Flexible Lab-Tailored Cut-Offs for Suitability of Formalin-Fixed Tumor Samples for Diagnostic Mutational Analyses

Sara Mariani1, Cristiana Di Bello3*, Lisa Bonello3*, Fabrizio Tondat3, Donatella Pacchioni2, Luca Molinaro2, Antonella Barreca1, Luigia Macrì2, Luigi Chiusa2, Paola Francia di Celle2, Paola Cassoni1, Anna Sapino1*

1 Department of Medical Sciences; University of Torino, Torino, Italy, 2 Azienda Ospedaliera Universitaria Città della Salute e della Scienza; Presidio Ospedaliero Molinette of Torino, Torino, Italy, 3 Department of Molecular Biotechnology and Health Sciences, University of Torino, Torino, Italy

☯ These authors contributed equally to this work.
* anna.sapino@unito.it

Abstract

The selection of proper tissues from formalin-fixed and paraffin-embedded tumors before diagnostic molecular testing is a responsibility of the pathologist and represents a crucial step to produce reliable test results. The international guidelines suggest two cut-offs, one for the percentage and one for the number of tumor cells, in order to enrich the tumor content before DNA extraction. The aim of the present work was two-fold: to evaluate to what extent a low percentage or absolute number of tumor cells can be qualified for somatic mutation testing; and to determine how assay sensitivities can guide pathologists towards a better definition of morphology-based adequacy cut-offs. We tested 1797 tumor specimens from melanomas, colorectal and lung adenocarcinomas. Respectively, their BRAF, K-RAS and EGFR genes were analyzed at specific exons by mutation-enriched PCR, pyrosequencing, direct sequencing and real-time PCR methods. We demonstrate that poorly cellular specimens do not modify the frequency distribution of either mutated or wild-type DNA samples nor that of specific mutations. This observation suggests that currently recommended cut-offs for adequacy of specimens to be processed for molecular assays seem to be too much stringent in a laboratory context that performs highly sensitive routine analytical methods. In conclusion, new cut-offs are needed based on test sensitivities and documented tumor heterogeneity.

Introduction

The identification of biologically active genes and pathways that are disrupted in various cancer types has led to the development of clinically relevant diagnostic requirements. Several oncogenes have become therapeutic targets [1–3]. Thus, oncology is rapidly moving towards an era of precision medicine, where patients receive tailored treatments based on the biology and the
The genetic makeup of their tumors [4]. In the last few years, the Food and Drugs Administration (FDA), the European Medicines Agency (EMA) and the Italian Medicines Agency (AIFA) have introduced many biological drugs in therapeutic schemes of various cancers. Among others, anti-epidermal growth factor receptor (anti-EGFR) monoclonal antibodies (Cetuximab and Panitumumab) were approved for the treatment of wild-type RAS metastatic colorectal cancers (mCRCs) and anti-tyrosine kinases (Erlotinib and Vemurafenib) were admitted in therapeutic schemes to treat EGFR mutated non-small cell lung cancers (NSCLCs) and BRAF-mutated metastatic melanomas (mMEls), respectively. The conditions of their clinical uses were determined by specific trials (CRYSTAL, IPASS, BRIM-1, PEAK and PRIME), designed on the basis of results from retrospective studies [5–8].

The development of such targeted therapies has contributed to establishing a strong collaboration between pathologists and molecular biologists to provide clinicians with the most accurate results. The histo-morphological revision and the selection of formalin-fixed and paraffin-embedded (FFPE) tumor tissues are essential prerequisites of the analytical phase to allow discrimination of patient eligibility for targeted therapies. The Italian guidelines suggest that for mutational analyses tissue samples should contain an absolute number of at least 100 tumor cells and a tumor cell percentage representation over 50% (https://testbiomolecolari.it/). The suitability of tumor specimens with low percentage and/or absolute number of cancer cells represents a grey zone for molecular analyses, still debated within the scientific community [9–12]. The present work focuses on DNA sequencing analyses applied to patient specimens with a poor tumor cellularity. The frequency distributions of mutated and wild-type DNA sequences in hotspot regions of K-RAS, BRAF and EGFR genes were compared in a large series of morphologically acceptable and not acceptable samples in order to understand the relationship between morphology-driven tissue selection and the pertinence to molecular biology analyses in the era of highly sensitive sequencing methods.

Materials and Methods

Patients

This study was performed on FFPE tumor samples subjected to DNA extraction and sequencing. Tissue specimens from oncologic patients were collected at our institution, which is one of the regional reference laboratories for the study of somatic mutations as predictive markers of response to targeted therapy-based regimens. The case series included the tumor tissues of 1084 mCRCs, 572 never-, low- or ex-smoking NSCLC patients at stages IIIb and IV, and 141 mMEls or inoperable melanomas at stages IIIc and IV. Cytological specimens and histological surgical samples were 190 and 1607, respectively. Fine-needle aspiration biopsies (FNAB) and brushing techniques were the most commonly used methods for cytological sampling. The paraffin-embedded samples had been stored for a variable period of time (from few months up to eight years) before molecular analyses were performed.

