NEPHROBLASTOMA ANALYSIS IN MRI IMAGES

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ABSTRACT

The annotation of the tumour from medical scans is a crucial step in nephroblastoma treatment. Therefore, an accurate and reliable segmentation method is needed to facilitate the evaluation and the treatments of the tumour. The proposed method serves this purpose by performing the segmentation of nephroblastoma in MRI scans. The segmentation is performed by adapting and a 2D free hand drawing tool to select a region of interest in the scan slices. Results from 24 patients show a mean root-mean-square error of 0.0481 ± 0.0309 , an average Dice coefficient of 0.9060 ± 0.0549 and an average accuracy of $99.59\% \pm 0.0039$. Thus the proposed method demonstrated an effective agreement with manual annotations.

Keywords: Continuous Max-Flow, Graph Segmentation, Kernel Induced Space, MRI images, Nephroblastoma, Wilms tumour.

INTRODUCTION

Nephroblastoma or Wilms' tumour is a form of kidney cancer that affects around 80-85 children in the UK each year (Macmillan Cancer Support, 2016). Nephroblastoma generally affects children below the age of five and it may begin to develop in the womb when the baby is still unborn. The most common symptom is a painless swelling in the abdomen. Nephroblastoma rarely affects adults. In most cases it affects only one kidney (unilateral), but it can also affect both (bilateral). Nephroblastoma is a fast growing tumour with a median tumour size of around 400 ml at the time of diagnosis. Fortunately due to prospective clinical trials and effective treatments more than 90% of patients can be cured today. Imaging analysis tools play an important role in diagnosis and treatment planning. The assessment of the tumour during preoperative chemotherapy is of prognostic relevance and is crucial for postoperative treatment stratification (David et al., 2012).

Over the past years, many segmentation methods of different types of tumours such as glioblastoma, prostate and Wilms) have been proposed (Bauer *et al.*, 2013a; Gordillo *et al.*, 2013; Pham *et al.*, 2000). These methods include full automated and semi-automated methods and they are applied to interpret scans such as MRI, CT or ultrasound. Each technique has its own advantages and disadvantages. Many of these methods fail due to different acquisition conditions of resolution, illumination and field of view, and the variability of body shapes and positions.

Overlapping tissue and organs and the irregular size of the tumours can also cause a significant degradation to the performance of the segmentation methods, particularly the fully-automated methods. For the above reasons, there is a need for reliable and robust semi-automated segmentation methods for Wilms' tumour studies. Semi-automated tumour segmentation techniques are computational methods that perform an extraction of the tumour tissue from multi-sequence Magnetic Resonance Imaging (MRI) with the interaction of human experts. These methods can provide important time-saving for neuroradiologists in tumour assessment, and their accuracy can enhance tumour characterisation for radiotherapy, surgical planning and drug development assessment. The aim of this document is to report findings during the CHIC project for semi-automated Nephroblastoma tumour segmentation in multi-sequence MRI imaging. In the following section, we briefly review some previously-proposed tumour segmentation methods.

RELATED WORKS

Image segmentation aims to partition an image into sub-regions. These regions can be classified according to the functional areas, tissue types, structures of interest etc. (Bauer *et al.*, 2013a).

Advance innovation in hardware and software is leading to a continuing increase in computing power. This has made supervised classification techniques an attractive research area (Menze et al., 2015) in the medical image analysis. The performance of these methods relies heavily on a large set of training data, which include manually annotated images for learning features in the medical scans. Neural Networks (Goceri and Goceri, 2015; Ramrez et al., 2018; Zhong et al., 2018), Support Vectors Machines (Lee et al., 2008; Amiri et al., 2016) and Random Forests (Shah et al., 2017; Bauer et al., 2012; Conze et al., 2016) are amount the popular methods used for medical images segmentation. Unlike these supervised methods, unsupervised segmentation techniques are very flexible as they can be applied on a small set of data from which they can learn classification rules based some similarity criterion.

The segmentation of Wilms tumour is not a trivial task due the complexity, inhomogeneity and variability of its structures Farmaki *et al.* (2010) and may require inputs such as seed-points, location and shapes from human experts. For this reason, a fully automated segmentation method may not be a good option for clinical use (Farmaki *et al.*, 2010; David *et al.*, 2012). Thus, a semi-automatic tumour segmentation method can be a good alternative option for Wilms' tumour analysis in multi-sequence MRI scans.

