

High-Throughput Detection of Actionable Genomic Alterations in Clinical Tumor Samples by Targeted, Massively Parallel Sequencing

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ABSTRACT

Knowledge of “actionable” somatic genomic alterations present in each tumor (e.g., point mutations, small insertions/deletions, and copy-number alterations that direct therapeutic options) should facilitate individualized approaches to cancer treatment. However, clinical implementation of systematic genomic profiling has rarely been achieved beyond limited numbers of oncogene point mutations. To address this challenge, we utilized a targeted, massively parallel sequencing approach to detect tumor genomic alterations in formalin-fixed, paraffin-embedded (FFPE) tumor samples. Nearly 400-fold mean sequence coverage was achieved, and single-nucleotide sequence variants, small insertions/deletions, and chromosomal copy-number alterations were detected simultaneously with high accuracy compared with other methods in clinical use. Putatively actionable genomic alterations, including those that predict sensitivity or resistance to established and experimental therapies, were detected in each tumor sample tested. Thus, targeted deep sequencing of clinical tumor material may enable mutation-driven clinical trials and, ultimately, “personalized” cancer treatment.

SIGNIFICANCE: Despite the rapid proliferation of targeted therapeutic agents, systematic methods to profile clinically relevant tumor genomic alterations remain underdeveloped. We describe a sequencing-based approach to identifying genomic alterations in FFPE tumor samples. These studies affirm the feasibility and clinical utility of targeted sequencing in the oncology arena and provide a foundation for genomics-based stratification of cancer patients. *Cancer Discovery*; 2(1);82–93. ©2011 AACR.

INTRODUCTION

The maturation of cancer genome characterization efforts has fueled the notion that many treatment decisions might ultimately be guided by the genetic makeup of individual tumors (1). Moreover, the rapid proliferation of targeted agents in development has called specific attention to the importance of molecular profiling approaches that pinpoint *in situ* those tumors most likely to respond. Knowledge of such alterations in the clinical and translational arenas—including mutations, somatic copy-number alterations, and polymorphisms affecting drug metabolism—should ultimately facilitate individualized approaches to cancer treatment. However, systematic genetic profiling of cancers remains underdeveloped in the

clinical setting. Because many targeted agents in development are designed to intercept proteins and/or pathways commonly perturbed by tumor genetic changes, an urgent need exists to implement robust approaches that determine the “actionable” genetic profiles of individual tumors. If widely obtained, such information might better identify those patients most likely to respond to existing and emerging anticancer regimens.

We and others have developed tumor mutation–profiling platforms that use mass-spectrometric genotyping (2, 3) or allele-specific PCR-based technologies (4). Each of these approaches interrogates known oncogene or tumor suppressor gene mutations present in DNA obtained from either frozen or formalin-fixed, paraffin-embedded (FFPE) tumor tissue. However, genotyping-based platforms have certain limitations that may preclude their applicability as definitive cancer diagnostic modalities. These include the finite number of prespecified point mutations that can be assayed (designated *a priori* from a restricted subset of known cancer genes), difficulties in detecting small insertions or deletions (“indels”), insensitivity to most tumor suppressor gene mutations (which may occur anywhere within the gene), inability to detect gene amplifications or deletions, and decreased sensitivity in tumor samples with high stromal admixture. At the present time, no systematic mechanism exists whereby clinical tumor specimens might be interrogated *in situ* for a fully comprehensive panel of actionable cancer gene alterations.

The advent of massively parallel sequencing is transforming the cancer genomics landscape by enabling

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comprehensive cancer genome characterization at an unprecedented scope (1, 5, 6). Concomitantly, hybrid selection-based methods that enrich for coding sequences prior to sequencing (“exon capture”; refs. 7, 8) are routinely being implemented in discovery-oriented settings (5). Here, we describe an adaptation of exon capture and massively parallel sequencing for robust detection of somatic genomic alterations in FFPE samples. The approach leverages a targeted exon capture technique to enrich for a cancer-relevant genomic territory consisting of 137 genes (~400,000 coding bases), thereby allowing multiple barcoded samples to be pooled into a single sequencing reaction while preserving deep (e.g., >300- to 400-fold) sequencing coverage of targeted regions. This approach simultaneously identifies mutations and chromosomal copy-number alterations in clinical tumor material and may inform a comprehensive means to achieve DNA-based patient stratification in the clinical and translational oncology arena.

RESULTS

We generated a list of 137 “druggable” or potentially actionable genes known to undergo somatic genomic alterations in cancer (Supplementary Table S1). These include targets of existing and novel therapeutics, prognostic markers, and other oncogenes and tumor suppressors that are frequently mutated in cancer. In addition, we included 79 pharmacogenomic polymorphisms in 34 genes that may predict heightened sensitivity/resistance or toxicity to conventional cancer therapies (Supplementary Table S2). Altogether, these genes comprise 2,372 exons encoding 433,159 bases. We then designed and synthesized 7,021 unique biotinylated RNA baits corresponding to these genomic regions.

We leveraged a solution-based exon capture/massively parallel sequencing approach in which a pool of long oligonucleotides complementary to these exons of interest were used to reduce the complexity of tumor genomic DNA for clinically-oriented sequencing. Here, a 6-nucleotide DNA barcode was appended to the ends of DNA fragments during library construction, thus allowing multiple samples to be pooled before hybrid selection to expand the scope of genomic profiling (9). The approach is illustrated schematically in Supplementary Figure S1.

Capture Performance and Reproducibility

We first optimized the approach by using genomic DNA from normal samples and tumor cell lines known to harbor mutations and/or chromosomal copy-number alterations affecting multiple cancer genes represented in our hybrid capture baits. Ten cancer cell lines with well-characterized, mutually exclusive cancer gene mutations were chosen (Supplementary Table S3) as well as control diploid genomic DNA. Equimolar amounts of the resulting sequencing libraries were pooled together with an additional library from the HT-29 cell line, which was added at a 50% molar ratio compared with the other libraries. This pool of 12 libraries was subjected to a single hybrid

selection reaction and sequenced in a single Illumina lane with 100-bp paired end reads.

The 11 equimolar DNAs were evenly represented, with 12 to 17 million purity-filtered reads generated per sample (average of ~14.6 million purity-filtered reads; Supplementary Table S4), whereas the sample present at 50% concentration (HT-29, index 2) had ~7.8 million purity-filtered reads, as expected. The percent of bases mapping “on-target” averaged 60% (range, 56%–64%) across all samples in the pool, yielding a mean 527× target coverage (range, 441× to 593×) for the 11 equimolar samples. More than 95% of target exons exhibited more than 30× coverage after sequencing (sufficient to call “high-confidence” variants in a sample with 70%–80% tumor purity), whereas only 1% had no coverage (Supplementary Fig. S2A and B). In general, poorly captured exons had greater than 70% GC content, although GC content did not account for all of the poorly captured targets (Supplementary Fig. S2C). The capture performance for a particular target exon was highly reproducible from sample to sample (Supplementary Figure S2D–F).

Detection of Single-Nucleotide Variants, Insertions/Deletions, and Copy-Number Alterations

In total, 102 single-nucleotide variants and 6 indels (excluding known germline polymorphisms) were detected in coding sequences across the 10 cell lines, including all 21 single-nucleotide variants and 3 of 4 indels reported for these lines in the Catalogue of Somatic Mutations in Cancer database (COSMIC) (Supplementary Table S5; ref. 10). The single indel that was not initially identified—a 9-bp deletion in *PIK3CA* in the NCI-H69 cell line—was readily detected by manual inspection of the raw sequencing data. Therefore, all previously reported point mutations and indels for this small collection were detectable by this approach. (A complete listing of all alterations identified in these cell lines can be found in the Supplementary Appendix.)

