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## Automatic Retrieval of Anatomical Structures in 3D Medical Images

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INSTITUT NATIONAL DE RECHERCHE EN INFORMATIQUE ET EN AUTOMATIQUE

***Automatic Retrieval of Anatomical Structures in  
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## Automatic Retrieval of Anatomical Structures in 3D Medical Images

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Programme 4 — Robotique, image et vision  
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**Abstract:** This paper describes a method to automatically generate the mapping between a completely labeled reference image and the 3D medical image of a patient. To achieve this, we combined three techniques: the extraction of 3D feature lines, their matching using 3D deformable line models, the extension of the deformation to the whole image space using warping techniques.

We present experimental results for the segmentation of structures in Magnetic Resonance images of the brain of different patients; the segmentation of the cortical and ventricle structures. We emphasize the advantages of using crest lines deformable models prior to surface based models. This gives a sparser representation of the data, easier to manipulate, and which makes the convergence of the model much less sensitive to initial positioning.

In the future, we hope to use this method to generate anatomical atlases, by the automatic interpretation of large sets of 3D medical images.

**Key-words:** deformable model, electronic atlas, feature line, non-rigid matching, warping.

*(Résumé : tsvp)*

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# Détection Automatique de Structures Anatomiques dans les Images Médicales 3D

**Résumé :** Le présent rapport décrit une méthode pour identifier automatiquement des structures anatomiques dans une image 3D d'un patient à partir d'une image où ces mêmes structures sont complètement référencées. Pour ce faire, nous avons combiné trois techniques : l'extraction de lignes caractéristiques, leur mise en correspondance en utilisant des modèles déformables de ces lignes, l'extension de la déformation ainsi obtenue à l'image entière grâce à des techniques de déformation d'images.

Nous présentons des résultats expérimentaux sur des images par résonance magnétique de cerveaux de patients différents, en identifiant plus précisément les structures corticales et ventriculaires. Nous insistons sur les avantages d'utiliser des modèles déformables de lignes de crête plutôt que des modèles de surfaces : en effet, cette représentation des données est plus concise et plus facile à manipuler. De plus, la convergence du modèle est moins sensible au positionnement initial.

A l'avenir, nous espérons appliquer ce procédé pour générer des atlas anatomiques électroniques par l'interprétation automatique de grands ensembles d'images médicales 3D.

**Mots-clé :** modèles déformables, atlas électronique, ligne caractéristique, mise en correspondance non-rigide, déformation.

# 1 Introduction

It becomes needless to emphasize the advantages of electronic atlases versus conventional paper atlases. Many examples of such atlases are known, such as the Voxel-Man ([28], [17], [24]), following the pioneering work of Bajcsy [2] and Evans [21].

However, even if such atlases are available, and even if Computer Graphic techniques are sufficiently developed to manipulate and display those atlases in real time, there remains a crucial need for a theoretical framework and *automatic* tools to :

- Generate atlases from large data sets,
- Average among features and models to create reference patients,
- Analyze the variability of features between patients,
- Find correspondences between the image of any patient and the atlases.

This paper presents one possible approach to achieve some of those goals, usually referred to as a segmentation problem, for which a strong a-priori knowledge of the human anatomy can be used.

As in many image processing problems, there are usually two dual and complementary ways to explore, which are the region based and the feature based techniques. The first kind of methods concentrates on the voxel values inside the regions (see for example [10]), whereas the second one concentrates on the boundaries of those regions (see for example [13] for 2D cases), such as the interfaces between organs, or specific lines or points of those surfaces (see [38], [36]).

In the present paper, we concentrate on a feature line based technique to segment fully automatically the same organ in the images of different patients. We give first a global description of the method, which is then detailed into feature lines extraction using differential geometry, registration of lines using deformable models, and at last 3D space deformation using warping techniques. Finally, we present a practical example, which is the automatic extraction of the cortical and ventricle surfaces from the 3D Magnetic Resonance image of a patient.

## 2 A global description of the method

We believe very much in bootstrapping techniques for atlas building: from an approximate description of a patient anatomy, or from existing electronic atlases or from a set of manually segmented 3D reference images, we hope to build automatically better average representations of the studied organs, usable for the re-interpretation of the reference images, or the analysis of a larger set of images. The key point here is of course the automatic inter-patient registration.

To simplify, let us start with a single reference image  $I_r$ , with an associated fully labeled image  $M_r$ , called map (figure 1): each voxel value of the map  $M_r$  specifies the type of a corresponding structure in the reference image  $I_r$ . We call *structure* a set of connected voxels of  $M_r$  having the same label.  $I_p$  is the image of a new patient to process.

We will suppose also that images  $I_r$  and  $I_p$  have been acquired with the same modality and parameter settings: their intensities are very similar.

To find the correspondence between  $I_p$  and the reference map  $M_r$ , we propose to find first the correspondence between  $I_p$  and  $I_r$  and then to deform the reference map  $M_r$  into a new map  $M_p$ , exactly superimposable to  $I_p$ .

### 2.1 The general scheme

More precisely, what we propose is:

- to automatically find and label crest lines in  $I_r$ , corresponding to the structures of  $M_r$ ,
- to find automatically the corresponding lines in  $I_p$ ,
- to find the correspondence  $D_{p,r}$  between those sets of lines,
- to either deform individual structures, or the global map  $M_r$  into new structures or into a new map  $M_p$ , which is exactly superimposable to  $I_p$ . This step is achieved by applying a warping technique to  $M_r$  using the found correspondences (see [6], [14]).



