

# 1 Experiment

Throughout the following section the results and configuration details of our trained model are described and explained.

## 1.1 Configuration Details

Our model only consists of a single hourglass. The hourglass consists of 4 down- and upsamples, with 1 residual module between each down- and upsample. In each skip-connection a single residual module is used, and in the bottleneck 3 residual modules are used. Newell *et al.* [3] and Olsen [4] experiment with different amount of hourglasses and with different amount of residual modules in each hourglass. They both come to the conclusion, that stacking multiple hourglasses or using hourglasses with multiple residual modules between each down- and upsample increases the performance of the model. However, as the main purpose of this thesis is not to create a model with state-of-the-art results, but instead to create a model that can be interpreted and explored, we have chosen to reduce the size of the model. For the same reasons, we will not be developing and testing various configurations of the architecture. Likewise, the purpose is neither to improve the model developed by Newell *et al.* [3], hence why we will be making the same configuration choices as Newell *et al.* [3] and Olsen [4], which are described in the following.

To prevent the model from overfitting we use batch normalization. Newell *et al.* does not describe where to perform the batch normalization, so we follow Olsen [4] and perform the batch normalization before each convolutional layer in each residual module, after the first convolutional layer of the entire network and before the last convolutional layer of the entire network. For the choice of activation function we use the *ReLU*-function after each batch normalization. Each max pooling and nearest neighbor upsampling uses a kernel size of 2, which halves and doubles the size of the input, respectively. The full network has been visualized in Figure 1.

For the initial values of the weights we initialize each weight by sampling from a *Glorot normal distribution* (also known as a *Xavier normal distribution*), described as

$$\mathcal{N}\left(0, \frac{2}{fan_{in} + fan_{out}}\right)$$

where  $fan_{in}$  is the amount of input connections and  $fan_{out}$  is the amount of output connections to the layer of the weight [1]. By doing so we make all layers have the same activation variance and gradient variance, essentially helping the model to converge [2].

We make use of a mini-batch size of 16 and no data augmentation, since the dataset is already rather large and captures a lot of the variances. To optimize the network we make use of *MSE* as our loss function, *RMSPROP* as our optimizer, as well as use a initial learning rate of  $2.5e-4$ .

After each epoch we find the validation accuracy of the model by computing the *PCK* between the predictions and the ground truth of the validation dataset, as described by Olsen [4] and the source code of the Stacked Hourglass Network [5], where it is called *heatmap accuracy*. The pseudocode of PCK has been visualized in Algorithm 1. The algorithm works by iterating over each annotated ground truth heatmap and the corresponding predicted heatmap. It then finds the Euclidean distance between the maximum activation of a ground truth heatmap and the corresponding predicted heatmap. The distance is then normalized by a constant  $c$  and compared to a threshold radius  $r$ . The ratio of normalized distances that are less than the threshold  $r$  are then computed and returned, yielding the PCK accuracy between the ground



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**Algorithm 1** PCK [4][5]

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**Require:** Ground truth heatmaps  $heatmaps_{gt}$  of keypoints

**Require:** Predicted heatmaps  $heatmaps_{pred}$  of keypoints

**Require:** Threshold radius  $r$

**Require:** Normalization constant  $c$

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1: Let  $n \leftarrow 0$  be the running total of correctly predicted keypoints
2: Let  $N$  be the amount of annotated heatmaps
3: for each annotated ground truth heatmap,  $heatmap_{gt}$ , in  $heatmaps_{gt}$  do
4:   Let  $(x_{gt}, y_{gt})$  be the 2D index of the maximum activation of  $heatmap_{gt}$ 
5:   Let  $(x_{pred}, y_{pred})$  be the 2D index of the maximum activation of the predicted heatmap
   corresponding to  $heatmap_{gt}$ 
6:   Let  $dist$  be the Euclidean distance between  $(x_{gt}, y_{gt})$  and  $(x_{pred}, y_{pred})$ .
7:   Normalize  $dist$ :  $dist \leftarrow \frac{dist}{c}$ 
8:   if  $dist < r$  then
9:      $n \leftarrow n + 1$ 
10: Let  $ratio \leftarrow \frac{n}{N}$  be the ratio of correctly annotated heatmaps
11: return  $ratio$ 
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truth heatmaps and the corresponding predicted heatmaps [4] [5]. The aim is thus to maximize the PCK accuracy. To produce the final PCK accuracy of the model, the PCK accuracy is computed for each image in the validation dataset. The mean PCK accuracy is then used as the PCK accuracy of the model. For the two constants,  $c$  and  $r$ , we let  $c$  be one tenth of the heatmap resolution size (that is,  $\frac{64}{10} = 6.4$ ) and  $r$  be 0.5.

While training the model the PCK accuracy of the model is computed after each epoch, keeping track of the best PCK accuracy. The first time the best PCK accuracy has not improved for 5 continuous epochs, the learning rate is dropped by a factor of 5 permanently, helping the training loss reach a minimum.

## 1.2 Results

In Figure 2 the evolution of the training loss, validation loss and validation PCK accuracy has been visualized. The model were initially set to train for 100 epochs, however, we decided to stop the training early, as the model clearly started to overfit after 30 epochs, as seen by comparing the training and validation loss.

