Why the Therapeutic Impact of RAS Mutation Clearance in Plasma ctDNA Deserves to Be Further Explored in Metastatic Colorectal Cancer

Chiara Nicolazzo 1, Francesca Belardinilli 2, Salvatore Caponnetto 3, Angela Gradilone 1, Enrico Cortesi 3, Giuseppe Giannini 2,4 and Paola Gazzaniga 1*

1 Liquid Biopsy Unit, Department of Molecular Medicine, Sapienza University of Rome, Rome, Italy, 2 Department of Molecular Medicine, Sapienza University of Rome, Rome, Italy, 3 Department of Radiological, Oncological and Pathological Sciences, Sapienza University of Rome, Rome, Italy, 4 Institute Pasteur-Cenci Bolognetti Foundation, Rome, Italy

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INTRODUCTION

An increasingly large body of evidence supports the assertion that the analysis of circulating tumor DNA (ctDNA) in metastatic colorectal cancer (mCRC) patients allows the temporal heterogeneity of cancer during the course of targeted therapies to be monitored. From liquid biopsy-guided genomic studies in mCRC, we have learned that there may be a rise in RAS mutant clones in the plasma of patients before the onset of secondary resistance to anti-EGFR therapy, generating new hypotheses for blood-guided adaptive therapeutic strategies (1). Similarly, the clearance of RAS mutant clones in plasma has been more recently suggested, supporting, for the first time, an unexpected negative selection of RAS mutant clones during the clonal evolution of mCRC (2–4). This phenomenon had previously been described in leukemia patients, with some showing a loss of RAS mutations during disease progression, supporting the hypothesis that the evolutionary pressure of therapies can result in positive but also negative selection of RAS mutant clones at relapse (5). The temporary prevalence of wt RAS clones at relapse in mCRC raises the question of whether liquid biopsy testing might expand the population of anti-EGFR-eligible patients by including those with primary RAS-mutant mCRC that “convert” to wild type in plasma at the time of disease progression (PD). Whether undetectable RAS mutations in plasma might really reflect wt RAS status or might simply mirror a low analytic sensitivity in the adopted assays is still a matter of debate (6).

CLEARANCE OF RAS MUTATION IN PLASMA: TUMOR-SPECIFIC MUTATIONS TO CONFIRM/EXCLUDE THE PRESENCE OF CTDNA

It is undeniable that correct interpretation of the clearance of RAS mutations in plasma must be supported by the demonstration of detectable ctDNA; therefore, the crux of the matter is the urgent need for appropriate methods to confirm (or to exclude) the presence of ctDNA in plasma samples. For this purpose, plausible options would be (1) to monitor a somatic mutation other than RAS that was previously detected in the primary tissue, (2) to provide evidence of wt RAS status in metastatic sites at the time of PD. To date, two groups have investigated the clearance of RAS mutations in plasma at the time of PD for therapeutic purposes in patients with primary RAS-mutant mCRC, even if in small sample series. The first study reported a clinical benefit achieved by some patients with RAS-mutant primary mCRC treated with anti-EGFR, based on detection of wt RAS status in plasma at the time of PD (2). In this series, 11
patients were screened via liquid biopsy at PD, and in 5/11 cases (45%) no RAS mutation was detected in plasma. Unfortunately, for only one patient did the authors provide evidence that the disappearance of RAS mutation was concomitantly associated with the persistence of one somatic mutation previously detected in the primary tumor. In the other four cases, the absence of any somatic mutation other than RAS that could be monitored in the blood led to “inconclusive” results. Globally, the real “RAS clearance rate,” calculated after excluding the inconclusive samples from the whole population, was 1/7 (14%). Owing to these preliminary data, a phase II prospective study was planned (7, 8) with the aim of investigating whether targeting the plasma wt RAS window with EGFR-inhibitors might represent on a large scale an exploitable second-line option in RAS-mutant mCRC. In 2019, a similar phase II prospective trial was aimed at evaluating the efficacy and safety of FOLFIRI plus panitumumab as a second-line treatment in mCRC patients with RAS mutan primary mCRC with no evidence of RAS mutation in plasma at the time of PD (9). Preliminary results have shown that of the 16 patients analyzed, only two had wt RAS mCRC after first-line treatment. Both were screen failures, so no patient has been included to date (9).

