

## *Outline:*

- Motivation.
- Bayesian Learning.
- Neural Networks.
- Combination.

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## Science & Environment

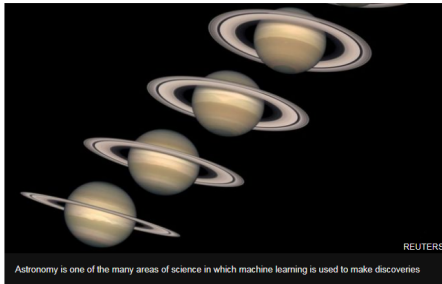
### AAAS: Machine learning 'causing science crisis'

By Pallab Ghosh  
 Science correspondent, BBC News, Washington

© 16 February 2019

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American Association for the Advancement of Science Meeting



Machine-learning techniques used by thousands of scientists to analyse data are producing results that are misleading and often completely wrong.

## Keith Baggerly: Forensic Bioinformatics

**Using the NCI60 to Predict Sensitivity**

Genomic signatures to guide the use of chemotherapeutics

Jeff Potti<sup>1</sup>, Holly K. Dressman<sup>2,3</sup>, Andrew Rhee<sup>4,5</sup>, Richard E. Black<sup>6,7</sup>, Gina Chen<sup>1</sup>, Robert Saxe<sup>8</sup>, David Casper<sup>9</sup>, Hope Conant<sup>10</sup>, Michael J. Keller<sup>1</sup>, Rebecca Peterson<sup>1</sup>, David Hargreaves<sup>1</sup>, Jeffrey Marko<sup>1</sup>, Andrew Beckman<sup>11,12</sup>, Geoffrey A. Gilchrist<sup>13</sup>, Philip Haber<sup>14,15</sup>, Jonathan Lammert<sup>16</sup>, Joseph B. Sparano<sup>17</sup>

Potti et al (2006), Nature Medicine, 12:1294-1300.

The main conclusion is that we can use microarray data from cell lines (the NCI60) to define drug response "signatures", which can be used to predict whether patients will respond.

They provide examples using 7 commonly used agents.

This got people at MDA very excited.

The Brad Efron Honorary Symposium on LARGE-SCALE INFERENCE - Keith Baggerly

2006

**A Catalyzing Event: July 16, 2010**

**THE CANCER LETTER**

PO Box 9905 Washington DC 20016 Telephone 202-362-1809

**Prominent Duke Scientist Claimed Prizes He Didn't Win, Including Rhodes Scholarship**

*By Paul Goldberg*

Jul 19/20: Letter to Varmus; Duke resumes trials.  
Oct 22/9: First call for paper retraction.  
Nov 9: Duke terminates trials.  
Nov 19: call for Nat Med retraction, Potti resigns

The Brad Efron Honorary Symposium on LARGE-SCALE INFERENCE - Keith Baggerly

2010

Playing with people's lives.

# Confusion matrix

- N: number of negative samples, P: number of positive samples.
- *True negatives* TN: number of negative samples correctly classified.
- *False positives* FP: number of negative samples misclassified.
- *True positives* TP: number of positive samples correctly classified.
- *False negatives* FN: number of positive samples misclassified.
- *Confusion table*:

	N	P
classified negative	TN	FN
classified positive	FP	TP

# Sensitivity and Specificity

- *Sensitivity*, aka *true positive rate*, *recall* and *probability of detection*: fraction of positive samples correctly classified:  $\text{TPR} = \frac{\text{TP}}{P}$ .
- *Specificity* or *true negative rate*: fraction of negative samples correctly identified:  $\text{TN} = \frac{\text{TN}}{N}$ .
- *False negative rate* or *miss rate*:  $\text{FNR} = \frac{\text{FN}}{P} = 1 - \text{TPR}$ .
- *False positive rate* or *fall-out*:  $\text{FPR} = \frac{\text{FP}}{N} = 1 - \text{TN}$ .

# Likelihood Ratios

- *Likelihood ratio for positive results*: how much more likely is positive classification in positive samples compared to in negative samples:

$$LR+ = \frac{TPR}{FPR}.$$

- *Likelihood ratio for negative results*: how much more likely is negative classification in positive samples compared to in negative samples:

$$LR- = \frac{FNR}{TNR}.$$

- All so far independent of prevalence.