The study was submitted to and approved by the ethic institutional review board for “Biobanking and use of human tissues for experimental studies” of the Pathology Service of the Azienda Ospedaliera Universitaria Città della Salute e della Scienza di Torino, Torino, Italy. The project provided a verbal and not written informed consent from patients due to the retrospective approach of the study, which did not impact on their treatment. All the cases were anonymously recorded. The IRB approved this consent procedure.

Material selection criteria

The selection of the tumor area was performed on Hematoxylin and Eosin (H&E) archival slides originally used for cytological examinations or histo-pathological diagnosis, classification
and grading, according to World Health Organization criteria. On the basis of clonal evolution trends, primitive or relapsed tumors or synchronous distant metastases were preferred in mCRCs, NSCLCs and mMELs [13–15]. The pathologist marked the areas of tumor enrichment on the slide from either cytological or histological specimens and recorded the percentage and absolute number of cancer cells in the request form (see below). Accordingly, five serial sections were cut from the ethanol-fixed or FFPE blocks. The first four sections were 10 μm-thick and were collected in two sterile tubes (Eppendorf, Milano, Italy) for DNA extraction. The last section, 3 μm-thick and H&E-stained, was used to confirm the correspondence with the original marked areas and to further check the presence of tumor cells in the samples.

The criteria of morphological suitability for performing DNA mutational analyses were derived from the Biogate portal (https://testbiomolecolari.it/) of the Association of Italian Medical Oncologists (AIOM). In particular, specimens satisfying the guideline criteria were recorded as acceptable (A), whereas specimens submitted to tumor enrichment but with less than 50% (70% until 2011) or 100 tumor cells had to be considered as not acceptable (NA). We named these specimens NA1 and NA2, respectively.

DNA extraction

Enriched tumor sections from NA and A specimens were submitted to DNA extraction (Maxwell16 FFPE Tissue LEV DNA Purification Kit, Promega, Madison, WI, USA) on a Maxwell16 Instrument (Promega, Madison, WI, USA). DNA was eluted in a final volume of 50 μL, and the concentrations/purity was measured by a Nanodrop 1000 spectrophotometer (Thermo Fisher Scientific, Wilmington, DE, USA).

Restriction enzyme-mediated selective polymerase chain reaction (REMS-PCR) for the analysis of the K-RAS gene mutations in codon 12

The K-RAS procedure was optimized by Fuery et al. [16]. Due to its high sensitivity, close to 0.1%, the assay lets to investigate nucleotide substitutions affecting even low copies of genomic DNA at the codon 12. REMS-PCR was assessed as previously described [17] with 120 ng of patient and control DNA in a final volume of 25 μL. The original protocols were modified as follow: BstNI was substituted with 30 U of the more thermo stable PspGI restriction endonuclease (EuroClone, Pero, Milano, Italy), and the reverse primer (DIA) of the K-RAS codon 12 was changed to avoid the amplification of the K-RAS pseudogene (the new sequence is TTTACCTCTATTGTTGGATCATATTC). The amplified samples were visualized by electrophoresis through a Criterion Tris-HCl precast 10% polyacrylamide gel (Biorad, Segrate, Milano, Italy). The results were interpreted as reported by Mixich et al. [17]. The reverse DIA primer changed the size of the expected K-RAS mutated amplicons from 82 to 107 bp.

Mutational analysis of the K-RAS and BRAF genes by Pyrosequencing

K-RAS codons 12 and 13 or BRAF codon 600 mutations were detected by Pyrosequencing [18] in mCRC or mMEL samples according to the manufacturer’s instructions (Diatech Pharmacogenetics, Jesi, Italy). Briefly, DNA from patients and controls were amplified in a Rotor-Gene Q 2plex HRM instrument (Qiagen, Manchester, United Kingdom) by specific PCR protocols (Anti-EGFR MoAb Response-KRAS status or Anti-EGFR MoAb Response-BRAF status). The endpoint PCR products were sequenced in a PYROMARK Q96 instrument and analyzed according to the manufacturer’s instructions.
Mutational analysis of the K-RAS, BRAF and EGFR genes by Sanger Sequencing

The Sanger Sequencing method [19] was applied to samples with doubtful K-RAS and BRAF analyses after Pyrosequencing, and as reference test for the analysis of mutations at the exons 18–21 of the EGFR gene.

