The genetics of uveal melanoma: current insights

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Abstract: Uveal melanoma (UM) is the most common malignant eye tumor in adults affecting ~7,000 individuals per year worldwide. UM is a rare subtype of melanoma with distinct clinical and molecular features as compared to other melanoma subtypes. UMs lack the most typical cutaneous melanoma-associated mutations (BRAF, NRAS, and NF1) and are instead characterized by a different set of genes with oncogenic or loss-of-function mutations. By next-generation sequencing efforts on UM tumors, several driver genes have been detected. The most frequent ones are BAP1, EIF1AX, GNA11, GNAQ, and SF3B1. In many cases, mutations in these genes appear in a mutually exclusive manner, have different risk of metastasis, and are consequently of prognostic importance. The majority of UM cases are sporadic but a few percentage of the cases occurs in families with an inherited predisposition for this malignancy. In recent years, germline mutations in the BAP1 gene have been found to segregate in an autosomal dominant pattern with numerous different cancer types including UM in cancer-prone families. This cancer syndrome has been denoted as the tumor predisposition syndrome.

Keywords: uveal melanoma, driver genes, oncogenes, tumor suppressor genes, familial cancer

Introduction

Uveal melanoma (UM) is the most common neoplasms of eye that develop in adults displaying a high propensity for metastasis. It is a rare subtype of melanoma, representing ~5% of all melanoma tumors. UM can appear in the choroid, ciliary body, or iris of the eye and is by far the most common ocular tumor in adults. Ocular melanomas can also rarely arise in melanocytes in the conjunctiva – melanoma of the conjunctiva accounts for ~2%–3% of all eye neoplasms. The incidence rate of UM ranges from 0.2 to 0.3 per million individuals in African/Asian populations to up to 6 per million individuals in white populations. The average age of diagnosis is ~60 years and it affects both sexes equally or slightly more frequently males as per some reports. It is more common among light-skinned individuals, but it can affect individuals from any ethnicity. Cutaneous melanoma (CM) and UM share some risk factors such as fair skin color, blue eyes, red/blond hair, and freckling/many nevi. Whereas the incidence of CM has been rising in many Caucasian populations, the incidence of UM has been stable over the years. Also in contrast to CM, the impact of ultraviolet (UV) light exposure is less clear for UM. UM is molecularly diverse from CM and shows a different pattern of driver mutations. Compared to CM that shows one of the highest mutational-load among different cancer types, UM displays a low mutational burden. It has, in many cases, a poor prognosis since about half of all patients develop...
metastatic disease, predominantly in the liver. By using gene expression profile (GEP) classification, UM can be stratified into two distinct molecular classes with a significant difference in prognosis. Class 1 tumors can be further divided into two subgroups (class 1A and 1B) and has in general a good prognosis and low metastatic risk, whereas class 2 tumors have high metastatic risk and thereby a worse prognosis. The risk of metastasis has been determined to be 2% for class 1A tumors, 21% for 1B tumors, and finally 72% for class 2. The different molecular classes are also associated with mutations in different UM driver genes. The most frequently mutated genes that are considered to be drivers in UM development and progression are BAP1, EIF1AX, GNA11, GNAQ, and SF3B1 (Table 1).

Some UM cases occur in families with an inherited predisposition for UM. The only high-penetration susceptibility gene for familial UM identified so far is BRCA1-associated protein 1 (BAP1). The frequency of predicted pathogenic germline BAP1 mutations observed in different cohorts of UM patients ranges from 1.6% to 3%, with mutations predominantly found in patients with a family history of UM. The cancer risk mediated by germline BAP1 mutations is inherited in an autosomal dominant pattern with incomplete penetrance. Cancer types associated with BAP1 germline mutations include UM, CM, mesothelioma, meningioma, renal cell cancer, basal cell carcinoma and melanocytic BAP1-mutated atypical intradermal tumors, and possibly additional cancer types. While carrying a germline mutation in the CDKN2A gene is the strongest known inherited risk factor for CM, such mutations do not seem to increase the risk of UM.