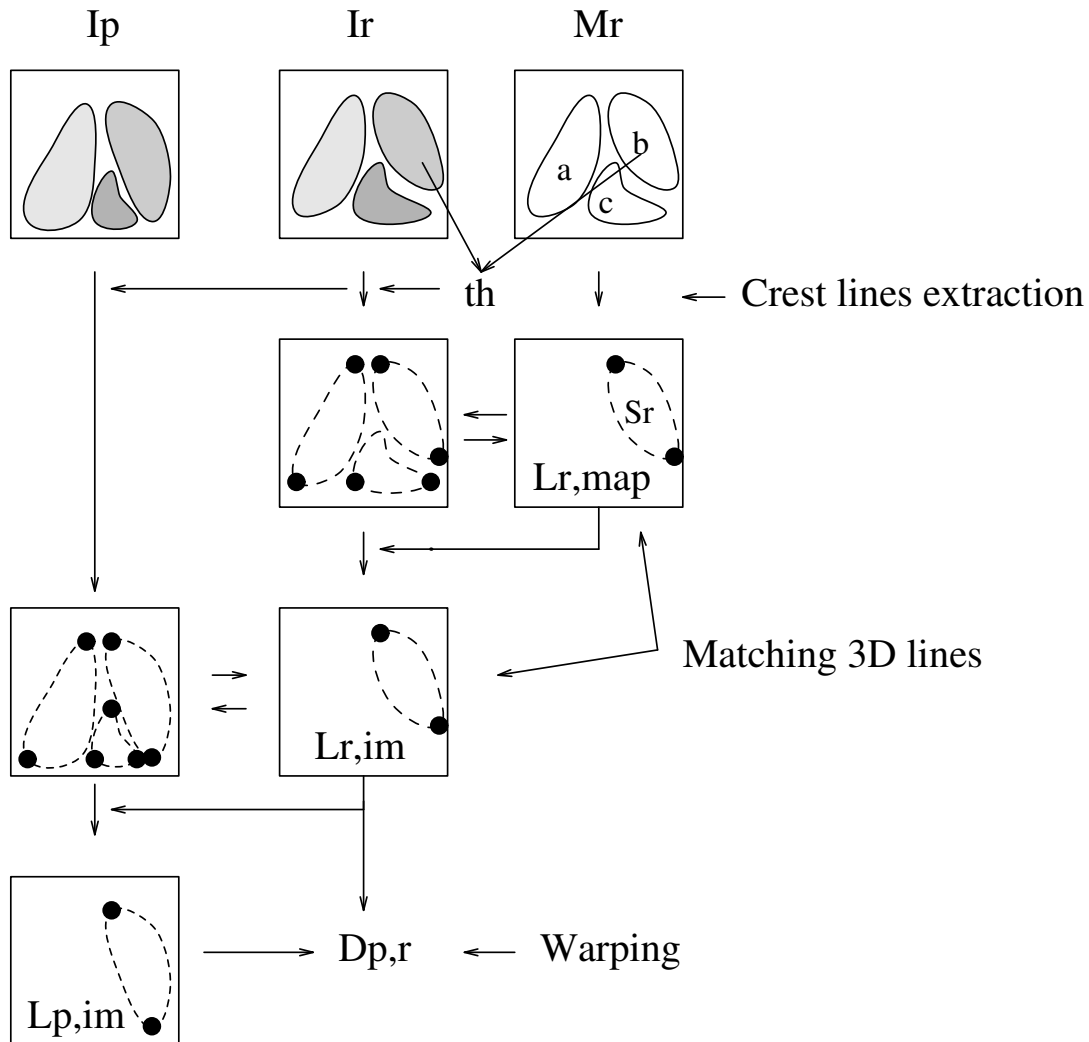


Figure 1: Crest lines  $L_{r,map}$  are extracted from the reference map and used to find the corresponding lines  $L_{r,im}$  in the reference image  $I_r$ . The crest lines  $L_{p,im}$  are extracted from the new image  $I_p$ , using  $L_{r,im}$ . This gives a set of correspondence and warping is applied to obtain the final deformation  $D_{p,r}$ .

## 2.2 The automatic labeling of crest lines

It is achieved structure by structure, fully automatically, in the following way (see also figure 1):

- Select one structure  $S_r$  of  $M_r$  (for example the ventricles of the brain).
- Extract the crest lines  $L_r^{map}$  of  $S_r$  (see [38]).
- Find the threshold  $th$  in  $I_r$  representative of the interface between  $S_r$  and the other structures, for example by computing the average value of the voxels of  $I_r$  in a region defined by the subtraction of the dilated and eroded versions of  $S_r$  (see [30]) for the definitions of those words).
- Extract the crest lines  $L_r^{im}$  in  $I_r$  imbedded in the iso-surface  $th$ .
- Register the two sets of crest lines  $L_r^{im}$  and  $L_r^{map}$  (see [18], [32]), to filter out from  $L_r^{im}$  the lines which have no correspondence in  $L_r^{map}$ .

The set of crest lines  $L_r^{im}$  is labeled with the name of the structure  $S_r$ .

## 2.3 The automatic retrieval of corresponding lines

We now search the corresponding lines  $L_p^{im}$  in the new image  $I_p$ :

- Extract the crest lines  $L_p^{im}$  of  $I_p$  within the iso-surface  $th$  (the hypothesis of similar dynamic of images is used here).
- Register  $L_p^{im}$  and  $L_r^{im}$ , and filter out from  $L_p^{im}$  the lines which have no correspondence in  $L_r^{im}$ .

At this point, we can compute a 3D B-spline approximation of a deformation  $D_{p,r}$  between the two image spaces  $I_p$  and  $I_r$ . This can be achieved on one hand for each individual structure  $S_r$ , using the corresponding points between  $L_p^{im}$  and  $L_r^{im}$ .  $D_{p,r}$  is then applied to  $S_r$  to obtain the structure  $S_p$  which is superimposable to  $I_p$ . On the other hand, this can be applied to the global map  $M_r$ , using simultaneously all structures, to deform  $M_r$  into  $M_p$ , the new map of image  $I_p$ .

### 3 Feature based non-rigid registration

This part describes the automatic extraction and deformable registration of crest lines.

#### 3.1 The feature type

Raw medical images are stored in a discrete 3D matrix  $I = f(x, y, z)$ . By thresholding  $I$ , isosurfaces of organs are computed (for instance, the surface of the skull for CT-Scan, of the brain or the face for MRI). The problem, then, is to compute specific features of these surfaces. Several methods have been proposed to achieve this:

- *surface features* : the mean and Gaussian curvatures are used to segment the isosurface into patches of some fundamental types. Such a decomposition permits to study the deformations of the left ventricle [16] or to describe the faces [9].
- *line features* : Hosaka [20] reports a wide range of characteristic lines based on differential geometry. The 3D Medial Axis Transform gives also sets of lines, charting for instance the gyral anatomy [33].
- *point features* : the “extremal points” [37], based on geometric invariants are used to perform 3D rigid registration.

A first example of clinical application can be found in Cutting et al. [12], where line and point features are both used to compute “an average” skull. In that study, however, the features are manually extracted.