To amplify K-RAS or BRAF, 70 ng of genomic DNA were mixed in 50 μL to 1x buffer, 1.5 mM of MgCl₂, 200 μM of dNTP, 1.2 U of AmpliTaq Gold DNA Polymerase (Invitrogen, Carlsbad, CA) and 0.5 μM of primers. The K-RAS primers (GTACTGGTGAGTAGTATTGTAATGTGT and TTTACCTCTATTTGGGATCATATTC) were designed using the software Primer3 (http://frodo.wi.mit.edu/primer3/) on the human K-RAS genomic sequence (http://www.ncbi.nlm.nih.gov/gene) (ID: 3845) and produced amplicons 212 bp long. The BRAF primers were derived from Moroni et al. [20], generating amplicons 228 bp.

To amplify exons 18–21 from EGFR, 35 ng of genomic DNA were mixed in 25 μL with the same reagents previously described, except for the substitution of the AmpliTaq Gold DNA Polymerase with 0.5 U of Platinum Taq DNA Polymerase (Invitrogen, Carlsbad, CA). The primer concentrations were 0.4 μM, and their sequences were defined in accordance with Lynch TJ et al. [21]. The exon-specific PCR products were 400, 372, 408 and 415 bp long.

PCR conditions consisted of a unique touchdown scheme (an annealing decrease of 3°C each 3 cycles starting from 64°C up to 57°C) that was repeated for 44 cycles.

The PCR products were then purified following the manufacturer’s instructions (AMPure PCR Purification Kit protocol, Agencourt, Beckman Coulter, Brea, CA, USA) and submitted to cycle sequencing reactions (BigDye Terminator v1.1 Sequencing kit, Applied Biosystems, Foster City, CA, USA) in the presence of 0.1 μM of forward and reverse primers (as well as for the PCRs), separately. Capillary electrophoresis of purified sequences (CleanSeq System kit, Agencourt, Beckman Coulter, Brea, CA, USA) was performed at 60°C through a POP-7 polymer in a 3130 four-capillary sequencer (Applied Biosystems, Foster City, CA, USA). The Sequencing platform (Applied Biosystems, Foster City, CA, USA) was used to acquire samples, which were then analyzed with the Variant Reporter software (Applied Biosystems, Foster City, CA, USA).

Mutational analysis of the EGFR gene by Therascreen

The Therascreen EGFR RGQ PCR kit detects 29 somatic mutations frequently occurring within exons 18–21 of the EGFR gene. This PCR kit applies the Scorpions and ARMS technologies to a real-time PCR running in a Rotor-Gene Q 2plex HRM instrument (Qiagen, Manchester, United Kingdom).

PCR conditions and criteria of interpretation of the results were adopted according to the manufacturer’s protocol (Qiagen, Manchester, United Kingdom). Briefly, samples with a cycle threshold (Cₚ) of the HEX control signal ≥ 37 were rejected, and samples with a Cₚ of FAM mutation signal ≥ 40 were scored as negative or indeterminate, as they were below the sensitivity limit of the assay. To define EGFR positivity, specific ranges of acceptable ΔCₚ values were defined in the manual.

Sensitivities of the assays

Home-made reactions were designed to determine the sensitivities of the assays by using mutated DNA from specific cell lines progressively diluted in blood-derived DNA of a healthy donor. Briefly, the A549 cell line was homozygous for the K-RAS p.Gly12Ser mutation, the HT29 was heterozygous for the BRAF p.Val600Glu mutation, the H1650 was heterozygous for the EGFR deletion at the exon 19 (p.Glu746_Ala750del), and the H1975 was heterozygous for...
the EGFR p.Leu858Arg mutation at the exon 21. The cell lines were purchased from the ATCC (LGC Standards S.r.l., Sesto San Giovanni, Milan, Italy). K-RAS mutations were detected by REMS-PCR, Pyrosequencing and Sanger Sequencing. Their sensitivities for heterozygous mutations were near 0.1%, 5% and 20%, respectively. BRAF mutations were detected by Pyrosequencing or Sanger Sequencing at sensitivities near 5%. EGFR mutations were detected by Sanger Sequencing up to 6% of sensitivity (data not shown). As reported (Therascreen protocol, Qiagen, Manchester, United Kingdom), the sensitivities of the Therascreen assay for the EGFR mutations ranged from 0.5% (p.Leu861Gln) to 7.02% (p.Thr79Met).

External Quality Assurance (EQA)

As reference laboratory for the diagnostic assessment of mutational screening in solid tumors since 2010, we have been participants in several EQA schemes of the European Society of Pathology (ESP) for K-RAS (four times) and EGFR (one time) mutations and of the AIOM-Italian Society of Pathological Anatomy and Cytology schemes (SIAPEC) for K-RAS (one time), BRAF (one time) and EGFR (two times). We successfully passed all the EQA procedures (data not shown).

Statistics

Pearson’s Chi square test (with the application of a Yates’s correction when required) or Fisher’s exact test were alternatively applied to Contingency 2x2 Tables comparing mutation frequencies or tissue characteristics in samples that were classified acceptable and not acceptable on the basis of morphologically defined tumor cell contents (www.openepi.com). The Chi square P value was calculated on a 2x3 Contingency Table to compare the frequency of mutation variants in mutated samples belonging to different categories of patients. Significance was set at P<0.05.

