Variation of mutant allele frequency in NRAS Q61 mutated melanomas

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Abstract

Background: Somatic mutations of BRAF or NRAS activating the MAP kinase cell signaling pathway are present in 70% of cutaneous melanomas. The mutant allele frequency of BRAF V600E (M%BRAF) was recently shown to be highly heterogeneous in melanomas. The present study focuses on the NRAS Q61 mutant allele frequency (M%NRAS).

Methods: Retrospective quantitative analyze of 104 NRAS mutated melanomas was performed using pyrosequencing. Mechanisms of M%NRAS imbalance were studied by fluorescence in situ hybridization (FISH) and microsatellite analysis.

Results: M%NRAS was increased in 27.9% of cases. FISH revealed that chromosome 1 instability was the predominant mechanism of M%NRAS increase, with chromosome 1 polysomy observed in 28.6% of cases and intra-tumor cellular heterogeneity with copy number variations of chromosome 1/NRAS in 23.8%. Acquired copy-neutral loss of heterozygosity (LOH) was less frequent (19%). However, most samples with high M%NRAS had only one copy of NRAS locus surrounding regions suggesting a WT allele loss. Clinical characteristics and survival of patients with either <60% or ≥60% of M%NRAS were not different.

Conclusion: As recently shown for M%BRAF, M%NRAS is highly heterogeneous. The clinical impacts of high M%NRAS should be investigated in a larger series of patients.

Keywords: Melanoma, M%NRAS, Imbalance, Pyrosequencing, WT allele loss

Background
Cutaneous melanoma is a highly aggressive and treatment-resistant human cancer. The most frequent genetic alterations involve genes of the MAP kinase signaling pathway [1–3]. Activating hot-spot mutations are mainly found in BRAF (codon V600) and in NRAS (codon Q61, and less frequently in the codons G12 and G13) genes, in 35–50% and 15–25% of cutaneous melanoma, respectively [4, 5]. Among BRAF alterations, the BRAF V600E mutation in exon 15 is predominant (85%) and due to a substitution of a valine to a glutamic acid (c.1799 T > A, p.V600E) [6, 7]. BRAF and NRAS mutations are almost always mutually exclusive [8, 9].

Mutant NRAS melanomas have been reported to have more aggressive clinical features than other subtypes, with thicker lesions, elevated mitotic activity, and higher rates of lymph node metastasis [10–12]. Additionally, NRAS mutation status was reported as a predictor of poorer outcomes with lower median survival compared to non-NRAS mutated melanoma [10, 13].

The discovery of BRAF mutations led to the development of targeted treatments [14, 15]. However despite major clinical benefit in melanomas with BRAF mutation, secondary resistance occurs in most patients during the first year of treatment. Thus combinations of BRAF and MEK inhibitors have been developed, and were shown to induce longer progression free survivals (PFS) of patients with BRAF mutated melanomas [16–18]. By contrast, targeted treatment of patients with NRAS mutated melanomas is still a challenge, although an international phase 3 prospective study with the MEK inhibitor binimetinib recently provided promising results [19].

We recently studied the frequency of BRAF mutant alleles (M%BRAF) and showed that M%BRAF is highly...
heterogeneous and frequently increased in BRAF mutated melanomas [20]. Interestingly, a recent clinical study showed that the increased BRAF V600 mutation level was significantly associated with a better response rate to vemurafenib during the first 10 months of treatment [21]. These observations highlighted the importance of quantitative evaluation of BRAF mutation before melanoma treatment.

Although biological and clinical implication of the frequency of mutant alleles of BRAF in melanomas are currently under investigation, no data are available concerning the variation of M%NRAS. Accordingly, we conducted this study to investigate NRAS Q61 mutations and M%NRAS in a series of 199 melanomas wild type for BRAF V600. The mechanisms of the M%NRAS variations were then studied by fluorescence in situ hybridization (FISH) and by amplified fragment length polymorphism (AFLP).