The semi-automated tumour segmentation methods can be divided into three major groups (Farmaki et al., 2010): region-based approaches, deformable model-based approaches and graph-based approaches. The region-based approaches perform the tumour segmentation using a single seed point and expand the pixel extraction from that seed point to fill a defined coherent region according to a similarity measure of neighbouring pixels within the image region (Adams and Bischof, 1994). The region-based approaches include region-growing and watershed methods. A typical region-growing approach was proposed in (Kim and Park, 2004) by Kim and Park. This combined the region-growing algorithm with a texture-based analysis operation, which was performed on sample images of the tumour to define a seed region as a starting point of the regiongrowing algorithm. Mancas (Mancas and Gosselin, 2003) proposed a semi-automatic segmentation on head and neck tumours in CT, MRI and PET scans. The approach used a marker-based watershed technique, by incorporating a fuzzy region with a tumour probability from 0 to 1. The watershed algorithm was used iteratively. This combination improved the watershed algorithm and avoided the over-segmentation problem. In (Sauwen et al., 2016),

Nicolas et al., presented a semi-automated framework for segmenting brain tumour from multi-sequence MRI scans using regularisation non-negative matrix factorisation. The technique makes use of the tumour regions seeds selected by the user to initialise the algorithm. A morphological post-processing technique is used to improve the segmentation results, which combined the special location voxels with some adjacency constraints defined in the process.

methods The region-based have specific advantages over other segmentation approaches: they are easy to implement and produce coherent regions. However they often perform poorly because they cannot find objects that span several disconnected areas, and decisions regarding region membership can be difficult. Methods presented in (Kim and Park, 2004), (Mancas and Gosselin, 2003), and (Sauwen et al., 2016), relied heavily on the selection of the seeds in the pathology region (i.e., tumours voxels) as a starting point to initialise the algorithm. The intensity variation in the tumour region can led to a misclassification of tumour pixels. Our method addresses this issue by interactively selecting a wider region of the tumour area which include different variation of the tumours voxels intensity.

The deformable model-based approaches such as active contours (snakes), level sets (LS), or geodesic active contours have been extensively used in medical image segmentation applications. They make use of regional properties or edge detection in the image to extract the tumour regions (Bauer et al., 2013b; McInerney and Terzopoulos, 1996). Methods such as level-set evolve toward the tumour region by searching in the image the largest gradient or by using region characteristics in the image. Wang et al. (Wang et al., 2009) proposed a fluid vector flow algorithm to evolve a contour toward the boundary of the tumour in T1-weighted images. In (Linguraru et al., 2009), Linguraru presented a semi-automated renal tumour quantification and classification method in a 3D size volume. The model combined a fast-marching operation and geodesic level-sets to define the shape of the lesions. In (Sachdeva et al., 2012), Sachdeva used a content-based active contour (CBAC) texture and intensity information to evolve an active contour toward the tumour boundary edge in MRI scans. Gu et al. (Gu et al., 2006) proposed a multistage method for a 3D segmentation of CT and MRI images and a new radial distance-based segmentation validation approach. Gu's algorithm is based on level sets and it incorporates an improved fast marching method and a morphological reconstruction model. In (Somaskandan and Mahesan, 2012), Suthakar and Sinnathamby proposed a tumour segmentation technique based on level set deformable algorithm. The method uses both the images gradient information and the reginal data information to deform the level set algorithm. They estimated the velocity forces based the tumour voxels statistical measure and surrounding health tissues information. The above deformable segmentation methods can perform well on medical images. However, their implementations can be very complex and costly. Any sharp intensity variation, artefacts or noises in the medical images can led to misclassification. Our method avoids these issues using a bias correction operation as a pre-processing step to remove all the imaging artefacts, noise and corrected the intensity inhomogeneity in the scans. In addition, the bias correction, we applied an energy function which includes both boundary and regional terms allowing the segmentation of heterogeneous objects, which not possible with deformable segmentation methods due to local minima. Furthermore, deformable model-based methods (Kainmüller et al., 2007; Liu et al., 2010; Zhan and Shen, 2006) use prior knowledge of the scan (i.e. intensities, shapes, colour, texture, positions etc.) to apply constraints on the segmentation algorithms. These constraints can lead to segmentation errors when the algorithms are used on different image modalities (Kaba et al., 2015). Our segmentation method addresses these issues by using a prior knowledge including intensities variation of the tumour, which is manually selected by the user.

