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Original Research Article

Automatic Three-Dimensional Magnetic Resonance-based measurements of tumour prominence and basal diameter for treatment planning of uveal melanoma

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ABSTRACT

Background and Purpose: Three-dimensional (3D) Magnetic Resonance Imaging (MRI) is increasingly used to complement conventional two-dimensional ultrasound in the assessment of tumour dimension measurement of uveal melanoma. However, the lack of definitions of the 3D measurements of these tumour dimensions hinders further adaptation of MRI in ocular radiotherapy planning. In this study, we composed 3D MR-based definitions of tumour prominence and basal diameter and compared them to conventional ultrasound.

Materials and methods: Tumours were delineated on 3DT2 and contrast-enhanced 3DT1 (T1gd) MRI for 25 patients. 3D definitions of tumour prominence and diameter were composed and evaluated automatically on the T1gd and T2 contours. Automatic T1gd measurements were compared to manual MRI measurements, to automatic T2 measurements and to manual ultrasound measurements.

Results: Prominence measurements were similar for all modalities (median absolute difference 0.3 mm). Automatic T1gd diameter measurements were generally larger than manual MRI, automatic T2 and manual ultrasound measurements (median absolute differences of 0.5, 1.6 and 1.1 mm respectively), mainly due to difficulty defining the axis of the largest diameter. Largest differences between ultrasound and MRI for both prominence and diameter were found in anteriorly located tumours (up to 1.6 and 4.5 mm respectively), for which the tumour extent could not entirely be visualized with ultrasound.

Conclusions: The proposed 3D definitions for tumour prominence and diameter agreed well with ultrasound measurements for tumours for which the extent was visible on ultrasound. 3D MRI measurements generally provided larger diameter measurements than ultrasound. In anteriorly located tumours, the MRI measurements were considered more accurate than conventional ultrasound.

1. Introduction

Uveal melanoma (UM) is the most frequently occurring primary intraocular malignancy [1]. For UM, tumour location and dimensions are major factors in treatment decision making, as only tumours up to a certain size can be treated with brachytherapy [1–3], whereas larger and juxtapapillary UM are preferably treated with external beam radiotherapy, such as proton beam therapy (PBT) or Cyber Knife [4–6]. Furthermore, tumour prominence and diameter measurements are of importance for radiotherapy planning: in brachytherapy, prominence and largest basal diameter (LBD) determine the application time and applicator size, while for PBT, these two-dimensional (2D) tumour measurements, together with the second basal diameter (SBD) are important factors in the definition of three-dimensional (3D) clinical target volume [2,7]. Additionally, prominence, LBD and SBD are used in tumour staging, prognosis and follow-up [8–12].

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Conventionally, tumour prominence and diameters are determined using 2D B-scan ultrasound [1]. For a correct prominence measurement, the transducer has to be positioned perpendicular to the tumour base, as oblique cuts through the tumour can result in overestimation of the true tumour prominence [13]. For anterior tumours, correct transducer placement might be hindered due to anatomical structures around the eye [14]. Additionally, to correctly determine the LBD, the longest tumour axis has to be identified, which can be difficult as no 3D visualization is available.

Magnetic resonance imaging (MRI) is increasingly used for the diagnosis, therapy planning and follow-up of UM [7,15–20]. Various studies suggest that MRI's 3D tumour visualisation and better tissue contrast might be beneficial in determining UM tumour dimensions compared to conventional 2D ultrasound, corresponding to findings in other malignancies [16–24]. Nevertheless, several uncertainties in MR-based dimensions measurements for ocular tumours remain that hinder broader application of MRI in ocular radiotherapy planning, for example regarding the optimal method to measure in 3D and the differences related to the various MRI contrasts.

