MemToolbox tutorial

(This tutorial uses a beta version of the toolbox and was last updated on 8/11/2012.)

The MemToolbox is a collection of MATLAB functions designed for analyzing data from studies of visual working memory. In particular, it contains functions that fit models to continuous report data (as made popular by Wilken & Ma, 2004 and Zhang & Luck, 2008), and that plot these model fits in order to understand where they capture the data and where they fail to capture the data. The toolbox comes loaded with a number of models, including those made popular by Zhang & Luck (2008), Bays et al. (2009), among others.

Through a series of demonstrations, code for which can be found in the MemDemos folder, the following tutorial covers most of the toolbox's core functionality:

Demo 1: Mixture modeling in just two lines of code.

The simplest way to use the MemToolbox is through MemFit, a function that houses much of the toolbox's functionality under one roof. To use MemFit, first specify your data as a vector of errors, one for each trial, in units of degrees on the color wheel between -180 and 180.

```
>> errors = [-89, 29, -2, 6, -16, 65, 43, -12, 10, 0, 178, -42, 52, 1, -2];
```

Next, call MemFit on the error vector.

```
>> MemFit(errors);
```

The toolbox will now run through its analysis of your data, showing you a histogram of the errors, the name and parameterization of the model it is fitting to the data, and the maximum a posteriori and credible intervals of the model parameter values inferred from the data.

```
Error histogram: -180 ______.'_._____ +180
Model: Standard mixture model
Parameters: g, sd

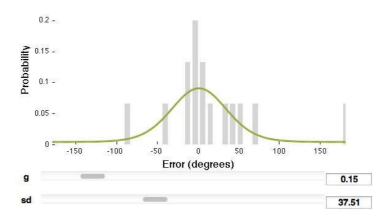
Just a moment while MTB fits a model to your data...
...finished. Now let's view the results:

parameter MAP estimate lower CI upper CI
g 0.151 0.027 0.623
sd 37.507 13.831 65.613
```

Then it will ask whether you'd like to see the fit:

```
Would you like to see the fit? (y/n):
```

If you respond in the affirmative (by entering the letter y and hitting the return key), you'll find an interactive visualization that lets you manipulate the values of the model parameters and see the impact on the predicted distribution of data.



Hooray! In just two lines of code, you've performed mixture modeling using the MemToolbox.

Demo 2: Choosing a different model.

In the previous example, the toolbox picked a model on your behalf, but you can choose a different model if you would like. A full listing of available models can be found by entering help MemModels at the command line. One of the available models extends the standard mixture more by allowing a bias term, such that the central tendency of the data isn't fixed at zero. You can access this model by using WithBias() together with the StandardMixtureModel():

```
>> model = WithBias(StandardMixtureModel);
```

To use this model, first specify your data:

```
>> errors = [-89, 29, -2, 6, -16, 65, 43, -12, 10, 0, 178, -42, 52, 1, -2];
```

Then call MemFit:

```
>> MemFit(errors, model);
```

The toolbox will then walk through the same analysis as before, but using the model that you picked.

Demo 3: Specifying your data as a structure with auxiliary variables.

The two models we have seen cared only about the errors made by the participant on the task. Other models require auxiliary data to make their predictions. For example, the swap model advocated by Bays & Husain (2009), which holds that participants sometimes mistakenly report the value of the other items in the display instead of the target item, requires not only the errors made on each trial but also the values of the non-target items. (The swap model is available in the toolbox under the name <code>SwapModel</code>,) These data, the errors and the non-target values, should be bundled together into a structure array:

```
>> data.errors = [-89, 29, -2, 6, -16 ...
>> data.distractors = [-10, 2, -100, 163, 42 ...
```

The help file for each model lists its required fields:

```
>> help SwapModel
SWAPMODEL returns a structure for a three-component model
```

```
with guesses and swaps. Based on Bays, Catalao, & Husain (2009) model.

Data struct should include:
  data.errors: errors, e.g., distance of response from target
  data.distractors, Row 1: distance of distractor 1 from target
  ...
  data.distractors, Row N: distance of distractor N from target
```

Let's test the model. Rather than making you type out a long vector of errors and a full matrix of distractor values, we'll use one of the datasets that comes loaded with the toolbox. These can be accessed through the function MemDataset, which returns a structure array with fields containing different aspects of the data. For example, data set #3 includes a vector of errors and distractor values for each trial. This is precisely what you need to use the swap model:

Now, instead of feeding MemFit the error vector, you can give it the entire data structure at once:

```
>> MemFit(data, SwapModel);
```

The toolbox will now run through the usual analysis. You may notice some new things in the output of MemFit. We'll explain these in demo #4.

