New perspectives in imaging of uveal melanoma

Malgorzata Solnik¹,², Natalia Paduszynska¹,², Anna M. Czarnecka²,³, Kamil J. Synoradzki⁴, Yacoub A. Yousef⁵, Tomasz Choragiewicz⁶, Robert Rejdak⁶, Mario Damiano Toro⁷,⁸, Katarzyna Dyndor⁹ and Michal Fiedorowicz⁴,*

¹ Faculty of Medicine, Medical University of Warsaw, 02-091 Warsaw, Poland; m.solnik98@gmail.com (M.S.), s073975@student.wum.edu.pl (N.P.)
² Department of Experimental Pharmacology, Mossakowski Medical Research Institute, Polish Academy of Sciences, 5 Pawinskiego Str., 02-106 Warsaw, Poland; anna.czarnecka@gmail.com (A.M.C.)
³ Department of Soft Tissue/Bone Sarcoma and Melanoma, Maria Sklodowska-Curie National Research Institute of Oncology, 5 Roentgen Str., 02-781 Warsaw, Poland; am.czarnecka@pib-nio.pl (A.M.C.)
⁴ Small Animal Magnetic Resonance Imaging Laboratory, Mossakowski Medical Research Institute, Polish Academy of Sciences, 5 Pawinskiego Str., 02-106 Warsaw, Poland; mfiedorowicz@imdik.pan.pl (M.F.), ksynoradzki@imdik.pan.pl (K.J.S.)
⁵ Department of Pediatric Surgery, Ministry of National Guard Health Affairs, King Abdulaziz Medical City-Jeddah (KAMC), King Saud Bin Abdulaziz University for Health Sciences, COM-J, King Abdullah International Medical Research Center, Riyadh 11481, 1513 Jeddah, Saudi Arabia; yamyousef@gmail.com (Y.A.Y)
⁶ Department of General Ophthalmology and Pediatric Ophthalmology Service, Medical University of Lublin, Lublin, Poland; rejdak@yahoo.com (R.R)
⁷ Department of General Ophthalmology, Medical University of Lublin, 1 Chmielna Str., 20-079 Lublin, Poland; toro.mario@email.it (M.D.T.)
⁸ Eye Clinic, University of Catania, Via S. Sofia 78, 95123 Catania, Italy
⁹ Department of Radiography, Medical University of Lublin, 8 Jaczewskiego Str., 20-090 Lublin, Poland; dyndorka@interia.pl (K.D.)
# These authors contributed equally
* Correspondence: mfiedorowicz@imdik.pan.pl

Simple Summary: Uveal melanoma is the most prevalent intraocular tumor in adults, derived from melanocytes, and the liver is the most common site of metastases. Due to troublesome tumor localization, different imaging techniques are utilized in diagnostics, i.e., ultrasonography (US), optical coherence tomography (OCT), single-photon emission computed tomography (SPECT), positron emission tomography/computed tomography (PET/CT), magnetic resonance imaging (MRI), fundus fluorescein angiography (FFA), indocyanine green angiography (ICGA), fundus autofluorescence (FAF). Specialists eagerly use these techniques, but sometimes precision and quality of obtained images are imperfect, which raises diagnostic doubts, which prompts the search for new ones. In this review, in addition to analyzing the currently utilized methods, we also introduce experimental techniques that may be adapted to clinical practice in the future. Moreover, we raise the topic and present a perspective for personalized medicine in uveal melanoma treatment.

Abstract: Uveal melanoma is the most common primary intraocular malignancy in adults characterized by insidious onset and poor prognosis strongly associated with tumor size and the presence of distant metastases, most commonly in the liver. Contrary to most tumor identification, biopsy followed by pathological exam is not recommended in ophthalmic oncology. Therefore, early and non-invasive diagnosis is essential to enhance patients’ chances for early treatment possibilities. We reviewed imaging modalities currently used in the diagnosis of uveal melanoma, i.e., fundus imaging, ultrasonography (US), optical coherence tomography (OCT), single-photon emission computed tomography (SPECT), positron emission tomography/computed tomography (PET/CT), magnetic resonance imaging (MRI), fundus fluorescein angiography (FFA), indocyanine green angiography (ICGA), fundus autofluorescence (FAF). The principle of imaging techniques was briefly explained, along with their role in the diagnostic process and a summary of their advantages and limitations. Further, the experimental data and the advancements in imaging modalities were searched. We described their innovations, showed current usage and research, and explained the possibilities of utilizing them to diagnose uveal melanoma and their potential application in personalized medicine such as theranostics.

Keywords: uveal melanoma; diagnosis; imaging; MRI; PET; CT; SPECT; OCT; ultrasonography
1. Introduction

The uveal melanoma is a neoplastic lesion that originates from melanocytes [1]. Although the disease is relatively uncommon, it is the most prevalent primary intraocular malignancy in adults. The annual prevalence varies between 2 per million in southern Europe to 8 per million in northern Europe [2]. Differences may be connected with increased ocular pigment content in southern populations [1]. The average incidence among Caucasians worldwide is 5.1 per million [3]. The symptoms appear in middle age, about 60 years old, and are more common in men [4]. The tumor can occur in the anterior uveal tract that contains the iris or posterior, containing the choroid and ciliary body [5]. Among uveal melanomas, even 90% derives from the choroid, 7% from the ciliary body, and 3% from the iris [3,6,7]. The main risk factors include light eye color, fair skin, and the ability to tan [1,8,9]. Fair hair color [1,8] and exposure to ultraviolet light other than welding were not statistically significant risk factors [10]. Established association between oculodermal melanocytosis, iris, and choroidal nevus, and uveal melanoma were found. Patients with present atypical and common cutaneous nevi and cutaneous freckles are also at risk [1].

At the time of diagnosis, clinically evident metastases are detected in 2-4% of all patients [11,12]. Due to hematogenous spread, the liver is the primary and the most common site of metastasis - 91% [13-15]. Other typical sites include lung (28%), bone (18%) and skin (12%) [15]. It is indicated that micrometastases are present since the diagnosis because of frequent systemic recurrences and the identification of circulating tumor cells in the bloodstream of patients who do not have clinically detected metastases [16,17]. Even with successful local treatment, over 30% of UV patients will develop metastases within ten years [18].

The overall prognosis of uveal melanoma is rather poor. Once the metastatic stage is diagnosed, overall survival varies from 3 to 16 months, with a mortality rate of 92% within two years [19]. Tumor size is one of the most important prognostic factors [1,20-22]. Small tumors (<3mm tumor thickness and <10mm basal diameter), medium tumors (3-8mm tumor thickness and <15mm basal diameter), and large tumors (>8mm tumor thickness and >15mm basal diameter) have estimated 5-year mortality of 16%, 32%, and 53%, respectively. With every one-millimeter increase in thickness, patients have a 5% bigger risk of metastasis at ten years [22]. Older age at the time of diagnosis [23], the male gender is also associated with unfavorable prognosis [20,24]. Based on tumor location, iris melanoma has the best prognostication than ciliary body melanoma [1,22]. Genetic factors such as chromosome 3, 1p loss, or 8p gain correlate with increased risk for metastasis [1,20]. When it comes to the histopathologic features, tumors dominated by epithelioid cells, with high mitotic activity and a large mean diameter of ten largest nucleoli, are linked to poor prognosis [1,20,25,26]. In the American Joint Committee on Cancer classification (AJCC), uveal melanoma is graded based on tumor basal diameter and thickness, extraocular extension, and involvement of the ciliary body. Poor prognosis was associated with the advanced AJCC stage [20].
The therapeutic methods undertaken after the diagnosis of uveal melanoma and recommended by National Comprehensive Cancer Network (NCCN) Guidelines based on radiotherapy (brachytherapy, fractionated stereotactic irradiation), particle beam radiation, and surgical treatment (Figure 1) [27]. Other techniques like laser and photodynamic therapy, cryotherapy possess occasional use [1, 18, 27, 28]. Radiation therapy is most commonly used for the treatment of UM. Brachytherapy is a form of radiotherapy containing plaques with isotopes emitting gamma radiation towards the tumor [3, 29]. During the proton beam therapy, tantalum rings are positioned at the tumor margin and constitute the border for radiation [30]. The stereotactic radiosurgery is based on radiation of gamma rays directed to circumscribed areas saving adjacent structures [31, 32]. Apart from the mentioned, conservative treatment includes transpupillary thermal therapy and photodynamic therapy [33, 34]. The extensive and advanced tumors are treated with radical methods as enucleation (a surgical extraction of the globe), exenteration (removal of an eye and ambient tissue), and uvectomy (selective removal of anterior uveal tract tumors) [1, 35, 36]. Surgical technique (enucleation) is recommended for tumors too large for brachytherapy, difficult to treat only by radiation, extensive extraocular extension, or optic nerve involvement [27]. The efficiency of chemotherapy using cytotoxic substances in hepatic metastatic lesions is insignificant [3, 35]. There are also known chemo-, immune- and radioembolization, the isolated and percutaneous hepatic perfusion [3]. Several clinical trials analyze the potential possibilities of using the monoclonal antibodies as uveal melanoma immunotherapy [37, 38].

Table 1. The summary of advantages and limitations of currently available diagnostic techniques