Although the 3D visualisation of the tumour of MRI allows for a more comprehensive assessment of tumour geometry [18], tumour dimension measurements can be performed in different planes, resulting in different possible prominence and diameter measurements, especially for complexly shaped tumours (Fig. 1) [19,25]. Additionally, manual measurement of prominence and LBD on MRI is reported to be difficult and time-consuming [26]. Therefore, the current clinical practice might benefit from automated tumour prominence and diameter measurements, using an unambiguous geometrical definition. Furthermore, studies showed an 11–44 % larger tumour volume on contrast-enhanced T1-weighted images (T1gd) compared to T2-weighted images [21,27]. Although such a difference between MRI sequences is not uncommon [28,29], these differences should be kept in mind when defining optimal measurements.

In this study, we aimed to compose and evaluate a geometrical 3D definition of prominence and diameters.

2. Materials and methods

Data of 25 patients, who received an MRI as part of clinical care (n = 17) or in the context of a scientific study (n = 8), were analysed retrospectively after approval of the local ethics committee, in accordance with the Declaration of Helsinki. Patients were diagnosed by an ocular oncologist based on fundus photography, ocular ultrasound and fluorescein angiography. The patients were selected to provide a clinically representative range in tumour size, shape and location. Patients were

 62 ± 10 years old and 57 % were male. A wide range of tumour sizes was present, ranging from The American Joint Committee on Cancer (AJCC, 8th edition) stage T1 up to T4, with most patients in stages T2 (40 %) and T3 (40 %) [30]. 28 % of patients had an anteriorly located tumour and most tumours were dome-shaped (68 %), followed by mushroom-shaped (16 %) and flat (16 %). Patients were treated with either 106-Ruthenium brachytherapy (32 %) or PBT (68 %). The average time between ultrasound and MRI was 14 ± 11 days.

2.1. Image acquisition

3DT1gd and 3DT2-weighted images, both with acquisition voxel size of $0.8 \times 0.8 \times 0.8 \text{ mm}^3$, were acquired to an earlier described protocol [15,16]. Ultrasound images were obtained by an ocular oncologist [25]. Details on image acquisition are further described in the Supplementary Materials.

2.2. Tumour delineation

Delineations were performed semi-automatically on the 3D T1gd and T2-weighted images using an in-house developed analysis pipeline in MeVisLab (MeVis Medical Solutions, Bremen, Germany) using a subdivision surface fit [26,31] (Fig. 1a-b). Areas which were hypointense compared to the vitreous on T2 and enhancing on T1gd compared to the native T1 were considered tumour.

The globe, defined as a combination of the vitreous, lens and tumour, was delineated in a similar fashion (Fig. 2a-b, yellow contour). Since clinical ultrasound measurements include the sclera, the outer sclera boundary at the tumour base was delineated as well (Fig. 2c). Based on the clinical evaluation of the MR-images by a neuro-radiologist with 20 years of experience (BV), an ophthalmic MRI expert with 9 years of experience (JWB) verified, and if necessary corrected, all contours.

2.3. Automatic 3D MRI measurements

In a pilot study, further described in the Supplementary Materials, we developed 3D definitions of tumour prominence and base. In short, three different 3D definitions of the tumour prominence and three different LBD definitions were evaluated in a multidisciplinary tumour board, consisting of ocular oncologists, radiation oncologists, clinical physicists and radiologists (BV, TV, MM, GL, CR, JWB). As these definitions were based on the 3D tumour contours, the resulting dimensions were not restricted to an individual slice of the MR-images. These definitions provided different approaches on how to define the tumour base and apex, e.g. point closest to the eye centre or point most distant from



Fig. 1. Several measuring planes on T1gd-weighted MRI for a mushroom shaped UM. (a) Three possible prominence measurements in the axial plane for a complexly shaped tumour: starting with the point closest to the centre of the eye (solid, the definition used in this study), parallel with the main tumour axis (dotted) and including the overhanging part of the tumour (dashed). (b) An evaluation of the same tumour in the sagittal plane resulted in a different measurement. (c) For obliquely oriented tumours, several base definitions are possible depending on whether the overhanging part is included as tumour base.



