Multiplex gene quantification as digital markers for extremely rapid evaluation of chemo-drug sensitivity

Graphical abstract

Highlights
- Non-targeted multiplex genes are screened as digital markers for drug sensitivity
- Transcription level cohort of 10s to 100s genes predicts drug sensitivity
- Digital markers are quantified using qRT-PCR within 1–3 h
- Digital markers guide extremely rapid chemo-drug uses after patient hospitalization

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In brief
Fan et al. developed computational algorithms to establish multiplex digital markers that can evaluate chemotherapy drugs for personalized medicine. The evaluation is completed in up to 3 h and satisfies the needs for immediate chemotherapy after patient hospitalization.
Article

Multiplex gene quantification as digital markers for extremely rapid evaluation of chemo-drug sensitivity

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https://doi.org/10.1016/j.patter.2021.100360

SUMMARY

Current administrations for precision drug uses are limited in evaluation speed. Here, we propose the use of multiplex gene-based digital markers for the extremely rapid personalized prediction of individual sensitivity to cancer drugs. We first screen the transcriptional profiles by applying two to three gene filters and scoring genes by their impact on drug sensitivity and finalize the gene lists by K-nearest neighbors cross-validation. The digital markers are cancer type dependent, are composed of tens to hundreds of gene expressions, and are rapidly quantified by reverse transcription quantitative real-time PCR (qRT-PCR) within 1–3 h after tumor sampling. The area under the receiver operating characteristic curve reached 0.88 when testing the performance of digital markers on organoids derived from colorectal cancer patient tumors. The algorithm and corresponding graphic user interface were developed to demonstrate the promise of digital markers for extremely rapid and personalized drug recommendation.

INTRODUCTION

Cancer is an extremely intractable disease, due to its dynamic, metastatic,1 and heterogeneric2 properties. Therefore, there is no standard therapeutic regimen for most cancer types, and even for the same cancer type, effective therapies for different tumors at different developmental stages are highly variable due to inter- and intra-tumor heterogeneity.2–4 Apart from targeted therapy that can be precisely chosen by detecting known gene mutation(s) or identifiable biomarkers, most chemotherapies are empirical. Precision or personalized drug uses are expected to improve cancer prognosis by increasing the rate of effective treatment, reducing the magnitude of side effects, and avoiding the waste of a therapeutic window.

In recent years, the availability of multi-omics big data and data mining techniques has enabled us to decipher the highly complicated interactive networks among the genomics, transcriptional identities, and underlying therapeutics.5,6 Several public datasets have been established to include genomic, transcriptomic, and drug responsive cohorts for cancer cell lines, including the Genomics of Drug Sensitivity in Cancer (GDSC)7–9 and the Cancer Cell Line Encyclopedia (CCLE),10,11 and for cancer patients, such as The Cancer Genome Atlas (TCGA).12

THE BIGGER PICTURE  In clinical cancer medicine, many patients require immediate chemotherapy after hospitalization. Current administrations for precision drug uses are limited in evaluation speed, including genomic sequencing and tumor organoid evaluation. An extremely rapid evaluation protocol is in high demand to realize drug recommendation within a few hours after tumor sampling. In this work, we have proposed an approach for extremely rapid and personalized drug recommendation.
Many computational algorithms and multiplex models have been developed to decipher the connection between multiomics information and drug sensitivity. For example, linear prediction models, including univariate regression, multivariate regression, and elastic net, have been used to select genomic predictors for anticancer drug sensitivity. Network-based approaches have been attempted to integrate heterogeneous information and learn low-dimensional feature vectors to predict drug-target interactions, as well as identify biomarkers for drug selection using protein-protein interaction networks.

Deep neural networks can integrate the information of gene mutations and transcriptional alterations and improve the prediction of drug responses for large clinical cohorts. The accessibility to such public datasets has expanded the scope of computation-based precision medicine. Sanchez-Vega et al. first proposed oncogenic signaling pathways as a tumor classification strategy, stratified tumors into 64 subtypes, and suggested new therapeutic opportunities by studying the large cohort in TCGA. Recently, Paul et al. discovered common regulatory knobs in gene networks across different tumor types and proposed a new mechanism of tumor classification that is independent of organ specification. A similar finding was further examined in non-tumor diseases. Computational models have also reported successes in probing drug tolerance transition and synergistic drug effects in cancer cells based on their transcriptional data. It is a fact that most tumors lack specific mutations for targeted therapy; however, they possess transcriptional alternations. The aforementioned studies support the investigation of transcriptional identities as markers for therapeutic prediction and personalized drug use.