<table>
<thead>
<tr>
<th>Imaging method</th>
<th>Advantages</th>
<th>Limitations</th>
<th>Resolution</th>
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<tr>
<td>US</td>
<td>non-invasive wide availability elementary tool in the initial diagnostic visualization of lesion shape, structure, and vascularization good imaging of retinal detachment, vascular malformations, and hemorrhage</td>
<td>operator-dependent method diagnostic difficulties in case of muscles atypical localization or vortex veins enlargement conditions mimicking uveal melanoma like choroidal naevus, retinal hamartoma, tuberculoma, neurilemmoma tumors less than 1mm in thickness could be unnoticed</td>
<td>150 and 450 µm, 30 and 60 µm (UBM)</td>
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<tr>
<td>Method</td>
<td>Description</td>
<td>Note</td>
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<td>Ultrasound biomicroscopy</td>
<td>Good assessment of extrascleral extension and adjacent structures involvement. Ultrasound biomicroscopy as a developed method in diagnostics. Relatively low price between $155 and $721.</td>
<td>Overestimation of the tumor dimension compared to MR.</td>
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<td>OCT</td>
<td>Non-invasive, radiation-free, high spatial resolution, can provide images of tissue microstructure, can visualize small lesions in the anterior eye segment, low price of around $200 on average.</td>
<td>Small tissue depth that can be visualized, not suitable for imaging of pigmented lesions as image shadowing occur, limited use for detection and measurements of posterior eye segment tumors. 10 µm or up to 1 µm in UHR-OCT.</td>
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<td>FFA/ICGA/FAF</td>
<td>Non-invasive, radiation-free, easily accessible, possible differential diagnosis, use in treatment follow-up (FAF) or assessing the proper placement of radioactive plaque (FAF).</td>
<td>Motion artifacts, subjective and non-quantitative interpretation of images, low diagnostic accuracy when used alone. 7.4 to 5 µm.</td>
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<td>SPECT</td>
<td>Non-invasive, more sensitive than PET in primary tumor diagnosis, suitable for small uveal melanoma diagnosis, possible diagnosis in atypical manifestation or ocular complications use of melanic specific radionuclides (123I-IMP), less expensive ($1900 on average) and more available than PET/CT.</td>
<td>Accurate diagnosis is highly time-dependent (up to 48h), not suitable for amelanotic melanoma diagnosis, exposure to radiopharmaceuticals and X-ray radiation with the use of SPECT and CT hybrid. 9.3 mm full width at half maximum (FWHM).</td>
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<td>MRI</td>
<td>Non-invasive without exposition to ionizing radiation, favorable soft-tissue contrast, a convenient method to visualize intra- and extraocular involvement, good to measure the tumor used to detect distant metastases, especially in the liver used in choosing the therapeutic method.</td>
<td>Difficult to distinguish amelanotic melanoma from melanic ones choroidal metastases could mimic a uveal melanoma. Difficult to distinguish tumor from vitreous hemorrhage. Movement artifacts relatively long duration time about 20 minutes. The relatively high price of €200–€1000. 3T MRI 800 µm, 7T 500 to 650 µm and 32 µm obtained by 9.4T MRI.</td>
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<td>PET/CT</td>
<td>CT</td>
<td>FI</td>
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<td>non-invasive diagnose of the medium and large primary tumor sensitive in the diagnosis of distant metastases, especially in the liver suitable for the metastatic disease prognostication can evaluate early treatment response with higher sensitivity than MRI</td>
<td>High sensitivity, whole-body imaging, a large area covered during acquisition, moderate/high resolution, less movement artifact, scanning of a broader range of patients (with metal depositions, pacemaker), short time of scan (anxiety patients or with claustrophobia)</td>
<td>Noninvasive, safe, easy to perform, cost-effective, digital photos can be easily stored and transmitted for consultation</td>
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<td>not widely available risk of falsely positive results in inflammation, infection, trauma not suitable for diagnosis of the small primary tumor exposure to radiation resulting in increased cancer risk very expensive ($2227 on average)</td>
<td>false positives, radiation exposure, high cost of replacement X-ray tubes, time-consuming analysis of data</td>
<td>inability to access deeper layers of the retina, optical aberrations or cataract influence on image quality, 2D representation of the spatial structure (possible artifacts, peripheral aberrations)</td>
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<td>PET cameras can provide images with a spatial resolution of around 2.4 mm full width at half maximum (FWHM)</td>
<td>500–625 µm</td>
<td>Depends on CCD or phone camera resolution (14 µm per pixel)</td>
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The gold standard of tumor confirmation is biopsy followed by histopathological examination. However, this does not apply in the diagnosis of ocular tumors, including uveal melanoma, since fine-needle aspiration biopsy could lead to the seeding of tumor cells. Additionally, non-specific clinical presentation and poor prognosis, especially after diagnosing metastatic disease, show the importance of early and non-invasive identification of uveal melanoma [39,40]. In this article, we analyzed the current diagnostic options for uveal melanoma, highlighted their limitations, and reviewed the recent experimental data.
2. Imaging techniques currently used in the diagnosis of uveal melanoma

2.1. Fundus imaging (FI)

Indirect ophthalmoscopy is the primary and most important method to diagnose UM (Figure 3) [41,42]. In this technique, a direct ophthalmoscopy image of the patients’ retina appears directly on the observer’s retina [43]. This observation allows determining several defining features like the size of the tumor, retinal detachment (in tumors greater than 4 mm in thickness), presence of pigmentation, its location (distance to the foveola and optic
disc), extrascleral extension towards the anterior direction, involvement of ciliary body [42,44].

Fundus photography (FP) performed initially allows assessing retinal pathology. By FP ophthalmologist may examine the central retina, macula, optic nerve, and after mydriatics, peripheral retinal area. It is a primary method of retinal imaging [45]. Choroidal melanoma looks like a dome, mushroom-shaped or diffuse form. The color of the tumor is gray to greenish brown. Retinal detachment, necrosis, or atrophy of retinal pigment epithelium cells may occur [46]. Fundus photography allows to detection of small lesions with a basal diameter lower than 3 mm [47].

Nowadays, wide-angle systems can capture up to 200° by utilizing montage images. Wide-angle systems could observe shallow margins of tumors and are precisely like USG, but these systems are characterized by low sensitivity in detecting lesions localized anterior to the equator [46,48-50].

Imaging of fundus by photography evolve from analog film-based to digital pictures. Nowadays, images are obtained by ocular fundus cameras with digital photography. Recently, smartphone-based fundus imaging is developed (smartphone connected with digital fundus camera). Compared to conventional digital fundus photography, smartphone-based fundus imaging is a low-cost alternative to imaging retina. This equipment requires less training time for a person performing the diagnosis. Obtained images possess a good quality comparable to computers [51,52].

Due to portability, better availability, non-infrastructure requirements (continuous power supply), and obtaining by non-specialists personnel (images can be sent for further analysis for retina specialists), this technique may improve the study of more patients in the future, especially in communities with less developed medical infrastructure in primary care [53].

Photographs of the retina let to observe small lesions in diagnosing uveal melanoma but consists of a baseline. To diagnose these tumors, additional techniques like OCT or US are additionally used [54]. Alone fundus photography may be useful to monitor tumor growth, assessing therapy treatment, or arise recurrences [46,55,56].

2.2. Fundus fluorescein angiography, indocyanine green angiography, and fundus autofluorescence

Fundus fluorescein angiography (FFA) is an imaging technique used to examine circulation in the retina and choroid and provide information about the retinal pigment's blood-retina barrier integrity and characteristics of epithelium (Figure 5). Intravenous administration of fluorescein dye (sodium fluorescein) is followed by illuminating the retina with blue light at a wavelength of 465 nm. A photon of light is absorbed by an electron at its resting stage, causing its excitation. Further, as the electron relaxes to the resting stage, a photon on yellow-green light is released. This process is captured by a series of quickly obtained photographs or a film taken after dye injection [57,58].

In large uveal melanomas, intrinsic tumor circulation is paired with choroidal circulation. This double circulation, late leakage, and hot spots resulting from pinpoint leakage from retinal pigment epithelium (RPE) are characteristic features of FFA in uveal melanomas. However, this imaging modality alone lacks diagnostic accuracy [42,55]. Meyer et al. [59] assessed the diagnostic accuracy of FFA in the diagnosis of choroidal melanoma between 17% and 75%, with 58% on average. Nevertheless, bundling multiple images acquired from FFA, OCT, fundus autofluorescence could help increase the accuracy of diagnosis [60]. On the other hand, FFA is a valuable method for differential diagnosis, as it can detect hemangioma, hemorrhage, or choroidal detachment [55,61]. FFA is also used in follow-up after brachytherapy to detect complications arising from radiotherapy, such as radiation retinopathy and radiation maculopathy [41,56].
Figure 5. 30-year-old female with left melanotic choroidal melanoma (A). Clinical exam showed RPE hyperplasia and metaplasia indicating underlying previous choroidal nevus (B). The lesion has orange pigment lipofuscins that showed fundus auto fluorescence (C), and the mass was associated with intrinsic vascularity in FFA (D).

Indocyanine green angiography (ICGA) is similar to FFA, except that the fluorescein dye used in this method is indocyanine green. Its peak absorption is at around 790 nm, with a peak emission of 835 nm. These properties allow penetration of retinal layers, melanin, macular pigment making it possible to visualize structures beneath RPE. Indocyanine green binds to plasma protein in about 98%, making it impossible for the dye to exit the bloodstream through choriocapillaris fenestrations, allowing visualization of not only large and medium vessels but choriocapillaris as well [60,62].

ICGA provides greater visualization of tumor vasculature as it can imagine microcirculation of uveal melanoma, especially when there is overlying blood [42,44]. Variable patterns of ICGA (hypofluorescent, isofluorescent, and hyperfluorescent) depend on tumor thickness and degree of pigmentation [61]. These non-specific patterns disqualify ICGA as an accurate diagnostic modality; however, ICGA can potentially differentiate pigmented and non-pigmented choroidal melanomas and choroidal hemangiomas. Non-pigmented tumors compared to pigmented tumors show earlier onset of fluorescence (<1 min vs. 3 min), and choroidal hemangiomas reveal characteristic findings on ICGA, including early onset of fluorescence and early maximum fluorescence followed by “washout” of the dye in later frames [55,63]. Application antireflection coated lens and indium iodide lamp allow to obtain 7.4 µm resolution [64]. It should also be mentioned that ICGA could induce phototoxicity on RPE cells, and unnecessary use should be avoided [65].

Fundus autofluorescence (FAF) is an in vivo imaging technique that allows visualization of stimulated emission of light coming from fluorescence chemical compounds (fluorophores) found in the ocular fundus with fundus camera containing specific filters [60].
The most common fluorophore is lipofuscin that accumulates in RPE cells and is seen as orange pigment on indirect ophthalmoscopy [66]. Secondary epitheliopathy that is common in malignant choroidal tumors causes the shedding of photoreceptor outer segments. Combined with high phagocytic activity, this process increases lipofuscin accumulation seen in uveal melanomas [60,66].

One of the predictive risk factors of small melanoma is an overlying orange pigment visualized with FAF. Therefore, FAF could play a role in the early detection of small choroidal melanomas and differentiation between choroidal melanoma and nevus [67]. Choroidal melanomas demonstrate clumps of hyper-autofluorescence, whether most of the choroidal nevus is iso-autofluorescence or hypo-autofluorescence [42,44,66]. Additionally, blue light FAF can be used to confirm the proper location of radioactive plaque used in local treatment [44]. Finally, all of these fluorescence-based modalities are dependent on the patient’s ability to stay still to avoid motion artifacts and experience in the interpretation of acquired images [57,60]. Fundus autofluorescence images can be obtained by confocal scanning laser ophthalmoscope with blue light excitation or fundus camera with appropriate filters. These methods allow to achieve 5 or 14 µm per pixel resolution, respectively [68].

2.3. Ultrasonography

Ultrasonography (US) is a diagnostic tool based on sending and receiving sound waves (with a frequency greater than 20000 Hz, inaudible to humans) with a probe (containing piezoelectric transducer) which is both a transmitter and receiver of waves (Figures 3-4, 6-8) [69]. In ophthalmology, there are two types used, A-mode scan to carry out measurements or movability assessment and B-mode scan to visualize intraocular structures of different echogenicity as a view grayscale created based on reflected waves [70]. These modes use different frequencies, the A-mode 8 MHz and the B-mode 10 MHz [71].

