TractoFlow pipeline is developed by the Sherbrooke Connectivity Imaging Lab (SCIL) in order to process diffusion MRI dataset from the raw data to the tractography. The pipeline is based on Nextflow and Singularity. The goal with this pipeline is to be fast and reproducible.
To run the pipeline you must install Nextflow. To use our Singularity container, you must install the Singularity package.

### 1.1 Nextflow

#### 1.1.1 Local Computer

1. Before installing check your current version `java -version`. If return something as `java version "1.\nX"` and X is 8 or later, you can skip this step else install java.

2. Install Nextflow:

   ```bash
   $> curl -s https://get.nextflow.io | bash && chmod +x nextflow && \
   echo 'export PATH=$PATH:'$(pwd) >> ~/.bashrc && source ~/.bashrc
   ```

#### 1.1.2 High Performance computer (HPC)

1. Try `module load java/1.8.0_121` or check with your administrator or on the HPC website.

2. Use `wget` to install the last Nextflow edge-all release, change the name, add execution rights and add the nextflow path in the bashrc.

   ```bash
   # Example with 19.01.0 version
   $> wget https://github.com/nextflow-io/nextflow/releases/download/v19.01.0/nextflow-\n   →19.01.0-all && \
   mv nextflow-19.01.0-all nextflow && \
   chmod +x nextflow && echo 'export PATH=$PATH:'$(pwd) >> ~/.bashrc && source ~/.bashrc
   ```
1.2 Singularity

Our Singularity container currently works on Linux. We highly recommend to use Singularity on a Linux local computer or on a HPC.

If you want to use Docker on Windows or MacOS, please see the Docker section.

1.2.1 Local Computer

Install `singularity-container`. Our current singularity container works only on Linux. A macOS version will be released soon.

If you are Debian/Ubuntu, you can get neurodebian:

```
$> sudo wget -O- http://neuro.debian.net/lists/xenial.us-ca.full | sudo tee /etc/apt/sources.list.d/neurodebian.sources.list && \
   sudo apt-key adv --recv-keys --keyserver hkp://pool.sks-keyservers.net:80 --keyserver-p每月=0xA5D32F012649A5A9 && \
   sudo apt-get update && sudo apt-get install -y singularity-container
```

1.2.2 High Performance computer (HPC)

Please try `module load singularity/2.6` or check with an administrator or on the HPC website.

1.3 Docker

1.3.1 MacOS

To install Docker on your MacOS computer, please check the following link:

https://hub.docker.com/editions/community/docker-ce-desktop-mac

1.3.2 Windows

To install Docker on your Windows computer, please check the following link:

https://hub.docker.com/editions/community/docker-ce-desktop-windows
2.1 TractoFlow pipeline

2.1.1 Release

Download the last release of TractoFlow pipeline:

```
$> wget https://github.com/scilus/tractoflow/archive/2.0.1.zip && unzip 2.0.1.zip
```

2.1.2 For developers

Clone TractoFlow pipeline repository:

```
# Clone with HTTPS
$> git clone https://github.com/scilus/tractoflow.git

# Clone with SSH
$> git clone git@github.com:scilus/tractoflow.git
```

2.2 Singularity for TractoFlow

2.2.1 Release

Download the last release of the Singularity container for TractoFlow:

```
```
2.2.2 For developers

Clone the singularity repository for TractoFlow pipeline:

```
# Clone with HTTPS
$> git clone https://github.com/scilus/singularity-tractoflow.git

# Clone with SSH
$> git clone git@github.com:scilus/singularity-tractoflow.git
```

Then, you can build the singularity image:

```
$> singularity build singularity_name.img singularity_tractoflow.def
```

2.3 Docker for TractoFlow

2.3.1 Release

Download the last release of the Docker container for TractoFlow:

```
$> wget http://scil.dinf.usherbrooke.ca/containers_list/docker_tractoflow_2.0.0_→a0cacfb_2019-04-25.tar.gz
```

Install the Docker container:

```
$> docker image load -i "docker_tractoflow_2.0.0_a0cacfb_2019-04-25.tar.gz"
```
Processing steps

TractoFlow pipeline consist of 23 different steps: 14 steps for the diffusion weighted image (DWI) processing and 8 steps for the T1 weighted image processing.
Chapter 3. Processing steps

