### cbm manual

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This document serves as the manual of cbm (computational and behavioral modeling) toolbox. cbm provides tools for hierarchical Bayesian inference (HBI). See the corresponding manuscript (here) for more information about the HBI and its comparison with other methods. You can download cbm here. In this manual, I present two well-known examples and explain how cbm tools should be used to perform Bayesian model comparison, parameter estimation and inference at the group level. To be able to follow this manual, you need to be familiar with matlab syntax (cbm will be published soon in other languages, particularly python).

### 1 Why a new method?

Althoguh I don't aim here to demonstrate benefits of the HBI over other methods, I would like to give a hint, which might motivate you to use the HBI (see the paper for theoretical and experimental observations). Bayesian methods are typically used for both parameter estimation as well as model comparison. Regarding the former, methods either perform heirarchical parameter estimation (i.e. building empirical priors based on group data) or non-hierarchical parameter estimation (i.e. fitting each subject individually). Of course, hierarchical parameter estimation is more accurate (and much less prone to subjective prior tuning). Regarding the model comparison, methods either take the model identity as a fixed effects (essentially making a very strong assumption that the same model is the true model for all subjects, so any difference between subjects is mereley due to noise) or as a random effects (different models might be true for different subjects). Of course, the random effects model comparison is better (particularly because of its robustness against outliers). The problem is, there is no method that does both! The HBI performs hierarhical parameter estimation and random effects model comparison, which improves both parameter estimation (even over previous hierarchical methods) and model comparison (over both previous hierarchical methods, or non-hierarchical but random effects methods). If you would like to know more about these issues, please take a look at the paper (in particular, figures 1-3).

## 2 Example 1: bandit task

I assume that the current directory contains this manual and "codes" directory, as well as "example\_RL\_task" and "example\_2step\_task" directories. The codes directory contains all cbm matlab functions.

For using cbm tools, you need to know what is your model-space and code your own models. You can then use cbm to make inference. For this example, models code and data have been stored in the example\_RL\_task directory.

Suppose you have 40 subjects' choice data in a 2-armed bandit task, in which subjects chose between two actions and received binary outcomes.

All data have been stored in a mat-file called all\_data.mat in a cell format, in which each cell contains choice data and outcomes for one subject. First, enter the example\_RL\_task directory and then load those data:

For example, subj1 contains information of subject 1: subj1.actions are all actions and subj1.outcome are the corresponding outcomes.

Also suppose you have 2 candidate models. An RL model, which has a learning rate (alpha) and a decision noise (beta) parameter. The other model is a dual-alpha RL model with two separate learning rates for positive and negative prediction errors (alpha1 and alpha2, respectively) and a decision noise (beta) parameter. See matlab functions model\_RL and model\_dualRL as examples.

It is important to remember that cbm does not care how your model works! It assumes that your models take parameters and data (of one subject) as input and produce a log-probability of data given the parameters (i.e. log-likelihood function). Tools in cbm only require that the input and output of models follow a specific format:

```
loglik = model(parameters,subj)
For example, for the model_RL, we have:
```

loglik = model\_RL(parameters,subj1)

here parameters is a vector, which its size depends on the number of parameters of the model\_RL (for model\_RL, its size is 2). subj1 is the structure containing data of subject 1 as indicated above. loglik is the log-likelihood of subj1 data given parameters, as computed by model\_RL.

We now use cbm tools to fit models to data. For this example, we use the two models implemented in model\_RL and model\_dualRL together with the data stored in all\_data.mat. Data stored in all\_data is a synthetic dataset of 40 subjects. The data of the first 10 subjects are generated by model\_RL and the data of the next 30 subjects are generated by model\_dualRL. Note, however, that cbm is not meant to provide a collection of different computational models. You should code your models for questions of your interest yourself. The tools in cbm fit your models to your data and compare the models given the data.

First, make sure that cbm is added to your matlab path and then load the data:

```
fdata = load('all_data.mat');
data = fdata.data;
```

Before using cbm tools, it is always good to check whether the format of the models is compatible with the cbm. To do that, create a random vector parameters and call the models:

Note that because parameters were randomly drawn, when you run the same code, it generates different values for F1 and F2. Also note that parameters are drawn from a normal distribution. In theory, RL models require constrained parameters (e.g. alpha between 0 and 1). In model\_RL and model\_dualRL, some transformations applied to the normally-distributed parameters to meet their theoretical bounds (see model\_RL and model\_dualRL). I'll explain later a bit more about transforming (normally-distributed) parameters in your models.

We checked the models with some random parameters. It is also good to check them with more extreme parameter values:

Again, F1 and F2 should be real negative scalers.