Demo 4: Digging deeper into MemFit()

Let's peek under the hood at the functions that underlie MemFit. By default, the MemToolbox uses Markov Chain Monte Carlo (MCMC) to do model fitting. MCMC samples parameter values in proportion to how well they describe the data and how well they match your prior beliefs. The distribution of these parameter values is used to estimate the true underlying model and to express our confidence in that estimate. In this demo, we'll show you how to read the various plots produced by MemFit that related to MCMC. Once you've mastered the basics from demos 1–3, we recommend the following as a standard workflow when using the toolbox.

Start by loading in a dataset, either yours or one from the toolbox:

```
>> data = MemDataset(1);
```

Now run MemFit using the StandardMixtureModel. This time, assign the output of MemFit to a variable and don't suppress printing with a semicolon.

```
>> fit = MemFit(data, StandardMixtureModel())
```

First, we see the usual histogram, model name, and parameterization:

There should be no surprises here. In the off chance that you have specified your data in an incorrect format (e.g., by using radians instead of degrees or by coding errors in the range [0,360]), the toolbox will try its best to massage your data into the correct format, always throwing a warning letting you know what it has done. Then the toolbox announces:

```
Just a moment while MTB fits a model to your data...

Running 3 chains...
```

This refers to the "chains" of Markov Chain Monte Carlo, which you can read about on Wikipedia (http://en.wikipedia.org/wiki/Markov_chain_Monte_Carlo). The default MCMC algorithm used by MemToolbox uses multiple chains whose starting values are specified in the model file (and modifiable), and it continues running them until they have converged to a similar range of parameters. Every 200 steps, the toolbox will update you on its progress:

```
... not yet converged (200)
```

and then at some point, depending on the model, the chains will converge:

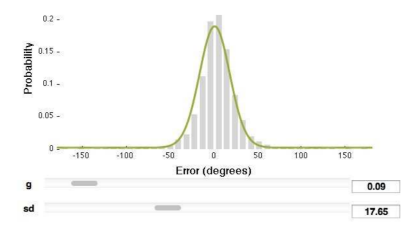
```
... chains converged after 1000 samples!
```

The final estimate is based on samples taken after convergence:

```
... collecting 15000 samples from converged distribution
```

The toolbox then analyzes these samples, returning the MAP (maximum a posteriori) estimate and credible intervals:

Depending on your interests, the maximum a posteriori (MAP) estimates and credible intervals might constitute an answer to your question. Even so, the next step is to visualize the model fit:



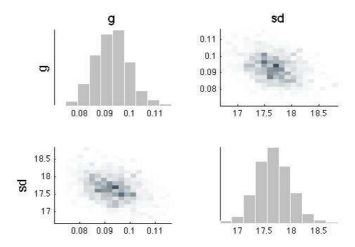
This is mostly useful for familiarizing yourself with the behavior of the model and for recognizing gross inconsistencies between the model and data. At first pass, this one looks okay.

Next, the toolbox will ask:

```
Would you like to see the tradeoffs between parameters, samples from the posterior distribution and a posterior predictive check? (y/n): y
```

Answering in the affirmative will bring up three plots:

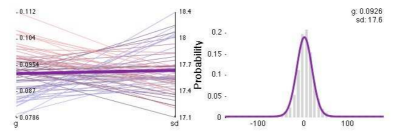
1. The first shows the full posterior distribution for each parameter and a heat map for each pair of them. This can be useful for diagnosing correlations between parameters and for understanding how much the data constrain the parameters:



For example, in the standard Zhang & Luck (2008) mixture model, there is a correlation between the guess rate parameter (*g*) and the standard deviation parameter (*sd*). The data is equally consistent with a slightly higher guess rate and slightly lower standard deviation, or with a slightly lower guess rate and slightly higher standard deviation. This is apparent in the negative slope of the heat map plots in the bottom left and upper right quadrant.

