

Preliminary Communication

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Mutation Detection in Patients With Advanced Cancer by Universal Sequencing of Cancer-Related Genes in Tumor and Normal DNA vs Guideline-Based Germline Testing

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Key Points

Question How many additional individuals with inherited cancer-predisposing mutations might be detected by DNA sequencing of tumor and normal tissue in patients with advanced cancer compared with restricting genetic testing to clinical guideline–directed testing?

Findings In this case series of 1040 patients with advanced cancer, 101 of 182 patients with clinically actionable inherited mutations detected by tumor-normal sequencing would not have been detected by guideline-directed testing based on family history, age, and tumor type.

Meaning In selected populations of patients with advanced cancer, universal sequencing of germline and tumor DNA for a broad panel of cancer-related genes may detect more potentially clinically significant heritable mutations than a targeted approach based on current clinical guidelines. It is not known if this will result in improved outcomes.

Abstract

Importance Guidelines for cancer genetic testing based on family history may miss clinically actionable genetic changes with established implications for cancer screening or prevention.

Objective To determine the proportion and potential clinical implications of inherited variants detected using simultaneous sequencing of the tumor and normal tissue (“tumor-normal sequencing”) compared with genetic test results based on current guidelines.

Design, Setting, and Participants From January 2014 until May 2016 at Memorial Sloan Kettering Cancer Center, 10 336 patients consented to tumor DNA sequencing. Since May 2015, 1040 of these patients with advanced cancer were referred by their oncologists for germline analysis of 76 cancer predisposition genes. Patients with clinically actionable inherited mutations whose genetic test results would not have been predicted by published decision rules were identified. Follow-up for potential clinical implications of mutation detection was through May 2017.

Exposure Tumor and germline sequencing compared with the predicted yield of targeted germline sequencing based on clinical guidelines.

Main Outcomes and Measures Proportion of clinically actionable germline mutations detected by universal tumor-normal sequencing that would not have been detected by guideline-directed testing.

Results Of 1040 patients, the median age was 58 years (interquartile range, 50.5-66 years), 65.3% were male, and 81.3% had stage IV disease at the time of genomic analysis, with prostate, renal, pancreatic, breast, and colon cancer as the most common diagnoses. Of the 1040 patients, 182 (17.5%; 95% CI, 15.3%-19.9%) had clinically actionable mutations conferring cancer susceptibility, including 149 with moderate- to high-penetrance mutations; 101 patients tested (9.7%; 95% CI, 8.1%-11.7%) would not have had these mutations detected using clinical guidelines, including 65 with moderate- to high-penetrance mutations. Frequency of inherited mutations was related to case mix, stage, and founder mutations. Germline findings led to discussion or initiation of change to targeted therapy in 38 patients tested (3.7%) and predictive testing in the families of 13 individuals (1.3%), including 6 for whom genetic evaluation would not have been initiated by guideline-based testing.

Conclusions and Relevance In this referral population with selected advanced cancers, universal sequencing of a broad panel of cancer-related genes in paired germline and tumor DNA samples was associated with increased detection of individuals with potentially clinically significant heritable mutations over the predicted yield of targeted germline testing based on current clinical guidelines. Knowledge of these additional mutations can help guide therapeutic and preventive interventions, but whether all of these interventions would improve outcomes for patients with cancer or their family members requires further study.

Trial Registration [clinicaltrials.gov Identifier: NCT01775072](https://clinicaltrials.gov/ct2/show/study/NCT01775072)

Introduction

In addition to guiding therapy, simultaneous DNA sequence analysis of tumor-normal pairs (“tumor-normal sequencing”) reveals inherited cancer predisposition mutations in 3% to 12.6% of pediatric and adult patients with cancer.¹⁻⁷ It remains unknown, however, how many inherited mutations would be detected by multigene tumor-normal analysis compared with a traditional family history–based approach to genetic counseling and testing, and what the clinical implications of these findings would be. Selection for genetic testing is traditionally based on pathologic features of the tumor, age at diagnosis, family history of cancer, and other factors represented in clinical

practice guidelines.⁸⁻¹⁰ Studies have not determined whether inherited mutations found by tumor-normal sequencing would have been detected by traditional approaches to selection for genetic testing.¹⁻⁵