After making sure that model\_RL and model\_dualRl work fine, we now use cbm tools to fit models to data.

First, we should run cbm\_lap, which fits every model to each subject data separately (i.e. in a non-hierarchical fashion). cbm\_lap employs Laplace approximation, which needs a normal prior for every parameter. We set zero as the prior mean. We also assume that the prior variance for all parameters is 6.25. This variance is large enough to cover a wide range of parameters with no excessive penalty (see supplementary materials of the reference article for more details on how this variance is calculated).

```
In [6]: v = 6.25;
    prior_RL = struct('mean',zeros(2,1),'variance',v); % note dimension of 'mean'
    prior_dualRL = struct('mean',zeros(3,1),'variance',v); % note dimension of 'mean'
```

We also need to specify a file-address for saving the output of each model:

```
In [7]: fname_RL = 'lap_RL.mat';
    fname_dualRL = 'lap_dualRL.mat';
```

Now we run cbm\_lap for each model. Note that model\_RL and model\_dualRL are both in the current directory.

```
In [8]: cbm_lap(data, @model_RL, prior_RL, fname_RL);
        % Running this command, prints a report on your matlab output
        % (e.g. on the command window)
cbm_lap
                                                   18-Jun-2019 22:04:28
Number of samples: 40
Number of parameters: 2
Number of initializations: 14
Subject: 01
Subject: 02
Subject: 03
Subject: 04
Subject: 05
Subject: 06
Subject: 07
Subject: 08
Subject: 09
Subject: 10
Subject: 11
```

```
Subject: 12
Subject: 13
Subject: 14
Subject: 15
Subject: 16
Subject: 17
Subject: 18
Subject: 19
Subject: 20
Subject: 21
Subject: 22
Subject: 23
Subject: 24
Subject: 25
Subject: 26
Subject: 27
Subject: 28
Subject: 29
Subject: 30
Subject: 31
Subject: 32
Subject: 33
Subject: 34
Subject: 35
Subject: 36
Subject: 37
Subject: 38
Subject: 39
Subject: 40
done :]
  Also run cbm_lap for model_dualRL
In [9]: cbm_lap(data, @model_dualRL, prior_dualRL, fname_dualRL);
       % Running this command, prints a report on your matlab output
       % (e.g. on the command window)
                                               18-Jun-2019 22:04:35
cbm_lap
______
Number of samples: 40
Number of parameters: 3
Number of initializations: 21
Subject: 01
Subject: 02
Subject: 03
```

```
Subject: 04
Subject: 05
Subject: 06
Subject: 07
Subject: 08
Subject: 09
Subject: 10
Subject: 11
Subject: 12
Subject: 13
Subject: 14
Subject: 15
Subject: 16
Subject: 17
Subject: 18
Subject: 19
Subject: 20
Subject: 21
Subject: 22
Subject: 23
Subject: 24
Subject: 25
Subject: 26
Subject: 27
Subject: 28
Subject: 29
Subject: 30
Subject: 31
Subject: 32
Subject: 33
Subject: 34
Subject: 35
Subject: 36
Subject: 37
Subject: 38
Subject: 39
Subject: 40
done :]
   Let's take a look at the file saved by the cbm_lap:
In [10]: fname = load('lap_RL.mat');
              = fname.cbm;
         % look at fitted parameters
         cbm.output.parameters
```

ans =

-0.9099	-0.1596
-2.2959	0.2281
-1.8879	0.0641
-2.9416	0.6446
0.1626	-1.7082
-2.3983	-0.2423
-2.2385	-0.3139
-1.1035	-0.1045
-0.1508	-1.0967
-3.1319	0.3766
-1.6312	2.8806
-1.9657	1.4585
1.3684	1.3809
0.2462	1.2869
0.5822	1.6409
0.1195	0.7961
0.2869	1.3080
-0.9798	2.4429
-0.2855	1.3087
0.6944	2.0278
1.3887	1.4371
0.0819	1.4371
-0.5053	1.9617
0.4317	
	1.4263
0.1695	0.8347
0.6204	-0.8997
0.0504	2.2729
-0.2048	-0.2258
0.4743	1.2425
0.5359	0.7142
1.0363	0.0334
0.7106	0.2387
-1.4016	1.7652
0.0332	2.4478
-0.8369	1.4053
-0.0262	0.6052
0.1145	1.2742
0.3787	1.6922
0.0006	2.0756
1.6417	0.6821

Note that these values are normally-distributed parameters (I'll explain later how to obtain bounded parameters). The order of parameters depend on how they have been coded in the corresponding model. In model\_RL, the first parameter is alpha and the second one is beta. Therefore, here the first column corresponds to alpha and the second one to beta.