- A perfect classifier would be 100% sensitive (all positives are correctly identified) and 100% specific (no negatives are incorrectly classified).
- $LR+ \rightarrow \infty$  and  $LR- \rightarrow 0$ .
- $LR+ > 10$  and  $LR- < 0.1$  make a useful classifier according to Jaeschke R, Guyatt G, Lijmer J, '*Diagnostic tests*' in Guyatt G, Rennie D, eds. '*Users guides to the medical literature*' Chicago: AMA Press, (2002).

## Opinion

OP-ED CONTRIBUTOR

# When an Algorithm Helps Send You to Prison

By Ellora Thadaneey Israni

Oct. 26, 2017



In 2013, police officers in Wisconsin arrested a man driving a car that had been used in a recent shooting. The man, Eric Loomis, pleaded guilty to attempting to flee an officer, and no contest to operating a vehicle without the owner's consent. Neither of his crimes mandates prison time.

At Mr. Loomis's sentencing, the judge cited, among other factors, Mr. Loomis's high risk of recidivism as predicted by a computer program called COMPAS, a risk assessment algorithm used by the state of Wisconsin. The judge denied probation and prescribed an 11-year sentence: six years in prison, plus five years of extended supervision.

No one knows exactly how COMPAS works; its manufacturer refuses to disclose the proprietary algorithm. We only know the final risk assessment score it spits out, which judges may consider at sentencing.

1,389 views | Jan 24, 2018, 11:47am

## Management AI: Bias, Criminal Recidivism, And The Promise Of Machine Learning

David A. Teich Contributor

Tirias Research Contributor Group

B2B technology analyst and consultant

- f** Criminal recidivism is when a released criminal goes back to crime. From charging crimes through probation, the criminal justice system is constantly looking for ways to better predict which criminals are more likely to remain legal on release and who is a risk of recidivism.
- in** Bias can create inaccuracies through weighing variables incorrectly, and machine learning might provide a way of limiting bias and improving recidivism predictions.



Shutterstock

A recent study by Julia Dressel and Hany Farid, [published in Science Advances](#), points to the limitations of deterministic algorithms with fixed parameters for the task of such predictions. The study analyzes the Correctional Offender Management Profiling for Alternative Sanctions (COMPAS) software, a package used by court systems to predict the likelihood of recidivism in



Larson J, Mattu S, Kirchner L and Angwin J (2016) at Pro Publica Inc. obtained data on the re-offending risks as returned by the COMPAS algorithm and the actual occurrences of re-offending within two years after release.

	N	P	
low risk	2681	1216	3897
high risk	1282	2035	3317
	3963	3251	7214

sensitivity:  $\text{TPR} = 0.63$      $\text{FNR} = 0.37$      $\text{LR+} = 1.97$

specificity:  $\text{TNR} = 0.68$      $\text{FPR} = 0.32$      $\text{LR-} = 0.54$

	Black		
	N	P	
low risk	990	532	1522
high risk	805	1369	2174
	1795	1901	3696

$$\text{TPR} = 0.72$$

$$\text{TNR} = 0.55 \quad \text{FNR} = 0.28$$

$$\text{LR+} = 1.60 \quad \text{FPR} = 0.45$$

$$\text{LR-} = 0.51$$

	White		
	N	P	
low risk	1139	461	1600
high risk	349	505	854
	1488	966	2454

$$\text{TPR} = 0.52$$

$$\text{TNR} = 0.77 \quad \text{FNR} = 0.48$$

$$\text{LR+} = 2.26 \quad \text{FPR} = 0.23$$

$$\text{LR-} = 0.62$$

## *Challenges:*

- Modeling data, keeping models simple while explaining the data adequately.
- New data arriving.
- Confidence in model predictions.
- Choice of model space.

## *The data prediction problem*

- We make the assumption that the data are a result of an underlying process which we do not know.
- Given measurements  $t_1, \dots, t_D$ , each measurement depends on parameters we know  $\mathbf{x}_1, \dots, \mathbf{x}_D$ .
- $D$  is the dimension of the data space.
- These are quantities which can be measured with more or less effort.

## *Unknowns*

- The measurements also depend on parameters we do not know.
- A real world application depends on factors which cannot be measured (or these measurements would be disproportionately difficult).
- For example the physics of waves are well understood. However, they depend on the medium the wave travels in, the material and its properties. These are the unknown parameters of the process.

