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## Chromosomal aberrations in iris melanomas

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#### Abstract

Background. Uveal melanomas can develop in the choroid, ciliary body and in the iris. In choroidal and ciliary body melanomas specific chromosomal changes correlate with metastatic disease. Iris melanomas have a better prognosis than choroidal melanomas and it would be interesting to know if they share chromosomal changes. In addition, iris melanomas might harbour UV-induced mutations of tumour suppressor genes, such as PTEN and CDKN2A.

Methods. Twenty iris melanomas were analyzed for chromosome 1p, 3, 6, 8, 9p and 10q abnormalities using fluorescence in situ hybridization. These results were correlated to clinical follow-up data using statistical analyses.

Results. (Partial) loss of chromosome 3 was observed in 9 iris melanomas and gain of 8q was present in 7 tumours. Loss of chromosome 9p was demonstrated in 7 tumours but no deletions of the PTEN region on chromosome 10 were found. Three patients died of metastatic disease and one patient developed liver metastases, but is still alive. Univariate analysis indicated a lower disease free survival for patients with diffuse growing melanomas (p=0.01), melanomas that lost a copy of chromosome 3 (p=0.03), or invading the ciliary body (p=0.01). In multivariate analysis none of the correlations were significant.

Conclusion. Loss of chromosome 3 as well as loss of chromosomal region 9p21 (that entails tumour suppressor gene CDKN2A) play a role in iris melanoma. A firm correlation with disease-free survival could not be established, possibly due to the small sample size,

Key words: iris melanoma, FISH, monosomy 3, CDKN2A

## Introduction

Iris melanomas are the least common of uveal melanomas (< 6%) and tend to present at a smaller size than posterior melanomas.<sup>1</sup> They can cause drainage blockage of the anterior chamber angle and secondary elevation of the intraocular pressure. They carry a better prognosis compared to other uveal melanomas with a metastatic rate of 0.5% at 3 and 7% at 10 years, respectively.<sup>2</sup> Melanocytic lesions of the iris are usually managed by a 'wait and see policy' with regular follow-ups to detect possible growth. Biomicroscopy, gonioscopy and measurements with the slit lamp are standard tests. Ancillary tests such as fluorescein angiography and ultrasound biomicroscopy are sometimes needed to assess growth more accurately. Iris melanomas can be treated by local excision, enucleation or radiotherapy. Metastases are more likely to develop in those patients who are older or with diffuse iris melanomas, compared to well-defined nodular ones. Other warning signs are tumour features of iris root location and ciliary body invasion with elevated intraocular pressure and extraocular extension.<sup>3</sup> The differential diagnosis of iris melanoma, cysts and pigment dispersion syndrome.<sup>4</sup>

The favourable outcome of iris melanomas compared to choroidal and ciliary body melanomas is poorly understood. To determine the prognosis on only histological classification proved to be difficult.<sup>5</sup> Therefore, more accurate tests are needed. The different anatomical locations of the intraocular melanomas result in different microenvironments. Aqueous humour and vitreous contain various soluble factors that might influence cell growth.<sup>6</sup> Another differential factor is UV-radiation. Its role in uveal melanoma is inconclusive, but arguably the iris is more exposed to the sun than the choroid. <sup>7-13</sup> While UV-radiation might induce changes in the genome, intrinsic genetic differences may also play a role.<sup>14</sup>

To further elucidate the genetic differences between choroidal and iris melanomas, the following study was performed. Twenty iris melanomas were analyzed for chromosomal aberrations well known in choroidal and ciliary body melanomas, such as loss of

chromosome 3 and gain of chromosome 8q<sup>15</sup>, in addition to common changes in skin melanomas, such as loss of 9p21 and 10q23 (regions that harbour the tumour suppressor genes CDKN2A and PTEN, respectively).<sup>16</sup>

#### Materials and methods

The research methodology followed the tenets of the Declaration of Helsinki. Informed consent was obtained for the use of human tissue from the patients. The research was approved by the local research ethics committee. All cases of iris melanomas were collected from the files from the departments of pathology in Rotterdam (1993-2009) and Leiden (1980 to 2009) and reviewed retrospectively. Cases that originated from the ciliary body were excluded as well as patients who had no available follow-up information. This resulted in 22 iris melanomas available for our study. FISH (Fluorescence in situ Hybridisation) was unsuccessful in 2 cases.

