Conservative management of retinoblastoma: Challenging orthodoxy without compromising the state of metastatic grace. “Alive, with good vision and no comorbidity”

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Dedicated to A. Linn Murphy, my paragon and mentor, and to Akhiro Kaneko who pioneered modern targeted chemotherapy of intraocular retinoblastoma

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Complication
Metastasis
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ABSTRACT

Retinoblastoma is lethal by metastasis if left untreated, so the primary goal of therapy is to preserve life, with ocular survival, visual preservation and quality of life as secondary aims. Historically, enucleation was the first successful therapeutic approach to decrease mortality, followed over 100 years ago by the first eye salvage attempts with radiotherapy. This led to the empiric delineation of a window for conservative management subject to a “state of metastatic grace” never to be violated.

Over the last two decades, conservative management of retinoblastoma witnessed an impressive acceleration of improvements, culminating in two major paradigm shifts in therapeutic strategy. Firstly, the introduction of systemic chemotherapy and focal treatments in the late 1990s enabled radiotherapy to be progressively abandoned. Around 10 years later, the advent of chemotherapy in situ, with the capitalization of new routes of targeted drug delivery, namely intra-arterial, intravitreal and now intracameral injections, allowed significant increase in eye preservation rate, definitive eradication of radiotherapy and reduction of systemic chemotherapy.

Here we intend to review the relevant knowledge susceptible to improve the conservative management of retinoblastoma in compliance with the “state of metastatic grace”, with particular attention to (i) reviewing how new imaging modalities impact the frontiers of conservative management, (ii) dissecting retinoblastoma genesis, growth patterns, and intraocular routes of tumor propagation, (iii) assessing major therapeutic changes and trends, (iv) proposing a classification of relapsing retinoblastoma, (v) examining treatable/preventable disease-related or treatment-induced complications, and (vi) appraising new therapeutic targets and concepts, as well as liquid biopsy potentiality.

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1. Introduction

Retinoblastoma is the archetype of developmental tumors and is responsible for 17% of neonatal (Halperin, 2000), 13% of infantile (Vasilatou-Kosmidis, 2003), 6% and 3% of all pediatric cancers under age 5 and 15 years, respectively (Broadus et al., 2009; Group et al., 2013). Left untreated, this malignancy is rapidly recognizable as a protruding mass. Within the limitations of retrospective diagnosis (Kivela and Polkunen, 2003), it is thus not surprising to find retinoblastoma already mentioned in the medical literature at the turn of the XVIth and XVIIth centuries by Pawius in the Netherlands (Pawius, 1657) and Hildanus in Switzerland (Hildanus, 1682) respectively. According to the Hippocratic theory of four humors still in use at that time, ocular tumors were thought to result from a causal chain, of which the patient’s constitution and hygiene was the first link (causa primitiva), triggering black bile overflow (causa antecedens) pouring into the eye after its transfer to the brain (causa coniuncta) (Koelbing, 1954). The advocated treatment was a combination of poultice, leech and purge to eliminate the flood of melancholic humor. This approach was supplanted by enucleation, made possible by the advent of anesthesia in 1903 (Hilgartner, 1903), allowing for the first time a conservative (globe-preserving) management of the disease. Since then, and for the next 90 years, external beam irradiation became the first line treatment, complemented by focal therapeutic modalities such as cryotherapy, xenon photocoagulation and brachytherapy for less advanced diseased eyes. In 1996, following the recognition of a radio-induced neoplasm predisposition in germline carriers, chemoreduction replaced radiotherapy, which use was relegated to treat chemo-resistant disease in only eyes as an alternative to enucleation. The introduction of this whole-body therapy failed, however, to improve eye preservation in advanced disease, while exposing children to systemic toxicity. This was explained by the pharmacokinetics of the perfused drugs not reaching tumoricidal concentrations in the various intraocular seeding compartments. To overcome these limitations, the concept of targeted chemotherapy using intra-arterial and intravitreal drug delivery was pioneered in Japan by Akihiro Kaneko already 30 years ago (Kaneko and Takayama, 1990; Yamane et al., 2004). In 2008, the technique of intra-arterial chemotherapy, revisited by Abramson (Abramson et al., 2008), became rapidly adopted and proved its efficacy to control retinal and subretinal tumors, but not vitreous seeds. The latter fully surrendered only to intravitreal chemotherapy, which became available in 2012 following the description of a safety-enhanced injection technique (Munier et al., 2012a). Aqueous seeding, the very last intra-ocular sanctuary to resist virtually all available treatment modalities, became manageable only since 2015 with the description of a technique adapted to safely inject into the anterior and posterior chambers (Munier et al., 2015), now successfully implemented in the current management of aqueous disease (Munier et al., 2017a, 2018). To date, the appropriate combination of all these modalities has enabled unprecedented eye survival rates, even for advanced retinoblastoma, contributing at the same time to the eradication of external beam irradiation in the treatment of intraocular retinoblastoma without jeopardizing patient survival.

2. Epidemiology

Retinoblastoma is a cancer of early childhood, habitually described as a rare tumor. Statistical evidence, however, suggests that it is likely the most common eye cancer worldwide (Kivela, 2009). As a developmental cancer, its incidence is highly dependent on age (Table 1), and therefore difficult to establish. To date, the most stable estimates are derived by assigning incident retinoblastomas to the year the patient was born, a method known as birth cohort analysis (Li et al., 2016; Park et al., 2014; Seregard et al., 2004).

2.1. Incidence

The incidence of retinoblastoma depends on the size of the population and the birth rate (Kivela, 2009). Around 7980 retinoblastomas are predicted to have been diagnosed in 2015 (Table 2). Predictions for Africa, representing 16% of the world population, are 2,293, or 29% of...
the total. Almost double the number, 4258 or 53%, is predicted for Asia, with 60% of the world population. Latin America, North America, and Europe with 8%, 3% and 6% of retinoblastomas also have fewer incident cases relative to their percentage of the world population of 9%, 5% and 10%, respectively. These discrepancies stem from differences in birth rates that range from 33.6 per 1000 population in Africa to 10.5 in Europe (Table 2).

Comprehensive registration of retinoblastoma has succeeded mainly in North America, Europe, and Japan, all of which have low birth rates and easy access to health care. For the rest of the world, registration has been incomplete. Mathematical estimates based on birth rates and population size remain more accurate than published incidences, which are derived from regional rather than national registers and subject to cultural, economic and logistic factors. Published incidence rates have at best approached, and more often fallen short of, those derived mathematically (Li et al., 2016; Usmanov and Kivela, 2014). Registration of childhood cancers in general is incomplete, although coverage increases with advances in diagnosis and treatment (Kroll et al., 2012).

Mathematical estimates are important in order to study and predict trends in incidence of retinoblastoma across countries and regions. Calculations based on World Population Prospects (United Nations, 2017) suggest that the world incidence of retinoblastoma will peak at 8300 cases in 2045 and then level off (Fig. 1). In Asia, Europe, and Latin America, the incidence has been declining, although recent waves of refugees from regions in which birth rates are higher may slow down the decline in Europe. The incidence increases in North America and especially in Africa where the annual incidence of retinoblastoma is still predicted to rise by 1300 cases, approaching that of Asia by 2060. Such long-term predictions might be affected by shifts in birth control policies and natural or other disasters such as famine and war, both directly and through refugees and other migrants.

Despite annual variation in crude incidences, especially in small populations and over short observation periods, the incidence of retinoblastoma has remained stable over the last fifty years wherever reliable records exist (Broadus et al., 2009; Gatta et al., 2012; Gregersen et al., 2016; Li et al., 2016; MacCarthy et al., 2009; Moreno et al., 2014), especially when the analysis is based on birth cohorts rather than on live births or on children of specified age in a given year (Park et al., 2014; Seregard et al., 2004).

Unilateral retinoblastoma represents 61–75% of all cases, with 25–39% bilateral according to recent population-based series (Andreoli et al., 2017; Azar et al., 2006; Gregersen et al., 2016; Khandekar et al., 2004; MacCarthy et al., 2009; Moreno et al., 2014). Part of this variation is explained by the fact that some initially unilateral hereditary retinoblastomas will progress to bilateral disease (Moreno et al., 2014), while the contribution from familial cases also varies between countries (Usmanov and Kivela, 2014). The incidence of trilateral tumors is estimated to be 3.5% in patients with hereditary disease (de Jong et al., 2015b).

2.2. Prevalence

Data on survivors of retinoblastoma are not readily available (Gatta et al., 2012). In the absence of a national registry, a crude prevalence estimate can be derived as follows: for every child born with retinoblastoma, the number of survivors will be roughly the number corresponding to their average life expectancy in that country or region, deduced by the average retinoblastoma mortality, then multiplied by the average number of retinoblastoma cases per year. To give an example, for a country with a mean life expectancy at birth of 73 years between 1950 and 2015 (from the World Population Prospects), a mean mortality from retinoblastoma of 3%, and 4 new patients per year, the estimate would be (73-(0.03x73))x4 or 283 survivors.

2.3. Predisposing factors for sporadic retinoblastoma

**Age.** Seven to 10% of retinoblastomas are neonatal (Kivela and Hadjistilianou, 2017), being diagnosed during the first month of life and occasionally at birth. In general, retinoblastoma is diagnosed in children under 3 years of age, is rare after 8 years of age and typically not found after the age of 15 (Gatta et al., 2012; Park et al., 2014; Rangamani et al., 2015). It is virtually unknown in adults (Kaliki et al., 2015a; Sengupta et al., 2016), accounting for less than 0.1% of all retinoblastomas.

**Sex.** In two large population-based registries of 1601 and 1452 patients, the ratio was 1.07 and 1.09 respectively (Andreoli et al., 2017; MacCarthy et al., 2009), while others report a preponderance of boys over girls with a ratio of 1.12–1.64 (Li et al., 2016; Moreno et al., 2014; Park et al., 2014; Rangamani et al., 2015), which is higher than the world statistic of 1.06 at birth (United Nations, 2017) (Andreoli et al., 2017; MacCarthy et al., 2009). The reasons for this discrepancy are not known, but it might be that in some cultures boys are preferentially brought to treatment (Li et al., 2016).

**Ethnicity and latitude.** No consistent difference in the incidence of retinoblastoma by ethnicity or latitude has been reported. However, racial and ethnic disparities in the incidence of retinoblastoma were recently reported in the United States (Friedrich et al., 2017; Steliarova-Foucher et al., 2017), with a significantly higher incidence of bilateral retinoblastoma in the Hispanic population, which remains as yet unexplained.

**Parental influence.** The risk of sporadic retinoblastoma increases with parental and especially maternal age (Moll et al., 1996; Saremi et al., 2014; Yip et al., 2006), although not consistently (Heck et al., 2012), and with maternal active smoking during pregnancy (Azary et al., 2016).

**Precursor lesions.** Retinoma has emerged as a frequent intermediary in the genesis of both hereditary and nonhereditary retinoblastoma. In one study, 20% of enucleated eyes with retinoblastoma had a retinoma component in the tumor, generally located towards the tumor base (Eagle, 2009). Like retinoblastomas, retinomas carry two mutated RB1 alleles, but they will remain quiescent unless additional hits drive them into retinoblastoma (Dimaras et al., 2008). The apparent high propensity of retinomas to degenerate into retinoblastoma likely explains their rarity as a true benign counterpart of retinoblastoma. It is widely presumed that transformation of long-standing retinomas is responsible for the exceptional adult retinoblastoma (Sengupta et al., 2016).

### Table 2

<table>
<thead>
<tr>
<th>Population</th>
<th>Birth at Risk Rate/1000</th>
<th>Median</th>
<th>95% Confidence Interval</th>
</tr>
</thead>
<tbody>
<tr>
<td>Africa</td>
<td>1,194 million</td>
<td>33.6</td>
<td>2.939–2.936</td>
</tr>
<tr>
<td>Asia</td>
<td>4,420 million</td>
<td>16.5</td>
<td>3.359–4.353</td>
</tr>
<tr>
<td>Europe</td>
<td>741 million</td>
<td>10.5</td>
<td>4.643–4.711</td>
</tr>
<tr>
<td>Latin America</td>
<td>632 million</td>
<td>16.4</td>
<td>0.552–5.677</td>
</tr>
<tr>
<td>North America</td>
<td>356 million</td>
<td>12.4</td>
<td>0.264–2.293</td>
</tr>
<tr>
<td>World</td>
<td>7,383 million</td>
<td>18.6</td>
<td>7.983–7.170–8.795</td>
</tr>
</tbody>
</table>

*a Based on total population, crude birth rate, and infant mortality rate as reported in the World Population Prospects: The 2017 Revision (United Nations, 2017), and the reported retinoblastoma incidence of 1 in 16,642 live births (95% confidence interval 15.105–18.528) by birth cohort analysis (Seregard et al., 2004), as described (Usmanov and Kivela, 2014).
Fig. 1. Incidence and predicted incidence of retinoblastoma from 1960 to 2060. A. In the world and by region (B) Africa, (C) Asia, (D) Europe, (E) Latin America, and (F) North America. The population, birth rate and infant mortality are taken from the World Population Prospects, 2017 Revision (United Nations, 2017), and the number of retinoblastomas per live born children from a North European birth cohort estimates (Seregard et al., 2004). Vertical lines indicate 2015. Undulations and peaks in the graphs are primarily explained by changes in birth rates.

Fig. 2. Gaze-dependent leukocoria as presenting sign of a retinoblastoma in the left eye of a 14-month-old boy not visible on straight gaze but only from the temporal side (A) due to the nasal location of the tumor (B).
educational program in Honduras allowed to significantly reduce the proportion of patients with extracocular spread at diagnosis within less than a decade (Leander et al., 2007). This unique feature of retinoblastoma has the potential to improve patient and ocular survival by earlier detection, not only in developing countries where more than 90% of retinoblastoma patients reside (see 2), but also in high-income countries where there is still room for increased sensitization.

In high-income countries, it is estimated that less than 3% of retinoblastoma patients are diagnosed with extra-ocular disease at the time of presentation (Chantada et al., 2011). Leukocoria, a white pupillary reflex, is the inaugural sign in 60% of the cases (Abramson et al., 1998). According to the size and location of the tumor(s) (Fig. 2), this sign can be initially intermittent, with detection also influenced by the pupil diameter, and frequently misinterpreted as an optical illusion by the parents or health care providers until the growing tumor elicits a permanent leukocoria. It is not rare to identify leukocoria with flash photography (photoleukocoria), which can precede the diagnosis by several months, especially when the anti-red eye reflex mode of the camera is inactivated, preventing pupil constriction from the pre-photograph flash.

Strabismus occurs when central vision is lost secondary to macular tumoral involvement. It is the second most frequent presenting sign, found in 20% of the cases, manifesting earlier and associated with less advanced disease and better ocular survival than leukocoria (Abramson et al., 1998, 2003; Shields et al., 2004a; Wallach et al., 2006).

Atypical signs in the remaining 20% of cases usually manifest in older children, where they can masquerade other conditions such as uveitis, glaucoma, cataract, and presumed trauma with intra-ocular hemorrhage. These manifestations in a patient with unsuspected retinoblastoma often lead to invasive diagnostic or curative procedures and are associated with iatrogenic risk of tumor spread. Finally, midfacial dysmorphic features suggestive of a syndromic form of retinoblastoma with developmental delay and birth defects due to a deletion of the long arm of chromosome 13, can prompt screening of the fundus in otherwise asymptomatic patients (see 5.1.2.2).

In developing countries, the diagnosis can be significantly delayed, with extra-ocular disease present at diagnosis, occurring in less than 10% of cases in middle-income countries (Chantada et al., 2004, 2011; Kaliki et al., 2019) and up to 45% in low-income countries (Bowman et al., 2008; Waddell et al., 2015). In this context, late presenting signs such as buphthalmic eyes with neovascular glaucoma, hyphema, non-infective orbital cellulitis and proptosis predominate.

3.2. Clinical examination and ancillary testing

Retinoblastoma is the only pediatric cancer not requiring pathology to confirm the diagnosis, due to its pathognomonic funduscopic features. Indirect ophthalmoscopy with scleral depression under anesthesia is the gold standard for detection of retinoblastoma and is usually sufficient to establish the diagnosis and grouping of eye disease. Fundus photography provides the necessary documentation for monitoring conservative therapy as well as for telemedicine purposes. Ancillary investigations, such as ultrasonography and MRI, are instrumental for the differential diagnosis in eyes with opaque media and for staging purposes, by revealing possible extraocular extension of the disease. During conservative management, the therapeutic strategy is continuously challenged by tumor’s response and treatment-related complications. The fine tuning in decision making increasingly relies on ancillary imaging techniques such as fluorescein angiography (FA), optical coherence tomography (OCT) and ultrasonic biomicroscopy (UBM). These imaging modalities have become pivotal in the subclinical detection and monitoring of tumoral and non-tumoral events, and thus have a profound impact on both the treatment strategy and the final clinical outcome. As such, these diagnostic tools should be part of the equipment of any tertiary referral center for retinoblastoma.

3.2.1. Fundus documentation and fluorescein angiography

Iconographic documentation of the fundus is mandatory at presentation and at each new exam, to provide precise and detailed documentation of each tumor comprising location, size and aspect, together with their relation to the optic nerve, macula or ora serrata. Rupture of the internal limiting membrane on the tumor apex, presence of vitreous seeding, retinal detachment (past or present) or subretinal dissemination are signs of more extensive disease and thus higher risk of recurrence and therapeutic failure. All of these clinical details must be documented accurately to better anticipate the disease course and to adapt the treatment strategy accordingly.

Fundus drawing is the simplest and also the most instructive method of documentation for the examiner, requiring a rigorous sense of observation and faithful interpretation. It is, however, time consuming and examiner-dependent. The schematic representation according to a conventional code may easily fail to depict small changes of the tumor, or slow progression of growth. In addition, exchange with other experts or physicians with this kind of support can be problematic. For these reasons, fundus photography has taken on an important role in the documentation of retinoblastoma, providing objective representation and easier comparison of images. Several models of hand-held digital cameras with corneal contact lenses (30–130°) are commercialized for use in the operating theater, namely the RetCam™ (Clarity Medical Systems, Inc., Pleasanton, CA, USA), the PanoCam™ Pro (Visunex Medical Systems, Fremont, CA, USA), and ICON™ (Phoenix Technology Group, Pleasanton, CA, USA). FA modules are available on all 3 devices. For children over 4 years of age and examined without anesthesia, Optos™ (Marlborough, MA, USA) is an ultrawide-angle imaging system providing a 200° retinal field view in one non-contact capture, even with a pupil diameter as small as 1 mm.

FA of both the anterior and posterior segment plays an essential role in the evaluation of retinoblastoma patients, especially in the context of intra-arterial chemotherapy, by providing undeniable information regarding tumor activity or vascular complications under the following circumstances:

(i) At presentation, to detect subclinical iris neovascularization (Kim et al., 2014) and assess the tumor and retinal vascularization status (Ohnishi et al., 1982).

(ii) After each intra-arterial course, to detect vascular complications such as ischemic retinopathy, associated or not with retinal and/or iris neovascularization (see 7.3.4).

(iii) At completion of first line and consolidation therapy, to provide a baseline angiogram to be used for future comparison (Shields et al., 1982).

(iv) In case of suspicion of relapse, especially in the context of type II and III regression patterns, by revealing tumor neovascularization (Ohnishi et al., 1982; Shields et al., 1982).

(v) In case of new retinal detachment, neovascularization and/or intraoculard hemorrhage during conservative treatment, to assess the intraocular vascularization status (Stathopoulos et al., 2018a).

(vi) To monitor treatment response to intraocular vascular complications such as focal treatment of ischemic retina, retinal detachment repair and/or anti-Vascular Endothelial Growth Factor (VEGF) therapy (Stathopoulos et al., 2018a).

3.2.2. Ultrasonic biomicroscopy (UBM)

UBM yields high resolution images of the anterior segment and structures of the posterior chamber that are hidden from direct visualization, such as the ciliary body, zonules or anterior hyaloid. Over the last decade, UBM has proved to be instrumental in evaluating any anterior extension of retinoblastoma (Moulin et al., 2012; Vasquez et al., 2011). Using a 35 MHz transducer with an acoustic 68 μm axial resolution (OTT Scan 2000; Ophthalmic Technologies, North York, Ontario, Canada), we have shown that tumoral invasion of the posterior chamber can be detected with high sensitivity (81%) and specificity (100%), even in the absence of anterior chamber involvement (Moulin et al., 2012).
et al., 2012). It is now possible to obtain scans of 12 mm at 50 MHz (Aviso™, Quantel Medical, Clermont-Ferrand, France) with a 35 μm axial resolution, providing unprecedented details of the micro-anatomy of the anterior segment (Fig. 3) and improvement in terms of tumor detection sensitivity.

In clinical practice, UBM has become a fundamental tool for:

(i) Assessing the anterior extent of retinoblastoma at diagnosis or at relapse with respect to the integrity of the anterior hyaloid and the anterior uvea (ciliary body and iris) (see 6.2.4).
(ii) Exploring the extreme periphery and posterior chamber in diffuse anterior retinoblastoma for the identification of the primary tumor, which can be located in the retinal rim under the vitreous base or anterior to the ora serrata (see 6.1.2.3).
(iii) Identifying a safe pars plana meridian, devoid of parietal tumor growth, as entry site for intravitreal chemotherapy, especially when pupil dilation is compromised or in the presence of opaque media (see 7.3.2).

3.2.3. Optic coherence tomography (OCT)

Commercialization of hand-held spectral-domain OCT apparatus (HH-SD OCT), (Biophtigen™, Durham, North Carolina, USA or Optovue iVue, Fremont, CA, USA) specifically adapted for pediatric use, has enabled OCT imaging in children lying in the supine position. The integration of HH-SD OCT in the management of retinoblastoma is attested by an increasing number of publications allowing better understanding and evaluation of both tumoral and non-tumoral abnormalities.

In practice, OCT provides an invaluable tool to improve retinoblastoma care in the following situations:

(i) Early detection of tumors or tumor growth allowing prompt therapeutic response, particularly beneficial in case of juxta-papillary or juxta-foveal lesions. Specific cases include the following presentations:
   a. Cryptic retinal tumors (Berry et al., 2016a; Rootman et al., 2013; Seider et al., 2016; Soliman et al., 2017), appearing as round hyperreflective lesions located in the inner or outer retina (Fig. 4A and B). Such subclinical findings challenge the classification if present at diagnosis, possibly changing unilateral to bilateral involvement and/or grouping (see 3.3.1).
   b. Cryptic incipient retinal recurrences (Gaillard et al., 2018) (Fig. 4C).
   c. Epipapillary lesions at a pre-invasive stage (Fabian et al., 2017b) (Fig. 4D and E).
   d. Choroidal invasion (Stathopoulos et al., 2018b) (Fig. 4I).
   e. Precise delineation of the invisible front line of tumor growth in diffuse infiltrating retinoblastoma (Stathopoulos et al., 2019) (Fig. 9G–I).
   f. Cluster of cells (anterior chamber seeding) in the irido-corneal angle and/or placoid growth on the corneal endothelium that could otherwise go unnoticed (anterior chamber OCT) (Fig. 4J and K).

(ii) Monitoring of the treatment response in the vitreous (Fig. 4G and H) or retina (Fig. 4J and K). In the specific case of cryptic type 0 regressed tumors (Saktanasate et al., 2015), OCT enables documentation of a residual hyperreflective lesion that should either receive focal therapy to result in a type 4 regression, or be carefully monitored to prove its inactivity (absence of growth) (Fig. 4J and K).

(iii) Evaluation of the retinal and optic nerve status behind dense retrohyaloid or epiretinal seeding (Hasanreisoglu et al., 2015) (Fig. 4F).

(iv) Assessment of the foveal integrity to evaluate visual potential. In the specific case of retinal detachment, OCT can be used to monitor foveal detachment while awaiting complete spontaneous or

surgical reapplication before intensifying amblyopia treatment (see 7.3.1).
(v) Documentation of non-tumoral treatment-related alterations such as choroidal thinning post intra-arterial chemotherapy (Maidana et al., 2014) or radio-induced macular edema (Miller et al., 2005).
(vi) Differential diagnosis of other retinal tumors (Fig. 4L–N).
3.2.4. Magnetic resonance imaging (MRI)
High-resolution contrast-enhanced MRI is currently considered to be the most sensitive technique to evaluate any extraocular extension of retinoblastoma. In middle and high-income countries, it has now completely supplanted computer tomography (CT) imaging, which carries an increased risk of radiation-induced oncogenesis without
Fig. 4. Role of optical coherence tomography (OCT) in the management of retinoblastoma. A. Detection of an invisible parapapillary tumor in the left eye of a 2-month-old girl referred for a familial retinoblastoma (scT1bN0M0), (B) which regressed to a hyperreflective scar on OCT after Argon laser treatment. C. Early diagnosis of a recurrence over a parapapillary scar. D. Diagnosis of an invisible epipapillary vitreous seed. Note the intact demarcation line between the seed and the papilla, supporting the absence of prelaminar invasion, before (D) and after (K) treatment with intravitreal melphalan. F. Documentation of an intact fovea behind dense epiretinal seeding predictive of a good visual prognosis. G. Assessment of the treatment response of prehyaloidal vitreous seeds (spheres) before (G) and after (H) intravitreal melphalan. The appearance of hyperreflective dots in spheres indicates complete response implying treatment completion. I. Early diagnosis of a secondary choroidal invasion. J. Posterior pole retinoblastoma (arrow heads) at diagnosis and (K) after 6 cycles of systemic chemotherapy in the context of bilateral disease, showing regression type 0 on indirect ophthalmoscopy, but persistent lesion on OCT necessitating appropriate follow-up. L-N. Two-week-old girl with leukocoria, neovascular glaucoma and total exudative retinal detachment in the right eye (L) referred for suspicion of retinoblastoma. M, N. Left eye examination revealed multiple small retinal lesions involving the nerve fiber layer on OCT, corresponding to retinal astrocytic hamartomas. Tuberculous sclerosis was confirmed on magnetic resonance imaging one week later, and by the presence of a germline TSC2 mutation. Pathology of the enucleated right eye revealed a giant cell astrocytoma. NB: Lines on fundus photography correspond to the adjacent optical coherence tomography scan.

adding any diagnostic value. The success of MRI in the context of retinoblastoma relies, however, on specific requirements which differ from those used for routine brain and orbit examinations. A consensual standardized protocol for the evaluation of retinoblastoma at diagnosis has recently been proposed by members of the European Retinoblastoma Imaging Collaboration (ERIC) (de Graaf et al., 2012). In order to achieve high resolution imaging, two techniques using 1.5 or 3 T scanners can be employed. With the 1.5 T scan, the use of a surface coil to increase the signal-to-noise ratio is necessary. Surface coils have, however, the disadvantage of requiring more time for accurate positioning, allow the examination of only one orbit at a time, and are less precise in imaging the posterior part of the orbit due to rapid signal loss. Surface coils are not needed with a 3 T scanner equipped with a 32-channel coil (de Jong et al., 2015a).

MRI plays a critical role in assessing the following:

(i) Detection of tumor calcifications, which is important in the differential diagnosis especially in cases where the fundus view is obscured. Calcifications can be observed with high sensitivity and specificity on T2 images without the help of CT, as proved by an ex vivo MRI study (Rodjan et al., 2015).

(ii) Patient staging by enabling detection of extra-ocular invasion at
the level of the:

a. **Optic nerve**. In the presence of a normal-sized optic nerve sheath, the diagnosis of retrolaminar invasion relies on contrast enhancement after gadolinium injection in T1. However, high resolution-MRI shows limited sensitivity in depicting early-stage optic nerve invasion and therefore cannot substitute for pathologic analysis. These limitations are highlighted in a 1.5 T surface coil study with histological verification, which proved to be disappointing in terms of sensitivity, although high specificity (range 0.64–1) and high negative predictive values (range 0.81–0.97) could be confirmed (Brisse et al., 2015). In the case of massive optic nerve invasion, an intra-cranial mass can form and cerebrospinal fluid invasion leading to leptomeningeal enhancement can be visualized.

b. **Choroid**. The normal choroid enhances after gadolinium injection, whereas disruption of the choroid blush is a sign leading to the suspicion of choroidal invasion (de Graaf et al., 2012).

c. **Sclera**. The sclera does not enhance and is seen as a hypointense line on T2 images. Disruption of this line and the deformation of the globe are signs of scleral invasion.

(iii) Screening for trilateral retinoblastoma (see 3.6) (Fig. 5).
(iv) Diagnosis of second primary neoplasms, especially in hereditary retinoblastoma survivors treated by external beam irradiation (see 3.8).

### 3.3. Classifications

Classification schemes in cancer aim to provide a universal language for the evaluation of the disease extent, which is necessary to determine and compare different treatment strategies and to enable prognostication. Unlike cancers affecting other organs, where tumor excision or biopsy is usually performed for diagnosis confirmation and histopathologic grading, the diagnosis of retinoblastoma is clinical. Classifying retinoblastoma poses, however, a particular challenge as both intraocular and extraocular tumor extension need to be assessed in order to predict not only the likelihood of eye preservation, but also the

<table>
<thead>
<tr>
<th>Category</th>
<th>Subcategory</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>cTX</td>
<td>Unknown evidence of intraocular tumor</td>
<td></td>
</tr>
<tr>
<td>cT0</td>
<td>No evidence of intraocular tumor</td>
<td></td>
</tr>
<tr>
<td>cT1</td>
<td>Intraretinal tumor(s) with subretinal fluid ≤ 5 mm from base of any tumor</td>
<td></td>
</tr>
<tr>
<td>cT1a</td>
<td>Tumors ≤ 3 mm and further than 1.5 mm from disc and fovea</td>
<td></td>
</tr>
<tr>
<td>cT1b</td>
<td>Tumors &gt; 3 mm or closer than 1.5 mm from disc or fovea</td>
<td></td>
</tr>
<tr>
<td>cT2</td>
<td>Intraocular tumor(s) with retinal detachment, vitreous seeding, or subretinal seeding</td>
<td></td>
</tr>
<tr>
<td>cT2a</td>
<td>Subretinal fluid &gt; 5 mm from the base of any tumor</td>
<td></td>
</tr>
<tr>
<td>cT2b</td>
<td>Vitreous seeding and/or subretinal seeding</td>
<td></td>
</tr>
<tr>
<td>cT3</td>
<td>Advanced intraocular tumor(s)</td>
<td></td>
</tr>
<tr>
<td>cT3a</td>
<td>Phthisis or pre-phthisis bulbi</td>
<td></td>
</tr>
<tr>
<td>cT3b</td>
<td>Tumor invasion of choroid, pars plana, ciliary body, lens, zonules, iris, or anterior chamber</td>
<td></td>
</tr>
<tr>
<td>cT3c</td>
<td>Raised intraocular pressure with neovascularization and/or buphthalmos</td>
<td></td>
</tr>
<tr>
<td>cT3d</td>
<td>Hypoplasia and/or massive vitreous hemorrhage</td>
<td></td>
</tr>
<tr>
<td>cT3e</td>
<td>Aseptic orbital cellulitis</td>
<td></td>
</tr>
<tr>
<td>cT4</td>
<td>Extraocular tumor(s) involving orbit, including optic nerve</td>
<td></td>
</tr>
<tr>
<td>cT4a</td>
<td>Radiologic evidence of retrobulbar optic nerve involvement or thickening of optic nerve</td>
<td></td>
</tr>
<tr>
<td>cT4b</td>
<td>Extraocular tumor(s) clinically evident with proptosis and/or an orbital mass</td>
<td></td>
</tr>
<tr>
<td>cNX</td>
<td>Regional lymph nodes cannot be assessed</td>
<td></td>
</tr>
<tr>
<td>cN0</td>
<td>No regional lymph node involvement</td>
<td></td>
</tr>
<tr>
<td>cN1</td>
<td>Evidence of preauricular, submandibular, and cervical lymph node involvement</td>
<td></td>
</tr>
<tr>
<td>cM1</td>
<td>Distant metastasis without microscopic confirmation</td>
<td></td>
</tr>
<tr>
<td>cM1a</td>
<td>Tumor(s) involving any distant site (e.g., bone marrow, liver) on clinical or radiologic tests</td>
<td></td>
</tr>
<tr>
<td>cM1b</td>
<td>Pathological evidence of tumor in any distant site (e.g., bone marrow, liver) on clinical or radiologic tests</td>
<td></td>
</tr>
<tr>
<td>pM1</td>
<td>Distant metastasis with microscopic confirmation</td>
<td></td>
</tr>
<tr>
<td>pM1a</td>
<td>Pathological evidence of tumor in the cerebrospinal fluid or CNS parenchyma</td>
<td></td>
</tr>
<tr>
<td>pM1b</td>
<td>Pathological evidence of tumor in the CNS parenchyma</td>
<td></td>
</tr>
<tr>
<td>H</td>
<td>Hereditary Trait</td>
<td></td>
</tr>
<tr>
<td>HX</td>
<td>Unknown or insufficient evidence of a constitutional RB1 gene mutation</td>
<td></td>
</tr>
<tr>
<td>H0</td>
<td>Normal RB1 alleles in blood tested with demonstrated high-sensitivity assays</td>
<td></td>
</tr>
<tr>
<td>H1</td>
<td>Bilateral retinoblastoma, retinoblastoma with an intracranial primitive neuroectodermal tumor (i.e., trilateral retinoblastoma), patient with family history of retinoblastoma, or molecular definition of a constitutional RB1 gene mutation</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Pathological Definitions (pTNM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>pTX</td>
</tr>
<tr>
<td>pT0</td>
</tr>
<tr>
<td>pT1</td>
</tr>
<tr>
<td>pT2</td>
</tr>
<tr>
<td>pT2a</td>
</tr>
<tr>
<td>pT2b</td>
</tr>
<tr>
<td>pT3</td>
</tr>
<tr>
<td>pT3a</td>
</tr>
<tr>
<td>pT3b</td>
</tr>
<tr>
<td>pT3c</td>
</tr>
<tr>
<td>pT3d</td>
</tr>
<tr>
<td>pT4</td>
</tr>
</tbody>
</table>
risk of dissemination, which in turn correlates with patient survival.

