

**Functional** Magnetic Resonance **Imaging** 

GitHub <a href="https://github.com/dcdace/fMRI\_training">https://github.com/dcdace/fMRI\_training</a>

Dace Apšvalka Winter, 2024

### Outline

- Introduction
- Experimental design
- Data management
- Pre-processing
- Statistical analysis
- Practical demo

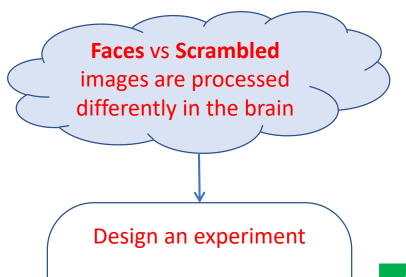


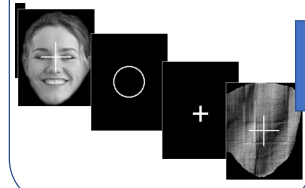
**Data**Organise & Manage



Pre-process

Analyse





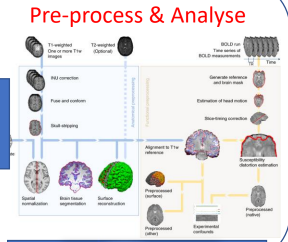
Data

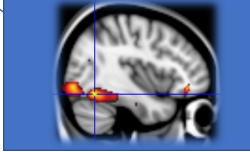
Stimuli Timing Collect the MRI data



Data

Anatomical image Functional images Event details





# fMRI Aanalysis

The General Linear Model (GLM)

Largely based on Idan Blank's materials

https://cbmm.mit.edu/videos?field\_video\_grouping\_tid%5B%5D=770

### Finding a Face area in the brain

Which brain regions are engaged when people look at faces



- With fMRI the meaningful questions are questions that compare two conditions
  - We need some sort of control condition -> Scrambled condition

- Which brain regions respond more to looking at face images than scrambled images
  - The control question hopefully helps to wash out all the regions we are not interested in. Because regions that we are NOT interested in should activate both conditions to the same extend (e.g. visual areas)

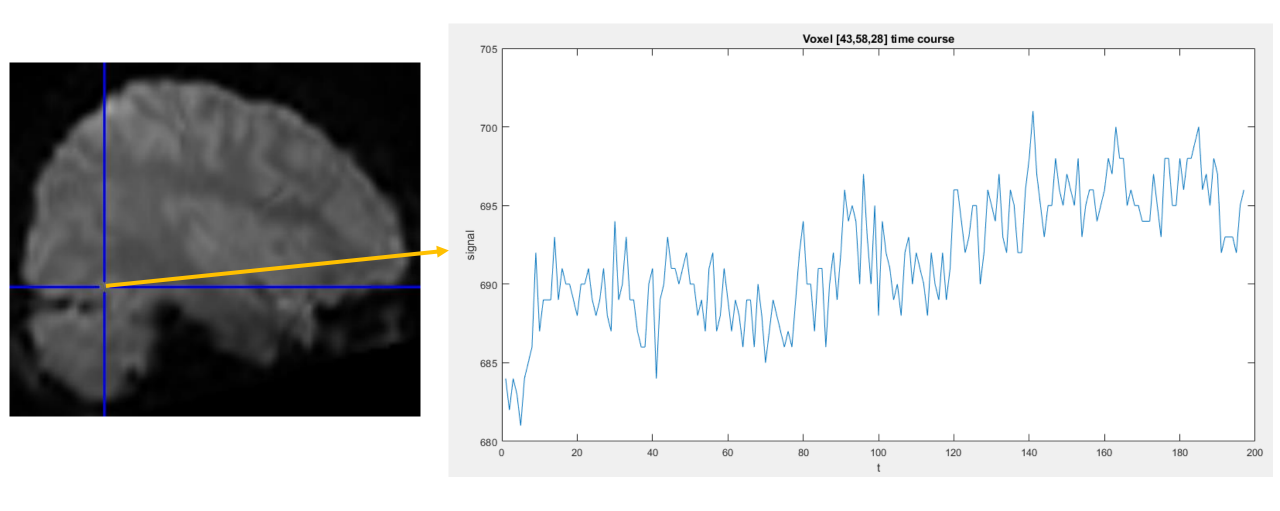
### The structure of our data

• Question: Which brain regions respond more to face images than to scrambled

images Experiment Time The brain data that we collect are a stack of 2D

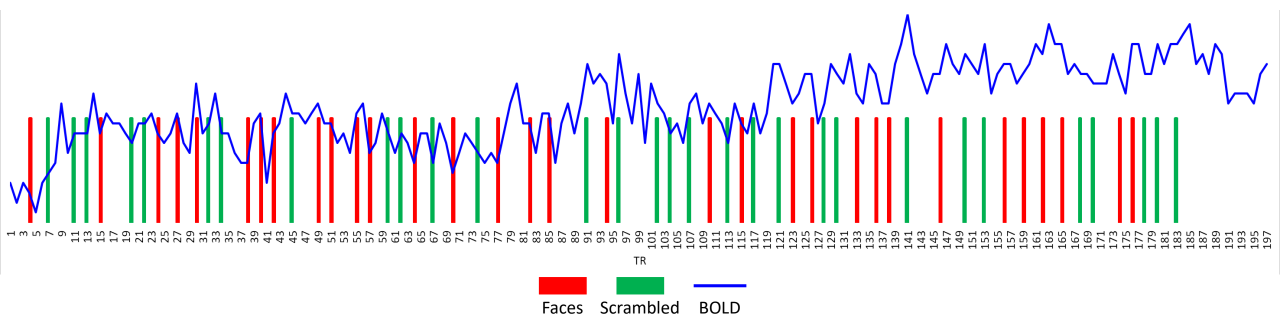
slices that form a 3D volume -> a large 3D cube with it's smallest unit -> a small 3D cube -> a voxel

### Voxel time course



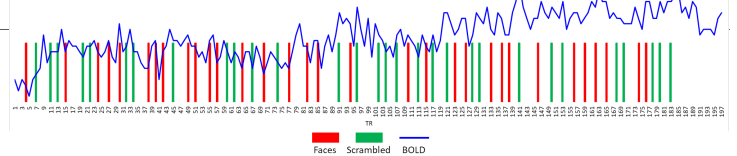
## Question: Which brain regions respond more to face images than to scrambled images

- Event onsets and BOLD signal in a sample voxel
- How do we analyse this?

