A 41-Gene Pair Signature for Predicting the Pathological Response of Locally Advanced Rectal Cancer to Neoadjuvant Chemoradiation

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Background and Purpose: Pathological response status is a standard reference for the early evaluation of the effect of neoadjuvant chemoradiation (nCRT) on locally advanced rectal cancer (LARC) patients. Various patients respond differently to nCRT, but identifying the pathological response of LARC to nCRT remains a challenge. Therefore, we aimed to identify a signature that can predict the response of LARC to nCRT.

Material and Methods: The gene expression profiles of 111 LARC patients receiving fluorouracil-based nCRT were used to obtain gene pairs with within-sample relative expression orderings related to pathological response. These reversal gene pairs were ranked according to the mean decrease Gini index provided by the random forest algorithm to obtain the signature. This signature was verified in two public cohorts of 46 and 42 samples, and a cohort of 33 samples measured at our laboratory. In addition, the signature was used to predict disease-free survival benefits in a series of colorectal cancer datasets.

Results: A 41-gene pair signature (41-GPS) was identified in the training cohort with an accuracy of 84.68% and an area under the receiver operating characteristic curve (AUC) of 0.94. In the two public test cohorts, the accuracy was 93.37 and 73.81%, with AUCs of 0.97 and 0.86, respectively. In our dataset, the AUC was 0.80. The results of the survival analysis show that 41-GPS plays an effective role in identifying patients who will respond to nCRT and have a better prognosis.

Conclusion: The signature consisting of 41 gene pairs can robustly predict the clinical pathological response of LARC patients to nCRT.

Keywords: rectal cancer, neoadjuvant chemoradiation, relative expression orderings, signature, survival
INTRODUCTION

Neoadjuvant chemoradiation (nCRT) followed by surgery is an effective treatment for locally advanced rectal cancer (LARC) (1). This treatment is considered a safe and effective strategy for locally advanced colon cancer (2). The pathological response to nCRT determines surgical plan and post-operative quality of life of LARC patients (3). Pathological responders can benefit from nCRT, whereas non-pathological responders can choose not to undergo nCRT to avoid the risk and pain associated with this type of treatment. Nevertheless, among patients with LARC, the proportion of patients with pathological complete response and partial pathological response is only ~10–25% and 40%, respectively (4). Therefore, predicting the nCRT response before treatment may substantially improve the choice of patients for pre-operative chemotherapy.

Several prediction models based on tumor tissue expression profiles have shown high accuracy on their respective datasets (5–12), but the high variability and batch effects make it difficult to apply these predictive models to independent data (13). In addition, the standardization process of adjusting batch effects in gene expression profiling requires the collection of a certain number of samples, which delays the subsequent treatment of patients in clinical practice (14). The accuracy of model prediction will also be affected by sample normalization (15). Our previous evidence showed that the within-sample relative expression orderings (REOs) of gene pairs (16) can robustly resist batch effects, and we have successfully screened gene pairs with marked differences between responders and non-responders of LARC (17). However, the selection of gene pairs is still a problem worthy of discussion, as the accuracy varies among predictive signatures composed of different gene pairs. The random forest algorithm (18) can effectively analyze high-dimensional data (the number of variables is more than a hundred times the number of observations), and at the same time provide a variable importance measure (VIM), which makes random forest especially suitable for the study of expression profile to identify potential biomarkers (19). Compared to our previous research, we can measure how important a gene is by the size of the VIM value, rather than saying that the gene is useful or useless. This can prevent the screening of genes that have a classification effect.

This study aimed to screen gene pairs using the VIM values of random forest, thereby exploring the gene pair signature (GPS) in predicting the response of LARC to nCRT.

MATERIALS AND METHODS

Data and Pre-processing

A total of 232 rectal cancer patients were included in this study. They were from three public cohorts of LARC, namely GSE87211 (20), GSE35452 (21), and GSE45404 (22), and a cohort measured at our laboratory. The surgical plan for all patients was neoadjuvant fluorouracil (5-FU)-based chemoradiation combined with 50.4 Gy radiotherapy.

In GSE87211, 363 samples were measured using the Agilent-026652 whole human genome microarray 4 × 44K v2 (GPL13497) platform, including 203 patients with rectal cancer. We screened 111 patients who received nCRT and provided pathological response information. The pathological response of these patients was determined according to the American Joint Committee on Cancer tumor regression grade (TRG) (23).

The validation cohorts GSE35452 and GSE45404 were measured using the Affymetrix Human Genome U133 Plus 2.0 Array (GPL570) platform. These cohorts comprised 46 and 42 rectal cancer patients who received nCRT, respectively. The response status of these patients was assessed using the Mandard TRG system. Although the classification standards of the Mandard and AJCC TRG systems are slightly different, the difference between them is negligible (23). The last validation cohort with 33 LARC patients was from Fujian Medical University Union Hospital (17).