Uveal melanoma on A-scan has low to medium reflectivity and presents a positive angle kappa sign, which means that the spikes are high and decrease toward the sclera [1]. On B-scan uveal melanoma is a homogenous mass with a low acoustic profile [72]. Because the eye has high fluid content, it provides an acoustic window and facilitates relatively detailed visualization. If the tumor presents rich vascularization, the pulsations can be registered [73]. The mass can resemble a mushroom shape, and it is synonymous with Bruch’s membrane disruption [54]. Ultrasound can facilitate the assessment of adjacent structures involvement because the extraocular area has a higher reflectivity than neoplasm, so infiltration is visible as regions of hyporeflectivity beyond the physiological limits of the sclera, which are marked by echoes from orbital fat [56,74,75]. Thus, the US is suitable for assessing the tumor extrascleral extension and visualizing lesions even 2 mm [74]. Moreover, it enables the determination of lesion shape and general structure [76]. The average cost of the US in the world is estimated between $155 to $721 [77].

The classic US allows to evaluation neoplasm of the anterior segment. Still, ultrasound biomicroscopy (UBM), which uses higher frequencies (from 35MHz to 50MHz), facilitates the visualization of lesions derived from the ciliary body, tumor progression to posterior regions to the iris, and differentiation of cystic or solid nature of the lesion [55]. UBM with application of a higher frequency compare to B-scanners, allow to obtain better axial and lateral resolution 30 and 60 µm (150 and 450 µm in B-scanner) [78]. Due to increased resolution in UBM, it is possible to reveal the posterior margin of the tumor and evaluate the disease extension [79]. The disadvantage is the relatively low availability of that method.
Scott et al. [80] also compared the effectiveness of US, magnetic resonance imaging (MRI), and computed tomography (CT) and revealed 100% sensitivity for an ultrasound. Martin et al. [81] suggested that the US should be an elementary tool in the initial diagnosis of choroidal melanoma and control during the treatment. Because of high ultrasound penetration compared to light through the ocular tissues, B-mode imaging is considered a more effective diagnostic process tumor than anterior-segment optical coherence Tomography (AS-OCT) [1]. The retinal detachment, a concomitant condition with uveal melanoma, could be successfully detected by the US because of the accumulation of subretinal fluid which splits the layers (choroid and retina), which generally are seen as integrated and that also enables the visualization of choroidal growth located behind the damaged retina [82]. Other conditions apart from tumors or retinal detachment as hemorrhage and vascular malformations, can be successfully detected with US [83].

The diagnostic process of the neoplastic lesion could be complicated when it is located close to the optic nerve if the ocular muscles are atypically situated or in the case of vortex veins enlargement [75,81]. The ocular disorders which sonographic appearance is similar to melanoma are choroidal nevus, retinal hamartoma, tuberculoma, or neurilemmoma [55]. The tumor thickness is a relevant limitation of the B-scan method because lesions <1mm could be unnoticed [54]. The size measurements are performed with a high frequency of about 10 kHz probe [84]. The significant limitation is that the higher frequency used, the higher the image resolution, the greater wave attenuation, and the reduced penetration depth. Due to the accuracy and precision of the method, the tumor sizes could be overestimated by the 2D US compared to MR with the average difference value of 1mm [84]. In research studies, three-dimensional US facilitates the analysis of extent scale, but it is not performed in clinical practice [85].
Figure 7. This left eye had amelanotic choroidal melanoma (A) and B-scan showed a 7.7 mm thick dome shape tumor with moderate low internal reflectivity. 6 months after treatment by I-125 Radioactive plaque, the tumor showed regression clinically (C), and thickness decreased to 4.2 mm (D).

Color Doppler (CDI) is a non-invasive method that shows blood flow in vessels considering flow direction. Doppler function (CDUS) could serve as a tool to evaluate the response to radiation treatment because the untreated lesions characterize richer vascular patterns than untreated ones [55]. Although ultrasound is widely available, it is operator-dependent method, and only two-dimensional images can be obtained while other methods allow making reconstructions [84]. The other disadvantage is no penetration of tissues like bones and poor imaging of tissues around the perimeter of the wave beam [86].

2.4. Optical coherence tomography

Optical coherence tomography (OCT) is a diagnostic method analogous to the US, which essence, is receiving and processing light waves (with a near-infrared wavelength) to create cross-sectional images (Figures 3-4) [87]. It uses the interferometry technique; the light is split into a reference beam and sample beam and then reassembled. Echo time delay and reflected or backscattered light intensity are being measured to provide images of tissue microstructure. With depths of 1–2.5mm into the tissue, OCT can provide images with a spatial resolution of 10 µm or up to 1 µm in Ultra-High Resolution OCT (UHR-OCT), the most advanced form of OCT used currently only for academic and research purposes [88]. With a low cost of around $200, this method is also safe as no adverse events occur and no radiation is used [89].
Figure 8. Patient with oculo-dermal melanosis (A) was found to have retrolenticular melanotic mass (B), which was found to be large choroidal melanoma (13 mm thick) with ciliary body involvement (C). This eye was enucleated and shown to have scleral melanosis (D).

The two main technologies used to create an OCT image are the time-domain (TD-OCT) and the Fourier domain (FD-OCT), also known as the frequency domain. The TD-OCT was developed first, and its main limitation came from the need to move the reference mirror with increasing speed to measure the light echoes coming in sequence. Later developed, FD-OCT has a reference arm with a static mirror. It allows fast scanning as well as improved image quality. Spectral-domain OCT (SD-OCT) evolved from FD-OCT and significantly increased acquisition speed up to 100-fold time than TD-OCT, which in consequence can reduce motion artifacts and enhance spatial resolution [90].

The OCT is a non-invasive and non-contact method used in the diagnostic process, choosing therapeutic decisions and treatment control [56]. AS-OCT allows real-time imaging of anterior ocular segment elements such as the cornea, anterior chamber angle, sclera, iris, and lens. When it comes to pigmented tumor detection, iris melanoma, iris nevi, and iris melanocytoma can be detected by this method [55]. The conjunctival melanoma is visible on OCT as a hyperreflective structure covered with hyperreflective epithelium regardless of whether the lesion is melanocytic or not [91]. AS-OCT is a convenient technique to differentiate conjunctival melanoma from the other types of tumor and ocular surface squamous neoplasia [92]. OCT can also efficiently differentiate melanoma and melanocytoma (retinal compound and special grow pattern) [93]. According to studies, OCT could serve as a tool for differentiation conditions such as choroidal metastasis, choroidal osteoma, hemangioma, and retinal tumors [94].

Optical coherence tomography angiography (OCTA) is a new non-invasive imaging technique with the ability for both, structural and angiographic analysis in vivo, without the use of dye injection [95-98]. Recently, Neroev et al. [99] have shown the role of OCTA may be used in complex diagnosis of early UM and circumscribed choroidal hemangioma for detection of tumor’s own vessels and the nature of their branching, as well as for vessel caliber comparison. Additionally, an increase in the frequency of detection of tumor’s own
vessels will make possible early differential diagnostics of a malignant or benign tumor and may help in establishing an adequate conserving therapy. In OCT angiography, the choroidal vascular flow rate in choroidal melanoma was significantly lower than that in choroidal nevus. Ghassemi et al. [100] found a decreased flow rate of the surface microvasculature (SMV) of choroidal melanoma cases when compared with nevi. In their study, Valverde-Megías et al., have shown that eyes with choroidal nevus demonstrate similar central macular thickness (CMT), foveal avascular zone (FAZ) area and capillary vascular density (CVD) when compared to the controlateral eye. In contrast, eyes with melanoma show increased CMT, enlarged FAZ, and reduced CVD, particularly related to increasing tumor thickness. [101].

According to some authors [102-104] OCTA measurements can provide quantitative biomarkers for early detection of radiation retinopathy (RR) and/or radiation optic neuropathy, following brachytherapy in patients with UM. OCTA provided a quantitative measurement of retinal capillary changes associated with ischemia that correlated with visual acuity and radiation dose and may predict future development of radiation-induced retinal toxicity or as a quantitative endpoint to address visual prognosis. Therefore, Matet et al. [105] have shown as visual acuity of eyes with radiation maculopathy is influenced by structural and microvascular factors identified with OCTA, including FAZ area and DCP integrity.

Jing Yan Yang et al. [106] have investigated macular microvascular characteristics imaged by optical coherence tomography angiography (OCTA) in uveal melanoma (UM). Patients were treated with Conbercept injections precede by plaque radiotherapy. According to the results showed by the authors, OCTA may provide a quantitative evaluation of early retinal microvascular changes following radiotherapy. Despite its recent progress, image artifacts on OCTA are commonly encountered and appear to be more frequent in eyes with pathology and poor visual acuity, so limiting its use. Recognition of these artifacts might help improve image interpretation and decision making [95].

The important limitation is image shadowing occurring in the case of lesions that contain pigment [79]. The light penetration is limited by the sclera or iris epithelium, and AS-OCT is preferable for lesions located superficially and without pigmentation. In other cases, ultrasound biomicroscopy could be a more beneficial choice [107]. However, in a study by Torres et al. [94], all the lesions thicker than 1mm, which was not detected by the US, were revealed using Enhanced Depth Imaging (EDI-OCT) technique. Pavlin et al. [108] described AS-OCT as a valuable tool for detecting small hypopigmented iris tumors. However, with pigmented iris tumors, OCT visualized only the surface of lesions; their posterior margin was not detected, making thickness measurement impossible. The same happened with large hypopigmented iris tumors as their image faded out with deeper layers of the lesion. Irido-ciliary tumors could not be thoroughly examined with AS-OCT as with provided image. One could only assume the presence of a tumor, but no details can be discerned.

2.5. Magnetic resonance imaging

Magnetic resonance imaging involves using the magnetic properties of atomic nuclei with odd protons or neutrons numbers, most frequently protons [109].

The images obtained by this method depend on the imaging modality, e.g., PD-weighted (PDw, proton density-weighted), T1-weighted (T1w, longitudinal relaxation time-weighted), T2-weighted (T2w, transverse relaxation time-weighted), or diffusion-weighted imaging [110,111]. This method is also highly suitable in experimental settings (Figure 9), e.g., for studying animal models of uveal melanoma, allowing qualitative evaluation of the images and quantitative measurements of the ocular dimensions [112]. The value of the static magnetic field generated by the devices in clinical practice is typically 1.5 or 3 Tesla, but in experimental settings, the values may exceed 7 Tesla [39]. The greater the magnetic field used, the more profitable the signal-to-noise ratio and the shorter acquisition time [110,113]. MRI was shown to provide a more accurate measurement of the
tumor dimensions than US; in some cases, MR diagnosis would allow eye-preserving therapy [114].