Now we can do hierarchical Bayesian inference using cbm\_hbi. cbm\_hbi needs 4 inputs. The good news is that you already have all of them!

```
In [11]: % 1st input: data for all subjects
         fdata = load('all_data.mat');
         data = fdata.data;
         % 2nd input: a cell input containing function handle to models
         models = {@model_RL, @model_dualRL};
         % note that by handle, I mean @ before the name of the function
         % 3rd input: another cell input containing file-address to files saved by cbm_lap
         fcbm_maps = {'lap_RL.mat', 'lap_dualRL.mat'};
         % note that they corresponds to models (so pay attention to the order)
         % 4th input: a file address for saving the output
         fname_hbi = 'hbi_RL_dualRL.mat';
   Now, we are ready to run cbm_hbi:
In [12]: cbm_hbi(data,models,fcbm_maps,fname_hbi);
         % Running this command, prints a report on your matlab output
         % (e.g. on the command window)
                                                   18-Jun-2019 22:04:43
cbm_hbi_hbi
Running hierarchical bayesian inference (HBI)...
HBI has been initialized according to
        lap_RL.mat [for model 1]
        lap_dualRL.mat [for model 2]
Number of samples: 40
Number of models: 2
Iteration 01
Iteration 02
        model frequencies (percent)
        model 1: 50.5 | model 2: 49.5 |
                                                            dL:
                                                                   6.62
                                                                  49.48
                                                            dm:
                                                                   0.22
                                                            dx:
Iteration 03
        model frequencies (percent)
        model 1: 48.6 | model 2: 51.4 |
                                                            dL:
                                                                   2.14
                                                            dm:
                                                                   1.91
                                                            dx:
                                                                   0.12
Iteration 04
        model frequencies (percent)
```

	4 40 01 1 1 0 50 71		
model	1: 46.3   model 2: 53.7	17	2 00
		dL:	3.09
		dm:	2.30
T++ 0F		dx:	0.13
Iteration 05	fraguancias (nament)		
	frequencies (percent) 1: 43.3   model 2: 56.7		
model	1. 43.3  Model 2. 30.7	dL:	4.28
		dm:	3.04
		dm:	0.16
Iteration 06		ux.	0.10
	frequencies (percent)		
	1: 39.4 model 2: 60.6		
modol	1. 66.1; model 2. 66.6;	dL:	4.91
		dm:	3.84
		dx:	0.18
Iteration 07			0.10
	frequencies (percent)		
	1: 35.6  model 2: 64.4		
		dL:	4.10
		dm:	3.81
		dx:	0.17
Iteration 08			
model	frequencies (percent)		
model	1: 32.7  model 2: 67.3		
		dL:	3.42
		dm:	2.93
		dx:	0.15
Iteration 09			
model	frequencies (percent)		
model	1: 31.2  model 2: 68.8		
		dL:	2.08
		dm:	1.47
		dx:	0.11
Iteration 10			
	frequencies (percent)		
model	1: 30.5  model 2: 69.5		
		dL:	1.40
		dm:	0.72
		dx:	0.12
Iteration 11			
	frequencies (percent)		
model	1: 30.0  model 2: 70.0	ar .	0.00
		dL:	0.99
		dm:	0.53
Iteration 12		dx:	0.10
	frequencies (percent)		
moder	ireductes (hercent)		

model	1: 29.6   model 2: 70.4		
		dL:	0.65
		dm:	0.34
		dx:	0.08
Iteration 13			
	frequencies (percent)		
model	1: 29.4   model 2: 70.6	dL:	0.44
		dm:	
		dx:	
Iteration 14			
model	frequencies (percent)		
model	1: 29.3   model 2: 70.7		
		dL:	0.30
		dm: dx:	
Iteration 15		ux:	0.04
	frequencies (percent)		
	1: 29.2  model 2: 70.8		
		dL:	0.21
		dm:	
		dx:	0.03
Iteration 16	f		
	frequencies (percent) 1: 29.2   model 2: 70.8		
moder	1. 23.2  model 2. 70.0	dL:	0.15
		dm:	
		dx:	0.02
Iteration 17			
	frequencies (percent)		
model	1: 29.1   model 2: 70.9		0 11
		dL: dm:	0.11
		dm: dx:	0.02
Iteration 18		<b>411</b>	0.01
model	frequencies (percent)		
model	1: 29.1   model 2: 70.9		
		dL:	0.08
		dm:	0.01
Iteration 19		dx:	0.01
	frequencies (percent)		
	1: 29.1   model 2: 70.9		
		dL:	0.06
		dm:	0.00
		dx:	0.01
		Conver	ged :]

Runnig cbm\_hbi writes a report on your standard output (often the screen). On every iteration, cbm\_hbi reports model frequency, which is the estimate of how many individual datasets (i.e. subjects) is explained by each model (in percent). Furthermore, on every iteration, there are 3 metrics showing the changes relative to the previous iteration: dL is the change in the log-likelihood of all data given the model space (more specifically a variational approximation of log-likelihood); dm is the (percentage of) change in model frequencies; dx indicates changes in (normalized value of) parameters. Although either of these measures or their combination can be used as stopping criteria, cbm\_hbi uses dx as the stopping criteria. By default, cbm\_hbi stops when dx<0.01.

