Predictive Biomarkers of Dicycloplatin Resistance or Susceptibility in Prostate Cancer

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Background: Prostate cancer (PCa) is among the leading causes of cancer mortality. Dicycloplatin is a newer generation platinum-based drug that has less side effects than cisplatin and carboplatin. However, its effects in PCa is mixed due to lack of appropriate stratifying biomarkers. Aiming to search for such biomarkers, here, we analyze a group of PCa patients with different responses to dicycloplatin.

Methods: We carried out whole-exome sequencing on cell-free DNA (cfDNA) and matched leukocyte DNA from 16 PCa patients before treatment with dicycloplatin. We then compared the clinical characteristics, somatic mutations, copy number variants (CNVs), and mutational signatures between the dicycloplatin-sensitive (nine patients) and dicycloplatin-resistant (seven patients) groups and tested the identified mutations, CNV, and their combinations as marker of dicycloplatin response.

Results: The mutation frequency of seven genes (SP8, HNRNPCL1, FRG1, RBM25, MUC16, ASTE1, and TMBIM4) and CNV rate of four genes (CTAGE4, GAGE2E, GAGE2C, and HORMAD1) were higher in the resistant group than in the sensitive group, while the CNV rate in six genes (CDSN, DPCR1, MUC22, TMSB4Y, VARS, and HISTCH2AC) were lower in the resistant group than in the sensitive group. A combination of simultaneous mutation in two genes (SP8/HNRNPCL1 or SP8/FRG1) and deletion of GAGE2C together were found capable to predict dicycloplatin resistance with 100% sensitivity and 100% specificity.

Conclusion: We successfully used cfDNA to monitor mutational profiles of PCa and designed an effective composite marker to select patients for dicycloplatin treatment based on their mutational profile.

Keywords: prostate cancer, dicycloplatin, whole-exome sequencing, biomarker, anti-cancer (anticancer) drugs

INTRODUCTION

Prostate cancer (PCa) is one of the most common cancers in men worldwide and has the second highest mortality (Siegel et al., 2020). It was estimated that there would be about 190,000 new cases of PCa (21% of all are male cancers) and 30,000 deaths (10% of all are male cancer deaths) in the United States alone in 2020 (Siegel et al., 2020). Hormone therapy is an effective therapy that
can improve the survival time and clinical benefits for early stage PCAs. However, about 10–20% of patients will inevitably develop into drug resistance within 5 years during the course of treatment, leading to castration-resistant prostate cancer (Kirby et al., 2011). Platinum-based chemotherapies, such as cisplatin and carboplatin, are attracting more and more attention in the treatment of cancer (Apps et al., 2015). These drugs mainly function through induction of DNA cross-links, therefore inhibiting DNA synthesis, mitosis, and induce apoptosis (Lokich and Anderson, 1998; Ozols et al., 2003). However, the clinical benefits from these drug therapies are still low, they can prolong patients’ overall survival for only 3–6 months (Fortin et al., 2013). Moreover, their clinical application is limited by severe adverse effects, including ototoxicity, neurotoxicity, and myelosuppression (Rossi et al., 2012). Therefore, there is an urgent need to develop more effective drugs to PCa with lower side effects.

Dicycloplatin is a derivative of carboplatin in which a carboxylic acid ligand is bound to the carboplatin moiety through hydrogen bonds. Therefore, it has a more stable chemical structure and better aqueous solubility than carboplatin (Yang et al., 2010). Previous studies showed that dicycloplatin has a better anticancer activity and much lower toxicity than cisplatin and carboplatin in patients with non-small cell lung cancer (Wang and Yu, 2004). In vivo and in vitro studies showed that dicycloplatin can induce cell cycle arrest and apoptosis and inhibit cell proliferation through reactive oxygen species stress-mediated death receptor pathway and mitochondrial pathway (Yu et al., 2014). Furthermore, a phase II clinical trial in non-small cell lung cancer has demonstrated the safety and efficacy of dicycloplatin in combination with paclitaxel (Liu et al., 2014). However, a considerable number of patients did not benefit from dicycloplatin treatment with some unknown reasons. Effective and reliable prognostic factors, therefore, are desperately needed in order to target dicycloplatin to the subset of patients who would benefit most from the treatment.