However, the current RNA sequencing (RNA-seq)-based big data mining has limitations in temporal efficiency. The RNA-seq operation takes 4–6 weeks and requires costly gene sequencers. In many clinical practices, chemo-drug administration takes 4–6 weeks and requires costly gene sequencing. The accessibility to such public datasets has expanded the scope of computation-based precision medicine. The accessibility to such public datasets has expanded the scope of computation-based precision medicine. Sanchez-Vega et al. first proposed oncogenic signaling pathways as a tumor classification strategy, stratified tumors into 64 subtypes, and suggested new therapeutic opportunities by studying the large cohort in TCGA. Recently, Paul et al. discovered common regulatory knobs in gene networks across different tumor types and proposed a new mechanism of tumor classification that is independent of organ specification. A similar finding was further examined in non-tumor diseases. Computational models have also reported successes in probing drug tolerance transition and synergistic drug effects in cancer cells based on their transcriptional data. It is a fact that most tumors lack specific mutations for targeted therapy; however, they possess transcriptional alternations. The aforementioned studies support the investigation of transcriptional identities as markers for therapeutic prediction and personalized drug use.

In this work, we aim to establish an approach that recommends drug use within a few hours after accessing patient tumors. It starts from computational screening of the whole transcriptome to build small gene libraries for prediction of drug response. The transcriptional profiles of screened multiplex gene libraries compose digital (virtual) markers for prediction of chemo-drug sensitivity, which is analogous to the biomarkers beyond a certain transcription level and with a stronger correlation to certain drug sensitivities were retained after being filtered (Figures 1B and 1C). After gene filtering, the genes were scored, and used to compose the gene candidate list. For cell lines, as the RNA-seq data from CCLE and GDSC do not match well, in order to make full use of the data and improve robustness, we performed gene filtering on both datasets separately and synthesized the scores of genes as the rating criteria to finalize the list of gene candidates (see experimental procedures for details).

The second step was K-nearest neighbors (KNN) cross-validation and establishment of digital markers (Figure 1A). The KNN algorithm determined the top m0 genes to compose digital markers and the parameter k0 to reach the highest area under the receiver operating characteristic (ROC) curve (AUC) values of receiver operating characteristic curves (see experimental procedures for details).

The last step was prediction of drug sensitivity using the KNN model. For cell lines, drug sensitivity data are continuous; KNN regression was therefore used to predict the Z scores of the IC50 values. For patient samples, drug sensitivity values were provided as labels according to treatment outcomes; we therefore used the KNN classification here.

In “cross-validation of digital marker-based prediction of drug sensitivity on cancer cell lines and patient tumors,” we describe the gene expression value conversion and experimental validation. In “digital markers accurately predict drug sensitivity of cell lines and patient tumor organoids,” we perform experimental validation of the prediction of drug sensitivity using cell lines and patient tumor-derived organoids cultured in the laboratory.

Cross-validation of digital marker-based prediction of drug sensitivity on cancer cell lines and patient tumors
The transcriptional profiles and drug sensitivity of cancer cell lines and patient tumors were shown to be cancer type specific (Figures S1 and S2; Table S2). Correspondingly, we built the prediction models in a cancer type-specific manner. For each cancer type, a set of digital markers was established by computational screening of individual drugs. Figure 2A shows the comparison of the whole transcriptome and the screened digital markers in when evaluating the treatment efficacy of fluorouracil (S-FU) in colorectal cancer cell lines in the GDSC database. The
Cell lines are indicated by the yellow dots and mapped to the two-dimensional (2D) space by principal-component analysis (PCA). The background colors represent the drug sensitivity of 5-FU, rendered by KNN classification in the 2D feature space. The prediction using the digital markers aggregates the drug-sensitive and non-sensitive cell lines into two distinct groups in the feature map, whereas the whole transcriptome group (all genes) fails to prove its prediction ability. It suggests that the digital marker-based prediction can better distinguish the drug-sensitive and non-sensitive cell lines.