The reduction of the learning rate happened after 21 epochs. By looking at the validation accuracy in Figure 2 we can see, that the accuracy rapidly increases shortly after the reduction of the learning rate, hinting at the effectiveness of dropping the learning rate.

Comparing the training loss, validation loss and the validation accuracy from Table 1 we see, that there is not an overlap between the models yielding the best training loss, validation loss and validation accuracy. As we in section REFERENCE MANGLER want to explore a model that performs decently well, we will be using the model with the highest validation accuracy as our model going forward. Thus, our model is the model from epoch 47, which has a training loss of  $4.19 \cdot 10^{-5}$ , a validation loss of  $5.43 \cdot 10^{-5}$  and a validation accuracy of 0.433.

## 1.3 Training Details

The stacked hourglass was implemented in Python 3.8.2 using PyTorch version 1.7.1 and Cuda version 10.2 on a machine using Windows 10 version 20H2, build 19042. The network was

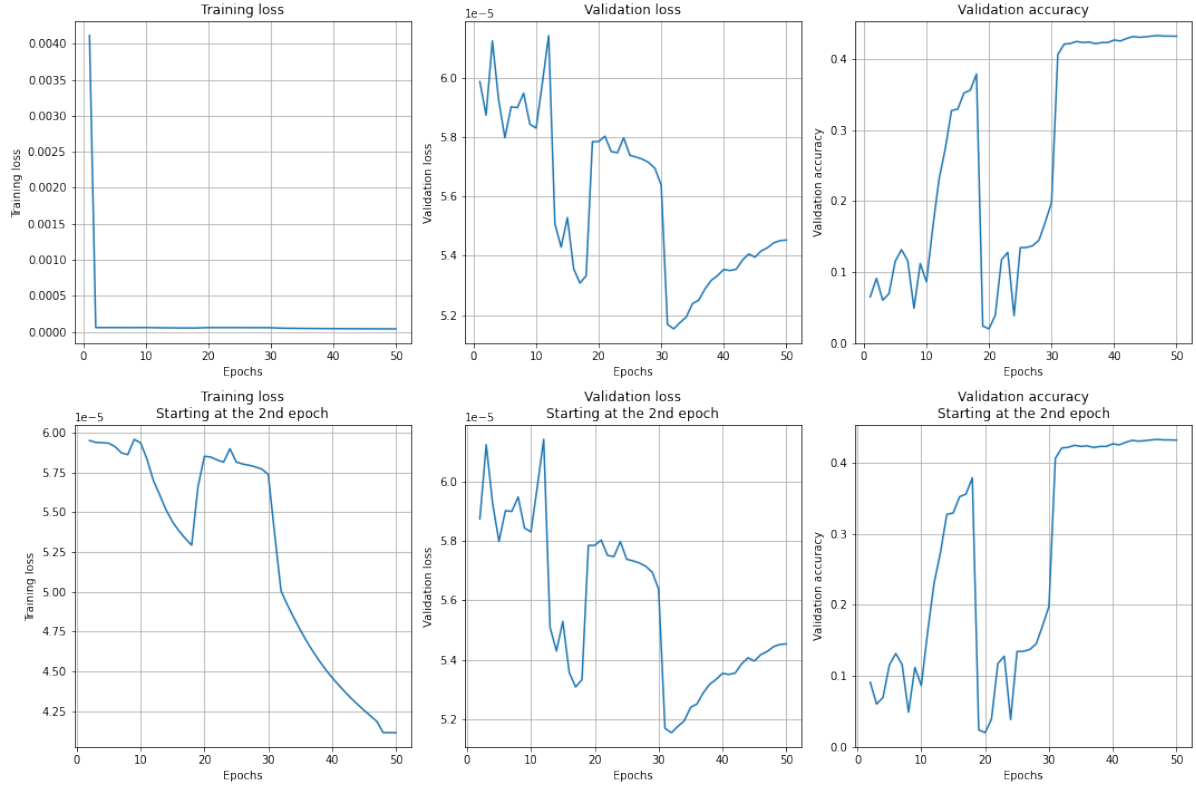


Figure 2: Visualization of the evolution of the training loss, validation loss and validation PCK accuracy of the trained model. Top row shows all of the 50 epochs. Bottom row shows epoch 2 and forward to ease the reading of the training loss

Description	Epoch	Training loss	Validation loss	Validation accuracy
Best training loss	50	$4.11 \cdot 10^{-5}$	$5.45 \cdot 10^{-5}$	0.43
Best validation loss	32	$5.01 \cdot 10^{-5}$	$5.15 \cdot 10^{-5}$	0.42
Best validation accuracy	47	$4.19 \cdot 10^{-5}$	$5.43 \cdot 10^{-5}$	0.433

Table 1: Comparison of the the epochs yielding the best training loss, validation loss and validation accuracy

trained on an 8 GB NVIDIA GeForce GTX 1070 GPU using a Samsung 840 EVO SSD for data storage. Training the network takes about 70 minutes per epoch, totalling to about 58 hours for 50 epochs.