#### 3.2 The crest lines

As explained before, we choose to use only line features: the crest lines introduced in [22]. They are defined as the successive loci of a surface for which the largest principal curvature is locally maximal in the principal direction (see figure 2). Let  $k_1$  be the principal curvature with maximal curvature in

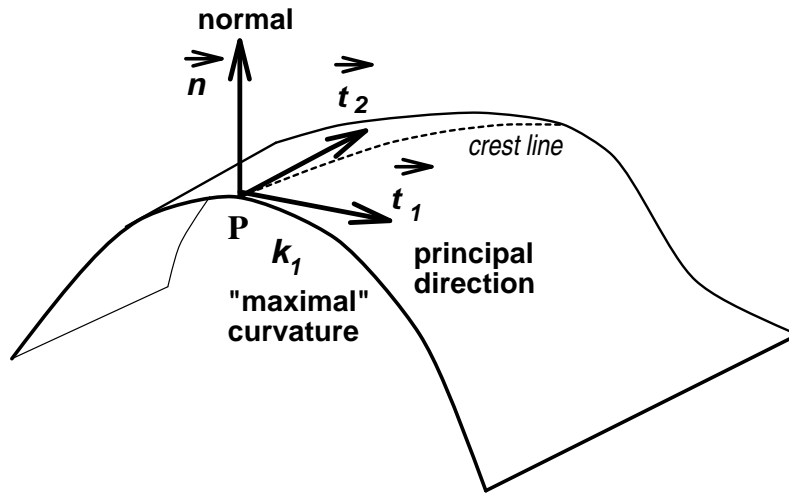


Figure 2: *Differential characteristics of a surface and the crest line.*

absolute value and  $\vec{t}_1$  the associated principal direction, each point of a crest line verifies:  $\vec{\nabla} k_1 \cdot \vec{t}_1 = 0$ .

These lines are automatically extracted from an isosurface by the “marching lines” algorithm [38].

As we can see in the figure 5, crest lines are anatomically meaningful: on the skull, the crest lines represent the salient lines (the orbits, the nose, the mandible or the temples) as emphasized in [7]; on the brain, the crest lines follow the convolutions, the sulci and the gyri patterns described in Ono et al. [25].

### 3.3 The 3D registration algorithm

The 3D curves registration algorithm is a key point in our scheme: given two sets  $A$  and  $A'$  composed of the crest lines  $L_i$  and  $L'_j$  extracted from images of two different patients, we want to find which lines  $L_i$  of  $A$  (or portions  $P_{i,k}$ ) correspond to which lines  $L'_j$  (or portions  $P_{j,l}$ ) of  $A'$  (see figure 3). Two difficulties arise: the number of lines of each set is quite large (several hundreds,

sometimes more than a thousand) and the registration between  $A$  and  $A'$  is not rigid.

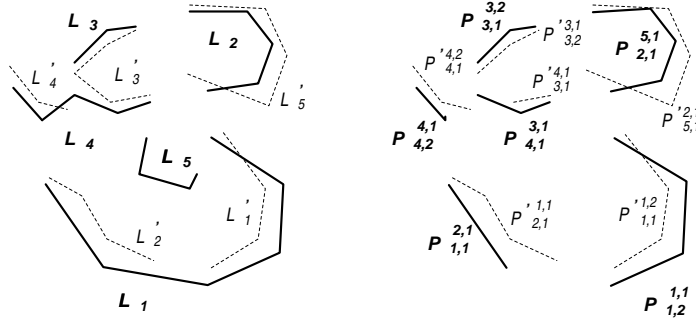


Figure 3: The registration algorithm has to find the portions  $P_{i,k}^{j,l}$  and  $P_{j,l}^{i,k}$ , respectively the  $k^{\text{th}}$  portion of  $L_i$  which corresponds to the  $l^{\text{th}}$  portion of  $L_j$  and vice versa.

In [3] and [29], 3D curves matching enables to recognize rigid synthetic objects. First, boundary curves are smoothed and then matched with prestored models but the registration is only rigid. In [18], the algorithm smoothes curves by using non-uniform B-splines. Then the two sets of curves are matched with a hashing table indexed by euclidean differential invariants. Results are very good [1] especially with sets of crest lines but the method only succeeds in finding a rigid displacement and cannot be generalized easily to the non-rigid case. Zhang in [39] and independently Besl [4] introduced an “iterative closest points” matching method (also generalized in [34]). It consists in three steps: for each point  $M_i$  of  $A$ , find the closest point  $M'_i$  of  $A'$ . Then, compute the global rigid displacement between the two sets of matched points  $(M_1 \dots M_n)$  and  $(M'_1 \dots M'_n)$  by a least-squares technique. Apply this motion to  $A$  and iterate until the motion is “small”. Both authors use the algorithm to register free-form curves but once again for the rigid case. Nevertheless, we can improve and generalize this method to our problem.

Our algorithm follows the steps of the “iterative closest point” method.

### 3.3.1 Points matching

Each point of  $A$  is linked with its closest neighbour in  $A'$  according to the euclidean distance. We plan also to use in the distance computation the differential curve parameters as the tangent, normal, curvature and torsion [18] or surface parameters as the normal, the principal directions and principal curvatures as described in [15]. This gives two lists of registered points,  $\mathcal{C}_1$ .

But as we have curves, i.e. an *ordered* list of points, we can apply some topological constraints in order to remove no-consistent couples of linked points and to avoid the configurations of the figure 4 top where the line  $L_1$  belongs to  $A$  and  $L_2$  to  $A'$ .

With these couples of points  $\mathcal{C}_2$ , two coefficients are computed:  $p_i^j$  and  $p_j^i$  which are the proportion of the curve  $i$  of  $A$  matched with the curve  $j$  of  $A'$  and vice versa. Thus, by thresholding,  $p_i^j \geq thr$  and  $p_j^i \geq thr$ , we can determine the curves “registered” at  $thr$  percent. For instance, curves can be considered completely registered when  $p_i^j \geq 0.5$  and  $p_j^i \geq 0.5$ .

### 3.3.2 Least-squares transformation

We register  $A$  and  $A'$  with polynomial transformations. The 0<sup>th</sup>-order is a rigid transformation and 1<sup>st</sup>-order an affine transformation but they are not sufficient for satisfying non-rigid registration. So, we use 2<sup>nd</sup>-order polynomial transformations defined by:

$$\begin{cases} x' &= a_1x^2 & +a_2y^2 & +a_3z^2 & +a_4xy & +a_5yz & +a_6xz & +a_7x & +a_8y & +a_9z & +a_{10} \\ y' &= b_1x^2 & +b_2y^2 & +b_3z^2 & +b_4xy & +b_5yz & +b_6xz & +b_7x & +b_8y & +b_9z & +b_{10} \\ z' &= c_1x^2 & +c_2y^2 & +c_3z^2 & +c_4xy & +c_5yz & +c_6xz & +c_7x & +c_8y & +c_9z & +c_{10} \end{cases}$$

As these polynomials are linear in their coefficients, we can use the least-squares method [26], [5] to compute  $a_i$ ,  $b_i$  and  $c_i$ .