Results

Frequency of NA specimens

From January 2010 to August 2013, 1797 FFPE tumor samples, derived from patients with mCRC, mMEL and NSCLC were selected to detect K-RAS, BRAF and EGFR somatic mutations, respectively. All samples were submitted to DNA sequence analyses irrespective of their A or NA status.

First, we compared the frequency of NA in the mCRC and NSCLC specimens that were analyzed with different morphological criteria of unsuitability (in 2010, the tumor cell cut-off was 70%, but from 2011–2013 the tumor cell cut-off was 50%). The ratio for NA tumor specimens was 10/332 and 41/752 (3.0% vs 5.4%) in mCRC samples and 15/74 and 106/498 (21.3% vs 20.3%) in NSCL tumor specimens within the 2010 and 2011–2013 periods, respectively. The difference was not statistically significant (p>0.05). Therefore, we evaluated as a single cohort the specimens and the results collected from 2010–2013.

As shown in Fig 1, the NA tumor specimens were significantly more frequent (121/572) (21.1%) in NSCLCs than in mCRCs (51/1084) (4.7%) and in mMELs (7/141) (5.0%) (p<0.01).

Frequency of NA specimens in cytological and histological tumor samples

DNA was extracted from 190 (10.6%) cell blocks (from cytological sampling) and 1607 (89.4%) tissue blocks (biopsy or surgical specimens). Cytological specimens represented 1.3% of samples in mCRCs, 2.1% in mMELs and 30.2% in NSCLCs (p<0.0001). The frequencies of NA
specimens in cytological and histological samples were 14.3% vs 4.6% in mCRCs, 33.3% vs 4.3% in mMELs and 35.3% vs 15.0% in NSCL tumors (Table 1), respectively. The frequency of the NA samples was significantly higher in cytological specimens of NSCLC (p<0.01).

Distribution of wild-type and mutated K-RAS gene sequences among NA and A specimens from mCRC patients

The main purpose of this work was to establish if NA and A specimens, derived from stringent criteria of microscopically revision of tissue sections, corresponded to a statistically significant different number of wild-type (WT) and mutated (MUT) samples for the genes analyzed. As the analytical sensitivities were different for each sequenced gene (see materials and methods), we separately calculated the frequencies of WT and MUT samples in NA and A specimens belonging to the three different diseases.

Table 1. Frequencies of acceptable and not acceptable tumor samples in cytological and histological specimens from mCRC, mMEL and NSCLC patients.

|                  | CYTLOGICAL SPECIMENS (n = 190) | HISTOLOGICAL SPECIMENS (n = 1607) |
|------------------|---------------------------------|-----------------------------------|
|                  | n (%)                           | n (%)                             |
| mCRCs (n = 1084) |                                 |                                   |
| A                | 12 (85.7)                       | 1021 (95.4)                       |
| NA               | 2 (14.3)                        | 49 (4.6)                          |
| mMELs (n = 141)  |                                 |                                   |
| A                | 2 (66.7)                        | 132 (95.7)                        |
| NA               | 1 (33.3)                        | 6 (4.3)                           |
| NSCLCs* (n = 572)|                                 |                                   |
| A                | 112 (64.7)                      | 339 (85.0)                        |
| NA               | 61 (35.3)                       | 60 (15.0)                         |

A: Acceptable samples
NA: Not acceptable samples

*p<0.01
Fig 2A shows the frequencies of WT and MUT samples in NA and A specimens of mCRC patients. The number of WT and MUT specimens were 28 and 23 out of 51 (54.9% and 45.1%, respectively) in the NA samples and 575 and 458 out of 1033 (55.7% and 44.3%, respectively) in the A samples. In NA1 samples WT and MUT specimens were 20 and 18 out of 38 (52.6% and 47.4%, respectively) and in NA2 samples 8 and 5 out of 13 (61.5% and 38.5%, respectively) (Fig 2B). The NA2 group showed a higher percentage (not statistically significant) of WT specimens than both the NA1 or the A group.

Three of the NA samples had a second acceptable (A) specimen available. One of these cases, which contained less than 10% tumor cells and resulted in WT K-RAS, was MUT in the A specimen.

Distribution of wild-type and mutated BRAF gene sequences among NA and A specimens from mMEL patients

As shown in Fig 2C, in the 7 NA samples the number of WT and MUT specimens was 2 and 5 (28.6% and 71.4%, respectively), and in the 134 A samples the WT and MUT distribution was 59 and 75 (44.0% and 56.0%, respectively). In the NA1 samples WT and MUT specimens were 20 and 18 out of 38 (52.6% and 47.4%, respectively) and in NA2 samples 8 and 5 out of 13 (61.5% and 38.5%, respectively) (Fig 2B). The NA2 group showed a higher percentage (not statistically significant) of WT specimens than both the NA1 or the A group.

