Clinical Cancer Research

Individualized Molecular Analyses Guide Efforts (IMAGE): A Prospective Study of Molecular Profiling of Tissue and Blood in Metastatic Triple-Negative Breast Cancer

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Abstract

Purpose: The clinical utility of next-generation sequencing (NGS) in breast cancer has not been demonstrated. We hypothesized that we could perform NGS of a new biopsy from patients with metastatic triple-negative breast cancer (TNBC) in a clinically actionable timeframe.

Experimental Design: We planned to enroll 40 patients onto a prospective study, Individualized Molecular Analyses Guide Efforts (IMAGE), to evaluate the feasibility of obtaining a new biopsy of a metastatic site, perform NGS (FoundationOne), and convene a molecular tumor board to formulate treatment recommendations within 28 days. We collected blood at baseline and at time of restaging to assess cell-free circulating plasma tumor DNA (ptDNA).

Results: We enrolled 26 women with metastatic TNBC who had received ≥ 1 line of prior chemotherapy, and 20 (77%) underwent NGS of a metastatic site biopsy. Twelve (60%) evaluable patients

received treatment recommendations within 28 days of consent. The study closed after 20 patients underwent NGS, based on protocol-specified interim futility analysis. Three patients went on to receive genomically directed therapies. Twenty-four of 26 patients had genetic alterations successfully detected in ptDNA. Among 5 patients, 4 mutations found in tumor tissues were not identified in blood, and 4 mutations found in blood were not found in corresponding tumors. In 9 patients, NGS of follow-up blood samples showed 100% concordance with baseline blood samples.

Conclusions: This study demonstrates challenges of performing NGS on prospective tissue biopsies in patients with metastatic TNBC within 28 days, while also highlighting the potential use of blood as a more time-efficient and less invasive method of mutational assessment. *Clin Cancer Res;* 23(2); 379–86. ©2016 AACR.

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Introduction

As patients and oncologists seek additional relevant data to help direct treatment in metastatic breast cancer, demand for genomic profiling of tumors is rapidly increasing. Therapeutic options for patients with metastatic disease for estrogen receptoralpha (ER), progesterone receptor (PR), and HER2-negative, or "triple negative" breast cancer (TNBC), are limited to chemotherapy. It is becoming increasingly clear that TNBC can be further divided into subtypes and that individual patients might benefit from targeted treatments (1, 2). Multiple genomic panels are now commercially available. However, in order for genomic testing to be clinically meaningful for patients with TNBC, who often have rapidly progressive disease, these tests must be performed in a timely fashion to allow for patients and physicians to implement therapies that might be rationally directed from patient-specific alterations.

Despite these challenges, there is strong interest in next-generation sequencing (NGS) approaches to provide molecular profiles of tumors from individual patients for direction of treatment. One recent single-institution experience of 2,000 patients with

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Note: Supplementary data for this article are available at Clinical Cancer Research Online (http://clincancerres.aacrjournals.org/).

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doi: 10.1158/1078-0432.CCR-16-1543

Translational Relevance

Molecular profiling of tumors engenders hope that specific molecular alterations can be matched with targeted therapies. Due to tumor heterogeneity, one would prefer to obtain the mutational status of a new tissue biopsy for informing decisions regarding next line therapy. We prospectively enrolled patients with metastatic triple-negative breast cancer and obtained a recent or new biopsy, and subjected this to nextgeneration sequencing (NGS) with FoundationOne. Our goal was to perform molecular profiling on a recent biopsy and convey recommendations back to the referring physician in 28 days or less. We met our interim analysis endpoint for futility, and the study was stopped after 26 patients were enrolled. However, we also showed that NGS of plasma DNA had high concordance with mutations found in tissue biopsies, and allowed for subsequent follow-up/monitoring using blood. These results demonstrate the potential of "liquid biopsies" for mutational profiling and serial monitoring.

advanced solid tumors demonstrated that comprehensive implementation of NGS tumor profiling for patients was feasible (3). However, the authors found that 23% of patients enrolled in the study were unable to undergo testing due to inadequate tissue or DNA. Notably, the majority of patients were enrolled on the study not to direct the immediate next course of therapy, but to direct future treatments that had no timeline. Presumably, this is due to the time associated with testing and the fact that many patients with rapidly progressive cancer require therapy urgently. One concern with using NGS tumor profiling results to guide future treatment is that the mutational landscape of cancer evolves over time, so an isolated biopsy of metastatic cancer may not be representative of that cancer months or years in the future (4, 5). A single biopsy may also not be genetically representative of other sites of metastatic disease, although the oncogenic drivers of the primary tumor seem to be conserved in metastatic sites (6, 7).

