# PsPM: Psychophysiological Modelling

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# incorporating SCRalyze

# by the PsPM team<sup>1</sup>:

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 $<sup>^1</sup>$ If you have comments on or error corrections to this documentation, please send them to the PsPM team or post them on: pspm.sourceforge.net

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# Part I Background

# 1 What is model based analysis?

# 1.1 Conventional analysis

Psychophysiology is concerned with inferring psychological states from physiological (often peripheral) measurements such as skin conductance responses (SCR), heart beats, respiration, and others. Conventional approaches in psychophysiology seek to identify data features ("indices") that closely follow a psychological state of interest. Conventional data analysis algorithms extract these data features, to "index" central states. For example, to infer stimulus-evoked sympathetic arousal (SA) from skin conductance responses (SCR), one may filter the SCR data to reduce observation noise, define a response window after the stimulus, and define some criteria do detect peaks within this window. The amplitude of such peaks may then be taken as "index" of SA. The development of such indices has been a major endeavour in the field of psychophysiology. It is usually based on qualitative or semi-quantitative models of how the index relates to the central state of interest. Model-based analysis provides a formalisation of this process drawing on strict, quantitative causal models - see [1] for a review.

# 1.2 Model-based analysis

Model-based approaches start with explicit, mathematical models that formulate how observed data are generated by central processes. For example, a model may formulate the relation between sympathetic arousal (SA) and skin conductance responses (SCR),  $SA \mapsto SCR$  in mathematical form. This kind of model, in general, is often termed a "forward model": it predicts a data time series (SCR) from a known central process (SA). Another term for this kind of model is "generative", because it describes how a central process generates the data. When we analyse experimental data, we are faced with the opposite situation: we know the observed data data but not the central process, and seek to infer the central process from the data. In order to do so, one has to turn the forward

model backwards, to arrive at the relation  $SA \leftarrow SCR$ . In statistics, this process is often termed "model inversion". It provides the most likely estimates of the central process, given the observed data and the model. Both conventional and model-based analysis aim to infer central process from observed data. The difference is that model-based methods use a stringent mathematical language and probabilistic model inversion to do so. Note that probabilistic models acknowledge the existence of noise in the data, and they do not seek to explain that noise with variation in central states. Hence, different from what may be a common misconception in the psychophysiological literature, the criterion for the quality of a model-based method is not how well the model "fits the data", but how well it can recover the central states of interest.

## 1.3 Evaluation

How can we infer that one method of inferring a central state is better than another? Each method returns indices or estimates of the central state, but which ones are more precise? Often we are interested in psychological states which can not be measured directly. We propose to use experimental manipulations to create two central states that are known to differ. We can then evaluate how well a method recovers that difference. We have termed the sensitivity to distinguish these central states "predictive validity". Simply speaking, when we compare the two central states with a t-test on their estimates, the test statistic should be as high as possible. In a more rigorous fashion, we can frame this evaluation as comparison of non-nested predictive models, and use a measure of model evidence to assess the difference between methods. This approach also allows a principled statement whether methods are significantly different. All methods included in this software have empirically been shown to provide better or equal predictive validity than the best available corresponding conventional index.

# 2 Skin conductance (SCR) models

#### 2.1 General

Skin conductance responses are elicited via the sympathetic nervous system. Hence, SCR models relate a psychological construct termed "sympathetic arousal" (SA) to observed SCR [2]. All SCR methods split up the relation between SA and skin conductance responses  $SA \mapsto SCR$  into two relations:  $SA \mapsto SN \mapsto SCR$ , where SN is the sudomotor nerve activity. The peripheral model  $SN \mapsto SCR$  is a biophysical model that specifies how SN activity generates SCR. The central or neural model  $SA \mapsto SN$  embodies knowledge and assumptions about the kind of SN impulses that are generated by a central (psychological) state, SA.

# 2.2 Peripheral model

Peripheral SCR models use a standard convolution model, also termed linear time-invariant (LTI) model, introduced in [3, 4]. This model is assumed to describe all processes between initiation of a SN burst in the CNS, and the measured SCR. All these processes are not described in biophysical detail but instead lumped into a phenomenologically derived impulse response function which is convolved with the sudomotor input:

$$SN \mapsto SCR$$

$$SCR\left(t\right) = SN \otimes SCRF + \varepsilon = \int_{0}^{t} SCRF\left(\tau\right) SN\left(t - \tau\right) d\tau + \varepsilon, \ t \geq 0$$

where  $\varepsilon$  describes the observation error.

#### **Properties**

The peripheral convolution model has two specific testable properties. There is direct (invasive) and indirect evidence for each of these properties.

Time invariance: the system's response does not explicitly depend on time, or in other words, a given input always produces the same output.

- Direct evidence: The amplitude of individual SN bursts is linearly related to the amplitude of ensuing SCR, with a considerable scatter [5]; to the maximal rate of sweat expulsion; and, somewhat more weakly, to the integrated sweat production during the skin response [6]. After initial dishabituation, constant SN stimulation leads to SCR, with constant amplitude and latency [7]. Repeated SN stimulation yields slightly different SCR shapes [8], although this could be due to variation in elicited neural responses that were not measured downstream. In summary, these findings are consistent with (but do not prove or disprove) the time invariance principle.
- Indirect evidence: We have shown that for short events (< 1 s duration) that are separated by at least 30 s, more than 60% of the variance in (high pass filtered) SCR can be explained under an LTI model. We have established the evidence for linear models using aversive white noise bursts, aversive electric stimulation, aversive pictures, auditory oddballs, and a visual detection task [4]. Because only 60% of the variance can be explained under this model, there is substantial residual variance 40% of total variance. Is this residual variance due to violations of the LTI assumptions? We have shown that in the absence of any event for 60 s,

signal variance amounts to more than 60% of total variance during stimulus presentation. That is to say, by introducing events, residual baseline variance is reduced by 20%. This suggests that residual variance is not caused by violations of the time invariance assumption but variation of the input (for example, spontaneous fluctuations), and observation error.

## Linearity: the system's response does not depend on previous inputs.

- Direct evidence: Latency of sweat expulsion is slightly shortened when sweat expulsion rate is very high (i. e. for very strong SN bursts) but not when SN firing is frequent [6]. SCR to individual SN stimulations depend linearly on skin conductance level, and are slightly suppressed upon repetition of the stimulation after 1 s [9]. This suggests that the linearity principle does not hold under all conditions.
- Indirect evidence: The linearity principle amounts to saying that responses are not influenced by the (preceding) baseline signal. We tested this assumption by presenting two subsequent events with an inter stimulus interval (ISI) of either 2 s, 5.5 s, or 9 s, such that the baseline signal at these three time points differed markedly. Violations of the linearity principle in the peripheral system would imply that the amplitude of the subsequent response is dependent on the baseline signal, and thereby, upon the ISI. The second response was always smaller than the first. However, this was not dependent on the ISI, and hence not on the baseline signal. We interpret this effect as central repetition suppression (of the inputs into the peripheral system) and conclude that linearity is appropriate for the peripheral system. In other words, the peripheral response to one input is not modulated or affected by the response to another, even when they overlap in time [4].

In summary, it appears that both properties are good approximations to reality with the expectation that non-linearities may occur if the time interval between two SN firing bursts is very short (< 1 s) or when the SN firing burst is very strong. It is in principle possible to model non-linearities in a linear model, using Volterra kernels. This approach is implemented for analysis of functional magnetic resonance images in the software SPM [10, 11]. It is not implemented for SCR models.

# Skin Conductance Response Function (SCRF)

The peripheral model in its general form does not specify a particular form of the SCRF. There are three principled ways of defining a SCRF:

1. Individual response function. The most precise option is probably to measure the SCRF for each individual by providing brief inputs into the peripheral system and measuring the response. PsPM allows specifying in-

dividual response functions. The benefit of this approach has not been empirically tested.

- 2. Canonical response function. The most parsimonious method is to use a so-called canonical SCRF which is assumed to be constant across individuals. We have previously shown that a canonical SCRF can explain 50% of the variance in individual SCR elicited by various stimuli, while individual SCRFs explain about 75% of the variance [4]. Using a canonical SCRF, linear and non-linear models show a good predictive validity, superior to conventional methods [12], see also (Staib et al., submitted).
- 3. Constrained response function. One can estimate an SCRF from the data of an actual experiment and use this for analysis. It is usually necessary to constrain the shape of the SCRF to make the model estimable. PsPM provides two options for this:
- In GLM, one can estimate parameters of the (orthogonalised) time derivative of the SCRF. In order to combine this with the canonical SCRF, the response can afterwards be reconstructed for each experimental condition, and the peak amplitude taken as estimate of SA. This approach has higher predictive validity than using the canonical SCRF alone. Less constrained options for estimating individual SCRFs are also available (in particular, using the dispersion derivative together with the time derivative) but have been shown inferior [12].
- In the non-linear model for event-related SCR, one can use the combined data from all evoked responses from one individual to estimate parameters of a third-order linear ordinary differential equation (ODE) for each participant. This approach only considers data within the inter-trial interval. In validation experiments, the ITI was not long enough to include the tail of the SCRF, and this approach has empirically been shown inferior to using a canonical response function in most cases [Staib et al., submitted].

# Canonical Skin Conductance Response Function

The canonical SCRF that is currently implemented was derived from a large dataset of 1278 SCRs from 64 individuals subjected to different experimental manipulations (white noise 85 dB and 95 dB, painful electric stimulation, aversive IAPS pictures, detection of auditory oddballs, detection of a visual target in a stimulus stream). We extracted and mean-centred the 30 s following each stimulus onset and performed a principal component analysis. The first principal component served as empirical SCRF [4].

Formulation for linear models The first PCA component of the empirical SCRF was approximated was fitted by a Gaussian smoothed bi-exponential function (also named bi-exponentially modified Gaussian function), and this provided better fit than other function categories, in particular Gaussian smoothed

monoexponential function, exponential-Gaussian hybrid function, or smoothed sigmoid-exponential function. The function is thus defined as

$$SCRF_{can}(t) \propto \int_{0}^{t} N(t-\tau) \left(E_{1}(\tau) + E_{2}(\tau)\right) d\tau, \ t \geqslant 0$$

$$N(t) = \frac{1}{\sqrt{2\pi\sigma}} e^{-(t-t_{0})^{2}/2\sigma^{2}}$$

$$E_{i}(t) = e^{-\lambda_{i}t}$$
(2.1)

$$SCR(t) = SN(t) \otimes SCRF_{can}$$

with estimated parameters:  $\hat{t}_0 = 3.0745$  seconds for peak latency;  $\hat{\sigma} = 0.7013$  for definition of the rise time; and  $\hat{\lambda}_1 = 0.3176$  and  $\hat{\lambda}_2 = 0.0708$  to define the two decay components. This function is normalised to its maximum such that an infinitely short SN impulse with unit amplitude elicits an SCR with unit amplitude. This renders parameter estimates from linear models easily interpretable in relation to conventional (peak-scoring) analysis.

Historical remarks:

- (a) The appendix of [12] contains a typographic error in the definition of the canonical response function, in particular in the integration limits of eq. (2.1). However, in all versions of the software (PsPM 1.0-2.x) the canonical SCRF was constructed using the Matlab function conv() which defines vector convolution analogous to the integration operation defined above. This implementation has not changed since the initial formulation of the model.
- (b) The initial formulation of the SCRF [3] used a Gaussian-smoothed Gamma distribution, based on a smaller dataset. In the extended data set published in [12], the polynomial part of the Gamma distribution was estimated to be one, resulting in a Gaussian smoothed exponential function.

**Formulation for non-linear models** An alternative formulation is provided for non-linear models which require a definition of the SCRF in the form of an ordinary differential equation (ODE). We use an inhomogeneous third-order linear ODE

$$\ddot{x} + \vartheta_1 \ddot{x} + \vartheta_2 \dot{x} + \vartheta_3 x + u = 0$$

$$u(t) = SN(t)$$

$$x = SCR(t)$$
(2.2)

where dot notation is used for time derivatives, and this formulation embeds convolution of the SCRF with the SN time series. Parameters were estimated from the empirical SCRF by using a Gaussian bump as canonical SN input (see section on SN below) as  $\hat{\vartheta}_1 = 0.122505$ ,  $\hat{\vartheta}_2 = 1.411425$ ,  $\hat{\vartheta}_3 = 1.342052$ . In this formulation, the delayed response (conduction latency) is not embedded in the SCRF but modelled separately as delay of the input with a parameter estimate of  $\hat{\vartheta}_4 = 1.533879$  seconds. Amplitude of the response function is rescaled such that a canonical Gaussian bump SN impulse with unit amplitude elicits an SCR with unit amplitude.

Formulation for spontaneous fluctuations For modelling spontaneous fluctuations (SF), a different database was used that contained 1153 semi-automatically detected SF from 40 male participants [13] The first PCA component was taken as empirical SCRF which has slightly different shape than the empirical SCRF for evoked responses. A number of reasons may account for this difference, from participant selection over data pre-conditioning to differences in the canonical SN input. In the absence of further knowledge about the reason for this difference, we propose to use this empirical response function for SF. We estimated parameters for the non-linear model formulation  $\hat{\vartheta}_1 = 0.9235$ ,  $\hat{\vartheta}_2 = 3.9210$ ,  $\hat{\vartheta}_3 = 2.1594$ . The delay of the response is modelled explicitly as delay of the SN input with parameter estimated from the model for event-related SCR,  $\hat{\vartheta}_4 = 1.533879$  seconds. Amplitude of the response function is rescaled such that a canonical Gaussian bump SN impulse with unit amplitude elicits an SF with unit amplitude.

**Recommendation** For GLM, it is recommended to use a constrained SCRF by combining the canonical SCRF with time derivative and reconstruct the response after estimation of amplitude parameters [1]. For non-linear models, it is recommended to use the ODE formulation of the canonical SCRF [Staib et al., submitted]. The benefit of individual response functions in either scheme has not yet been tested.

#### 2.3 Neural model

The neural model specifies how event-related or spontaneous sympathetic arousal elicits SN activity. There are three types of neural models. All models contain a latent error term  $\omega$  which absorbs spontaneous SN bursts and influences of SA unrelated to the experiment and is added to the experimentally induced SN time series,  $SN_{exp}$ :

$$SN(t) = SN_{exp} + \omega$$

# 2.3.1 General Linear Model (GLM) for evoked SA

This is the simplest neural model in which evoked SA causes an infinitely short central SN firing burst with zero latency after some event. This model is a good approximation to situations in which short external stimuli elicit SA. For each experimental event:

$$SN_{exp}(t) = \begin{cases} a & t = 0\\ 0 & \text{otherwise} \end{cases}$$

$$a = SA$$

# 2.3.2 Non-linear model (DCM) for event-related SA

This model [14] assumes that event-related SA causes a Gaussian bump like SN firing burst after some delay and with a specified dispersion. For each experimental event:

$$SN_{exp}(t) = a \exp \frac{-(t-\mu)^2}{2\sigma^2} + \omega, \ 0 < \mu < \mu_{max}, \ \sigma_{min} < \sigma < \sigma_{max}$$

$$a = SA > 0$$

For situations in which short external stimuli elicit SA, this model uses a canonical central burst latency of zero and dispersion of  $\sigma_e=0.3$  seconds. For situations in which central processes translate SA into SN with unknown parameters, latency and dispersion are estimated from the data, alongside the SA amplitude, using the following constraints:  $\sigma_{min}=0.1, \ \sigma_{max}=0.5\mu_{max}$  where  $\mu_{max}$  is the maximally allowed response latency.

#### 2.3.3 Non-linear model for tonic SA (spontaneous fluctuations, SF)

Spontaneous SN bursts cause spontaneous fluctuations (SF) in skin conductance. The number of spontaneous SN bursts is thought to depend on SA. Each spontaneous SN burst is modelled as a Gaussian bump with unknown timing and amplitude and fixed dispersion. For each spontaneous SF burst:

$$SN_{SF} = a \exp \frac{-(t-\mu)^2}{2\sigma^2} + \omega, \ \sigma = \sigma_0, \ a > 0$$

The dispersion is set to  $\sigma_0 = 0.3$  seconds.

#### 2.4 Model inversion

For all models we only consider one error term:  $\varepsilon' = \varepsilon + \omega \otimes SCRF$  which collapses observation error and latent error.

## 2.4.1 General linear model (GLM) inversion

In the GLM for SCR, only the amplitude of evoked SN is being estimated. Both latent error and observation error are lumped into one error term  $\varepsilon$ . This renders the model linear, and the estimation can be performed by simple matrix inversion. Essentially, predicted SCR functions are created per experimental

condition by modelling events in an experimental condition as stick functions, and convolving them with the SCRF. The result then forms one column in a design matrix:

$$Y = X\beta + \varepsilon'$$

Here,  $\beta$  is the vector of amplitude parameters for each experimental condition which relates to the mean evoked SA in condition i:  $\beta_i = SA_i$ . Finally, each column of X is defined as

$$X_{ij} = SN_i \otimes SCRF_i$$

where  $SN_i$  is the expected time series of SN firing for an experimental condition i with unit amplitude, and j is the index of the SCRF basis set. Finally, X contains columns for the intercept. It is possible to concatenate data from different experimental sessions, and an intercept is modelled for each session. The model is inverted using a maximum-likelihood approach to estimating the Moore-Penrose pseudoinverse  $X^+$  of X as implemented in the matlab function pinv(). This allows to deal with situations in which X is not of full rank.

$$\hat{\beta} = X^+ Y$$

# 2.4.2 Non-linear model (DCM) inversion for event-related SA

Models in which timing and/or dispersion of SN bursts need to be estimated are non-linear and are inverted using an efficient Variational Bayes algorithm [15]. The full model can be written as:

$$SCR = SCR_{exp} + SCR_{SF} + SCL + \varepsilon'$$

The SCR components are defined by eq. (2.2) with respective parameters and a respective input function u(t). Specifically the experimental SCR component models evoked and event-related responses, and the SF component models spontaneous fluctuations between trials (part of the latent error). The SCL term is an explicitly modelled error term that absorbs baseline fluctuations and is defined as

$$SN_{SCL} = a \exp \frac{-\left(t - \mu\right)^2}{2\sigma^2}, \ \sigma = \sigma_0$$

where dispersion is set to  $\sigma_0=1.0$  seconds, and  $\mu$  is estimated from the data.

We propose the following limits for modelling SF and SCL between trials: SN bursts occur more than 5 s after the previous trial, and more than 2 s before the next trial. These limits ensure that the modelled SCL and SF do not absorb variance caused by the experiment. The use of different limits has not empirically been tested.

In this model, the parameter space is too large to invert it at once. Inversion is done in a trial-by-trial approach: a number of trials,  $T_{1..n}$  is inverted. The ensuing parameter estimates for  $T_1$  are extracted, and the parameter estimates for  $T_{2..n}$  are used as prior values for the next inversion for trials  $T_{2..n+1}$  where the estimated hidden state values at the start of trial 2 are taken as starting values of the hidden states x. This is continued until all trials have been estimated. This scheme ensures that inversion is kept tractable, and at the same time overlapping responses are taken into account. It also means that estimation errors accumulate over trials which is why part of the latent error is explicitly modelled as SF and SCL. At an inter trial interval of around 10 seconds, inverting 2 or 3 trials at the same time yields comparable predictive validity [Staib et al., submitted].

# 2.4.3 Non-linear model inversion for tonic SA (spontaneous fluctuations, SF)

DCM This model is inverted using a Variational Bayes algorithm [16, 15]. Because the number of SN bursts is difficult to estimate directly, the SF model inversion uses a fixed rate of SN bursts (set to f=0.5 Hertz by default) the amplitude of which is estimated from the data. An amplitude threshold is defined to separate "true" SN bursts from noise. An amplitude threshold of 0.1  $\mu$ S rendered the predictive highest validity in a validation paper [17]. The results from this paper have been replicated in an unpublished re-analysis using AIC as objective function in a manner similar to [12]. Re-analysing the second data set described in [17], the optimal threshold was between 0.15 - 0.20  $\mu$ S. Further systematic research is needed to fine tune the optimal threshold.

MP This is a fast (< 1 s per minute of data) approximation to the DCM inversion, by using a matching pursuit algorithm with the same amplitude threshold as the DCM inversion. In theory, this method is less precise, but empirically it is comparable to DCM results when analysing empirical data [18].

# 2.5 Data conditioning

#### 2.5.1 Filtering

The currently implemented peripheral model requires high pass filtering of the data because it does not account for slow fluctuations of the skin conductance level which render the peripheral system infinite. In case of a signal with precisely known RF, the matched filter theorem provides a way of theoretically deriving a filter that maximises the signal-to-noise ratio. In our case, the true RF is not precisely known, and also varies between individuals, such that we empirically determined the filter characteristics that maximise predictive validity of SN estimates. A unidirectional 1st order Butterworth high pass filter with cut off frequency 0.05 Hz was optimal for GLM [12]. The design matrix of the GLM is filtered with the same frequency to account for distortions of response

shape. For the non-linear model (DCM) for event-related responses, an optimal filter frequency was found at 0.0159 Hz when using a canonical SCRF and inversion of 2 trials at the same time [Staib et al., submitted]. Filter characteristics of SF models have not been investigated. In order to reduce computing load, data are resampled to 10 Hz sampling rate for all data analyses which requires a low-pass filter cut off frequency of 5 Hz.

