Japanese version of The Cancer Genome Atlas, JCGA, established using fresh frozen tumors obtained from 5143 cancer patients

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Abbreviations: CCP, comprehensive cancer panel; CDG, cancer driver gene; CNV, copy number variation; GEP, gene expression profiling; TMB, tumor mutation burden; WES, whole-exome sequencing.
Abstract
This study aimed to establish the Japanese Cancer Genome Atlas (JCGA) using data from fresh frozen tumor tissues obtained from 5143 Japanese cancer patients, including those with colorectal cancer (31.6%), lung cancer (16.5%), gastric cancer (10.8%) and other cancers (41.1%). The results are part of a single-center study called “High-tech Omics-based Patient Evaluation” or “Project HOPE” conducted at the Shizuoka Cancer Center, Japan. All DNA samples and most RNA samples were analyzed using whole-exome sequencing, cancer gene panel sequencing, fusion gene panel sequencing and microarray gene expression profiling, and the results were annotated using an analysis pipeline termed “Shizuoka Multi-omics Analysis Protocol” developed in-house. Somatic driver alterations were identified in 72.2% of samples in 362 genes (average, 2.3 driver events per sample). Actionable information on drugs that is applicable in the current clinical setting was associated with 11.3% of samples. When including those drugs that are used for investigative purposes, actionable information was assigned to 55.0% of samples. Germline analysis revealed pathogenic mutations in hereditary cancer genes in 9.2% of samples, among which 12.2% were confirmed as pathogenic mutations by confirmatory test. Pathogenic mutations associated with non-cancerous hereditary diseases were detected in 0.4% of samples. Tumor mutation burden (TMB) analysis revealed 5.4% of samples as having the hypermutator phenotype (TMB ≥ 20). Clonal hematopoiesis was observed in 8.4% of samples. Thus, the JCGA dataset and the analytical procedures constitute a fundamental resource for genomic medicine for Japanese cancer patients.

KEYWORDS
actionable alteration, cancer genome, driver alteration, individualized medicine, multi-omics analysis platform

1 | INTRODUCTION
PanCancer genome analysis has provided important information, including on tumorigenic mechanisms, actionable genetic alterations to select molecular-targeted drugs, tumor mutation burden (TMB) to select immune checkpoint inhibitors (ICIs), hereditary cancer syndromes and clonal hematopoiesis.1-7 Racial and ethnic population-specific cancer driver alterations and single nucleotide polymorphisms (SNPs) may complicate the evaluation and interpretation of the results of genetic analyses. Several studies have reported obvious racial differences in cancer-specific driver alterations among Japanese individuals.8-10 Furthermore, 15% of SNPs are reportedly specific to Japanese individuals,11 thus affecting the evaluation of the clinical significance of genetic alterations. Therefore, for accurate annotation of these genetic aberrations, race-specific analysis pipelines are required.

In the present study, we evaluated 5521 fresh frozen tumor tissues obtained from 5143 Japanese cancer patients through whole-exome sequencing (WES), cancer gene panel sequencing, fusion gene panel sequencing and microarray-based gene expression profiling (GEP). Furthermore, we developed an in-house analytical pipeline termed “Shizuoka Multi-omics Analysis Protocol,” which includes SNPs specific to Japanese individuals and data on cancer-specific gene alterations obtained from Japanese cancer patients. The present results are a part of the single-center study called “High-tech Omics-based Patient Evaluation” or “Project HOPE”12 conducted at the Shizuoka Cancer Center from 2014, which aims to establish the Japanese version of The Cancer Genome Atlas (JCGA).

2 | MATERIALS AND METHODS
2.1 | Ethical statement
All experimental protocols were approved by the institutional review board at the Shizuoka Cancer Center (Authorization Number:
25-33). Written informed consent was obtained from all patients for their participation in the study. All experiments using clinical samples were carried out in accordance with the approved guidelines.13

2.2 Clinical samples

To identify somatic and germline genomic alterations in patients, tumor and matched normal samples were subjected to WES, comprehensive cancer panel (CCP) sequencing, fusion gene panel sequencing and GEP. These samples were obtained from patients receiving surgical treatment at the Shizuoka Cancer Center Hospital. Fresh surgical specimens were assessed by a pathologist, and samples with tumors weighing >100 mg were analyzed. Samples were not further filtered in accordance with pathophysiological features or cancer type. Blood samples from the same patients were used as the control for WES and CCP, and tumor-adjacent normal tissue specimens were used as the control for GEP. WES, CCP and fusion gene panel sequencing were performed using an Ion Proton System (Thermo Fisher Scientific) and transcriptome profiling was performed using an Agilent system (Agilent Technologies). Detailed experimental protocols have been previously described.14-17 Sequence data analysis procedures are described in Doc. S1. The mean depth of coverage in WES and CCP was 130 and 1169, respectively.