Historically, the classification of retinoblastoma was divided into multiple intraocular retinoblastoma eye grouping (see 3.3.1) and patient staging (see 3.3.2) systems, still variously used and therefore making comparison between studies difficult. The 8th edition of the American Joint Committee on Cancer “Tumor, Node, Metastasis and Heritability” (TNMH) classification has recently been proposed to more accurately predict the salvage of the eye(s), metastasis and patient survival and holds the promise to provide a single universal classification (Mallipatna et al., 2017) (Table 3).

Surprisingly, despite the broad use of OCT in retinoblastoma patients (see 3.2.3), none of the existing classification systems have been adapted to consider possible subclinical findings at diagnosis. Moreover, although heavily-pretreated eyes are being increasingly offered salvage therapies, a classification of intraocular retinoblastoma at relapse is still missing and precludes the possibility to compare study outcomes of salvage treatments. This prompted us to add subclinical features at diagnosis in the TNMH classification system (see 3.3.1) as well as to propose a classification at relapse (see 3.3.3).

3.3.1. Grouping of the eye at presentation (RE, IIRC, ICRB, COG, TNMH)

A system to evaluate the intraocular extent of the disease was first introduced in the 1960s by Reese and Ellsworth, who stratified intraocular retinoblastoma into five groups (RE I to V), predicting eye survival in response to external beam radiotherapy (Reese and Ellsworth, 1963). The therapeutic shift from external beam radiotherapy to chemoreduction motivated the introduction by Linn Murphree, in 2005, of a novel International Intraocular Retinoblastoma Classification (IIRC), recapitulating the natural history of retinoblastoma into five groups (A to E) (Linn Murphree, 2005), with the presence of group E features being an absolute indication for enucleation. In 2006, this classification was revisited with some modifications by the group of Philadelphia (Shields et al., 2006a), followed in 2011 by the Children’s Oncology Group (COG) giving rise to the International Classification of Retinoblastoma (ICRB) and the COG, respectively. At the present time, the parallel use of 3 coexisting A to E classifications confuses the literature, especially regarding advanced eyes and impairs comparison of different clinical reports (Scelfo et al., 2017). For example, in a study of more than 2000 eyes (Kaliki et al., 2019), up to 56% of ICRB group E eyes would have been classified IIRC group D eyes, since IIRC group D eyes with tumor occupying 2/3 of the eye volume are reclassified ICRB group E eyes.

Although the TNMH classification includes the use of radiologic imaging to assess any optic nerve involvement (Mallipatna et al., 2017) (Table 3), it fails, like the previous ones, to allow grouping of invisible tumors only detected by OCT, as may be the case in the context of screening RB1 mutant carriers with a positive family history. We therefore propose to revise the TNMH classification by adding categories scT1a and b for subclinical staging: scT1a if at least further than 1.5 mm from the optic nerve head or 3 mm from fovea, and scT1b if closer (Fig. 4A).

3.3.2. Patient staging at presentation (IRSS, TNMH)

The first classification system for retinoblastoma aiming to cover the whole spectrum of the disease (IRSS) was introduced in 2006 by a multidisciplinary team of physicians from different countries and based on clinical evaluation, imaging and histopathology (Chantada et al., 2006a) (Table 5). Two years later, the same group standardized the processing of the enucleated eyes in order to obtain reliable histopathologic results and fresh tumor for molecular testing to define pathologic risk factors for extraocular relapse (Sastre et al., 2009). The recently introduced TNMH classification considers not only intraocular, extraocular and pathological disease but also, for the first time, the hereditary status and aims to become the gold standard reference in the future (Mallipatna et al., 2017).

The recommended procedures for assessing extraocular dissemination in retinoblastoma include contrast-enhanced MRI of the brain and orbits (de Graaf et al., 2012) as well as bone marrow and cerebrospinal fluid cytopathologic evaluation, with or without PCR-based detection of specific molecular biomarkers (Laurent et al., 2016) (see 3.7.1).

Staging investigations, such as lumbar puncture and bone marrow aspiration/biopsy are relatively invasive and require general anesthesia. They are not commonly performed in early stages (IRSS stage 0 or 1) where the risk of harboring disseminated disease is considered minimal. These investigations, as well as skeletal scintigraphy are, however, indicated in case of symptoms (pain, neurologic manifestations), exteriorization of the disease or high-risk pathology features reported after enucleation (Azar et al., 2003; Barai et al., 2004; Karciglu et al., 1997; Moscinski et al., 1996; Pratt et al., 1989). If conservative treatment is attempted, some groups advocate cytological evaluation of the cerebrospinal fluid in cases where the optic nerve cannot be visualized (Mallipatna et al., 2017). At the present time, however, a universal consensus especially for the metastatic work-up concerning conservatively-treated advanced intraocular disease is still missing. The development of more precise or less invasive techniques, such as molecular biomarkers in cerebrospinal fluid and bone marrow (Torbidoni et al., 2015), or functional imaging, would be ideal for diagnosis and follow-up of disseminated disease (Kobe et al., 2015).

3.3.3. Classification of retinoblastoma at relapse (RSU classification)

Similar to the classification of retinoblastoma at presentation, the classification of retinoblastoma at relapse aims to standardize appropriate treatment for relapse, achieve a better comprehension of treatment outcomes, predict prognosis for ocular salvage and define enucleation criteria for secondary enucleation. However, because recurrences do not necessarily involve the retina (R), but can be exclusively extraretinal with isolated seeding (S) or uveal involvement (U), it is not possible to classify them by applying one of the grouping

Table 4

<table>
<thead>
<tr>
<th>Group</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>Small tumors (≤3 mm) confined to the retina (no seeding), at least 3 mm from the fovea and 1.5 mm from optic nerve.</td>
</tr>
<tr>
<td>B</td>
<td>Tumors (&gt;3 mm) confined to the retina in any location, with clear subretinal fluid ≤ 5 mm from the tumor margin</td>
</tr>
<tr>
<td>C</td>
<td>Localized vitreous and/or subretinal seeding (&lt; 5 mm in total from tumor margin). If there is more than 1 site of subretinal/vitreous seeding, then the total of these sites must be ≤ 6 mm. Up to 1 quadrant subretinal seeding may be present.</td>
</tr>
<tr>
<td>D</td>
<td>Massive retinoblastoma with anatomic or functional destruction of the eye with one or more of the following:</td>
</tr>
<tr>
<td></td>
<td>• Neovascular glaucoma</td>
</tr>
<tr>
<td></td>
<td>• Massive intravitreal hemorrhage</td>
</tr>
<tr>
<td></td>
<td>• Aseptic orbital cellulitis</td>
</tr>
<tr>
<td></td>
<td>• Tumor anterior to anterior vitreous face</td>
</tr>
<tr>
<td></td>
<td>• Tumor touching the lens</td>
</tr>
<tr>
<td></td>
<td>• Diffuse infiltrating tumor</td>
</tr>
<tr>
<td></td>
<td>• Phthisis or pre-phthisis</td>
</tr>
<tr>
<td>E</td>
<td>Diffuse vitreous and/or subretinal seeding (≥5 mm in total from tumor margin). Seeding more extensive than Group C. Retinal detachment &gt; 1 quadrant</td>
</tr>
</tbody>
</table>
systems used at diagnosis. Therefore, we propose a specific classification for relapsing retinoblastoma based on its localization which is detailed in Table 6. In this classification, the recognition of tumor relapse does not only rely on clinical observation but also on imaging devices such as UBM (Chhablani et al., 2010; Stathopoulos et al., 2018b) or SD-OCT (Fabian et al., 2017b; Gaillard et al., 2018). The definition of relapse or recurrence refers to any tumor regrowth or new growth occurring after a minimum of 2 months progression-free and treatment-free follow-up at a monthly examination. In contrast, persistent, due to absence of response, or progressive disease despite ongoing treatments, as well as new primary retinal tumors, have to be distinguished from recurrence, and as such are not concerned by the proposed RSU-classification. Moreover, caution should be taken not to label as relapse, active disease diagnosed without an intermediate progression-free follow-up interval of at least two months since the last treatment, as in such cases disease persistence/progression cannot be excluded.

Prompt secondary enucleation is mandatory for relapse in the presence of persisting vitreous hemorrhage obscuring the fundus, neovascular glaucoma, phthisis bulbi or suspicion of extra-ocular extension.

### 3.4. Differential diagnosis

The list of conditions simulating retinoblastoma includes a broad spectrum of both acquired and congenital diseases (Balmer and Munier, 2007; Shields et al., 2013) (Table 7). In children seen in non-ocular oncology specialized centers, leukocoria, the main presenting sign of retinoblastoma, was due to congenital cataract in 25–65% of the cases and retinoblastoma in only 6–18% (Haider et al., 2008; Meier et al., 2006). In a retrospective review of nearly 2800 cases specifically

### Table 5

International retinoblastoma staging system (IRSS).

<table>
<thead>
<tr>
<th>Stage 0</th>
<th>Stage I</th>
<th>Stage II</th>
<th>Stage III: Regional extension</th>
<th>Stage IV: Metastatic disease</th>
</tr>
</thead>
<tbody>
<tr>
<td>Conservative treatment</td>
<td>Eye enucleated,</td>
<td>Eye enucleated,</td>
<td>A: overt orbital disease</td>
<td>A: Hematogenous metastases</td>
</tr>
<tr>
<td></td>
<td>Completely resected</td>
<td>Microscopic residual</td>
<td>B: preauricular or cervical lymph node extension</td>
<td>1. Single lesion</td>
</tr>
<tr>
<td></td>
<td>histologically</td>
<td>tumor</td>
<td></td>
<td>2. Multiple lesions</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>B: CNS extension</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>1. Prechiasmatic lesion</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>2. CNS mass</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>3. Leptomeningeal disease</td>
</tr>
</tbody>
</table>

NB: In cases of bilateral disease, the staging depends on the eye with more advanced disease. Abbreviation: CNS = central nervous system.


### Table 6

Retinoblastoma classification at relapse (RSU-Classification).

<table>
<thead>
<tr>
<th>Category</th>
<th>Subcategory</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>RX</td>
<td>Retinal relapse not assessable due to opaque media</td>
<td></td>
</tr>
<tr>
<td>R0</td>
<td>No retinal relapse</td>
<td></td>
</tr>
<tr>
<td>R1</td>
<td>Intra-retinal relapse</td>
<td></td>
</tr>
<tr>
<td>R1a</td>
<td>Localized (accessible to focal therapy including brachytherapy) retinal relapse at least &gt; 3 mm from the foveola and 1.5 mm from the optic nerve head</td>
<td></td>
</tr>
<tr>
<td>R1b</td>
<td>Diffuse retinal relapse (any retinal relapse which is not localized) or any retinal relapse contiguous to the foveola (≤3 mm) or the optic nerve head (≤1.5 mm)</td>
<td></td>
</tr>
<tr>
<td>SX</td>
<td>Seeding not assessable due to opaque media</td>
<td></td>
</tr>
<tr>
<td>S0</td>
<td>No seeding relapse</td>
<td></td>
</tr>
<tr>
<td>S1</td>
<td>Subretinal seeding relapse</td>
<td></td>
</tr>
<tr>
<td>S1x</td>
<td>Subretinal seeding not assessable due to opaque media</td>
<td></td>
</tr>
<tr>
<td>S1a</td>
<td>Localized subretinal seeding (≤1 quadrant), at least &gt; 3 mm from the foveola and 1.5 mm from the optic nerve head</td>
<td></td>
</tr>
<tr>
<td>S1b</td>
<td>Diffuse (&gt; 1 quadrant) subretinal seeding or any subretinal seeding contiguous to the foveola (≤3 mm) and/or the optic nerve head (≤1.5 mm)</td>
<td></td>
</tr>
<tr>
<td>S2</td>
<td>Intravitreal seeding relapse</td>
<td></td>
</tr>
<tr>
<td>S2x</td>
<td>Intravitreal seeding not assessable due to opaque media</td>
<td></td>
</tr>
<tr>
<td>S2a</td>
<td>Localized vitreous and/or retrohyaloid seeding (≤3 mm from retinal tumor)</td>
<td></td>
</tr>
<tr>
<td>S2b</td>
<td>Diffuse vitreous and/or retrohyaloid seeding (any vitreous and/or retrohyaloid seeding which is not localized)</td>
<td></td>
</tr>
<tr>
<td>S3</td>
<td>Aqueous seeding relapse</td>
<td></td>
</tr>
<tr>
<td>UX</td>
<td>Uveal relapse not assessable due to opaque media and no UBM/MRI evaluation</td>
<td></td>
</tr>
<tr>
<td>U0</td>
<td>No uveal relapse</td>
<td></td>
</tr>
<tr>
<td>U1</td>
<td>Choroidal relapse</td>
<td></td>
</tr>
<tr>
<td>U1a</td>
<td>Focal choroidal relapse (≤3 mm in largest diameter)</td>
<td></td>
</tr>
<tr>
<td>U1b</td>
<td>Massive choroidal relapse (&gt; 3 mm in largest diameter)</td>
<td></td>
</tr>
<tr>
<td>U2*</td>
<td>Intraciliary relapse (* = number of clock hours involved)</td>
<td></td>
</tr>
<tr>
<td>U3</td>
<td>Iris relapse</td>
<td></td>
</tr>
</tbody>
</table>
referred for suspected retinoblastoma in a tertiary center in the United States of America, 78% had a confirmed retinoblastoma whereas 22% suffered a different disease, dominated by Coats’ disease (Shields et al., 2013). The latter has notably the particularity of mimicking retinoblastoma at different stages, including endophytic macular retinoblastoma group B in Coats’ disease stage IIb, characterized by the presence of a subfoveal pseudo-tumor measuring up to 2.5 mm and sometimes associated with pseudo-feeding vessels (Darui et al., 2017), or exophytic retinoblastoma group D and E in Coats’ disease stages IIIb and IV, respectively (Gaillard et al., 2014). In the majority of cases, a comprehensive history with thorough clinical examination and appropriate ancillary tests is usually sufficient to avoid misdiagnosis and undesirable treatments (Shields et al., 2013). The indication for fine needle biopsy should be limited to extraordinarily unusual circumstances (Karcioglu, 2002).

3.5. Prenatal diagnosis of retinoblastoma

Early screening for retinoblastoma is advised for children with a positive family history of the disease. Intra-uterine and early postnatal retinoblastoma lesions are predominantly located in the posterior pole, where they are more likely to compromise vision (Kivela and Hadjistilianou, 2017; Soliman et al., 2016) and known for their rapid growth with an estimated doubling time of 2 weeks (Shah et al., 2010).

Before genetic testing and the use of modern imaging techniques (high frequency ultrasound probes and MRI), diagnostic screening for retinoblastoma was performed postnatally with indirect ophthalmoscopy (Abramson et al., 2002). With the introduction of RB1 mutation testing, offspring of parents with a known mutation can now be screened during pregnancy. Currently, DNA extraction for mutational analysis can be proposed either by chorionic villus sampling at the end of the first trimester (11 weeks of gestation) or by amniocentesis during the second trimester (16 weeks of gestation). Parents should be informed of a procedure-related potential miscarriage risk of 0.22% and 0.11% after chorionic villus sampling and amniocentesis, respectively (Akolekar et al., 2015). Confirmation of a fetal RB1 mutation allows parents to make an informed decision as to whether or not to proceed with the pregnancy. Standards for the follow-up of a fetus with high-risk to develop the disease do not currently exist. In Lausanne, for parents who decide to proceed with the pregnancy despite confirmation of a fetal RB1 mutation, for or those who refused amniocentesis, fetal ultrasonography is performed at 26 weeks of gestation and repeated every 3 weeks until 32 weeks, and then every 2 weeks until 36 weeks of gestation. The ultrasonography includes standard 2D and 3D images of fetal anatomy acquired through a magnified axial view of both eyes (Fig. 6). The 3D acquisition of volume is an integral part of our evaluation as it allows systematic analysis of the posterior wall of the eye. An alternative imaging detection method is fetal MRI, which, compared to ultrasonography, has a higher contrast resolution but is more affected by fetal motion (Staffieri et al., 2015). In case of tumor detection, a consultation with an ocular oncologist, perinatologist and neonatologist is needed to balance the burden of an early term delivery, preceded or not by pulmonary maturation therapy, against the oncologic risks of management of a full-term delivery.

In practice, only few cases of fetal retinoblastoma have been published. Two were very large lesions that deformed not only the globe but also the face (Maat-Kievit et al., 1993; Salim et al., 1998). Five occurrences of small intraocular retinoblastomas were reported either by ultrasonography (Paquette et al., 2012; Soliman et al., 2016; Stathopoulos et al., 2018c; Toi et al., 2003) or by MRI (Staffieri et al., 2015). Ultrasonography is not sensitive enough to detect tumors measuring less than 2 mm in thickness. In a fetus carrying a RB1 mutation, two lesions of 1.2 and 1.9 mm, respectively, were detected at the posterior pole in each eye by MRI performed at 35 weeks of gestation (Staffieri et al., 2015). In a prospective study of the use of both ultrasonography and MRI in 6 high risk fetuses, fetal MRI did not detect any tumors, whereas one could be seen on ultrasonography, possibly because of the limitation of slice thickness (Paquette et al., 2012).

A recent retrospective observational study suggested that induced early delivery (by 37 weeks of gestation), which remains a controversial procedure (Gombos, 2012), might benefit the child carrying a mutated RB1 in terms of treatment-related morbidity and ultimate salvage of vision (Soliman et al., 2016). Noteworthy, when retinoblastoma is diagnosed during the neonatal period, at least half of the children initially have a unilateral rather than bilateral tumor, typically group B, but almost all germline mutation carriers later progress to bilateral involvement (Kivela and Hadjistilianou, 2017; Soliman et al., 2016). On the other hand, all infants with neonatal retinoblastoma have at least one eye with a tumor close to the foveola, whereas the macula of the fellow eye is often spared and loss of reading vision from both eyes is consequently infrequent (Imhof et al., 2006; Kivela and Hadjistilianou, 2017; Soliman et al., 2016).

Fig. 6. A. Prenatal detection of a macular tumor in the left eye by transabdominal ultrasonography (5 MHz) at 33 weeks of gestation. B. Postnatal ultrasonography (10 MHz) obtained after provocation at 36 weeks of gestation showing the left macular tumor on ultrasonography and (C) fundus photography. D-F. Corresponding imaging of the right eye showing a tiny posterior pole lesion sparing the fovea detectable only post-natally on both ultrasound and fundus photography (infero-temporal to the fovea). (Images courtesy of: Dr. Ants Toi, Medical Imaging, Mt. Sinai Hospital and Dr. Brenda Gallie, Ophthalmology and Vision Sciences. Hospital for Sick Children. University of Toronto. Canada).
3.6. Trilateral retinoblastoma

Approximately 3.5% of patients with hereditary retinoblastoma will develop an intracranial tumor, located most often in the pineal gland and less frequently in the suprasellar or parasellar region (de Jong et al., 2014a) (Fig. 5A and B). Such a primitive neuroectodermal tumor associated with hereditary retinoblastoma was first described in 1977 (Jakobiec et al., 1977) and in 1982, named trilateral retinoblastoma (Bader et al., 1982). Originally considered to have a dismal prognosis (Kivela, 1999; Paulino, 1999), sophisticated imaging allowing earlier detection and improved chemotherapy regimens have, since 1995, significantly increased 5-year survival rates from 6 to 44% for pineal tumors and from 0 to 57% for non-pineal tumors (de Jong et al., 2014a). On the other hand, understanding that children frequently have a cystic, non-neoplastic pineal gland with radiological features that differ from pineoblastoma, has helped to avoid over-diagnosis of trilateral retinoblastomas (Barboriak et al., 2001; Beck Popovic et al., 2006).

3.6.1. Incidence, early detection and screening

The risk period of developing trilateral retinoblastoma extends from birth to 7 years of age (de Jong et al., 2014a; Kivela, 1999), with 95% of the cases diagnosed before the age of 5 (de Jong et al., 2015b). Patients with hereditary retinoblastoma have a higher risk of developing trilateral disease compared to those with non-hereditary disease, the latter carrying the same risk as the normal population. A recent meta-analysis restricted to cohorts with at least 100 retinoblastoma patients estimated the risk of developing trilateral disease to be 3.8% in bilateral disease (2.9% for pineal tumors) and 3.5% (3.2% for pineal tumors) when considering all hereditary retinoblastoma, including unilateral cases with a positive family history (de Jong et al., 2015b). This naturally leads to the question of the need and the frequency at which one should screen for trilateral retinoblastoma with MRI.

Trilateral retinoblastomas detected synchronously with intraocular retinoblastoma are usually smaller in size, asymptomatic and have a better prognosis compared to those detected metachronously (de Jong et al., 2014a; Kivela, 1999). Five-year survival is also more favorable if the trilateral tumor is pineal and less than 15 mm in size (de Jong et al., 2014a). Baseline screening is therefore currently considered justified by most authors (De Ioris et al., 2014; de Jong et al., 2014a; Rodjan et al., 2012), but cost-effectiveness of subsequent imaging remains controversial (Moll et al., 2002).

3.6.2. Influence of the retinoblastoma treatment on the incidence of trilateral retinoblastoma

Previous radiotherapy for retinoblastoma, especially before the age of 12 months, has been associated with a potentially higher incidence of pineal trilateral retinoblastoma in patients with hereditary retinoblastoma, even though the pineal gland is usually not (directly) within the field of radiation (Moll et al., 2002). The question of whether previous systemic chemotherapy is protective of developing trilateral retinoblastoma is still debated with reports that favor (Shields et al., 2001) or challenge such a preventive effect (Chantada et al., 2014). In the current era of intra-arterial and intravitreal chemotherapy, the likelihood to develop trilateral retinoblastoma does not seem to have increased, with a reported incidence of 2.7% at 5 years follow-up (Habib et al., 2018).

3.6.3. Improved treatment of trilateral retinoblastoma: high-dose chemotherapy

Trilateral retinoblastoma used to be treated with surgery, chemotherapy, and radiotherapy (Blach et al., 1994; Jubran et al., 2004). In 2010, a multimodality approach to metastatic retinoblastoma with intensive induction of chemotherapy, high-dose chemotherapy and stem cell reinfusion was reported to provide a 5-year survival of 67% in a small series of patients with non-central nervous system (CNS) metastases (Dunkel et al., 2010c) and a cure for only 1 out of 6 patients with CNS dissemination from trilateral retinoblastoma (Dunkel et al., 2010b). This led to a collaborative, prospective clinical trial for metastatic or trilateral retinoblastoma by the Children’s Oncology Group (COG) and Grupo de America Latina de Oncología Pediátrica (GALOP). The treatment consisted of four courses of intensive chemotherapy and, according to response, a three-drug high-dose chemotherapy regimen with stem cell reinfusion, with or without radiotherapy. The results recently confirmed a 3-year event-free survival of 79% after non-CNS metastases, whereas the survival rate remained at an unsatisfactory 8% after CNS metastases or trilateral retinoblastoma (Dunkel et al., 2017).

3.6.4. Differential diagnosis: pineal cysts

Technological progress in imaging has contributed to an increased observation of pineal cysts, a benign finding reported in more than half of the pediatric population using 3 T MRI (Whitehead et al., 2013). These lesions must be distinguished from trilateral retinoblastomas in order to avoid the administration of unnecessary chemotherapy. On 3 T MRI, typical benign cysts are characterized by an enlarged pineal gland with a hypointense central region on T1-weighted images, isointense on T2-weighted images with a thin wall that enhances after gadolinium injection, and no nodular structures (Barboriak et al., 2001; Whitehead et al., 2013). They can increase in size during the first year of observation as the child grows, and then stabilize or decrease in size (Fig. 5E–G) until the age of 4–5 years (Sirin et al., 2016). Pineal cysts in retinoblastoma patients have a reported incidence ranging between 5 and 37%, with some authors indicating an association with hereditary disease (Beck Popovic et al., 2006; Ruiz Del Rio et al., 2014), whereas others do not (Ramasubramanian et al., 2013). They are comparable in size with cysts in age-matched healthy controls (Pham et al., 2015), and do not progress to a tumor (Barboriak et al., 2001; Beck Popovic et al., 2006; Karatza et al., 2006; Ramasubramanian et al., 2013).

Contrary to typical pineal cysts, nodular pineal cysts can transform to pineoblastoma (de Jong et al., 2014b) (Fig. 5 C, D). Moreover, in some instances pineoblastomas can also show a cystic part. Any unusual feature in a pineal cyst calls therefore for close follow-up (Rodjan et al., 2012). A retrospective analysis (de Graaf et al., 2012) by the European Retinoblastoma Imaging Collaboration (ERIC) recently resulted in a description, terminology and classification that allows clinicians to distinguish benign pineal cysts from those that need to be followed because of risk of transformation to pineoblastoma (Sirin et al., 2016).

3.7. Metastatic retinoblastoma

Metastatic retinoblastoma at presentation is rare in high-income countries, but represents the most frequent cause of death for this tumor worldwide (Canturk et al., 2010). In high-income countries, the few cases seen are those developing metastasis after initial treatment with enucleation or conservative therapy. The median time between diagnosis of intraocular retinoblastoma primarily enucleated with high pathologic risk factors and first evidence of metastases is 10 months (range: 2–24 months) in patients receiving adjuvant chemotherapy, but appears to occur earlier at 5 months in untreated ones (Antonelli et al., 2003; Chantada et al., 2010; Cozza et al., 2009; Lu et al., 2018; MacKay et al., 1984; Rodriguez-Galindo et al., 2003). In low-income countries, metastatic dissemination is generally already observed at diagnosis in older children who present with extensive orbital masses or after parents declined enucleation of an eye with intraocular disease (Canturk et al., 2010), and is attributed to delayed diagnosis/treatment rather than a difference in tumor biology (Chawla et al., 2016; Leal-Leal et al., 2006).

3.7.1. Diagnosis of metastatic retinoblastoma

Retinoblastoma may metastasize to the CNS, bone, bone marrow or lymph nodes and, rarely, to other sites such as the liver. The diagnosis of metastatic disease can occur in asymptomatic patients during a work-
up motivated by clinical features at presentation, presence of classic high-risk pathologic findings (Sastre et al., 2009), or possibly by the degree of anaplasia (Mendoza et al., 2015) after enucleation, or in symptomatic patients during/after conservative treatment. In most cases of metastatic disease at diagnosis, massive orbital extension occurs along with distant metastatic dissemination (Kaliki et al., 2017). The detection of minimally disseminated disease in bone marrow and cerebro-spinal fluid at diagnosis and follow-up can be now evaluated by means of quantitative polymerase chain reaction of cone-rod homeobox (CRX) transcription factor messenger RNA (Laurent et al., 2016).

**CNS metastasis.** CNS involvement occurs mainly by direct extension of the tumor via the optic nerve or the sub-arachnoidal space (leptomeningeal dissemination), but can also be parenchymal or paraspinal through hematogenous dissemination, and rarely as a direct CNS extension through a facial bone. In the case of trilateral retinoblastoma, the tumor may also disseminate to the cerebrospinal fluid from a midline mass (Popovic et al., 2007), typically resulting in a leptomeningeal dissemination (Laurent et al., 2013). Interestingly, molecular studies using polymerase chain reaction for detecting minimally disseminated disease showed that cerebrospinal fluid relapses occurred more commonly as an isolated event with negative bone marrow, suggesting that the route of dissemination to the cerebrospinal fluid in retinoblastoma is mainly through the optic nerve and independent of systemic dissemination (Torbidoni et al., 2015) or due to trilateral seeding (Torbidoni et al., 2018). However, seeding by subclones arising from the systemic circulation at diagnosis, which persists during therapy, cannot be ruled out. Affected subjects may show increased intracranial pressure resulting in headache, irritability, vomiting, vision loss, and less commonly, focal neurological signs. Clinical examination includes evaluation of the orbit and facial bones as well as evaluation of the whole neuroaxis. CNS dissemination is investigated by both cerebrospinal fluid cytology (Bakhshi et al., 2011) and imaging, with the latter being performed prior to diagnostic lumbar puncture in order to exclude intracranial hypertension. A cytospin of cerebrospinal fluid cytology typically shows retinoblastoma cells in clusters. If isolated, these cells can, however, be missed or confused with normal lymphocytes in young children. For this reason, the cytospin should always be examined microscopically, and malignant cells confirmed with immunocytochemistry for GD2 or CRX. If this procedure is not available, flow cytometry examination, usually available in centers that also treat pediatric leukemia, can be used to detect the non-hematopoietic origin of the cells by showing the lack of expression of molecules such as CD45 antigen with positivity for CD56 (Shen et al., 2013).

**Bone and bone marrow metastasis.** Bone metastasis may present as a painful and palpable mass in the long bones, but facial bones may also be affected, as well as the diploe of the neurocranium. Bone marrow invasion may occasionally be suspected by detecting abnormal blood counts, especially thrombocytopenia, but is more commonly detected in asymptomatic patients by bone marrow biopsy or on imaging such as MRI or positron-emission tomography (Chantada et al., 2006b). Bone marrow evaluation needs at least 2 aspirates and 2 biopsies and, if available, immunocytochemistry using markers like GD2 (Chantada et al., 2006b) or reverse transcriptase quantitative PCR for CRX mRNA should be performed (Torbidoni et al., 2015).

### 3.7.2. Treatment of metastatic retinoblastoma

Although retinoblastoma is a chemosensitive tumor, metastatic disease is not curable with conventional treatment and is therefore considered to have a poor prognosis, especially in cases of CNS involvement (Kaliki et al., 2017; Pant et al., 2017). High dose chemotherapy, followed by autologous stem cell rescue with radiation to the sites of bulky disease, has been the only treatment reported to date to potentially cure these patients (Dunkel et al., 2010c; Palma et al., 2012), allowing remission in close to 70% of children in a series of 15 patients with metastatic retinoblastoma not involving the CNS (Dunkel et al., 2010c), whereas those with CNS involvement have only anecdotally benefitted from it (Dunkel et al., 2010a). Focusing on intrathecal chemotherapy either with topotecan (Laure et al., 2005) or radio-immunotherapy with radiolabeled anti-GD2 antibody (Kramer et al., 2010) may improve the currently dismal outcome.

#### 3.7.3. Influence of conservative retinoblastoma treatment in the occurrence of metastasis

While reducing the enucleation rate, the transition from one therapeutic era to the next also brought great concerns regarding a potential negative impact on the “state of metastatic grace”, an empiric window for globe preserving therapies, where the risk for metastasis is not higher than the risk encountered following enucleation with adjuvant therapy in case of histopathologic high risk factors. Fear of inappropriate management of secondary enucleated eyes due to downstaging after chemotherapy on the one hand (Zhao et al., 2011), and of the possible consequences of lower systemic chemotherapy exposure in patients treated with intra-arterial chemotherapy alone on the other hand (Yousef et al., 2016), is nourished by the lack of well-designed studies aimed at capturing metastatic events and related deaths.

Among the few studies which have addressed this crucial issue, two compared the incidence of metastatic disease and orbital recurrence in advanced retinoblastoma group D/E treated with either systemic or intra-arterial chemotherapy versus primary enucleation and concluded to the absence of variation by treatment modality (Berry et al., 2017b; Yannuzzi et al., 2015). In another paper comparing patients primarily treated by external beam radiotherapy versus first line chemotherapy (Chantada et al., 2014), the authors failed to detect a change in the probability of extraocular relapse, despite the prediction of a putative positive effect of chemoreduction in preventing systemic metastasis in children with microscopically disseminated disease. The controversy regarding the rate of metastases following first line intra-arterial versus intravenous chemotherapy for advanced disease clashes with biases linked to the fact that these two administration routes influence the ratio of CNS versus non-CNS dissemination (Brennan et al., 2015).

Neoadjuvant systemic chemotherapy may offer a less favorable coverage in patients with high risk for CNS involvement when compared to the unmatched chemotherapeutic optic nerve concentration achieved by intra-arterial drug delivery (Taich et al., 2016), while intravenous chemotherapy may be more efficient against distant minimally disseminated disease. Although the end result in terms of survival remains unknown to date, patient survival following the introduction of first line intra-arterial chemotherapy appears, however, unimpaired compared to previous treatment regimens, with no increase in the metastasis rate, ranging between 2 and 2.5% (Funes et al., 2018) and death rate from metastatic disease around 1% (Abramson et al., 2017b). Noteworthy, in all cases, the cause of metastases resulting in death was parental refusal of enucleation of a single remaining eye (Abramson et al., 2017b).

### 3.8. Second primary neoplasms

Retinoblastoma patients with a germline RB1 mutation are at lifetime risk to develop various types of second primary (Abramson et al., 1979; Eng et al., 1993; Wong et al., 1997) and even third, fourth and fifth malignancies (Abramson et al., 2001; Marees et al., 2010) (see 3.8.1). This risk is further increased after external beam irradiation and, to a lesser extent, chemotherapy (see 3.8.3.2). In middle and high-income countries where the majority of patients survive retinoblastoma, second primary tumors represent the leading cause of death in patients with hereditary retinoblastoma (Kleinerman et al., 2005; Marees et al., 2009; Möll et al., 1997, 2001). Despite various follow-up strategies for retinoblastoma survivors (Baker et al., 2016), specific screening protocols are still missing (see 3.8.4).

