

# Image Registration Driven by Combined Probabilistic and Geometric Descriptors <sup>★</sup>

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**Abstract.** Deformable image registration in the presence of considerable contrast differences and large-scale size and shape changes represents a significant challenge for image registration. A representative driving application is the study of early brain development in neuroimaging, which requires co-registration of images of the same subject across time or building 4-D population atlases. Growth during the first few years of development involves significant changes in size and shape of anatomical structures but also rapid changes in tissue properties due to myelination and structuring that are reflected in the multi-modal Magnetic Resonance (MR) contrast measurements. We propose a new registration method that generates a mapping between brain anatomies represented as a multi-compartment model of tissue class posterior images and geometries. We transform intensity patterns into combined probabilistic and geometric descriptors that drive the matching in a diffeomorphic framework, where distances between geometries are represented using currents which does not require geometric correspondence. We show preliminary results on the registrations of neonatal brain MRIs to two-year old infant MRIs using class posteriors and surface boundaries of structures undergoing major changes. Quantitative validation demonstrates that our proposed method generates registrations that better preserve the consistency of anatomical structures over time.

## 1 Introduction

Image registration is a basic task in defining a standard space for analyzing populations that change over time, which is essential to determine development in normal growth and neurological disorders. The growth process can involve large-scale size and shape changes, as well as changes in tissue properties and appearance. These factors pose significant challenges in image registration, as image intensities need to be interpreted differently at different stages. A strong example is the human brain at early development stages.

A driving research question is to determine the process of white matter myelination, which manifests as two distinct white matter appearance patterns primarily during the first year of development. Other clinical research questions are

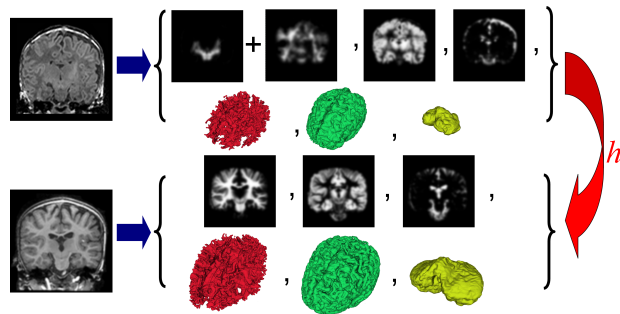
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related to finding a link between cognitive development and the rapid, locally varying growth of specific anatomical structures. To approach these questions, a robust registration method is necessary for mapping longitudinal brain MRI to a reference space so that we can perform reliable analysis of the tissue property changes reflected in the MR measurements. Knickmeyer *et al.*[1] showed that the total brain volume grows by 100% the first year and 15% the second year, whereas the cerebellum shows 220% volume growth for the first and another 15% for the second year, indicating the very different growth rates of different anatomical structures. Through regression on shape representations, Datar *et al.*[2] illustrated that the rapid volume changes are also paralleled by significant shape changes which describe the dynamic pattern of localized, nonlinear growth. These challenges require a method that does not rely on raw intensity measurements, while also being capable of estimating large structural deformations. Xue *et al.*[3] addressed these issues by proposing a registration scheme for neonatal brains by registering inflated cortical surfaces extracted from the MRI.

We propose a new registration framework for longitudinal brain MRI that makes use of the underlying anatomies, which are represented by both class posteriors and boundary surfaces. This framework is able to match internal anatomical regions and simultaneously preserving a consistent mapping for the boundaries of relevant anatomical objects. We show results of registering neonatal brain MRI to 2-year old brain MRI of the same subjects obtained in a longitudinal neuroimaging study. Our method consistently provides transformations that better preserve time-varying structures than obtained by intensity-only registration.

## 2 Registration between Anatomies



**Fig. 1.** Overview of the proposed registration method that can handle large deformations and different contrast properties, applied to mapping brain MRI of neonates to 2-year olds. We segment the brain MRIs and then extract equivalent anatomical descriptors by merging the two different white matter types present in neonates. The probabilistic and geometric anatomical descriptors are then used to compute the transformation  $h$  that minimizes the distance between the class posterior images, as well as the distance between surfaces represented as currents.