In the absence of paired normal samples, the majority of variants detected are germline alterations. Nonetheless, previously unreported variants were still informative in several instances. For example, 12 single-nucleotide variants were detected in the breast cancer cell line MDA-MB-231, including all 4 alterations in the COSMIC database (*BRAF*, *TP53*, *KRAS*, and *NF2*; Fig. 1A and B, Supplementary Table S6; ref. 10). One of the additional alterations was a 1-bp frameshift insertion involving the *NF1* tumor suppressor predicted to generate a truncated protein product (Fig. 1C). This *NF1* insertion likely represents a *bona-fide* cancer-associated mutation. The MDA-MB-231 cell line has previously been shown to lack both an *NF1* mRNA isoform and the neurofibromin protein (the product of *NF1*); thus, these findings may provide a genetic basis for neurofibromin loss in this setting (11).

Although detection of point mutations and indels by targeted, massively parallel sequencing has become increasingly common, the simultaneous detection of chromosomal copy-number alterations by this approach is less well-established, particularly in the clinical arena. To determine copy-number

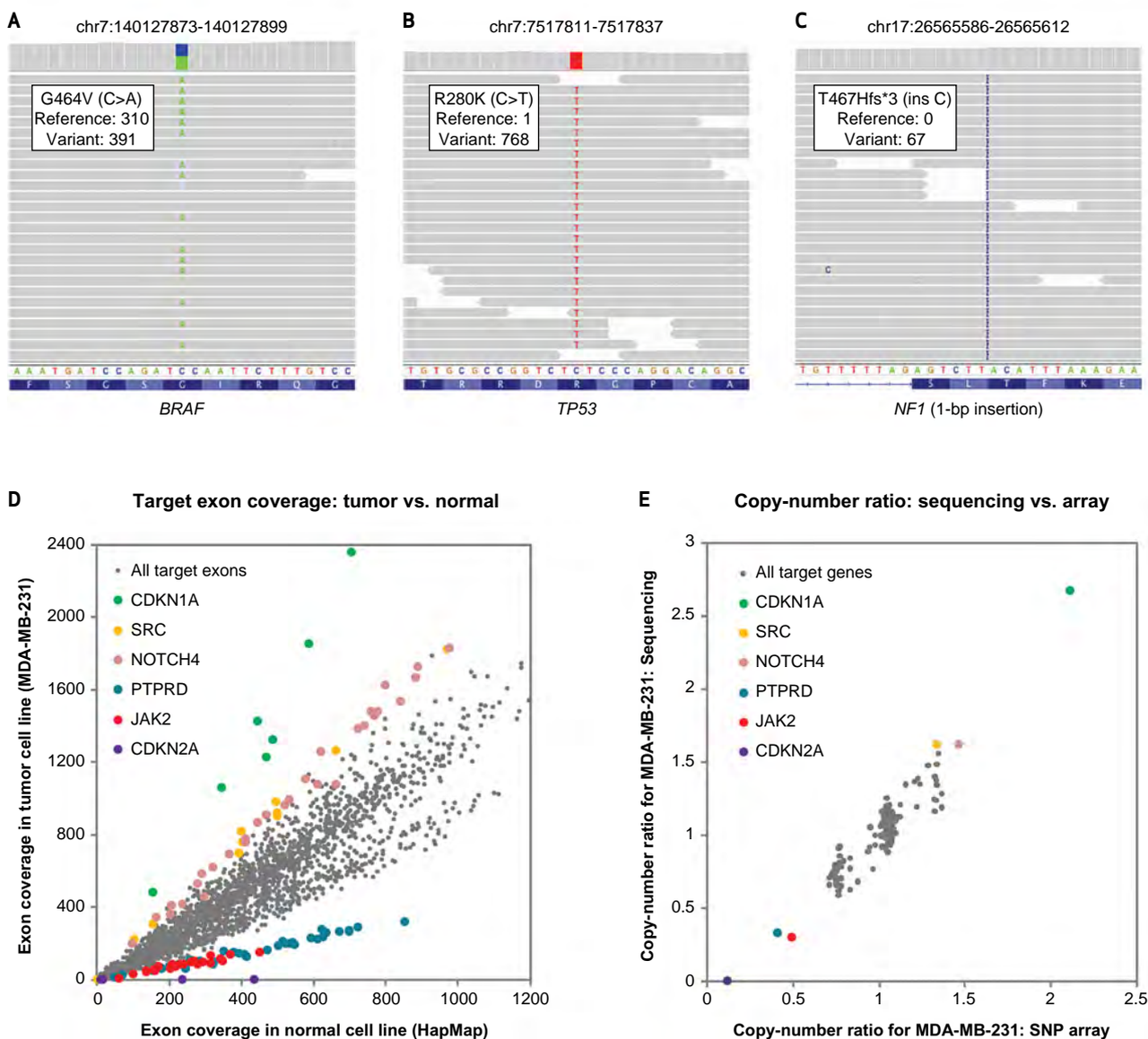


Figure 1. Genomic alterations in breast cancer cell line MDA-MB-231. **A–C**, representative genome images from the Integrated Genome Viewer (IGV) for several alterations found in the breast cancer cell line MDA-MB-231. The number of reads for the reference allele and the variant allele are shown for each alteration. **A**, *BRAF* oncogene point mutation. **B**, point mutation in the *TP53* tumor suppressor gene. **C**, a 1-bp insertion in tumor suppressor *NF1*. **D**, sequence coverage for each target exon in breast cancer cell line MDA-MB-231 compared with a normal diploid sample. Targets from several genes with copy-number gains and losses are highlighted. **E**, comparison of gene-level copy-number alterations as detected by exon capture and copy-number data previously obtained with a high-density single-nucleotide polymorphism (SNP) array (Affymetrix SNP 6.0 platform). Several genes with copy-number gains and losses are highlighted. Copy-number data are highly correlated, with a correlation coefficient of 0.94.

alterations, the accumulated sequence coverage for each exon in the tumor sample was compared with the coverage obtained for the same exon in the diploid normal control (after normalization for global differences in “on-target” sequence coverage). When tumor and normal reads are displayed as a scatter plot (normal = x-axis and tumor = y-axis), exons with a neutral copy-number across the 2 samples should be distributed along a diagonal with a slope

of 1. Amplified exons present in the tumor should have a greater number of relative reads and therefore fall above the diagonal, whereas deleted exons should have fewer reads and fall below the diagonal.

Guided by this framework, we determined relative copy-number ratios for all targeted exons across the cell line collection. An example for the MDA-MB-231 breast cancer cell line (compared with a normal diploid sample) is shown in

Figure 1D. In total, 8 genes with amplifications (defined as mean sequence coverage >3-fold greater than the reference normal) and another 8 with deletions (mean sequence coverage >3-fold lower than the reference normal) were seen across the cell lines. Comparison of overall copy-number values derived by sequencing to those obtained from high-density single-nucleotide polymorphism (SNP) array data (Affymetrix SNP 6.0 platform) demonstrated a robust correlation at the gene level, with correlation coefficients ranging from 0.89 to 0.98 (Supplementary Table S7). As an example, the correlation for the MDA-MB-231 cell line ($r^2 = 0.94$) is shown in Figure 1E.

Profiling of Archival Tumor Samples by Massively Parallel Sequencing

Having established a robust approach for high-throughput exon capture and massively parallel sequencing of 137 cancer genes, we next sought to determine whether this approach might prove useful in the clinical setting. As a proof-of-principle, we characterized a pilot collection of 10 FFPE tumor samples from patients with breast or colon cancer. As was the case with the aforementioned cell line experiment, each of the 12 barcoded samples was evenly represented, with a mean coverage of 391× (Table 1). There was greater variation in the tumor samples compared with the cell lines, with coverage ranging from 116× to 537×. This variance may reflect differences in quality of FFPE-derived input DNA. For 11 samples, 94% of

exons targeted had more than 30× coverage after sequencing and 1% had no coverage. In one sample (FFPE 9; Table 1), 86% of exons showed more than 30× coverage and 2% had zero coverage—this sample also had the lowest mean coverage of the group (116×). The tumor purity for 8 samples was greater than 50%, whereas 2 samples had tumor purities of 20% or less (FFPE 2 and FFPE 3; Table 1).

