# **Autoencoders for Anomaly Detection**

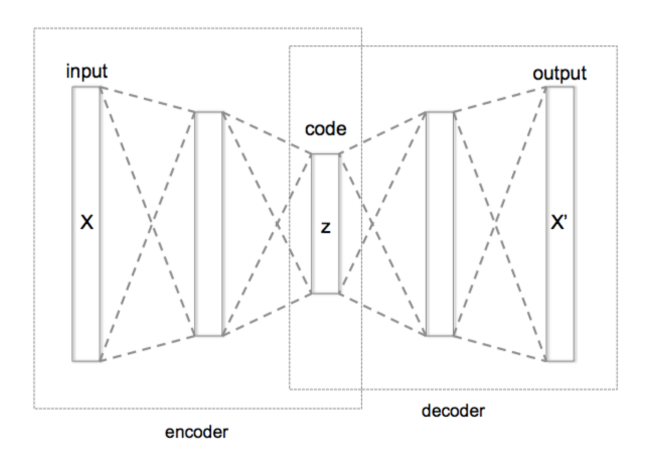




### **Autoencoders**

### An autoencoder is a type of neural network

- The network is designed to reconstruct its input vector
- lacktriangle The input is a tensor  $oldsymbol{x}$  and the output should be similar to the same tensor  $oldsymbol{x}$



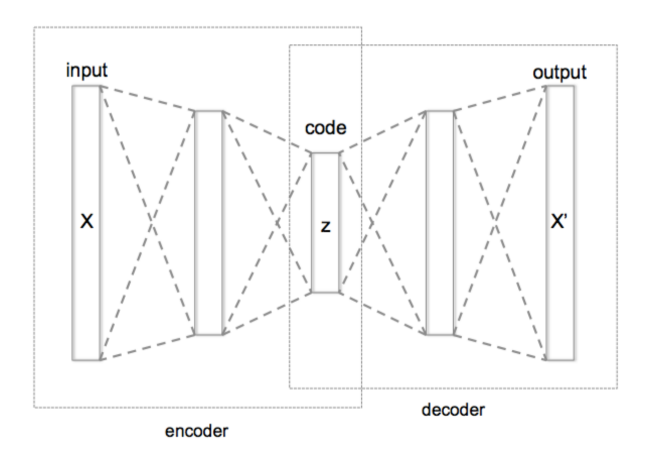




#### **Autoencoders**

#### Autoencoders can be broken down in two halves

- An encoder, i.e.  $e(x, \theta_e)$ , mapping x into a vector of latent variables z
- A decoder, i.e.  $d(z, \theta_d)$ , mapping z into reconstructed input tensor







# **Training an Autoencoder**

### Autoencoders are typically trained for minimum MSE:

$$\underset{\theta_e,\theta_d}{\operatorname{arg \, min}} \|d(e(x_i,\theta_e),\theta_d) - x_i\|_2^2$$

- I.e. d, when applied to the output of e
- ...Should approximately return the input vector itself

A nice tutorial about autoencoders can be found on the Keras blog

### There is a risk that an autoencoder learns a trivial transformation (x' = x)

This is obviously undesired, and it can be avoided by:

- Choosing a small-dimensional latent space (compressing autoencoder)
- By encouraging sparse encodings with an L1 regularizer (sparse autoencoder)





# **Autoencoders for Anomaly Detection**

#### Autoencoders can be used for anomaly detection

...By using the reconstruction error as an anomaly signal, e.g.:

$$||x - d(e(x, \theta_e), \theta_d)||_2^2 \ge \theta$$

#### This approach has some PROs and CONs compared to KDE

- The size of a Neural Network does not depend on the size of the training set
- Neural Networks have good support for high dimensional data
- ...Plus limited overfitting and fast prediction/detection time
- However, input reconstruction can be harder than density estimation

### Let's prepare the data to test the approach





Shall we standardize/normalize the data? And why?