#### 3.8.1. Incidence and mortality of second primary neoplasms

According to long-term retinoblastoma survivor cohort studies from
around the world (Marees et al., 2010), the risk of second primary neoplasms is much higher for hereditary than for non-hereditary retinoblastoma survivors, for whom the standardized incidence ratio is comparable to that of the general population (Table 8). This risk also increases significantly with a longer follow-up, in particular for epithelial cancers such as lung, bladder or breast carcinomas. Thus, according to a Dutch study, the 40-year absolute excess risk of second primary neoplasms in hereditary retinoblastoma survivors was 261 excess cases per 10,000 survivors per year with a cumulative incidence of 28% (95% confidence interval = CI (21.0–35.0%)) (Marees et al., 2008).

The cumulative mortality from second primary malignancies 50 years after retinoblastoma diagnosis was 25.5% (95% CI (21–30%)) in hereditary versus 1% (95% CI (0.2–1.8%)) in non-hereditary retinoblastoma survivors (Marees et al., 2009; Yu et al., 2009).

3.8.2. Characteristics of secondary primary neoplasms

Second primary tumors can develop in diverse anatomic locations, including the skull, bones, soft tissues, nasal cavity, skin, orbit, brain, breast, bladder and lung. In a review of 676 secondary primary malignancies in 602 retinoblastoma survivors, the most prevalent neoplasms were sarcoma (68%), carcinoma (14%), melanoma (8%), leukemia and lymphoma (4%) (Woo and Harbour, 2010). The latency to develop a second tumor depends on the secondary primary tumor type and previous treatment. The median age at diagnosis was 13 years for sarcoma, 27 years for melanoma and 29 years for carcinomas (Woo and Harbour, 2010). In a cohort of German pediatric retinoblastoma survivors, the most prominent second primary neoplasms were sarcoma and leukemia with a standardized incidence ratio of 148 (95% CI (39.81–378.87)) and 41 (95% CI (11.13–105.95)) respectively (Temming et al., 2015). In this study, a different latency in the diagnosis was observed for solid versus non-solid second primary neoplasms, diagnosed in children older or younger than 5 years of age, respectively (Temming et al., 2015). However, in this cohort, the increased risk of leukemia could be due to the adjunct of systemic alkylating agents in the German protocol (Kunkele et al., 2015).

The diagnosis of a second primary neoplasm needs to be confirmed by cytopathologic analysis of a tumor biopsy. Small round blue cell tumors such as lymphoma, rhabdomyosarcoma or nephroblastoma, however, can be hard to distinguish from metastatic retinoblastoma. In such cases, the presence or absence of previously identified somatic RB1 mutations and/or retinoblastoma-specific somatic copy number changes can help differentiate second cancers from metastatic disease (Racher et al., 2016).

3.8.3. Factors influencing the risk of secondary primary neoplasms

3.8.3.1. Genetic predisposition. Retinoblastoma survivors carrying a germline mutation, treated with enucleation or focal treatment with no additional radiation or chemotherapy, are clearly at increased risk to develop second primary tumors (Abramson et al., 1979; Eng et al., 1993; Wong et al., 1997).

In a study investigating variations in second cancer risk by family history of retinoblastoma among long-term survivors, the authors concluded that carriers of an inherited germline mutation are at slightly higher risk compared with those with a de novo germline mutation (Kleinerman et al., 2012), which may be partially attributed to mosaicism or reduced penetration (Moll et al., 2012). Preliminary research also suggested an increased second primary malignancy risk in a group with a nonsense RB1 mutation (hazard ratio or HR = 3.53 (95% CI (1.82–6.84)) compared to bearers of a low penetrance RB1 mutation (HR = 0.19 (95% CI (0.05–0.81))) (Dommering et al., 2012). However, this observation was recently challenged, at least in irradiated hereditary patients, suggesting no protective effect in low penetrant RB1 mutation carriers (Chaussade et al., 2019).

3.8.3.2. Influence of therapy

3.8.3.2.1. Radiotherapy. External beam irradiation increases the risk of developing second primary malignancies in patients with hereditary retinoblastoma threefold (Kleinerman et al., 2005; Marees et al., 2008; Temming et al., 2017), especially bone cancers and soft tissue sarcomas in a dose-dependent manner (Fletcher et al., 2004; Wong et al., 1997), or if given before the age of one year (Abramson and Frank, 1998; Moll et al., 2001). More recent radiotherapeutic modalities with proton radiotherapy and/or stereotactic conformal radiotherapy are expected to induce fewer second primary neoplasms in hereditary retinoblastoma survivors compared to conventional radiotherapy (Munier et al., 2008; Pica et al., 2011; Sethi et al., 2014).

3.8.3.2.2. Chemotherapy. There is still little long-term information about the contribution of chemotherapy alone on the risk of second primary malignancies in hereditary retinoblastoma patients. The cumulative incidence ratio in hereditary retinoblastoma treated with chemotherapy alone (3.0, CI 0.3–10.8) appears, however, to be comparable to patients treated with enucleation or focal treatment alone and 3 times lower compared to patients receiving external beam irradiation (Temming et al., 2017). Some have reported a potential acute risk to develop at an early time point acute myelogenous leukemia after systemic chemotherapy (Chantada et al., 2014; Gombos et al., 2007; Temming et al., 2015), while this was not confirmed by others for chemoreduction with vincristine, etoposide and carboplatin (Turaka et al., 2012). However, etoposide-related secondary leukemias with 11q23 involvement have been reported in the literature even in patients who received what are considered safe cumulative dosages for other pediatric tumors (Chantada et al., 2014; Gombos et al., 2007).

Targeted chemotherapy (periocular, intra-arterial and intravitreal) does not seem to increase the rate of secondary primary malignancies (Ishida et al., 2016; Suzuki et al., 2011), although follow-up time is too short to draw definite conclusions. Recently, a review of 214 patients with hereditary retinoblastoma treated by intra-arterial chemotherapy over a 10-year period revealed a comparable rate of second primary neoplasm to previously published ones (Habib et al., 2018).

3.8.4. Long-term follow-up in heritable retinoblastoma survivors

While some institutions routinely perform annual MRI of the head or total body MRI screening (Friedman et al., 2014), others perform complementary exams only when indicated. In general, retinoblastoma survivors undergo as many, but not more, cancer screening tests compared to the general population (Sheen et al., 2008). Regular skin examination for melanoma is important, especially for patients with impaired vision. Overall, the most important aspect of screening starts by educating the patient, family and (primary) doctors on the increased risk and on common signs of second primary tumors (Dimaras et al., 2015).

Table 8

Incidence of second primary neoplasms in different countries.

<table>
<thead>
<tr>
<th></th>
<th>Standardized incidence ratio (95% Confidence interval)</th>
<th>Absolute excess risk of second primary tumors per 10,000 survivors per year</th>
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<tbody>
<tr>
<td>Hereditary retinoblastoma survivors</td>
<td>Non-hereditary retinoblastoma survivors</td>
<td></td>
</tr>
<tr>
<td>MacCarthy et al. (2013)</td>
<td>Great Britain 13.7 (11.3–16.5) 1.5 (0.9–2.3)</td>
<td>58</td>
</tr>
<tr>
<td>Kleinerman et al. (2005)</td>
<td>USA 19 (16-21) 1.2 (0.7–2.0)</td>
<td>97</td>
</tr>
<tr>
<td>Marees et al. (2008)</td>
<td>Holland 20.4 (15.6–26.1) 1.85 (0.96–3.24)</td>
<td>68</td>
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</table>
4. Retinoblastoma genesis

Like any cancer, the biological and clinical behavior of retinoblastoma is dictated by the unique features of the cancer cells and their interactions with the host. In turn, the cancer cell features are largely determined by the intrinsic gene expression program of the retinoblastoma cell-of-origin in combination with changes to the cell-of-origin signaling circuitry that are induced by oncogenic mutations. However, in contrast to most cancers, the genetic changes that initiate retinoblastoma are well defined, and the mechanisms through which the retinal cell-of-origin transforms into a cancer are coming into focus. This chapter describes how our understanding of retinoblastoma genesis has come about, including the identification of the RB1 gene, genomic alterations beyond RB1 mutations, the origin of retinoblastoma from cone photoreceptor precursors, and collaboration of RB1 mutations with the underlying cone precursor circuitry.

4.1. Mutational inactivation of the RB1 gene initiates retinoblastoma tumorigenesis

Based on a statistical study of the distribution of the number of tumors and age at diagnosis in 48 patients with retinoblastoma, Alfred Knudson hypothesized that retinoblastoma is caused by two mutational events (Knudson, 1971). It was not part of Knudson’s original hypothesis that these alterations target two alleles of an autosomal gene locus. However, analysis of genotype changes at genetic polymorphisms in tumors, specifically the pattern of loss of constitutional heterozygosity (LOH), indicated that two mutational events target a gene on chromosome 13q (Cavenee et al., 1983).

In 1986, fifteen years after Knudson’s initial paper, RB1 was identified as the gene that is the target of two-step mutational inactivation in retinoblastoma (Friend et al., 1986; Fung et al., 1987; Lee et al., 1987). Fulfilling the predictions of Knudson’s hypothesis, heritable and non-heritable retinoblastomas are distinct with respect to the timing of the two mutational events that are required to alter both RB1 alleles (see 5.1).

In more recent years, comprehensive genomic analysis has confirmed that, as far as genetic alterations are concerned, bi-allelic inactivation of RB1 may be sufficient for retinoblastoma genesis (Grobner et al., 2018; Zhang et al., 2012). Indeed, somatic chromosomal copy number alterations are absent in most retinoblastomas from children who are diagnosed at an early age (Herzog et al., 2001), and exome sequencing analyses revealed that some early tumors lack gene mutations other than RB1 as well as chromosomal copy number alterations (Kooi et al., 2016a). However, the question of whether this implies that all retinoblastomas initially form without further genomic alterations remains open.

4.2. Most retinoblastomas show more than two-hits

Although early tumors may develop with biallelic RB1 loss as the sole genomic alteration, cytogenetic meta-analyses of several hundred retinoblastomas have shown recurrent chromosome 1q, 2p, and 6p gains and 16q losses (Corson and Gallie, 2007; Kooi et al., 2016b). Genomic gains on the short arm of chromosome 2 appear to target the MYCN locus. The targets of genomic gains at 1q and 6p are not well defined, but initial candidates included MDM4, KIF14 on 1q and DKK and E2F3 on 6p (Corson and Gallie, 2007). A meta-analysis identified other genes in the peak gain regions with significant gene-dosage effects, including ZBTB41, CRB1 and NEK7 on 1q and SOX4 on 6p (Kooi et al., 2016b). Numerous genes have been proposed to underlie the frequent 16q loss such as CDH11, encoding a cadherin possibly involved in cell-cell attachments and RB1-like 2 (RB2L2) encoding the pRB-related p130 protein, which have decreased expression in tumors with 16q loss (Kooi et al., 2016b). These chromosome 16q changes appear to be associated with vitreous seeding, yet a causal link has not been established (Gratias et al., 2007). Exome analyses also revealed recurrent focal deletion or mutational inactivation of BCOR, a gene on the X chromosome that encodes a non-canonical polycomb repressor complex 1 component in about 10% of retinoblastomas and a recurrent mutation of CREBBP at even lower frequency (Kooi et al., 2016a; Zhang et al., 2012).

At present, it is unclear whether the recurrent genomic changes (1q, 2p, and 6p gains, 16q losses, and BCOR and CREBBP inactivation) are required for the initial development of the retinoblastomas in which they are found, or if they are selected after the initial appearance of a tumor in order to mediate more rapid growth. Motivated by the observation that genomic gains and LOH are often present in addition to bi-allelic RB1 gene inactivation, a multitstep model of development of retinoblastoma was suggested (Corson and Gallie, 2007; Gallie et al., 1999). Further support for such a model was drawn from the observation that samples from tumor areas that were classified as retinoblastomas based on histomorphologic findings, showed bi-allelic RB1 inactivation and relatively few chromosomal copy number changes, whereas adjacent areas with histomorphologic features of retinoblastoma had more chromosomal gains and losses (Dimaras et al., 2008; Sampieri et al., 2009). Likewise, the larger tumors typical of later diagnosis have more chromosomal alterations as well as decreased differentiation-related gene expression, suggesting that retinoblastomas progress via sequential acquisition of chromosomal and gene expression changes (Cobrinik, 2015; Kooi et al., 2015). However, subsequent analyses indicated that chromosomal change did not appear in a particular order (Kooi et al., 2016b). Thus, the multistep evolution of retinoblastoma may differ from the prototypical adenoma-carcinoma sequence in colon cancer in which chromosomal alterations appear in a specific sequence (Vogelstein et al., 1988).

A better understanding of retinoblastoma progression may come from studies of retinoblastoma tumor DNA obtained via paracentesis of aqueous humor. Recently, it was shown that DNA in the aqueous humor resembles that of enucleated tumors and can change over time (Berry et al., 2017c). In one case, a recurrent chromosomal change (6p +) was not detected in the earliest sample but appeared in subsequent aqueous humor taps, suggesting that this change was selected during tumor evolution, albeit under chemotherapy selection. Further analyses of copy number alterations in multiple aqueous humor samples beginning at the time of diagnosis, may reveal the order(s) in which they occur during conservative treatment (see 8.4).

4.3. RB1 mutations initiate tumorigenesis in the retinoblastoma cell-of-origin – the cone photoreceptor precursor

The malignant behavior of retinoblastoma depends on the effects of RB1 loss and other genomic alterations superimposed on the cell signaling circuitry of the retinoblastoma cell-of-origin. Here, we recount efforts to identify the cell-of-origin, the evidence favoring or disfavoring a cone photoreceptor origin, and the potential contributions of the cell-of-origin circuitry for retinoblastoma initiation and clinical behavior.

4.3.1. First hints pointing towards a photoreceptor cell-of-origin

The identity of the retinoblastoma cell-of-origin has long been debated, in part because early inferences were based on analyses of tumor cells that were far removed from tumor initiation events. Briefly, the tumors were initially referred to as “glioma of the retina” based on use of imperfect stains that suggested a glial composition. In the 1890s, Flexner and Wintersteiner independently recognized that a subset of the tumors had symmetrically arranged structures – later termed “Flexner-Wintersteiner rosettes” – that appeared to be composed of photoreceptor-like cells and were suggestive of a rod or cone origin (Flexner, 1891; Wintersteiner, 1897). However, most tumors lacked Flexner-Wintersteiner rosettes while others had “Homer Wright rosettes” that are also seen in diverse non-retinal tumors (Albert, 1987), which argued against a photoreceptor origin. Eventually, the term “retinoblastoma”
was adopted based on the perceived similarity of the tumor cells to embryonic retinal progenitor cells (Albert, 1987; Verhoef and Jackson, 1926), yet this hardly settled the cell-of-origin question.

Decades later, a growing body of evidence emerged suggesting a predominant cone photoreceptor phenotype in retinoblastoma. Specifically, researchers detected an intact cone but not rod phototransduction cascade as well as high-level expression of RNAs and proteins mediating cone phototransduction in both differentiated and undifferentiated tumors (Bogenmann et al., 1988; Hurwitz et al., 1990; Rodrigues et al., 1992). However, the cone phenotype remained arguable, with evidence that retinoblastoma cells might simultaneously express non-cone markers and recognition that some retinoblastoma cell phenotypes could be adopted after transformation of an unrelated retinal cell type. Further evidence of a cone origin came from the finding that retinoblastoma foci are distributed over the retinal surface in a pattern similar to that of L/M cones (Curcio et al., 1990; Munier et al., 1994), although the cone-like distribution could also have other explanations (King et al., 2015).

4.3.2. Efforts to identify the retinoblastoma origin using genetically engineered mice

The lack of progress in identifying the cell-of-origin based on characteristics of retinoblastoma tumors prompted the adoption of mouse modeling approaches that held potential to reveal inaugural steps in retinoblastoma initiation. However, this approach also ran into difficulties, as no retinal tumors formed in Rb1± mice (Clarke et al., 1992; Jacks et al., 1992; Lee et al., 1992). Furthermore, mice with retinal-targeted mutation of Rb1 and constitutive germline mutation of the Rb1-related p107 or p130 formed retinal tumors with a predominant amacrine or horizontal cell phenotype, rather than the cone photoreceptor phenotypes that seemed to predominate in human retinoblastoma (Cobrinik, 2013; Macpherson, 2008). Perhaps, the different human vs. mouse phenotypes could have been reconciled if tumors in both species derived from retinal progenitor cells yet took on different retinal cell-type-specific features. However, the data showed that retinal progenitor cells in Rb1−/− as well as in Rb1−/−/p107−/− retinai were largely unaffected, whereas diverse post-mitotic retinal neurons re-entered the cell cycle, proliferated, and underwent apoptosis to varying extents (Chen et al., 2004). Concordantly, mice with retin-specific inactivation of Rb1 in a p130−/−/p107−/− background appeared to initiate tumorigenesis in horizontal cell interneurons only after terminal differentiation and synaptogenesis had begun (Ajikoka and Dyer, 2008). Thus, models with combined Rb1 family mutations revealed that retinal tumors can develop from post-mitotic interneurons that aberrantly re-enter the cell cycle, rather than from retinal progenitor cells that fail to exit the cell cycle. Moreover, the resulting mouse tumors retained features of their interneuron cell-of-origin, at odds with the notion that retinal tumors dramatically alter their phenotype during tumorigenesis.

4.3.3. Circumstantial and direct evidence for a cone precursor cell-of-origin

The disparate retinal phenotypes that had been observed in retinoblastoma tumors prompted a re-examination of the retinal cell-type-specific protein expression (Xu et al., 2009). The studies focused on retinal cell-type-specific transcription factors and distinguished neoplastic retinoblastoma cells, which are deficient in the RB1-encoded pRB protein, from pRB+ non-neoplastic cell infiltrates. Analyses of 40 retinoblastomas revealed that the vast majority of pRB− tumors expressed the cone-specific cell lineage factors TRβ2 and RXRγ, the cone + rod + bipolar cell-specific CRX transcription factor, and the cone-specific phototransduction proteins cone arrestin and L/M-opsin. Rare pRB− cells co-expressed S-opsin and L/M-opsin, consistent with the occasional S- and L/M-opsin co-expression in developing L/M cones (Cornish et al., 2004). All of these proteins were expressed at levels comparable to the cone precursors in developing retina. In contrast, only rare and non-neoplastic pRB+ cells in rare tumors expressed markers of rods, retinal ganglion cells, amacrine or horizontal cells, or retinal progenitor cells at levels comparable to the developing retina (including NRL, rhodopsin, Brn3b, syntaxin, Prox1, Chx10, and Pax6). Moreover, combined immunostaining and in situ hybridization revealed that rare cells that lacked pRB and cone marker expression retained two RB1 alleles, implying that only cone-marker + cells had biallelic RB1 loss (Xu et al., 2009). These findings indicated that pRB-deficient and RB1-mutant retinoblastoma cells express numerous proteins that are characteristic of L/M-cones but not markers specific to other retinal cell types.

In keeping with the predominant cone protein expression profile of the pRB− cells, the Muller cell and astrocyte markers Nestin and GFAP were consistently detected only in non-neoplastic pRB + cells (Xu et al., 2009). The Nestin+ and GFAP + cells also expressed Pax2, indicative of derivation from non-neoplastic retinal astrocytes, and expressed the astrocyte and neural stem cell determinant SOX2 (Xu et al., 2010), consistent with the detection of other neural stem cell markers in a subset of cells in retinoblastoma tumors (Seigel et al., 2007). Importantly, SOX2 was only detected in cells expressing pRB protein and/or retaining RB1 alleles, establishing the lack of this stem cell marker in neoplastic retinoblastoma cells (Xu et al., 2010).

The neoplastic retinoblastoma cells’ cone protein expression profile provided one of several lines of circumstantial support for a cone precursor cell-of-origin. For example, a cone origin was also consistent with the high-level pRB expression observed in maturing human cone precursors (Lee et al., 2006) but not in maturing mouse cone precursors (Spencer et al., 2005; Xu et al., 2009). Moreover, maturing human but not mouse cone precursors prominently expressed the proto-oncoproteins MDM2 and MYCN (Xu et al., 2009). As most tumors require the inactivation of the p53 pathway as well as the RB pathway (Sherr and McCormick, 2002), the cone precursors’ high-level MDM2 expression provided a way to suppress p53 function without mutation of the TP53 gene or MDM2 amplification. However, the cone precursor’s expression of these oncogenes provided only circumstantial evidence of a cone origin, as their expression could have been acquired during the development of retinoblastoma from an alternative cell type.

More recent studies provided direct evidence for the cone precursor cell-of-origin. Specifically, pRB knockdown elicited cell cycle entry and proliferation of human cone precursors but not other retinal cell types (Xu et al., 2014). The cone precursor response to pRB knockdown was evident in dissociated retinal cultures, in prospectively isolated cone precursors, and in non-dissociated retinal tissue. Moreover, cell cycle entry depended on the cone cell lineage factors TRβ2 and RXRγ and on the cone-precursor-expressed MYCN, whereas the survival of the aberrantly proliferating cells depended on the endogenous MDM2. Notably, co-depletion of 130 along with pRB enhanced proliferation, consistent with evidence that the RBL2/p130 gene on 16q is often lost in retinoblastoma. Finally, pRB-depleted and pRB/p130-depleted cone precursors formed retinoblastoma-like tumors in orthotopic xenografts, documenting the cone precursors’ capacity to serve as a retinoblastoma cell-of-origin.

4.3.4. Reconciling the cone precursor cell-of-origin with apparently contradictory observations

Despite the direct evidence of the cone precursors ability to form retinoblastoma-like tumors (Xu et al., 2014), the cone origin theory - if true – must be reconciled with all relevant biologic and clinical observations. In this regard, two observations that initially seemed to contradict the cone cell-of-origin theory are addressed here.

First, the earliest lesion detected in retinoblastoma patients via OCT appears to be centered in the inner retina, either in the inner nuclear layer or in the outer plexiform layer, rather than in the outer aspect of the outer nuclear layer where cones reside (Berry et al., 2016a; Rootman et al., 2013). This location suggested that the tumors either derive from a non-cone cell in the inner nuclear layer (which might subsequently adopt a cone phenotype) or derive from cones that are
displaced towards the inner nuclear layer, perhaps through a rare but normal phenomenon (Semo et al., 2007) or in response to pRB loss. Recently, the movement of pRB-depleted cone precursors from the outer edge of the outer nuclear layer to the outer plexiform layer were detected via live imaging of intact cultured retinas, in support of the cone precursor cell-of-origin concept (Singh et al., 2018a).

A second concern was that retinoblastoma tumor cells express RNAs that encode proteins that are characteristic of diverse retinal cell types (McEvoy et al., 2011). The expression of non-cone-related RNAs appeared to conflict with evidence that retinoblastoma cells mainly express cone-related proteins (Xu et al., 2009) and suggested that the retinoblastoma cell-of-origin might have adopted a hybrid gene expression program, with expression of cone as well as non-cone genes, after the transformation of non-cone cell type. However, the data is also consistent with the expression of a hybrid program after transformation of cone precursors in combination with preferential translation of cone-related RNAs. More studies are needed to resolve the discrepancy in the expression of non-cone RNAs with the predominant cone-related protein profile in retinoblastoma cells.

Finally, if cone precursors represent the cell-of-origin, pRB depletation may be expected to induce not only cone precursor proliferation but also the different tumorigenesis stages that have been inferred from clinical observations. With regard to this hypothesis, non-proliferative but also the different tumorigenesis stagesthat have been inferred from this hypothesis, non-proliferative but also the different tumorigenesis stages that have been inferred from clinical observations.

5. Retinoblastoma genetics and genetic counseling

In this part are described the different variants of heritable (50%) and non-heritable (50%) (MacCarthy et al., 2009) presentations of retinoblastoma, as well as their implications for the patient's family members, based on family history, laterality at presentation, and current genetic tests.

5.1. Genetic presentations of retinoblastoma

Retinoblastoma can be classified into four distinct genetic presentations according to the timing of the first Rb1 mutation:

1. Familial, or inherited heritable form, in which the retinoblastoma
A patient has inherited a predisposition to retinoblastoma from an affected parent;
2. **Isolated (or sporadic) heritable form**, in which retinoblastoma is typically bilateral with no family history resulting from a de novo pre-zygotic germline mutation;
3. **Mosaic** form, in which retinoblastoma is usually unilateral, caused by a post-zygotic mutation, and heritable only if the germline is also involved;
4. **Non-heritable** forms, in which retinoblastoma is always unilateral, unifocal and sporadic, resulting from a somatic mutation.

The heritable form of retinoblastoma – either familial or isolated – is caused by germline genetic alterations that disable the RB1 gene functions, also referred to as oncogenic variants or alleles. Individuals with familial or isolated heritable retinoblastoma are heterozygous for an oncogenic RB1 allele (the “first mutation”) throughout their body (Fig. 8A). Each individual retinoblastoma focus that develops in such individuals is initiated by an additional genetic alteration targeting the other RB1 allele (the “second mutation”) during retinal development. Thus, expressivity varies according to the number of independent second mutations that give rise to tumor foci. In addition, expressivity can also vary according to the nature of the first mutation, which determines the complete or incomplete penetrance of the inherited or de novo predisposition, as described below.

5.1.1. **Inherited heritable retinoblastoma (familial retinoblastoma)**

Familial occurrence of retinoblastoma is a form of heritable retinoblastoma in which predisposition to this tumor is transmitted as an autosomal dominant trait. In the familial setting, retinoblastoma screening may be initiated pre- or perinatally and therefore usually associated with a presymptomatic diagnosis (see 3.5). In high-income countries, the fraction of familial cases among heritable retinoblastoma is estimated around 25%, but is rapidly rising due to improved life prognosis.

5.1.1.1. **Familial retinoblastoma with complete retinoblastoma**. In most pedigrees with familial retinoblastoma, all family members who are heterozygous for an oncogenic RB1 allele develop at least one retinoblastoma focus (complete penetrance, Fig 8Ba). Typically, patients develop several tumors and, consequently, bilateral retinoblastoma is prominent in most families. As a rule, a high probability of tumor development is caused by genetic alterations that result in complete or almost complete loss of pRB protein function, which are most often nonsense alterations, frameshift-indels, and alterations at canonical splice signals that result in frameshift exon skipping. In these pedigrees, the mean number of tumors per patient is around 6 (Abramson and Gombos, 1996; Munier et al., 1994).

5.1.1.2. **Familial retinoblastoma with incomplete penetrance**. In some pedigrees, heritable retinoblastoma shows incomplete penetrance with some family members free of any tumor, despite being heterozygous carriers of an oncogenic RB1 allele (Fig 8Bb). Incomplete penetrance is a consequence of lower probability of tumor development and those patients typically develop fewer retinoblastoma foci compared to those with complete penetrance, associated with genetic alterations that retain some of the pRB protein functions or affect regulation of transcription of this gene (reviewed in Lohmann and Gallie, 2004). Consequently, patients in families with incomplete penetrance also tend to show only unilateral retinoblastoma. Most of the difference in penetrance of heritable retinoblastoma can be explained by differences in the functional consequences of oncogenic RB1 alterations. The most prominent examples of variant RB1 alleles associated with a low penetrance phenotype are certain missense variants (e.g. LRG_517p1:p.R661W), in-frame indels, alterations at remote splice signals, specific variants in exon 1, and alterations of transcription-factor binding sites in the RB1 promoter.

Fig. 8. A. Schematic timing of the first mutation in retinoblastoma. a. Inherited heritable. b. Isolated heritable. c. Mosaic. d. Non-heritable. The timing of the first mutation determines both the presence of a constitutional tumor predisposition and the risk of transmission to offspring. In familial retinoblastoma (a) an oncogenic variant allele (rb) is inherited from a parent. The individual is heterozygous (RB rb) and half of the individual's gametes will carry the variant allele. Inactivation of the other allele in the retinoblast by a second mutation (red arrow) will trigger the development of retinoblastoma. In isolated heritable retinoblastoma (b) the first mutation has occurred de novo prior to conception (pre-zygotic mutation). As the individual is heterozygous (RB rb) the situation regarding tumor predisposition and transmission to offspring is as in familial disease. A first post-zygotic mutation may result in somatic mosaicism (c). The extension of the mutant sector and the risk of transmission to offspring depend on the timing and targeted cell type of this mutation. Risk of retinoblastoma oncogenesis is proportional to the extent of mutated retina. Likewise, the proportion of gametes carrying the mutant allele is determined by the contribution of the mutant sector to the individual's germline. A somatic first mutation in a retinoblast (d) is expected to predispose to one single tumor focus with no possible transmission to offspring (isolated non-heritable retinoblastoma). B. Pedigrees/modes of presentation of retinoblastoma. a. Multigenerational, complete penetrance (bilateral retinoblastoma) b. Multigenerational, incomplete penetrance (uni/bilateral and unaffected) c. Two-generational, founder unilateral d. Isolated bilateral e. Isolated unilateral.
Families with lower probability of tumor development provide a good opportunity to identify factors in trans to the RB1 locus that may affect intra-familial variation of phenotypic expression. The minor allele of rs2279744 (NM_001145337:2c.-291T > G), a single-nucleotide polymorphism in the MDM2 promoter known to be linked to enhanced MDM2 expression, has been reported to act as a modifier of phenotypic expression in heritable retinoblastoma (Castara et al., 2010). Parent-of-origin effects have been identified as another source of intra-familial variation of phenotypic expression (Klutz et al., 2002), found later to be related to human RB1 gene imprinting (Eloy et al., 2016; Kanber et al., 2009).

Identification of the mechanisms underlying phenotypic expression, namely low-penetrance RB1 alleles, trans-acting modifiers, and epigenetic factors, is not an end in itself. It can be anticipated that some of the mechanisms mediating milder phenotypic expression will be used to reduce the number of tumor foci, in family members at risk.

5.1.2. Isolated heritable retinoblastoma

In this presentation, retinoblastoma occurs in a child of unaffected parents. Isolated heritable retinoblastoma results from a de novo pre-zygotic mutation in the gametes of one of the healthy parents. According to the nature of the first mutation, expressivity – defined as the number of primary tumor foci – may be attenuated or not.

5.1.2.1. Isolated bilateral retinoblastoma (full expressivity). Most patients with newly diagnosed bilateral retinoblastoma have no relatives with retinoblastoma and are referred to as having isolated sporadic bilateral disease (Fig. 8 Bd). Most patients (95%) with isolated bilateral disease are heterozygous for an oncogenic RB1 allele (Lohmann and Gallie, 2004) and, consequently, also have heritable disease, but occasionally (5%) may be mosaic (see 5.1.3). Typically, oncogenic alleles identified are the result of de novo mutations (Fig. 8 Ab). They are usually indistinguishable from RB1 alleles underlying familial retinoblastoma with complete penetrance, and result in complete or almost complete loss of pRB protein functions (see 5.1.1.1).

Due to the non-familial occurrence, patients with isolated bilateral retinoblastoma will not have been formally screened for retinoblastoma during infancy, and the age at diagnosis reflects the mean time to elicit symptoms, which is usually around 9 months (Goddard et al., 1999). In this population, the mean number of tumors per patient is 6, which is different from that observed in familial retinoblastoma patients with complete penetrance (personal observation).

5.1.2.2. Isolated retinoblastoma with genomic 13q14 deletion (reduced expressivity). Some patients with germline deletions include RB1 as well as neighboring genes on chromosome 13q. Those have a lower probability of tumor development and thus show an excess of unilateral presentations (40%) (Munier et al., 1989) compared to patients carrying other types of loss of function alleles. The mean age at diagnosis in this population is around 17 months (Baud et al., 1999; Munier et al., 1989), which is intermediate between isolated heritable and non-heritable RB1-related retinoblastoma. It appears that the risk to develop retinoblastoma in carriers of deletions larger than 1 Mb containing the MDM4 gene is reduced compared to non-chromosomal mutation carriers (Mitter et al., 2011) with a mean number of tumors per patient of 2.5 (FLM personal observation) versus 6.3 (Munier et al., 1994) respectively. The observed 61% reduction of tumor number in 13q-patients actually closely corresponds to the 63% predicted from peripheral leucocytes is low, this may result in false negative findings in genetic testing. The absence of detectable mosaicism, however, does not rule out somatic mosaicism, since the cells carrying the mutant RB1 allele might not be detected in the sampled cells (blood, buccal etc), but might be present in other tissues such as the retina or germ cells. Thus, there is no test that excludes risk of retinoblastoma in the unaffected eye or, if the mutant sector includes germine cells, in offspring of the patient. However, risk estimates are lower in unilateral retinoblastoma patients who test negative for constitutional RB1 mutation compared to the pretest state of knowledge.

5.1.3. Mosaic retinoblastoma

In familial retinoblastoma with complete penetrance it is not unusual to find that the first affected family member (i.e. the founder on the phenotype level), presents with unilateral retinoblastoma while affected family members in subsequent generations show bilateral disease (Fig. 8 Bc). Genetic analysis has shown that some of these founders are somatic mosaics for an oncogenic RB1 allele (Sippel et al., 1998). In these individuals, the first mutation that led to the oncogenic RB1 allele was not received via the germline but occurred somatically after conception, giving rise to a mutant sector of variable extent according to the timing of the event. A mutational event that occurs during early stages of intrauterine development may result in a mutant sector in the individual. The number of tumor foci in those patients is expected to be lower because the probability of initiation of retinoblastoma development in an individual with somatic mosaicism is reduced inasmuch as the mutant sector contributes only partially to the developing retina.