The graph-based approach is one of the most attractive segmentation methods in Computer-aided diagnosis (CAD). Like the deformable model-based approaches, the graph-based image segmentations are also based on energy optimisation. They combine boundary regularisation with regularisation of regional proprieties in the same approach as Mumford-Shah (Boykov and Funka-Lea, 2006). The segmentation is performed by dividing the image into subclasses known as Foreground (targeted tumour) and Background. The segmentation process is performed using a graph, which consists of a set of nodes (i.e. pixels) and a set of edges (i.e. weights on the nodes) that connect the nodes using their degrees of similarity (intensity) (Kaba *et al.*, 2015).

In (Freedman and Zhang, 2005), Daniel Freedman and Tao Zhang presented an interactive graph cut algorithm, which incorporates shape priors into the energy function. The segmentation is performed by the shapes template and integrating the shape information in the weights on the edges in the graph. Xinjian Chen et al., (Chen *et al.*, 2012) proposed a segmentation of method for abdominal 3-D organ. The segmentation is performed by combining the active appearance model, live wire, and graph cuts. The overall method included model building, object recognition, and delineation. In (van der Lijn et al., 2008) Fedde van der Lijn et al., presented a graph cut segmentation algorithm for the extraction of the hippocampus from MRI scan. The graph energy functional contains the intensity energy and the prior energy which included a spatial and a regularity term. A 3D semi-automated graph cut-based segmentation algorithm of liver cancer is proposed in (Esneault et al., 2007). The liver cancer in a contrast-enhanced 3D CT volume was segmented with a user interaction in the selection of initial seeds for the foreground or background. These seeds are then used as a training base for the final segmentation of cancer tissue. Wei Ju et al., presented in (Ju et al., 2015) a lung tumour segmentation technique on PET-CT scans combining random walk with a graph cut algorithm. The random walk is used to initialise seeds for the graph cut algorithm and shape penalty cost is combined into the graph energy function to constrain the tumour area during segmentation. The graph-based segmentation methods are very powerful techniques because they allow the incorporation of prior knowledge (shapes, positions, sizes, seeds, textures) into the graph energy formulation to guide the algorithm defining the optimal segmentation results (Salazar-Gonzalez et al., 2014). The performance of graph-based techniques above depend highly on the selection of initial seed points (Esneault et al., 2007; Chen et al., 2012), shape constraints (Ju et al., 2015; Freedman and Zhang, 2005) and intensity variation in the images (Kaba et al., 2015; van der Lijn et al., 2008). As results they required parameters turning to be applied in different images data. We have addressed these problems, by incorporating a bias correction operation to correct corrected the intensity inhomogeneity and a graph cut algorithm, which allows the interaction of human expert to guide the algorithm.

The rest of the paper is organized as follows. Our segmentation method is discussed in Section 3. Section 4 highlights information about the data-set used in the study. Section 5 shows the experimental results of the proposed method and the conclusion is drawn in Section 6.

METHOD

An automated segmentation of the nephroblastoma tumour is not trivial. To produce an acceptable segmentation results, the segmentation algorithm will rely heavily on the prior knowledge such as shape, colour, texture, position. Therefore, a semiautomated segmentation method appears to be the best choice for the analysis of nephroblastoma scans in the clinical domain. This allows the interaction of expert physicians in order to guide the segmentation algorithm.

Our method serves this purpose, it allows the radiologists to interact with a 2D image slice. The first step of the segmentation algorithm consists of removing some common artefacts of MRI medical imaging from the scans (i.e. noise, intensity inhomogeneity). This pre-processing operation is performed using a bias correction algorithm (Tustison et al., 2010). The tumour extraction in multi-sequence MRI scans is performed by adapting a continuous max-flow min-cut approach (Yuan et al., 2010). This includes a free 2D hand drawing tool, which allows human experts to select a region of interest (i.e. tumour seeds) and a kernel-induced segmentation functional (Salah et al., 2011; 2014). The proposed method could provide time and cost saving in nephroblastoma annotation and it can be used for high-throughput studies, such as clinical trials. It will be incorporated in the segmentation tools for the Computational Horizons in Cancer (CHIC) project Stamatakos and Graf (2017), funded under the European Commission FP7 programme. Fig. 1 shows the illustration of the proposed method.

