic neural network. While we require the model to be deterministic here, this only affects the inference stage of a model. Thus, one is free to use any stochastic regularisation methods at training time. Moreover, we relate this surprise to aleatoric and epistemic uncertainty estimates used by prior work.

Given a dataset $\mathcal{D} = \{\mathbf{X}, \mathbf{Y}\}$, the goal of a discriminative task is to model the conditional distribution $p(\mathbf{y}|\mathbf{x})$, where $\mathbf{x} \in \mathbf{X}$ and $\mathbf{y} \in \mathbf{Y}$. Let $\hat{\mathbf{y}}$ correspond to the output of a neural network trained to approximate $p(\mathbf{y}|\mathbf{x})$, which induces the

conditional distribution $p_\theta(\hat{\mathbf{y}}|\mathbf{x})$. The conditional entropy

$$\begin{aligned} H(\hat{\mathbf{y}}|\mathbf{x}) &= E_{p_\theta(\hat{\mathbf{y}}|\mathbf{x})} [-\log(p_\theta(\hat{\mathbf{y}}|\mathbf{x}))] \\ &= E_{p(\mathbf{x})} \left[-\int d\hat{\mathbf{y}} p_\theta(\hat{\mathbf{y}}|\mathbf{x}) \log(p_\theta(\hat{\mathbf{y}}|\mathbf{x})) \right] \end{aligned} \quad (1)$$

corresponds to the total uncertainty about $\hat{\mathbf{y}}$ given knowledge of \mathbf{x} across the dataset \mathcal{D} . Both quantities under the expectation have been successfully used to quantify aleatoric uncertainty in deterministic neural networks for classification (Hendrycks & Gimpel, 2016; Alemi et al., 2018)¹ and regression (Kendall & Gal, 2017)². The negative log-likelihood $-\log(p_\theta(\hat{\mathbf{y}}|\mathbf{x}))$ is called *surprisal* of a particular value $\hat{\mathbf{y}}$ given an input \mathbf{x} . Further, $-\int d\hat{\mathbf{y}} p_\theta(\hat{\mathbf{y}}|\mathbf{x}) \log(p_\theta(\hat{\mathbf{y}}|\mathbf{x}))$ is the *expected surprisal* for a given input \mathbf{x} .

However, eq. 1 only quantifies the uncertainty/surprise about predictions given a specific input \mathbf{x} and not about observing \mathbf{x} in the first place. To account for this, consider the entropy of the joint probability distribution

$$\begin{aligned} H(\hat{\mathbf{y}}, \mathbf{x}) &= H(\mathbf{x}) + H(\hat{\mathbf{y}}|\mathbf{x}) \\ &= E_{p(\mathbf{x})} \left[-\log(p(\mathbf{x})) - \int d\hat{\mathbf{y}} p_\theta(\hat{\mathbf{y}}|\mathbf{x}) \log(p_\theta(\hat{\mathbf{y}}|\mathbf{x})) \right]. \end{aligned} \quad (2)$$

In eq. 2 the negative log-likelihood $-\log(p(\mathbf{x}))$ quantifies the surprise about observing \mathbf{x} and requires learning the density $p(\mathbf{x})$. Estimating $-\log(p(\mathbf{x}))$ directly would provide an uncertainty estimate independent of the neural network used to parameterize $p_\theta(\hat{\mathbf{y}}|\mathbf{x})$ and its generalisation. It can be considered as a model-agnostic distributional uncertainty. Moreover, this distributional uncertainty incorporates the entire information in the distribution of \mathbf{X} - also information irrelevant for the discriminative task. Additionally, a high-dimensional \mathbf{X} induces further complications, since learning such densities is an active field of research.

Consider a neural network comprised of L layers with $L - 1$ latent representations $(\mathbf{z}_0, \dots, \mathbf{z}_{L-2})$ with a joint distribution factorizing according to $p_\theta(\mathbf{x}, \hat{\mathbf{y}}, \mathbf{z}_0, \dots, \mathbf{z}_{L-2}) = p_\theta(\hat{\mathbf{y}}|\mathbf{z}_{L-2})p(\mathbf{z}_{L-2}|\mathbf{z}_{L-3})\dots p_\theta(\mathbf{z}_0|\mathbf{x})p(\mathbf{x})$. Exploiting its deterministic nature and the data processing inequality allows us to make the following statements:

$$H(\hat{\mathbf{y}}|\mathbf{z}_i) = H(\hat{\mathbf{y}}|\mathbf{x}) \quad \forall i \in [0, \dots, L-2] \quad (3)$$

$$H(\mathbf{x}) \geq H(\mathbf{z}_0) \geq \dots \geq H(\mathbf{z}_{L-2}) \quad (4)$$

A detailed derivation is in the supplement. These inequalities relate the distribution of the latent representations to the uncertainties in eq. 2. Inequality 4 implies that uncertainty

¹In fact, (Hendrycks & Gimpel, 2016) uses the maximum softmax probability. This does not change the ordering of the uncertainty values, since the logarithm is a strictly monotonic function.

²The authors parameterize the output with a unimodal Gaussian and use its variance as aleatoric uncertainty. In this case the entropy is proportional to the logarithm of the variance.

estimates based on deeper layers are less conservative. On the other side, since the distribution of the latent space at a particular layer is a functional of all the previous layers, we expect these uncertainty estimates to incorporate some degree of model dependence. We verify this intuition empirically in section 4.1 by detecting OOD samples using density estimate of different layers.

We also investigate whether the latent distribution contains information about aleatoric uncertainty. Therefore, we learn the joint density $p(\mathbf{z}_i, \hat{\mathbf{y}})$ at layer i . This can be achieved by separately estimating the output-conditional distribution $p(\mathbf{z}_i|\hat{\mathbf{y}})$ and $p(\hat{\mathbf{y}})$. Using Bayes’ formula and marginalization over $\hat{\mathbf{y}}$, we can compute both values under the expectations in eq. 1. In section 4.1 we find empirically that deeper layers enable better aleatoric uncertainty estimates. We hypothesize that this results from the neural network filtering out task-irrelevant information, which makes it easier to estimate the output-conditional density.

To summarise, given a novel input \mathbf{x}^* with corresponding latent vector \mathbf{z}^* , we identify the epistemic uncertainty with the surprise of observing that latent representation $-\log p(\mathbf{z}_i^*)$ and the aleatoric uncertainty with the expected surprisal of $\hat{\mathbf{y}} \sim p(\hat{\mathbf{y}}|\mathbf{z}_i^*)$ which we subsequently denote with $h(\hat{\mathbf{y}}|\mathbf{z}_i^*)$:

$$-\log p(\mathbf{z}^*) = -\log \left(\int d\hat{\mathbf{y}} p(\mathbf{z}^*|\hat{\mathbf{y}}) p(\hat{\mathbf{y}}) \right) \quad (5)$$

$$h(\hat{\mathbf{y}}|\mathbf{z}_i^*) = - \int d\hat{\mathbf{y}} p_{\theta}(\hat{\mathbf{y}}|\mathbf{z}_i^*) \log(p_{\theta}(\hat{\mathbf{y}}|\mathbf{z}_i^*)) \quad (6)$$

3.3. Pitfalls

Feature Collapse. One of the strengths of discriminative neural networks is to ignore task-irrelevant features of the data. However, this can lead to activations of OOD data becoming indistinguishable from activations of the training data. This phenomenon is also called **feature collapse** (van Amersfoort et al., 2020) and poses a practical challenge for uncertainty quantification based on hidden activations. Different prior works have addressed this problem by tuning additional hyperparameters enforcing adequately structured latent spaces. Hereby, the network is made more sensitive to changes in the input by constraining its Lipschitz constant. This is achieved by using a distance-based loss on the hidden representations (Mandelbaum & Weinshall, 2017), a gradient penalty (van Amersfoort et al., 2020) or spectral normalization (Liu et al., 2020). Further, although (Aleml et al., 2018) does not mention it explicitly, training with the VIB framework regularises the distribution of hidden representations according to a prior distribution.