Let's now take a look at the saved file:

Almost all useful parameters are stored in cbm.output, which we explain them here.

First, let's look at model\_frequency, which is the HBI estimate of how much each model is expressed across the group:

```
In [14]: model_frequency = cbm.output.model_frequency
model_frequency =
    0.2913    0.7087
```

Note that the model\_frequency is normalized here (so it sums to 1 across all models). Also note that the order depends on the order of models fed into to HBI as input. Therefore, in this example, HBI estimated that about 29% of all subjects are explained by the model\_RL and about 71% by the model\_dualRL.

Now let's take a look at estimated group mean stored in cbm.output.group\_mean.

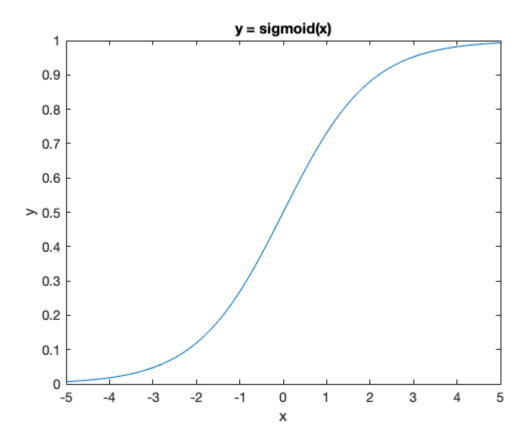
Note that these are normally distributed parameters. That's the reason that group learning rate (which is typically constrained to be between zero and one) is negative here. This is because HBI (and other tools in the cbm) assume that parameters are normally distributed. Therefore, if you have a model with some constraints on parameters (e.g. an RL), you should transform the normally distributed parameter in your model function. To make this point clear, take a look at the model\_RL function. On the first two lines of this function, you see these codes:

```
nd_alpha = parameters(1); % normally-distributed alpha alpha = 1/(1+exp(-nd_alpha)); % alpha (transformed to be between zero and one)
```

Here, nd\_alpha is the normally-distributed parameter passed to the model (for example by cbm tools). Before using it, the model transformed it to alpha, which is bounded between 0 and 1 and is served as the effective learning rate. To do this, a sigmoid function has been used:

```
sigmoid(x) = \frac{1}{1+e^{-x}} which is illustrated here:
```

```
In [16]: x = -5:.1:5;
    y = 1./(1+exp(-x));
    plot(x,y);
    title('y = sigmoid(x)'); xlabel('x'); ylabel('y');
```



As you see, sigmoid is a monotonically increasing function, which transforms its input (x) to an output (y) between 0 and 1. Therefore, if you want to obtain the parameters of your model in their theoretical range (e.g. a learning rate between 0 and 1), you should apply the same transformation (e.g. the sigmoid) to the normally distributed parameter (e.g. to the group\_mean\_RL(1)).

The second parameter of the model\_RL is the decision noise parameter, which is theoretically constrained to be positive. For transforming this one, we did an exponential-transformation to the second parameter passed to the model\_RL:

```
nd_beta = parameters(2);
beta = exp(nd_beta);
```

The HBI also quantifies (hierarchical) errorbar of the group\_mean parameters, which is saved in cbm.output.group\_hierarchical\_errorbar:

You can use the group\_mean and group\_hierarchical\_errorbar values to plot group parameters, or use cbm\_hbi\_plot to plot the main outputs of the HBI.

```
model_names = {'RL', 'Dual RL'};
% note that they corresponds to models (so pay attention to the order)

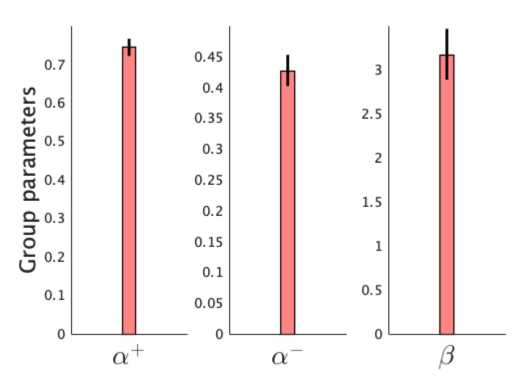
% 3rd input: another cell input containing parameter names of the winning model
param_names = {'\alpha^++', '\alpha^--', '\beta'};
% note that '\alpha^++' is in the latex format, which generates a latin alpha

% 4th input: another cell input containing transformation function associated with each
transform = {'sigmoid', 'sigmoid', 'exp'};
% note that if you use a less usual transformation function, you should pass the handle
cbm_hbi_plot(fname_hbi, model_names, param_names, transform)
% this function creates a model comparison plot (exceednace probability and model frequency)
```