In total, 155 sequence variants and 14 indels were detected across the samples. In addition, 2 gene amplifications (>3-fold increase in mean sequence read counts compared with a reference normal sample) and 2 gene-level deletions (3-fold decrease in mean sequence read counts) were seen. (Summary information for all 10 samples is shown in Supplementary Table S8; a complete listing of all alterations can be found in the Supplementary Appendix.)

Detection of Clinically Actionable Genomic Alterations in FFPE Tumor Samples

Next, we developed an initial framework to segregate genetic alterations on the basis of their predicted clinical utility. Toward this end, we designated 3 categories of alterations. One category, termed “actionable in principle,” includes variants that predict tumor sensitivity or resistance to U.S. Food and Drug Administration (FDA)-approved (tier 1) or experimental therapies (tier 2). Another category contains prognostic or diagnostic variants. The remaining alterations are termed “variants of unclear significance,” which may

Table 1. Summary of sequencing results for FFPE samples

Sample	Tumor type	Tumor purity, %	Percent of total			Mean target coverage	Percent of target bases with at least 30× coverage
			PF reads	PF reads in pool	Percent selected bases		
HAPMAP	N/A	N/A	9,655,996	7	46	394	96
FFPE 1	Colon	60	11,161,868	8	46	457	96
FFPE 2	Colon	10	8,841,660	7	48	353	94
FFPE 3	Colon	20	13,047,230	10	44	498	96
FFPE 4	Colon	60	10,144,562	8	38	300	95
FFPE 5	Breast	80	16,450,558	12	36	472	95
FFPE 6	Breast	70	15,188,624	11	42	532	96
FFPE 7	Colon	50	8,480,282	6	39	250	94
FFPE 8	Breast	80	15,758,604	12	41	537	96
FFPE 9	Colon	60	3,640,236	3	42	116	86
FFPE 10	Colon	50	14,429,284	11	36	410	96
HT-29 (cell line)	Colon	N/A	7,519,880	6	53	369	96

Abbreviations: N/A, not available; PF, purity filtered.

NOTE: Barcoded and pooled genomic DNA from FFPE tumor samples was subjected to exon capture and sequenced in a single 100-bp paired-end Illumina HiSeq2000 lane. PF sequence reads for each sample are shown; the percent of total PF reads shows the relative representation of each sample within the pool. “Percent selected bases” indicates bases that mapped within 250 bp of a target exon, including both on- and near-target sequence. “Mean target coverage” represents the average number of unique reads in which each base was sequenced.

include biologically important mutations without known therapeutic implications as well as uncharacterized mutations in genes with presumed clinical relevance.

We detected biologically or clinically meaningful alterations in all 10 FFPE samples, including the 2 samples that contained only 10% to 20% tumor cells. These include known somatic mutations in *KRAS*, *BRAF*, *PIK3CA*, and *CTNNB1*; nonsense mutations in the tumor suppressors *APC*, *MSH2*, *SMAD2*, *TSC1*, and *TP53*; and a 2-bp deletion in *BRCA1*. In particular, 12 of the 155 single-nucleotide variants and 1 of the 14 indels were deemed plausibly actionable (“actionable in principle” or “prognostic/diagnostic”; Table 2). *KRAS* mutations in colon cancer predict resistance to cetuximab (12, 13) and exemplify tier 1 actionable alterations. In addition, mutations in *PIK3CA* have been shown in some studies to promote resistance to cetuximab in patients with colon cancer (13–16) and trastuzumab in breast cancer (17, 18), and therefore may conceivably represent tier 1 alterations (although this has not been shown definitively). Multiple tier 2 actionable alterations (targeted by drugs currently in clinical development) were also seen, including mutations in *PIK3CA* [phosphoinositide 3-kinase (PI3K) pathway inhibitors; ref. 19], *KRAS* (MEK inhibitors; ref. 20), *TSC1* (TOR inhibitors; ref. 21), *BRAF* (MAPK pathway inhibitors; ref. 22, 23) and *BRCA1* (PARP inhibitors; ref. 24). Other noteworthy

alterations included a nonsense mutation in *MSH2*, which is diagnostic for hereditary nonpolyposis coli and is a prognostic marker in colon cancer, and a nonsense mutation in *SMAD2*, which has been suggested to be associated with advanced disease and decreased survival in colon cancer (25).

Plausibly actionable amplifications of both *FGFR1* and *CCND1* were observed in a breast tumor sample (Fig. 2A). In preclinical studies, *FGFR1* amplification was shown to predict resistance to hormonal therapy in breast cancer (26) and thus may be considered a candidate tier 1 copy-number event for this FDA-approved indication. Clinical trials are currently underway to test FGFR inhibitors against tumors with amplified or overexpressed FGFR1, making *FGFR1* amplification a tier 2 actionable variant as well. Amplification of *CCND1* (which encodes the cyclin D1 cell-cycle regulator) has also been suggested to predict resistance to hormonal therapy (27, 28). Moreover, this alteration may predict sensitivity to cyclin-dependent kinase inhibitors (tier 2 actionable event; ref. 29), as well as overall disease prognosis in patients with breast cancer (prognostic alteration; refs. 27, 28, 30). Lower-level copy-number alterations (between 2- and 3-fold relative changes) were observed in several known or putative cancer genes, including *CDK8*, *GNAS*, *MYC*, and *SRC*. Although these events are most likely to reflect aneuploidy, some may represent higher level copy-number alterations in samples with low tumor purity.

Table 2. Actionable or prognostic genomic alterations in 10 FFPE tumor samples

Sample	Tumor type	Mean target coverage	Actionable in principle		Prognostic/diagnostic
			Tier 1	Tier 2	
FFPE 1	Colon	457	<i>KRAS</i> (Q61H) ^b	<i>KRAS</i> (Q61H)^a	
FFPE 2	Colon	353	<i>KRAS</i> (G13C)^a	<i>KRAS</i> (G13C)^a	
FFPE 3	Colon	498	<i>KRAS</i> (G13C)^a <i>PIK3CA</i> (H1047R) ^b	<i>KRAS</i> (G13C)^a <i>PIK3CA</i> (H1047R)^a	<i>MSH2</i> (R680*)^a
FFPE 4	Colon	300		<i>BRAF</i> (D594G) ^d <i>TSC1</i> (E258*) ^c	
FFPE 5	Breast	472	<i>CCND1</i> amp ^b <i>FGFR1</i> amp ^d	<i>CCND1</i> amp ^d <i>FGFR1</i> amp^a	<i>CCND1</i> amp ^b
FFPE 6	Breast	532		<i>BRCA1</i> (2-bp del)^a	<i>BRCA1</i> (2-bp del)^a
FFPE 7	Colon	250	<i>KRAS</i> (G13D)^a <i>PIK3CA</i> (E545K) ^b	<i>KRAS</i> (G13D)^a <i>PIK3CA</i> (E545K)^a	
FFPE 8	Breast	537	<i>PIK3CA</i> (H1047R) ^b	<i>PIK3CA</i> (H1047R)^a	
FFPE 9	Colon	116			<i>SMAD2</i> (S306*) ^b
FFPE 10	Colon	410	<i>KRAS</i> (Q61H) ^b	<i>KRAS</i> (Q61H)^a	

Abbreviations: amp, amplification; del, deletion.

The level of evidence for each actionable alteration is denoted by the following footnotes:

^aClinically validated and approved alterations (for tier 1 or prognostic/diagnostic) or specifically targeted alterations (for tier 2), shown in bold.

^bLimited clinical evidence.

^cClinical evidence in a different tumor type only.

^dPreclinical evidence only.