Whole exome sequencing (WES) is a high-throughput sequencing technology that can explore the whole functional DNA sequence and genetics variations of each patient to uncover novel molecules that may be related to therapies. In this retrospective study, we carried out WES on cell-free DNA (cfDNA) from blood- and patient-matched leukocyte DNA in 16 PCA patients before they received dicycloplatin monotherapy. A comprehensive analysis was performed to search for the association between the clinical outcomes of dicycloplatin treatment and molecular characterization of the patients, such as somatic mutations, copy number variants, and mutational signatures.

MATERIALS AND METHODS

Patients, Clinical Evaluation, and Sample Collection

Sixteen PCA patients were retrospectively enrolled from Chinese PLA General Hospital with the following criteria: (Siegel et al., 2020) Eastern Cooperative Oncology Group performance status of 0–2; (Kirby et al., 2011) patients with distant metastasis who have received surgical castration or medical castration (serum testosterone ≤ 50 ng/dl or 1.7 nmol/ml); (Apps et al., 2015) at least 4 weeks after antiandrogenic therapy; and (Lokich and Anderson, 1998) the value of prostate specific antigen (PSA) was more than 2 ng/ml, and sustained increase more than 50% one week. The patients were on dicycloplatin treatment during 2016–2019. The study protocol was approved by the Ethics Committee of Chinese PLA General Hospital (approval number S2017-032-02). All patients provided a written informed consent.

The pathological diagnosis was performed by experienced pathologists of the hospital. Tumor response to the treatment was evaluated based on the patients’ radiological images [computed tomography (CT) and magnetic resonance imaging (MRI)] according to the Response Evaluation Criteria in Solid Tumors, version 1.1 (Tirkes et al., 2013).

Whole-Exome Sequencing

Ten milliliter of blood specimens was collected before dicycloplatin treatment and used to prepare cfDNA by MagMAX(CELl-Free DNA Isolation Kit (Applied Biosystems)(A29319) and leukocyte DNA by Maxwell® RSC Blood DNA Kit (Promega AS1400). cfDNA for WES needed to meet the following conditions: there is an obvious peak in the region of 100–300 bp, and the regional molarity of the peak was >0.7 of all contents between 100 bp and 42 kb (indicating that there is no contamination from large genomic DNA). The purified DNA was sonicated using a Covaris L220 sonicator and hybridized to the probes in SureSelect Human All Exon V5 kit (cat. # 5190-6209 EN, Agilent Technologies, Sta. Clara, CA, United States) to capture exonic DNA, then prepared to libraries using the SureSelectXT Low Input Target Enrichment and Library Preparation system (cat. # G9703-90000, Agilent Technologies). Paired end reads of 150 × 150 bp were generated from the libraries using an Illumina NovaSeq-6000 sequencer (Illumina, San Diego, CA, United States). Image analysis and base calling were done using the onboard RTA3 software (Illumina). After removing adapters and low-quality reads, the reads were aligned to National Center for Biotechnology Information (NCBI) human genome reference assembly hg19 using the Burrows–Wheeler Aligner alignment algorithm and further processed using the Genome Analysis Toolkit (GATK, version 3.5), including the GATK Realigner Target Creator to identify regions that needed to be realigned. Somatic mutations, including single-nucleotide variants (SNVs), indel, and copy number variation (CNV) were determined by a comparison between the aligned sequences from cfDNA and patient-matched leukocyte DNA using the MuTect/ANNOVAR/dbNSFP31, VarscanIndel, and CNVnator software, respectively, as previously reported (Zang et al., 2019). The mutational signature classification was based on COSMIC Mutational Signature (version 2—March 2015), which was generated from studies performed by others (Maitra et al., 2013; Alexandrov et al., 2015; Nik-Zainal et al., 2016). Tumor mutation burden (TMB) was defined as the total number of somatic non-synonymous mutations in each sample according to a previous method for WES data (Chalmers et al., 2017). All autosomal
microsatellite tracts containing 1–5 bp repeating subunits in length and comprising five or more repeats in GRCh37/hg19 were identified using MISA\(^1\) and used to calculate microsatellite instability score (MSI). MSI score was calculated by the number of unstable microsatellite sites/total valid sites.

The WES data of each patients were submitted in NCBI with submission number SUB9593847 and accession number PRJNA727718.

