ACCURATE AND FAST 3D INTERACTIVE SEGMENTATION SYSTEM APPLIED TO MR BRAIN QUANTIFICATION

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Abstract

This work presents an efficient interactive segmentation system for volumetric data-sets based on advanced 3D morphological analyses and an interaction paradigm that allows a good match with user intentions. This system has been designed to produce accurate results under the complete control of the user, to minimize the interaction time and to address a generality of 3D segmentation tasks. The system has been tested and compared with other softwares on normal MR brain structure quantification and on a challenging clinical setting pointed to the detection of the presence of subtle brain atrophy associated to primitive immunodeficiency (PID).

1 Introduction

Due to the hard separation between high-level human recognition power and machines' computational power, interactive (user-guided) approaches are adopted in a predominant part of medical image segmentation techniques. Interactive segmentation systems (ISS) have been classified regarding their computational [1,2] rather than interactive part [3]. Different parts are designed (and/or selected) and put together in order to make the overall segmentation technique more or less suited and usable for certain applications [4]. Here we focalise on segmentation of volumetric anatomical data-sets and present the design and realization of a technique which implements a successful combination of visual interaction and non trivial morphological analysis tools. In our view, interactive segmentation of medical data-sets requires used technologies to be well matched with user intentional and semantic message exchange. centred Morphological techniques can be seen as well suited for such a cognitive matching in that they inherently deal with some image attributes that reflect significant image coherency features. Moreover 3D extension of morphological tools does not involve particular complexity issues (contrarily to what usually emerge in extending other kind of segmentation technologies, e.g. deformable models or level set methods). implemented system presents good accuracy, The repeatability and efficiency capabilities, which are the evaluation features highlighted for ISSs by Olabarriaga and Smeulders [3]. Furthermore we aim to guarantee an additional feature, namely generality, which has been used as a design criterion and constitutes a strength point of our system. Generality can be defined as the capability to deal with all problems of a certain, suitably wide, class (characterized by a an a priori knowledge about the segmentation problem nature) despite the inherent degree of unpredictability (due to anatomic and pathological variety) and discrimination difficulties (due to the physical signal and noise limitations on the image information content). Generality is a valuable feature because techniques that are conceived to deal with general tasks are usually well adapted to be specialized for dedicated tasks (specialization can be modelled in terms of additional a-priori knowledge). The opposite is not generally true.

1.1 Interactive segmentation model

In Fig.1 we present our reference ISS user-centred model. The physician is represented with the smiling face and its actions with dashed lines. Segmentation and refinement interaction tools (SIT and RIT) are conceived as elements of segmentation and refinement toolboxes respectively. SIT and RIT should be designed, presented and selected according to data and working environment requirements and to the user skills. They should be also complete in representing different tasks and robust in providing different ways to communicate the same information. Relevance and adequacy of the ITs stem from the possibility of being translated in a set of quantitative parameters representing what the physician would like to communicate (which could be different from one tool to another). These parameters are estimated and calculated in the PEC blocks and conveyed as inputs to the segmentation and refinement computational tools (SCT, RCT) which should be conceived to be capable to fully exploit such information. A preprocessing analysis, PA computational block also appears into the scheme. This block is separated from the SCT because of its peculiar role into the segmentation pipeline. PA is in general allowed to receive some PEC parameters as well. SIT and RIT govern the data rendering engines (Ro and Rs).



Figure 1 Interactive Segmentation System model

2 Segmentation Interaction Tools

Interactive tools and their usage must be easy and selfexplanatory with respect to the task to accomplish and the user culture. Different tools could differ by specific vocation but they should have usage intersections so that the physician could normally obtain desired results by choosing alternative tools. In our solution a set of visual interaction tools $S_k \in SIT$, k=1,...,4 have been designed. The S_k are presented to the user by an expressive icon. Each SIT is associated to a dedicated PEC translator. Not necessarily the set of computational parameters that can be derived is the same for all S_k , and the computational part must be aware of this fact. In the proposed ISS, the user selects one (or more) volume slices (along a certain slicing plane) and a SIT among the following ones (see Fig. 2):

- S_1 Elementary shape (ellipse/rectangle) with markers: the physician draws an elliptical (rectangular) shape which encompasses the structure of interest (SoI). Markers are then positioned to further specify the SoI.