Historical remarks:

All models were developed using a high pass filter with cut off frequency of 0.0159 Hz as commonly employed in conventional analysis (bidirectional for phasic responses, unidirectional for spontaneous fluctuations). However, this turned out to not be the optimal filter for GLM in follow-up analyses of optimal data conditioning. Note that the canonical SCRF was developed from data filtered with the original settings, and this has not changed since.

#### 2.5.2 Normalisation

For within-subjects analysis, we propose data normalisation after filtering to remove SCR scaling differences due to peripheral factors such as skin property. This increases predictive validity (unpublished analyses). For methods that yield trial-by-trial estimates, post-hoc normalisation of amplitude estimates before averaging within conditions yields even better predictive validity (Staib et al., in preparation). For between-subjects analysis, this is not an option as it removes between-subjects variance.

# 2.6 Implicit estimates of tonic sympathetic arousal

Our model for SF implies that

$$\int_{I} SCR(t) - SCL_{0}dt \propto \sum_{i=1}^{n} a_{i}$$

where I is a time interval, and  $a_{1..n}$  are the amplitudes of the n sudomotor bursts in this interval and  $SCL_0$  is the skin conductance level at rest. That is, by simply taking the time integral, or area under the curve (AUC), of the baseline-corrected data, we get a measure of the number x amplitude of spontaneous SN bursts [13]. This measure is implemented in the software as well. However, we have subsequently shown that the number of spontaneous SN bursts is a better predictor of tonic SA than AUC, and hence we recommend to use the non-linear model for tonic SA.

## 2.7 Relation of SCR models to other approaches

Historically, three more approaches to model-based analysis of SCR have been introduced in the literature [19, 20, 21, 22] - one of them is implemented in the software package Ledalab. In this method, SN is calculated from SCR using a deterministic inverse filter. Peaks in the SN time series are then identified and

used as operational index of SA. In a head-to-head comparison, predictive validity to identify states with known SA was compared for experiments involving short stimuli and evoked SA. PsPM had better predictive validity than Ledalab in 4 out of 5 tested contrasts, and equal validity in the fifth. Ledalab did not consistently perform superior to classical peak scoring analysis [2].

# Part II User Guide

# 3 Installation

You can find and download the PsPM package at the following URL:

http://pspm.sourceforge.net/

Decompress and copy the files to any directory of your choice. Then add the PsPM files to the Matlab search path by entering the following command in the Matlab command line:

addpath([your Matlab path]) where [your Matlab path] is a string of the file location on your hard drive, e.g.

addpath('C:\Program Files\MATLAB\R2013a\toolbox\PsPM\').

Do not add subfolders of the PsPM directory to the matlab path.

Type pspm in the command line to start the package. Alternatively, each function can be called individually from the command line. In that case, you can use scr\_init to load the settings, and scr\_quit to remove PsPM subfunctions from the path and delete the settings.

The GUI requires Matlab 2007 (R14) or higher for the graphical interface. If you are using command line functions only, it should be possible to use earlier Matlab versions, but this has not been tested. Display of non-linear model inversion may not work properly under Matlab 2014 due to limitations of an external code.

# 4 Import

Related function: scr\_import

Import external data files for use by PsPM. First, specify the data type. Then, other fields will come up as required for this data type. The imported data will be written to a new .mat file, prepended with 'scr\_'.

Data Type CED Spike 4 IMPORT 19

**Text** Text files can only contain numbers (i.e. no header lines with channel names) and one data column per channel. Make sure you use the decimal point (i.e. not decimal comma as used in some non-English speaking countries). At the moment, no import of event markers is possible

Matlab Each input file must contain a variable called data that is either a cell array of column vectors, or a data points × channels matrix. At the moment, no import of event markers is possible. Data structures containing timestamps cannot be imported at the moment; rather, PsPM will ask you for a sample rate.

Biopac AcqKnowledge (up to v. 3.9)

exported Biopac AcqKnowledge

Labchart exported (up to v. 7.1) Export data to matlab format (plugin for the LabChart software, available from www.adinstruments.com)

Labchart exported (v. 7.2 and higher)

VarioPort

**Biograph Infiniti exported** Export data to text format, both "Export Channel Data" and "Export Interval Data" are supported; a header is required

Mindmedia Biotrace exported

**BrainVision** 

Windaq (wdq) Requires an ActiveX Plugin provided by the manufacturer and contained in the subfolder Import/wdq for your convenience. This plugin only runs under 32 bit Matlab on Windows.

Windaq (PsPM Version) Windaq import written by the PsPM team. Is platform independent, thus has no requirements for ActiveX Plugins, Windows or 32bit Matlab. Imports the \*.wdq files. Up to now the import has been tested with files of the following type: Unpacked, no Hi-Res data, no Multiplexer files. A warning will be produced if the imported data-type fits one of the yet untested cases. If this is the case try to use the import provided by the manufacturer (see above).

Observer XT compatible

NeuroScan

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#### **BioSemi**

**Eyelink** Imports files which have been converted to ascii format (.asc files).

### European Data Format (EDF)

# Data File(s)

Specify which data file(s) to import. You can import multiple files that all have the same structure.

# **New Channel**

Define each channel that you want to import.

### Channel Type

Specify the type of data in this channel. Currently supported data types: SCR, ECG, heart beat markers, interpolated heart rate, interpolated heart period, respiration traces, interpolated respiration period, pupil size, marker channels.

Channel /Column Number Specify where in the original file to find the data - depending on your data type this will be a channel or column number. You can either specify a number (i. e. the n-th channel in the file), or (for many data types) search for this channel by its name. Note: the channel number refers to the n-th recorded channel, not to its number during acquisition: if you did not save all recorded channels, these might be different for some data types.

- Search [if available]: Search for channel by its name this only works if the channel names are unambiguous, and not for all data types.
- Specify Channel Number [if required, otherwise a default channel number will be assigned]: Specify the n-th channel. This counts the number of channels actually recorded.

**Sample Rate [if required]** Sample rate in Hz (i. e. samples per second). For most data types, this is read from the original data file.

**Transfer Function [for SCR data only]** Enter the conversion from recorded data to Microsiemens.

• File: Enter the name of the .mat file that contains the transfer function constants. This file needs to contain the following variables: 'c' is the transfer constant: data = c \* (total conductance in mcS); 'Rs' is the series resistance in Ohm (usually 0), and 'offset' any offset in the data (usually 0).

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• Input: Enter the transfer constants manually.

The relation between measured conductance G to recorded voltage U is given by

$$G = \frac{1}{\frac{c}{U - offset} - R}$$

- 1. Transfer Constant c: Constant by which the measured conductance is multiplied to give the recorded signal (and by which the signal needs to be divided to give the original conductance).
- 2. Offset: Fixed offset (signal at 0 conductance).
- 3. Series Resistor R: Resistance of any resistors in series with the subject.
- None: No transfer function. Use this only if you are not interested in absolute values, and if the recording settings were the same for all subjects.

# 5 Trim

Related function: scr\_trim

Trim away unnecessary data, for example before an experiment started, or after it ended. Trim points can be defined in seconds with respect to start of the data file, in seconds with respect to first and last marker (if markers exist), or in seconds with respect to a user-define marker. The resulting data will be written to a new file, prepended with 't'.

#### Data file

Choose the data file you want to trim.

#### Reference

Choose your reference for trimming: file start, first/last marker, or a user-defined marker. All trimming is defined in seconds after this reference – choose negative values if you want to trim before the reference.

#### File

Trim from xx seconds after file start to xx seconds after file start.

- From (seconds after file start): Choose a positive value.
- Trim To (seconds after file start): Choose a positive value larger than the 'from' value.

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## First/Last Marker

Trim from xx seconds after first marker to xx seconds after last marker.

• From: (seconds after first marker): Choose a positive (after first marker) or negative (before first marker) value.

- To: (seconds after last marker): Choose a positive (after last marker) or negative (before last marker) value.
- Marker Channel: If you have more than one marker channel, choose the reference marker channel (default: use the fist marker channel in the file)
  - Default (first marker channel=0)
  - Number

#### Any Marker

Trim from xx seconds after any marker of your choice to xx seconds after any marker of your choice.

- From: Choose marker number and trimming point in seconds after this marker.
  - Marker Number x: Choose an integer value.
  - Seconds after Marker x: Choose a positive (after this marker) or negative (before this marker) value.
- To: Choose marker number and trimming point in seconds after this marker.
  - Marker Number y: Choose an integer value.
  - Seconds after Marker y: Choose a positive (after this marker) or negative (before this marker) value.
- Marker Channel: If you have more than one marker channel, choose the reference marker channel (default: use the fist marker channel in the file)
  - Default (first marker channel=0)
  - Number

# Overwrite Existing File

Choose "yes" if you want to overwrite existing file(s) with the same name. Default: no.

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# 6 GLM for SCR

 $Related\ function:\ \mathtt{scr\_glm}$ 

General linear convolution models (GLM) are powerful for analysing evoked responses that follow an event with (approximately) fixed latency. This is similar to standard analysis of fMRI data. The user specifies events for different conditions. These are used to estimate the mean response amplitude per condition. These mean amplitudes can later be compared, using the contrast manager. References: [3, 4, 12][2].

# Model Filename & output directory

Specify file name and directory where the \*.mat file with the resulting model will be written.

#### SCR Channel

Indicate the channel containing the SCR data. By default the first SCR channel is assumed to contain the data for this model. If the first SCR channel does not contain the data for this model (e. g. there are two SCR channels), indicate the the channel number (within the SCR file) that contains the data for this model.

#### Time Units

Indicate the time units on which the specification of the conditions will be based. Time units can be specified in 'seconds', number of 'markers', or number of data 'samples'. Time units refer to the beginning of the data file and not to the beginning of the original recordings e. g. if data were trimmed.

#### Option for marker channels:

 Marker Channel: Indicate the marker channel. By default the first marker channel is assumed to contain the relevant markers. Markers are only used if you have specified the time units as 'markers'.

## Data & Design

Add the required number of data files or sessions here. These will be concatenated for analysis.

# Data File

Specify the data file containing the SCR data (and potential marker information). If you have trimmed your data, add the file containing the trimmed data.

# Missing Epochs

Indicate epochs in your data in which the SCR signal is missing or corrupted (e.g., due to artifacts).

- No Missing Epochs: The whole time series will be analyzed.
- Define Missing Epochs: Define the start and end points of the missing epochs either as epoch files or manually. Missing epochs will be excluded from the design matrix. Start and end points have to be defined in seconds starting from the beginning of the session.
  - Missing Epoch File: Indicate an epoch file specifying the start and end points of missing epochs. The mat file has to contain a variable 'epochs', which is an m x 2 array, where m is the number of missing epochs. The first column marks the start points of the epochs that are excluded from the analysis and the second column the end points.
  - Enter Missing Epochs Manually (m: nr. of epochs): Enter the start and end points of missing epochs manually. Specify an m x 2 array, where m is the number of missing epochs. The first column marks the start points of the epochs that are excluded from the analysis and the second column the end points.

## Design

Specify the timing of the events within the design matrix. Timing can be specified in 'seconds', 'markers' or 'samples' with respect to the beginning of the data file. See 'Time Units' settings. Conditions can be specified manually or by using multiple condition files (i.e., an SPM-style mat file).

- Condition File: Create a file with the following variables:
  - names: a cell array of string for the names of the experimental conditions
  - onsets: a cell array of number vectors for the onsets of events for each experimental condition, expressed in seconds, marker numbers, or samples, as specified in time units. If seconds, do use precise values according to the data type (e. g. at least 0.1 s time resolution for SCR)
  - durations (optional, default 0): a cell array of vectors for the duration of each event. You need to use 'seconds' or 'samples' as time units
  - pmod: this is used to specify regressors that specify how responses in an experimental condition depend on a parameter to model the effect e.g. of habituation, reaction times, or stimulus ratings. pmod is a struct array corresponding to names and onsets and containing the fields

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\* name: cell array of names for each parametric modulator for this condition

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- \* param: cell array of vectors for each parameter for this condition, containing as many numbers as there are onsets
- \* poly (optional, default 1): specifies the polynomial degree
- e.g. produce a simple multiple condition file by typing: names =
   {'condition a', 'condition b'}; onsets = {[1 2 3], [4 5 6]};
   save('testfile', 'names', 'onsets');
- Enter conditions manually:
  - Name: Specify the name of the condition.
  - Onsets: Specify a vector of onsets. The length of the vector corresponds to the number of events included in this condition. Onsets have to be indicated in the specified time unit ('seconds', 'markers', or 'samples').
  - Durations: Typically, a duration of 0 is used to model an event onset. If all events included in this condition have the same length specify just a single number. If events have different durations, specify a vector with the same length as the vector for onsets.
  - Parametric Modulator(s) [optional] (n: nr. of pmods): If you want to include a parametric modulator, specify a vector with the same length as the vector for onsets. For example, parametric modulators can model reaction times, ratings of stimuli, or habituation effects over time. For each parametric modulator a new regressor is included in the design matrix. The normalized parameters are multiplied with the respective onset regressors.
    - 1. Name: Specify the name of the parametric modulator.
    - 2. Polynomial Degree: Specify an exponent that is applied to the parametric modulator. A value of 1 leaves the parametric modulator unchanged and thus corresponds to a linear change over the values of the parametric modulator (first-order). Higher order modulation introduces further columns that contain the non-linear parametric modulators [e.g., second-order: (squared), third-order (cubed), etc].
    - Parameter Values: Specify a vector with the same length as the vector for onsets.

#### Nuisance File (optional)

You can include nuisance parameters such as motion parameters in the design matrix. Nuisance parameters are not convolved with the canonical response function. The file has to be either a .txt file containing the regressors in columns, or a .mat file containing the regressors in a matrix variable called R. There must be as many values for each column of R as there are data values in your data file. PsPM will call the regressors pertaining to the different columns R1, R2,

. . .

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# **Basis Function**

Basis functions for the peripheral model. Standard is to use a canonical skin conductance response function (SCRF) with time derivative for later reconstruction of the response peak.

#### SCRF0

SCRF without derivatives.

#### SCRF1

SCRF with time derivative (default). This is the recommended option based on the publication [2].

#### SCRF2

SCRF with time and dispersion derivative.

#### FIR

Uninformed finite impulse response (FIR) model: specify the number and duration of time bins to be estimated.

- Arguments
  - N: Number of Time Bins: Number of time bins.
  - D: Duration of Time Bins: Duration of time bins (in seconds).

## Normalize

Specify if you want to z-normalize the SCR data for each subject. For withinsubjects designs, this is highly recommended, but for between-subjects designs it needs to be set to false.

# Filter Settings

Specify how you want filter the SCR data.

## Default

Standard settings for the Butterworth bandpass filter. These are the optimal settings from [2]: unidirectional Butterworth band pass filter with cut off frequencies of 0.05 and 5 Hz, downsampling to 5 Hz.

## **Edit Settings**

Create your own filter settings (not encouraged).

- Low-Pass Filter
  - Enable
    - 1. Cutoff Frequency: Specify the low-pass filter cutoff in Hz.
    - 2. Filter Order: Specify the low-pass filter order.
  - Disable
- High-Pass Filter
  - Enable
    - 1. Cutoff Frequency: Specify the high-pass filter cutoff in Hz.
    - 2. Filter Order: Specify the high-pass filter order.
  - Disable
- New Sampling Rate: Specify the sampling rate in Hz to down sample SCR data. Enter NaN to leave the sampling rate unchanged.
- Filter Direction {uni, bi}, (default: uni): A unidirectional filter is applied twice in the forward direction. A 'bidirectional' filter is applied once in the forward direction and once in the backward direction to correct the temporal shift due to filtering in forward direction.

# Overwrite Existing File

Specify whether you want to overwrite existing mat files.

# 7 Non-Linear Model for SCR

Related function: scr\_dcm

Non-linear models for SCR are powerful if response timing is not precisely known and has to be estimated. A typical example are anticipatory SCR in fear conditioning – they must occur at some point within a time-window of several seconds duration, but that time point may vary over trials. Dynamic causal modelling (DCM) is the framework for parameter estimation. PsPM implements an iterative trial-by-trial algorithm. Different from GLM, response parameters are estimated per trial, not per condition, and the algorithm must not be informed about the condition. Trial-by-trial response parameters can later be summarized across trials, and compared between conditions, using the contrast manager. References: [14], [Staib et al., submitted]

# Model Filename & Output directory

Specify file name for the resulting model. Specify directory where the mat file with the resulting model will be written.

#### SCR channel

Indicate the channel containing the SCR data. By default the first SCR channel is assumed to contain the data for this model. If the first SCR channel does not contain the data for this model (e. g. there are two SCR channels), indicate the channel number (within the SCR file) that contains the data for this model.

# Data & design

Add the appropriate number of data files or sessions here. Results will be concatenated.

#### Data File

Add the data file containing the SCR data. If you have trimmed your data, add the file containing the trimmed data.

## Design

Specify the timings of individual events from your design either by creating a timing file or by entering the timings manually. The DCM framework allows you to specify two types of events:

Fixed latency (evoked): An event is assumed to elicit an immediate response. The amplitude of the sympathetic arousal will be estimated, while the latency and duration of the response are fixed. This event type is meant to model evoked responses.

Flexible latency and duration (event-related): An event is assumed to elicit sympathetic arousal within a known response window, but with unknown amplitude, latency and duration. For each event of this type, specify a time window in which the response is expected. PsPM will then estimate the amplitude, duration and latency of the response. An example for this type of event is an anticipatory response which might vary in timing between trials (e. g. in fear conditioning).

• Timing File: The timing file has to be a .mat file containing a cell array called 'events'. For a design with n trials and m events per trial (n × m events in total), the cell array has to be structured in the following way: Create m cells in the cell array (one per event type that occurs in each trial). Each cell defines either a fixed or a flexible event type. A cell that defines a fixed event type has to contain a vector with n entries, i.e. one time point per trial, in seconds, samples or markers. A cell that defines a flexible type has to contain an array with n rows and two columns. The

first column specifies the onsets of the time windows for each trial, while the second column specifies the offsets of the time windows. It is assumed that all trials have the same structure, i.e. the same number of fixed and flexible event types. For individual trials with a different structure you can enter negative values as time information to omit estimation of a response. For later comparison between trials of different conditions, it is absolutely mandatory that they contain the same types of events, to avoid bias. Hence, if one condition omits an event (e. g. unreinforced trials in conditioning experiments), the omitted event needs to be modelled as well.

- Enter Timing Manually: In the DCM framework, a session is structured in n individual trials. Each trial contains m fixed and/or flexible event types. All trials need to have the same structure. Create event types to define the structure of a trial and enter the timings for all events.
  - Name (optional): Enter a name of the event. This name can later be used for display and export.
  - Onsets: For events with a fixed response, specify a vector of onsets. The length of the vector corresponds to the number of trials (n). For events with a flexible response, specify a two column array. The first column defines the onset of the time window in which the response occurs. The second column defines the offset. The number of rows of the array corresponds to the number of trials (n). All timings are specified in seconds.

#### Condition names (optional)

Specify the conditions that the individual trials belong to. This information is not used for the parameter estimation, but it allows you to later access the conditions in the contrast manager.

- Name: Specify the name of the condition.
- Index: Specify a vector of trial indices between 1 and n. The length of the vector corresponds to the number of events included in this condition.

# **Data Options**

# Normalization

Specify if you want to normalize the SCR data for each subject. For withinsubjects designs, this is highly recommended. During analysis, data are always z-transformed, but the parameter estimates are transformed back if this is set to false.

# Filter Settings

Specify if you want filter the SCR data.

- Default: Standard settings for the Butterworth bandpass filter. These are the optimal settings from [Staib et al., submitted]: bidirectional Butterworth band pass filter with cutoff frequencies of 0.0159 Hz and 5 Hz, downsampling to 10 Hz.
- Edit Settings: Create your own filter (discouraged).
  - Low-Pass Filter
    - 1. Enable
      - Cutoff Frequency: Specify the low-pass filter cutoff in Hz.
      - Filter Order: Specify the low-pass filter order.
    - 2. Disable
  - High-Pass Filter
    - 1. Enable
      - Cutoff Frequency: Specify the high-pass filter cutoff in Hz.
      - Filter Order: Specify the high-pass filter order.
    - 2. Disable
  - -New Sampling Rate: Specify the sampling rate in Hz to down sample SCR data. Enter NaN to leave the sampling rate unchanged.
  - -Filter Direction {uni, bi}, (default: bi): A unidirectional filter is applied twice in the forward direction. A 'bidirectional' filter is applied once in the forward direction and once in the backward direction to correct the temporal shift due to filtering in forward direction.