We propose a registration method that makes use of the underlying anatomy in the MR images. Fig. 1 shows an overview of the registration process. We begin by extracting anatomical descriptors from the images, followed by computing a transformation that minimizes the distance between the anatomical descriptors.

## 2.1 Anatomical Descriptors

We represent brain anatomy as a multi-compartment model of tissue classes and manifolds. We associate each position  $x$  with a vector of tissue probability densities. In a given anatomy, we capture the underlying structures by estimating, for each image, the class posterior mass functions associated with each of the classes. Given  $\Omega$  as the underlying coordinate system of the brain anatomies, each anatomy  $\mathcal{A}_{i=1,\dots,N}$  is represented as

$$\mathcal{A}_i = \{p_{i,c=1}(x), \dots, p_{i,c=N_c}(x), \mathcal{M}_{i,j=1}(2), \dots, \mathcal{M}_{i,j=N_s}(2) \subset \Omega\} \quad (1)$$

where  $N_c$  is the number of probability images,  $N_s$  is the number of surfaces,  $p_c(x)$  is the class posterior for tissue  $c$  at location  $x$ , and  $\mathcal{M}_j(2)$  are 2-dimensional submanifolds of  $\Omega$  (surfaces).

The classification of brain MR images with mature white matter structures into class posteriors are well studied. We extract the posteriors from 2-year old brain MR images using the segmentation method proposed by van Leemput *et al.* [4]. The method generates posterior probabilities for white matter (wm), gray matter (gm), and cerebrospinal fluid csf. These probabilities can then be used to generate surfaces from the maximum a posteriori tissue label maps.

The classification of neonatal brain MR images is challenging as the white matter structure undergoes myelination, where the fibers are being covered in myelin sheathes. Several have proposed methods that make use of prior information from an atlas or template that takes into account the special white matter appearance due to myelination [5] [6]. We use the method described by Prastawa *et al.* [6] for extracting the tissue class posteriors of neonatal brain MRI which includes for myelinated wm, non-myelinated wm, gm, and csf. These can then be used to create an equivalent anatomy to the 2-year old brain by combining the two white matter class probabilities and surfaces.

For the results in this paper, we use the probabilities  $\{p_{wm}(x), p_{gm}(x), p_{csf}(x)\}$  and we use the surfaces of white matter, gray matter, and cerebellum. The cerebellum surfaces are generated from semi-automated segmentations that are obtained by affinely registering a template image followed by a supervised level set segmentation. The cerebellum has a significant role in motor function and it is explicitly modeled as it undergoes the most rapid volume change during the first year of development and thus presents a localized large-scale deformation.

## 2.2 Registration Formulation

Given two anatomies  $\mathcal{A}_1$  and  $\mathcal{A}_2$ , the registration problem can be formulated as an estimation problem for the transformation  $h$  that minimizes

$$\hat{h} = \arg \min_h E(h \cdot \mathcal{A}_1, \mathcal{A}_2)^2 + D(h, e)^2 \quad (2)$$

where  $h \cdot \mathcal{A}_1$  is the transformed anatomy,  $E(\cdot, \cdot)$  is a metric between anatomies and  $D(\cdot, e)$  is a metric on a group of transformations that penalizes deviations from the identity transformation  $e$ .