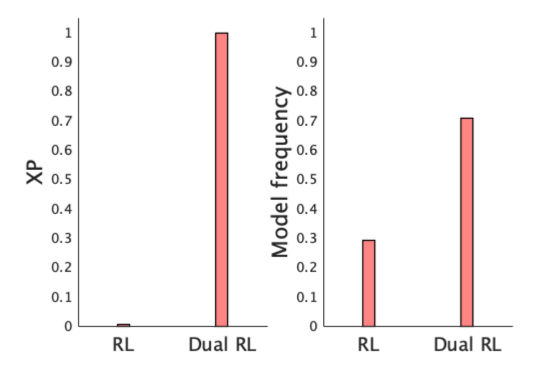
plotting the group parameters of the most frequenct modelThere is no protected exceedance probable Plotting exceedance probability instead...

## Parameters of Dual RL

% a plot of transformed parameters of the most frequent model.



# **Bayesian model comparison**



Similar to a t-test, you can use the hierarchical errorbars to make an inference about a parameter at the population level. We explain that feature in the next example.

The value of individual parameters are saved in the cbm.output.parameters

```
In [19]: parameters_RL = cbm.output.parameters{1};
    parameters_dualRL = cbm.output.parameters{2};
```

Also you can look at the estimated responsibility that each model generated each individual dataset. Across models, responsibilities sum to 1 for each subject.

```
In [20]: responsibility = cbm.output.responsibility
```

responsibility =

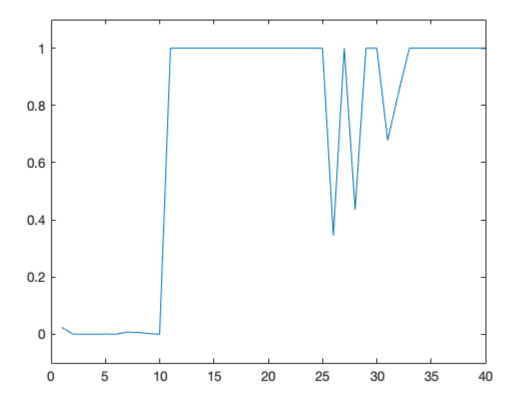
```
0.97550.02450.99920.00080.99930.00070.99960.00040.99900.00100.99950.00050.99230.0077
```

```
0.9934
          0.0066
0.9972
          0.0028
1.0000
          0.0000
0.0000
          1.0000
0.0000
          1.0000
0.0000
          1.0000
0.0000
          1.0000
0.0000
          1.0000
0.0002
          0.9998
0.0000
          1.0000
0.0000
          1.0000
0.0000
          1.0000
0.0000
          1.0000
0.0000
          1.0000
0.0000
          1.0000
0.0000
          1.0000
0.0000
          1.0000
0.0007
          0.9993
0.6553
          0.3447
0.0000
          1.0000
0.5644
          0.4356
0.0000
          1.0000
0.0007
          0.9993
0.3221
          0.6779
0.1548
          0.8452
0.0000
          1.0000
0.0000
          1.0000
0.0000
          1.0000
0.0002
          0.9998
0.0000
          1.0000
0.0000
          1.0000
0.0000
          1.0000
0.0012
          0.9988
```

The first and second columns indicate the responsibility of model\_RL and model\_dualRL in generating the corresponding subject data, respectively.

Look at the estimated responsibility of model\_dualRL:

```
In [21]: plot(responsibility(:,2)); ylim([-.1 1.1])
```



As you see, the estimated responsibility of model\_dualRL for the first 10 subjects is almost zero. These results make sense as this is a synthetic dataset and the first 10 subjects are actually generated using model\_RL. The next 30 individual datasets are generated using model\_dualRL.

Let's also take a look at exceedance probability, a metric typically used for model selection.

The exceedance probability indicates the probability that each model is the most likely model across the group.

A more useful metric is called protected exceedance probability, which also takes into account the null hypothesis that no model in the model space is most likely across the population (i.e. any difference between model frequencies is due to chance).

```
In [23]: pxp = cbm.output.protected_exceedance_prob
```

```
pxp = NaN NaN
```

As you see, this is currently only NaN values. This is because for computing protected exceedance probabilities, the HBI should be re-run under the (prior) null hypothesis.

