Is CT Still Useful in the Study Protocol of Retinoblastoma?

BACKGROUND AND PURPOSE: Intralesional calcium deposition is considered a key element for differentiating retinoblastoma from simulating lesions. Our aim was to assess whether MR imaging associated with ophthalmologic investigations (ophthalmoscopy and ultrasonography) could replace CT in the detection of diagnostic intralesional calcifications in retinoblastoma.

MATERIALS AND METHODS: Ophthalmoscopic findings, MR images, CT scans, and histologic examination of 28 retinoblastomas from 23 consecutive children (11 males, 12 females; age range at admission, 1–35 months; mean age, 11 months; median age, 9 months) were retrospectively evaluated. Ultrasonography was performed in 18 patients with 21 retinoblastomas. MR imaging included T2-weighted spin-echo and gradient-echo images, fluid-attenuated inversion recovery images, and T1-weighted spin-echo images with and without contrast enhancement. Clinical data were integrated with MR imaging data to evaluate the utility of both approaches to discover calcifications; particularly, a correlation between intralesional signal-intensity void spots on MR imaging and hyperattenuating areas on CT scans was performed.

RESULTS: Ophthalmoscopy detected calcifications in 12 of 28 eyes (42.85%). Ultrasonography detected calcifications in 20 of 21 eyes (95.23%). CT showed hyperattenuating intralesional areas consistent with calcifications in 27 of 28 eyes (96.42%). MR imaging showed intralesional signal-intensity void spots in 25 of 28 eyes (89.28%). All spots detected with MR imaging matched the presence of calcifications on CT scans. Gradient-echo T2*-weighted and fast spin-echo T2-weighted images showed the highest degree of correlation with CT. When we put together ophthalmoscopy, ultrasonography, and MR imaging data, no calcifications detected on CT were missed, and the differential diagnosis was thorough.

CONCLUSIONS: A combination of clinical data and MR images may remove potentially harmful ionizing radiation from the study protocol of retinoblastoma.

Early diagnosis and treatment of retinoblastoma greatly improve the patient’s survival rate and quality of vision. Because fine-needle aspiration biopsy carries a significant risk of procedure-related tumor seeding,1,2 patients with retinoblastoma are treated on the basis of clinical and imaging data, without a histologic diagnosis. Thus, noninvasive diagnostic accuracy is crucial. A decisive additional support of the ophthalmoscopic diagnosis is given by matching intralesional calcifications.3,4 The detection of intraocular calcium deposition is a key element to differentiate retinoblastoma from simulating lesions. Histology reports show that retinoblastomas calcify in ≥95% of cases;5 CT has a designated sensitivity in detecting calcifications in retinoblastoma of 81%–96%;6,10 whereas the sensitivity of sonography has been reported to be ≤92.5% of cases.4 In children younger than 3 years of age, an intrabulbar calcified mass is most likely a retinoblastoma.11,12 However, in children older than 3 years of age, other intraocular lesions, such as retinopathy of prematurity, toxocariasis, Coats’ disease, retinal astrocytoma, and optic nerve drusen may appear as a calcified masses.3,12 Retinoblastoma may very rarely occur in older patients, but we have had a new diagnosis in a child older than 15 years of age (personal observation).13

CT or MR imaging or both are usually required to detect optic nerve, choroid, and scleral infiltration stages; disease spread beyond the eye; or intraocular disease. Imaging studies are used for diagnostic purposes only in very selected instances, as in cases in which ophthalmoscopy is impossible, such as with the presence of opaque ocular refractive media and massive retinal detachment or for differentiating lesions simulating retinoblastoma.11,14–19

The purpose of this study was to assess the utility of the association of ophthalmologic evaluation and MR imaging in the detection of intralesional calcifications in retinoblastoma, to evaluate whether CT is still mandatory in the protocol study of retinoblastoma and simulating lesions.

Materials and Methods

The population of this prospective study included 23 consecutive children (11 males, 12 females). Patients’ ages ranged from 1 to 35 months (mean age, 11 months; median age, 9 months). Fifteen patients had unilateral retinoblastoma, and the remaining 8 had bilateral retinoblastoma. Patients’ clinical data were collected from inpatient and outpatient medical records of the referral center for retinoblastoma of our institution, including sex, age, affected eyes, and ophthalmoscopic data. Ultrasonographic data were available in 18 patients with 21 affected eyes. Three patients with bilateral retinoblastoma had already been submitted to enucleation of 1 eye before the admission to the department of ophthalmology. Parents gave informed consent for MR imaging and CT scanning in all the cases. The
study protocol was approved by the ethics committee of our institution.

Two experienced ophthalmologists (T.H., S.D.F.) performed ophthalmoscopy, which always included careful examination of the ocular fundus and a search for intralesional calcifications.