The ROC curves of cross-validation of prediction of drug sensitivity are shown in Figures 2B and S3–S8 and Figures 2C and S9 for cancer cell lines and patient tumors, respectively. Table 1 summarizes the evaluation scores in Figures 2B and 2C. For both cancer cell lines and patient tumors, the AUC for cross-validation reached scores above 0.9.

To test the generalizability of our model, we performed “cross-prediction” for cell lines by training the KNN model in one dataset (GDSC/CCLE) and predicting drug sensitivity in the other dataset (CCLE/GDSC). We compared the performance of three prediction methods: (1) performing gene filtering on both GDSC and CCLE and predicting drug sensitivity by 5-fold cross-validation in GDSC and CCLE separately, (2) using GDSC/CCLE as the training dataset and predicting drug sensitivity by 5-fold cross-validation in the same dataset, (3) “cross-prediction,” using GDSC/CCLE as the training dataset and predicting drug sensitivity by 5-fold cross-validation in CCLE/GDSC. Except for a few instances, the performance of the cross-prediction method was satisfactory, reaching accuracy grades similar to that of the controls (Figures 2B and S3–S8). The digital markers of different cancer type-drug pairs are finalized by method 1 and are listed in Tables S3 and S4 for cancer cell lines and patient samples, respectively. Most digital marker genes were proven essential for the progression in certain cancers. For example, in the digital markers of patient samples (Table S4), the epithelial membrane protein 2 (EMP2) occurred in patient samples of colorectal cancer in response to 5-FU or oxaliplatin administration (Table S4), representing a novel therapeutic target for endometrial cancer stem cells, while the fact that the BAG cochaperone 3 (BAG3) occurred in colorectal cancer cell lines exposed to 5-FU or oxaliplatin correlates with apoptosis in treated colon cancer cell lines and chemoresistance.

Venn diagrams of cancer genes and digital markers of cancer cell lines and tumors show both similarity and distinction between digital marker genes and cancer genes (Figures S12 and S13; Table 2). Many screened genes have oncotherapeutic implications. For example, the Fas cell surface death receptor is related to apoptosis, activating T cell killing, and resistance in tumor-immune conflict in colorectal cancer, while the BCL2-associated X, an apoptosis regulator, is key to apoptosis and p53 transcription pathways in lung cancer and also correlates with chemoresistance to cisplatin.

We also analyzed the robustness of the digital markers and the top 200 genes filtered by the 2 datasets, respectively. We compared the top 200 genes filtered by CCLE and GDSC, respectively.
Figure 2. Digital marker-based prediction of drug sensitivity on GDSC and CCLE datasets

(A) The two-dimensional PCA dimensionality reduction maps of colorectal cancer cell lines. The digital marker separates the sensitive and non-sensitive cell lines to 5-FU, whereas the complete RNA-seq data fails to separate the two classes. The color represents the drug sensitivity of 5-FU, rendered by KNN classification.

(B) ROC curves for different methods on GDSC and CCLE datasets. The AUC values are shown for each method.

(C) ROC curves for COAD+READ and STAD+ESCA for 5-FU. The AUC value is 0.947 for COAD+READ and 0.904 for STAD+ESCA.

(legend continued on next page)
individually, and confirmed the overlapped genes (Table S5). For example, there were 30 overlapped genes in the top 200 genes for lung cancer cell lines and cisplatin and 26 for lung cancer cell lines and cyclophosphamide. For other cancer type-drug pairs, the overlapped genes were fewer in number, probably due to the limited data availability in the two datasets for training. It also implied the importance of data accessibility in proceeding with computation-based precision medicine.