This study presents the results of analyses of inherited (“germline”) DNA performed in a prospective analysis of patients with advanced cancer tested by a targeted tumor-normal sequencing panel as previously described.^{11,12} The goals of the study were to determine the incremental proportion of clinically actionable mutations detected by concurrent germline analysis in patients with cancer undergoing universal tumor profiling compared with selective germline testing based on existing practice guidelines and to assess the association of identified mutations with therapeutic management and targeted cancer prevention in family members.

Methods

Patient Cohort

The cohort comprised patients with advanced cancer at Memorial Sloan Kettering Cancer Center undergoing tumor and normal DNA sequencing using MSK-IMPACT (Memorial Sloan Kettering-Integrated Mutation Profiling of Actionable Cancer Targets), a 410-gene panel.^{1,11,12} From May 2015, in the context of an institutional review board–approved protocol, patients with selected tumor types were prospectively offered secondary germline analysis after consenting to tumor genetic analysis. Germline analysis included 76 genes on the MSK-IMPACT panel associated with hereditary cancer predisposition, including all cancer-predisposing genes identified in the American College of Medical Genetics and Genomics guidelines.¹³⁻¹⁵ Nine mutation carriers were included in a prior study,¹ and 23 and 51 mutation carriers were included in recent series of prostate cancer,^{16,17} although clinical annotation including family history and treatment information was not available for those articles.

Ascertainment, Consent, Sequencing, Variant Calling, and Results Reporting

Through their physicians, patients were offered participation in a research study, using a video consent aid explaining risks and benefits of testing for inherited mutations (germline testing). Eligibility was restricted to those who also consented to tumor sequencing, with emphasis on patients with advanced (stage III-IV) disease; however, physicians had discretion for patient referral. Genetic testing reports were issued to the medical record, and

individuals with pathogenic or likely pathogenic variants (henceforth referred to as “pathogenic variants”) were invited for genetic counseling where at-risk family members were identified (eAppendix 1 in [Supplement 1](#)). Tumor DNA and nontumor DNA were sequenced and variants were reported as described previously¹¹ (eAppendix 2 in the [Supplement 1](#)). Variants were interpreted based on American College of Medical Genetics and Genomics criteria¹⁸ by a clinical molecular geneticist or molecular pathologist; variants of unknown significance (eTable in [Supplement 2](#)) were not included in clinical reports. Loss of heterozygosity (LOH), defined as loss of the normal allele in the tumor at the locus of the inherited mutation, was assessed by the FACETS algorithm as published previously¹⁹; hypermutated status was defined as 20 or more mutations; and microsatellite instability high (MSI-H) was defined as more than 10% of loci on the MSK-IMPACT panel demonstrating microsatellite instability.²⁰

In this study, not all pathogenic mutations (associated with disease causation) were considered clinically actionable. Clinical actionability of pathogenic variants was defined by evidence for their utility in cancer prevention (eAppendix 3 in [Supplement 1](#)) and/or potential utility as therapeutic targets. For this study, all germline mutations of established low, moderate, or high risk (penetrance) were considered clinically actionable^{8-10,16,21,22} (eTable 1 in [Supplement 1](#)). Mutations were classified as having high penetrance (relative risk >4), moderate penetrance (relative risk 2-4), or low penetrance (relative risk <2) as well as being recessive or of uncertain clinical actionability based on known disease-associated risks and current modeling.^{8-10,16,21}

Comparison of Conventional Family History–Based and Agnostic Testing for Cancer Predisposition Syndromes