Good soft tissue contrast creates a favorable condition to visualize intra- and extra-orbital compounds, tumor localization to assess tumor dimension, and the involvement of adjacent tissues [115,116]. Nowadays, the acquired images influence therapeutic decisions, evaluating tumor advancement, prognosis, and response to treatment [117]. The irrefutable advantage of MR is that patient is not exposed to ionizing radiation and avoids its potential complications. GrechFonk et al. estimated the average cost of MRI at €200–€1000 (the price is lower in Asia, Australia, Europe, and higher in the USA; Table 1, Figure 1). They suggested that, although the price is not lower than ultrasonography, it is still significantly lower than the possible costs of therapy [84].

![Figure 9. T2-weighted MR images of the eyes of experimental animals – (A) mouse and (B) rat. High-resolution images enable detailed evaluation of the ocular anatomy.](image)

The presence of orbital adipose tissue enables relatively good tissue contrast [110]. The characteristic features of uveal melanoma obtained in MR (Figure 10) compared to the vitreous (because of the high-water content in the vitreous body, it is hypointense in T1w and hyperintense in T2w) include hyper-intensity on T1-weighted images, hypo-intensity on T2-weighted images, hyper-intense on PD images, and homogenous enhancement with gadolinium contrast [117-119]. Tissue contrast results from T1- and T2 – times shortening by unpaired electrons of melanin, and this property correlates with the quantity of melanin [110,120-122]. An uneven melanin distribution can contribute to inhomogeneous intensity character, similar to the disturbed vascular system of the tumor [110,116]. However, in the case of amelanotic melanoma, the image analysis can be troublesome because the T2 relaxation time will not be shortened, so the amelanotic melanoma cannot be distinguished from melanotic melanoma by this technique [75,123].

![Figure 10. T2-weighted MR images of the eyes of experimental animals – (A) mouse and (B) rat. High-resolution images enable detailed evaluation of the ocular anatomy.](image)
In certain situations, the recognition of uveal melanoma may be problematic due to conditions that look similar. For example, the choroidal metastases could mimic a uveal melanoma [117]. Furthermore, the differentiation between uveal melanoma and other pathologies as vitreous hemorrhage also is challenging basing on T1w or T2w imaging [117]. Fresh blood could be isointense or hypo-intense on the T1w or T2w image, and the hyper-intensity will occur after a few weeks [123]. The differences of blood in MRI appearance depend on the oxygenation of hemoglobin, oxy-, deoxy- or methemoglobin present at different stages of hemorrhage. Because of the neoplasm extension, a common concomitant condition is a retinal detachment, which could be confirmed by no contrast enhancement reaction (unlike the tumor). Still, without the use of a contrast agent, these lesions look very similar [117].

In a study by Mafee et al. [110], among 21 patients with suspected uveal melanoma and other complications as hemangioma (less contrast enhancement than melanoma), choroidal or retinal detachment, magnetic resonance imaging was an effective technique. Stroszczynski et al. [124] suggested that MR could be an even more favorable method than the computed tomography. But compared to CT, MRI is less appropriate in the visualization of calcifications [125]. According to Rescan et al., MRI is characterized by 100% sensitivity in detecting scleral involvement and extrascleral propagation with 50% and 89% specificity, respectively [126,127].3T magnetic resonance allows to acquire resolution 800 µm approx., 7T apparatus 500 to 650 µm [84,128]. More accurately (32 µm) was obtained by high-spatial-resolution MRI at 9.4 T [129].

The movement artifacts resulting from involuntary eye movements are a significant obstacle in deteriorating the image quality [130]. During the procedure, drifts, tremors, saccades or blinking may occur, thus it is important to restrict the movements and one of the possible counter measure is the application retro- or parabulbar anesthesia [131]. Different devices that aim to ensure the immobilization of the globe contribute to magnetic field disturbances in the path between air and eye surface due to the permanent opening of the eyelids during the imaging process [132]. The best solution to limit artifacts would be to shorten the acquisition time, but it would negatively affect the image quality [117]. The surface coil is placed on the patient’s head to improve the resolution properties but with relatively low accuracy in visualizing the deeper orbital tissues because the signal intensity decreases with the distance from the coil [117].

Outside of assessment of primary tumor, MR imaging is used to detect distant metastases, especially in the liver. Marshall et al. [133] showed that a six-monthly MRI could detect around 90% of hepatic lesions before they become symptomatic, enhancing early treatment possibilities in the group of patients with high-risk uveal melanoma. Similarly, Servois et al. [134] assessed MRI and 18F-FDG PET applicability in preoperative staging of hepatic metastases from uveal melanoma. The study showed the superiority of MRI over 18F-FDG PET in this regard, with differences marked especially in lesions smaller than 10 mm. Correspondingly, in the study conducted by Francis et al. [135], MRI presented accuracy in detecting liver lesions and determining their nature in newly diagnosed uveal melanoma patients.

Additionally, MRI serves as an important tool in the choice of therapeutic method and provides information about anatomical relationships before brachytherapy, proton beam, and stereotactic radiotherapy [54,136]. But considering patients with less than 6mm or more than 8,5mm tumor thickness, it turned out that MRI does not introduce any significant changes into their therapeutic course [84]. The duration of a single test depends on the parameters of the device and the protocol that was used.

MR imaging can also be essential in patients who underwent vitrectomy. In these cases, the replacement of the vitreous body with silicon oil (SiOil) tamponade, causes difficulties for UM diagnosis and assessing treatment follow-up with imaging methods such us US, where sound waves reflect at the SiOil-water interface [137]. SiOil by principle does...
not hinder MR imaging [137,138], although some strong artifacts may appear in the conventional protocols as the result of the off resonance of the SiOil. Dedicated MRI protocols such as 3 T and 7 T might be able to provide high-resolution images of vitrectomized eyed with SiOil tamponade [137].

The basic protocol needed to assess the tumor dimensions includes T1- and T2-weighted images before and after the application of gadolinium contrast agent and lasts about 20 minutes [139]. To gain some additional and more precise information, the other sequences could be used. The GE (Gradient Echo) sequences characterize the neoplasm structure, and the SE (Spin Echo) and fat suppression sequences image the scleral boundary [117]. The multiplanar reconstructions provide measurement possibilities in various planes and reveal possible propagation beyond the sclera similar to post-contrast T1-weighted scans with fat suppression, and the isotropic sequences visualize the tumor boundaries [117]. The reconstructions of 3D TSE (turbo spin-echo) sequences are desirable in disturbances due to eye motion. Another modality is Diffusion-Weighted Imaging (DWI) which is based on measuring water molecules’ Brownian motions depending on cellularity or the presence of edema to generate contrast on MR images. DWI may differentiate benign or malignant tumors and use ADC (Apparent Diffusion Coefficient) to analyze prognostic response to treatment by the proton beam [136,140]. A significant drawback of DWI is high susceptibility to motion artifacts. So future studies are desirable to develop a solution reducing artifacts and shortening the acquisition time.

Some other promising MR modalities include perfusion-weighted imaging with gadolinium CA clinically used in almost all oncological MR exams and has a specific UM value. Such an approach can differentiate between UM and retinal detachment [141,142]. 3D imaging may also be superior to US [143] and be used for radiotherapy planning [144]. Another remarkable approach is diffusion-weighted imaging used for differential diagnosis [140] and treatment follow-up [145].

2.6. Computed tomography

CT uses X-ray beams that cross the human body and are received by the detectors. Computer algorithms align obtained scans. The patient is surrounded by a ring of X-ray detectors opposite to them, localize the X-ray tube. The X-ray tube emits N photons per unit of time. Emitted X-ray beams passed by biological tissues attenuate a portion of radiation (this also causes a decrease in photons and energy). Detectors receive the signal and convert it into electrical pulses sent and analyzed by a computer. In the final result, data obtained by the scanning process contains cross-sectional images of the human body. Each image consists of volume picture elements called voxels. Each voxel contains information on attenuation and localization. The attenuation is measured by Hounsfield units (HU), and on the image, it corresponds to gray, but grayscale cannot be converted directly to HU [146,147].

Modern CT systems develop to several special techniques. MDCT methods (image segmentation using manual or semi-automatic approaches) allow determining the volume of objects on CT scans. It may apply in the diagnosis of osteoporosis, loss of the skeletal muscle, measuring adipose tissue, or organ volumes. CT allows localization of the lesion and its impact on adjacent tissues. Knowing of precise location facilitates planning surgical approach. CT angiography (CTA) allows assessing a vascular tree applicable in stroke evaluation, aneurysm location, size, or neurosurgery. MDCT combined with PET can highlight amyloid depositions in imaging Alzheimer’s disease [146,148,149].

Computed tomography is rarely used in UM diagnosis. Other techniques as MRI or US possess a better precision and are commonly used [44]. By CT scans, it is difficult to distinguish UM from retinal detachment or exudative macular degeneration. CT images give unambiguous results of examining epi- and extrascleral growth (e.g., in one case not lesion but macular degeneration type Junius Kulmt reveal by fundoscopy). CT enables easy distinguish vitreous from the lesion (possess high radiographic density), but small uveal melanoma lesion was impossible to detect. It is difficult to precisely determine the
size of the tumor and the minimum measurement of a lesion. Measurements of the tumor were very inaccurate (too small, too big, or tumor was not visible) compared to the US or histological examinations [150]. In CT scanning systems for eye and orbit, approximately 1 mm thick slice scan, in most CT scanners spatial resolution is 500–625 µm despite the thinner than 2 mm tumors, could not be observed [149,151,152].

CT is nonspecific. Lesions are difficult to recognize due to the high attenuation of the signal by the choroid. MRI imaging is better than CT because melanin has intrinsic T1 and T2 shortening effects, allowing to distinguish the presence of tumor lesions [122].

CT has been used to confirm diagnostic performed previously by echography or the US to confirm the extra-scleral and extraocular extension [80,81,153]. It also may be useful to observe bony orbital expansion or scalloping. It also may show small calcifications [154]. CT may help in the diagnostic process in eyes with secondary vitreous hemorrhage or cataract [54].

Computed tomography may be used for accurate staging when metastasis occurs. CT and PET are used to assess liver and extra-liver (e.g., bone, lymph nodes) metastatic choroidal melanoma [155].

2.7. Single-photon emission computed tomography

Single-photon emission computed tomography (SPECT) uses the gamma-emitting radiopharmaceuticals delivered intravenously into patients’ bodies and records their emission from many angles by the nuclear camera to provide 3D images [156]. Radionuclides used in SPECT emit one gamma-ray photon, which differentiates them from radionuclides used in positron emission tomography (PET), which emit two photons created after positron-electron encounter [157]. The nuclear cameras rotate around the patient while gamma photons radiating from the patient body pass through the collimator and finally hit the detector plane. After detectors collect all the data (location of the interaction, photon energy), they can create an image of the tissues [158]. Spatial resolution is dependent mainly on the type of used collimator as it can improve with the use of a collimator with smaller and longer holes. A 9.3 mm (FWHM) resolution can be achieved using all/general-purpose collimator type [156].

SPECT using Technetium-99m-MIBI accumulation in tumors for evaluating ocular malignant lesions in patients allows detection of lesions in sizes 9.5 to 12 mm [159].