- S_2 Free-hand closed curve with markers: similar to S_1 , but it is expected that the line corresponds to an accurate contouring of the SoI in the selected slice.

- S_3 Line with crossing line: a first line is traced on the SoI and a second one is traced to cross the first one while passing through nearby structures.

- S_4 Multiple seeds: a freely chosen number of point markers are placed on SoI.

SIT vocations are summarized as follows: S_1 can be used for objects of various dimensions that can be easily identified and approximately encompassed by an ellipse or a rectangle (e.g. lesions, tumours), S_2 requires more accurate interaction which is justified when object shape is more involved and/or the object is not so easily detectable (e.g. inner organs), S_3 is mainly indicated for thin structures on image slices (e.g. vessels, grey matter) or to indicate a representative portion of extended structures (e.g. white matter), S_4 can be used when desired objects have nearly uniform intensity values (e.g. bones, CSF) or to indicate several points belonging to an extended and complex structure (e.g. grey matter). From each S_k interaction the corresponding data portion is collected and a local intensity histogram lH(i) is estimated. Direct lH(i)computation is possible with S_2 and S_3 , while an estimation is performed with S_4 and S_1 by using sampled intensity information and main histogram moment selection (associated to local peaks) respectively. From lH(i), SoI maximum and minimum intensity (I_{min}, I_{max}) can be estimated. SoI markers are directly acquired with S_1 , S_2 and S_4 while they can be derived from the first traced line with S_3 . Another meaningful information to be captured from the interaction can be expressed in terms of partitioning strength (PS) and is derived by a localized SoI analysis. The smoother the interesting structure boundaries the weaker (conservative) the *PS* should be. *PS* are directly estimated when S_2 and S_3 are selected, this is done with a signal intensity profile analysis across the boundaries of S_2 or along the second crossing line for S_3 . When using S_1 or $S_4 PS$ estimation is not so direct but it can be performed as well using suitable heuristics or set to a default value and left to be tuned by the user.



Figure 2 Segmentation Interaction Tools

3 Computational Tools

As already stated, SIT and SCT should be coherent in sharing common segmentation abstractions. Accordingly, non trivial mathematical morphology tools has been designed and implemented in our ISS.

3D connected operators analysis preprocessing It is known that a good denoising is beneficial because it reduces oversegmentation problems. However, denoising should not alter the information content and in particular the contour definition and other relevant image features. To this ends a 3D morphological filter-by-analysis family of operators, conceived as a volumetric extension of 2D connected operator filtering [5], has been realized [6]. With these operators a simplification of the original volume can be performed by discarding 3D connected components with respect to different distinguishing criterions (e.g. volume, surface, complexity, etc). 3D connected operators denoising is a suitable PA tool in our ISS in that it guarantees good denoising performance and works definitely better compared to similar 2D operators applied on single slices. In fact, while in 2D slices anatomic structures could be confused with noise in their peripheral or highly textured parts, in the 3D case even a thin and intricate structure will survive to the filter by analysis step because it is fully explored in its volumetric extension [5]. On the contrary, noise conserves its granular structure and can be effectively filtered out. In this work only volume based simplification (more suited for denoising purposes) has been considered, i.e. 3D connected components that do not reach a given minimum volume (e.g. 10^3 voxels) are removed. Moreover our 3D connected operator filtering works on grey level connected flat components and preserve the continuity of the signal intensity. This makes this processing a good allied of the subsequent gradient-based water flooding analysis. 3D connected operator filtering is a quite computationally expensive task but it can be easily done offline with respect to the interactive segmentation session.

Fast 3D water flooding (pseudo-watershed) segmentation Volume segmentation is performed by a fast, interaction steered, and computationally optimized marker based 3D "water flooding" technique. This can be seen as a simplified version of a marker based watershed analysis and share the same water immersion interpretation of the geodesic version of the signal under analysis [7]. For segmentation purposes, a useful signal to work with is the volume gradient. We use the morphological gradient G(v) of the original MR volume MR(v) (where v = (x, y, z) is the voxels' coordinate set) defined by $G(v)=\max_n (abs(MR(v)-MR(\eta_v)))$, being η_v the v related neighbouring voxels according to a 6-connectivity kernel. Moreover we propose an original way to produce and exploit prior knowledge coming from the user interaction in order to improve the convergence towards the desired result and minimize the computational complexity. Parameters coming from the PEC modules are converted into SoI related bounding thresholds: (I_{min}, I_{max}) restricts voxels that are considered, seed markers determine the catchments basins from which to start the merging, PS is converted in a region merging threshold (rmt) which regulates the region merging strength by determining a flooding limit on the gradient, i.e. a level under which different basins are merged. A pictorial view of how the bounded water flooding works is given in Fig. 3 for a one-dimensional case. The described 3D water flooding technique allows quick visual feedbacks (one to few seconds using a normal PC on a 256^3 voxels volume). Finally, note that a set of parameters derived from the interaction is associated to any segmentation result. This ensure a perfect reproducibility of results, i.e. the possibility to automatically repeat the segmentation without having to archive the segmentation results as additional volumetric data-sets.