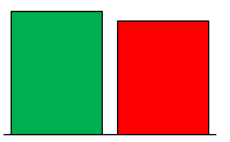


Question: Which brain regions respond more to face images than to

scrambled images



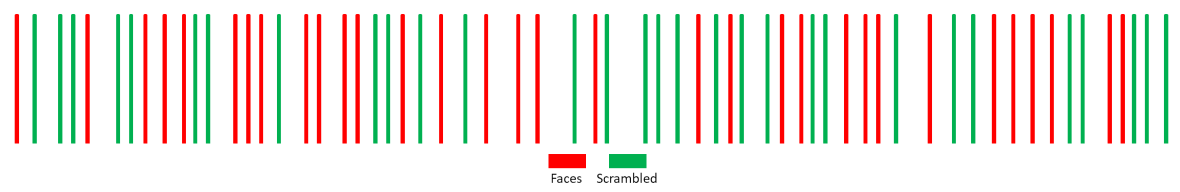
- A raw BOLD signal is noisy
- Let's start with an intuitive way, and that's how people started to analyse fMRI data
- Analysis: an intuitive approach
  - 1. For each voxel, look at its signal time-series (activity across time)
  - Average the signal across volumes that were collected while participants were looking at Faces
    - Mean Faces = 691.92, Mean Scrambled = 692.35, F-S = -0.43



Why does the averaging method does not work?

Question: Which brain regions respond more to face images than to scrambled images

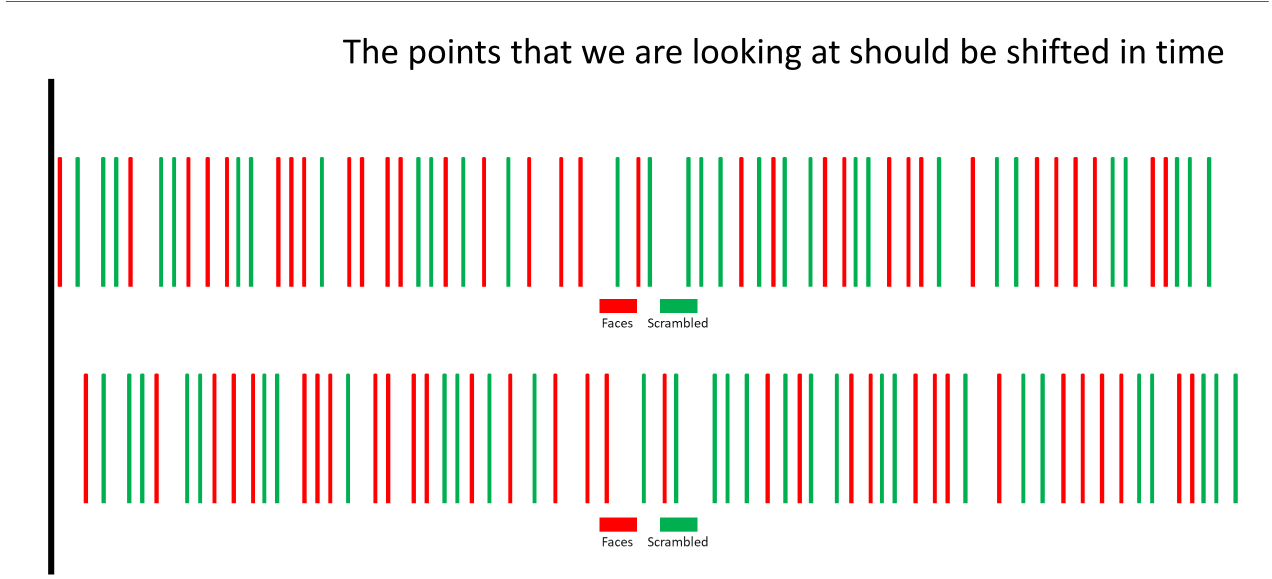
Stimuli



- Neural activity
  - Starts briefly after the onset of the stimulus
- BOLD signal (not a direct measure of neural activity)
  - When neurons are active, the fMRI BOLD signal will rise \_\_\_\_\_
    - A. Immediately
    - B. 1-3 seconds after the neural activity
    - C. 6-12 seconds after the neural activity
    - D. 25-30 seconds after the neural activity

We were averaging the wrong points! The points in time when the 'Face area' would respond are actually little later than the onset of the stimulus; about 6-12 s later then the onset.

Question: Which brain regions respond more to face images than to scrambled images



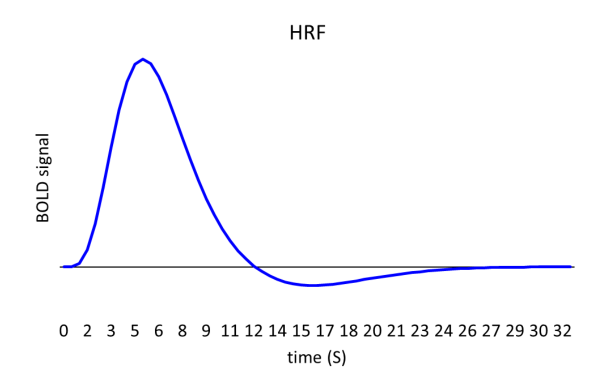
Question: Which brain regions respond more to face images than to scrambled images +6s shift