A. Input files
- DWI
- bval/bvec
- T1
- Reverse B0

B. DWI processes
- Preliminary brain extraction DWI
  - Denoise DWI
  - Eddy Topup
  - Brain extraction DWI
  - N4 DWI
  - Crop DWI
  - Normalize DWI
  - Resample DWI
  - Extract DTI Shell
  - DTI metrics
  - Compute FRF
  - Mean FRF
  - fODF metrics

C. T1 processes
- Denoise T1
- N4 T1
- Resample T1
- Brain extraction T1
- Crop T1
- Register T1 on DWI
- Segment T1
- Tracking maps

Tracking
3.1 Input

- Diffusion weighted image (DWI)
- b-values
- b-vectors
- T1 weighted image
- Reverse phase encoding B0 (Optional)

3.2 DWI processes

- Brain extraction (FSL)
- Denoising (Mrtrix3)
- Topup (FSL)
- Eddy (FSL)
- N4 bias correction (ANTs)
- Resample (Dipy)
- DTI metrics (Dipy)
- fODF metrics (Dipy)

3.3 T1 processes

- Brain extraction (ANTs)
- Denoising (Dipy)
- N4 bias correction (ANTs)
- Resample (Dipy)
- Registration (ANTs)
- Tissue segmentation (FSL)

3.4 Tractography

The particle filter tractography is performed. Two types of seeding is available: WM-GM interface or WM mask.
4.1 Root parameter

The input root parameter is called using --root and require the following file structure:

```
[root]
  S1
    dwi.nii.gz
    bval
    bvec
    rev_b0.nii.gz (optional)
    t1.nii.gz
  S2
    dwi.nii.gz
    bval
    bvec
    rev_b0.nii.gz (optional)
    t1.nii.gz
```

The root folder must contains subjects folders (e.g. S1, S2, . . .). Each subject folder contains the required images:

- *dwi.nii.gz* are the diffusion weighted images.
- *bval* is the b-value file in the FSL format.
- *bvec* is the b-vector file in the FSL format.
- *rev_b0.nii.gz* (optional) is the reversed phase encoded b0 image also called blip-up/blip-down. Used to correct distortion due to diffusion acquisition (Documentation).
- *t1.nii.gz* is the T1 weighted image.
4.2 BIDS parameter

Work in progress.
To display the options of Tractoflow, please use `nextflow run tractoflow/main.nf --help`.

### 5.1 Options list

**--b0_thr_extract_b0** _MAX_VALUE_ (default: 10) All b-values below a maximum value are considered b=0 images.

**--dwi_shell_tolerance** _TOLERANCE_ (default: 20) All b-values to +-tolerance are considered as the same b-value.

**--bet_prelim_f** _THRESHOLD_ (default: 0.16) Fractional Intensity threshold (-f for the bet FSL command) for preliminary DWI brain extraction. See FSL bet documentation for more info.

**--dilate_b0_mask_prelim_brain_extraction** _FACTOR_ (default: 5) Dilation factor to keep the whole brain and be more robust to the geometric distortions. This is only applied to the preliminary BET. Not the final extraction.

**--run_dwi_denoising** _BOOL_ (default: true) Run dwi denoising (dwidenoise from Mrtrix3). See Mrtrix3 dwidenoise documentation for more info.

**--extent** _SIZE_ (default: 7) Denoising block size. Recommended block size should follow the following rule of thumb: extent^3 >= # directions. See Mrtrix3 dwidenoise documentation for more info.

**--run_topup** _BOOL_ (default: true) Run Topup. If TractoFlow find any reversed phase encoded b=0 images. Topup will be automatically ignored. See FSL Topup documentation for more info.

**--encoding_direction** _DIRECTION_ (default: y) Encoding direction of the DWI [x, y, z]. See FSL Topup documentation for more info.

**--dwell_time** _VALUE_ (default: 0.062) Dwell-time value.

**--run_eddy** _BOOL_ (default: true) Run Eddy.

**--eddy_cmd** _COMMAND_ (default: eddy_openmp) Eddy command to use [eddy_openmp, eddy_cuda].
TractoFlow-documentation Documentation

--bet_topup_before_eddy_f THRESHOLD (default: 0.16) Fractional Intensity threshold (-f for the bet FSL command) for intermediate BET operation on topup corrected images.

--use_slice_drop_correction BOOL (default: true) If set, will use the slice drop correction option (--repol) from Eddy.