# Response Function Options

#### Estimate the Response Function from The Data

A response function can be estimated from the data and used instead of the canonical response function (default 0 for fully flexible and 1 for fix and flex/fix paradigms). This is not normally recommended unless you have long inter trial intervals in the range of 20-30 s [Staib et al., submitted].

# Only Estimate RF (Do Not Do Trial-Wise DCM)

This option can be used to estimate an individual response function to be used in analysis of another experiment.

#### Call External File to Provide Response Function

Call an external file to provide a response function, which was previously estimated using the option "only estimate RF".

# **Inversion Options**

### Number of Trials to Invert at The Same Time (default: 2 s)

The iterative DCM algorithm accounts for response overlap by considering several trials at a time. This can be set here.

### SF-Free Window Before First Event [s] (default: 2 s)

The DCM algorithm automatically models spontaneous fluctuations in inter trial intervals. Here, you can define a time window before the first event in every trial in which no spontaneous fluctuation will be estimated.

## SF-Free Window After Last Event [s] (default: 5 s)

The DCM algorithm automatically models spontaneous fluctuations in inter trial intervals. Here, you can define a time window after the last event in every trial in which no spontaneous fluctuation will be estimated.

### Maximum Frequency of SF in ITIs [Hz] (default: 0.5 Hz)

The DCM algorithm automatically models spontaneous fluctuations in intertrial intervals. Here you can define the minimal delay between two subsequent spontaneous fluctuations.

#### SCL-Change-Free Window Before First Event [s] (default: 2 s)

The DCM algorithm automatically models baseline drifts in inter trial intervals. Here, you can define a window before the first event in every trial in which no change of the skin conductance level will be assumed.

#### SCL-Change-Free Window After Last Event [s] (default: 5 s)

The DCM algorithm automatically models baseline drifts in inter trial intervals. Here, you can define a window after the last event in every trial in which no change of the skin conductance level will be assumed.

# **Display Options**

#### Display Progress Window

Show a on-line diagnostic plot for each iteration of the estimation process.

## Display Intermediate Windows

Show small plots displaying the progress of each iteration in the estimation process.

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# 8 SF models

Related function: scr\_sf

This suite of models is designed for analysing spontaneous fluctuations (SF) in skin conductance as a marker for tonic arousal. SF are analysed over time windows that typically last 60 s and should at least be 15 s long. PsPM implements 3 different models: (1) Skin conductance level (SCL): this is the mean signal over the epoch (2) Area under the curve (AUC): this is the time-integral of the above-minimum signal, divided by epoch duration. This is designed to be independent from SCL and ideally represents the number x amplitude of SF in this time window. (3) Number of SF estimated by DCM (Dynamic Causal Modelling) or MP (Matching Pursuit): this is a non-linear estimation of the number of SF, and is the most sensitive indicator of tonic arousal. It relies on absolute data values as it implements and absolute threshold for data peaks. References: [17][18]

# Data File

Add the data file containing the SCR data (and potential marker information). If you have trimmed your data, add the file containing the trimmed data.

# Model Filename & Output directory

Specify file name for the resulting model. Specify directory where the mat file with the resulting model will be written.

#### Method

Choose the method for estimating tonic sympathetic arousal: AUC (equivalent to number x amplitude of spontaneous fluctuations), SCL (tonic skin conductance level), DCM or MP. The latter two estimate the number of spontaneous fluctuations, requiring absolute data units as both methods implement an absolute amplitude threshold. In theory, DCM provides highest sensitivity but is slow[17]. MP is a very fast approximation to the DCM results, and comparable in sensitivity for analysis of empirical data [18].

#### Time Units

Indicate the time units on which the specification of the conditions will be based. Time units can be specified in 'seconds', number of 'markers', or number of data 'samples'. Time units refer to the beginning of the data file and not to the beginning of the original recordings e. g. if data were trimmed. Choose 'seconds', 'markers', 'samples', or 'whole' to analyse the entire data file as one epoch.

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# Seconds/Markers/Samples

- Epochs define data epochs to analyse
  - Epoch File
  - Enter Epochs Manually

#### Markers

• Marker Channel (default: 0): Indicate the marker channel. By default the first marker channel is assumed to contain the relevant markers. Markers are only used if you have specified the time units as 'markers'.

# Filter Settings

Specify if you want filter the SCR data.

#### Default

Standard settings for the Butterworth bandpass filter: unidirectional Butterworth bandpass filter with cut off frequencies of  $0.0159~\mathrm{Hz}$  and  $5~\mathrm{Hz}$ , down sampling to  $10~\mathrm{Hz}$ .

#### **Edit Settings**

Create your own filter (discouraged).

- Low-Pass Filter
  - Enable
    - 1. Cutoff Frequency: Specify the low-pass filter cutoff in Hz.
    - 2. Filter Order: Specify the low-pass filter order.
  - Disable
- High-Pass Filter
  - Enable
    - 1. Cutoff Frequency: Specify the high-pass filter cutoff in Hz.
    - 2. Filter Order: Specify the high-pass filter order.
  - Disable
- New Sampling Rate: Specify the sampling rate in Hz to down sample SCR data. Enter NaN to leave the sampling rate unchanged.
- Filter Direction): A unidirectional filter is applied twice in the forward direction. A 'bidirectional' filter is applied once in the forward direction and once in the backward direction to correct the temporal shift due to filtering in forward direction.

## Channel

Indicate the channel containing the SCR data. By default the first SCR channel is assumed to contain the data for this model. If the first SCR channel does not contain the data for this model (e. g. there are two SCR channels), indicate the channel number (within the SCR file) that contains the data for this model.

# Overwrite Existing File

Specify whether you want to overwrite existing mat file.

# Threshold (used for DCM and MP only)

Threshold for SN detection - default  $0.1~\mu S$ .

# Display Progress Window (used for DCM and MP only)

Show a on-line diagnostic plot for each iteration of the estimation process (DCM) or for the result of the estimation process (MP).

# Display Intermediate Windows (used for DCM only)

Show small plots displaying the progress of each iteration in the estimation process.

# 9 Review First-Level Model (via Batch editor)

Related function: scr\_review

This module allows you to look at the first-level (within-subject) model to investigate model fit and potential estimation problems. This is not necessary for standard analyses. Further processing can be performed directly on the second level after first-level model estimation.

#### Model File

Choose model file to review.

# Model Type

Specify the type of model.

## GLM

Specify the plot that you wish to display:

• Design matrix

- Orthogonality
- Predicted & Observed
- Regressor names
- Reconstructed

#### DCM

- Inversion results for one trial: Non-linear SCR model (DCM) Review individual trials or sequences of trials inverted at the same time.
  - Session Number (Default 1): Data session. Must be 1 if there is only one session in the file.
  - Trial Number: Trial to review.
- Predicted & Observed for all trials: DCM for event-related responses Review summary plot of all trials.
  - Session Number
  - Figure Name [optional]: Adding a figure name saves the figure as .fig file.
- SCRF: DCM for event-related responses review SCRF (useful if SCRF was estimated from the data).

#### $\mathbf{SF}$

DCM for spontaneous fluctuations: Show inversion results for one episode

• Episode Number: Episode to review.

#### Contrasts

Display contrast names for any first level model.

# 10 First-Level Model Review Manager (via GUI)

Related function: scr\_review

Display information stored in a model file through an easily accessible interface. Add a model file to the menu via Add Models. Available content of a highlighted model is listed on the right hand side under Figure Selection. For non-linear models, you can select the session number by entering its index in the field Session number; GLMs are inverted and displayed across all sessions. Click on any Display or Show button to visualize information. Details for each model type are listed in Chapter 9.

# 11 First-Level Contrasts (via Batch editor)

Related function: scr\_con1

Define within-subject contrasts here for testing on the second level. Contrasts can be between conditions (GLM), epochs (SF) or trials (Non-linear SCR models). Contrast weights should add up to 0 (for testing differences between conditions/epochs/trials) or to 1 (for testing global/intercept effects across conditions/epochs/trials). Example: an experiment realises a 2 (Factor 1: a, A) x 2 (Factor 2: b, B) factorial design and consists of 4 conditions: aa, aB, Ab, AB. Testing the following contrasts is equivalent to a full ANOVA model: Main effect factor 1: [1 1 -1 -1], Main effect factor 2: [1 -1 1-1], Interaction factor 1 x factor 2: [1 -1 -1 1]. To test condition effects in non-linear models, assign the same contrast weight to all trials from the same condition.

# Model File(s)

Specify the model file for which to compute contrasts.

# Specify Contrasts on Datatype

Contrasts are usually specified on parameter estimates. For GLM, you can also choose to specify them on conditions or reconstructed responses per condition. In this case, your contrast vector needs to take into account only the first basis function. For DCM, you can specify contrasts based on conditions as well. This will average within conditions. Use the review manager to extract condition names and their order. This argument will be ignored for other first-level models.

- Parameter: Use all parameter estimates.
- Condition: Contrasts formulated in terms of conditions in a GLM, automatically detects number of basis functions and uses only the first one (i.e. without derivatives), or based on assignment of trials to conditions in DCM.
- Reconstructed: Contrasts formulated in terms of conditions in a GLM, reconstructs estimated response from all basis functions and uses the peak of the estimated response.

# Contrast(s)

# Contrast Name

This name identifies the contrast in tables and displays.

#### **Contrast Vector**

This is a vector on all included conditions (GLM), trials (DCM for event-related responses), or epochs (DCM for SF). Shorter vectors will be appended with zeros. To specify a condition or trial difference, the contrast weights must add up to zero (e. g. was sympathetic arousal in condition A larger than in condition B: c = [1 - 1]). To specify a summary of conditions or trials, the contrast weights

should add up to 1 to retain proper scaling (e. g. was non-zero sympathetic arousal elicited in combined conditions A and B:  $c = [0.5\ 0.5]$ ).

#### **Delete Existing Contrasts**

Deletes existing contrasts from the model file.

# 12 First level contrast manager (via GUI)

Related function: scr\_con1

The Manager guides you through the setup of pre-defined within-subject contrasts applied to the parameter estimates of your model.

#### Load Model

Import a model file from a subject containing parameter estimates for which to compute contrasts.

#### Define Contrast Name

Enter a name to identify a contrast. By pressing *New Contrast*, the contrast is added to the box *Contrasts*. You can reset existing contrasts by highlighting a contrast and pressing *Clear Contrast* or you can remove contrasts from your model file by pressing *Delete Contrast*.

#### Define Stats Type

Depending on the model (GLM, spontaneous fluctuation or non-linear model), different data types are available (see Chapter 11 for details). Choose the type you want to be entered into a statistical test.

#### Define Test

Three pre-defined types of statistics can be computed on your estimates. After choosing the test, you can add or remove conditions/trials to a group by clicking on them in the *Names* window. The color will indicate to which group they belong.

- Test of intercept: Compute the average of conditions entered in *Group*1. This option is suitable if you are interested in the main effect of a condition.
- Test of condition differences: Compute the difference between the averages of *Group 1* and *Group 2*.
- Test of quadratic effects: Here you compute the combined effect of *Group 1* and *Group 3* against *Group 2*. Choose this option if you have three experimental conditions.

#### Run

Execute computation of statistics. The contrasts are stored in your model files and can be reviewed any time using the Review Manager (Chapter 10).

# 13 Export Statistics

Related function: scr\_exp

Export first level statistics to a file for further analysis in statistical software, or to the screen.

# Model File(s)

Specify file from which to export statistics.

# Stats type

Normally, all parameter estimates are exported. For GLM, you can choose to only export the first basis function per condition, or the reconstructed response per condition. For DCM, you can specify contrasts based on conditions as well. This will average within conditions. This argument cannot be used for other models.

- Parameter: Export all parameter estimates.
- Condition: Export all conditions in a GLM, automatically detects number of basis functions and export only the first one (i.e. without derivatives), or export condition averages in DCM.
- Reconstructed: Export all conditions in a GLM, reconstructs estimated response from all basis functions and export the peak of the estimated response.

#### Target

Export to screen or to file?

# Delimiter for Output File

Select a delimiter for the output file. Default is 'tab', you can select from a choice of several options or specify as free text.

#### 14 Define Second-Level Model

Related function: scr\_con2

Define one-sample and two-sample t-tests on the between-subject (second) level. A one-sample t-test is normally used to test within-subject contrasts and is equivalent to t-contrasts in an ANOVA model A two-sample t-test is required for between-subjects contrasts or interactions. This module sets up the model but does not report results.

# Test Type

Specify the test type.

# One Sample T-Test

• Model File(s)

## Two Sample T-Test

- Model File(s) 1: Model files for group 1.
- Model File(s) 2: Model files for group 2.

# Output directory

Specify directory where the mat file with the resulting model will be written.

# Filename for Output Model

Specify name for the resulting model.

#### **Define Contrasts**

Define which contrasts to use from the model files, and the second level contrast names. Names can be read from the first model file, or just be numbered.

#### Read from First Model File

- All Contrasts
- Contrast Vector: Index of first-level contrasts to use.

#### **Number Contrasts**

- All Contrasts
- Contrast Vector: Index of first-level contrasts to use.

# Overwrite Existing File

Specify whether you want to overwrite existing mat file.

# 15 Report Second-Level Results

 $Related\ function:\ \mathtt{scr\_rev2}$ 

Result reporting for second level model.

#### Model File

Specify 2nd level model file.

# Contrast Vector (only accessible from Batch Editor)

Index of contrasts to be reported (optional).

# 16 Display Data

Related function: scr\_display

Display PsPM data file in a new figure.

#### Data File

Specify data file to display.

## 17 Rename File

 $Related\ function:\ \mathtt{scr\_ren}$ 

Rename PsPM data file. This renames the file and updates the file information.

#### File

Choose how many files to rename.

#### File Name

Choose name of original file.

#### New File Name

Choose new file name.

# 18 Split Sessions

Related function: scr\_split\_sessions

Split sessions, defined by trains of markers. This function is most commonly used to split fMRI sessions when a (slice or volume) pulse from the MRI scanner has been recorded. The function will identify trains of markers and detect breaks in these marker sequences. The individual sessions will be written to new files with a suffix '\_sn', and the session number. You can choose a marker channel if several were recorded.

#### Data File

Choose the data file, in which you want to split sessions.

## Marker Channel

If you have more than one marker channel, choose the marker channel used for splitting sessions (default: use first marker channel).

- Default
- Number

## Overwrite Existing File

Choose "yes" if you want to overwrite existing file(s) with the same name.

## 19 Artifact Removal

Related function: scr\_pp

This module offers a few basic artifact removal functions. Currently, a median filter and a Butterworth low pass filter are implemented. The median filter is useful to remove short "spikes" in the data, for example from gradient switching in MRI. The Butterworth filter can be used to get rid of high frequency noise that is not sufficiently filtered away by the filters implemented on-the-fly during first level modelling.

#### Data File

Choose the data file.

#### Channel Number

Select the channel to work on.

#### Filter Type

Currently, median and Butterworth filters are implemented. A median filter is recommended for short spikes, generated for example in MRI scanners by gradient switching. A Butterworth filter is applied in most models; check there to see whether an additional filtering is meaningful.

#### Median Filter

• Number of Time Points: Number of time points over which the median is taken.

#### Butterworth Low pass Filter

• Cutoff Frequency: Cutoff frequency has to be at least 20Hz.

#### Overwrite Existing File

Choose "yes" if you want to overwrite existing file(s) with the same name.

# 20 Downsample Data

Related function: scr\_down

This function downsamples individual channels in an PsPM file to a required sampling rate, after applying an anti-aliasing Butterworth filter at the Nyquist frequency. The resulting data will be written to a new .mat file, prepended with 'd', and will contain all channels – also the ones that were not downsampled.

#### Data File

Name of the data files to be downsampled.

#### New Frequency

Required sampling frequency.

#### Channels To Downsample

Vector of channels to downsample, or all channels.

#### Overwrite Existing File

Choose "yes" if you want to overwrite existing file(s) with the same name.

# 21 Convert ECG to Heart Beat

Related function: scr\_ecg2hb

Detect QRS complexes in ECG data and write timestamps of detected R spikes into a new heart beat channel. This function uses an algorithm by Pan & Tompkins adapted from [23]. Hidden options for minimum and maximum heart rate become visible when the job is saved as a script and should only be used if the algorithm fails.

#### Data File

Specify data file. The detected heart beat data will be written to a new channel in this file.

#### Channel

Number of heart beat channel (default: first heart beat channel).

# 22 Convert Heart Beat to Heart Period

Related function: scr\_hb2hp

Interpolate heart beat time stamps into continuous heart period data and write into new channel. This function uses heart period rather than heart rate because heart period varies linearly with ANS input into the heart.

#### Data File

Specify data file. The interpolated heart period data will be written to a new channel in this file.

#### Sample Rate

Sample rate for heart period channel. Default: 10 Hz.

#### Channel

Number of heart beat channel (default: first heart beat channel).

# 23 Preprocess respiration data

Related function: scr\_resp\_pp

Convert a continuous respiration channel into various respiration data types. The function detects respiration cycles for bellows and cushion systems, computes respiration period, amplitude and RLL, assigns these measures to the start of each cycle and linearly interpolates these (expect for rs [= respiration time stamps]). The results are written to new channels in the same file.

#### Data File

Specify data file. The processed respiration data will be written to a new channel in this file.

#### Sample Rate

Sample rate for the new channel. Default: 10 Hz. Will be ignored for datatype "respiration time stamps".

#### Channel

Number of respiration channel (default: first respiration channel).

## **Options**

#### Systemtype

Type of the measuring system: bellows or cushion (default: bellows).

#### **Datatype**

Choose for each possible process datatype either yes or no (default: yes):

- Respiration period Create a channel with interpolated respiration period
- Respiration amplitude Create a channel with interpolated respiration amplitude
- Respiration line length Create a channel with interpolated respiration line length
- Respiration time Create a channel with respiration time stamp

#### Diagnostic plot

Specify whether a respiratory cycle detection plot should be created (Yes) or not (No) (default: No).

# 24 Troubleshooting and known problems

#### 24.1 Path

If the PsPM folder is added to the path with all subfolders, the batch editor will not work. Only put the PsPM main folder (containing the function pspm.m) to the path.

#### 24.2 Diagnostic graphics for non-linear model inversion

During non-linear model inversion, a diagnostic window is displayed for each trial. Between two trials, there is a short period of time during which several tabs appear in that window. If you click on one of the hidden tabs, model inversion will fail due to a display error.

Diagnostic plots do not work in Matlab versions 2014 and upwards. This is due to a limitation of an external software. We are working on a fix.

# 24.3 Export statistics into text format with tab delimiter

The tab delimiter in the exported files is not correctly interpreted by text editors like MS Word. Nonetheless, the data can be imported into softwares such as Excel or SPSS.

# Part III Tutorial

Contributed by Christoph Korn & Matthias Staib.

# 25 Appraisal data

Here, we analyze SCR data using a general linear model (GLM). The example data set comprises SCR data from 15 participants and can be downloaded from pspm.sourceforge.net (see: sample data). Results from this data set have been previously published [12, 2]. Each participant saw 45 neutral and 45 aversive pictures within one session that included two short breaks. SCR data were recorded using a 0.5 V coupler, optical (wave to pulse) transducer, and CED Spike, with a minimum sampling rate of 100 Hz.

# 25.1 Import

We import the SCR data and the corresponding marker data. We first import data from one subject.

- In the batch editor go to  $PsPM \rightarrow Data\ Preparation \rightarrow Import.$
- Specify the Data Type. Several data formats are available. The current data set is in the smr format.
- Select the SCR Data File of the first subject trsp\_1\_25.smr.
- Specify the Channels. We need one SCR channel and one marker channel.
  - The Channel Number of the SCR channel is 2.
  - For the Transfer Function choose None.
  - For the *Channel Number* of the marker channel put in *Search* to test the automatic search (otherwise you can specify the number of the marker channel manually. Here, the marker channel is 0).

See Figure 25.1 for a picture of the final batch editor. We recommend saving the batch and script so that you can easily re-run your analyses at a later time point (see Chapter 27). Run the batch. A mat file with the name scr\_trsp\_1\_25.mat will be created in the same folder as the original data file.

You can have a look at the imported data by loading this file into MATLAB. The cell array data contains two structures. The first one contains the SCR data and the second one the marker data (this data set contains 96 markers).

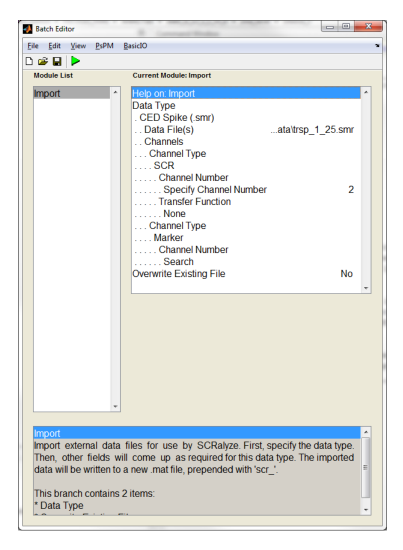


Figure 25.1: Batch editor for importing raw SCR data.