PRE-PROCESSING

Generally, the fuzziness of tumour boundary (lack of clear edges) in multi-sequence MRI scans is caused by the presence of imaging artefacts such as noise, inhomogeneity and similarity to surrounding tissues. The bias correction operation (Tustison *et al.*, 2010) is used to correct these issues in the images to allow a correct segmentation of the tumour in the scan. The operation is performed by estimating the residual bias field. This residual is then subtracted from the corrupted scans to enhance the tumour. The bias correction operation can also reduce false positives during the segmentation pro

TUMOUR EXTRACTION

To perform the segmentation of nephroblastoma tumour from MRI scans, we represent each pixel point in a scan as a graph $G(v,\varepsilon)$, consisting of a set nodes v (i.e. pixels) and a set of edges ε connecting neighbouring nodes or pixels $\{p,q\}$ Boykov and Jolly (2001). The nodes also include two special nodes known as terminals S (foreground:tumour seeds) and T (background seeds). Each pair of neighbouring pixels $\{p,q\}$ in the image grid is connected by n-links (i.e. edge between pixels). Each pixel p in the scan is also connected to the foreground terminal through the slinks with $\{p, s\}$ and to the background terminal with t-links with $\{p, t\}$.

The edges in the graph including both n-links and t-links are assigned some positive weights $W_e > 0$ based on the similarity measured between pixels (i.e. intensities). After creating the graph, the segmentation is performed by producing a cut through the graph. The preferred cut or path will have the minimum total weights of edges for travelling from a start node to an end node. The selected cut separates the MRI scan into two disjoint regions, one representing the tumour tissue (foreground) and the other one the rest of the image (background) as shown in Figure 3 2.

If s - t is the cut that separates the tumour tissue $(F_g \text{ or foreground pixels})$ from the rest of the image $(B_g \text{ or background pixels})$, then s - t represents a subset of edges $C \in \varepsilon$ where

$$|C| = \sum_{e \in C} W_e \tag{1}$$

To formulate the energy function of the graph, let *A* be an indexing function, whose components assign each pixel in the MRI image to a region $R_l = \{p \in \Omega \mid p \text{ is labelled } l\}$:

$$A: p \in \Omega \longrightarrow A_p \in \eta \tag{2}$$

where η is a finite set of binary labels (i.e., l = 1 if I_p is a foreground pixel (F_g) and l = 0 if I_p is a background pixel). Therefore, each label l in the MRI image grid represents either the tumour pixel F_g (foreground) or the background pixel B_g dividing the image in two regions. From this the energy function of the segmentation can be defined as:

$$E(A) = B(A) + \lambda \cdot R(A).$$
(3)

where B(A) represents the boundary term, it describes the coherence between neighborhood pixels (base on their intensities). R(A) is the data term or regional term. It represents the likelihood of a pixel belonging to a region in a scan and it is used to specify the labelling preference for pixels based on their intensities. λ is a non-negative coefficient specifying the importance of the data term relative to the boundary term Boykov and Jolly (2001) and it has a value of 0.2 in our case. The boundary term is derived as:



Fig. 1: Illustration of different processes of the proposed method. Top left: 3D MRI Image. Top right: 2D slice for user selection of ROI. Bottom left right: segmentation results of 2D slice. Bottom right: segmentation result in 3D.