This is how you can do it:

```
In [24]: fdata = load('all_data.mat');
        data = fdata.data;
        fname_hbi = 'hbi_RL_dualRL';
        % 1st input is data,
        % 2nd input is the file-address of the file saved by cbm_hbi
        cbm_hbi_null(data,fname_hbi);
        % Running this command, prints a report on your matlab output
        % (e.g. on the command window)
cbm_hbi_hbi
                                              18-Jun-2019 22:05:01
Running hierarchical bayesian inference (HBI)- null mode...
HBI has been initialized according to
       lap_RL.mat [for model 1]
       lap_dualRL.mat [for model 2]
Number of samples: 40
Number of models: 2
_____
Iteration 01
Iteration 02
                                                       dL:
                                                             5.07
                                                       dx:
                                                             0.15
Iteration 03
                                                       dL:
                                                              1.14
                                                       dx:
                                                             0.05
Iteration 04
                                                       dL:
                                                             -0.08
                                                       dx:
                                                             0.02
Iteration 05
                                                       dL:
                                                             0.06
                                                       dx:
                                                             0.01
Iteration 06
                                                       dL:
                                                             0.08
                                                       dx:
                                                             0.01
```

cbm\_hbi\_null saves a new file 'hbi\_RL\_dualRL\_null.mat' and it also updates the cbm in hbi\_RL\_dualRL.mat.

Load again hbi\_RL\_dualRL.mat and look at the protected exceedance probability

Note that here values of xp and pxp are not really different (their difference is very small). In many datasets, however, their difference might be quite substantial.

### 3 Example 2: Two-step Markov decision task

We now use cbm tools for computational modeling in the two-step Markov decision task introduced by Daw et al. (2011). This task is a well-known paradigm to distinguish two behavioral modes, model-based and model-free learning. Daw et al. have proposed three reinforcement learning accounts, a model-based, a model-free and their hybrid (which nests the other two and combines their estimates according to a weight parameter), to disentangle contribution of these two behavioral modes on choices.

For this example, we use an empirical dataset reported in Piray et al. (2016) (20 subjects for this example), which is stored in the example\_2step\_task directory.

The hybrid, model-based and model-free algorithms contain 7, 4 and 6 parameters, respectively. Please see Daw et al. for formal description of the models.

loglik = model(parameters,subj);

Now enter the example\_2step\_task directory and load data.

#### struct with fields:

```
choice1: [201x1 double]
  transit: [201x1 double]
  state2: [201x1 double]
  choice2: [201x1 double]
  outcome: [201x1 double]
description: {5x1 cell}
```

See subj1.description for a description of information stored for each subject. Now, you should call cbm\_lap with each model separately.

If you have many subjects or a model with many parameters, the cbm\_lap can take a long time to fit all of them. If you have access to cluster computing, however, you can run cbm\_lap in parallel for subjects. For example, here we fit model\_mf to the data of only subject 1:

```
In [28]: % create a directory for individual output files:
        mkdir('lap_subjects');
        % 1st input: data
        % now the input data should be the data of subject 1
        data_subj = data(1);
        % 2nd input: function handle of model (i.e. @model_mf)
        % 3rd input: a prior struct. The size of mean should
        % be equal to the number of parameters
        prior_mf = struct('mean',zeros(6,1),'variance',6.25);
        % 4th input: output file
        % note that here the output is associated with subject 1
        % we save all output files in the lap_subjects directory
        fname_mf_subj = fullfile('lap_subjects', 'lap_mf_1.mat');
        cbm_lap(data_subj, @model_mf, prior_mf, fname_mf_subj);
Warning: Directory already exists.
                                               18-Jun-2019 22:05:04
cbm_lap
               _____
Number of samples: 1
```

```
Number of parameters: 6
Number of initializations: 42
Subject: 01
done :]
   After all jobs finished, you should call cbm_lap_aggregate to aggregate individual files:
In [29]: % first make a list of lap_mf_* files:
         fname_subjs = cell(20,1);
         for n=1:length(fname_subjs)
             fname_subjs{n} = fullfile('lap_subjects',['lap_mf_' num2str(n) '.mat']);
         fname_subjs
fname_subjs =
  20x1 cell array
    {'lap_subjects/lap_mf_1.mat' }
    {'lap_subjects/lap_mf_2.mat' }
    {'lap_subjects/lap_mf_3.mat' }
    {'lap_subjects/lap_mf_4.mat' }
    {'lap_subjects/lap_mf_5.mat' }
    {'lap_subjects/lap_mf_6.mat' }
    {'lap_subjects/lap_mf_7.mat' }
    {'lap_subjects/lap_mf_8.mat' }
    {'lap_subjects/lap_mf_9.mat' }
    {'lap_subjects/lap_mf_10.mat'}
    {'lap_subjects/lap_mf_11.mat'}
    {'lap_subjects/lap_mf_12.mat'}
    {'lap_subjects/lap_mf_13.mat'}
    {'lap_subjects/lap_mf_14.mat'}
    {'lap_subjects/lap_mf_15.mat'}
    {'lap_subjects/lap_mf_16.mat'}
    {'lap_subjects/lap_mf_17.mat'}
    {'lap_subjects/lap_mf_18.mat'}
    {'lap_subjects/lap_mf_19.mat'}
    {'lap_subjects/lap_mf_20.mat'}
   Now specify the final output file-address and call cbm_lap_aggregate
In [30]: fname_mf = 'lap_mf.mat';
```