In our main contribution (section 4.1 & 4.2) we do not explicitly address feature collapse, since it requires altering the training procedure, adds architectural constraints and, most importantly, risks overfitting epistemic uncertainty to

the task of OOD detection. All mentioned prior works focus on OOD detection and it remains unclear whether such approaches translate into good epistemic uncertainty, especially close to the training data distribution. We demonstrate in section 4.1 that, *without additional regularisation*, the density of the hidden representations of deeper layers as a proxy for epistemic uncertainty is similarly calibrated as other well established - but computationally more expensive - methods, e.g. deep ensembles (Lakshminarayanan et al., 2017), while maintaining good OOD detection performance.

Furthermore, in order to link these results to prior works above and illustrate the risk of overfitting epistemic uncertainty to OOD detection, we explore within the framework of eq. 4 how regularisation techniques that yield \mathbf{z}'_l such that $H(\mathbf{x}) \geq H(\mathbf{z}'_l) \geq H(\mathbf{z}_l)$ influence the estimated epistemic uncertainty. Therefore, we augment the original loss by a weighted reconstruction objective to achieve more informative latent representations \mathbf{z}'_l :

$$\tilde{\mathcal{L}} = \mathcal{L}_{orig} + \lambda \text{MSE}(\mathbf{x}, \hat{\mathbf{x}}) \quad (7)$$

where \mathbf{x} is the input and $\hat{\mathbf{x}} = D(\mathbf{z}'_l)$ is the reconstruction from an additional decoder network D , which is a symmetric version of the encoding network (see supplement for details). We evaluate the impact of this regularisation on epistemic uncertainty in section 4.3.

High-Dimensional Densities. The hidden representations are high-dimensional for large neural networks or shallow layers. This renders estimating their density impractical. Prior work has also faced the **curse of dimensionality**. For example, (Lee et al., 2018) reduced the dimensionality of hidden representations by performing principal component analysis (PCA) on the training data and, subsequently, reusing the transformation at inference time. Moreover, (Papernot & McDaniel, 2018) uses **locality-sensitive hashing** in combination with **random projections** for nearest neighbour search. Here, we follow (Lee et al., 2018) applying PCA and find that it yields satisfying results in practice. However, we note that PCA as well as random projections are not optimal solutions in the case of convolutional neural networks whose hidden representations entail translational invariances that get lost by applying dense projection matrices.

3.4. Implementation

In order to then estimate epistemic (eq. 5) and aleatoric uncertainty (eq. 6), we extract the latent representations of a particular layer i and learn the output-conditional density $p(\mathbf{z}_i|\hat{\mathbf{y}})$ and the marginal distribution of the neural network’s outputs $p(\hat{\mathbf{y}})$ on the training data.

We learn $p(\mathbf{z}_i|\hat{\mathbf{y}})$ using an explicit generative model. For image classification (section 4.1), we use one GMM with $k \in \mathbb{N}_{\setminus \{0\}}$ components for each class. For regression (section 4.2) which entails continuous predictions, we found

that conditional normalizing flows (CNFs) and in particular the conditioning scheme used in (Ardizzone et al., 2019) yielded good results. We refer the reader to the supplementary material for details.

Estimating $p(\hat{y})$ for a classification task simply involves counting the predicted labels on the training data. For a regression task, however, it is necessary to estimate the density of the predictions in the output space. This is relatively simple, since the output space is low dimensional for most common discriminative tasks. We found it sufficient to approximate it with common parametric distributions - a uniform distribution in section 4.2 and a betaprime distribution for the predicted depth in the supplement.

We then obtain the epistemic uncertainty in eq. 5 by marginalizing \hat{y} . To solve the integral, we use summation (classification) or numerical integration (regression). We found the latter to be sufficient for the one dimensional output spaces in our regression experiments (see section 4.2 and the supplement). Then, we calculate the aleatoric uncertainty in eq. 6 by applying Bayes’ formula.

4. Experiments

While there exists no ground truth for uncertainty, we follow the approach of other works (Gal & Ghahramani, 2016; Lakshminarayanan et al., 2017; Malinin & Gales, 2018) and examine epistemic uncertainty over input transformations and distributional shifts and aleatoric uncertainty via the correlation with the network’s predictive performance on in-distribution data. We first investigate the behaviour of uncertainties extracted from the output-conditional density over varying depth of the representations on image classification. We then validate this approach for the task of regression on a toy dataset and finally investigate whether additional training losses can improve the extracted uncertainty. We refer to the supplementary material for additional experiments with semantic segmentation and depth regression, as well as all experimental details and parameters.

4.1. Image Classification

We train a multilayer perceptron (FCNet) consisting of fully-connected layers and ReLU activations on MNIST (LeCun et al., 1998) and FashionMNIST (Xiao et al., 2017) and ResNet18 (He et al., 2016) on CIFAR10 (Krizhevsky et al., 2009) and SVHN (Netzer et al., 2011).

4.1.1. DEPTH DEPENDANT UNCERTAINTY QUALITY

We estimate the output-conditional density of the hidden representations on the training data of FCNet trained on MNIST and of ResNet18 trained on SVHN using one GMM with five components per class. Each training and subsequent density estimation is conducted five times independently

yielding mean and standard deviation. We quantify epistemic and aleatoric uncertainty according to eq. 5 and eq. 6. We compare our results to MC dropout (Gal & Ghahramani, 2016) and deep ensembles (Lakshminarayanan et al., 2017) for which we compute aleatoric and epistemic uncertainty according to (Gal et al., 2017) (see supplement).

First, we evaluate the behaviour of the epistemic uncertainty estimate when the input is shifted away from the training data distribution in small discrete steps. We expect epistemic uncertainty to correlate with model generalisation, which is the observed behaviour of established methods such as MC dropout and deep ensembles. We chose to compare with these methods due to their popularity. We compute the area under the ROC curve (AUROC) between the unperturbed testset and each perturbation magnitude. The AUROC measures how well a binary classifier identifies the perturbed test samples based on the estimated epistemic uncertainty. On MNIST we rotate test samples from 0° to 180° in 20° steps and on SVHN we add independent Gaussian noise to the pixel values of input images, while increasing the standard deviation from 0 to 80 with step size 10. Fig. 1 shows the results.

We observe that the epistemic uncertainty obtained from density estimates from shallow layers behaves more conservatively in the sense that they are quicker to label perturbed data as OOD. On the other side, epistemic uncertainty estimates from deeper layers behave less conservatively and similar to MC Dropout and Deep Ensembles, while only requiring training one model and a single forward pass.

Fig. 2 analyzes the quality of the estimated aleatoric uncertainty using calibration curves. These show the accuracy of the neural network depending on the uncertainty. We use a threshold sliding through increasing percentiles of the aleatoric uncertainty and plot the remaining accuracy. Ideally this yields a strictly monotonically decreasing curve. We compare aleatoric uncertainty from various layers, MC dropout, deep ensembles and the softmax entropy.

Our experiments show that aleatoric uncertainty from density estimates of deeper layers demonstrates similar calibration curves to established approaches. However, aleatoric uncertainty based on shallow layers can perform poorly (e.g. ResNet18 on SVHN). As discussed in section 3.3, we argue that this results from difficulties in estimating the density of very high-dimensional hidden representations.

