## **Ironsmith QSM Toolkit**

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## This software has been developed for research purposes only and is not a clinical tool.

#### **Description:**

Ironsmith is a toolkit for creating and processing Quantitative Susceptibility Maps (QSM) and for extracting QSM based iron concentrations from subcortical and cortical brain regions.

Ironsmith can perform the following tasks in a fully automated pipeline:

- a) Create QSM maps from GRE DICOM images using the MEDI Toolbox (see section 7 for details) .
- b) Align MPR/MEMPR T1 images to QSM maps and then segment them into 87 ROIs (ROI list in section 8) using freesurfer.
- c) Filter outlier voxels from these ROIs, extract QSM based iron concentration, and format the output into easy to read tables.
- d) Calculate SNR (magnitude image based) for each ROI as a measure of quality control for QSM in an easy to read table.
- e) Warp QSM maps and aligned MPR/MEMPR to MNI152 1mm space for voxelwise QSM analyses.
- f) Process single or multiple participants at a time (multiple instances and nohup supported).

## 1) Software requirements:

#### a) Operating system

#### Unix

Any Linux distribution that supports Singularity (see point c).

Ironsmith tested on:

Red Hat Enterprise Linux Workstation release 7.8 (Maipo)

#### Windows 10 WSL2

Ironsmith tested on:

Ubuntu 18.04 running on Windows 10 Subsystem for Linux V2 (WSL2)

Ubuntu 16.04 running on Windows 10 Subsystem for Linux V2 (WSL2)

#### b) MATLAB

Ironsmith requires Matlab to run MEDI Toolbox and supports versions R2017b to R2019b. Matlab is not needed if MEDI is not required.

#### c) Singularity

Ironsmith tested on Singularity versions 3.5.2 and 3.5.3 Installation guide:

https://sylabs.io/guides/3.5/admin-guide/installation.html

- d) Bash UNIX shell version 4.2.46(2) or later.
- e) MEDI Toolbox version 01/15/2020

Ironsmith requires MEDI Toolbox if QSM maps need to be generated Download from:

http://pre.weill.cornell.edu/mri/pages/qsm.html

MEDI Toolbox is not required if QSM maps already available Currently only MEDI Toolbox version 01/15/2020 is supported

## 2) Installation:

#### a) Download Ironsmith QSM Toolkit

Option 1: download from github

https://github.com/vzachari/IronSmithQSM

Option 2: using git

git clone https://github.com/vzachari/IronSmithQSM.git && cd IronSmithQSM && git checkout v1.0

#### b) Download QSM Container.simg (8.8GB)

from: https://drive.google.com/file/d/1wPdd2Xa0oLV2wwpHneXZ7nllZB3XoKFb/view?usp=sharing

Or

from: https://tinyurl.com/QSMContainer

- c) Place QSM Container.simg in IronSmithQSM/Functions
- d) Download MEDI Toolbox version 01/15/2020 (~7MB)

From: http://pre.weill.cornell.edu/mri/pages/gsm.html

- e) Unzip MEDI Toolbox (typically MEDI\_toolbox.zip)
- f) Place MEDI\_toolbox folder (folder with README.m, UPDATES.m etc) into IronSmithQSM/Functions

NOTE: Make sure the MEDI\_toolbox folder in IronSmithQSM/Functions does not have another MEDI\_toolbox folder in it (e.g MEDI\_toolbox/MEDI\_toolbox)

g) Edit IronSmithQSM/Matlab\_Config.txt with the path to the matlab executable on your system.

(e.g. /usr/local/MATLAB/R2019b/bin/matlab) Supported versions R2017b to R2019b.

#### h) Add the IronSmithQSM directory to \$PATH

Guide: https://opensource.com/article/17/6/set-path-linux

## 3) Syntax:

#### Ironsmith [MyInputFile] [absolute path to output folder]

Example: Ironsmith File.csv /home/data/MyAmazingExp/QSM\_Analysis"

- a) The output folder does not need to exist.
- b) The output folder does not need to be empty but Ironsmith will skip any participant specified in MyInputFile that has a corresponding folder inside the output folder.
- Ex. if S0001 is specified in MyInputFile and folder S0001 exists in output folder, S0001 will be skipped.

c) Freesurfer\_Skip is a reserved folder name under output folder and may be used by Ironsmith. See section #5 on quality of life features below.