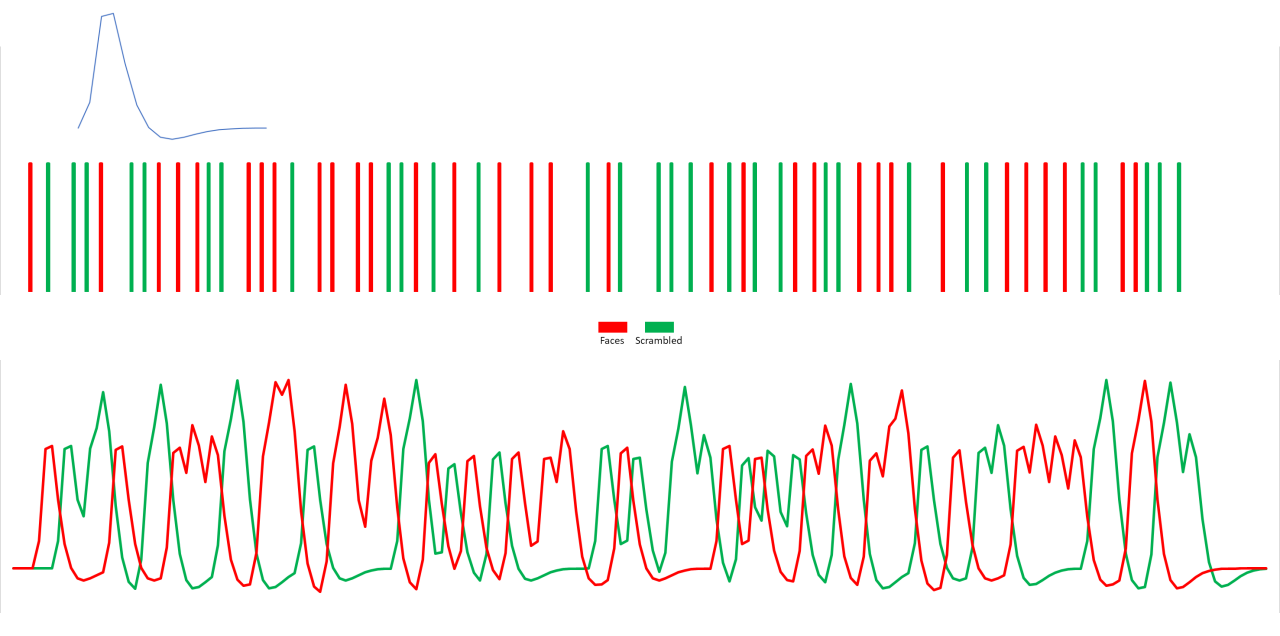
Bold signal does not change from 0 to 1 (on or off)!

Looks slightly better, but still not working!

## Question: Which brain regions respond more to face images than to scrambled images

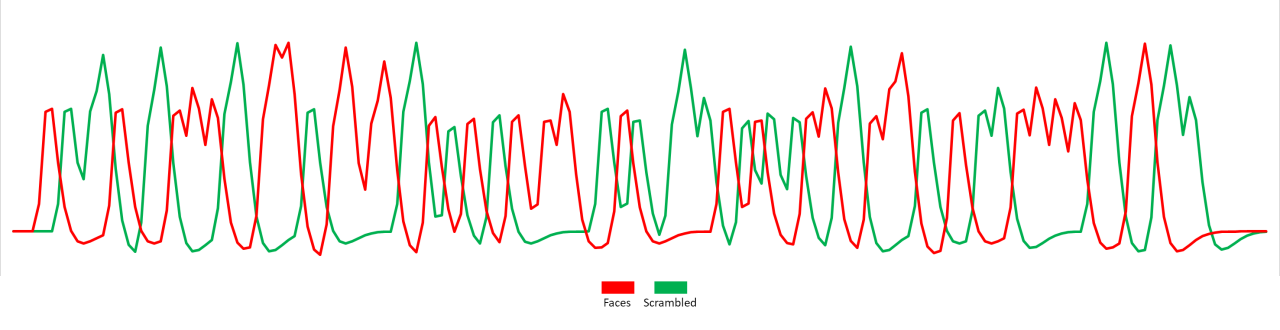
- The haemodynamic response function looks like this
- It is not binary, but smooth
- We need to make our prediction to look more like this
- How do we do that? This is done by mathematical process called convolution
- It is filtering the signal to look more like this shape



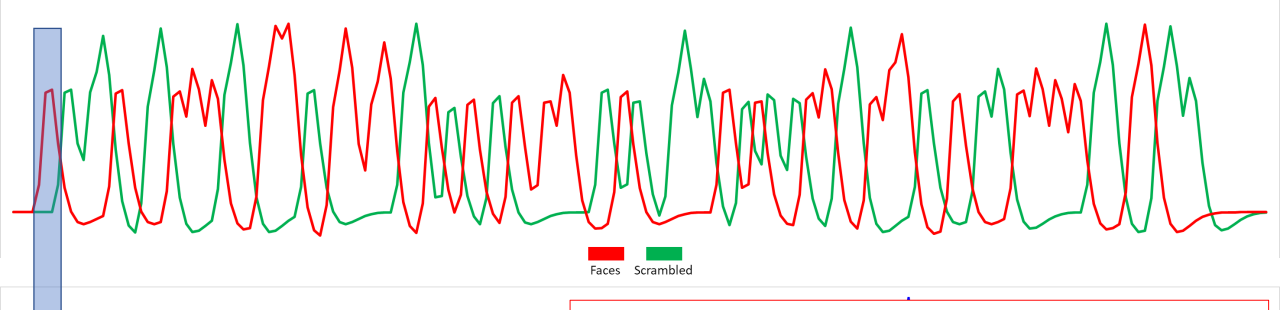


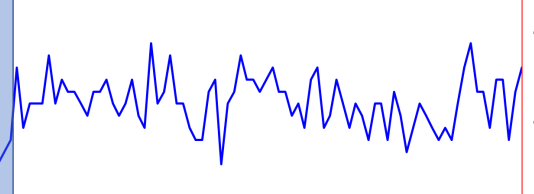
These are our BOLD predictions. This is how we should expect our signal to look like!