## Dictionaries

- If we had a set of candidate functions  $d_1(\mathbf{x}), \dots, d_M(\mathbf{x})$ , which all are solutions to the process for different parameters, we could try which fits the measurements and thus infer the underlying structure.
- We say the functions  $d_1(\mathbf{x}), \dots, d_M(\mathbf{x})$  form a dictionary and assume

$$f(\mathbf{x}) = \sum_{m=1}^M c_m d_m(\mathbf{x}),$$

where  $c_1, \dots, c_M$  are coefficients and these need to be determined.

- The basis functions of the dictionary are the **building blocks** which build a model for the data.
- $M$  is the dimension of the model space.

## Noise

- The relationship to the measurements is

$$t_i = f(\mathbf{x}_i) + \epsilon_i.$$

- $\epsilon_i$  is noise intrinsic to the measurement process and assumed to be independent and identically, normally distributed,  $\mathcal{N}(0, \sigma^2)$ .

## Mathematical model

$$t_i = f(\mathbf{x}_i) + \epsilon_i = \sum_{m=1}^M c_m d_m(\mathbf{x}_i) + \epsilon_i,$$

- Let  $\mathbf{D}$  be the matrix with entries  $\mathbf{D}_{i,m} = d_m(\mathbf{x}_i)$  and let  $\mathbf{t}^T = (t_1, \dots, t_D)$ ,  $\mathbf{c}^T = (c_1, \dots, c_M)$  and  $\boldsymbol{\epsilon}^T = (\epsilon_1, \dots, \epsilon_D)$ , then

$$\mathbf{t} = \mathbf{D}\mathbf{c} + \boldsymbol{\epsilon}.$$

- $\mathbf{D}$  is an  $D \times M$  matrix. However,  $D$  and  $M$  are not static.  $D$  varies with the number of measurements of the same process, while  $M$  varies with the dictionary of basis functions.



## *Sparse Bayesian Learning*

- Central idea is that the coefficients  $\mathbf{c}$  follow a distribution.
- Each coefficient  $c_m$  is a priori normally distributed with mean zero and variance  $\alpha_m^{-1}$ .
- $\alpha_m$  is the **precision** of the distribution.
- If the precision is very large, the distribution becomes peaked at its mean and we have more confidence in the value of  $c_m$  than if it is small and the width of the distribution large.

- Multivariate prior distribution:

$$p(\mathbf{c}|\boldsymbol{\alpha}) = (2\pi)^{-M/2} \sqrt{|A|} \exp(\mathbf{c}^T A \mathbf{c}),$$

where  $A$  is a diagonal matrix with entries  $A_{mm} = \alpha_m$

- Multivariate posterior distribution is normal with mean  $\boldsymbol{\mu}$  and variance  $\boldsymbol{\Sigma}$  given by

$$\boldsymbol{\Sigma} = (A + \sigma^{-2} D^T D)^{-1} \quad \boldsymbol{\mu} = \sigma^{-2} \boldsymbol{\Sigma} D^T \mathbf{t}.$$

- Since  $\mathbf{t} = \mathbf{D}\mathbf{c} + \boldsymbol{\epsilon}$ , the data is viewed as being drawn from a normal distribution with mean  $\mathbf{D}\boldsymbol{\mu}$  and variance  $\sigma^2 \mathbf{I} + \mathbf{D}\boldsymbol{\Sigma}\mathbf{D}$ .

- The **marginal likelihood** is the probability of the data given the model specified by  $\mathbf{D}$ ,  $\alpha$  and  $\sigma^2$  after integrating out the coefficients  $\mathbf{c}$ .

$$\mathcal{L}(\mathbf{t}|\mathbf{D}, \alpha, \sigma^2) = (2\pi)^{-N/2} |\sigma^2 \mathbf{I} + \sum_{m=1}^M \frac{1}{\alpha_m} \mathbf{d}_m \mathbf{d}_m^T|^{-1/2} \exp \left( -\frac{1}{2} \mathbf{t}^T (\sigma^2 \mathbf{I} + \sum_{m=1}^M \frac{1}{\alpha_m} \mathbf{d}_m \mathbf{d}_m^T)^{-1} \mathbf{t} \right).$$

- We maximize the likelihood.