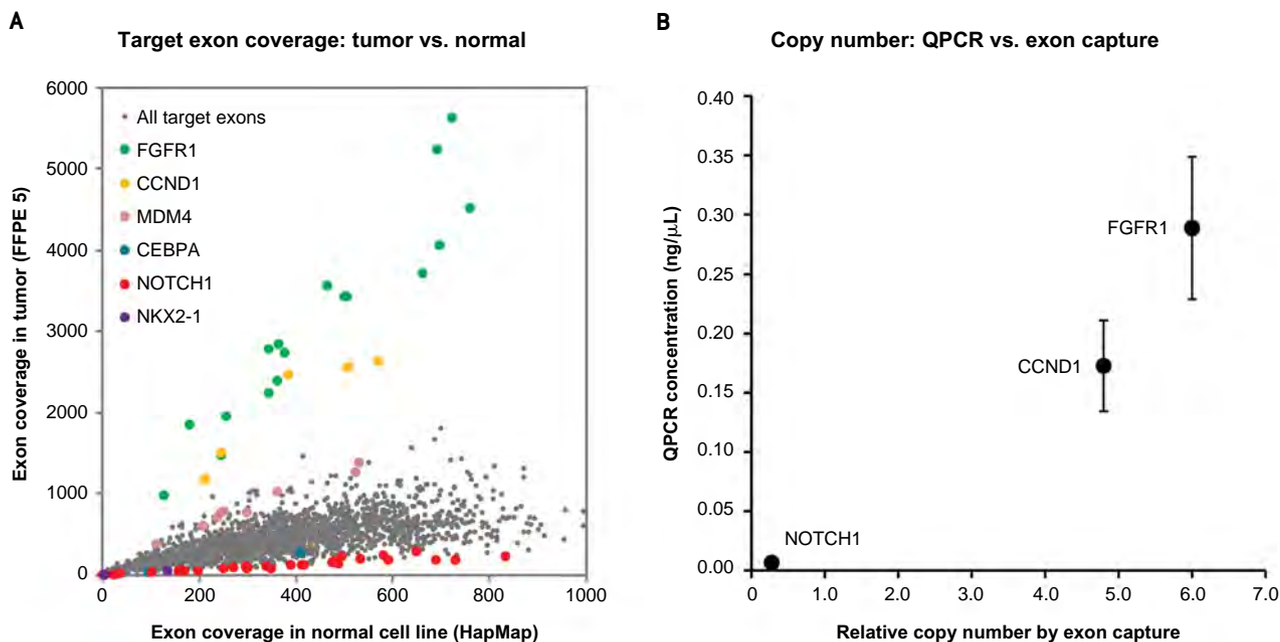


Figure 2. Copy-number alterations in an archival breast cancer sample. **A**, sequence coverage is shown for each target in the tumor sample compared with a normal diploid sample. Exon targets from several genes with copy-number gains and losses are highlighted. **B**, copy-number correlation between exon capture and QPCR in sample FFPE 5. Quantitative PCR of *FGFR1*, *CCND1*, and *NOTCH1* with 3 independent sets of primers was performed and average values for each gene were compared to exon capture copy-number.

Examination of 79 pharmacogenomic loci facilitated inspection of plausibly actionable polymorphisms (Supplementary Table S9). The *ERCC2-K751QC* allele, associated with increased risk of FOLFOX-induced grade 3 or 4 hematologic toxicity (31), was present in 2 samples (i.e., FFPE 2 and FFPE 9). The *UGT1A1-G3156A* allele was found to be heterozygous in 5 samples but homozygous in none of them. This allele is associated with irinotecan-related neutropenia when present as a homozygous event (32).

To validate these findings, a representative subset of alterations (31 nonsynonymous variants and 2 indels; samples 4–7) were independently queried by mass spectrometric genotyping (2, 3). All 31 single-nucleotide variants and 2 indels tested were confirmed, demonstrating 100% specificity of the targeted exon capture approach in the small subset examined. Copy-number alterations involving 3 genes that were amplified or deleted in sample FFPE 5 (*FGFR1*, *CCND1*, *NOTCH1*) were also tested by quantitative PCR with the use of 3 independent primer pairs for each gene. As shown in Fig. 2B, the quantitative PCR results were highly correlated to the copy-number ratios detected by targeted exon capture/sequencing in FFPE 5 ($r^2 = 0.94$). The correlation coefficient (r^2) for these same genes in sample FFPE 9—which has a 2.3-fold amplification of *FGFR1* but no copy-number changes in *CCND1* or *NOTCH1*—was 0.99 (Supplementary Fig. S3).

Comparison with an Existing Mutation Profiling Platform

We next wished to compare the sensitivity and specificity of targeted hybrid capture/sequencing to an existing mass spectrometric genotyping-based platform because this type of approach is currently being used in several clinical and translational oncology settings (2, 33–35). We thus performed OncoMap, a mass-spectrometric genotyping technology that interrogates more than 400 known mutations in 33 cancer genes. Of the 155 single-nucleotide variants seen by hybrid capture/sequencing of the FFPE samples described previously, 13 were also interrogated by assays present in OncoMap (Table 3). However, when OncoMap was performed on these samples, only 10 of these 13 mutations were detected. To determine the basis for this discrepancy, we assayed all 13 mutations by an orthogonal genotyping approach that uses distinct reagent chemistry (hME genotyping; see Methods). All 13 mutations were confirmed by this orthogonal genotyping method, suggesting that the 3 mutations not detected by OncoMap were false-negative results by mass spectrometric genotyping (Table 3; shown in bold). All mutations seen by OncoMap were also detected by targeted exon capture.

DISCUSSION

We have developed a targeted, massively parallel sequencing platform to detect actionable genomic alterations in

Table 3. Comparison of OncoMap and targeted exon capture profiling in FFPE samples

FFPE Sample	Sample type	Gene	Mutation	Seen via OncoMap	Seen via exon capture	Validated by hMe
FFPE 1	Colon	<i>KRAS</i>	Q61H	Yes	Yes	Yes
FFPE 2	Colon	<i>KRAS</i>	G13C	Yes	Yes	Yes
FFPE 3	Colon	<i>KRAS</i>	G13D	Yes	Yes	Yes
FFPE 3	Colon	<i>PIK3CA</i>	H1047R	Yes	Yes	Yes
FFPE 4	Colon	<i>BRAF</i>	D594G	No	Yes	Yes
FFPE 4	Colon	<i>APC</i>	Q1367*	No	Yes	Yes
FFPE 4	Colon	<i>APC</i>	Q1378*	Yes	Yes	Yes
FFPE 6	Breast	<i>TP53</i>	R248Q	Yes	Yes	Yes
FFPE 7	Colon	<i>KRAS</i>	G13D	Yes	Yes	Yes
FFPE 7	Colon	<i>PIK3CA</i>	E545K	Yes	Yes	Yes
FFPE 7	Colon	<i>CTNNB1</i>	S45F	No	Yes	Yes
FFPE 8	Breast	<i>PIK3CA</i>	H1047R	Yes	Yes	Yes
FFPE 10	Colon	<i>KRAS</i>	Q61H	Yes	Yes	Yes

NOTE: Mutations that were not detected by OncoMap are shown in bold.

clinical tumor samples. In this initial proof-of-concept effort, we sequenced 137 cancer genes from 10 pooled FFPE tumor DNA samples (plus 2 control samples) and achieved 391× mean coverage per sample within a single paired-end sequencing lane. This depth of coverage afforded robust, simultaneous detection of base mutations, indels, amplifications, and deletions. Thus, targeted massively parallel sequencing provides a unifying approach for detection of multiple categories of actionable genetic alterations.

In our pilot study, all of the tumor samples profiled contained biologically or clinically meaningful genomic alterations, including several that might predict sensitivity or resistance to targeted agents or provide useful prognostic information. In particular, 15 alterations (at least one per sample) were plausibly actionable, and might thus be predicted to impact clinical decision-making or clinical trial enrollment if identified as part of an experimental therapeutics or phase I trial program. Several actionable somatic alterations (*KRAS*, *PIK3CA*, and *MSH2*) were detected in samples with tumor purity as low as 10% to 20%, highlighting the utility of this approach in “real-world” clinical tumor samples.