The feasibility of gene screening as we demonstrated was further proved by analyzing the biological representations of some screened genes. For example, trefoil factor 1 (TFF1), a small cysteine-rich protein that is frequently expressed in breast tumors under estrogen control treatment, is highly related to the migration of breast cancer cell lines; TFF3 is a predictive biomarker of endocrine response in breast cancer and is associated with breast cancer phenotypes. TFF1 and TFF3 overlapped frequently among the top 200 genes for breast cancer cell lines and 8 cancer drugs used in our study. Similarly, BCL9L plays a critical role in stem cell maintenance in epithelial homeostasis and carcinogenesis through the canonical Wnt signaling pathway in colon cancer. It is apparent that the overlapping genes were either known cancer therapeutic targets or their role is otherwise worth further exploration.

**qPCR-based extremely rapid prediction of drug sensitivity**

qRT-PCR data can be used to quantify the transcription levels of small gene libraries (e.g., containing tens to hundreds of genes) within a few hours, thus satisfying the needs of nearly instantaneous or point-of-care evaluation. To prove the concept, we first investigated the conversion of gene transcriptional values acquired by qRT-PCR (as the model testing data) to TPM (transcripts per million, see Table 2) values by RNA-seq (as the model training data). The principles of qRT-PCR and RNA-seq suggest that the $\triangle C_q$ values (the difference of quantification cycles between the target genes and the reference gene) in qRT-PCR are linearly correlated to log(TPM).

We first demonstrated this conversion on the colorectal cancer cell line SW620 by using RNA-seq and qRT-PCR data acquired in our laboratory. log(TPM) was replaced with log(TPM + 1) to avoid the error when TPM = 0 and GAPDH was set as the reference gene. Figure 3A shows a linear fit of the screened digital marker genes. Using RNA-seq data from the GDSC database resulted in a nearly identical graph (Figure 3B), demonstrating a robust conversion between qRT-PCR and RNA-seq data. The fitted parameters were then imported to the computational model with an embedded conversion formula: $\log(\text{TPM} + 1) = -0.3995 \times \triangle C_q + 5.6974$ (to convert $\triangle C_q$ to TPM).

To experimentally validate the use of digital markers, half of the cells were treated with drugs of serially increased concentrations and their viability rates to obtain the IC50 values were tested (Figures S10 and S11). The other half of the cells were lysed and used in the qRT-PCR quantification of the digital marker genes. The gene transcription values were then imported into the computational model. The predicted drug sensitivity values were compared with the experimental values for validation.

**Digital markers accurately predict drug sensitivity of cell lines and patient tumor organoids**

Seven colorectal cancer cell lines (HCT-116, HT-29, SW620, SW480, Caco-2, LoVo, and DLD-1) were cultured and administered three chemotherapy drugs, cisplatin (Figure 4A), paclitaxel (Figure 4B), and 5-FU (Figure 4C). The experimental and the predicted IC50 values presented nearly linear and positive correlations. The wrong prediction is marked by a cross in Figure 4B.

For drug effectiveness in cancer patients, clinical tracking and prognosis of cancer chemotherapy take a few years to accomplish. Therefore, to shorten the course of validation, we used patient-derived organoids to recapitulate gene expression and mimic the responses of their parental tumors. Tumor organoids have been proven to recapitulate tumor heterogeneity and personalized responses to chemotherapies with the prediction accuracy generally higher than 90%.

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Table 1. Evaluation scores of the cross-validation performance

| Drug and sample | Accuracy | Precision | Recall | Specificity | F1 weighted | AUC |
|-----------------|----------|-----------|--------|------------|-------------|-----|
| Cisplatin, colorectal cancer cell line in GDSC | 0.931 | 0.960 | 0.960 | 0.750 | 0.931 | 0.990 |
| 5-FU, colorectal cancer cell line in GDSC | 0.839 | 0.833 | 1.000 | 0.167 | 0.788 | 0.900 |
| Paclitaxel, colorectal cancer cell line in GDSC | 0.929 | 1.000 | 0.920 | 1.000 | 0.936 | 0.960 |
| 5-FU, COAD, and READ in TCGA | 0.900 | 1.000 | 0.526 | 1.000 | 0.887 | 0.839 |
| Capecitabine, STAD, and ESCA in TCGA | 0.756 | 0.737 | 0.167 | 0.970 | 0.697 | 0.904 |