## 25.2 Trim

Most of the time, the SCR recording is switched on some time before the actual experiment starts and is switched off some time after it ends. These parts of the time series may contain large artifacts from subject motion, setting up the equipment, etc. Trimming excludes these parts from the analyses.

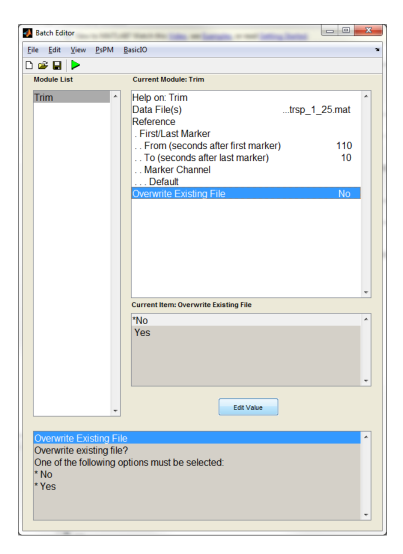


Figure 25.2: Batch editor for trimming SCR data.

- In the batch editor go to  $PsPM \rightarrow Data\ Preparation \rightarrow Trim.$ 
  - Select the currently imported file as Data File. Alternatively, you can use the *Dependency* button.
- $\bullet\,$  You can choose different References to specify how data should be trimmed. Here, select First/Last~Marker and
  - From (seconds after first marker): +110 (In the current data set the first seconds are not task-relevant and are thus trimmed. Note that a positive number will delete at least the first marker of the original

data. Thus, make sure that a possible specification of the First-Level takes this into account.)

- To (seconds after first marker): 10 (The specified time window should be large enough to completely include the last trial.)

You can also specify time points before the markers by using negative numbers (see above). See Figure 25.2 for a picture of the final batch editor. Again, it is recommended to save the batch and script for future use. Run the batch. A mat file with the name tscr\_trsp\_1\_25.mat will be created (i.e., a "t" will be prepended).

#### 25.3 First-Level

Now, we specify the GLM on the first (i.e., the subject-specific) level, estimating average responses per experimental condition (this equivalent to averaging responses over trials in classical "peak scoring" analysis, or to performing a first level analysis in SPM).

- In the batch editor go to  $PsPM \to First\ Level \to SCR \to GLM$ .
- *Model Filename*: This is the name of the resulting mat file containing the GLM. Here, we call it glm\_25.
- Output Directory: The resulting mat file containing the GLM will be saved in this folder.
- *Time Units* on which the specification of the conditions will be based. Here, we want to analyze with respect to Markers.
- Session(s): Similar to SPM, SCRalyze allows the specification of multiple sessions per subject. Here, we only have one session.
  - Add the trimmed data (tscr\_trsp\_1\_25.mat) as Data File. Alternatively, you can use the *Dependency* button.
  - Missing Epochs allows the specification of time periods which you do not want to include in the analysis for example because they contain corrupted data. In the example data set, there are no missing epochs.
  - Data & Design: This option allows specifying the different conditions of the experiment. You need to provide the information when the different conditions started. This time information has to be in the unit specified above (see Time Units). In the current data set, condition information is provided in Markers and is stored in the mat file trSP\_appraisal\_01\_25.mat, which you should select as Condition File. When you open this file in MATLAB, you will see that the file contains a 1x3 cell "names" and a 1x3 cell "onsets" specifying three conditions. The numbers in "onsets" correspond to the nth marker when the respective trial started.

 Nuisance File allows the inclusion of nuisance confounds such as motion or respiration parameters. Here, we leave it empty.

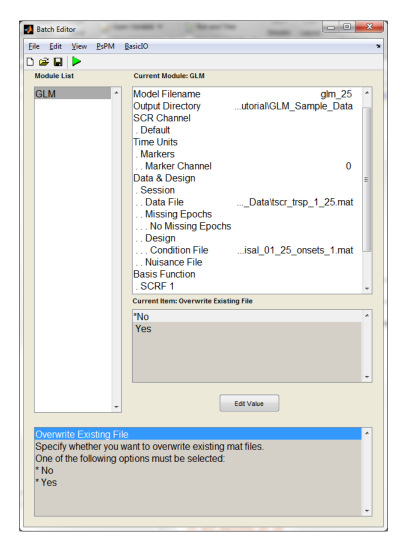


Figure 25.3: Batch editor for First-Level analysis.

- Basis Function: We chose the default SCRF 1, which is the version with time derivative. The default basis set follows the recommendations by [12].
- *Normalize*: The design of the current data set is "within-subjects." Therefore, we want to z-normalize the data.

- Filter Settings: We choose the default filter settings, which follow the recommendations by [12]:
  - first-order low-pass Butterworth filter with a cutoff frequency of 5 Hz
  - first-order high-pass Butterworth filter with a cutoff frequency of 0.05  $\,\mathrm{Hz}$
  - Data are resampled to 10 Hz.
  - The filter direction is unidirectional.

See Figure 25.3 for a picture of the final batch editor. Again, we recommend saving the batch and script for future use. Run the batch. A mat file with the specified name will be created in the specified folder.

#### 25.4 Review First-Level Model

You can have a look at plots related to the first-level model in the batch editor.

- In the batch editor go to  $PsPM \rightarrow First\ Level \rightarrow Review\ First-level\ Model.$
- Specify the Model file that you want to review. Select the mat file containing the GLM (e.g., glm\_25.mat).
- The *Model Type* is of course GLM in the current example. You can set the following options to get different types of information:
  - 1. Design matrix: On the x-axis, you see the number of regressors in the model. Here we have specified 7 regressors in total. On the y-axis, you see the time course of the experiment. Note that there are 7 regressors because the specification for the Basis Function was SCRF with time derivative. That is, 3 (conditions times) x 2 (basis functions) + 1 (constant) = 7 regressors. You can see that neutral and aversive are intermixed and fall into three blocks. See Figure 25.4.
  - 2. Orthogonality: On the x- and y-axes you see the specified regressors. Regressor names are additionally specified along the y-axes. The plot depicts the collinearity between regressors. See Figure 25.5.
  - 3. Predicted & Observed: You see the model fit including the observed response over the whole time series, which you can visually compare to the predicted time series by the specified model. Note that the predicted data will never follow the observed data very closely, because we estimated the average response per condition which will deviate from the response in each individual trial. See Figure 25.6.
  - 4. Regressor names: The names of the regressors as specified in the Conditions File under Data & Design are written to the MATLAB command window. The labels bf 1 and bf 2 refer to the two used basis functions (SCRF with time derivative).

5. Reconstructed: Here you see the estimated responses per condition for the first basis function. These are estimates of the true responses which contain noise. The noise term can be negative but if all estimated responses across an entire group are negative it is a good idea to look for confounds or correlations in the design. See Figure 25.7.

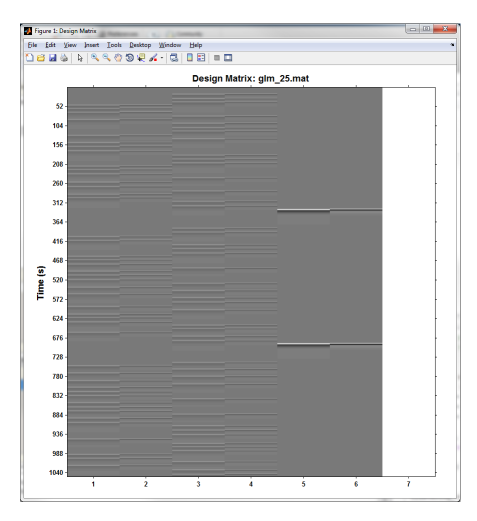


Figure 25.4: Design matrix for sample participant. On the x-axis, you see the number of regressors in the model. There are 7 regressors (i.e., 3 (conditions times)  $\times$  2 (basis functions)  $\times$  1 (constant)).

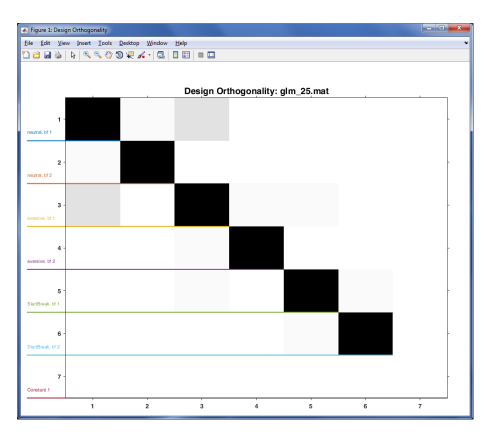


Figure 25.5: Design orthogonality plot for sample participant. On the x- and y-axes you see the 7specified regressors.

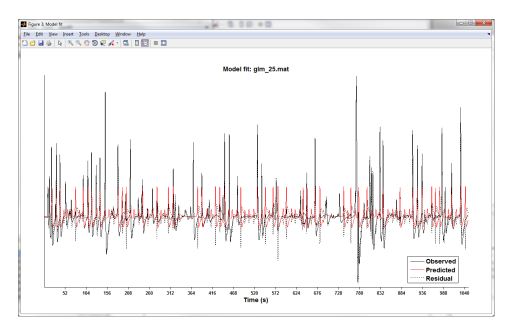


Figure 25.6: Model fit for sample participant. You see the observed response and the predicted response according to the model over the whole time series.

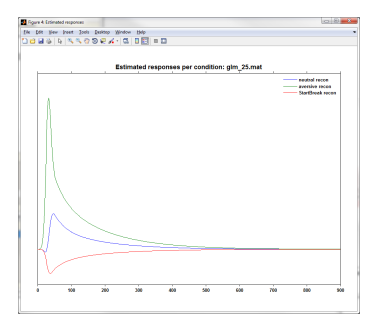


Figure 25.7: Estimated responses per condition for sample participant. You see the estimated responses per condition for the first basis function.

# 25.5 First-Level Contrast Manager

We want to compare the difference between aversive and neutral pictures across participants (i.e., on the second level). This can be tested in a paired t-test. A paired t-test is essentially a one-sample t-test on difference values between two conditions. The contrast manager allows you to compute such difference values, for each participant (i.e., on the first level).

- In the batch editor go to  $PsPM \rightarrow First-Level \rightarrow First-Level Contrasts$ .
- *Model File(s)*: Select the mat file containing the GLM (here glm\_25.mat).
- Stats type: You can choose between 3 different options to set up the contrasts. Here, we choose the option Condition, which specifies contrasts formulated in terms of conditions and automatically uses only the first basis function (i.e., without derivatives).
- Contrast
  - Contrast Name: Chose a name so that you can easily refer to the contrast (e.g., aversive>neutral)
  - Contrast Vector: Specify a vector of the contrast you want to test for. Here, we are interested in the difference between aversive and neutral pictures, i.e., the first and the second condition. Therefore, specify the vector [-1, 1, 0]. You could also leave out the last zero because trailing zeros are automatically appended. (But you could not leave out the first zero in a contrast such as [0 -1 1].)

#### 25.6 Export Statistics

You can use this functionality to export all necessary parameters on the screen or into an extra file, which allows you to analyze the data in the statistics program of your choice. For details see  $PsPM \to First\ Level \to Export\ Statistics$ .

# 25.7 Adapting scripts for multiple participants and secondlevel analyses

For didactic purposes, we have so far only considered one participant. But in most experiments, you want to analyze data from multiple subjects (i.e., a second-level analysis). For a description on how to do this see Chapters 27 and 28. You can find a script that runs the above described analyses for multiple participants at pspm.sourceforge.net (import\_trim\_glm\_contr\_job\_loop).

# 26 Delay fear conditioning data

The example data set comprises SCR data from 20 participants, with 40 trials each, which can be downloaded from pspm.sourceforge.net together with the

result files of the model inversion. Participants underwent a differential delay fear conditioning paradigm where coloured circles were presented for 4 seconds each. Either the blue or orange coloured circles (balanced over participants) were probabilistically paired with an electric stimulation (unconditioned stimulus, US) that coterminated with the stimulus (conditioned stimulus, CS+) while the CS- were always presented alone.

Presenting the CS+ causes phasic sympathetic arousal in anticipation of an electric shock. This anticipatory reaction typically occurs with an unknown and variable latency after stimulus onset. For such cases, a GLM should not be used because it assumes that sympathetic arousal follows the stimulus with a fixed latency. In contrast, the non-linear model allows estimating (i) a variable onset within a user-specified time window and (ii) a variable duration of a sympathetic response. Additionally, the non-linear model includes estimation of spontaneous fluctuations that occur between trials.

In this tutorial, the non-linear model is estimated. The estimated parameters can then be used to test the within-subject hypothesis that the sympathetic arousal between CS+ and CS- are different on a group level.

#### 26.1 Setup the Non-Linear Model

In this step you prepare the model estimation. For this, you need to provide timing information about the experimental events which can either be entered manually or as a Matlab file. For large numbers of events as in this experiment, a timing file can easily be created using a few lines of Matlab code. Here we use timing information saved in the trimmed PsPM file. In this file, a trigger was saved at every CS onset. For timing information from other sources, the code has to be adapted accordingly. Go to the Matlab command window and enter the following lines, but replace [insert path!] by the correct path where the filtered SCR data from pspm.sourceforge.net is saved on your hard drive, e.g. C:\PsPM\_Tutorial\nonlinear\.

```
data_path = [insert path!]; % set the correct PsPM file path here!

p = 1; % the first participant
SOA = 3.5; % delay between CS and US onset in seconds

% load events of CS onsets from trimmed scr data
pname = fullfile(data_path,sprintf('tscr_s%i.mat',p))
[sts, infos, data, filestruct] = scr_load_data(pname,'events');

CS_onset = data{1, 1}.data; % recorded triggers of CS onset in seconds
US_onset = data{1, 1}.data + SOA; % US starts 3.5 seconds after CS

% variable latency for CS response between CS onset and US onset/omission
events{1} = [CS_onset US_onset]; % define an interval of 3.5 seconds for each
trial
% constant onset for US response
```

```
events{2} = US_onset;
save(fullfile(data_path,sprintf('event_timings_s%i.mat',p)),'events')
```

The resulting file event\_timings\_s1.mat contains the variable events with two cells.

- The first cell events {1} is a 40 x 2 matrix with the onset and offset of a time window for each trial in which we expect the response to the CS (both CS+ and CS- are treated the same here). We expect the response to the CS to occur with a variable onset in time. Note that the window for the CS response ends when the US starts (or is omitted), i.e. after 3.5 seconds.
- The second cell events {2} is a 40 x 1 vector that marks the onset of the electrical stimulation. For this event we expect an evoked response with constant latency after US occurrence. Importantly, we provide timings both for the occurrence and the omission of the US. Modelling an unconditioned response in all trials is necessary to get unbiased estimates for the CS responses. In general, the trial structure should always be the same for all trials, across experimental conditions.

Now the non-linear model estimation can be set up (Figure 26.1).

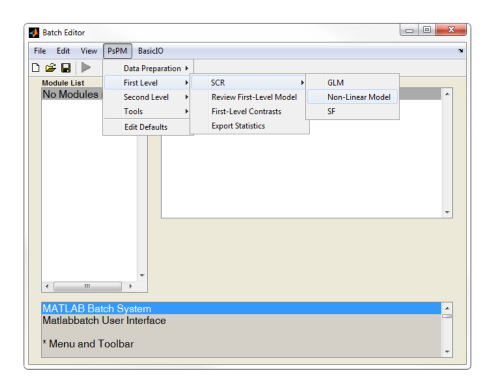


Figure 26.1: Prepare a batch to set up a non linear model for SCR analysis.

- In the batch editor go to  $PsPM \to First\ Level \to SCR \to Non-Linear\ Model$
- Model Filename: dcm s1
- Output Directory: Specify a folder where the model will be saved
- Channel: Default
- $Data \ \ \ \ Design: \rightarrow New: \ Session$ 
  - Data File: choose tscr s1.mat from your hard drive
  - $Design \rightarrow Timing\ File \rightarrow Select$  the previously created event\_timings\_s1.mat which was saved to your scr folder
  - Condition names: New Condition here we can define conditions that each trial belongs to. Create three new conditions. Note that we split CS+ trials in reinforced (CS+|US+) and non-reinforced (CS+|US-) trials because CS+|US+ trials are excluded from some analyses.
    - \* Condition
      - · Name: CS+/US-
      - · Index: 6 7 11 14 16 18 26 29 38 40

- \* Condition
  - · Name: CS+/US+
  - · Index: 13 17 19 20 21 24 28 31 32 37
- \* Condition
  - · Name: CS-
  - · Index: 1 2 3 4 5 8 9 10 12 15 22 23 25 27 30 33 34 35 36 39
- Display Options
  - Display Progress Window: If you plan to run the estimation in the background, you can turn off the progress window with this option
- For the remaining fields you can keep the default settings

Save the batch and script as batch\_model\_s1.m and press run. A window will be displayed that shows the progress of model estimation for each trial. The estimation for an entire session takes several minutes depending on your system. The file dcm\_s1.mat will be created. If you want to, you can review the first-level model by clicking *Review model* in the PsPM user interface. In the new window (Figure 26.2) add the non-linear model dcm\_s1.mat and choose *Display All trials for one session*. A graphics window (Fig X) will display the original data (blue) together with the model estimation (green).

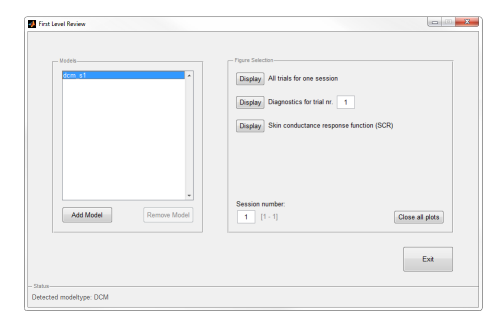


Figure 26.2: Use *Review Model* to inspect parameter estimation for one or multiple trials.

#### 26.2 First-level Contrasts

In this step you specify the hypothesis that a CS+ elicits a stronger sympathetic arousal than a CS- for an individual participant by providing a contrast weight vector.

- In the batch editor go to  $PsPM \rightarrow First\ Level \rightarrow First\ level\ Contrasts$
- Model File(s): Select dcm\_s1.mat
- Specify Contrasts on Datatype: param
- Contrasts: New Contrast
  - Contrast
    - \* Name: Fear Learning

    - \* Name: Electric Stimulation We include this contrast optionally to check if the electrical stimulation caused an increase in sympathetic arousal which is a premise for successful fear conditioning.
    - \* Contrast Vector: -1 -1 -1 -1 -1 -1 -1 -1 -1 -1 -1 -1 3 -1 -1 3 3 3 -1 -1 3 -1 -1 3 -1 -1 3 3 3 -1 -1 3 -1 -1 3 -1 -1 3 -1 -1 3 3 3 -1 -1 3 -1 -1 -1
- Delete: existing contrasts: No (default)

Save the batch and script as batch\_contrast\_s1.m and press run. The model file of participant s1 now contains the contrast that can be forwarded to a statistical test on group level. Preparing analyses for more than one scr file is explained in Chapter 27.

#### 26.3 Exporting statistics

PsPM provides basic tools for statistical analysis on the estimates of the non-linear model. Alternatively you can export the results as a text file to use them in different software applications.

- In the batch editor go to SCR $\rightarrow$ First Level $\rightarrow$ Export Data
- Model File(s): Select dcm s1.mat
- Specify Export on Datatype: param
- Target
  - Filename Enter dcm\_s01
- Specify Delimiter for Output File

- Semicolon

The text file will be saved to Matlab's current working directory.

# 27 Adapting scripts for multiple participants

In most experiments, you want to analyze data from multiple subjects, which implies that you have to repeat model estimation several times. There are two simple strategies to make this easier.

- Use dependencies: You can specify all steps for one subject in one go. At some points, you need to specify a file from a previous step (e.g., when trimming you need the file containing the imported data). In these cases, you can use the button "Dependency" to specify the corresponding file.
- Loop over participants in the script: You can easily adapt the scripts for many subjects by including a loop and adapting the file names. This is independent of whether or not you use dependencies. To show you how this works, see the examples below.

#### 27.1 Importing and trimming data

Here, we create an example script for importing and trimming data. We chose these steps since they are required in all types of analysis. The current example is based on the Appraisal dataset described in Chapter 25.