Similar to PET, gamma photons are being emitted from sites with high radionuclide accumulation. Radiotracers used in SPECT include$^{99m}$Tc, $^{201}$Tl, $^{67}$Ga, or $^{123}$I. All of them have a long half-life time that varies between several hours and days, allowing a long imaging time and making the diagnostic accuracy highly time-dependent [158].

N-isopropyl-p-$^{123}$Iodoamphetamine ($^{123}$I-IMP) is a radiopharmaceutical used initially as a tool for blood perfusion imaging. Later it was found $^{123}$I-IMP accumulates in areas with increased melanin production [160]. Goto et al. [161] found $^{123}$I-IMP SPECT is a suitable diagnostic method for uveal melanoma with atypical manifestations or for its diagnosis when ocular complications (cataract, hemorrhage, etc.) are present. In the study, a significantly high intake of $^{123}$I-IMP in an adequate tumor location was observed in 25 out of 27 (92.6%) patients with suspected uveal melanoma who did not undergo any treatment. The smallest found lesion was 3mm x 4mm. In the same study, 35 out of 36 (97.2%) patients without significant $^{123}$I-IMP accumulation were diagnosed with different intraocular disorders. In another study, $^{123}$I-IMP SPECT was done 24h after the intravenous administration of radionuclide in a group of 19 patients. In all the 12 (100%) patients with increased uptake of $^{123}$I-IMP, uveal melanoma was diagnosed. However, only 2 of them were SPECT positive three hours after the $^{123}$I-IMP application. In the remaining seven results, one was falsely negative (85.7%). It was also shown that the $^{123}$I-IMP SPECT method was more sensitive and accurate than 18-fluoro-2-deoxy-D-glucose ($^{18}$F-FDG) PET in uveal melanoma diagnosis [162].

Abe et al. [160] evaluated the degree of usability of $^{123}$I-IMP SPECT using the uptake index (UI) and increased rations (IRs) as the defining tool. The accuracy of the $^{123}$I-IMP
SPECT method improved with time after radionuclide administration. At 48h time point and with a UI cut-off value of 1.90, sensitivity and accuracy were 100% and 95%, respectively. Moreover, calculated UI and IRs in uveal melanoma patients were significantly higher than in the group of patients without uveal melanoma. In a recent study, Kamazaki et al. [163] assessed SUV in SPECT/CT hybrid images to provide comparable UI evaluation results. Moreover, the SUV-based method evaluated at a six h time point can forecast the degree of $^{123}$I-IMP uptake at a 24h time point and is more predictable than the UI evaluation method.

SPECT is superior to PET in costs, as SPECT has valued in the US $1900 [164] on average and its availability [165]. However, due to the use of radionuclides in SPECT and even additional radiation coming from X-rays in SPECT/CT, increased cancer risk is a significant hazard [166].

2.8. Positron emission tomography/computed tomography

Positron emission tomography is a diagnostic imaging technique that visualizes areas based on their affinity and biodistribution of administered radiotracers [167]. After administering radiopharmaceutical is distributed in the tissue, it emits a positron that interacts with an electron, creating two photons emitted in virtually opposites directions. This interaction is detected and estimated within a ring of detectors placed around the patient [168]. It allows visualization as well as the quantification of said biodistribution in a three-dimensional reconstruction [167]. PET cameras can provide images with a spatial resolution of around 2.4mm full width at half maximum (FWHM) [169]. The most commonly used radiotracer, $^{18}$F-FDGallows detecting tumor cells because of their high glucose uptake [170].

Consequently, PET has been widely used in oncology to detect tumors, metastases, evaluate treatment response, and detect any recurrences due to its current availability and effectiveness [11]. However, high glucose avidity is also related to inflammation, infection, or trauma. Some treatment options, such as radiotherapy, can also cause high glucose uptake up to 3 months after the procedure is performed [171]. With the combination of computed tomography and PET, information about metabolic abnormalities is anatomically accurate, making this method much more accurate [39,170].

For semiquantitative image analysis, standardized uptake volume (SUV) is used. It shows the correlation between tracer activity in tissue (microcuries per gram) and injected radiotracer dose (millicuries) as well as patient weight (kilograms). Typically, the SUV value above 2.5-3.0 indicates malignant tumors. The SUV varies based on the time between radiotracer administration and PET imaging, making it important to standardize the time interval in PET/CT scans, especially when used for treatment response evaluation [172].

For primary tumor detection, Reddy et al. [173] found 33%of the AJCC T2 staged melanomas,75% of T3 staged, and none of the T1 stages (with SUV cut-off value of 2.5). SUV correlated positively with the tumor volume [174]. In a different study, Matsuo et al. [175] observed high glucose uptake in nodular choroidal melanomas while diffusely infiltrating tumors were not detected. It was also shown that higher SUV$_{\text{max}}$correlated with greater thickness of the tumor. McCannel et al. [176] found correspondence between metabolic activity on PET/CT scan and monosomy 3 (54% sensitivity, 100% specificity). Correlation between positive PET/CT imaging and increasing tumor size along with chromosome 3 loss was also described by Papastefanou et al.[177].High SUV value was found in older patients with lesions presenting larger diameters linked to poor prognosis and greater metastatic risk [174]. According to those findings, PET/CT seems to be a useful tool for predictive purposes, but not for primary tumor diagnosis, as if it may not be able to detect small uveal melanomas or differentiate them from uveal nevi [167,173].

The main role for uveal melanoma patients $^{18}$F-FDG PET/CT is detecting distant metastases [170]. Both Klingenstein et al. [178] and Kurli et al. [155] reported 100% sensitivity, 100% specificity of this method in detecting liver metastases. However, in a different
study, Strobel et al. [179] reported PET/CT identified only 16 of 27 (59%) known liver metastases. Freton et al. [17] described whole-body 18F-FDG PET/CT as a valuable method for initial staging in patients with UM in a retrospective study. Hepatic metastases observed on PET/CT scans were confirmed via biopsies, suggesting a 100% positive predictive value. Extrahepatic lesions (lungs, bone, lymph node, brain, spleen) also were found even though blood tests and other imaging modalities did not indicate the presence of metastatic disease in those sites. Furthermore, PET/CT seems to be valuable in assessing early therapy response. In response to the administration of chemotherapy, PET/CT showed decreased metabolic activity in lesions even though their size remained the same, according to MRI [180]. However, DW MRI can also show an early treatment response [181].

PET/CT is an expensive imaging technique as its cost in the US varies between $429 and $2933 (average of $2227; Table 1, Figure 1) [182]. As well as other imaging methods using radiotracers, PET/CT scan is associated with a significant radiation dose and can increase cancer risk, so the risk-benefit ratio should be evaluated beforehand [183].

3. New and experimental imaging techniques with the potential for diagnosing uveal melanoma

3.1. New radiolabeled radiotracers in PET

As of now, 18F-FDG is the most common radiotracer used in PET in oncology imaging. However, false-negative results are seen in well-differentiated tumors or lesions with low metabolic rates, and false-positive results can be observed in areas with active inflammation. Over the years, the understanding of cancer cell biology was becoming more advanced, and processes outside the glucose uptake have been discovered. Therefore, new non-FDG radiolabeled radiotracers have emerged to overcome 18F-FDG limitations and better imaging capabilities for certain types of cancer [184,185].

The oldest non-FDG tracer is bone-specific sodium fluoride labeled with fluorine 18 (18F-NaF). Fluorine ions bind to the surface of hydroxyapatite proportionally to the bone remodeling and bone blood flow, indicating undergoing osteolytic or osteoblastic processes. [184,186].

All of this led to the use of this radiotracer in PET alone or PET/CT combination to detect and assess skeletal metastases (Table 2). Most studies were conducted on 18F-NaF-PET concerning detecting bone metastases in prostate, breast, lung, and thyroid cancer [186]. In a meta-analysis, Liu et al. [187] compared 18F-NaF PET/CT and 99mTc-methylene diphosphonate (MDP) planar bone scintigraphy (BS) in bone metastases diagnosis. On the patient basis, sensitivity was higher when equivocal results were described as positive (96% vs. 93%), but specificity was higher when equivocal results were considered as negative (95% vs. 93%). On the lesion basis, both sensitivity and specificity showed similar results to those based on the patient basis. Compared to 99mTc-MDP BS, 18F-NaF PET/CT showed higher diagnostic accuracy in bone metastases detection with superior sensitivity and specificity. Similar results were shown in a meta-analysis by Shen et al. [188] as well. Although this modality has not been used yet directly in the imaging of uveal melanoma, it can be applied for bone metastases detection, which is the most common site outside of liver metastases, or for assessment of extrascleral extension of the lesion.

In melanoma, available imaging methods have limited capability to detect stage I and II melanoma and small distant metastases. Consequently, new specific radiotracers that can bind to melanin have been developed, including radiolabeled antibodies, benzamide (BZA), and benzamide analogs [189,190]. Although presented below studies focus on cutaneous melanoma, highly specific melanin tracers should also be considered for uveal melanoma diagnosis and to detect distant metastases.

One of the more recently introduced melanin-targeted radiotracers is 5-bromo-N-(-2-[diethylamino]ethyl) picolinamide labeled with 18F (18F-5-FPN). This benzamide derivative is characterized by a high affinity to melanin and quick renal clearance [189,190].
the preclinical study conducted by Feng et al. [189] potential of 18F-5-FPN PET in diagnosis and staging of melanoma was evaluated in vivo in B16F10 tumor-bearing mice. 18F-5-FPN PET displayed high effectiveness in detecting primary tumors and simulated small metastases (1-2mm) in the lungs.

Compared to 18F-FDG PET, the tumor-to-background ratio in 18F-5-FPN PET was proven to be ten times higher. Nevertheless, a significant disadvantage of 18F-5-FPN PET is the hepatic retention that can reduce the efficacy of this method in detecting hepatic metastases. Similar results were obtained in the study by Wang et al. [190]. Models of lymph node and pulmonary metastases were created in C57BL/6 mice and then evaluated with both 18F-5-FPN PET and 18F-FDG PET. On 18F-5-FPN PET scans, small metastases in both lymph nodes and lungs were more visible, and the tumor-to-background ratio was significantly higher than 18F-FDG PET.

Evaluation of 18F-6-fluoro-N-[2-(diethylamino)ethyl] pyridine-3-carboxamide (18F-MEL050), melanin-specific benzamide analog for the imaging of primary and metastatic melanoma was conducted by Denoyer et al. [191]. In that study, an 18F-MEL050 PET scan was performed on melanoma model - C57BL/6 mice inoculated with cell lines of B16-F0 and B16-BL6 murine melanotic melanoma. 18F-MEL050 showed a clear delineation of melanoma lesions with rapid background washout and higher tumor-to-background than 18F-FDG. Rizzo-Padoin et al. [192] confirmed this data and reported that 18F-MEL050 PET/CT could detect sub-millimeter pulmonary metastases not found on 18F-FDG PET/CT. Additionally, Denoyer et al. [193] reported that lymph node metastases could be identified with perilesional 18F-MEL050 administration.