Comparison with OncoMap, a mass spectrometric genotyping platform in current translational use, confirmed robust performance of targeted massively parallel sequencing, even when applied to FFPE tumor specimens. In our previous study, the sensitivity and specificity of OncoMap in FFPE tissue was 89.3% and 99.4%, respectively, based on a focused comparison with massively parallel sequencing of *KRAS* (codon 12) in 93 FFPE samples. In the current study, OncoMap detected 10 of 13 mutations (79% sensitivity) that

were seen by sequencing at multiple loci (including *KRAS*). The OncoMap approach involves iPLEX genotyping of >500 mutations followed by hMe validation of all candidates (see Methods)—the iPLEX method allows increased multiplexing, but in our hands has proved somewhat less sensitive than hMe genotyping. The fact that all 13 mutations were subsequently confirmed by hMe chemistry suggests that massively parallel sequencing to several hundred-fold mean coverage affords enhanced sensitivity compared to mass spectrometric genotyping. Moreover, most alterations found by sequencing are not assayed by genotyping or allele-specific PCR-based mutation profiling platforms. Thus, the sequencing-based approach may uncover more actionable options for patients than allele-specific approaches.

Hybrid selection approaches have been widely used to promote gene discovery by reducing genome complexity before sequencing (5). In this study, we adapted this technique to capture a highly restricted genomic territory composed of 137 known cancer genes and 400,000 coding bases. This afforded an expanded depth of coverage (to >400-fold) while also enabling multiple barcoded samples to be pooled within a single sequencing lane, thereby increasing throughput and lowering costs. We previously used a similar approach to characterize a frozen tumor sample from a patient with metastatic melanoma who developed resistance to the RAF-inhibitor vemurafenib, and identified an activating mutation in *MEK1* that caused resistance to RAF- and MEK-inhibition (36). Here, we have adapted the approach to capture and sequence multiple barcoded samples and to identify distinct categories of genomic alterations simultaneously.

An advantage of solution-phase hybrid capture is that redesign and synthesis of long oligonucleotides for bait generation is a straightforward process that may be performed iteratively until an optimal set of baits has been developed. Thus, prioritized genomic regions can be readily amended as new knowledge of cancer gene mutations becomes available. Furthermore, DNA barcoding and pooling decreases the sequencing cost per sample in a manner proportional to the number of pooled samples present within a sequencing lane. Achieving deep sequencing coverage increases the sensitivity of mutation detection—particularly in the setting of high stromal admixture, which can pervade clinical tumor tissue. As such, this study extends earlier barcoding and hybrid capture/sequencing efforts (36–49) by identifying multiple types of actionable somatic alterations in archival (i.e., FFPE) tumor specimens. Because most clinical samples are stored as FFPE material, this approach may prove suitable for many translational and clinical applications.

At the same time, variations in FFPE sample quality may adversely affect library construction, hybrid selection, or sequencing. Potential solutions include the incorporation of additional pre-processing steps to enrich for high-quality FFPE DNA, pooling of fewer samples prior to hybrid selection, and/or increasing the overall depth of sequencing if the starting library complexity is sufficiently high (50). The use of orthogonal technologies such as direct genotyping, quantitative PCR, or FISH to validate actionable alterations may prove useful in the short term because these techniques are used widely in existing clinical laboratories. However, if the superior sensitivity and specificity is confirmed in independent clinical studies, massively parallel sequencing may become increasingly used in diagnostic or Clinical Laboratory Improvement Amendments (CLIA) laboratory settings.

Several additional areas for technical and analytical optimization remain. Although we generally achieved robust sequence coverage of targeted regions, genomic territory with very high or very low GC content presents certain challenges. Options to improve coverage of these regions include redesign or inclusion of additional baits targeting regions that are difficult to capture. On the analytical side, detection of longer indels (such as the 9-bp *PIK3CA* deletion in the NCI-H69 cell line) remains difficult with current algorithms. Because actionable indels occur in multiple genes, including *EGFR*, *ERBB2*, and *KIT*, supplemental assays may be needed to ensure sensitive indel detection. Moreover, exon-directed capture approaches do not detect clinically relevant gene rearrangements such as those involving *ALK*, *ABL*, and *PDGFR*. One potential strategy to detect known rearrangements would involve design of baits tiled across common translocation breakpoints. Furthermore, whereas both amplifications and deletions could be detected in cell line DNA, such events were only observed in a single FFPE sample, which had 80% tumor purity. Detection of copy-number aberrations by targeted sequencing may be more problematic in samples with significant stromal contamination. Future analytical methods that incorporate allelic information to infer tumor purity may enhance detection of copy gains and losses in samples with variable tumor purity.

Emerging frameworks for clinical interpretation of genome sequencing data typically categorize alterations based on “actionability” or prognostic utility. Potentially actionable alterations may be further subdivided depending on the level of evidence about a particular alteration, ranging from those with established therapies to others with sound preclinical evidence. Plausibly actionable alterations may also include those for which the predictive implications within a particular cancer type are not known (e.g., *BRAF* mutations in lung cancer), or for which there is no established clinical proof of concept (e.g., *RET* mutations in lung cancer) even though a particular therapy against the target (sorafenib) may be commercially available. This category may also include mutations in tumor suppressor genes (e.g., *PTEN*) hypothesized to predict vulnerability to targeted agents (e.g., PI3K inhibitors).

More than 160 variants of unclear significance were identified in our sample set. Undoubtedly, many such variants represent uncharacterized germline polymorphisms. Differentiating somatic from germline alterations is readily accomplished by including matched normal samples (36), although paired normal material is not always available in research settings. Even among alterations that are clearly somatic, additional approaches to interpret their potential significance and communicate the results to clinicians and patients will be needed. Development of a rigorous formalism for clinical interpretation of complex genomic data will likely become an active research area, with the goal of enabling optimal, genomics-driven decision making for therapy or clinical trial enrollment.

Potential applications for targeted hybrid capture/massively parallel sequencing in translational and clinical oncology research include both retrospective and prospective profiling of tumor cohorts. Here, the goal may be to identify predictive and prognostic genes or validate pharmacogenomic polymorphisms. Ultimately, similar approaches may be used for prospective genomic profiling of cancer patients to guide clinical decision making. Toward this end, the potential turnaround time for the current approach is ~2 weeks. Emerging sequencing instruments promise vast reductions in turnaround time. Cost, a significant consideration in clinical sequencing, can also be reduced dramatically by sample pooling. Indeed, it is likely that a combination of multiplexing together with falling sequencing costs may ultimately eliminate cost as a limiting barrier to sequencing data generation.

In conclusion, the results described herein suggest that targeted, massively parallel sequencing offers a promising method to detect actionable genetic alterations across a large panel of cancer genes in the clinical diagnostic arena. If widely deployed, such implementation may open new opportunities to link cancer genomics with molecular features, clinical outcomes, and treatment response in a manner that empowers multiple directions in molecular cancer epidemiology. In addition, this approach may ultimately impact clinical practice by offering a categorical means to identify genetic changes affecting genes and pathways targeted by existing and emerging drugs, thereby speeding the advent of personalized cancer medicine.

METHODS

High-Throughput, Targeted Deep Sequencing: Overview

Massively parallel sequencing libraries (Illumina) that contain bar-coded universal primers (9) were generated with the use of genomic DNA from formalin-fixed, paraffin-embedded tumor material. After preamplification and DNA quantification, equimolar pools were generated consisting of 12 barcoded tumor DNAs. These DNA pools were subjected to solution-phase hybrid capture with biotinylated RNA baits targeting all exons from 137 actionable cancer genes. Each hybrid capture reaction was sequenced in a single paired-end lane of an Illumina flow cell. Subsequently, the sequencing data were deconvoluted to match all high-quality barcoded reads with the corresponding tumor samples, and genomic alterations (single-nucleotide sequence variants, small insertions/deletions, and DNA copy-number alterations) were identified. The approach is illustrated schematically in Supplementary Figure S1.