## 4) MyInputFile format:

- a) MyInputFile has to be CSV formatted (entries separated by commas ',').
- b) Each row corresponds to a different participant.

see Example\_File.csv in IronSmithQSM folder:

### If MEDI Toolbox is required to create QSM images/maps:

Column1 = Subj (nominal subject variable e.g. S0001 or 01 or Xanthar\_The\_Destroyer)

**Column2** = MEDI\_Yes <-- this is case sensitive

Column3 =

**Either** absolute path to directory with MPR/MEMPR files **OR** absolute path to a single NIFTI (.nii or .nii.gz) MPR/MEMPR file

(e.g. /home/subjects/S01/MPR OR /home/subjects/S01/MPR/S01\_MEMPR.nii.gz)

#### If a single NIFTI (.nii or .nii.gz) MPR/MEMPR file is provided:

File can have any name.

File can have multiple time-points, each corresponding to a different echo (RMS will be calculated).

File can have a single echo/time-point.

#### If path to directory with MPR/MEMPR files is provided:

MPR/MEMPR files in provided folder can be:

a) DICOMS

Only MPR/MEMPR DICOMS should be present in the MPR folder if you want Ironsmith to process DICOMS. If NIFTI files exist together with DICOMS, they will be selected instead (see "b" below).

b) Multiple .nii/.nii.gz files each corresponding to a different echo (will be catenated and RMS will be calculated).

To make sure the correct files are selected, each NIFTI file needs to have \_e# in the file name, where # is the echo number.

 $(e.g.\ S0001\_MEMPR\_e1.nii.gz,\ S0001\_MEMPR\_e2.nii.gz...)$ 

This is the default **dcm2niix** output format for multiple echos.

- c) Single .nii/nii.gz file with multiple time-points, each corresponding to a different echo (RMS will be calculated). This single NIFTI file can have any name.
- d) Single .nii/nii.gz file with a single echo/time-point.

This can be rms/averaged across echos or just a single echo T1 MPRAGE.

This single NIFTI file can have any name and will be used as is.

**Column4** = Absolute path to folder with QSM DICOM files

(e.g. /home/subjecs/S01/QSM\_Dicom)

QSM DICOM folder must include DICOMS for both GRE magnitude and phase.

Preferably only QSM DICOMS should be present in the QSM\_Dicom folder. However, Ironsmith can filter out the following filetypes .nii .json .txt .nii.gz .HEAD .BRIK .hdr .img

All 4 columns need to be provided, otherwise Ironsmith will exit with errors.

#### If QSM maps and GRE magnitude images are already available:

Column1 = Subj (nominal subject variable e.g. S0001 or 01 or Xanthar\_The\_Destroyer)

Column2 = MEDI No <-- This is case sensitive

Column3 =

**Either** absolute path to directory with MPR/MEMPR files **OR** absolute path to a single NIFTI (.nii or .nii.gz) MPR/MEMPR file

(e.g. /home/subjects/S01/MPR OR /home/subjects/S01/MPR/S01\_MEMPR.nii.gz)

#### If a single NIFTI (.nii or .nii.gz) MPR/MEMPR file is provided:

File can have any name.

File can have multiple time-points, each corresponding to a different echo (RMS will be calculated). File can have a single echo/time-point.

#### If path to directory with MPR/MEMPR files is provided:

MPR/MEMPR files in provided folder can be:

a) DICOMS

Only MPR/MEMPR DICOMS should be present in the MPR folder if you want Ironsmith to process DICOMS. If NIFTI files exist together with DICOMS, they will be selected instead (see "b" below).

b) Multiple .nii/.nii.gz files each corresponding to a different echo (will be catenated and RMS will be calculated).

To make sure the correct files are selected, each NIFTI file needs to have \_e# in the file name, where # is the echo number.

(e.g. S0001\_MEMPR\_e1.nii.gz, S0001\_MEMPR\_e2.nii.gz...)

This is the default **dcm2niix** output format for multiple echos.

- c) Single .nii/nii.gz file with multiple time-points, each corresponding to a different echo (RMS will be calculated). This single NIFTI file can have any name.
- d) Single .nii/nii.gz file with a single echo/time-point.

This can be rms/averaged across echos or just a single echo T1 MPRAGE.

This single NIFTI file can have any name and will be used as is.

**Column4** = Absolute path including filename to QSM magnitude image (e.g. /home/subjecs/S01/QSM/QSM\_Magnitude.nii.gz)

**Column5** = Absolute path including filename to QSM map (e.g. /home/subjects/S01/QSM/QSM\_Map.nii.gz)

All 5 columns need to be provided, otherwise Ironsmith will exit with errors

## 5) Quality of life features:

## Skipping freesurfer segmentation

If a participant already has a completed freesurfer recon-all -all segmentation folder and you would like Ironsmith to skip the freesurfer segmentation step, do the following:

a) Copy the freesurfer recon-all folder (the one containing the *label, mri, scripts, stats, surf...* folders) into /OutputFolder/Freesurfer\_Skip, where OutputFolder is the one specified/to be specified in the Ironsmith command.