**Statistical Analysis**

All statistical analyses were performed using R\(^2\) or SPSS 25 for Windows (SPSS Inc., Chicago, IL, United States). Differences in the distribution of somatic mutations, mutational signatures, and clinical characteristics between patient subgroups were evaluated by the Fisher’ exact test and Mann–Whitney U test for categorical and continuous parameters, respectively, and events of \(p < 0.05\) were considered statistically significant.

**RESULTS**

**Clinicopathological Characteristics of the Patients**

In total, 16 patients treated with dicycloplatin were enrolled, and their baseline characteristics are shown in **Table 1**. Nine of the patients were sensitive to the treatment, including six with partial response (PR) and three with complete response (CR) evaluated by CT and MRI scans. The remaining seven patients were resistant to the treatment, including six with progressive disease (PD) and one with stable disease (SD). The overall tumor response rate was 56.3%. The representative diagnostic images are shown in **Figure 1**. The PSA and free PSA (fPSA) levels of each patients during treatment course were analyzed; both PSA and fPSA were decreased in patients with PR and CR, while they were increased in patients with PD and SD (**Supplementary Figure 1**). Patients who received endocrine therapy before dicycloplatin treatment were more likely resistant to dicycloplatin (**Table 1**, Fisher’s exact test, \(p = 0.003\)). Other factors, such as age, smoking and drinking history, tumor stage, Gleason score, pre-dicycloplatin treatment history of surgery, chemotherapy, and radiotherapy, all showed no significant difference between the dicycloplatin-sensitive and dicycloplatin-resistant groups.

**The Mutational Landscape of the Patients**

The whole-exome DNAs of the patients were captured and sequenced on an Illumina platform. The average sequencing depth was \(\times 925\) for cfDNA libraries and \(\times 117\) for leukocyte DNA libraries. The sequences from cfDNA were compared to matched leukocyte DNA to give somatic genetic changes including SNV, indel, and CNV (see section “Materials and Methods”). The median TMB was similar between treatment-sensitive group (3.3 mutations/Mb) and treatment-resistant group (4.4 mutations/Mb), with no statistical difference (Mann–Whitney U test, \(p = 0.680\)). There were also no differences in MSI score, the proportions of gene amplifications, and deletions

**TABLE 1 | Comparison of baseline characteristics between dicycloplatin-sensitive and dicycloplatin-resistant patients.**

| Baseline characteristic | Sensitive (\(n = 9\)) | Resistant (\(n = 7\)) | \(p\) value | All patients (\(n = 16\)) |
|-------------------------|-----------------------|-----------------------|-------------|--------------------------|
| Age, median (range), years | 66 (61–79) | 63 (51–72) | 0.210 | 66 (51–79) |
| Smoking history, no. (%) | | | | 0.633 |
| Yes | 4 (44.4) | 2 (28.6) | | 6 (37.5) |
| No | 5 (55.5) | 5 (71.4) | | 10 (62.5) |
| Drinking history, no. (%) | | | | 0.596 |
| Yes | 2 (44.4) | 3 (42.9) | | 6 (37.5) |
| No | 7 (55.5) | 4 (57.1) | | 10 (62.5) |
| Tumor stage at diagnosis, no. (%) | | | | 1 |
| II | 1 (11.1) | 0 (0.0) | | 1 (6.3) |
| IV | 8 (88.9) | 7 (100.0) | | 15 (93.8) |
| Gleason sum at diagnosis, no. (%) | | | | 1 |
| \(\leq 7\) | 6 (66.6) | 4 (57.1) | | 10 (62.5) |
| \(\geq 8\) | 3 (33.3) | 3 (42.9) | | 6 (37.5) |
| Prior treatment for PCa, no. (%) | | | | 3 (18.8) |
| Surgery | 2 (22.2) | 1 (14.3) | | 3 (18.8) |
| Radiotherapy | 1 (11.1) | 2 (28.6) | | 3 (18.8) |
| Chemotherapy | 3 (33.3) | 5 (71.4) | | 0.157 |
| Endocrine therapy | 2 (22.2) | 7 (100.0) | | 0.003 |
| Pretreatment PSA level, median (range), ng/ml | 87.6 (9.9–775.6) | 79.3 (2.9–800) | 0.837 | 86.0 (2.9–800) |
| Pretreatment-free PSA level, median (range), ng/ml | 14.0 (0.5–48.6) | 10.8 (0.9–29.8) | 0.681 | 11.2 (0.5–48.6) |

PSA, prostate-specific antigen.