#### **NNs and Standardization**

#### Normalization is important for NNs, due to the use of gradient descent

The performance of SGD depends a lot on its starting point

- DL libraries all come with robust weight initialization procedures
  - ...And robust default parameters for the gradient descent algorithms
- ...But those are designed for data that is:
  - Reasonably close to zero
  - Mostly contained in a  $[-1, 1]^n$  box

#### You can use NNs with non standardize data

...But expect far less reliable results

- In addition, vector output should always be standardized/normalized
- We'll see why in a short while





### **Data Preparation**

#### We'll prepare our data as we did for KDE

First we apply a standardization step:

```
In [3]: tr_end, val_end = 3000, 4500
hpcs = hpc.copy()
tmp = hpcs.iloc[:tr_end]
hpcs[inputs] = (hpcs[inputs] - tmp[inputs].mean()) / tmp[inputs].std()
```

The we separate a training, validation, and test set

```
In [4]: trdata = hpcs.iloc[:tr_end]
valdata = hpcs.iloc[tr_end:val_end]
tsdata = hpcs.iloc[val_end:]
```





### **Building an Autoencoder**

#### The we can build an autoencoder (we'll use tensorflow 2.0 and keras)

First, we build the model using (e.g.) the functional API

```
In [5]: import keras
from keras import layers, callbacks

input_shape = (len(inputs), )
ae_x = keras.Input(shape=input_shape, dtype='float32')
ae_z = layers.Dense(64, activation='relu')(ae_x)
ae_y = layers.Dense(len(inputs), activation='linear')(ae_z)
ae = keras.Model(ae_x, ae_y)
```

- Input builds the entry point for the input data
- Dense builds a fully connected layer
- "Calling" layer A with parameter B attaches B to A
- Model builds a model object with the specified input and output





#### **Autoencoders in Keras**

#### Then we can prepare our model for training

In keras terms, we compile it

```
In [6]: ae.compile(optimizer='Adam', loss='mse')
```

We are using the Adam optimizer (a variant of Stochastic Gradient Descent)

#### Then we can start training:

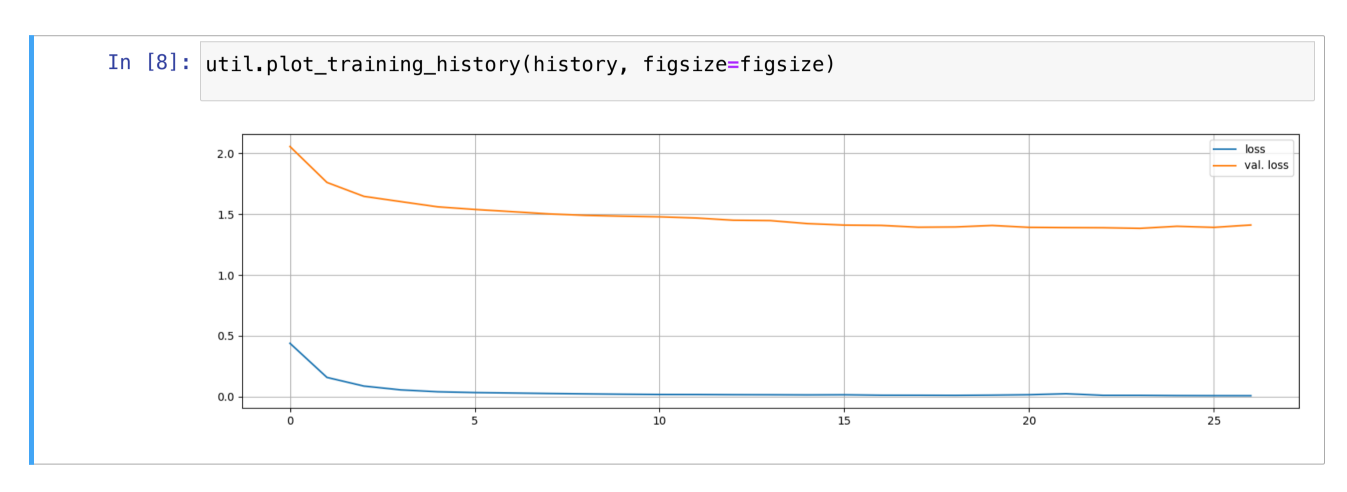
- We are using a callback to stop training early
- ...If no improvement on the validation set is observed for 3 epochs





### **Autoencoders in Keras**

### Let's have a look at the loss evolution over different epochs







### **Autoencoders in Keras**

#### Finally, we can obtain the predictions

In [9]: preds = pd.DataFrame(index=hpcs.index, columns=inputs, data=ae.predict(hpcs[inputs], verbose
 preds.head()