There has been great interest in the use of "liquid biopsies" for cancer sequencing as an alternative to biopsies of metastatic tissues (8). It has been demonstrated that blood is an easily accessible source of circulating cell-free tumor DNA (9–16). Both normal and malignant cells shed DNA into the circulation, and NGS technologies can be used to detect circulating cell-free plasma tumor DNA (ptDNA), making blood a source for real-time genomic tumor profiling.

We hypothesized that we could efficiently create a molecular profile from tumors from patients with metastatic TNBC, and make treatment recommendations based on these results in a clinically actionable timeframe. We initiated a prospective clinical trial to investigate the feasibility of molecular profiling of patients' tumors within 28 days from enrollment to treatment recommendations. Secondarily, we also investigated the use of ptDNA to genomically profile patients' cancers, hypothesizing that blood offers an easily accessible source of tumor DNA.

Patients and Methods

Participants

We initiated a prospective study designated Individualized Molecular Analyses Guide Efforts in Breast Cancer (IMAGE). Women with newly progressing metastatic TNBC with Eastern Cooperative Oncology Group (ECOG) performance status of 0 to 2 who had received at least one line of prior chemotherapy were eligible. Patients were enrolled from clinics within the Johns Hopkins Medical Institutions. We obtained a new biopsy from a metastatic site for molecular profiling at study entry. Archived metastatic biopsy specimens were allowed only if patients had not commenced on a new systemic therapy. The protocol was approved by The Johns Hopkins Institutional Review Board. All patients provided written informed consent prior to enrollment onto the study.

Tumor tissue analysis

Surgical specimens were reviewed by the study pathologist and stained for ER, PR, HER2, and androgen receptor (AR) by immunohistochemistry in a Clinical Laboratory Improvement Amendments (CLIA)-approved laboratory at Johns Hopkins. Specimens then underwent hybrid-capture-based NGS (FoundationOne; Foundation Medicine Inc.). Methods of the clinical cancer gene assay have been previously published, and assay performance has been rigorously validated (17). In brief, DNA was extracted from formalin-fixed, paraffin-embedded tissue ($\geq 1 \text{ mm}^3$) containing no less than 20% tumor nuclei by enzymatic digestion and subsequent purification. DNA was fragmented by sonication to 200 base pair segments. Indexed sequencing adapters were ligated to the DNA fragments and PCR amplified to yield >500 ng of sequencing library. Hybridization selection was performed using individually synthesized baits targeting 236 cancer-related genes and 47 introns of 19 genes frequently rearranged in cancer. The Illumina HiSeq 2500 platform was used in 49×49 paired end sequencing. Sequence data were mapped to the human genome (hg19) using BWA aligner v0.5.9. Sequence data were analyzed through a computational analysis pipeline to call variants present in the sample, including substitutions, short insertions and deletions, rearrangements, and copy-number variants.

Treatment recommendations

Clinical data and genomic profiling reports were reviewed within a week of receipt by the Genomic Alterations In Tumors With Actionable Yields (GAITWAY) molecular profiling tumor board at Johns Hopkins, which interprets genetic alterations found in a patient's tumor sample to identify potentially "actionable" genes and/or proteins and was previously described (18). A potentially actionable alteration was defined as a mutation that (1) has a FDA-approved therapy for the given tumor type, (2) has an FDA-approved therapy for a different tumor type, (3) may provide rationale for participation in a clinical trial, or (4) may lead to recommendations for genetic counseling and germline mutation testing. Recommendations were provided to the treating oncologist, and patients were followed for treatment decision and clinical outcomes.