Fig. 2. Tumour (white contour) and globe (defined as combination of vitreous, lens and tumour, yellow contour) were delineated on T2-weighted (a) and contrastenhanced T1-weighted (T1gd) (b) images, with the native T1-weighted image used as a reference (inset). Anatomical structures are marked as follows: (1) tumour, (2) retinal detachment, (3) lens, (4) sclera (hypointense layer), (5) vitreous, (6) optic nerve. Tumour prominence (c, red line) and largest basal diameter (d, blue line) were automatically determined. (c) Tumour prominence was measured from the tumour apex (white dot) to the sclera (dashed yellow line) (centre-based definition, Appendix A). (d) LBD was the largest Euclidean distance between two points in the tumour base (base: dashed line, measurement: solid line). In this image, the axial plane was shown as an example, whereas the tumour delineation was evaluated in the axial, sagittal and coronal plane. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

the sclera. The evaluation of these definitions in 25 patients showed comparable measurements for patients with rotational-symmetric tumours. However, in the asymmetric tumours, some of the definitions resulted in measurement which did not conform the current clinical consensus. For example, defining the apex as the point most distant from the sclera resulted in an oblique prominence measurement which could not be used for radiotherapy planning purposes (Supplementary Fig. 1). The multidisciplinary evaluation of the definitions resulted in one prominence and one tumour base definition which provided visually acceptable measurements in all patients.

Based on the results of this pilot study, the tumour apex was defined as the point of the tumour contour closest to the centre of the globe (Fig. 2c). The prominence was measured along the line between the centre of the globe and the tumour apex (Fig. 1a, solid red line) and included the sclera to match the clinical convention at our centre.

LBD was defined as the largest Euclidean distance between two points in the tumour base (Fig. 2d, base: dashed line, LBD: solid line), which was defined as all points of the tumour contour less than 2 mm distant from the inner sclera contour, in order to create a robust tumour base. SBD was the largest distance between two points in the tumour base, perpendicular to the LBD.

For all patients the prominence, LBD and SBD were automatically determined using the T1gd- and T2-based 3D contours in Python (version 3.7.6) (Supplementary Materials).

2.4. Manual MRI measurements

In order to assess how these automatic 3D measurements relate to the currently used manual assessments, the prominence and LBD were measured manually on the T1gd-weighted images. For the prominence, the clinical measurement was used, measured perpendicular to the sclera [25]. For the LBD, the measurements were repeated on the delineated tumour contour, to avoid the confounding effect of a different tumour boundary interpretation [27]. For each patient, the angle between the lines of manual and the automatic measurement was determined.

2.5. Statistical analysis

Automatic T1gd prominence and LBD measurements were compared to manual MRI measurements. Additionally, automatic prominence and LBD were compared between T1gd and T2. Furthermore, the automatic T1gd and clinical ultrasound measurements were compared for prominence, LBD and SBD. For the prominence, differences > 0.5 mm and for the diameters, differences > 1.0 mm were considered clinically significant [25]. Medians and 25th-75th percentiles of the automatic-manual, T1gd-T2 and T1gd-ultrasound differences were assessed, as nonnormality was demonstrated with a Shapiro-Wilk test (e.g. p = 0.0003for the ultrasound-T1gd prominence comparison). Results for anterior tumours, defined as having a centre of mass in the anterior 50 % of the globe, were assessed separately. Additionally, tumours where the full extent could not be accurately determined on MRI or ultrasound, e.g. due to a flat tumour or a limited penetration depth or limited field of view (FOV) of the ultrasound transducer, were marked (Supplementary Fig. 2) [25].

3. Results

Representative examples of automatic prominence and LBD measurements on T2 and T1gd, and the corresponding ultrasound measurements, are shown in Fig. 3, with an overview of the primary outcomes. All comparisons and individual measurements for all patients can be found in the Supplementary Materials.