4.1.2. EPISTEMIC UNCERTAINTY ON OOD DETECTION

This experiment evaluates the OOD detection performance of epistemic uncertainty. Table 1 reports average performance over 5 independent trainings, with standard deviations in the supplementary material. We measure the AUROC of the OOD detection using epistemic uncer-

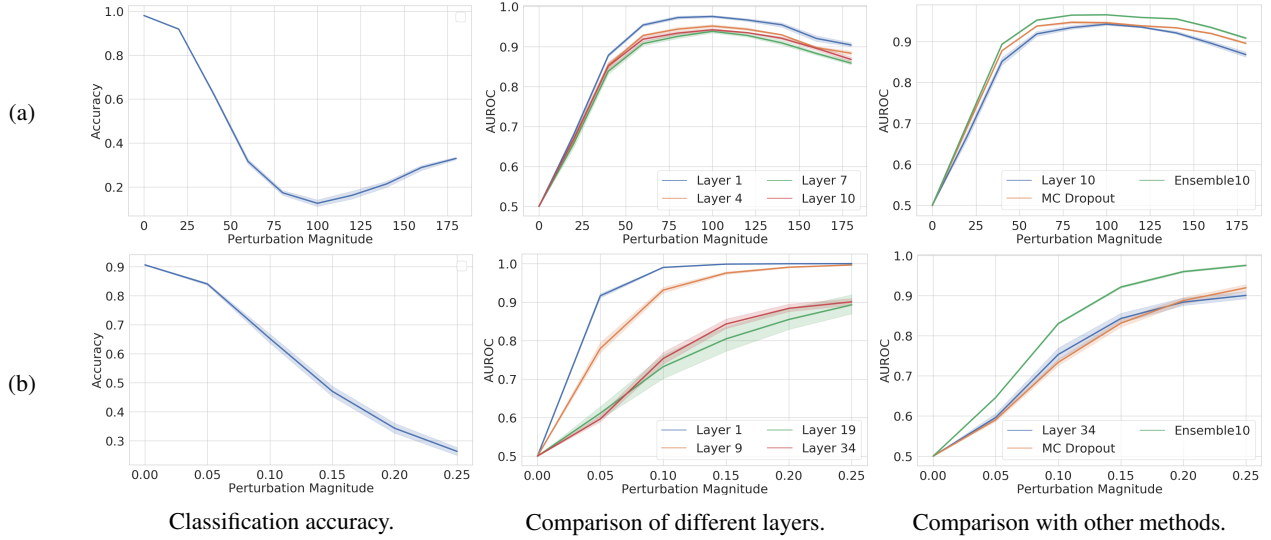


Figure 1: Analysis of the quality of estimated epistemic uncertainty over different layers for FCNet on MNIST (a), and ResNet18 on SVHN (b). We plot the accuracy of the classification network (**left**) and the area under the ROC curve (AUROC) (obtained using epistemic uncertainty as threshold) against the perturbation magnitude on the test data (**center** and **right**). We compare epistemic uncertainty obtained from hidden representations of varying depth (**center**) and other methods (**right**). We observe that (i) density estimates based on deeper layers behave similar as other established - but computationally more expensive - methods (i.e. MC Dropout and Deep Ensembles) and (ii) density estimates based on shallow layers yield more conservative estimates.

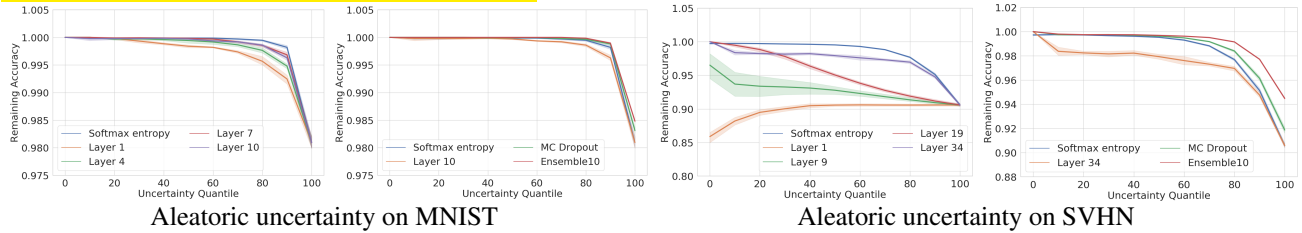


Figure 2: Analysis of the calibration of aleatoric uncertainty on MNIST and SVHN. We plot the remaining accuracy when selecting test samples using incrementally increased aleatoric uncertainty threshold. We observe that aleatoric uncertainty based on deeper layers tends to behave similar as other approaches (softmax entropy, deep ensembles, MC dropout).

tainty of different layers and compare with deep ensembles. When training on MNIST/FashionMNIST we evaluate on FashionMNIST/MNIST, OMNIGLOT (Lake et al., 2015) and when training on CIFAR10/SVHN we evaluate on SVHN/CIFAR10, STL10 (Coates et al., 2011). Additionally we evaluate on test data perturbed with additive Gaussian noise, rotated by 90° and flipped horizontally/vertically.

We generally observe a strong performance of epistemic uncertainty obtained from shallow layers which coincides with the results of section 4.1.1, where we found that shallow layers yield more conservative estimates. However, we find that, when using a convolutional architecture (ResNet18), epistemic uncertainty obtained from shallow layers fails to detect OOD data generated by globally transforming the test data. We argue that this is expected behaviour since the hidden representations of shallow layers in convolutional networks denote low-level features that are likely to not

represent such global transformations. Finally, we note the poor performance of shallow layers when training on CIFAR10 and evaluating on SVHN. These results suggest that recently found difficulties of explicit generative models on the task of OOD detection (Nalisnick et al., 2019a) also apply to shallow layers. However, epistemic uncertainty from deeper layers does not suffer from this problem.

4.2. Regression

As a simple regression experiment, we train regressors on data sampled from $f(x) = \frac{1}{2}(\sin(4\pi x - \frac{\pi}{2}) + x)$ for $x \in [-1, 1]^3$. The training data and methods are illustrated in fig. 3. We train on the gray outlined data with a simple four-layer perceptron, extract embeddings at the penultimate layer (following our findings from section 4.1) and fit the density with a CNF. For inference, we perform numerical

³Designed to be naturally normalised to $[-1, 1] \rightarrow [-1, 1]$

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	OOD Data	L 1	L 4	L 7	L 10	Ensemble
Trained on MNIST	FashionMNIST	0.975	0.922	0.855	0.811	0.896
	OMNIGLOT	0.972	0.937	0.892	0.893	0.979
	white noise	1.000	0.972	0.903	0.841	0.785
	Rotated 90°	0.978	0.976	0.950	0.935	0.965
	HFlip	0.902	0.907	0.883	0.864	0.905
	VFlip	0.887	0.868	0.851	0.830	0.881
Trained on FashionMNIST	MNIST	0.985	0.991	0.975	0.978	0.962
	OMNIGLOT	0.971	0.987	0.960	0.967	0.960
	white noise	1.000	0.985	0.971	0.930	0.840
	Rotated 90°	0.884	0.780	0.804	0.835	0.670
	HFlip	0.719	0.696	0.701	0.693	0.657
	VFlip	0.898	0.891	0.891	0.901	0.845

	OOD Data	L 1	L 9	L 19	L 34	Ensemble
Trained on SVHN	CIFAR10	0.991	0.974	0.934	0.907	0.976
	STL10	0.999	0.991	0.951	0.912	0.982
	white noise	1.000	1.000	0.986	0.903	0.992
	Rotated 90°	0.615	0.646	0.689	0.918	0.957
	HFlip	0.500	0.503	0.495	0.500	0.501
	VFlip	0.506	0.520	0.551	0.708	0.736
Trained on CIFAR10	SVHN	0.042	0.029	0.091	0.736	0.723
	STL10	0.790	0.871	0.821	0.651	0.806
	white noise	1.000	1.000	1.000	0.681	0.999
	Rotated 90°	0.553	0.517	0.543	0.757	0.824
	HFlip	0.500	0.500	0.499	0.500	0.501
	VFlip	0.519	0.513	0.537	0.714	0.789

Table 1: OOD performance in terms of AUROC on OOD datasets when **estimating epistemic uncertainty** using the hidden activations. We evaluate the epistemic uncertainty computed from the distribution of the output of several affine layers. We evaluate layers (L) 1, 4, 7, 10 for a multilayer perceptron trained on MNIST or FashionMNIST (**left**) and layers 1, 9, 19, 34 for a ResNet18 trained on SVHN or CIFAR10 (**right**). For comparison, we further report the performance of an ensemble of 10 identical networks. We observe that uncertainty estimates based on shallow layers demonstrate strong OOD performance.