Plasma DNA analysis

Peripheral blood was obtained at study entry, and whenever possible, every 3 to 4 months and at time of progression. Blood samples and plasma DNA collection and preparation were performed as previously described (11). DNA was extracted from 1.5 to 3.0 mL plasma and used as input into sample preparation without sonication. Indexed sequencing adapters were ligated to the DNA fragments and PCR amplified to yield >2 μ g of sequencing library. Hybridization selection was performed using individually synthesized baits targeting full exons of 27 cancer-related genes (BRCA1, BRCA2, CCND1, CD274, CDH1, CDK4, CDK6, CDKN2A, CRKL, EGFR, ERBB2, ERRFI1, FGFR1, FGFR2, KRAS, MDM2, MET, MYC, NF1, PDCD1LG2, PTEN, PTPN11, SMO, TP53, VEGFA, FOXL2, MYCN) and partial exons of 34 cancerrelated genes (ABL1, AKT1, ALK, ARAF, BRAF, BTK, CTNNB1, DDR2, ESR1, EZH2, FGFR3, FLT3, GNA11, GNAQ, GNAS, HRAS, IDH1, IDH2, JAK2, JAK3, KIT, MAP2K1, MAP2K2, MPL, MTOR, MYD88, NPM1, NRAS, PDGFRA, PDGFRB, PIK3CA, RAF1, RET, and TERT) and introns of 6 genes frequently rearranged in cancer (ALK, EGFR, FGFR3, PDGFRA, RET, and ROS1). The Illumina HiSeq 2500 platform was used in 176 \times 176 paired end sequencing, and >5,000X unique coverage was generated for most samples. Sequence data were mapped to the human genome (hg19) using BWA MEM aligner v0.7.10. Sequence data were analyzed through a computational analysis pipeline which performs error corrections and calls variants present in the sample, including substitutions, short insertions and deletions, and rearrangements.

Mutational concordance for genes was then analyzed for patient samples where NGS of both tissue and blood were successfully performed and was qualitatively recorded as present in both or present in either tissue or blood. Only mutations (not amplifications) in these 27 genes were analyzed for concordance studies.

Statistical methods

The primary objective was to assess feasibility of completing the process from consent to GAITWAY recommendations-including consent, biopsy, molecular profiling of the tumor sample, convening of a tumor board to discuss the results, and reporting of treatment suggestion based on molecular profiling-within 28 days. Secondary objectives included demonstrating the ability to make treatment suggestions based on the molecular profile of patients' tumors, and to prospectively follow ptDNA in all patients who took part. The primary endpoint of this study was feasibility, defined as accomplishing the aforementioned steps within 28 days from consent, for at least 80% of patients. We planned to enroll 40 women, which would lead to a 90% confidence interval (CI) with maximum width of $\pm 14\%$, which we judged would provide usefully precise estimates of the primary endpoints. If the primary objective of feasibility was met in 32 of 40 patients (80%), the 90% CI would be 67% to 90%. The protocol included an interim futility analysis of the first 20 patients who completed the process. The study would close for futility if the analysis predicted there was 25% or less probability of success at the end of the trial.

Results

From September 2013 to April 2015, we enrolled 26 eligible women. The median age was 55 years (range, 25–78); 13 (50%) patients identified as white, and 11 (42.3%) as black; the median number of prior lines of chemotherapy in the metastatic setting was one (range, 0–4); and 65.4% of patients had visceral disease (Table 1). At the time of this analysis, the median duration of follow-up for the 20 patients who underwent NGS was 7.5 months.

Twenty (77%) eligible patients underwent successful NGS of a metastatic site biopsy (Table 2). These included 12 new biopsies, and archival tissue from 8 recent biopsies. For patients with archival specimens, the median number of days from biopsy to registration was 13.5 (range, 7–40), and these patients did not receive any new therapy during this period as specified by our

Characteristic	
Age, years	
Median (range)	55 (25-78)
Race	
White	13 (50%)
Black	11 (42.3%)
Asian	1 (3.8%)
Other	1 (3.8%)
ECOG	
0	12 (46.2%)
1	11 (42.3%)
2	3 (11.5%)
BRCA status (germline)	
BRCA1/2 positive	1 (3.8%)
BRCA negative	15 (57.7%)
Unknown/not done	10 (38.5%)
Prior (neo)adjuvant chemotherapy	
Yes	24
Anthracycline and taxane based	16
Anthracycline based	2
Taxane based	5
Other/unspecified	2
No	2
Prior (neo)adjuvant hormonal therapy ^a	5
Other	1
Number of prior systemic therapy for metastatic dis	sease
All regimens, median (range)	1.5 (0-4)
Chemotherapy	1 (0-4)
Hormonal ^a	1.5 (1-2)
Other/unknown	1 clinical trial
	1 PDL-1 trial
AR status ($N = 21$)	
Positive	5 (23.8%)
Negative	16 (76.2%)

