Oncogenic BRAF mutations and p16 expression in melanocytic nevi and melanoma in the Polish population

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Abstract

Introduction: Twenty-five – fifty percent of skin melanomas arise from nevi. Melanocyte proliferation is activated by BRAFV600E, then is arrested, but single nevi transform to melanomas. p16 controls arrest, and p16 loss may promote transformation.

Aim: To analyze BRAFV600E, p16 expression and melanocyte proliferation in dermal, compound and dysplastic nevi, cells of primary and metastatic melanoma in the Polish population.

Material and methods: One hundred and thirty-two nevi (dermal, compound, dysplastic) and 41 melanomas (in situ, primary, metastatic) were studied. BRAF was assessed by cobas® 4800 BRAFV600E Mutation Test, High Resolution Melting Assay validated with: pyrosequencing and immunohistochemistry. p16 and Ki67 expression was analyzed by IHC.

Results: Eighty-two percent of nevi and 57% of melanomas display BRAFV600E expression. Most dermal and compound nevi had > 50% of p16(+) cells. BRAFV600E dysplastic nevi had a low number of p16(+) cells. Nevi without BRAFV600E (WT), had 90% of cells p16(+). In 60% of in situ and primary melanomas, there was a low number of cells of p16(+). Fifty percent of WT metastatic melanoma and 33% of BRAFV600E showed a high level of p16. The number of Ki67(+) cells in dysplastic nevi was very low. In 25% of BRAFv600E melanomas in situ and 55% of WT, > 10% cells were Ki67(+). All BRAFV600E primary melanomas and 66% of WT had > 10% Ki67(+) cells. Twenty percent of BRAFV600E and WT metastases had > 10% of Ki67(+), however, 62% of BRAFV600E and 32% of WT samples had > 50% of Ki67(+).

Conclusions: BRAFV600E and p16 are more frequent in nevi than in melanoma in vivo. A significantly higher p16 expression was observed in mutated nevi than in WT, while in melanoma it was just the opposite. The proliferation rate of melanoma cells negatively correlated with p16 expression.

Key words: melanoma, nevi, BRAF, CDKN2A, p16, Ki67.

Introduction

Melanoma is a very heterogeneous disease. It shows a significant diversity in presentation, incidence and mortality in various countries, ethnic groups or patients with socioeconomic status [1]. The diversity is in part related to both histopathological [1] and genomic [2] melanoma subtypes, germline genetic factors predisposing or affecting melanoma development and progression [3] or environmental factors such as ultraviolet (UV) exposure [4]. Approximately 25–50% of skin melanomas arise or are associated with pre-existing benign melanocytic nevi [5]. On the other hand, nevi rarely transform to melanoma either exposed or not exposed to UV [6]. Recently, significant progress in understanding of the nevi progression to melanoma has been made in vitro, in animal models or human in vivo clinical studies [7, 8].

BRAF is serine/threonine protein kinase B-raf, constituent of the ERK1/2-signaling pathway involved in regulation of cells proliferation, differentiation and survival [9]. Active mutations in BRAF most frequent V600E...
(BRAF\textsuperscript{V600E}) are driver mutations in melanoma pathogenesis [9]. Moreover, a high frequency of BRAF\textsuperscript{V600E} is observed in nevi [10], which mostly develop during the first 25 years of life. Paradoxically, BRAF\textsuperscript{V600E} also induces p16 response in melano-cytic nevi that in response halts proliferating melanocytes [11]. Nevi melanocytes proliferation is transiently activated by BRAF\textsuperscript{V600E}. It is usually stopped when nevi reach 3–5 mm in diameter. Then nevi melanocytes proliferation is arrested or some nevi regress in the advanced age despite the expression of the BRAF\textsuperscript{V600E} for decades [11]. However, the arrest is reversible since a small fraction of nevi transforms to skin melanomas [12]. Originally it was proposed that nevi are senescent tumors [10]. However, later studies analyzing expression of effectors of senescence including p16\textsuperscript{INK4A}, p53 and DNA damage (γ-H2AX) and predictive markers of senescence such as Ki67, PML, senescence-associated β-galactosidase showed that nevi cells could not be distinguished from their precursors (skin melanocytes) and transformed melanocytes (melanoma cells). Accordingly, the concept that human nevi are arrested by senescence was challenged [13].