Somatic alterations
Risk of metastatic disease and UM-associated survival is strongly correlated with the molecular subtype. Poor prognosis and high risk of metastatic disease are often accompanied with loss of chromosome 3 in the tumor, while tumors with intact chromosome 3 correlates with good prognosis and rarely leads to disseminated disease. The metastatic rate

<table>
<thead>
<tr>
<th>Mutated gene</th>
<th>Chr</th>
<th>Gene function</th>
<th>Frequency</th>
<th>Characterized by</th>
<th>Type of mutation(s)</th>
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<tr>
<td>BAP1</td>
<td>3p21</td>
<td>Deubiquitinating hydrolase involved in tumor suppressor activity, DNA damage response, and proliferation</td>
<td>18%–45%</td>
<td>Almost mutually exclusive with SF3B1 and EIF1AX mutations. Associated with class 2 GEP tumors, monosomy 3, metastasis, older patients, and poor prognosis</td>
<td>Inactivating mutations. Often truncating. No hotspot mutations</td>
</tr>
<tr>
<td>GNAQ</td>
<td>9q21</td>
<td>Mediating signaling between G-protein-coupled receptors and downstream effectors and upregulating MAPK pathway</td>
<td>28%–50%</td>
<td>Mutually exclusive with GNA11 mutations. Considered as an early event. No correlation with prognosis</td>
<td>Oncogenic mutations at codons Glu209 and Arg183</td>
</tr>
<tr>
<td>GNA11</td>
<td>19p13</td>
<td>Mediating signaling between G-protein-coupled receptors and downstream effectors and upregulating MAPK pathway</td>
<td>32%–50%</td>
<td>Mutually exclusive with GNAQ mutations. Considered as an early event. No correlation with prognosis</td>
<td>Oncogenic mutations at codons Glu209 and Arg183</td>
</tr>
<tr>
<td>EIF1AX</td>
<td>Xp22</td>
<td>Involved in eukaryotic translation initiation</td>
<td>14%–21%</td>
<td>Almost mutually exclusive with BAP1 and EIF1AX mutations. Associated with disomy 3, class 1A GEP tumors, and good prognosis</td>
<td>Heterozygous mutations mainly in exons 1 and 2</td>
</tr>
<tr>
<td>SF3B1</td>
<td>2q33</td>
<td>Essential for pre-mRNA splicing</td>
<td>10%–24%</td>
<td>Almost mutually exclusive with BAP1 and EIF1AX mutations. Associated with disomy 3, younger patient age, and development of late metastasis</td>
<td>Heterozygous mutations. Hot spot mutation at codon Arg625</td>
</tr>
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Abbreviations: Chr, chromosome; GEP, gene expression profile.
for tumors with partial loss of chromosome 3 has shown a great variation depending on the study (ranging from 0% to 48%). However, later studies indicate that partial monosity of chromosome 3 often associates with a good prognosis. Other frequent chromosomal aberrations in UM include gain of chromosome 8q which, similar to the loss of chromosome 3, has been detected in about one-third of the tumors, often in the same tumor. This abnormality is usually associated with better patient survival, possibly because it rarely occurs in tumors with monosity of chromosome 3. Inactivation of CDKN2A may be part of UM pathogenesis, either through methylation of the CDKN2A promoter region or through loss of chromosome 9p or a smaller region surrounding the 9p21, harboring the CDKN2A locus. Both promoter methylation and chromosomal loss affect up to one-third of the tumors each.

Other important pathways often altered in UM, as in many cancer types, are the retinoblastoma (Rb) and p53 pathways. Mutations in the genes encoding for these proteins, RB1 and TP53, are infrequent in UM tumors suggesting other ways of inactivation. Cyclin D1 overexpression or CDKN2A promoter methylation are two plausible explanations for hyperphosphorylation and inactivation of the Rb-protein, while inactivation of p53 may be caused by MDM2 overexpression. Also, constitutively activation of the PI3K/AKT pathway and inactivation of the tumor suppressor PTEN (mainly by LOH of the PTEN locus) are common events in UM tumors.

