# Measuring baseline whole-brain perfusion on GE 3.0T using arterial spin labeling (ASL) MRI with FAIR Sequence

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## **Overview**

This document describes the procedure for measuring baseline whole-brain perfusion in humans using arterial spin labeling (ASL) MRI on GE 3.0T using a modified flow-sensitive inversion recovery (**FAIR**) pulse sequence with single-shot or interleaved **spiral** read outs. Reconstruction of the data into AFNI briks is described. Additional calibration scans are required in order to allow quantification of the perfusion signals via a MATLAB script.

A separate manual is available for measuring baseline whole-brain perfusion using the 8-Phase Multiphase PCASL (8P-MPPCASL) sequence, which yields higher SNR and more precise control of temporal bolus.

## **Acquiring ASL data**

There are two protocols available: **Brain\_perfusion\_4S** and **Brain\_perfusion\_SS**, saved in the **head** section of the protocol list. The CMFRI webpage lists the pros and cons of these two protocols. Each of the protocols consists of 3 scans, **ASL-FAIR**, **CSF** and **MinContrast**. The ASL-FAIR scan collects perfusion-weighted data, and the CSF and MinContrast are calibration scans that are needed for perfusion quantification. A GE 8-channal phased array coil is used for all the scans.

#### **ASL-FAIR**

The ASL-FAIR acquisition uses a modified FAIR sequence. The global inversion pulse used to acquire the tag image in standard FAIR is replaced by a spatially selective inversion pulse, which extends 10cm in either direction outside the imaging slab. QUIPSSII saturation pulses are applied. Some important pulse sequence parameters are: TI1=600 msec, TR=2.5sec, TE=minimum, FOV=22x22cm.

The **Brain\_perfusion\_4S** sequence uses a spiral read out with 4 interleaves to reduce signal dropout due to susceptibility effects, and 20 tag+control image pairs are acquired.

The **Brain\_perfusion\_SS** sequence uses a single shot spiral acquisition, collecting 50 tag+control image pairs. Below are the instructions on how to prescribe and acquire the ASL-FAIR scan.

- 1. View Edit the ASL-FAIR series.
- 2. Click **Graphical Rx** button and prescribe slices (Default: 20x5mm with 1mm gap, axial slices)

Considerations for slice prescription: The slice orientation is defaulted to axial and the slice number increments from bottom to top. Please make sure not to change these default settings when prescribing slices. The default 20x5mm (1mm gap) imaging slab will provide full brain coverage for most subjects. When prescribing, place the lowest slice (i.e. slice No.1) at the very bottom edge of the brain region that you are interested in, and verify that the top slice covers the top edge of the desired region. If necessary, a few more slices can be added, but only to the **top** of the slab. We do not recommend using more than 22 slices. It is fine if a few top slices fall outside of the brain.

#### 3. Save series and Prepare to scan

#### 4. Scan

Verify that the scan time is 6:40 minutes (**Brain\_perfusion\_4S**) or 4:10 minutes (**Brain perfusion SS**).

#### **CSF**

- 1. View Edit
- 2. **Copy Rx** from the ASL-FAIR run (Note: The FOV and slice thickness/gap have to match those of the ASL\_FAIR run in order to use Copy Rx.)
- 3. Save series and Prepare to scan
- Click *Manual Prescan*, and once the clicking sound is heard, click *Done*. (this step ensures that no prescan is performed and all the gain settings are preserved from the ASL-FAIR scan).
- 5. Make sure that the scan time is 36sec and Click Scan

#### MinContrast

- 1. View Edit
- 2. Copy Rx from the ASL-FAIR scan.
- 3. Save series and Prepare to scan
- 4. Click *Manual Prescan*, once hearing clicking sound, click *Done*.
- 5. **Scan**

#### Data transfer and reconstruction

Unlike data acquired with GE product sequences, ASL data are stored as raw data in P-files in directory /usr/g/mrraw. Image reconstruction into AFNI briks is done automatically offline after the scan. Please allow time for reconstruction to finish (~5 minutes after the scan finishes) before transferring your data. Open a command tool and follow the instructions below:

- 1. cd/usr/g/mrraw
- 2. Is -Itr P\* to find the latest P-files.
- 3. To transfer the P-file and reconstructed brik:

asigecopy -s server -r raid# -d datadir Pfilename brikname usrname

(EXAMPLE: aslgecopy –s cfmri –r raid3 –d 060601Pilot P01208.7 scan1 guest)

#### **Important Notes:**

The P-files created during ASL scans are stored in directory /usr/g/mrraw. The space available in this directory is NOT reflected in the storage bar on the GE GUI. It is recommended that you check the space available before running any ASL scans. Do this by typing df –k . in the command line after cd-ing to /usr/g/mrraw.