2. The second plot produced by MemFit shows the parameters of the model in a parallel coordinates plot (http://en.wikipedia.org/wiki/Parallel_coordinates), along with a visualization of their fit to the data. Each line on the parallel coordinates plot corresponds to a reasonable set of parameters for fitting the data (e.g., a sample from the posterior on the parameters given the data). The lines are colored such that higher values on the first parameter (g) get red lines, and lower values get blue lines. Thus, the negative correlation between the parameters of this particular model is indicated by the fact that on the right side of the plot, the red lines tend to be lower than the blue lines (the opposite of the left side). A positive correlation would be shown by the lines being mostly parallel between the left and right sides, meaning high values of one parameter give high values of the other.

With only two parameters, the parallel coordinates plot does not provide as much information as the scatter plots and heat maps in the first figure. However, when more parameters are present, parallel coordinates plots allow you to simultaneously understand how all of the parameters relate to each other, rather than just how pairs of parameters relate; we will explore this later in the tutorial.

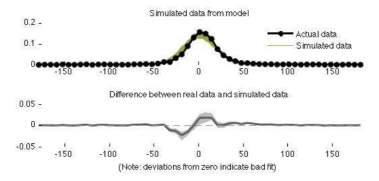


This plot is also interactive. Try clicking one of the lines with a high value of guess rate (g). The plot of the data on the right will redraw with the new parameter values you chose. Now try clicking

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on one of the lines with a low value of guess rate (g). Can you see why both provide equally good fits?

3) The third plot shows the *posterior predictive distribution*—the model's residual. The first two plots provide ways of examining which parameter values of the model provide the best fits. This figure instead shows whether or not the model—with its best parameter values—provides a good fit to the data. In particular, we simulate 'fake' data from the posterior of the model and ask whether it looks like the real data. If the model provides a good fit, then the data that is simulated using the best fit parameters (the posterior) will closely resemble the actual data. If the model is systematically wrong in some way, this should be (visually) evident as a mismatch between the simulated and actual data. The top of the plot shows the data in black, and the simulated data in green. They should be on top of each other if the model provides a good fit. The bottom of the plot shows the difference between the two, both on average (line) and with 95% confidence intervals (gray shading). Any regions where the gray shading does not include zero are regions where the model is systematically wrong.

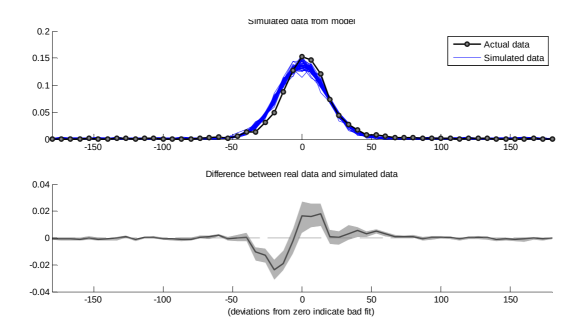


Demo 5: Model comparison

In this demo we'll cover model comparison. Let's start by using the standard mixture model on the first built-in dataset:

```
>> data = MemDataset(1);
>> model1 = StandardMixtureModel();
>> MemFit(data,model1)
```

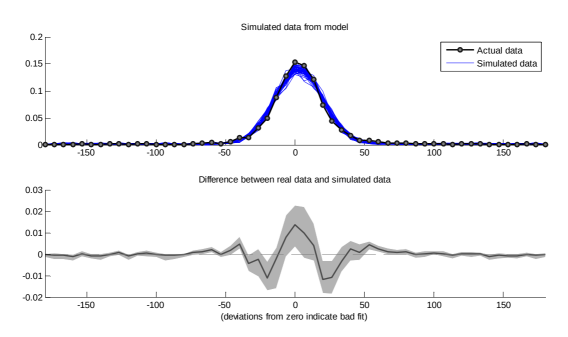
We can see that the residuals for this model show that it does not fit well:



In particular, it looks like the model overestimates counterclockwise error and underestimates clockwise errors. This particular subject had a clockwise-bias in their responses, and the <code>StandardMixtureModel</code> is always centered around zero. To fix this, we can use a different model. In particular, we can use the <code>StandardMixtureModel</code> model, but allow a bias parameter to shift the model's center point:

```
>> model2 = WithBias(StandardMixtureModel);
>> MemFit(data,model2)
```

This model fits much better than the one with no bias.