- In the batch editor specify all parameters in PsPM  $\rightarrow$  Data Preparation  $\rightarrow$  Import.
- You do not need to specify the SCR Data File. This will later be put in the script that loops over subjects.
- In the batch editor specify all parameters in PsPM  $\rightarrow$  Data Preparation  $\rightarrow$  Trim.
- Use the dependency button to select the imported Data File.
- Save batch and script.
- Open the saved job script (e.g., batch\_import\_trim\_job.m) and build a loop around the lines of code that were saved. In this loop, create a subject-specific filename (and make sure to include the whole path to the files in the variable data path).

```
data_path = [insert path!];
for p = 25:39; % participants
  % load participant-specific raw data
  matlabbatch{1}.pspm{1}.prep{1}.import.datatype.spike.datafile = ...
```

```
{fullfile(data_path,sprintf('trsp_1_%i.smr',p))};
  spike.importtype{1}.scr.chan_nr.chan_nr_spec = 2;
  spike.importtype{1}.scr.transfer.none = true;
 spike.importtype{2}.marker.chan_nr.chan_nr_spec = 0;
 matlabbatch{1}.pspm{1}.prep{1}.import.datatype.spike = spike;
 matlabbatch{1}.pspm{1}.prep{1}.import.overwrite = false;
 matlabbatch{2}.pspm{1}.prep{1}.trim.datafile(1) = ...
    cfg_dep('Import: Output File', substruct('.', ...
    'val', '{}',{1}, '.','val', ...
    '{}',{1}, '.','val', '{}',{1}), substruct('()',{':'}));
 matlabbatch{2}.pspm{1}.prep{1}.trim.ref.ref_mrk.from = 0;
 matlabbatch{2}.pspm{1}.prep{1}.trim.ref.ref_mrk.to = 15;
 matlabbatch{2}.pspm{1}.prep{1}.trim.ref.ref_mrk.mrk_chan.chan_def = 0;
 matlabbatch{2}.pspm{1}.prep{1}.trim.overwrite = false;
 % run batch
 scr_jobman('initcfg');
 scr_jobman('run', matlabbatch);
end
```

Run the script. The folder data\_path now contains trimmed scr files for all subjects.

# 27.2 Inverting non-linear models

Here we adapt the model inversion from Chapter 26 to multiple subjects. We use similar loops as above to create timing files and PsPM batches.

First, open the MATLAB script that created the timing file event\_timings\_s1.mat and construct a loop around it for subjects 2 to 20. Execute the script to obtain files containing timing information for subjects 2 to 20. Alternatively you can use the timing files for subjects 2 to 20 downloaded from pspm.sourceforge.net.

Next, open the saved job script (e.g., batch\_model\_s1\_job.m) created in Chapter 26 and build a loop around the lines of code that were saved. Make sure to include the whole path to the files in the variable data\_path. You can use the following code as guideline.

```
% set the path where you saved the scr data files
data_path = [insert path!];
% set the path where you saved the model file of participant 1
model_path = [insert path!]; % initiate PsPM scr_init % loop through participants 2 to 20
% initialize PsPM
scr_init
scr_jobman('initcfg')
```

```
for p = 2:20
  clear matlabbatch dcm
  dcm.modelfile = ['dcm_s' num2str(p)];
  dcm.outdir = {'D:\SCRalyze_tutorial\dcm_hra'};
  dcm.chan.chan_def = 0;
  dcm.session.datafile = ...
    {fullfile(data_path,['tscr_s' num2str(p) '.mat'])};
  dcm.session.timing.timingfile = ...
    {fullfile(data_path,['event_timings_s' num2str(p) '.mat'])};
  dcm.session.condition(1).name = 'CS+|US-';
  dcm.session.condition(1).index = [6 7 11 14 16 18 26 29 38 40];
  dcm.session.condition(2).name = 'CS+|US+';
  dcm.session.condition(2).index = [13 17 19 20 21 24 28 31 32 37];
  dcm.session.condition(3).name = 'CS-';
  dcm.session.condition(3).index = ...
    [1 2 3 4 5 8 9 10 12 15 22 23 25 27 30 33 34 35 36 39];
  dcm.data_options.norm = 0;
  dcm.data_options.filter.def = 0;
  dcm.resp_options.crfupdate = 0;
  dcm.resp_options.indrf = 0;
  dcm.resp_options.getrf = 0;
  dcm.resp_options.rf = 0;
  dcm.inv_options.depth = 2;
  dcm.inv_options.sfpre = 2;
  dcm.inv_options.sfpost = 5;
  dcm.inv_options.sffreq = 0.5;
  dcm.inv_options.sclpre = 2;
  dcm.inv_options.sclpost = 5;
  dcm.inv_options.ascr_sigma_offset = 0.1;
  dcm.disp_options.dispwin = 0;
  dcm.disp_options.dispsmallwin = 0;
  matlabbatch{1}.pspm{1}.first_level{1}.scr{1}.dcm = dcm;
  % run batch
  scr_jobman('run', matlabbatch);
end
```

Similarly to the previous step, the batch of the first-level Contrast of participant 1 is adapted to participants 2 to 20. Run the following code in MATLAB, but replace [insert path!] with the path on your hard drive where you saved dcm\_s1.mat:

```
% set the path where you saved the estimated non-linear model
model_path = [insert path!];
% initialize PsPM
```

```
scr_init
scr_jobman('initcfg')
% loop through participants 2 to 20
for p = 2:20
  clear matlabbatch
  matlabbatch{1}.pspm{1}.first_level{1}.contrast.modelfile = ...
    {fullfile(model_path,['dcm_s' num2str(p) '.mat'])};
  matlabbatch{1}.pspm{1}.first_level{1}.contrast.datatype = 'param';
  matlabbatch{1}.pspm{1}.first_level{1}.contrast.con(1).conname = 'Fear Learning';
  matlabbatch{1}.pspm{1}.first_level{1}.contrast.con(1).convec = ...
    [-1 -1 -1 -1 -1 2 2 -1 -1 -1 2 -1 0 2 -1 2 0 2 ...
    0 0 0 -1 -1 0 -1 2 -1 0 2 -1 0 0 -1 -1 -1 -1 0 2 -1 2];
  matlabbatch{1}.pspm{1}.first_level{1}.contrast.con(2).conname = ...
    'Electric Stimulation';
  matlabbatch{1}.pspm{1}.first_level{1}.contrast.con(2).convec = ...
    [-1 -1 -1 -1 -1 -1 -1 -1 -1 -1 -1 -1 3 -1 -1 -1 ...
    3 -1 3 3 3 -1 -1 3 -1 -1 -1 3 -1 -1 3 3 -1 -1 -1 3 -1 -1 -1];
  matlabbatch{1}.pspm{1}.first_level{1}.contrast.deletecon = true;
  % run batch manually
  scr_jobman('run', matlabbatch);
end
```

Run the script. The statistics for each contrast are now added to the model file.

# 28 Second-level Contrast Manager

In most cases, we want to compare the difference between conditions across participants on a group level (i.e., the second level in a hierarchical model). Each participant represents one data point that goes into a one sample t-test.

Below we demonstrate how to set-up Second-level contrasts for GLM and non-linear models.

#### 28.1 GLM

For the appraisal data of Chapter 25.

#### 28.1.1 Define Second-Level Model

- In the batch editor go to  $PsPM \rightarrow Second\ Level \rightarrow Define\ Second\ Level\ Model.$ 
  - Test Type: One Sample T-Test
  - Model File(s): Select the model files from all participants
  - Output Directory: Choose a location to save the group result

- Filename for Output Model: Chose a name so that you can easily refer to the contrast
- Define Contrast: Read from First Model File: Choose All Contrasts.
   This will select the contrast(s) that we created in the first level (i.e., in our case aversive>neutral)

#### 28.1.2 Report Second-Level Results

- In the batch editor go to PsPM → Second Level→Report Second-Level Results.
  - Model File: Select the second level model file that you created in the previous step.
  - Contrast Vector: Leave empty because we want all contrasts to be reported.

Run the batch. A figure appears that shows you a bar graph of the mean of the parameter estimates and the statistics will be printed to the MATLAB command window.

#### 28.2 Non-linear models

Here, we test the difference in anticipatory sympathetic arousal elicited by a CS+ vs CS- on a group level (see Chapter 26). As a supplementary hypothesis we want to check if the US+ indeed elicited a stronger response than the US-. From the 1st level contrast, eight statistics were generated for each model file. Use the *Review Model* utility from the PsPM main GUI to get a list of all contrasts. In the *First Level Review Manager*, click on *Add Model*, choose any model file created previously and press *Done*. Now press the Show button next to *Contrast names in command window* (Figure 28.1) to get a list of all contrasts:

- ullet Contrast 1: Fear Learning Flexible response # 1: amplitude
- Contrast 2: Electric Stimulation Flexible response # 1: amplitude
- ullet Contrast 3: Fear Learning Flexible response # 1: peak latency
- Contrast 4: Electric Stimulation Flexible response # 1: peak latency
- ullet Contrast 5: Fear Learning Flexible response # 1: dispersion
- Contrast 6: Electric Stimulation Flexible response # 1: dispersion
- Contrast 7: Fear Learning Fixed response # 1 response amplitude
- ullet Contrast 8: Electric Stimulation Fixed response # 1 response amplitude

Our hypothesis is concerned only with the response amplitude of the flexible event (anticipation of the CS) and the response amplitude of the fixed event (electrical stimulation), which are the first and eighth contrasts. The indices of these two contrasts are used for the setup of the Second-Level model below.

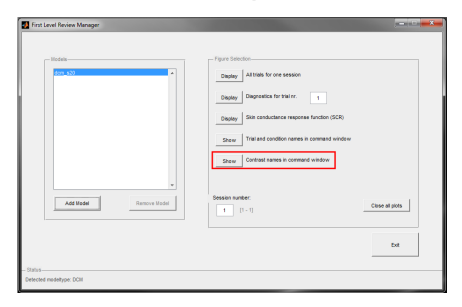


Figure 28.1: The First Level Review Manager displays all contrasts that are stored in a model file.

- In the batch editor go to PsPM→Second Level→Define Second-Level Model
- Test Type: One Sample T-Test
- Model File(s): Select the model files from all participants
- Output Directory: Choose a location to save the group result
- Filename for Output Model: Fear Learning
- Define Contrast: Read from First Model File
  - Contrast Vector: [1, 8]
- Overwrite Existing File: No (default)

Run the batch. You can display the results of the second level contrasts by pressing *Report 2nd level result* in the PsPM user interface and choosing Fear\_Learning.mat. A Figure displays the mean differences and standard error for the hypothesis that the CS+ elicits a stronger sympathetic arousal than CS- and that an electric stimulation results in an increase in sympathetic arousal (Figure 28.2).

Statistical parameters are printed in the MATLAB command window (Figure 28.3).

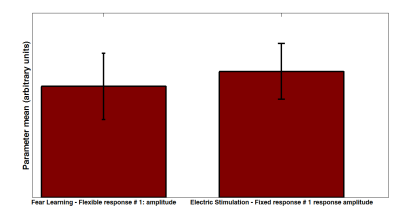


Figure 28.2: Display of results from a group analysis (2nd level). Left bar: difference in amplitude of sympathetic arousal (aSA) to CS+|US- minus CS-. Right bar: difference in aSA to US+ minus US-

Statistics:									
mean	sem	t	р	df	Contrast name				
5.16	1.54	3.34	0.0035	19	Fear Learning - Flexible response # 1: amplitude				
5.85	1.30	4.52	0.0002	19	Electric Stimulation - Fixed response # 1 response amplitude				

SCRalyze (c) Dominik R. Bach, Wellcome Trust Centre for Neuroimaging, UCL London UK

Figure 28.3: Statistics from one-sampled t-tests CS|US- minus CS- and US+ minus US- for 20 participants.

# Part IV

# Developer's Guide

# 29 General

# 29.1 Data files: General structure

In PsPM the data is saved in mat-files. Each file contains two variables:

- infos A struct variable with general infos
- data A cell array with a cell for each channel.

The cells contain a struct with channel specific infos and data. The structs have the mandatory fields:

- infos.duration (in seconds)
- data{n}.header
  - data{n}.header.chantype (as defined in the settings)
  - data{n}.header.sr (sample rate in 1/second, or timestamp units in seconds)
  - data{n}.header.units (data units, or 'events')
  - data{n}.data (actual data)

Additionally, a typical file contains the optional infos:

- infos.sourcefile
- infos.importfile
- infos.importdate
- infos.sourcetype
- infos.recdate
- infos.rectime

Some data manipulation functions (in particular, scr\_trim) update infos to record some file history.

### 29.2 How to add a new import data type

#### 29.2.1 Add function

Function name: scr get xxx (where xxx is the data type name).

Format:

```
[sts, import, sourceinfo] = scr_get_xxx(datafile, import)
```

The function needs to take an import job and add, for each job, fields

- .data the actual data for this channel (column vector)
- .sr the sample rate for this channel (only if .autosr enabled in scr init)

optional fields

- .marker for marker channels (timestamps or continuous, see scr get marker)
- .markerinfo optional, see scr\_get\_marker

- .minfreq minimum frequency for pulse channels
- .units if data units are defined by the recording software
- sts: -1 if import is unsuccessful

sourceinfo: contains information on the source file, with field

• .chan - a cell of string descriptions of the imported source channels, e. g. names, or numbers any optional fields that will be added to infos.source (e. g. recording date & time, and others)

Notes for multiple blocks: file formats that support multiple block storage within one file can return cell arrays import{1:blkno} and sourceinfo{1:blkno}; PsPM will save individual files for each block, with a filename 'scr fn blk0x.mat'.

#### 29.2.2 Add information to settings

The file scr\_init contains a block that defines possible import data types. Add a new field here

```
% Description of data type
% -------
defaults.import.datatypes(1) = ...
struct('short', 'xxx', ... % short name for internal purposes
   'long', 'Datatype description', ... % long name for GUI
   'ext', '*', ... % data file extension
   'funct', @scr_get_xxx, ... % import function
   'chantypes', {{defaults.chantypes.type}}, ... % allowed channel
types
   'chandescription', 'channel', ... % description of channels for
GUI
   'multioption', 1, ... % import of multiple channels for GUI
   'searchoption', 1, ... % allow channel name search for GUI
   'automarker', 0, ... % marker not stored in separate channel
   'autosr', 1); % sample rate automatically assigned
```

#### Good to know:

- the "long" definition is used in the GUI make sure it's readable
- if no event channels can be imported, change .chantypes
- ullet if channels have searchable names in the import file, set .searchoption = 1
- $\bullet$  if no channel number needs to be assigned for the marker channel, set .automarker = 1

- $\bullet$  if sample rate is contained in import file and determined during import, set .autosr = 1
- if you need external functions put them into a folder in the 'import' subdirectory and add/remove this path within the scr get xxx function

#### 29.3 How to add a new channel type

#### 29.3.1 Add function

Function name: scr get xxx (where xxx is the channel type)

Format:

```
[sts, data] = scr_get_channeltype(import)
```

data: data cell of structure readable by scr\_load\_data

Good to know: for event channels, use the function scr\_get\_events to convert various event formats into time stamps (see scr\_get\_marker or scr\_get\_hb as an example)

#### 29.3.2 Add information to settings

Add information on the new channel type and import function to

#### 29.4 How to add a new GLM type

#### 29.4.1 Add information to settings (Example SCR)

#### 29.4.2 Add default basis function

Function name: scr bf xxx

Function arguments: vector of arguments, first element is time resolution, further arguments as defined in defaults.glm(n).cbf.args

# 29.5 Warning IDs in PsPM

#### 29.5.1 General

- invalid\_input
- $\bullet$  invalid\_channeltype
- $\bullet \ \ nonexistent\_file$
- $\bullet \ \ channel\_not\_contained\_in\_file$
- ullet obsolete\_function
- $\bullet \ \, {\rm not\_allowed\_channel type}$

## 29.5.2 Function specific

 $scr_load_data$ 

 $\bullet$  invalid\_data\_structure

scr trim

• marker\_out\_of\_range

 $scr\_find\_channel$ 

- $\bullet \ \ multiple\_matching\_channels$
- no\_matching\_channels

 $scr\_get\_scr$ 

• no\_conversion\_constant

 $\operatorname{scr} \_\operatorname{pp}$ 

• invalid freq

 $scr\_prepdata$ 

- no\_low\_pass\_filtering
- $\bullet \ \ downsampling\_failed$
- nonint sr

 $scr\_get\_timing$ 

• invalid\_vector\_size

- $\bullet \ \, event\_names\_dont\_match \\$
- $\bullet \ \ no\_numeric\_vector$
- $\bullet$  no\_integers

 $scr\_down$ 

 $\bullet \ \ {\rm rate\_below\_minimum}$ 

# 30 List of data formats

# 30.1 Supported Channel types

Data format	SCR	ECG	Heart Rate	Heart Beat	Heart Period	Respiration	Pupil Size	Marker	Custom
CED Spike	<b>√</b>	<b>√</b>	<b>√</b>	<b>√</b>	<b>√</b>	<b>√</b>	<b>√</b>	<b>√</b>	<b>√</b>
Matlab	<b>√</b>	<b>√</b>	<b>√</b>	<b>√</b>	<b>√</b>	<b>√</b>	<b>√</b>	<b>√</b>	<b>√</b>
Text	<b>√</b>	<b>√</b>	<b>√</b>		<b>√</b>	<b>√</b>	<b>√</b>		<b>√</b>
Biopach	<b>√</b>	<b>√</b>	<b>√</b>	<b>√</b>	<b>√</b>	<b>√</b>	<b>√</b>	<b>√</b>	<b>√</b>
AcqKnowledge (≤									
v3.9.0)									
Biopac AcqKnowledge	✓	<b>√</b>	<b>√</b>	<b>√</b>	<b>√</b>	<b>√</b>	<b>√</b>	<b>√</b>	<b>√</b>
(exported)									
Labchart exported	✓	✓	✓		✓	✓	✓	✓	✓
$(\leq v7.1)$									
Labchart exported	<b>√</b>	✓	✓		✓	✓	✓	✓	✓
$(\geq v7.2)$									
VarioPort	✓	<b>√</b>	<b>√</b>		<b>√</b>	<b>√</b>	<b>√</b>	✓	✓
Biograph Infiniti	✓			<b>√</b>		<b>√</b>			
(exported)									
Mindmedia Biotrace	<b>√</b>	✓	✓		✓	✓	✓	✓	✓
(exported)									
Brain Vision	✓	✓	✓		✓	<b>√</b>	<b>√</b>	✓	<b>√</b>
Windaq (wdq)	✓	<b>√</b>	<b>√</b>	<b>√</b>	<b>√</b>	<b>√</b>	✓	<b>√</b>	<b>√</b>
Observer XT	✓	<b>√</b>	<b>√</b>	<b>√</b>	<b>√</b>	<b>√</b>	<b>√</b>	<b>√</b>	<b>✓</b>
compatible									
NeuroScan	✓	<b>√</b>	<b>√</b>		<b>√</b>	<b>√</b>	<b>√</b>	<b>√</b>	<b>√</b>
BioSemi	<b>√</b>	<b>√</b>	<b>√</b>		<b>√</b>	<b>√</b>	<b>√</b>	<b>√</b>	<b>√</b>
Eyelink							<b>√</b>	<b>√</b>	<b>√</b>
European Data Format	<b>√</b>	<b>√</b>	<b>√</b>	<b>√</b>	<b>√</b>	<b>√</b>	<b>√</b>	<b>√</b>	<b>√</b>

# 30.2 Further settings

Data format	Datatype	File extension	Manufacturer	Import multiple channels	Search channel names	Automarkers	Ask for sampling rate
CED Spike	spike	.smr	CED	<b>√</b>	<b>√</b>		
Matlab	mat	.mat		<b>√</b>			<b>√</b>
Text	txt	.txt		<b>√</b>	<b>√</b>		
Biopach	acq	.acq	Biopac	<b>√</b>	<b>√</b>		
AcqKnowledge ( $\leq$ v3.9.0)							
Biopac AcqKnowledge (exported)	acqmat	.mat	Biopac	<b>√</b>	<b>√</b>		
Labchart exported (\le v7.1)	labchartmat_ext	.mat	ADInstruments	✓	<b>√</b>	<b>√</b>	<b>√</b>
Labchart exported $(\geq v7.2)$	labchartmat_in	.mat	ADInstruments	<b>√</b>		<b>√</b>	
VarioPort	vario	.vpd	Becker MediTec	✓	<b>√</b>	<b>√</b>	
Biograph Infiniti (exported)	biograph	.txt	Thought Technology				
Mindmedia Biotrace (exported)	biotrace	.txt	MindMedia			<b>√</b>	
Brain Vision	brainvision	.eeg	BrainProducts	<b>√</b>	<b>√</b>	<b>√</b>	
Windaq (wdq)	windaq	.wdq	Dataq	<b>√</b>			
Observer XT compatible	observer	.any	Noldus	✓	✓		
NeuroScan	cnt	.cnt		<b>√</b>	<b>√</b>	<b>√</b>	
BioSemi	biosemi	.bdf		<b>√</b>	<b>√</b>	<b>√</b>	
Eyelink	txt	.asc		<b>√</b>		<b>√</b>	
European Data Format	edf	.edf	European	<b>√</b>	<b>√</b>	<b>√</b>	
			Data Format				

# 30.3 How to

# 30.3.1 CED Spike

Import the original acquisition files

#### 30.3.2 Text

Text files can only contain numbers (i.e. no header lines with channel names) and one data column per channel. Make sure you use the decimal point (i.e. not decimal comma as used in some non-English speaking countries). At the moment, no import of event markers is possible

#### 30.3.3 Matlab

Each input file must contain a variable called data that is either a cell array of column vectors, or a data points × channels matrix. At the moment, no import of event markers is possible. Data structures containing timestamps cannot be imported at the moment; rather, PsPM will ask you for a sample rate.