$$B(A) = \sum_{p,q \in N} B_{p,q} \cdot \phi(A_p, A_q)$$
(4)

where

$$\phi(A_p, A_q) = 1 \text{ for } A_p \neq A_q$$
(5)
$$\phi(A_p, A_q) = 0 \text{ otherwise}$$

$$B_{p,q} = \exp\left(-\frac{(I_p - I_q)^2}{2\sigma^2}\right) \cdot \frac{1}{\mathrm{d}(p,q)} \tag{6}$$

where $B_{p,q}$ evaluates the discontinuity between neighbouring pixels in Ω . The value of $B_{p,q}$ is large when the pixels intensities I_p and I_q are similar and it is close to zero otherwise Boykov and Jolly (2001). σ is the standard deviation and d is the distance between two pixels (*p* and *q*) Boykov and Jolly (2001). The values of both parameters were defined experimentally where $\sigma = 5e^{-4}$ and d = 0.11 Similarly the data term is defined as:

$$R(A) = \sum_{p \in \Omega} R_p(A_p) = \sum_{l \in \eta} \sum_{p \in R_l} -\log P(I_p \mid R_l). \quad (7)$$

where R_l represents the scan region that has label l and $P(I_p | R_l)$ is the conditional probability that measures the likelihood of a pixel intensity (I_p) in the scan grid given a model distribution in each image region R_l . Due to the complexity of the MRI scan, the piecewise Gaussian distribution, which is often used as piecewise constant segmentation model is not always sufficient to extract the tumour tissue from a nonlinearly separable MRI image. For this reason, the data term of the graph formulation is implicitly transformed by mapping the original MRI data into a higher dimensional feature space using a kernel function (Salah *et al.*, 2011; 2014), where the piecewise constant model of the graph cut formulation is applicable. This process allows a linear separation in the image data.

Let κ_l be the piecewise constant model parameter of an image region R_l , then the data term (7) can be defined as:

$$R(A) = \sum_{p \in \Omega} R_p(A_p) = \sum_{l \in \eta} \sum_{p \in R_l} (\kappa_l - I_p)^2.$$
(8)

If $\psi(.)$ is a nonlinear mapping function from the MRI data space Θ to a higher dimensional feature space Δ . The graph cut formulation becomes:

$$E(\{\kappa_l\}, A) = \sum_{p,q \in N} B_{p,q} \cdot \phi(A_p, A_q) + \lambda \cdot \sum_{l \in \eta} \sum_{p \in R_l} (\psi(\kappa_l) - \psi(I_p))^2.$$
⁽⁹⁾

After assigning a unique label to each pixel, the MRI image can be divided into two regions with each region containing one label. In this case a kernel induced space image segmentation with the graph cut would simply result in finding the labelling which minimises the graph formulation (9). By applying Mercer's theorem and simplifying Salah *et al.* (2011), $(\psi(\kappa_l) - \psi(I_p))^2$ in (9), we obtain the following kernel-induced segmentation functional

$$E(\{\kappa_l\}, A) = \sum_{p,q \in N} B_{p,q} \cdot \phi(A_p, A_q) + \lambda \cdot \sum_{l \in \eta} \sum_{p \in R_l} R(I_p, \kappa_l).$$
(10)

The expression in (10) depends on both the labelling function *A*, the image region parameter κ_l and it is optimised using max-flow algorithm Yuan *et al.* (2010). This optimisation process allows a separation of the tumour tissue from the MRI scan.

The interactive segmentation is performed as follows: The slices in a scan are displayed as a single image object as shown in Fig. 2. This operation allows the user to visualize all the slices that contain the tumour. The expert user can select tumour seeds region in one of the tumour slices (red circle in Fig. 2). The intentisty values of the selected tumour seeds is used as the foreground seeds of the gragh. Then the likelihood of each pixel p in the scan belonging to the tumour pixels is meseaued using (10).

The rest of the slices are automatically segmented in the scan by the proposed method as shown in (Fig. 2 (b)). One of the advantages of this interactive segmentation is that it allows radiologists to interactively select tumour seeds in one of the scan slices to allow a complete extraction of the tumour and avoid the segmentation of health tissues or other tissues similar to the tumour. It is also allows a global optimal solution for a graph segmentation, which is not possible in a fully automated segmentation methods.

DATA SET

The multi-sequence MRI scan data used in this study was obtained from ongoing pioneering research conducted on Wilms tumour in the department of Paediatric Oncology and Haematology at Saarland University Medical Centre in Homburg, Germany. The data set contains a sequence of T2 weighted multisequence MRI scans with transversal cut of 24 cancer patients. The acquisition parameters of multi-sequence MRI are set differently by the operators to allow a better visualisation of different scans as seen a Table 1.