We define distance between anatomies  $E$  by defining a norm on an anatomy as a combination of the  $L^2$  norm on the class posteriors and a Reproducing Kernel Hilbert space norm on the manifolds defined as “currents” through Glaunes [7]. The currents norm does not require geometric correspondence and thus can be used to register manifolds with different resolutions. For an oriented surface  $\mathcal{M}(2)$  in  $R^3$  the norm  $[\mathcal{M}(2)]$  is the vector valued Borel measure corresponding to the collection of unit-normal vectors to  $\mathcal{M}(2)$ , distributed with density equal to the element of surface area  $ds$  and can be written as  $\eta(x)ds(x)$ , where  $\eta(x)$  is the unit normal and  $ds(x)$  is the surface measure at point  $x$ . When  $\mathcal{M}(2)$  is a discrete triangular mesh with  $N_f$  faces, a good approximation of the norm can be computed by replacing  $[\mathcal{M}(2)]$  by a sum of vector-valued Dirac masses:

$$\|[\mathcal{M}(2)]\|_k^2 = \sum_{f=1}^{N_f} \sum_{f'=1}^{N_f} \langle \eta(f), \eta(f') \rangle k(c(f), c(f')), \quad (3)$$

where  $k(\cdot, \cdot)$  is a shift-invariant kernel (e.g., Gaussian or Cauchy),  $N_f$  is the number of faces of the triangulation, and for any face  $f$ ,  $c(f)$  is its center and  $\eta(f)$  its normal vector with the length capturing the area of each triangle.

Having defined the norm on probability images and surfaces, the dissimilarity metric between anatomies  $\left\| [\mathcal{A}_1] - [\mathcal{A}_2] \right\|_k^2$  is given by:

$$\sum_{c=1}^{N_c} \int_{\Omega} |p_{1,c}(x) - p_{2,c}(x)|^2 dx + \sum_{j=1}^{N_s} \|[\mathcal{M}_{1,j}(2) \cup (-\mathcal{M}_{2,j}(2))]\|_k^2 \quad (4)$$

where the distance between two surface currents  $\|[\mathcal{M}_{1,j}(2) - \mathcal{M}_{2,j}(2)]\|_k = \|[\mathcal{M}_1(2) \cup (-\mathcal{M}_2(2))]\|_k$  is computed as the norm of the union between surface  $\mathcal{M}_1(2)$  and surface  $\mathcal{M}_2(2)$  with negative measures.

We use the large deformation framework [8] that generates dense deformation maps in  $R^d$  by integrating time-dependent velocity fields. The flow equation is given by  $\frac{\partial h^v(t,x)}{\partial t} = v(t, h^v(t,x))$ , with  $h(0, x) = x$ , and we define  $h(x) := h^v(1, x)$ , which is a one-to-one map in  $\mathbb{R}^d$  (diffeomorphism). We define an energy functional that ensures the regularity of the transformations on the velocity fields:  $\|v(t, \cdot)\|_V^2 = \int_{\mathbb{R}^d} \langle Lv(t, x), Lv(t, x) \rangle dx$ , where  $L$  is a differential operator acting on vector fields. This energy also defines a distance in the group of diffeomorphisms:

$$D^2(h, e) = \inf_{v, p^v(1, \cdot) = h} \int_0^1 \|Lv(t)\|_V^2 dt. \quad (5)$$

The registration optimizations in this paper are performed using a greedy approach by iteratively performing gradient descent on velocity fields and updating the transformations via an Euler integration of the O.D.E. At each iteration of

the algorithm the velocity field is calculated by solving the p.d.e  $Lv = F(h)$  where  $F(h)$  is the variation of  $\left\| [h \cdot \mathcal{A}_1] - [\mathcal{A}_2] \right\|_k^2$  with respect to  $h$ . This variation is a combination of the variation of the  $L^2$  norm on the class posteriors and of the currents norm; computed using the gradient  $\frac{\partial \|\mathcal{M}(2)\|_k^2}{\partial x_r}$  :

$$\sum_{f|x_r \in f} \left[ \frac{\partial \eta(f)}{\partial x_r} \right] \sum_{f'=1}^{N_f} k(c(f'), c(f)) \eta(f') + \frac{2}{3} \sum_{f'=1}^{N_f} \frac{\partial k(c(f), c(f'))}{\partial c(f)} \eta(f')^t \eta(f) \quad (6)$$

