Tumor Mutational Burden Associated With Response to Hyperthermic Intraperitoneal Chemotherapy

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Background: Gastric cancer (GC) is one of the most common cancer types, especially in Asian countries. Hyperthermic intraperitoneal chemotherapy (HIPEC) has been shown to improve the progression-free survival among gastric cancer patients with peritoneal metastases; however, not all patients demonstrate response to HIPEC.

Methods: Biomarkers are needed to select patients for effective treatment of HIPEC. Here, we performed whole-exome sequencing on tumor samples from 18 gastric cancer patients who received HIPEC treatment and assessed the association between genomic mutation features and progression-free survival. Exome sequencing was further conducted on tumor samples from additional 15 gastric cancer patients as a replication study.

Results: The tumor mutational burden (TMB) was significantly higher in the group of patients with a better response to HIPEC treatment than that of the others. Kaplan–Meier survival curve showed that patients with high TMB had a significantly longer survival time than that in patients with low TMB. This discovery was validated in the replication cohort. Genes bearing mutations recurrently and selectively in patients with better response to HIPEC were found in the two cohorts.

Conclusion: We found that higher TMB is significantly associated with better response to HIPEC. Our results provide useful hints for prognostic stratification of HIPEC treatment.

Keywords: gastric cancer, hyperthermic intraperitoneal chemotherapy, survival, tumor mutational burden, biomarker

INTRODUCTION

Gastric cancer is among the most common cancer types and a leading cause of cancer death in both men and women worldwide (1), especially in Asian countries, which imposes a considerable global health burden. In addition, when gastric cancer is diagnosed, patients are often already at an advanced stage. Surgery with subtotal gastrectomy or total gastrectomy is the current mainstay of...
treatment (1). Surgical resection is a curative therapeutic approach for gastric cancer and confers good outcome for early-stage gastric cancer. However, early gastric cancer often remains asymptomatic unless detected by endoscopy and biopsy. The survival rate for patients with metastatic gastric cancer is very low, ranging from 4 to 12 months (1), despite the successful application of modern chemotherapy to other solid tumors.

Hyperthermic intraperitoneal chemotherapy (HIPEC) is carried out by perfusing the abdominal cavity with 43°C circulating hyperthermic saline containing anticancer drugs. HIPEC treatment has been applied to gastric cancer, colorectal cancer, ovarian cancer, pancreatic cancer, and other abdominal cancer types. Poor peritoneal penetration of chemotherapy drugs is an outstanding limitation in the treatment of gastric cancer. HIPEC has thus been administered to patients in combination with cytoreductive surgery to achieve better therapeutic effects. Cumulative evidence has shown the beneficial treatment effect of HIPEC. Studies showed that a combination of cytoreductive surgery (CRS) and HIPEC can reduce the incidence of peritoneal recurrence of advanced gastric cancer and improve the median survival (2–4). A systematic review (5) on 7 studies (6–12) has been done to compare the prophylactic HIPEC after surgery for patients with gastric cancer without clinically evident metastases or positive peritoneal cytology sign. Their analysis results suggested that, compared to surgery alone, the combination with HIPEC may decrease peritoneal recurrence and increase survival rate without affecting the morbidity and mortality of patients despite the overall risk of bias in these studies that is likely due to the non-standard of care in the studies that was carried out more than 10 years ago. A meta-analysis of randomized controlled trials in patients with advanced gastric cancer and peritoneal metastases also showed a beneficial effect of HIPEC in terms of 3-year survival rate and complete response rate (13). Continuous breakthroughs have been made in developing the theoretical basis and technical execution in HIPEC, which further improved the treatment effect (14, 15). Because of the promising treatment of HIPEC, the International Conference on Peritoneal Cancer in Amsterdam (Netherlands) and the American Anti-Cancer Association adopted CRS combined with HIPEC as the standard treatment for gastric cancer with peritoneal metastasis (16, 17).