Table 1: Details information of the dataset use in the study. P = Patient, R.T = Repetition Time, E.T = Echo Time, I.S = Inslice Sampling, S.T = Slice Thickness, S.S = Spacing between Slices.

Р	R.T	E.T	I.S	S.T	S.S
P1	2879.42	120.00	1,19x1,19	5.00	6.50
	3644.50	120.00	1x1	5.00	6.50
P2	434.26	80.00	0.63x0.63	4.00	6.00
	1726.45	80.00	0.98x0.98	4.00	4.40
P3	2219.72	80.00	0.9x0.9	4.00	4.40
	2219.72	80.00	0.9x0.9	4.00	4.40
P4	2470.00	104.00	0.51x0.51	5.00	5.50
	2470.00	104.00	0.45x0.45	5.00	5.50
P5	3407.29	72.00	0.99x0.99	7.00	9.10
	1000.00	96.00	0.78x0.78	4.50	5.85
P6	1994.36	80.38	0.63x0.63	5.00	5.50
	2000.00	80.13	0.63x0.63	5.00	5.50
P7	3563.53	98.00	1.13x1.13	6.00	6.60
	1618.53	98.00	1.19x1.19	6.00	7.80
P8	4.30	2.15	0.59x0.59	6.00	7.20
	4.30	2.15	0.59x0.59	6.00	7.20
P9	4500.00	65.11	1.72x1.72	5.00	6.00
	5460.00	72.02	0.94x0.94	5.00	6.00
P10	1410.00	114.00	0.58x0.58	7.00	8.40
	1410.00	114.00	0.58x0.58	7.00	9.10
P11	7058.80	88.00	0.94x0.94	5.00	5.50
	9230.77	84.00	0.47x0.47	5.00	5.50
P12	1800.00	100.00	0.78x0.78	6.00	7.00
	3808.47	99.00	0.33x0.33	3.00	3.60
P13	1200.00	62.00	0.39x0.39	5.00	6.00
	6261.80	145.00	0.25x0.25	3.29	3.80
P14	7740.00	9.60	0.98x0.98	6.00	8.10
	1000.00	82.00	1.37x1.37	6.00	7.20
P15	4633.50	105.00	1.15x1.15	4.00	5.20
	3760.00	105.00	1.17x1.17	4.00	5.20
P16	307.00	1.27	1.17x1.17	5.00	6.50
	4.63	2.32	0.98x0.98	5.00	8.00
P17	369.95	80.00	1.09x1.09	6.00	7.00
	351.39	60.00	0.49x0.49	5.00	6.00
P18	4868.15	109.00	0.63x0.63	4.00	4.80
	4377.06	109.00	0.63x0.63	4.00	4.80
P19	4333.09	105.00	0.78x0.78	4.00	4.80
	4160.83	105.00	0.78x0.78	4.00	4.80
P20	6432.04	109.00	0.65x0.65	4.00	4.80
	6360.66	109.00	0.65x0.65	4.00	4.80
P21	5091.47	80.00	0.39x0.39	4.00	4.40
	5826.67	96.00	1.15x1.15	4.00	4.40
P22	7852.03	69.00	0.94x0.94	5.00	5.50
	12590.94	69.00	0.94x0.94	5.00	5.50
P23	1000.00	83.00	1.19x1.19	5.00	5.50
	4852.50	92.00	0.72x0.72	4.00	4.40
P24	1300.00	80.00	0.94x0.94	5.00	5.50
	1875.00	80.00	0.91x0.91	5.00	5.50

Each patient has two scans, one before radiochemotherapy and the other one after the radiochemotherapy to determine the effectiveness of the treatment. The age of the patients ranged from 1 to 8 year old including intermediate risk tumour (stromal



Fig. 2: User Selection of ROI. (a) the user expertise is needed to select a tumour tissue (red circle) in one of the 3D image slices, the rest of the tumour tissues are segmented automatically by the proposed algorithm. (b) the segmentation results of the method. (c) manually labelled image slices.

predominant, mixed histology or regressive) and high risk tumour (blastemal predominant). In total we have 48 scans from the 24 patients including before and after chemotherapy. The tumour area of each scan was manually annotated by an experienced radiologist with a total hand labelled of 48 images. These expert hand labelled images were used as benchmark (ground truth) to evaluate the performance of our segmentation method. This study has not measured the inter and intra observer variability due to the unavailability of additional hand labelled data-set.