Figure 3 Marker based water flooding by immersion

Refinement tools Basic RIT and RCT such as local cuts and local erosions have been implemented. These can be used to exclude, from the segmented volume, regions that have been erroneously included due to pixel intensity contiguity. Also a local and global gradient ascending constrained dilation of the obtained structures can be performed to compensate for eventual border underestimation. RIT derived parameters constitute additional constrains for the bounded water flooding which can be repeated by comprising them.

Despite their simple nature, and thanks to the satisfactory "first shot" performance of the SCT phase, RIT and RCT usually let the user drive the segmented data-set to the desired result with a minimal effort. This was the case of all experimental results in this work. Of course our model could include e.g. more intelligent scissors in order to make the convergence to the desired result faster by shortening the interaction. Such tools are object of current investigations [8].

5 Results on brain quantification and atrophy

Our interest now focuses on a specific clinical application which consists in a quantitative analysis of brain atrophy in a small population of patients affected by a rare disease compared to a population of healthy subjects. A recent study reported progressive neuro-degeneration in patients affected by primary immunodeficiency disease (PID) [9]. The aetiology of this syndrome is unknown. Neuroimaging findings are represented by brain atrophy with prominent sulci and enlarged ventricles. Since a reliable estimation of atrophy critically depends on the accuracy of the segmentation procedure, the applicability of the presented ISS for atrophy assessment depends on its volume quantification performance. For these reasons in order to evaluate if the proposed ISS is capable to detect brain atrophy, a preliminary system validation has been performed by comparing gray matter (GM), white matter (WM) and cerebrospinal fluid (CSF) parameters with available literature [10-12]. Thirteen patients affected by PID and 9 healthy controls selected from a population of volunteers were considered. Inclusion criteria for control group: (1) normal brain MRI as determined by an experienced neuroradiologist, (2) no history of severe head trauma requiring medical attention. Subjects were matched according to gender and age. A 3D structural MRI scan was acquired for each subject by using a T1-weighted MPRAGE sequence (1mm isotropic voxels). The proposed ISS has been used to segment WM, GM and CSF and to produce volume measurements (see Tab.1). Accurate interactive segmentation of each structure took from 2 (simpler cases) to 10 minutes (more complex cases and structures). A visual example of resulting GM and WM volumes is shown in Fig.4 (GM and WM extracted from the same subject). Our findings in compatible subject classes were compared with those obtained by Ge [10], Chard [11] and Blatter [12] in normal volunteers1 (see Tables 1 and 2). These studies reported normative brain volumetry in healthy segmented subjects. It should be noted that the three studies [10-12] used different techniques and tools: 3DVIEWNIX [13], SPM99 [14], Analyze [15] respectively. The cited segmentation tools are widely used as benchmark in medical computing literature.

Normal brain *volume quantification* Our findings concerning WM, GM and CSF volumes in normal subjects are comparable with the available data in literature [10-12]and they are coherent in terms of age and gender differences. As it can be observed, average values are intermediate between other studies while standard deviation values obtained by the proposed method are definitely the lowest. These values obviously depends on the sample, however being our sample less populated with respect to the others we feel qualified to positively interpret the obtained results. Moreover, the analysis on healthy subjects demonstrated that there is a statistically significant difference in TIV between males (1.47 L) and females (1.30 L) (T test p < 0.001), and that both TIV and BPV in healthy females are statistically lower compared to healthy males (TIV $1.30L \pm 0.02$ vs 1.47L \pm 0.02, p < 0.001; BPV 1.13 \pm 0.04 vs 1.28 \pm 0.04), confirming the anatomic evidence reported by Blatter et al. [12]. Confirmation of available results and a better standard deviation let us conclude that the proposed method is suitable for quantitative brain analysis.