This table lists the evaluation scores of cross-validation of prediction of drug sensitivity for colorectal cancer cell lines and patient tumors (the same instances as in Figures 2B and 2C). For cancer cell lines, the threshold of the Z score was $-1.0$ to distinguish sensitivity and non-sensitivity.
Table 2. The terminology table

| Terminology                        | Abbreviation | Meaning                                                                 |
|------------------------------------|--------------|-------------------------------------------------------------------------|
| Digital marker                     | –            | the genes (and their transcription quantification) screened to predict drug sensitivity for each cancer type-drug pair |
| Gene list                          | –            | the list of digital marker genes                                        |
| Gene library                       | –            | the assembly of gene lists or all drug candidates for a certain cancer type |
| Chemo-drug                         | –            | the non-targeted cancer drugs                                           |
| Cancer gene                        | –            | genes in the COSMIC Cancer Gene Census                                  |
| Gene filter                        | –            | the filters used to screen digital markers                               |
| Half maximal inhibitory concentration | IC$_{50}$   | IC$_{50}$ is a measure of antagonist drug potency in pharmacological research; IC$_{50}$ values represent the concentration of a drug that is required for 50% inhibition in vitro |
| Transcripts per million TPM         |              | A normalized quantification of transcription of RNA-seq results          |

Therefore, we used tumor organoids to approximate patient tumor gene expression and drug responses. After in vitro culture for 4–5 days, a group of tumor organoids were treated with different concentrations of 5-FU for 72 h to obtain the IC$_{50}$ values (Figure S11). Another group was lysed and used for qRT-PCR. The experimentally measured drug sensitivity, represented by the IC$_{50}$ values, and the predicted responses by the computational model, were compared.

Figure 4D shows the 2D PCA dimensionality reduction map of the training data and the experimental data. The squares and circles are the training data possessing the effective and ineffective treatment outcomes. The blue and red stars are the experimental testing data of the effective and ineffective cases. The background colors are filled by the probability estimation (the probability to be ineffective) of KNN classification in the 2D feature space, green for effectiveness and red for ineffectiveness. From left to right, the black dashed lines separate the space into "effective zone," "intermediate zone," and "ineffective zone."

From the organoids derived from ten patient tumors we observed that 5-FU administration proved effective (IC$_{50} < 100$ µM) in four cases and ineffective (IC$_{50} > 100$ µM) in six cases. Figure 4E shows the experimental and predicted values for the ten tumor samples. The prediction probability was positively correlated with the experimentally measured IC$_{50}$ values (in log form), except for two false-positive predictions (Figures 4D and 4E); the AUC score of the experimental validation reached 0.88 (Figure 4F).

**DISCUSSION**

In conclusion, we have proposed and validated a multiplex gene-based digital marker system to realize extremely rapid evaluation of drug sensitivity. For potential clinical practice, we have suggested a pipeline of standard uses involving tumor sampling, RNA extraction, qRT-PCR (or potentially gene chip) quantification, and intelligent readout (i.e., drug recommendation).

Based on our model validation on organoids derived from patient tumors, we propose a clinical application pipeline for extremely rapid and personalized drug recommendation (Figure 5). We will screen gene libraries, establish digital markers for all the common cancers and their clinical therapeutic drugs, and construct a user-friendly algorithm. When a patient is hospitalized, the doctor should obtain a sample of the tumor (e.g., via surgery, gastrointestinal endoscopy, or biopsy) to extract the transcriptomic materials. The transcriptional profile of the predictive digital marker should be then quantified using qRT-PCR and input to the prediction algorithm to be processed. The algorithm then outputs the predicted drug recommendation. The whole process can be completed within 1–3 h post sampling, satisfying the clinical need for immediate administration of chemotherapy following hospitalization. The process can be further improved by developing customized multiplex gene microfluidic chips that can quickly quantify transcriptomics in one small and portable device.

By using our customized computational model comprised of gene filters and a KNN algorithm, we have demonstrated the construction of small gene libraries containing digital markers that can best predict drug sensitivity. For a given cancer type, a small group (e.g., 10–20) of genes can predict a patient’s response to a common chemotherapy drug; a larger library (e.g., 50–200) of genes can predict the effects of several common drugs used in clinical practice to a specific cancer type. Therefore, it may be feasible to establish gene libraries that cover most clinical chemotherapy drugs and use them to construct cancer type-specific digital markers.