Oncogene-induced arrest is promoted by a high expression of cyclin-dependent (CDK) kinase inhibitors CDKN2A encoded p16 and CDKN2B encoded p15 [8]. p15 and p16 were highly expressed in isolated nevus melanocytes, while in isolated epidermal melanocytes from adjacent tissue displayed a low expression. In experimental models, primary melanocytes transduced with p15 halted proliferation, while transduced with p16 continued to proliferate but more slowly than controls [8]. In melanoma, both p15 and p16 expression was decreased [14]. Deletion of chromosome arm 9p, which contains the CDKN2A and CDKN2B genes, is common in primary melanoma [15]. In 90% of deletions of CDKN2A gene, the deletion of adjacent CDKN2B gene is also observed [16].

Mutations or polymorphic variants of CDKN2A are linked with predisposition to sporadic and hereditary melanoma [17, 18]. Variation in the phenotype and the nevi number within population is mostly genetically determined, for example A148T variant showed a protective role on the number of atypical nevi in the family sample [17]. In the Polish population, germ-line mutations are rare though there are three common polymorphic variants of CDKN2A, A148T, N1500C>G and N540C>T. Variant A148T was shown to be associated with an increased risk of melanoma [19–21].

Aim

The aim of the study was to analyze oncogenic BRAF\textsuperscript{V600E} and p16 expression as well as melanocyte proliferation in dermal, compound and dysplastic nevi, and melanoma cells of primary and metastatic melanoma in the Polish population.

Material and methods

Patient samples

The study was carried out at the Poznan Medical University (PMU) and Greater Poland Cancer Centre (GPCC). It was a retrospective study approved by the Bioethics Committee of PMU. Melanoma and nevi formalin-fixed, paraffin-embedded tissue blocks (FFPE) were obtained from archives of the Department of Dermatology, PUM and Department of Cancer Pathology, GPCC, Poznan, Poland. Samples were collected from patients in 2013–2016. There were FFPE obtained from 173 subjects (41 melanoma patients, 132 healthy individuals with nevi), which were randomly selected for studies and evaluated by two independent investigators (pathologists, E.B. and A.Mar.). Nevi were diagnosed as dermal (89 samples), compound (30) and dysplastic (13). Melanoma samples were described as melanoma in situ (20 samples), primary melanoma Clark III/IV (7) and metastatic (14; lymph node metastases – 10, other metastases – 4 samples).

DNA isolation

For genomic DNA isolation, the COBAS® DNA Sample Preparation Kit (Roche) was used according to the protocol provided by the manufacturer. Briefly, one 5 μm thick section from FFPE was deparaffinized and then lysed in DNA Tissue Lysis Buffer supplemented with Proteinase K. DNA Binding Buffer and isopropanol were added to the samples and the lysates were placed on a High Pure Filter Tube. Samples were washed twice and the DNA was eluted with DNA Elution Buffer. The concentration was measured with NanoDrop™ 2000 Spectrophotometer (Thermo Fisher). DNA was stored in –80°C.

Determination of BRAF status

Two different methods were utilized for BRAF status determination: (i) cobas® 4800 BRAF V600 Mutation Test and (ii) High Resolution Melting (HRM) Assay [22], which was further validated with two tests: pyrosequencing of BRAF exon 15 and immunohistochemistry (IHC) with anti-BRAF V600E (VE1) Mouse Monoclonal Primary Antibody (Ventana). Out of 173 FFPE samples, 126 were tested with cobas® 4800 BRAF V600 Mutation Test according to the protocol provided by the manufacturer. Forty-eight samples and the lysates were placed on a High Pure Filter Tube. Samples were washed twice and the DNA was eluted with DNA Elution Buffer. The concentration was measured with NanoDrop™ 2000 Spectrophotometer (Thermo Fisher). DNA was stored in –80°C.

Immunohistochemistry

From nevi and melanoma, FFPE sections of 4–5 μm were cut, mounted on the adhesive slides and subjected to IHC using CINtec p16 Histology (E6H4 clone) to detect p16\textsuperscript{INK4A}. The IHC used RoView’s ultraView Universal DAB Detection Kit. Immunoperoxidase staining was
performed on BenchMark Ultra from Ventana Medical System (Roche Diagnostics). As an external p16 positive control, an earlier validated squamous carcinoma specimen was used. For semiquantitative assessment of p16, 12-point score was calculated. It was based on the 4-point scale of staining intensity (0 – indicates no staining, 1 – weak positive, 2 – positive, 3 – highly positive) and on a 4-point percentage scale of the number of stained cells (1 – 10% and below of stained cells, 2 – 11–50%, 3 – 51–75% of stained cells and 4 – 76–100%). The final score was obtained by multiplying the degree of intensity by percentage of stained cells. Two independent investigators scored the samples (V.F. and A.Mar).