Some sequence information is stored in the header of the AFNI briks. This information (including sequence parameters, ASL parameters and Rx info) can be accessed using the command (from within the directory containing the brik): **3dNotes scan1brik+orig** 

The briks can be viewed in AFNI. Running the **ASL a3/d3** plug-in on a brik will create three additional briks, labeled as follows:

**A3**: running average of the data (using the 2 neighboring images)

**D3**: running subtraction of the data (using the 2 neighboring images)

avgD3: Average D3 (average perfusion image).

## Data reconstruction using SENSErc program v2.0 (for multishot data only)

Multi-shot data, such as that acquired using the **Brain\_perfusion\_4S** sequence, is sensitive to modulations between shots caused by physiological fluctuations or motion. This can lead to significant image artifacts. The use of a SENSE reconstruction rather than a simple Fourier transform (as performed in the automatic reconstruction described above) significantly reduces sensitivity to these modulations and produces cleaner images. The **SENSErc** program is designed to reconstruct fully-sampled multi-shot spiral data using the SENSE algorithm.

#### System Requirements

- Redhat linux 8.0 or above (Windows and Mac OS not supported).
- Matlab R2007a or above.
- 512Mb or more available system memory recommended.
- AFNI v2.55j or above.

• **FSL** 3.x or **FS**L 4.0 (FSL 4.1 and above are not supported at this time).

**NOTE**: FSL output type needs to be set to **ANALYZE**. To do this, in a command terminal type:

setenv FSLOUTPUTTYPE ANALYZE

or, if using a bash shell:

export FSLOUTPUTTYPE=ANALYZE

You can check your environment variable settings by typing:

env

#### Program Installation

Download SENSErc2.0.tar from the cfmri website:

http://cfmriweb.ucsd.edu/Howto/asl.shtml

Unlike previous versions, the version 2.0 can be used to process data acquired on the 11x, 14x software platforms in addition to those acquired on the 20x platform currently running on the MR750 3TW scanner.

- Untar the file (in Linux: tar -xvf SENSErc2.0.tar).
- Add the SENSErc folder to Matlab path.

(e.g. add the line "addpath yourpath/SENSErc2.0" in your startup.m )

#### Required Data

- ASL P-file
- If CBF quantification is required, a CSF P-file is also needed.

#### Reconstruction steps

- Start Matlab
- In Matlab command window, type:

```
senserc({'Pfilename1',' Pfilename2'},{'AFNIprefix1',' AFNIprefix2'});
e.g.
```

senserc({'P02056.7', 'P03895.7'}, {'senseASL', 'senseCSF'});

Pfilenames are the names of the P files to be reconstructed

AFNIprefixes are the prefixes of the AFNI briks that are output

#### **Important Notes:**

- If CBF quantification is required, both the ASL and the CSF scans should be reconstructed using the SENSErc program; the minimum contrast data does not require this, and the brik created automatically at the scanner console should be used.
- The reconstruction is slow and may take several hours depending on the size of the raw data and the speed of the computer.
- This data reconstruction approach is NOT necessary or advised for data acquired using the single shot sequence (Brain\_perfusion\_SS).

## Perfusion Quantification using CBF program (v3.4)

#### System Requirements

- Redhat linux 8.0 or above (Windows and Mac OS not supported).
- Matlab R2007a or above, plus Matlab toolboxes.
- AFNI v2.55j or above.
- FSL 3.x or FSL 4.0 (FSL 4.1 and above are not supported at this time).

**NOTE:** FSL output type needs to be set to **ANALYZE**. To do this, in a command terminal type:

setenv FSLOUTPUTTYPE ANALYZE

or, if using a bash shell:

export FSLOUTPUTTYPE=ANALYZE

You can check your environment variable settings by typing:

env

512Mb or more available system memory recommended.