Methods

Patients and samples

Melanoma samples were obtained from the bank of biological resources of Ambroise Paré Hospital in Boulogne-Billancourt. The research was performed in compliance with the ethical principles of the Helsinki Declaration (1964) and with the French ethics laws. Patients were informed and approved the use of their samples for research purpose. Tumor samples collection was declared to the French Ministry of Research (DC 2009–933). Melan-Cohort study was approved by CPP IDF 8 Ethics committee (030209) and registered with https://www.clinicaltrials.gov/ct2/search (NCT00839410). Clinical and survival data were collected from clinical records of the Dermatology Department of Ambroise Paré Hospital.

The frequency of NRAS Q61 mutations was evaluated in a consecutive series of melanomas received for diagnosis from March 2013 until May 2015. Additionally, a second series of patients, whose samples were received earlier to March 2013, mutated for NRAS were also included for the evaluation of M%NRAS. In our previous paper concerning the BRAF mutant allele frequency in melanoma [20], we observed a distinct distribution of the percentage of mutated allele according to the percentage of tumor cells. However, the inter-pathologist reproducibility for the evaluation of tumor cell content was substantial for the 80% cut-off (κ = 0.79). Therefore, we excluded samples with less than 80% of melanoma cells from further NRAS molecular analysis.

NRAS mutant allele detection and quantification

Before DNA extraction, HES slides were reviewed to confirm the presence of melanoma cells and to select areas with highest density of tumor cells for macrodissection. For microsatellite analysis, DNA from corresponding normal tissue section was extracted. Genomic DNA was extracted from formalin-fixed and paraffin-embedded (FFPE) fragments of melanoma as previously described [22].

Pyrosequencing was performed as already described [23]. Profiles for different NRAS mutations were established and confirmed by Sanger sequencing method (Additional file 1: Figure S1). Three different assays for detection of NRAS Q61 mutation were designed and primers used for DNA amplification, pyrosequencing, and Sanger method are presented in Table 1. Pyrosequencing Assay 1 allows the quantification of all but one NRAS Q61 mutation (c.183A > T p.Q61H) [24]. Pyrosequencing Assay 2 is an edited version of Assay 1, in which the order of injected nucleotides was modified to allow the quantification of all NRAS Q61 mutations. Pyrosequencing Assay 3 was developed to rescue some cases with a very bad FFPE DNA quality. For patients with several samples available, the M%NRAS used was that obtained from the first metastasis.

Microsatellite analysis

Eight microsatellite markers were selected from the NCBI dbSNP short genetic variations database and analyzed by AFLP. Their positions are showed in Fig. 1, and the primers used in Table 1. No highly heterozygous microsatellite was identified within NRAS gene; thus the genetic status of this gene was evaluated on the basis of two microsatellites close to NRAS locus (rs3220698 and rs3220987). PCR reactions were performed with fluorescent-labeled forward primers and the amplified PCR products were analyzed by capillary array electrophoresis on the ABI PRISM 3100 sequencer (Applied Biosystems, Foster City, USA) and GeneScan software (Applied Biosystems). Detection of loss of heterozygosity was performed as detailed in Loss of Heterozygosity Analysis Getting Started Guide. Two independent experiments were performed to confirm LOH results. The probability of one NRAS allele loss was evaluated as very high if only one allele was detected in each NRAS locus surrounding microsatellites (e.g. Fig. 1, case Y11.136). If only one of NRAS surrounding markers presented one microsatellite allele, this probability was evaluated as mean (e.g. Fig. 1, case Y14.711). The analysis was considered as inconclusive if both NRAS locus surrounding microsatellites gave non informative results (e.g. Fig. 1, case Y10.1471).

