

# Baby Brain Toolkit

Fbrain ERC project: Computational Anatomy of Fetal Brain

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## 1 Introduction

BTK stands for Baby Brain Toolkit. This toolkit is developed in the context of the Fbrain ERC project: “Computational Anatomy of Fetal Brain”<sup>1</sup>. Studies about brain maturation aim at providing a better understanding of brain development and links between brain changes and cognitive development. Such studies are of great interest for diagnosis help and clinical course of development and treatment of illnesses. Several teams have begun to make 3D maps of developing brain structures from children to young adults. However, working out the development of fetal and neonatal brain remains an open issue. This project aims at jumping over several theoretical and practical barriers and at going beyond the formal description of the brain maturation thanks to the development of a realistic numerical model of brain aging.

### 1.1 Copyright

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<sup>1</sup><http://lsiit-miv.u-strasbg.fr/miv/index.php?contenu=erc>

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## 1.2 Installation

### 1.2.1 Dependencies

Baby Brain Toolkit (BTK) depends on:

- CMake ([www.cmake.org](http://www.cmake.org)): this library can be installed for debian-based distribution using the following command line : `apt-get install cmake cmake-curses-gui`
- Insight Toolkit (ITK) version 3.20 (it can be download at [www.itk.org](http://www.itk.org)). Extract this file (the directory InsightToolkit-3.20 is created). Make another directory where all the build files will be placed, e.g. InsightToolkit-3.20-build:

```
mkdir InsightToolkit-3.20-build
cd InsightToolkit-3.20-build
cmake ../InsightToolkit-3.20/
```

This will bring up the CMake configuration screen. Press [c] for configure and then use [t] to toggle the advanced mode. Make the following changes:

```
BUILD_TYPE = Release
ITK_USE_OPTIMIZED_REGISTRATION = ON
ITK_USE_REVIEW = ON
```

Then press [c] to configure and [g] to generate the make file. Finally, type `make` at the prompt to obtain the final build of ITK.

- Tclap library: this library can be installed for debian-based distribution using the following command line : `apt-get install libtclap-dev`
- OpenMP library: this library can be installed for debian-based distribution using the following command line : `apt-get install libgomp1`
- VTK library: this library can be installed for debian-based distribution using the following command line : `apt-get install libvtk5-dev`
- ANN library ([www.cs.umd.edu/~mount/ANN](http://www.cs.umd.edu/~mount/ANN)): this library can be installed for debian-based distribution using the following command line : `apt-get install libann-dev`

### 1.2.2 Download and compile the BTK sources

- Install Git: this library can be installed for debian-based distribution using the following command line : `apt-get install git-core`
- Get the BTK sources: `git clone https://github.com/rousseau/fbrain.git`
- Then:

```
mkdir fbrain-build
cd fbrain-build
cmake ../fbrain
make
```

Most of the programs of the BTK suite use the OpenMP library for multi-threading purpose. The number of cores used can be tuned using the following command line (in this example, 4 cores will be used): `export OMP_NUM_THREADS=4`

## 2 The whole pipeline

BTK allows to implement the whole pipeline for the processing of fetal images, i.e. the reconstruction of anatomical and diffusion data, and the final tractography, all expressed in the same local coordinate system. This processing can be summarized in the following steps:

1. Image conversion
2. Anatomical image reconstruction
3. Reconstruction of the diffusion sequence
4. Registration of diffusion to anatomical data
5. Tractography

In the following we explain the applications and utilities to use for each of the aforementioned steps.

### 2.1 Image conversion

BTK supports and has been tested by using images in Nifti format (<http://nifti.nimh.nih.gov/nifti-1>). However, images are frequently available in DICOM format and an image conversion is required. This can be performed by using `dcm2nii`, a converter freely available at <http://www.cabiatl.com/mricon/mricon/dcm2nii.html>. In this site you can find other options that can be best suited for your data.

### 2.2 Anatomical image reconstruction

This can be performed by using `btkImageReconstruction` (Section 3.2) followed by `btkReorientImageToStandard` (Section 4) to transform the result to a standard orientation.