Our model screened many genes possessing important drug predictive abilities (Tables S6 and S7). Most of these digital marker genes are not yet included in the recognized cancer gene list but have been repeatedly reported in the literature because of their importance in cancer progress or therapy. For example, the screened genes, such as EMP2 and BAG3, have become prevalent targets for cancer therapeutic studies, which have been discussed in the Results, but they were not included in the current cancer gene list. The double check from both molecular biology study and computational study may guide the exploration of unspecific genes as targets for drug development and personalized therapy.

Unlike traditional biomarkers that have defined biological functions, (e.g., proteins, mRNA, or exosomes) digital biomarkers do not have a defined biological significance. However, experimental and computational evaluation support their powerful predictive capacity for applications such as drug sensitivity. Further studies on large datasets may allow us to look into the black box. The strategy of establishing digital virtual markers may also reduce the cost and shorten the time course of new biomarker development.

Limited data accessibility, high diversity, and lack of standardization in biomedicine are common issues for computational-based diagnosis and prognosis. Although this panorama is quickly changing, learning how to extract useful information from “small” data remains a valuable goal to achieve in the...
This study aims to shorten the time from patient tumor sampling to the start of administration of an effective treatment. We therefore adopted a computational-based strategy to reduce the library of genes to construct the predictive digital markers. Other reported state-of-the-art methods can predict drug sensitivity using RNA-seq data or the whole transcriptome as the input for the algorithm. However, this results in a longer time before obtaining a prediction. Therefore, the direct comparison of this study with those different algorithms was not performed, as it can be misleading. The main focus of our method was on providing rapid recommendations. Nonetheless, our method provides a delightful solution toward extremely rapid evaluation of personalized drug sensitivity, which might inspire more computational work to serve this purpose with improved robustness and accuracy.

**EXPERIMENTAL PROCEDURES**

**Resource availability**

**Lead contact**
Further information and requests should be directed to and will be fulfilled by the lead contact, Shaohua Ma (ma.shaohua@sz.tsinghua.edu.cn).

**Materials availability**
This study did not generate new unique reagents.

**Data and code availability**
Data and python codes of prediction of drug sensitivity and the graphic interface are available from the GitHub repository at [https://github.com/qftan77/cancer_digital_marker.git](https://github.com/qftan77/cancer_digital_marker.git).

Any additional information required to reanalyze the data reported in this paper is available from the lead contact upon request.

**Tumor sampling**
Patient tissues for tumor fabrication of organoids were collected under the approval of The First Affiliated Hospital of Zhengzhou University Ethics Committee and The Affiliated Cancer Hospital & Institute of Guangzhou Medical University Ethics Committee, following both national and local regulations. Tissues were stored in “ready-to-use” solution (Aqix, UK) with 100 μg/mL primocin (InvivoGen, USA) and transported to the laboratory under incubation on ice.

**Culture of cell lines**
The colorectal cancer cell lines presented in this study, including HCT-116, HT-29, SW620, SW480, Caco-2, LoVo, and DLD-1, were provided by the Peter E. Lobie Group at the Tsinghua Shenzhen International Graduate School (SIGS). Cell lines SW620 and SW480 were grown in DMEM medium supplemented with 10% FBS (v/v) (Gibco, USA) and 1% penicillin/streptavidin (v/v) (Gibco). DLD-1 was grown in RPMI medium with 10% FBS and 1% penicillin/streptavidin. HCT-116 and HT-29 were grown in McCoy’s 5A medium with 10% FBS and 1% penicillin/streptavidin. LoVo was grown in DMEM/F12 medium with 10% FBS and 1% penicillin/streptavidin. All cell lines were maintained at 37°C in a humidified incubator at 5% CO₂.