Similar to the fact that cancer patients respond distinctly to immunotherapy, patients also have different responses to HIPEC treatment. The different results observed regarding the efficacy of HIPEC imply that not all patients can benefit from the treatment and proper stratification may be necessary before HIPEC treatment (14, 15). Few studies have been carried out to examine the biomarkers that can effectively predict the efficacy of HIPEC and stratify patients for HIPEC treatment. Through a candidate gene approach, three studies identified biomarkers, the expression level of which can be predictive for the efficacy or resistance to HIPEC treatment on patients with colorectal peritoneal metastasis, including Bloom syndrome protein (BLM) (18), Vascular endothelial growth factor (VEGF), Versican (VCAN) (19), PAX interacting protein 1 (PAXIP1), and Single stranded DNA binding protein 2 (SSBP2) (20). A recent study focusing on candidate genetic variants reported the association of NQO1*3 allele with poor peritoneal recurrence rate and low disease-free survival (21) among colorectal cancer patients who received cytoreductive surgery plus hyperthermic intraperitoneal mitomycin C. There was no study examining the genetic association with efficacy of HIPEC on gastric cancer. Furthermore, an unbiased study via a genomic approach may be more fruitful in identifying biomarkers to predict HIPEC efficacy.

**METHODS**

**Tumor Samples and Clinical Characteristics**

The study was approved by the institutional review boards of the Affiliated Cancer Hospital of Guangzhou Medical University. All tissue samples were obtained with the approval of patients’ consents from 2010 to 2017. The recruited patients met the following criteria. They had proven gastric cancer with histopathology and received CRS and closed HIPEC treatment after surgery. HIPEC was administered intraperitoneally with chemotherapeutic agents in 4–6 L of perfusate at a temperature of 43°C for 90 min with a flow rate of 400–600 ml/min. The resected tumor samples were examined by the pathology department to confirm the stage of tumor tissue. The tumor tissues and adjacent regions were prepared as formalin-fixed paraffin-embedded (FFPE) tissue blocks following the standard protocol of the diagnosis laboratory.

Approximately 30% of GC patients have regional spread at diagnosis; the local regional progression of gastric cancer generally results in peritoneal metastases (PMs), which have a significant negative impact on the overall survival (OS) and quality of life as a result of refractory ascites, progressive intestinal obstruction, and uncontrollable abdominal pain (22). CRS+HIPEC could obviously decrease the volumes of ascites for a long time compared to only CRS for advanced GC patients. We checked the OS as the criteria of HIPEC effect. Patients with survival time of more than 1 year after HIPEC treatment were classified as the durable clinical benefit (DCB) group, and those with less than 1 year were included in the no durable benefit (NDB) group.

**Statistical Analysis**

The data analysis was conducted using IBM SPSS Statistics 24.0 (SPSS, Inc., Chicago, IL, USA). Continuous data were expressed as mean ± standard deviation (SD), and inter-group comparison was conducted using an independent-samples t-test. Count data were expressed by percentage or constituent ratio, and the comparison between groups was carried out by chi-square test or exact probabilities method. The level of statistical significance was set at p < 0.05 and α = 0.05. Specifically, the comparison of baseline characteristics of patients was conducted with the following statistical methods: age between groups was examined by unpaired t test; the comparison of sex, Eastern Cooperative Oncology Group (ECOG), and degree of
differentiation between groups was examined by chi-square test; and the comparison of the degree of peritoneal metastasis (P degree), ascites, and number of chemotherapy in half a year between groups was examined by Fisher exact test. The Kaplan–Meier (K–M) method and the log-rank test were used to evaluate OS.

**Whole-Exome Sequencing**

Tumor samples and para-cancer control samples from FFPE tissue blocks were analyzed by whole-exome sequencing (WES). The para-cancer control samples were taken from the normal tissue within the 1–5-cm distance from the visible tumor area. Library preparations were performed with KAPA Hyper Prep Kit (KAPA Biosystems, USA). Target enrichment was performed using the xGen Exome Research Panel and Hybridization and Wash Reagents Kit (Integrated DNA Technology, USA) according to manufacturer’s protocol. Standard WES was performed with paired-end sequencing on the Illumina HiSeq2000 platform to generate reads of 2 × 100 bp with an average of 200× mean target coverage for tumor samples and 20× mean coverage for controls.

**Exome Analysis Pipeline**

Quality control and filtering steps were performed on the raw sequencing data, and the data were further checked using software FastQC, including per sequence quality scores, GC content, per base sequence quality, per base sequence content, sequence duplication levels, and overrepresented sequence. Then, the Quality Control (QC)’ed sequencing data were aligned to the Grch37 genome built using Burrows–Wheeler Aligner (BWA) (v0.7.12) (23). Picard Tool was used to mark duplicates, and Genome Analysis Tool Kit (GATK) was used to conduct indel realignment, base-quality score recalibration, and duplicate-read removal (24). Variant calls for both Single Nucleotide Variation (SNV) and Insertion Deletion (INDEL) were generated using software VarDict (25) based on the paired tumor–normal variant calling algorithm. Variant annotation was performed using ANNOVAR (26) and vcf2maf on the VCF file, and further filtering was described below.