Question: Which brain regions respond more to face images than to scrambled images



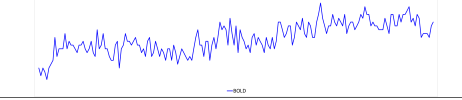
The time series of a voxel that responds more to Faces than to Scrambled should look like a combination of the two signals above, with Faces having more weight than Scrambled.





#### Which points should we average?

- Averaging does not seem to be the best idea anymore.
- We need a more principled solution, not an arbitrary choice of points!
- Voxels in the 'Face area' should respond
   a lot to Faces, but very little to Scrambled
   images.



We can approximate the signal time-series of a voxel by combining these 3 signals:

- Baseline signal:
  - an average activity in that voxel when you are doing nothing
- Response to Faces:



Response to Scrambled:

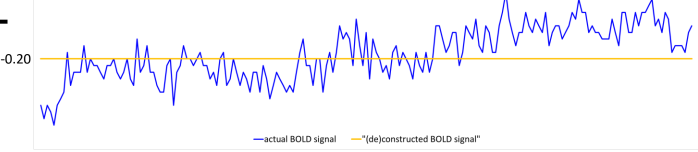


The true BOLD signal

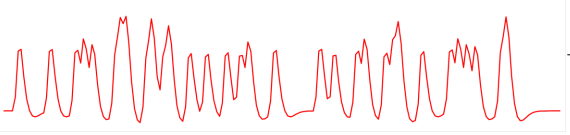
Deconstructed BOLD signal

**1.** Let's say the average signal at baseline was -0.2. We add this baseline measure to our deconstructed BOLD signal and see how it compares with the actual BOLD signal.

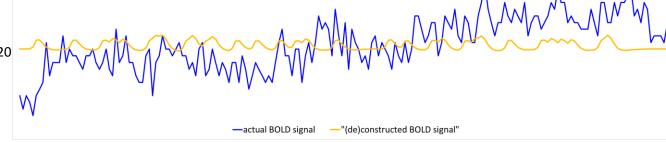








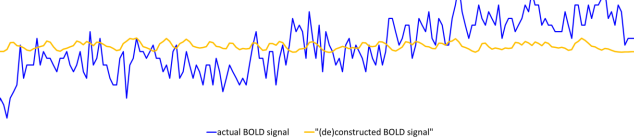
2. We predict that in this voxel Faces condition contributes quite a lot to the true BOLD signal. Let's assign it a weight 0.89 (we will find out later how exactly we find these weights), multiply the predicted response by it and add to the deconstructed signal.



#### + Scrambled 0.36



3. We predict that in this voxel Scrambled condition has little contribution to the true BOLD signal. Let's assign it a weight 0.36, multiply the predicted response by it and add to the deconstructed