Out[9]:
 ambient\_temp cmbw\_p0\_0 cmbw\_p0\_1 cmbw\_p0\_10 cmbw\_p0\_11 cmbw\_p0\_12 cmbw\_p0\_13 cmbw\_p0\_14 cmbw\_p0\_2 cmbw\_p0

	ambient_temp	cmbw_p0_0	cmbw_p0_1	cmbw_p0_10	cmbw_p0_11	cmbw_p0_12	cmbw_p0_13	cmbw_p0_14	cmbw_p0_2	cmbw_p(
0	-1.414166	-1.399190	0.241891	1.693468	2.040897	1.790998	1.716491	1.900154	-0.967087	-1.04827
1	-0.802001	-0.534030	0.324981	2.182271	2.325772	2.268217	2.234467	2.220423	0.365094	-0.70460
2	-1.004940	-0.994740	-0.386334	2.368342	2.337509	2.426816	2.356327	2.266239	0.613221	0.709304
3	-0.777538	-0.840914	-0.509004	2.256818	2.314422	2.294407	2.215047	2.178997	0.818741	0.677289
4	-0.986344	-0.957748	-0.516556	2.290415	2.334587	2.300023	2.264766	2.237829	0.806280	0.861571

5 rows × 159 columns

■ These are the reconstructed values for all the input features





### **Alarm Signal**

### We can finally obtain our alarm signal, i.e. the sum of squared errors

```
In [10]: labels = pd.Series(index=hpcs.index, data=(hpcs['anomaly'] != 0), dtype=int)
          sse = np.sum(np.square(preds - hpcs[inputs]), axis=1)
          signal_ae = pd.Series(index=hpcs.index, data=sse)
          util.plot signal(signal ae, labels, figsize=figsize)
           1.4
           1.2
           1.0
           0.8
           0.6
           0.4
           0.2
           0.0
                              1000
                                           2000
                                                        3000
                                                                      4000
                                                                                   5000
                                                                                                 6000
```





#### Let's try to understand what we have just done

When we train an autoencoder (renamed here as h), we solve:

$$\underset{\theta}{\arg\min} \|h(x,\theta) - x\|_2^2$$

By expanding the L2 norm, we get:

$$\underset{\theta}{\operatorname{arg\,min}} \sum_{i=1}^{m} \sum_{j=1}^{n} \left( h_{j}(x_{i}, \theta) - x_{i,j} \right)^{2}$$

By introducing a  $\log$  and  $\exp$  transformation we obtain:

$$\underset{\theta}{\operatorname{arg \, min \, log \, exp}} \left( \sum_{i=1}^{m} \sum_{j=1}^{n} \left( h_{j}(x_{i}, \theta) - x_{i,j} \right)^{2} \right)$$





#### Then, from the last step:

$$\underset{\theta}{\operatorname{arg \, min \, log \, exp}} \left( \sum_{i=1}^{m} \sum_{j=1}^{n} \left( h_{j}(x_{i}, \theta) - x_{i,j} \right)^{2} \right)$$

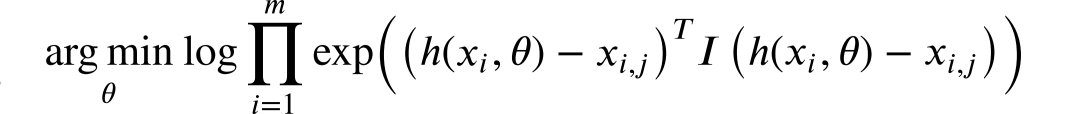
We rewriting the outer sum using properties of exponentials:

$$\arg\min_{\theta} \log \prod_{i=1}^{m} \exp \left( \sum_{j=1}^{n} \left( h_{j}(x_{i}, \theta) - x_{i,j} \right)^{2} \right)$$

Then we rewrite the inner sum in matrix form:







### **Starting from the last step:**

$$\underset{\theta}{\operatorname{arg \, min}} \log \prod_{i=1}^{m} \exp \left( \left( h(x_{i}, \theta) - x_{i,j} \right)^{T} I \left( h(x_{i}, \theta) - x_{i,j} \right) \right)$$

We make a few adjustment that do not change the optimal solution:

- We negate the argument of exp and swap the arg min for a arg max
- We multiply exponent by  $1/2\sigma$  (for some constant  $\sigma$ )
- We multiply the exponential by  $1/\sqrt{2\pi}\sigma$

$$\arg\max_{\theta} \log \prod_{i=1}^{m} \frac{1}{\sqrt{2\pi\sigma}} \exp\left(-\frac{1}{2} \left(h(x_i, \theta) - x_{i,j}\right)^T (\sigma I) \left(h(x_i, \theta) - x_{i,j}\right)\right)$$





#### Let's look at our last formulation:

$$\arg\max_{\theta} \log \prod_{i=1}^{m} \frac{1}{\sqrt{2\pi}} \exp\left(-\frac{1}{2} \left(h(x_i, \theta) - x_{i,j}\right)^T (\sigma I) \left(h(x_i, \theta) - x_{i,j}\right)\right)$$

