

Sysmex OncoBEAM™ circulating tumor DNA testing in clinical practice

Ultra-high sensitivity
diagnostics to enable
clinical decisions

NRAS

BRAF

PIK3CA

KRAS

EGFR

AKT1

ESR1

AR

Introduction

Evidence for “liquid biopsy” in clinical oncology continues to increase. Clinical practice guidelines now recommend plasma analysis alongside, and in specific situations in place of, tumor tissue^{1,2}. With the introduction of diverse new liquid diagnostics, it is now more complex than ever for physicians to select the right test for the right patient. A key starting point is to recognize that different types of assays excel for different clinical intended uses — matching the performance characteristics of a test with the clinical context of each patient is necessary to appropriately inform medical decisions.

History

Cell-free DNA (cfDNA) was first discovered in 1948³ and is now known to originate from many different sources including infectious organisms, fetal DNA during pregnancy, genomic DNA from white blood cells, and tumor cells. Tumor-derived cfDNA originating from necrotic and apoptotic tumor and deposited into peripheral circulation is known as circulating tumor DNA (ctDNA) and was first described about 40 years after the initial discovery of cfDNA⁴. The first studies relating disease burden and cfDNA levels were completed in the early 2000s, and in landmark studies conducted in 2008, investigators at Johns Hopkins University (Baltimore, MD) showed that ctDNA levels in patients with colorectal cancer change in response to changes in tumor burden⁵. Discrimination of ctDNA from normal DNA is achieved by the presence of mutations. However, due to the fact that ctDNA typically represents a very small fraction of cfDNA present in the blood, use as a biomarker for evaluating tumor dynamics requires a quantitative assay with high analytical and clinical sensitivity to characterize accurately the relatively low number of mutant ctDNA fragments in a sample.

Different technologies have different strengths

Quantitative PCR (qPCR), next-generation sequencing (NGS), and digital PCR (dPCR) are the three most common technologies used for ctDNA analysis. Well designed assays based on any of these methods can serve as useful diagnostic tools depending on the specific clinical needs of the patient.

■ qPCR is an established technology that focuses on detection of one or a few mutations at a time with moderate sensitivity, which is best suited for cases where sample input is not limited. It has been adapted for in vitro diagnostic kits, as well as complete “sample-to-answer” instruments which may be able to offer patients improved access.

■ NGS provides coverage across many genomic targets and unique mutation types. High sensitivity is also possible, however panels that will be validated for clinical use must be carefully designed to optimize the balance between sensitivity, coverage, cost, and sample requirements.

■ dPCR is widely regarded for possessing the highest accuracy, precision, and consistency of genetic diagnostic techniques. Its ability to cover known clinical indications at extremely high sensitivity makes it ideal for cases where detection and quantification of low-frequency mutations can deliver key clinical information.

Clinical evidence shows that very low frequency ctDNA mutations (<0.1% allele frequency in plasma) may have important clinical implications across a variety of different cancer types. This evidence continues to accumulate at a rapid pace⁶⁻⁸. In these cases, ultra-high sensitivity is essential to ensure vital information is not missed so that patient samples are appropriately characterized.

OncoBEAM analytical sensitivity

Sysmex OncoBEAM uses BEAMing technology (Beads, Emulsion, Amplification, Magnetics), a modified digital PCR method that interrogates millions of unique molecules within a sample to ensure detection of rare mutant molecules in the presence of many wildtype copies. The lower limit of detection is therefore consistent with low plasma mutant allele frequencies that are present for a significant proportion of cancer patients. Table 1 provides a comparison of analytical sensitivities for several leading ctDNA tests based on different technologies.