Detailed clinical annotation, including self-reported religion and race, was abstracted for each patient record. Religion and race information was collected to analyze genetic effects of population stratification. Ashkenazi ancestry was determined by self-report of religious preference, via a list of choices, at the time of registration or by specific report of Eastern European Jewish background of relatives at the time of genetic counseling. Race was self-reported by fixed categories at the time of registration. Three-generation family history information was assessed at the time of results communication or from records at the time of genetic testing. Published guidelines and syndrome-specific genetic testing algorithms^{8-10,23-25} were used to determine which genetic tests would be indicated based on tumor histologic features, bilaterality, multiple metachronous cancers, age at onset of cancer, family

history of cancer (including age at which relatives were affected), and self-reported Ashkenazi ancestry. Where indicated, multigene panels (eTable 2 in [Supplement 1](#)) were considered standard of care and were applied using decision rules based on published guidelines (eAppendix 4 in [Supplement 1](#)). A pathogenic variant in secondary analysis was considered incremental if it would not have been detected by testing that would have been ordered based on application of these decision rules, with additional decision rules for cases harboring 2 variants (eAppendix 5 in [Supplement 1](#)).

Adjustment for Founder Mutations and DNA Repair Genes

To adjust for effects of founder mutations, a set of Ashkenazi and European founder mutations in *BRCA1/2*, *APC*, *MSH2*, *MSH6*, *CHEK2*, or *MUTYH* were included in the overall analysis but also identified so as to allow subset analysis (eBox in [Supplement 1](#)).

Mutations

in *ATM*, *BAP1*, *BARD1*, *BRCA1/2*, *BRIP1*, *CHEK2*, *FAM175A*, *MLH1*, *MSH2*, *MSH6*, *NBN*, *PALB2*, *PMS2*, and *RAD51D* recently associated with advanced prostate cancer¹⁶ were coded for separate analysis of patients with advanced prostate cancer, and to assess patients potentially amenable to targeted therapies.

Statistical Analysis

Variant frequencies in cases were compared with allele frequencies in noncancer controls from public databases and stratified by European and Ashkenazi subsets.²⁶⁻³⁰ Allele frequencies were compared by Fisher exact 2-sided binomial test in R version 3.3 software (R Foundation) using RStudio version 0.99.903 (RStudio). To estimate findings resulting from a different case mix, tumor type-specific rates of pathogenic variants were multiplied by cancer rates in the general population³¹ as well as rates of discordant variants observed in a separate ascertainment¹ (eAppendix 6 in [Supplement 1](#)). Where proportions are presented, 95% confidence intervals were derived. Clinical variables were compared with regard to genetically defined subsets, mutational load as measured by number of somatic mutations, and time from diagnosis to tumor-normal analysis for metastatic disease, using 2-sample *t* test for independent samples, with 2-tailed *P* values significant at *P* < .05. Rates of incremental findings were compared between subsets by Fisher exact test, and proportions of germline findings were compared by stage of disease using χ^2 test.

Results

Cohort Characteristics

As part of an institutional review board–approved protocol, from January 1, 2014, until May 31, 2016, 10 336 patients consented to genetic analysis of their tumors. The distribution of cancer diagnoses, sex, age, and stage of disease in the 1040 patients who consented to secondary germline testing from May 1, 2015, through May 31, 2016, is shown in [Table 1](#). Of the 1040 patients, 65.3% were male and 34.7% female. The median age of those who consented to genetic analysis of their tumors was 58 years (interquartile range [IQR], 47-67 years), and the median age of those who consented to secondary germline testing was 58 years (IQR, 50.5-66 years). The median age was similar for all tumor subsets except prostate cancer, for which patients consenting to germline analysis were slightly younger than those who underwent tumor variant calling only (median age, 61 [IQR, 55-67] vs 63 [IQR, 56-68] years, respectively; $P = .04$). Of the 1040 patients, the proportions with stage 0, I, II, III, and IV disease at the time of genomic analysis were 0.3%, 3.3%, 7.9%, 7.2%, and 81.3%, respectively. Self-identification of Jewish ancestry was more common in those consenting to secondary germline analysis than in those undergoing tumor variant calling only (26.9% vs 18.1%, respectively).