2.3 Evaluation of driver and actionable genomic alterations

Genomic aberrations were evaluated taking into consideration driver and actionable alterations; the former refers to genomic changes contributing to tumorigenesis and the latter to those attacked by molecular-targeted drugs. These aberrations were evaluated using an analysis pipeline called “Shizuoka Multi-omics Analysis Protocol (SMAP)” developed in-house (Figures 1 and 2). Three types of genomic changes were analyzed: (a) SNV and indel via WES and CCP; (b) fusion genes via a fusion gene panel; and (c) oncogene amplification and tumor suppressor gene (TSG) deletion via integrative analysis of GEP and copy number variations (CNVs). Drivability of somatic and germline mutations was classified into five tiers in accordance with the reliability of supporting information by sequentially comparing alterations among multiple databases (Figure S1).18-27 Variants classified under tier 1 were considered for drivability assessment, as described below. When classifying genomic alterations based on drivability, exact matches with database entries were required. A combination of (a) chromosome number, genomic coordinates and base substitution patterns or (b) gene symbols, amino acid positions and amino acid substitution patterns were used for database matching.

Actionable mutations were classified into five evidence levels in accordance with those proposed by the Center for Cancer Genomics and Advanced Therapeutics.28 Confidence levels of drugs (approved by a regulatory agency, clinical trial, case report or pre-clinical analysis) and the matching status of cancer types between the database and the query were used in the classification (Figure S2). Such alterations, classified as level A, were defined as druggable alterations.

2.4 Construction of a catalogue of cancer-related genes and pathway assignment

To focus on cancer driver genes, data on 914 genes including (a) oncogenes and TSGs and (b) genes harboring somatic pathogenic mutations were compiled. The former were obtained from COSMIC Cancer Gene Census,29 OncoKB Cancer Gene List,30 and the literature30-32 and the latter were obtained from integrating genes with somatic pathogenic mutations reported in CGI,18 ClinVar,26 DoCM,22 and OncoKB25 and non–functional mutations in IARC-TP53.24 Furthermore, in our analysis pipeline SMAP, 1074 cancer-related genes were compiled from 27 resources including cancer gene panels. To evaluate biological processes affected by driver and actionable alterations, genes were annotated with pathway classification in accordance with the KEGG pathway,33 UniProt34 and the literature2,31,32 followed by manual curation. Twenty-seven pathways classified into 11 categories were assigned to 1462 genes (Table S1). The list of 1988 genes including oncogene/TSG classification and pathway information is provided in Table S2.

3 RESULTS

3.1 The HOPE cohort

The HOPE cohort comprised 5521 tumor specimens (5020 primary tumors and 501 metastatic tumors) derived from 5143 patients treated at the Shizuoka Cancer Center Hospital from January 2014 to March 2019. The types of cancers are summarized in Figure 3. The most prominent cancer type was colon cancer (1014 samples, 18.4%), followed by lung (905, 16.5%) (including 658 lung adenocarcinoma and 178 lung squamous cell carcinoma), rectal (733, 13.3%), stomach (599, 10.8%), head and neck (344, 6.2%), breast (288, 5.2%), and liver (242, 4.4%) cancer, accounting for 74.7% of total samples. In contrast, 9.5% of samples (523 out of 5521) were derived from 50 rare cancers,35 the most prominent being gastrointestinal stromal tumor (GIST) (86 samples), followed by brain tumor (79 samples), head and neck cancer (62 samples), and sarcoma (58 samples). Our dataset included 39 pathologically proven benign tumor samples, which were excluded in subsequent analyses.
3.2 | Germline mutations for hereditary cancer syndromes

