# rapidtide2

The central program in this package is rapidtide. This is the program that quantifies the time strength and time delay of pervasive signals in a BOLD fMRI dataset.

## Description:

At its core, rapidtide is simply performing a crosscorrelation between a "probe" timecourse and every voxel in an fMRI dataset. As with many things, however, the devil is in the details, and so rapidtide provides a number of features which make it pretty good at this particular task. A few highlights:

- O) There are lots of ways to do something even as simple as a cross-correlation in a nonoptimal way (not windowing, improper normalization, doing it in the time rather than frequency domain, etc.). I'm pretty sure what rapidtide does is, if not the best way, at least a very good and very fast way.
- 1) The sample rate of your probe regressor and the fMRI data do not have to match rapidtide resamples the probe regressor to an integral multiple of the fMRI data rate automatically
- 2) The probe and data can be temporally prefiltered to the LFO, respiratory, or cardiac frequency band with a command line switch, or you can specify any low, high, or bandpass range you want.
- 3) The data can be spatially smoothed at runtime (so you don't have to keep smoothed versions of big datasets around). This is quite fast, so no reason not to do it this way.
- 4) rapidtide can generate a test regressor from the global mean of the data itself no externally recorded timecourse is required. Optionally you can input both a mask of regions that you want to be included in the mean, and the voxels that you want excluded from the mean (there are situations when you might want to do one or the other or both).
- 5) rapidtide can do an iterative refinement of the probe regressor by aligning the voxel timecourses in time and regenerating the test regressor.
- 6) rapidtide fits the peak of the correlation function, so you can make fine grained distinctions between close lag times. The resolution of the time lag discrimination is set by the length of the timecourse, not the timestep this is a feature of correlations, not rapidtide.
- 7) Once the time delay in each voxel has been found, rapidtide outputs a 4D file of delayed probe regressors for using as voxel specific confound regressors or to estimate the strength of the probe regressor in each voxel. This regression is performed by default, but these outputs let you do it yourself if you are so inclined.
- 8) I've put a lot of effort into making the outputs as informative as possible lots of useful maps, histograms, timecourses, etc.
- 9) There are a lot of tuning parameters you can mess with if you feel the need. I've tried to make intelligent defaults so things will work well out of the box, but you have the ability to set most of the interesting parameters yourself.

#### Inputs:

At a minimum, rapidtide needs a Nifti file to work on (space by time). This can be Nifti1 or Nifti2; I can currently read (probably) but not write Cifti files, so if you want to use grayordinate files you need to convert them to nifti in workbench, run rapidtide, then convert back. As soon as nibabel finishes their Cifti support, I'll add that.

#### Outputs:

Outputs are space or space by time Nifti files, and some text files containing textual information, histograms, or numbers. Output spatial dimensions and file type match the input dimensions and file type (Nifti1 in, Nifti1 out). Depending on the file

## A note on coding style:

This code has been in active development since June of 2012. This has two implications. The first is that it has been tuned and refined quite a bit over the years, with a lot of optimizations and bug fixes - most of the core routines have been tested fairly extensively to get rid of the stupidest bugs. I find new bugs all the time, but most of the showstoppers seem to be gone. The second result is that the coding style is all over the place. When I started writing this, I had just moved over from C, and it was basically a mental port of how I would write it in C, and was extremely unpythonic. Over the years, as I've gone back and added functions, I periodically get embarassed and upgrade things to a somewhat more modern coding style. I even put in some classes - that's what the cool kids do, right? But the pace of that effort has to be balanced with the fact that when I make major architectural changes, I tend to break things. So be patient with me.