To overcome significant hepatic retention of 18F-5-FPN, its modification 18F-N-[2-diethy laminoethyl]-4-(2-[2-fluoroethoxy]ethoxy)ethoxy)pyridine (18F-PEG3-FPN) was recently introduced. 18F-PEG3-FPN was proved to have lower liver uptake than 18F-5-FPN while still demonstrating a high affinity to melanin. Hepatic and pulmonary metastases in a mouse model were detected with 18F-PEG3-FPN PET with a high tumor-to-background ratio, suggesting this imaging method for diagnosing primary and metastatic malignant melanoma [194]. Clinical application of 18F-PEG3-FPN PET/MR and PET/CT is evaluated in patients with clinically suspected or confirmed melanoma in the recruiting clinical trial with a study completion date estimated at the end of 2023 [NCT04747561]. Other radiolabeled melanin-specific tracers assessed in studies include 18F-labeled picolinamide [NCT03033485] and a new experimental drug 18F-SKI-248380, a dasatinib-derivative [NCT01916135]. With the potential shown in cutaneous melanoma cases, studies should be expanded to uveal melanoma patients.

Table 2. Radiopharmaceuticals for PET imaging with potential for use in uveal melanoma diagnosis.

<table>
<thead>
<tr>
<th>Feature</th>
<th>18F-NaF</th>
<th>18F-5-FPN</th>
<th>18F-MEL050</th>
<th>18F-PEG3-FPN</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Molecular target</strong></td>
<td>Bone remodeling</td>
<td>Melanin</td>
<td>Melanin</td>
<td>Melanin</td>
</tr>
<tr>
<td><strong>Current use</strong></td>
<td>Detection of bone metastases in prostate, breast, lung, thyroid cancer, diagnosis of fibrous dysplasia</td>
<td>Preclinical studies – detecting small metastases from malignant melanoma in lymph nodes and lungs</td>
<td>Preclinical studies – detecting small metastases from malignant melanoma in lymph nodes and lungs</td>
<td>Preclinical studies – detecting small metastases from malignant melanoma in lungs and liver</td>
</tr>
</tbody>
</table>
Potential use in uveal melanoma

Detection of bone metastases, assessing extrasceral extension of the primary tumor.

Detection of submillimeter metastases in lungs, lymph nodes.

Detecting submillimeter metastases in lungs, lymph nodes.

Detecting submillimeter metastases in liver, lungs.

3.2. PET/MRI

Discussions about combining PET and MR imaging started in the 1990s alongside the proposal of PET/CT. Great soft tissue contrast, a low dose of radiation, various imaging sequences, and possible simultaneous image acquisition suggested its promising role in oncological, neurological, and cardiovascular applications [195].

PET/MRI showed superiority over PET/CT to screen for liver metastases, especially for detecting lesions smaller than 1cm [196]. Donati et al. [197] compared liver metastases detection in PET/CT, gadoxetic-acid-enhanced MRI, and retrospectively fused images of PET and MR in 37 patients. The sensitivity was significantly higher for PET/MRI (93%) compared to PET/CT (76%), however in gatoxetic-acid-enhanced MRI (91%), sensitivity was proven to be not significantly different than PET/MRI. In the study conducted on 70 patients by Beiderwellen et al. [198], PET/MRI showed better diagnostic confidence and higher lesion conspicuity than PET/CT regarding the depiction of liver lesions in integrated whole-body PET/MRI. Additionally, Reiner et al. [199] proved higher accuracy of PET/MRI compared to contrast-enhanced PET/CT, but only in subcentimeter lesions without abnormal FDG uptake.

We have found no studies on PET/MRI applicability in ophthalmic oncology. However, great soft-tissue contrast and functional MRI sequences combined with a functional element of PET can provide accurate tumor information at the time of diagnosis and in therapy response evaluation in regions with complex anatomy such as ocular space [167]. Low radiation dose is another advantage, and it suggests using this modality while searching for liver metastases or in follow-up imaging. However, some challenges need to be addressed, including high costs, long acquisition time. Moreover, the research proving PET/MRI superiority over other imaging modalities was studied on only small patient groups suggesting the need for more research [195].

3.3. Contrast agents for MRI

The aim of contrast agents (CA) application in magnetic resonance imaging (MRI) and magnetic resonance angiography (MRA) is to improve the tissue contrast and visualize signal differences between contrast-containing and non-contrast-containing tissues. CA can be divided into positive which shortens T1 relaxation time and increases the signal on T1-w images, and negative which shortens T2 relaxation time and decreases the signal on T2-w images. The other classification distinguishes paramagnetic agents (metal ion with unpaired electrons: $^{24}\text{Cr}^{3+}$, $^{25}\text{Mn}^{2+}$, $^{26}\text{Fe}^{3+}$, $^{63}\text{Cu}^{2+}$, $^{64}\text{Gd}^{3+}$, $^{66}\text{Dy}^{3+}$ and chelating ligand) and super-paramagnetic agents (iron oxide or Fe/Mn compound and polymer) [200].

Commonly used CA contain gadolinium (III) compounds, have paramagnetic properties, shorten the T1 relaxation time of water protons, i.e., they act as positive agents [201]. Gadolinium is a lanthanide that is used in the form of chelates bonded to ligands. To increase the sensitivity, special molecules are combined with gadolinium as liposomes, micelles, apoferritin, lipoproteins [200].

As previously mentioned, besides Gd$^{3+}$, the other metal ions could be exploited as CA. Regarding Mn$^{2+}$, both chelates (dipyridoxyl diphosphate Mn-DPDP) and manganese nanoparticle scans are used. After dissociation of DPDP, free Mn is absorbed by liver cells (intracellular agent) and excreted by the bile and DPDP ligand (which lowers toxicity) by the kidneys [202]. This compound is dedicated to the hepatic cells, although it is also taken
up by other tissues (containing a large number of mitochondria) [202]. Pathological lesions composed of liver cells (e.g., HCC, FNH, adenoma) could be visualized on T1-w [203]. Because Mn\textsuperscript{2+} gets inside the neuron by voltage-gated calcium canal, it can act as a brain-imaging tool [204]. One of the manganese compounds, Tesla scan/ mangafodipirtrisodium, has been used in clinical practice but has been withdrawn. The other magnetic ion is dysprosium Dy\textsuperscript{3+} which has a high magnetic moment [205]. It undergoes an encapsulation process and is incorporated into the liposomes, thus serving as a tool of cell surface structures (e.g., transporters) imaging [200]. Furthermore, dysprosium-based CA could enable myocardial and brain perfusion visualization [205]. None of the compounds containing dysprosium have been registered for use.

The super-paramagnetic magnetic nanoparticles (SPMNPs) are the second group of CA, which generally influences T2 relaxation time [206]. Within the MNP class, we can distinguish superparamagnetic iron oxide (SPIO) and ultrasmall superparamagnetic iron oxide (USPIO). SPIO molecules are more than 40nm in diameter, USPIO is smaller in dimension (less than 40 nm) and gives higher contrast on T2 images than SPIO [207]. Intravenous administration of that CA results in their phagocytosis by macrophages and further signal reduction of the T2 signal from tissues infiltrated by cells containing CA [208]. Therefore, considering that the accumulation of macrophages increases in inflamed and infected tissue, SPIO may indicate its localization of inflammation and enable visualization of transplanted stem cells [209].

The purpose of other modifications of iron oxide nanoparticles is to target the molecules at those receptors confirmed to be present on tumor cells Yang et al. [210] proposed on the mouse model combination with of urokinase-type plasminogen activator (u-PA) and called it ATF-IO (amino-terminal fragment conjugated-iron oxide). ATF-IO binds specifically to the receptor u-PAR, which is abundant on changed cells. The paramagnetic nanoparticles directed against Robo4 and α\textsuperscript{v}β\textsuperscript{3}-integrin – markers associated with angiogenesis, were studied in imaging of melanoma in mice [211]. Indeed, they have not yet been analyzed in any research of uveal melanoma, but this makes it possible to test their usefulness in other types of cancer in the future.

Due to special coatings (SiO\textsubscript{2}, Al\textsubscript{2}O\textsubscript{3}, FeO\textsubscript{3}), the metal-alloy nanoparticles are not toxic than metallic CA [212]. Among others, this group includes Co, Fe, FeCo, and FePt [200]. The last of the mentioned compound is also called superparamagnetic iron platinum particles (SIPP). Taylor et al. [213] suggested that because more convenient relaxation features than conventional CA give the possibility to smaller cancer imaging at lower CA concentration. Encapsulation of SIPP with phospholipids into micelles and binding to the PSM antibody provides the prospect of prostate neoplasm diagnosis [213].

Magnetosomes (MS) are nano-sized crystals that are surrounded by a membrane and synthesized by Gram-negative bacteria (Magnetotactic Bacteria MTB) [214]. Those compounds could contain magnetite Fe\textsubscript{3}O\textsubscript{4}, greigite Fe\textsubscript{3}S\textsubscript{4} and are located inside the cell [215]. They are promising experimental tools as carriers for various substances as drugs, chemotherapeutics, or antibodies [216]. Due to bacterial movement ability provided by flagella and appropriate labeling of these organisms, it is possible to monitor their activity and the efficiency of delivery, especially that magnetosomes have an influence on T1 and T2 relaxation time (greater impact on T2 time; thus, they are negative CA) [217]. Boucher et al. [218] used magnetosomes with biogenic iron oxide and RGD peptide to visualize the contrast distribution in mice. They noticed that CA deposition in brain tumor cells occurred after 24 hours after administration. Zhang et al. [219] utilized the method to equip magnetosomes (by protein MamC) with antibodies against HER2 in the diagnostic process of breast cancer on an animal model. This method allows for selective targeting of CA to precisely defined neoplastic cells and will certainly be the research subject in the future. However, imaging is significantly more difficult when the receptors on the surface of tumor cells are not known. Xiang et al. [220] enriched magnetosomes with the P75 protein that binds selectively to MDA-MB468, SKBR3 cells (on HER2 and EGFR positive cells), and xenograft tumors.
CA specifically targeting uveal melanoma has not been developed so far, but attempts are underway to use them differently. One of the studies aims to diagnose Crohn’s disease with a new oral contrast agent (MRI) and comparison to CT imaging [NCT00587210]. The efficiency of MR lymphangiography after administration of ferumoxtran-10 (USPIO) was analyzed in the detection of the neoplasm in pelvic lymph nodes [NCT00147238][221]. Gadoxetate disodium in MRI was compared to CT in imaging of hepatocellular carcinoma and cirrhosis [NCT01341132]. In the other research, the efficiency of MRI with Gadofosveset and MR elastography are investigated in the diagnostics of prostate cancer [NCT01761812].