The prevalence of germline mutations causing hereditary cancer syndromes was evaluated from 3022 blood samples for which confirmatory results were available. Pathogenic mutations in 49 hereditary cancer genes with a minor allele frequency of <1% were extracted. Germline driver mutations were detected in 9.2% of cases (279 out of 3022) in 25 genes (Table 1A). The top 5 genes, MSH2, BRCA1, CDH1, SDHD, and APC, accounted for 57.7% of the total. Among them, 12.2% (1.1% of total) were confirmed as pathogenic mutations using a confirmatory test and 6.1% were diagnosed as hereditary cancer syndromes. Driver mutations in these cases were detected in 14 genes, with BRCA1 being the most prominent gene in 9 cases, followed by MLH1 (5 cases), BRCA2 (4 cases), CHEK2 (3 cases), PTEN (2 cases), and SDHB (2 cases). Among 25 genes with driver mutations, 22 genes were matched to those reported by The Cancer Genome Atlas (TCGA) group. These genes covered 97.5% (285 out of 293) and 39.5% (335 out of 853) of mutations in HOPE and TCGA cohorts, respectively. BRCA1, CHEK2, BRCA2, TP53, NF1, and MSH6 were commonly identified in the top 20 genes. Distinct distribution of cancer type and analysis pipeline used in both cohort contribute dataset specific mutations. No and one kinship were observed in 34 and 279 cases, respectively, by means of family history and germline SNP analysis. In the latter, APC p.Gln2322Arg were shared by one colon and one rectal tumor sample, indicating that they belong to the same family. In the remaining 277 cases, no kinship was detected.
3.3 | Germline mutations for non–cancerous hereditary diseases

Herein, we focused on 12 non–cancerous hereditary diseases resulting from mutations in 34 genes that are recommended to be disclosed to patients according to the guidelines of the American College of Medical Genetics and Genomics (ACMG). Accordingly, 3022 blood samples were analyzed and 11 pathogenic mutations for non–neoplastic genetic diseases in 11 cases (0.4% out of 3022) were confirmed (Table 1B), including 3 cases of familial hypercholesterolemia (LDLR mutation), 3 cases of hypertrophic cardiomyopathy (MYH7 mutation), 1 case of Marfan syndrome (FBN1 mutation), 1 case of Fabry’s disease (GLA mutation), 1 case of long QT syndrome (KCNH2 mutation), 1 case of Familial hypertrophic cardiomyopathy (MYL2 mutation), and 1 case of cardiomyopathy (TNNT2 mutation).

3.4 | Prevalence of clonal hematopoiesis

Clonal hematopoiesis results from a mutation in hematopoietic stem cells, wherein the mutant clone undergoes de novo mutagenesis and potentially progresses to myelodysplastic syndrome and leukemia. Because clonal hematopoiesis occurred in blood cells used as matched controls in this study, this may affect the evaluation of somatic mutations. Therefore, 3751 blood samples derived from patients with primary tumors without cancer anamnesis were subjected to analysis. Germline SNVs identified in 880 known oncogenes and TSGs were analyzed using a previously reported method. Consequently, clonal hematopoiesis was identified in 8.4% of cases (Table 2). Upon stringent comparison, clonal hematopoiesis in 41 cases (1.1%) was identical to that reported in other studies. Mutated genes and their frequencies in these 41 cases included DNMT3A (18 cases), SRSF2 (5 cases), TET2 (5 cases), ASXL1 (4 cases) IDH2 (3 cases), SF3B1 (3 cases), GNAS (2 cases), JAK2 (2 cases), NRAS (1 case), TP53 (1 case), and U2AF1 (1 case). Consistent with previous reports, clonal hematopoiesis was more frequently observed among the elderly.

3.5 | Tumor mutation burden and mutation signatures

The TMB, also referred to as mutation load, representing the number of somatic mutations per megabase, has received increasing attention owing to its potential to estimate the efficacy of responses to ICIs. The TMB was determined for 5395 samples for which WES results were available. PanCancer analysis revealed bimodal distribution, as shown in Figure 4A. When the cutoff value was set to 20, which corresponded to 663 mutations on WES, 5.4% of samples (292 out of 5395)
were classified as having the hypermutator phenotype (Figure 4A, B). Further classification revealed distinct TMB distributions among cancer types (Figure 4C). Bimodal distribution was observed in several cancers, including brain, stomach, colon, and uterine cancer and melanoma, and the cutoff varied among these cancers. However, others revealed no clear signal in the TMB distribution to classify hypermutator phenotypes.