**Tumor dissociation and organoid culture**
Tumor tissues were washed with cold 1× PBS with 5% penicillin/streptavidin (v/v) (Gibco) and cut into sub-millimeter sized small pieces. Tissues were then digested by 1 mg/mL collagenase IV (Gibco), 0.1 mg/mL DNase I (Sigma-Aldrich, USA), 0.1 mg/mL dispase type II (Sigma-Aldrich), 100 μg/mL primocin (InvivoGen) and 1% FBS (v/v) in DMEM/F12 medium on an orbital shaker at 37°C, incubated for 20–40 min. After tissue digestion, dissociated cells, cell clusters, and microtissues were passed through a 100-μm cell strainer (Corning, USA). Then the dissociated cells and cell clusters were resuspended in DMEM/F12 and centrifuged at 1,000 rpm for 5 min.

The cells dissociated from tumor tissues were suspended in Matrigel (Corning) at a density of 10⁶ cells/mL. Organoids were formulated by following the reported protocol developed by Jiang et al. Organoids were cultured in medium containing 20% FBS (v/v), 1% penicillin/streptavidin, 1X Glutamax.

![Figure 3. Comparison between ΔCq and TPM for gene transcription quantification](image-url)

The fitted linear relationship between log(TPM + 1), derived from RNA-seq of the cancer cell line SW620 from (A) laboratory experiments and (B) the GDSC dataset, and ΔCq measured by laboratory qRT-PCR. The reference gene for qRT-PCR was GAPDH.
supplement (Thermo Fisher Scientific, USA), 1× B-27 supplement (Thermo Fisher Scientific), 1.25 mM N-acetylcysteine (Sigma-Aldrich), 100 ng/mL noggin (MedChemExpress [MCE], USA), 100 mg/ml R-spondin 1 (MCE), 5 ng/mL EGF (R&D, USA), 5 mM nicotinamide (Sigma-Aldrich), 5 μM Y-27632 (R&D), 100 μg/mL primocin (InvivoGen) and 1× FibrOut (CHI Scientific, USA) in DMEM/F12 medium. The medium was changed every 2 days.

Use of datasets GDSC, CCLE, and TCGA
We obtained the transcriptional and drug sensitivity (or prognosis) data from the public datasets. The GDSC data were downloaded from https://www.cancerrxgene.org/. We used data from GDSC2, a newer version of GDSC containing 899 cell lines and 198 compounds, in this study. The CCLE data were downloaded from https://portals.broadinstitute.org/ccle. TCGA data were downloaded from https://portal.gdc.cancer.gov/.

Gene filters
The gene filters were slightly different for cell lines and patient samples, but the essential idea was the same: the genes were first filtered by their transcriptional level, and then by their correlation level, to drug sensitivity. In all the filtering steps, TPM values across all samples were considered to calculate the mean value, Pearson correlation coefficient, or the dispersion scores.

For gene filters applied to cell lines
First, genes with mean expression level log(1 + TPM) > 1 (across all samples) were retained. After this step, about one-third of the genes remained (10,000–20,000 genes, the number was different for different cancer type-drug pairs) (Figure 1B). Second, genes with a Pearson correlation coefficient (between TPM and Z scores of IC50) > 0.25 were retained. After this step, about 3,000 genes were retained. Third, genes were scored and rated by the absolute values of the Fisher’s linear discriminant of the gene transcription magnitudes between cell lines with the highest and lowest 15% of the IC50 values. Genes were enumerated in the subsequent KNN algorithm by order. KNN algorithm cross-validation determined the top m (m is the number of genes, m ≤ 50) genes to be selected as digital markers to reach the best performance.
Gene selection and KNN enumeration

After applying the gene filters, the remaining genes were scored and ranked according to their importance for prediction of drug sensitivity. The KNN algorithm was used to finalize the gene list for each drug. The top $m$ ($m = 1$ to 30) genes were enumerated in the subsequent KNN 5-fold cross-validation with order. For each $m$, the parameter $k$ of KNN was also enumerated to reach the best cross-validation performance. Finally, the composition of the digital markers and parameter $k_0$ were determined with the desirable AUC score in the 5-fold cross-validation process.

Gene selection was performed for each cancer type-drug pairs in most of the situations. For cancer cell lines, to finalize the digital markers, we performed gene filtering on both datasets separately and synthesized the scores of genes as the rating criterion to finalize the gene candidate list. Then we performed KNN cross-validation on CCLE/GDSC datasets according to the gene candidate list to determine the digital markers for a certain cancer type-drug pair. To test the robustness of the digital markers, we also screened digital markers separately on CCLE and GDSC.