The enucleation and biopsy specimens were fixed in standard 10%, buffered formalin for 24 h. They were embedded in conventional, paraffin wax and sections were cut at 5  $\mu$ m. Sections were stained with Haematoxylin and Eosin (H&E) in order to obtain a histopathological diagnosis. The specimens were reviewed and classified according to the Jakobiec histopathologic classification of iris melanocytic tumours <sup>5</sup>: group 1 melanocytosis, group 2 melanocytoma, group 3 epithelioid cell nevus, group 4 intrastromal spindle cell nevus, group 5 spindle cell nevus with surface plaque, group 6 borderline spindle cell nevus, group 7 spindle cell melanoma, group 8 mixed spindle and epithelioid cell melanoma.

Paraffin-embedded iris melanoma slides were used for FISH (chromosome 1p, 3, 6, 8, 9p and 10q). The pathologist located the tumour and dual colour FISH (digoxygenin and biotin labelled probes) was performed with the following probes: RP11-48E9 (1p36), RP11-522N9 (3p13), Pα3.5 (centromere 3), RP11-64F6 (3q25), RP11-356B3 (6p22),

RP11-787I22 (6q21), RP11-24P4 (8p21), D8Z2 (centromere 8), RP11-88J22 (8q22), P1.063 (9p21) and PAC190D6 (10q23).<sup>17</sup> The concentration for centromeric probes was 5 ng per slide and for BAC probes 50 to 75 ng per slide was used. As a control for aneuploidy a probe on chromosome 5 was used: RP11-1059N10 (5q12). The tissue sections were pretreated by dewaxing with xylene, permeabilizing with sodium thyocyanate, proteolysis and denaturation. After hybridization and washing, slides were counterstained with 4', 6-diamidino-2-phenylindole and mounted in anti-fade solution (Dabco-Vectashield 1:1). The amount of scored nuclei depended on the size of the tumour, at least 50 nuclei and till a maximum of 200 nuclei were scored. Cut-off limits for deletion (>25% of the nuclei with one signal) or amplification (>10% of the nuclei with 3 or more signals) were adapted from own experiments and the available literature.<sup>18</sup>

The primary endpoint for disease free survival (DFS) was the time to development of metastatic disease, whereby death due to other causes was being censored. Statistical analyses were performed with SPSS software, release 15.0. Actuarial probabilities of disease free survival (with an event defined as development of metastatic disease or metastasis-related death) were estimated according to the Kaplan–Meier method. To examine the possibility that other clinical, histopathological or chromosomal variations affected the prognosis, we performed Cox proportional hazard analysis for each confounding variable. An effect was considered significant if the p-value was 0.05 or less. The following factors were evaluated: age, cell type (spindle, mixed or epithelioid cell), Jakobiec classification, growth pattern (nodular, multiple nodules or diffuse), involvement of ciliary body, tumour invasion of the chamber angle, chromosomal aberrations (chromosomes 1, 3, 6p, 6q, 8q, 9p and 10q).

## Results

#### Tumour characteristics

Twenty iris melanomas were included in our study. The specimens were classified according to the Jakobiec histopathologic classification of iris melanocytic tumours <sup>5</sup>: 3 borderline spindle cell nevi, 4 spindle cell melanomas, 6 mixed spindle and epithelioid cell melanomas and 7 epithelioid cell melanomas (Figure 1).

Fifteen melanomas grew in the chamber angle resulting in secondary glaucoma in 5 cases caused by angle closure. Nine iris melanomas that spread into the angle also spread into the ciliary body, but primarily originated from the iris and in all of them epithelioid cells were present (Table 1).

#### Clinical data

The median age of the patients at the time of diagnosis was 62 years. Three patients had a trabeculectomy for glaucoma in their medical history (melanoma # 2, 9 and 14). Surgery or Ruthenium plaque irradiation was chosen to treat the iris melanoma. The median follow-up time was 42 months (range 2-204 months). Three patients died of metastases. One patient had been treated with a trabeculectomy 12 years earlier (histopathological diagnosis at that time: Cogan Reese syndrome) and died of metastatic disease (melanoma # 9). The second patient presented with metastases, located in the bones and submandibular lymph nodes, 3 years after a trabeculectomy (histopathological diagnosis at that time: iris nevus without signs of malignancy) (melanoma #1). The third patient died of liver metastases 3 years after the iris melanoma, treated by Ruthenium plaque irradiation, recurred. This recurrence was treated by enucleation (melanoma #13). Another patient presented with liver metastases 2,3 years after enucleation as the primary treatment of her iris melanoma (melanoma #10).

#### FISH results

All 20 iris melanoma tissue slides could be hybridized with the chromosome 3p and centromere 3 probe. For the other FISH probes it was not always possible to obtain data,

because often the material was too sparse to give a definite answer. See table 2 for the results.