However, overall the extent of genomic instability and chromosomal aberrations is relatively low in UM tumors compared to many other cancer types such as in CM. Also the mutational load in UM tumors is low, and the mean mutation rate of UM tumors has been determined to be around 0.5 per Mb sequence, both concerning genomic and protein coding regions. In UM tumors, several frequent driver mutations have been described, none of them being described as key drivers in other melanoma subtypes. The most commonly mutated genes are BAP1, EIF1AX, GNA11, GNAQ, and SF3BI. In addition, there are numerous other genes with rare mutations. The list of rarely mutated genes will most probably increase with time due to ongoing and future sequencing studies. BAP1, EIF1AX, and SF3BI often occur in a mutually exclusive manner as do GNA11 and GNAQ. Some of these driver mutations have also been shown to be of importance for the prognosis since they mediate a variable risk of metastatic disease. BAP1 is associated with monosity 3, poor prognosis, and class 2 GEP tumors, while EIF1AX is associated with class 1 GEP tumors and good prognosis. SF3BI has been associated with younger patient age and good prognosis.

**BRCA1-associated protein 1**

Loss of chromosome 3 was for a long time the best predictor for metastatic disease in UM patients. Later, the identification of different GEPs, which led to the development of the GEP classification, has improved the prognostic accuracy. The class 2 tumors that are aggressive with high metastatic potential were found to be accompanied by loss of chromosome 3. Using next-generation sequencing, it was discovered that a vast majority of the class 2 tumors carried a mutation in the BAP1 gene, mapped to chromosome 3p21.1, while very few of the class 1 tumors harbored a mutation in this gene. Thus, inactivating hemizygous mutations of BAP1 leads to protein inactivation and loss of BAP1 expression. This implicate BAP1 to function as a tumor suppressor gene, with loss of one copy of chromosome 3 and mutation in the other allele, fulfilling the Knudsen two hits hypothesis definition of a tumor suppressor gene. Indeed, BAP1 has previously been shown to display tumor suppressor capacity by binding to the BRCA1 protein and thereby enhancing BRCA1-mediated tumor suppression. BAP1 mutations strongly correlate with metastatic disease in UM; over 80% of metastasizing UM has been found to carry a mutation in this gene. The frequency of BAP1 mutations in primary UM has been estimated to be approximately 30%–40%. Most of the BAP1 mutations are truncating variants or missense variants affecting the ubiquitin carboxyl-terminal hydrolase domain. In some cases, BAP1 is not altered by a sequence mutation but by hemizygous deletion of one or more exons. Such alterations may be missed by traditional Sanger sequencing because of the presence of normal DNA in the sample. Thus, in some cases, immunohistochemistry (IHC) might be a better choice of detection, if tissue samples are available. Loss of BAP1 expression using IHC has strongly been correlated with risk of metastasis, BAP1 mutation status, and loss of chromosome 3 and has, therefore, been proposed as a valid prognostic test. BAP1 is also frequently mutated in other tumor types, including cholangiocarcinoma, renal cell carcinoma, mesothelioma, and bladder cancer (www.cbioportal.org). Several of these cancer types are part of the hereditary cancer syndrome known as tumor predisposition syndrome...
that is characterized by germline mutations of \( BAP1 \) in patients belonging to cancer-prone families (discussed in detail in section Inherited susceptibility). \( BAP1 \) encodes a deubiquitinating hydrolase with multiple cellular functions, except tumor suppressor activity, such as regulation of chromatin dynamics, DNA damage response, cell cycle regulation, and cell growth. For example, \( BAP1 \) is involved in the polycomb multiprotein repressor complex that is critical for transcriptional silencing of target genes by removing ubiquitin molecules from histone H2A. As a consequence of this functional loss, an accumulation of monoubiquitinated histone H2A has been revealed, which in turn was found to cause a more dedifferentiated phenotype.\(^{45} \) \( BAP1 \) seems to be involved in other important cellular functions as well, for example, in cell proliferation by deubiquitinating the cell cycle regulator host factor 1.