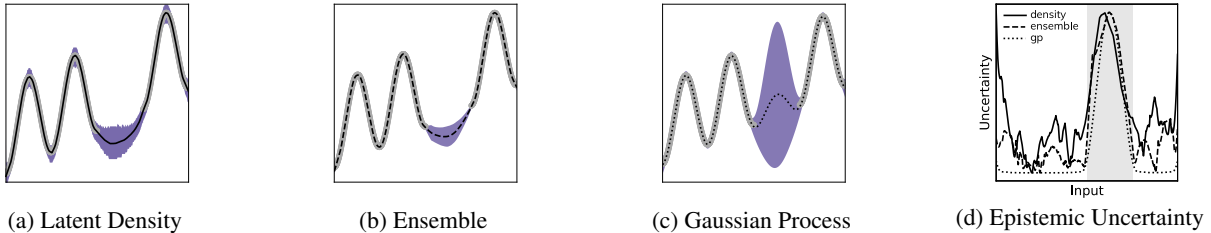


Figure 3: Simple regression task. The models are trained on the data outlined in gray. **(a-c)**: Model prediction is given in black and the high-confidence area in blue. In **(d)**, we compare the normalised estimated epistemic uncertainty of latent density (solid), deep ensemble (dashed) and a Gaussian process (GP) (dotted). The gray background indicates the gap in the training data. **Also for regression we find that the epistemic uncertainty estimated by density estimation matches established methods.**

integration to evaluate the likelihood of the latent vectors. The same architecture is used for the ensemble. The Gaussian process (GP) has a RBF kernel with parameters fit to the training data.

Fig. 3 compares the latent density approach (a) to a Deep Ensemble (b) and a GP (c). All methods show growing uncertainty with further distance to the training data, indicating that latent densities also contain information about epistemic uncertainty in regression networks. For a better evaluation, we compare $-\log p(\mathbf{z}_i^*)$ against other epistemic uncertainty methods in fig. 3d. We observe that all methods follow the expected behaviour of high uncertainty in the gray data gap and low uncertainty in the region of the training data, while the estimated epistemic uncertainty is slightly increasing towards the boundaries of the training distribution for the ensemble and the latent density.

4.3. Effects of Regularisation

To **confirm that more informative hidden representations translate into better OOD detection**, we apply an additional reconstruction loss (section 3.3). We show this on FCNet trained on MNIST and evaluated FashionMNIST and on

ResNet18 trained on SVHN and evaluated on STL10. We reconstruct the input from layer 10/19 (FCNet/ResNet18)⁴.

Fig. 4 shows that in both cases the AUROC increases with the weight of the reconstruction loss and the reconstruction quality (PSNR). Most importantly, the improvement in AUROC is already achieved with low-weighted reconstruction losses that do not cause a noticeable reduction in accuracy.

Fig. 5 shows how the accuracy and AUROC evolve when the original in-distribution dataset is perturbed with increasing magnitude⁵. Following section 4.1 we apply rotations to MNIST and additive Gaussian noise to SVHN. We can see that the reconstruction weight has no impact on the shape of the accuracy curve, however we generally observe larger AUROCs with increasing reconstruction weights. Note, that this ambiguity complicates the hyperparameter choice since it is difficult to determine which AUROC behavior close to the training data distribution is optimal. Training without

⁴Unlike for FCNet we do not choose the last layer in tab. 1 for ResNet18 since layer 34 has only 10 dimensions which we consider too small for reconstructing the input image.

⁵We choose the maximum reconstruction weight such it does not negatively impact best accuracy on the validation set.

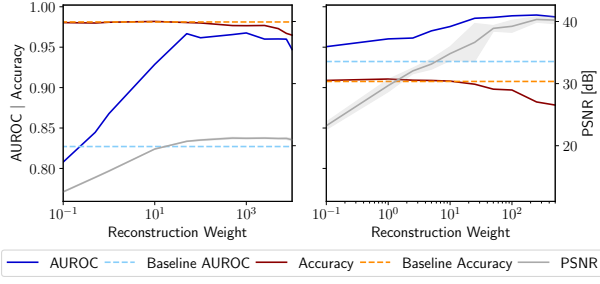


Figure 4: Detecting FashionMNIST/STL10 (left/right) when training on MNIST/SVHN with varying weight of the reconstruction loss. We use layer 10/19 for reconstruction and AUROC computation. Baseline Accuracy & AUROC show the performance of a model trained without additional reconstruction loss. More informative representations boost OOD detection significantly.

the reconstruction loss tends induce strong correlation with the behavior MC dropout while higher weights tend to be close to deep ensembles (compare with fig. 1).

5. Discussion & Conclusion

This work introduced a holistic framework for uncertainty estimation based on the density of latent representations and verified its effectiveness empirically on the task of classification (section 4.1) and regression (section 4.2). In the absence of additional regularisation losses, we found that latent representations of shallow layers yield more conservative uncertainty estimates, while epistemic uncertainty obtained from deeper layers behaves similarly to MC Dropout and Deep Ensembles (section 4.1.1). By applying perturbations of increasing magnitude to the test data (section 4.1.1), we showed that latent densities denote an effective proxy for epistemic uncertainty and, thus, quantify a model’s ability to generalise. This has promising practical implications since evaluating the latent density is faster than other state-of-the-art approaches and can be applied to any neural network post-training. Moreover, section 4.1.1 showed that the calibration of aleatoric uncertainty from deeper layers is similar to other well established methods.

Section 4.1.2 investigated the OOD performance of epistemic uncertainty and concludes that epistemic uncertainty from shallow layers has superior performance, which is in line with the results in section 4.1.1. They even outperform Deep Ensembles for various OOD datasets, while adding negligible computational overhead and leaving the training procedure unchanged. This comes at the cost of worse uncertainty calibration close to the training data distribution (section 4.1). We note that this result is achieved without applying an additional regularisation loss. However, we also find that certain global perturbations of the data distribution are better detected using the density of deeper layers.

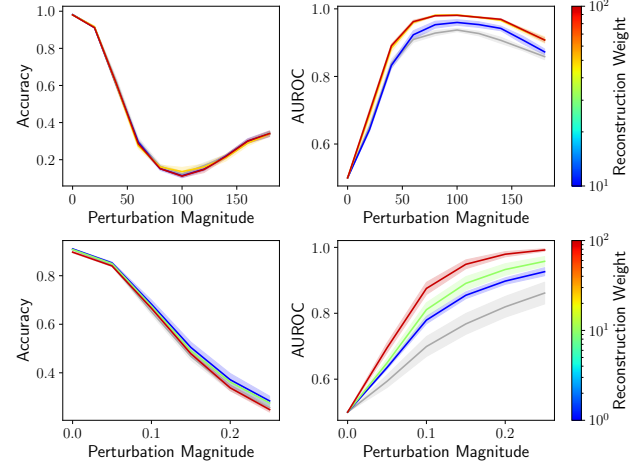


Figure 5: OOD detection performance on rotated MNIST (top) and SVHN with additive Gaussian noise (bottom) for various perturbation magnitudes and reconstruction weights. We reconstruct and estimate uncertainty from layers 10 (MNIST) and 19 (SVHN). The baseline without reconstruction loss is shown in gray. The AUROC increases with higher reconstruction losses while the Accuracy does not deviate much from the baseline.

This suggests that the task of OOD detection can benefit from aggregating uncertainty information from several layers. We leave this exploration for future work. In fact, (Lee et al., 2018) utilizes density estimates from several layers. However, thus far they require logistic regression trained on OOD data for merging the results from several layers.