# Maximizing the Marginal Likelihood

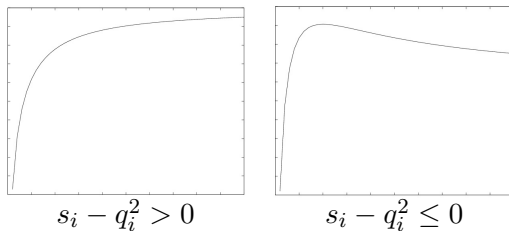
- If the derivative is positive, we move towards a maximum,  
negative, we move away from the maximum,  
zero, we are at the maximum.
- Defining

$$\mathbf{C} = \sigma^2 \mathbf{I} + \sum_{m=1}^M \frac{1}{\alpha_m} \mathbf{d}_m \mathbf{d}_m^T, \quad \mathbf{C}_{-i} = \mathbf{C} - \frac{1}{\alpha_i} \mathbf{d}_i \mathbf{d}_i^T,$$
$$s_i = \mathbf{d}_i^T \mathbf{C}_{-i}^{-1} \mathbf{d}_i, \quad q_i = \mathbf{d}_i^T \mathbf{C}_{-i}^{-1} \mathbf{t}.$$

- The derivative with respect to  $\alpha_i$  of the logarithm of the marginal likelihood is

$$\underbrace{\frac{1}{2}(\alpha_i + s_i)^{-2}}_{>0} (s_i - q_i^2 + \underbrace{\frac{s_i^2}{\alpha_i}}_{\geq 0}).$$

# Sparse Bayesian Learning



In practice many  $\alpha_m$  become infinite during maximization, meaning that the posterior distribution of the corresponding  $c_m$  is **infinitely peaked at 0** and the corresponding building block can be removed from the model.

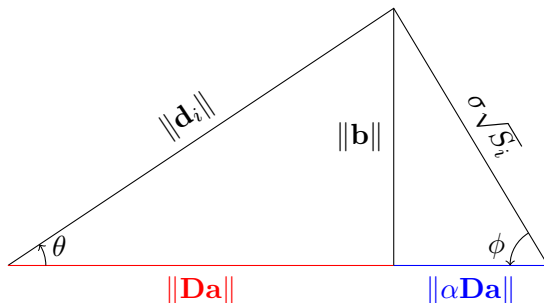
## *Sparsity and Quality Factor*

- Let  $S_i = \mathbf{d}_i^T \mathbf{C}^{-1} \mathbf{d}_i$ , then  $s_i = \frac{\alpha_i S_i}{\alpha_i - S_i}$ .
- Let  $Q_i = \mathbf{d}_i^T \mathbf{C}^{-1} \mathbf{t}$ , then  $q_i = \frac{\alpha_i Q_i}{\alpha_i - S_i}$ .
- It can be shown that

$$\mathbf{C}^{-1} \mathbf{t} = \sigma^{-2} (\mathbf{t} - \mathbf{D} \boldsymbol{\mu}).$$

- $Q_i$  quantifies how well aligned the building block is with this error.
- If it is orthogonal, then  $\mathbf{d}_i$  will not help in removing this error.

# Sparsity and Quality Factor



- By the law of sines  $\frac{\sin \theta}{\sin \phi} = \frac{\sigma \sqrt{S_i}}{\|d_i\|} = \sigma \sqrt{\frac{d_i^T C^{-1} d_i}{\|d_i\| \|d_i\|}}$ .
- $S_i$  measures, how different the building block is to the others.

## *Model Generation*

- **Initializing** the model with a **single building block**.
- All other hyper-parameters are notionally infinity.
- The basis function  $d_m$ , where setting its hyper-parameter  $\alpha_m$  to its optimal value (given the current model) gives the largest increase in the marginal likelihood, is found and the model updated accordingly.
- Note that the optimal value of  $\alpha_m$  can be finite or infinite.
  - **Addition**: If  $d_m$  is not in the model and the optimal  $\alpha_m$  is finite.
  - **Deletion**: If  $d_m$  is in the model and the optimal  $\alpha_m$  is infinite.
  - **Re-estimation**: If  $d_m$  is in the model and the optimal  $\alpha_m$  is finite.
- This addresses the **first challenge**.