RESULTS AND DISCUSSIONS

The proposed method is implemented on MATLAB R2011b and the computation time of our algorithm is less than 80 seconds for each multi-sequence MRI scan on a MAC OX X running at 2.66 GHz, with 4G of RAM memory. To evaluate the accuracy of our method, we compared our results against the experts manual annotations. The evaluation is performed using statistical metrics such as the sensitivity (true positive rate), the specificity (true negative rate), the false negative rate and the average accuracy rate are shown in Table 2. Others metrics including the Dice coefficient, the root-mean-error and the Jaccard index are also used for the validation of spatial tumour shape representation in Table 3.

Table 2 shows the evaluation results of the proposed method using the above metrics on all the 48 scans. The proposed method achieves a high performance with average accuracy of 99.59%, TPR = 94.52% and FPR = 0.34%. The same performance is also observed in Table 3 with a Dice coefficient of 0.9060, Jaccard index of 0.8326 and an average RMSE of 0.0481. An overview of the performance results shows that the proposed method achieved a good agreement with experts manual labelled tumour boundaries.

To facilitate the performance comparison between our method and the experts hand-labelled tumours, we

also tested our method separately on scans before and after radio-chemotherapy. Table 4 and Table 5 compare the performance of the proposed method on the scans before and after treatment. An overview of the results shows that in both scans, our proposed method is in good agreement with experts manual labelled. However the segmentation of the scans after radio-chemotherapy performs slightly better than the one before radio-chemotherapy: around 0.18% better on average accuracy. In addition to the tables, Fig.3 shows the box plots of Dice coefficient, RMSE, TNR and TPR of the all scans. Fig.4 shows the segmentation results comparison of the proposed method against the experts hand-labelled tumour. Fig.5 shows the segmentation results of a scan before and after the radio-chemotherapy.

For additional illustration, we performed the box plots of the Dice coefficient, RMSE, TNR and TPR for the evaluation of our method as shown in Fig 3.

Fig.6 illustrates the importance of the semi-automated segmentation method for Wilms' tumour. In Fig.6 (a) without the expert knowledge, it is almost impossible to define the tumour region due the non-uniformity of the scan intensity and the irregular shape of the tumour. Therefore the need of expert input (i.e. selection of tumour seeds) is essential for the extraction of the tumour from the scan. This shows the importance of semi-automatic segmentation methods in Wilms' tumour segmentation, which is not possible in fully automated segmentation methods.

Table 2: Performance evaluation of the tumour tissue segmentation using true positive rate (TNR), false positive rate (FNR) and the accuracy ACC) with their respective standard deviation (Std) and 95% confidence interval (Conf). - 48 scans

Metrics	Average	Std	Conf
Sensitivity (TPR)	0.9452	0.0429	0.0077
Specificity (TNR)	0.9966	0.0039	0.0007
FPR	0.0034	0.0039	0.0007
FNR	0.0548	0.0429	0.0077
ACC	0.9959	0.0039	0.0007

Table 3: Performance evaluation of the tumour tissue segmentation using the Dice coefficient, the root-mean-square (RMSE) and the Jaccard index with their respective standard deviation and 95% confidence interval (Conf). - 48 scans

Metrics	Average	Std	Conf
Dice	0.9060	0.0549	0.0098
Jaccard	0.8326	0.0898	0.0161
RMSE	0.0481	0.0309	0.0055

Table 4: Performance evaluation of the tumour tissue segmentation using true positive rate (TNR), false positive rate (FNR) and the accuracy ACC) with their respective standard deviation (Std) and 95% confidence interval (Conf). - 24 scans before and after radio-chemotherapy