FISH analysis

FISH analysis was performed on tissue microarray (TMA) containing 94 melanomas and on tissue sections from 7 melanoma patients. FISH probe preparation and FISH
technique was performed as already described [20, 25]. All samples were analyzed with RP11-245I3 and RP11-269F19 probes covering \(\text{NRAS}\) and a region of chromosome 1 telomeric to \(\text{NRAS}\) gene (chr1:45.142.760–45.303.288, (2009 GRCh37/hg19)), respectively. Chrome- 

\[ \text{NRAS disomie was concluded if two FISH signals for each probe were observed and polysomy if three or more FISH signals were detected in the majority of cell.} \]

Chromosome 1/\(\text{NRAS}\) signal ratios 2:1 and 1:2 were described as \(\text{NRAS}\) monosomy and \(\text{NRAS}\) gain, respectively. Innumerable \(\text{NRAS}\) FISH signals were interpreted as amplification.

**Statistical analysis**

Overall survival (OS) was defined as the period between the date of the primary melanoma diagnosis to the date of death (all causes) or last follow-up evaluation. Survival was censored at the last follow-up evaluation. Distant metastasis free survival (DMFS) was defined as the period between the date of the primary melanoma diagnosis to the date of onset of stage IV melanoma or death. The date of onset of stage IV melanoma was defined as the date of the clinical examination or imaging procedure that provided an unequivocal diagnosis of distant visceral metastasis. Progression-free survival was defined as the period between the date of the primary melanoma diagnosis (with or without lymph node sentinel biopsy procedure) to the date of the first regional (node or cutaneous) recurrence (stage IIIB minimum). Progression-free survival, distant metastasis-free survival and overall survival curves were estimated using the Kaplan-Meier method, and differences between PFS, DMFS, and OS curves were assessed using the log-rank test.

Survival and histoprognostic markers (age, gender, Breslow index, ulceration and mitotic activity) of \(\text{NRAS}\) Q61 mutated primary melanomas were compared between two groups: one group of 48 melanomas with <60% of M%\(\text{NRAS}\) and another group of 24 tumors with ≥60% of M%\(\text{NRAS}\).

Student tests were performed for quantitative values, and Chi\(^2\) tests for qualitative values. The results were considered significant when \(P < 0.05\).

**Results**

**NRAS mutation frequency and allele quantification**

The frequencies of \(\text{BRAF}\) V600 and \(\text{NRAS}\) Q61 mutations were evaluated in 267 FFPE melanoma patients of the series (flow chart in Additional file 2: Figure S2) diagnosed between March 2013 and May 2015. \(\text{NRAS}\) Q61
mutation was detected in 48 melanomas, corresponding to 18% (48/267) of all melanomas and 33.8% (48/142) of \textit{BRAF} \textit{V600} wild type cases. Additionally, 63 \textit{NRAS} \textit{Q61} mutated melanomas, diagnosed before this period, were included into the molecular analyses. In total, 111 \textit{NRAS} \textit{Q61} mutated tumors were collected with 60–95\% tumor cells.

Characterization of M\%\textit{NRAS} were performed in a larger group of 104 \textit{NRAS} \textit{Q61} mutated melanomas containing ≥80\% tumor cells. The corresponding mutations were c.182A > G p.Q61R in 58.6\% (61/104), c.181C > A p.Q61K in 23.1\% (24/104), c.182A > T p.Q61L in 13.5\% (14/104), c.183A > T p.Q61H in 2.9\% (3/104) and c.183A > C p.Q61H in 1.9\% (2/104) of samples. M\%\textit{NRAS} was highly heterogeneous, ranging from 15.5 to 94\% (Fig. 2). The majority of cases (60.6\%, 63/104) had ≥30 to 60\% M\%\textit{NRAS} and was thus considered as heterozygous (HET). The remaining 41 cases were considered as having non-heterozygous M\%\textit{NRAS}: 11.5\% (12/104) had <30\% of M\%\textit{NRAS} (Low non-HET) and 27.9\% (29/104) had ≥60\% of M\%\textit{NRAS} (High non-HET).