Further characterization of hypermutator phenotypes was performed by analyzing mutation signatures. Distinct signature profiles were observed in different cancer types (Figure 5). For instance, tobacco smoking-associated signatures in lung adenocarcinoma and mismatch repair deficiency-associated hypermutation in colon and uterine cancer were identified as expected (Figure S3). More detailed investigation of signatures in these cases revealed distinct predisposing factors causing this phenotype among cancer types. Such samples were characterized by an extremely high TMB and POLE signature (Figure 5). Samples under the POLE category were limited to colorectal and endometrial cancer and numerous mutations had accumulated owing to damaging mutations in DNA repair enzyme polymerase epsilon and these phenotypes were considered different from those in Lynch syndrome.39

### 3.6 Characteristics of driver somatic alterations

To evaluate genomic alterations contributing to tumorigenesis, multi-omics profiling results were analyzed using SMAP (Figure 1). Among 5521 samples, 4131 were available for all of the following datasets and used in the following analyses: (a) mutation dataset: WES for tumor and blood samples and CCP for tumor samples; (b) fusion dataset: fusion gene panel sequencing for tumor samples; and (c) expression dataset: GEP for tumor and normal samples. Consequently, 6817 driver events were detected in 2982 samples (72.2% out of 4131) (Figure 6). On average, 2.3 driver events were detected. SNV and indel yielded the highest sample coverage (62.2%), followed by expression aberrations (21.0%) and gene fusions (12.9%). Two or three types of alterations were observed in 13.0% of samples. The frequency of samples with driver alterations varied among cancer types, with the highest detection rate observed in rectal cancer (95.3%), followed by colon cancer (92.2%), uterine cancer (86.9%), and GIST (84.3%), and the lowest rate observed in kidney cancer (11.7%) (Figure 6). Furthermore, the relative frequency of the three alteration types differed

| Gene | Number of cases | Cancer type |
|------|-----------------|-------------|
| BRCA1 | 9 | Breast (3), rectum (2), GIST (1), lung (1), ovary (1), pleura (1) |
| MLH1 | 5 | Colon (4), stomach (1) |
| BRCA2 | 4 | Breast (2), colon (1), head and neck (1) |
| CHEK2 | 3 | Breast (1), lung (1), rectum (1) |
| PTEN | 2 | Breast (1), pancreas (1) |
| SDHB | 2 | Breast (1), head and neck (1) |
| MSH6 | 2 | Colon (1), uterus (1) |
| MSH2 | 1 | Uterus (1) |
| EXT2 | 1 | Colon (1) |
| CDH1 | 1 | Breast (1) |
| NF1 | 1 | Stomach (1) |
| NTRK1 | 1 | Stomach (1) |
| TP53 | 1 | Sarcoma (1) |
| VHL | 1 | Rectum (1) |

### Table 1 Summary of germline mutations in the High-tech Omics-based Patient Evaluation (HOPE) cohort

| Number of cases | Prevalence |
|-----------------|------------|
| Examined 3022   |            |
| Detected 279    | 9.2%       |
| Confirmed 34     | 1.1%       |

| Gene       | Number of cases | Disease                      |
|------------|-----------------|------------------------------|
| LDLR       | 3               | Familial hypercholesterolemia |
| MYH7       | 3               | Hypertrophic cardiomyopathy  |
| FBNI       | 1               | Marfan syndrome              |
| GLA        | 1               | Fabry’s disease              |
| KCNH2      | 1               | Long QT syndrome             |
| MYL2       | 1               | Familial hypertrophic cardiomyopathy |
| TNNT2      | 1               | Cardiomyopathy               |

| Gene with reported clonal hematopoiesis | Number of cases | Prevalence |
|-----------------------------------------|-----------------|------------|
| DNMT3A (18), SRSF2 (5), TET2 (5), ASXL1 (4), IDH2 (3), SF3B1 (3), GNAS (2), JAK2 (2), NRAS (1), TP53 (1), U2AF1 (1) | 41 | 1.1% |