Mosaicism for an oncogenic RB1 allele is estimated to account for 5% of patients with isolated bilateral disease (Fig. 8 Ac) and 20% of the cases with isolated unilateral retinoblastoma (Lohmann D personal communication). If the proportion of the mutant allele present in DNA from peripheral leucocytes is low, this may result in false negative findings in genetic testing. The absence of detectable mosaicism, however, does not rule out somatic mosaicism, since the cells carrying the mutant RB1 allele might not be detected in the sampled cells (blood, buccal etc), but might be present in other tissues such as the retina or germ cells. Thus, there is no test that excludes risk of retinoblastoma in the unaffected eye or, if the mutant sector includes germine cells, in offspring of the patient. However, risk estimates are lower in unilateral retinoblastoma patients who test negative for constitutional RB1 mutation compared to the pretest state of knowledge.

5.1.4. Non-heritable retinoblastoma

5.1.4.1. Isolated unilateral retinoblastoma with somatic biallelic RB1 mutation. In about half of newly diagnosed retinoblastoma patients only one eye is affected and family history is negative (isolated unilateral disease, Fig. 8 Be). In nearly 90% of these patients, genetic analysis of DNA from retinoblastoma shows genetic or epigenetic alterations of both RB1 alleles that, upon targeted genetic testing, are not detected in DNA from blood (Schuler et al., 2005). This shows that both alterations are the result of mutation events that occurred in somatic cells but does not rule out somatic mosaicism instigated by the first mutation. In the remaining 10% of isolated unilateral retinoblastoma a de novo germline mutation can be identified.

In non-heritable retinoblastoma, there is only one tumor in the affected eye (i.e., there is a unilateral unifocal phenotype). The mean age at diagnosis in this population is 24 months (Goddard et al., 1999), although an earlier age at presentation is not necessarily suggestive of a germline or mosaic RB1 mutation carrier (Schuler et al., 2005).

5.1.4.2. Retinoblastoma without alterations of the RB1 gene. In less than 3% of patients with isolated unilateral retinoblastoma, analysis of DNA from the tumor shows no alteration at the RB1 locus. However, nearly half of these tumors (1.4%) show high amplification levels of a genomic region on the short arm of chromosome 2 that includes the MYCN gene (Rushlow et al., 2013). Although some retinoblastomas with RB1 gene inactivation also show increased MYCN copy numbers, tumors without RB1 mutations have gained a far higher number of MYCN copies. The distinct genetic makeup of MYCN-amplified tumors is reflected by a clinical phenotype marked by very early age at diagnosis of less than 6 months and histopathologic features reminiscent of neuroblastoma. Thus, it might be appropriate to regard this genetic variant as a distinct entity, possibly linked to a distinct genesis. However, a recent report suggests that tumors from MYCN transduced retinal cells not only appear earlier than those with RB-depletion, coinciding with earlier age at diagnosis in MYCN-induced retinoblastoma patients, but also derive from a cone precursor cell of origin (Singh et al., 2019).
5.2. Genetic counseling and testing

Initial counseling and the testing approach for each individual patient depends on the clinical presentation, including familial versus isolated background and bilateral versus unilateral appearance. The three presentations do not necessarily reflect the genetic types of retinoblastoma described in 5.1, as sporadic bilateral or unilateral retinoblastoma can result from several genetic mechanisms.

5.2.1. Familial retinoblastoma

For pre-test genetic counseling and recurrence risk prediction in typical situations of familial bilateral retinoblastoma it is reasonable to assume a probability of transmission of 50% (autosomal dominant inheritance) and a probability of tumor development of 100% (complete penetrance in heterozygous individuals). Consequently, the estimated pretest recurrence risk of heritable retinoblastoma in offspring of a family member with bilateral retinoblastoma is 50% at birth. Although indirect genetic testing by linkage analysis of marker alleles linked to the \textit{RB1} gene may be available, most laboratories prefer direct genetic testing by determining the presence or absence of the oncogenic heritable predisposition in a given family.

A wide spectrum of genetic alterations can result in oncogenic \textit{RB1} alleles (allelic heterogeneity) (Valverde et al., 2005). Various methods are required to cover the relevant genomic regions and the different types of potential oncogenic alterations (small mutations, copy number changes, rearrangements).

In almost all pedigrees with apparent complete penetrance and bilateral retinoblastoma, current routine methods will identify a pathogenic allele (> 95%), provided that the mutation screening analysis is performed on a sample from an affected family member who has inherited the retinoblastoma predisposition.

In rare pedigrees with complete penetrance retinoblastoma, no oncogenic \textit{RB1} allele is identified, due to inactivating alterations not covered by current routine technology (e.g. in frame copy number, neutral inversions), but destined to be diagnosed in the near future with diagnostic sensitivity improvement, especially with the implementation of genomic sequencing technology for routine testing.

In pedigrees where incomplete penetrance is obvious, diagnostic sensitivity of genetic testing is somewhat lower compared to families with complete penetrance. This is because there are fewer constraints regarding the spectrum of alterations that result in only partial inactivation of pRB protein functions compared to alterations that result in full inactivation. As current routine methods used for mutation screening do not cover all regions that may be affected by oncogenic alterations with incomplete penetrance (e.g. deep intronic variants, regulatory variants outside of known transcription factor binding sites), some oncogenic alterations are missed.

Another point that contributes to reduced diagnostic sensitivity of genetic testing in familial retinoblastoma with incomplete penetrance is the uncertainty regarding the pathogenetic role of variants of unknown significance, even if cosegregating with the heritable predisposition.

Although there is no valid report of familial retinoblastoma not genetically linked to the \textit{RB1} locus, locus heterogeneity cannot be formally ruled out. What has been reported, however, are observations of fortuitous co-occurrence of retinoblastoma in relatives (Dryja et al., 1993; Munier et al., 1993).

5.2.2. Isolated bilateral retinoblastoma

The empirical recurrence risk for siblings of a sporadic bilaterally-affected patient is 2% (Draper et al., 1992) and this figure reflects both the low frequency of germinal mosaicism in one of the parents and the lower than 50% probability of transmission of a mutant allele in these situations. It is rare to find an unaffected parent heterozygous for a variant that confers high probability to tumor formation. Consequently, recurrence risk in more remote family members (e.g. cousins) is low. Pretest recurrence risk in offspring of a patient with isolated bilateral retinoblastoma is expected to be 50% at the time of birth, assuming a 50% probability of transmission of 100% probability of tumor development. However, as some patients with isolated bilateral retinoblastoma have somatic mutational mosaicism, the overall probability of transmission is less than 50%.

In principle, the strategies of genetic testing are analogous to those in familial retinoblastoma. With current routine methods, an oncogenic variant is identified in DNA from blood of almost 95% of patients with isolated bilateral retinoblastoma, which is lower compared to familial retinoblastoma, due to somatic mosaics with low proportions of mutation-carrying leukocytes. Although this limitation can be overcome by deep sequencing technology, at least for some types of alterations such as single nucleotide variants, robust strategies for technical validation of low-level findings remains to be established. In most patients, the signal ratio of wild type and variant alleles in DNA from blood suggests a heterozygous genotype. However, checking a second source of constitutional DNA (e.g. DNA from buccal swabs or finger nails) is a worthwhile adjunct analysis as it may detect possible somatic mosaicism. If informed consent is granted, parents should be tested for the presence of the variant allele. Most loss of function-type alleles appear to have occurred \textit{de novo}, and are preferentially found on the allele of paternal origin (Hagstrom and Dryja, 1999; Zhu et al., 1989).

In some patients, the signal ratio of wild type and variant alleles in DNA from blood is imbalanced. Such a finding may be due to somatic mosaicism, but further genetic testing is required to validate this interpretation. However, if somatic mosaicism is confirmed this excludes an increased risk in siblings and relatives of the patient's parents.

Therefore, the additional efforts required to confirm somatic mosaicism in a patient with isolated bilateral retinoblastoma are often justified.

Another option to overcome the limitations of diagnostic sensitivity due to somatic mosaicism is to include analysis of tumoral DNA. Analytically, the most reliable source of tumor DNA is a fresh frozen sample obtained at the time of enucleation. If the tumor mutations are not identified, constitutionally heterozygous intragenic polymorphic \textit{RB1} markers can be used looking for a loss of heterozygosity (LOH) in the tumor, as seen in 63–70% of the cases (Cavenee et al., 1983; Zhu et al., 1992). Whenever LOH is present, the recurrence risk is linked to the transmission of the allele retained in the tumor. If tumor DNA markers are not informative, or in the absence of enucleation, the known paternal preferential origin of \textit{de novo} germline \textit{RB1} mutants, seen in 90% of the cases, can be used to modulate the recurrence risk, which is maximized if the paternal or grand-paternal allele is passed to a sib of an offspring respectively (Tran et al., 2012).

If the oncogenic \textit{RB1} variant responsible for the heritable predisposition is known in the index patient, genetic counseling in relatives is as straightforward as in familial retinoblastoma (see above), with two notable exceptions: i) If testing of DNA from blood of parents did not show the variant allele identified in the child, this will reduce but not exclude increased risk in siblings. The remaining risk is due to germline mosaicism, i.e. the presence of cells with the mutant allele in germline cells of a parent. Unfortunately, empirical data required to quantify the risk due to germline mosaicism are still missing. ii) The second caveat relates to the origin of chromosomal aberrations such as genomic deletions at 13q14 (see 5.1.2.2). If a chromosomal alteration identified in a child is not present in either parent, this does not exclude the presence of aberrations causing unbalanced ones. Chromosome analysis in parents can detect such aberrations in some families.

5.2.3. Isolated unilateral retinoblastoma

Pretest genetic counseling has to take into account that various genetic mechanisms may result in isolated unilateral retinoblastoma. The empirical recurrence risk figures provided by Draper, which integrate all alternative causes, are still valid for risk estimation in siblings and offspring of patients with isolated unilateral retinoblastoma (Draper et al., 1992). The low empirical recurrence risk for siblings (1% at time of birth) reflects that heterozygosity with incomplete
penetrance or germline mosaicism for an oncogenic allele in a parent is rare. Moreover, although the probability of transmission of an oncogenic allele from a rare heterozygous parent is 50%, recurrence risk is further reduced because of the lower probability of tumor development caused by alleles associated with incomplete penetrance. Risk for the patient’s first child is higher (6% at time of birth) and, in most cases, is due to somatic mosaicism in the unilaterally affected parent. Because of somatic mosaicism, probability of transmission is reduced (overall < 50%). However, the oncogenic alleles transmitted by such parents are typically associated with high probability of tumor development. This explains why tumors in children of patients with isolated unilateral retinoblastoma typically show bilateral disease.

The results of genetic testing can help to identify which of the alternative genetic mechanisms has resulted in isolated unilateral disease in a given patient. In fewer than 15% of patients, constitutional heterozygosity for an oncogenic RB1 allele is detected. Genetic counseling of these families must take into account estimates of disease penetrance as indicated by the functional type of mutation. The most frequent genetic mechanism (in over 80% of isolated unilateral patients) is somatic inactivation of both alleles of the RB1 gene. This diagnosis excludes an increased recurrence risk in siblings. If testing shows no evidence of somatic mosaicism then pretest risk is reduced (possibly less than 1%), with the option to exclude an increased risk after predictive genetic testing for heritable oncogenic alterations that had been identified in the parent’s tumor. Of note, although some somatic alterations that cause RB1 inactivation in retinoblastoma are

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**Fig. 9. Retinoblastoma growth patterns.**

A. Endophytic retinoblastoma with class 1 (dust) and 2 (spheres) vitreous seeding. B. Exophytic retinoblastoma with peritumoral retinal detachment. C. Exophytic growth with total retinal detachment causing complete spontaneous vitrectomy (kissing retinal bubbles). D. Mixed, predominantly exophytic growth with extended class 2 subretinal seeding and class 3 (cloud) vitreous seeding. E. Cavitary retinoblastoma with exophytic growth and coexisting endophytic retinoma before (E) and after (F) intra-arterial chemotherapy. G. Diffuse infiltrating retinoblastoma in a 14-year-old boy. White arrowheads delineating the posterior front line of the tumor as documented by OCT before (H, H′) and 3 weeks after (I, I′) the first course of systemic chemotherapy. The white asterisk in H shows the posterior limit of the tumor front line. NB: Lines on fundus photograph correspond to the adjacent ultrasonography image or optical coherence tomography scan.
not heritable (e.g. chromothripsis (McEvoy et al., 2014)) and epigenetic silencing (Gregor et al., 1994), it is still possible that the propensity to acquire such alterations is influenced by heritable genetic variation.

In rare cases, genetic analysis of the patient’s retinoblastoma shows no alteration at the RB1 locus but high-level amplification of the MYCN gene. Current data and models strongly suggest that these tumors are distinct entities without recurrence risk for the unaffected eye of the patient, nor for his or her relatives. However, to date, the genetic diagnosis of MYCN-retinoblastoma has not been validated medically. Moreover, prior to utilizing the results for predictive testing, it would be reassuring to have results from longitudinal observations of patients and their families. Recognition of this specific entity is likely to be possible even in conservatively-treated eyes based on liquid biopsy, by paracentesis of aqueous humor as shown for copy number variations (Berry et al., 2018b).

6. Clinical growth and seeding patterns

The retinoblast undergoing malignant transformation initiates a primary intraretinal tumor, which has not only the capacity to metastasize but also, unlike other cancers, to generate secondary tumors by seeding. Clinical features of retinoblastoma retinal growth and seeding patterns are reviewed below.

6.1. Retinoblastoma growth patterns

Primary tumors generated by retinal oncogenesis are characterized clinically by 6 distinct growth patterns, namely endophytic, exophytic, cavitary, diffuse infiltrating, anterior diffuse and spontaneously regressed retinoblastoma. Note that on top of this phenotypic retinal repertoire, retinoblastoma can also result from extra-retinal oncogenesis leading to ectopic growth in the pineal gland and supra/parasellar region (trilateral retinoblastoma) (see 3.6).

6.1.1. Classic growth patterns of primary retinoblastoma: endophytic, exophytic, mixed growth

Incipient retinoblastoma can be traced by OCT to the inner nuclear layer or in the outer plexiform layer (see 4.3.4). By definition, tumor progression towards the inner or outer retina is associated with endophytic or exophytic retinoblastoma respectively (Fig. 9 A, B), but the link with the OCT variants remains unknown.

Endophytic retinoblastoma presents as a whitish mass rapidly interrupting the retinal vasculature and protruding into the vitreous cavity secondary to alterations of the vitreoretinal interface by two distinct mechanisms (Fig. 9 A):

(i) At the tumor apex, by tumoral infiltration of the internal limiting membrane and hyaloid, leading to vitreous seeding (see 6.2.1).
(ii) At the tumor base, by tracional detachment of the hyaloid, creating a new ocular compartment for retrohyaloid seeding (see 6.2.2).

Finally, note that in case of endophytic growth, the retina contiguous to the tumor base can also undergo a localized circumferential tractional detachment with well-demarcated boundaries.

Exophytic retinoblastoma presents as a mass beneath the retina, sparing the retinal surface, which retains unaltered retinal vasculature (Fig. 9 B). This growth variant impacts the retinal pigment epithelium-photoreceptor complex by causing an exudative retinal detachment, under which subretinal seeding can occur. The detachment, initially intermittent, position-dependent, and leaving poorly defined demarcation lines on the retina, can become total and bullous with further tumor growth, inducing complete spontaneous vitrectomy when the retina enters in contact with the lens (Fig. 9 C).

Mixed endophytic and exophytic growth is a feature of more advanced retinoblastoma (Fig. 9 D), often associated histopathologically with choroidal infiltration across the RPE and Bruch’s membrane. This combination may occur more often with exophytic than endophytic retinoblastoma (Nawaiseh et al., 2015; Palazzi et al., 1990; Shields et al., 1993a). Endophytic and exophytic retinoblastoma can also coexist in the same eye.

6.1.2. Rare variants of retinoblastoma

In addition to the common endophytic and exophytic growth patterns, 3 other variants of retinoblastoma can be encountered, accounting for about 5% of the cases at presentation, namely cavitary (2.7–4%) (see 6.1.2.1), diffuse infiltrating (1–2%) (see 6.1.2.2), and diffuse anterior retinoblastoma (< 1%) (see 6.1.2.3). All these subtypes are characterized by an atypical presentation, often causing a delayed diagnosis or even life-threatening invasive investigations or therapeutic procedures. Awareness of characteristic features described below help the early recognition, crucial to provide appropriate management.

6.1.2.1. Cavitary retinoblastoma

Cavitary retinoblastoma is a rare low-grade endophytic growth variant with intratumoral pseudo-cysts visible as grey lucent cavities at the tumor surface on ophthalmoscopy, appearing hypofluorescent on fluorescein angiography, and as empty spaces on ultrasonography (Palamara et al., 2008) (Fig. 9 E, F). Cavitary retinoblastoma typically displays a mean number of 2–3 cavitations of an average diameter of 3 mm (Chaudhry et al., 2018; Rojanaporn et al., 2012). Diagnosis is made at a mean age of about 15 months and predominates in bilaterally affected patients in approximately 2/3 of the cases (Chaudhry et al., 2018; Rojanaporn et al., 2012). These tumors may co-exist with non-cavitary retinoblastoma tumor foci in the same or fellow eye. Secondary intracavitary seeding has been observed in 10% (n = 2/20) of the cases followed in Lausanne (unpublished data) (see 6.2). Noteworthy, cavitary retinoblastoma may develop new cavities during treatment, and secondary cavities may become visible in an initially non-cavitary tumor after an average of two courses of chemotherapy (Chaudhry et al., 2018). On the other hand, cavities can collapse under treatment as seen in 44% of the eyes after a mean follow-up of 18 months (Chaudhry et al., 2018).

In contrast to non-cavitary retinoblastoma where systemic or intra-arterial chemotherapy induces a mean tumor reduction of approximately 30–35% in basal diameter and 50–56% in thickness (Shields et al., 1996, 2011), cavitary tumors typically show shrinking only by half these values as a mean (Rojanaporn et al., 2012), and sometimes not at all (Fig. 9E and F). In addition, relapse of cavitary retinoblastoma appears to be exceptional, occurring in none of 18 eyes (0%) after a mean follow-up of 40 months in a series from London (Chaudhry et al., 2018), and in only one of 26 eyes (4%) in the series from Philadelphia after a mean follow-up of 4 years (Rojanaporn et al., 2012), again very different from the relapse rate of 20–28% characterizing non-cavitary retinoblastoma (Gombos et al., 2002; Shields et al., 2004b, 2011).

In general, despite apparent resistance, cavitary retinoblastoma does not require aggressive or prolonged treatment. However, considering a recurrence risk estimate of about 4%, similar to that of retinoma, life-long follow-up is advised (Abouzeid et al., 2012; Balmer et al., 1991; Singh et al., 2000) (see 6.3).

6.1.2.2. Diffuse infiltrating retinoblastoma

Diffuse infiltrating retinoblastoma is another rare variant found in 1–2% of the cases, characterized by a planar horizontal growth rather than the vertical growth seen in endophytic and exophytic retinoblastoma (Fig. 9 G). Typically, the tumor infiltrates the retina which appears thickened on B-scan, UBM or MRT, without visible mass or calcifications. Histopathologic reports have shown diffuse tumoral invasion of the ganglion cell layers (Fernandez et al., 2017), appearing as a hyper-reflective homogenous thickening on OCT (Stathopoulos et al., 2019) (Fig. 9 H, I). This OCT signature was shared in all four diffuse infiltrating tumors investigated so far in Lausanne (Stathopoulos et al., 2019; FLM unpublished data). Interestingly, a type 0 regression with complete restoration of the retinal microanatomy was observed on
Compartments are characterized by the yellow mass and yellow arrows respectively. Invaded aqueous and vitreous compartments are indicated in light yellow.

The schematic en-face view and longitudinal sections of the fundus extremus please refer to the legends given in Fig. 5. Tumor and tumor spread into the different compartments are characterized by the yellow mass and yellow arrows respectively. Invaded aqueous and vitreous compartments are indicated in light yellow.

OCT after intra-arterial and intravenous chemotherapy (Stathopoulos et al., 2019) (Fig. 9 H', F').

Compared to classic retinoblastoma, the diagnosis of diffuse infiltrating retinoblastoma is usually made in older children (mean age 5.7 years) following atypical presenting symptoms such as vision loss, ocular redness or pain, in the context of misleading signs most frequently masquerading inflammation or pointing to other diagnoses, including neovascular glaucoma and cataract (Traine et al., 2016). In addition to frequent diagnostic delay, slow tumor growth could explain the older age at presentation (Bhatnagar and Vine, 1991; Traine et al., 2016). Most cases (> 90%) are unilateral and sporadic. Diffuse infiltrating retinoblastoma and typical retinoblastoma have, however, been observed in the same patient (FLM personal observation) or within family members carrying the same mutation (Kao, 2000; Scedler et al., 2016), raising the question of a possibly distinct genesis in this atypical disease subtype. As retinal specific micro-environmental features are known to contribute to retinoblastoma tumorigenesis (Xu et al., 2010), a specific tropism of infiltrating retinoblastoma tumor cells for the ganglion cells favoring this atypical growth over the inner plexiform layer can be hypothesized.

Until recently, all cases have been managed with primary enucleation (Bhatnagar and Vine, 1991; Fernandez et al., 2017; Foster and Mukai, 1996; Shields et al., 2008). Last year, for the first time, one successful case of eye salvage with vision preservation was reported after focal treatment and intra-arterial chemotherapy at an updated 38-month follow-up (Stathopoulos et al., 2019). Optical coherence tomography provides instrumental information to monitor the tumor front line following treatment in such cases.

6.1.2.3. Diffuse anterior retinoblastoma. The term diffuse anterior retinoblastoma was first proposed in 1998 to describe an exceedingly rare variant of the disease (Grossniklaus et al., 1998), totaling only 13 published cases up to now (Jijelava and Grossniklaus, 2013; Kelly et al., 2016; Munier et al., 2017a; Shields et al., 2016c). Diagnosis occurs usually later than the other forms (mean age 6.4 years). Half of the cases present with a cellular infiltration of the anterior chamber without retinal or vitreous involvement, while in the other half, a small retinal lesion is found at the ora serrata, typically with concomitant vitreous seeding (Jijelava and Grossniklaus, 2013; Kelly et al., 2016; Munier et al., 2017a; Semb et al., 1961; Shields et al., 2016c).

Diffuse anterior retinoblastoma has been considered a variant of diffuse infiltrating retinoblastoma, originating either from a very small focus close to the ora serrata (Jijelava and Grossniklaus, 2013) or directly from the anterior segment (Shields et al., 2016c). In our opinion, however, this subtype of retinoblastoma constitutes a unique, separate entity, originating from two distinct sources found at the extreme fundus periphery (Fig. 10), explaining the two different clinico-pathologic manifestations observed in this subtype of retinoblastoma, namely without or with minimal peripheral retinal involvement. Hence, in the case of diffuse anterior retinoblastoma without retinal tumor, we think that the tumor arises from ectopic retina found anteriorly to the ora serrata, corresponding either to retinal islands embedded in the pars plana (Eisner, 1973) or to cristae retinæ (Fig. 10). As both structures lack the internal limiting membrane (Daicker, 1972), a tumor originating from this ectopic retina is expected to seed immediately into the posterior chamber without growing a mass (Fig. 10A–C). Conversely, in the case of anterior diffuse retinoblastoma with a minimal peripheral retinal involvement, we believe that the tumor originates from the retinal rim found under the vitreous base (Fig. 10), where L/M cones are known to out-number rods (Ahnelt, 1998; Williams, 1991). Since the retinal rim under the vitreous base is devoid of hyaloid and has a thin fenestrated inner limiting membrane (Le Goff and Bishop, 2008), a tumor arising from this location will initially grow as an endophytic mass which can rapidly invade both posterior chamber and vitreous cavity (Le Goff and Bishop, 2008) (Fig. 10D–F).

6.2. Intra-ocular seeding

Seeding of tumor cells is a characteristic feature of retinoblastoma. It can, however, be observed in other primary intra-ocular tumoral conditions such as melanocytoma of the optic nerve (Shields et al., 2006c), ciliary body medulloepithelioma (Kaliki et al., 2013a), retinal astrocytoma (Cohen et al., 2008), and uveal melanoma (Metz et al., 2016).

Seeding of the primary tumor engenders no less than 12 recognizable phenotypes, comprising all possible combinations of the 3 known classes of seeds, i.e. dust, spheres and clouds, across the 4 intraocular seeding compartments, i.e. the vitreous, aqueous, as well as the sub-retinal and retrohyaloid spaces (Francis et al., 2015a; Munier, 2014).

Fig. 10. Primary aqueous humor invasion mechanisms in anterior diffuse retinoblastoma. A-B. Schematic illustration of anterior diffuse growth. In this case, the tumor presumably arises from the peripheral retina where it can grow both trans-orally directly into the posterior chamber (Petit’s canal) and/or into the vitreous through the fenestrated limiting membrane, as illustrated in (C) by fundus photography and UBM (50 MHz) of a primary oral tumor in an 11-year-old patient. D-F. Anterior diffuse growth with aqueous seeding (no retinal tumor) arising from ectopic ciliary retina (E) or from an oral retinal tooth (F). NB: For the microanatomy of the schematic en-face view and longitudinal sections of the fundus extremus please refer to the legends given in Fig. 5. Tumor and tumor spread into the different compartments are characterized by the yellow mass and yellow arrows respectively. Invaded aqueous and vitreous compartments are indicated in light yellow.

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6.2.1. Vitreous seeding

Vitreous seeding occurs following the disruption of the inner limiting membrane of the retina and hyaloid either by spontaneous or by iatrogenic mechanisms. Interestingly, transmission electron microscopy measurements showed that the thickness of the internal limiting membrane is characterized by huge topographic variations ranging from 20 nm at the foveola to 2500 nm in the macular region and decreasing progressively with greater eccentricity down to 50 nm at the vitreous base (Foos, 1972), where the inner limiting membrane is perforated (Le Goff and Bishop, 2008). The inner limiting membrane opposes therefore less resistance to seeding in the periphery, accounting for the preferential anterior location of class 3 seeds as recently reported (Francis et al., 2016). Schematic representation and peripheral clinical limits of intravitreal seeding in endophytic retinoblastoma growth pattern is shown in Fig. 11 A-C.

Spontaneous vitreous seeding can be present at diagnosis (primary seeding) as a pathognomonic feature of endophytic retinoblastoma, or complicate the disease course at tumor relapse (secondary seeding) independently of the initial growth pattern (Munier, 2014). In the first scenario, by far the most frequent, the passage of tumor cells into the vitreous is part of an active infiltration at tumor apex, discharging individual cells (class 1 seeds). The ability of these cells to survive and grow under the hypoxic vitreous conditions will depend on metabolic reprogramming and/or resistance to anoikis. In other words, only a fraction of these cells will acquire the clonal capability to form spheres (class 2 seeds). According to their adhesion-independent or dependent properties, they will give rise to free floating spheres or hemispheres anchored to the inner face of the hyaloid. In the second scenario, the passage into the vitreous follows a passive mechanism by mechanical or necrotic disruption of the tumor apex, leading to the formation of a cloud (class 3 seeds). The translocation of the tumor content by gravity into the vitreous body is sometimes endowed of gravitational motility, thus masking or unmasking the retina depending on the head position (Fig. 12A-C). Here again the translocated tumor mass is mostly composed of dead cells (Amram et al., 2017) and only a minority will survive to produce spheres (class 2 seeds).

Iatrogenic (secondary) vitreous seeding is a well-known complication of inappropriate energy transfer by photocoagulation or diode-mediated hyperthermia (Gombos et al., 2006), followed by dispersion of tumor cells (class 1 seeds) into the vitreous. It can also occur in the context of intravenous (Parness-Yossifon et al., 2009) or intra-arterial chemotherapy (De Francesco et al., 2014), with the sudden formation of class 3 seeds consecutive to internal limiting membrane disruption following rapid tumor regression. Finally, indentation during ophthalmoscopic examination of endophytic tumors at diagnosis, or during positioning of a plaque, may also be at the origin of iatrogenic tumor dispersion.

6.2.2. Retrohyaloid seeding

Retrohyaloid seeding is underdiagnosed and often confounded with vitreous seeding. The retrohyaloid space is generated secondary to rapid endophytic growth and formation of a tractional hyaloid detachment, or to the inability of the contracted vitreous to re-expand and compensate the collapsed tumor volume after treatment. The extent of the retrohyaloid volume can be evaluated by ultrasonography, occupying up to half of the vitreous volume. The recognition of a hyaloid detachment is crucial in order to avoid grade 4 or 5 melphalan-induced retinopathy linked to inadvertent retrohyaloid injection of intravitreal chemotherapy (see 7.3.5.1.2.2).

The retrohyaloid seeds can float freely (Fig. 12D-F), or attach to the inner limiting membrane of the retina and/or the outer hyaloid face. Class 3 retrohyaloid seeding typically appears as a circular position-

Fig. 11. Schematic representation and peripheral clinical limits of intravitreal, retrohyaloid and subretinal seeding in endophytic and exophytic retinoblastoma growth patterns. A-C. Vitreous seeding from endophytic growing peripheral retinal tumor secondary to the rupture of the internal limiting membrane and hyaloid. C. Fundus picture and UBM (35 MHz) showing vitreous seeding respecting the anterior hyaloid with a tumor-free posterior chamber. D-F. Retrohyaloid class 3 seeding from an endophytic growing tumor, secondary to internal limiting membrane rupture but with an intact hyaloid (D, E). In this context, (F), anterior seeding extension is limited by the vitreous base, posteriorly to the ora serrata (F). G-H. Subretinal seeding from an exophytic growing tumor. (I) Fundus photography showing subretinal seeding reaching the ora serrata by gravity. NB: For the microanatomy of the schematic en-face view and longitudinal sections please refer to the legends given in Fig. 3. Tumor and tumor spread into the different compartments are characterized by the yellow mass and arrows respectively.
dependent cloud, masking the optic nerve and fovea only in dorsal decubitus (Munier, 2014) (Fig. 12G-K). Schematic representation and peripheral clinical limits of retrohyaloid seeding in endophytic retinoblastoma growth pattern is shown in Fig. 11D-F.

6.2.3. Subretinal seeding

Subretinal seeding is the hallmark of exophytic retinoblastoma. Two distinct presentations of class 2 subretinal seeding can be distinguished, i.e. spheres usually anchored to the detached outer retina, or flat pla- cloid lesions with a tendency to confluence in a shallow subretinal space (Fig. 11G-I). The area of initial retinal detachment remains at risk for persistent or recurrent subretinal seeding following first line treatment. It is important to keep in mind the full extent of the detached retinal territory, sometimes difficult to identify due to the on/off phenomenon seen in exudative retinal detachment, in order to distinguish new peripheral tumors from recurrent subretinal seeds.

By gravity, class 2 seeds tend to accumulate inferiorly just posterior to the ora serrata (Fig. 11G-I). In the case of bullous retinal detachment, the seeds attach to the outer retina and can project anterior to the ora serrata. When the retina is folded over the pars plana, UBM is the only way to differentiate these subretinal ante-oral tumors from an invasion of the posterior chamber.

6.2.4. Aqueous seeding

Aqueous seeding refers not only to the presence of retinoblastoma tumor cells in the anterior chamber on biomicroscopy (Fig. 13), but also to its invisible posterior chamber counterpart. Chronologically, invasion of the posterior chamber typically occurs first, leaving the anterior
chamber in a transient cryptic seeding state before converting to an overt involvement. In case of infiltration of the iris root by (choroido)-ciliary or supraciliary mass, however, the passage of retinoblastoma cells to the anterior chamber can bypass the posterior chamber (see 6.2.4.3).

Ultrasonographic biomicroscopy is instrumental to document anterior segment tumor involvement, enabling in particular to determine whether the aqueous seeding is isolated or associated with an anterior uveal infiltration of the ciliary body and/or iris (Fig. 14).

6.2.4.1. Prevalence of anterior chamber seeding. Anterior chamber seeding at diagnosis is rarely seen in high-income countries, being present in approximately 1% of the cases (Haik et al., 1987), except in two rare subtypes of retinoblastoma, namely diffuse infiltrating (Shields et al., 2008) and anterior diffuse retinoblastoma (Jijelava and Grossniklaus, 2013), where it is found in 65% and 100% of the cases, respectively. The presence of anterior chamber seeding automatically grades disease as group E according to the International Intraocular Retinoblastoma Classification (IIRC), with primary enucleation recommended (Linn Murphree, 2005).

Secondary anterior chamber seeding appears at a mean interval of 28 months (range 3–99) after diagnosis (Munier et al., 2018), and was found in 6% of secondary enucleated eyes with advanced retinoblastoma treated with chemoreduction (Shields et al., 2009), and 23.5% (Kiratli et al., 2017) to 50% (Pavlidou et al., 2015) of those treated with intra-arterial chemotherapy.

Until recently (see 7.2.4), secondary anterior chamber seeding has always been managed by enucleation (De Francesco et al., 2014; Parness-Yossifon et al., 2009).

6.2.4.2. Prevalence of anterior uvea invasion. Anterior uvea (iris and ciliary body) invasion, isolated or combined with aqueous seeding is reported in 2–7% of primary enucleated eyes in developed countries (Brennan et al., 2015; Kaliki et al., 2013b; Uusitalo et al., 2001), versus 8–12% of primary enucleated eyes in developing countries (Kaliki et al., 2015b). In a recent series directly comparing primary and secondary enucleations, the authors described a significantly higher incidence of anterior segment involvement (17% vs 0%, p < 0.05) following Fig. 13. Aqueous seeding. (A) Class 1, (B, C) 2 and (D) 3 seeding. E. Anterior segment optical coherence tomography showing presence of hyper-reflective material over the corneal endothelium, iris and angle.