Tumor Tissue and Cell Line DNA

Discarded and de-identified tumor specimens were obtained from the Cooperative Human Tissue Network. An exemption from the Institutional Review Board was obtained for all samples from the Dana-Farber/Partners Cancer Care Office for the Protection of Research Subjects (Protocol 10-380). Genomic DNA was extracted from tumor tissue using methods previously described (2). Cell line genomic DNA was purchased directly from the American Type Culture Collection (ATCC). Authentication of cell line genomic DNA was performed by ATCC by the use of short tandem repeat profiling, which uses multiplex PCR to simultaneously amplify the amelogenin gene and 8 of the most informative polymorphic markers in the human genome. Control genomic DNA was from the HapMap consortium, which was purchased from the Coriell Institute for Medical Research.

Barcoded Genomic DNA Library Construction

Genomic DNA was quantified by the use of Quant-iT PicoGreen® dsDNA Assay Kit (Invitrogen, Carlsbad, CA). A total of 1 µg of genomic DNA from each sample was sheared by sonication with the following conditions: duty cycle 10%, intensity 5, cycles per burst 200, and 135 seconds (Covaris S2 instrument). Paired-end adapters for massively parallel sequencing (Illumina) were added as previously described (51), with the following modifications to the paired end library preparation step (basic protocol 2). First, the multiplex adapter provided with the Multiplex Paired-End Library Sample Preparation Kit (Illumina) was used instead of the standard paired-end adapter. Second, PCR enrichment was conducted in 150 µL of total volume with 3 primers from the Multiplexing Sample Preparation Oligonucleotide Kit (Illumina). Each PCR enrichment reaction contained 75 µL of Phusion polymerase (Finnzymes), 3 µL of Multiplexing PE Primer 1.0 (25 µM), 3 µL of Multiplexing PE Primer 2.0 (0.5 µM), 3 µL of an Index primer (25 µM), 36 µL of paired-end library, and 30 µL of nuclease-free water. Samples were denatured for 5 minutes at 95°C; 18 cycles of 10 seconds at 95°C, 30 seconds at 65°C, and 30 seconds at 72°C; and a final 5 minutes at 72°C before cooling to 4°C. PCR primers were removed by using ×1.8 volume of Agencourt AMPure PCR Purification Kit (Agencourt Bioscience Corporation).

Selection of Targeted Genes

We identified 137 genes that are biologically or clinically relevant in cancer, including targets of new and existing therapies, genes that predict sensitivity or resistance to therapies, genes that are prognostic markers, and oncogenes and tumor suppressors that are known to undergo recurrent somatic genomic alterations in cancer (Supplementary Table S1). These genes were identified by mining

existing databases including the Catalogue of Somatic Mutations in Cancer (10) and The Cancer Genome Atlas (52). In addition, we identified 79 pharmacogenomic polymorphisms described in the literature, which might predict sensitivity or resistance to conventional cancer therapies (Supplementary Table S2).

Biotinylated RNA Baits

The Agilent SureSelect E-array program was used to design 7,021 unique RNA baits corresponding to the coding sequence of the 137 genes described previously, as well as to the 79 pharmacogenomic polymorphisms and to 24 SNPs for fingerprinting. Target loci were covered with a tiling density of ×2. Baits were replicated 8 times on the 55,000-bait library array. The sequences of all 7,021 baits are listed in the Supplementary Appendix. Biotinylated RNA baits were synthesized by Agilent for the SureSelect Target Enrichment system.

Pooling and Hybrid Capture

DNA libraries were pooled by mixing 300 ng of each library in a single 1.5-mL polypropylene sample tube, lyophilizing by the use of a speedvac evaporator, and resuspending in 4 µL of nuclease-free water. This entire amount (3,600 ng DNA in 4 µL) was used for hybrid selection. Solution-phase hybrid capture was performed as previously described (51) with 3 modifications to the hybrid selection step (Basic Protocol 3). First, instead of 1.5 µL of Blocking Oligo 2.0, 0.125 µL of each of 12 additional 200 µM blocking oligonucleotides with sequences complementary to the barcodes were added to the hybridization reaction (see the Supplementary Methods for sequences). Second, the biotinylated oligonucleotide baits were diluted 1:8 with nuclease-free water from a concentration of 100 ng/µL to 12.5 ng/µL immediately before hybridization and 5 µL of this solution was added to the hybridization reaction.

The final volume of the hybridization reaction was 19 µL, consisting of the following components: 4 µL of pooled DNA libraries, 2.5 µL of 1.0 mg/mL human Cot-1 DNA, 2.5 µL of 10.0 mg/mL salmon sperm DNA, 1.5 µL of 200 µM blocking oligo 1.0, 1.5 µL of total of the twelve 200 µM blocking oligonucleotides, 5.0 µL of 12.5 ng/µL biotinylated oligonucleotide baits, 1.0 µL of 20 U/µL Superase-In RNase inhibitor, and 1 µL of nuclease-free water. Third, during PCR enrichment of the captured DNA (“the catch”), PCR was performed with primers P5 (5'-AAT GAT ACG GCG ACC ACC GA-3') and P7 (5'-CAA GCA GAA GAC GGC ATA CGA-3'), both at 100 µM, instead of PCR primers PE1.0 and PE2.0. PCR conditions remained as described. All custom primers were obtained from Integrated DNA Technologies (IDT).

Sequencing and Analysis

We sequenced 100 bases from both ends of library DNA fragments by using an Illumina HiSeq 2000 instrument. The sequence reads were aligned to human reference genome hg18 with the Burrows-Wheeler Alignment tool (53) with use of the following parameters: -q 5 -l 32 -k 2 -o 1. Artfactual duplicate read pairs were removed with Picard tools (picard.sourceforge.net). An average of 450 megabases of aligned sequence was generated for each library.

Single-nucleotide variants and small insertions/deletions were identified by the use of algorithms from the Genome Analysis Toolkit developed at the Broad Institute (54). A local multiple sequence alignment was performed on intervals suspected to harbor indels to derive the most probable underlying genomic structure of the query sample. Single-nucleotide variants were called separately on each sample with UnifiedGenotyper and annotated with GenomicAnnotator. Variants were discarded if they were present in dbSNP and not in the COSMIC database (10), they exhibited an unfavorable strand balance score (> -20), or they were detected in the HapMap normal control. Novel recurrent single-nucleotide variants were manually reviewed

to eliminate additional systematic artifacts. Indels were called with IndelGenotyperV2 and were retained if they occurred in protein-coding exons and on both DNA strands, in <2% of reads in the HapMap normal control, and were absent from dbSNP.

To calculate relative copy-number levels of the 137 target gene loci, we computed the mean sequence coverage for each gene across all protein-coding exons by using the DepthOfCoverage tool in the Genome Analysis Toolkit. All bases in reads with mapping quality <5 were ignored, as were any additional bases with base quality <5. Gene-level coverage in each tumor was normalized by the gene-level coverage for an indexed HapMap diploid cell line included in the same pooled hybrid selection experiment (after adjusting for differences in the overall amount of aligned sequence per sample). Sequence-derived estimates of copy number were then compared to SNP array-derived estimates of copy number for the cancer cell lines.

Mass Spectrometric Genotyping

Mass spectrometric genotyping was performed with the OncoMap 3.0 platform as previously described (2, 3) with iPLEX chemistry for initial mutation profiling and validation by multibase hME extension chemistry. Genomic DNA from all tumor samples was quantified using Quant-iT PicoGreen® dsDNA Assay Kit (Invitrogen).

To validate alterations detected by massively parallel sequencing that were not included in the OncoMap assay collection, base substitutions and indels were queried using multi-base hME extension chemistry with plexing of ≤6 assays per pool. Conditions for hME validation were implemented as described previously (2, 3). Primers and probes used for hME validation were designed with the Sequenom MassARRAY Assay Design 3.0 software, applying default multi-base extension parameters but with the following modifications: maximum multiplex level input equal to 6; maximum pass iteration base adjusted to 200.