**Eukaryotic translation initiation factor 1A, X-linked**

\( EIF1AX \), located at chromosome Xp22, was identified as a UM driver gene by whole-exome sequencing.\(^{15} \) Approximately 14%–20% of all UM carries a mutation in this gene, with most mutations found in exons 1 and 2.\(^{15,39,42} \) \( EIF1AX \) mutations usually occur in nonmetastatic cases, are associated with class 1 GEP tumors and good prognosis, and are inversely associated with metastasis.\(^{40,46} \) \( EIF1AX \) mutations are usually mutually exclusive with \( BAP1 \) mutations and to a large extent also to \( SF3B1 \) mutations. As expected most \( EIF1AX \) mutations are identified in tumors with disomy 3 (48%) and rarely occur in monosomy 3 tumors (3%).\(^{15} \) In contrast to, for example, \( BAP1 \) mutations, which mainly are truncating and loss-of-function variants, the majority of the \( EIF1AX \) mutations are heterozygous nonsynonymous variants, or in some cases splicing variants, leading to deletions of one or two amino acids. Thus, in most cases, the core protein remains unchanged. \( EIF1AX \), located on the X-chromosome, encodes the eukaryotic translation initiation factor 1A (eIF1A). This factor is essential in the initiation phase of translation of eukaryotic cells by the transfer of methionyl initiator tRNA to the small (40S) ribosomal unit.\(^{47} \) This stabilizes the formation of the ribosome around the AUG start codon, which enables translation. \( EIF1AX \) mutations are usually seen as heterozygous mutations in the tumor-DNA, suggesting that \( EIF1AX \) serves as a dominant acting oncogene. However, it has been reported that UM tumors carrying an \( EIF1AX \) mutation only express the mutant allele, which indicates that \( EIF1AX \) also may function in a recessive manner.\(^{15} \) Mutations in this gene have also been described in other cancers such as thyroid and ovarian cancers and in the rare melanocytic neoplasm primary leptomeningeal melanocytic neoplasms (LMNs). LMNs are also prevalent for mutations in \( GNAQ, GNA11 \) and \( SF3B1.\(^{58-30} \)

**Guanine nucleotide-binding protein subunit alpha-Qan
dguanine nucleotide-binding protein subunit alpha-11**

\( GNAQ \) encodes the alpha subunit (G\( \alpha q \)) and \( GNA11 \) the alpha subunit 11 (G\( \alpha 11 \)), both being guanine nucleotide-binding proteins belonging to the heterotrimeric protein family, which are of importance in transmembrane signaling systems. The alpha subunits serve as a switch between the G-proteins active state – when bound to guanosine triphosphate (GTP) – and the inactive state – when GTP is hydrolyzed to guanosine diphosphate.\(^{51,52} \) Activating mutations in \( GNAQ/GNA11 \) were the first described driver mutations in UM. \( GNAQ \) and \( GNA11 \) mutations occur in a mutual exclusive pattern and are exclusively found in codon 209 and in some cases in codon 183. Mutations at these positions lead to a constitutive activation of the G\( \alpha q \) and G\( \alpha 11 \) subunits by abolishing their intrinsic GTPase activity, thereby preventing the return to an inactive state. In total, ~85% of all UM carry a mutation in either of these genes. Both \( GNAQ \) and \( GNA11 \) have been found to upregulate the MAP kinase pathway when constitutively activated in a similar fashion as \( BRAF \) and \( NRAS \) mutations. In CM activating mutations in \( BRAF \) are a very common event, whereas UM rarely carries any mutation in \( BRAF \).\(^{53-55} \) The activation of the MAPK pathway in the absence of \( BRAF/NRAS \) mutations in UM was at first unforeseen until the identification of \( GNAQ \) and later \( GNA11 \) mutations that had the same effect as the \( V600EBRAF \) mutation. Interestingly, \( BRAF \) mutations have been seen in up to nearly half of all iris melanomas.\(^{16} \) This could be explained by the iris being more anterior and therefore more exposed to UV radiations than the ciliary body or the choroid. Cell lines with a \( GNAQ \) Q209L mutation have also been found to be highly sensitive to mitogen-activated protein kinase (MEK) inhibition.\(^{13} \) Mutations in these genes have not been associated with the two different molecular classes of UM tumors. In addition, \( GNAQ/GNA11 \) mutations have not been reported to be of prognostic value and they occur at similar frequencies in metastatic and nonmetastatic lesions. Furthermore, they have not been linked to patient outcomes. Taken all this data into consideration is supportive of \( GNAQ/GNA11 \) being early events.\(^{12} \) The hotspot mutations in \( GNAQ \) or \( GNA11 \) are also commonly found in benign nevi such as
blue nevi.14,57 Actually, GNAQ Q209 was most frequently found in blue nevi, observed in 55% of the lesions, whereas 45% of the primary UM and 22% of the metastatic UM, respectively, carry this mutation.14 Inverse relationship was seen for the GNA11 Q209 mutation where metastatic lesions showed the highest number (56%) followed by primary UM tumors (32%) and lastly blue nevi (6%).14 Mutations affecting codon R183 are less frequent, present in 2% of the blue nevi and 5% of primary UM tumors, GNAQ and GNA11 mutations combined.14