Because the pathological response status has a positive effect on disease-free survival (DFS) (24), the 41-GPS should be used to predict the survival benefit of 5-FU-based nCRT. First, we used the signature to evaluate the survival benefit of 105 patients with rectal cancer treated with 5-FU-based nCRT in the GSE87211 cohort. To further verify our signature, we used the 41-GPS to predict the survival benefits of 285 patients with colorectal cancer. The information of these patients was in reference (17). Because rectal and colon cancers are almost genetically indistinguishable (25), we expected that the 41-GPS would also be applicable to patients with colon cancer. We also collected 158 The Cancer Genome Atlas Rectum Adenocarcinoma (TCGA-READ) to predict DFS benefits.

In addition, nine colorectal cancer (CRC) cell lines in GSE20298 (26) were used in this study, including Caco-2, LS513, LS1034, SW403, SW480, SW620, SW837, SW1116, and SW1463. We used these to evaluate the resistance of the cell lines, which is measured by the survival rate of tumor cells under 3 μM/L 5-FU and 2 Gy X-ray radiation exposure.

TCGA cohorts were obtained from the Genomic Data Commons Data Portal, whereas others were obtained from the Gene Expression Omnibus repository (27). All data are standardized by the Robust Multi-Array Average algorithm (28) and are shown in Table 1. Data normalization of the inter-sample was not performed. When multiple probes were mapped to a gene symbol, the arithmetic mean of the probe values was used (on the log 2 scale).

Procedure of Discovering Signature

The signature discovery process is illustrated in Figure 1. First, according to the Tumor Regression Classification (TRG) of the American Joint Cancer Council of the United States, patients in the training group were divided into two groups, namely the response and the non-response group. Then the REO algorithm was used to obtain the reversal gene pairs of the training set (Binomial test, \( p < 0.05 \)). The reversal gene pair includes two genes, gene \( a \) and \( b \), and the relationship was recorded as \( G_a > G_b \) or \( G_a < G_b \), where \( G_a \) and \( G_b \) represent expression values. The reversal gene pairs were then inputted into the random forest algorithm to obtain the VIM ranking according to the mean decrease Gini (MDG) index. Finally, the best top-N gene pairs were selected as the signature, where best top-N is the top N gene...
TABLE 1 | Datasets analyzed in this study.

| GEO ID     | Describe         | Platform  | Sample size | Tissue sample type |
|------------|------------------|-----------|-------------|--------------------|
| GSE87211   | Expression profile | GPL13497  | 111         | Rectal cancer      |
| GSE35452   | Expression profile | GPL570    | 46          | Rectal cancer      |
| GSE45404   | Expression profile | GPL570    | 42          | Rectal cancer      |
| GSE39582   | Expression profile | GPL570    | 200         | Colorectal cancer  |
| GSE14333   | Expression profile | GPL570    | 85          | Colorectal cancer  |
| TCGA       | Expression profile | -         | 158         | READ               |
| GSE20298   | Cell lines       | GPL4133   | -           | Colorectal cancer  |
| Our dataset | Expression profile | GPL15207  | 33          | Rectal cancer      |

FIGURE 1 | Flowchart of signature discovery process and validation in this study.

pairs having the highest classification accuracy with the majority voting rule in the training dataset.

Statistical Analysis

The open-source statistical analysis software R4.0.2, with R packages randomForest, rms, limma, and survival, was used for statistical analysis. The reversal gene pair obtained using the REO algorithm was evaluated using a Binomial test ($p < 0.05$). The online software DAVID Bioinformatics Resources 6.8 was used for gene ontology (GO) enrichment analysis. Term with a $P$-value $< 0.05$ was considered statistically significant. The R package limma is used to identify differentially expressed genes. The accuracy of the signature is defined as the ratio of the correctly predicted samples to all samples in the cohort of responders and non-responders, and the 95% confidence interval was calculated by the Clopper-Pearson method ($29$). The parameters of the best random forest model in the seed 20200914 were “best mtry” $= 88$ and “ntree” $= 2,500$. Receiver operating characteristic (ROC) curve and area under the curve (AUC) were used to assess the predictive ability of the signature. The Kaplan–Meier method and log-rank test were used to evaluate the survival analysis of the training dataset and TCGA dataset. DFS was defined as the time from surgery to relapse or the date of final recording ($24$). In all statistical analyses, $P < 0.05$ was considered statistically significant.

RESULTS

Establishing the 41 Gene Pair Signature for Response Prediction in LARC

The discovery workflow is shown in Figure 1. Using GSE87211 as training data and REO algorithm, we identified 13,388 reversal gene pairs that were significantly related to the pathological
response status of LARC patients receiving nCRT (Binomial test, \(p < 0.05\)). We then introduced all reversal gene pairs into the random forest model to obtain the VIM value. The