## Predictions

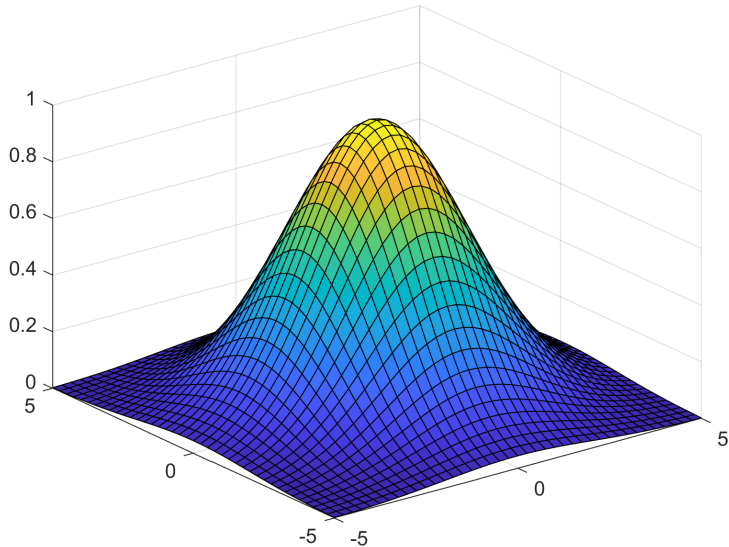
- For a new  $\mathbf{x}_*$ , the predictions  $t_* = \mathbf{c}^T \mathbf{d}_*$ , where  $\mathbf{d}_*^T = (d_1(\mathbf{x}_*), \dots, d_M(\mathbf{x}_*))$ , follow a univariate normal distribution with

$$\text{mean} \quad m_* = \boldsymbol{\mu}^T \mathbf{d}_*,$$

$$\text{variance} \quad \sigma_*^2 = \mathbf{d}_*^T \boldsymbol{\Sigma} \mathbf{d}_*,$$

where  $\boldsymbol{\mu}$  and  $\boldsymbol{\Sigma}$  are the mean and variance of the posterior distribution of the coefficients.

# Predictions



- A new measurement  $(\mathbf{x}_*, t_*)$  means adding a row to  $D$

$$D_* = \begin{pmatrix} D \\ \mathbf{d}_*^T \end{pmatrix}.$$

- The logarithm of the marginal likelihood  $\log \mathcal{L}(\mathbf{t}_* | \boldsymbol{\alpha}, \sigma^2)$  is  $\log \mathcal{L}(\mathbf{t} | \boldsymbol{\alpha}, \sigma^2) + \Delta \mathcal{L}$ , where

$$\Delta \mathcal{L} = \log \frac{1}{\sqrt{2\pi}\sigma_*} \exp \left( -\frac{(m_* - t_*)^2}{2\sigma_*^2} \right).$$

- Thus the change is the **logarithm of the likelihood of the new measurement  $t_*$  at  $\mathbf{x}_*$  given the model.**

- Since  $\sigma_* \geq \sigma$ , the change lies between  $-\infty$  and  $\log \frac{1}{\sqrt{2\pi}\sigma}$ .
- If it is **positive**, the new measurement affirms the model.
- If it is **negative**, the model is not good enough and should be updated.
- This can be done following the previous method.
- This addresses the **second challenge**.

## *Confidence in predictions*

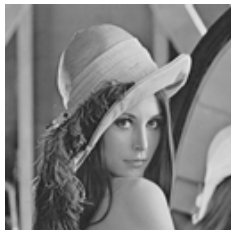
- Note, that the predictive distribution depends on the choice of basis functions. In particular, if  $\mathbf{d}_* = 0$ , then the mean  $m_*$  is zero, while the variance  $\sigma_*^2 = \sigma^2$ . The model fails completely.
- Let  $\mathcal{S}$  be a subset of the samples. This could be all samples or a suitable set of neighbours of  $\mathbf{x}_*$ .
- We **estimate the probability distribution of  $t_*$  to be normal** with mean and variance

$$\begin{aligned}\bar{m} &= \text{mean}_{\mathbf{x}_i \in \mathcal{S}} \{t_i\}, \\ \bar{\sigma} &= \text{var}_{\mathbf{x}_i \in \mathcal{S}} \{t_i\}.\end{aligned}$$

- The expected change in the logarithm of the marginal likelihood is estimated as

$$E[\Delta\mathcal{L}] = \log \frac{1}{\sqrt{2\pi}\sigma_*} - \frac{\bar{\sigma}^2 + (\bar{m} - m_*)^2}{2\sigma_*^2}.$$

- If the predictive probability distribution agrees well with the estimated distribution, the change is positive and we have confidence in our predictions.
- If it does not match well, the expected change is negative, indicating that here more data should be gathered.
- This addresses the **third challenge**.