MR imaging of the orbit was performed on a 0.5T unit (Gyroscan NT5; Philips Medical Systems, Best, the Netherlands) in 2 patients with 3 retinoblastomas and on a 1.5T unit (Gyroscan ACS-NT15; Philips Medical Systems) in 21 patients with 25 retinoblastomas.

In both patients studied on the 0.5T unit, unenhanced transverse spin-echo T1*-weighted images (TR, 500 ms; TE, 25 ms; thickness, 3 mm; NEX, 2) and transverse fast-field echo T2* gradient-echo images (TR, 415 ms; TE, 27 ms; thickness, 3 mm; flip angle, 15°; NEX, 3) were obtained; in 1 patient, sagittal spin-echo T1*-weighted images after gadolinium injection (TR, 500 ms; TE, 25 ms; thickness, 3 mm; NEX, 2) were obtained.

In the patients studied on the 1.5T unit, unenhanced transverse spin-echo T1*-weighted images (TR, 500 ms; TE, 20 ms; thickness, 3.3 mm; NEX, 2) and transverse turbo spin-echo (TSE) T2*-weighted images (TR, 4000 ms; TE, 120 ms; echo-train length, 11; thickness, 3 mm; NEX, 2) were obtained in all of them (25 eyes). Unenhanced transverse fast-field echo T2* gradient-echo images (TR, 376 ms; TE, 23 ms; thickness, 3.3 mm; flip angle, 15°; NEX, 3), transverse fat-suppressed turbo fluid-attenuated inversion recovery images (TR, 5000 ms; TE, 100 ms; echo-train length, 18; thickness, 2.2 mm; NEX, 3), transverse 3D TSE T2*-weighted images (TR, 3000 ms; TE, 150 ms; echo-train length, 17; thickness, 1.5 mm; gap, −0.8 mm; NEX, 1), and transverse 3D fast-field echo T2* gradient-echo images (TR, 80 ms; TE, 23 ms; thickness, 2 mm; gap, −1 mm; flip angle, 15°; NEX, 2) were obtained in 19 patients (21 eyes), 18 patients (22 eyes), 10 patients (10 eyes), and 4 patients (4 eyes), respectively. After intravenous injection of gadolinium, transverse spin-echo T1*-weighted images (TR, 500 ms; TE, 20 ms; thickness, 3.3 mm; NEX, 2) and transverse fat-suppressed spin-echo T1*-weighted images (TR, 550 ms; TE, 14 ms; thickness, 3.3 mm; NEX, 2) were obtained in 15 patients (19 eyes).

In all patients, orbit examination was performed with a surface coil (used as a receiver coil) positioned over both orbits, parallel to the coronal plane. In most patients, MR imaging was performed by using chloral hydrate for sedation (100 mg/kg of body weight).

MR images were qualitatively evaluated for evidence of intralesional signal-intensity-void spots by 2 experienced neuroradiologists (P.G., A.C.), unaware of ophthalmologic, CT, and histopathologic results. Intrallesional spots were recorded as present, absent, or uncertain. In cases of initial disagreement, the final decision was made by consensus. CT scans of the orbits were obtained in all 23 patients (Sytec and BrightSpeed; GE Healthcare, Milwaukee, Wis). Unenhanced CT of the orbits was performed in all 23 patients with 28 retinoblastomas. Contiguous 1.25- to 1.5-mm-thick sections parallel to the canthomeatal line through the orbits were obtained. An experienced neuroradiologist (C.V.), blinded to MR imaging, ophthalmologic, and histologic findings, examined the CT scans to evaluate the presence of hyperattenuating intraocular areas consistent with calcifications. Correspondence between intraocular intralesional spots on MR imaging and intraocular hyperattenuating areas on CT was then evaluated.

Histologic examination was performed in all 28 enucleated eyes. The eyes were fixed in saline-buffered formalin, sampled, embedded in paraffin, sectioned, and stained with hematoxylin-eosin.

Results

In all the enucleated eyes, the diagnosis of retinoblastoma was histologically confirmed, with evidence of calcifications in 27 of 28 eyes.

Calcifications were ophthalmoscopically detected in 12 of 28 eyes (42.85%). Ultrasonography showed posterior shadowing, consistent with calcifications, in 20 of 21 eyes (95.23%). CT showed hyperattenuating intraocular areas consistent with calcifications in 27 of 28 eyes (96.42%). Comparison between signal-intensity voids detected on each MR image and calcifications on CT is summarized in the Table.