FISH Analysis of paraffin sections demonstrated (partial) loss of chromosome 3 in nine tumours. Seven melanomas with loss of chromosome 3 had concomitantly gained chromosome 8q (see Figure 2). Loss of the chromosomal region 9p21 was present in seven tumours, and six of those had concomitant loss of chromosome 3 and 9p. Five tumours had all three chromosome aberrations (loss of chromosome 3, 9p and gain of 8q) mentioned above. Chromosome 10q was normal in all cases.

The tumours of the patients who died of metastases revealed loss of chromosome 3. Two of them also lost 9p21, and two melanomas had gained 8q. Two of the progressively growing melanomas were composed of mixed spindle and epithelioid cells and one was composed of epithelioid cells only. All three grew into the angle of the chamber and into the ciliary body.

Three melanomas in our group were spindle cell nevi with borderline malignancy (Jakobiec group 6). Interestingly, two of them did not have any chromosomal aberrations, but the third one did lose one copy of chromosome 3q and 9p simultaneously with gain of chromosome 8q.

## Statistical analysis

Univariate analysis was performed for all parameters, showing a lower disease free survival for patients with diffuse growing melanomas (p=0.01), melanomas that lost a copy of chromosome 3 (p=0.03) or invading the ciliary body (p=0.01). The other chromosomal aberrations did not show significant correlations. In a multivariate Cox proportional hazard analysis including growth pattern, ciliary body involvement and chromosome 3, they all lost significance.

#### Discussion

Our study reports on chromosomal abnormalities in the largest group of iris melanomas described thus far. We demonstrated that in iris melanomas (partial) loss of chromosome 3 tends to occur often, in 9/20 tumours (45%), but only correlates with progressive disease in a univariate analysis. The frequency of chromosome 3 loss is almost similar to choroidal melanomas, in which case it strongly associates with patients at risk of metastatic disease.<sup>19</sup>

Similar to common changes in skin melanomas, such as loss of 9p21 and 10q23, 7/ 20 iris melanomas (35%) revealed loss of chromosome 9p. UV-radiation induces mutations in the CDKN2A gene that encodes for the p16 protein that acts as a negative regulator of the cell cycle and is located on chromosome 9p21.<sup>23</sup> Alterations of chromosome 9 occur frequently in cutaneous melanomas and relate to progressive disease.<sup>24-26</sup>. Conversely, CDKN2A mutations are rarely observed in UM.<sup>20, 21</sup> Our results indicate a difference between iris and posterior melanomas, because loss of 9p is found in 7/20 iris melanomas.

Yet, we could not demonstrate aberrations of chromosome 10q in iris melanomas. Previously, it has been reported that the tumour suppressor gene PTEN (located on chromosome 10q) is not involved in uveal melanomas, whereas mutations of PTEN are present in up to 40% of cutaneous melanomas.<sup>20-22</sup> The same holds true for N-ras mutations. Mutations of the N-ras gene, located on chromosome 1p13, correlate with sunlight exposure and occur frequently in cutaneous melanoma arising in sun exposed areas.<sup>7</sup> Nevertheless, N-ras mutations are usually absent in uveal melanomas, further questioning the role of sunlight in the pathogenesis of uveal melanoma.<sup>8, 15</sup> In 8/20 iris melanomas the ciliary body was invaded by tumour cells, and four of these tumours spread systemically. Ciliary body involvement was only found to be a significant prognostic parameter in univariate analysis.

Little is known about somatic mutations and chromosomal aberrations in iris melanomas. To date only 8 iris melanomas have been cytogenetically analyzed and described worldwide. The first karyogram of an iris melanoma revealed extra copies of chromosomes 2, 7 and 18, and loss of chromosome 15. Additionally, rearrangements on chromosomes 8q, 4q, 12p and marker chromosomes were observed.<sup>27</sup> Sisley et al. karyotyped 3 iris melanomas and did not find specific chromosomal changes common to all tumours. Clonal abnormalities of chromosomes 3, 5, 6, 7, 8, 9, 12, 15, 17, 18, 19, and Y were found, and in one case a large number of marker chromosomes were observed.<sup>14</sup> They concluded that iris melanomas experience relatively high levels of chromosomal alterations compared to choroidal melanomas, which is perhaps reminiscent of cutaneous melanoma. The third study included 4 iris among other uveal melanomas that were analyzed by comparative genetic hybridization.<sup>28</sup> Gains of chromosome 6p and 8q were identified twice and gains of chromosome 8p and 13q were identified once. One melanoma was completely diploid. These three studies did not identify critical chromosomal regions and, furthermore, the results were not compared to disease free survival.