On T1gd, manual and automatic prominence measurements were comparable (median absolute difference 0.3 mm, Fig. 4a), with differences > 0.5 mm for 8/25 patients. The median absolute difference between automatic and manual LBD measurements was 0.5 mm, with differences > 1.0 mm for 14/25 patients (Fig. 4b). Larger differences were found in patients where the manual measurement was performed in a different plane than the automatic measurement (Fig. 4c). The mean angle between the manual and automatic measurement was 26 degrees (Fig. 4c, Supplementary Fig. 3).

Due to a limited penetration depth of the UBM transducer, the tumour apex was not visible in 2/25 patients, both of which had an anteriorly located tumour (Supplementary Fig. 2a). The full extent of the tumour base did not fit within the ultrasound's field of view for 9/25 patients, of which 7 had an anteriorly located tumour (Supplementary Fig. 2b). In 3/4 patients with an ultrasound prominence < 4 mm, the extent of the tumour base was difficult to assess on MRI.

The difference between ultrasound- and automatic T1gd-based prominence was > 0.5 mm in 6/25 patients, with a median absolute difference of 0.4 and 0.3 mm for anterior and posterior tumours, respectively (Fig. 5a). A retrospective re-evaluation of the ultrasound and fundus images of the patient with the largest difference (1.3 mm), showed that a haemorrhage at the tumour apex was erroneously



e	Median (max) absolute difference [mm]		T2 – T1gd	US – T1gd	US – T2	Manual – automatic (MRI T1gd)
	Prominence	Entire group	0.3 (0.9)	0.3 (1.6)	0.4 (1.3)	0.3 (0.9)
		Anterior	0.3 (0.6)	0.4 (1.6)	0.6 (1.3)	0.2 (0.7)
		Posterior	0.3 (0.9)	0.3 (1.3)	0.4 (0.7)	0.3 (0.9)
	LBD	Entire group	1.7 (6.7)	1.1 (7.6)	1.1 (6.8)	0.5 (5.3)
		Anterior	1.0 (4.1)	1.6 (4.5)	1.6 (4.4)	0.5 (1.4)
		Posterior	1.8 (6.7)	1.0 (7.6)	1.1 (6.8)	0.5 (5.3)

Fig. 3. Four typical examples of ultrasound (upper images) and MRI measurements (lower images) from four different patients and an overview of the main comparisons. (a) T2 and ultrasound measurements were similar. (b) T1gd prominence was larger for an anteriorly located tumour for which the extent was not entirely visible on UBM. (c) T1gd LBD was larger than on ultrasound for a tumour whose extent was visible on both modalities. (d) T1gd LBD was larger than on ultrasound and extent was not entirely visible on ultrasound. (e) Overview of the main comparisons performed in this study: automatic T2 – automatic T1gd, ultrasound (US) – automatic T1gd, ultrasound – automatic T2, manual T1gd – automatic T1gd.



Fig. 4. (a) Differences between automatic and manual prominence measurements. (b) Differences between automatic and manual LBD measurements. (c) Manually determining the plane of the largest basal diameter proved to be difficult due to the curved tumour base. As a result, manual measurements generally underestimated the LBD.

included as tumour on ultrasound (Supplementary Fig. 4). For the remaining patients the largest absolute difference of 0.9 mm was found in a patient with an anteriorly located tumour, for which the apex was visible on ultrasound, with a prominence of 13.3 mm on ultrasound and 12.4 mm on MRI. Differences between T1gd- and T2-based prominence were small: for 6/25 patients, the difference was > 0.5 mm, with a median absolute difference of 0.3 mm and a maximum of 0.9 mm (Fig. 5b).

Compared to ultrasound, 16/25 of LBD measurements were > 1.0 mm larger on the automatic T1gd measurement (Fig. 3c). Median absolute differences between ultrasound and T1gd (Fig. 5c) were 0.9 mm for posteriorly located tumours and 1.6 mm for anteriorly located tumours. Slightly larger differences were found for the SBD, which resulted in a median absolute difference of 1.3 mm (maximum 4.6 mm, Supplementary Fig. 5).