ctDNA Test	Technology	Target Allele Frequency	Analytical Sensitivity (Single Nucleotide Variants)	Limit of Detection (SNV)
Competitor R	qPCR	Not reported	Not reported	25-100 copies/ mL (LoD95) ⁹
Competitor F	NGS, broad panel	>0.5%	>98.9% (95% CI 98.4-99.4%) ¹⁰	Not reported
		0.1-0.49%	67.3% (95% CI 61.7-72.5%) ¹⁰	
		<0.1%	Not reported	
Competitor G	NGS, broad panel	>0.25%	100% ¹¹	0.3% (LoD95) ¹¹
		0.05-0.25%	63.8% ¹¹	
Competitor I	NGS, disease-specific panel (NSCLC)	≥0.5%	100% ¹²	0.25% (LoD90) ¹²
		0.25-0.33%	99.48% ¹²	
		0.13-0.16%	88.93% ¹²	
		0.06-0.08%	56.25% ¹²	
OncoBEAM Lung	BEAMing	0.04-0.11%	100% (95% CI 97.8-100%)	0.04% (LoD95)

Table 1. Analytical sensitivities for several leading ctDNA tests. Notes:

• For Competitor R, LoD95 is reported on a per-mutation basis as copies/ mL. Assuming an average of 5 ng DNA/ mL plasma and 3.3 pg/ genomic equivalent (GE) = ~1500 genomic equivalents per mL; 25 mutant copies/ 1500 GE = 1.7% MAF; 100 mutant copies/ 1500 GE = 6.7% MAF.









• OncoBEAM Lung: DNA input in validation studies was ≥40 mutant molecules in order to minimize random sampling error. Analytical sensitivity and CI were calculated for LoD samples according to CLSI EP12-A2a. LoD95 was calculated according to CLSI EP17-A2.

Non-small cell lung cancer anti-EGFR therapy resistance – an ideal case for liquid biopsy

Acquired resistance to first-line tyrosine kinase inhibitors (TKI) for NSCLC patients who harbor epidermal growth factor receptor (EGFR) sensitizing mutations (exon 19 deletions, L858R point mutation) presents a common clinical problem. Resistance often develops after 10-12 months and is most commonly driven by an acquired mutation in EGFR, T790M, which presents in up to 60% of patients^{13,14}. Third-generation TKIs such as osimertinib (approved by the FDA in Nov 2015) have been shown to be effective in patients with T790M-mediated resistance and disease progression.



Though tissue is the preferred sample type for EGFR analysis for NSCLC, obtaining a tissue biopsy in patients with advancing disease presents significant challenges. In addition to some patients being unwilling or unable to undergo secondary biopsy, the complication rate for intrathoracic biopsies is nearly 20%^{15,16}. Furthermore, even if secondary biopsy is feasible, the combined turn-around time for tissue acquisition and subsequent molecular analysis can be too long, which can significantly delay the administration of appropriate therapy^{17,18}. Tissue sampling in the setting of secondary resistance is further confounded by molecular heterogeneity, wherein T790M may be present in only a subset of tumor cells. Thus, sampling of a single region of single metastatic lesion by tissue biopsy may fail to capture the T790M cells that are driving resistance/progression¹⁹.

Because ctDNA analysis is minimally-invasive, faster than tissue analysis, easily repeatable via additional blood draws, and may better represent disease heterogeneity, it is ideally suited for EGFR analysis for NSCLC patients who have progressed on first-line therapy. However, a ctDNA test used in this setting must have high analytical sensitivity since T790M may be present in a small number of tumor cells, as well as at low concentration in the blood. Most importantly, clinical data must demonstrate that the diagnostic test can be used to predict meaningful patient outcomes for second-line EGFR therapy.

Tissue Biopsy	ctDNA Liquid Biopsy
 Invasive	 Non-invasive
 14+ days	 <7 days
 Single sample one time	 Multiple samples over time
 Only captures genotype at biopsy site	 Better captures disease heterogeneity

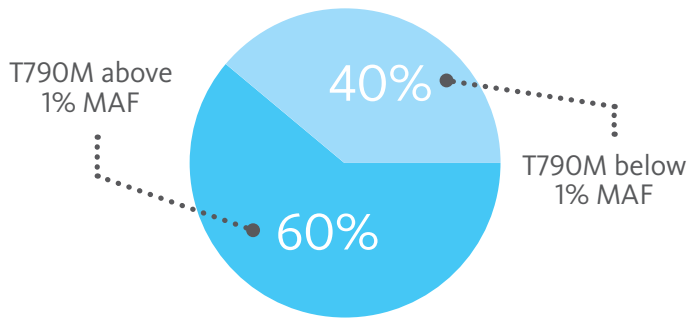
BEAMing demonstrates clinical utility for NSCLC

