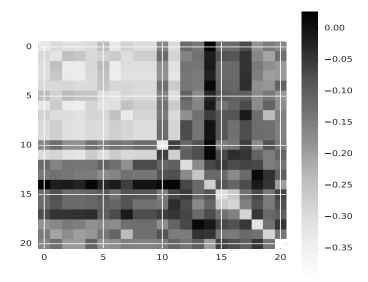


# **NeuroDesign: optimalisation report**

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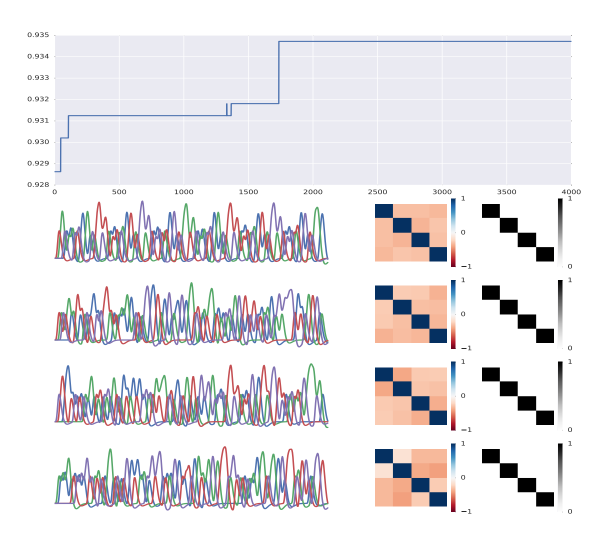
## Correlation between designs

During the optimisation, the designs are mixed with each other to find better combinations. As such, the designs can look very similar. Actually, the genetic algorithm uses natural selection as a basis, and as such, the designs can be clustered in families. This is the covariance matrix between the final 20 designs



### **Selected designs**

The following figure shows in the upper panel the optimisation score over the different generations. Below are the expected signals of the best designs from different families, more specific and in relation with the covariance matrix, designs 0, 12, 15, 17. Next to each design is the covariance matrix between the regressors, and the diagonal matrix with the eigenvalues of the design matrix.



### **Experimental settings**

Repetition time (TR):	0.68
Number of trials:	128
Number of scans:	542
Number of different stimuli:	4

Stimulus probabilities:	0.25	0.25	0.25	0.25
Stimulus probabilities:	0.23	0.25	0.25	0.20

Duration of stimulus (s)	22
Duration of stillialas (s)	2.2
Seconds before stimulus (in trial):	0.0
Seconds after stimulus (in trial)	0.4
Duration of trial (s):	2.6
Total experiment duration(s):	368

Number of stimuli between rest blocks 0
Duration of rest blocks (s): 0.0

	0.5	0.5	-0.5	-0.5
Contrasts:	-0.5	0.5	-0.5	0.5

ITI model: exponential

minimum ITI: 0.0 mean ITI: 0.26 maximum ITI: 6.0

Hard probabilities: False
Maximum number of repeated stimuli: 6
Resolution of design: 0.136

Assumed autocorrelation: 0.3

## **Optimalisation settings**

Optimalisation weights (Fe,Fd,Fc,Ff):	.0 (	J.1 (	J.4	0.5
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Aoptimality? True Number of designs in each generation: 20 Number of immigrants in each generation: 4 Confounding order: 3 Convergence criterion: 1000 Number of precycles: 1000 Number of cycles: 4000 Percentage of mutations: 0.01 Seed: 214013809