Original

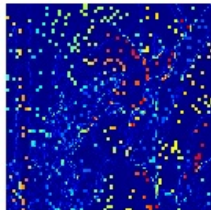


Decimation by 55%

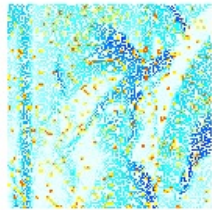
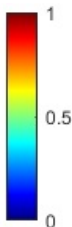
# Experiments



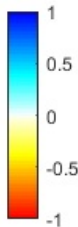
FSIM = 0.74



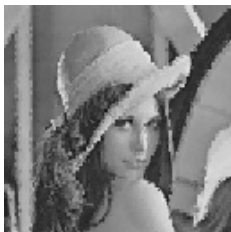
Scaled absolute difference



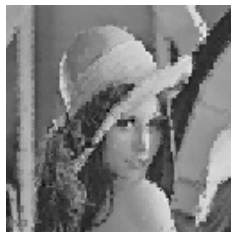
Scaled  $E[\Delta\mathcal{L}]$







5% more samples  
as informed by  $E[\Delta\mathcal{L}]$   
FSIM = 0.93



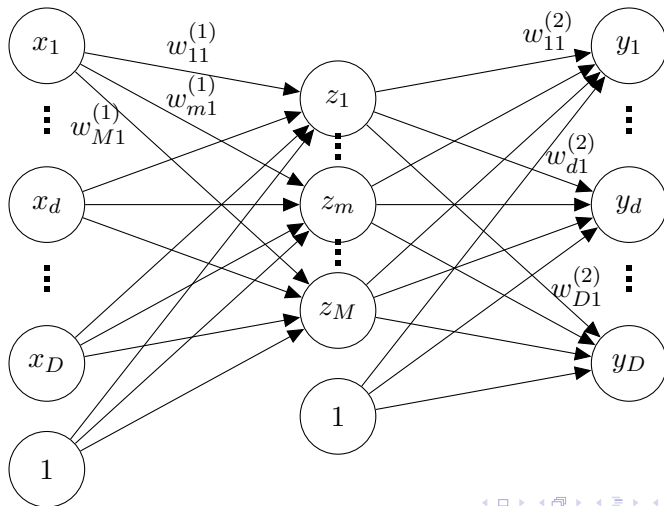
Improvements with different  
basis functions  
FSIM = 0.91

# MNIST

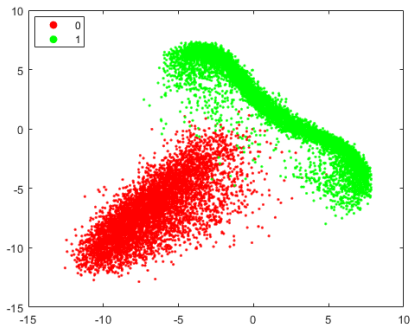
60,000 images of handwritten digits of size  $28 \times 28 = 784$



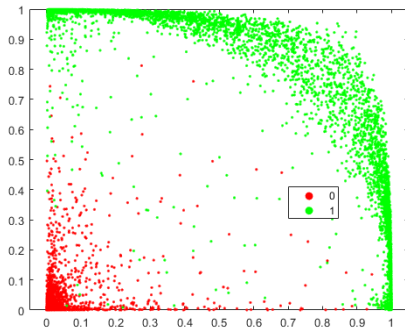
# Autoencoder



# 2 Hidden Neurons, 2 Digits

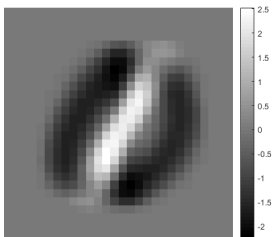


Spatial separation of activations.