**Fig. 1** Chromosome 1 microsatellite analyses in \textit{NRAS} mutated melanomas. Summary of AFLP analysis of eight chromosome 1 microsatellite markers performed in 29 \textit{NRAS} \textit{Q61} mutated melanomas. Each row contains following information for one patient: identification n°, \textit{NRAS}/chromosome 1 FISH result, \textit{NRAS} mutant allele percentage, results of microsatellite markers LOH. \textit{NRAS} gene and centromere localization are indicated. NA – non analysable, NI – non informative, NR – not realized. FISH: 1 - disomy, 2 - disomy with rares polysomic cells, 3 polysomy (A:3–4 copy, B: >4 copy), 4 - amplification, 5- monosomy, 6- high intra-tumor copy number variations of \textit{NRAS}/chromosome 1.

**Fig. 2** \textit{NRAS} \textit{Q61} mutant allele burden in melanomas. Histogram representation of \textit{NRAS} \textit{Q61} mutant allele quantity (in percentage) in 104 \textit{NRAS} mutated melanomas. The X and Y axis correspond to the percentage of \textit{NRAS} mutant and to the number of cases, respectively.
We then compared our results with database of the Cancer Genome Atlas (TCGA) analyzed on the CBioPortal platform. Among the 85 NRAS mutated cases that were available only 50.6% (43/85) had a heterogeneous status of NRAS mutation, while 34.1% (29/85) was High non-HET and 15.3% (13/85) was Low non-HET (Additional file 3: Figure S3).

**FISH analysis**
Among 101 samples (41 NRAS WT and 60 NRAS Q61) analyzed by FISH with NRAS locus and chromosome 1 specific fluorescent probes, different types of alternations were observed: no alteration of NRAS/chromosome 1 (disomy), disomy but rare cells with polysomy, polysomy and monosomy, which were detected in 23.1% (24/101), 19.2% (20/101), 9.6% (10/101) and 3.8% (4/101) of cases, respectively. NRAS amplification was a rare alteration and observed in 6.7% of melanoma samples (7/101). In 23.1% of samples (24/101), we observed intra-tumor cellular heterogeneity in NRAS/chromosome 1 copy numbers. Finally, FISH analysis of NRAS gene was non informative in 14.4% of cases (15/101).

To better understand the chromosomal mechanisms leading to M%NRAS increase, we compared M%NRAS and NRAS/chromosome 1 copy number status between 32 BRAF/NRAS WT and 57 NRAS Q61 mutated melanomas (Fig. 3). Disomy of NRAS/chromosome 1 (with or without polysomy in few cells) were detected in 59.4% (19/32) of BRAF/NRAS WT, 50% (15/30) of HET, but in only 28.6% (6/21) of High non-HET melanomas (P = 0.08). Polysomy of NRAS/chromosome 1 was detected in 13.3% (4/30) of HET, and 28.6% (6/21) of High non-HET samples, but was absent in BRAF/NRAS WT melanomas (P < 0.05). Amplification of NRAS was detected in 6.3% (2/32) and 9.5% (2/21) of BRAF/NRAS WT and High non-HET melanomas, respectively. Additionally, in 3.3% (1/30) of HET and 9.5% (2/21) of High non-HET cases, a gain of NRAS gene was observed. Deletion of NRAS was also a rare event and was observed in 9.4% (3/32) of BRAF/NRAS WT and 3.3% (1/30) of HET melanomas. A high intra-tumor copy number variation of NRAS/chromosome 1 was observed in 25% (8/32) of BRAF/NRAS WT, 30% (9/30) of HET and 23.8% (5/21) of High non-HET. FISH results of Low non-HET group were excluded from the comparative analysis because of insufficient numbers of samples (only 6 samples).

**Chromosome 1 microsatellite analysis**
Eight polymorphic microsatellite markers were analyzed in 6 HET and 23 High non-HET tumors melanoma samples. In all HET melanomas, non LOH of NRAS locus surrounding microsatellites was detected but one for which microsatellites presented a low LOH, near the upper LOH detection threshold. In the majority of samples of High non-HET group, the microsatellite analysis revealed the presence of only one marker in the regions surrounding NRAS gene (Fig. 1). The probability of one NRAS allele loss was evaluated as very high in 30% (7/23) and mean in 26.1% (6/23) of tumors. In 26.1% (6/23) of tumors, the LOH results were non informative. In 17.4% (4/23), LOH with the presence of two alleles rather than one allele loss was detected.