3.4. Hyperpolarized MRI

The hyperpolarized MR imaging (HP MRI) is an innovative imaging technique of nuclear spin polarization that gives information about metabolic processes in particular tissues (including tumors) in vivo in real-time using unique injected substrates, usually $^{13}$C pyruvate, which participates in different metabolic pathways and converts into isotope-labeled products [222,223]. Apart from $^{13}$C, the other nuclei may also be hyperpolarized and serve as markers of different molecules, e.g., $^1$H, $^{29}$Si, $^{15}$N, $^{89}$Y, $^{129}$Xe, $^{31}$P [224-226]. Every nucleus to be hyperpolarized must have spin-$1/2$ [227]. The high polarization of specific probes is obtained by external nuclear polarization and provides more enhanced signal than conventional MRI [227]. The methods to reach the effect of hyperpolarization of the molecules include, e.g., the dynamic nuclear polarization (DNP) – when spin polarization is transferred from electrons to nuclei using microwave radiation (the Overhauser effect).

Substances used in HP MRI should meet several conditions to be used as markers. The applicability of the molecule is determined by its essential role in a metabolic pathway, small dimension, ease of passing to a different cellular compartment, long T1 relaxation time, and the isotope $^{13}$C is the most common marker used in hyperpolarized probes [227]. Appropriate distribution of the injected hyperpolarized substance and sufficiently long half-life time of hyperpolarized magnetization are challenging aspects of that method that require future research. The lifetime of $^{13}$C-labeled markers is in the order of several tens of seconds [228].

The pyruvate is a significant component of cellular metabolism. As a product of glycolysis, it could participate in aerobic reactions (pyruvate may convert into acetyl CoA in the process of decarboxylation in which also carbon dioxide is formed, CO$_2$ then transform into bicarbonate) and in anaerobic reactions (when pyruvate is converted into lactate in the process of dehydrogenation by LDH-lactate dehydrogenase) [229]. The characteristic feature of neoplasms associated with pyruvate transformation is the Warburg effect, in which tumor cells in aerobic conditions preferably convert pyruvate to lactate (what is called oxygen glycolysis) [230]. Visualization of the Warburg effect by HP MRI confirms the presence of neoplasm, and the assessment of that process intensity after treatment could correlate with the therapy effectiveness [231].

In addition to mentioned substances, the other markers are developed. $[1^{13}$C$]$acetate as a probe is dedicated to tissues whose metabolism is based on fatty acids. Acetate is converted into acetyl-CoA, which could be included in the trichloroacetic acid cycle or transform into acetylcarnitine [232]. $[1^{13}$C$]$alanine could serve as a precursor of pyruvate (then lactate) and facilitate monitoring that metabolite without signal interference derived from extracellular pyruvate [233]. Xiao Ji et al. suggested that alanine and glycine have more favorable parameters of level and duration of hyperpolarization, giving the possibility to long storage of the marker before application [228]. $[1,4^{13}$C$]$fumarate transformation to malate is associated with cell necrosis and positive therapy response [234]. $[^{13}$C$]$urea (metabolically inert compound) has a dual character because it may be a probe or contrast agent [235]. $[1^{13}$C$]$dehydroascorbic acid (DHA) – oxidized ascorbic acid gives information about cellular redox state (oxidative stress may coexist with cancer). It could serve as a tool to control treatment response, but the uptake of DHA varies significantly.
between different tissues, so its role as a probe is not reliable [227]. [1-13C] succinate is a substrate of succinyl-CoA or fumarate and may give information about deregulation of oxidative phosphorylation [227]. The disorder of pH balance may occur in the course of cancer disease and could be detected using the following probes [13C] bicarbonate, [13C α-ketoisocaproate, [13C,15N]N-(2-acetamido)-2-aminoethanesulfonic acid, [15N] pyridine [227,236-238].

A significant disadvantage of this method is the high concentration of injected polarized (by DNP) substances and their potentially toxic character. In one of the clinical trials [NCT01229618], it was observed that the dose at which only mild side effects occur is 0.43 ml/kg of HP pyruvate. The other limitation is the signal loss due to T1 relaxation and RF pulse excitation, so hyperpolarization enhances the signal only for a short time after the advertisement of the HP agent [239]. That is why the development of the techniques preventing signal losses could constitute the field of studies.

Future research should also focus on looking for more selective and targeted markers concerning different cancers. Moreover, specifically selected probes will also allow the image to detect enzymatic disruptions resulting in reaction failure and substrate accumulation typically involved in the reaction. Like conventional MR, some technical conditions need to be improved, for example, the acquisition time. The subject of further study may concern new polarization techniques (the SPINlab (automatic) polarizer with 5T magnetic field is used in patient studies [240]) or transparent differentiation of signal from normal and neoplastic tissue. Furthermore, the HP substances with relatively long half-life time should be investigated to enable more effective distribution to specific compartments. The other aspect that could be developed is receiving qualitative information about the presence of cancer and quantitative data relating the signal intensity to the stage of cancer.

Although HP MRI is not yet applicable in the diagnostics of uveal melanoma, the studies currently carried out reveal more and more new applications of that method that may be introduced into clinical practice in the future. The ongoing first phase study [NCT04772456] focuses on the safety of HP MRI and efficiency of brain tumors (gliomas) imaging after administration of HP [1-13C] pyruvate in comparison to conventional MRI [241]. The visualization of lactate accumulation after injection of HP 13C-Pyruvate in locally advanced cervical cancer (LACC) gives a more specific metabolic characterization of the neoplasm. It is researched if co-imaging with DWI and PET would be possible to identify the areas less sensitive to radiotherapy or brachytherapy using HP MRI [NCT03129776]. Lactate levels evaluation in tumor cells is also performed in intracranial metastasis treated with stereotactic radiosurgery [NCT03324360]. There are also attempts to visualize advanced prostate cancers after administration of hyperpolarized marker (13C-pyruvate) [NCT04346225] and control the response to chemotherapy in pancreatic cancer after injection of HP pyruvate [EudraCT Number: 2016-004491-22].

4. Personalized medicine

Personalized medicine is promising and dynamically developing field covering both aspects of patient predispositions, diagnostics, clinical management, prognosis determination, and treatment [242]. On a more individual basis, the development of medicine will only be possible thanks to improved diagnostic methods that allow detecting specific differences influencing the choice of therapeutic decision. Genetic tests provide the most detailed and thorough information. Shields et al. [243] analyzed the cytogenetic tumor profile of 1059 patients to individualize the diagnosis. The occurrence of uveal melanoma metastases mainly was related to specific mutations (3 complete monosomy, 6 disomy, 8q gain, 8p loss, and three complete monosomy, 6 disomy, and 8q gain). It was also noticed that the other mutations as GNAQ, GNA11, SF3B1, EIF1AX, BAP-1, and PRAME are important in assessing the prognosis [243].
Indeed, extending the diagnostics with genetic changes in cancer cells significantly increases the scope of information, precision, and accuracy of diagnosis and generates further challenges to effectively use the obtained data.

4.1. Theranostics is a step towards personalized medicine

A promising strategy that should be mentioned here is to combine non-invasive diagnostic methods with therapy to apply personalized treatment called theranostics. The new imaging methods presented previously may be utilized in that field because they allow the use of selectively targeted agents (which may be drug-coated), control their transport to destination and therapy response, and, importantly, offer the possibility of real-time imaging. However, these methods also have limitations, e.g., reaching the target tissue by the probe. Still, various solutions are being developed to ensure the probe penetration into tumor tissue, including the use of small-sized particles (nanoparticles, liposomes, micelles), coverage with appropriate particles binding to special receptors, drug release technology induced by hypoxia, temperature, or specific pH ranges [244].

Due to their beneficial properties, proteins, antibodies, and aptamers could be used as theranostic tools which bind to molecules that are present in excess on tumor cells or in its environment, e.g. insulin-like growth factor 1 receptor (IGF1R), folate receptor (FR), epidermal growth factor receptor (EGFR), human epidermal growth factor receptor 2 (HER2), vascular endothelial growth factor (VEGF), integrins [245]. Combining these particles with markers allows for visualization, and the conjugation with anti-cancer drugs allows for extremely precise and localized therapy. Dal Corso et al. [246] proposed small-molecule−drug conjugates (SMDC) which contain tripeptide Arg-Gly-Asp (RGD) sequence and are ligands for αvβ3 integrin (present on tumor vessels). These conjugates can transport various drugs, e.g., doxorubicin, paclitaxel, cisplatin, or camptothecin used in systemic chemotherapy, but their toxic effect on the body is significantly reduced. It is also possible to utilize fluorescent probes activated by extracellular enzymes or intracellular lysosomal transformations [247]. This method improves the image quality as the marker only accumulates in the target tissue, and background signals are eliminated. Sun et al. [248] introduced aminopeptidase N (APN)-activated theranostic prodrug (NBFMel), which is a PET probe for diagnosis and therapy. In this case, the toxic substance to cancer cells is melphalan, which acts selectively and saves normal tissues.

An attractive solution in novel theranostic particles is the practical use of a particular feature of neoplasms, which maintains a slightly acidic pH (ca. 6.8) [249]. The iron oxide nanoparticles with pH-sensitive magnetic nanogrenades (PMNs) can self-organize under the influence of acidic reactions. The tumors of minimal size (<3mm in diameter) are detectable through the combination of MRI and fluorescence imaging. Ling et al. [249] suggested that pH-targeted photodynamic therapy (light radiation affects the photosensitizers to produce reactive oxygen species) could be utilized in the treatment of resistant tumors. Kim et al. [250] proposed multifunctional magnetite nanoparticles (AHP@MNPs). Apart from FeOx, they consist of photosensitizer, hyaluronic acid (AHP) and are synthesized from amines. Due to the surface charge, they can be covered with different coatings. These particles enter the tumor cells through CD44 and have dual nature. They may serve as a tool of magnetic hyperthermia therapy (magnetic nanoparticles convert electromagnetic energy to heat) and photodynamic therapy.

There are also nanoparticles which activation is enzymatically catalyzed [251]. Such an activation process is possible when the enzyme concentration in the tumor environment is increased. The molecule must be adequately designed; it should confirm easy bonding with the catalyst. One of the concepts assumes that the nanoparticle releases the drug after an enzymatic decomposition of the coating structure. The particle surface is properly modified, making it susceptible to enzymatic catalysis, or the drug is placed inside the liposome and released after the carrier’s destruction. Several types of enzymes might be involved in these nanoparticle decompositions, e.g., proteases, lipases, glycosidases (e.g., cathepsin B, peroxidase, urease, glucose oxidase, α-amylase). This technology
is a part of targeted cancer therapy, allows for assessing enzymatic activity, and poses a challenge to understand the tumor’s enzymatic arsenal further.