#### Program Installation

Download CBFv3.4.tar from the cfmri website:

http://cfmriweb.ucsd.edu/Howto/asl.shtml

- Untar the file (in Linux: tar –xvf CBFv3.4.tar).
- Add the CBFv3.4 folder to your Matlab path.

(e.g. add the line "addpath yourpath/CBFv3.4" in your startup.m )

#### Required Data

- ASL-FAIR brik, CSF brik, MinContrast brik
- Optional: High resolution anatomical brik.

**NOTE:** The anatomical brik must be created from the dicom files using the **d2afni** command rather than **to3d**. d2afni is available on the servers and simply calls to3d, additionally saving Rx information into the brik header.

#### **CBF** Quantification Steps

- Start Matlab
- In Matlab command window, type:

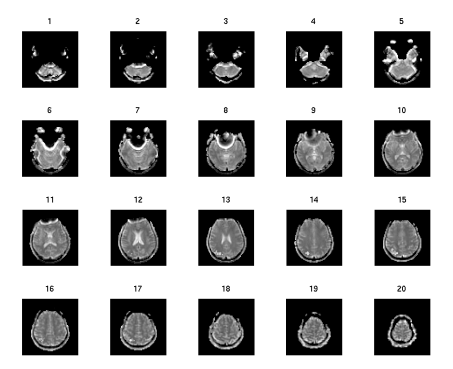
```
calcCBF('ASL-FAIR','CSF','MinCon');
```

OR, if anatomical is available:

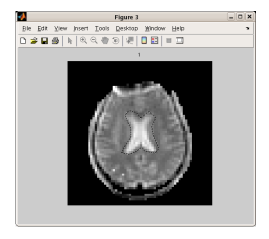
calcCBF('ASL-FAIR','CSF','MinCon','anat');

**Note**: Pass only the prefix of the Afni brik names (no '+orig' needed).

- If the SENSE reconstruction was used, take care to pass the SENSE ASL and CSF briks to calcCBF, rather than the brik automatically transferred at the scanner console using 'aslgecopy'.
- If anatomical is supplied:
  - o The anatomical is segmented into gray, white and CSF (takes ~6mins).
  - All brain slices will then pop up in a matlab figure window; select those containing ventricles by left-clicking.
- If anatomical is NOT supplied:
  - A matlab figure window will pop-up. Identify a slice that best defines a region of cerebral-spinal fluid (CSF) (e.g. slice No. 12 in the figure below). Type the slice # in the matlab command window when prompted. Another window pops up containing the chosen CSF slice.



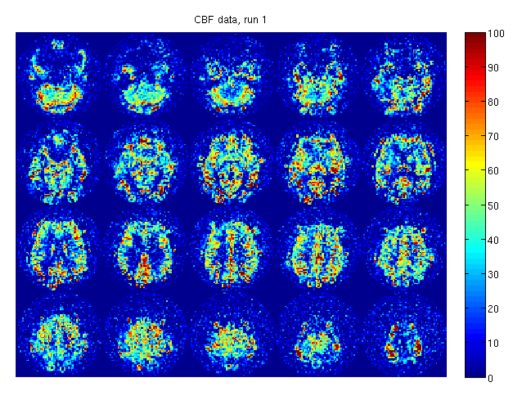
 Use left mouse button to carefully trace around the edge of CSF region and double click to finish. If anatomical is supplied, this is performed for all chosen slices. (You may find double click does not work for certain version of matlab. In this case, please refer to the help file for *roipoly* in your matlab.)



**Note**: this step creates a new directory called '**processed**' under your current working directory. A text file called '**CSFvalue.txt**' is saved under the new directory. This file contains the calculated CSF value, which will be automatically retrieved should you process the same dataset again (you will be prompted to choose whether or not to use this saved value in the repeated analysis). Note also that if the anatomical was supplied,

this directory will also contain the rotated anatomical that aligns with the ASL slices, as well as partial volume maps created from the anatomical.

 The calculated CBF maps are displayed in a new figure. If the anatomical was supplied, a histogram showing the CBF values within gray matter voxels is also shown.



An AFNI brik containing the calibrated CBF values is also saved. Look for the suffix
 CBF' in the brik name.

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Please contact David Shin at <a href="mailto:ddshin@ucsd.edu">ddshin@ucsd.edu</a> with questions/comments