Splicing factor 3b subunit 1
SF3B1 located at chromosome 2 is another driver gene identified by whole-exome sequencing of UM tumors. SF3B1 is essential in pre-mRNA splicing by encoding the unit of the splicing factor 3b protein complex that is a critical part of both major (U2-like) and minor (U12-like) spliceosomes.58 The spliceosomes are part of the splicing machinery, which bind to the intron near the branchpoint.59,60 SF3B1 has recently also been designated as a factor involved in DNA-damage repair.61 Missense mutations in specific regions of the SF3B1 gene have been found to alter the splicing of many target genes.62,63 These mutations predominantly alter codon Arg625 in exon 14 of the SF3B1 gene and have been identified in UM with a reported mutation rate between 10% and 21%.11,15,64 Some studies report an association between SF3B1 mutations and good prognosis, lower age at diagnosis (a favorable prognostic factor), and tumors with disomy 3.11,64 However, in a study with longer follow-up time, tumors with disomy 3 and a SF3B1 mutation showed significant worse prognosis and development of late metastasis compared to wild-type tumors. In patients with a SF3B1 mutation, most metastasis occurred more than 5 years after diagnosis (median 8.2 years, range 23–145 months).55

In a study by Martin et al,15 29% of the tumors with disomy 3 carried a heterozygous mutation in SF3B1 compared to only 3% in tumors with monosomy 3. Furthermore, in tumors with partial monosomy 3, preferentially with loss of 3q and retention of 3p, 54% were found to carry Arg625 mutation in SF3B1.15 In hematological and lymphoid malignancies, mutational hotspots have been detected in specific codons, for example, codon 700, coding for the HEAT repeats (HD) 4–9.62,66,67 Resequencing of the exons encoding for this region in UM tumors revealed a mutation rate of 15% in UM tumors (n=66).15

SF3B1 mutations often occur in tumors that express the oncogene PRAME.68 Expression of PRAME has been found to be associated with class 1 tumors with an intermediate risk of metastasis, suggesting that there is a risk class of tumors lying between the high-risk tumors characterized by BAP1 mutations and low-risk tumors frequently harboring EIF1AX mutations.

Other rare mutations/alterations in UM tumors
By whole-genome and whole exome sequencing of UM tumors, a recurrent gain-of-function mutation in the phospholipase C, beta 4 (PLCB4), gene was identified.60 This was the only gene with a recurrent mutation (two out of 28 samples) that was found above the known driver genes in UM (BAP1, EIF1AX, GNA11, GNAQ, and SF3B1). The mutation (c.G1888T, p.D630Y) lies in the Y-domain of the highly conserved catalytic core of PLCB4 and was predicted to be deleterious/probably damaging using the prediction tools SIFT and PolyPhen. Interestingly, the PLCB4 protein is a downstream target of GNA11/GNAQ and the p.D630Y PLCB4 mutation was mutually exclusive with mutations in GNA11 and GNAQ.

In CM, recurrent mutations in the core promoter of the telomerase reverse transcriptase (TERT) gene are common. TERT is part of the telomerase enzyme and maintains the telomere ends by adding the telomere repeat TTAGGG. Deregulation of telomerase and aberrant expression of TERT have been found in several different cancer forms such as thyroid and bladder cancers (www.cbioportal.org). Approximately 70% of CM tumors have been reported to carry any of the two mutual exclusive recurrent mutations in the TERT promoter, both being consistent with the typical UV-damage signature. These mutations affect the expression levels of TERT by creating a novel binding site for the transcription factor E-twenty-six.69,70 A germline TERT promoter mutation with the same functional effect as the described somatic mutations has been found to segregate with high penetrance in two large melanoma-prone families in two separate studies.69,71 In UM tumors, TERT promoter mutations are very rare.72 In a study by Dono et al,42 one out of 50 patients carried one of the previously described TERT promoter mutations.42 Here, the promoter mutation was observed in combination with mutations in GNA11 and EIF1AX as well as two normal copies of chromosome 3. TERT promoter mutation thus seems to be infrequent in UM; however, an elevated level of TERT expression has been observed in a subset of UM tumors with wild-type TERT promoters42 suggesting that telomere maintenance might be an important factor also in UM etiology, although with less impact than for CM.
**Inherited susceptibility**