cbm\_lap\_aggregate(fname\_subjs,fname\_mf);

```
% Running this command prints a report on your matlab output % (e.g. on the command window)
```

Aggregation is done over 20 subjects :]

You see that lap\_mf.mat is saved by cbm\_lap\_aggregate. Similarly, you can fit model\_hybrid to data using cbm\_lap. I did that and saved lap\_hybrid.mat.

Now that we have fitted models to data using cbm\_lap, we can run cbm\_hbi. Note that you can configure the algorithm using optional inputs:

```
In [31]: % 1st input: data for all subjects
         fdata = load('all_data.mat');
         data = fdata.data;
         % 2nd input: a cell input containing function handle to models
         models = {@model_hybrid, @model_mb, @model_mf};
         % note that by handle, I mean @ before the name of the function
         % 3rd input: another cell input containing file-address to files saved by cbm_lap
         fcbm_maps = {'lap_hybrid.mat', 'lap_mb.mat', 'lap_mf.mat'};
         % note that they corresponds to models (so pay attention to the order)
         % 4th input: a file address for saving the output
         fname_hbi = 'hbi_2step.mat';
         cbm_hbi(data,models,fcbm_maps,fname_hbi);
         % Running this command prints a report on your matlab output
         % (e.g. on the command window)
   Now, we look at the hbi_2step.mat file saved by the cbm_hbi
In [32]: fname_hbi = load('hbi_2step.mat');
         cbm
               = fname_hbi.cbm;
   First, take a look at model frequencies:
In [33]: cbm.output.model_frequency
ans =
    0.6531
           0.3469
                     0.0000
```

As you see the hybrid model takes about 65% of responsibility. Now let's take a look at the exceedance probability

```
In [34]: cbm.output.exceedance_prob
```

```
ans = 0.9102 0.0897 0.0000
```

Note that for computing the protected exceedance probability, the HBI should be re-run under the (prior) null hypothesis cbm\_hbi\_null.

Next, we look at the parameters of the hybrid model:

```
In [35]: cd(fullfile('...','example_2step_task'))

% 1st input is the file-address of the file saved by cbm_hbi
fname_hbi = 'hbi_2step.mat';

% 2nd input: a cell input containing model names
model_names = {'Hybrid', 'MB', 'MF'};

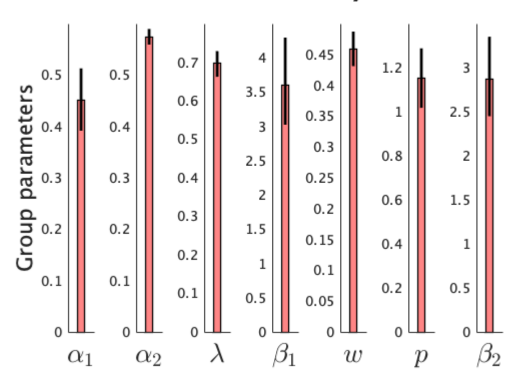
% note that they corresponds to models (so pay attention to the order)

% 3rd input: another cell input containing parameter names of the winning model
param_names = {'\alpha_1', \alpha_2', \lambda', \beta_1', \w', \p', \beta_2'};

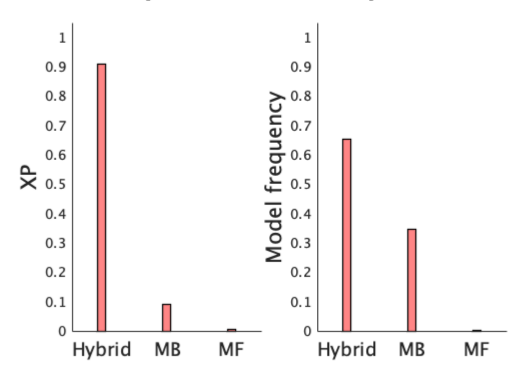
% 4th input: another cell input containing transformation function associated with each
transform = {'sigmoid', 'sigmoid', 'sigmoid', 'exp', 'sigmoid', 'none', 'exp'};
% note that no transformation applied to parameter p (i.e perserveration) in the hybrid
cbm_hbi_plot(fname_hbi, model_names, param_names, transform)
% this function creates a model comparison plot (exceednace probability and model frequent
% a plot of transformed parameters of the most frequent model.
```

plotting the group parameters of the most frequenct modelThere is no protected exceedance probable Plotting exceedance probability instead...