**Correlation of M%NRAS and clinical data**
Information on primary tumor was available for 72 patients. The main histological subtypes were SSM and nodular melanomas, in 44.4% (32/72) and 38.9% (28/72) of cases, respectively. Median Breslow was 2.84 mm [0.4–10]. An ulceration was present in 47.2% (34/72) of melanomas, and 58.3% (42/72) had a mitotic activity (> 1 mitosis / mm²). Among these cases, we found NRAS Q61R mutation in 63.9% (46/72) of melanomas, and 58.3% (42/72) had a mitotic activity (> 1 mitosis / mm²). Among these cases, we found NRAS Q61R mutation in 63.9% (46/72) of melanomas, NRAS Q61K in 25% (18/72), NRAS Q61L in 6.9% (5/72) and NRAS Q61H in 4.2% (3/72). M%NRAS was quantified by analysis of primary melanoma in 18.1% (13/72) of cases and of metastasis (lymph node, cutaneous or visceral) in 81.9% (59/72). Patients were divided into two groups depending on % of NRAS mutant allele. M%NRAS was ≥60% in the first group (n = 24, 33.3%) and <60% in the second cohort (n = 48, 66.7%). The second group contained 41 HET M%NRAS and 7 low not-HET M%NRAS cases. Clinical and pathological features of primary NRAS mutated melanomas were compared between these two groups and they are
summarized in Table 2. No statistically significant differences were observed between these two groups with baseline criteria ($P > 0.05$). The median follow-up of the patients was 40 months (range [1–445]). PFS, DMFS and OS in both groups were not different (Fig. 4).

Table 2 Clinicopathologic characteristics of studied subjects. Comparison of clinical and pathological features of NRAS mutated primary melanomas according to the NRAS mutant allelic burden (<60% or ≥60%)

| Origin of pyrosequencing sample | ≥60% of M% NRAS n = 24/72 (33.3%) | <60% of M% NRAS n = 48/72 (66.7%) | P-value (statistic test) |
|---------------------------------|---------------------------------|---------------------------------|------------------------|
| From primary melanoma           | 5                               | 8                               | 0.91 (Chi2 with Yates' correction) |
| From metastasis                 | 19                              | 40                              |
| Age                             | 63.1 ± 17.4                     | 65.2 ± 14.7                     | 0.61 (Student)         |
| Median, year [range]            | 60 [37–93]                      | 66 [32–97]                      |
| Gender                          | 13/11                           | 30/18                           | 0.49 (Chi2)           |
| Breslow index                   | 3.0 ± 2.1                       | 3.3 ± 2.2                       | 0.57 (Student)        |
| Median, mm [range]              | 3.00 [0.40–10.00]               | 2.61 [0.50–9.00]                | 0.14 (Chi2)           |
| Ulceration                      | 14                              | 20                              | 0.45 (Chi2)           |
| Mitotic activity                | 11                              | 31                              |
| Initial AJCC stage              | I 5                             | 10                              | 0.7 (Chi2)            |
|                                | IA 4                            | 2                               |
|                                | IB 1                            | 8                               |
|                                | II 11                           | 26                              |
|                                | IIA 4                           | 10                              |
|                                | IIB 6                           | 5                               |
|                                | IIC 1                           | 11                              |
|                                | III 7                           | 9                               |
|                                | IIIA 0                          | 2                               |
|                                | IIIB 4                          | 2                               |
|                                | IIIC 3                          | 6                               |
|                                | IV 1                            | 0                               |
|                                | NA 0                            | 2                               |
| Histologic subtype             | Nodular melanoma 9              | 19                              | 0.63 (Chi2)           |
|                                | SSM 10                          | 22                              |
|                                | Acral melanoma 1                | 0                               |
|                                | Lentigo maligna melanoma 0      | 1                               |
|                                | Mucosal melanoma 0              | 1                               |
|                                | On congenital naevus 0          | 1                               |
|                                | Inclassable 1                   | 2                               |
|                                | NA 3                            | 2                               |