Moreover, section 4.3 demonstrated that using a simple heuristic for increasing the information stored in the hidden representations further improves the OOD detection performance significantly, which is in line with prior work. However, our experiments in general showed that for uncertainty estimation OOD detection performance is not a sufficient metric. Different weights of the regularisation term lead to identical accuracy curves with very different AUROC curves close to the training data distribution. This leads to difficulties in choosing the optimal regularisation weight. Overall, if OOD detection is the primary objective of epistemic uncertainty additional regularisation can boost performance. However, note that a similar boost can also be achieved with density estimates of shallow layers while leaving the training procedure unchanged. When taking different aspects of epistemic uncertainty into account leaving out additional regularisation and using density estimates of deeper layers is a strong baseline given its simplicity and decent performance.

In conclusion, we find that the hidden activations of neural networks contain information about both aleatoric and epistemic uncertainty. Uncertainty proxies derived from density estimates of deep layers closely follow those of es-

established - but computationally more expensive - methods. The finding that these uncertainties are available in already trained models enables a wide range of applications and rapid deployment.

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Appendix

The supplementary material is structured in the following way: Section A provides additional explanations and proofs regarding the proposed method. Section B lists all experimental details and additional results to facilitate reproduction and reimplementaion of our experiments. Finally, section C shows two case-studies on deployment of latent activation based uncertainty estimation to real-world autonomous driving scenarios.

A. Background

A.1. Process of Estimating the Output-Conditional Distribution of Hidden Representations

Algorithm 1 show schematically how to estimate the output-conditional density of hidden representations which is needed to quantify uncertainty with the proposed approach.

Algorithm 1 Estimating the output-conditional distribution of hidden representations.

Input: Dataset $\{X, Y\}$, Index l of layer used for uncertainty estimation, d maximum dimension of hidden representations.

Result: Trained model M , output-conditional density model M_{gen} of hidden representations at layer l , estimate of $P(\hat{Y})$

Train model M on $\{X, Y\}$

$Z_l \leftarrow$ activations at layer l on $\{X, Y\}$

$\hat{Y} \leftarrow M(X)$

if $\dim(Z_l) > d$ **then**

$Z_l \leftarrow$ reduce dimension of Z_l using PCA

end

Initialize generative model M_{gen}

Train M_{gen} on Z_l given \hat{Y} to estimate $P(Z_l|\hat{Y})$

if *classification* **then**

Estimate marginal categorical distribution of network predictions $P(Y)$ by counting frequencies

else

Estimate marginal distribution of network predictions $P(\hat{Y})$ with univariate parametric distribution (e.g. Gaussian, beta prime)

end

A.2. Proof of Equation 3 and 4

Consider a deterministic neural network comprised of L layers with $L - 1$ latent representations $(\mathbf{z}_0, \dots, \mathbf{z}_{L-2})$ and some input \mathbf{x} . Since the network is deterministic we know

that $p(\mathbf{z}'_i|\mathbf{x}) = \delta(\mathbf{z}'_i - f_i(\mathbf{x}))$ and consequently:

$$\begin{aligned} p(\hat{\mathbf{y}}|\mathbf{x}) &= \int p(\hat{\mathbf{y}}|\mathbf{z}'_i)p(\mathbf{z}'_i|\mathbf{x})d\mathbf{z}'_i \\ &= \int p(\hat{\mathbf{y}}|\mathbf{z}'_i)\delta(\mathbf{z}'_i - f_i(\mathbf{x}))d\mathbf{z}'_i \\ &= p(\hat{\mathbf{y}}|\mathbf{z}_i) \end{aligned} \quad (8)$$

where $\mathbf{z}_i = f_i(\mathbf{x})$ denotes the evaluation of the neural network until layer i . Eq. (3), $H(\hat{\mathbf{y}}|\mathbf{z}_i) = H(\hat{\mathbf{y}}|\mathbf{x}) \forall i \in [0, L - 1]$, follows directly from that. Furthermore, according to the **data processing inequality**, any deterministic function f applied to a random variable x can only decrease its (differential) entropy: $H(x) \geq H(f(x))$. Considering that the z_i are deterministic functions of the z_{i-1} proves eq. (4): $H(\mathbf{x}) \geq H(\mathbf{z}_0) \geq \dots \geq H(\mathbf{z}_{L-2}) \forall i \in [0, L - 1]$.

A.3. Conditional Normalizing Flows

Normalizing Flows (NFs) (Dinh et al., 2014; 2017) are a family of explicit generative models based on **invertible neural networks**. Assume we wish to model the distribution p_X of some data X . Given some parametric distribution p_Z and a NF f_θ comprised of weights θ , likelihood evaluation of a data point $x \in X$ in NFs is based on the change of variable formula

$$p_X(x) = p_Z(f_\theta(x)) \left| \det \left(\frac{\partial f_\theta(x)}{\partial x} \right) \right| \quad (9)$$

and made feasible by designing invertible layers that allow evaluating the determinant of their Jacobian efficiently. One such layer is the **coupling layer** introduced by (Dinh et al., 2017). Each coupling layer splits its input activations into two sets $u_{1/2}^{in}$ and uses one set u_1^{in} to compute scaling and translation of the other set u_2^{in} using **trainable, non-invertible functions g_s and g_t** , while it applies the identity function to u_1^{in} .

$$\begin{aligned} u_1^{out} &= u_1^{in} \\ u_2^{out} &= (u_2^{in} + g_t(u_1^{in})) \odot g_s(u_1^{in}) \end{aligned}$$

This transformation renders the Jacobian a lower triangular matrix where the determinant is simply the product of the main diagonal.

Several ways have been proposed to use NFs for modeling the conditional distribution $p_{X|C}$ of $x \in X$ conditioned on some random variable $c \in C$. We adapt the conditioning scheme used in (Ardizzone et al., 2019), where the authors use a conditional version of the above coupling layer. Consequently, the coupling layers in the conditional normalizing flow $f_\theta(x|c)$ become:

$$u_1^{out} = u_1^{in}$$

$$u_2^{out} = (u_2^{in} + g_t(u_1^{in}; c)) \odot g_s(u_1^{in}; c)$$

g_s and g_t are implemented as multi-layer perceptrons. To feed the conditional information into g_s and g_t , we apply a separate multi-layer perceptron to c and add the result to $g_t(u_1^{in})$ and $g_s(u_1^{in})$.

A.4. Estimating Aleatoric and Epistemic Uncertainty with Deep Ensembles and Monte-Carlo Dropout

In the case of MC dropout and deep ensembles, following (Gal et al., 2017) aleatoric uncertainty u_{al} is computed as the average entropy of the predicted distribution (for several samples from the weight distribution):

$$u_{al} = E_{w \sim p(w)} [E_{y \sim p(\hat{y}|x, w)} [\log(p(\hat{y}|x, w))]]$$

Here, $p(\hat{y}|x, w)$ is the predicted distribution given an input x and a set of weights w . $p(w)$ is the weight distribution (i.e. approximated posterior distribution). Here, we omit the conditioning of the weight distribution on the data. The epistemic uncertainty u_{ep} is then estimated by approximating the mutual information between the weights w and the predictions \hat{y} conditioned on the input x :

$$u_{ep} = I(\hat{y}, w|x)$$

$$= H(\hat{y}|x) - H(\hat{y}|w, x)$$

$$= E_{y \sim p(\hat{y}|x)} [\log(p(\hat{y}|x))] - u_{al}$$

Here, $p(\hat{y}|x) = \int dw p(w) p(\hat{y}|x, w)$ is evaluated using a finite set of samples/ensemble members.

B. Experimental Details

B.1. Image Classification

B.1.1. ARCHITECTURES

The architectures of the neural networks used in our image classification experiments can be found in fig. 6. We applied the multi-layer perceptron (fig. 6) to MNIST and Fashion-MNIST and ResNet18 (He et al., 2016) to CIFAR10 and SVHN. Note that the dropout layer in FCNet only has $p > 0$ when using MC dropout.