Higher order polynomials may create large unexpected undulations as emphasized in [8]. 2<sup>nd</sup>-order polynomial transformations give accurate registration but we are not able to decompose them into intuitive physical meaning transformations such as rotation, translation or scaling. Notice that, at each iteration, we compose the transformation with a 2<sup>nd</sup>-order polynomial and so, we obtain after  $n$  iterations, a 2<sup>n</sup>-order polynomial transformation. However,

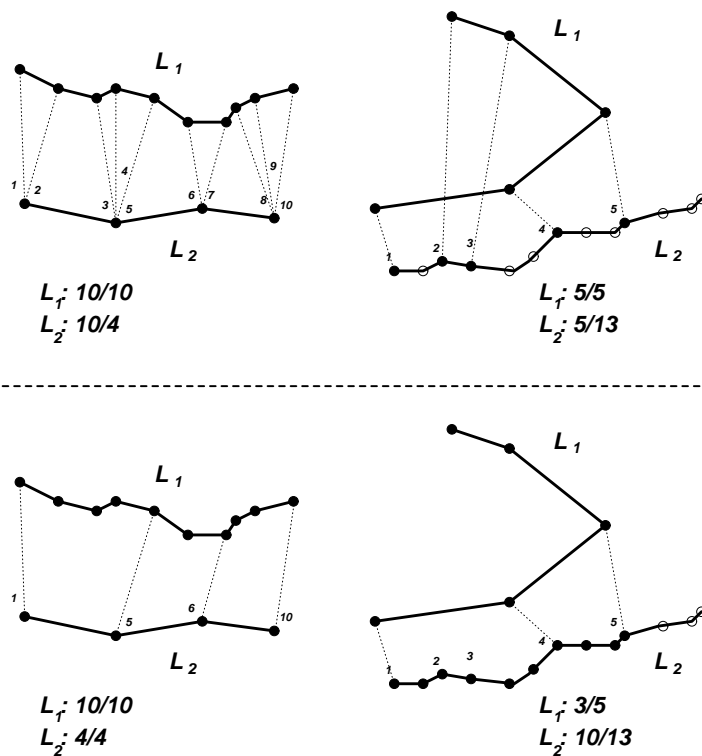


Figure 4: *The topological constraints help to remove inconsistent couples of linked points: left, some points have too many correspondents, right, a part of  $L_1$  have a erroneous correspondent on  $L_2$ .*

such a iterative composition does not create the undulations [8] emphasizes: it seems that those undulations do not appear for such a class of polynomial transformations.

Such transformations are also used by Greitz et al. in [17] to model natural deformations as brain bending called scoliosis.

### 3.3.3 Updating

The transformation is applied, then the algorithm iterates again or stop according to several criteria (mean value of the distance distribution between matched points, stability of the registration coefficients  $p_i^j$  and  $p_j^i$ , threshold on the matrix norm  $\|T - I_d\|$  where  $T$  is the transformation and  $I_d$  the identity matrix).

### 3.3.4 Parameters adaptation

By incrementing the threshold value *thr* at each iteration, for instance, from 0 to 0.5 by step of 0.025 and by taking only into account the matched point couples  $(M, M')$  belonging to “registered” curves at *thr* percent, the algorithm tends to improve the registration of already matched curves and to discard isolated ones. Moreover, we can begin to apply rigid transformations to align the two sets of lines, then affine transformations to scale them and, at last, quadratic transformations to refine the registration.

At the end, we obtain a good registration between the two sets of lines and a point to point correspondence between lines; however, the transform is global, we need then to use a B-spline based warping technique to obtain a more local and a better approximation of the deformation.

## 4 Warping with B-splines

This section describes the method that has been developed to get a full superposition of two images ( $I_r$  and  $I_p$ ), having a sparse set of corresponding points.

### 4.1 The problem

The process detailed in the previous section gives a set of pairs of points, each pair contains two matched points in both images. The aim of the following technique is to establish a matching on the whole images. Let us consider  $F$  as the matching function, this means a geometric transformation that, taking



a point in  $I_r$ , gives an anatomically equivalent point in  $I_p$ :

$$\begin{array}{ccc} F : I_r & \longrightarrow & I_p \\ & & P_r \longmapsto P_p \end{array}$$

This function has obvious regularity properties.

A pair of points obtained with the above algorithm is hence an estimation of the pair  $(P_r, F(P_r))$ . Given the set of such pairs of landmarks, we have a partial knowledge of  $F$ , which will help us to define an estimation  $\phi$  of it on the whole image. Having  $\phi$ , it will be possible to warp the first image on the second.

## 4.2 Calculation of the warping function

A similar study has been proposed by Bookstein and Green ([6]), they calculate  $\phi$  as a thin-plate spline interpolating function. We adopted a similar approach, but defining  $\phi$  as an approximation of  $F$  rather than an interpolation.

### 4.2.1 The B-spline approximation

Let us consider  $(u, v, w)$  the coordinate functions of  $\phi$ . We define them as a three-dimensional tensor product of B-spline basis functions:

$$\begin{aligned} u(x, y, z) &= \sum_{i=0}^{n_x-1} \sum_{j=0}^{n_y-1} \sum_{k=0}^{n_z-1} \alpha_{ijk} B_{i,K}^x(x) B_{j,K}^y(y) B_{k,K}^z(z) \\ v(x, y, z) &= \sum_{i=0}^{n_x-1} \sum_{j=0}^{n_y-1} \sum_{k=0}^{n_z-1} \beta_{ijk} B_{i,K}^x(x) B_{j,K}^y(y) B_{k,K}^z(z) \\ w(x, y, z) &= \sum_{i=0}^{n_x-1} \sum_{j=0}^{n_y-1} \sum_{k=0}^{n_z-1} \gamma_{ijk} B_{i,K}^x(x) B_{j,K}^y(y) B_{k,K}^z(z) \end{aligned}$$

with the following notations (for the  $x$  coordinate, for instance):

- $n_x$  : the number of control points in the  $x$  direction. It controls the accuracy of the approximation.

- $\alpha$  : the 3D matrix of the control points abscissae. This is what we are looking for.
- $B_{i,K}^x$  : the  $i^{\text{th}}$  B-spline basis function. Its order is  $K$ . These  $B_{i,K}^x$  generate the vectorial space of piecewise  $K^{\text{th}}$  degree polynomials. (see [27]).  $u$  is then a piecewise  $K^{\text{th}}$  degree polynomial in each variable  $x$ ,  $y$  and  $z$ .