Distribution of wild-type and mutated EGFR gene sequences among NA and A specimens from NSCL adenocarcinoma patients

In the 121 NA samples, the number of WT and MUT specimens was 98 and 23 (81.0% and 19.0%, respectively) and in 451 A samples was 369 and 82 (81.8% and 18.2%, respectively) (Fig 2D). The number of WT and MUT specimens was 88 and 21 out of 109 (80.7% and 19.3%, respectively) in the NA1 samples and 10 and 2 out of 12 (83.3% and 16.7%, respectively) in the NA2 samples (Fig 2E) (all differences statistically not significant). In 8 NA samples, two of which (with less than 5% and 100 tumor cells) EGFR MUT, the analysis performed in a second A specimen confirmed the 2 MUT and the 6 WT (data not shown).
Distribution of K-RAS and EGFR mutations between NA and A samples compared to results collected from published datasets

We measured the frequency of the principal mutation variants occurring at the codon 12 of K-RAS and at the exons 18, 19, 20 and 21 of EGFR among NA and A samples and compared the results to specific online databases, as reported below.

The mutation frequencies related to codon 12 of K-RAS within our dataset were compared to the Cosmic database (Sanger Institute database; [www.sanger.ac.uk/genetics/CGP/cosmic/](http://www.sanger.ac.uk/genetics/CGP/cosmic/)) and were calculated as number of specimens with each specific mutation divided by the totality of the mutated specimens (Cosmic n = 12489, NA n = 21 and A n = 365). The frequencies are shown in Fig 3A and reported in Table 2. To obtain a Chi square P value, a 2x3 Contingency Table was applied to compare the frequency of each mutation variant (measured on the total mutations) in the three groups, with the exception of the “CGT” mutated samples, which did not meet the Cochrane recommendations [22] and to which a Fisher’s exact test was applied in a classical 2x2 Contingency Table. The mutation frequencies analyzed within the NA, A and database samples did not result in statistically significant differences (Fig 3A).

The distribution of the EGFR gene mutations occurring at the exons 18, 19, 20 and 21 in the NA, A and online dataset samples (obtained from the NSCL SM-EGFR-database, [www.somaticmutations-egfr.org/overall_mutation_frequency.html](http://www.somaticmutations-egfr.org/overall_mutation_frequency.html)) is shown in Fig 3B and reported in Table 3. Because the EGFR mutations were too complex for detailed analysis, we grouped them on the basis of the exons involved. The frequency was calculated as the number of specimens with each exon-specific mutation, divided for the sum of the mutated specimens (SM-EGFR-database: 3303 cases, NA:23 cases and A:82 cases). In some samples from our dataset, multiple coexisting mutations were found. The different frequencies of samples with each mutation among the three categories were not statistically significant, except for exon 20,

![Graph A](https://example.com/graphA.png)

*Fig 3A. Specific somatic mutation frequencies in samples morphologically acceptable, not-acceptable and derived from published data. The frequencies of each specific mutation at codon 12 of K-RAS in not acceptable (NA) or acceptable (A) mCRC tumor specimens compared to the corresponding online dataset (Database Cosmic) were not statistically different (p>.05). Mutation frequencies at the exons 18–21 of EGFR were statistically different only for the exon 20 between NA or A specimens versus the online dataset (B). X-axis: specific K-RAS or EGFR mutations, divided into NA, A and the online dataset samples; Y-axis: frequency distributions of mutated samples.*

* doi:10.1371/journal.pone.0121815.g003

![Graph B](https://example.com/graphB.png)
which was highly mutated in both the NA and A specimens compared to the online dataset group (p<0.002) (Fig 3B).

**Discussion**

In the present work we demonstrated that the frequency of mutated and wild-type DNA sequences in hotspot regions of K-RAS, EGFR and BRAF genes were poorly affected by the current definition of cell or tissue sample suitability. In particular, by analyzing the frequencies of activating mutations and wild-type gene sequences in a total number of 179 NA and 1618 A specimens grouped for tumor pathology, we demonstrated that: 1) the cytological specimens derived from bronco-aspirate in the NSCLC set were more frequently NA due to hypocellularity; 2) the proportion of NA specimens is not modified by lowering the cut-off from 70% to 50%; 3) the frequencies of WT and MUT sequences were similar between NA and A specimens in every patient group, and neither the percentages nor the number of tumor cells present in the samples affected the WT frequency; and 4) the frequency of mutation variants occurring at specific hotspots (codon 12 of K-RAS and exons 18 to 21 of EGFR) was equivalent in NA and A samples.

Focusing on the NA1 category, which refers to the percentage of tumor cells, the cut-off value of 70% of tumor cells per section, than lowered to 50%, was originally set based on the sensitivity of the direct sequencing, generally reaching 25%-30% to detect homozygous mutation pattern (50–60% for heterozygous genotypes) [23, 24]. In the present study, we could not establish an appropriate lab-tailored cut-off, because the percentage of tumor cells present in each sample was reported as within or below the cut-off and not as defined number.