### 2.3 Reconstruction of the diffusion sequence

To reconstruct diffusion data, you want to follow the steps described in Section 3.3. The use of two applications is required here: `btkGroupwiseS2SDistortionCorrection` and `btkRBFInterpolationS2S`.

### 2.4 Registration of diffusion to anatomical data

This can be performed by using `btkRegisterDiffusionToAnatomicalData` (Section 3.2)

### 2.5 Tractography

If you have followed the previous steps correctly, at this point you should have the reconstructed anatomical and diffusion data spatially aligned, and ready to perform the tractography. To do this, BTK provides `btkTractography` (Section 4)

## 3 Applications

### 3.1 Denoising

**btkNLMDenoising** This program applies a non-local mean filter to a 3D image for denoising purpose.

Usage: `-i input_image_filename -o output_image_filename`. The best results are usually obtained by using a mask (or a padding value).

**btkNLMDenoising4DImage** This program applies a non-local mean filter to each 3D image of a 4D image, for denoising purpose. Usage: `-i input_image_filename -o output_image_filename`. The best results are usually obtained by using a mask (or a padding value).

### 3.2 Anatomical reconstruction

**btkImageReconstruction** This program allows to obtain a high-resolution image from a set of low-resolution images, typically axial, coronal, and sagittal acquisitions [2].

Minimal usage: `btkImageReconstruction -i image1 ... -i imageN -o output --box`.

Recommended usage: `btkImageReconstruction -i image1 ... -i imageN -m mask1 ... -m maskN -o output --mask`. The use of a mask provide better results since it allows an accurately estimation of the initial transform, and constrains the registration to the region of interest.

The full list of optional parameters of the method can be obtained by `btkImageReconstruction --help`

### 3.3 Reconstruction of DW sequences

The reconstruction of diffusion-weighted (DW) sequences aims at obtaining a sequence corrected for fetal moving and eddy-current distortions. This can be performed in BTK by using the two following applications.

**btkGroupwiseS2SDistortionCorrection** It performs a groupwise slice-by-slice registration of the image components of the sequence.

Minimal usage: `btkGroupwiseS2SDistortionCorrection -i input.nii ... -g gradients_in.bvec -o output.nii -c gradients_out.bvec -t pathToTransforms`.

- i input sequence
- g gradient table for the input sequence
- o output sequence
- c gradient table for the output sequence
- t folder to save the transformation files

The slice-by-slice transform for a given DW image is saved as a set of  $N$  transforms in ITK format, with  $N$  the number of slices in the image.

NOTE: For slice registration, 20% of the samples are used for computing the image metric. As the region of interest of the slice can be small, this number of samples might be insufficient to compute the metric accurately. However, this percent of samples has been sufficient for the tested sequences.

**btkRBFInterpolationS2S** It performs an interpolation of scattered data generated from the application of slice-by-slice transforms. To this end, radial basis functions are used.

Minimal usage: `btkGroupwiseS2SDistortionCorrection -i input.nii ... -g gradients_in.bvec -r reference.nii -o output.nii -t pathToTransforms`.

- i input sequence
- g gradient table for the input sequence
- r a reference sequence providing the resampling grid
- o output sequence
- c gradient table for the output sequence
- t folder to save the transformation files

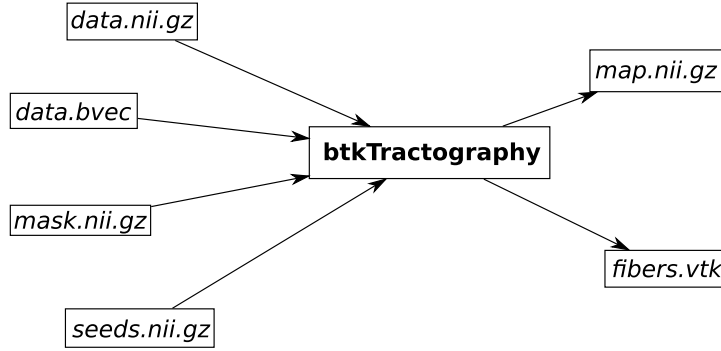


Figure 1: Standard pipeline of the btkTractography program.