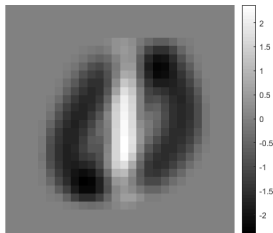


Spatial separation of latent variables.

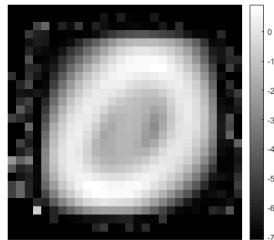
# 2 Hidden Neurons, 2 Digits



First basis.



Second basis.

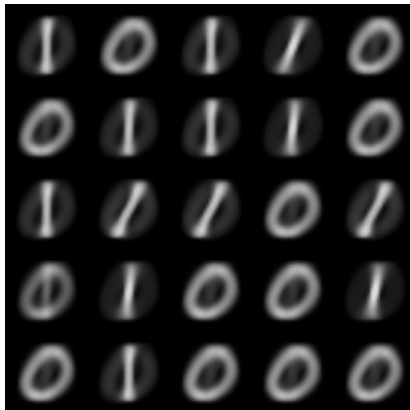


Bias.

## 2 Hidden Neurons, 2 Digits

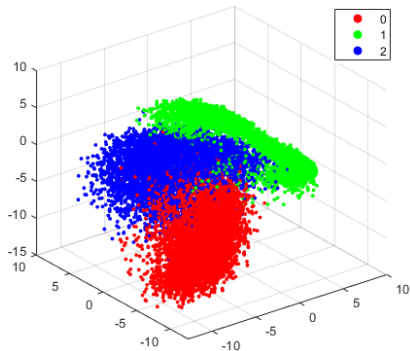


Original.

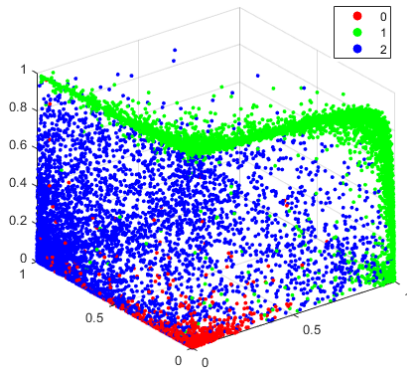


Reconstruction.

# 3 Hidden Neurons, 3 Digits

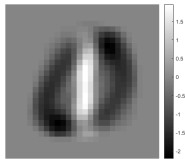


Spatial separation of activations.

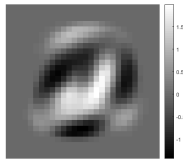


Spatial separation of latent variables.

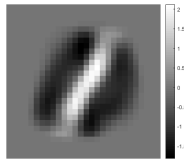
# 3 Hidden Neurons, 3 Digits



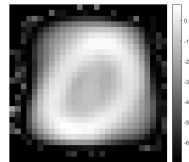
First basis.



Second basis.



Third basis.



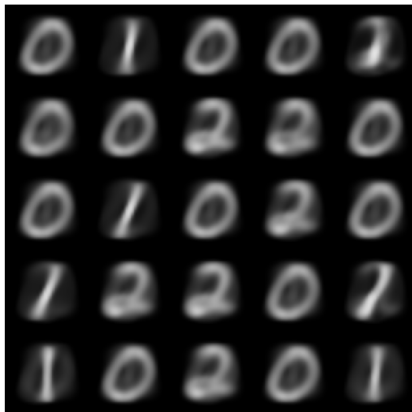
Bias.



# 3 Hidden Neurons, 3 Digits

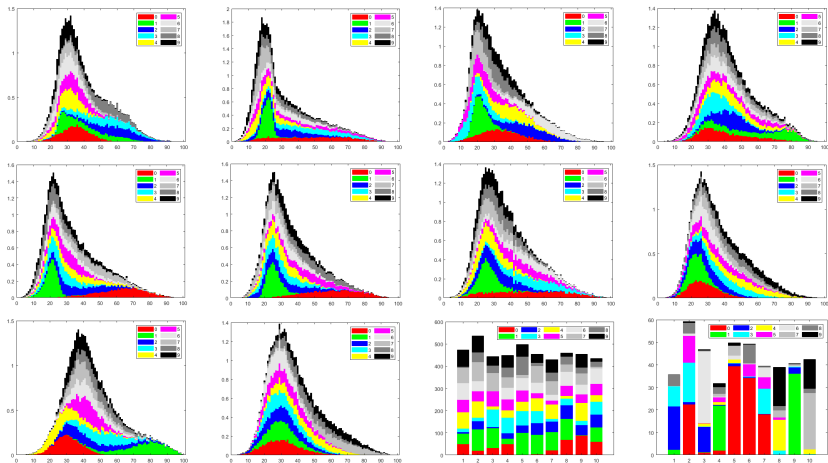


Original.