\(p\) values are based on Fisher’ exact test and Mann–Whitney U test for categorical and continuous parameters, respectively.

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\(^{1}\)http://pgrc.ipk-gatersleben.de/misa/misa.html

\(^{2}\)https://cran.r-project.org
between the two groups (Mann–Whitney U test, $p > 0.05$; Figure 2).

The most common somatic mutated genes were ANKRD36C (56.3%), KIAA2018 (56.3%), MUC4 (56.3%), TMBIM4 (50.0%), and RNFI45 (50.0%) among all 16 patients (Figure 3A). We identified seven genes whose mutation rates were significantly higher in the resistant group than in the sensitive group: SP8, HNRNPC1L1, FRG1, RBM25, MUC16, ASTE1, and TMBIM4 (Fisher’s exact test, $p < 0.05$; Figure 3B). The mutation rates of SP8, HNRNPC1L1, and FRG1 were 57.1% (4/7) in the sensitive group and 0% in the resistant group. The mutation rates of ASTE1, MUC16, and RBM25 were 71.4% (5/7) in the sensitive group and 11.1% (1/9) in the resistant group.

All mutated genes were screened for a possible link to signaling pathways associated with platinum metabolism curated from the literature, such as cell cycle dependence, bicyclic platinum molecule activation, DNA damage repair, tumor cell apoptosis, drug transmembrane transport, platinum metabolizing drugs, DNA homeostasis disorders, and potential secondary drug resistance. The result showed that the mutation rates of these genes were similar in two groups, except for MUC16 (Supplementary Figure 2).

Copy Number Variations
To further explore events that are related to dicycloplatin treatment, we analyzed CNVs harbored by the patients. LAMTOR5-AS1, JUND, and SCAND1 were the most common genes with CNV deletion, and the number of CNV deleted genes was greater than the number of CNV-amplified genes (Figure 3C). The variation rates of CNV of four genes (CTAGE4, GAGE2E, GAGE2C, and HORMAD1) were higher in the resistant group than in the sensitive group (Fisher’s exact test, $p < 0.05$). In contrast, the variation rates of CNV of six genes (VARS, TMSB4Y, MUC22, DPCR1, CDSN, and HISTCH2AC) were lower in the resistant group than in the sensitive group (Fisher’s exact test, $p < 0.05$; Figure 3D).

Mutational Spectrum and Mutational Signatures
A scan of the observed mutations in the patients showed that C>T was the most common substitution in the cfDNA samples (Figure 4A). T>G substitution was the most common of the six base substitutions in the sensitive group. Its proportion decreased in the resistant group, but there was no statistical difference (Mann–Whitney U test, $p > 0.05$). The COSMIC mutational signatures of the patients are presented in Figure 4B. Signatures 3 and 1 were dominant in all samples with median percentages at 40.1 and 8.1%, respectively. The median percentages of the other mutational signatures ranged from 0 to 2.1%. Signature 12 had a higher proportion in the resistant group than in the sensitive group (Mann–Whitney U test, $p = 0.018$), while the other signatures did not show statistical differences between these two groups (Mann–Whitney U test, $p > 0.05$).
FIGURE 2 | Comparison of the genomic profiles between dicycloplatin-sensitive and dicycloplatin-resistant groups. (A) Tumor mutational burden (TMB). (B) Microsatellite instability (MSI) score. (C) Percentage of exons with amplification (defined as copy number > \(2 \times\) average ploidy). (D) Percentage of exons with deletion (defined as copy number < \(0.5 \times\) average ploidy).