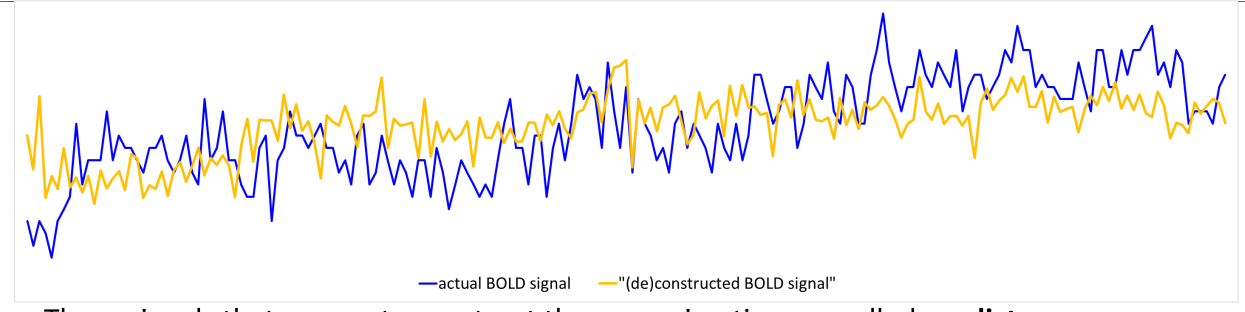


mp5-0.01 + mp6 0.64 +

—"(de)constructed BOLD signal

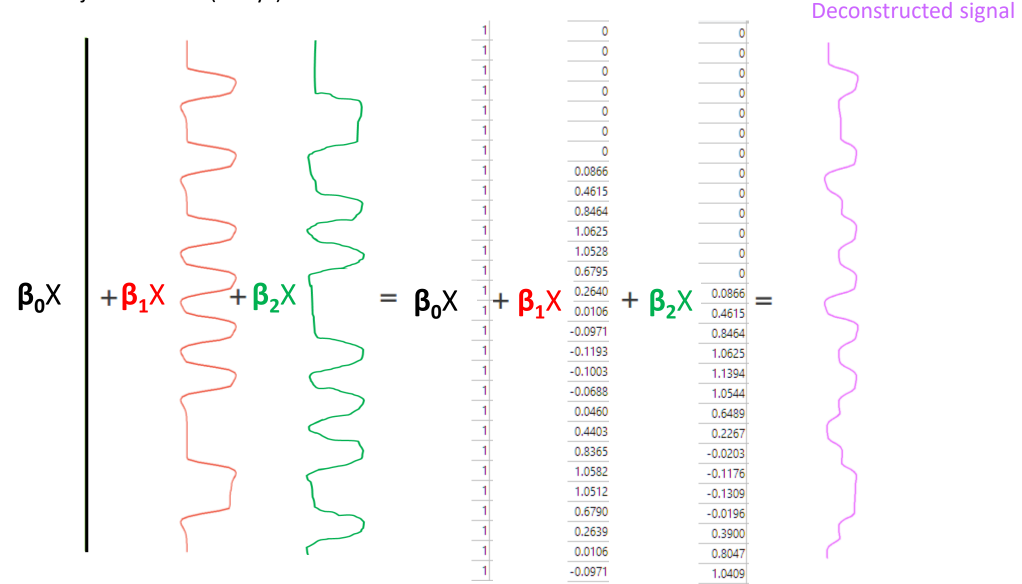


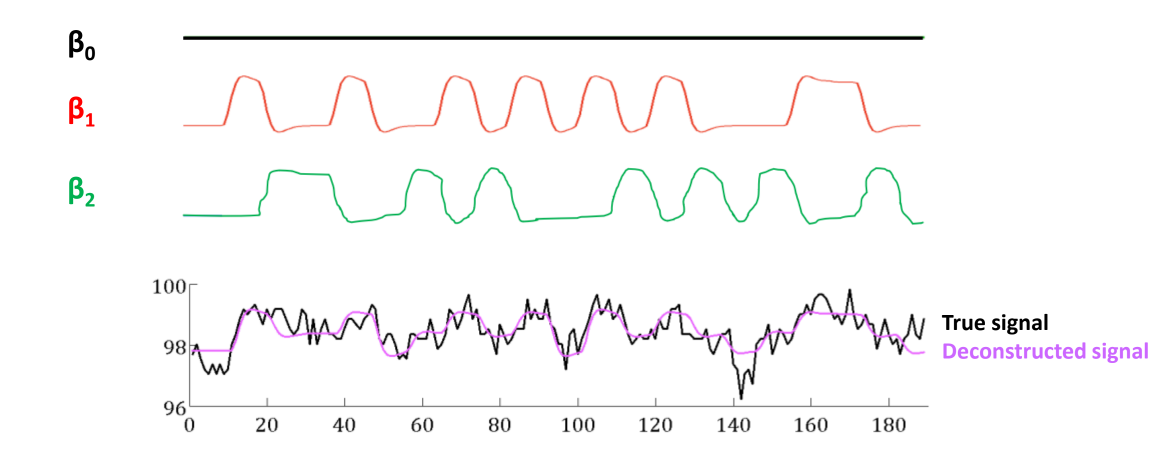
The deconstructed signal matches the true signal much better now!



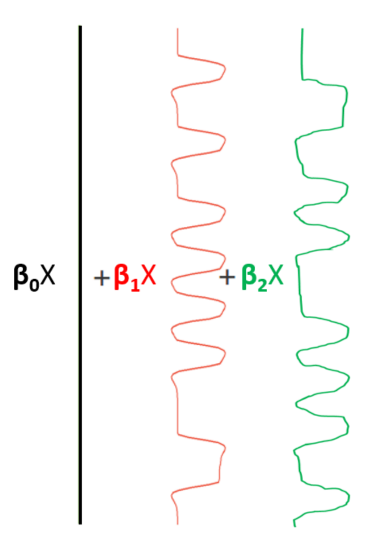
- These signals that we use to construct the approximation are called predictors
- One predictor predicts a constant response, just baseline
- Another predictor predicts how you should respond to the task
- Each predictor is associated with a weight called a beta-weight
- To create a **linear combination** of predictors, which **approximates a true signal**, we multiply each predictor by its beta-weight and then sum the results.

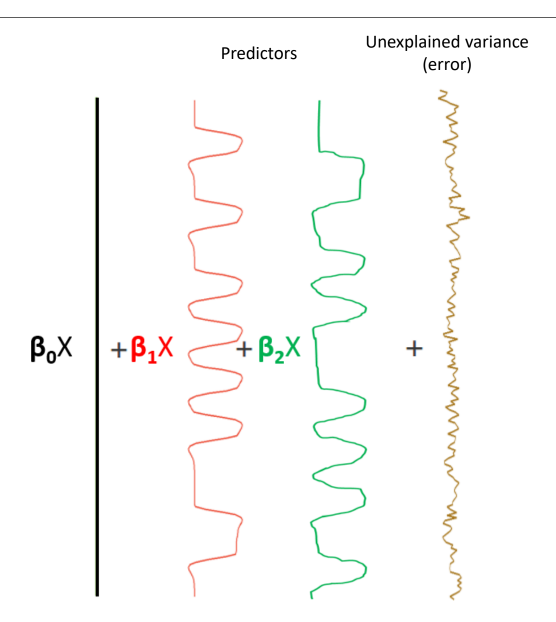
These signals are just columns (arrays) of numbers stored in a matrix.