# **Parameters of Hybrid**



# **Bayesian model comparison**



A critical parameter of the hybrid model is the weight parameter, which indicates the degree to which choices influenced by model-based and model-free values. Since the weight parameter is also constrained to be between 0 (i.e. pure model-free) and 1 (i.e. pure model-based), the normally distributed parameter has been transformed in model\_hybrid using the sigmoid function.

Similar to a t-test, you can use the hierarchical errorbars to make an inference about a parameter at the population level. For example, suppose you are interested to test whether the subjects show significantly more model-based than model-free behavior. In terms of the hybrid model, this can be examined by testing whether the (transformed) weight parameter is significantly different from 0.5 (which indicates equal contribution of model-based and model-free values). Since sigmoid function transforms 0 to 0.5, we should test whether the normally distributed weight parameter is significantly different from 0. cbm\_hbi\_ttest performs this inference according to a Student's t-distribution:

```
% 4th input: the index of the parameter of interest
i = 5; % here the weight parameter is the 5th parameter of the hybrid model
[p,stats] = cbm_hbi_ttest(cbm,k,m,i)

p =
    0.1667

stats =
    struct with fields:
    tstat: -1.4585
    pval: 0.1667
    df: 14.0626
```

We see that the p-value is not smaller than 0.05, so there is no significant evidence that the weight is different from 0. In other words, there is no significant evidence that subjects are more influenced by model-based or model-free values.

As another example, let's see whether the perseveration parameter is significantly different from 0 (parameter p in the above plot). This parameter indicates whether subjects repeat their choices (or avoid if p<0) regardless of the estimated values. This parameter has not been transformed, so the test should be against m=0.

```
stats =
struct with fields:
```

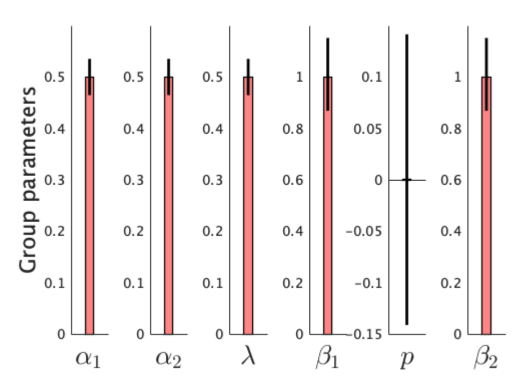
tstat: 8.5369 pval: 6.1737e-07 df: 14.0626

Therefore, the perseveration parameter is significantly larger than zero (p<0.001).

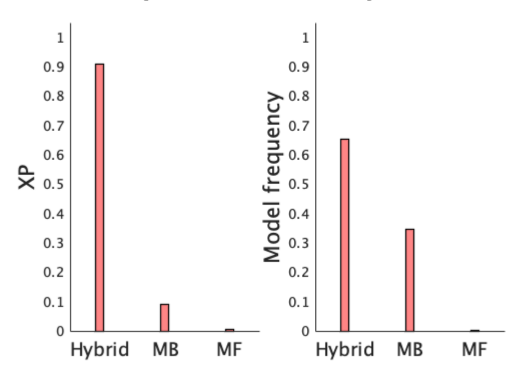
In this example, the model-free model does not take any responsibility. In this situation, the HBI assigns prior values to the individual parameters (with a large variance). This prior value is 0 before transformation. Therefore, for parameters bounded in the unit range (e.g. learning rate), this prior value (after transformation) is 0.5, for decision noise parameter is 1 and for the reservation parameter is 0.

There is no protected exceedance probability as cbm\_hbi\_null has not been executed Plotting exceedance probability instead...

# Parameters of MF



# **Bayesian model comparison**



### 4 Reference

If you use cbm, please cite this paper:

Piray P, Dezfouli A, Heskes T, Frank MJ, Daw ND. (2019) "Hierarchical Bayesian inference for concurrent model fitting and comparison for group studies", PLoS Comp Biol. Link

For a more formal description of the HBI algorithm, please see this paper.