Differences between automatic T2 and automatic T1gd LBD were > 1.0 mm in 15/25 patients, with 13/15 being larger on T1gd. A median absolute difference of 1.6 mm was found between the two sequences (Fig. 5d). Often, peritumoral choroidal enhancement was visible on T1gd, which consequently was included as tumour, while this was not considered tumour on T2 and ultrasound (Fig. 6b-c). T2-based LBD measurements were more in agreement with ultrasound (median absolute difference 0.4 mm), compared to T1gd-based measurements (Supplementary Fig. 6). These smaller diameters on T2 were also reflected in the tumour volumes: tumour volume was larger on T1gd for 20/25 patients (average volume difference 18 %).

4. Discussion

With this study, we proposed an unambiguous 3D-imaging based prominence and base definition for treatment planning of uveal melanoma, which takes the requirements for brachytherapy and PBT planning into account [25,32,33], and which poses an extension of an earlier study [34], which assumed a dome shaped tumour with a circular base.

The proposed prominence and base definitions were considered appropriate by the multidisciplinary tumour board in our cohort of 25 patients. Although the cohort contained a variety of tumour sizes and shapes, we noted that in none of the obliquely oriented tumours, the LBD contained the overhanging part. As a result, the effects from overhanging parts were not reflected in the LBD assessments. Accurately incorporating overhanging parts in directions other than the LBD measurement axis, will however be beneficial for PBT, where similar discrepancies have been reported describing the tumour-marker relation [25]. We anticipate that the proposed definition may result in erroneous measurements for very prominent UM, with the centre of the eye within the mass. Nonetheless, the proposed prominence definition proved to be robust for variations in the location of the centre of the eye (Supplementary Fig. 7).

As the proposed automatic measurement methods are independent of field strength they are also applicable for patients scanned at lower field strengths than 3 T. In this study, subjects were scanned with closed eyes, as a previous comparison between different protocols showed that, while the use of a cued blinking protocol [35] results in less motion artefacts in some patients, not all patients are able to adhere to such a protocol, resulting in overall degraded image quality [36].



Fig. 5. Comparison of prominence and LBD between ultrasound and MRI. (a) For the prominence, ultrasound-T1gd differences were largest in anteriorly located tumours that were not fully visible on ultrasound. The largest prominence difference of 1.3 mm was caused by the erroneous inclusion of haemorrhage in the tumour prominence on ultrasound (Supplementary Figure 3). (b) Absolute differences between T1gd and T2 prominence measurements were < 0.5 mm in 19/25 patients. (c) Comparing ultrasound and T1gd MRI, LBD was larger on T1gd MRI in 22/25 patients. (d) Comparing LBD between T2 and T1gd, LBD was larger on T1gd in 22/25 patients.

As part of the data was acquired as part of regular clinical care, the MRI and ultrasound were not acquired on the same day for most patients. However, we expect the impact of this delay to be limited, as UM are known to be slow growing tumours [37,38], which is confirmed by the accordance in prominence measurements between both modalities.

In general, tumour prominence measurements agreed well across modalities if tumour extent was entirely visible, with all interquartile ranges within half the MRI acquisition voxel size. This corresponds to the recently reported interobserver delineation variation of 0.4 mm of ocular MRI [27] and is smaller than the reproducibility of B-mode ultrasound of 0.6 mm [13,14]. In our study, the largest prominence differences were found in patients for whom the tumour apex was not visible on ultrasound. This particularly occurred in patients with anteriorly located tumours imaged with a UBM probe, which has a limited penetration depth due to its higher frequency (50 MHz). For these patients, MRI was considered to be more accurate. Previous studies by Ferreira and Schueller [16,39] reported similar prominence measurements, although measurement methods differed.