We choose cubic B-splines in our examples ( $K = 3$ ), for their regularity properties. For the knots, we took the classic regular mesh:

$$\begin{aligned} t_0^x &= \dots = t_K^x = \min_x \\ t_i^x &= \min_x + (\max_x - \min_x) \frac{i - K}{n_x - K} \quad \text{for } K < i < n_x \\ t_{n_x}^x &= \dots = t_{n_x+K}^x = \max_x \end{aligned}$$

where  $\min_x$  and  $\max_x$  are the boundaries of the definition domain (the image domain). The knots values can be optimized to get an accurate approximation, but the data are not that precise.

#### 4.2.2 The constraints

We try to determine the best  $\phi$ , with respect to our data. We then define three criteria  $J^x$ ,  $J^y$  and  $J^z$ , one for each coordinate. For instance, for  $u$ ,  $J^x$  splits in two parts:

- position term. For each data point,  $u$  taken on the point in the first image must be as close as possible to the abscissa of the corresponding point in the second image. We choose a least square criterion:

$$J_{\text{position}}^x(u) = \sum_{l=1}^N \left( u(x_1^l, y_1^l, z_1^l) - x_2^l \right)^2$$

which is developed as:

$$J_{\text{position}}^x(u) = \sum_{l=1}^N \left( \sum_{i=0}^{n_x-1} \sum_{j=0}^{n_y-1} \sum_{k=0}^{n_z-1} \alpha_{ijk} B_{i,K}^x(x_1^l) B_{j,K}^y(y_1^l) B_{k,K}^z(z_1^l) - x_2^l \right)^2$$

$\alpha_{ijk}$  is the 3D matrix of the control points abscissae,  $x_1^l$  the abscissa of the  $l^{\text{th}}$  data point of the first image, etc...

- smoothing term. B-splines have intrinsic rigidity properties, but it is sometimes not enough. We choose a second order Tichonov stabilizer: it measures how far from an affine transformation the deformation is.

$$J_{smooth}^x(u) = \rho_{smooth} \int \int \int_{\mathbb{R}^3} \left[ \frac{\partial^2 u^2}{\partial x^2} + \frac{\partial^2 u^2}{\partial y^2} + \frac{\partial^2 u^2}{\partial z^2} + \frac{\partial^2 u^2}{\partial x \partial y} + \frac{\partial^2 u^2}{\partial x \partial z} + \frac{\partial^2 u^2}{\partial y \partial z} \right]$$

where  $\rho_{smooth}$  is a balance coefficient. It is manually defined, some solutions to choose it automatically is currently studied. The integrals are calculated with the Gauss-Legendre algorithm, which gives exact results for polynomials with very few evaluations of the integrand.

The criterion to minimize is the sum of those two:

$$J^x(u) = J_{position}^x(u) + J_{smooth}^x(u)$$

### 4.2.3 The linear systems

$J$  is a positive quadratic function of the  $\alpha_{ijk}$  variables. To find the coefficients that minimizes  $J^x$ , we derive its expression with respect to all the  $\alpha_{ijk}$ : it gives  $n_x \cdot n_y \cdot n_z$  linear equations which are written, for  $0 \leq a < n_x$ ,  $0 \leq b < n_y$  and  $0 \leq c < n_z$ :

$$\begin{aligned} & \sum_{i,j,k} \alpha_{ijk} \left[ \sum_{l=1}^N B_a^x(x_1^l) B_b^y(y_1^l) B_c^z(z_1^l) B_i^x(x_1^l) B_j^y(y_1^l) B_k^z(z_1^l) \right. \\ & \quad \left. + \rho_{smooth} \left( I_{abc,ijk}^{xx} + I_{abc,ijk}^{yy} + I_{abc,ijk}^{zz} + 2 I_{abc,ijk}^{xy} + 2 I_{abc,ijk}^{xz} + 2 I_{abc,ijk}^{yz} \right) \right] \\ & = \sum_{l=1}^N B_a^x(x_1^l) B_b^y(y_1^l) B_c^z(z_1^l) x_2^l \end{aligned}$$

with shortening notations for the smoothing term:

$$\begin{aligned} I_{abc,ijk}^{xx} &= \int \int \int_{\mathbb{R}^3} B_a''^x B_b^y B_c^z B_i''^x B_j^y B_k^z \\ I_{abc,ijk}^{xy} &= \int \int \int_{\mathbb{R}^3} B_a'^x B_b'^y B_c^z B_i'^x B_j'^y B_k^z \\ &\dots \end{aligned}$$

Each integral is separable in a product of 3 simple integrals, they are hence easy to compute.

#### 4.2.4 The resolution of the systems

The assembling of the matrix of the linear system is easy because this matrix is sparse: when all  $B_{i,K}^x$  are evaluated for a given  $x$ , a maximum of  $K$  functions are non-zero (and then a minimum of  $n_x - K$  is equal to zero). Moreover, the matrix is symmetric and positive, because the criterion is positive.

We use then a conjugate gradient method to solve our three systems (one for each coordinate).

### 4.3 Advantages of this warping method

The main advantages to compute B-splines are threefold:

- the B-splines functions are easy to evaluate with the Casteljau algorithm. The assembly of the matrices and the evaluation of the  $\phi$  are then fast.
- the intrinsic rigidity properties of B-splines gives a regular function.
- a data point has a local influence : to evaluate an image of a point, we need only  $(K + 1)^3$  control points (to be compared with the  $n_x \cdot n_y \cdot n_z$  that have been calculated), those which are controlling the area around this point. Hence, the influence of outliers is very local.

## 5 Results and discussion

The data are presented on figures 6 and 7. On top, the reference brain extracted from  $I_r$ . Bottom, the patient brain extracted from  $I_p$ . Notice the differences of shapes and orientations; the patient brain is more compact than the reference brain, and it is rotated by a few degrees. The figure 7 shows the crest lines of the surfaces of the brains. The thin lines are those of the brain, the thick ones are those of the ventricle of each brain. Notice how different they are. These crest lines were the lines used in the registration algorithm.

On figure 8, the top line is the reference image  $I_r$  with the reference cortical surface  $S_r$ . The middle line is the image of the new patient  $I_p$  with the reference cortical surface  $S_r$  before deformation. The bottom line is the image  $I_p$  with the result  $S_p$  of the found deformation applied on  $S_r$ : see how it follows the

convolutions. The next figure shows a similar sequence with the surface of the ventricle.

The figure 10 helps to have an overall view of the deformation : it shows different 3D views of the reference brain (left) and the same brain after warping (right). The brains have been cut so that we can see the ventricles (the dark-grey structure) in their respective position.

The results are very good on that specific example, and they are encouraging in the perspective of a completely automatic registration process, given reference and patient images.

## 6 Conclusion and perspectives

The proposed method allows us to build fully automatically the maps associated to the 3D images of new patients, from manually designed maps of reference patients. It can be used to efficiently initialize 3D surface snakes if a more precise final segmentation of the organs is needed (see [11], [35]). The advantage of using crest lines prior to surface models is to have a much more compact representation, more easy to manipulate, that allows us to explore more numerous deformation hypotheses. Also, finding point to point correspondences between 3D lines is less ambiguous than between 3D surfaces, because lines are much sparser, and often correspond to anatomical landmarks. By having a very good starting point, and 3D structures whose topology is inferred directly from the reference maps, 3D surface snakes are more likely to converge toward the desired solution.

We are currently studying the integration of such 3D surface snakes into our method ([11]), in order to improve the quality of the automatically generated maps, and we develop also tools for averaging between patient features, and measuring variability (see [23], [32]). In the long run, we shall validate more thoroughly this study with a larger number of cases, and we plan to build tools for the automatic generation of anatomical atlases, using bootstrapping.

## **7 Acknowledgements**

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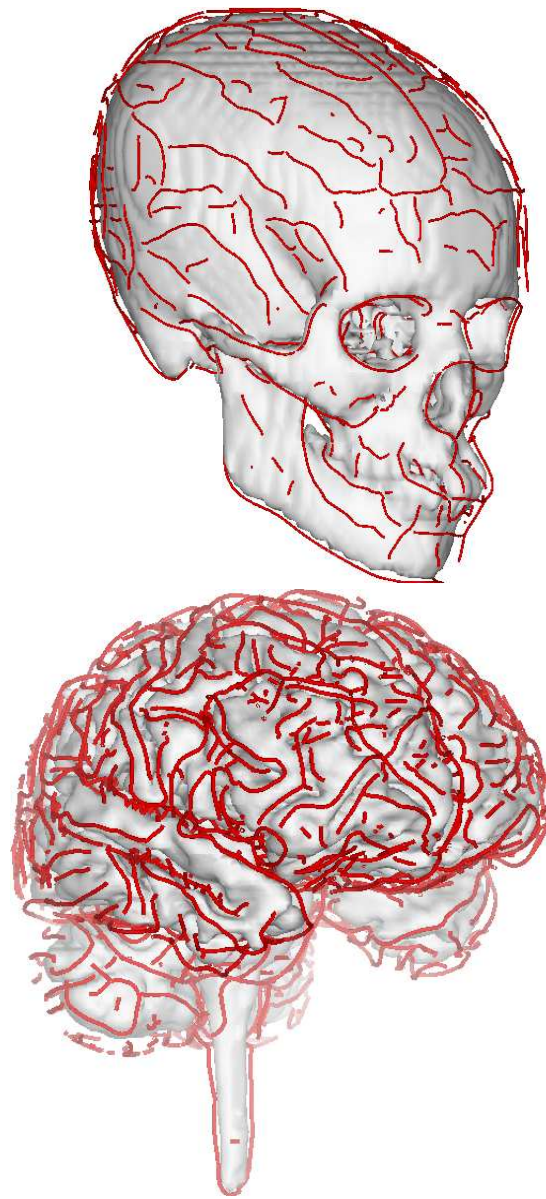


Figure 5: *Crest lines of a skull and of a brain.*

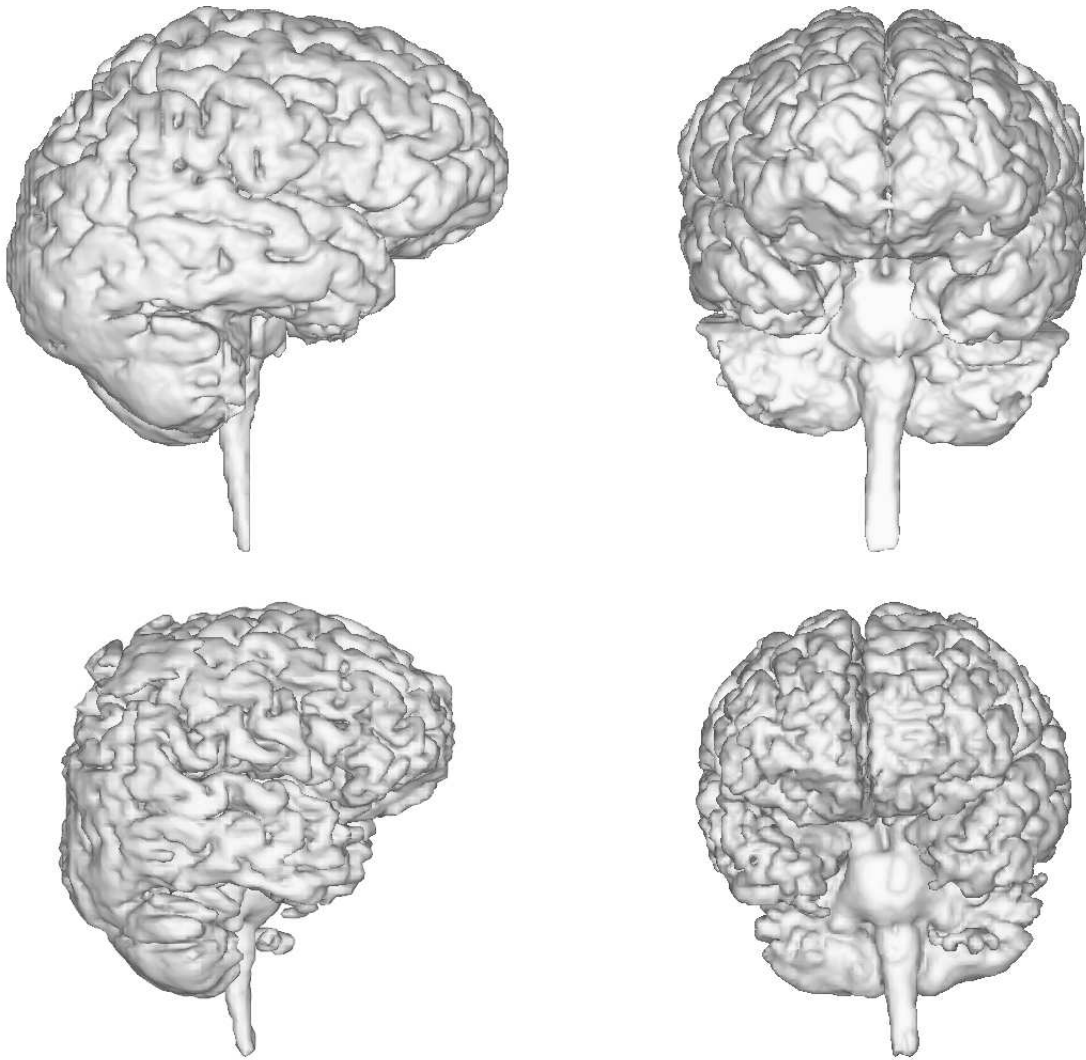


Figure 6: *The data : top, the reference brain (the one which is warped), bottom, the patient brain. Notice the differences in shapes and orientations.*



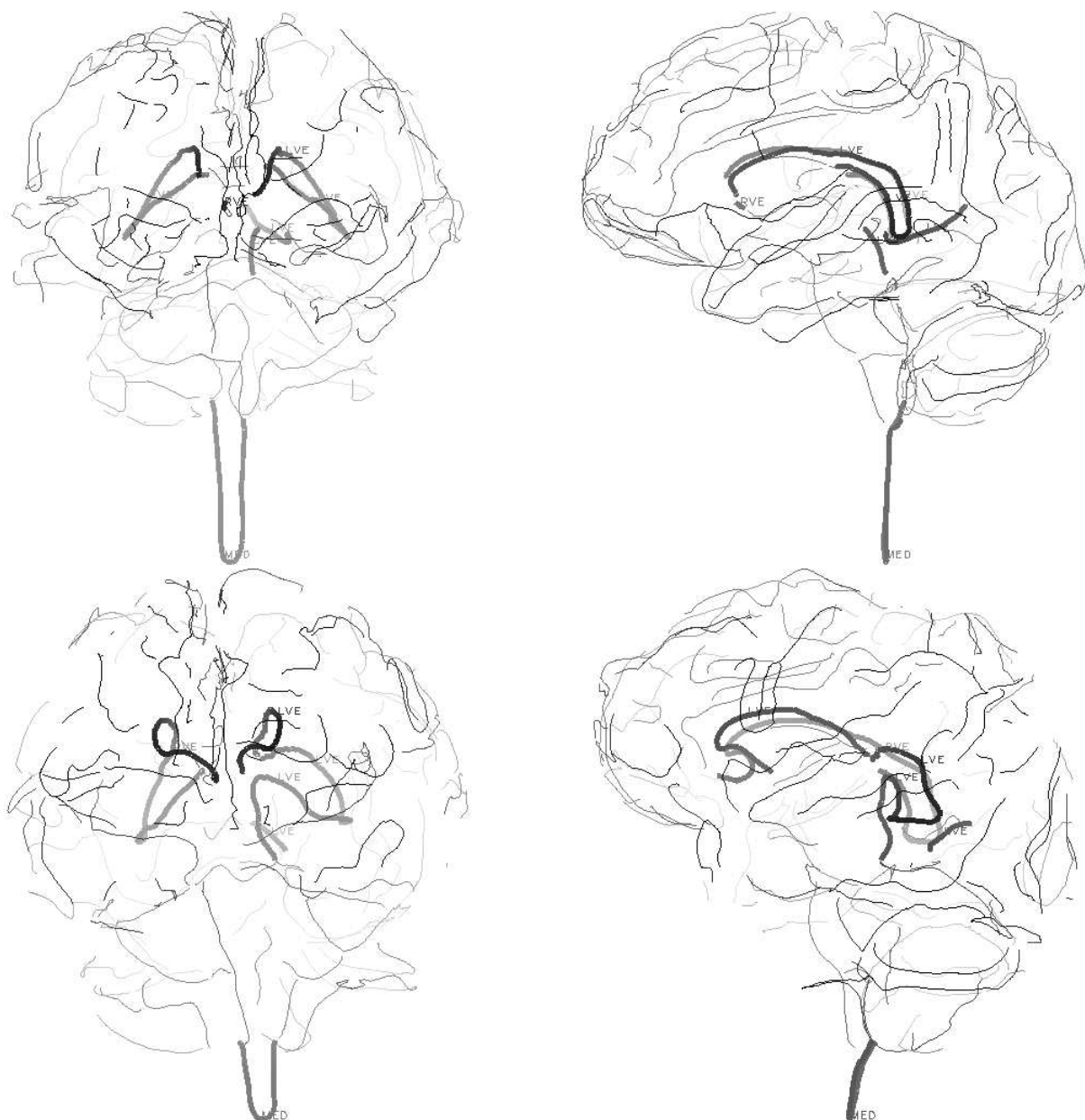


Figure 7: The crest lines automatically extracted and labeled from the reference brain (top) and from the patient brain (bottom). The thick lines are those of the ventricle and of the medulla of each brain.

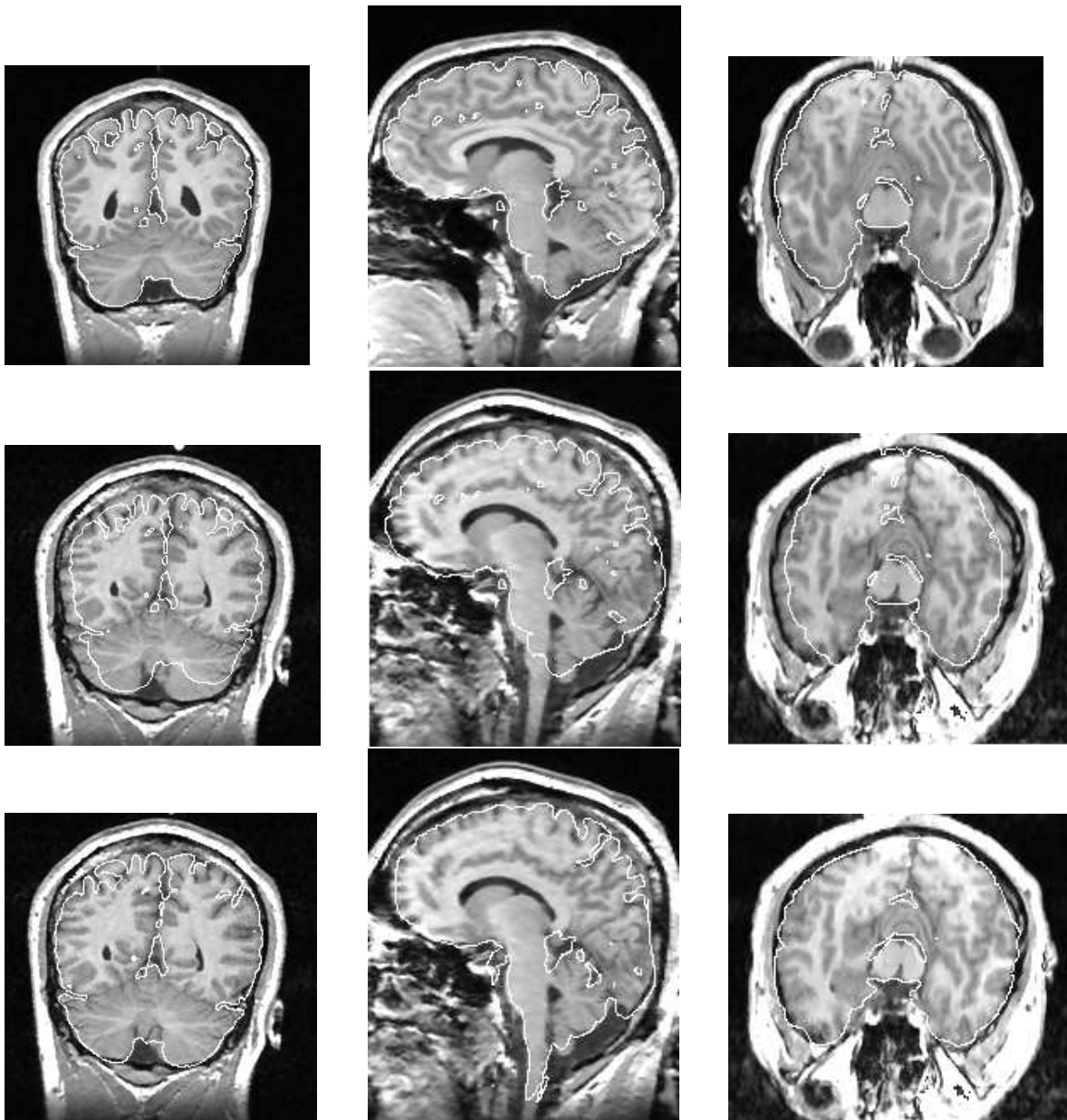


Figure 8: *Top line, the reference image  $I_r$  with the cortical surface  $S_r$ . Middle line, the patient image  $I_p$  with  $S_r$  before deformation. Bottom line,  $I_p$  with the result  $S_p$  of the found deformation applied on  $S_r$ .*

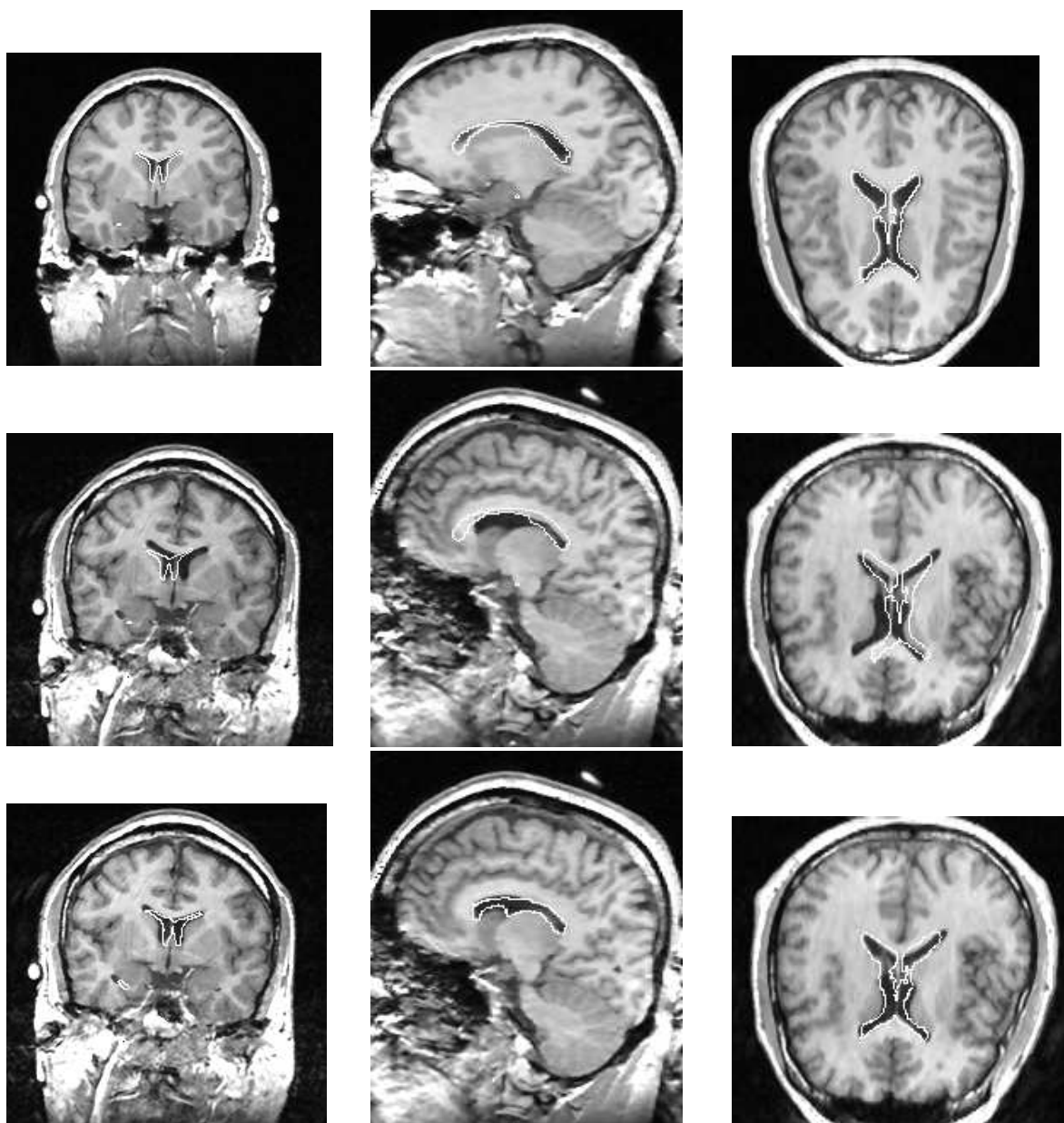


Figure 9: *Top line, the reference image  $I_r$  with the ventricle surface  $S_r$ . Middle line, the patient image  $I_p$  with  $S_r$  before deformation. Bottom line,  $I_p$  with the result  $S_p$  of the found deformation applied on  $S_r$ .*



Figure 10: *Different 3D views of the brains with their ventricle: left, the reference image, right, after warping.*

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