UM predominantly occurs in a sporadic fashion, with approximately only 1% of the cases considered to be hereditary cases. However, patients with an inherited predisposition are likely to be more common than initially believed due to the discovery of UM being part of cancer syndromes. Between 2% and 5% of the UM cases have, therefore, been proposed to be familial cases.\(^\text{71}\) The first report of a germline mutation in \textit{BAP1} was published in 2010.\(^\text{10}\) One year later, two independent groups described inactivating germline mutations segregating in cancer-prone families, mainly characterized by distinct melanocytic neoplasms and mesothelioma in combination with UM and other cancers.\(^\text{74,75}\) Several other studies have, thereafter, described the link between familial UM and \textit{BAP1} germline mutations.\(^\text{76,77}\) The neoplasms associated with \textit{BAP1} germline mutations, also called the tumor predisposition syndrome, have been expanded to include CM, renal cell carcinoma, meningioma, and basal cell carcinoma.\(^\text{76,78-83}\) Additional cancer types are continuously being linked to this syndrome. In a review by Rai et al.,\(^\text{19}\) it was reported that 56 out of 57 families with a reported \textit{BAP1} germline mutation had one or more family members diagnosed with any of the main cancer types associated with this cancer syndrome (UM, CM, mesothelioma, or renal cell carcinoma).\(^\text{19}\) Still, only a subset of the families with an inherited predisposition for UM carries a germline mutation in \textit{BAP1} suggesting the presence of other, yet to be identified, high penetrance susceptibility genes. Several attempts identifying such genes have been done through large-scale sequencing approaches (ie, whole exome- and whole-genome sequencing) with little success so far, indicating that these genes are very rare and collaborations between research groups are needed. Another plausible explanation for the absence of additional UM susceptibility genes could be the presence of phenocopies in families that often comprise two relatives affected by UM, and where no clear aggregation of cancer cases is seen in the pedigree. Whether there is a polygenic component behind the inherited susceptibility for UM, as has been suggested for CM, is unclear. To date, no low to intermediate risk genes have been associated with UM.

**Clinical implications**

About half of all UM patients will subsequently suffer from metastatic disease, by hematogenous spread mainly affecting the liver. The survival for these patients is poor since UM is resistant to standard treatments with chemotherapy. Novel targeted therapies and immunotherapies have revolutionized the treatment of metastatic CM during the past years. In contrast, activating \textit{BRAF} mutations are very rare in UM and, consequently, \textit{BRAF} inhibitory treatment is not applicable. In CM, treatment with antibodies against immune checkpoint molecules has improved both disease-free survival and overall survival. A high burden of coding mutations has been associated with a better response to immune checkpoint inhibitors, such as CTLA-4.\(^\text{54,85}\) A high mutational load has also been correlated to a greater repertoire of neoantigens, which is associated with a better treatment response. Thus, the low burden of mutations found in UM might, therefore, be an explanation for the lower response to immunotherapies as compared to other melanoma subtypes.\(^\text{7}\) Due to the recent advances in sequencing technologies, multiple driver genes for UM have been discovered that hopefully will improve the understanding of the carcinogenesis behind this neoplasm and subsequently lead to prolonged survival of patients suffering from metastatic UM. The identification of driver genes has led to the identification of novel treatment targets and several clinical trials are ongoing investigating these targets in UM therapy.