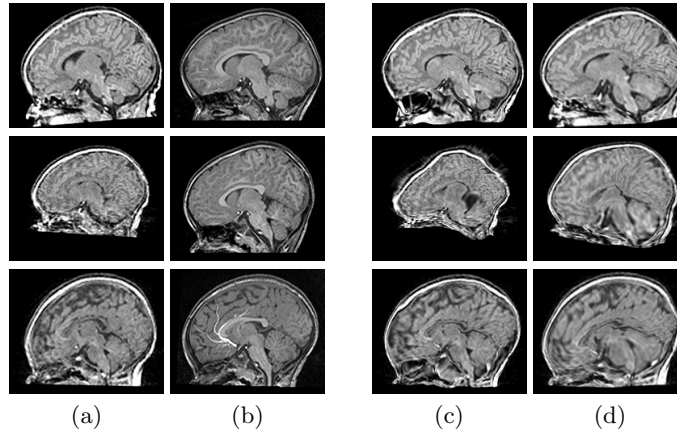
given that points  $\{x_r, x_s, x_t\}$  form the triangular face  $f$  and its center  $c(f) = \frac{x_r + x_s + x_t}{3}$  and its area-weighted normal  $\eta(f) = \frac{1}{2}(x_s - x_r) \otimes (x_t - x_r)$ .

### 2.3 Efficient Norm Computation using Particle Mesh Approximation on GPU

The major challenge of implementing the currents norm (Eq. 3) for realistic brain surfaces is the high computational cost to compute the dissimilarity metric of all pairs of surface elements, which is  $O(N_f^2)$  where  $N_f$  is the number of faces (can be up to millions). For computational tractability, Durrleman *et al.* [9] used a sparse representation of the surface based on matching pursuit algorithm. An efficient framework based on the standard fast Gauss transform [10] requires the construction and maintenance of the kd-tree structure on the fly, which is slow on the current CPU model. Our method, however, exploits the Particle-Mesh approximation to reduce the complexity to  $M \log M$  where  $M$  is the volume size of the embedded grid. The grid size is chosen as the size of the input images to limit the approximation error to the same order of matching term for the class posteriors. This approximation have been extensively studied in the cosmological N-body simulation literature (see Hockney and Eastwood [11] for details). Particle mesh framework shares the same computational grid with the class posteriors, moving the computation to the grid level and enabling an efficient parallel computation.

Even with the particle mesh approximation of the norm computation, the total complexity of the method is still very high. On a high-end workstation with 8-CPU cores, a highly optimized multi-threaded implementation in C++ takes several hours for one matching pair, hence can not be used for parameter exploration and real-time analysis. Fortunately, this computational cost can be amortized using the massive parallel power and bandwidth provided by the graphics processing unit (GPU). Based on the GPU framework by Ha *et al.* [12], we developed an implementation that runs entirely on the GPU. The main benefit of a GPU implementation is the ability to exploit parallel efficiency of regular grid presentation.

Computing the currents  $u$  and gradient between a surface with 160535 triangular faces and another with 127043 faces takes approximately 504 seconds on an AMD Phenom II X4 955 CPU, while it takes 0.33 seconds on an NVIDIA GTX 260 GPU. The speed gain is in order of three magnitudes over the equivalent CPU implementation using particle mesh, while the computing time for the



**Fig. 2.** Registration results of neonates mapped to two-year olds. From left to right: (a) neonatal T1 image after affine registration, (b) reference T1 image at 2-years, followed by (c) neonatal T1 after deformable mutual information registration using B-splines, and (d) after combined probabilistic and geometric registration. From top to bottom: subject 0012, 0102, 0106.

exact norm on CPU is difficult to measure since it takes significantly longer. The proposed algorithm typically converges in 1000 iterations, so on average it takes less than eight minutes to register two anatomies. This allows us to perform parameter exploration and real-time analysis on a single desktop with commodity GPU hardware. The efficiency of the GPU method also provides an opportunity to apply the algorithm for high quality atlas formation using our framework on a GPU cluster, which gives us the ability to perform statistical tests that are previously impossible due to excessive time requirements.