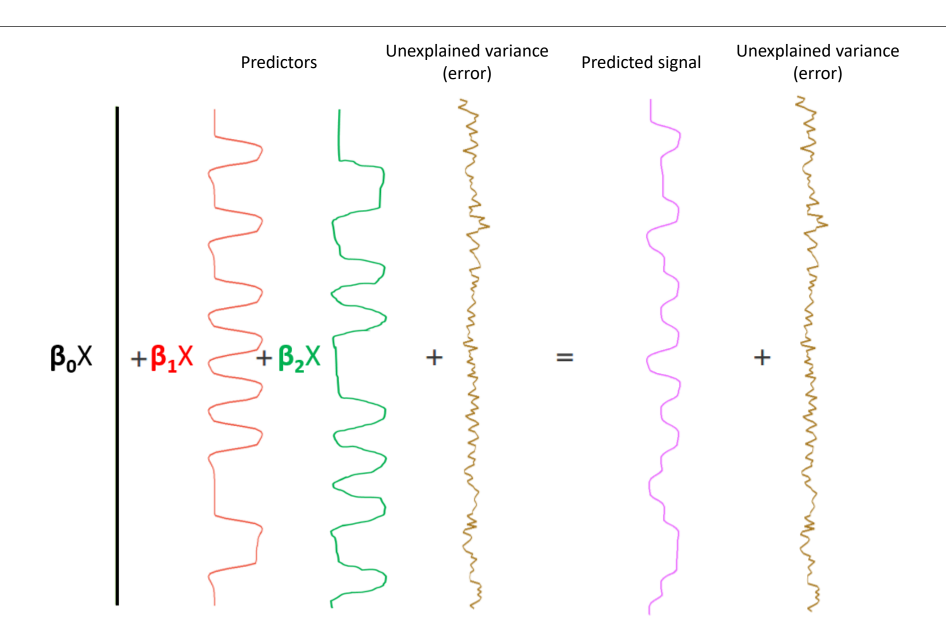


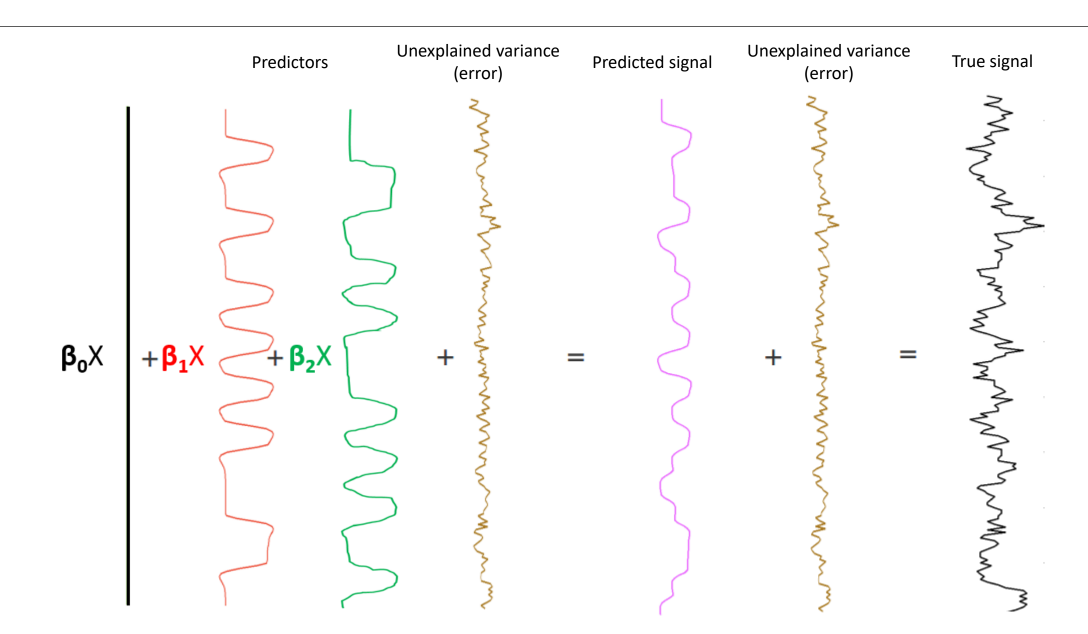


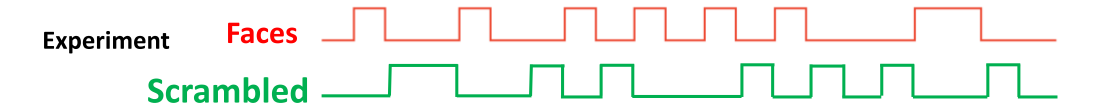
For every time point: signal(t) - prediction(t) = error(t)











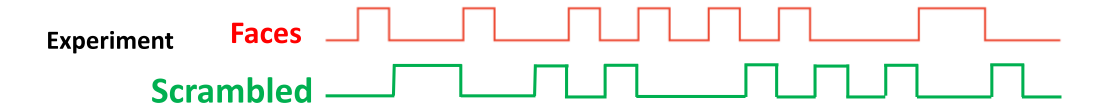
**Analysis:** the approach that works

- Find the beta-weights that best approximate the voxel's signal time-series. The best approximation is the one with the least errors
- Compare the beta-weights for estimated response to Faces to betaweights for estimated response to Scrambled

```
BOLD signal = task-related activity changes + noise (other changes)
explained variation unexplained variation

Linear combination of predictors (deconstructed signal)

+ errors
```



**Analysis:** the approach that works

- Find the beta-weights that best approximate the voxel's signal time-series. The best approximation is the one with the least errors
- Compare the beta-weights for estimated response to Faces to betaweights for estimated response to Scrambled

To find beta-weights, we use General Linear Model (GLM)

### How to find betas: GLM $Y = X\beta + \epsilon$

#### What we know?

- BOLD signal: we collect this from the brain (functional data)
- X: the design matrix (each column is a predictor that we built ourselves)

#### What we want to find?

 b: vector of beta-weights (one weight for predictor in X) that give the best approximation of the BOLD signal

#### How we find it?

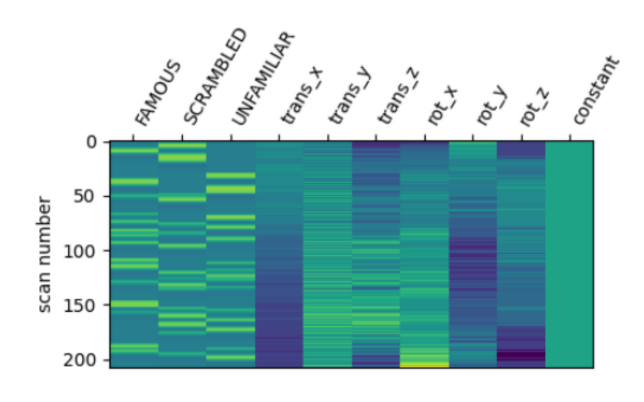
 By minimising the sum of squared errors. In practice, the GLM has a formula, which guarantees to find these beta-weights

### How to find betas: GLM $Y = X\beta + \varepsilon$

- Any predictor that can help approximate the BOLD signal will decrease the Sum of Squared Errors
- Therefore, we include additional predictors:
  - The 6 head-motion parameters

An example design matrix. Each column is a predictor:

- events
- 6 movement parameters
- constant



### Multiple regression, to find beta-weights

0.064

voxe	cel	S	F	rp1	rp2	rp3	rp4	rp5	rp6				Dependent	
-1.98	8409	0	0	0.744	-0.968	-0.815	-0.263	2.13	-0.0816			Þ	voxel1	
-2.49	9237	0	0	1.15	-0.399	-1.42	-0.365	1.95	0.0903				Metho	od Enter 🔻
-1.98	8409	0	0	-1.33	0.482	-1.81	-0.729	0.666	-0.304				Covariates	
-2.23	3823	0	0	2.11	0.35	-1.81	-0.748	0.177	-0.157			$\triangleleft$	<b>♦</b> S <b>F</b>	^
-2.74	4651	0	0.0865661	1.49	0.722	-1.99	-0.63	-0.0958	-0.202				<b>♦</b> rp1	
-1.98	8409	0	0.374888	2.35	1.17	-1.98	-0.718	0.0911	-0.509					
-1.729	2996	0	0.384923	0.989	-0.953	-2.12	-1.12	-0.369	-0.378				<b>№</b> rp4	· •
-1.47	7582	0.0865661	0.216117	2.11	1.49	-2	-1.41	-0.556	-0.0755					
icients												1.3		
Model		Un	Unstandardized		Standard Error		Standardized		t	р		1.1		Ţ
1	intercept		-0.199		0.106				-1.877	0.062				
			0.359		0.329		0.070							
	S		0.3	59	0.3	329	0.0	70	1.091	0.277	Fe	0.9		
	S F		0.3 0.8			329 338	0.0° 0.1°				alue	0.9	Т	
				91	0.3			71	1.091	0.277	value	0.7	Ţ	
	F '		0.8	91 02	0.3	38 )77	0.1	71 02	1.091 2.637	0.277 0.009	value			
	rp1		0.8 -0.3	91 02 29	0.3 0.0 0.0	38 )77	0.1 -0.3	71 02 29	1.091 2.637 -3.916	0.277 0.009 < .001	beta value	0.7		
	rp1 rp2		0.8 -0.3 -0.1	91 02 29 26	0.3 0.0 0.0 0.0	338 )77 )51	0.11 -0.36 -0.12	71 02 29 27	1.091 2.637 -3.916 -2.544	0.277 0.009 < .001 0.012	value	0.7 0.5		

A comparison of beta-weights is called a **contrast**.

0.643

rp6

Formally, a contrast is a vector indicating which beta-weights we are testing.  $\beta(F) - \beta(S)$ : [-1 1 0 0 0 0 0 0] -0.359 + 0.891 = 0.532

9.978

< .001

-0.1

0.359

0.891

0.643

### How to find betas: GLM

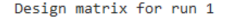
- Extract the signal time-series from a given voxel
- 2. Create a design matrix (predictors)
- 3. Run GLM (the signal and your design matrix are the inputs) to find beta-weights that best approximate the true signal
- 4. Define your **contrast** and test it
- 5. Repeat for all voxels
  - Produces
    - An image file for each predictor with beta values for each voxel: beta-maps
    - An image file with contrast values for each voxel: contrastmaps
    - An image file with contrast-specific t-values for each voxel:
       t-maps

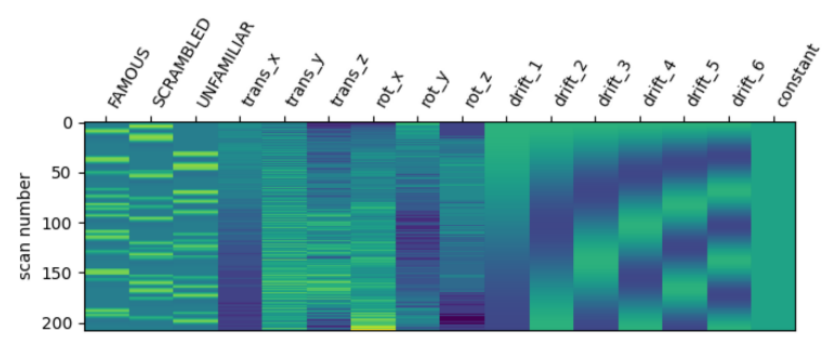
### First-level analysis: model specification

```
fmri_glm = FirstLevelModel(
    t_r = TR,
    slice_time_ref = 0.5,
    hrf_model = 'spm',
    drift_model = 'Cosine',
    high_pass = 1./128,
    smoothing_fwhm = 6,
    noise_model = 'ar1'
)
```

- **t\_r**, Repetition Time (TR)
- **slice\_time\_ref**: This parameter indicates the time of the reference slice used in the slice timing pre-processing step
- hrf\_model: defines the HRF model to be used
- drift\_model: specifies the desired drift model for the design matrices. It can be 'polynomial', 'cosine' or None. Default = 'cosine'.
- high\_pass: specifies the cut frequency of the high-pass filter in Hz for the design matrices. Used only if drift\_model is 'cosine'. Default=0.01 (1/128, as in SPM). Slow signal drifts with a period longer than 128 s will be removed. It is a way to remove possible confounds.
- **smoothing\_fwhm**: the full-width at half maximum in millimeters of the spatial smoothing to apply to the signal (smoothing was not done in fMRIPrep!).
- noise\_model: {'ar1', 'ols'} The temporal variance model. Default='ar1'. Accounting for serial correlations in fMRI time series due to aliased biorhythms and unmodelled neuronal activity. Accounts for non-independency in the BOLD response.