For Ki67, monoclonal antibody RTU Clone MIB-1 and IHC kit En VisionTM FLEX GV800 (DAKO, Denmark) were used. The IHC staining was carried out using automated system OMNIS (DAKO). As positive control for Ki67 tonsil tissue was applied. The result obtained was expressed as a number (in percent) of stained cells: 1–10%, 11–50%, 51–75% and 76–100%.

**Statistical analysis**

All statistical analyses were performed with GraphPad Prism 6. Continuous variables are given as means ± SD unless otherwise specified. Chi-squared test was used for comparisons of proportions. We used Mann Whitney test and Kruskal-Wallis test to compare means of two or more than two groups, respectively. Spearman correlation was used to analyze the p16 and Ki67 expression. All given \( p \)-values are two-tailed and a \( p \)-value < 0.05 was regarded statistically significant.

**Results**

**Expression of \( \text{BRAF}^{\text{V600E}} \)**

Expression of \( \text{BRAF}^{\text{V600E}} \) was found in 108 out of 132 nevi (82%) and in 24 out of 41 melanomas studied (58%) (Figure 1 A). The frequency of \( \text{BRAF}^{\text{V600E}} \) mutation in our melanoma samples corresponds to the previously reported high level of \( \text{BRAF}^{\text{V600E}} \) mutation in TCGA data (163 samples out of 366 sequenced melanomas (44.5%) display \( \text{BRAF}^{\text{V600E}} \) mutation according to www.cbioportal.org) [16, 24].

Eighty-two out of 89 dermal nevi (92%), 21 out of 30 compound nevi (70%), 5 out of 13 dysplastic nevi (38%), 12 out of 21 melanomas in situ (57%), 4 out of 7 primary melanomas Clark III/IV (57%) and 8 out of 14 metastatic melanomas (57%) were \( \text{BRAF}^{\text{V600E}} \) positive. There was a statistically significant difference in \( \text{BRAF}^{\text{V600E}} \) frequency between nevi and melanoma (\( p < 0.0001 \)) (Figure 1 B). Intensity of positive anti-\( \text{BRAF}^{\text{V600E}} \) staining in IHC was significantly higher in melanomas than in nevi (Figure 2).

**Expression of p16**

Expression of p16 was observed in all but 3 cases, which included one compound nevus, one dysplastic nevus and one melanoma case. In dermal and compound nevi, there are 90% and 80% of samples with more than 50% of p16 positive cells, respectively (Figure 3 A). In compound nevi with \( \text{BRAF} \) mutation, more than half of cases had a significantly lower percentage of p16 positive cells (below 50%) as opposed to compound nevi without \( \text{BRAF} \) mutation (referred to as wild type – WT), which

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**Figure 1.** The frequency of \( \text{BRAF}^{\text{V600E}} \) point mutation in nevi and melanoma samples. A – The \( \text{BRAF}^{\text{V600E}} \) mutation was detected in 108 out of 132 nevi samples (82%) and 24 out of 42 melanoma samples (57%). B – Dermal nevi are characterized by the highest frequency of \( \text{BRAF}^{\text{V600E}} \) mutation (92%) when compared to compound or dysplastic nevi. The frequency of \( \text{BRAF}^{\text{V600E}} \) mutation among melanoma samples is similar. Error bars = SD, **\( p < 0.01 \)**
Figure 2. The p16 immunohistochemistry score differs significantly between nevi and melanoma samples. A – The frequency of p16-positive cells in nevi and melanoma samples according to the pathological classification and the BRAF mutation status. 5-point percentage scale of the number of positive cells was used (0 = no positive cells, 1 = 1–10% of positive cells, 2 = 11–50%, 3 = 51–75% and 4 = 76–100%). B – The average p16 IHC score is significantly higher in nevi when compared to melanoma samples (\( p < 0.0001 \)). C – The average p16 IHC score in nevi and melanoma according to the BRAF mutation status. D – The average p16 IHC score in nevi and melanoma samples according to the pathological classification. Melanoma in situ is characterized by the lowest level of p16 IHC score. E – The frequency of BRAF\(^{V600E}\) mutation in nevi and melanoma samples according to the level of p16 IHC score (low score \( \leq 6 \), high score > 6). BRAF\(^{V600E}\) mutation is more frequent in nevi with high p16 (\( p < 0.05 \)) in comparison to low p16 nevi; in contrast, BRAF\(^{V600E}\) mutation is slightly less frequent in melanoma with high p16 when compared to low p16 samples. F – The frequency of BRAF\(^{V600E}\) mutation in nevi and melanoma samples according to the level of p16 IHC score (low score \( \leq 6 \), high score > 6) and the pathological classification. Error bars = SD, ***\( p < 0.0001 \)
have shown more than 50% of p16 expressing cells in 90% of cases (Figure 3 A). In melanoma in situ and primary melanoma (Clark III/IV), about 60% of cases showed a low (below 50%) number of cells expressing p16. In metastatic melanoma WT about 50%, while in BRAFV600E only 33%, of cases showed a high p16 expression (above 50% cells positive) (Figure 3 A).