You can create the /OutputFolder/Freesurfer\_Skip folder or Ironsmith will create it for you if you have run it at least once previously for OutputFolder.

b) Rename the recon-all folder to **Subj\_FreeSurfSeg\_Skull**. Subj should match the one provided in MyInputFile and should correspond to the participant you want the segmentation step skipped.

**Note:** if Ironsmith runs freesurfer it will create **Subj\_FreeSurfSeg\_Skull** and place it under /**OutputFolder/Subj/MPR**. This helps reduce processing time if for any reason one would like to repeat the analysis on a given participant (e.g. due to crash or errors). Just copy/move this folder over to /**OutputFolder/Freesurfer\_Skip**, delete the problematic participant folder (e.g. /**OutputFolder/Subj**) and rerun Ironsmith.

#### Viewing output NIFTI files

If you do not have a NIFTI viewer, AFNI can be launched from within the QSM\_Container.simg by using the Ironsmith\_AFNI command. Just type Ironsmith\_AFNI from within the folder you would like to view NIFTI files from.

AFNI viewer documentation:

https://afni.nimh.nih.gov/pub/dist/edu/latest/afni\_handouts/afni03\_interactive.pdf

## 6) Outputs:

Each participant processed by Ironsmith will have a corresponding folder in **OutputFolder**. For example, if "/home/QSM\_Analysis" is the OutputFolder and "S0001" is one of the participants processed, then /home/QSM\_Analysis/S0001 will be created and populated with data.

a) All masks/ROIs are placed under:

S0001/QSM/Freesurf\_QSM\_Masks/Cort\_Masks\_AL\_QSM\_RS\_Erx1 S0001/QSM/Freesurf\_QSM\_Masks/SubC\_Masks\_AL\_QSM\_RS\_Erx1

b) All QSM maps/images created are placed under S0001/QSM/Freesurf\_QSM\_Masks and are labelled as:

Subj\_QSM\_Map\_FSL.nii.gz <-- Default MEDI
Subj\_QSM\_Map\_New\_CSF\_FSL.nii.gz <-- Lateral ventricles as the QSM reference structure
Subj\_QSM\_Map\_New\_WM\_FSL.nii.gz <-- White matter as the QSM reference structure</pre>

c) All QSM maps warped to MNI space are placed under

S0001/QSM/Freesurf QSM Masks/MNI152 QSM

d) QSM per ROI means (87 ROIs) are under /QSM\_Analysis/Group as follows:

Group\_QSM\_Mean.csv Group\_QSM\_Mean\_CSF.csv Group\_QSM\_Mean\_WM.csv Group\_QSM\_ADJ\_Mean.csv Group\_QSM\_ADJ\_Mean\_CSF.csv Group\_QSM\_ADJ\_Mean\_WM.csv Group\_QSM\_SNR.csv

Mean = Using only positive QSM voxels

\_ADJ\_Mean = Using only positive QSM voxels and adjusting for ROI size

\_CSF = Lateral ventricles as the QSM reference structure

\_WM = White matter as the QSM reference structure

SNR is calculated as follows:

mean signal intensity of magnitude image within an ROI / standard deviation of magnitude signal outside the head.

The outside the head mask can be found here:

/QSM\_Analysis/S0001/QSM/Freesurf\_QSM\_Masks/Subj\_QSM\_Mag\_FSL\_rms\_OH\_Mask.nii.gz

# 7) Ironsmith uses the following software, provided in the form of a Singularity image:

#### **AFNI**

RW Cox. AFNI: Software for analysis and visualization of functional magnetic resonance neuroimages. Computers and Biomedical Research, 29:162-173, 1996.

RW Cox and JS Hyde. Software tools for analysis and visualization of FMRI Data. NMR in Biomedicine, 10:171-178, 1997.

S Gold, B Christian, S Arndt, G Zeien, T Cizadlo, DL Johnson, M Flaum, and NC Andreasen. Functional MRI statistical software packages: a comparative analysis. Human Brain Mapping, 6:73-84, 1998.

#### dcm2niix

Li, Xiangrui, et al. "The first step for neuroimaging data analysis: DICOM to NIFTI conversion." Journal of neuroscience methods 264 (2016): 47-56.

#### **Freesurfer**

Dale, A.M., Fischl, B., Sereno, M.I., 1999. Cortical surface-based analysis. I. Segmentation and surface reconstruction. Neuroimage 9, 179-194.

Dale, A.M., Sereno, M.I., 1993. Improved localization of cortical activity by combining EEG and MEG with MRI cortical surface reconstruction: a linear approach. J Cogn Neurosci 5, 162-176.