Variants Detected

Of the 1040 patients undergoing secondary germline analysis, 205 patients (19.7%) harbored pathogenic variants conferring cancer predisposition. Of the 205 patients with pathogenic variants, 182 carried clinically actionable mutations of high ($n = 97$), moderate ($n = 52$), or low ($n = 33$) penetrance, 8 were carriers of variants associated with recessive syndromes, and 15 carried variants in genes of unproven clinical actionability. [Table 2](#) shows the distribution of 220 variants in these 205 patients, listed by penetrance class and by tumor type. For tumor types with more than 10 cases tested, the incidence of patients with inherited pathogenic variants ranged from 56.3% (9 of 16 patients) for bladder cancers (including urothelial carcinomas) to 25.0% (44 of 176 patients) for pancreatic cancer, 19.6% (71 of 362 patients) for prostate cancer, 16.4% (23 of 140 patients) for renal cancer, and 9.2% (8 of 65 patients) for colon cancer. Each patient had an average of 1.8 variants of uncertain significance in the 76 genes tested, with 833 of 1040 patients (80%) having at least 1 variant of uncertain significance (eTable in [Supplement 2](#)). Of the 1040 patients, 15 had more than 1 variant (eTable in [Supplement 3](#)).

Pathogenic variants of genes involved in DNA repair pathways¹⁶ were observed in 49 of 362 patients (13.5%) with prostate cancer, all of whom had advanced disease, but also in 87 of 678 patients (12.8%) with cancers other than prostate cancer. Of the 220 mutations detected in 205 patients, 83 mutations (37.7%; in 79 patients) were known founder mutations (including Ashkenazi founder mutations, and European founder mutations in *MUTYH* and the *CHEK2* c.1100delC). Cases with founder mutations represented 75.8% of low-penetrance, 40.4% of moderate-penetrance, and 34% of high-penetrance pathogenic alleles (eTable 3 in [Supplement 1](#)).

Comparisons With Phenotype-Directed Approaches

A total of 101 cases had incremental clinically actionable findings that would not have been detected by phenotype-directed testing using current clinical criteria, representing 9.7% (95% CI, 8.1%-11.7%) of the 1040 cases overall and 55.5% (95% CI, 48.2%-62.5%) of the 182 patients with clinically actionable mutations ([Figure](#)). Had the case mix conformed to population cancer incidence rates,³¹ the 55.5% proportion of incremental findings would have been 49.1%. Of the 101 patients with incremental clinically actionable findings, 27, 38, and 36 patients carried high-, moderate-, or low-penetrance mutations, respectively (eTable 4 in [Supplement 1](#)). Thus, of the 101 incremental findings, 65 were moderate- or high-penetrance mutations. There was no difference in the proportion of patients with incremental findings in the 85.2% of probands with family history assessed by genetic counselors or the 14.8% assessed from physician records.

Had all individuals been screened for population-specific founder mutations in addition to guideline-directed testing, the proportion of incremental clinically actionable findings would have declined to 57 of 182 patients (31.3%; 95% CI, 25.0%-38.4%). Had patients with prostate cancer also been screened with a panel of DNA repair genes, the resulting proportion of findings considered incremental would be 35 of 182 patients (19.2%; 95% CI, 14.2%-25.6%) or 3.4% (95% CI, 2.4%-4.7%) of the 1040 cases overall (eTable 4 in [Supplement 1](#)). The highest proportion of actionable findings that would not have been detected based on clinical guidelines were observed in biliary, prostate, colorectal, renal, and pancreatic tumors (eTable 4 in [Supplement 1](#)). [Table 3](#) shows all cases with incremental actionable findings, excluding known founder mutations, which would not have been detected using guideline-directed approaches for high-penetrance mutations (*BRCA2*, *CDKN2A*, *PALB2*, *VHL*, *SDHA*, *MSH2*, *MSH6*, and *BAP1*), moderate-penetrance mutations (*CHEK2*, *ATM*, *MITF*, *BRIP1*, and *RAD51D*), and low-penetrance mutations (*APC*, *MUTYH*). Twenty-two percent of

germline *BRCA1/2* mutations (13 of 59 mutations) and 42.8% of mismatch repair gene mutations (6 of 14 mutations), including founder mutations, were seen in patients who would not have been referred for testing using existing guidelines.