**Variant Filtering**

Variants were filtered out in any of the following conditions: (allele frequency × read depth <6) and (mean number of mismatches >1.0 and mean mapping quality <55.0); or (mean number of mismatches >2.0 and mean mapping quality <60.0); or (read depth <10) and (alternate allele quality <45); or alternate allele quality <55 and allele frequency <0.2 and somatic variant p-value >0.06. In addition, variants were filtered out if the reference genotype likelihood of their para-cancer control were >3.5. To keep stringent data quality, we only kept strong somatic variants. Then, variants were further filtered based on their predicted effect on protein structure and function and variant allele frequency in large population databases. Variants that might have an impact on protein function (e.g., in_frame_del, in_frame_ins, missense_mutation, nonsense_mutation, nonstop_mutation, splice_site, and translation_start_site) and allele frequency <0.01 in the following databases were kept, including ExAC_EAS, gnomAD_exom_ALL, gnomAD_exom_EAS, gnomAD_genome_ALL, gnomAD_genome_EAS, 1000g2015aug_all, and 1000g2015aug_eas.

**Analysis of Tumor Mutational Burden on HIPEC Efficacy**

Tumor mutational burden (TMB) was defined as the total number of strong somatic non-synonymous mutations in each tumor exome that passed our QC filtering. Mann–Whitney test was used to compare the number of somatic mutations in DCB and NDB groups. Patients were divided into the high mutation group and the low mutation group based on the median of somatic mutation number of all patients, which is 373. It is about 9.8 mutations/Mb, which is similar to the TMB threshold adopted in other studies (27, 28). Log-rank test was used to compare K-M survival curves between the two groups.

**Association Analysis at Single-Variant Level**

We first evaluated the association of variants with HIPEC efficacy at the single-variant level by Fisher’s exact test, and the statistical significance threshold was p-value <0.05.

**Association Analysis at the Gene Level**

RVTESTS (29) were used to perform correlation analysis of rare variants at the gene level. We used the VCF file generated from the above pipeline as the input file for RVTESTS, followed by Fisher’s exact test (CMC Fisher) to evaluate genes being positively or negatively associated with the efficacy of HIPEC.

**Pathway Enrichment Analysis of Genes Associated With HIPEC Efficacy**

Pathway enrichment analysis was performed via WEB-based GEne SeT AnaLysis Toolkit (30) (http://www.webgestalt.org/) that is based on the hypergeometric test.

From International Cancer Genome Consortium (ICGC) databases (https://dcc.icgc.org/), whole-genome sequencing data and follow-up data of Chinese population were obtained, and the TMB value of non-synonymous mutations in the exon regions was calculated for each patient. The patients were divided into high TMB group and low TMB group by the median value of TMB, and K-M survival curve was performed to examine the association between TMB and survival.

**RESULTS**

**High Tumor Mutational Burden Was Associated With Improved Patient Survival**

We conducted a retrospective study to identify genetic determinants for HIPEC response involving 18 gastric cancer patients, among whom 8 patients showed DCB (patients with survival time of more than 1 year) and 10 patients had NDB. The baseline characteristics of these patients are shown in Table 1. The average age of the NDB group is significantly lower than that of the DCB group (43.8 ± 9.7 vs. 54.1 ± 8.1, p = 0.030), while the other
Table 1 | The baseline characteristics of patients who received hyperthermic intraperitoneal chemotherapy (HIPEC).