Furthermore, table 2 and 3 show the correspondences between layer indices used in the main work and the layer type. Moreover, in the main work we primarily estimate the output-conditional distribution at the "Add" layers of ResNet18. This is done for convenience since it allows estimating the density from a single layer output instead of

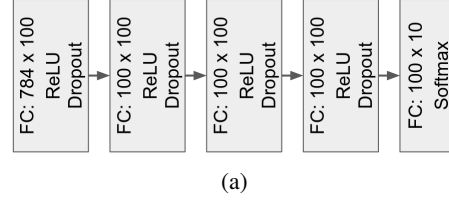


Figure 6: Architecture used in our image classification experiments. Multi-layer perceptron. FC denotes a fully-connected layer.

Index	Layer
0	Flatten
1	Linear
3	Dropout
4	Linear
5	ReLU
6	Dropout
7	Linear
8	ReLU
9	Dropout
10	Linear
11	ReLU
12	Dropout
13	Linear
14	Softmax

Table 2: Layer index to layer type for FCNet.

taking to parallel branches of the computational graph into account.

B.1.2. DEEP ENSEMBLE AND MC DROPOUT

All deep ensembles in section 4.1.1 consist of ten neural networks of the same architecture as used by our approach. When training MC Dropout, we place an additional dropout layer after every dense layer (FCNet) or after every Res-Block (resnet).

B.1.3. TRAINING

We describe the training of the neural networks on image classification and the output-conditional densities on the latent representations separately.

Neural Network Training

We train both, FCNet and ResNet18, using Adam (Kingma & Ba, 2014) ($\beta_1 = 0.9$, $\beta_2 = 0.999$) and a batch size of 32. We use a learning rate of 0.001 and weight decay of 0.0001. We train the networks for 200 epochs, while performing early stopping with a patience parameter of 20. For all datasets, CIFAR10, SVHN, MNIST and FashionMNIST, we use fixed validation set containing 20% of the training data. All reported results are on the test set of the corresponding dataset.

Output-conditional Density Training

After training the neural networks, we extract the activations of the training data at layers of varying depth with the cor-

Index	Layer
0	Input
1	Conv2D
2	Batchnorm2D
3	ReLU
4	Conv2D
5	Batchnorm2D
6	ReLU
7	Conv2D
8	Batchnorm2D
9	Add
10	Dropout
11	ReLU
12	Conv2D
13	Batchnorm2D
14	ReLU
15	Conv2D
16	Conv2D
17	Batchnorm2D
18	Batchnorm2D
19	Add
20	Dropout
21	ReLU
22	Conv2D
23	Batchnorm2D
24	ReLU
25	Conv2D
26	Conv2D
27	Batchnorm2D
28	Batchnorm2D
29	Add
30	Dropout
31	ReLU
32	AveragePooling2D
33	Flatten
34	Linear
35	Softmax

Table 3: Layer index to layer type for ResNet18.

responding prediction of the neural network. Subsequently, we estimate the distribution of the latent representations of each class prediction with a separate GMM of five components. We use the sklearn GMM implementation⁶. Since the shallow layers of ResNet18 are high-dimensional ($\gg 100$), density estimation - especially in the output-conditional case - becomes challenging. We counter this by reducing the dimensionality of the activations of each layer to 512 using a principal component analysis. Our multi-layer perceptron does not require dimensionality reduction, since each hidden layer has only 100 hidden units.

We also experimented with using NFs for this task. However, we found that a GMM is sufficient to fit the output-conditional distribution of latent representations throughout all layers.

B.1.4. ERROR MARGINS

We report the error margins over multiple runs on the OOD experiments in the tables 4 and 5.

B.2. Regression

B.2.1. MODEL TRAINING

For the regression prediction, we train a multilayer perceptron with 100 hidden units and 4 hidden layers until

convergence. We then extract latent activations z from the penultimate layer and train a CNF conditioned on the predictions of the multilayer perceptron over the whole training data. The CNF is based on GLOW (Kingma & Dhariwal, 2018) and consists of 4 layers with GLOW coupling blocks, each conditioned on the regression prediction, and random permutations. Since our training data is relatively small (750 samples), we do not use minibatches in both trainings.

For the ensemble, we take the same perceptron architecture and train 10 models with different random seeds.

The GP has a simple RBF kernel and is fit (with the sklearn implementation) to the same training data as the perceptron. The fit parameters are $0.601 \times 2 \times \text{RBF}(\text{length_scale}=0.164)$.

B.2.2. EVALUATION

To evaluate eq. 5, we take a range of 1000 support points of $y \in [-10, 10]$ and estimate $p(z|y)$ with the CNF. We further assume a uniform prior $p(y)$ and numerically integrate over the support points to yield $\log p(z)$.

For the visualised confidence regions, we start the integration at the prediction of the regression network and integrate separately in positive and negative direction until the integral reaches the set probability mass. In fig. 3, we visualize the integration boundaries to reach 20% of the probability mass from the CNF.

B.3. Effects of Regularisation

B.3.1. MODEL ARCHITECTURE

To explore the effects of regularization, we augment the model with a reconstruction network. The input to the reconstruction network is the latent vector extracted from the layer of our choice. The mapping from the input to the latent vector can be seen as the encoder. We choose the decoder for reconstruction to be symmetrical to the encoder such that their combination forms a symmetrical autoencoder. In the case of the fully connected network, we reconstruction the latent vector from layer k using another k fully connected layers as shown in fig. 7. The reconstruction network for the ResNet18 architecture follows a similar design. The difference is that the output of the convolutional layers has spatial dimensions, which we retain when feeding the latent tensor into the reconstruction network. The decoder is a ResNet18 model itself where the strided convolutions are replaced with a transpose convolution. We build the decoder with stacks of residual units where each stack upsamples the spatial resolution by a factor of two. The number of stacks can be directly inferred from the spatial resolution of the latent tensor.

⁶<https://scikit-learn.org/stable/modules/generated/sklearn.mixture.GaussianMixture.html>

The Hidden Uncertainty in a Neural Network’s Activations

	OOD Dataset	Layer 1	Layer 9	Layer 19	Layer 34	Ensemble
Trained on SVHN	CIFAR10	0.991 ± 0.001	0.974 ± 0.004	0.934 ± 0.008	0.907 ± 0.003	0.976 ± 0.002
	STL10	0.999 ± 0.000	0.991 ± 0.002	0.951 ± 0.009	0.912 ± 0.003	0.982 ± 0.002
	Gaussian noise	1.000 ± 0.000	1.000 ± 0.000	0.986 ± 0.011	0.903 ± 0.016	0.992 ± 0.002
	Rotated (90°)	0.615 ± 0.001	0.646 ± 0.010	0.689 ± 0.024	0.918 ± 0.002	0.957 ± 0.002
	HFlip	0.500 ± 0.000	0.503 ± 0.005	0.495 ± 0.010	0.500 ± 0.002	0.501 ± 0.002
	VFlip	0.506 ± 0.000	0.520 ± 0.004	0.551 ± 0.010	0.708 ± 0.006	0.736 ± 0.001
Trained on CIFAR10	SVHN	0.042 ± 0.001	0.029 ± 0.004	0.091 ± 0.016	0.736 ± 0.055	0.723 ± 0.048
	STL10	0.790 ± 0.010	0.871 ± 0.019	0.821 ± 0.038	0.651 ± 0.008	0.806 ± 0.004
	Gaussian noise	1.000 ± 0.000	1.000 ± 0.000	1.000 ± 0.000	0.681 ± 0.057	0.999 ± 0.002
	Rotated (90°)	0.553 ± 0.003	0.517 ± 0.020	0.543 ± 0.005	0.757 ± 0.001	0.824 ± 0.003
	HFlip	0.500 ± 0.000	0.500 ± 0.000	0.499 ± 0.001	0.500 ± 0.002	0.501 ± 0.001
	VFlip	0.519 ± 0.003	0.513 ± 0.006	0.537 ± 0.006	0.714 ± 0.001	0.789 ± 0.004

Table 4: OOD performance in terms of AUROC on various datasets when estimating epistemic uncertainty using the hidden activations of a resnet 18 on SVHN and CIFAR10. We evaluate AUROC using the epistemic uncertainty computed from the distribution of the output of several affine layers (1, 9, 19, 34). For comparison, we further report the performance of an ensemble containing 10 identical networks. We observe that uncertainty estimates based on shallow layers demonstrate strong OOD performance.