#### 30.3.4 Biopac AcqKnowledge

Import the original acquisition files

# 30.3.5 exported Biopac AcqKnowledge

# 30.3.6 Labchart exported (up to v. 7.1)

Export data to matlab format (plugin for the LabChart software, available from www.adinstruments.com)

#### 30.3.7 Labchart exported (v. 7.2 and higher)

# 30.3.8 VarioPort

Import the original acquisition files

#### 30.3.9 Biograph Infiniti exported

Export data to text format, both "Export Channel Data" and "Export Interval Data" are supported; a header is required

#### 30.3.10 Mindmedia Biotrace exported

#### 30.3.11 BrainVision

Import the original acquisition files

#### 30.3.12 DATAQ Windaq (.wdq; read with ActiveX-Lib)

Requires an ActiveX Plugin provided by the manufacturer and contained in the subfolder Import/wdq for your convenience. This plugin only runs under 32 bit Matlab on Windows. Import the original acquisition files.

# 30.3.13 DATAQ Windaq (.wdq)

Windaq import written by the PsPM team. Is platform independent, thus has no requirements for ActiveX Plugins, Windows or 32bit Matlab. Imports the original acquisition files files. Up to now the import has been tested with files of the following type: Unpacked, no Hi-Res data, no Multiplexer files. A warning will be produced if the imported data-type fits one of the yet untested cases. If this is the case try to use the import provided by the manufacturer (see above).

#### 30.3.14 Observer XT compatible

30.3.15 NeuroScan

30.3.16 BioSemi

30.3.17 Eyelink

Import files converted into ascii files (.asc).

### 30.3.18 European Data Format

Import the original acquisition files.

# 31 GUI

Contributed by Gabriel Gräni.

## 31.1 Matlabbatch: Getting started

- 1. Add the trunk folder to the matlab path
- 2. Type scr\_init into the command window (after the execution of the command the folders scr\_cfg and matlabbatch should be added to the matlab path)
- 3. Start matlabbatch by the typing cfg\_ui into the command window
- 4. If the item SCR exists in the menu bar of matlabbatch you can skip steps 5 to 7 and continue at step 8
- 5. Select -> File -> Add Application
- 6. Navigate to the folder scr\_cfg on the left hand side of the window and select the file scr\_cfg.m on the right hand side -> Press the button Done
- 7. A new item, called SCR, will appear in the upper menu bar.
- 8. By selecting SCR the desired action can be selected (at the moment, there is only Data Preparation -> {Import, Trim} available)

#### 31.1.1 Example Function: Trim

This example demonstrates how matlabbatch can be used to execute a function. For all other functions matlabbatch behaves in the same manner.

- Select a file by pressing the Select Files Button (under Datafile)
- Select Reference and choose an item in the lower part of the window
- Fill in the desired values in the fields which are marked with "<-X"
- After you have chosen a file and filled in all values correctly, you will see a green arrow on the upper left part of the window
- By pressing on the green arrow the selected file will be trimmed according to the filled in values

#### 31.2 Matlabbatch: How to

#### 31.2.1 Preliminaries

- Add folder of matlabbatch to the matlab path
- Add first application and then load the batch in order to execute a function

#### 31.2.2 Some notes for creating a new application

- Leafs (items) are specified first
- Assigning child items to .val or .values fields of their parent items
- Root node of a tree is specified last
- Some examples of items:

```
- cfg item:
  item1= cfg_item; % Defines generic configuration item
  item1.name = 'Def 1'; % The display name
  item1.tag
              = 'def1'; % The name appearing in the harvested
  job
 % structure. This name must be unique
 % among all items in the val field of the
 % superior node
  item1.val = {true}; % Value of item (optional)
 item1.help = {'Help...'}; % Help text
- cfg entry:
  entry1 = cfg_entry; % Defines entry configuration item
  entry1.name = 'Input';
 entry1.tag = 'input';
 entry1.strtye = 'r'; % Type of values which can be entered
```

```
entry1.num = [1 1]; % Expected dimension of the input
 entry1.help = {'Help...'};
- cfg choice:
 choice = cfg_item; % Defines choice configuration item
  choice.name = 'Choice';
  choice.tag = 'choice';
 choice.values = {item1, entry1}; % Defines which items will
 % selectable in the choice menu.
 choice.help = {'Help...'};
- cfg exbranch:
 fct = cfg_exbranch; % Defines the branch that has information
 % about how to run this module
 fct.name = 'Trim';
 fct.tag = 'trim';
 fct.val = {choice}; % The items that belong to this branch.
 % All items must be filled before this
 % branch can run or produce virtual
 % outputs
 fct.prog = @cfg_run_fct; % A function handle that will be
 \% with the harvested job to run the
 % computation
 trim.vout = @cfg_vout_fct; % A function handle that will be
 % called with the harvested job to
 % determine virtual outputs
 trim.help = {Help...'};
- There exists a number of other item classes. Here is a list of the
 most important classes: cfg item, cfg entry, cfg choice, cfg menu,
 cfg exbranch, cfg files, cfg branch, cfg repeat
 For more information call the help function in matlab (e.g. help
 help cfg item)
```

# • Note:

The inputs to each module have to be described in a tree-like structure. During data entry, there is no way to change the tree structure based on input data. Add application to the configuration tree by default

# 31.2.3 Add application to the configuration tree by default

In the following it is shown how an application can be added to the menu bar of matlabbatch by default (without adding it every time matlabbatch is started)

- Start matlabbatch and add the appliaction cfg\_confgui in the folder matlabbatch/cfg\_confgui
- Put Generate code into the Module list by selecting ConfGUI -> Generate code in the menu bar
- Fill out all the input fields on the right side:
  - Output filename: This file will contain the whole menu structure, validity constraints and links to run time code of the application.
  - Output directory: All files which are created by the ConfGUI will be stored into this directory (chose a directory which is added to the matlab path)
  - Root node of config: Name of the root node of the application's configuration tree
  - Options:
    - 1. Create Defaults File: Yes
    - 2. Create mlbatch appcfg File: Yes
    - 3. Create Code for addpath(): No
- Finally press the green arrow on the upper left side of the batch editor
- As no error occurred 3 new files ({Output filename}.m, {Output filename}\_def.m, cfg\_mlbatch\_appcfg.m) should be created and added into the folder {Output directory}.
- Each time matlabbatch is started, it will search for any cfg\_mlbatch\_appcfg.m file (this file contains the names of the configuration files) and will add the corresponding application to the batch editor.

#### 31.2.4 Add modules to module list

Example: Module Import and Trim will be added to the module list

```
arg1 = 'scr.prep.import_data';
arg2 = 'scr.prep.trim';
mod_cfg_id1 = cfg_util('tag2mod_cfg_id',arg1);
mod_cfg_id2 = cfg_util('tag2mod_cfg_id',arg2);
cjob = cfg_util('initjob');
mod_job_id1 = cfg_util('addtojob', cjob, mod_cfg_id1);
mod_job_id2 = cfg_util('addtojob', cjob, mod_cfg_id2);
cfg_util('harvest', cjob, mod_job_id1);
cfg_util('harvest', cjob, mod_job_id2);
cfg_ui('local_showjob', findobj(0,'tag','cfg_ui'), cjob);
```

#### 31.2.5 Changes

• In the function cfg\_onscreen at line 36 figure(fg); is commented out in order to prevent the appearance of the GUI for a short time if the function cfg\_ui('Visible', 'off') is called.

# 31.3 Matlabbatch: changing help texts and fieldnames

#### 31.3.1 File structure of matlabbatch GUI

There exist two files per function: 1 configuration file and 1 run file. The configuration file defines the structure of the corresponding function in the matlabbatch GUI whereas the run file firstly gathers all entered values and secondly calls the corresponding SCR function. Both types of files are located in the subfolder scr\_cfg. The name of a configuration or a run file consists of two parts. The prefix of a configuration filename is called scr\_cfg\_ whereas the filename of a run file begins with scr\_cfg\_run. The second part of the filename is named after the function name (eg. for the function scr\_import.m -> scr\_cfg\_import.m, scr\_cfg\_run\_import.m).

#### 31.3.2 Edit help texts and fieldname

In order to change any help text or fieldname in a matlabbatch GUI function the corresponding configuration file has to be opened. For each item in a matlabbatch GUI function a struct variable which contains several struct fields is defined in the configuration file.

- Help text The field .help defines the help text of the item which can be edited in order to change the help text. As soon as matlabbatch has been closed and opened again, the changes in the help text will be visible in matlabbatch GUI.
- Fieldname The fieldname of an matlabbatch GUI item is defined by the struct field .tag . In case a fieldname of an item should be changed be careful to verify if no other item, which has the same root node, hold the same fieldname. Otherwise matlabbatch will not work properly. After the fieldname of an item has been changed the run file (scr\_cfg\_run\_functionname.m) of the corresponding function has to be adapted as well in order to ensure that the function call in the run file is done properly.

# 32 Test Environment

Contributed by Linus Rüttimann & Tobias Moser.

# 32.1 Unittest: General implementation

In PsPM the Matlab Unit Testing Framework is used for testing of functions. For each tested function there is a Matlab class with the name 'functionname\_test',

which contains the unittests for that specific function. Additionally there is a documentation page for each of the test classes, where information about the unittests can be found.

To run the unittests of a test class, an object of the class has to be created:

```
testCase = functionname test
```

where 'testCase' is an arbitrary object name and 'funtionname\_test' is the name of a test class. Then all the unittest that are contained in the test class can be run with:

testCase.run

A specific unittest can be run with:

```
testCase.run('unittest_name')
```

Remember that a new test class object must be generated each time the test class has been changed.

# 32.2 Testcases: scr add channel

#### 32.2.1 Information

```
Testclass: scr find channel test
```

Function:  $sts = scr_add_channel(fn, newdata, msg)$ 

#### 32.2.2 Testcases

#### Invalid input arguments

Function name: invalid input(this)

Description: Checks for warnings, if the input arguments are invalid.

Setup:

newdata.header.sr = 100;

Input	Expected warning
scr_add_channel('fn') [invalid fn, no newdata]	ID:invalid_input
scr_add_channel('fn', 'foo') [invalid fn and newdata]	ID:invalid_input
scr add channel(newdata, newdata) [no fn]	ID:invalid input

#### Valid input arguments

Function name: valid input(this)

Description: Generates a test channel and passes it to scr add channel.

#### Tests:

- 1. Test if function runs through without any warnings
- 2. Test if 'sts' is equal 1
- 3. Test if 'newdata' really has been added to the dataset
  - (a) Test if 'scr\_load\_data' is successful
  - (b) Test if 'msg' has been added to infos.history
  - (c) Test if 'data' has 2 channels

# 32.3 Testcases: scr ecg2hb

#### 32.3.1 Information

Testclass: scr ecg2hb test

Function: [sts,pt debug] = scr ecg2hb(fn, chan, options)

# 32.3.2 Setup

#### Constants

- chan = struct('nr', 3, 'name', 'ecg');
- fn = 'ImportTestData\ecg2hb\tscr s102 s1.mat';
- backup suffix = ' backup';
- options = struct('semi', 0);

#### Variables

• backup\_file

#### 32.3.3 Testcases

# Invalid input arguments

Function name: invalid\_input(this)

Description: Pass invalid input arguments and check if the warnings are as expected.

# Tests:

Input	Ex
F ***	ĽA
sc_ecg2hb() [no arguments]	ID
scr_ecg2hb(1) [invalid file name]	ID
scr_ecg2hb(this.fn, 'bla') [invalid channel (text)]	ID
scr_ecg2hb(this.fn, 1) [invalid channel type]	ID:not_a
o.twthresh = 'bla'; scr_ecg2hb(this.fn, this.chan.nr, o) [invalid twthresh (text)]	ID
o.minHR = 202; scr_ecg2hb(this.fn, this.chan.nr, o) [invalid minHR (> default_maxHR)]	ID
o.minHR = 202; o.maxHR = 19; scr_ecg2hb(this.fn, this.chan.nr, o) [invalid minHR > maxHR]	ID
o.maxHR = 19; scr_ecg2hb(this.fn, this.chan.nr, o) [invalid maxHR (< default_minHR)]	ID
o.debugmode = 5; scr_ecg2hb(this.fn, this.chan.nr, o) [invalid debugmode (not in [0,1])]	ID
o.semi = 5; scr_ecg2hb(this.fn, this.chan.nr, o) [invalid semi (not in [0,1])]	ID

# Valid input arguments

Function name: valid\_input(this)

Description: Pass valid input arguments and check if there are no warnings.

#### Tests:

Input	Expected warning
scr_ecg2hb(this.fn, this.chan.nr, this.options)	-
scr_ecg2hb(this.fn, this.chan.name, this.options)	-
this.test_added_data()	-

#### 32.3.4 Other Methods

#### Test for added data

Function name: test\_added\_data()

Description: Check if added hb channels show an expected behaviour.

Tests (for each channel):

Tested Value	Expected Value
Sampling rate	1
Unit	'events'
Channel type	'hb'
Amount of data points in data	> 1
Data is distributed equally (standard deviation)	< 2s
Time between end of recording and last data point	< 60s

# 32.4 Testcases: scr find channel

#### 32.4.1 Information

 $Testclass: \ scr\_find\_channel\_test$ 

 $Function: \ chan = scr\_find\_channel(headercell, \ chantype)$ 

#### 32.4.2 Testcases

# Invalid input arguments

Function name: invalid inputargs(this)

Description: Checks for warnings, if the input arguments are invalid.

Setup:

```
headercell = {'heart', 'scr', 'pupil'};
```

Tests:

Input	Expected warning
scr_find_channel('str','scr') [no headercell]	ID:invalid_input
scr_find_channel(headercell, 'str')	ID:not_allowed_channeltype
scr_find_channel(headercell, 4) [no string chantype]	ID:invalid_input

# Valid Input Arguments

Function name: valid\_inputargs(this)

Description: Checks for correct return value if the input arguments are valid

Setup:

```
headercell = {'heart', 'scr', 'pupil', 'mark', 'gsr', 'eda'};
```

Input	Exp. Output	Expected warning
scr_find_channel(headercell, 'pupil')	3	
scr_find_channel(headercell, 'resp')	0	ID:no_matching_channels
scr_find_channel(headercell, 'scr')	-1	ID:multiple_matching_channels
scr_find_channel(headercell, {'mark',	4	
'str', 'bla'})		
scr_find_channel(headercell, {'call',	0	no matching channel, but no warning
'str', 'me'})		
scr_find_channel(headercell, {'scr',	-1	multiple matching channels, but no
'gsr', 'eda'})		warning

# 32.5 Testcases: scr\_get\_ecg

#### 32.5.1 Information

 $Testclass: scr\_get\_ecg\_test$ 

Function:  $[sts, data] = scr\_get\_ecg(import)$ 

#### 32.5.2 Testcases

#### Test

Function name: test(this)

Description: Test if all fields are returned correctly

#### Tests:

- 1. Test if 'sts' is equal 1.
- 2. Test if data.data is equal import.data
- 3. Test if data.header.chantype is 'ecg'
- 4. Test if data.header.units is equal import.units
- 5. Test if data.header.sr is equal import.sr

# 32.6 Testcases: scr get events

#### 32.6.1 Information

 $Testclass: \ scr\_get\_events\_test$ 

Function: [sts, import] = scr get events(import)

#### 32.6.2 Testcases

# Check warnings

Function name: check warnings(this)

Description: Checks for warnings, if the field '.markers' is missing or contains

invalid content.

#### Tests:

Input	Expected warning	
Missing marker field	ID:nonexistent_field	
import.marker = 'foo'	ID:invalid_field_content	

#### **Timestamps**

Function name: timestamps(this)

Description: Checks for correct output if the input is timestamp data

#### Tests:

- 1. Test if 'sts' is equal 1.
- 2. Test if the length of the output data is equal to the length of the input data

#### Continuous

Function name: continuous(this)

Description: Checks for correct output if the input is continuous data

#### Tests:

1. Perform three tests with different settings

#### Tests:

- (a) Test if 'sts' is equal 1.
- (b) Test if the length of the field 'markerinfo' is equal to the length of the output data.
- (c) Test if the length of the output data is equal to the expected number of pulses in the input data.

#### Settings:

- (a) flank = 'both' (default)
- (b) flank = 'both' & data offset 50
- (c) flank = 'ascending'
- (d) flank = 'descending'
- (e) inverted input signal
- (f) signal with angular flanks
- (g) check with

- 2. Additional test for setting (b): Test if data offset has been removed in the output data.
- 3. Additional test for setting (c) and (d): Test if positions returned by output data correspond to flank changes in the input data.
- 4. Test if markerinfo is not set if it has been set before.

# 32.7 Testcases: scr get eyelink

#### 32.7.1 Information

Testclass: scr\_get\_eyelink\_test

Function: [sts, data] = scr get eyelink(import)

#### **32.7.2** Methods

# $verify\_basic\_data\_structure$

Function name: verify\_basic\_data\_structure(this, data, sourceinfo, channel\_types) Description: Tests if the returned data structure is valid and match a given expected pattern.

#### Tests:

- 1. Test if all channels are numeric
- 2. Test if recorded time and date have a valid format
- 3. Test if blink channels have correct unit
- 4. Test if pupil channels have either 'diameter' or 'area' as unit
- 5. Test if channels labeled with 'position' have unit 'pixel'
- 6. Test if channels labeled with 'blink' have unit 'blink'

#### 32.7.3 Testcases

# test\_two\_eyes

Function name: test\_two\_eyes(this)

Description: Test if the returned data structure fits into the pattern of a two eyes data set.

#### Tests:

1. Create an import data set and the expected channel data set an pass it to 'verify basic data structure()'

#### test\_one\_eye

Function name: test one eye(this)

Description: Test if the returned data structure fits into the pattern of a one eye data set.

#### Tests:

1. Create an import data set and the expected channel data set an pass it to 'verify\_basic\_data\_structure()'

# 32.8 Testcases: scr get hb

#### 32.8.1 Information

 $Testclass: scr\_get\_hb\_test$ 

Function:  $[sts, data] = scr\_get\_hb(import)$ 

#### 32.8.2 Testcases

# Test

Function name: test(this)

Description: Test if all fields are returned correctly

#### Tests:

- 1. Test if 'sts' is equal 1.
- 2. Test if data.data is equal import.data
- 3. Test if data.header.chantype is 'hb'
- 4. Test if data.header.units is 'events'
- 5. Test if data.header.sr is 1

# 32.9 Testcases: scr\_get\_hr

#### 32.9.1 Information

 $Testclass: \ scr\_get\_hr\_test$ 

Function:  $[sts, data] = scr\_get\_hr(import)$ 

#### 32.9.2 Testcases

#### $\mathbf{Test}$

Function name: test(this)

Description: Test if all fields are returned correctly

- 1. Test if 'sts' is equal 1.
- 2. Test if data.data is equal import.data
- 3. Test if data.header.chantype is 'hr'
- 4. Test if data.header.units is equal import.units
- 5. Test if data.header.sr is equal import.sr

# 32.10 Testcases: scr get marker

#### 32.10.1 Information

Testclass: scr get marker test

Function: [sts, data] = scr get marker(import)

#### 32.10.2 Testcases

#### Test

Function name: test(this)

Description: Test if all fields are returned correctly

#### Tests:

- 1. Test if 'sts' is equal 1.
- 2. Test if data.data is equal import.data
- 3. Test if data.header.chantype is 'marker'
- 4. Test if data.header.units is 'events'
- 5. Test if data.header.sr is 1

# 32.11 Testcases: scr get pupil

#### 32.11.1 Information

Testclass: scr get pupil test

Function: [sts, data] = scr get pupil(import)

#### 32.11.2 Testcases

#### Test

Function name: test(this)

Description: Test if all fields are returned correctly

- 1. Test if 'sts' is equal 1.
- 2. Test if data.data is equal import.data
- 3. Test if data.header.chantype is 'pupil'
- 4. Test if data.header.units is equal import.units
- 5. Test if data.header.sr is equal import.sr

# 32.12 Testcases: scr get resp

#### 32.12.1 Information

```
Testclass: scr_get_resp_test
Function: [sts, data] = scr_get_resp(import)
```

#### 32.12.2 Testcases

#### Test

Function name: test(this)

Description: Test if all fields are returned correctly

#### Tests:

- 1. Test if 'sts' is equal 1.
- 2. Test if data.data is equal import.data
- 3. Test if data.header.chantype is 'resp'
- 4. Test if data.header.units is equal import.units
- 5. Test if data.header.sr is equal import.sr

# 32.13 Testcases: scr get scr

#### 32.13.1 Information

```
Testclass: scr_get_scr_test
Function: [sts, data] = scr_get_scr(import)
```

#### 32.13.2 Testcases

There are three test functions. One for the case that no transfer parameters are defined, one for the case that the transfer parameters are defined in a struct and one for the case that they are defined in a .mat file. They are all performing the following tests, plus eventually some individual tests

- 1. Test if 'sts' is equal 1.
- 2. Test if the field data.data exists
- 3. Test if the field data.data is not empty
- 4. Test if the field data.header.units exists
- 5. Test if the field data.header.sr exists
- 6. Test if the field data.header.chantype exists
- 7. Test if data.header.sr is equal import.sr
- 8. Test if data.header.chantype is 'scr'

# No transfer parameters

Function name: no transferparams(testCase)

Description: Test if all fields are returned correctly, if no transfer parameters are defined.

