Baby Brain Toolkit

Fbrain ERC project: Computational Anatomy of Fetal Brain

January 31, 2011

Contents

1	Introduction]
	1.1 Copyright	
		•
2	Applications	2
	2.1 Denoising	4
	2.2 Anatomical reconstruction	
	2.3 Tractography	,
3	Utilities	_

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



Figure 1: Example of an anatomical reconstruction of a fetal brain by using btkImageReconstruction. (a) axial, (b) coronal, and (c) sagital view.

1.2 Dependencies and Compilation

Baby Brain Toolkit (BTK) depends on:

- The most recent version of the Insight ToolKit (ITK) www.itk.org.
- 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

2 Applications

2.1 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).

2.2 Anatomical reconstruction

btkImageReconstruction This program allows to obtain a high-resolution image from a set of low-resolution images, typically axial, coronal, and sagital acquisitions [1].

 $\label{eq:minimal usage} \mbox{Minimal usage: btkImageReconstruction -i image1} \mbox{ \cdots -i imageN -o output.}$

Recommended usage: btkImageReconstruction -i image1 \cdots -i imageN -m mask1 \cdots -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

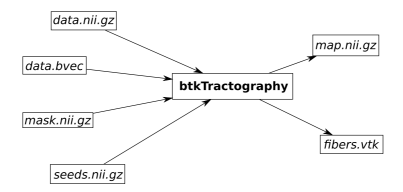


Figure 2: Standard pipeline of the btkTractography program.

2.3 Tractography

Theory

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 repsectively for instance data.nii.gz, data.bvec, mask.nii.gz and seeds.nii.gz. The tractography is accomplished by the command below.

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. 2.

Advanced usage

In addition to standard arguments of btkTractography program, there are some other parameters that let you to alter algorithm's behaviour. 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

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 <n>,
```

for $n \in \{2, 4, 6, 8\}$. The default value is 4.

Model's regularization

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

```
--model-regularization <r>,
```

for $r \in \mathbb{R}$. The default value is set as 0.006.

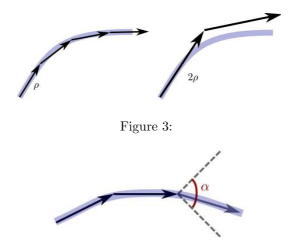


Figure 4:

Displacement step size

Angular threshold

Rigidity

Number of particles

Resampling threshold

3 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

Acknowledgment

The research leading to these results has received funding from the European Research Council under the European Communitys Seventh Framework Programme (FP7/2007-2013 Grant Agreement no. 207667).

References

[1] 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.

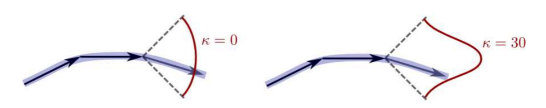


Figure 5: