ARTÍCULO DE INVESTIGACIÓN Vol. 36, No. 2, May-Ago 2015, pp. 121-129

dx.doi.org/10.17488/RMIB.36.2.4pdf

REVISTA MEXICANA DE INGENIERÍA BIOMÉDICA

Automatic Segmentation of the Cerebellum in Ultrasound Volumes of the Fetal Brain

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ABSTRACT

The size of the cerebellum in ultrasound volumes of the fetal brain has shown a high correlation with gestational age, which makes it a valuable feature to detect fetal growth restrictions. Manual annotation of the 3D surface of the cerebellum in an ultrasound volume is a time consuming task, which needs to be performed by a highly trained expert. In order to assist the experts in the evaluation of cerebellar dimensions, we developed an automatic scheme for the segmentation of the 3D surface of the cerebellum in ultrasound volumes, using a spherical harmonics model. In this work we present our validation results on 10 ultrasound volumes in which we have obtained an adequate accuracy in the segmentation of the cerebellum (mean Dice coefficient of 0.689). The method reported shows potential to effectively assist the experts in the assessment of fetal growth in ultrasound volumes.

Keywords: 3D fetal ultrasound segmentation, fetal cerebellum, statistical shape models, spherical harmonics

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Fecha de recepción: 15 de enero de 2015

Fecha de aceptación: 20 de abril de 2015

RESUMEN

El tamaño del cerebelo, en un volumen de ultrasonido del cerebro fetal, ha mostrado una alta correlación con la edad gestacional, lo que hace importante a esta medición para la detección de restricciones del crecimiento del feto. La anotación manual de la superficie 3D del cerebelo en un volumen de ultrasonido es una tarea demandante, que debe ser realizada por un experto. Con el propósito de apoyar a los expertos en la evaluación de las dimensiones del cerebelo fetal, hemos desarrollado un método automático para la segmentación de la superficie 3D del cerebelo en volúmenes de ultrasonido, utilizando un modelo de harmónicos esféricos (spherical harmonics). En este trabajo presentamos los resultados de una evaluación del método automático en 10 volúmenes de ultrasonido con los que hemos obtenido un valor adecuado de exactitud (coeficiente promedio de Dice de 0.689). El método reportado tiene potencial para asistir de manera efectiva a los expertos en la evaluación del crecimiento fetal, utilizando volúmenes de ultrasonido.

Palabras clave: segmentación 3D de ultrasonido fetal, cerebelo fetal, modelos estadísticos de formas, harmónicos esféricos.

INTRODUCTION

Cerebellar volumes obtained by multiplanar, and the Virtual Organ Computer Aided Analysis (VOCAL) techniques have shown a high correlation with gestational age and might have statistically significant differences between different ethnic groups Cerebellar volumes evaluated by |1|.multiplanar and VOCAL techniques have shown differences between growth restricted fetuses and also adequacy for the assessment of the gestational age [2]. In [3],and [4] is evaluated the growth of the cerebellum in normal pregnancy, using the multiplanar technique. Both techniques (VOCAL and multiplanar evaluation) are useful for fetal evaluation of brain structures, however both have the necessity of manual delineation by an experienced operator, which is considered a disadvantage respect to automated segmentation techniques due to the low reproducibility and agreement obtained by manual segmentation techniques. In order to ease the task for the experts and, more importantly, to contribute to improved reproducibility in the measurement of the cerebellar volume, we have developed two different schemes for automatic segmentation of the cerebellum in ultrasound volumes.

Previous work on the automatic segmentation of fetal brain structures has been reported in Yaqub et al., [12] were a random decision forest classifier was trained to automatically segment: the choroid plexus, the lateral posterior ventricle cavity, the cavum septum pellucidi, and the cerebellum in 3D ultrasound. A mean Dice coefficient of 0.63 was reported for the segmentation of the cerebellum.

In Gutierrez et al., [5] was reported the development of a point distribution model (PDM) of the cerebellum which is automatically adjusted to an ultrasound volume. The model is able to automatically segment the 3D surface of the cerebellum with good accuracy (mean Dice coefficient of 0.8) with a mean processing time of 50 s. Although good results have been achieved with the PDM, in this work we explore the use of a spherical harmonic (SPHARM) model of the cerebellum for automatic segmentation of the structure in ultrasound volumes. SPHARMs have methodological advantages for the construction of statistical deformable models of 3D shapes: There is no need

for extensive annotation of landmark points; accurate shape modeling is achieved through principal components analysis of a set of normalized 3D shape examples. In the rest of this work is reported the development of our SPHARM of the cerebellum and its use for the automatic segmentation of the structure in ultrasound volumes. In section 2 we report the construction of the SPHARM of the cerebellum trained from a set of 9 annotated ultrasound volumes. In section 3 we describe the method used to automatically adjust the SPHARM model to an ultrasound volume. In section 4 are reported the results from automatic adjustment of the SPHARM of the cerebellum to a set of 10 different nontraining ultrasound volumes, using the leaveone-out method. In section 5 we present the discussion and conclusions of the work reported.

SPHARM MODEL OF THE CEREBELLUM

Our spherical harmonics (SPHARM) model of the cerebellum was trained with 9 cerebellum shapes annotated by an expert in the corresponding ultrasound volumes. Each shape was discretized as shown in Fig. 1.

The corresponding triangular mesh was generated for each shape using marching cubes [6]. An example of a mesh is shown in Fig.2.

In order to build the SPHARM model, each mesh of the training set is modeled with spherical harmonic functions as follows [7]:



Figure 1. Discretized cerebellum example.



Figure 2. Triangular mesh of a cerebellum example.

Spherical harmonic functions of order mand degree $l: Y_l^m, -l \le m \le l$ are defined in $\theta \in [0, \pi], \varphi \in [0, 2\pi)$ as:

$$Y_l^m(\theta,\varphi) = \sqrt{\frac{2l+1}{4\pi} \frac{(l-m)!}{(l+m)!}} P_l^m(\cos(\theta)) e^{im\varphi}$$
(1)

where: P_l^m is the associated Legendre polynomial:

$$P_l^m(w) = \frac{(-1)^m}{2^l l!} (1 - w^2)^{\frac{m}{2}} \frac{d^{m+1}}{dw^{m+1}} (w^2 - 1)^l$$
(2)

where: w is $\cos(\theta)$.

To define a surface three SPHARM functions are used, one for each coordinate of the points in the surface:

$$v(\theta,\varphi) = (x(\theta,\varphi), y(\theta,\varphi), z(\theta,\varphi))^T \qquad (3)$$

The surface is then described by the vectorvalued function:

$$v(\theta,\varphi) = \sum_{l=0}^{L_{\text{max}}} \sum_{m=-l}^{l} c_l^m Y_l^m(\theta,\varphi) \qquad (4)$$

where the coefficients are 3D vectors corresponding to the 3 coordinate functions in (3):

$$c_l^m = (c_l x^m, c_l y^m, c_l z^m)^T$$
(5)

The coefficients are obtained using a least squares adjustment to a given shape mesh, as described in [7].



Figure 3. SPHARM of the cerebellum for an increasing degree (L_{max}) : A) $L_{\text{max}} = 1$; B) $L_{\text{max}} = 7$; C) $L_{\text{max}} = 15$.

Using its decomposition in spherical harmonics we can model the 3D surface corresponding to each cerebellum volume in the training set. In Fig. 3 is illustrated an SPHARM model of one cerebellum, we can observe how shape details increase with the degree (L_{max}) of the SPHARM.

Once we have a set of SPHARM coefficients for each shape of the cerebellum training set, a principal component analysis (PCA) of the shape parameters is performed through registration of all SPHARM models, using a few corresponding landmark points on each shape [7], for the cerebellum we used 6 landmark points. The Iterative Closest Point method [11] is used to register the shapes in Euclidean space, followed by registration in parameter space through minimization of the Root Mean Square Distance (RMSD) [7]:

$$RMSD = \sqrt{\frac{1}{4}\pi \sum_{L=0}^{L_{\text{max}}} \sum_{m=-1}^{l} \|c1 - c2\|^2} \quad (6)$$

where: c1 and c2 are two SPHARM parameterizations to be registered.

Principal Component Analysis is performed on the parameters $(c_i, \text{ eq. 5})$ of all the examples in the registered training set, to calculate the mean shape and the main modes of shape variation. From which it is possible to generate new shapes of the class of the training set (i.e. cerebellums) using Eq. 7.

$$S_{\alpha} = S_{\mu} + \sum_{i=1}^{M} \alpha_i S_i \tag{7}$$

where: S_{μ} is the mean shape.

 S_i , are the principal component vectors

of the training set and α_i are the shape adjustment parameters, with each α_i within the range:

$$3\sqrt{\lambda_i} < \alpha_i < 3\sqrt{\lambda_i}$$

 λ_i is the corresponding eigenvalue of each principal component S_i .

In this work we used the Matlab toolbox SPHARM-MAT [7] to construct the SPHARM model of the cerebellum. The

toolbox generates the parameterizations of each shape on a training set, performs the registration and calculates the mean shape and the main modes of variation (principal components of the covariance matrix). In Fig. 4 is shown our SPHARM model of the cerebellum.

Our SPHARM model was used to automatically segment the surface of the cerebellum in fetal ultrasound volumes, as described in the following section.



Figure 4. SPHARM based deformable model of the cerebellum. Shape variations are shown for the 3 main modes within the range $\pm 3\sqrt{\lambda_i}$.

AUTOMATIC ADJUSTMENT OF THE SPHARM MODEL TO THE CEREBELLUM IN AN ULTRASOUND VOLUME

We used the method reported by Ahmadi et al [8] for the automatic adjustment of our SPHARM model to the fetal cerebellum in an ultrasound volume. The method is based on the optimization of the following local (gray level) energy function, through gradient descent.

$$E(S) = \int_{int S} f_i dx + \int_{ext S} fedx \qquad (8)$$

where:

$$f_i = (I - c_i(x)^2), where \tag{9}$$

$$c_{i}(x) = \frac{\int_{int \ S} B_{\epsilon}(x)I(x)dx}{\int_{int \ S} B_{\epsilon}(x)dx}$$

$$f_{e} = (I - c_{e}(x)^{2}), where$$

$$c_{e}(x) = \frac{\int_{ext \ S} B_{\epsilon}(x)I(x)dx}{\int_{ext \ S} B_{\epsilon}(x)dx}$$
(10)

 B_{ϵ} is an sphere of radius ϵ used to calculate the values c_i and c_e for each position x of I. The function E(S) is minimum at the surface which separates two homogeneous regions with significantly different mean grey levels. The gradient of E(S) is given by Eq. 11.

$$\frac{\partial}{\partial \alpha_j} E(S_\alpha) = \frac{\partial E}{\partial S} \frac{\partial S}{\partial \alpha_j} = \int_S (f_i - f_e) N \cdot S_j ds$$
(11)

where: N is surface normal and S_j the vertexwise cartesian deformations for the *j*-th shape parameter.

Together, these partial derivatives yield the gradient of E with respect to the shape vector α , which we denote by $\nabla_{\alpha}E$. The corresponding discrete equation is:

$$\int_{S} (f_i - f_e) N S_j ds \approx \sum_{k=1}^{N} [f_i - f_e]_k [N]_k \cdot [S_j]_k$$
(12)

where: k denotes the vertex number and $[\cdot]_k$

denotes the evaluation at vertex k.

Eq. 12 is evaluated for all the vertices of the mesh to find optimum values of the shape parameters (α_i) - which accurately fit the cerebellum in an ultrasound volume - through gradient descent.

TESTS AND RESULTS

We used 10 different ultrasound volumes acquired in an axial plane using a Voluson 730 Expert from General Electric, with a 4-8 MHz 3D probe. All volumes were acquired with informed consent of the patients at the National Institute of Perinatology in México City. The cerebellum was annotated on 10 corresponding volumes by an expert sonographer.

The automatic segmentation of the cerebellum was evaluated on the training set using the leave-one-out method. Nine volumes were used for training the SPHARM of the cerebellum with validation of automatic segmentation on the volume left out. This was repeated 10 times, validating the automatic segmentation on each of the 10 fetal ultrasound volumes. The Dice Similarity Coefficient (DSC) [9] was used to measure the accuracy of the automatic SPHARM segmentation as compared against manual expert annotations.

$$DSC = \frac{2TP}{(2TP + FP + FN)} \qquad (13)$$

where: TP corresponds to the number of true cerebellum voxels,

FP corresponds to the number of voxels wrongly included as cerebellum and

FN corresponds to the number of cerebellum voxels wrongly left out of the segmentation.

For each automatic segmentation test, the SPHARM was initialized with the mean cerebellum shape and size, located within the central plane of the cerebellum in an ultrasound volume, as shown in Fig. 5. The optimization parameters were: step size



Figure 5. SPHARM vertices initialization.





Figure 6. Automatic segmentation results, shown on the SRAD filtered images: Expert annotations are shown in red; automatic annotations are shown in yellow.

of 0.05 for gradient descent, sphere radius of 5 voxels and 30 iterations. Speckle reduction anisotropic diffusion filtering (SRAD) was applied to all ultrasound volumes [10] with 20 iterations and a diffusion coefficient of 0.75. In table 1 are shown the DSC coefficients for our 10 validation ultrasound volumes. In Fig. 6 are shown the results of automatic

Test volume	DSC
1	0.7258
2	0.7186
3	0.7022
4	0.6405
5	0.6942
6	0.7043
7	0.6966
8	0.6552
9	0.6793
10	0.6742
Mean	0.6890
Standard deviation	0.0269





Figure 7. 3D Automatic segmentation result.

segmentation on three crossections of an ultrasound volume. In Fig. 7 is shown an example of the automatic 3D segmentation of the cerebellum.

DISCUSSION AND CONCLUSIONS

From table 1 it can be observed that our method is able to automatically segment the cerebellum in an ultrasound volume with adequate accuracy (mean DSC of 0.689). DSC values starting from 0.7 are considered in good agreement with the corresponding expert annotations. In Yaqub et al., [12] a mean DSC of 0.63 was reported for the segmentation of the cerebellum. This, lower, segmentation accuracy is most likely due to the lack of shape models [12].

However higher DSC values have been previously obtained with an automatically adjusted point distribution model (PDM) of the cerebellum, as reported in [5] where 20 ultrasound volumes were used for training and validation using leave-one-out, with a mean DSC for automatic segmentation of 0.8. The gains in accuracy are likely to be due to an improved objective function constructed with grey level voxel profiles, normal to the deformable model surface. In the following stage of development we will explore the use of this objective function with an SPHARM model for automatic segmentation of the cerebellum.

ACKNOWLEDGEMENTS

The authors are grateful to the Mexican National Science and Technology Council (CONACYT) and to the former Institute of Science and Technology of the Federal District in México City, for their financial support to the Biomedical Imaging Lab. at CCADET, UNAM. Gustavo Velásquez is grateful to CONACYT, for the financial support in the form of a PhD scholarship.

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