|                | DCB (n = 8) | NDB (n = 10) | P      |
|----------------|-------------|--------------|--------|
| Age (yr)       | 54.1 ± 8.1  | 43.8 ± 9.7   | 0.030* |
| Sex            |             |              |        |
| Male           | 3 (37.5)    | 4 (40.0)     | 0.999  |
| Female         | 5 (62.5)    | 6 (60.0)     |        |
| ECOG (score)   |             |              |        |
| 0–1            | 7 (87.5)    | 9 (90.0)     | 0.999  |
| 2–4            | 1 (12.5)    | 1 (10.0)     |        |
| Degree of differentiation |          |              |        |
| G1+G2          | 2 (25.0)    | 2 (20.0)     | 0.999  |
| G3+GX          | 6 (75.0)    | 8 (80.0)     |        |
| P (degree)     |             |              |        |
| p1x            | 1 (12.5)    | 0 (0.0)      | 0.999  |
| p1             | 1 (12.5)    | 2 (20.0)     |        |
| p2             | 3 (37.5)    | 4 (40.0)     |        |
| p3             | 3 (37.5)    | 4 (40.0)     |        |
| Ascites        | No          | 6 (75.0)     | 5 (50.0)| 0.552  |
| A small amount of chemotherapy in half a year | 2 (25.0) | 5 (50.0)     |        |
| 1–3            | 7 (87.5)    | 7 (70.0)     | 0.751  |
| 4–6            | 1 (12.5)    | 3 (30.0)     |        |

ECOG, the grade of Eastern Cooperative Oncology Group. They were classified into durable clinical benefit (DCB) group and no durable benefit (NDB) group based on their response to treatment.

*p-value < 0.05.

characteristics were similar between DCB and NDB groups. We analyzed the WES data of these tumor samples with their matched normal tissues for germline references. The mean target coverage in the WES data of tumor samples is 200× and, on average, >90% of the target sequence was covered to a depth of >10×. The number of non-synonymous somatic mutations in all patients ranged from 90 to 3,027, with a median of 373.

We first analyzed the overall TMB of non-synonymous strong somatic mutation with low frequency in a large population database (variant allele frequency <0.01). The TMB was defined as the total number of strong somatic non-synonymous mutations in the coding regions of the human genome. The TMB in the DCB group ranges from 113 to 3,027, with a median of 799.5; and the TMB in the NDB group ranges from 90 to 569, with a median of 278. Therefore, patients with a partial or stable response to HIPEC had a significantly longer survival time than that of patients with low TMB (HR = 4.82, 95% CI = 1.46–15.95, p = 0.001; Figure 1B). In addition, analyzing the data of Chinese gastric cancer patients in the ICGC database, we found no significant association between TMB and patient survival time (p > 0.05; Figure 1C). In addition, a previous research with comparable sequencing approach among 262 GC patients after excluding patients who received routine preoperative chemoradiotherapy or biological immunotherapy showed that the prognosis of GC and OS are better among patients with higher TMB than those with lower TMB (28). Thus, the association of high TMB and better response to HIPEC observed in our study is not likely due to the relationship between TMB and prognosis of gastric cancer per se. The above evidence suggests an association between high TMB and longer survival for patients who received HIPEC treatment.

We further conducted a replication study. The replication cohort is composed of an independent set of 15 GC samples from patients with similar HIPEC treatment, including 10 patients in the DCB group and 5 patients in the NDB group according to the same criteria. The baseline clinical characteristics of the replication cohort were similar to those of the discovery cohort (Table 2). The median TMB was 364.5 in tumors from patients in the DCB group compared to 233 in those in the NDB group (Mann–Whitney p = 0.02) (Figure 1D). With the same TMB criteria, the GC samples were classified as high TMB group (n = 7) and low TMB group (n = 8). K-M survival analysis was similarly conducted to measure the effect of TMB on the OS of GC patients in the replication cohort. The results confirmed that patients with high TMB had a significantly longer survival time than that in patients with low TMB (HR = 5.842, 95% CI = 1.179–28.94, p = 0.031; Figure 1E).

Single Variant Associated With HIPEC Therapy

We then sought to identify mutations associated with efficacy of HIPEC therapy. The results based on Fisher’s exact test did not suggest any strong somatic mutations surpassing statistical significance, which is likely due to the limitation of a small sample size. Three mutations occurred in either the DCB or the NDB group specifically without carriers in the other group (Table 3).

Genes Correlated With HIPEC Treatment

We then focused on identifying genes harboring mutations recurrently and selectively associated with response or resistance to HIPEC. We found that genes GPI, PCDH9, and C21orf140 harbored mutations in three or more NDB patients but none in the DCB group, and further statistical analyses of collapsing variants to the gene level showed a significant negative correlation with HIPEC response (p < 0.05). On the other hand, a total of 25 genes bearing mutations are recurrently and selectively enriched in the DCB group but none in the NDB group (Figures 2A, B). Several of these genes were known to be involved in regulating cancer proliferation, metastasis, or invasion (Supplementary Table S1). Analyzing the association
between non-synonymous mutations in these genes and OS of Chinese gastric cancer patients in the ICGC database, we did not find any significant association with OS. We checked the distribution of the above enriched single variants and genes in the replication cohort. Gene MUC16 harbored mutations in three patients of the DCB group but none in the NDB group; gene DNAH3 contained mutations in four patients of the DCB group but none in the NDB group; the other genes did not exhibit such enrichment. It is not unexpected for discrepancies to appear at the variant and gene levels between the discovery and replication cohorts, considering the sample size. Thus, studies with larger sample sizes are certainly required to identify and validate predictive markers for HIPEC treatment.