In nevi, the p16 expression (average IHC score = 9.129 ±2.883) was statistically higher (p < 0.0001) than in melanoma (average IHC score = 6.122 ±3.480) (Figure 3 B). In WT nevi, p16 expression (average IHC score = 7.750 ±3.505) was lower than in BRAF V600E nevi (average IHC score = 9.435 ±2.684), while in WT melanoma it was higher (average IHC score = 6.588 ±3.063) than in BRAFV600E melanoma (average IHC score = 5.792 ±3.776), although with no statistical significance (Figure 3 C). However, in mutated BRAFV600E nevi, p16 expression was significantly higher than in BRAFV600E melanoma (p < 0.0001). In WT and BRAFV600E dermal nevi, p16 expression was at the same level (average IHC score in WT = 7.944 ±2.725 and in BRAFV600E = 8.857 ±2.268), however it slightly differs in compound nevi (average IHC score in WT = 6.333 ±3.841 and in BRAFV600E = 6.095 ±3.150) but with no statistical significance (Figure 3 D).

Due to variations in p16 expression within groups studied we decided to divide patients in two subgroups, one with high and the other with low p16 expression (Low and High p16 IHC score with cutoff expression of 6, respectively) and evaluate the frequency of BRAFV600E in nevi and melanoma separately (Figure 3 E). In nevi with low p16 and nevi with high p16, there were 57% and 85% of BRAFV600E carriers, respectively (p < 0.01). On the other hand, in melanoma with low p16, there were 70% of BRAFV600E cases and in melanoma with high p16, – 47% of BRAFV600E cases, however, the difference was not statistically significant. There was a very significant difference between the frequency of BRAFV600E samples in high p16 nevi (85%) and high p16 melanoma (47%) (p < 0.0001). We further subdivided samples according to the pathological classification and observed that in compound nevi, all samples showed a high p16 score (Figure 3 F). In melanoma in situ and primary melanoma Clark III/IV, the frequency of BRAFV600E is higher in low p16 samples, however with no statistical significance. In contrast, metastatic melanoma samples with high p16 display a slightly higher frequency of BRAFV600E cases in comparison to the low p16 subgroup (60% vs. 50%) (Figure 3 F).

Expression of Ki67

Expression of Ki67 was evaluated in dysplastic nevi and melanoma cases. The number of dysplastic nevi cases expressing Ki67 in more than 10% of cells was very low (Figure 4 A). 80% of BRAFV600E and 83% of WT cases demonstrated a similar number of stained cells, which did not exceed 10% of Ki67 positive cells. In 44% of melanoma in situ with BRAFV600E and in 70% of WT melanoma samples, more than 10% of Ki67 positive cells were seen. All cases of primary melanoma Clark III/IV with BRAFV600E and 66% of WT had more than 10% of Ki67 positive cells. In melanoma metastases with BRAFV600E and WT, the fraction of cases with less than 10% of Ki67 positive cells was similar, approximately 20%, however, there were 62% of BRAFV600E samples and 32% of WT cases with more than 50% of Ki67 positive cells (Figure 4 A).