### 3.4 Tractography

#### Standard usage

Suppose you want to perform a tractography on a diffusion weighted MRI dataset. You should have a dwi image, the corresponding gradient vectors' coordinates, a mask of the brain white matter and a label image of the seeds. Assume this data is stored in files named respectively for instance `data.nii.gz`, `data.bvec`, `mask.nii.gz` and `seeds.nii.gz`. The tractography is accomplished by the command below.

```
btkTractography -d data.nii.gz -g data.bvec -m mask.nii.gz -l seeds.nii.gz
```

When the program terminates its task, the probability connection map and the fibers estimation are saved in files respectively named `map.nii.gz` and `fibers.vtk`. The connection map is a volume image of probability intensities (i.e. intensities between 0 and 1) with the same origin, orientation and spacing as the diffusion weighted image. The fibers are polygonal data of VTK library in world coordinates. The standard pipeline of the program is shown in Fig. 1.

#### Advanced usage

In addition to standard arguments of `btkTractography` program, there are some other parameters that let you to alter algorithm's behaviour. These options can be classified into three groups : model's options, constraints on trajectory and filter's options. The first group options allow you to tweak the model (for more details about it, please refer to [1]). The second group options let you to control the particle's trajectory. These options provide prior informations to the algorithm. The last group options are dedicated to the particle filter control.

Since the default parameters values may work in the most of cases, they are optional. A list is of optional features is available by using the command

```
btkTractography --help
```

and program's arguments are much more described below.

#### Model's order

The model's order (i.e. the spherical harmonics' order) can be specified by the option

```
--model_order <order> ,
```

for  $\text{order} \in \{2, 4, 6, 8\}$ . The default value is 4. For more details, please refer to [1].

#### Model's regularization

A Laplace-Beltrami regularization coefficient is used to assume a better estimation of the model. This coefficient can be manually modified by the option

```
--model_regularization <coefficient> ,
```

for  $\text{coefficient} \in \mathbb{R}$ . The default value is set as 0.006. For more details, please refer to [1].

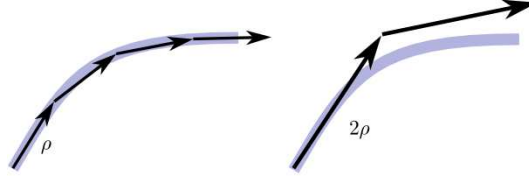


Figure 2: Effect of the step size option on a particle's trajectory. With a large step size (right), the particle may overshoot the trajectory of the ground truth.

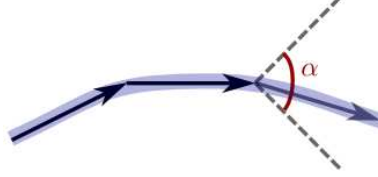


Figure 3: An angle threshold allows the algorithm to sample successive direction only in the cone defined by this angle. This illustration show the principle in two dimensions.

### Displacement step size

The displacement step size of a moving particle can ben adjusted as you want by using the option

```
--step_size <length> ,
```

where `length`  $\in \mathbb{R}_+^*$ . Note that this option is expressed in `mm`. The default value is fixed at `0.5 mm`. By setting a big step size, the particles will move quickly. So the bigger is the step, the faster the algorithm will finish, but as shown by Fig. 2, some informations may be missed and the particle's trajectories may overshoot the ground truth, resulting in a bad estimation.

### Angular threshold

An angular threshold prevent a particle to return back. This option has to be expressed in radian and can bet set by

```
--angular_threshold <angle> ,
```

where `angle`  $\in ]0, 2\pi[$ . The default value is set as a  $\frac{\pi}{3}$  angle. As illustrated in two dimensions in Fig. 3, an angle threshold is used to define an allowed area for successive sampled directions. This can be seen as a global curvature parameters on trajectories. A small angle defines trajectories with a small curvature. This is a prior information on ground truth trajectory.

### Rigidity

The rigidity option controls how much you want the particles to have straight trajectory. You can adjust it by

```
--curve_constraint <rigidity> ,
```

where `rigidity`  $\in \mathbb{R}_+^*$ . The default value is fixed at `30`. This value correspond to a concentration parameter of a von Mises-Fisher density probability used in the prior density of the system. As Fig. 4 illustrates locally in two dimensions, a high value leads to a straight trajectory.