^aEligible patients could have previously had ER-positive and/or HER2-positive disease, but must have had confirmation of TNBC on their most recent tissue biopsy.

protocol. Six patients did not undergo NGS due to either absence of a metastatic site amenable for biopsy (N = 3) or inadequate tissue for NGS from archival (N = 2) or new (N = 1) biopsies. Of

Table 2.	Outcomes
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Signed consent (<i>N</i>)	28
Registered	26
Screen failures (N)	6
Tissue not available	5
Tissue not usable	1
Time from consent to tissue availability, days (range)	
All	2.5 (0-21)
New specimen ($N = 12$)	0 (0-13)
Archival ($N = 8$)	4.5 (1-21)
Time from biopsy to specimen shipment, days (range; $N = 12$)	4 (2-7)
Time from specimen shipment to report from FM, days (range)	15.5 (12–30)
Time from report to tumor board, days (range)	5 (3-10)
Recommendation within 28 days of biopsy	
Yes	12 (60%)
No	8 (40%)
Successful NGS of metastatic biopsy (N)	20
Potentially actionable mutation identified (N)	15
Tumor board recommended targeted therapy as possible next	13
treatment (N)	
Received targeted therapy as next treatment (4) ^a	4

^aA patient with an AR⁺ tumor received bicalutamide on study, a patient with a *BAP1* mutation received carboplatin/PARP inhibitor on study, a patient with a *MAP2K1* amplification received trametinib off study, and a patient with an *ERBB2* mutation received trastuzumab off study.

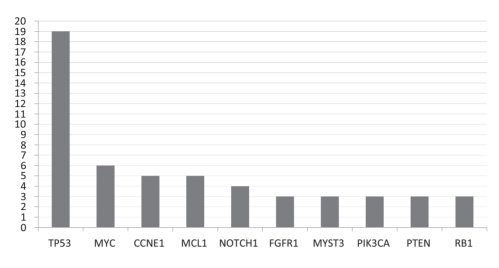


Figure 1.

Most frequent somatic genomic alterations in metastatic biopsies of 20 TNBCs. Shown are the frequencies of genetic alterations identified by NGS in the 20 metastatic lesions from TNBC patients. Alterations include point mutations and copy-number alterations.

the 20 patients who underwent NGS, 12 (60%) patients received treatment recommendations within 28 days of consent (90% CI, 39%–78%). The study met the predefined statistical endpoint for futility and closed after 20 patients underwent NGS. Failure to meet this timeframe was due to difficulties in accessing archival tumor tissue (N = 5) and need to retrieve additional tissue for molecular analysis (N = 3).

All 20 patients who underwent NGS of a metastatic site had at least one identifiable somatic alteration (Fig. 1; Supplementary Fig. S1; Supplementary Table S1). Members of the GAITWAY tumor board reviewed the clinical data, family history, and genomic profile for each of the 20 participants and determined that 15 had at least one potentially actionable alteration. The board recommended targeted treatment as a possible next line of therapy in 13 patients. The recommendations were reported to the treating physician, who ultimately decided with the patient on the next administered therapy. Three patients went on to receive a NGS-identified targeted therapy as their next treatment (Table 2). One patient received AR-directed treatment as her next line of therapy (Table 2). The primary reasons 9 patients did not receive targeted treatment were lack of geographically accessible or available clinical trials, deterioration of patient's performance status precluding enrollment on recommended studies, and initiating another therapy prior to receiving the GAITWAY tumor board recommendations.

The GAITWAY tumor board regarded AR expression as a potential therapeutic target, based on previously published work showing the AR antagonist bicalutamide has clinical activity in ARexpressing breast cancer (19). Five of 21 (23.8%) evaluable patients had AR-expressing tumors, and AR-directed treatment was recommended as a possible next line of therapy in 4 patients. All four of these patients had additional targetable alterations identified by NGS (Table 2).