### Correspondence between signal-void spots on MR imaging and calcifications on CT

| Sequence | True-Positive (%) | True-Negative (%) | False-Positive (%) | False-Negative (%) | Uncertain (%) |
|----------|-------------------|-------------------|--------------------|--------------------|---------------|
| SE T1    | 13/28 (46.42)     | 1 (3.57)          | 7 (25)             | 7 (25)             |               |
| TSE T2   | 17/28 (60.71)     | 1 (3.57)          | 5 (17.85)          | 5 (17.85)          |               |
| FS FLAIR | 15/22 (68.18)     | 1 (4.54)          | 4 (18.18)          |                    | 2 (9.09)      |
| GE T2*   | 21/24 (87.5)      | 1 (4.16)          |                    | 3 (12.5)           |               |
| 3D TSE T2 | 10/10 (100)      | 1 (4.16)          |                    |                    |               |
| 3D GE T2*| 4/4 (100)         | 1 (4.16)          |                    |                    |               |
| SE T1 gd | 11/20 (55)        | 1 (5.26)          | 5 (25)             | 4 (20)             |               |
| FS SE T1 gd | 9/19 (47.36) | 1 (5.26)          | 5 (25)             | 4 (20)             |               |

Note.—SE indicates spin-echo; GE, gradient-echo; TSE, turbo-spin-echo; FS, fat-suppressed; FLAIR, fluid-attenuated inversion recovery; gd, gadolinium.

Discussion

Accurate and expedient diagnosis of retinoblastoma by a careful ophthalmoscopic examination through clear optical media is usually quite easy. When refractive media are clear, the presence of large calcified areas is easily documented by ophthalmoscopy.

Sonography is performed to assess the presence of vitreal or subretinal masses, the configuration and acoustic reflectivity of such masses, and the possible presence of retinal detachment. Sonography may be also used to detect the presence of intraocular calcifications successfully; particularly, calcium has been detected in ≤92.5% of cases in the most frequent retinoblastoma forms of growth (≤95.23% in our personal records). However, sonographic capacity for detecting small calcified masses is limited by the presence of complex intraocular interfaces associated with vitreous opacities, retinal masses, subretinal fluid, and retinal detachment.
Studies such as CT and MR imaging provide valuable help in the differential diagnosis between retinoblastoma and the simulating lesions, also supplying a detailed study of intra- and extraocular structures (including the brain). Moreover, they are not impaired by those conditions that weaken sonography and ophthalmoscopic examinations.

CT and MR imaging represent the primary modalities for visual-pathway imaging in children. MR imaging is the method of choice in depicting ocular lesions, however, CT is still the method of choice for detecting intraocular calcium and investigating orbital pathologies.

The disadvantages of CT are reduced soft-tissue contrast and spatial resolution compared with MR imaging; the presence of the potential risk of iodinated contrast administration; and radiation exposure, especially in those patients with the hereditary form of the disease, who have elevated radiosensitivity. MR imaging is considered the method of choice for follow-up imaging in already diagnosed cases, especially to detect preclinical intracranial localization of the neoplastic disease. The MR imaging appearance of retinoblastoma is usually specific enough to differentiate retinoblastoma from simulating lesions; however, currently, MR imaging is not considered as specific as CT in detecting intraocular calcification.

The effect of calcium on tissue signal intensity in MR imaging is variable. Calcified tissues are far more frequently hypointense (due to diamagnetic properties of the calcium), but isointensity or hypointensity is also possible, depending on the concentration of the surface area of the calcium particles and on the MR imaging technique used. Isohyperintensity has been shown in particles with \( \approx 30\% \) calcium by weight, but it decreases at high concentrations, as in the retinoblastoma-associated calcifications.

On spin-echo T2-weighted images, detectable calcifications are far more frequently hypointense, especially when a high-field magnet and thin sections are used. One of the deficiencies of the spin-echo sequence lies in its well-recognized limited ability to detect calcification consistently.

Recent developments have introduced a very high-resolution 3D TSE T2, which allows very thin sections with high signal intensity-to-noise ratios; though this sequence allows a more accurate detection of orbital pathology, it could be more prone to motion artifacts than a conventional multisection sequence and, to the best of our knowledge, the usefulness of such a sequence in calcium detection has not been demonstrated. In our study group, the 3D TSE T2 sequence showed higher specificity in showing intralesional spots than the conventional fast spin-echo T2-weighted sequence (Fig 1).

Gradient-echo T2*-weighted imaging has been shown to be more effective than spin-echo techniques in the detection of calcified structures, typically as very hypointense regions. Because retinoblastoma is a heterogeneous tumor, the partially calcified areas may appear as hypointense foci within the higher intensity of the soft-tissue mass.

In our study, this behavior has been a significant advantage: the higher degree of signal-intensity loss of the calcifications and the lower hypointensity of the whole tumor on gradient-echo T2*-weighted images in comparison with routine TSE T2-weighted spin-echo images, has led to a superior detectability of intralesional spots in the former sequence; moreover, gradient-echo T2*-weighted images increase the sensitivity in intralesional signal intensity–loss spot detection.