Performance of Gene Classifiers in Predicting Resistance or Sensitivity to Dicycloplatin

There were differences in the frequency of multiple gene mutations and CNV variants between the resistant group and the sensitive group. The mutation frequency of seven genes (SP8, HNRNPCL1, FRG1, RBM25, MUC16, ASTE1, and TMBIM4) and the CNV variation frequency of four genes (CTAGE4, GAGE2E, GAGE2C, and HORMAD1) were higher in the resistant group than in the sensitive group. In order to search for markers of dicycloplatin treatment effects, we first calculated the sensitivity and specificity of these genes individually in assessing dicycloplatin resistance (Table 2, Nos. 1–11). Individually, none of these genes can satisfy a high sensitivity and high specificity at the same time. The sensitivity of three mutated genes (SP8, HNRNPCL1, and FRG1) and two CNV genes (GAGE2E and GAGE2C) was 57.1%, and their specificity was 100.0%. The sensitivity and specificity of three other mutated genes (RBM25, MUC16, and ASTE1) were 71.4 and 88.9%, respectively. Of the 11 genes tested, TMBIM4 had the highest sensitivity (85.7%), but its specificity was the lowest (77.8%). The sensitivity of the remaining two copy number variant genes (CTAGE4 and HORMAD1) was 71.4%, and their specificities were 100 and 88.9%, respectively.

The sensitivity and specificity of the CNV of six genes (VARS, TMSB4Y, MUC22, DPCR1, CDSN, and HISTCH2AC) in assessing dicycloplatin susceptibility were also calculated and shown on Table 2 (Nos. 12–17). The sensitivity of five copy number variant genes (VARS, TMSB4Y, MUC22, DPCR1, and CDSN) was 55.6%, and their specificity was 100%. The sensitivity and specificity of HISTCH2AC were 100 and 71.4%, respectively.

We next calculated sensitivity and specificity of genes in different combinations to assess dicycloplatin resistance and susceptibility. We limited the number of genes in the combination within 3. Among all the tested combinations, one configuration stood out. It consisted of mutations of two genes (either SP8/HNRNPCL1 or SP8/FRG1) and CNV loss of GAGE2C and successfully detected resistant cases with 100% sensitivity and 100% specificity (Table 3, Nos. 1–14). Another configuration of CNV loss of DPCR1 and TMSB4Y together could detect sensitive cases with 88.9% sensitivity and 100% specificity (Table 3, Nos. 15–18).

DISCUSSION

In this study, we found several genes with mutation or CNV to predict the resistance or susceptibility to dicycloplatin.
in PCa and evaluated the patients’ response to a novel platinum drug. To our knowledge, this is the first predictive study in its category. Patients stratification by molecular subtyping would be helpful to improve the effectiveness of dicycloplatin in PCa.

Whole exome sequencing analysis in solid tumors is normally done with tissue samples. However, in the case of PCa, due to both the nature of the organ and surgical manipulation involved, tissue sample is often unavailable. Therefore, we extracted cfDNA from the blood of the PCa patients and used that in WES analysis instead of tumor tissue. Similar application of WES using cfDNA has been reported before and detected SNV, CNV, mutational signatures, TMB, and other genomic parameters in a reasonable accuracy, as verified by a positive correlation between the analysis results from cfDNA and tumor tissue samples (Bos et al., 2020).

In this study, all the patients were classified as stage IV, except one patient who was classified as stage II. This implies that the samples used in the current study were mostly from advanced tumors with high circulating tumor DNA (ctDNA) fractions. Therefore, the sensitivity of identified prediction factors in WES is expected to match to that from tissue samples. Our result indicates that cfDNA is a promising surrogate of tumor tissue because cfDNA can be obtained through a minimal-invasive procedure (blood drawing) and at the same time has the advantage to overcome tumor heterogeneity (Gonzalez-Billalabeitia et al., 2019). We expect that the surrogate strategy may even have expanded clinical uses beyond screening biomarkers for dicycloplatin response.

Among the potential predictive genes of dicycloplatin response that we identified, MUC16 has been extensively studied...
MUC16 is known to promote the progression and metastasis of a variety of malignant tumors, and the abnormal expression of MUC16 can lead to drug resistance to cytotoxic drugs and inhibition of apoptosis (Das et al., 2015). This is consistent with our findings that the mutation frequency of MUC16 was higher in the dicycloplatin-resistant group. Most of the remaining genes that we identified have been found involved in tumorigenesis and disease progression. SP8, which encodes specificity protein 1/Klf-like zinc-finger transcription factor, inhibits KARS-mediated transformation and is also a tumor suppressor by itself (Fernandez-Zapico et al., 2011). The expression of SP8 was decreased in primary gastric cancer compared with normal gastric mucosa in a recent study (Chang et al., 2009). FRG1 expression was decreased in PCa tissues and that affected the migration and invasion of cancer cells (Tiwari et al., 2019). Deleterious mutations of FRG1, which had been identified in calcified pleura fibrous tumor and follicular thyroid cancer, were suggested to contribute to tumorigenesis (Erinjeri et al., 2018; Mehrad et al., 2018). Splicing regulator RBM25 was identified as a tumor suppressor in acute myeloid leukemia, and the low level of RBM25 was associated with high MYC activity and poor prognosis of patients (Ge et al., 2019). In PCa, p53 regulates EMT by activating RBM25, thus promoting tumor progression and metastasis (Yang et al., 2019). However, the roles of the mentioned genes above in platinum metabolism and drug resistance remain unclear and need further investigation.