The term inside the product is the PDF of a <u>multivariate normal distribution</u>

$$\arg \max_{\theta} \log \prod_{i=1}^{m} f(x_i, h(x_i), \sigma I)$$

- In particular a distribution centered on  $h(x_i)$
- ...With independent Normal components
- ...All having uniform variance





### Let's discuss some implications

When we use a MSE loss, we are training for maximum likelihood

- ...Just like density estimators!
- This is actually true for many ML approaches





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The output of a (MSE trained) regressor has a probabilistic interpretation

- Specifically, the output is the mean of a conditional distribution
- The distribution represents the variability of the target
- ...Once the effect of the input is taken into account
- Another way to think of it: noise around the prediction





### Let's discuss some implications

We are implicitly assuming that the noise is normally distributed

- This true in many cases, but not always
- E.g., sometimes large values are under-represented
- ....Leading to log-normal distributions
- In this cases, applying a log transformation to the output can be very effective





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- In this cases, applying a log transformation to the output can be very effective

We are also assuming that the all output components have the same variance

This is another (very) good reason to standardize the output





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- This might be true even if the output components themselves are correlated
- ...But still it is not true in all cases





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All these implicit assumption can make the problem harder

This is why error reconstruction can be harder than density estimation





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This is why error reconstruction can be harder than density estimation

Finally, our alarm signal can be interpreted as a density:

- To see why, just apply the transformation to  $\|x d(e(x, \theta_e), \theta_d)\|_2^2$
- This fact explains why the signal is similar to the KDE one





### **Threshold Optimization**

### The threshold can be optimized as usual

```
In [11]: c_alarm, c_missed, tolerance = 1, 5, 12
         cmodel = util.HPCMetrics(c alarm, c missed, tolerance)
         th range = np.linspace(1e4, 2e5, 200)
         th ae, val cost ae = util.opt threshold(signal ae[tr end:val end], hpcs['anomaly'][tr end:val end]
         print(f'Best threshold: {th ae:.3f}')
         tr cost ae = cmodel.cost(signal ae[:tr end], hpcs['anomaly'][:tr end], th ae)
         print(f'Cost on the training set: {tr_cost_ae}')
         print(f'Cost on the validation set: {val cost ae}')
         ts cost ae = cmodel.cost(signal ae[val end:], hpcs['anomaly'][val end:], th ae)
         print(f'Cost on the test set: {ts cost ae}')
         Best threshold: 119798.995
         Cost on the training set: 0
         Cost on the validation set: 263
         Cost on the test set: 265
```

The performance is similar to KDE (not surprisingly)





### **Mutiple Signal Analysis**

#### But autoencoders do more than just anomaly detection!

- Instead of having a single signal we have many
- So we can look at the individual reconstruction errors

```
In [12]: se = np.square(preds - hpcs[inputs])
          signals_ae = pd.DataFrame(index=hpcs.index, columns=inputs, data=se)
          util.plot_dataframe(signals_ae, labels, vmin=-5e4, vmax=5e4, figsize=figsize)
            20
            40
            80
           100
           120
           140
                           1000
                                         2000
                                                                                                    6000
                                                        3000
                                                                       4000
                                                                                     5000
```





### **Mutiple Signal Analysis**

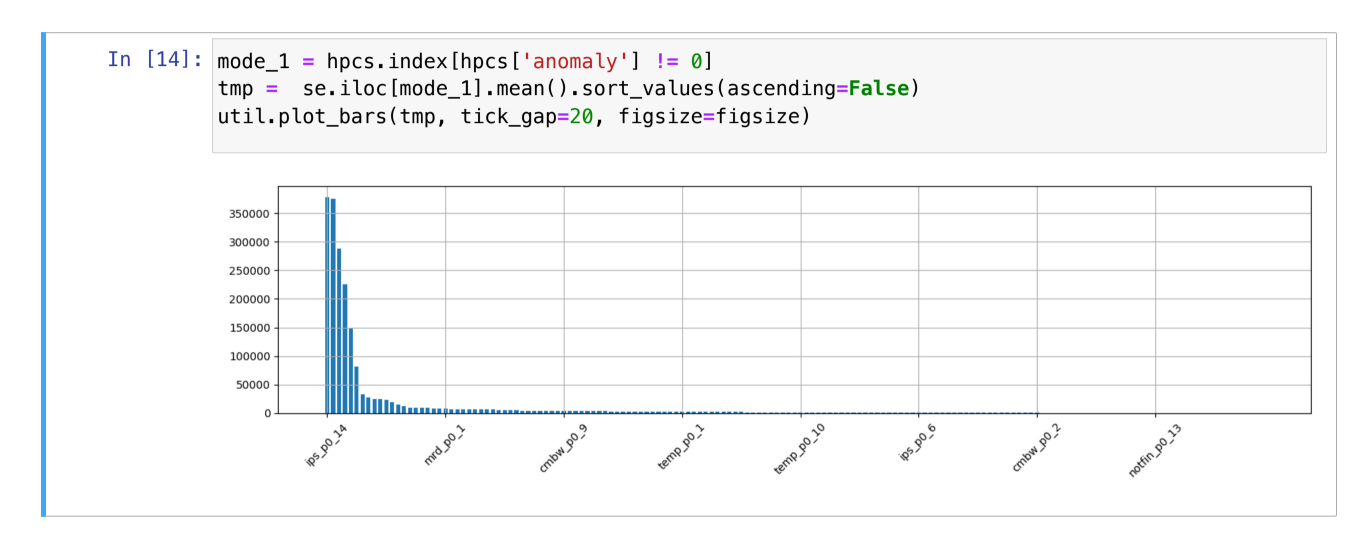
#### Reconstruction errors are often concentrated on a few signals