### Table 2. Frequency distribution of single point mutations occurring at codon 12 of K-RAS in morphologically not acceptable, acceptable and dataset mCRC-derived tumor specimens.

| MUTATED mCRC SAMPLES | AGT | CGT | GAT | GCT | GTT | TGT |
|----------------------|-----|-----|-----|-----|-----|-----|
| **NA** (n = 21)      | 2   | 1   | 8   | 2   | 7   | 1   |
| **A** (n = 365)      | 34  | 3   | 152 | 40  | 99  | 37  |
| **COSMIC DB** (n = 12489) | 930 | 193 | 5582 | 1002 | 3499 | 1283 |

NA = Not acceptable samples  
A = Acceptable samples  
COSMIC DB = Cosmic database of somatic mutations

doi:10.1371/journal.pone.0121815.t002

### Table 3. Frequency distributions of mutations within exons 18–21 of EGFR in morphologically not acceptable, acceptable and dataset NSCLC-derived tumor specimens.

| MUTATED NSCLC SAMPLES | EXON 18 | EXON 19 | EXON 20 | EXON 21 |
|-----------------------|--------|--------|--------|--------|
| **NA** (n = 23)       | 1      | 9      | 5      | 9      |
| **A** (n = 82)        | 5      | 49     | 10     | 31     |
| **NSCL SM-EGFR DB** (n = 3303) | 137   | 1662   | 213    | 1291   |

NA = Not acceptable samples  
A = Acceptable samples  
NSCL SM-EGFR DB = Database of EGFR somatic mutations from NSCL cancer patients

doi:10.1371/journal.pone.0121815.t003
Nevertheless, when the appropriate size of NA specimens in each pathological patient group is achieved, it can be subdivided in five categories of decreasing tumor enrichment (from 50% to 0%). Therefore, the distribution of gene mutations between each category and the A series becomes comparable. Lab-tailored cut-offs will then be comprised in the last category of tumor enrichment behaving as the adequate samples. Although our lab-tailored cut-off was not yet determined in the present study, mutated specimens in the NA1 series most likely contain a percentage of mutant DNA cells higher than 20%, corresponding to the minimal rate detectable by our less sensitive assay (Sanger Sequencing) applied to tumor DNA heterozygous for the examined mutation. New generation assays with higher sensitivity further reduces the need for high percentages of tumor cells.

The cut-off issue is further complicated by the occurrence of intrinsic tumor heterogeneity. In this regard, Yu and co-authors [10] demonstrated that the percentage of K-RAS mutant DNA in enriched tumor areas from FFPE blocks of 47 mCRC showed a continuous distribution from 10.8% (high heterogeneity) to 98.3% (low heterogeneity). Assuming, for example, a sensitivity limit of 5% to detect K-RAS mutations by Pyrosequencing, the starting percentage of morphologically enriched tumor cells should vary between 46.3% and 5.1% for tumors highly or barely heterogeneous for the tested mutation, respectively. Thus, the tumor enrichment required before molecular biology testing should be directly proportional to the tumor heterogeneity for mutant alleles and inversely proportional to the assay sensitivities. Since tumor heterogeneity varies among patient samples, histology-driven tissue multisampling combined with cell pooling and cell sorting techniques should be applied to a large series of specimens upstream, in order to define the best morphological cut-off in relation to both the extent of tumor heterogeneity and the specific analytical methods used for mutational testing. We cannot exclude the possibility that in our series few NA specimens presented tumor heterogeneity or whenever homogeneous for the mutations analyzed, their tumor abundance was below the detection level of the assays. Nevertheless, our data suggest that the number of these specimens was low enough to maintain unchanged the overall results. One example is the NA mCRC specimen, which resulted in a false WT when compared with an A sample derived from the same patient three years later. In this case, tumor heterogeneity or clonal evolution [25] cannot be excluded. Conversely, in the NSCLC group one NA1 specimen coupled with its A counterpart was truly mutated, although the former contained less than 5% enriched tumor cells, in line with the sensitivity cut-off level of the assay used and with a presumed complete homogeneity of cancer cells.

About the NA2 cut-off, the amount of genomic DNA corresponding to 100 diploid tumor cells in G1 phase is comprised between 500 and 1000 pg, as one cell contains near to 7 pg of DNA [26]. This range is slightly below the optimal input of good quality DNA suggested to obtain PCR products reliable for forensic genotyping [27]. The results obtained in the present study show that the collection of more tissue sections in specimens defined as NA2 (less than 100 tumor cells per section) may be a good solution for recovering a sample suitability for molecular analysis.