**CLEARANCE OF RAS MUTATION IN PLASMA: METHYLATION TO CONFIRM/EXCLUDE THE PRESENCE OF cTDNA**

The detection of tumor-specific DNA methylation alterations in ctDNA has recently been suggested as a specific tool for confirming the tumoral origin of ctDNA. Moati et al. (4) investigated the clearance of RAS mutant clones under chemotherapy pressure by ctDNA analysis in patients with RAS mutant mCRC. By monitoring ctDNA with methylated markers (WIFI and NPY genes), the authors concluded that the clearance of RAS mutations by chemotherapy is actually a rare event. Moati et al. define the clearance of RAS mutation as “the disappearance of RAS mutation concomitantly associated with the persistence of ctDNA proven by the detection of either (i) at least one mutation in other genes by NGS, (ii) or methylation of WIFI and NPY genes determined by ddPCR (met-ddPCR).” All samples with no mutation and no WIFI/NPY methylation were considered as “inconclusive.” In their study, Moati et al. described 8/36 (22%) patients who converted to a wt-RAS status in blood at the time of disease progression, but only in two patients was the presence of ctDNA in the sample confirmed by methylation test, while for the other six cases, the results were inconclusive. Thus, taking samples with inconclusive results out of the whole series, Moati et al. found a real switch from mutant to wt-RAS at progression in 2/30 patients (6.6%). Although an increasing amount of evidence has been provided that DNA methylation markers may be employed to track response during therapy in mCRC (10), the use of a WIF-1/NPY-specific methylation test to confirm the presence of ctDNA unfortunately relies on two proof of concept/exploratory studies, strongly impairing the generalization of their conclusions. Specifically, these two studies have evaluated WIF-1/NPY methylation dynamics in mCRC cases on treatment: (1) Garrigou et al. (11), who failed to detect methylation by liquid biopsy in 20% of mCRC, and (2) Garlan et al. (12), who, using the same test, showed only 69.2% positivity in KRAS/BRAF/TP53 wild-type mCRC. A more recent study (13), using high genome-coverage methods, demonstrated that other methylated loci are more compliant to liquid biopsy analyses, suggesting that their use in combination might improve the positivity of ctDNA detection. In this study, NPY was discarded due to positivity in normal mucosa and in whole blood samples (which contain white blood cells as the main contaminant of ctDNA), and WIFI was found to have limited differential methylation between normal healthy/normal adjacent mucosa and tumor. Other recently published studies found NPY methylation in the plasma of healthy donors (14) and non-cancer patients (15). Therefore, to date, we are far from true agreement on the use of WIF-1/NPY methylation testing to exclude or confirm the presence of ctDNA. Table 1 is a summary of the studies that have to date investigated the clearance rate of RAS mutations in plasma at the time of disease progression in patients with metastatic colorectal cancer.

**DISCUSSION**

To date, few studies have reported the clearance of RAS mutations in the plasma of mCRC patients. Comparing these studies, the clearance rate of RAS mutation was globally higher in Raimondi’s series, which could be consistent with the lower sensitivity of the method used (Idylla™, Biocartis), although no definitive conclusions can be drawn due to the small sample size of both studies. In this context, a recent comparison between the OncoBEAM RAS CRC mutation test and Idylla ctKRAS assay from paired plasma samples of mCRC patients identified a “gray zone” below a 1% mutant allele fraction (MAF) where Idylla shows reduced RAS mutation detection accuracy vs. OncoBEAM (16). On the other hand, an increased sensitivity of RAS testing under 1%, which is undoubtedly useful for monitoring the early onset of mutations, in this specific case might risk excluding from treatment patients with low MAF, who might benefit from anti-EGFR (17). In this respect, the use of an excessively sensitive method to detect RAS mutations in blood could represent one plausible explanation for the missed enrollment of patients in the Convertix trial. To sum up, our opinion is that the collective effort must be aimed at the search for one or more methods that allow the presence of ctDNA in plasma samples to be confirmed, thus decreasing the unacceptably high percentage of “inconclusive” results. Possibly, the use of NGS panels covering a broader spectrum of mutations (better if specific for colorectal cancer) will improve the detection rate of somatic mutations in tumor tissues to be longitudinally monitored in plasma. Nevertheless,
the few datasets that are available suggest that the percentage of patients with proven RAS metastatic colorectal cancers is approximately 4% (6.6% in the Italian and French study) respectively, and can be derived from the findings of the CIRCOS study. This study indicates that if we can treat a few patients, or even one patient, then personalized medicine will have achieved its main purpose.

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**AUTHOR CONTRIBUTIONS**

All authors listed have made a substantial, direct and intellectual contribution to the work, and approved it for publication.

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### TABLE 1 | Studies that investigated the clearance of RAS mutations in the plasma of metastatic colorectal cancer patients at the time of progression of disease.

| References                                      | N patients | ctDNA assay used | Patients with clearance of RAS mutations in plasma at PD (%) | Patients with confirmed ctDNA presence in plasma at PD (%) | Method to confirm ctDNA presence |
|-------------------------------------------------|------------|-----------------|-------------------------------------------------------------|-------------------------------------------------------------|----------------------------------|
| Raimondi et al. (5)                             | 11         | Idylla          | 5/11 (45%)                                                   | 1/7 (14%)                                                   | NGS                              |
| Moati et al. (4)                                | 36         | ddPCR/NGS       | 8/36 (22%)                                                  | 2/30 (6.6%)                                                  | Methylation test                  |
| Fernández Montes et al. (9)                     | 15         | Not specified   | 2/15 (13%)                                                   | 2/15 (13%)                                                   | Not specified                     |

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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