Automatic LBD measurements were generally larger than manual MRI measurements, which were often performed in a plane not containing the largest diameter. Since the LBD can, by definition, not be overestimated, the automatic measurement was considered more accurate. However, since 68 % of the differences were < 1 mm, manual measurements are generally sufficient, especially given the known interobserver variability of the tumour-choroid interface on MRI of 0.6 mm [27]. On T1gd, peritumoral enhancement was included, which was invisible on T2, resulting in larger measurements on T1gd (median absolute difference 1.7 mm). Although the enhancement could be inflammatory of origin, it could also correspond to a small flat tumour extension. We therefore recommend to include these areas as tumour, until histopathology proves otherwise.

Generally, we found larger LBDs on T1gd than on ultrasound, with differences largest in anteriorly located tumours, especially when the full tumour extent was not visible on ultrasound. These differences can partly be explained by manual definition of the plane of the LBD of the ultrasound exam, which similar to the manual MRI measurements, might miss the actual largest diameter, with the added difficulty that no 3D visualisation can be made with ultrasound. In some patients the peritumoral choroidal enhancement observed on T1gd was included on ultrasound (Fig. 6c), while in other patients it was not (Fig. 6d). This inconsistency leads to uncertainties in the LBD measurement of flat UM as well, with differences between T1gd and ultrasound up to 7.6 mm. In 3/4 patients with flat UM, the MRI-report explicitly mentioned that the tumour extent was difficult to assess, confirming previous findings that MRI of flat melanoma can be difficult [16,25,34,40,41]. However, on ultrasound, the extent of these flat tumours is also not clearly visible. Therefore, optical validation of the flat tumour extent, which can be measured on some fundus and OCT imaging modalities [10], or increased margins, is necessary. For the SBD larger discrepancies with ultrasound were observed than for the LBD, however, the resulting ellipsoid model is more accurate than a circular model based on the LBD alone [8,9,42-45].

The clinical impact of the observed differences between ultrasound and MRI will depend on the application, and therefore an MRI-scan will not be needed for all patients. In general, we recommend to perform an MRI when the tumour extent is not fully visible on ultrasound, which was most often seen in anteriorly located tumours. For treatment selection, the effect of incorporating the MRI-based measurements



Fig. 6. Prominence and LBD measurement on T2-weighted MRI (left column), T1gd-weighted MRI (middle column) and US (right column). (a) Prominence was generally similar on T1gd and T2. (b) The largest difference between T2 and T1gd prominence was 0.9 mm. (c,d) LBD was generally larger on T1gd.

proposed in this study will depend on the type of treatments that are available. When a 0.5 mm change in prominence or 1 mm change in LBD can result in a shift of optimal treatment modality [19,46], we advise to perform an MRI with automatic T1gd measurements to confirm the ultrasound measurements. Although the benefit of including MRI in radiotherapy planning requires further study, we expect less benefit for patients treated with brachytherapy, due to the limited degrees of freedom with the currently available applicators. For PBT, however, the currently relatively large margins of up to 3 mm are in part needed for the uncertainties introduced by the ultrasound measurements [47–49]. Given the earlier reported benefits of including MRI for ocular PBT planning, we would recommend to perform an MRI for these patients and use the proposed 3D prominence and basal diameter definitions to resolve the discrepancies observed for complexly shaped tumours between MRI and ultrasound [25], especially until fully image-based 3D

treatment planning systems for UM are clinically available [21,50-52].

In conclusion, 3D MRI-based tumour measurements were comparable to 2D ultrasound-based measurements. For anteriorly located tumours, MRI was more accurate. Furthermore, the methods proposed in this paper can contribute to a more personalized, image-based, radiotherapy planning for ocular oncology patients.

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Declaration of Competing Interest

The authors declare the following financial interests/personal relationships which may be considered as potential competing interests: J. W.M. Beenakker received research support from Philips Healthcare. The other authors do not have any interests to declare.

Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.phro.2022.11.001.

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