--bet_dwi_final_f THRESHOLD (default: 0.16) Fractional Intensity threshold (-f for the bet FSL command) for the final DWI BET.

--run_resample_dwi BOOL (default: true) Run resample DWI. Resampling is done at the resolution given by --dwi_resolution.

--dwi_resolution RESOLUTION (default: 1) DWI resolution (in mm).

--dwi_interpolation METHOD (default: lin) Interpolation method [nn, lin, quad, cubic].

--number_of_tissues NUMBER (default: 3) Number of tissue classes (-n for the fast FSL command).

--fa THRESHOLD (default: 0.7) Initial FA threshold to compute the fiber response function (FRF).

--min_fa MIN_THRESHOLD (default: 0.5) Minimum FA threshold to compute the FRF.

--roi_radius RADIUS (default: 20) Region of interest radius to compute the FRF. This ROI starts from the center of the 3D volume (sizeX/2, sizeY/2, sizeZ/2).

--set_frf BOOL (default: false) Set manually the FRF.

--manual_frf FRF (default: “15,4,4”) FRF set manually. The FRF must be at 10^-4 scaling in mm^2/s. This corresponds to an elongated symmetric diffusion tensor with eigenvalues (15, 4, 4) x 10^-4 mm^2/s along the principal axis and radial axes respectively.

--mean_frf BOOL (default: true) Mean the frf of all subjects. USE ONLY IF ALL OF SUBJECTS COME FROM THE SAME SCANNER AND HAVE THE SAME ACQUISITION.

--sh_order ORDER (default: 8) Spherical harmonics order.

    Suggested rule of thumb:
    --sh_order=8 for 45+ directions
    --sh_order=6 for 20+ directions
    --sh_order=4 otherwise

--basis BASIS (default: descoteaux07) fODF spherical harmonics (SH) basis type [descoteaux07, tournier07].

--fodf_metrics_a_factor FACTOR (default: 2.0) Multiplicative factor for AFD max in ventricles. As recommended in [Dell’Acqua et al HBM 2013].

--relative_threshold THRESHOLD (default: 0.1) Relative threshold on fODF amplitude in ]0,1].

--max_fa_in_ventricle THRESHOLD (default: 0.1) Maximal threshold of FA to be considered as ventricle voxel. Used to compute the ventricles mask and find the maximum fODF amplitude in the ventricles.

--min_md_in_ventricle THRESHOLD (default: 0.003) Minimal threshold of MD in mm^2/s to be considered as ventricle voxel. Used to compute the ventricles mask and find the maximum fODF amplitude in the ventricles.

--wm_seeding BOOL (default: true) If ‘--wm_seeding true’, use the WM-GM interface and the WM mask as seeding mask, else use the WM-GM interface as seeding mask.

--algo ALGO (default: prob) Tracking algorithm [prob, det].
prob: streamline probabilistic.
det: streamline deterministic.

--seeding TYPE (default: npv) Seeding type [npv, nt].

npv: number of seeds per voxel of the seeding mask
nt: total number of seeds randomly placed in the seeding mask

--nbr_seeds NUMBER (default: 10) Number of seeds related to the seeding type param.
--random SEED (default: 0) Random seed. Fixed for reproducible seeds
--step SIZE (default: 0.1) Step size in mm.
--theta ANGLE (default: 20) Maximum angle between 2 steps in degrees.
--min_len LENGTH (default: 20) Minimum length in mm.
--max_len LENGTH (default: 200) Maximum length in mm.
--compress_streamlines BOOL (default: true) Compress streamlines.
--compress_value THRESHOLD (default: 0.2) Compression error threshold in mm. See [Presseau et al Neuroimage 2015] and [Rheault et al Front Neuroinform 2017]
--template_t1 PATH (default: /human-data/mni_152_sym_09c/t1) Path to the template T1 directory for antsBrainExtraction. The folder must contain t1_template.nii.gz and t1_brain_probability_map.nii.gz. The default path is the human_data folder in the Singularity/Docker container.
--processes_brain_extraction_t1 NUMBER (default: 4) Number of processes for T1 brain extraction task.
--processes_denoise_dwi NUMBER (default: 4) Number of processes for DWI denoising task.
--processes_denoise_t1 NUMBER (default: 4) Number of processes for T1 denoising task.
--processes_eddy NUMBER (default: 1) Number of processes for eddy task.
--processes_fodf NUMBER (default: 4) Number of processes for fODF task.
--processes_registration NUMBER (default: 4) Number of processes for registration task.
--output_dir PATH (default: ./results) Directory where to write the final results.
--processes NUMBER (default: Maximum number of threads) The number of parallel processes to launch. Only affects the local scheduler.
6.1 Local computer