#### Usage:

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usage: rapidtide2 fmrifilename outputname
[-r LAGMIN, LAGMAX] [-s SIGMALIMIT] [-a] [--nowindow] [-G] [-f GAUSSSIGMA] [-O
oversampfac] [-t TRvalue] [-d] [-b] [-V] [-L] [-R] [-C] [-F]
LOWERFREQ, UPPERFREQ[, LOWERSTOP, UPPERSTOP]] [-0 OFFSETTIME] [-T] [-P] [-A
ORDER] [-B] [-h HISTLEN] [-i INTERPTYPE] [-I] [-Z DELAYTIME] [-N NREPS][--
refineweighting=REFINETYPE] [--refinepasses=NUMPASSES] [--
excludemask=MASKNAME] [--includemask=MASKNAME] [--lagminthresh=LAGMINTHRESH]
[--lagmaxthresh=LAGMAXTHRESH] [--ampthresh=AMPTHRESH][--
sigmathresh=SIGMATHRESH] [--refineoffset] [--pca] [--ica] [--refineupperlag]
[--refinelowerlag] [--tmask=MASKFILE][--limitoutput] [--
timerange=STARTPOINT,ENDPOINT]
[--numskip=SKIP] [--sliceorder=ORDER] [--regressorfreq=FREQ] [--
regressor=FILENAME] [--regressorstart=STARTTIME]
required arguments:
                   - the BOLD fmri file
    fmrifilename
    outputname
               - the root name for the output files
preprocessing options:
                     - override the TR in the fMRI file with the value
TRvalue
                     - disable antialiasing filter
    -a
                     - disable linear trend removal
    --nodetrend
                     - invert the sign of the regressor before processing
    -I
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-i	<pre>- use specified interpolation type (options are 'cubic',    'quadratic', and 'univariate (default)')</pre>
-o	- apply an offset OFFSETTIME to the lag regressors
-b	<ul> <li>use butterworth filter for band splitting instead of trapezoidal FFT filter</li> </ul>
-F	<ul> <li>filter data and regressors from LOWERFREQ to</li> </ul>
UPPERFREQ.	
	LOWERSTOP and UPPERSTOP can be specified, or will be
	calculated automatically
-V	- filter data and regressors to VLF band
-L	- filter data and regressors to LFO band
-R	- filter data and regressors to respiratory band
-C	- filter data and regressors to cardiac band
-N	<ul> <li>estimate significance threshold by running NREPS null correlations (default is 10000, set to 0 to disable)</li> </ul>
nowindow	- disable precorrelation windowing
-f GAUSSSIGMA	- spatially filter fMRI data prior to analysis using
••	GAUSSSIGMA in mm
-M	<ul> <li>generate a global mean regressor and use that as the reference regressor</li> </ul>
-m	- mean scale regressors during global mean estimation
sliceorder	- use ORDER as slice acquisition order used (6 is
Siemens	
	interleave, default is 0 (do nothing))
numskip SKIP	<ul> <li>SKIP tr's were previously deleted during preprocessing (default is 0)</li> </ul>
correlation options:	
-O OVERSAMPFAC	- oversample the fMRI data by the following integral
	factor (default is 2)
regressor	- Read probe regressor from file FILENAME (if none
rogragarfrag	specified, generate and use global regressor)
regressorfreq	<ul> <li>Probe regressor in file has sample frequency FREQ (default is 1/tr)</li> </ul>
rogrossorstart	- First TR of fmri file occurs at time STARTTIME
regressors care	in the regressor file (default is 0.0)
–G	- use generalized cross-correlation with phase alignment
-0	transform (GCC-PHAT) instead of correlation
	orangeon (ood ram), rabeedd or oorretaeron
correlation fitting	options:
	- don't fit the delay time - set it to DELAYTIME seconds
	for all voxels
-r LAGMIN,LAGMAX	- limit fit to a range of lags from LAGMIN to LAGMAX
-s SIGMALIMIT	- reject lag fits with linewidth wider than SIGMALIMIT
regressor refinement	options:
refineweightin	g - apply REFINETYPE weighting to each timecourse prior
	to refinement (valid weightings are 'None', 'R', 'R2' (default)
refinepasses	- set the number of refinement passes to NUMPASSES (default is 1)
includemask	<ul> <li>only use voxels in MASKNAME for global regressor generation and regressor refinement</li> </ul>
excludemask	<ul> <li>do not use voxels in MASKNAME for global regressor</li> </ul>
excrudemask	- do not use voxels in MASKNAME for global regressor generation and regressor refinement
lagminthresh	<ul><li>for refinement, exclude voxels with delays less</li></ul>
raymrnchresh	than LAGMINTHRESH (default is 1.5s)
lagmaxthresh	- for refinement, exclude voxels with delays greater
raymaxtiir esii	Tor retriement, exerude voxers with derays greater

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than LAGMAXTHRESH (default is 1000s)
    --ampthresh
                       - for refinement, exclude voxels with correlation
                         coefficients less than AMPTHRESH (default is 0.3)
                       - for refinement, exclude voxels with widths greater
    --sigmathresh
                         than SIGMATHRESH (default is 50s)
    --refineoffset
                       - adjust offset time during refinement to bring peak
                         delay to zero
                       - only use positive lags for regressor refinement
    --refineupperlag
    --refinelowerlag
                       - only use negative lags for regressor refinement
                       - use pca to derive refined regressor (default is
    --pca
                         averaging)
    --ica
                       - use ica to derive refined regressor (default is
                         averaging)
output options:
    --limitoutput
                     - don't save some of the large and rarely used files
                     - save a table of lagtimes used
    -h
                     - change the histogram length to HISTLEN (default is
100)
    --timerange
                     - limit analysis to data between timepoints STARTPOINT
                       and ENDPOINT in the fmri file
                     - turn off GLM filtering to remove delayed regressor
    --noglm
                       from each voxel (disables output of rCBV)
miscellaneous options:
                     - data file is a converted CIFTI
    -c
    -S
                     - simulate a run - just report command line options
    -d
                     - display plots of interesting timecourses
experimental options (not fully tested, may not work):
    --tmask=MASKFILE - only correlate during epochs specified in
                       MASKFILE (NB: each line of MASKFILE contains the
                       time and duration of an epoch to include
                     - prewhiten and refit data
    -p
                     - save prewhitened data (turns prewhitening on)
    -P
                     - set AR model order to ORDER (default is 1)
    -A, --AR
                     - biphasic mode - match peak correlation ignoring sign
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