$SMM$ superficial spreading melanoma  
$NA$ data not available  
$SD$ standard deviation  
$M/F$ male/female

**Discussion**

In this study, we reported the prevalence of NRAS Q61 mutation and, for the first time, the variations of NRAS mutant alleles (M%NRAS), in a large series of human melanoma samples. We have demonstrated that M%NRAS
was highly heterogeneous; indeed, only 61% of NRAS mutated melanomas were heterozygous, while 30% of cases had a significantly increased M%NRAS (≥60%). Our results were confirmed by analysis of the cases of the TCGA database.

NRAS pyrosequencing assays used in this study were developed to identify all hot-spot mutations in the codon 61 of NRAS gene. The specificity of these assays for different mutations was confirmed by Sanger sequencing. Additionally, the genotyping accuracy of 40 NRAS mutated melanomas, 27 of which were p.Q61R was confirmed by immunohistochemistry with an antibody against Q61R [27]. In a recent study, we have demonstrated that pyrosequencing was a robust molecular technique for oncogenic mutant allele quantification, by comparing it with quantitative real time PCR and picodroplet digital PCR [20].

Mutation in one allele of an oncogene is sufficient for activation of its targets and M%NRAS is expected to be around 50% in diploid cells. However, in tumours with high chromosome instability, chromosome number is rarely disomic and M%NRAS could widely exceed 50%. To better understand the chromosome mechanisms leading to NRAS mutant allele increase in the proportion of NRAS mutated melanoma, we firstly performed FISH analyses with 2 BAC probes covering NRAS region and another region of chromosome 1, telomeric to this gene, in a large series of 104 melanomas. Different types of NRAS/chromosome 1 status were observed. Polysomy was mainly observed in NRAS mutated tumours and disomic and/or disomic but rare polysomic cells were less frequent in High non-HET M%NRAS than in M%NRAS WT tumours. Amplification and deletion of NRAS gene were rarely observed and were seen in both NRAS WT and NRAS mutated melanomas. Genomic analysis of human cutaneous melanoma genomes have been described in several studies. However, in most of them, only melanoma cell lines were studied. The analysis of 60 melanoma cell lines by Gast [33] have revealed targeted focal amplifications of NRAS genes in 11% of them (n = 7/60) and amplification were detected in both NRAS mutated and NRAS WT melanomas. This frequency is higher that the frequency of NRAS amplification detected in the present series and in other reports. In a subset of cutaneous melanocytic lesions,
NRAS amplification was found to be restricted to a few cases with NRAS mutations [34]. Additionally, Stark and colleagues reported rare instances of focal amplification including NRAS gene in two cell lines with NRAS mutation; however, a poor correlation between copy number increase and concomitant mutation in this oncogene was described [35]. Polysomy of chromosome 1 and intra-tumour NRAS/chromosome 1 heterozygosity were frequently found in our series and was preferentially observed in NRAS mutated cancers. Correlation between mutant burden and gene copy gains have already been described for KRAS [36] and BRAF gene [20]. To our knowledge, NRAS/chromosome 1 copy number variations have never been described in melanomas with regard to the NRAS mutant allele burden.