Based on the study by Oxnard et al. using BEAMing²⁰, NCCN guidelines now recommend plasma testing for EGFR T790M for NSCLC patients who have progressed on a first or second-generation TKI¹. Equivalent clinical outcomes were observed between patients treated with osimertinib who were plasma-positive for T790M, and patients for whom T790M was detected via tissue analysis.

OncoBEAM Liquid Biopsy	Tissue Biopsy
 164 plasma samples positive for T790M	 173 tissue samples positive for T790M
63% response to osimertinib	62% response to osimertinib

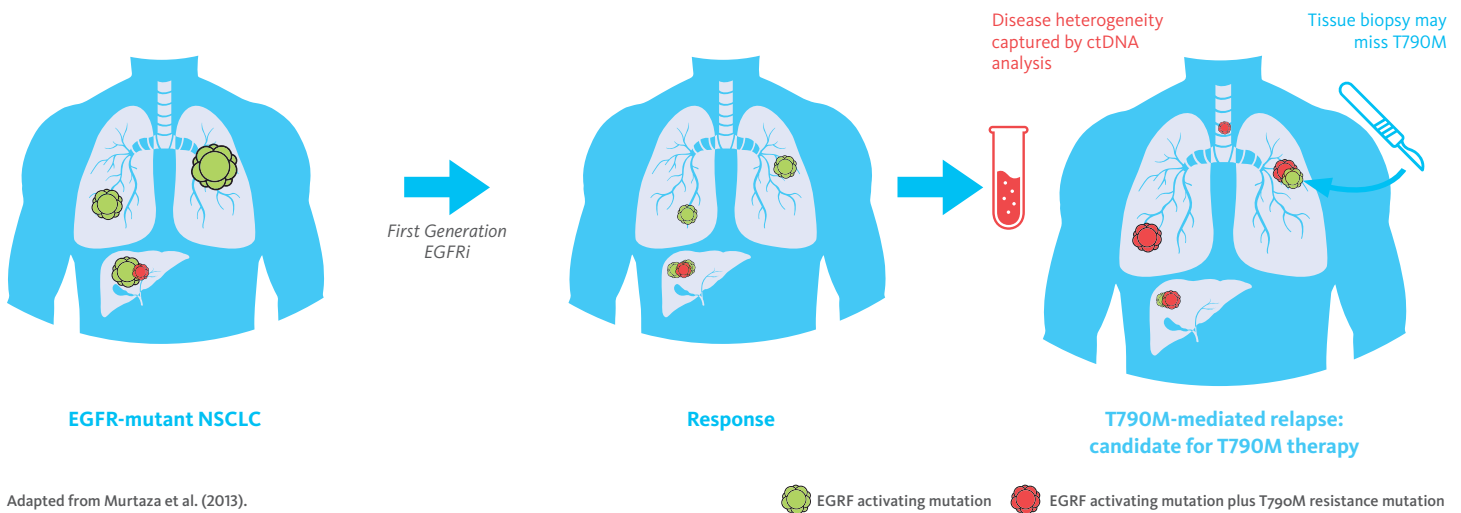
Importantly, for patients who were positive for T790M in tissue, the median mutant allele frequency for T790M detected in plasma by BEAMing was <1%, with a number of patients exhibiting the mutation at <0.1%. This is below the threshold for reliable detection for many other ctDNA assays, including broad NGS panels that excel at generating data across many genomic locations, rather than very high-resolution data focused on regions with established clinical significance.

Mutant allele fraction distribution for EGFR T790M



For 40% of samples, T790M was below 1% MAF and may not be reliably detected by conventional NGS testing.²¹

N=158 plasma samples



Adapted from Murtaza et al. (2013).

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