The sensitivity and specificity to predict dicycloplatin response by the individual actionable genes did not reach 100% (Figure 5).
From a clinical point of view, when the specificity of a screen is <100%, the patients who are false-positively identified can be tested by complementary methods and further cleared with their drug response status. However, if the sensitivity of a screen is <100%, the patients who are false-negatively identified will be missed in the screen. For oncologists and patients, the cost of suboptimal specificity is lower than the cost of suboptimal sensitivity. Therefore, it is preferable to evaluate patients’ drug response with a high-sensitivity screen. One way to increase the sensitivity is through the combination of multiple marker genes. We test several configurations and found the combinations of either \( SP8/\text{HNRNPL1}/\text{GAGE2C} \) or \( SP8/\text{FRG1}/\text{GAGE2C} \) reached 100% in both sensitivity and specificity (Table 3). Although the sample size of this study is
small, the putative drug response marker genes identified should provide a preliminary but critical assessment of the clinical value of dicycloplatin in PCa.

COSMIC mutational signature 25 had the higher proportion in the sensitive group than in the resistant group (Supplementary Figure 3). It was found highly presented in Hodgkin’s lymphoma, although its etiology remains unknown and needs further study.

In addition to the molecular markers, we also found that patients who had prior endocrine therapy were more likely to develop resistance to dicycloplatin (Table 1). It is possible that endocrine therapy and dicycloplatin resistance are mechanistically linked, probably due to clonal selection. However, another possibility is that the endocrine-treated patients were already in a more advanced stage when they received dicycloplatin, therefore had a worse clinical performance overall.

In conclusion, our study identifies that mutation or CNV in several genes are putatively predictive to dicycloplatin response in PCa. However, further tests in cell and animal models are necessary to search for and verify the possible action mechanism of these genes in platin drug resistance. The prediction method that we postulated should be valuable to screen patients suitable for dicycloplatin treatment, therefore reducing the suffering of PCa patients who are predicted as not good responders of the therapy.

DATA AVAILABILITY STATEMENT

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found below: https://www.ncbi.nlm.nih.gov/bioproject/PRJNA727718.

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ETHICS STATEMENT

Written informed consent was obtained from the individual(s) for the publication of any potentially identifiable images or data included in this article.

AUTHOR CONTRIBUTIONS

ML, CL, and GZ carried out carried on the study design and contributed to sample preparation. ML, XZ, and JZ analyzed the clinical data and wrote manuscript. JL and SJ revised the manuscript and approved final manuscript. All authors contributed to the article and approved the submitted version.

SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fgene.2021.669605/full#supplementary-material

Supplementary Figure 1 | Dynamics of PSA and free PSA (fPSA) of patients during treatment course. PSA (A) and fPSA (B) levels were decreased in 6 PR patients and 3 CR patients, were increased in 6 PD patients and 1 SD patient.

Supplementary Figure 2 | Mutational landscape of selected genes associated with tumor and platinum metabolism. (A) Oncoplot of the top 20 mutated gene in dicycloplatin-treated samples. The percentages and the bar plots on the right represent the fraction and the number of samples with variations in the corresponding gene. (B) Differentially somatic mutated genes between dicycloplatin-sensitive and dicycloplatin-resistant group. The dots and horizontal bars denote the hazard rate and 5–95% CI. *P-value < 0.05. NS: not significant.

Supplementary Figure 3 | Comparison of 30 cosmic signatures between dicycloplatin-sensitive and dicycloplatin-resistant patients.
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Conflict of Interest: XZ, JL, and CL were employed by company GenomiCare Biotechnology Co., Ltd., Shanghai, China.

The remaining authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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