Gene Set Enrichment Analysis of Significantly Correlated Genes

To identify potential signaling pathways that may contribute to HIPEC sensitivity, we conducted pathway enrichment on 348 genes that bear mutations occurring only in the DCB group but not the NDB group. Among the 134 genes that were mapped to Kyoto Encyclopedia of Genes and Genomes (KEGG) pathways, the pathways of extracellular matrix (ECM)–receptor interaction, Notch signaling pathway, focal expression, and between non-synonymous mutations in these genes and OS of Chinese gastric cancer patients in the ICGC database, we did not find any significant association with OS. We checked the distribution of the above enriched single variants and genes in the replication cohort. Gene MUC16 harbored mutations in three patients of the DCB group but none in the NDB group; gene DNAH3 contained mutations in four patients of the DCB group but none in the NDB group; the other genes did not exhibit such enrichment. It is not unexpected for discrepancies to appear at the variant and gene levels between the discovery and replication cohorts, considering the sample size. Thus, studies with larger sample sizes are certainly required to identify and validate predictive markers for HIPEC treatment.

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### TABLE 3 | The three variants that are specifically enriched in the durable clinical benefit (DCB) group or the no durable benefit (NDB) group.

| Chr | Pos   | Variant          | AA alteration | Cases | Controls |
|-----|-------|------------------|---------------|-------|----------|
| 11  | 1092910 | NM_002457:exon31:c.G4729A | MUC2:p.G1577S | 3     | 0        |
| 19  | 23158703 | NM_001267:exon4:c.T1436C | ZNF728:p.L479P | 3     | 0        |
| 13  | 10396322 | NM_001146:exon4:c.C7725A | CCDC168:p.N2575K | 0     | 3        |

### FIGURE 2 | The genes bearing mutations specifically enriched in the durable clinical benefit (DCB) group or no durable benefit (NDB) group. (A) The location of the mutations in the top response-associated genes with respect to the structural domains of each protein product encoded by these genes. (B) The genes carrying mutations that only occurred in patients in the DCB group but none in the NDB group. The yellow color indicates a patient in the DCB group carrying a mutation in the corresponding gene. The lower panel showed the total number of mutations in each patient. The number on the right shows the p-value of Fisher exact test comparing the number of GC patients carrying mutation(s) in each gene between the DCB and NDB groups. (C) The pathways enriched by genes with at least one mutation specifically occurring in patients in the DCB group.
human papillomavirus infection showed significant enrichment (q-value <0.05) (Figure 2C).

**DISCUSSION**

Though HIPEC has been widely administered to patients of gastric cancer, it has been controversial about its efficacy likely due to tumor heterogeneity; therefore, it is important to identify genomic determinants of response to HIPEC treatment. To our knowledge, this is the first study examining molecular features of HIPEC efficacy using an unbiased high-throughput sequencing approach. Within our discovery cohort and a replication cohort, we found a significant association between high TMB and patients’ beneficial treatment effect.

HIPEC may induce immune response activation. It has been reported that focal thermal ablation of tumor may stimulate a systemic antitumor immune response (31). From the tumor cells and surrounding tissues that were damaged by heat, a variety of immune molecules were released, including cytokines and chemokines. It has been reported that the serum levels of tumor necrosis factor-α (TNFα) and interleukin-1β (IL-1β), IL-6, and IL-8 were elevated after thermal ablation (32–35). In addition, heat shock proteins (HSPs) such as HSP70, which is highly expressed by tumor cells, were detected to be increased following thermal ablation. HSPs play important roles in inhibition of apoptosis inside cells, as well as antigen-presenting and activating dendritic cells and modulating the activity of T-regulatory cells in the microenvironment of tumor cells. HSPs are also involved in other antitumor immune responses (36–39). Increased activation of dendritic cells and decreased activity of T-regulatory cells have been observed in multiple studies (32, 33). Immune cell infiltration was also observed, such as B and T lymphocytes, dendritic cells, neutrophils, natural killer (NK) cells, and macrophages. Such intense immune and inflammatory response occurring after focal thermal ablation may also be able to explain the positive correlation between TMB and response to HIPEC. We hypothesize that tumor cell damage caused by HIPEC results in the release of various intracellular molecules to the tumor microenvironment, and the higher the TMB, the more neo-antigens may be captured by immune cells, therefore leading to better response after treatment.