Four patients in the study received treatment consistent with GAITWAY tumor board recommendations. Patient 1-005 had AR^+ TNBC and was treated with single-agent bicalutamide as an off-label use. Bicalutamide is a nonsteroidal antiandrogen with evidence of activity in $ER^-/PR^-/AR^+$ breast cancer (19). After 2 weeks of treatment, she had progression of disease in the skin and the drug was stopped. Patient 1-017 had a *BAP1* E498fs*38 mutation and *BAP1* loss in her tumor. Bap1 is a BRCA1-associated tumor suppressor protein. A clinical trial of a PARP inhibitor was

recommended as a possible next step in treatment, as PARP inhibitors have demonstrated efficacy in patients with BRCA mutations (20). She was treated with a PARP inhibitor in combination with carboplatin. After two cycles of treatment, she had stable disease, with eventual disease progression after four cycles. Patient 1-021 had a MAP2K1 amplification in her tumor and was recommended a clinical trial with a MEK inhibitor as a possible next therapeutic choice. Trametinib is one such oral tyrosine kinase inhibitor approved to treat BRAF V600E or V600K-mutated melanoma in combination with dabrafenib. Her treating oncologist then started off-label, single-agent trametinib and documented clinical response in the breast and clavicle. After several weeks of treatment, she had disease progression. Finally, patient 1-023 had two known ERBB2 kinase domain mutations in her tumor (D769H and V777L) that in preclinical models confer sensitivity to HER2-directed treatment (21). Off-label HER2-directed treatment was recommended as a potential next treatment for her. She received trastuzumab alone for 10 months and then had progression of disease. Capecitabine was then added to trastuzumab, with stable disease for 9 months.

Members of the GAITWAY tumor board also considered whether particular genetic alterations were suggestive of potential germline mutations, for example, *BRCA1*, *BRCA2*, or *TP53* mutations. In 8 (40%) patients, genetic counseling was recommended to further investigate family history and potential need for formal germline genetic testing.

We successfully collected baseline blood samples from all 26 patients. Two of 26 baseline patient samples had insufficient cellfree DNA (cfDNA) for NGS, yielding inadequate coverage for accurate mutation identification. NGS of baseline ptDNA samples was successfully performed and detected at least one mutation in all remaining 24 patients, including 6 patients who did not have successful NGS of tumor tissue (Table 3). For patients who had NGS analysis of both tumor and blood (N = 18; Fig. 2), 23 of 33 mutations (70%) were concordant. A total of 5 patients had discordant results between tumors and tissues. In 3 patients (patients 14, 17, and 19), 4 mutations found in tumor tissues were not identified in the blood including two separate TP53 mutations found only in the metastatic biopsy for patient 19. Conversely, 4 mutations found in blood were not found in corresponding tumor tissues in 4 patients (patients 10, 13, 14, and 19), though none were deemed targets for therapy.

Patient	Time	-	Alteration	ptDNA-allelic
ID	point	Gene	(AA)	frequency (%)
1	Om	TP53	R342	20.1
1	fu1	TP53	R342 [°]	48.0
2	Om	TP53	Y163C	31.66
2	fu1	TP53	Y163C	7.32
2	fu2	TP53	Y163C	41.95
3	0m	TP53	M66fs 57	33.93
3	fu1	TP53	M66fs 57	0.15
3	fu2	TP53	M66fs 57	7.50
4	0m	TP53	L206	1.64
4	0m	BRCA1	E1849	1.21
5	0m	TP53	R209fs 6	0.63
5	0m	PTEN	Y188fs 2	0.08
5	0m	PTEN	T319fs [°] 6	0.10
6	0m	TP53	R175H	16.96
6	0m	AKT1	E17K	8.27
7	0m	TP53	R248Q	1.27
8	0m	TP53	Y234C	47.05
9	0m	TP53	P219fs [°] 2	0.076
9	0m	PIK3CA	V344G	20.12
10	0m	TP53	E294fs [*] 51	0.29
10	fu1	TP53	E294fs [*] 51	0.39
10	fu2	TP53	E294fs [*] 51	0.20
10	0m	TP53	R249W	3.24
10	fu1	TP53	R249W	0.71
10	fu2	TP53	R249W	1.08
11	0m	TP53	R110P	2.02
12	0m	TP53	I195T	59.28
13	0m	TP53	R273L	2.44
13	fu1	TP53	R273L	1.86
13	0m	TP53	Y163C	0.6
13	fu1	TP53	Y163C	10.22
14	0m	TP53	R213fs [°] 34	0.21
15	0m	TP53	R248Q	27.98
15	0m	TP53	splice site 376-1G>A	0.65
16	0m	TP53	Y234del	34.06
17	0m	TP53	A276G	10.51
17	fu	TP53	A276G	24.48
17	fu	BRCA2	S2001C	0.79
18	0m	TP53	R213 [*]	15.38
19	0m	JAK2	V617F	1.31
20	0m	TP53	E221fs [*] 4	0.1077
21	0m	TP53	T253N	25.17
21	Om	TP53	N247K	25.15
21	0m	TP53	R175H	0.73
22	Om	TP53	P151R	1.65
22	Om	BRCA1	E23fs [*] 17	30.00
24	Om	NF1	Q83 [°]	2.97
24	fu	NF1	Q83°	23.97
24	Om	CDH1	S111fs [°] 6	18.95
24	fu	CDH1	S111fs 6	27.21
24 24	Om	PIK3CA	H1047R	18.12
24	fu	PIK3CA	H1047R	25.58
24 26	0m	PTEN	C296fs 12	0.101
26	Om	PIEN PIK3CA	M1043I	0.081
26	0m	PIK3CA	N1044K	0.082