Before therapy	Average	Std	Conf
Sensitivity (TPR)	0.9446	0.0355	0.0064
Specificity (TNR)	0.9960	0.0044	0.0008
FPR	0.0040	0.0044	0.0008
FNR	0.0554	0.0355	0.0064
ACC	0.9950	0.0042	0.0008
After therapy	Average	Std	Conf
After therapy Sensitivity (TPR)	Average 0.9458	Std 0.0500	Conf 0.0089
After therapySensitivity (TPR)Specificity (TNR)	Average 0.9458 0.9972	Std 0.0500 0.0033	Conf 0.0089 0.0006
After therapy Sensitivity (TPR) Specificity (TNR) FPR	Average0.94580.99720.0028	Std 0.0500 0.0033 0.0033	Conf 0.0089 0.0006 0.0006
After therapySensitivity (TPR)Specificity (TNR)FPRFNR	Average0.94580.99720.00280.0542	Std 0.0500 0.0033 0.0033 0.00500	Conf 0.0089 0.0006 0.0006 0.0089

Table 5: Performance evaluation of the tumour tissue segmentation using the Dice coefficient, the root-mean-square (RMSE) and the Jaccard index with their respective standard deviation (Std) and 95% Confidence Interval (Conf). - 24 scans before and after radio-chemotherapy

Before therapy	Average	Std	Conf
Dice	0.9046	0.0425	0.0076
Jaccard	0.8285	0.0705	0.0126
RMSE	0.0551	0.0290	0.0052
After therapy	Average	Std	Conf
Dice	0.0074	0.0660	0.0119
Dicc	0.9074	0.0000	0.0110
Jaccard	0.8367	0.0000	0.0118



Fig. 3: Box plots of Dice coefficient, RMSE, TNR and TPR of the all scans.



Fig. 4: Row (a): left-hand side 2D medical image, right-hand side 3D scan (input image). Row (b): left-hand side segmentation result of the proposed method in 2D (green colour), right-hand side segmentation result in 3D (green colour). Row (c): left-hand side 2D hand-labelled image (red colour), right-hand side corresponding 3D hand-labelled (red colour). Row (d): overlap 2D image of the segmentation result and the hand-labelled image, right-hand side corresponding 3D overlapping of both scans.



Fig. 5: Segmentation results. Column (a): 2D images before radio-chemotherapy, green the segmentation result, red the hand-labelled and the overlap of both images. Column (b): corresponding 3D images before radio-chemotherapy, green the segmentation result, red the hand-labelled and the overlap of both images. Column (c): 2D images after radio-chemotherapy, green the segmentation result, red the hand-labelled and the overlap of both images. Column (d): corresponding 3D images after radio-chemotherapy, green the segmentation result, red the hand-labelled and the overlap of both images. Column (d): corresponding 3D images after radio-chemotherapy, green the segmentation result, red the hand-labelled and the overlap of both sans.



Fig. 6: Illustration of the importance of our semi-automatic segmentation method from selected slices in the 3D scan: (a) Input scan with ill-defined boundary. (b) our segmentation result, (c) the hand-labelled image.

CONCLUSION

In summary, we have developed a semi-automated method to extract Wilms' tumour from multi-sequence MRI scans. The method has been designed, tested and validated in conjunction with medical experts, enabling its deployment and exploitation in the clinical setting. The overall procedure includes a pre-processing step that enhances the contrast of the tumour in a multi-sequence scan using a bias correction operation, and a tumour extraction step that includes kernel mapping and continuous maxflow algorithm. The method was validated on 48 multi-sequence MRI scans and it is proved to be accurate, robust, flexible, and fast, leading to a successful extraction of the tumour tissue. Our method has the advantage over fully automated segmentation methods because it allows an easy interaction of the user with different slices and facilitates the

segmentation of irregular tumour tussies as well as enabling the radiologist to make decisions regarding tumour size and shape. During the research, we have identified challenges and opportunities that constitute the basis for future improvement and utilisation of the technique. Accuracy, robustness and flexibility of the algorithm is crucial for the assessment of the tumour for surgical planning and post-surgical assessment.

Radiomics is an emerging field where both imaging and non-imaging information are combined for noninvasive imaging biomarkers that can be used for assessing tumour phenotyping, association to genetic biomarkers (radiogenomics) using machine learning methods. These advances in imaging allow to further improve the current response to tumour analysis criteria that currently rely on over-simplistic metrics of tumour size and disease progression. Combining these methods with advanced Radiomics, and using machine learning methodologies would refine and personalise the evaluation of patient imaging data.

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