Fig. 14. Invasion of the anterior uvea and aqueous seeding class 3. A, B. Iris and intra-ciliary body invasion documented by slit lamp photography and both longitudinal and transverse ultrasound biomicroscopy (35 MHz) in a 26-month-old boy with previous enucleation of the contralateral eye, before (A) and after (B) successful treatment. C. Fundus of the same patient at 18-month event-free follow-up (Snellen visual acuity 0.5).
secondary enucleation (Fabian et al., 2017c). Others reported similar findings with a higher frequency of ciliary body invasion in secondary enucleated eyes having received neoadjuvant chemotherapy (13%) compared to primary enucleation (7%), and hypothesized that this increase in anterior histopathologic features may reflect a differential targeting of chemotherapy against ciliary body compared to posterior uvea, due its poorer vascular supply (2 long posterior vs 20 short posterior ciliary arteries) (Brennan et al., 2015).

6.2.4.3. Invasion mechanisms of the aqueous humor. Aqueous humor invasion can occur via five different pathways, as documented on clinical, ultrasonographic (UBM) and histopathologic observations (Fig. 15):

(i) **Trans-hyaloid invasion** from tumor cells originating from vitreous seeds anchored to the anterior hyaloid, or from peripheral parietal tumors growing over the pars plana (Fig. 15E–H).

(ii) **Trans-ciliary invasion** via growth of a ciliary/choroido-ciliary tumor directly into the posterior or anterior chamber and/or by infiltration of the iris root (Fig. 15 A, B).

(iii) **Iatrogenic invasion** by tumor cells freely invading the posterior chamber from the posterior segment after mechanical disruption of the anterior hyaloid, e.g. post pars plana vitrectomy, endcapsular lensectomy or posterior capsulorhexis.

(iv) **Epiciliary invasion** by tumor cells originating from an oral or ectopic retinal source, as seen in diffuse anterior retinoblastoma (see 6.1.2.3) (Fig. 10).

(v) **Supra-ciliary invasion** by anterior chamber angle infiltration of a supra-ciliary mass (Fig. 15 C, D).

Recognition of the invasion mechanism underlying aqueous humor invasion is determinant for the adequate treatment strategy, as the seeding source must be sterilized concomitantly to the aqueous seeding, which became treatable only recently with intracameral chemotherapy (see 7.2.6) (Fig. 16). Whereas iris invasion can be controlled by intra-arterial melphalan (Munier et al., 2018), ciliary body involvement needs to be treated by brachytherapy alone (Chhablani et al., 2010) if isolated, or by combining brachytherapy and intracameral chemotherapy if associated with aqueous seeding (Munier et al., 2018).
6.3. Retinoma/retinocytoma and phthisis bulbi

Retinoma, also called retinocytoma, retinoblastoma group 0, and spontaneously arrested or regressed retinoblastoma, is used to designate a benign phenotypic counterpart of retinoblastoma. The pathologic nature of these lesions was described in 1983 as benign photoreceptor proliferations with numerous fleurettes and absence of necrosis or mitotic activity (Margo et al., 1983). More recently, review of histopathologic sections of enucleated eyes showed areas with features of retinoma in 15–20% of them, suggesting evolution from retinoma to retinoblastoma (Dimaras et al., 2008; Eagle, 2009).

6.3.1. Prevalence and relationship with retinoblastoma

The exact frequency of retinoma in the population is not known. Half of the reported cases are diagnosed by routine examination of first degree relatives of retinoblastoma probands, especially in familial cases (Abouzeid et al., 2012; Gallie et al., 1982). In a series of 171 sporadic retinoblastomas, retinoma was diagnosed in only 2 asymptomatic progenitors (FLM personal observation). The reported proportion of retinoma among the population with retinoblastoma varies from 1.8% (17/920) to 3.2% (16/505) (Abouzeid et al., 2012; Singh et al., 2000). Retinoma can also be suspected retrospectively in late-onset diagnosed retinoblastoma in older children or adults (Eagle et al., 1989; Matafsi...
et al., 2012), or in tumors with no or minimal initial response to chemotherapy and absence of growth after discontinuation of all treatments (Chung et al., 2010; Shields et al., 2002).

Like retinoblastoma, retinoma can be present in one or both eyes. The lesion can also co-exist with retinoblastoma in the same or fellow eye. In retinoma germline carriers, the underlying disease-predisposing R81 mutations are indistinguishable from those found in retinoblastoma patients (Abouzeid et al., 2009), indirectly confirming that R81 double hits are necessary but not sufficient to cause retinoblastoma (Dimaras et al., 2008).

6.3.2. Clinical features and risk of malignant transformation

The majority of the lesions correspond to spontaneously arrested and/or regressed proliferations, appearing typically as a translucent retinal mass with or without calcifications and/or retinal pigment epithelial alterations (Gallie et al., 1982). In a report on 86 retinoma foci in 49 eyes, 40% of the retinoma foci showed regression patterns types I and IV, and 60% types II and III, including 5 tumors with pseudo-cyst, among which 2 displayed benign cavity enlargement over time (Abouzeid et al., 2012).

Phenotypic variability also includes spontaneously regressed vitreous seeds (Abouzeid et al., 2012; Hadjistilianou et al., 2006; Lueder et al., 1995; Shah et al., 2011), regressed subretinal seeds, clinical signs of past retinal detachment and histopathologic evidence of full thickness choroidal and prelaminar optic nerve head invasion (Abouzeid et al., 2012) (Fig. 17 A, B). Such findings suggest the existence of a more aggressive retinoma subtype prior regression than typical ones. Finally, phthisis bulbi has been reported in 6.6–8.3% in the two largest series of patients with spontaneously regressed retinoblastoma (Abouzeid et al., 2012; Gallie et al., 1982).

The mechanisms underlying regression/growth arrest of retinoma remain to be determined. Considering the occasional multifocal and/or bilateral disease, however, such a coordinated growth stop of millions of cells across different ocular compartments is more likely to be caused by a still unknown humoral or circulating factor, than explained by a cell-intrinsic mechanism or vascular collapse.

As retinoma can undergo secondary malignant transformation with an estimated lifelong risk of about 5%, regular control of the lesions are needed (Abouzeid et al., 2012; Singh et al., 2000) (Fig. 17C-E).

7. Conservative management of intraocular retinoblastoma

With the emergence of new therapies over the last 20 years, the management of retinoblastoma has also become more complex. The first rule remains the defense of life, which presupposes a permanent arbitration between pursuit of conservative therapy and secondary enucleation. This being said, one of the most important keys to success is continuous adaptation of the therapeutic strategy to the different eye compartments involved. The right decision-making implies a solid knowledge of the strengths and limits of the different modalities that have come to enrich the therapeutic armamentarium of retinoblastoma, but also the awareness of their potential complications. Efforts to implement eye-conservative strategies in less developed countries should consider local socio-economic and cultural factors as well as availability of resources for treatment and patient support. In many of these settings, compliance with follow-up visits and timely acceptance of enucleation when conservative therapies have failed may not be comparable to higher income countries, causing an increased risk for disease progression and death. In addition, even mild chemotherapy regimens may be associated to increased risk of toxic death in these settings (Chantada et al., 2013).

This part deals first with an overview of the pharmacokinetics of the most commonly used chemotherapeutic drugs in retinoblastoma treatment according to the different routes of delivery (see 7.1), illustrating why the development of targeted routes of chemotherapy administration directly into the eye was essential to achieve control of tumor seeding. We then review the different available therapeutic modalities and their contribution in reducing the enucleation rate and eradicating the need for external beam radiotherapy (see 7.2). Furthermore, major disease- and treatment-related intraocular complications encountered during conservative retinoblastoma management are highlighted to bring into perspective one of the biggest remaining challenges of disease management, namely vision preservation (see 7.3). Finally, this section ends with a review on the quality of life (see 7.4).

7.1. Pharmacokinetics

To achieve the intended goal, chemotherapeutic drugs must become available in a sufficient concentration of their active form at the intended tumor site. In parallel, for retinoblastoma patients, it is also
important to take into account the amount of drug in the bloodstream to counterbalance efficacy with systemic toxicity. The vitreous-to-plasma and the retina-to-plasma drug exposure ratios are important concepts to define the most selective route for ocular drug delivery and to avoid exposure of normal tissues to antineoplastic agents.

Despite the use of chemotherapy in the conservative treatment of retinoblastoma for more than 20 years, knowledge about the intraocular bioavailability of the different drugs remains incomplete. Moreover, information, mainly based for ethical reasons on animal studies, is often characterized by discrepancies with the clinical observations secondary not only to differences in the anatomy between animal and human eyes, but probably also between non-tumor bearing and tumor-bearing-eyes, whose integrity of the blood-ocular barrier is compromised.

The following overviews the main pharmacokinetic studies with emphasis on the different possible administration routes of the chemotherapeutic drugs used against retinoblastoma. A schematic representation of previously published data of chemotherapy ocular exposure in different animal models after several local and systemic routes of drug delivery is presented in Fig. 18.

7.1.1. Intravenous drug delivery

To penetrate the different ocular compartments after intravenous delivery, the drugs have first to make it through the first-pass metabolism and, once in the choroidal and retinal vessel, to overcome the blood-ocular barrier. The latter consists posteriorly of the tight junctions of the retinal pigment epithelium and the retinal endothelial cells and, anteriorly, of the inner non-pigmented ciliary epithelium, the posterior iris epithelium, and the endothelium of the iris and ciliary body vessels. High doses of chemotherapy are consequently required intravenously to attain pharmacologically active concentrations in the ocular tissues, carrying simultaneously the risk of severe systemic adverse events (Del Amo et al., 2017; Edelhauser et al., 2010).

Systemic chemotherapy as conservative retinoblastoma treatment (typically with carboplatin, etoposide, and vincristine) was introduced in the 1990s, without preclinical studies on intraocular pharmacokinetics and remains the most frequently used treatment worldwide even though only a small fraction of the administered drugs is thought to make its way through the first-pass metabolism and the blood-ocular barrier.

Specifically, studies in tumor-bearing juvenile rats showed that intravenous carboplatin, a cell-phase nonspecific alkylating agent, achieved an intravitreal exposure of 60% of the systemic area under the curve (AUC) (Laurie et al., 2005), whereas in a non-human primate model, the maximum achieved concentration in the vitreous (Cmax) was only 1% of the plasma Cmax but, surprisingly, the aqeous humor Cmax was 20% of the systemic level (Mendelsohn et al., 1998). Interestingly, the concentration of carboplatin in the vitreous in enucleated eyes from retinoblastoma patients was 13-fold higher than that attained after the same dosage in non-human primates (Abramson et al., 1999a), but similar to the levels obtained by others in rabbits subjected to cryotherapy (Wilson et al., 1996).

Pharmacokinetic studies with intravenous etoposide, a topoisomerase II inhibitor, performed in rats and non-human primate models have shown a very low to undetectable vitreous exposure compared to the systemic exposure (Laurie et al., 2005; Mendelsohn et al., 1998).

Intravenous vincristine, a cell-mitosis inhibitor from the vinca alkaloid family, exhibited good retinal and vitreous exposure in rats, almost equal to plasma exposure, but the assays performed were not able to distinguish between a degraded or protein-bound form of the molecule (Laurie et al., 2005).

The low selectivity of the intravenous route for chemotherapy delivery into the vitreous was also reported for topotecan, a topoisomerase type I inhibitor, in non-tumor-bearing rats and rabbits with a vitreous-to-plasma exposure ratio of less than 0.4 (Carcaboso et al., 2007; Laurie et al., 2005). Nonetheless, intravenous topotecan is still used in the clinics for retinoblastoma treatment (Brennan et al., 2017).

From the above, it appears that other administration routes had to be explored to improve tumor control, especially in advanced disease with seeding into the different ocular compartments.

7.1.2. Intra-arterial drug delivery

The technique of the super-selective catheterization of the ophthalmic artery was developed to allow the injection of high drug concentration directly into the artery that irrigates the ocular tissues, thus avoiding the first-pass elimination and reducing at the same time systemic exposure to the cytotoxic drug. The hypothesis of an increased bioavailability in the ocular tissues with greater selectivity compared to systemic exposure was confirmed four years after the initial publication on 9 cases treated this way (Abramson et al., 2008), in pharmacokinetic animal studies using non-tumor bearing pigs (Schaquievhich et al., 2012a, 2012b).

The most frequently injected drug in the ophthalmic artery is melphan, a nitrogen mustard agent found to be the most efficient drug among the 12 tested against retinoblastoma cells in vitro (Inomata and Kaneko, 1987). Its cytotoxic activity is mediated by alkylating nucleic
acids and proteins, which alters DNA function and ultimately leads to cell death. Rapid spontaneous hydrolysis of the reconstituted drug from its commercial vial implies infusion within 1.5 h in order to avoid the loss of the active moiety.

Another frequently injected drug is topotecan as its active moiety of lactone operating during the S phase of the cell cycle, is commonly used in combination with melphalan to take advantage of synergistic effects (Kaufmann et al., 1996; Schaiquevich et al., 2012b).

### 7.1.2.1. Intraocular distribution of intra-arterial melphalan and topotecan

Pharmacokinetics studies after intra-arterial melphalan in non-tumor bearing pigs, found that attained vitreous Cmax barely reached the concentration that inhibits 50% of the retinoblastoma cell growth in vitro (IC50), whereas the vitreous AUC was 3-fold higher than the plasma AUC (Schaiquevich et al., 2012b) (Fig. 19). The relatively low intravitreal levels were thought to be related to a poor affinity of the melphalan to the L-amino acid transporter 1 (LAT1), the mediator of its transport through the blood-retina barrier (Hosoya et al., 2008). Comparatively, intravitreal concentrations of topotecan after intra-arterial injection using the same animal model were above IC50 for at least 4 h (Schaiquevich et al., 2012a) (Fig. 20) and the AUC in the vitreous was 29-fold higher than that in the bloodstream of the study pigs (Schaiquevich et al., 2012a). Another study also performed in pigs showed 143- and 246-fold higher retina and vitreous levels respectively after intra-arterial topotecan than after intravenous infusion of the same dose (Taich et al., 2016), with comparable systemic exposure. In the same study, topotecan exposure in the optic nerve was 80-fold higher after intra-arterial injection than after intravenous infusion at the same dose of 4 mg.

Despite the melphalan vitreous-to-plasma ratio of almost 10-fold less than the ratio obtained for topotecan in non-tumor pigs studies (Schaiquevich et al., 2012a, 2012b), it was clear from clinical experience that melphalan alone was efficient to treat retinal and subretinal tumors, whereas this was not the case with intra-arterial topotecan as monotherapy. On the other hand, intra-arterial melphalan, alone or in combination with topotecan, was insufficient to manage all cases with vitreous disease.

Of note, melphalan was detected in the RPE-choroid in almost all the eyes of the pigs after intra-arterial chemotherapy but barely quantified in the retina, showing the selective accumulation of the drug in the RPE-choroid tissue.

#### 7.1.2.2. Systemic adverse effects of intra-arterial delivered drugs

Although intra-arterial chemotherapy is a local route for ocular drug delivery, a fraction of the administered drug becomes systemically available. This implies that there is a limitation to the maximum dose that can be given in order to avoid systemic adverse effects, and supplementary caution should be made in cases of tandem therapy. Normalizing the dose based on body weight was found to minimize inter-patient variability in melphalan pharmacokinetics. Thus, patients exceeding the threshold dosage of 0.48 mg/kg showed a 50% chance of presenting severe neutropenia compared to less than 1% in children receiving lower doses (Dunkel et al., 2014; Schaiquevich et al., 2012b).

Finally, a clinical pharmacokinetic study on intra-arterial chemotherapy with combined melphalan (0.48 mg/kg) and topotecan (0.5–1 mg) in patients with retinoblastoma, failed to show severe hematologic toxicity secondary to a potential drug-drug interaction (Taich et al., 2014). Total topotecan systemic exposure after intra-arterial chemotherapy was 95 ng*h/ml per mg of topotecan with about one-third of this exposure corresponding to the active lactone moiety (Taich et al., 2014), which is way below the threshold for severe neutropenia after intravenous delivery (Brennan et al., 2017).

### 7.1.3. Intravitreal drug delivery

Compared to all the other administration routes, drug injection directly into the vitreous bypasses the ocular barriers and the first-pass elimination, and provides therefore the highest intraocular exposure in the vitreous humor, ideal for the management of vitreous disease. After the intravitreal injection, elimination takes place anteriorly by drug flow from the vitreous into the aqueous humor and thereafter Schlemm's canal and posteriorly through the components of the blood-retinal barriers to the choroidal blood flow and the systemic circulation.

Similar to intra-arterial chemotherapy, the most commonly injected drugs are melphalan and topotecan. Other drugs, such as carboplatin and etoposide phosphate, are under investigation in preclinical studies with rabbits as a potential alternative to melphalan, but will not be discussed here as they are not currently used clinically (Mohney et al., 2017).

#### 7.1.3.1. Intravitreal distribution of intravitreal melphalan

Intravitreal melphalan (15 μg-dose) in rabbits resulted in vitreous concentrations above the IC50 for up to 5 h, while undetectable in plasma throughout the 12-h period of the study (Buitrago et al., 2016) (Fig. 19). The high selectivity for the vitreous and the low systemic exposure of melphalan after intravitreal injection in the animal study are in agreement with the notable efficacy for vitreous seeds control and the lack of hematologic toxicity reported in patients treated this way (see 7.2.5.4). On the other hand, the drug aqueous humor exposure was less than 1% of that in the vitreous, making it inadequate for targeting anterior chamber seeding (Munier et al., 2017a).

Although data are limited, binding of drugs to vitreous proteins as well as drug interaction with melanin has been hypothesized to lead to prolonged residence time in the vitreous humor and pigmented tissues (iris, ciliary body, retinal pigment epithelium, choroid) respectively, resulting in prolonged pharmacological toxicity (Rimpela et al., 2018). This may explain, at least in part, the retinal and iris toxicity reported after intravitreal melphalan in patients or in a preclinical rabbit model, as well as the greater reduction in electroretinogram (ERG) components observed in more pigmented human eyes (Francis et al., 2014, 2015b, 2017b).

#### 7.1.3.2. Intravitreal distribution of intravitreal topotecan

In rabbits, the intravitreal administration of only 5 μg of topotecan led to vitreous concentrations above the IC50 for up to 16 h after the injection (Buitrago et al., 2010), with no retinal toxicity (Buitrago et al., 2010).

![Fig. 19. Melphalan concentration versus time profile in intravitreal administration to rabbits.](image-url)
and outer blood-retinal barrier, as well as by the orbital and choroidal bioavailability of the chemotherapeutic drugs. After periocular injection, to be successful in treating aqueous seeding, although insufficient to control intravitreous disease (Munier et al., 2015, 2017a).

7.1.5. Periocular drug delivery

Periocular delivery was developed before the use of intra-arterial and intravitreal chemotherapy in an attempt to increase the intraocular bioavailability of the chemotherapeutic drugs. After periocular injection, the intraocular bioavailability is limited by the sclera, choroidal and outer blood-retinal barrier, as well as by the orbital and choroidal clearance (Del Amo et al., 2017; Edelhauser et al., 2010). The diffusion of periocular injected drugs can be stimulated by transcleral iontophoresis, as shown for low-molecular-weight charged molecules, such as carboplatin (Hayden et al., 2006).

Preclinical studies in rabbits showed a 6-fold increase in vitreous-to-plasma AUC after periocular carboplatin compared to the same ratio after intravenous infusion (Hayden et al., 2004; Simpson et al., 2002) and, in the non-human primate, the vitreous and aqueous humor \(C_{\text{max}}\) was over 2-fold the levels attained in plasma (Mendelsohn et al., 1998).

Similar studies with periocular topotecan showed vitreous drug exposure of only 30% of the systemic AUC after periocular injection in rabbits (Carcaboso et al., 2007) (Fig. 19), whereas in non-tumor-bearing rats and pigs, topotecan vitreous AUC was 2- and 3-fold the systemic exposure, respectively (Nemeth et al., 2011; Schaiquevich et al., 2012a), possibly due to inter-species differences in the orbital and ocular vasculature. Lastly, pharmacokinetic data on topotecan after periocular delivery in humans obtained from a Phase I study in retinoblastoma patients, revealed a non-toxic systemic exposure to lactone topotecan below 55 ng/mL/h (Chantada et al., 2009).

7.1.6. Suprachoroidal drug delivery

Despite not being explored for targeting retinal diseases, suprachoroidal administration may provide an alternative route for drug delivery to the posterior segment. The drug is placed in the space between the sclera and choroid and could be used to localize drug delivery to the choroid, retinal pigment epithelium and, to a lesser extent, the retina (Patel et al., 2011). A rapid diffusion of the injected solution around the eye has been shown in animal models (Chiang et al., 2018). An interesting aspect is the use of microneedles for performing the injection that allows minimal damage of the sclera and avoids penetration into the choroid and retina (Hartman and Kompella, 2018; Patel et al., 2011).

Although suprachoroidal drug delivery may be advantageous for targeting the choroid and retinal pigment epithelium, less evidence is available to demonstrate adequate and prolonged exposure in the retina (Chiang et al., 2018; Del Amo et al., 2017).

7.2. Treatment modalities

Retinoblastoma has always been credited with the highest 5-year disease-free survival rate among pediatric cancers, presently reaching 97–99% in high-income countries (Fernandes et al., 2018; Group et al., 2013; Jenkinson, 2015). Until the recent introduction of new targeted treatment modalities, namely intra-arterial and intravitreal chemotherapy, survival was often only achieved using mutilating and disabling therapies. During the irradiation era, 96% of bilaterally affected children had their worst eye enucleated at diagnosis, with primary enucleation of both eyes necessary in 3% and secondary enucleation of an only remaining eye due to treatment failure leading to total blindness in 25–29% (Abramson et al., 2004; Epstein et al., 2003)

Despite not being explored for targeting retinal diseases, suprachoroidal administration may provide an alternative route for drug delivery to the posterior segment. The drug is placed in the space between the sclera and choroid and could be used to localize drug delivery to the choroid, retinal pigment epithelium and, to a lesser extent, the retina (Hartman and Kompella, 2018; Patel et al., 2011).

Although suprachoroidal drug delivery may be advantageous for targeting the choroid and retinal pigment epithelium, less evidence is available to demonstrate adequate and prolonged exposure in the retina (Chiang et al., 2018; Del Amo et al., 2017).

7.1.4. Intracameral drug delivery

Intracameral injections were specifically developed to target the anterior and posterior chambers as aqueous seeding had remained untreatable with other delivery routes. Thus, despite drug penetration to the anterior chamber after intravenous (see 7.1.1) or periocular delivery (see 7.1.5) achieving an even higher aqueous Cmax concentration compared to the intravitreal one in both human and animal eyes (see 7.1.3), the obtained doses were not high enough to reach tumoricidal levels. On the other hand, neither intravitreal melphalan (Buitrago et al., 2016) nor topotecan (Buitrago et al., 2010) achieved pharmacologically active levels in the aqueous humor (see 7.1.3) in a non-tumor bearing rabbit model. Finally, although preclinical studies have not been designed to study aqueous concentrations after intra-arterial chemotherapy, the latter has never been reported to be successful in the management of aqueous seeding (see 7.2.4).

To date, very limited data are available on the drug pharmacokinetics after intracameral injections, but clinical evidence showed them to be successful in treating aqueous seeding, although insufficient to control intravitreous disease (Munier et al., 2015, 2017a).

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is also expected to reduce the incidence of binocular visual impairment and blindness in this population (Stacey et al., 2019).

In Lausanne, the simultaneous availability since 2008 of both intra-arterial and intravitreal chemotherapy as additional options to the traditional chemoreduction and focal treatment, contributed to achieve 5-year ocular survival estimates of almost 100% for treatment-naïve IIRC group A-C (98%, 99% CI 86.7%–99.7%) (Fig. 21), and significantly improved the ocular survival of treatment-naïve IIRC group D and selected E eyes, with Kaplan-Meier estimates of 85.7% (95%CI 75.7%–91.8%) at 2-years and 79.4% (95% CI 67.0%–87.6%) at 5-years (very similar to the 80.2% for COG group D and E after Abramson (Abramson et al., 2017a)), with no metastasis in the cohort of naïve cases. This compares favorably to the era prior to 2008, where ocular survival estimates were 55.0%, (95% CI 38.7%–68.7%) at 2- and 49.2% (95% CI 33.0–63.6%) at 5-years (p = 0.0007) (Fig. 22, data shown for group D only).

7.2.1. Intravenous chemotherapy

The indications for intravenous chemotherapy in the treatment of retinoblastoma have progressively changed over the years. During the era of external beam radiotherapy (Ellsworth, 1969), systemic chemotherapy was limited to extracocular (Doz et al., 1995; Pratt et al., 1985) or metastatic disease (Saleh et al., 1988). In the 1990s, its role extended to first line treatment with the concept of chemoreduction, aiming to render the tumor volume amenable to focal treatment (Greenwald and Strauss, 1996; Kingston et al., 1996; Murphree et al., 1996; Shields et al., 1996). Systemic chemotherapy contributed to increase globe salvage rate, significantly reducing the need for external beam radiotherapy and the incidence of secondary primary malignancies. To date, systemic chemotherapy has been supplanted in part by the introduction of intra-arterial (see 7.2.4) and intravitreal (see 7.2.5) chemotherapy (Abramson et al., 2015b).

7.2.1.1. Drug regimen and number of cycles. Initiation of systemic chemotherapy usually takes place within 24–48 h after the first ophthalmological examination to take advantage of the focal cryorupture performed to improve drug penetration into the vitreous and/or subretinal fluid (Wilson et al., 1996). Before each new chemotherapy cycle, the child undergoes an examination under anesthesia in order to evaluate the tumor response. After completion of the second course of chemotherapy, sequential intensive focal treatment is applied for every lesion that is not flat or calcified, including advanced group IIRC D and selected E eyes, have been now replaced by intra-arterial, combined or not with intravitreal chemotherapy (Abramson et al., 2015b).

Kaplan-Meier for eye retention in Rb pre and post 2008 A-C eyes

![Kaplan-Meier for eye retention in Rb pre and post 2008 A-C eyes](image)

**Fig. 21.** Five-year Kaplan-Meier survival curves for retinoblastoma IIRC group A-C eyes initially treated in Lausanne before and after 2008.
7.2.2. Treatment outcomes

7.2.2.1. Relapse rate and role of consolidation by focal treatments. Since its introduction in the conservative treatment of retinoblastoma, systemic chemotherapy was commonly used in combination with focal treatment by most groups (see Table 9). The role of consolidation was specifically pointed out in a study which demonstrated that chemotherapy alone (without consolidation) failed complete inactivation of the disease in 92% of treated eyes, even after completion of 6 cycles (Wilson et al., 2001). When retinal tumors were treated by combined chemoreduction and focal therapy, disease control was shown to improve substantially (p = 0.05), with a 2.5 reduction of the relapse risk (Shields et al., 2004b). In addition, focal treatment given following chemoreduction as prophylactic consolidation (Wilson et al., 2005) rather than deferred until documented disease progression (Rodriguez-Galindo et al., 2003) was credited to decrease the need for external beam radiotherapy for salvage after chemoreduction from 44 to 25–30% (Wilson et al., 2005).

7.2.2.2. Eye preservation without external beam radiotherapy. Despite the concomitant use of diverse grouping classifications, and the variety of protocols in terms of chemotherapy regimen and number of cycles making comparison between studies difficult, major studies with at least 2-year mean follow-up claim an overall tumor control of 70–100% for RE I to III or IIRC group A to C and 23–64% for RE IV/V or IIRC group D/E (Table 9). Despite an unsatisfactory outcome in advanced eyes, intravenous chemotherapy contributed, however, overall to a significant decrease in the use of irradiation and consequently the incidence of secondary radiation-induced tumors (see 3.8).

7.2.3. Chemotherapy-related toxicity

Chemotherapy-related toxicity is one of the major concerns in the treatment of pediatric cancers. Even if common and relatively easily managed in centers of high-income countries, chemotherapy-related adverse effects may sometimes result in fatal toxicities in less developed countries (Luna-Fineman et al., 2019).

7.2.3.1. Expected toxicities. Systemic chemoreduction with standard carboplatin-etoposide ± vincristine regimens usually causes mild acute toxicity, mostly involving alopecia, nausea/vomiting, and moderate myelotoxicity, occasionally requiring hospitalization for fever, neutropenia or blood product transfusions. The use of more intense regimens is predictably associated with higher myelotoxicity. Febrile neutropenia requiring hospitalization has been variably reported in 12.7–38% of the patients, depending on the specific chemotherapeutic regimen (Bartuma et al., 2014; Berry et al., 2013; Chung et al., 2008; Kunkele et al., 2013). The need for transfusion of blood products is also inconsistently reported, ranging from 0% (Berry et al., 2015) to 78% (Brennan et al., 2017) of the patients depending on the intensity of treatment. Support with Granulocyte-colony stimulating factor was used by some groups for 100% of the patients (Brennan et al., 2017), whereas others completely avoided its use (Chantada et al., 2014).

7.2.3.2. Long-term toxicities. As previously discussed (see 3.8.3.2.2), intravenous etoposide may induce secondary leukemias, even in patients who received what are considered safe cumulative dosages (Chantada et al., 2014; Gombos et al., 2007). Permanent ototoxicity is an important late effect of carboplatin-based chemoreduction regimens reported in up to 5–25% of the cases, especially if given at a younger age and with prolonged exposure (Jehanne et al., 2009; Qaddoumi et al., 2012; Soliman et al., 2018).

7.2.4. Intra-arterial chemotherapy

The success of intra-arterial chemotherapy lies in the selective delivery of high dose drug(s) to the eye with minimal systemic absorption. The modern technique of drug injection directly into the ophthalmic artery through a trans-femoral microcatheter placed just beyond its ostium, also called super-selective intra-arterial chemotherapy or ophthalmic artery chemoembolization, was introduced in 2008 by Abramson et al. (2008), inspired by previous work by Kaneko et al. (Kaneko and Takayama, 1990) and Yamane et al. (2004).

In the hands of well-trained interventional neuro-radiologists, the procedure is safe and reaches a technical success rate of around 96–99% (Francis et al., 2018a; Munier et al., 2011; Suzuki et al., 2011),

![Kaplan-Meier for eye retention in Retinoblastoma pre and post 2008 D eyes](image-url)
Table 9
Summary of the major studies on systemic chemotherapy as first line treatment for retinoblastoma.

<table>
<thead>
<tr>
<th>Author /year</th>
<th>N eyes/patients</th>
<th>Drug regimen</th>
<th>Mean n Cycles (range)</th>
<th>Focal Treatments</th>
<th>Globe salvage rate without EBR (%)</th>
<th>Overall globe salvage rate (%) (including EBR)</th>
<th>Mean follow-up (months) (range)</th>
</tr>
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<tbody>
<tr>
<td>Greenwald and Strauss (1996)</td>
<td>11/6</td>
<td>CE</td>
<td>6 (6-7)</td>
<td>11 (100)</td>
<td>3 (75) RE: II = 1/1 (100) III = 2/3 (67)</td>
<td>8 (73) RE: II = 1/1 (100) III = 3/3 (100) IV = 0/1 (0) V = 0/6 (67)</td>
<td>26 (12-40)</td>
</tr>
<tr>
<td>Murphree et al. (1996)</td>
<td>35/na</td>
<td>CEV</td>
<td>3 (1-6)</td>
<td>35 (100)</td>
<td>10 (100) RE: I = 3/3 (100) II = 7/7 (100)</td>
<td>15 (43) RE: IV = 1/4 (25) V = 4/21 (19)</td>
<td>na</td>
</tr>
<tr>
<td>Gallie et al. (1996)</td>
<td>28/21</td>
<td>CEV+ Ga</td>
<td>6 (2-12)</td>
<td>40 (100)</td>
<td>15 (100) RE: I = 6/6 (100) II = 5/5 (100) III = 4/4 (100)</td>
<td>26 (93) RE: I = 6/6 (100) II = 5/5 (100) III = 4/4 (100) IV = 1/1 (100) V = 10/12 (83)</td>
<td>3 (1-57)</td>
</tr>
<tr>
<td>Beck et al. (2000)</td>
<td>33/24</td>
<td>CE</td>
<td>3 (1-5)</td>
<td>33 (100)</td>
<td>18 (100) RE: I = 5/5 (100) II = 10/10 (100) III = 3/3 (100)</td>
<td>29 (88) RE: I = 5/5 (100) II = 10/10 (100) III = 3/3 (100) IV = 1/1 (100) V = 10/12 (83)</td>
<td>31 (4-41)</td>
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<td>Gombos et al. (2002)</td>
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<td>CEV</td>
<td>6</td>
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<td>22 (69) RE: I = 1/4 (25) II = 17/19 (89) III = 4/9 (44)</td>
<td>34 (81) * RE: I = 4/4 (100) II = 17/19 (89) III = 7/9 (78) IV = 1/2 (50) V = 5/8 (63)</td>
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<td>CEV</td>
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<td>15 (75)</td>
<td>8 (100) RE: I = 3/3 (100) II = 3/3 (100) III = 2/2 (100)</td>
<td>16 (80) * * RE: I = 3/3 (100) II = 3/3 (100) III = 2/2 (100)</td>
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<td>Rodriguez-Galindo et al. (2003)</td>
<td>43/25</td>
<td>CV</td>
<td>8 (6-8)</td>
<td>0 (0)</td>
<td>15 (63) RE: I = 6/7 (86) II = 7/12 (64) III = 2/5 (40)</td>
<td>30 (70) RE: I = 7/7 (100) II = 10/12 (83) III = 3/5 (60) IV = 0/3 (0) V = 10/16 (63)</td>
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<td>75 (71)</td>
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<td>73 (70) RE: I = 4/4 (100) II = 22/22 (97) III = 6/7 (86) IV = 28/35 (80) V = 13/36 (36)</td>
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<tr>
<th>Author /year</th>
<th>N eyes/patients</th>
<th>Drug regimen</th>
<th>Mean n Cycles (range)</th>
<th>Focal Treatments N = eyes (%)*</th>
<th>Globe salvage rate without EBR (%)</th>
<th>Overall globe salvage rate (%) (including EBR)</th>
<th>Mean follow-up (months) (range)</th>
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<td>CE</td>
<td>(4-8)</td>
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<td>CE (98% patients) Cyclophosphamide-vincristine (2% patients)</td>
<td>5 (1-15)§ yes (no further details)</td>
<td>na</td>
<td>na</td>
<td>na</td>
<td>92 (80)§§ IIRC: A=19/19 (100) B=46/48 (96) C=15/19 (79) D=12/29 (41)</td>
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<td>249 (100)</td>
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<td>51 (47) ICRB: D = 51/109 (47)</td>
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<tr>
<td>Shin et al. (2010)</td>
<td>65/52</td>
<td>cisplatin, etoposide, vincristine or CEV or cisplatin cyclophosphamide adriamycin, etoposide ± vincristine, or ifosfamide, etoposide, vincristine</td>
<td>13 (1–29)</td>
<td>45 (69)</td>
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Table 9. continued

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<th>Author</th>
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<th>Number of cycles</th>
<th>Mean n</th>
<th>Focal Treatments</th>
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<td>64 (100)</td>
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Mean follow-up (months) (range)

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<th>Group I-III/A-C</th>
<th>Group IV-V/D-E</th>
</tr>
</thead>
<tbody>
<tr>
<td>23/23 CE</td>
<td>101 (19–219)</td>
</tr>
</tbody>
</table>

Overall globe salvage rate (%)(including EBR)

| Munier et al. (2017b) | 78 |
| Brennan et al. (2017) | 100 |

Globe salvage rate without EBR (%) Overall globe salvage rate (%)(including EBR)

| Munier et al. (2017b) | D = 13/23 (57) |
| Brennan et al. (2017) | D = 18/25 (72) |

7.2.4.1. Drug regimen and number of cycles. There is no consensus to date either on the minimal drug dose required, or on any advantage to start the treatment with a single versus a combination of drugs. The drug most commonly used is melphalan. Topotecan and/or carboplatin can be used in combination with melphalan (2 or 3-drug regimen) to capitalize on their synergistic effects. One to 3 cycles of melphalan are generally given on a 3–4-week basis in intra-arterial chemotherapy-naive eyes, and a combination of drugs (usually melphalan and topotecan) for tumor relapse or poor response following previous treatment with melphalan only.