The average Ki67 IHC score is higher in melanoma (average IHC score = 5.225 ±3.092) than in nevi (average IHC score = 2.364 ±0.809) (p-value = 0.001), however it does not differ in pathological subgroups of melanoma, even when subdivided into BRAFV600E and WT-expressing cases (Figures 4 B–D). However, when samples were divided into a low and a high Ki67 expressing group and the frequency of BRAFV600E mutation was tested in each pathological subgroup, we observed a higher frequency of BRAFV600E in high Ki67 expressing subgroups of melanoma in situ (71% vs. 42% in low Ki67 group), primary melanoma Clark III/IV (67% vs. 50% in low Ki67 group) and metastatic melanoma (63% vs. 50% in low Ki67 group) (Figure 4 E).

The correlation of p16 and Ki67 expression

The correlation of p16 and Ki67 expression in melanoma samples was determined using Spearman rank correlation (Figure 5). We observed a statistically significant (p < 0.05) negative correlation between p16 IHC score and Ki67 IHC score in melanoma samples (r = –0.3234). The presence of BRAFV600E or WT does not affect the p16 and Ki67 correlation (data not shown). To present in the graph melanoma samples with identical p16 and Ki67 IHC scores, the color legend was utilized.

Discussion

In the last decade significant progress in genomic classification [2] and genetic evolution of skin melanoma from precursor lesions [3, 5, 6] has been achieved. Most of the current knowledge of melanoma is derived from the study of patients from populations of European descent [1]. However, there are no reports of BRAFV600E driven evolution melanomas in the Polish population. Previous reports indicated CDKN2A variants associated with melanoma and other cancers in Poland [19–21, 25]. Moreover, our functional in vitro studies using vectors carrying 3′UTR sequence variants 500C>G and 540C>T together with luciferase reporter gene transfected into multiple human melanoma cell lines demonstrated a very significant reduction (up to 4000 times) of the luminescence emitted by transfected cells [26]. This indicates that polymorphisms may modulate CDKN2A gene expression, thus the level of p16 in melanoma cells.

Beyond in vitro and mouse models [7, 8, 27], there is only a single clinically focused report [28] on onco- genic BRAFV600E mutation and p16 expression impact.
Figure 3. The Ki67 immunohistochemistry score differs significantly between dysplastic nevi and melanoma samples. 
A – The frequency of Ki67-positive cells in nevi and melanoma samples according to the pathological classification and the BRAF mutation status. 4-point percentage scale of the number of positive cells was used (1 = 0–10% of positive cells, 2 = 11–50%, 3 = 51–75% and 4 = 76–100%). B – The average Ki67 IHC score is significantly higher in melanoma when compared to dysplastic nevi samples ($p = 0.001$). C – The average Ki67 IHC score in dysplastic nevi and melanoma according to the pathological classification. D – The average Ki67 IHC score in dysplastic nevi and melanoma samples according to the pathological classification and BRAF mutation differs significantly. E – The frequency of BRAF$^{V600E}$ mutation in dysplastic nevi and melanoma samples according to the level of Ki67 IHC score (low score ≤ 6, high score > 6). F – The frequency of BRAF$^{V600E}$ mutation in dysplastic nevi and melanoma samples according to the level of Ki67 IHC score (low score ≤ 6, high score > 6) and the pathological classification. In dysplastic nevi there are no samples with high Ki67 IHC score. Error bars = SD, *$p < 0.05$, **$p < 0.01$, ***$p = 0.001$
on the growth rate of nevi and early melanoma in vivo. However, the growth rate in later phases of $\text{BRAF}^{\text{V600E}}$ mutated melanoma development (metastases) in the context of p16 expression remains unexplored. Recently Tschandl et al. [29] has measured the growth rate of early melanoma and nevi in vivo using digital dermoscopy. They found that melanomas grew faster than nevi. Melanomas with $\text{BRAF}^{\text{V600E}}$ mutation grew significantly faster than WT. The effect of $\text{BRAF}^{\text{V600E}}$ was not observed in melanocytic nevi. Moreover, they found no difference in p16 expression between nevi and melanoma however, they saw a higher growth rate of $\text{BRAF}^{\text{V600E}}$ melanoma in vivo with negative p16. In our studies we found a significant difference between $\text{BRAF}^{\text{V600E}}$ frequency in nevi and melanoma as well as p16 lower expression in $\text{BRAF}^{\text{V600E}}$ mutated melanoma than nevi. Moreover, p16 expression level (score) negatively correlated with melanoma cells proliferation (Ki67). However, the final conclusion of Tschandl et al. [29] report related to negative correlation between p16 and in vivo growth rate of $\text{BRAF}^{\text{V600E}}$ melanoma variant was identical with our conclusions. In addition, Tschandl et al. admitted a limitation of the inclusion of cases in their study, which a priori were pre-selected for digital monitoring.