Atrophy assessment for PID disease Accepting the fact that the small sample size requires cautious interpretation, the proposed ISS detected both presence and degree of regional GM, WM, and CSF changes in patients with PID. We detected a loss of brain parenchyma in patients affected by PID, which was expressed by a statistically significant reduction of fractional GM volumes (51.9 ± 2.3 vs 55.2 ± 2.3 , p=0.008) paralleled by a significant increase in CSF volumes (CSF 0.24±0.09 vs 0.18±0.04 p=0.006), without changes in TIV, thus confirming the observations of Ziegner et al. in their pilot study [9].

5 Conclusions

A new ISS has been presented where user knowledge and a properly designed interaction language are effectively exploited to guide fast and accurate morphological analysis and segmentation tools. The computational pipeline is composed by 3D connected-operators preprocessing, fast 3D water-flooding segmentation and means to quickly refine the results. Accuracy of the proposed ISS has been evaluated in comparison with other technologies and a clinical validation has been performed based on a small population of healthy subjects and patients affected by a rare neurodegenerative disease. Obtained results confirmed technical and clinical expectations and are meaningful indices of good accuracy, repeatability, efficiency and generality properties. In particular, in all considered cases we obtained rapid convergence to the desired result (i.e. without or with only few refinement steps). This demonstrates the good match between the interaction and the computational parts of the system, and distinguishes our user-guided method from many others which also use fast (often localized) operators but require, especially on complex or noisy structures, many refinement loops (usage cycles) to obtain satisfactory results.

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Figure 4 WM and GM results (3D volume rendering of 4 different views for each segmented structure).

	$\mathrm{GM}(\mathrm{mL})$	WM(mL)	$\operatorname{CSF}(\operatorname{mL})$	TIV(mL)	BPV(mL)	BPF	GM F	WM F	CSF F	GM/WN
C.F.20	731, 191	457,715	151,792	1.340,698	1.188,906	0,887	0,545	0,341	0, 113	1,597
C.M.28	762,353	528,954	211,562	1.502,869	1.291, 307	0,859	0,507	0,352	0, 141	1,441
C.F.28	765, 627	354,850	184,797	1.305, 274	1.120,477	0,858	0,587	0,272	0,142	2,158
C.M.21	777, 248	443,750	240, 261	1.461, 259	1.220,998	0,836	0,532	0,304	0,164	1,752
C.M.31	786,938	512,377	134, 144	1.433,459	1.299, 315	0,906	0,549	0,357	0,094	1,536
C.F.28	697, 541	390, 524	192,708	1.280,773	1.088,065	0,850	0,545	0,305	0,150	1,786
C.M.26	832, 982	444,630	191, 154	1.468,766	1.277, 612	0,870	0,567	0,303	0,130	1,873
C.M.22	843, 198	495,685	154, 534	1.493, 417	1.338,883	0,897	0,565	0,332	0,103	1,701
C.F.20	739,808	394,160	159,725	1.293,693	1.133,968	0,877	0,572	0,305	0, 123	1,877
P.M.28	767, 166	438,954	245, 432	1.451,552	1.206, 120	0,831	0,529	0,302	0,169	1,748
P.M.25	672,721	354,937	267, 214	1.294,872	1.027,658	0,794	0,520	0,274	0,206	1,895
P.M.37	689, 445	588,022	155, 851	1.433, 318	1.277, 467	0,891	0,481	0,410	0,109	1,172
P.M.39	700,405	469,696	325, 273	1.495,374	1.170, 101	0,782	0,468	0,314	0,218	1,491
P.M.19	650,984	405,638	266, 485	1.323, 107	1.056,622	0,799	0,492	0,307	0,201	1,605
P.M.25	762,405	481,564	323, 116	1.567,085	1.243,969	0,794	0,487	0,307	0,206	1,583
P.F.20	707,076	427,658	282,921	1.417,655	1.134,734	0,800	0,499	0,302	0,200	1,653
P.M.33	777,748	474,520	203, 649	1.455,917	1.252,268	0,860	0,534	0,326	0,140	1,639
P.M.19	833, 037	438,954	245, 432	1.517, 423	1.271,991	0,838	0,548	0,289	0,161	1,898
P.F.40	737,886	430,615	185,969	1.354,470	1.168,501	0,863	0,545	0,318	0,137	1,714
P.M.23	812, 361	460, 229	205, 613	1.478,203	1.272,590	0,861	0,550	0,311	0,139	1,765
P.M.25	798, 217	484,233	190, 356	1.472,806	1.282,450	0,871	0,542	0,329	0, 129	1,648
P.M.24	877,737	513, 171	208,927	1.559,835	1.390,908	0,869	0,548	0,320	0,130	1,710