Carboplatin is usually given to minimize systemic side-effects in the less advanced eye when combined (or so-called tandem) intra-arterial chemotherapy is performed. When doing so, the more advanced eye is usually treated with melphalan (with or without topotecan). Drugs between the eyes are then switched during the second or third session (Abramson et al., 2010, 2016b).

7.2.4.2. Indications

7.2.4.2.1. Current indications. Initially used as salvage treatment for eyes that did not resolve with systemic chemotherapy and/or radiotherapy, indications for intra-arterial chemotherapy extended to first line treatment for naïve unilateral eyes (group B with macula and/or papilla involvement, and group C and D). Other indications include the more affected eye in bilateral cases with asymmetric presentation, provided that the fellow eye is accessible to focal therapy. Simultaneous tandem therapy is performed by some groups for bilateral cases with any combination of group B with macula/papilla involvement, C, D or selected E eyes, namely those characterized by tumor touching the lens with or without total retinal detachment (Abrahamson et al., 2015b; Rowlands et al., 2018) (Table 10), whereas some centers prefer to start with chemoreduction, followed by intra-arterial chemotherapy if needed for salvage treatment (Abramson et al., 2015b).

Group E eyes with neovascular glaucoma, buphthalmos, orbital cellulitis, anterior chamber invasion or phthisis undergo enucleation. In cases where enucleation cannot be performed safely, neoadjuvant systemic chemotherapy can be advised prior to surgery, especially in eyes with buphthalmos and/or radiological evidence of optic nerve invasion (Bellaton et al., 2003).

7.2.4.2.2. Potential future indications. Intra-arterial chemotherapy with combined melphalan and topotecan was reported to successfully treat one case of a choroidal relapse that showed complete regression on UBM three weeks after the first injection. The child received two more intra-arterial injections and has since then remained metastasis- and recurrence-free at an updated follow-up of more than 4 years (Stathopoulos et al., 2018b). In another recent report, two courses of intra-arterial melphalan enabled to control iris invasion in a monophthalmic patient, but not a concomitant intra-ciliary involvement responding only to brachytherapy with an updated disease-free follow-up of more than 20 months (Munier et al., 2018).

The differential efficacy of intra-arterial chemotherapy on iris and choroid vs ciliary body may be explained by the lower vascular supply provided that it is performed in children with sufficient vascular maturation (children older than 3 months and with a body weight of over 5 kg) (Gobin et al., 2012). Anatomic variants such as departure of the ophthalmic artery from the meningeal artery, or unfavorable angle with the internal carotid artery compromising direct access to the ophthalmic artery for the delivery of the chemotherapy (around 16% of the procedures) (Abramson et al., 2012a), can usually be overcome with the use of alternative non-ophthalmic routes and/or of a temporary balloon-occlusion of the internal carotid (Klufas et al., 2012). When assessing treatment response one should be aware that anastomoses between the external and internal carotid or stenosis of its ostium, as a consequence of a vasospasm or micro-dissection from a previous procedure, may alter the local hemodynamic flow of the ophthalmic artery, and lead to insufficient drug perfusions (Francis et al., 2018a).
of the latter consisting of 2 long posterior ciliary arteries vs 7 anterior and 20 short posterior ciliary arteries vascularizing iris and choroid respectively.

7.2.4.3. Treatment outcomes. Treatment outcomes of the major studies obtained with first-line and salvage intra-arterial chemotherapy are summarized in Table 11.

7.2.4.3.1. Recurrence rate and salvage therapy. Intra-arterial chemotherapy appears to be associated with an increased risk of recurrence if administered more than 4 weeks apart (p < 0.05) and if more than 50% of the injections are delivered via a non-ophthalmic route of delivery (p < 0.001) (Francis et al., 2018a). The role of focal consolidation remains to be determined. Recurrences after first line or salvage intra-arterial chemotherapy have been observed within the first year following treatment completion in 24–25% of the cases receiving also focal consolidation (Francis et al., 2018a; Munier et al., 2017b), and in 29% of those without consolidation (Tuncer et al., 2016).

Relapse after intra-arterial chemotherapy can be successfully managed with focal treatments, including intravitreal chemotherapy in more than half of the cases (Francis et al., 2018a; Munier et al., 2017b). A second course of intra-arterial chemotherapy can, however also be given if needed, after first-line or salvage intra-arterial chemotherapy, with a reported globe salvage rate of about 76% (Francis et al., 2015a; Shields et al., 2015).

7.2.4.3.2. Eye preservation without external beam radiotherapy. Comparison of the studies on globe salvage rate with intra-arterial chemotherapy is not only challenged by the lack of randomized clinical trials and the use of different versions of the retinoblastoma classification (see 3.3.1), but also by the availability of intravitreal chemotherapy at the time of the treatment. A survey of the literature during the four years where only intra-arterial was available, i.e. before the 2012 publication of the safety enhanced technique for intravitreal chemotherapy, reveals that intra-arterial chemotherapy as first line treatment achieved globe salvage without the need of external beam irradiation in 96% group B and C eyes at a median follow-up of 16 months (Abramson et al., 2012a), 81% at 2 years of group D (Reese-Ellsworth V) (Gobin et al., 2011), but only 64% at 2 years for those with vitreous seeding (including group C, D and E) (Abramson et al., 2012c).

After 2012, first line intra-arterial chemotherapy associated with intravitreal chemotherapy as necessary, achieved globe salvage without the need of external beam irradiation in 100% ICRB/IIRC group B and C at a mean follow-up of 19 months (Shields et al., 2014), a 5 year-ocular survival of 85–100% COG/IIRC group D (Abramson et al., 2017a; Munier et al., 2017b), and 36% in ICRB group E at a mean follow-up of 19 months (Shields et al., 2014).

7.2.4.4. Adverse effects. Intra-arterial chemotherapy is safe, with no reported fatal complications and fewer systemic adverse effects compared to intravenous chemotherapy and external beam irradiation. Only one case of delayed neurological complication that spontaneously resolved has been described to date (De la Huerta et al., 2016). An intra-operative autonomous reflex characterized by hypoxia, reduced lung compliance, systemic hypotension and bradycardia, that can be easily managed by an experienced anesthesiologist, develops in

**Table 10**

<table>
<thead>
<tr>
<th>Disease in first eye</th>
<th>Treatment</th>
<th>Disease in second eye</th>
<th>Treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Group A and B macula/papilla spared</td>
<td>Laser photocoagulation</td>
<td>Group A and B macula/papilla spared</td>
<td>Laser photocoagulation</td>
</tr>
<tr>
<td>Group A and B macula/papilla spared</td>
<td>Cryocoagulation</td>
<td></td>
<td>Cryocoagulation</td>
</tr>
<tr>
<td>Group A and B macula/papilla spared</td>
<td>Plaque</td>
<td></td>
<td>Plaque</td>
</tr>
<tr>
<td>Group B macula/papilla involved</td>
<td>Laser photocoagulation</td>
<td>Group B macula/papilla involved</td>
<td>IAC ± IVIC</td>
</tr>
<tr>
<td>Group C, D and selected E</td>
<td>Cryocoagulation</td>
<td>Group C, D and selected E</td>
<td>Plaque</td>
</tr>
<tr>
<td>Groups B macula/papilla involved</td>
<td>Plaque</td>
<td>Group B macula/papilla involved</td>
<td>IAC ± IVIC</td>
</tr>
<tr>
<td>Group C, D and selected E</td>
<td>IAC ± IVIC or IVC</td>
<td>Group C, D and selected E</td>
<td>IAC ± IVIC or IVC</td>
</tr>
</tbody>
</table>

Table 11
Summary of the major studies on first line and salvage intra-arterial chemotherapy for retinoblastoma.

<table>
<thead>
<tr>
<th>Author (year)</th>
<th>Eyes/ Patients</th>
<th>Classification Group: n (%eyes)</th>
<th>Treatment Prior IAC, n (%eyes)</th>
<th>Drug used, n (%eyes)</th>
<th>Mean cycles (range)</th>
<th>Additional treatments post IAC, n (%eyes)</th>
<th>Globe salvage, n (%group)</th>
<th>Mean/ median follow-up, months (range)</th>
<th>Extracocular extension, n (%patients)</th>
<th>Pineoblastoma, n (%patients)</th>
<th>Metastasis, n (%patients)</th>
<th>Deaths, n (% patients)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Abramson et al. (2012a)</td>
<td>30/na</td>
<td>1° = 30 (100)</td>
<td>COG: B = 19 (63) C = 11 (37)</td>
<td>None = 30 (100) M = 30 (100) T = na (na)</td>
<td>Laser = 15 (50) cryo = 1 (3) cryo + laser = 1 (10) Focal trt = 17 (46) Plaque = 4 (11)</td>
<td>1° = 24 (89)</td>
<td>2° = 10 (27)</td>
<td>1° = 24 (89)</td>
<td>0 (0)</td>
<td>0 (0)</td>
<td>0 (0)</td>
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</tr>
<tr>
<td>Palirou et al. (2012)</td>
<td>37/37</td>
<td>1° = 27 (73) 2° = 10 (27)</td>
<td>COG: B = 11 (41) C = 16 (59) D = 10 (34) E = 6 (16)</td>
<td>None = 27 (100) M = na (na) T = na (na) IVC = 9 (90) ERRT = 3 (30)</td>
<td>1° = 24 (89)</td>
<td>2° = 10 (27)</td>
<td>1° = 24 (89)</td>
<td>2° = 10 (27)</td>
<td>1° = 24 (89)</td>
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<td>1 (3)</td>
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<tr>
<td>Thampi et al. (2013)</td>
<td>12/10</td>
<td>1° = 12 (100)</td>
<td>ICRB: B = 3 (25) C = 1 (8) D = 6 (50) E = 2 (17)</td>
<td>None = 12 (100) M = 12 (100)</td>
<td>Local therapy = 2 (17)</td>
<td>1° = 24 (89)</td>
<td>2° = 10 (27)</td>
<td>1° = 24 (89)</td>
<td>0 (0)</td>
<td>0 (0)</td>
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</tr>
<tr>
<td>Venturi et al. (2013)</td>
<td>39/36</td>
<td>1° = 17 (44) 2° = 22 (56)</td>
<td>TNM: 1a = 1 (6) 1b = 1 (6) 2a = 7 (41) 2b = 4 (24) 3a = 4 (24) 3b = 11 (50) 2a = 1 (5) 2b = 10 (45)</td>
<td>None = 17 (100) M = 22 (100)</td>
<td>Local therapy = 2 (17)</td>
<td>1° = 24 (89)</td>
<td>2° = 22 (56)</td>
<td>1° = 24 (89)</td>
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<tr>
<td>Bracco et al. (2013)</td>
<td>52/48</td>
<td>1° = 22 (42) 2° = 30 (58)</td>
<td>INC: B = 4 (18) C = 1 (8) D = 18 (82) E = 2 (17)</td>
<td>None = 22 (100) M = 22 (100)</td>
<td>Local therapy = 2 (17)</td>
<td>1° = 24 (89)</td>
<td>2° = 30 (58)</td>
<td>1° = 24 (89)</td>
<td>2° = 30 (58)</td>
<td>1° = 24 (89)</td>
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<td>0 (0)</td>
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<tr>
<td>Shields et al. (2014)</td>
<td>70/70</td>
<td>1° = 36 2° = 34</td>
<td>ICRB: B = 1 (3) C = 4 (11) D = 17 (47) E = 14 (39)</td>
<td>None = 36 (100)</td>
<td>Local therapy = 2 (17)</td>
<td>1° = 24 (89)</td>
<td>2° = 34 (56)</td>
<td>1° = 24 (89)</td>
<td>2° = 34 (56)</td>
<td>1° = 24 (89)</td>
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<tr>
<td>Ong et al. (2015)</td>
<td>17/12</td>
<td>1° = 6 (35) 2° = 11 (65)</td>
<td>ICRB: B = 1 (17) C = 1 (9) D = 1 (17) E = 4 (66)</td>
<td>None = 0 (100)</td>
<td>Local therapy = 2 (17)</td>
<td>1° = 24 (89)</td>
<td>2° = 11 (65)</td>
<td>1° = 24 (89)</td>
<td>2° = 11 (65)</td>
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(continued on next page)
<table>
<thead>
<tr>
<th>Author (year)</th>
<th>Eyes/ Patients</th>
<th>1° = first line IAC</th>
<th>Classification Group: n (%eyes)</th>
<th>Treatment Prior IAC, n (%eyes)</th>
<th>Drug used, n (%eyes)</th>
<th>Mean cycles (range)</th>
<th>Additional treatments post IAC, n (%eyes)</th>
<th>Globe salvage, n (%group)</th>
<th>Mean/median follow-up, months (range)</th>
<th>Extraocular extension, n (%eyes)</th>
<th>Pineoblastoma, n (%patients)</th>
<th>Metastasis, n (%patients)</th>
<th>Deaths, n (%patients)</th>
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</thead>
<tbody>
<tr>
<td>Yannuzzi et al. (2015)</td>
<td></td>
<td>COG: C = 2 (Gebulla et al.) D = 52 (68) E = 23 (30)</td>
<td>None = 77 (100)</td>
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<td>TTT = 45 (58) Cryo = 34 (44) ERBT = 1 (13) Plaque = 12 (15.6) POC = 3 (3.9) IV/C = 7 (9.1) IV/C = 5 (6.5)</td>
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<td>Akyuz et al. (2015)</td>
<td>56/46</td>
<td>1° = 12 (21)</td>
<td>ICRB: C = 4 (33) D = 6 (50) E = 2 (17) A = 7 (16) B = 6 (14) C = 12 (27) D = 13 (30) E = 6 (13)</td>
<td>None = 12 (100)</td>
<td>M = 12 (100)</td>
<td>3 (1-5)</td>
<td>IVC = 1 (1) Local therapy = 1 (1) Radiotherapy = 3 (5)</td>
<td>1° = 9 (75)</td>
<td>21 (1-27)</td>
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<td></td>
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<td>2° = 44 (79)</td>
<td>IVC = 44 (100)</td>
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<td>M = 44 (100)</td>
<td>2 (1-7)</td>
<td></td>
<td>2° = 28 (64)</td>
<td>12 (1-28)</td>
<td>1 (3)</td>
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<td>2 (6)</td>
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<td>112/103</td>
<td>1° = 54 (48)</td>
<td>COG: D = 54 (100)</td>
<td>None = 47 (87) Bridge IVC = 7 (13) IVC = 51 (100) EBRT = 15 (26) Plaque = 4 (7) POC = 9 (16) Cryo only = 1 (2)</td>
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<td>2° = 58 (52)</td>
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<td>4 (1-9)</td>
<td>Laser + cryo = 54 (100) POC = 3 (3) Plaque = 12 (11) IVC = 2 (2) ERRT = 2 (2)</td>
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<tr>
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<td>66/66</td>
<td>1° = 66 (100)</td>
<td>ICRB: B = 3 (4) C = 4 (6) D = 36 (55) E = 23 (35) A = 7 (16) B = 6 (14) C = 12 (27) D = 13 (30) E = 6 (13)</td>
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<td>M = 34 (52) M + T = 32 (48)</td>
<td>3 (1-6)</td>
<td>h/t/C = 11 (17) Focal trt not mentioned</td>
<td>1° = 48 (73)</td>
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<td>ICRB: D = 17 (71) E = 7 (29)</td>
<td>None = 24 (100)</td>
<td>M = 20 (84) M + T = 2 (8) M + T + C = 2 (8)</td>
<td>3 (2-5)</td>
<td>TTT = 2 (8) Cryo = 4 (17) Plaque = 2 (8) h/t/C = 3 (13) Cryo ± trt ± Laser h/t/C IAC = 1 (4) Plaques, TTT, POC, cryo, IV/C given in some</td>
<td>1° = 25 (100)</td>
<td>42 (20-90)</td>
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<td>M = 25 (100)</td>
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<tr>
<td>Abramson et al. (2017a)</td>
<td>106/na</td>
<td>1° = 106 (100)</td>
<td>COG: D = 74 (70) COG: E = 32 (30)</td>
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<td>na</td>
<td>na</td>
<td>na</td>
<td>5-year Kaplan-Meier estimates: 80%</td>
<td>28 (0.5-101)</td>
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<td>na</td>
<td>3 (2)</td>
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<td>ICRB: B = 11 (10) C = 11 (10) D = 56 (53) E = 29 (27)</td>
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<td>M = 40 (37) M + T = 67 (63)</td>
<td>3 (2-5)</td>
<td>na</td>
<td>1° + 2° = 84 (79)</td>
<td>B = 11 (100) C = 11 (100) D = 44 (79) E = 18 (62)</td>
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<td>2° = 77 (72)</td>
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</tbody>
</table>
Deaths, n(% patients)

Pineoblastoma, n(% patients)

IVC=12 (34) M-containing

Funes et al.

97/81 1°=35 (36) IIRC:

5 (1–8) IvitC=1 (3)

Table 1 (continued)

<table>
<thead>
<tr>
<th>Classification</th>
<th>Patients</th>
<th>IAC n (%eyes)</th>
<th>Total n (%eyes)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1°=first line</td>
<td>173/209</td>
<td>1°=82 (41)</td>
<td>2°=62 (36)</td>
</tr>
<tr>
<td>± subretinal</td>
<td>86/209</td>
<td>1°=23 (31)</td>
<td>2°=33 (40)</td>
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<tr>
<td>± external</td>
<td>84/209</td>
<td>1°=36 (39)</td>
<td>2°=27 (32)</td>
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<tr>
<td>± systemic</td>
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<tr>
<td>± chemotherapy</td>
<td>84/209</td>
<td>1°=30 (36)</td>
<td>2°=24 (29)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Eyes/ Patients</th>
<th>1 = first line</th>
<th>2 = salvage</th>
<th>Group n (%eyes)</th>
<th>Drug used, n (%eyes)</th>
<th>Mean/median follow-up, years</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 = first line</td>
<td>173/209</td>
<td>1°=82 (41)</td>
<td>2°=62 (36)</td>
<td>1°=26 (16)</td>
<td>1°=24 (69)</td>
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<tr>
<td>± subretinal</td>
<td>86/209</td>
<td>1°=23 (31)</td>
<td>2°=33 (40)</td>
<td>1°=19 (18)</td>
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<td>± external</td>
<td>84/209</td>
<td>1°=36 (39)</td>
<td>2°=27 (32)</td>
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</tr>
<tr>
<td>± systemic</td>
<td>84/209</td>
<td>1°=29 (35)</td>
<td>2°=25 (30)</td>
<td>1°=19 (18)</td>
<td>1°=24 (69)</td>
</tr>
<tr>
<td>± chemotherapy</td>
<td>84/209</td>
<td>1°=30 (36)</td>
<td>2°=24 (29)</td>
<td>1°=19 (18)</td>
<td>1°=24 (69)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Authors</th>
<th>Year</th>
<th>Eyes/ Patients</th>
<th>Classification</th>
<th>Treatment</th>
<th>Group n (%eyes)</th>
<th>Drug used, n (%eyes)</th>
<th>Mean/median follow-up, years</th>
</tr>
</thead>
<tbody>
<tr>
<td>Funes et al.</td>
<td>2012</td>
<td>173/209</td>
<td>1°=82 (41)</td>
<td>2°=62 (36)</td>
<td>1°=26 (16)</td>
<td>1°=24 (69)</td>
<td>1°=24 (69)</td>
</tr>
<tr>
<td>± subretinal</td>
<td>86/209</td>
<td>1°=23 (31)</td>
<td>2°=33 (40)</td>
<td>1°=19 (18)</td>
<td>1°=24 (69)</td>
<td>1°=24 (69)</td>
<td>1°=24 (69)</td>
</tr>
<tr>
<td>± external</td>
<td>84/209</td>
<td>1°=36 (39)</td>
<td>2°=27 (32)</td>
<td>1°=19 (18)</td>
<td>1°=24 (69)</td>
<td>1°=24 (69)</td>
<td>1°=24 (69)</td>
</tr>
<tr>
<td>± systemic</td>
<td>84/209</td>
<td>1°=29 (35)</td>
<td>2°=25 (30)</td>
<td>1°=19 (18)</td>
<td>1°=24 (69)</td>
<td>1°=24 (69)</td>
<td>1°=24 (69)</td>
</tr>
<tr>
<td>± chemotherapy</td>
<td>84/209</td>
<td>1°=30 (36)</td>
<td>2°=24 (29)</td>
<td>1°=19 (18)</td>
<td>1°=24 (69)</td>
<td>1°=24 (69)</td>
<td>1°=24 (69)</td>
</tr>
</tbody>
</table>

Legend: M = melphalan; T = topotecan; C = carboplatin; MTX = methotrexate; ma = not available; COG = Children’s Oncology Group; IVIC = intravitreal chemotherapy; IVAC = intra-arterial chemotherapy; TA = temozolomide; POC = periocular chemotherapy; Sr C = subretinal carboplatin.

1°=first line; 2°=second line; IAC = intra-arterial chemotherapy; POC = periocular chemotherapy; Sr C = subretinal carboplatin.

Locally, adverse effects consist of redness (29%), transient eyelid edema (5%), loss of eyelashes (5%), blepharoptosis (5%), forehead hyperemia (2%), and rarely third or sixth nerve palsy, each of them resolving within 2–3 months (Wyse et al., 2016). Intraocular vascular complications, such as ophthalmic artery obstruction (2%), vitreous hemorrhage (2%), choroidal occlusive vasculopathy (7%), are of greater concern (see 7.3.5.2), especially in eyes with visual potential, as they can irreversibly compromise vision (Dalvin et al., 2018; Shields et al., 2014).

7.2.5. Intravitreal chemotherapy

The concept of intravitreal chemotherapy was confined for decades to isolated heroic attempts with thiopeta in desperate cases (Ericson and Rosengren, 1961; Seregard et al., 1995), due to the risk of extraocular tumor spread. In 1994, Akihiro Kaneko started using this route of drug delivery, injecting 8μg of melphalan combined with whole-eye hyperthermia for vitreous disease and reported a 51.3% eye survival rate at more than 50 months follow-up (Kaneko and Suzuki, 2003), comparable to the results achieved with external beam radiotherapy (Abramson and Scheffer, 2004b) and chemoreduction (47%) (Shields et al., 2002). After the introduction of intra-arterial chemotherapy in 2008, the probability of globe preservation in eyes with vitreous seeding increased to 64% in treatment-naive eyes at 2 years follow-up, but enucleation was still necessary in about 30% of the eyes due to uncontrollable vitreous disease (Abramson et al., 2012c). Similarly, in 2011, Shields et al. published a 67% control of the vitreous disease at approximately one year follow-up (Shields et al., 2011).

In 2012, the development of a safety-enhanced technique for intravitreal injection through a tumor-free pars plana site using anti-reflux measures and needle tract sterilization (Munier et al., 2012b) enabled vitreous disease to be specifically addressed and became rapidly adopted worldwide, leading not only to the reduction of the enucleation rate, but also definitively eradicating the need for external beam irradiation. Since then, long-term studies have confirmed safety and efficacy for both primary and recurrent vitreous seeding, approaching 100% of vitreous disease control (see 7.2.5.3). On the other hand, clinical observations and a better knowledge of the cellular composition of the seeding allowed us to re-evaluate the treatment response and redefine the treatment intensity according to the seed classification. Finally, increased efforts have been made to understand and minimize intravitreal melphalan-related toxicity (see 7.3.5.1.2).

7.2.5.1. Drug regimen and number of injections. As for intra-arterial chemotherapy, the most commonly used drug for intravitreal injections is melphalan. Preclinical studies have established that melphalan of at least 4μg/ml is required to fully suppress retinoblastoma cells in vitro (Inomata and Kaneko, 1987), and that at a vitreous concentration of 5.9μg/ml is functionally and structurally non-toxic to the retina of albino rabbits (Ueda et al., 1995). To date, there is no international consensus regarding the starting dose, but we recommend 20–30μg depending on eye size and tumor volume (see 7.3.5.1.2.1). A lower dose (8μg) has been reported to be ineffective,
<table>
<thead>
<tr>
<th>Classification</th>
<th>Type</th>
<th>Clinical features</th>
<th>Response to intravitreal melphalan</th>
<th>Regression types</th>
<th>Histopathologic features</th>
</tr>
</thead>
<tbody>
<tr>
<td>Active vitreous seeds</td>
<td>Class 1</td>
<td>Dust</td>
<td>Loose cellular spread usually at the edge of a retinal tumor</td>
<td>2–3 weeks to regress, receives least drug (median 20μg)/injections (median 3)</td>
<td>100% Regression Type 0: Complete disappearance</td>
</tr>
<tr>
<td></td>
<td>Class 2</td>
<td>Sphere</td>
<td>Two types: - translucent - whitish center (necrosis) surrounded by translucent mono- or multi-layered tumor cells</td>
<td>6–7 weeks to regress receives medium amount of drug (median 30μg)/injections (median 5)</td>
<td>90% Regression Type 0 10% Regression Type I: Refrangent/and or calcified vitreous seeding or/and Regression Type II: Amorphous often non-spherical inactive residues with or without pigment</td>
</tr>
<tr>
<td></td>
<td>Class 3</td>
<td>Cloud</td>
<td>Dense, cumulus-like spread infiltrates</td>
<td>30–32 weeks to regress receives most drug (median33μg)/injections (median 8)</td>
<td>55%: Regression Type 0 45%: Regression Type I Regression Type II Regression Type III: combination of Type I and II</td>
</tr>
</tbody>
</table>


Table 13
Summary of the main studies on intravitreal chemotherapy for retinoblastoma.

<table>
<thead>
<tr>
<th>Author (year)</th>
<th>Eyes/Patients</th>
<th>Drug used, n (%eyes)</th>
<th>Dose (μg), n (%eyes)</th>
<th>Mean number of injections (range)</th>
<th>Vitreous seed control, n (% eyes)</th>
<th>Recurrence, n (% of VS with control)</th>
<th>Globe salvage n (%eyes)</th>
<th>Median/mean follow-up, months (range)</th>
<th>Extraocular extension, n (%eyes)</th>
<th>Metastasis, n (% patients)</th>
<th>Deaths, n (% patients)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Munier et al. (2012a)</td>
<td>23/23</td>
<td>M = 23 (100)</td>
<td>M 20–30 = 23 (100)</td>
<td>M = 5 (2–12)</td>
<td>21 (91)</td>
<td>3 (14)§</td>
<td>20 (87)</td>
<td>22 (9–31)</td>
<td>0 (0)</td>
<td>0 (0)</td>
<td>0 (0)</td>
</tr>
<tr>
<td>Ghassemi et al. (2014)</td>
<td>9/9</td>
<td>M = 9 (100)</td>
<td>T = 9 (100)</td>
<td>M 40 = 9 (100)</td>
<td>M = 2 (1-3)</td>
<td>9 (100)</td>
<td>6 (67)</td>
<td>16 (7–25)</td>
<td>0 (0)</td>
<td>0 (0)</td>
<td>0 (0)</td>
</tr>
<tr>
<td>Francis et al. (2014)</td>
<td>16/16</td>
<td>M = 16 (100)</td>
<td>M 30 = 16 (100)</td>
<td>M = 7 (5–8)</td>
<td>15 (94)</td>
<td>1 (7)</td>
<td>14 (88)</td>
<td>5 (1–11)</td>
<td>0 (0)</td>
<td>0 (0)</td>
<td>0 (0)</td>
</tr>
<tr>
<td>Suzuki et al. (2015)</td>
<td>239/na</td>
<td>M = 239 (100)</td>
<td>M 8–24 = 24 (100)</td>
<td>M = 4 (1-25)</td>
<td>163 (68)</td>
<td>31 (19)</td>
<td>132 (55)</td>
<td>118 (33–256)</td>
<td>1 (&lt; 1)</td>
<td>11 (5)§</td>
<td>5 (2)¥</td>
</tr>
<tr>
<td>Ji et al. (2016)</td>
<td>19/17</td>
<td>M = 19 (100)</td>
<td>M 20 = 19 (100)</td>
<td>M = 6 (1-15)</td>
<td>16 (84)</td>
<td>0 (0)</td>
<td>14 (74)</td>
<td>26 (17–42)</td>
<td>0 (0)</td>
<td>0 (0)</td>
<td>0 (0)</td>
</tr>
<tr>
<td>Shields et al. (2016b)</td>
<td>40/38</td>
<td>M = 40 (100)</td>
<td>T = 11 (27)</td>
<td>M 30 = 14 (35)</td>
<td>40 (100)</td>
<td>1 (3) ±</td>
<td>35 (88)</td>
<td>36 (6–88)</td>
<td>0 (0)</td>
<td>0 (0)</td>
<td>0 (0)</td>
</tr>
<tr>
<td>Berry et al. (2017a)</td>
<td>28/25</td>
<td>M = 28 (100)</td>
<td>M 20–40 = 28 (100)</td>
<td>M = 3 (1-10)</td>
<td>28 (100)</td>
<td>0 (0)</td>
<td>19 (68)</td>
<td>33 (9–51)</td>
<td>0 (0)</td>
<td>0 (0)</td>
<td>0 (0)</td>
</tr>
<tr>
<td>Kiratli et al. (2017)</td>
<td>39/37</td>
<td>M = all</td>
<td>M 20–30 = 19 (49)</td>
<td>M = 2 (1-5)</td>
<td>27 (69)</td>
<td>0 (0)</td>
<td>22 (56)</td>
<td>12 (6–38)</td>
<td>0 (0)</td>
<td>0 (0)</td>
<td>0 (0)</td>
</tr>
<tr>
<td>Rao et al. (2018b)</td>
<td>17/17</td>
<td>T = 17 (100)</td>
<td>T 30 = 17 (100)</td>
<td>T = 3 (2-6)</td>
<td>17 (100)</td>
<td>0 (0)</td>
<td>16 (94)</td>
<td>24 (15-34)</td>
<td>0 (0)</td>
<td>0 (0)</td>
<td>0 (0)</td>
</tr>
</tbody>
</table>

Legends: N = number; M = melphalan; T = topotecan; § one was true recurrence, 7 months after first course of 5 injections, 2 were iatrogenic nature from tumor apex rupture at plaque application; ¥ 11 patients had distant metastasis or intracranial invasion (5 of them also suffered orbital recurrence), all of whom had high-risk pathological factors for metastasis but refused adjuvant chemotherapy. ± vitreous relapse with concomitant retinal recurrence. ¥ One patient lost to follow-up before completion of adjuvant chemotherapy.
whereas a higher dose (50 μg) was associated with phthisis bulbi (Ghassemi and Shields, 2012).

The second most frequently injected drug is topotecan, usually combined with melphalan in case of failure of tumor control, potentially due to melphalan resistance, or recurrences. Noteworthy, in the 10-year experience with intravitreal melphalan in Lausanne (more than 160 eyes and 750 injections), there was only a single case of true resistance to melphalan requiring topotecan to control the recurrent vitreous disease (personal observation). Compared to melphalan, topotecan has a longer intra-ocular half-life, with low ocular toxicity and also the advantage of a lower cost (Schaigevich et al., 2017).