| Gene with reported clonal hematopoiesis | Number of cases | Prevalence |
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Cancer type-specific driver genes such as APC in colorectal cancer, BRAF, EGFR, KIT, and KRAS in melanoma, lung adenocarcinoma, GIST, and multiple cancer types, respectively, were confirmed (Figure S5A). These driver events were detected in 362 genes, enriched in pathways involved in cell growth (MAPK, PI3K and RTK pathways), differentiation (WNT pathway), and genome maintenance (TP53 pathway) (Figure 7).

Racial differences in driver genes and mutations were investigated by comparing the observed frequencies in HOPE and TCGA cohorts in 18 cancer types for which ≥20 samples were analyzed using both datasets. Among multi-ethnic patients’ data included in the TCGA dataset, those samples that were derived from Caucasians were used in this analysis. The prevalence of driver events in 8 genes differed between these datasets (Figure S6). KRAS in lung adenocarcinoma and pancreatic cancer and PTEN in gastric cancer were enriched in Caucasian individuals, while EGFR in lung adenocarcinoma, AKT1 in breast cancer, and CTNNB1, KRAS, and PIK3CA in ovarian cancer were enriched in Japanese individuals. Mutation level analysis indicated differences in hotspot mutations. PTEN p.K267fs in gastric cancer and KRAS p.G12C in lung adenocarcinoma were more frequent in Caucasian and EGFR p.E746_A750del and p.L858R in lung adenocarcinoma and KRAS p.G12D and p.G12V and PIK3CA p.E542K were enriched in Japanese individuals. To examine the effect of a low tumor content sample on driver mutation detectability, samples with estimated tumor content ≥0.25 in the HOPE dataset were compared with TCGA. As a result, no differences were observed for AKT1 in breast cancer and KRAS in pancreatic cancer. For ovarian cancer, enrichment of these genes in the HOPE dataset reflected a distinct subtype composition (Table S3). Driver mutations in CTNNB1 were observed in endometrioid cancer and KRAS and PIK3CA were identified in multiple subtypes. In contrast, driver mutations in these genes were observed less frequently in serous ovarian cancer, which constitutes 100% and 18% (15 out of 83 samples) of ovarian cancers in TCGA and HOPE, respectively.

(Figure S4).
3.7 | Actionable genomic alterations associated with molecular-targeted drugs

The association between somatic alterations and molecular-targeted drugs was classified into five evidence levels (levels A-E). Figure 8 shows the proportion of 4131 tumor specimens associated with molecular-targeted drugs. Druggable alterations (level A) were identified among 11.3% of samples (467 out of 4131, red bar in Figure 8) associated with drugs potentially applicable in the clinical setting. Among them, 107 samples (2.6% of total) were associated with ICIs. Druggable alterations were enriched in 6 genes, including fusion genes and MSI-high samples. KIT was most enriched (81.4% of GIST, 13.6% of melanoma), followed by EGFR (40.8% of lung...
adenocarcinoma, 3.4% of lung cancer other than adenocarcinoma and squamous cell carcinoma), BRAF (31.8% of melanoma), MSI-high samples (9.5% of uterus, 7.5% of colon and 5.6% of stomach), fusion genes (8.2% of sarcoma), PDGFRA (2.9% of GIST), ERBB2 (2.7% of breast, 2.4% of stomach), and MET (0.7% of lung adenocarcinoma) (Figure S5B).
In the remaining samples, 18.7% were annotated as second (level B, blue bar in Figure 8) and had possibility for clinical trial enrollment. Moreover, 14.0% of samples were classified as third (level C, green bar in Figure 8), serving as candidates for application expansion of molecular-targeted drugs. A total of 7.9% of samples were assigned to drugs with level D for which the efficacy was determined through a small-scale study or through case reports. Finally, 3.2% of samples were classified as the lowest (level E) for which in vivo/in vitro experimental results were reported. Overall, data on actionable alterations were obtained from 55.0% of samples (2274 out of 4131) when including drugs with evidence levels A-E. Cancer types with the highest sample coverage were GIST (84.3%), followed by melanoma (77.3%) and uterine (75.0%), colon (71.6%), lung adenocarcinoma (71.4%), and rectal (69.1%) cancers (Figure 8). Resistant mutations were identified in 18.3% of samples (756 out of 4131), with the highest rate observed in 45.3% of rectal cancers, followed by 41.9% of colon cancer, 13.5% of lung cancer, and 1.4% of GIST.