NOTE: Allelic frequency is the percentage of mutant DNA relative to total DNA as assessed by NGS.

*Stop codons.

Interestingly, patients 14 and 19 did not have any mutations found in both tissue and blood.

In addition to baseline blood samples, we obtained 12 followup blood samples from 9 patients every 3 to 4 months for analysis. Two of nine initial follow-up samples (FU1) were unevaluable due to insufficient input cfDNA. In seven of seven (100%) evaluable FU1 samples and in three of three (100%) subsequent follow-up samples (FU2), NGS identified all mutations found in baseline samples. In the patient unable to undergo NGS tissue analysis with follow-up blood collected (patient 24), NGS identified three of three mutations seen at a baseline blood sample. In 1 of 7 patients, NGS identified a new mutation at FU1 not seen at baseline or in the tumor sample (patient 17).

In general, patients with follow-up blood samples displayed changing allelic fractions of ptDNA for any given mutation (Table 3), reflective of disease progression and in some cases response to therapy. Notably for patient 3, who had brain and other organ metastases, an initial baseline TP53 mutation was detected in both tumor and plasma, with an allelic frequency in blood of 33%. In fact, this percentage was high enough that we tested her germline DNA and confirmed that this was indeed a somatic mutation. The ptDNA sample at baseline was taken prior to removal of a metastatic brain lesion, and the brain lesion itself was used for NGS on this study. A subsequent follow-up blood sample demonstrated a remarkably lower allelic fraction (0.15%) of mutant TP53. Ultimately, the patient progressed systemically and succumbed to her disease, and blood samples obtained at the time of progression demonstrated an increase in mutant TP53 ptDNA (7.5%).

Discussion

In this prospective study of patients with advanced TNBC, we obtained a new biopsy of metastatic disease, performed genomic analysis and clinical interpretation, and returned information to the referring physician and patient. Our goal was to complete these steps within a clinically relevant timeframe of 28 days, but the study was terminated after meeting prespecified criteria for futility. NGS in tissue could not be performed in 8 of 20 (40%) patients within 28 days due to sampling issues, including difficulty accessing archival tumor tissue, and need for repeat submission of tumor tissue due to inadequate initial sample. In contrast to NGS of tissue samples, we successfully obtained and performed NGS on plasma samples in 24 of 26 patients, and were able to identify mutations in all 24 patients' ptDNA. Because levels of plasma DNA are variable, we found that there were inadequate quantities of cfDNA for NGS in 2 of 26 patients, using approximately 1.5 mL of plasma. Although we now obtain additional blood tubes to mitigate this issue, in the current study, plasma exhaustion accounted for the inability to sequence these two patient samples. Our results demonstrate the clinical challenges of obtaining prospective tissue biopsies for sequencing in patients with metastatic TNBCs within 28 days, while also highlighting the potential use of blood as a more time-efficient method of cancer gene sequencing.