Finally, we considered the test sensitivity, which may highly impact on the overall results, showing equivalent distributions of the mutation variants between NA and A samples. A significantly higher percentage of NSCLCs that contain a mutated exon 20 in EGFR was found in our series when compared to the specimens belonging to the lung database collection. Although we cannot exclude the hypothesis of different assay sensitivities for that mutation, we have also considered a bias in the data collection between our series and the published results, or the coexistence of multiple mutations as demonstrated in some specimens of our series, which may have been less prevalent or detectable in the online lung dataset.
In conclusion, the molecular methods recently introduced in diagnostic practice are more sensitive than in the past. These new procedures may need less strict pre-analytical purification of the tumor tissue areas in order to identify the attended mutations. Our data demonstrate that the frequency of mutated or wild-type specimens is identical in samples within and below the international cut-offs of suitability. This observation suggests extending the DNA sequencing analyses also to the NA specimens, thus leaving the final definition of true positive (mutated) or putative false negative (wild-type) cases to the "post-analytical" phase. New and more flexible lab-tailored cut-offs need to be considered, depending on the molecular biology methods used and the extent of the tumor heterogeneity.

**Author Contributions**

Conceived and designed the experiments: SM. Performed the experiments: CDB LB FT. Analyzed the data: SM PFDC. Wrote the paper: SM PC AS. Made histo-pathological diagnoses and determined the morphologically-based adequacy of the specimens: DP L. Molinaro AB L. Macri LC.

**References**

1. Marks JL, Broderick S, Zhou Q, Chitale D, Li AR, Zakowsky MF, et al. Prognostic and therapeutic implications of EGFR and KRAS mutations in resected lung adenocarcinoma. J Thorac Oncol. 2008; 3: 111–116. doi: 10.1097/JTO.0b013e318160c607 PMID: 18303429

2. Dancey JE, Bedard PL, Onetto N, Hudson TJ. The genetic basis for cancer treatment decisions. Cell. 2012; 148: 409–420. doi: 10.1016/j.cell.2012.01.014 PMID: 22304912

3. Vucic EA, Thu KL, Robison K, Rybaczyk LA, Chari R, Alvarez CE, et al. Translating cancer ‘omics’ to improved outcomes. Genome Res. 2012; 22: 188–195. doi: 10.1101/gr.124354.111 PMID: 22301133

4. Awada A, Attimos PG. Targeted therapies of solid cancers: new options, new challenges. Curr Opin Oncol. 2013; 25: 296–304. doi: 10.1097/CCO.0b013e32835ff318 PMID: 23493919

5. Kaneda H, Tamura K, Kurata T, Uejima H, Nakagawa K, Fukuoka M, et al. Retrospective analysis of the predictive factors associated with the response and survival benefit of gefitinib in patients with advanced non-small-cell lung cancer. Lung Cancer. 2004; 46: 247–254. PMID: 15474673

6. Lee JT, Li L, Braddock PA, Halloran MB, Sproesser K, Haass NK, et al. PLX4032, a potent inhibitor of the B-Raf V600E oncoprotein, selectively inhibits V600E-positive melanomas. Pigment Cell Melanoma Res. 2010; 23: 820–827. doi: 10.1111/j.1755-148X.2010.00763.x PMID: 20973932

7. Schwartzberg LS, Rivera F, Karthaus M, Fasola G, Canon JL, Hecht JR, et al. PEAK: a randomized, multicenter phase II study of panitumumab plus modified fluorouracil, leucovorin, and oxaliplatin (mFOLFOX6) or bevazucimab plus mFOLFOX6 in patients with previously untreated, unresectable, wild-type KRAS exon 2 metastatic colorectal cancer. J Clin Oncol. 2014; 32: 2240–2247. doi: 10.1200/JCO.2013.53.2473 PMID: 2468633

8. Douillard JY, Siena S, Cassidy J, Tabernero J, Burkes R, Barigel M, et al. Randomized, phase III trial of panitumumab with infusional fluorouracil, leucovorin, and oxaliplatin (FOLFOX4) versus FOLFOX4 alone as first-line treatment in patients with previously untreated metastatic colorectal cancer: the PRIME study. J Clin Oncol. 2010; 28: 4697–4705. doi: 10.1200/JCO.2009.27.4860 PMID: 20921465

9. Smits AJ, Kummer JA, de Bruin PC, Bol M, van den Tweel JG, Seldenrijk KA, et al. The estimation of tumor cell percentage for molecular testing by pathologists is not accurate. Mod Pathol. 2014; 27: 168–174. doi: 10.1038/modpathol.2013.134 PMID: 23887293

10. Yu S, Xiao X, Lu J, Qian X, Liu B, Feng J. Colorectal cancer patients with low abundance of KRAS mutation may benefit from EGFR antibody therapy. PLoS One. 2013; 8:e68022. doi: 10.1371/journal.pone.0068022 PMID: 23874486

11. Hlinkova K, Babal P, Berzinec P, Majer I, Mikle-Barathova Z, Piacova B, et al. Evaluation of 2-year experience with EGFR mutation analysis of small diagnostic samples. Diagn Mol Pathol. 2013; 22: 70–75. doi: 10.1097/PPM.0b013e31827e6984 PMID: 23628817