Clinical Implications of Results Transmitted

At a median time to transmission of results to patients after testing of 1 month (range, 0-16 months), as of May 2017, germline results have been communicated to 193 of 205 patients (94.1%) with pathogenic variants detected and 175 of 182 patients (96.2%) with clinically actionable findings. For 29 patients (probands) found to have actionable mutations, genetic testing had been offered to relatives; for 13 of these families, the index patient's genetic findings would not have been discovered through guideline-based genetic evaluation. In half (15 of 29) of the families tested, the mutation has been detected in at least 1 relative; for 20 relatives with mutations, increased surveillance or risk-reducing surgical procedures have been recommended, with oophorectomy documented in a daughter of a proband with advanced prostate cancer. Of the 182 probands with actionable findings, 132 had mutations in DNA repair genes, resulting in discussion or initiation of a change to US Food and Drug Administration–approved or off-label use of targeted therapy in 38 patients. Of these 38 patients, 11 received treatment with poly adenosine diphosphate–ribose polymerase (PARP) inhibitors and/or platinum-based chemotherapy as of May 2017, and such treatments were discussed or in planning in the remaining 27 patients (Table 4 and eTable 5 in Supplement 1). In an additional patient in his mid-60s with advanced prostate cancer with more than 20 somatic mutations and a *PMS2* germline mutation, immunotherapy was planned but not yet administered. Of the subset of 101 cases with actionable findings not predicted by phenotype, 59 involved DNA repair genes. This resulted in discussion or initiation of targeted therapy in 17 patients, with targeted treatment being administered in 6 of these 17 individuals (Table 4 and eTable 5 in Supplement 1).

Comparison of Incremental Findings by Stage of Disease

Among 194 stage 0 to III cases, 16 (8.3%; 95% CI, 5.1%-13.0%) had likely pathogenic germline variants, compared with 189 of 846 patients (22.3%; 95% CI, 19.7%-25.3%) with stage IV disease ($P < .001$). Among 166 clinically actionable pathogenic variants in stage IV cases, 93 (56.0%; 95% CI, 48.4%-63.4%) were incremental, compared with 8 of 16 incremental actionable variants (50.0%; 95% CI, 28.0%-72.0%) among patients with stage 0 to III disease ($P = .64$). This analysis yielded similar results by all tumor types

except colon cancer, in which Lynch mutations were more common in patients with earlier-stage disease (eTable 6 in [Supplement 1](#)). The median number of tumor (somatic) mutations was 4 (IQR, 2-6) in 205 cases with pathogenic variants and 3 (IQR, 2-5) in 835 cases without pathogenic variants ($P = .31$), with a median of 3 mutations observed in both stage 0 through III disease (IQR, 1-5) and stage IV disease (IQR, 2-6) ($P = .07$). The median time from diagnosis to tumor-normal testing for metastatic disease was 2 years for both the 181 patients with pathogenic variants (IQR, 1-6 years) and the 648 patients without such variants (IQR, 1-5 years) ($P = .91$).

Analysis of Cohorts of Ashkenazi and Non-Ashkenazi Background

Of 205 patients with pathogenic variants, 192 self-reported ancestry (68 Ashkenazi, 124 non-Ashkenazi). Had guideline-directed approaches been augmented with testing for population-specific founder mutations, 5 of 68 patients (7.4%; 95% CI, 3.2%-16.1%) of Ashkenazi ancestry with clinically actionable variants would have had incremental findings not predicted by phenotype or ancestry, compared with 46 of 124 patients (37.1%; 95% CI, 29.1%-45.9%) who were not of Ashkenazi ancestry ($P < .001$) (eTable 7 in [Supplement 1](#)).