Kim et al. [252] suggested making use of the fact of the increased level of H$_2$O$_2$ in cancer metastases (as an adaptation to increased amounts of reactive oxygen species) for imaging and simultaneous stimulation of the anti-tumor effect of the transported drug (prodrug 7). The fluorescence induction and the substance SN-38 detachment were due to the change of boronate moiety to a hydroxyl group in ‘prodrug 7’ by H$_2$O$_2$. Thus, such an approach ensures the selectivity of the therapy and allows controlling the efficiency of the delivery of the molecules to the target site. Similar properties can be attributed to nanoparticles which functionality is dependent on tissue hypoxia. Zhou et al. [253] studied azo-derivative (AzP1) irinotecan analog. This compound consists of an inhibitor of topoisomerase I and aryl-azo benzyl alcohol (azo-derivate substances undergo a reduction in conditions of low oxygen concentration and serve as hypoxia markers). The visualization of fluorescence also allows for the quantification of the released chemotherapeutic agent.

The theranostic concept is a hope for the development in the field of oncology in the context of precise diagnostics, which will enable timely treatment. The appropriate structure of modified compounds using long-known anti-cancer substances allows for a highly selective action of these substances at the tumor site while limiting the harmful effect on other tissues and persistent side effects for the patient [254]. However, once introduced into clinical practice, the high cost of these methods remains an important issue due to their cost-effectiveness, time-consuming, and advanced technology.

Several studies are currently underway in this area (admittedly in diseases other than uveal melanoma). Somatostatin-expressing neoplasms like neuroendocrine tumors, medulloblastoma, meningioma, and neuroblastoma are treated with $^{90}$Y-DOTATOC, and the usefulness of $^{68}$Ga-DOTATOC PET/CT in diagnosis and treatment is assessed [NCT02441088]. Additionally, circulating tumor DNA (CTD) will be evaluated as a theranostic marker in glioblastomas [NCT03115138]. Combining radioactive particles such as $^{123}$Iodine, $^{89}$Yttrium with appropriate proteins: MIBG or DOTATOC ensures targeting drugs to tumor cells and minimizing side effects of therapy. Concurrent application of $^{123}$I-MIBG and $^{90}$Y-DOTATOC in mid-gut neuroendocrine tumors (NETs) and SPECT/CT control will be performed in a study [NCT03044977]. In turn, the study [NCT04769817] will analyze the results of patients previously treated with Lutetium 177 ($^{177}$Lu) –PSMA with metastatic castration-resistant prostate cancer (mCRPC) and evaluation the analogy between PSMA and ($^{18}$F-FDG) PET/CT images.

4.2. Immunotherapy as a component of personalized medicine

If the theranostics of uveal melanoma develops in the future, it would be worth adopting immunotherapy (Table 3). Although the development of adequately modified particles and their production is demanding and time-consuming, it will be a promising element of effective cancer therapy [255]. Thus, increasing attention is focused on using immunotherapy in uveal melanoma, even though the eye is considered an immunologically privileged structure. It is worth noting that cancer cells use many of these mechanisms, and some even mimic them, to avoid elimination by the immune system at the sites of metastases [256]. These include the expression of many particles with immunosuppressive activity against T lymphocytes or NK cells, the ability to inactivate complement, inhibit proliferation and induce apoptosis of T cells via PD-L1 (programmed death ligand-1) [256]. Therefore, monoclonal antibody anti-PD1, pembrolizumab was studied in Phase 2 clinical trials to treat advanced uveal melanoma (stages IIIA, IIIB, IIIC, IV) [NCT02359851]. Pembrolizumab with entinostat (histone deacetylase), which aims to increase the effects of immunotherapy, is also being evaluated in metastatic disease [NCT02697630]. Nivolumab is another tested anti-PD 1 antibody. Combination of nivolumab and relatlimab (anti-LAG3 -lymphocyte activation gene 3 is found on NK, B, activated T cells and limits their proliferation and activation [257]) is analyzed in patients with unresectable or metastatic uveal melanoma [NCT03743766]. Apart from that, the syn-
A different approach involves targeting the drug molecule to a compound that is in excess in this tumor, i.e., c-kit, with the participation of a monoclonal antibody such as imatinib [258]. Sunitinib also acts as an inhibitor of this kinase, and the SUAVE study compares its effects to decarbazine in metastatic uveal melanoma [NCT01551459]. While in a trial [NCT01005472] with sunitinib, temozolomide was used in patients with stages III and IV of the disease. Another kinase that was tried to be inhibited was c-MET; cabozantinib was used for this purpose. The effectiveness of this drug was assessed in relation to the compounds used in chemotherapy (temozolomide and dacarbazine) [NCT01835145] in a randomized trial. However, cabozantinib did not show the expected advantage. In turn, crizotinib is currently in the second phase of clinical trials, used as adjuvant treatment in patients at high risk of relapse after therapeutic intervention [NCT02223819]. This monoclonal antibody functions as an inhibitor of anaplastic lymphoma kinase (ALK) and c-ros oncogene 1 (ROS1).

The following molecules are also in the second phase of clinical trials: a combination of cediranib and durvalumab in metastatic UM [NCT04184518], axitinib in metastatic or unresectable UM [NCT0133948], defactinib and VS-6766 in metastatic UM [NCT04720417], niraparib in UM and mutation BAPI and with mutation BAPI other DNA damage [NCT03207347].

Mention should also be made of Immune mobilizing monoclonal T cell receptors (TCRs) against cancer ImmTAC™ molecules. These particles can redirect lymphocytes against cancer cells, and interestingly, activation affects both effector and memory cells. They are bifunctional, binding to CD3 on T cells and gp100 on UM cells [259]. Currently, the efficacy and safety of IMCGP100 (belonging to the group ImmTAC™) in advanced UM therapy is being studied in two clinical trials [NCT03070392], [NCT02570308].

### Table 3. Investigational compounds in the treatment of uveal melanoma or metastatic disease

<table>
<thead>
<tr>
<th>Evaluated treatment</th>
<th>Mechanism</th>
<th>Clinical trial</th>
</tr>
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<tbody>
<tr>
<td>pembrolizumab</td>
<td>anti PD-1</td>
<td>NCT02359851</td>
</tr>
<tr>
<td>pembrolizumab + entinostat</td>
<td>anti PD-1 + histone deacetylase</td>
<td>NCT02697630</td>
</tr>
<tr>
<td>nivolumab + relatimab</td>
<td>anti PD-1 + anty LAG-3</td>
<td>NCT03743766</td>
</tr>
<tr>
<td>nivolumab + ipilimumab</td>
<td>anti PD-1 + anti CTLA-4</td>
<td>NCT02626962</td>
</tr>
<tr>
<td>sunitinib</td>
<td>c-kit inhibitor</td>
<td>NCT01551459</td>
</tr>
<tr>
<td>sunitinib + temozolomide</td>
<td>c-kit inhibitor + DNA alkylating agent</td>
<td>NCT01005472</td>
</tr>
<tr>
<td>cabozantinib</td>
<td>c-MET inhibitor</td>
<td>NCT01835145</td>
</tr>
<tr>
<td>crizotinib</td>
<td>ALK and ROS1 inhibitor</td>
<td>NCT02223819</td>
</tr>
<tr>
<td>cediranib + durvalumab</td>
<td>VEGF inhibitor + anti-PD-L1</td>
<td>NCT04184518</td>
</tr>
<tr>
<td>axitinib</td>
<td>tyrosine kinase inhibitor</td>
<td>NCT01533948</td>
</tr>
<tr>
<td>defactinib + VS-6766</td>
<td>focal adhesion kinase inhibitor + RAF/MEK Inhibitor</td>
<td>NCT04720417</td>
</tr>
<tr>
<td>niraparib</td>
<td>PARP1 and PARP2 enzymes inhibitor</td>
<td>NCT03207347</td>
</tr>
<tr>
<td>IMCGP100</td>
<td>gp100-specific TCR fused to anti-CD3</td>
<td>NCT03070392, NCT02570308</td>
</tr>
<tr>
<td>selumetinib</td>
<td>MEK inhibitor</td>
<td>NCT02768766</td>
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### Table of Investigational Treatments

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Description</th>
<th>Clinical Trial ID</th>
</tr>
</thead>
<tbody>
<tr>
<td>bevacizumab + teozolomide</td>
<td>VEGF inhibitor + DNA alkylating agent</td>
<td>NCT01217398</td>
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<tr>
<td>glembatumumabvedotin</td>
<td>Antibody-drug conjugate targeted to transmembrane glycoprotein NMB</td>
<td>NCT02363283</td>
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<tr>
<td>bortezomib + paclitaxel + carboplatin</td>
<td>26S proteasome inhibitor + cytostatic alkaloid + complex compound that cross-links with DNA</td>
<td>NCT00288041</td>
</tr>
</tbody>
</table>

Data as of the date 23.04.2021. Clinical trials were searched on [https://www.clinicaltrials.gov/](https://www.clinicaltrials.gov/) with the terms “uveal melanoma” and “uveal melanoma treatment”.

### 5. Conclusions

Uveal melanoma is a rare intraocular malignancy with unfavorable diagnosis closely connected to tumor size, time of diagnosis, and presence of distant metastases. Tumor biopsy followed by pathological examination is not recommended in ophthalmic oncology; therefore, non-invasive imaging modalities are necessary to evaluate patients with suspected uveal melanoma [40]. US and OCT are the elementary imaging technique in the assessment of primary tumors. Both methods are cheap, widely available, and provide information about the tumor’s size, shape, structure, and vascularization, especially in the anterior segment of the eye. SPECT could also be used in the initial diagnosis, although exposure to radiopharmaceuticals and the high cost of this method make it more of an additional rather than primary imaging technique [39]. Additionally, FAF/ICGA/FAF are helpful in differential diagnosis [44]. MRI, as well as PET/CT, have limited value in the diagnosis of the primary tumor; however, they are necessary for tumor staging and detection of distant metastases [39,40].

The introduction of new advancements in imaging techniques could broaden the horizons in the diagnosis of uveal melanoma. The development of new radiolabeled radiotracers in PET allows extended differential diagnosis and detection of submillimeter metastases in the liver using melanin-specific radiotracers. Multimodal imaging with PET and MRI combined offers exceptional soft-tissue contrast that could deliver primary tumor characteristics and be used for staging and metastases identification. Visualization of specific metabolic processes is possible with HP MRI, which could allow early detection of uveal melanoma and any metastases. Similarly, new contrast agents for MRI maximize tissue contrast and can help to detect intraocular and hepatic lesions. On the other hand, new improvements in non-invasive imaging techniques can also be utilized in personalized medicine, such as theranostics. Although the applicability of those diagnostic advancements has been yet to be proven in the group of patients with uveal melanoma, their irrefutable potential needs to be tested in future studies.

### Author Contributions:

Conceptualization, M.F., and A.M.C.; writing—original draft preparation, M.S. and N.S.; writing—review and editing, M.F. and A.M.C.; supervision, M.F., and A.M.C.; funding acquisition, M.F. All authors have read and agreed to the published version of the manuscript.

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