- These correspond to the properties of the input vector that were harder to reconstruct
- ...And often they are useful clues about the nature of the anomaly

```
In [13]: se = np.square(preds - hpcs[inputs])
          signals_ae = pd.DataFrame(index=hpcs.index, columns=inputs, data=se)
          util.plot dataframe(signals ae, labels, vmin=-5e4, vmax=5e4, figsize=figsize)
           20
            40
            60
           80
           100
           120
           140
                           1000
                                          2000
                                                        3000
                                                                       4000
                                                                                      5000
                                                                                                     6000
```

# Multiple Signal Analysis

Here are the average errors for all anomalies (sorted by decreasing value)



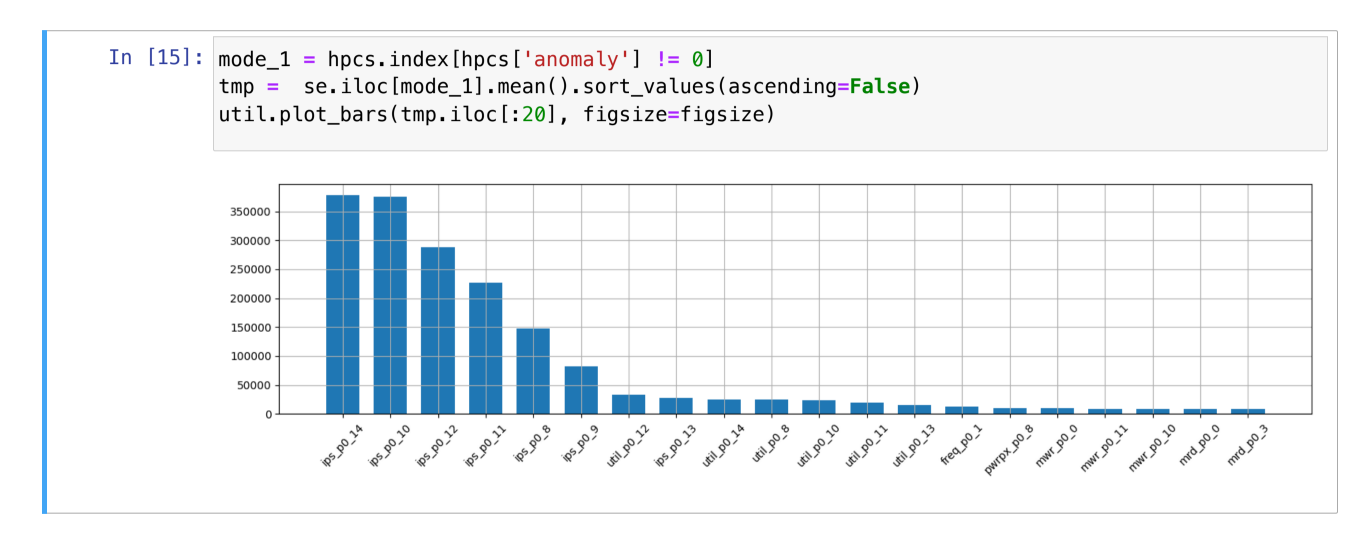
Errors are concentrated on 10-20 features





### Multiple Signal Analysis

### These are the 20 largest average errors for all anomalies



- The largest errors are on "ips", then on "util" (utilization)
- This kind of information can be very valuable for a domain expert!