In contrast, we observed involvement of chromosome 9p in seven tumours and loss of chromosome 3 in nine iris melanomas. These results suggest that the tumour suppressor gene CDKN2A, located on chromosome 9p, might play a role in iris melanomas, and requires analyses of the gene itself. The role of chromosome 3 loss in iris melanomas needs further investigation. The prognostic significance is not as strong as in choroidal melanomas, but possibly this is the result of the small sample size. The differences between choroidal and iris melanomas suggest that environmental influences such as UV-radiation, aqueous and vitreous humour may contribute to the metastatic potential of the tumour.

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## Table 1.

Table 1. Tumour characteristics										
Iris melanoma no#	Age years	Jakobiec group	Growth pattern	Ciliary body involved	Cell type	Angle invasion	Survival	Metastases		
1	53	9	diffuse	yes	epithelioid	yes	48	yes †		
2	63	9	diffuse	no	epithelioid	yes	112	no		
3	41	9	multiple	yes	epithelioid	yes	20	no		
4	55	9	diffuse	yes	epithelioid	yes	16	no		
5	77	6	multiple	no	spindle	yes	62	no		
6	76	6	nodular	no	spindle	no	10	no		
7	71	6	nodular	no	spindle	no	2	no		
8	42	9	multiple	no	epithelioid	yes	144	no		
9	52	8	diffuse	yes	mixed	yes	37	yes †		
10	59	8	diffuse	yes	mixed	yes	29	yes		
11	71	8	diffuse	no	mixed	yes	23	no		
12	46	7	nodular	no	spindle no		96	no		
13	49	8	nodular	no	mixed	ixed no		no		
14	62	7	multiple	no	spindle	yes	62	no		
15	69	9	diffuse	no	epithelioid	yes	5	no		
16	66	8	diffuse	yes	mixed	yes	35	yes †		
17	56	8	multiple	yes	mixed	yes	86	no		
18	69	7	nodular	no	spindle	no	24	no		
19	83	9	multiple	yes	epithelioid	yes	157	no		
20	67	7	nodular	no	spindle	no	204	no		

† Death due to metastases

# Table 2.

Table 2. FISH results											
Iris melanoma	Copy number per chromosomal region										
no#	1p36	3p13	cen3	3q25	6p22	6q21	8p21	cen8	8q22	9p21	10q23
1	1	1	1	1	2	2	2	3	3	1	2
2	2	1	1	1	2	1	3	3	3	1	2
3	2	2	2	2	2	2	2	2	2	2	2
4	2	2	2	2	2	2	2	3	2	1	2
5	2	2	2	1	2	2	2	3	3	1	2
6	2	2	2	2	2	2	2	2	2	2	2
7	2	2	2	2	2	2	2	2	2	2	2
8	1	2	2	2	2	2	2	2	2	2	2
9	2	1	1	1	2	2	2	3	2	1	2
10	2	1	1	1	2	2	1	3	3	2	2
11	2	1	1	1	2	2	4	3	3	1	2
12	2	2	2	2	2	2	2	2	2	2	2
13	n/a	2	2	n/a	n/a	n/a	n/a	3	n/a	n/a	n/a
14	1	1	1	n/a	n/a	n/a	n/a	n/a	3	1	2
15	1	1	1	1	2	2	1	3	3	2	2
16	1	1	1	n/a	2	1	n/a	n/a	3	2	2
17	n/a	2	2	n/a	n/a	n/a	n/a	n/a	n/a	2	2
18	2	2	2	2	2	2	2	2	2	2	2
19	1	1	1	n/a	n/a	n/a	n/a	n/a	3	2	2
20	2	2	2	n/a	n/a	n/a	n/a	n/a	2	2	2
Total aberrations	6/18	9/20	9/20	6/14	0/15	2/15	4/14	9/15	9/18	7/19	0/19

# **Figure legends**

**Figure 1.** Histology (H&E stain) of A and C) no.12, shows a spindle cell iris melanoma, and B and D) no. 1, shows a epithelioid cell iris melanoma with invasion of the angle chamber .



**Figure 2.** FISH analysis of iris melanoma no.1, EOM 149, demonstrates 2 signals for the biotin probe (red signal) on chromosome 10q and 1 signal for the digoxenin probe (green signal) on chromosome 9p (A), and 1 signal for both bio and dig probes on chromosome 3 (B).