However, we can see that this model also seems to be systematically incorrect. In particular, the data appear to be 'peakier' than this model suggests they should be. Even so, this model is still an improvement on the model that assumes no bias. How can we quantify this? MemToolbox has built-in model comparison metrics. In particular, simply calling MemFit with more than one model automatically compares them:

```
>> MemFit(data, {model1, model2})
```

This gives the following output:

```
You've chosen to compare the following models:
       Model 1: Standard mixture model
    Parameters: g, sd
MLE: 0.0925789,
                              17.6412
       Model 2: Standard mixture model with bias
    Parameters: mu, g, sd
MLE: 2.83124, 0.0925, 17.4251
Just a moment while MTB fits these models to your data...
model
         Bayes factor
1:2
          -11.0993
         Log likelihood
model
          -14429.6
          -14396.7
1:2
          -32.9079
model
          AIC
          28863.2
          28799.4
        63.8158
1:2
       AICc
model
          28863.2
          28799.4
1:2
          63.8119
model
         BIC
1
          28875.3
          28817.5
1:2
          57.7702
model
         Log posterior odds
----
1
          -6270.1
2.
          -6259
          -11.0993
```

It compares the models using six different metrics, including the Bayes factor, log Likelihood, AIC, AICc, BIC, and Log posterior odds.

Demo 6: Using the parallel toolbox to speed up the analysis

The MemToolbox is compatible with the Parallel Computing toolbox. Using it can achieve a big speedup. To use it, run the command matlabpool open. You can automate this within your program by opening the pool if it isn't already open:

```
if(~(matlabpool('size') > 0))
matlabpool open;
```

end

If your machine has multiple processors or multiple cores, then all of the fitting functions — MCMC, MLE, etc. — will work many times faster once you run matlabpool.

Demo 7: Prettier plots of the posterior

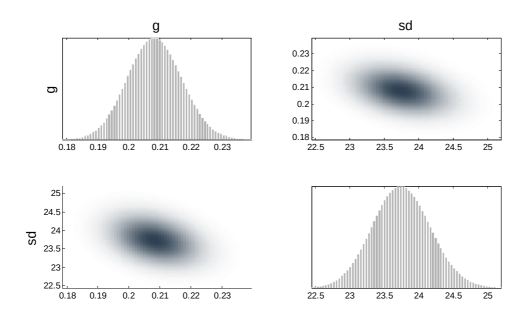
Using MCMC means that we are sampling from the posterior rather than evaluating it directly. Thus, even though the true posterior is almost always a smooth function of the parameters, the plots obtained from the standard MemFit call will not always be smooth. We can fix this by using GridSearch rather than MCMC to refine our graph. In particular, given the results of a standard call to MemFit:

```
>> data = MemDataset(3);
>> model = StandardMixtureModel();
>> fit = MemFit(data, model);
```

We can make a more refined version of the first figure from Demo 4:

```
>> fullPosterior = GridSearch(data, model, ...
    'PosteriorSamples', fit.posteriorSamples)
>> PlotPosterior(fullPosterior, model.paramNames)
```

Then, rather than the noisy sampling-based-plot, we can get a full posterior :



This example also gives you a little bit of a peak under the hood of MemFit — in particular, the separate fitting functions of MemToolbox (try help MemFitting) and plotting functions (help MemPlots). We first fit the data using GridSearch and obtain the posterior (which we call fullPosterior, since it came from GridSearch; the results from MCMC we call posteriorSamples). Then we call the plot function PlotPosterior, which takes either a fulPosterior or posteriorSamples as the first parameter, and the names of the model's parameters as the second parameter. More advanced uses of the MemToolbox will generally require separate calls to the MemFitting and MemPlots functions.

Demo N-1: Arranging your plots (..., 'NewFigure', false); subplot(); subfigure(); PalettableX()

Demo N: Creating your own model.