### Number of particles

The number of particles in the system is set by the option

```
--number_of_particles <number> ,
```

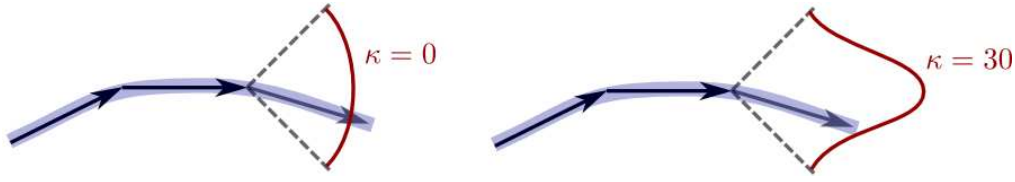


Figure 4: Local effect of rigidity parameter on a particle's trajectory. This parameter helps to “attract” the current displacement vector in the direction of the previous displacement vector of a particle. It correspond to a concentration paramter of a von Mise-Fisher density probability used in the prior density of the system. For instance, a rigidity of 0 leads to an equiprobable distribution, whereas a rigidity tending to infinity leads to a distribution focused on a point.

where **number**  $\in \mathbb{N}^*$ . By default, the algorithm will use 1000 particles. A poor number of particles leads to a short computation time and a poor estimation. A large number of particles leads to a long computation time and a good estimation. In general, the default number of particles is a good compromise between computation time and estimation.

### Resampling threshold

This option modify the resampling threshold of the system. When the number of effective particles in the system falls below this resampling threshold, the particles are resampled according a multinomial resampling. It can be adjust by

```
--resampling_threshold <percent> ,
```

where **percent**  $\in [0, 1]$  is the percent of minimal effective particles in the system. A low threshold value will result in an inefficient algorithm because the particles with low weight are not are not often eliminated. Conversely, a high threshold value leads to a bad estimation because the search space will not be explored enough.

## 4 Utilities

**btkModifyImageUsingLookUpTable** This program modifies one image using a look up table defined in a ascii file (2 columns, one for the original values, one for the final values). Usage: `-i input_image_filename -t input_table_filename -o output_image_filename`

**btkNrrdToNifti** This program convert an image from Nrrd file (\*.nhdr and \*.nrrd) to a Nifti file (\*.nii or \*.nii.gz). The conversion of a DWI image is possible by using the option `-d`. Usage: `-i input.nhdr -o output.nii.gz`. Usage for DWI sequence: `-i input.nhdr -o output.nii.gz -g gradients.bvec -d`.

**btkNiftiToNrrd** This program convert a diffusion sequence in nifti format<sup>2</sup> (\*.nii, \*.nii.gz) to the nrrd format (\*.nhdr).

Usage: `-i input.nii -b bvalues.bval -g gradients.bvec -o output.nhdr`

The list of optional parameters can be obtained by `btkNiftiToNrrd --help`

**btkSetStandardCoorSystem** It transforms an image to a coordinate system with identity direction and origin at the center of the image (by default, a linear interpolation is used).

Usage: `btkSetStandardCoorSystem -i image -o output -d 3`. The argument `-d` specifies the image dimension.

For diffusion sequences, the command line changes to provide the gradient table, which must be modified accordingly.

<sup>2</sup>Currently there is no nifti standard for DWI, so DW images are saved as a standard nifti sequence and two text files containing the b-values (.bval) and the gradient directions (.bvec).



Figure 5: Example of an anatomical reconstruction of a fetal brain by using `btkImageReconstruction`. (a) axial, (b) coronal, and (c) sagittal view.