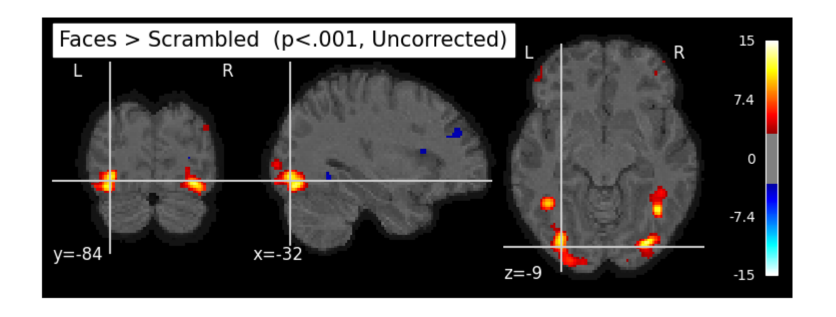
### First-level analysis: design matrix and contrasts





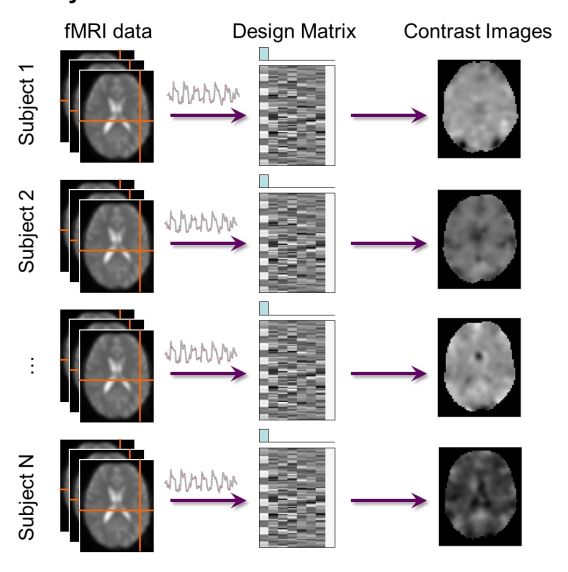
### Faces > Scrambled

Statistical brain map (z-map) of regions showing significant positive difference between Faces and Scrambled



### First-level analysis

• Run the GLM for each subject







## Data Organise & Manage



Notebook: nb05 Subject-Level-Analysis.ipynb

**Pre-process** 

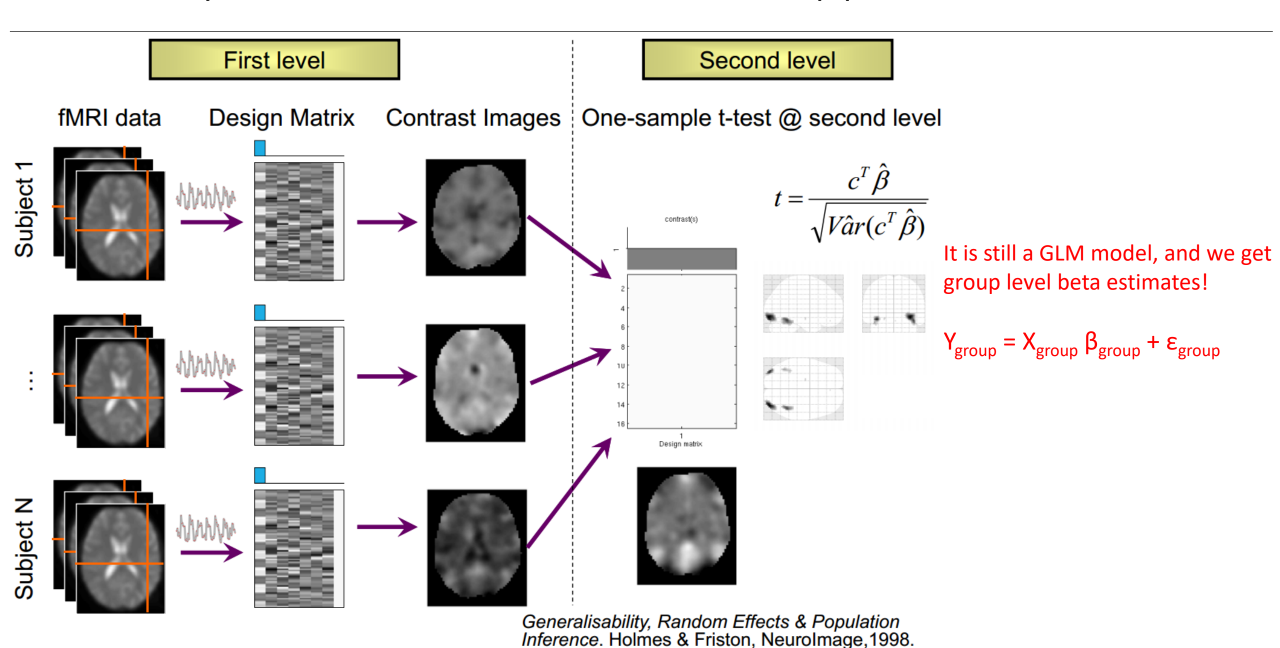
Analyse

### Group level (2nd level) analysis is across subjects

- Which voxels are showing a significant activation differences between our conditions consistently within a group
  - Average contrast effect across sample (e.g., one-sample t-test)

- Importantly, all subject brains need to be in common space, e.g. MNI, to perform voxel-wise group analyses
  - That was achieved by the Normalisation step in pre-processing

### Summary statistics, Random effects approach







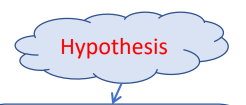
## **Data**Organise & Manage



Notebook: nb06 Group-Level-Analysis.ipynb

Pre-process

Analyse



#### Design an experiment

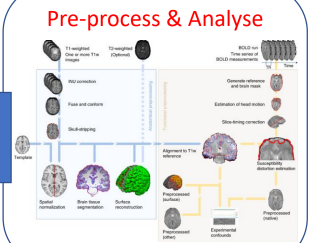


Stimuli Timing

#### Collect the data

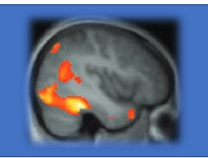


Anatomical image Functional images Event details













### Sharing & Reporting



• Share your **code** and notebooks on GitHub



- Make it citable with Zendono
  - <a href="https://docs.github.com/en/repositories/archiving-a-github-repository/referencing-and-citing-content">https://docs.github.com/en/repositories/archiving-a-github-repository/referencing-and-citing-content</a>



 If you have consent from participants, share the BIDS data on OpenNeuro. Before sharing, 'deface' the T1w images



Add your contrast maps to NeuroVault

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