Reconstruction.

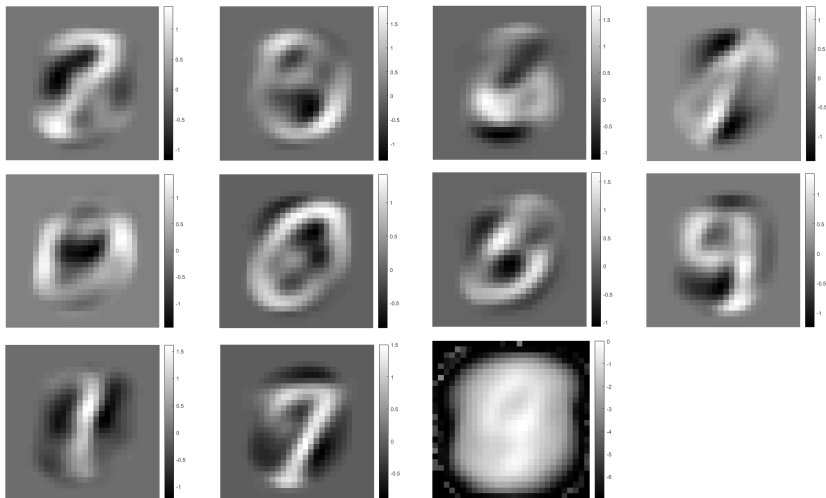
# 10 Hidden Neurons, 10 Digits



# 10 Hidden Neurons, 10 Digits

Neuron \ Digit										
	0	1	2	3	4	5	6	7	8	9
1			×	×					×	
2	×			×		×			×	
3			×				×			
4		×							×	
5	×							×		
6	×					×			×	
7	×			×		×	×			
8					×					×
9		×	×							
10								×		×

# 10 Hidden Neurons, 10 Digits

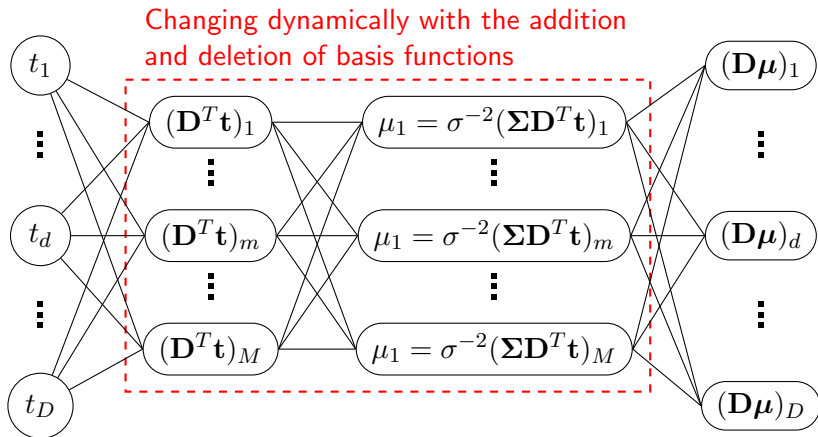


# 10 Digits, 10 Hidden Neurons

Ten digits, ten hidden neurons. ☹️



⇒ more hidden neurons 😊



## *Conclusions*

- Flexible framework.
- Giving probabilistic meaning to the relevance of the model components.
- Capable to generate and include new dictionaries if new insights are gained.
- Capable to incorporate new data.
- Confidence measure for predictions.
- Guidance for the data gathering process.

## Contact

- LinkedIn:  
<https://www.linkedin.com/in/anita-faul-123750104/>
- Forthcoming book: <https://www.amazon.co.uk/Concise-Introduction-Machine-Learning/dp/0815384106>