Secondly, we analyzed chromosome 1 microsatellite polymorphism in normal and tumor DNA in a group of 29 NRAS mutated cancers by ALFP method. As expected, LOH with WT allele loss was mostly restricted to the High non-HET M%NRAS group. However, unlike in haematological malignancy [37, 38], acquired copy-neutral LOH was not a predominant mechanism of mutant allele imbalance in NRAS Q61 mutated melanomas; indeed this aberration was detected in only 23% of our cases. Other mechanisms of High non-HET M%NRAS were amplification and gain of NRAS gene (14%) and polysomy of chromosome 1 (23.8%). In 9 melanomas (38.1%), an intra-tumor copy number variation of NRAS/chromosome 1 was detected. As most melanomas have copy number variations of whole chromosomes and of chromosome segment, NRAS mutant allele increase could be a consequence of chromosome instability and clonality in these tumors.

Correlation of M%NRAS with clinical data revealed no association with age, sex, histological melanoma subtypes, nor with histoprognostic markers of the patients with NRAS Q61 mutated melanomas. Moreover, no differences in patient survival outcomes were observed between patients with <60% and ≥60% of M%NRAS. However, this cohort is a retrospective monocentric study, and the analyses were limited by small number of patients. Furthermore, the value of M%NRAS has to be investigated for prediction of response to targeted therapy, as done for M%NRAS with promising results. We hypothesize that High non-HET M%NRAS could have an oncogenic addiction effect, which could improve the sensitivity of targeted therapy in this subgroup of NRAS Q61 mutated melanoma.

**Conclusion**

We report herein for the first time that 30% of cutaneous NRAS mutant melanomas have a high M%NRAS. Chromosome instability, (chromosome 1 polysomy, intra-tumor copy number variation of chromosome1/NRAS) rather than the acquired copy neutral LOH seems to be responsible for most of the cases with high M%NRAS. Histoprognostic markers and survivals were not different when comparing patients with <60% and ≥60% of M%NRAS; however this should be checked in a larger and multicentric series.

**Additional files**

**Additional file 1: Figure S1.** Identification of NRAS WT and different NRAS mutations. Sequence of NRAS wild type allele (A) and of different NRAS mutant alleles and the corresponding pyrosequencing profiles. Pyrosequencing profiles by assays 1 (A, B, D, E) and by assay 2 (C and F) are present (TIFF 178 kb)

**Additional file 2: Figure S2.** Flow chart. Flow chart for melanoma samples regarding the NRAS gene status and the type of molecular analysis carried out (TIFF 170 kb)

**Additional file 3: Figure S3.** NRAS Q61 mutant allele burden in ATGC melanomas. Histogram representation of NRAS Q61 mutant allele quantity (in percentage) in 85 ATGC NRAS mutated melanomas. The X and Y axis correspond to the percentage of NRAS mutant and to the number of cases, respectively (TIFF 102 kb)

**Abbreviations**

ALFP: Amplified fragment length polymorphism; DMFS: Distant metastasis free survival; FISH: Fluorescence in situ hybridization; HET: Heterozygous; LOH: Loss of heterozygosity; M%NRAS: NRAS mutant allele frequency; OS: Overall survival; PFS: Progression-free survival

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**Availability of data materials**

Please contact the corresponding author for all data requests.

**Authors’ contributions**

Conception and design: ZHR, EFB, JFE; Development of the methodology, Acquisition of data: ZHR, EFB, NT, CM, PS, JFE; Writing and revision of the manuscript: ZHR, EFB, JFE; Study supervision: JEF, PS. All authors have read and approved the final manuscript.

**Ethical approval and consent to participate**

The research was performed in compliance with the ethical principles of the Helsinki Declaration (1964) and with the French ethics laws. Patients were informed and approved the use of their samples for research purpose. Tumor samples collection was declared to the French Ministry of Research (DC 2009–933). Melan-Cohort study was approved by CPP IDF 8 Ethics committee (030209) and registered with https://www.clinicaltrials.gov/ct2/search (NCT00839410).

**Consent for publication**

Not applicable

**Competing interests**

JFE. received honoraria from Roche and Glaxo Smith Kline for counseling on patients with melanomas on the diagnosis and/or treatment with BRAF
inhibitors. P.S. received honoraria for counseling on diagnosis and/or treatment of patients with melanomas from Roche and Glaxo Smith Kline.

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