Microarray Analysis of Chromosomal Copy Number

Chromosomal copy-number information was obtained from the Broad-Novartis Cancer Cell Line Encyclopedia project, which has high-density SNP array data from the Affymetrix SNP 6.0 platform for all cancer cell lines profiled in this study (55).

Quantitative PCR Analysis of Chromosomal Copy Number

Quantitative PCR was performed with the SYBR Green PCR Master Mix Kit (Applied Biosystems) according to the manufacturer's instructions. To determine the chromosomal copy number of each gene, 3 sets of gene-specific primers were designed to interrogate the genetic locus. Primers recognizing LINE sequences were used for reference amplification/normalization as described previously (56). Primer sequences are provided in the Supplementary Methods. Male genomic DNA (Promega) was included as a standard, and HapMap DNA (Coriell) was used as a normal diploid control. Quantitative PCRs were performed in triplicate for each sample using an ABI 7300 instrument, in 25- μ L reactions containing 0.5 ng of genomic DNA and forward and reverse primers each at a concentration of 600 nM.

Disclosure of Potential Conflicts of Interest

Consultant/advisory role: Foundation Medicine (N. Wagle, M.F. Berger, M.J. Davis, M. Meyerson, L.A. Garraway), Novartis (W.C. Hahn, M. Meyerson, L.A. Garraway), Daiichi Sankyo (L.A. Garraway). Ownership interest: Foundation Medicine (N. Wagle, M. Meyerson, L.A. Garraway). Research support: Novartis (W.C. Hahn, M. Meyerson, L.A. Garraway). Patents: Laboratory Corporation of America (M. Meyerson). Honoraria: Illumina (M.F. Berger).

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REFERENCES

1. Macconail LE, Garraway LA. Clinical implications of the cancer genome. *J Clin Oncol* 2010;28:5219–28.
2. MacConaill LE, Campbell CD, Kehoe SM, Bass AJ, Hatton C, Niu L, et al. Profiling critical cancer gene mutations in clinical tumor samples. *PLoS One* 2009;4:e7887.
3. Thomas RK, Baker AC, Debiase RM, Winckler W, Laframboise T, Lin WM, et al. High-throughput oncogene mutation profiling in human cancer. *Nat Genet* 2007;39:347–51.
4. Dias-Santagata D, Akhavanfard S, David SS, Vernovsky K, Kuhlmann G, Boisvert SL, et al. Rapid targeted mutational analysis of human tumours: a clinical platform to guide personalized cancer medicine. *EMBO Mol Med* 2010;2:146–58.
5. Meyerson M, Gabriel S, Getz G. Advances in understanding cancer genomes through second-generation sequencing. *Nat Rev Genet* 2010;11:685–96.
6. Stratton MR, Campbell PJ, Futreal PA. The cancer genome. *Nature* 2009;458:719–24.
7. Gnirke A, Melnikov A, Maguire J, Rogov P, LeProust EM, Brockman W, et al. Solution hybrid selection with ultra-long oligonucleotides for massively parallel targeted sequencing. *Nat Biotechnol* 2009;27:182–9.
8. Hodges E, Rooks M, Xuan Z, Bhattacharjee A, Benjamin Gordon D, Brizuela L, et al. Hybrid selection of discrete genomic intervals on custom-designed microarrays for massively parallel sequencing. *Nat Protoc* 2009;4:960–74.
9. Craig DW, Pearson JV, Szelinger S, Sekar A, Redman M, Corneveaux JJ, et al. Identification of genetic variants using bar-coded multiplexed sequencing. *Nat Methods* 2008;5:887–93.
10. Forbes SA, Bindal N, Bamford S, Cole C, Kok CY, Beare D, et al. COSMIC: mining complete cancer genomes in the Catalogue of Somatic Mutations in Cancer. *Nucleic Acids Res* 2011;39(Database issue):D945–50.
11. Ogata H, Sato H, Takatsuka J, De Luca LM. Human breast cancer MDA-MB-231 cells fail to express the neurofibromin protein, lack its type I mRNA isoform and show accumulation of P-MAPK and activated Ras. *Cancer Lett* 2001;172:159–64.
12. Allegra CJ, Jessup JM, Somerfield MR, Hamilton SR, Hammond EH, Hayes DF, et al. American Society of Clinical Oncology provisional clinical opinion: testing for KRAS gene mutations in patients with metastatic colorectal carcinoma to predict response to anti-epidermal growth factor receptor monoclonal antibody therapy. *J Clin Oncol* 2009;27:2091–6.
13. Bardelli A, Siena S. Molecular mechanisms of resistance to cetuximab and panitumumab in colorectal cancer. *J Clin Oncol* 2010;28:1254–61.
14. Sartore-Bianchi A, Di Nicolantonio F, Nichelatti M, Molinari F, De Dosso S, Saletti P, et al. Multi-determinants analysis of molecular alterations for predicting clinical benefit to EGFR-targeted monoclonal antibodies in colorectal cancer. *PLoS One* 2009;4:e7287.
15. Sartore-Bianchi A, Martini M, Molinari F, Veronese S, Nichelatti M, Artale S, et al. PIK3CA mutations in colorectal cancer are associated with clinical resistance to EGFR-targeted monoclonal antibodies. *Cancer Res* 2009;69:1851–7.
16. De Roock W, Claes B, Bernasconi D, De Schutter J, Biesmans B, Fountzilias G, et al. Effects of KRAS, BRAF, NRAS, and PIK3CA mutations on the efficacy of cetuximab plus chemotherapy in chemotherapy-refractory metastatic colorectal cancer: a retrospective consortium analysis. *Lancet Oncol* 2010;11:753–62.
17. Berns K, Horlings HM, Hennessy BT, Madiredjo M, Hijmans EM, Beelen K, et al. A functional genetic approach identifies the PI3K pathway as a major determinant of trastuzumab resistance in breast cancer. *Cancer Cell* 2007;12:395–402.