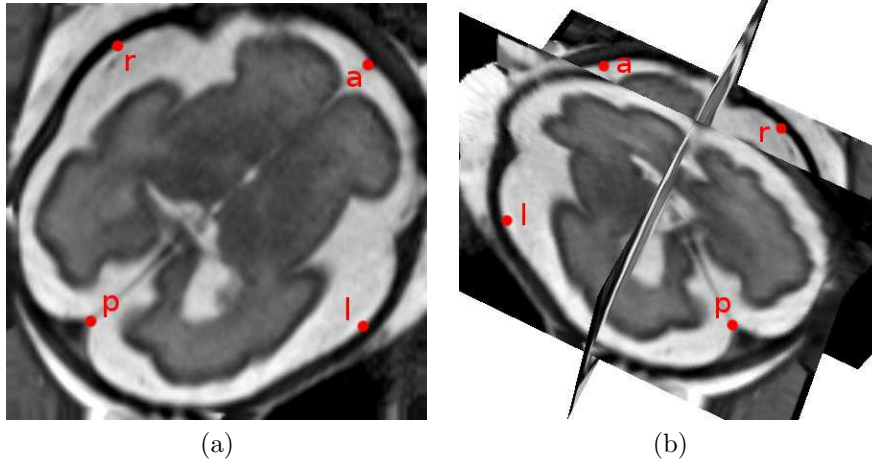


Figure 6: Placement of landmarks by using Slicer. (a) axial slice, (b) 3D view.

```
Usage: btkSetStandardCoorSystem -i image -o output -d 4 -g gradients.bvec
-c outputGradients.bvec
```

**btkReorientImageToStandard** Sometimes it is useful to reorient the image to the standard orientation. This is necessary with fetal images since in general the fetus is in a random orientation with respect to the scanner. To do this with BTK, before it is necessary to convert the image to a coordinate system by using `btkSetStandardCoorSystem`.

Usage: `btkReorientImageToStandard -i image -o output -l landmarks`. `landmarks` is a text file containing points that define the left-right and the posterior-anterior directions. The points *l* and *r* define the left → right direction, and the points *p* and *a* define the posterior → anterior direction. Such file can be easily generated by using Slicer<sup>3</sup> as follows:

1. Open the high-resolution image by using the *Volume* module.
2. Toogle on the visibility of all slices in the 3D view. This allows to identify the left and right sides of the brain in the 2D views.
3. Place the landmarks *l*, *r*, *p*, and *a* in this order by using [p].
4. Save the file (\*.fcsv) by using the menu File → Save.

**btkReorientDiffusionSequenceToStandard** Reorients a DW sequence to the standard orientation. This is necessary with fetal images since the fetus is in a random orientation with respect to the

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<sup>3</sup><http://www.slicer.org>



scanner. This is particularly important in DWI because colormaps lack of significance, which makes difficult the identification of specific bundles

Usage: `btkReorientDiffusionSequenceToStandard -i image -g gradients.bvec -l landmarks -o output -c gradients_corr.bvec`.

`landmarks` is a landmarks file obtained as explained above. `gradients_corr.bvec` is a text file containing the corrected gradient table.

**btkCropImageUsingMask** This program crops one (3D or 4D) image using a 3D mask. Usage: `-i input_image_filename -m input_mask_filename -o output_image_filename -d 3`, where '-d' is the dimension of the input image (by default 3).

**btkRegisterDiffusionToAnatomicalData** This program registers a DW sequence to an anatomical image.

Recommended usage: `btkReorientDiffusionSequenceToStandard -i input.nii -g gradients_in.bvec -r reference.nii -o output.nii -c gradients_out.bvec --mask mask.nii`.

- i input sequence
- g gradient table for the input sequence
- r reference image (anatomical image)
- o resampled sequence (by default, linear interpolation is used)
- c gradient table for the resampled sequence
- m image mask for the B0 image

The list of optional parameters can be obtained by `btkNiftiToNrrd --help`

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## References

- [1] M. Descoteaux, E. Angelino, S. Fitzgibbons, and R. Deriche. Regularized, fast, and robust analytical q-ball imaging. *Magnetic Resonance in Medicine*, 58(3):497–510, 2007.
- [2] Francois Rousseau, Orit A Glenn, Bistra Iordanova, Claudia Rodriguez-Carranza, Daniel B Vigneron, James A Barkovich, and Colin Studholme. Registration-based approach for reconstruction of high-resolution in utero fetal MR brain images. *Acad Radiol*, 13(9):1072–1081, Sep 2006.