We identified multiple genes carrying mutations enriched in the DCB or NDB group specifically. The expression level or SNPs in these genes have been reported as being significantly associated with several cancer types. Functional studies showed that these genes have been implicated in key signaling pathways in cancer, such as the proliferation, migration, and invasion of tumor cells, as being summarized in Supplementary Table S1. However, due to the limited sample size, the individual variant or gene is not robust enough to be identified as a definitive predictive marker for responses to HIPEC treatment. A larger cohort with more samples needs to be sequenced to validate these findings, and experimental studies need to be carried out to investigate the role of these genes in molecular and cellular responses to HIPEC.

**Limitations**

The limitation of our study lies in three aspects. First, the sample size is small; therefore, it is not unexpected that none of the enriched variants at the single-variant level is statistically significant. The results from gene level need to be further examined in studies with larger sample sizes. Second, an independent replication study sampled from other populations is needed to investigate whether TMB that was identified as a predictive mutation feature for HIPEC efficacy can be generalized to other populations. Third, functional studies are warranted to further examine how such a mutational feature contributes to response or resistance to HIPEC treatment. Nevertheless, our study demonstrated that a high mutational burden is significantly associated with better response to HIPEC, which is consistent with previous reports that HIPEC induces immune responses and high TMB confers better immunotherapy efficacy.

**Conclusion**

Though limited by the small sample size and single data type at the DNA level, we conducted the first biomarker study of HIPEC treatment to gastric cancer based on unbiased high-throughput sequencing data. We found that TMB is significantly associated with HIPEC efficacy. The genes identified are relevant to the activation and inhibition of signaling pathways in cancer cells. Our results demonstrated that high mutational burden confers a beneficial HIPEC treatment effect, which warrants further confirmation in independent cohorts and functional examination of the underlying mechanism of HIPEC sensitivity. Our study provides insights into the development of mutational features to select patients for effective HIPEC treatment.

**DATA AVAILABILITY STATEMENT**

According to national legislation/guidelines, specifically the Administrative Regulations of the People’s Republic of China on Human Genetic Resources (http://www.gov.cn/zhengce/content/2019-06/10/content_5398829.htm, http://english.www.gov.cn/policies/latest_releases/2019/06/10/content_281476708945462.htm), no additional raw data are available at this time. Data of this project can be accessed after an approval application to the China National Genebank (CNGB, https://db.cnbg.org/cnsa/). Please refer to https://db.cnbg.org or email CNGBdb@cnbg.org for detailed application guidance. The accession code CNP0002628 should be included in the application.

**ETHICS STATEMENT**

The studies involving human participants were reviewed and approved by the institutional review boards of the Affiliated Cancer Hospital of Guangzhou Medical University. The patients/
participants provided their written informed consent to participate in this study.

AUTHOR CONTRIBUTIONS

SC and JLi designed and supervised the study. LZ and YT prepared patients’ samples for sequencing and collected clinical data. LZ and XH performed data analysis. The other authors partially participated in the study and helped on data analysis and interpretation. LZ, XH, QX, JLi, and SC wrote the initial article. All authors have read and approved the article.

FUNDING

This work was supported by the National Natural Science Foundation of China (No. 81972918, 81771769, and 82172885), Natural Science Foundation of Guangdong Province (No. 2018A030310249, 2020A1515010818, and 2021A1515012392), and Key Clinical Technique of Guangzhou (No. 2019ZD16 and Guangzhou Key Medical Discipline Construction Project Fund.

ACKNOWLEDGMENTS

We are grateful to all the patients for their participation in the study and Nanjing Geneseeq Technology Inc. and 3D Medicines Inc. for conducting whole-exome sequencing.

SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fonc.2022.796263/full#supplementary-material

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Conflict of Interest: Authors KL and YS are employed by Nanjing Geneseeq Technology Inc.

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