18. Wang L, Zhang Q, Zhang J, Sun S, Guo H, Jia Z, et al. PI3K pathway activation results in low efficacy of both trastuzumab and lapatinib. *BMC Cancer* 2011;11:248.
19. Courtney KD, Corcoran RB, Engelman JA. The PI3K pathway as drug target in human cancer. *J Clin Oncol*. 2010;28:1075–83.
20. Duffy A, Kummar S. Targeting mitogen-activated protein kinase kinase (MEK) in solid tumors. *Target Oncol* 2009;4:267–73.
21. Wagner AJ, Malinowska-Kolodziej J, Morgan JA, Qin W, Fletcher CD, Vena N, et al. Clinical activity of mTOR inhibition with sirolimus in malignant perivascular epithelioid cell tumors: targeting the pathogenic activation of mTORC1 in tumors. *J Clin Oncol* 2010;28:835–40.
22. Paik PK, Arcila ME, Fara M, Sima CS, Miller VA, Kris MG, et al. Clinical characteristics of patients with lung adenocarcinomas harboring BRAF mutations. *J Clin Oncol* 2011;29:2046–51.
23. Smalley KS, Xiao M, Villanueva J, Nguyen TK, Flaherty KT, Letrero R, et al. CRAF inhibition induces apoptosis in melanoma cells with non-V600E BRAF mutations. *Oncogene* 2009;28:85–94.
24. Fong PC, Boss DS, Yap TA, Tutt A, Wu P, Mergui-Roelvink M, et al. Inhibition of poly(ADP-ribose) polymerase in tumors from BRCA mutation carriers. *N Engl J Med* 2009;361:123–34.
25. Xie W, Rimm DL, Lin Y, Shih WJ, Reiss M. Loss of Smad signaling in human colorectal cancer is associated with advanced disease and poor prognosis. *Cancer J* 2003;9:302–12.
26. Turner N, Pearson A, Sharpe R, Lambros M, Geyer F, Lopez-Garcia MA, et al. FGFR1 amplification drives endocrine therapy resistance and is a therapeutic target in breast cancer. *Cancer Res* 2010;70:2085–94.
27. Bostner J, Ahnstrom Waltersson M, Fornander T, Skoog L, Nordenskjold B, Stal O. Amplification of CCND1 and PAK1 as predictors of recurrence and tamoxifen resistance in postmenopausal breast cancer. *Oncogene* 2007;26:6997–7005.
28. Lundgren K, Holm K, Nordenskjold B, Borg A, Landberg G. Gene products of chromosome 11q and their association with CCND1 gene amplification and tamoxifen resistance in premenopausal breast cancer. *Breast Cancer Res* 2008;10:R81.
29. Finn RS, Dering J, Conklin D, Kalous O, Cohen DJ, Desai AJ, et al. PD 0332991, a selective cyclin D kinase 4/6 inhibitor, preferentially inhibits proliferation of luminal estrogen receptor-positive human breast cancer cell lines in vitro. *Breast Cancer Res* 2009;11:R77.
30. Roy PG, Pratt N, Purdie CA, Baker L, Ashfield A, Quinlan P, et al. High CCND1 amplification identifies a group of poor prognosis women with estrogen receptor positive breast cancer. *Int J Cancer* 2010;127:355–60.
31. Boige V, Mendiboune J, Pignon JP, Loriot MA, Castaing M, Barrois M, et al. Pharmacogenetic assessment of toxicity and outcome in patients with metastatic colorectal cancer treated with LV5FU2, FOLFOX, and FOLFIRI: FFOCD 2000–05. *J Clin Oncol* 2010;28:2556–64.
32. Lara PN, Jr., Natale R, Crowley J, Lenz HJ, Redman MW, Carleton JE, et al. Phase III trial of irinotecan/cisplatin compared with etoposide/cisplatin in extensive-stage small-cell lung cancer: clinical and pharmacogenomic results from SWOG S0124. *J Clin Oncol* 2009;27:2530–5.
33. Arcila M, Lau C, Nafa K, Ladanyi M. Detection of KRAS and BRAF mutations in colorectal carcinoma roles for high-sensitivity locked nucleic acid-PCR sequencing and broad-spectrum mass spectrometry genotyping. *J Mol Diagn* 2011;13:64–73.
34. Brevet M, Johnson ML, Azzoli CG, Ladanyi M. Detection of EGFR mutations in plasma DNA from lung cancer patients by mass spectrometry genotyping is predictive of tumor EGFR status and response to EGFR inhibitors. *Lung Cancer* 2011;73:96–102.
35. Beadling C, Heinrich MC, Warrick A, Forbes EM, Nelson D, Justusson E, et al. Multiplex mutation screening by mass spectrometry: evaluation of 820 cases from a personalized cancer medicine registry. *J Mol Diagn*. 2011;13:504–13.
36. Wagle N, Emery C, Berger MF, Davis MJ, Sawyer A, Pochanard P, et al. Dissecting therapeutic resistance to raf inhibition in melanoma by tumor genomic profiling. *J Clin Oncol* 2011;29:3085–96.
37. Bell CJ, Dinwiddie DL, Miller NA, Hateley SL, Ganusova EE, Mudge J, et al. Carrier testing for severe childhood recessive diseases by next-generation sequencing. *Sci Transl Med* 2011;3:65ra4.
38. Berg JS, Evans JP, Leigh MW, Omran H, Bizon C, Mane K, et al. Next generation massively parallel sequencing of targeted exomes to identify genetic mutations in primary ciliary dyskinesia: implications for application to clinical testing. *Genet Med* 2011;13:218–29.
39. Chmielecki J, Peifer M, Jia P, Socci ND, Hutchinson K, Viale A, et al. Targeted next-generation sequencing of DNA regions proximal to a conserved GXGXXG signaling motif enables systematic discovery of tyrosine kinase fusions in cancer. *Nucleic Acids Res* 2010;38:6985–96.
40. Grossmann V, Kohlmann A, Klein HU, Schindela S, Schnittger S, Dicker F, et al. Targeted next-generation sequencing detects point mutations, insertions, deletions and balanced chromosomal rearrangements as well as identifies novel leukemia-specific fusion genes in a single procedure. *Leukemia* 2011;25:671–80.
41. Johansson H, Isaksson M, Sorqvist EF, Roos F, Stenberg J, Sjoblom T, et al. Targeted resequencing of candidate genes using selector probes. *Nucleic Acids Res* 2011;39:e8.
42. Kiialainen A, Karlberg O, Ahlford A, Sigurdsson S, Lindblad-Toh K, Syvanen AC. Performance of microarray and liquid based capture methods for target enrichment for massively parallel sequencing and SNP discovery. *PLoS One* 2011;6:e16486.
43. Liao GJ, Lun FM, Zheng YW, Chan KC, Leung TY, Lau TK, et al. Targeted massively parallel sequencing of maternal plasma DNA permits efficient and unbiased detection of fetal alleles. *Clin Chem* 2011;57:92–101.
44. Meder B, Haas J, Keller A, Heid C, Just S, Borries A, et al. Targeted next-generation sequencing for the molecular genetic diagnostics of cardiomyopathies. *Circ Cardiovasc Genet* 2011;4:110–22.
45. Nijman IJ, Mokry M, van Boxtel R, Toonen P, de Bruijn E, Cuppen E. Mutation discovery by targeted genomic enrichment of multiplexed barcoded samples. *Nat Methods* 2010;7:913–5.
46. Shearer AE, DeLuca AP, Hildebrand MS, Taylor KR, Gurrola J, 2nd, Scherer S, et al. Comprehensive genetic testing for hereditary hearing loss using massively parallel sequencing. *Proc Natl Acad Sci U S A* 2010;107:21104–9.
47. Summerer D, Schracke N, Wu H, Cheng Y, Bau S, Stahler CF, et al. Targeted high throughput sequencing of a cancer-related exome subset by specific sequence capture with a fully automated microarray platform. *Genomics* 2010;95:241–6.
48. Teer JK, Bonnycastle LL, Chines PS, Hansen NF, Aoyama N, Swift AJ, et al. Systematic comparison of three genomic enrichment methods for massively parallel DNA sequencing. *Genome Res* 2010;20:1420–31.
49. Zang ZJ, Ong CK, Cutcutache I, Yu W, Zhang SL, Huang D, et al. Genetic and structural variation in the gastric cancer kinome revealed through targeted deep sequencing. *Cancer Res* 2011;71:29–39.
50. Fisher S, Barry A, Abreu J, Minie B, Nolan J, Delorey TM, et al. A scalable, fully automated process for construction of sequence-ready human exome targeted capture libraries. *Genome Biol* 2011;12(1):R1.
51. Blumenstiel B, Cibulskis K, Fisher S, DeFelice M, Barry A, Fennell T, et al. Targeted exon sequencing by in-solution hybrid selection. *Curr Protoc Hum Genet* 2010;Chapter 18:Unit 18.4.
52. The Cancer Genome Atlas. Available from: cancergenome.nih.gov.
53. Li H, Durbin R. Fast and accurate short read alignment with Burrows-Wheeler transform. *Bioinformatics* 2009;25:1754–60.
54. McKenna A, Hanna M, Banks E, Sivachenko A, Cibulskis K, Kernytsky A, et al. The Genome Analysis Toolkit: a MapReduce framework for analyzing next-generation DNA sequencing data. *Genome Res* 2010;20:1297–303.
55. Broad-Novartis Cancer Cell Line Encyclopedia. Available from: www.broadinstitute.org/ccle.
56. Garraway LA, Widlund HR, Rubin MA, Getz G, Berger AJ, Ramaswamy S, et al. Integrative genomic analyses identify MITF as a lineage survival oncogene amplified in malignant melanoma. *Nature* 2005;436:117–22.