### 3 Results

We have applied the registration method for mapping neonatal MRI scans to two-year MRI scans of the same subjects in ten datasets. The datasets are taken from an ongoing longitudinal neuroimaging study with scans acquired at approximately two weeks, one year, and two years of age. Due to rapid early brain development, each longitudinal MR scan shows significant changes in brain size and in tissue properties. For comparison, we also applied the standard intensity based deformable registration using mutual information (MI) metric and B-spline transformation proposed by Rueckert *et al.*[13] which has been applied for registering 1-year old and 2-year old infants. The T1 weighted images before and after registration using the different approaches for the first three subjects are shown in Fig. 2.

A quantitative study of the performance of the registration method is performed by measuring the overlap between the transformed segmentation maps of neonates to the segmentation maps of two-year olds. Since we consider the

	Subject	0012	0102	0106	0121	0130	0146	0156	0174	0177	0180
White matter	MI	0.329	0.155	0.281	0.384	0.379	0.230	0.257	0.300	0.350	0.301
	P+G	0.497	0.397	0.442	0.453	0.482	0.414	0.461	0.440	0.478	0.442
Cerebellum	MI	0.755	0.212	0.588	0.515	0.732	0.820	0.713	0.569	0.631	0.777
	P+G	0.881	0.821	0.875	0.878	0.858	0.899	0.907	0.885	0.896	0.892

**Fig. 3.** Overlap measures comparing the registered segmentation maps against the reference segmentation maps for the white matter and cerebellum structure, obtained through deformable mutual information registration (MI) and our proposed method (P+G).

Subject	0012	0102	0106	0121	0130	0146	0156	0174	0177	0180
MI on CPU	92	63	103	92	101	112	106	99	91	96
P+G on GPU	9	8	8	8	8	7	9	8	7	7

**Fig. 4.** Time elapsed, in minutes, for registration using deformable mutual information on CPU (MI) and our proposed approach (P+G) on GPU with 1000 iterations of gradient descent.

segmentation maps at two years of age to be the standard, we use the following overlap metric:

$$Overlap(h \cdot S0, S2) = \frac{|h \cdot S0 \cap S2|}{|S2|} \quad (7)$$

where  $h \cdot S0$  is the transformed neonate segmentation map,  $S2$  is the reference two-year segmentation map, and  $|\cdot|$  indicates the volume of a binary map. Please note that this metric gives considerably lower values than the standard Dice coefficient. Fig. 3 shows the quantitative analysis for the white matter and cerebellum segmentation maps. Registration using both probabilistic and geometric descriptors provides consistently better results and are generally more stable for the structures of interest. In particular, our method better preserves the shape of the cerebellum, which has weak intensity boundaries in regions where it touches the cerebrum and thus cannot be registered properly using only image based information. Another significant challenge is that the cerebellar growth is distinctly different from the growth of neighboring structures. Registrations using our approach on the GPU takes 8 minutes on average, while registration on the CPU using mutual information metric and B-spline transformation takes 100 minutes on average. Detailed time measures are listed in Fig. 4. We have also performed validation using only the probabilistic descriptor, which generates results that are less accurate compared to our method (particularly for the cerebellum) while more accurate than image registration using MI.

## 4 Conclusions

We have proposed a registration framework that makes use of the probabilistic and geometric structures of anatomies embedded in the images. This allows us to

enforce matching of important anatomical features represented as regional class posteriors and tissue boundaries. Our framework allows us to register images with different contrast properties by using equivalent anatomical representations, and we have demonstrated results for registering brain MRIs with different white matter appearances at early stages of growth. The overlap validation measures in Fig. 3 show that geometric constraints, particularly for the cerebellum, is crucial for registering structures undergoing significant growth changes. In the future, we plan to apply this framework in early neurodevelopmental studies for analyzing the effects of neurological disorders such as autism and Fragile X syndrome. The proposed registration framework is generic and independent of the application domain, it can thus be applied to any registration where one encounters large-scale deformation and different appearance patterns.

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