The wet laboratory validation adopted a different procedure for gene selection. For experimental validation of prediction of drug sensitivity, patients were divided into two groups for each cancer type-drug pair. Mann-Whitney U test was performed on the TPM of each gene across the two groups of patients. Scored and ranked genes were enumerated in the subsequent KNN algorithm with order. KNN algorithm cross-validation determined the top $m$ ($m$ is the number of genes, $m \leq 30$) genes to be selected as digital markers to reach the best performance (Figures 1C and S14).

For gene filters applied to patient samples

First, genes with mean expression level log(TPM) > 0 (across all samples) were retained. After this step, about 10,000–20,000 genes were retained (the number was different for different cancer type-drug pairs) (Figure 1C). Second, genes were scored and ranked by the p values from the Mann-Whitney U test. According to the drug sensitivity label, patients were divided into two groups for each cancer type-drug pair. Mann-Whitney U test was performed on the TPM of each gene across the two groups of patients. Scored and ranked genes were enumerated in the subsequent KNN algorithm with order. KNN algorithm cross-validation determined the top $m$ ($m$ is the number of genes, $m \leq 30$) genes to be selected as digital markers to reach the best performance (Figures 1C and S14).

**Prediction of drug sensitivity**

We performed the gene filtering process to screen gene candidates (which will feed into the KNN model to generate the final digital markers) for each cancer type-drug pair, which means that, for each cancer type-drug pair, we established a set of digital markers. In total, we studied three cancer types and eight drugs for cancer cell lines, and six cancer type-drug pairs for patient samples, listed in Table S2.
analysis. Total RNA of both cell lines and tumor organoids were extracted using the Eastep Super Total RNA Extraction Kit (Promega)/RNeasy Mini Kit (- QIAGEN, Germany), following the manufacturer’s instructions. The purity and concentration of RNA were determined by using a NanoDrop 2000 spectrophotometer (Thermo Fisher Scientific). RNA (1–2 μg) were used to prepare cDNA using the PrimeScript RT Reagent Kit (TaKaRa, Japan) with random primers and oligo dT primers.

**qRT-PCR assay**

qRT-PCR were performed in 96-well plates (Sangon, China) on a Real-Time PCR Detection system (Bio-Rad, USA). Primer sequences and characteristics are described in Tables S8 and S9. The qRT-PCR reaction mix was composed of 12.5 μL GoTaq Green Master Mix, 2X (Promega), 10 μM upstream primer, 10 μM downstream primer, 2 μL DNA template, and 25 μL nuclease-free water. Cycling reaction conditions were 95°C for 10 min, 40 cycles of 95°C for 15 s, and 60°C for 1 min. All qRT-PCR experiments were performed with three biological replicates. All qRT-PCR data are shown in the Figures S15 and S16.

**SUPPLEMENTAL INFORMATION**

Supplemental information can be found online at https://doi.org/10.1016/j.patter.2021.100360.

**ACKNOWLEDGMENTS**

The work was supported by the National Natural Science Foundation of China (grant no. 61971255), the National Science Foundation of Guangdong Province (grant no. 2021B1515020092), the Shenzhen Science and Technology Innovation Commission (grant nos. RCYX20200714114736146 and WDZC20200821114349001), and the Shenzhen Bay Laboratory Fund (grant no. SZBL2020090501014).

**AUTHOR CONTRIBUTIONS**

J.Y. and Y.F. contributed equally to this study. J.F. contributed to the algorithms and codes, drug screening, and manuscript drafting. Y.F. contributed to tumor organoid sampling and gene transcription quantification. Y.C. provided help on figure editing and paper revision. Z.W. and H.Z. provided guidance on biological experiments. E.A.G. provided help on text editing. Q.L., S.C., and W.Z. provided patient samples. S.M. conceived and advised on this work. All authors edited and approved the final version of the manuscript.

**DECLARATION OF INTERESTS**

The authors declare no competing interests.

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