To run the pipeline, use the following command:

```bash
# With Singularity
$> nextflow run tractoflow/main.nf --root input_folder --dti_shells "DTI_SHELLS" --fodf_shells "FODF_SHELLS" -with-singularity singularity_name.img
```

```bash
# With Docker
$> nextflow run tractoflow/main.nf --root input_folder --dti_shells "DTI_SHELLS" --fodf_shells "FODF_SHELLS" -with-docker tractoflow:docker
```

Where `DTI_SHELLS` are the shells used to compute the DTI metrics (typically b-value < 1200 e.g. “0 1000”) and `FODF_SHELLS` are the shells used to compute the fODF metrics (typically b > 700 e.g. “0 1000 2000”).

6.1.1 Mounted partition

If your data is not on the same storage disk than your OS (e.g. a mounting disk, a USB stick, an external disk,...), you must bind your disk to the singularity container. Create a file (e.g. `singularity.conf`) and write the following line:

```bash
singularity.runOptions="--bind PATH_TO_DATA"
```

Where `PATH_TO_DATA` is the path to your storage disk.

Then run the following command:

```bash
# With Singularity
$> nextflow -c singularity.conf run tractoflow/main.nf --root input_folder --dti_shells "DTI_SHELLS" --fodf_shells "FODF_SHELLS" -with-singularity singularity_name.img
```

(continues on next page)
# With Docker

```
$> nextflow -c singularity.conf run tractoflow/main.nf --root input_folder --dti_shells "DTI_SHELLS" --fodf_shells "FODF_SHELLS" -with-docker tractoflow:docker
```

## 6.2 High Performance Computer (HPC)

The following example is based on the SLURM executor:

If you want to use only one node, please use the same commands presented for the local computer. The following lines must be saved in `.sh` file (e.g. `cmd.sh`) to be executed with `sbatch`.

```
#!/bin/sh

#SBATCH --nodes=1
#SBATCH --cpus-per-task=32
#SBATCH --mem=0
#SBATCH --time=48:00:00

nextflow -c singularity.conf run tractoflow/main.nf --root input_folder --dti_shells "DTI_SHELLS" --fodf_shells "FODF_SHELLS" -with-singularity singularity_name.img -with-mpi -resume
```

To launch on multiple nodes, you must use the MPI option that uses Ignite executor. The following example uses 2 nodes with 32 threads on each node. The following lines must be saved in `.sh` file (e.g. `cmd.sh`) to be executed with `sbatch`.

```
#!/bin/sh

#SBATCH --nodes=2
#SBATCH --cpus-per-task=32
#SBATCH --mem=0
#SBATCH --time=48:00:00

export NXF_CLUSTER_SEED=$(shuf -i 0-16777216 -n 1)

srun nextflow -c singularity.conf run tractoflow/main.nf --root input_folder --dti_shells "DTI_SHELLS" --fodf_shells "FODF_SHELLS" -with-singularity singularity_name.img -with-mpi -resume
```

As a local computer, you must bind your storage disk to the singularity (Please see Mounted partition subsection above).

To launch the pipeline on the HPC:

```
$> sbatch cmd.sh
```
Results

The pipeline creates 2 folders: results and work. The files in results are symlinks in works. We highly recommend to not remove work folder.

To transfert or copy-paste the results folder, please use one of the following commands:

```
# On local computer
$> cp -rL results NEW_PATH/results

# On HPC
$> rsync -rL login@adress:/HPC_PATH/results NEW_PATH/results
```
8.1 2.0.1

Date: 8 May 2019
Modify normalization mask and change some default option values

8.2 2.0.0

Date: 27 Mar 2019
First release for public access
Github repositories

TractoFlow pipeline repository: TractoFlow
TractoFlow Singularity repository: Singularity-TractoFlow
END USER LICENSE AGREEMENT

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Maxime Descoteaux (maxime.descoteaux@usherbrooke.ca)
If TractoFlow is used in a publication, please cite the following references:
A NeuroImage Toolbox paper for TractoFlow is currently under review.
Download:
References  Bibtex