Comparisons With Public Databases

Among patients of non-Ashkenazi ancestry, a set of *CHEK2* mutations, predicted by family history in only a single patient, was overrepresented in prostate and pancreatic cancers compared with controls from the Exome Aggregation Consortium.^{26,27} *MUTYH* mutations were enriched in non-Ashkenazi patients with prostate cancer and not predicted by family history. Among patients of Ashkenazi ancestry, there was enrichment for *CHEK2* mutations in patients with pancreatic cancer, compared with a public database of 2177 controls of Ashkenazi ancestry³⁰ (eTable 8 in [Supplement 1](#)).

Tumor-Germline Correlations

In the 205 patients with a pathogenic variant, 93 of 170 tumors (54.7%) showed LOH or a pathogenic somatic second mutation in the same gene as the pathogenic variant. Of 180 evaluable pathogenic variants in 170 tumors, 82 variants showed LOH in the tumors and 13 tumors demonstrated a loss-of-function mutation or a previously reported deleterious missense variant.^{32,33} Concurrent somatic LOH or a second mutation at the same locus

accompanied the germline mutation in 37 of 48 evaluable tumors (77.1%) in patients with *BRCA1/2* germline mutations, 9 of 12 patients (75.0%) with mismatch repair gene variants, and 13 of 36 patients (36.1%) with germline *CHEK2* variants, including 5 of 10 patients (50.0%) with the founder *CHEK2* c.1100delC variant. Of the 103 pathogenic *BRCA* mutations identified in the tumors of the 1040 patients in this study, only 59 were germline in origin, whereas 44 were detected in the tumor.

A hypermutated tumor profile (defined as >20 somatic mutations) or MSI-H (defined as >10% of loci by MSIsensor²⁰) was observed in 51 of 1040 patients (4.9%). Germline pathogenic variants in mismatch repair genes were identified in 12 of these 51 patients (23.5%). Two additional patients with germline *MSH6* truncating mutations had tumors that were not hypermutated or MSI-H by MSIsensor (1 with renal cell cancer and 1 with prostate cancer). Among the hypermutated or MSI-H cases, 14 of 51 (27.5%) fulfilled clinical criteria for Lynch syndrome (revised Bethesda guidelines and/or Amsterdam II criteria); 6 of 12 patients (50.0%) with an MSI-H tumor and an identified germline pathogenic variant in a mismatch repair gene met these criteria. Of the 14 cases with germline mismatch repair mutations, 6 were incrementally detected by germline analysis in the absence of a family history diagnostic of Lynch syndrome, including 3 patients with bladder cancer and 3 with prostate cancer. Of these 6 cases, 3 demonstrated LOH or a second somatic mutation (Table 3).

Discussion

This study identified clinically actionable inherited mutations in 17.5% of patients with advanced cancer, compared with 3% to 12.6% in prior series.¹⁻⁷ Of the 1040 patients, 101 (9.7%) would not have been detected using clinical guidelines, which represented 55.5% of the total of 182 patients with actionable findings in the series. Germline findings led to discussion or initiation of change to targeted therapy in 38 (3.7%) of the 1040 patients tested and predictive testing in the families of 13 individuals (1.3%), including 6 for whom genetic evaluation would not have been initiated by guideline-based testing.

The prevalence of germline variants in a clinical setting will be affected by stage, case mix, ethnic ancestry of the population, and methods of variant classification. There was a significantly greater overall prevalence of germline mutations observed in patients with metastatic disease, although the proportion of germline findings that were incremental to predictions based on

family history (approximately 50%) was similar in metastatic and nonmetastatic disease. This interesting association may reflect more aggressive biological features (eg, via greater mutational load of deleterious variants) or improved chance of survival with metastases in those with germline mutations. However, in the cohort studied here, there was no association of inherited mutations with increased tumor mutational load or difference in time from diagnosis to metastatic disease; further molecular profiling and prospective studies of treatment response may provide an explanation for the increased prevalence of germline mutations in metastatic disease. In this series, there was an abundance of late-stage prostate, pancreatic, renal, and breast cancers in which germline variants were more frequent, probably accounting for the higher observed prevalence of pathogenic variants found here compared with prior studies. While these findings reflect the experience at a referral cancer center, had the case mix more closely resembled population cancer incidence rates, the proportion of incremental findings would have been 49.1%. Had the entire cohort been screened for ancestry-specific (Ashkenazi and northern European) mutations, the proportion of incremental actionable findings would have been 31.3%, approximating the rate in a hypothetical cohort with no population diversity. However, if family history assessment by clinicians is less complete than the 3-generation information used in this analysis, the proportion of apparent incremental findings may be higher in practice. Thus, guideline-based testing will fail to detect a third to half (31.3%-55.5%) of genetic findings found by tumor-normal sequencing, taking into account case mix, disease stage, and ethnic ancestry.