**FIGURE 9** Summary of genomic alterations identified in the High-tech Omics-based Patient Evaluation (HOPE) cohort

4 | DISCUSSION

Since January 2014, the Shizuoka Cancer Center has conducted a single-center study called "High-tech Omics-based Patient Evaluation" or "Project HOPE" to establish a Japanese version of The Cancer Genome Atlas (JCGA). We analyzed 5521 fresh frozen tumor tissues obtained from 5143 Japanese cancer patients, using WES, CCP sequencing for 409 cancer-related genes, fusion gene panel sequencing for 491 known fusion genes and microarray-based GEP. Because the patients included herein received surgical treatment to eliminate tumors and yielded fresh and enough quantity of cancer tissues, various types of cancers including rare cancers were evaluated. Furthermore, we developed a novel analytical methodology SMAP to identify and evaluate driver and actionable genomic aberrations in the genome of Japanese cancer patients. To identify somatic driver alterations, genetic alterations in 914 cancer driver genes (CDGs) were evaluated. This list comprised two subsets: (a) 880 oncogenes and TSGs compiled from COSMIC Cancer Gene Census, OncoKB, and the literature; and (b) 207 genes for which somatic mutations have been registered as pathogenic in mutation databases. Four types of genetic alterations in CDGs were defined as somatic driver alterations: (a) pathogenic somatic mutations in CGI, DoCM, OncoKB, IARC-TP53, and ClinVar; (b) oncogene amplification; (c) TSG deletion; and (d) fusion genes (Figure 1). Integrative microarray and CNV analyses were incorporated into SMAP to identify gene amplifications and deletions. The results of the fusion gene panel were also integrated. For germline analysis, 49 hereditary cancer-related genes were used. Pathogenic mutations according to ClinVar and HGMD were considered germline driver alterations. To make the results more understandable for medical staff, pathway classification of driver genes was integrated into SMAP; 92.7% of driver genes (847 out of 914) were classified into 26 pathways (Table S1).

Regarding SNPs in Japanese individuals, 15% of variants were specific to Japanese individuals among 26 populations, thus deterring the analysis of the clinical significance of genetic alterations. Therefore, for more reliable annotation, iJGVD40 and HGVD41 were integrated into SMAP. The usefulness of Japanese SNPs in the analysis of the Japanese cancer genome is attributed to the following features of the analytical methodology: (a) for germline analysis, SNPs were filtered out as non–pathogenic; (b) for somatic analysis, SNPs in variant of unknown significance were considered to have a low functional impact; and (c) for tumor only somatic analysis, SNPs were used as matched controls and non–SNPs were considered somatic mutations. Furthermore, the shared ratio of SNPs at a 1% allele frequency among different populations was 0.5-0.9.42 Therefore, analyses without data on matched racial SNPs yielded more false-positive findings (SNPs specific to Japanese individuals were identified as driver mutations) and false-negative findings (non-Japanese SNPs were eliminated from the final list).
Our major findings are summarized in Figure 9. This study reveals a Japanese version of The Cancer Genome Atlas and is applicable for cancer therapy in Japan. Somatic driver alterations were identified in 362 genes (out of 914 CDGs) in 2982 samples (72.2% of 4131) (Figure 6). On average, 2.3 driver events were detected (minimum = 1 event in 1120 samples, maximum = 15 events in 1 sample) and the number of driver events varied among cancer types. The co-occurrence of druggable and driver alterations among samples also differed among various cancers (Figure S7). For example: (a) both druggable and driver alterations were identified in most samples (ex. GIST); (b) driver alterations were identified in >80% of samples (however, molecular-targeted drug assignments were rare: eg, colon, rectum and uterus); (c) the driver detection rate exceeded 70% and half were associated with molecular-targeted drugs (eg, lung adenocarcinoma and melanoma); and (d) others.