Our study emphasizes several pragmatic limitations regarding the ability to use NGS of tumor tissue for clinical management, including obtaining a recent or new biopsy for analysis. Although other studies have used archival specimens for NGS, it is now well established that tumor heterogeneity and clonal evolution after therapy can alter the genomic landscape of metastatic lesions. For example, recent findings suggest that *ESR1* (estrogen receptoralpha) ligand binding domain mutations occur predominantly, if not exclusively, in metastatic breast cancer patients after endocrine therapies (22–24), and that ptDNA can detect additional mutations not found in tissue biopsies (13, 25, 26). Thus, for molecular tumor profiling, a new biopsy prior to initiating a new therapy would in theory yield the most relevant information to

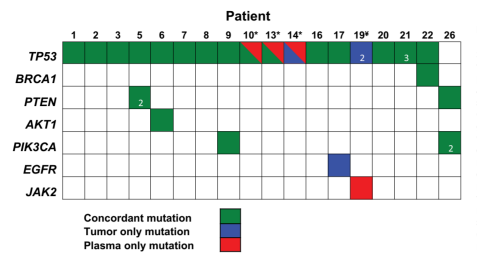


Figure 2

Plasma and tumor mutation concordance. Shown are concordance between mutations found in tumor tissue and mutations found in paired blood samples. Copy-number alterations were not performed on blood samples and were therefore not analyzed Concordance between tissue and blood for a given mutation is represented in green, whereas mutations found exclusively in tissue or blood are represented as blue and red, respectively, Patients 5, 21, and 26 had multiple distinct mutations within a given gene with the number of mutations shown for each gene (2 for PTEN, 3 for TP53, 2 for PIK3CA, respectively). Three patients had additional, separate TP53 mutations found only in blood (*), whereas 1 patient had two separate TP53 mutations found only in tumor (¥).

guide patient care. But obtaining tissue sufficient for NGS in patients with metastatic TNBC in a timely way was difficult. Six patients of 26 could not undergo NGS tumor testing due to either inaccessible tissue (N = 5) or insufficient tissue for analysis (N = 5)1). It is important to note that these 6 patients had already consented to this study, indicating that their treating physicians believed they had accessible tumors amenable to biopsy. Biopsy also added significant delay, as it requires scheduling with a specialized team. We purposefully chose to test our feasibility hypothesis in patients with TNBC since their rapidly progressive disease and limited standard-of-care treatment options represent an especially challenging group of patients for benchmarking timed molecular analyses. Moreover, a 28-day timeframe was chosen, as this is often a required "washout" period in clinical trials of new therapies. However, our results demonstrate the realworld challenges in performing such studies, including technical difficulties of obtaining tissue, DNA requirements for NGS, and the need for patients to start therapy if results are not delivered in a relevant clinical timeframe.

Currently, tumor NGS is used most often to identify genomic alterations that may provide rationale for targeted therapies. This feature depends on the frequency of actionable mutations, which varies widely depending on tumor type, and changes over time as new drugs come under investigation and become commercially available. For example, despite the TP53 gene being altered in 95% of patients in this study, our GAITWAY tumor board did not consider these mutations targetable. If future therapies that effectively target mutant p53 are successfully developed, this would greatly affect the use of NGS for TNBC patients. Of the 20 patients with successful NGS, only 4 (20%) went on to receive a targeted treatment guided by NGS, although 15 had GAITWAY-determined targetable aberrations. For the majority of those patients with targetable alterations, the only directed treatments available were via clinical studies. In most cases, these studies were not available locally. This issue of study availability is consistent with findings at other academic centers, and will require additional resources and careful study designs to accommodate numerous, low frequency mutations (3, 27, 28).

At the same time, we were able to successfully obtain and perform NGS on ptDNA from the majority of participants (24 of 26). Beyond the ease of obtaining blood versus tissue, liquid

biopsies may have other advantages over biopsy of a single metastatic site. For example, inter- and intratumor heterogeneity is an important consideration when using tissue biopsy specimens as the source of genomic testing (29). Emerging evidence suggests that ptDNA represents mutational burden from disparate clonal populations in patients with metastatic disease (13, 30) due to blood acting as a reservoir for all metastatic sites shedding ptDNA into the circulation. In accord with this notion, our analysis demonstrates that additional mutations were detected in ptDNA that were not found in the tumor biopsy. For example, 3 patients had a second TP53 mutation found exclusively in blood, possibly reflecting mutations from other metastatic sites. Interestingly, in 1 patient (patient 19) with two TP53 mutations found only in the tumor, a JAK2 V617F mutation was seen only in the blood at relatively low allelic frequency. Because JAK2 V617F mutations are almost exclusively found in hematologic diseases (e.g., polycythemia vera), there is a possibility that that this ptDNA mutation may reflect an undiagnosed hematologic disorder. Importantly, ruxolitinib is an approved JAK inhibitor for the treatment of JAK2 V617F hematologic diseases. Further verification of the origin of this patient's JAK2 mutation is ongoing.