Additional Tests:

No additional tests

# Struct transfer parameters

Function name: stuct transferparams(testCase)

Description: Test if all fields are returned correctly, if the transfer parameters are defined in a struct.

# Additional Tests:

- 1. Check for warning if the conversion constant (import.transfer.c) is not defined
- 2. Checks that there are no warnings if import.transfer.Rs or import.transfer.offset is not defined.

#### File transfer parameters

Function name: file transferparams(testCase)

Description: Test if all fields are returned correctly, if the transfer parameters are defined in a .mat file.

#### Additional Tests:

1. Check for warning if the transfer parameter file doesn't exist.

# 32.14 Testcases: scr get timing

#### 32.14.1 Information

```
Testclass: scr_get_timing_test
Function: [sts, multi] = scr_get_timing('onsets', intiming, timeunits)
[sts, epochs] = scr_get_timing('epochs', epochs)
[sts, epochs] = scr_get_timing('events', events)
```

#### 32.14.2 Testcases

### Invalid input arguments

Function name: invalid\_inputargs(this)

Description: Checks for warnings, if the input arguments are invalid.

#### Tests:

Input	Expected warning
scr_get_timing('epochs') [missing	ID:invalid_input
input var]	
scr_get_timing('onsets', 'str') [no	${ m ID:invalid\_input}$
timeunits var]	
scr_get_timing('foo') [unknown	${ m ID:invalid\_input}$
format]	
scr_get_timing('onsets', intiming,	ID:number_of_elements_dont_match
'samples') [two sessions with	
nonmatching number of conditions]	
scr_get_timing('onsets', intiming,	$ID:event\_names\_dont\_match$
'samples') [two sessions with	
nonmatching condition names]	
scr_get_timing('onsets', intiming,	ID:no_numeric_vector
'samples') [intiming.onsets{1} is no	
numeric vector]	
scr_get_timing('epochs', fn_mat,	ID:no_integers
'samples') [epochs is not an integer	
array]	

#### Case Epochs

```
Function name: case\_epochs(this)
```

Description: Checks the function in 'epochs' mode.

Function: [sts, epochs] = scr get timing('epochs', epochs)

# Test 1 (matfile input)

Input: mat file with variable: epochs = [1 4; 2 5; 3 6]

Check if sts==1 and if the return value is equal the input array.

# Test 2 (spm input)

```
Input: mat file with variable: onsets\{1\} = [1\ 2\ 3]';onsets\{2\} = [4\ 5\ 6]';
```

Check if sts==1 and if the return value is equal [onsets{1}, onsets{2}].

#### Test 3 (textfile input)

```
Input: textfile with variable: epochs = [1 \ 4; 2 \ 5; 3 \ 6]
```

Check if sts==1 and if the return value is equal the input array.

#### Test 4 (matrix input)

```
Input: matrix: epochs = [1 \ 4; \ 2 \ 5; \ 3 \ 6]
```

Input: mat file with the variables:

Check if sts==1 and if the return value is equal the input array.

#### Case onsets

```
Function name: case onsets(this)
Description: Checks the function in 'onsets' mode.
Function: [sts, multi] = scr get timing('onsets', intiming, timeunits)
```

#### Test 1

```
names = {'name1', 'name2'};
onsets = \{[1\ 2], [3\ 4]\};
pmod.name = {'name3', 'name4'};
pmod.param = \{[2\ 3], [4\ 5]\};
pmod.poly = \{2, 2\};
save(fn_mat, 'names', 'onsets', 'pmod');
Function call:
[sts, outtiming] = scr get timing('onsets', fn mat, 'samples');
Tests:
```

```
Check if sts==1, if onsets and names are unchanged and if
outtiming.pmod.param == \{[2\ 3], [4\ 9], [4\ 5], [16\ 25]\}
```

#### Test 2

```
Input:
```

```
mat file with the variables: names = {'name1', 'name2'};
onsets = \{[1\ 2\ 3], [3\ 4\ 5]\}; durations = \{[3\ 4\ 5]', [5\ 6\ 7]'\};
pmod.name = {'name3', 'name4'};
pmod.param = \{[2 \ 3 \ 4], [4 \ 5 \ 6]\};
```

```
pmod.poly = {2, 1};
Function call:
[sts, outtiming] = scr_get_timing('onsets', fn_mat, 'samples');
Tests:
Check if sts==1, if onsets,names and durations are unchanged and if outtiming.pmod.param == {[2 3 4], [4 9 16], [4 5 6]}
```

#### Case events

```
Function name: case_events(this)
Description: Checks the function in 'events' mode.
Function: [sts, epochs] = scr get timing('events', events)
```

Check the function if input is a one element cell array and a multiple element cell array. Check for warnings (ID:invalid\_vector\_size) if elements have more than two columns and if not all elements have the same number of rows.

# 32.15 Testcases: scr get <datatype>

#### 32.15.1 Information

The datatype import functions are all tested in a similar way. The individual testclasses must inherit the class 'scr\_get\_superclass', from which they inherit the main test function 'valid\_datafile'. They also have to implement the property 'fhandle', which is a function handle to the specific import function.

The tests are performed with the sampledata files that are listed in the SampleDataMasterList.docx file (as at 18.11.2013).

```
Testclasses: scr_get_acq_test
scr_get_acqmat_test
scr_get_biograph_test
scr_get_biosemi_test
scr_get_biotrace_test
scr_get_brainvis_test
scr_get_edf
scr_get_labchartmat_ext_test
scr_get_labchartmat_in_test
scr_get_mat_test
scr_get_obs_test
scr_get_spike_test
scr_get_superclass
```

Superclass: scr get superclass

```
scr_get_txt_test
scr_get_vario_test
scr_get_eyelink_test
```

Function: [sts, import, sourceinfo] = scr\_get\_<datatype>(datafile, import)

#### 32.15.2 Notes

#### 32.15.3 Setup

#### define testcases

In this method the testcases are defined and the testdata is generated (if needed). Each testcase is a cell in the cellarray 'testcases'. Each testcase has the following fields:

- .pth: the path to the samplefile
- .import: the input variable

For datatypes which support blocks there has to be an additional field:

• .numofblocks

#### 32.15.4 Testcases

#### Valid datafile

Function name: valid datafile(this)

Description: The main test function, for tests with valid inputdata. It tests all testcases equally.

- 1. Test if 'sts' is equal 1.
- 2. If the datatype supports blocks, test if the number of blocks is correct.
- 3. Test if number of elements of the returned 'import' variable is correct.
- 4. Test if each import job has a field 'data', that is a numeric vector.
- 5. Test if each import job has a field 'sr', that is a number.
- 6. Test if each import job has a field 'type'.
- 7. Test if all event importions have a field 'marker'.
- 8. Test if all import jobs have duration below 1h.
- 9. Test if all import jobs have a samplerate between 1 and 10000 for continuous channels or between  $10^-6$  and 1 for timestamp channels.

#### invalid datafile

Function name: invalid datafile(this)

Description: The main test function, for tests with invalid input data.

Tests:

If the datatype supports multiple channels: Check for warning when trying to import a channel, that is not contained in the file ('ID:channel not contained in file').

# 32.16 Testcases: scr glm

#### 32.16.1 Information

Testclass: scr glm test

Function: glm = scr glm(model, options)

There are seven testcase functions. One invalid input arguments test and test 1 to 6. Tests 1 to 5 are of the same kind. There are one or multiple testcases per test function, have a look at the testcase description for more information. In these tests only Kronecker delta functions are used as basis functions, furthermore all conditions, pmods and nuisance regressors are pairwise orthogonal. The data is also not down sampled and not filtered. With these limitations it's easy to calculate the data vectors and the expected stats. For each testcase it is then tested:

- If numel(glm.names) has the expected value.
- If numel(glm.stats) has the expected value.
- If glm.stats has the expected value (with a tolerance of 1%).

In test 6 the default basis functions are used, and not all conditions and pmods are orthogonal. The data is down sampled and low and high pass filtered. In exchange the stats are not tested for correct values, just for the correct number of elements.

#### 32.16.2 Testcases

#### Invalid input arguments

Function name: invalid input (this)

Description: Checks for warnings, if the input arguments are invalid.

Input	Expected warning
scr glm(model) [no timeunits field]	ID:invalid input
scr_glm(model) [no timeunits var]	ID:invalid_input
scr_glm(model) with model.timeunits	ID:invalid_input
= 'foo' [no valid timeunits field]	_
$scr_glm(model)$ with model.timing =	ID:invalid_input
zeros(10,2) [no valid timing field]	
scr_glm(model) with model.modality	ID:invalid_input
= 'foo' [no valid modality field]	
$scr_glm(model)$ with model.channel =	ID:invalid_input
'scr' [no valid channel field]	
$scr_glm(model)$ with model.norm =	ID:invalid_input
'no' [no valid norm field]	
scr_glm(model) with model.filt.down	${ m ID:invalid\_input}$
= 'none' [filt.down is not numeric]	
scr_glm(model) with model.bf.fhandle	ID:invalid_fhandle
= 'foohandle' [non existing bf]	
scr_glm(model) with	ID:number_of_elements_dont_match
numel(model.datafile) !=	
numel(model.timing)	
scr_glm(model) with model.missing is	ID:invalid_input
struct [non valid missing field]	
scr_glm(model) with	ID:number_of_elements_dont_match
numel(model.datafile) !=	
numel(model.missing)	
scr_glm(model) with model.nuisance	ID:invalid_input
is struct [non valid nuisance field]	
scr_glm(model) with	ID:number_of_elements_dont_match
numel(model.datafile) !=	
numel(model.nuisance)	
scr_glm(model) with no R variable in	ID:invalid_input
the nuisance file	
scr_glm(model) with R	ID:number_of_elements_dont_match
variable in the nuisance	
file that has not the same	
length as the datafile	

# $\mathbf{Test}\ \mathbf{1}$

Function name: test1(this)

Description: Basic test with one basis function, one session, no nuisance regressors, no missings and one condition. Timeunits are seconds.

# ${\bf Test cases:}$

1. no pmods

- 2. one pmod
- 3. two pmods

#### Test 2

Function name: test2(this)

Description: Test with one basis function, one session, no nuisance regressors, no missings and two conditions. Timeunits are seconds.

#### Testcases:

- 1. no pmods
- 2. first condition: no pmods; second condition: one pmod
- 3. first condition: one pmod; second condition: two pmods

#### Test 3

Function name: test3(this)

Description: Test with one basis function, one session, two nuisance regressors (1Hz cosinus, 1Hz sinus), no missings, one condition and no pmods. Timeunits are seconds.

#### Testcases:

Only one testcase.

#### Test 4

Function name: test4(this)

Description: Test with one basis function, two sessions, no nuisance regressors, no missings and one condition.

#### Testcases:

- 1. timeunits are seconds
- 2. timeunits are samples
- 3. timeunits are markers

#### Test 5

Function name: test5(this)

Description: Test with two basis functions, one session, no nuisance regressors and one condition. Timeunits are seconds.

#### Testcases:

- 1. no missings
- 2. with missings

#### Test 6

Function name: test6(this)

Description: Test with default basis function and non-orthogonal conditions and

pmods

#### Testcase:

Default basis functions, no nuisance regressors, no missings, two sessions and two conditions. Timeunits are seconds.

- first condition: two pmods (with pmod(1).poly{1} = 2 and pmod(1).poly{2} = 3)
- second condition: no pmods

# 32.17 Testcases: scr import

#### 32.17.1 Information

 $Testclass: \ scr\_import\_test$ 

Function: outfile = scr import(datafile, datatype, import, options)

#### 32.17.2 Testcases

#### Invalid input arguments

Function name: invalid inputargs(ths)

Description: Checks for warnings, if the input arguments are invalid.

Test No.	Input	Expected warning
1	scr_import(datafile, datatype) [no	ID:invalid_input
	import variable]	
2	scr_import(datafile, datatype, 'foo')	ID:invalid_input
	[no cell/struct import var.]	
3	scr_import(datafile, 'foo', import)	ID:invalid_channeltype
	[invalid channeltype]	
4	scr_import(5, datatype, import) [no	ID:invalid_input
	char filename]	

#### Invalid import variable structure

Function name: invalid import struct(this)

Description: Checks for warnings, if the structure of the import variable is invalid.

#### Tests:

Test No.	Input	Expected warning
1	Multiple channel, though not supported	ID:ivalid_import_struct
2	Not allowed channeltype	ID:ivalid_import_struct
3	No sr given, though autosr is not supported	ID:ivalid_import_struct
4	Nonexistent file	ID:nonexistent_file

#### One datafile

Function name: one datafile(this)

Description: Checks the function, if datafile is a string (import of one datafile) and all inputs are correct. The outfile is checked with the scr\_load\_data function. The tests are performed with a spike samplefile and a labchartmat\_in samplefile (to check the handling of blocks).

#### Multiple datafiles

Function name: multiple datafiles(this)

Description: Checks the function, if datafile is a cell array of strings (import of multiple datafiles) and all inputs are correct. The outfiles are tested with the scr\_load\_data function.

# 32.18 Testcases: scr load data

#### 32.18.1 Information

```
Testclass: scr_load_data_test
Function: [sts, infos, data, filestruct] = scr_load_data(fn, chan)
```

#### 32.18.2 Setup

If not otherwise declared, the input variable fn is referring to a datafile which was generated with scr testdata gen and consists out of the following channels:

```
data{1}.chantype = 'scr';
data{2}.chantype = 'marker';
data{3}.chantype = 'hr';
data{4}.chantype = 'hb';
data{5}.chantype = 'marker';
data{6}.chantype = 'resp';
```

data{7}.chantype = 'scr';

The duration of the channels is 10s.

# 32.18.3 Testcases

# Invalid input arguments

Function name: invalid\_inputargs(testCase)

Description: Checks for warnings, if the input arguments are invalid.

#### Tests:

Input	Expected warning
scr_load_data [no filename]	ID:invalid_input
scr_load_data(1) [no char filename]	ID:invalid_input
scr_load_data(fn, -1) [neg. channel no]	ID:invalid_input
scr_load_data(fn, 'foobar') [no allowed ch type]	ID:invalid_input
scr_load_data(fn, foo) [missing field in foo struct]	ID:invalid_input
scr_load_data(fn, {1}) [invalid channel option]	ID:invalid_input
scr_load_data(stuct) [struct has no infos field]	ID:invalid_input

# Invalid datafile

Function name: invalid datafile(testCase)

Description: Checks for warnings, if the datafile is invalid.

Test No.	Vo. Input Expected warnin	
1	non-existent datafile ID:nonexistent_	
2	missing 'infos' variable	ID:invalid_data_structure
3	missing 'data' variable	ID:invalid_data_structure
4	missing 'data' field in 'data{2}'	ID:invalid_data_structure
5	missing 'header' field 'data{3}'	ID:invalid_data_structure
6	missing 'sr' field in 'data{7}.header'	ID:invalid_data_structure
7	7 data{4} is a nx2 vector (instead of a ID:invalid_data	
	nx1 vector)	
8	the length of data{1}.data is	ID:invalid_data_structure
	incompatible with the duration	
9	An entry of data{2}.data is larger	ID:invalid_data_structure
	than 'duration'	
10	data{5} has an non-existent chantype	ID:invalid_data_structure
	('scanner')	
11	duplicates (9) with struct chan input	ID:invalid_data_structure

#### Return all channels

Function name: valid datafile 0(testCase)

Description: Checks the function, if all channels shall be returned (chan = 0).

#### Return all channels (struct input)

Function name: valid \_datafile \_1(testCase)

Description: Checks the function, if all channels shall be returned (chan = 0) and the input is a struct.

#### Return one channel

Function name: valid datafile 2(testCase)

Description: Checks the function, if only one channel shall be returned (chan = 2).

#### Return one channel

Function name: valid datafile 3(testCase)

Description: Checks the function, if multiple channels shall be returned (chan  $= [3 \ 5]$ ).

#### Return scr channels

Function name: valid datafile 4(testCase)

Description: Checks the function, if only the scr channels shall be returned.

#### Return event channels

Function name: valid datafile 5(testCase)

Description: Checks the function, if only the event channels shall be returned.

#### Save data

Function name: valid datafile 6(testCase)

Description: Checks the function, if data is to be saved (chan struct).

# 32.19 Testcases: scr\_pp

#### 32.19.1 Information

Testclass: scr pp test

Function: newfile = scr\_pp('median', datafile, n, channelnumber) or newfile =

scr pp('butter', datafile, freq, channelnumber)

#### 32.19.2 Testcases

#### Invalid input

Function name: invalid input(this)

Description: Checks for warnings, if the input arguments are invalid.

Tests:

Input	Expected warning	
scr_pp('butter', 'file') [no freq]	ID:invalid_input	
scr_pp('foo', 'file', 100) [no valid first argument]	ID:invalid_input	
scr_pp('butter', 'file', 19) [freq below 20]	ID:invalid_input	

#### Median test

Function name: median test(this)

Description: Checks medianfilter functionality

Setup:

Testfile with 3 channels (scr, hb, scr).

Tests:

- 1. Filter one channel [Input: newfile = scr\_pp('median', testfile, 50, 3)]
  - i. Check if sts == 1, when data is loaded with scr load data.
  - ii. Check if newfile has the same number of channels as testfile
- 2. Filter multiple channel [Input: newfile = scr pp('median', testfile, 50)]
  - i. Check if sts == 1, when data is loaded with scr\_load\_data.
  - ii. Check if newfile has the same number of channels as testfile

#### Butterworth filter test

Function name: butter test(this)

Description: Checks Butterworth filter functionality

Setup:

Testfile with 3 channels (scr, hb, scr).

- 1. Filter one channel [Input: newfile = scr\_pp('butter', testfile, 40, 3)]
  - i. Check if sts == 1, when data is loaded with scr\_load\_data.
  - ii. Check if newfile has the same number of channels as testfile

- 2. Filter multiple channel [Input: newfile = scr\_pp('butter', testfile, 40)]
  - i. Check if sts == 1, when data is loaded with scr\_load\_data.
  - ii. Check if newfile has the same number of channels as testfile

# 32.20 Testcases: scr prepdata

# 32.20.1 Information

```
Testclass: scr_prepdata_test
Function: [sts, outdata, newsr] = scr_prepdata(data, filt)
```

#### 32.20.2 Testcases

#### Invalid input

Function name: invalid input(this)

Description: Checks for warnings, if the input arguments are invalid.

#### Tests:

Input	Expected warning	
scr_prepdata([1 2 3]) [no filt variable]	ID:invalid_input	
scr_prepdata(data, filt) [filt has no hporder field]	ID:invalid_input	
scr_prepdata('foo', filt) [no numeric data]	ID:invalid_input	
scr_prepdata(data, filt) [with lpfreq = 'foo' (not valid)]	ID:invalid_input	

# Hipassfilter test

```
Function name: hipassfilter_test(this)
```

Description: Checks hipassfilter functionality (without downsampling)

#### Setup:

```
data = rand(1000,1);
filt.sr = 100;
filt.lpfreq = 'none';
filt.lporder = 1;
filt.hpfreq = 20;
filt.hporder = 1;
filt.down = 'none';
```

- 1. Unidirectional tests [filt.direction = 'uni']
  - i. Check if sts == 1
  - ii. Check if newsr == filt.sr
  - iii. Check if outdata is empty

```
iv. Check if length(outdata) == length(data)
2. Unidirectional tests [filt.direction = 'bi']
i. Check if sts == 1
ii. Check if newsr == filt.sr
iii. Check if outdata is empty
iv. Check if length(outdata) == length(data)
```

#### Lowpassfilter test

```
Function name: lowpassfilter_test(this)
Description: Checks hipassfilter functionality (without downsampling)
```

```
Setup:

data = rand(1000,1);

filt.sr = 100;

filt.lpfreq = 40;

filt.lporder = 1;

filt.hpfreq = 'none';

filt.hporder = 1;

filt.down = 'none';
```

Tests:

Same tests as in hipassfilter\_test. Additionally there is a check for a warning if filt.lpfreq is higher (or equal) than the nyquist frequency:

Input	Expected warning
$scr_prepdata(data, filt)$ [filt.sr = 100; filt.lpfreq = 60]	ID:no_low_pass_filtering

#### Bandpassfilter test

```
Function name: bandpassfilter_test(this)
Description: Checks bandpassfilter functionality (without downsampling)
```

```
Setup:

data = rand(1000,1);

filt.sr = 200;

filt.lpfreq = 99;

filt.lporder = 1;

filt.hpfreq = 20;

filt.hporder = 1;

filt.down = 'none';
```

Tests: Same tests as in hipassfilter test.