Intravitreal chemotherapy is carried out every 7–10 days. At each visit, the residual vitreous tumor burden is reassessed and usually an additional injection for consolidation is given once complete regression is observed. Complete response is established if the seeds either completely disappear (vitreous seeding regression type 0) or convert into refringent and/or calcified residues (vitreous seeding regression type I), amorphous often non-spherical inactive residues (vitreous seeding regression type II), or a combination of the latter two (vitreous seeding regression type III) (Munier, 2014). Another useful criterion of complete response is the presence of hyper-reflective dots within class 2 seeds as shown by OCT (see 3.2.3) (Munier, 2014). The timing to complete regression is a matter of weeks, months and up to one year for class 1, 2 and 3 seeds respectively (Table 12). Since the initial recommendations (Francis et al., 2015a), the number of injections necessary to achieve tumor control for the various classes of vitreous seeds can be reduced, namely two for class 1, three for class 2, and four for class 3.

7.2.5.2. Safety. Since the adoption of the safety-enhanced technique of injection in 2012, there has been no report, to our knowledge, of any tumor exteriorization as documented by a meta-analysis of the published literature (Smith and Smith, 2013) or by retrospective analysis of more than 3500 injections performed in 10 North and South American, European, Israeli, and Chinese centers (Francis et al., 2017a). More recently, the safety of intravitreal chemotherapy was further confirmed when assessing the expression of the cone-rod homeobox gene (CRX) by real-time quantitative polymerase chain reaction as a surrogate of retinoblastoma RNA, failed to detect any CRX-positive cycle from the entry site or from the cryoprobe tip (Winter et al., 2018).

7.2.5.3. Treatment outcomes. Intravitreal melphalan is efficient with a reported vitreous seeding control in 69–100% of the cases (Table 13). Two single-institution studies have compared the outcomes in the eras before and after the introduction of intravitreal chemotherapy. In the first, complete abandon of external beam radiotherapy was effective only after the introduction of intravitreal chemotherapy (Francis et al., 2018a), while the second documented a significant improvement of ocular survival from 77 to 87%, and from 25 to 73% in ICRB group D and E eyes respectively (p < 0.04) (Shields et al., 2016a), as well as a threefold decrease in the enucleation rate during the pre-intravitreal (44%) vs intravitreal (15%) era (p = 0.012). In a study comparing the outcome of eyes with class 3 seeding (cloud) treated with intra-arterial and intravitreal chemotherapy versus intra-arterial alone, the authors demonstrated a significantly shorter time to regression (5.7 months vs 14.6 months, p < 0.001) as well as significantly fewer recurrences and fewer enucleations with the concomitant use of intravitreal chemotherapy (Francis et al., 2017c). The same group also reported on their 4-year experience with intravitreal chemotherapy in 130 eyes with a 2-year Kaplan-Meier estimate for ocular survival of 94.2% (CI 89.2–99.4%), and an estimate for disease-free survival of 86.2% (Francis et al., 2017b). According to the literature, there also seems to be no significant difference between eyes with persistent versus recurrent vitreous seeding regarding time to regression, number of injections needed, or cumulative dose of melphalan (Berry et al., 2017a; Jī et al., 2016; Kaneko and Suzuki, 2003; Munier et al., 2012a).

Until now, additional topotecan was usually considered in the case of resistant vitreous seeding and injected with concomitant melphalan (Francis et al., 2017b; Ghassemi et al., 2014; Shields et al., 2016b). In a recent paper, control of vitreous disease was achieved with intravitreal topotecan alone in 100% of the case (n = 17) at a median follow-up of 24 months with no clinical ocular toxicity (Rao et al., 2018b).

Finally, although intravitreal chemotherapy is not primarily directed against retinal or subretinal tumors, partial or complete regression of subretinal seeds after intravitreal melphalan has been documented in isolated cases (Francis et al., 2015c; Munier, 2014; Shields et al., 2016b). The efficacy of intravitreal chemotherapy for indications other than active vitreous disease remains, however, to be established (Abramson et al., 2018, 2019).

7.2.5.4. Adverse effects. Intravitreal injections of melphalan have proved to have fewer adverse effects and no systemic toxicity compared to intra-arterial chemotherapy (Francis et al., 2014). The most common side effect after standard doses of intravitreal melphalan is localized salt-and-pepper retinopathy at the site of injection, reported in 0.8–43% of the cases (Smith et al., 2014), which significantly correlates with reduction in ERG amplitudes with an estimated 5.3 μV decrease for every intravitreal melphalan injection of 25–30 μg (Francis et al., 2017b) (see 7.3.6.1). Other rare side effects include subconjunctival hemorrhage, mild vitreous hemorrhage, cataract, transient hypopyon and retinal detachment (Smith et al., 2014). Iris depigmentation, iris recession with retinal necrosis and focal scleral malacia with localized pigmentation have been reported in isolated cases (Francis et al., 2015c).

7.2.6. Intracameral chemotherapy

Aqueous seeding (see 6.2.4.1) with or without concomitant anterior uveal involvement was until recently considered an intractable form of retinoblastoma. The prognostic value of aqueous seeding and/or anterior uveal invasion such as iris or corpus ciliaris is controversial, but is no longer considered to be high risk for metastasis and, consequently, does not represent an automatic indication for adjuvant chemotherapy (Baroni et al., 2014; Sreelakshmi et al., 2017; Suryawanshi et al., 2011).

Along the same lines as for intravitreal injections, a safety-enhanced technique was developed for intracameral injections and used for the first time in 2011 to successfully treat aqueous seeding in a patient with diffuse anterior retinoblastoma (Munier et al., 2015, 2017a). Unlike vitreous volume, the exact value of aqueous volume according to age is not known, which precludes the precise determination of the aqueous drug concentration following the injection of a given dose. In order to obtain in all injected eyes the same final aqueous concentration of melphalan irrespective to age, athalamia is first created by a paracentesis emptying both anterior and posterior chambers, followed by the reconstitution of the cameral volume by injecting a solution at the desired final melphalan concentration (15–20 μg/ml) (Munier et al., 2017a).

Alternatively, it is also possible to inject very small volumes of highly concentrated drug without prior anterior chamber tap, i.e. 0.015 ml at 500 μg/ml (7.5 μg) of topotecan (Paez-Escamilla et al., 2017), or 0.05 ml at 30 μg/ml (15 μg) of melphalan (Cassoux et al., 2017). This technique, however, is suboptimal regarding the drug concentration achieved in the aqueous humor, with a risk of under-exposure at the level of the posterior chamber.

7.2.6.1. Treatment outcomes. Following the first case report on intracameral chemotherapy (Munier et al., 2017a), a series of 11 eyes with secondary aqueous seeding treated with this new technique was published (Munier et al., 2018). In this study, intracameral melphalan injection contributed to a globe preservation of 55% of the treated eyes at a mean updated event-free follow-up of 29 months (range 16–48), including 5 only eyes. Despite initial complete regression of the
aqueous seeds, as assessed by cytopathology and cell culture (Munier et al., 2018), five cases were nevertheless enucleated. Overall, the eye retention rate appeared to be less favorable in eyes with iatrogenic disrupted anterior hyaloid (25% or 1/4, salvaged), compared to eyes with intact anterior hyaloid (71% or 5/7 salvaged). Since this publication, three additional patients with anterior chamber seeding benefited from intracameral chemotherapy in Lausanne, with a recurrence-free follow-up of 3, 6, and 14 months respectively (personal observation).

7.2.6.2. Treatment-related adverse effects. All 15 patients treated in the Lausanne cohort are alive with no extracocular tumor spread or metastasis at a mean follow-up of 18 months (range 3–84 months) after a mean of 6.5 injections (range 3–14). Most frequently observed ocular side effects were cataract in 6 eyes (40%), among which 5 eyes had received concomitant brachytherapy for a retinal or ciliary body relapses, and hypochromic heterochromia in 3 (20%). Noteworthy, no potential cytotoxic effect was detected in the three patients old enough to undergo endothelial cell count (personal observation). The technique of intracameral injection having been only recently developed further studies with more cases and longer follow-up will be needed to conclude on long-term adverse effects.

7.2.6.3. Potential alternatives to intracameral chemotherapy for aqueous seeding. Aqueous seeding control was also reported in isolated or limited case series by means of intravenous chemotherapy combined with intravitreal and periocular topotecan (Rao et al., 2018a) or brachytherapy (Shields et al., 2016c). In the case from India, it is not possible to determine how tumoricidal concentration was achieved to control aqueous disease and which of the 3 routes of delivery and 4 injected drugs or combinations of drugs contributed to the result (see Fig. 20). In the series from Philadelphia, 3 eyes with anterior diffuse retinoblastoma devoid of any retinal or vitreous involvement, achieved complete remission at a mean follow-up of 35 months by means of iodine plaque therapy delivered to the entire anterior segment. All eyes needed cataract surgery within a mean interval of 16 months. With such an approach, other delayed, potentially sight-threatening complications are to be feared, such as radio-induced damage to the trabeculum and/or corneal limbal stem cells.

7.2.7. Periocular chemotherapy

Periocular chemotherapy with carboplatin or topotecan was introduced in the 1990s, generally as an adjunct to systemic chemotherapy or cryotherapy in order to increase the intracocular drug concentration (see 7.1.5). The periocular delivery can be obtained using subconjunctival (Abramson et al., 1999b) or sub-Tenonian (Francis et al., 2015) injections with or without echographic localization (Cebulla et al., 2009). Different drug vehicles have been tested, including fibrin sealant (Mallipatna et al., 2011) or nanoparticles (Kalita et al., 2014).

In a long-term study on periocular carboplatin (20 mg in 2 ml) (Marr et al., 2012), only 2/13 eyes (15%) showed long-term complete response to periocular carboplatin alone, indicating inefficiency if used as monotherapy. Reported adverse local effects could also be quite severe, including ocular motility restriction (Mulvihill et al., 2003), orbital fat atrophy and fibrosis (Kim et al., 2010b) as well as optic nerve atrophy and subsequent blindness (Schmack et al., 2006), relegate periocular carboplatin to a marginal role, if any, in the conservative management of retinoblastoma.

A phase I study on periocular topotecan showed that 2 mg of the drug diluted in 2 ml normal saline could be administered without significant toxicity (Chantada et al., 2009). A retrospective study on periocular topotecan at a median dose of 0.18 mg/kg (range 0.5–2.8 mg) in 10 eyes, reported efficacy restricted to tumor reduction for group A and B eyes, which still required, however, complementary focal treatments to be controlled. Periocular topotecan combined with systemic chemotherapy was also found to be significantly inferior to intra-arterial melphalan for salvage treatment of relapsed retinoblastoma (Schaiquevich et al., 2013).

Since the availability of intra-arterial and intravitreal chemotherapy, indications for periocular chemotherapy are limited to consolidation therapy of recurrent resistant tumors at the posterior pole in combination with other focal therapies. The recent description of long-acting episceral chemotherapy implant (chemo-plaque) could revive the periocular route of drug delivery in advanced retinoblastoma (Carcaboso et al., 2010; Pontes de Carvalho et al., 2006). Two prospective studies are currently open in the United States using a device for sustained controlled-release of topotecan for primary unilateral group D and for eyes that have failed standard available therapy (http://grantome.com/grant/NIH/RC3-CA150730-01).

7.2.8. Focal treatments modalities

Focal treatments, including cryotherapy, hyperthermia, photo-coagulation and brachytherapy, are aimed at destroying small intraocular tumors confined to the retina without the need of treating the whole eye. Because of their efficacy with few adverse effects, they represent important modalities in the therapeutic arsenal against retinoblastoma. Current indications of focal therapies include first line use in group A, macula/or papilla-sparing group B tumors, second line or consolidation after chemoreduction/chemotherapy, and salvage treatment of accessible recurrent retinal tumors. There is presently no consensus regarding the need of consolidation focal treatments following intra-arterial chemotherapy and their potential role in preventing recurrence as observed post chemoreduction (Table 11). Finally, although successful treatment of localized intravitreal seeding has been occasionally reported with brachytherapy (Abouzeid et al., 2008; Francis et al., 2013b) or cryotherapy (Abramson and Schefer, 2004a), these treatment modalities are not, however, primarily aimed at the control of vitreous disease.

All focal treatments have in common to be performed under general anesthesia. The choice of the treatment modality depends mainly on the tumor size and location. The number of sessions is adapted to the tumor response to treatment, with regression to a flat scar or totally calcified tumor being the ideal endpoint of treatment. If needed, they can be repeated (usually every 2–4 weeks) or used in combination. Presence of fish-flesh tissue is associated with a higher recurrence rate compared to calcified tissue, but corresponds to a complete response if stable over the follow-up (Ghassemi et al., 2013; Schueler et al., 2006).

7.2.8.1. Cryotherapy. Cryotherapy destroys small tumors (usually up to 2 mm high and less than 3 mm diameter) by freezing them at ~80°C, resulting in intracellular ice formation and plasmatic membrane rupture of tumor and endothelial cells, visualized by the presence of petechias at the time of recirculation. The treatment is performed with a cryoprobe directly applied to the outer surface adjacent to the tumor with a triple freeze-thaw technique allowing the iceball to completely cover the tumor. Cryotherapy is ideally used to treat pre-equatorial tumors, but lesions located more posteriorly may be reached by placing the cryoprobe in the sub-Tenon space after opening the conjunctival cul-de-sac. The treatment leaves a chorioretinal scar and has been reported to eradicate up to 90% of the treated lesions (Abramson and Schefer, 2004b). The most frequently observed complication is transient conjunctival chemosis, whereas retinal tears and detachments, vitreous hemorrhage and transient subretinal fluid have been infrequently reported (Anagnoste et al., 2000; Hamel et al., 2000; Shields et al., 1993b).

7.2.8.2. Hyperthermia. Hyperthermia induces a cytotoxic effect on the tumors by heating them up to sub-coagulation temperatures of 45°C–60°C. It is applied using an 810-nm infrared diode laser applied transpupillary on a continuous mode, either by indirect ophthalmoscopy and a 20 Diopter magnifying lens or with an adaptor.
coupled to an operating microscope through a three-mirror lens contact. Compared to the indirect ophthalmoscopy where the spot size is variable, the microscope adaptor allows to set the spot size between 0.3 and 2.0 mm as well as to achieve uniform delivery of the laser on the lesion. In the absence of temperature monitoring during treatment, the laser power setting is entirely empirical and varies from 100 to 1200 mW adapted to the tumor size, location and underlying level of pigmentation. The duration of treatment depends on the observed clinical response (good signs being whitening of the treated lesion and micro-hemorrhage on its surface) and usually varies from 5 to 30 min. Small naive tumors usually respond quickly, whereas white avascular or calcified tumors show no changes due to limited energy transfer. The problems regarding the setting of energy and duration could be overcome in the future by the implementation of real-time temperature determination using opto-acoustic technology (Brinkmann et al., 2012). Ideal for posterior pole tumors, hyperthermia can also be used for peripheral tumors after appropriate indentation of the lesion.

Hyperthermia resolves over 90% of small tumors (≤ 1.5 mm in base diameter) without the need of any other treatment (Abramson and Schefler, 2004a). Indocyanine green (Grobner et al., which has an absorption peak similar to the emission of the diode laser and thought to promote a potentiated laser response in ICG retaining tissue, has been reported to be successful for tumors refractory to thermotherapy after intravenous or intra-arterial chemotheraphy (so called ICG-enhanced thermotherapy) (Francis et al., 2013a). Hyperthermia can also be delivered 1–2 h after intravenous carboplatin, so-called chemotheraphy, to potentiate its cytotoxic effect by virtue of a synergic effect between heat and drug at the time of its plasmatic peak (Murphree et al., 1996; Lumbroso et al., 2002). With the advent of intra-arterial chemotherapy, chemotheraphy is less frequently used.

Complications of transpupillary thermotherapy include iris atrophy with or without concomitant focal lens opacities, tumor seeding by rupture of the internal limiting membrane, retinal fibrosis, rhegmatogenous retinal detachment and vascular occlusion (Gombos et al., 2006; Shields et al., 1999; Tawansy et al., 2006).

7.2.8.3. Photocoagulation. Photocoagulation can be done with a green (532-nm) laser, continuous wave Nd:YAG (1064-nm) or infra-red (810 nm) transpupillary laser (Hamel et al., 2000) and enables tumor destruction indirectly by occluding tumor vessels (532-nm) or directly by generating heat with protein-denaturing temperature in excess of 65 °C (coagulation) within the treatment spot. Complications include retinal fibrosis, traction, and vitreous seeding secondary to rupture of the internal limiting membrane (Gombos et al., 2006).

Transscleral diode laser (810 nm) can be used to destroy small peripheral tumors with a power setting varying between 1000 and 2000 mW and duration of 1–2 s, applied until a whitening of the tumor is observed by indirect ophthalmoscopy. The laser probe has a focus of 1–2 mm of penetration into the eye and is thus only suitable to treat small peripheral tumors up to one disc diameter, but with the advantage to induce a much smaller chorioretinal scar than cryotherapy.

7.2.8.4. Brachytherapy. Brachytherapy is used for solitary, medium-sized tumors (up to 10 mm in diameter and 6 mm in height) and located more than 3 mm from the optic disc or fovea. Most commonly employed isotopes are iodine 125 (I125) or ruthenium 106 (Ru106) and calculations are made for a delivered dose of 40 Gy to the apex of the tumor. Both types of plaque allow a similar tumor control rate of up to 95% of the treated cases at a 5-year follow-up (Schueler et al., 2006; Shields et al., 2001). However, radiation-related complications, including proliferative and non-proliferative retinopathy, radiation-induced maculopathy/papillopathy, subcapsular cataract and neovascular glaucoma are less frequent with the beta-emitting Ru106 because of its limited axial and lateral dosimetry compared to the gamma-emitting I125 (Fluhs et al., 2004).

7.3. Disease and treatment-related complications

The conservative treatment of retinoblastoma can be complicated by various conditions, such as cataract, rhegmatogenous/tractional retinal detachment or secondary neovascularization which, left untreated, may compromise the active tumor-related management or the vision/globe preservation of eyes considered in remission. In the era of intra-arterial and intravitreal chemotherapy, such complications are actually representing up to 50% of the indications to secondary enucleation (Eagle et al., 2011; Pavlidou et al., 2015) compared to less than 5% in the era of systemic chemotherapy (Balaguer et al., 2009; Chantada et al., 2007), and this rate is expected to increase in the future as more eyes with very advanced disease are now offered salvage treatments.

This section deals with the most frequently encountered complications during conservative management of retinoblastoma. Whereas some of them (amblyopia, cataract, retinal detachment and secondary neovascularization) can be offered specific treatments, others, related to drug toxicity and/or drug delivery procedure (chorioretinal complications after intravitreal or intra-arterial chemotherapy) are irreversible, and efforts should be made to prevent them until less toxic therapies become available.

Among the treatable conditions, early amblyopia management is without doubt one of the most important active measures to achieve vision preservation. As there is now sufficient evidence that useful vision can be obtained in the majority of affected eyes, including those with macular involvement, each time visual potential is not given a chance to be realized can be considered a partial treatment failure (see 7.3.1).

Other conditions, such as cataract (see 7.3.2) or retinal detachment (see 7.3.3) may require invasive procedures, considered hazardous in the context of retinoblastoma. However, the introduction of safety-enhanced interventions can overcome the fear of potential tumor spread and should enable intraocular surgery to be considered in selected cases. As a rule, the decision to proceed with surgery should be based on the status of the contralateral eye, the estimated visual potential expected to be achieved after the intervention and, of course, full parental informed consent on the risk and benefits of the procedure.

7.3.1. Amblyopia

The raising concern in preservation of visual function while managing ocular salvage has become noticeable in the increasing incidence of recent publications in which visual acuity is considered as one of the success criteria of surviving eyes after treatment (Chan et al., 2009; Demirci et al., 2005; Hall et al., 1999; Kim et al., 2010b; Manjandavida et al., 2014; Narang et al., 2012; Schefler et al., 2007; Suzuki et al., 2011; Tsimpida et al., 2013; Weiss et al., 1994). Few authors, however, have addressed the importance of active visual reeducation in retinoblastoma eyes (Greenwald and Strauss, 1996; Watts et al., 2002). At Jules-Gonin Eye Hospital, a retrospective study was conducted to compare the visual outcome in 3 groups of children with a minimum of 4 years of age at last visit, all having fovea-involving tumors. The first group (n = 32) included patients with bilateral disease having undergone enucleation of one eye, these considered to be undergoing permanent occlusion. The second group (n = 27) included only unilateral retinoblastoma patients undergoing active patching of the unaffected eye. The third group (n = 13) included patients with unilateral retinoblastoma that failed to comply with amblyopia treatment. The median best corrected visual acuity was 0.7 logMAR, 1.1 logMAR and 1.7 logMAR for groups 1, 2 and 3 respectively. There was a highly significant difference of visual acuity between the monophtalmic group 1 and the treated group 2 (p = 0.001), and also between group 2 and the untreated group 3 (p = 0.016), demonstrating the benefit of active visual reeducation in all cases with foveal involvement. In addition, the significant statistical difference between groups 1 and 2 suggests that a more aggressive amblyopia treatment regimen could even increase the
visual gain (Fig. 24).

The visual deprivation resulting from disease or treatment-related foveal damage or obscuration following hemorrhage, seeding or cataract, can cause not only amblyopia but also strabismus and possibly nystagmus, these in turn having a potential negative impact on quality of life. In a report on the visual outcome in patients with macular involvement treated by chemoreduction and focal therapy, strabismus, mostly exotropia, was noted in 16% of the patients (Schefler et al., 2007).

More recently, in a study on the long-term functional results in patients with group D eyes, 22% had nystagmus, and 60% of bilaterally salvaged patients had strabismus with exotropia being the most common type (Fabian et al., 2017a). By univariate analysis, strabismus and foveal tumor at presentation were found to be significantly associated with strabismus at final follow-up (p = 0.043), while only foveal involvement was found to be significant (p < 0.001) on multivariate analysis (Fabian et al., 2018). It should be stressed that conservation of a misaligned eye can cause as much embarrassment as an artificial eye, especially in teenagers. Strabismus surgery can be envisaged once growth is completed. Similarly, an abnormal head posture due to nystagmus can also benefit from muscle translocation in order to redress the neutralized zone as close as possible to the primary position. Both procedures have been successfully performed in Lausanne (unpublished data).

7.3.2. Cataract

7.3.2.1. Incidence. During the irradiation era, cataract was a frequent treatment-induced complication, developing usually within two years following the treatment in up to 87% of the eyes treated with external beam irradiation (Blach et al., 1996; Fontanesi et al., 1996) and up to 37% of those treated with lens sparing technique or stereotactic conformal radiotherapy (Pica et al., 2011; Sahgal et al., 2006; Scott et al., 1999). Since retinoblastoma is now managed without external irradiation, cataract has become a less frequently encountered problem, but can still complicate brachytherapy (Abouzeid et al., 2008; Murakami et al., 2012; Schueler et al., 2006; Shields et al., 2006b), transpupillary thermotherapy (Shields et al., 1996), intravitreal (Francis et al., 2014; Ji et al., 2016; Munier et al., 2012a; Shields et al., 2016b) and/or intracameral chemotherapy (Munier et al., 2018). Radiation exposure during intra-arterial chemotherapy (from 5.55 to 191 mGy per eye) is below the reported doses for cataract formation (500 mGy) (Thampi et al., 2013).

Fig. 24. Influence of amblyopia treatment in fovea-involving retinoblastoma. Box and whisker plot shows the distribution of final visual acuity in three treatment groups. Group 1: only eye patients (permanent occlusion) Group 2: unilateral disease with active patching of unaffected eye. Group 3: unilateral disease with no patching. The median visual acuity of each group is denoted with a thick horizontal line that bisects the corresponding box-plot. The interquartile range is shown by the waist of the box-plot, when the waists of two box-plots do not overlap, this represents a significant difference, which is denoted with an asterisk (*p < 0.05, ***p < 0.001). The number of patient in each group is given on the top of the graphic.

7.3.2.2. Cataract surgery. The risk of tumor spread in eyes with active retinoblastoma undergoing cataract surgery has since long been recognized (Brooks et al., 1990; Honavar et al., 2001) and requires specific security guidelines. A minimum of 6–12 months disease-free interval before surgery is recommended (Brooks et al., 1990; Honavar et al., 2001; Miller et al., 2005; Osman et al., 2011), whereas eyes where fundus view is lost sooner are usually treated with enucleation (Berry et al., 2018a; Brennan et al., 2015). Providing that specific additional precautions are followed, however, such eyes, in selected cases, can still be managed conservatively.

General recommendations include performing clear cornea small incisions rather than pars plana or scleral tunnel to avoid any sub-Tenonian or sub-conjunctival contamination. The incisions should be secured with sutures at the end of the intervention and intraocular fluids sent for cytopathologic analysis to exclude retinoblastoma cells. Posterior capsulorhexis associated with anterior vitrectomy remains subject of controversy, since it can cause iatrogenic aqueous contamination (Munier et al., 2018). The dilemma of whether to leave the posterior capsule intact or not is particularly crucial in children younger than 5 years of age in whom it is difficult to perform YAG laser capsulotomy, but who at the same time are more likely to present a shorter time interval for treatment-free follow-up and relapses.

Specific recommendations prior surgery for eyes with a disease-free interval of less than 6 months should include an MRI to exclude extraocular disease extension, as well as two intravitreal chemotherapy injections one week apart in order to optimize safety prior the surgery. Cytology-negative results of the paracentesis performed at the time of the intravitreal injection should be confirmed before proceeding with the cataract surgery. Finally, in case of suspicion of active disease, posterior capsulotomy should only be done if the whole procedure is performed under infusion of melphalan at the concentration of 5 μg/ml. Following the above-mentioned recommendations, cataract surgery has been performed in 24 retinoblastoma eyes at Jules-Gonin Eye Hospital over the last 10 years, among which 9 had a disease-free follow-up of less than 6 months. No metastasis nor deaths were observed at a mean follow-up of 4.9 years (range 0.8–8.1 years) (unpublished data).

7.3.3. Rhegmatogenous retinal detachment

Retinal detachment, constitutive of exophytic growing tumor(s), is a frequently observed feature of retinoblastoma at presentation. Such detachments are purely exudative in nature and in the majority of cases
complete resolution is achieved, regardless of the first line therapy administered (Berry et al., 2016b; Rowlands et al., 2018; Shields et al., 1997, 2012). Rhegmatogenous retinal detachments, however, can appear as a treatment-related complication and require appropriate management to allow further tumor control and/or eye preservation.

7.3.3.1. Incidence. Little is known about the rate of rhegmatogenous retinal detachment occurring during the conservative treatment of retinoblastoma. It was found to complicate 6% (5/83) to 11% (5/45) of the eyes treated conservatively during the external beam irradiation era (Bovey et al., 1999; Lim and Robertson, 2000). All those eyes except one treated with plaque brachytherapy only had received EBRT (n = 9), with additional treatment including focal treatment (n = 7) and intravenous chemotherapy (n = 3). In the only report detailing the treatment of the eyes, those treated with external beam radiotherapy alone (0/16, 0%) or focal treatments alone (0/21, 0%) did not develop a retinal detachment (Bovey et al., 1999).

According to the single study focusing on vitreoretinal complications after systemic chemotherapy (Tawansy et al., 2006), the incidence of rhegmatogenous and/or tractional retinal detachment was limited to 1.2% (n = 5/404) and the role of concomitant focal therapy in the context of chemotherapy-related impaired wound healing pointed out (Anagnoste et al., 2000; Tawansy et al., 2006).

More recently, in a series of 75 eyes (68 ICRB group D/E) treated with first line intra-arterial chemotherapy without concomitant focal treatments between 2008 and 2015 (Shields et al., 2017), 6 eyes (8%) developed rhegmatogenous retinal detachment. Extensive endophytic pattern was associated with a significantly higher risk for rhegmatogenous retinal detachment (5/22, 23%) (p = 0.0073), while only 4% (1/24) of the eyes with mixed tumor growth and none of 29 eyes with exophytic tumor growth developed a rhegmatogenous retinal detachment. Similarly, in a series of 74 eyes (69 IIRC group D/E) receiving first line intra-arterial chemotherapy without concomitant focal treatments between 2008 and 2018 in Lausanne (unpublished data), 12 cases (16%) of rhegmatogenous retinal detachment were observed. The incidence of the complication was also found to be significantly higher in endophytic eyes (9/32, 28%) (p = 0.015), compared to eyes with mixed growth (3/24, 12%) or those with exophytic tumors (0/18, 0%). In both series, most cases (80%) of retinal detachment occurred within one month following the first intra-arterial chemotherapy, supporting the hypothesis that the complication develops from atrophic (often invisible) holes due to abrupt tearing forces caused by rapid necrotic tumor regression. Although specific data are missing, the lower frequency of rhegmatogenous retinal detachment observed after intra-venous chemotherapy, suggests a more gradual tumor regression compared to intra-arterial chemotherapy. If true, advanced endophytic tumors may benefit of an initial course of systemic chemotherapy, as done in bridge therapy, before intra-arterial chemotherapy to initiate a less aggressive regression and prevent this complication (see 7.2.1.2.2).

Finally, rhegmatogenous retinal detachment, known to occur in around 0.01% of intravitreal injections in adult eyes (mostly myopic) (Meyer et al., 2011), was only reported following intravitreal chemotherapy in one single case (Smith et al., 2014; Suzuki et al., 2015).

7.3.3.2. Surgical repair. The violation of the state of metastatic grace by invasive surgical procedures, such as vitrectomy, in eyes previously treated for retinoblastoma (Honavar et al., 2001) or with unsuspected retinoblastoma (Kaliki et al., 2018; Shields et al., 2000; Spaulding and Fuhs, 1968) remains a major concern. With this in mind, most authors prefer minimally invasive methods for repair, including non-drainage scleral buckling with retinopexy (Bovey et al., 1999; Mullaney et al., 1997; Yousef et al., 2016) or with post-operative laser photocoagulation of a visible retinal break (Buerk et al., 2006). To avoid high intraocular pressure due to tightening of the band potentially resulting in central retinal artery occlusion, different strategies have been proposed, including intravenous mannitol infusion 1 h before surgery, post-operative acetazolamide (Yousef et al., 2016), or per-operative anterior chamber paracentesis (Mullaney et al., 1997). Some authors recommend at least a 6- to 12-month disease-free interval before considering scleral buckling with external drainage or vitrectomy with cytopathologic evaluation of the intraocular fluids (Baumal et al., 1998; Lim and Robertson, 2000; Madreperla et al., 2000; Tawansy et al., 2006). In Lausanne, in the absence of signs of spontaneous retinal reattachment, an early ab externo approach with scleral buckling with no subretinal drainage is preferred, especially if the detachment occurs in a child's eye with visual potential, as a prolonged observation interval may compromise the functional outcome. An intra-operative anterior chamber paracentesis is performed systematically to allow sufficient tightening of the band. In case of inadvertent scleral perforation, aspiration and cytopathologic analysis of the fluid should be done and the operation field prophylactically immerged with 2 mg of topotecan (1 mg/ml). Following this strategy, complete retinal application was successfully achieved in 92% of the 12 (n = 11/12) above-mentioned eyes (see 7.3.3.1) and partially but with an applied fovea in 1 eye at a mean follow-up of 4.4 years (range 0.5–9.0). All patients are alive with no metastases (unpublished data).

If an intraocular procedure is preferred, tumor reactivation should be ruled out as the cause of the detachment. Pars plana vitrectomy (23 or 25G) with melphalan irrigation (5 μg/ml) with or without silicone oil tamponade is possible (Yarovoy et al., 2015). In the case of retinal detachment occurring concomitantly to proliferative vitreoretinopathy, with or without retinal dehiscence, an intravitreal anti-VEGF injection can be successfully given one week prior to vitrectomy under melphalan infusion combined with retinotomy (Stathopoulos et al., 2018a).

7.3.4. Secondary neovascularization

Secondary neovascularization can develop during the conservative management of retinoblastoma and affect both posterior and anterior segments of the eye. Its occurrence has been associated with retinal ischemia secondary to chronic retinal detachment (Stathopoulos et al., 2018a; Tawansy et al., 2006), focal treatments (Ghassemi and Amoli, 2014; Kase et al., 2008; Scheffer et al., 2007), external irradiation (Archer et al., 1993; Hwang et al., 2017), plaque radiotherapy (Abouezid et al., 2008; Albert, 1987; Francis et al., 2013b; Murakami et al., 2012), intra-arterial chemotherapy (Bianciotto et al., 2012; Stathopoulos et al., 2018a) as well as active tumor progression (Stathopoulos et al., 2018a) or recurrence (Kirali et al., 2017; Stathopoulos et al., 2018a).

The exact incidence of secondary neovascularization prior the era of intra-arterial chemotherapy is unknown. In a retrospective study including all eyes treated with primary or salvage intra-arterial chemotherapy over a 10-year period in Lausanne, 20% (n = 37/182) of the eyes, among which 30% (n = 11/37) had active tumor, developed secondary neovascularization, at a mean interval of 9.6 months since the first injection and after a mean of 3 injections. In that study, the incidence of secondary neovascularization was higher but not significantly after salvage compared to first-line intra-arterial chemotherapy (22%, 25/114 versus 18%, 12/68) (unpublished data), which is similar to the rates of 20% (17/85) after primary and 16% after salvage intra-arterial chemotherapy found by others over the same period of time (Ancona-Lezama et al., 2018). If heavily pre-treated eyes, particularly after external irradiation, are at higher risk to develop neovascular complications after intra-arterial chemotherapy remains to be established.