For the majority of patients, the same mutations were identified in tumor and blood (23/33, 70%) if both were successfully sequenced. Although we previously reported a higher degree of mutational concordance when tissue and blood were obtained concurrently (9), there were important differences in the current study, notably, the use of NGS for ptDNA detection, rather than digital PCR. It is known that allelic frequencies in ptDNA are often extremely low, and the current sensitivity of NGS may not allow for detection of mutations in plasma that are derived from lowlevel subclonal variants present in a metastatic site. Newer technologies and bioinformatics pipelines may enable improved sensitivity for future NGS studies. Another limitation of our ptDNA analysis is the inability to assess copy-number changes in blood. During this study period, copy-number analysis of ptDNA was still in development and therefore could not be reported. Finally, NGS generally requires a higher amount of input cfDNA compared with digital PCR. We had inadequate DNA quantity to obtain enough NGS coverage to confidently assess mutations in ptDNA for 2 of 26 patients. Future studies can avoid this limitation by obtaining additional tubes for each blood draw.

The ease of blood sampling also enables tracking mutational evolution of metastatic disease over time along with response to therapies. We collected serial measurements in 9 patients, and our data do indeed demonstrate that mutations can be monitored qualitatively and quantitatively in a serial fashion, supporting results by others that serial NGS of ptDNA is feasible (9-16). Small studies in breast and other cancers have shown that a change in ptDNA can predict for change in disease status in advance of clinical or imaging-based evidence (11, 16, 31, 32). Our study also suggests ptDNA can be used to track response to therapies, including neurosurgical removal of brain metastases as describe above. It will be important to confirm whether ptDNA can be used as a way to monitor disease burden and response to therapies, and determine if changing therapies in patients who do not have a decrease in ptDNA after starting treatment improves clinical outcomes

In summary, we have shown the clinical challenges of using new metastatic biopsies for molecular profiling of actionable mutations within a 28-day timeframe. This study highlights the need for easier, quicker mutational profiling methods in patients with metastatic TNBC, and suggests NGS of blood as a potential alternative to tissue. We envision that NGS of ptDNA could be used to identify more patients than tissue biopsy for enrollment onto clinical trials. Larger studies investigating the utility of NGS of ptDNA in tracking metastatic disease and predicting for response to treatment are needed.

Disclosure of Potential Conflicts of Interest

S.M. Ali, T.A. Clark, D. Lipson, and P.J. Stephens hold ownership interest (including patents) in Foundation Medicine. J.S. Ross reports receiving commercial research grants from and holds ownership interest (including patents) in Foundation Medicine. V. Stearns reports receiving commercial research grants from Abbvie, Celegene, Medimmune, Merck, Novartis, Pfizer, and Puma. B.H. Park is a consultant for Foundation Medicine, Inc. No potential conflicts of interest were disclosed by the other authors.

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Acknowledgments

Drs. Parsons and Park had full access to all of the data in the study and take responsibility for the integrity of the data and the accuracy of the data analysis.

Grant Support

This work was supported by The Conquer Cancer Foundation's Young Investigator Award, The Pearl M. Stetler Research Fund, NIH (CA088843, GM007309, CA168180, CA167939), The Avon Foundation, Commonwealth Foundation, the Santa Fe Foundation, The Canney Foundation, Cigarette Restitution Fund, The Cindy Rosencrans Fund for Triple Negative Breast Cancer Research, and the Marcie and Ellen Foundation. Foundation Medicine analyzed samples free of charge.

Role of the Funder/Sponsor

None of the funding sources influenced the design, interpretation, or submission of this manuscript.

The costs of publication of this article were defrayed in part by the payment of page charges. This article must therefore be hereby marked *advertisement* in accordance with 18 U.S.C. Section 1734 solely to indicate this fact.

Received June 17, 2016; revised July 24, 2016; accepted July 26, 2016; published OnlineFirst August 3, 2016.

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