	OOD Dataset	Layer 1	Layer 4	Layer 7	Layer 10	Ensemble
Trained on MNIST	FashionMNIST	0.975 ± 0.001	0.922 ± 0.013	0.855 ± 0.032	0.811 ± 0.035	0.896 ± 0.018
	OMNIGLOT	0.972 ± 0.002	0.937 ± 0.006	0.892 ± 0.006	0.893 ± 0.011	0.979 ± 0.001
	Gaussian noise	1.000 ± 0.000	0.972 ± 0.005	0.903 ± 0.005	0.841 ± 0.010	0.785 ± 0.005
	Rotated (90°)	0.978 ± 0.001	0.976 ± 0.004	0.950 ± 0.005	0.935 ± 0.006	0.965 ± 0.002
	HFlip	0.902 ± 0.002	0.907 ± 0.005	0.883 ± 0.007	0.864 ± 0.007	0.905 ± 0.001
	VFlip	0.887 ± 0.001	0.868 ± 0.002	0.851 ± 0.004	0.830 ± 0.012	0.881 ± 0.002
Trained on FashionMNIST	MNIST	0.985 ± 0.001	0.991 ± 0.003	0.975 ± 0.004	0.978 ± 0.005	0.962 ± 0.004
	OMNIGLOT	0.971 ± 0.001	0.987 ± 0.002	0.960 ± 0.012	0.967 ± 0.008	0.960 ± 0.003
	Gaussian noise	1.000 ± 0.000	0.985 ± 0.002	0.971 ± 0.006	0.930 ± 0.013	0.840 ± 0.004
	Rotated (90°)	0.884 ± 0.002	0.780 ± 0.045	0.804 ± 0.030	0.835 ± 0.013	0.670 ± 0.013
	HFlip	0.719 ± 0.003	0.696 ± 0.007	0.701 ± 0.010	0.693 ± 0.008	0.657 ± 0.010
	VFlip	0.898 ± 0.006	0.891 ± 0.014	0.891 ± 0.006	0.901 ± 0.013	0.845 ± 0.014

Table 5: OOD performance in terms of AUROC on various datasets when estimating epistemic uncertainty using the hidden activations of a multilayer perceptron on MNIST and FashionMNIST. We evaluate AUROC using the epistemic uncertainty computed from the distribution of the output of several affine layers (1, 4, 7, 10). For comparison, we further report the performance of an ensemble of 10 neural networks of identical architecture. We observe that uncertainty estimates based on shallow layers demonstrate strong OOD performance.

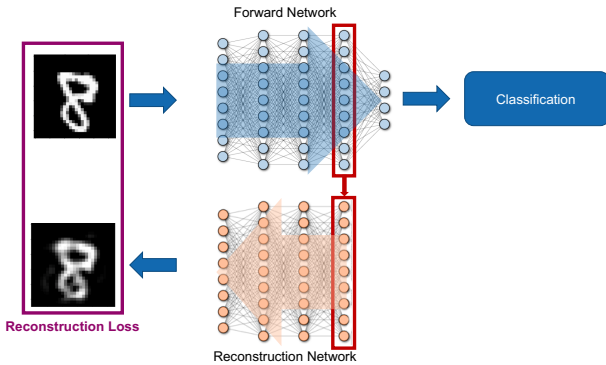


Figure 7: Overview of the model architecture with additional reconstruction network during training.

B.3.2. MODEL TRAINING

The combined model consisting of forward and reconstruction network are trained with the same parameters as described in section B.1.3. For fitting the output-conditional density after training, we only use the forward network. The reconstruction network is discarded after training as its sole purpose is regularization.

C. Additional Experiments on Real-World Models

In the following, we present two case studies on larger models that are trained to perform pixelwise predictions. Such models are often used in safety critical contexts like semantic segmentation for autonomous driving or segmentation of medical images. They also pose a specific implementation

method	Misclass. Detection AUROC	OOD Detection	
		AP	FPR ₉₅
softmax entropy	94	2.9	45
mc dropout	-	9.8	38
joint likelihood	70	3.1	24
conditional entropy	92	0.8	100

Table 6: Comparison against softmax entropy and monte-carlo dropout on the two downstream tasks of misclassification and OOD detection in semantic segmentation. All values in [%]. The aleatoric uncertainty estimated with the conditional entropy can detect misclassified pixel with high AUROC, matching the performance of the softmax entropy. For detecting anomalous objects, the epistemic uncertainty from mc dropout and the joint likelihood have very different performances in the two benchmark metrics. We conclude that aleatoric and epistemic uncertainty can be successfully extracted from latent activations and are separated as expected to indicate different uncertainty aspects.

problem for density based models since the convolutional latent features cannot be accumulated across the spatial dimension.

C.1. Semantic Segmentation

As a large-scale classification experiment, we evaluate our method on semantic segmentation for urban driving. We use the state-of-the-art DeepLabv3+ (Chen et al., 2018) network trained on the Cityscapes urban driving dataset (Cordts et al., 2016) and estimate the feature density in the last layer before the logits.

C.1.1. EXPERIMENTAL SETUP

We use the pre-trained DeepLabv3+ (Chen et al., 2018) provided by the authors⁷ as the base network for the experiments with semantic segmentation. To train the output-conditional GMMs, we extract latent embeddings from the last layer before the logits (decoder_conv1_0) before the relu activation. We found the embedding dimensionality of 256 to be low enough to fit densities directly, without further dimensionality reduction. To associate latent vectors with predicted classes, we interpolate them bilinearly to the output resolution of 1024x2048. To counter the correlations of latent vectors within the same image, we subsample a maximum of 500 vectors for every class and image. We then train the GMMs on a shuffled set of 100000 such vec-

⁷https://github.com/tensorflow/models/blob/master/research/deeplab/g3doc/model_zoo.md#deeplab-models-trained-on-cityscapes

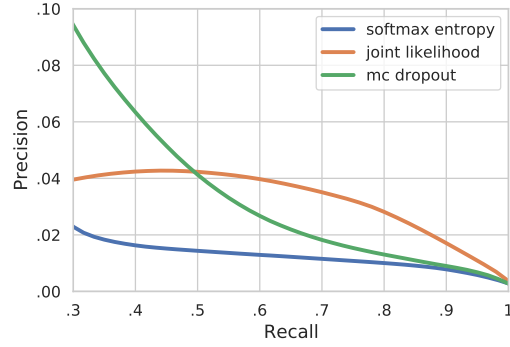


Figure 8: Precision-Recall curve for OOD detection in semantic segmentation on the Fishyscapes benchmark. The shown high-recall region is especially important for safety-critical applications. The curve illustrates the different characteristics that mc dropout and density based uncertainty exhibit on this task.

tors. We find suitable hyperparameters for the GMMs on the Fishyscapes Lost & Found validation set and use the same hyperparameters for all classes. The best average precision on the validation set was achieved with 2 mixture components per class, tied covariance matrices, 1000 iterations and covariance regularisation factor 0.0001. We show qualitative examples in figure 9.

For estimating aleatoric uncertainty, we experienced numerical issues when evaluating $p(\hat{y}|\mathbf{z}^*)$. This is due to the normalization $\sum_y p(\hat{y}|\mathbf{z}^*) = 1$ when some classes are much more likely than others, i.e. when the entropy is low. We counteract these numerical issues by evaluating $h(\hat{y}|\mathbf{z}^*)$ on underfit GMMs⁸ and on embedding vectors that were interpolated to output resolution (which was not found necessary for epistemic uncertainty and significantly increases memory usage).