In terms of TMB, 5.4% of samples were classified as hypermutator with a cutoff value ≥20 on PanCancer analysis (Figure 4A,B). Mutational signatures identified potential predisposing factors in samples with the most prominent hypermutator phenotype along with defective DNA mismatch repair (44.4%), followed by tobacco smoking (10.9%), POLE abnormality (8.9%), exposure to ultraviolet irradiation (2.1%), and unknown (27.6%) (Figure 5). Samples with the hypermutator phenotype with mismatch repair deficiency signatures could be a target for ICI. Thus, TMB along with signatures could be used to determine the effectiveness of ICI. A higher TMB indicates that driver genes should be carefully interpreted because causal driver alterations may be covered by a vast amount of passenger mutations in such case.

Genome-based selection of molecular-targeted drugs and ICI is one of the most significant findings in cancer genomic medicine. To simulate the amount of samples associated with these drugs using the current analytical methodology, somatic actionable alterations were evaluated. Consequently, 8.7% and 2.6% of samples were estimated as candidates for molecular-targeted drugs and ICI treatment, respectively, thus accounting for 11.3% of all samples. These drugs are used in the current clinical setting. High-frequency actionable alterations were concentrated in EGFR in lung adenocarcinoma, BRAF in melanoma, and KIT in GIST (Figure 5B). Nonetheless, there were substantial differences in actionable alterations among genes and cancer types, which reflected the availability of drugs for each cancer type in the current situation.

Through germline analysis, driver alterations for hereditary cancer syndromes and non–cancerous hereditary diseases in the ACMG list were identified in 1.1% of samples; in these cases, 0.4% were diagnosed as non–neoplastic and improvable genetic diseases (Table 1B) in accordance with the ACMG guidelines. Regarding clonal hematopoiesis, an analysis of 3751 blood samples revealed that clonal hematopoiesis was detected in 8.4% of samples. Clonal hematopoiesis is expected to be useful clinically for risk assessment upon progression to myelodysplastic syndrome and leukemia. In terms of cancer genome medicine for solid tumors, clonal hematopoiesis raises several issues related to the analysis of cancer driver mutations. Among these, the interpretation of the results of liquid biopsy may be affected; hence, careful examination is required to determine whether these genetic alterations occurred in circulating tumor cells, cell free DNA or during clonal hematopoiesis.

The present study shows that the general features of The Cancer Genome Atlas do not significantly differ between Japanese and Caucasian populations, except for racial SNPs. However, in some certain cancer types, clear racial differences in the incidence of driver genes were observed upon comparative analysis of HOPE and TCGA datasets (Figure S6): for example, a high frequency of EGFR mutations and a low frequency of KRAS mutations in lung adenocarcinoma in Japanese patients. Mutation analysis confirmed these results, as revealed through enrichment of EGFR p.E746_A750del and p.L858R in Japanese and KRAS p.G12C in Caucasian populations. Further analysis on other types of cancers is now evaluated and results will be presented in elsewhere. Furthermore, this study delineates the landscape of driver alterations in rare cancers in Caucasian individuals, including KIT in GIST, HRAS in skin cancer and KRAS in duodenal cancer.

The present study provides novel insights into the future of cancer genome research. Cancers without driver genetic alterations constitute an important issue. In our cohort, driver alterations were not detected in 27.8% of cases. Intensive analysis is required to clarify causal genomic alterations including other types of changes such as epigenetic and structural variations. Further studies are required to assess driver alterations for hereditary cancer syndromes. Herein, pathogenic mutations in hereditary cancer genes were identified in 9.2% of samples, although only 6.1% were diagnosed as hereditary cancer syndromes. The remaining cases may include those with variant of unknown significance, those with difficulties regarding family surveys, and probands of the hereditary cancer syndromes. Further studies on our cohort may elucidate the association between pathogenic genetic alterations and the development of hereditary cancer syndromes, because our cohort is established from a single institution and all clinical data could be matched to genetic alterations.

In conclusion, our dataset, JCIG, established from project HOPE, potentially serves as a fundamental resource for genomic medicine for Japanese cancer patients. Integrated analysis of patient outcomes and pharmacotherapeutic history would accelerate individualized treatment and drug development in the future in accordance with the HOPE dataset.
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SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section.

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