#### Integer samplerate ratio downsampling test

```
Function name: int sr ratio downsample test(this)
Description: Checks downsampling functionality, if the ratio between filt.sr and
filt.down is an integer.
Setup:
ratio = 2; %ratio between filt.sr and filt.down
filt.down = 100;
filt.sr = ratio
filt.down; filt.lpfreq = 40;
filt.lporder = 1;
filt.hpfreq = 'none';
filt.hporder = 1;
filt.direction = 'uni';
data = rand(filt.sr * 10,1);
Tests:
  1. Check if sts == 1
  2. Check if newsr == filt.down
  3. Check if outdata is empty
```

# 32.21 Testcases: scr pulse convert

4. Check if ratio\*length(outdata) == length(data)

#### 32.21.1 Information

```
\label{lem:convert_test} Testclass: scr_pulse\_convert\_test\\ Function: wavedata = scr_pulse\_convert(pulsedata, resamplingrate, samplingrate)
```

#### 32.21.2 Testcases

#### Invalid input

```
Function name: invalid input(testCase)
```

Description: Pass invalid input arguments and test if the error message is correct.

Input	Expected warning	
scr_pulse_convert()	ID:invalid_input	
scr_pulse_convert(10^-3 * (1:10000)')	ID:invalid_input	
scr_pulse_convert(10^-3 * (1:10000)', 10000)	ID:invalid_input	

#### Valid input

Function name: valid\_input(testCase)

Description: Pass generated, valid data and test if function issues no warning.

#### Tests:

1. Test function without downsampling the data

2. Test function with downsampling the data

# 32.22 Testcases: scr ren

# 32.22.1 Information

Testclass: scr ren test

Function: out newfilename = scr ren(filename, newfilename)

#### 32.22.2 Testcases

#### Invalid input

Function name: invalid input (this)

Description: Checks for warnings, if the input arguments are invalid.

# Tests:

$\sim$	05.		
	Input	Expected warning	
	scr_ren('fn') [no newfilename]	ID:invalid_input	
	scr_ren({'fn1', 'fn2'}, {'rfn1', 'rfn2', 'rfn3'}) [non same size cell arrays]	ID:invalid_input	

#### Char Valid Input

Function name: char\_valid\_input (this)

Description: Checks the function if the input variables are of type char. It uses scr load data to check the files.

- 1. Check if  $out_newefilename = newfilename$
- 2. Check if sts==1 (of scr load data output)
- 3. Check if the field 'infos.rendata' exists
- 4. Check if the field 'infos.newname' exists

5. Check if the original file has been deleted

#### Cell Valid Input

Function name: cell valid input (this)

Description: Checks the function if the input variables are of type cell. It uses scr load data to check the files.

Tests:

The inputs are two-element cell arrays. For both elements the same tests as in the char valid input function are performed individually.

# 32.23 Testcases: scr split sessions

#### 32.23.1 Information

Testclass: scr split sessions test

Properties:  $expected_number_of_files = 3;$ 

Function: newdatafile = scr split sessions(datafile, markerchannel, options)

#### 32.23.2 Setup

For the tests a testdatafile with three channels is used (duration is 100s). The markerchannel data is:

 $data = [1 \ 4 \ 9 \ 12 \ 30 \ 31 \ 34 \ 41 \ 43 \ 59 \ 65 \ 72 \ 74 \ 80 \ 89 \ 96]$ 

Hence if MAXSN=10 & BRK2NORM=3 (default values) the datafiles should be split into 3 files. If different values are being used, update the property 'expected number of files' of the testclass object accordingly.

#### 32.23.3 Testcases

# Invalid input

Function name: invalid\_input (this)

Description: Checks for warnings, if the input arguments are invalid.

Input	Expected warning	
scr_split_sessions() [no filename]	ID:invalid_input	
scr_split_sessions (2) [no string filename]	ID:invalid_input	
scr_split_sessions ('fn', 'foo') [no numeric marker channel no.]	ID:invalid_input	

#### One datafile

Function name: one datafile(this)

Description: Checks the function if the variable 'datafile' is of type char (one datafile). The markerchannel number is not assigned explicitly.

#### Tests:

- 1. Check if the file has been split into 'expected\_number\_of\_files' files For each output file the following tests are performed:
- 2. Check if sts == 1, when data is loaded with scr\_load\_data.
- 3. Check if number of channels is correct.
- 4. Check it the field infos.slitdate exists
- 5. Check if the field infos.splitsn exists
- 6. Check if the field infos.splitfile exists.

# Multiple datafiles

Function name: multiple datafiles(this)

Description: Checks the function if the variable 'datafile' is of type cell (two datafiles). The marker channel number is assigned explicitly.

#### Tests:

For both datafiles the same tests as in the one\_datafile function are performed individually. Additionally it is tested if the number of input files does match the number of output files.

# 32.24 Testcases: scr\_trim

# 32.24.1 Information

```
Testclass: scr_trim_test
Function: newdatafile=scr_trim(datafile, from, to, reference, options)
```

#### 32.24.2 Setup

If not otherwise declared, the input variable fn is referring to a datafile which was generated with scr\_testdata\_gen and consists of the following channels:

```
data{1}.chantype = 'scr';
data{2}.chantype = 'marker';
data{3}.chantype = 'hr';
data{4}.chantype = 'hb';
```

```
data{5}.chantype = 'marker';
data{6}.chantype = 'resp';
data{7}.chantype = 'scr';
```

The duration of the data recording is 10s.

#### 32.24.3 Testcases

#### Invalid input arguments

Function name: invalid inputargs(testCase)

Description: Checks for warnings, if the input arguments are invalid.

П	Pests

Input	Expected warning
scr_trim(testCase.fn, [1 2], 5,	ID:invalid_input
'marker') [invalid from parameter]	
scr_trim(testCase.fn, 0, 'bla',	ID:invalid_input
'marker') [invalid to parameter]	
scr_trim(testCase.fn, 0, '[]', 'marker')	ID:invalid_input
[invalid to parameter]	
$\operatorname{scr\_trim}(\operatorname{fn}, 0, 5)$ [no reference]	ID:invalid_input
$\operatorname{scr\_trim}(\operatorname{fn}, 0, 5, 6)$ [no char or	ID:invalid_input
2-element numeric reference]	
scr_trim(fn, 0, 5, 'bla') [invalid char	ID:invalid_input
reference]	
scr_trim(fn, 0, 5, [-1 5]) [invalid	ID:invalid_input
numeric start reference]	
scr_trim(fn, 0, 5, [5 4]) [invalid	ID:invalid_input
numeric start/end reference]	

#### 32.24.4 Reference = 'marker' tests

Function name: marker tests(testCase)

Description: A wrapper function for tests with reference = 'marker'. It executes the methods markertest  $\_k$ , where the testcases are defined.

#### markertest 1

Description: from and to are set so that the trimming points are out of the range [0,duration]. Hence the data should not be trimmed.

Expected warning: ID: marker\_out\_of\_range

Input: scr\_trim(fn, -20, 20, 'marker')

# $\mathbf{markertest} \_\mathbf{2}$

Description: from and to are set so that the trimming points are exactly (0, duration). Hence the data should not be trimmed.

Input: from = -1 \* marker(1) to = duration - marker(end) scr\_trim(fn, from, to, 'marker')

#### markertest 3

Description: from and to are set so that the trimming points in the range [0,duration].

Input: scr trim(fn, 1, -2, 'marker')

#### 32.24.5 Reference = 'file' tests

Function name: file tests(testCase)

Description: A wrapper function for tests with reference = 'file'. It executes the methods filetest\_k, where the testcases are defined.

#### filetest 1

Description: from and to are set so that the trimming points are out of the range [0,duration]. Hence the data should not be trimmed.

Expected warning: ID: marker out of range

Input: scr trim(fn, -12.5, 50, 'marker')

#### filetest 2

Description: from and to are set so that the trimming points are exactly (0, duration). Hence the data should not be trimmed.

Input: scr trim(fn, 0, duration, 'marker')

#### filetest 3

Description: from and to are set so that the trimming points in the range [0,duration].

Input: scr trim(fn,2.1, duration – 2.5, 'marker')

#### Numeric reference tests

Function name: num tests(testCase)

Description: A wrapper function for tests with reference = [a b] (a, b are two integers with a<br/>b). It executes the methods markertest\_k, where the testcases are defined.

#### numtest 1

Description: from and to are set so that the trimming points are out of the range [0,duration]. Hence the data should not be trimmed.

Expected warning: ID: marker out of range

Input: scr\_trim(fn, -20, 20, [2 14])

#### numtest 2

Description: from and to are set so that the trimming points are exactly (0, duration). Hence the data should not be trimmed.

Input: from = -1 \* marker(3) to = duration - marker(8) scr\_trim(fn, from, to, [3 8])

#### numtest 3

Description: from and to are set so that the trimming points in the range [0,duration].

Input: scr  $trim(fn, -1.5, 2, [2\ 7])$ 

#### numtest 4

Description: Second reference point is out of the marker range; from is set to 'none'. Hence the data should not be trimmed.

Expected warning: ID: marker out of range

Input: scr trim(fn, 'none', 0, [1 (numel(marker) + 1)])

#### Multiple file reference tests

 $Function\ name:\ multiple\_files(testCase)$ 

Description: The input variable datafile is either a cell array of two filenames or a cell array of two stucts. In both cases it is tested whether the return value is also a cell array of two filenames and whether both files are trimmed correctly.

# Options tests

#### Marker channel number option

Function name: marker chan num option test(testCase)

Description: Tests if the option marker\_chan\_num is working correctly. There are two tests: Test 1: Checks for a warning if the selected channel is no marker channel. Test 2: Checks if the selected channel is actually used.

# 33 External functions and tools

# 33.1 VB (Variational Bayes) inversion algorithm by Jean Daunizeau

Updated October 2014

Changes made for use in PsPM:

 $VBA\_ReDisplay.m$ , fixed try-catch syntax in various places by adding a comma after "try" to avoid warning in matlab > 2007

 $VBA\_inv.m$ , line 42: added warning off/on to suppress the warning "Matrix is singular, close to singular or badly scaled. Results may be inaccurate. RCOND = NaN."

# 34 List of functions

Name	Main author	Test exists	Test Doc
f_SCR	Dominik Bach & Jean	-	-
	Daunizeau		
f_SF	Dominik Bach	-	-
g_SCR	Dominik Bach	-	-
pspm	Dominik Bach	X	x
scr	Dominik Bach	X	x
scr_add_channel	Dominik Bach	X	-
scr_align_channels	Dominik Bach	-	-
scr_axpos	Dominik Bach	-	-
scr_bf_brf	Saurabh Khemka &	-	-
	Dominik Bach		
scr_bf_FIR	Dominik Bach	-	-
scr_bf_Fourier	Dominik Bach	-	-
scr_bf_hprf	Dominik Bach	-	-
scr_bf_scrf_f	Dominik Bach	-	-
$scr_bf_scrf$	Dominik Bach	-	
scr_butter	Dominik Bach	-	-
scr_con1	Dominik Bach	-	-

	D : 1 D 1	T	I
scr_con2	Dominik Bach	-	-
scr_contrast	Dominik Bach	-	-
scr_dcm_inv	Dominik Bach	-	-
scr_dcm	Dominik Bach	-	-
scr_denoise_spike	Dominik Bach	-	-
scr_display	Philipp C Paulus	-	-
scr_down	Dominik Bach	-	-
scr_downsample	Dominik Bach	-	-
scr_ecg2hb	Philipp C Paulus	X	x
scr_exp	Dominik Bach	_	_
scr_filtfilt	Dominik Bach	-	-
scr_find_channel	Dominik Bach	X	x
scr_get_acq	Dominik Bach	x	x
scr_get_acqmat	Dominik Bach	x	x
scr_get_biograph	Dominik Bach	x	Х
scr_get_biosemi	Dominik Bach	X	x
scr_get_biotrace	Dominik Bach	X	x
scr_get_brainvis	Dominik Bach	X	х
scr_get_cell	Dominik Bach	-	-
scr_get_cnt	Dominik Bach	-	-
scr get custom	Tobias Moser	-	-
scr_get_ecg	Dominik Bach	X	х
scr_get_edf	Tobias Moser	X	x
scr get events	Dominik Bach	X	х
scr get eyelink	Christoph Korn, Tobias	X	х
	Moser		
scr_get_hb	Dominik Bach	X	x
scr_get_hp	Dominik Bach	-	-
scr_get_hr	Dominik Bach	X	х
scr_get_labchartmat_ext	Dominik Bach	X	х
scr_get_labchartmat_in	Dominik Bach	X	х
scr_get_marker	Dominik Bach	X	х
scr_get_markerinfo	Dominik Bach	-	-
scr_get_mat	Dominik Bach	X	х
scr_get_obs	Linus Rüttimann	X	х
scr get pupil	Dominik Bach	X	х
scr get resp	Dominik Bach	X	х
scr get rf	Dominik Bach	-	-
scr_get_rp	Dominik Bach	-	-
scr_get_rr	Dominik Bach	-	-
scr_get_scr	Dominik Bach	X	X
scr get spike	Dominik Bach	X	X
scr_get_timing	Dominik Bach	X	Х

scr_get_txt	Dominik Bach	X	X
scr_get_vario	Dominik Bach	X	Х
scr_get_wdq	Dominik Bach	-	-
$scr\_get\_wdq\_n$	Tobias Moser	X	X
scr_glm_recon	Dominik Bach	-	-
scr_glm	Dominik Bach	X	X
scr_hb2hp	Dominik Bach	-	-
scr_hb2hr	Dominik Bach	-	-
scr_import	Dominik Bach	X	X
scr_init	Dominik Bach	-	-
scr_jobman	Gabriel Gräni	-	-
scr_job_create	Dominik Bach	-	-
scr_load_data	Dominik Bach	X	X
scr_load1	Dominik Bach	-	-
scr merge	Dominik Bach	-	-
scr_pp	Dominik Bach	X	X
scr predval	Dominik Bach	-	-
scr prepdata	Dominik Bach	X	X
scr pulse convert	Dominik Bach	X	-
scr quit	Dominik Bach	-	-
scr ren	Dominik Bach	X	X
scr resp2rp	Dominik Bach	-	-
scr resp2rr	Dominik Bach	-	-
scr_rev_con	Dominik Bach	-	-
$\operatorname{scr}\operatorname{\underline{-rev}}\operatorname{\underline{dcm}}$	Dominik Bach	-	-
scr_rev_glm	Dominik Bach	-	-
scr_rev2	Dominik Bach	-	-
scr review	Gabriel Graeni	-	-
scr rewrite channel	Samuel Gerster	-	-
scr sf auc	Dominik Bach	-	-
scr sf dcm	Dominik Bach	-	-
scr sf mp	Dominik Bach	-	-
scr sf scl	Dominik Bach	-	-
scr sf theta	Dominik Bach	-	-
scr sf	Dominik Bach	-	-
scr show arms	Dominik Bach	-	-
scr spike convert	Dominik Bach	-	-
scr split sessions	Linus Rüttimann	x	X
scr transfer function	Dominik Bach	-	-
scr_trim	Dominik Bach	X	X
scr_ledalab	Dominik Bach	-	-
scr_peakscore	Dominik Bach	-	-
scr_sf_get_theta	Dominik Bach	-	-
	I	_ I	I .

scr transfer fit	Dominik Bach	-	-	ĺ
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# Part V

# Release notes

# 35 PsPM Version 3.0.1

# **Import**

# New data types were implemented

- DATAQ Windaq (no ActiveX needed)
- European Data Format (EDF)

# **Tools**

'Preprocess respiration traces' replaces 'Convert Respiration to Respiration Period'. It supports conversions into the following datatypes:

- Respiration preriod (old funtionality)
- Respiration amplitude
- Respiration line length
- Respiration time stamp

# First level contrast

#### Z-score parameter estimates

The function scr\_con1 now supports z-scoring trial-by-trial parameter estimates. If selected, all parameter estimates for the same event type, across all trials, will be z-scored before computing the contrast. This option is currently only available for non-linear models.

# Review first level model

Next to the regressors on the x- and y-axes, orthogonality plots newly display regressor names along the y-axes.

# 36 PsPM Version 3.0

# **Import**

# New data types were implemented

- Noldus Observer compatible
- Eyelink

# Untested data types

CNT data import has not been not tested – please contact the developers with sample data files.

# Filtering for SCR models

Previous versions of PsPM have used a bi-directional high pass filter of 0.0159 for all SCR analyses. We have recently shown a better predictive validity for GLM with a unidirectional filter of 0.05 Hz. This also implies that different filters are used for different models. These are now set as defaults, and the way the default settings are implemented has changed. It is now possible to alter the filter settings in the model definition, although we discourage this.

# New SF method

A matching pursuit algorithm is now implemented to approximate the number of spontaneous fluctuations (SF) in skin conductance [18].

# General linear modelling (GLM)

#### Parametric modulators

Parametric modulators (pmods) are z-normalised before being entered into the design matrix. This is to account for possibly very large or very small numbers – a badly scaled design matrix can cause induced instability in the inversion. The parameter estimates of the pmods were not transformed back in previous versions, i. e. the parameter estimates of the pmods were independent of the scaling of the pmods. This is appropriate as long as they are the same for all datasets, or if analysis is done strictly on a within-subject level. Some researchers have reported designs in which inference was to be drawn on parameter estimates of pmods on a between-group level, and where the pmods systematically differed between these groups. To account for this possibility, the parameter estimates are now transformed back, such that they refer to the pmods in their original scaling/units.

#### Parametric confounds

Previous versions of PsPM contained an option to include a parametric modulator across all event types, to account for confounds across all conditions. For example, in an experiment with 5 conditions, one could have included 5 regressors, plus one reaction time confound across all events, without including an associated regressor that contains the event onsets for all these events. This option was removed.

# Design matrix filtering

Previous versions of PsPM filtered the design matrix after orthogonalisation of basis sets. This can introduce unwanted dependencies between regressors. PsPM 3.0 filters the regressors first, then orthogonalises the basis sets.

# File format

Some minor changes have been made to the data format. In particular, marker channels from previous versions can not be read with the current version - such data files have to be re-imported. Model files have changed drastically, and model files from previous versions can not be read with the current version of the software.

#### **VB** inversion

The VBA toolbox (http://code.google.com/p/mbb-vb-toolbox/) was updated in October 2014 [16]. This update incorporates bugfixes in this toolbox and slightly changed the model estimates in our test models. In terms of predictive validity, we noted that there was no consistent benefit of the old or new version of this code (Figure 36.1).

#### Part VI

# Acknowledgments

SCRalyze versions 1-2 were developed at the FIL, Wellcome Trust Centre for Neuroimaging, University College London, under the auspices of Karl J. Friston and Raymond J. Dolan. Guillaume Flandin contributed to GLM, and Jean Daunizeau to DCM. Eric Featherstone, Alfonso Reid, Oliver Josephs, and Chloe Hutton helped with the import functions. This development was funded by the Swiss National Science Foundation with a personal grant to Dominik R. Bach, and by the Wellcome Trust with a programme grant to Raymond J. Dolan.

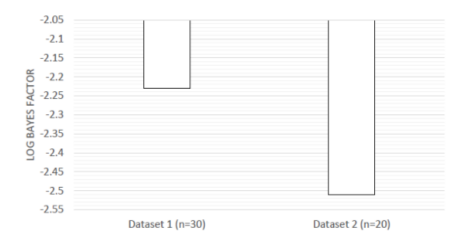


Figure 36.1: Model comparison between old and new versions of the VBA toolbox, based on two delay fear conditioning datasets. The log Bayes factor quantifies the difference between negative log likelihood (nLL) of parameter estimates obtained from model inversion using the old and new version of VBA. A difference in nLL above 3 indicates significant differences in model evidence which is not exceeded for either data set.

PsPM 3.0 was developed at the University of Zurich, funded by the University of Zurich and by the Swiss National Science Foundation with a project grant.

PsPM uses a number of SPM functions. For import of spike data, PsPM makes use of the SON library written by Malcolm Lidierth at King's College London, and of functions provided by the SPM physiology toolbox, written by Chloe Hutton and Eric Featherstone at the FIL. Biopac AcqKnowledge import uses a routine created by Sebastien Authier and Vincent Finnerty at the University of Montreal. The DCM inversion uses a variational Bayesian inversion scheme written by Jean Daunizeau.

The Matlabbatch GUI was written by Gabriel Gräni at University of Zurich. Help texts and tutorial were developed by Christoph Korn and Matthias Staib. The data display was written by Philipp C Paulus, University of Dresden. The test environment was contributed by Linus Rüttimann, and extended by Tobias Moser, both University of Zurich.

# Part VII

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