The complementary role of tumor and germline sequencing was exemplified by colon and breast cancer testing. Tumor-normal testing was able to diagnose Lynch syndrome in patients who would not otherwise have been tested. The sensitivity of tumor-derived hypermutation or MSI-H status for detection of Lynch syndrome was 85.7%, with germline sequencing resulting in the diagnosis of Lynch syndrome in an additional 2 patients, 1 of whom would not have been diagnosed by family history. Half of the 12 cases with pathogenic variants in mismatch repair genes in the setting of a hypermutated or MSI-H tumor would not have met guideline-based criteria for tumor-directed immunohistochemical analysis. For breast cancer cases, had tumor-only testing been performed, approximately 40% of patients with *BRCA1/2* variants detected with tumor sequencing would have been referred for a germline confirmation test that would have been negative; conversely “subtraction” of germline from tumor DNA sequence would have obscured 59

germline *BRCA1/2* cases. These observations support the rationale for combined tumor-normal sequencing.^{3,34}

Although epigenetic mechanisms of loss of the second allele may also exist, concurrent somatic alterations in the same gene or LOH in matched tumors support the pathogenic role of many of the germline variants observed in nonsyndromic settings. For example, germline *CHEK2* mutations were observed in 6 patients with renal cancer; the second allele was mutated in 3 of 5 evaluable tumors. Among novel nonsyndromic associations seen here, mutations in *CHEK2* were enriched in prostate and pancreas cancers, *MUTYH* heterozygous mutations were enriched in prostate cancer, and mutations in *MSH6*, *BARD1*, *PALB2*, *MITF*, and *SDHB*, were observed absent the typical family history associated with these genes. Other unanticipated findings reported here will require further functional and genetic epidemiologic genomic exploration, including the observation of the recurrent mutation *FH* c.1431_1433dupAAA (p.Lys477dup) and *RECQL4* loss-of-function variants.³⁵⁻³⁷

To our knowledge, this study marks the first large-scale effort to return germline findings in the context of tumor-normal sequencing to patients. Such an approach in patients with advanced cancer was found to uncover potentially actionable germline variants that would not be detected using existing guidelines for genetic risk assessment. Less resource-intensive strategies than tumor-normal sequencing could be applied but would result in lower sensitivity. For example, testing all patients for a dozen DNA repair genes and several *APC* and *MUTYH* founder mutations, combined with standard phenotypic assessment, would have detected 92.3% of patients with clinically actionable germline variants.

This study has several limitations. Among these is the lack of sufficient follow-up to assess the effect of the genetic information on patient or family outcomes, including potential harms due to false-positive results of screening. In addition, there was physician discretion for referral for tumor sequencing, potentially favoring enrollment of those who may have been eligible for targeted therapies. The usual care comparator in this study was synthetic and not the actual yield of testing a population according to guidelines. Also, interpretation of detailed family history information was by expert reviewers using reproducible but complex algorithms. In addition, the study had unique demographic characteristics and case mix. These factors will limit generalizability of findings to a community practice environment, where germline testing panels are also being introduced.³⁸

Conclusions

In this referral population with selected advanced cancers, universal sequencing of a broad panel of cancer-related genes in paired germline and tumor DNA samples was associated with increased detection of individuals with potentially clinically significant heritable mutations over the predicted yield of targeted germline testing based on current clinical guidelines. Knowledge of these additional mutations can help guide therapeutic and preventive interventions, but whether all of these interventions would improve outcomes for patients with cancer or their family members requires further study.

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