12. Lewandowska MA, Jozwicki W, Zurawski B. KRAS and BRAF mutation analysis in colorectal adenocarcinoma specimens with a low percentage of tumor cells. Mol Diagn Ther. 2013; 17: 193–203. doi: 10.1007/s40291-013-0025-6 PMID: 23606169
13. Jones S, Chen WD, Parmigiani G, Diehl F, Beerenwinkel N, Antal T, et al. Comparative lesion sequencing provides insights into tumor evolution. Proc Natl Acad Sci USA. 2008; 105: 4283–4288. doi: 10.1073/pnas.0712345105 PMID: 18337506

14. Sun L, Zhang Q, Luan H, Zhan Z, Wang C, Sun B. Comparison of KRAS and EGFR gene status between primary non-small cell lung cancer and local lymph node metastases: implications for clinical practice. J Exp Clin Cancer Res. 2011 Mar 17; 30:30. Available: http://www.jeccr.com/content/30/1/30. doi: 10.1038/jecr.2013.622 PMID: 24196789

15. Heinzerling L, Bailer M, Kuhnapfel S, Schuler G, Keikavoussi P, Agaimy A, et al. Mutation landscape in melanoma patients clinical implications of heterogeneity of BRAF mutations. Br J Cancer. 2013; 109: 2833–2841. doi: 10.1038/bjc.2013.622 PMID: 24196789

16. Fuery CJ, Impey HL, Roberts NJ, Applegate TL, Ward RL, Hawkins NJ, et al. Detection of rare mutant alleles by restriction endonuclease-mediated selective-PCR: assay design and optimization. Clin Chem. 2000; 46: 620–624. PMID: 10794742

17. Mixich F, Ioana M, Voinea F, Saftoiu A, Ciurea T. Noninvasive detection through REMS-PCR technique of K-ras mutations in stool DNA of patients with colorectal cancer. J Gastrointestin Liver Dis. 2007; 16: 5–10. PMID: 17410282

18. Ogino S, Kawasaki T, Brahmandam M, Yan L, Cantor M, Namgyal C, et al. Sensitive sequencing method for KRAS mutation detection by Pyrosequencing. J Mol Diagn. 2005; 7: 413–421. PMID: 16049314

19. Sanger F, Nicklen S, Coulson AR. DNA sequencing with chain-terminating inhibitors. Proc Natl Acad Sci. 1977; 74: 5463–5467. PMID: 271968

20. Moroni M, Veronese S, Benvenuti S, Marrapese G, Sartore-Bianchi A, Di Nicolantonio F, et al. Gene copy number for epidermal growth factor receptor (EGFR) and clinical response to antiEGFR treatment in colorectal cancer: a cohort study. Lancet Oncol. 2005; 6: 279–286. PMID: 15863375

21. Lynch TJ, Bell DW, Sordella R, Gurubhagavatula S, Okimoto RA, Brannigan BW, et al. Activating mutations in the epidermal growth factor receptor underlying responsiveness of non-small-cell lung cancer to gefitinib. N Engl J Med. 2004; 350: 2129–2139. PMID: 15118073

22. Sneeden GW, Cochran WG. Statistical Methods. Iowa State, Ames, Iowa; 1980.

23. van Krieken JH, Jung A, Kirchner T, Carneiro F, Seruca R, Bosman FT, et al. KRAS mutation testing for predicting response to anti-EGFR therapy for colorectal carcinoma: proposal for an European quality assurance program. Virchows Arch. 2008; 453: 417–431. doi: 10.1007/s00428-008-0665-y PMID: 18802721

24. Jancik S, Drabek J, Berkovcova J, Xu YZ, Stankova M, Klein J, et al. A comparison of Direct sequencing, Pyrosequencing, High resolution melting analysis, TheraScreen DxS, and the K-ras StripAssay for detecting KRAS mutations in non small cell lung carcinomas. J Exp Clin Cancer Res. 2012; 31: 79–92. doi: 10.1186/1756-9966-31-79 PMID: 22995035

25. Vignot S, Soria JC. Discrepancies between primary tumor and metastasis: impact on personalized medicine. Bull Cancer. 2013; 100: 561–568. doi: 10.1684/bdc.2013.1767 PMID: 23728079

26. Ranek L. Cytophotometric studies of the DNA, nucleic acid and protein content of human liver cell nuclei. Acta Cytol. 1976; 20: 151–157. PMID: 59501

27. Kirkham A, Haley J, Haile Y, Groot A, Kimpton C, Al-Marzougi A, et al. High-throughput analysis using AmpFISTR Identifier with the Applied Biosystems 3500xl Genetic Analyser. Forensic Sci Int Genet. 2013; 7: 92–97 doi: 10.1016/j.fsigen.2012.07.003 PMID: 22835329