| Hematoxylin + eosin | BRAF (Ventana VE1) | p16 | Ki67 |
|---------------------|--------------------|-----|------|
| Nevus $\text{BRAF}^{\text{V600E}}$ | | | |
| Dysplastic nevi $\text{BRAF}^{\text{V600E}}$ | | | |
| Melanoma $\text{BRAF}^{\text{V600E}}$ | | | |

Figure 4. Representative images of BRAF (VE1), p16 and Ki67 immunohistochemical staining in nevi and melanoma samples. Dermal nevus with $\text{BRAF}^{\text{V600E}}$ mutation, dysplastic nevi with $\text{BRAF}^{\text{V600E}}$ and WT status and primary melanoma Clark III/IV with $\text{BRAF}^{\text{V600E}}$ mutation immunostained for specific markers are depicted.

Figure 5. Spearman correlation of p16 and Ki67 IHC scores. The expression of p16 negatively correlates with the level of Ki67 in melanoma samples ($r = -0.3234$; $p = 0.0446$). Dots are colored as follows: white – single sample, light grey – 2 samples, dark grey – 3 samples, black – 4 samples with specific p16 and Ki67 IHC score values.
In our and Tschandl et al. [29] studies, only V600E BRAF mutation was investigated, while there are other BRAF mutations such as G466E, S467L, G469R/E, V559E, N581H/S/T, and others [2], which we did not study. Moreover, in our WT group there are other driver mutated genes such as NRAS, which are found in nevi and melanoma and similarly as BRAF activate mitogen-activated protein kinase (MAPK) pathway and are regulated by p16 [29]. In our earlier study (data not published), we detected BRAFV600E in 49% of metastatic melanoma, and various mutations of NRAS in 22% of the same cohort. Accordingly, the differences between BRAFV600E and WT study groups as referred by us could be underestimated. In addition, Tschandl et al. [29], in melanoma associated nevi and uninvolved nevi, found BRAFV600E mutation in 63% of melanomas, 65.2% of associated nevi and 50.0% of control nevi using the same VE1 anti-BRAFV600E antibody for IHC. Significant differences in distribution of BRAF and NRAS mutation were found. They observed the same difference between intensity of BRAFV600E IHC staining in nevi and melanoma as in our study. Based on the above results they challenged the current model of stepwise melanoma progression.

In cultured human melanocytes mutant BRAFV600E protein induced cells senescence by increasing expression of CDKN2A [15]. Accordingly, it has been hypothesized that to progress into melanoma, CDKN2A needs to be mutated or deleted. An experimental in vitro study demonstrated that simultaneous knockdown of BRAFV600E and CDKN2A in melanoma cells caused growth inhibition and apoptosis, while knockdown of BRAFV600E or CDKN2A alone had no effect. Accordingly, it became indicative of a role of multiple tumor suppressors in the prevention of BRAFV600E oncogenesis [11]. Indeed, recently another CDK inhibitor, namely CDKN2B gene (p15) was found to be involved in nevi proliferation arrest in which BRAFV600E activation induced expression of TFG-β signaling, which in turn up-regulated p15 and inhibited proliferation [8]. Taylor et al. [14] very recently reported on a random series of 29 nevi and 27 melanomas expression of p16 and p15. Both inhibitors demonstrated a high expression in nevi and substantially lower in melanoma on both protein and transcript levels. The differences were more significant for p15, which authors nominated for the biomarker candidate for distinguishing nevi from melanoma. However, they did not take into consideration BRAFV600E mutated melanomas and BRAFV600E subtypes of melanocytic nevi or melanoma in various stages of development.

Conclusions

Finally, we found that BRAFV600E and p16 are more frequent in nevi than in melanoma in vivo. A significantly higher p16 expression was observed in mutated nevi than in WT, while in melanoma it was just the opposite. The proliferation rate of melanoma cells negatively correlated with p16 expression. Our results confirm the hypothesis of the traditional model of stepwise tumor progression. However, further studies involving other driver mutations are required in order to fill the gap of our study, which is the better characterization of so called “WT” nevi and melanomas. Based on the obtained results and data available so far, just one study carried out in Austria [28], it is difficult to define the unique pattern of nevi-melanoma transition in the Polish population. Moreover, larger study groups are required.

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Conflict of interest

The authors declare no conflict of interest.

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