Neovascular complications in retinoblastoma, especially if concomitant with elevated intraocular pressure or intraocular hemorrhage, are usually followed by enucleation. Aneudotally, cases with no active tumor developing proliferative vitreoretinopathy associated with chronic retinal detachment have been treated with pars plana vitrectomy and membrane peeling (Tawansy et al., 2006; Warden and Mukai, 2006) and choroidoretinal neovascularization with Argon laser.
The use of intravitreal anti-VEGF in the treatment of 35 eyes with secondary retinal and/or iris neovascularization was recently reported (Stathopoulos et al., 2018a). In this study, intravitreal anti-VEGF provided the conditions for further conservative management allowing an overall globe preservation rate of 51% (n = 18/35) at a mean follow-up of 2.4 years. Eyes with no concomitant active tumor had a better globe salvage rate than eyes with active tumor (65% vs 25%) (Stathopoulos et al., 2018a).

7.3.5. Chorioretinal complications

Chorioretinal complications are the most frequently reported intraocular complications of both intravitreal and intra-arterial chemotherapy, leading to the so-called intravitreal melphalan-induced chorioretinopathy and choroidal occlusive vasculopathy respectively. Both are irreversible and can cause permanent visual loss if the macula is involved. Clinical features and risk factors of these adverse effects are discussed below. In addition, measures to reduce/prevent the intravitreal melphalan-induced retinal toxicity after intravitreal melphalan are proposed (see 7.3.5.1.2). Finally, a classification system to grade the as yet poorly described choroidal occlusive vasculopathy according to its extent is presented, in order to facilitate comparisons between studies reporting adverse effects and functional outcomes after intra-arterial chemotherapy in the future (see 7.3.5.2.1).

7.3.5.1. Intravitreal melphalan-induced chorioretinopathy

7.3.5.1.1. Clinical presentation, classification system and incidence. Intravitreal melphalan-induced retinopathy, also referred to as salt and pepper retinopathy, presents as an abrupt, irreversible, chorioretinopathy usually confined to the meridian of the needle entry site. The first visible signs are vascular in nature, resulting in an area of intra-retinal hemorrhages occurring within 2 days of the injection and evolving within one month to a well demarcated retinal atrophy with pigmented changes, associated with choriocapillaritis atrophy of various extents appearing progressively. A classification system of the intravitreal melphalan-induced chorioretinopathy was proposed to grade the retinal toxicity according to the location and extent of the retinal damage (Munier, 2014), with grade 1–3 referring to a pre- and retro-equatorial involvement sparing the macula, and grade 4 and 5 to a...
maculopathy and optic neuropathy respectively (Fig. 25).

Overall, the melphalan-induced chorioretinopathy has been variously reported to occur in 0.8–43% of the cases (Smith et al., 2014). In a retrospective review of all eyes (n = 90) receiving a first intravitreal treatment with melphalan at Jules-Gonin Eye Hospital over the last 10 years, 41% developed the complication, which was classified as grade 1 or 2 in over 90% of the cases. Complications grade 4 and 5 were never observed in the Lausanne cohort (unpublished data) (Table 14).

7.3.5.1.2. Risk factors associated with intravitreal melphalan-induced chorioretinopathy. There are several potential risk factors for retinopathy following intravitreal melphalan, namely vitreous drug concentration, eye pigmentation and localization of drug delivery. These aspects are discussed in detail below.

7.3.5.1.2.1. Intravitreal concentration

The intraocular toxicity of intravitreal melphalan was initially investigated in preclinical studies on the rabbit showing that a vitreous concentration of 5.9 μg/mL, corresponding to a human injected dose of 24 μg, was not causing any electroretinographic (ERG) perturbation (Ueda et al., 1995). This was further confirmed in a study describing the innocuity of melphalan at a vitreous concentration of 5 μg/mL, but a 52% decrease of both a- and b-wave amplitudes at a concentration of 10 μg/mL, with necrosis of the inner nuclear layer and thinning of the outer nuclear layer (Shimoda et al., 2008). In humans, an immediate and permanent 5.8 mV decrease in ERG amplitude per injection of 30 μg was reported (Francis et al., 2014).

According to the above, it became accepted that it was efficient to inject a dose of 20–30 μg melphalan (Munier et al., 2012a). As previously discussed, intravitreal injection of 8 μg melphalan has been reported to be ineffective, whereas 50 μg has been associated with phthisis bulbi (Ghassemi and Shields, 2012). Potential risk factors for the severity of the melphalan-induced chorioretinopathy (grade 3 versus grade 1 and 2) were studied in a series of 90 eyes treated for the first time with intravitreal melphalan in Lausanne (see 7.3.5.1.1), including time to develop the complication since the first injection, total intravitreal injected dose, and mean intravitreal drug concentration reached until the occurrence of the complication. Only intravitreal drug concentration integrating both age-related vitreous volume (Fledelius et al., 2017) was found to be significantly correlated with the severity of the melphalan-induced chorioretinopathy (p = 0.04) (Table 14). To determine the effective intravitreal melphalan concentration, the formula below was used, in which the vitreous was estimated to occupy 80% of the eye volume and the tumors to be ellipsoids:

\[
\text{Effective intravitreal drug concentration} = \frac{\text{Dose}}{(0.18 \times e^{0.15 \times \text{AL}}) \times 0.8 - \sum_{i=1}^{N} \frac{r_i \times h_i}{2} \times \frac{r_i^2 \times h_i}{3}}
\]

For this calculation, the axial lengths (AL, in mm) were estimated according to age or the corrected age in case of prematurity. Height (h), longitudinal (r1) and transverse (r2) radii (in millimeters) of each tumor (n) were measured with ultrasonography (12 MHz). The volume occupied by each tumor was summed up and subtracted from the estimated vitreous volume.

These observations imply that tumor volume and not only age-matched vitreous volume is an important factor to consider when choosing the dose to be injected. Recommended melphalan doses for intravitreal chemotherapy according to age and percentage of the vitreous volume occupied by the tumor(s) are presented in Table 15.

7.3.5.1.2.2. Para-vitreal injections

Para-vitreal injections can lead to a toxic exposition of the retina, despite adequate intraocular melphalan dose. Thus, in eyes with an extensive posterior hyaloid detachment, injection into the retro-hyaloid space may induce a grade 4 or 5 retinopathy (Aziz et al., 2017) (Fig. 25). Similarly, injection into an open Cloquet’s canal with subsequent filling of the virtual bursa premaculairis may induce a grade 5 retinopathy (Fig. 25).

To avoid paravitreal injections, ultrasonography should always be performed prior to the procedure, to inject preferentially in a tumor free quadrant where the posterior hyaloid is not detached. On the other hand, the correct positioning of the needle centrally behind the lens should be done under the operating microscope view, which also allows the visualization of a retro-lental bubble after initial injection of a small volume, indicating a smooth diffusion of the drug into the vitreous cavity. If no bubble can be seen, mis-injection into Cloquet’s canal should be feared, requiring a reposition of the needle elsewhere where a bubble can be observed.

7.3.5.1.2.3. Intraocular pigmentation

Since melphalan has a tropism for the melanin of the retinal pigment epithelium (Süsskind et al., 2016), it would be reasonable to expect that more pigmented eyes may be at greater risk of having the complication. Some authors showed a significant reduction in ERG findings in eyes with brown iris and pigmented fundus compared to those with blue iris and blond fundus (Francis et al., 2017b). In the retrospective analysis of the above-mentioned cohort of patients treated in Lausanne (see 7.3.5.1.1) considering the presence or absence of the toxic retinopathy according to the iris color, the opposite was, however, observed, with patients having blue or green irides (n = 28) presenting significantly more often the retinal complication than patients with brown irides (n = 61) (16/28 57% vs 21/61 34%, p = 0.04). In the eyes presenting intravitreal injection-induced chorioretinopathy, the total dose up to the complication was similar between the light- and dark iris groups (57 μg vs 53 μg), and average effective intravitreal drug concentration was within safety limits in both groups (7.9 μg/mL vs 6.6 μg/mL).

7.3.5.1.2.4. Concomitant drug interactions

It has also been suggested that concomitant intravitreal and intraarterial melphalan may increase the number of adverse effects (Francis et al., 2017b). In the Lausanne cohort however (unpublished data), there were fewer cases developing melphalan-induced

Table 14

<table>
<thead>
<tr>
<th>Classification</th>
<th>Number of eyes</th>
<th>Mean time to retinopathy from the 1st injection (months)</th>
<th>Total intravitreal melphalan dose received at time of retinopathy (μg)</th>
<th>Mean intravitreal melphalan concentration (μg/ml)</th>
<th>Evaluated without considering the intraocular tumor volume</th>
<th>Evaluated considering the intraocular tumor volume</th>
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<td>6.3</td>
<td>6.3</td>
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<td>59.4</td>
<td>6.6</td>
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<td>7.4</td>
</tr>
<tr>
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<td>0.53</td>
<td>30.0</td>
<td>7.5</td>
<td>10.6</td>
<td>10.6</td>
</tr>
</tbody>
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chorioretinopathy in the eyes which received intra-arterial melphalan two days apart from the intravitreal injection compared to those who were treated with intravitreal melphalan alone (19/52 (37%) vs 18/38 (47%), p = 0.39, Fisher test). Moreover, the eyes receiving combined treatment had also fewer cases with grade 2 and 3 intravitreal melphalan-induced chorioretinopathy (10/19 (52%) vs (12/18) 67%, p = 0.51, Fisher test) and a longer time to retinopathy development (3.1 vs 2.1 weeks, p = 0.08, Mann-Whitney U test).

7.3.5.2. Choroidal occlusive vasculopathy. The occurrence of intraocular adverse effects after intra-arterial chemotherapy was first reported in 2011, (Munier et al., 2011), with choroidal occlusive vasculopathy being the most frequently observed (Bianciotto et al., 2012; Bracco et al., 2013; Eagle et al., 2011; Muen et al., 2012; Munier et al., 2011).

7.3.5.2.1. Clinical presentation, classification system and incidence. Clinically, choroidal occlusive vasculopathy, also referred to as choroidal atrophy, choroidal ischemia or choroidal infarction, is an irreversible adverse effect of intra-arterial chemotherapy first recognizable by sectorial or diffuse retinal pigment epithelial alterations evolving towards partial or complete choriocapillary and retinal pigment epithelial atrophy. Like intravitreal melphalan-induced chorioretinopathy, the severity of the choroidal vasculopathy can be graded according to its extent and impact on the visual outcome. A classification based on clinical observations is proposed in Table 16 in the hope that it will help assessing the frequency and severity of this complication across groups (Fig. 27).

Its incidence varies substantially in the literature ranging from 7% (Dalvin et al., 2018) (Bianciotto et al., 2012) to 47% (Muen et al., 2012). This variability may be explained at least in part by a learning curve (Dalvin et al., 2018; Reddy et al., 2017), and/or by the fact that not all grades of the complication are necessarily reported (see Table 16). According to the Lausanne data collected over the last 10 years, this complication was observed in 21 of 560 injection procedures (3.8%) administered to 213 eyes (9.8%), and typically occurred after a mean number of 2.4 injections (range 1–5), distributed in 10% grade 1, 52% grade 3, 10% grade 4, and 28% grade 5.

7.3.5.2.2. Factors associated with choroidal occlusive vasculopathy. Different hypotheses have been proposed to account for the occurrence of choroidal occlusive vasculopathy, including the injection mode and technique of cannulation of the ophthalmic artery, the injected drug toxicity as well as intrinsic predisposing factors. In the literature, there is a decline in the reported vascular events over time, most likely due to a learning curve enabling complications of ophthalmic artery catheterization to be avoided, with the complication occurring in about 7–10% of the injected eyes (Dalvin et al., 2018; FLM unpublished data). However, when isolated choroidal atrophy (i.e. not associated with ophthalmic artery occlusion) and choroidal atrophy resulting from ophthalmic artery occlusion are cumulated, the overall choroidal vasculopathy incidence is between 10 and 15% of injected eyes (Ancona-Lezama et al., 2018; Dalvin et al., 2018).

Intra-arterial chemotherapy with melphalan or carboplatin has been shown in vivo to trigger vascular toxicity through endothelial cell inflammation and leukostasis, in a study without sham injection of saline to test the effect of the ophthalmic catheterization per se (Steinle et al., 2012). This was later tested in another animal model, demonstrating no vascular damage following saline injection (Daniels et al., 2018). The role of the micro-catheter positioning at the ostium versus occlusive engagement into the ophthalmic artery, as well as the continuous versus pulsatile injection mode, have also been debated. The issue of

**Table 15**

<table>
<thead>
<tr>
<th>Tumor Volume</th>
<th>Dose</th>
<th>At 6 months</th>
<th>At 36 months</th>
</tr>
</thead>
<tbody>
<tr>
<td>0%</td>
<td>20µg</td>
<td>6µg/ml</td>
<td>4µg/ml</td>
</tr>
<tr>
<td></td>
<td>30µg</td>
<td>9.5µg/ml</td>
<td>6.5µg/ml</td>
</tr>
<tr>
<td></td>
<td>40µg</td>
<td>13µg/ml</td>
<td>8.2µg/ml</td>
</tr>
<tr>
<td>10%</td>
<td>20µg</td>
<td>7.5µg/ml</td>
<td>5µg/ml</td>
</tr>
<tr>
<td></td>
<td>30µg</td>
<td>11µg/ml</td>
<td>7µg/ml</td>
</tr>
<tr>
<td></td>
<td>40µg</td>
<td>14µg/ml</td>
<td>9µg/ml</td>
</tr>
<tr>
<td>25%</td>
<td>20µg</td>
<td>9µg/ml</td>
<td>5.5µg/ml</td>
</tr>
<tr>
<td></td>
<td>30µg</td>
<td>13µg/ml</td>
<td>8µg/ml</td>
</tr>
<tr>
<td></td>
<td>40µg</td>
<td>17µg/ml</td>
<td>11µg/ml</td>
</tr>
<tr>
<td>50%</td>
<td>12µg</td>
<td>7µg/ml</td>
<td>5µg/ml</td>
</tr>
<tr>
<td></td>
<td>20µg</td>
<td>13µg/ml</td>
<td>8µg/ml</td>
</tr>
<tr>
<td></td>
<td>30µg</td>
<td>19µg/ml</td>
<td>13µg/ml</td>
</tr>
</tbody>
</table>

**Fig. 26.** Intravitreal drug concentration with respect to axial length and injection dose for an eye with completely regressed retinal tumor(s) (regression type IV), or 0% of vitreous volume occupied by tumor(s).
occlusive vs non-occlusive technique of injection appears not to be primarily responsible for the choroidal occlusive complications, at least in the rabbit model (Daniels et al., 2019).

A dose-related ocular toxicity of melphalan was hypothesized by the London group (Reddy et al., 2017), who reported that visual complications may be minimized with age-adjusted melphalan dosages. However, such a dosage effect, at least within the published dose range, could not be demonstrated in a subsequent study from Philadelphia (Dalvin et al., 2018).

The relative insolubility of melphalan is another potential source of toxicity linked to the embolization of micro-crystals up to 20 μm in diameter if not vigorously shaken (Tse et al., 2013). Filtration of melphalan after reconstitution using a 0.22 μm filter has been advised but its use is not widely reported in the intra-arterial chemotherapy literature.

Finally, genetic predispositions to thromboembolism have been reported in association with choroidal occlusive vasculopathy, including sickle cell trait, prothrombin mutation, plasminogen activator inhibitor-1 polymorphism (Abramson et al., 2012b; Francis et al., 2012), as well as methylenetetrahydrofolate reductase (MTHFR) polymorphic alleles (Böhm et al., 2018). In the latter case, 4 patients were recently reported who inhaled nitrous oxide pre-intra-arterial chemotherapy resulting in an increase of homocysteine in the blood (Böhm et al., 2018). The exact role of these inherited thrombophilic traits among the patients with choroidal occlusive vasculopathy is still unknown.

7.4. Evaluation of quality of life

This review is sub-titled: “Alive, with good vision and no co-morbidity”. Even fully achieved, this medical goal needs to be confronted with patients’ everyday life. How survivors themselves, as well as those around them, cope with the disease seems especially important (Hamama-Raz et al., 2012; van Dijk et al., 2009). To date, however, this issue has been only poorly addressed (Moll et al., 2013). Even if most of long-term retinoblastoma survivors and their parents report an overall good quality of life and limited psychosocial problems in comparison to peers (van Dijk et al., 2007a, 2007b; Weintraub et al., 2011; Sheppard et al., 2005), confrontation with the consequences of the disease still last for their entire lives. The perception of the disease as well as the psychosocial functioning and development of an affected child is influenced by the parents’ ability to cope with the stress associated with diagnosis and treatment (Hamama-Raz et al., 2012; van Dijk et al., 2007a, 2007b; Weintraub et al., 2011). In addition, patients not only have to withstand a delay in visuo-motor integration in comparison with their healthy peers (Ross et al., 2001), but also a life-long fear of developing recurrences, second primary malignancies, further loss of vision and passing on the disease to their offspring (van Dijk et al., 2010). When parents had to compare the quality of life of their affected children with healthy siblings, their report was significantly worse regarding the former (Batra et al., 2016a). A study comparing self (child)- versus parental evaluation of health-related quality of life showed that parents reported worse on emotional domains (Batra et al., 2016b).

Over the last few decades, a substantial number of patients, mostly those having received irradiation or experienced restrictions in daily life and bullying at school, needed special educational services and psychological guidance or treatment (Sheppard et al., 2005; van Dijk et al., 2010). Finally, fewer marriages and more divorces have been reported in retinoblastoma survivors treated by enucleation and/or external beam radiotherapy (Byrne et al., 1995).

A multi-disciplinary retinoblastoma team should ideally include a dedicated and fully integrated psychological and spiritual accompanying person. Such a presence benefits all families, regardless of religious, educational or cultural background. Over the last decade, virtually all steps forward were taken by challenging orthodoxy with new administration routes of known drugs subsequently evaluated in non-randomized retrospective studies. In the future, therapeutic breakthroughs should benefit from an increasing contribution of translational research and initiatives supporting rare disease consortium such as the European Retinoblastoma Group (EuRbG), favoring prospective multicentric clinical trials. In addition, the availability of new animal models for preclinical studies is expected to potentially improve treatment options (see 8.1). On the other hand, the advent of “omics” technology

8. Future directions and conclusion

Historically, therapeutic innovation in the conservative management of retinoblastoma was driven independently of randomized clinical trials, often in the absence of preclinical studies, mostly due to the orphan nature of the disease. Over the last decade, virtually all steps forward were taken by challenging orthodoxy with new administration routes of known drugs subsequently evaluated in non-randomized retrospective studies. In the future, therapeutic breakthroughs should benefit from an increasing contribution of translational research and initiatives supporting rare disease consortium such as the European Retinoblastoma Group (EuRbG), favoring prospective multicentric clinical trials. In addition, the availability of new animal models for preclinical studies is expected to potentially improve treatment options (see 8.1). On the other hand, the advent of “omics” technology

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**Table 16**

Clinical classification of the choroidal occlusive vasculopathy after intra-arterial chemotherapy.

<table>
<thead>
<tr>
<th>Grade</th>
<th>Description</th>
</tr>
</thead>
</table>
| Grade 1 | Any sectorial incomplete choroidopathy  
| | a. Sparing the fovea  
| | b. Not sparing the fovea  
| | c. Involvement of the fovea not assessable (due to tumor location or media opacities) |
| Grade 2 | Any sectorial complete choroidopathy  
| | a. Sparing the fovea  
| | b. Not sparing the fovea  
| | c. Involvement of the fovea not assessable (due to tumor location or media opacities) |
| Grade 3 | Any diffuse ( = involving the whole fundus) choroidopathy with complete choroidal atrophy involving less than 50% of the fundus  
| | a. Sparing the fovea  
| | b. Not sparing the fovea  
| | c. Involvement of the fovea not assessable (due to tumor location or media opacities) |
| Grade 4 | Mixed diffuse choroidopathy: fundus with 50% complete and 50% incomplete choroidal atrophy  
| | a. Complete atrophy sparing the fovea  
| | b. Complete atrophy not sparing the fovea  
| | c. Involvement of the fovea not assessable (due to tumor location or media opacities) |
| Grade 5 | Any diffuse choroidopathy with complete choroidal atrophy involving more than 50% of the fundus  
| | a. Complete atrophy sparing the fovea  
| | b. Complete atrophy not sparing the fovea  
| | c. Involvement of the fovea not assessable (due to tumor location or media opacities) |
(Golahchi et al., 2018; Hudson et al., 2018; Kapatai et al., 2013; Kooi et al., 2016b; Naru et al., 2017) is expected to allow not only a better understanding of the mechanisms underlying tumor biology (MYCN vs non-MYCN) and progression (somatic chromosomal copy-number alterations, epigenetic changes) including seeding, metastasis and relapse, but also novel diagnostic and therapeutic applications. Last but not least, this section also proposes to unveil some new potential therapeutic avenues, including gene therapy (see 8.2), cell-of-origin targeted therapy (see 8.3), personalized management (see 8.4) as well as new drugs (see 8.5) or new Galenic formulation of known drugs (see 8.6).

8.1 Animal models

There is a surprising paradoxical discrepancy between the evolutionary conserved RB-E2F pathway in most eukaryotes lineage species, including animals and plants (Cao et al., 2010) and the fact that RB1 loss of function causes preferential retinal oncogenesis as a human-specific condition, with no proven equivalent in the animal kingdom other than anecdotal cases of sporadic unilateral retinoblastoma-like tumors (RB1 status not investigated) in the dog and lama (Syed et al., 1997; Fugaro et al., 2005; Regan et al., 2013). This has imposed alternative models based in xenografts or genetically engineered mouse models (GEMMs) to conduct translational research in the field. Methods to monitor tumor growth and allow accurate evaluation of new treatments evaluation such as small animal imaging systems and diagnostic equipment adapted to the preclinical setting have been developed in parallel (Nemeth et al., 2011).

Intraocular xenografts using Y79 and WERI-Rb1 cell lines established more than 40 years ago from enucleated eyes of patients who received no previous treatment have been widely employed in immunodeficient mice where they can reproduce the inactivation of RB1 and metastasize (Y79) to the central nervous system (CNS) (Chevez-Barrios et al., 2000; Lee et al., 1988; McFall et al., 1977; Reid et al., 1974). Over the last years, the orthotopic transplantation of fresh tumor samples or early stage primary cultures to mice has been prioritized, producing very relevant xenografts with striking anatomic and genetic resemblance to the patient disease (Pascual-Pasto et al., 2016; Xu et al., 2009; Zhang et al., 2012). A few institutions have developed programs to systematize the sampling of viable tumor fragments or vitreous seeding from enucleated eyes at the pathology laboratory (Benavente and Dyer, 2015; Sastre et al., 2009). Serum-free, growth factor-supplemented culture medium is used to expand the small tumor fragments obtained from the enucleated eyes and maintain retinoblastoma cells in culture. This method selects stem-like cells that grow as floating tumor spheres resembling tumor seeding, as opposed to culture medium with bovine serum that promotes cell differentiation (Bond et al., 2013; Ma et al., 2011; Pascual-Pasto et al., 2016).

Alternatively, different genetically engineered mouse models (GEMMs) have been developed through manipulation of the genes in the Rb pathway (Benavente and Dyer, 2015). Inactivation of the Rb gene alone, however, is not sufficient to develop retinoblastoma in mice (Jacks et al., 1992), which need therefore concurrent inactivation of Rb-related genes. For instance, conditional deletion of Rb and p130, through the Pax6a promoter, induces bilateral intraocular tumors with 100% penetrance by one month postnatal, subsequent vitreous and aqueous seeding, as well as CNS and lymph node metastases (MacPherson et al., 2007). Other developed GEMMs include concurrent inactivation of Rb and p107 (Chen et al., 2004; Robanus-Maandag et al., 1998), Rb, p107 and p130 (McEvoy et al., 2011), or Rb and p107 combined with Arf deletion, or amplifications of MDMX and MYCN (Conkrite et al., 2012; McEvoy et al., 2011; Wu et al., 2017). Most of them, however, have the disadvantage of incomplete penetrance and/or long latency, making it difficult to interpret the efficacy of new therapies, or do not reproduce the patient outcome to treatments as compared to xenografts that replicate patient response closely (Brennan et al., 2011).

Finally, various tumor and non-tumor bearing animal models have been used to study the pharmacokinetic of different drugs depending on the intraocular delivery model. Large animals such as primates, pigs and rabbits are preferentially chosen because of their appropriate size, their similar proportion between ocular compartments compared to human eyes and the possibility they allow for repeated or continuous sampling of the ocular fluids and the cannulation of the ophthalmic arteries (Carcaboso et al., 2007, 2010; Daniels et al., 2018; Ditta et al., 2012; Schaiquevich et al., 2012c; Vézina, 2013). Macaques and rabbits have been successfully used to characterize the toxicity of intra-ophthalmic artery and intravitreal melphalan in the retina (Francis et al., 2009; Hudson et al., 2018; Kapatai et al., 2013; Kooi et al., 2016b; Naru et al., 2017).
2014; Shimoda et al., 2008; Wilson et al., 2011).

8.2. Gene therapy

The well-defined genetic origin of retinoblastoma and its confinement within a small, accessible and visible location nicely fit the conditions for a gene therapy approach. Transduced genes may generate therapeutic products to be released locally, or products that target dysfunctional oncogenic pathways within the tumor cells.

The first gene therapy to achieve a phase I clinical trial in retinoblastoma used a replication-deficient adenoviral vector (Adv-TK), carrying the herpes simplex thymidine kinase, a suicide gene activated by ganciclovir. The trial included eight retinoblastoma patients facing imminent enucleation due to active vitreous seeding (Chevez-Barrios et al., 2005; Hurwitz et al., 1999). Repeated intravitreal injections of Adv-TK and concomitant intravenous ganciclovir were proven feasible and safe. Treatment-related vitreal inflammation was controlled with corticosteroids. However, despite initial resolution of active vitreous seeds, all eyes were finally enucleated secondary to retinal tumor progression, likely because adenovirus 5 vectors have limited ability to penetrate, infect and transduce retinal tumors, or alternatively because the expression of the genes transduced by adenoviruses is only short-term. Subsequent studies on this vector found that a modified adenovirus 5 fiber gene achieved better cell infection and protein expression (Mallam et al., 2004). Another gene therapy approach explored local transduction of IFNβ, encoding interferon-β, an antineoplastic molecule with an adeno-associated virus, showing antitumor activity against Y79 xenografts in vivo (Shih et al., 2009).

Other viral vectors for retinoblastoma gene therapy induce an oncolytic effect, such as the adenoviruses H101 (Song et al., 2010) and Ad-TERTK p-E1 (Wang et al., 2013), which have been tested in preclinical studies. Currently, the most advanced oncolytic adenovirus for retinoblastoma is VCN-01, designed to replicate selectively in tumor cells with high abundance of free E2F-1 transcription factor activity due to the dysfunctional RB1 pathway (Pascual-Pasto et al., 2019). The genome of VCN-01 contains one deletion in the E1A gene that precludes viral replication in RB1-functional cells and one insertion of an E2F1 promoter under the E1A gene that favors replication in cells with free E2F-1 (Rodriguez-Garcia et al., 2015). In mice, the intravitreal administration of VCN-01 in orthotopic retinoblastoma xenografts infected and killed tumor cells, improved ocular survival significantly compared to standard of care chemotherapy, and reduced the frequency of CNS metastases (Pascual-Pasto et al., 2019). In immunocompetent rabbits, this treatment induced minor local inflammation and did not replicate in retinas (Pascual-Pasto et al., 2019). Intravitreal VCN-01 is currently undergoing a phase I clinical trial in children with refractory retinoblastoma (NCT03284268) and preliminary results of the 2 first patients treated have been published, showing the feasibility and safety of the procedure and evidence of viral replication markers in tumor cells (Pascual-Pasto et al., 2019).

8.3. Cell-of-origin targeted therapy

Two approaches that target intrinsic cell-of-origin features remain possible: First, if the cell-of-origin signaling circuitry that drives tumorigenesis represents an embryonic developmental stage, it may be possible to target this circuitry without affecting other cell types in retinoblastoma patients. For example, we know that the cone precursor proliferative response depends on intrinsically highly expressed proteins including RYR5, TRB2, MYCN, and MDM2 (Xu et al., 2014). Several preclinical agents are available to attack these troublemakers and could be considered for translational development. Second, it has been suggested that cancer pathways may be epigenetically deregulated as a direct or indirect result of RB1 loss (Zhang et al., 2012). It might thus be useful to target the cone precursors’ epigenetic responses to pRB loss.

8.4. Personalized therapy and liquid biopsy

Unprecedented opportunity for customized therapy is emerging by capitalization on individual liquid biopsy made possible since the introduction of the safety-enhanced intravitreal chemotherapy. Now that aqueous humor can be routinely and repeatedly sampled from one or both eyes in retinoblastoma patients as a non-invasive surrogate of tumor biopsy (Berry et al., 2017c), new insights into tumor biology can be undertaken based on all aqueous humor content, including cell-free DNA, exosomes and proteins.

The proof of principle of this approach has been recently established in a retrospective study, significantly correlating somatic chromosomal copy-number alterations (6p gain) in cell-free DNA (cf-DNA) from aqueous humor and secondary enucleation (Berry et al., 2018b). If clinically validated, this remarkable achievement may influence management, especially in terms of treatment monitoring and/or personalized therapies based on specific tumor profiles and to distinguish MYCN from non-MYCN retinoblastomas. In addition, this technology is opening an avenue to study the chronology of mutations associated with tumor progression.

Another seminal paper (Gerrish et al., 2019) now further expands the applications of cf-DNA analysis in liquid biopsy of retinoblastoma patients by demonstrating that i) somatic RB1 mutations can be successfully detected (even at concentrations < 0.1 ng/μl), including single-nucleotides and copy-number variants, ii) mutant RB1 allele frequency in cf-DNA tightly reflects tumor zygosity allowing the distinction of loss of heterozygosity (LOH) vs non-LOH events, iii) > 90% of cf-DNA in the aqueous humor is of tumoral origin, and iv) aqueous cf-DNA content appears to be proportional to the tumor burden. This outstanding advance was obtained by sequential targeted capture enrichment for the promoter and exonic regions of the RB1 gene, highly heterozygous single nucleotide polymorphisms and non-polymorphic regions across chromosome 13, followed by massively parallel sequencing. This breakthrough is expected to significantly impact the current clinical management of retinoblastoma, characterized by a dramatic reduction of enucleation rates even in advanced cases. In apparent non-hereditary retinoblastoma patients, eye preservation therapy is drying up the only existing source of tumor DNA for somatic mutation detection prior to genomic confirmation and thus restraining the identification of germline carriers (15%) in this population. Now that cf-DNA from the conserved eye is available for somatic mutational screening, accurate genetic counseling and clinical surveillance in the family at large can be re-enforced. Furthermore, the identification of compound heterozygote, homo- or hemizygote somatic hits in the aqueous humor of unilateral retinoblastoma, as well as one or more heterozygote somatic hits in germline patients with multifocal tumors, may provide biological markers for personalized detection of minimally disseminated disease in bone marrow biopsy and lumbar puncture.

8.5. New drugs or combinations of drugs

Among the increasing number of new potential targets against retinoblastoma, different therapeutic strategies can be envisaged targeting either the micro-environment (e.g. angiogenesis or glycolytic inhibitors) (Houston et al., 2013) or specific retinoblastoma pathways (e.g. inhibitors of SIK or MDM2/MDMX-p53 interaction) (Pritchard et al., 2016). Alternatively, new molecules potentiating the anti-tumor activity of known treatment modalities, including chemotherapy, radiotherapy or hyperthermia, may also lead to improved clinical outcome. For example, improvement of chemohyperthermia may result from combined administration of carboplatin and inhibitors of heat shock proteins, which are rapidly produced to buffer stress, including but not restricted to heat damage, and enable recovery in virtue of their anti-apoptotic properties. Interestingly, HSP70 and HSP90 have been identified in retinoblastoma cells (Jiang et al., 2008), justifying further investigations on the role of heat shock protein inhibitors as
potentiating molecules of hyperthermia.

8.6. New galenic formulation of known drugs

One of the limitations of chemohyperthermia is that less than 1% of the intravenous carboplatin reaches the tumor, while the entire body is exposed to toxic adverse effects. An alternative approach using small chemotherapeutic molecules activated by heat only could therefore be beneficial by reducing the dose without impacting efficacy (Fig. 28). Small-molecule thermoresponsive anticancer drugs that synergize with thermotherapy have been reported and validated in an adenocarcinoma-bearing murine model (Clavel et al., 2015).

8.7. Conclusion

Over the last decade, the advent of new techniques of chemotherapy delivery directly into the eye (namely intra-arterial, intravitreal and more recently intracameral injections) radically transformed the conservative treatment of retinoblastoma, allowing not only salvage of the majority of eyes, even those with advanced disease, but also eradication of external beam radiotherapy, as well as significantly reducing the rate of primary and secondary enucleations and the indications for systemic chemotherapy. These major advancements were obtained while maintaining a very low rate of metastatic disease, but our ability to detect minimally disseminated disease, as well as to treat CNS disease, still needs improvement to achieve zero-tolerance regarding metastatic death. Finally, in order to promote a better quality of life from diagnosis throughout the active therapeutic period and over the patients’ whole lifetime, further efforts should aspire to shorten the time to success, contain adverse side effects, improve visual outcome, reduce the incidence of second primary neoplasms, and better support the needs of the survivors and their families.

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Co-authors’ contributions

FLM: Design and redaction of the manuscript, iconography selection of and sketch conception; MBP: 3.6, 7.2; GC: 3.3, 3.7, 7.2; DC: 4; TK: 2, 3.5, 3.6; DL: 5; PM: 3.2.4; ACM: 3.8, 7.4; AM: 8.1, 8.2; AM: 6.1, 6.2, 6.3; PS: 7.1; CB: data analysis; PD: 8.5, 8.6; SH: data management - editing; FP: 7.2.2; YV: 3.5; MCG: 3.2, 7.3.1, 7.3.2; CS: Redaction and harmonization of the whole manuscript.

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