Desikan, R.S., Segonne, F., Fischl, B., Quinn, B.T., Dickerson, B.C., Blacker, D., Buckner, R.L., Dale, A.M., Maguire, R.P., Hyman, B.T., Albert, M.S., Killiany, R.J., 2006. An automated labeling system for subdividing the human cerebral cortex on MRI scans into gyral based regions of interest. Neuroimage 31, 968-980.

#### **FSL**

M.W. Woolrich, S. Jbabdi, B. Patenaude, M. Chappell, S. Makni, T. Behrens, C. Beckmann, M. Jenkinson, S.M. Smith. Bayesian analysis of neuroimaging data in FSL. NeuroImage, 45:S173-86, 2009

S.M. Smith, M. Jenkinson, M.W. Woolrich, C.F. Beckmann, T.E.J. Behrens, H. Johansen-Berg, P.R. Bannister, M. De Luca, I. Drobnjak, D.E. Flitney, R. Niazy, J. Saunders, J. Vickers, Y. Zhang, N. De Stefano, J.M. Brady, and P.M. Matthews. Advances in functional and structural MR image analysis and implementation as FSL. NeuroImage, 23(S1):208-19, 2004

M. Jenkinson, C.F. Beckmann, T.E. Behrens, M.W. Woolrich, S.M. Smith. FSL. NeuroImage, 62:782-90, 2012

## 8) ROI List:

- LR\_Occipital\_Lobe\_GM
- LR\_Temporal\_Lobe\_GM
- $L\_Caudal Anterior Cingulate\_GM$
- L\_CaudalMiddleFrontal\_GM
- L\_Cuneus\_GM
- L\_DLPFC\_GM
- L\_Entorhinal\_GM
- L\_Frontal\_GM
- L\_Fusiform\_GM
- $L\_InferiorParietal\_GM$
- $L\_InferiorTemporal\_GM$
- L\_Insula\_GM
- L\_IsthmusCingulate\_GM
- $L_LateralOccipital\_GM$
- $L\_Lateral Orbit of rontal\_GM$
- L\_Lingual\_GM
- $L\_MedialOrbitofrontal\_GM$
- $L_MiddleTemporal_GM$
- L\_Occipital\_GM\_Mask
- $L_Parietal\_GM\_Mask$
- $L\_Temporal\_GM\_Mask$
- L\_Parahippocampal\_GM
- L\_Pericalcarine\_GM
- L\_Postcentral\_GM
- L\_PosteriorCingulate\_GM
- L\_Precentral\_GM
- L\_Precuneus\_GM
- L\_RostalMiddleFrontal\_GM
- $L\_RostralAnterior Cingulate\_GM$
- $L\_SuperiorFrontal\_GM$
- L\_SuperiorParietal\_GM
- $L\_SuperiorTemporal\_GM$
- L\_TransverseTemporal\_GM
- $R\_Caudal Anterior Cingulate\_GM$
- $R_Caudal Middle Frontal_GM$
- R\_Cuneus\_GM
- R\_DLPFC\_GM
- $R\_Entorhinal\_GM$
- R\_Frontal\_GM\_Mask
- R\_Fusiform\_GM
- R\_InferiorParietal\_GM
- $R\_InferiorTemporal\_GM$
- R\_Insula\_GM
- $R\_IsthmusCingulate\_GM$
- $R_LateralOccipital\_GM$
- $R\_LateralOrbitofrontal\_GM$
- R\_Lingual\_GM
- $R\_MedialOrbitofrontal\_GM$
- R\_MiddleTemporal\_GM
- R\_Occipital\_GM\_Mask
- $R_Parietal\_GM\_Mask$
- R\_Temporal\_GM\_Mask
- $R\_Parahippocampal\_GM$
- R\_Pericalcarine\_GM
- $R_Postcentral\_GM$
- R\_PosteriorCingulate\_GM
- $R_Precentral_GM$

- R\_Precuneus\_GM
- R\_RostalMiddleFrontal\_GM
- $R\_RostralAnteriorCingulate\_GM$
- R\_SuperiorFrontal\_GM
- $R\_SuperiorParietal\_GM$
- R\_SuperiorTemporal\_GM
- $R\_TransverseTemporal\_GM$
- LR\_Accumbens\_area
- LR\_Amygdala
- LR\_Caudate
- LR\_Hipp
- LR\_Pallidum
- LR\_Putamen
- LR\_Thalamus\_Proper
- L\_Accumbens\_area
- L\_Amygdala
- L\_Caudate
- L\_Hipp
- L\_Pallidum
- L\_Putamen
- L\_Thalamus\_Proper
- R\_Accumbens\_area
- $R\_Amygdala$
- R\_Caudate
- R\_Hipp
- R\_Pallidum
- R\_Putamen
- R\_Thalamus\_Proper