Legend: ID interpretation =(C=control,P=patient).(M=male,F=female).(ag);total intracranial volume, TIV = WM+GM+CSF, brain parenchymal volume, BPV = WM+GM; brain parenchymal fraction, BPF = (WM+GM)/TIV; grey matter fraction, GM F = GM/TIV; white matter fraction, WM F = WM/TIV; and cerebrospinal fluid fraction, CSF F = CSF/TIV.

Table 1 Brain volumetry obtained from the proposed ISS

				20-49 y (32)		21-31 y (9) [†]			
				Mean	Std.Dev	Mean	Std.De	v	
		BP	V (mL)	1367,3	147,4	1285, 6	42,7		
		GM	(mL)	717,8	96,8	800,5	35,6		
		WM	(mL)	499,9	73,7	485,1	39,1		
		GM	F	0,525	0,041	0,544	0,025		
		WM	1 F	0,366	0,04	0,33	0,026		
		GM	/WM	1,5	0,3	1,7	0,2		
				males	males		fem		
	27.2-52		2.7 y (13	(13) 21-31 y (5) [†]		23.2-55.2 y (14)		20-28 y (4)	+
TIV	TIV (ml)		$1441, 7 \pm 128, 69$		$1472 \pm 27,49$		$\pm 73,89$	$1305, 1 \pm 25$,74
GM	GM (ml)		$\pm 65,88$	800	$800, 5 \pm 35, 55$		-44, 64	$733, 5 \pm 28,$	10
WM	WM (ml)		$414, 0 \pm 38, 95$		$485 \pm 39, 13$		23, 24	$399, 3 \pm 42,$	7
BPV	BPV (ml)		$1207, 7 \pm 96, 65$		$1285, 6 \pm 42, 71$		$\pm 60, 44$	$1132,9 \pm 42,04$	
GM	GM F		$0,55 \pm 0,018$		$0,54 \pm 0,024$		0,027	$0,56 \pm 0.026$	
WM	WM F		0.29 ± 0.018		0.33 ± 0.026		0.010	0.3 ± 0.028	
BPF	BPF		$0,84 \pm 0,024$		$0,84 \pm 0,029$		0,029	$0,87 \pm 0,017$	
			ma	ales				females	
	16-25 y	(24)	26-35 y (19)	21-31 y (5) †	16-25 y	(20)	26-35 y (24)	20-28 y (4) †
TIV (mL)	V (mL) 1594,1±		101,39 1549,43±8		$81,91$ $1472 \pm 27,49$		\pm 93, 09	$1329,67 \pm 112,37$	$1305, 1 \pm 25, 74$
GM (mL)	M (mL) 761,84 ±		106,99 749,09 ±		$800, 5 \pm 35, 55$	$694, 78 \pm 112, 71$		$647, 54 \pm 98, 61$	$733, 5 \pm 28, 10$
WM (mL)	M (mL) 746,45 ±		:89,25 687,56±		$485 \pm 39,13$ 179 98 ± 10 8	010 ± 109		580, 20 ± 79, 34 05 88 ± 99, 79	$399, 3 \pm 42, 7$
BPV (mL)	1508.29	±106.00 1436.66		-70.85 $1/2, 25 \pm 19, 5$ -70.85 1285.6 ± 42.7		$69, 50 \pm 30, 65$ 1310 99 \pm 79 64		90,00 ± 32,73 1233 70 ± 102 43	$130, 35 \pm 25, 7$ $1132, 9 \pm 42, 04$
BPF	PF $1,20 \pm 0$		0,42 1,27 ± 0,		31 0,84±0,029		, 53	$1,36 \pm 0,56$ $0,87 \pm 0,017$	

Table 2 Proposed ISS † vs 3D-Viewnix (top) SPM99 (mid) Analyze (bottom). Number of subjects belonging to the age-group in brackets.