C.1.2. RESULTS

We test the estimation of aleatoric uncertainty by measuring the performance for misclassification detection on the Cityscapes validation data. Due to the nature of misclassifications, which are caused by the network itself, we can only directly compare against other methods that work on the exact same network. For OOD detection, we evaluate on the Fishyscapes anomaly detection benchmark (Blum et al., 2019). We report the results in table 6.

We find that the aleatoric uncertainty estimated from output-conditional latent densities closely matches that of the softmax entropy. On the task of OOD detection the estimated epistemic uncertainty based on the log probability of the la-

⁸1 component, only 50 iterations, covariance regularisation with 0.1

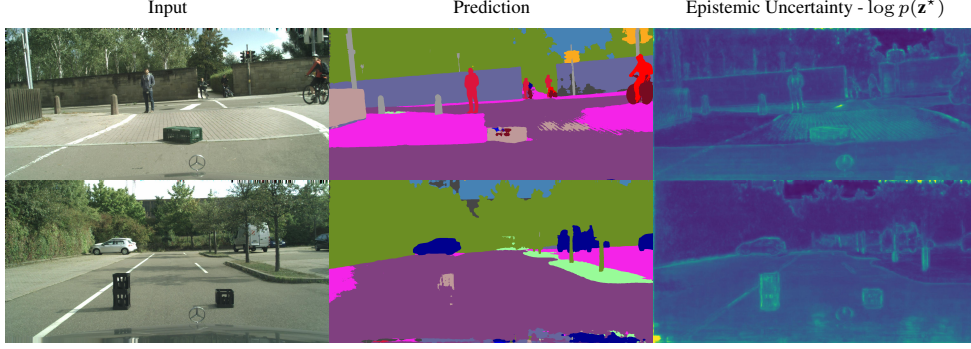


Figure 9: Qualitative examples of the estimated epistemic uncertainty in semantic segmentation. The examples are from the Lost & Found dataset that features images similar to the training set, but with OOD objects (Pinggera et al., 2016). We observe that the estimated epistemic uncertainty is high for anomalous objects and the uncommon street texture in the first row and low where the model correctly generalised.

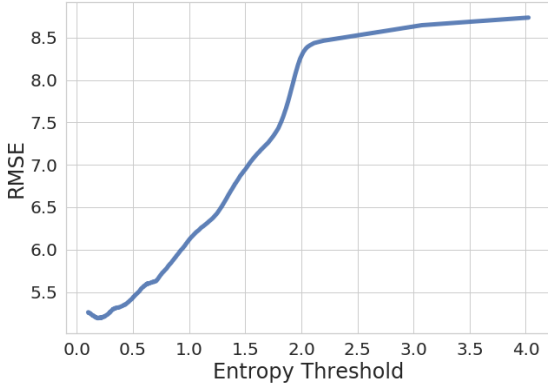


Figure 10: Average RMSE for a monocular depth prediction network (Godard et al., 2017) trained on KITTI (Geiger et al., 2012), accumulated below the given threshold of aleatoric uncertainty $h(\hat{\mathbf{y}}|\mathbf{z}^*)$. As expected of aleatoric uncertainty, it increases monotonically for larger errors.

tent features does however not directly match the behaviour of the MC dropout. In particular, figure 8 shows a very bad performance in low-recall areas and a good performance for the high-recall area, contrary to MC dropout. The qualitative examples in figure 9 show that outlier objects have a lower density, but there are also many other parts of the images with similar values. Further tests over different layers and architectures would be necessary to conclude whether this is an issue of e.g. feature collapse or poorly fitted densities.

Overall, we find that the holistic framework of aleatoric and epistemic uncertainty from latent densities scales with some adaptations to this complex task.

C.2. Depth Regression

As a large scale application in regression, we apply our framework to monocular depth regression on KITTI (Geiger et al., 2012).

C.2.1. EXPERIMENTAL SETUP

We use a pretrained model from (Godard et al., 2017)⁹ trained on KITTI and extract the activations \mathbf{z} from the penultimate layer (output of the penultimate convolution) on the training data and the corresponding depth prediction. We need to estimate two densities: the output-conditional density $p(\mathbf{z}^*|\hat{\mathbf{y}})$ and the density of the predictions $p(\hat{\mathbf{y}})$.

Output-conditional Density of Latent Representations

The conditional density $p(\mathbf{z}^*|\hat{\mathbf{y}})$ is shared across all pixels. To assemble a dataset for training the density model on the training data, we extract the depth predictions pixel-wise and all activations that are in the corresponding receptive field. This leads to a 144 dimensional distribution conditioned on a scalar depth prediction. We randomly extract 12800 samples from the 256×512 dimensional depth maps and shuffle the resulting dataset to ensure that correlations between adjacent pixels have a minimal impact on the i.i.d. assumption when training the CNF. Subsequently, we train a conditional version of the RealNVP (Dinh et al., 2017) on the resulting dataset, where we use a conditioning scheme similar to the one used by (Ardizzone et al., 2019) (see section A.3). Our RealNVP consists of 3 coupling layers where scaling and translation are each computed using a separate multi-layer perceptron with 2 hidden layers of each 100 dimensions. We refer to the original work (Dinh et al., 2017) for a general description of the RealNVP. We train the CNF using Adam optimizer with learning rate $1e - 6$ and weight decay $1e - 5$. We further use a batch size of 128

⁹<https://github.com/mrharicot/monodepth>

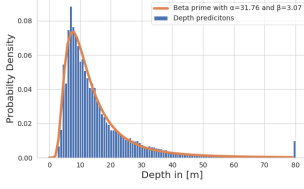


Figure 11: Fitting distribution of depth predictions with univariate beta prime distribution. We visualize normalized histogram of predicted depth values using a pretrained model from (Godard et al., 2017) and a beta prime distribution with parameters $\alpha = 31.76$ and $\beta = 3.07$. The small peak of the predicted depth values at 80m is an artifact of the code on which we base this experiment, since they clip depth predictions at 80m.

and use early stopping with a patience parameter of 20.

Parametric Distribution of Depth Predictions

We empirically found that the distribution of depth predictions on the training data is well represented by a univariate beta prime distribution with $\alpha = 31.76$ and $\beta = 3.07$. Fig. 11 shows the quality of this estimate.

Evaluation of Aleatoric Uncertainty

Evaluating aleatoric uncertainty requires computing $h(\hat{\mathbf{y}}|\mathbf{z}^*) = -\int d\hat{\mathbf{y}} p(\hat{\mathbf{y}}|\mathbf{z}^*) \log(p(\hat{\mathbf{y}}|\mathbf{z}^*))$ where $p(\hat{\mathbf{y}}|\mathbf{z}^*) = \frac{p(\mathbf{z}^*|\hat{\mathbf{y}})p(\hat{\mathbf{y}})}{p(\mathbf{z}^*)}$ and $p(\mathbf{z}^*)$ is obtained by marginalizing over $\hat{\mathbf{y}}$. Since the depth predictions in our experiment are only one-dimensional, computing $h(\hat{\mathbf{y}}|\mathbf{z}^*)$ is achieved in a straightforward manner using numerical integration. We discretize the interval of observed depth values $[3, 80]$ with 154 supporting points with equidistant spacing of 0.5. Note that for higher-dimensional $\hat{\mathbf{y}}$ above integral can be solved more efficiently using importance sampling.

C.2.2. RESULTS

We evaluate the quality of the aleatoric uncertainty. We compute $h(\hat{\mathbf{y}}|\mathbf{z}^*)$ according to eq. 6 and evaluate its correlation with the RMSE in fig. 10. As expected, larger errors have larger aleatoric uncertainty. We conclude that the aleatoric uncertainty estimate scales to large regression networks and allows extraction of aleatoric uncertainty post-training.