Targeting mutated \textit{GNAQ/GNA11} directly is difficult because of the molecular nature of the mutations causing an inactivation of intrinsic GTPase within the cell. However, for several downstream molecules of \textit{GNAQ/GNA11}, targeted therapies have become available. These include mitogen-activated protein kinase/extracellular signal-regulated kinase (MEK) that is shown to be upregulated in \textit{GNAQ/GNA11} mutated tumors.\(^\text{86,87}\) Inhibition of MEK has actually been found to decrease the proliferation of UM tumors both in vivo and in vitro.\(^\text{88,89}\) Furthermore, a clinical Phase II trial has shown a prolonged progression-free survival of nearly 9 weeks when treating patients with the MEK inhibitor selumetinib compared to chemotherapy (temozolamide).\(^\text{90}\) However, in another Phase II trial, there was no significant effect on overall survival when treating with semurafenib compared to chemotherapy, although there was a modest increase in response rate and progression-free survival.\(^\text{91}\) Other putative downstream targets of \textit{GNAQ/GNA11} mutated tumors are protein kinases C and molecules of the protein kinase B (AKT)/mammalian target of rapamycin pathway.\(^\text{92-94}\)

Also \textit{BAP1} mutations are difficult to target directly because of their recessive nature. However, the effects of the mutations are possible to target by the use of histone deacetylase (HDAC) inhibitors in tumors with loss of \textit{BAP1} function. The absence of functional \textit{BAP1} protein leads to hyperubiquitination of H2A in the cells.\(^\text{43}\) The use of HDAC inhibitors can reverse this phenotype, thereby causing a shift from aggressive, dedifferentiated class-2 UM cells.
to more differentiated and less aggressive cells. HDAC inhibitors have also been suggested as adjuvant treatment in high-risk patients.

In familial cancer combining clinical and genetic information can be used to improve prognostic estimates and to improve strategies for early diagnosis. By genetic testing of cancer-prone families, the clinical outcome can in many situations be improved, by detecting precursor lesions and tumors at an early stage in members of mutation-positive families. Often, however, this is not straightforward, as in familial melanoma, where a low frequency of mutations in high penetrance genes is seen and risk estimates for mutation carriers have not been well established at this point. Genetic testing is often only recommended when the result is of importance in the management of the patient and where there is a possibility of improving the clinical outcome. However, in families exhibiting the phenotype specific for BAP1 tumor predisposition syndrome, genetic testing should be offered. Additional research will be of importance to elucidate the penetrance and risk of developing different types of cancer in mutation carriers. Identifying the susceptibility factor in cancer-prone families will be of importance for choice of surveillance programs and follow-up of the patient and their relatives. For families with a high cancer burden but without mutation in any known high predisposing gene, next-generation sequencing will be the natural choice to search for novel susceptibility genes. This will subsequently increase the knowledge about genetic susceptibility and may in the future be the basis for improved early detection and prevention of UM as well as lead to the development of new targeted treatments.

**Conclusion**

Five driver genes have so far been found to be frequently mutated in UM. Two of these, GNAQ and GNA11, are considered to occur early in carcinogenesis and to be of no prognostic relevance. Mutations occurring in the other driver genes are likely to have arisen later in the tumor development and thus are of importance for patient outcome. EIF1AX-mutated tumors show in general a strong correlation with class 1 GEP tumors and increased patient survival; BAP1 mutations, in contrast, associate with GEP class 2 tumors and poor survival; and SF3B1-mutated tumors seem to fall in between, which associate with late-onset metastatic disease. As being a complement to the GEP classification, mutation status of UM driver genes will hopefully increase the prognostic accuracy and be of help for deciding different treatment regimens, such as MEK inhibition therapy in GNAQ- and GNA11-mutated tumors. Other, probably more infrequent, mutated genes are continuously being detected, which will help us add more details to resolve the puzzle. UM also occurs in families with an inherited predisposition. The only high penetrance gene for hereditary UM identified so far is the BAP1 gene. Germline mutations in BAP1, mainly truncating mutations, have been found to segregate with reduced penetrance in families with many different cancer diagnoses, including but not exclusively to UM, CM, mesothelioma, and renal cell carcinoma. The search for additional novel UM susceptibility genes in BAP1-mutation negative families are ongoing through large whole-exome and genome sequencing. Knowledge on highly segregating penetrant mutations in affected families is of great importance for the management and surveillance of the patients and their relatives and will hopefully have positive impact on prevention and early diagnosis in the future.

**Disclosure**

The authors report no conflicts of interest in this work.

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