Intralesional spots were detectable in 27 of the 28 eyes examined. Among these 27 eyes, in 2 patients, intralesional spots were detectable only on 3D TSE T2 sequences, whereas in 3 patients, they were detected only on gradient-echo T2*-weighted imaging. Putting together 3-mm-thick gradient-echo T2*, tridimensional TSE T2, and tridimensional gradient-echo T2*-weighted images, we found a correspondence of 100% between the presence of intralesional spots on MR imaging and the presence of calcifications on CT examinations. Even fine calcifications were detectable.

The criterion for calcifications as a localized marked hypointense signal intensity on both T1- and T2-weighted images without contrast enhancement has been sporadically described. Mihara et al.37 found spots of round signal-intensity drop-off in 3 of 7 retinoblastomas, considered calcifications on CT; recently, Lemke et al.7 showed MR imaging high sensitivity (91.7%) and specificity (88.9%) in detecting a marked amount of intraocular calcifications. Our series confirms the results of Lemke et al., adding the chance to reach 100% correspondence by using appropriate sequences and coupling neuroradiologic and clinical findings.

A limitation of our study was that the appearance of calcification at T2-weighted images is not specific: Actually, aside from calcification, hypointensity on T2-weighted images can be found at the boundaries or inside the substance of various stages of hemorrhage (ie, deoxyhemoglobin, methemoglobin, and, to the best of our knowledge, the usefulness of...
hemoglobin) and at the borders of melanocytic lesions. However, intratumoral hemorrhage in retinoblastoma is quite uncommon, whereas calcifications are present in a very high percentage of patients. Moreover, a real signal-intensity drop-off in gradient-echo images obtained on a high-field magnet is highly suggestive of calcifications, though hemosiderin or melanin may not be excluded; in our study, all areas suspected of calcifications at MR imaging were confirmed by CT examinations. Additionally, the correspondence between zones of signal-intensity loss on gradient-echo T2*-weighted images and calcifications documented on CT scans was recently demonstrated in 4 patients with Fabry disease. Finally, no areas of hyperintensity on T2-weighted images (possibly corresponding to oxyhemoglobin and extracellular methemoglobin stages of hemorrhage) have been documented in our series, also confirmed by histology in the 28 enucleated bulbs. These data strengthen the hypothesis of the correlation between drop-off signal intensity on MR imaging and calcifications.

Despite the fact that retinoblastoma and melanoma usually have different ages of presentation, both the diseases may present in children older than 7 years of age and in teenagers (personal observation). The MR imaging characteristics of retinoblastoma are quite similar to those of uveal melanoma; however, to our knowledge, MR imaging in uveal melanoma...
does not show spots of signal-intensity loss in spin-echo T2-weighted and gradient-echo T2*-weighted imaging. Furthermore, we never noticed such a sign in our personal case records. Finally, proton MR spectroscopy may allow a differential diagnosis, because the visual appearance of retinoblastoma and uveal melanoma are completely different.\(^1\)

Although CT is superior in calcium detection,\(^4\) the use of newer magnets (even if with the same magnetic field strength) allowing thinner sections may increase the specificity in calcium detection. We noticed that the use of very thin sections, both in TSE T2-weighted and in gradient-echo T2*-weighted techniques, may improve the detectability of intralesional spots. The development of a smaller diameter dedicated surface coil may allow the detection of very small intraocular lesions, even with a thickness of <1 mm.\(^2\)

Our study has clearly shown the possibility of detecting calcifications in retinoblastoma, when MR imaging, ultrasonographic, and ophthalmoscopic data are used jointly.

On the basis of the results of our study and considering the aims of orbital investigations in retinoblastoma, we think that a detailed neuroimaging study might comprise unenhanced T1-weighted, thin-section TSE T2, gradient-echo T2*-weighted (when available, 3D sequences), and fat-suppressed enhanced T1-weighted images. Fat-suppressed enhanced T1-weighted images are preferable to (in comparison with) non-fat-suppressed spin-echo T1-weighted enhanced images because of the superiority in detecting optic nerve invasion and intraorbital extension of a retinoblastoma.\(^4\)

**Conclusions**

Although intracranial calcifications have been extensively studied by MR imaging, there is not much detailed information about the capability of MR imaging in detecting intraocular calcifications.

We agree with the unquestionably superior specificity of CT in detecting calcifications; however, although brain calcification may be documented exclusively by means of neuroradiologic investigations, intraocular calcifications may also be discovered during ophthalmoscopic and ultrasonographic examination. In our 28 eyes with retinoblastoma, the combination of ophthalmologic investigations and MR imaging allowed us to make no mistakes in identifying calcifications; to the best of our knowledge, no study has coupled MR imaging and ophthalmologic investigations in detecting intraocular calcification.

We believe that if the ophthalmologist and the neuroradiologist work closely together, much more useful information on ocular structures may be obtained, including the detection...
of intraocular calcifications, without the use of potentially harmful ionizing radiation.

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