| Gene pairs | Gene pairs | Gene pairs |
|------------|------------|------------|
| \(G_a > G_b^*\) | \(G_a > G_b\) | \(G_a > G_b\) |
| 1. ABCC5 > FUT8 | 15. USP19 > HENMT1 | 29. TUBB2A > NENF |
| 2. EFNA1 > CDKN3 | 16. ECSIT > COA8 | 30. PRKCZ > TIMM9 |
| 3. AGRN > UBE2L6 | 17. AMBR A1 > SOCS4 | 31. BPI3N1 > TMEM39B |
| 4. SLC17A9 > SUP77L | 18. GLYCTK > ERP27 | 32. ASPHD1 > THAP1 |
| 5. EPS8L2 > PARN | 19. EB1 > ARHGEF39 | 33. SLC17A9 > APOL2 |
| 6. ITGA3 > MIES1B5P1 | 20. MIER3 > GSTT1 | 34. DHFR2 > LPCAT2 |
| 7. ZP1 > LTA | 21. MICAL1 > ZNF92 | 35. KMT2B > A/JUBA |
| 8. CRV2 > ROHOB7B1 | 22. ZNF460 > WDR66 | 36. LHFP12 > COA8 |
| 9. SEST21 > CCDC25 | 23. PREDL2 > PRMT5 | 37. SF1 > DCP1B |
| 10. GT1 > WASHC3 | 24. DCAF1 > TBC1D5 | 38. DCTPP1 > ENY2 |
| 11. NPHP4 > WDCP | 25. PRR1 > PPP1R2 | 39. HAGHL > KLHDC9 |
| 12. DHFR2 > NME7 | 26. STRP1 > KIN | 40. CRTC1 > GSTT1 |
| 13. EHBP1L1 > LPCAT2 | 27. GLYAT > DKK4 | 41. ASPHD1 > SMUG1 |
| 14. NWS1 > TSEN15 | 28. FZD9 > CYCL11 |

*\(G_a > G_b\) represents the response to neoadjuvant chemoradiation in locally advanced rectal cancer.

The 41-GPS contains 76 genes, nearly half of which (37 of 76) were differentially expressed between the responder and non-responder cohorts (\(t\)-test, \(p < 0.05\)) (Figure 3A). The overlapping genes in the signature were ASPHD1, COA8, DHFR2, GSTT1, LPCAT2, and SLC17A9. GO enrichment analysis revealed that 57.89% of the signature genes were enriched in the protein binding of molecular function pathway (\(p < 0.05\); Figure 3B).

**The 41-GPS Is a Strong Predictor in Independent Cohorts of LARC**

In the first verification cohort (GSE35452), the signature predicted nCRT responders with a sensitivity and specificity index of 95.83% (23/24) and 90.90% (20/22), respectively, and an accuracy of 93.37% (95% CI, 89.84–97.12%). In the second validation cohort (GSE45404), the sensitivity and specificity of the prediction of nCRT responders were 89.47% (17/19) and

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**FIGURE 2** Distribution of response gene pairs in patients of GSE87211 under 41-GPS prediction.

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**TABLE 2** The composition of 41-GPS.
FIGURE 3 | Composition and differentially expressed genes of 41-GPS. (A) Gene enrichment distribution map. The picture shows the six most significant enrichment pathways ($p < 0.05$). Dots represent the gene symbol, red denotes upregulated gene, and blue denotes downregulated genes. The color of the enrichment pathway indicates the size of the z-score. (B) The specific distribution of gene enrichment. Different colors represent different access pathways.
60.87% (14/23), respectively, and the accuracy was 73.81% (95% CI, 67.03–80.59%). The AUC of the two validation cohorts was 0.972 (95% CI, 0.929–1) and 0.849 (95% CI, 0.722–0.976) in that order (Figures 4B,C).

In our dataset, we observed that the prediction of nCRT responders had a sensitivity and specificity of 83.33% (10/12) and 61.90% (13/21), respectively, with an accuracy of 69.70% (95% CI, 64.53–74.87%). The AUC was 0.800 (95% CI: 0.647–0.952; Figure 4D).

The 41-GPS Predicts Survival and Treatment Response in LARC and Colorectal Cancer

We conducted a survival analysis on the DFS data of 105 patients in the training cohort, and tested whether 41-GPS could predict the benefit of post-operative chemotherapy of LARC. Among the 105 samples, 58 were predicted to be responders and 47 non-responders. After adjusting for stage, age, and
FIGURE 5 | Independence verification and survival analysis verification of 41-GPS. Stage*: The status of lymph node metastasis before surgery, 0 means no metastasis, 1 means metastasis. Stage+: The stage of colon cancer. 41-GPS+: In multivariate analysis of GSE87211, patients were divided into responders and non-responders. In multivariate analysis of GSE39582 and GSE14333, patients were divided into high-response groups and low-response groups according to the number of responding gene pairs. (A) Univariate analysis of GSE87211, GSE39582 and GSE14333. Hazard ratio and $P$-values were calculated using log-rank test. (B) Nomogram models of GSE87211. (C) Disease-free survival rate of 105 locally advanced rectal cancer patients in GSE87211. (D) Disease-free survival rate of 156 rectal cancer patients in TCGA-READ. (E) Disease-free survival rate of 200 colorectal cancer patients in GSE39582. (F) Disease-free survival rate of 85 colorectal cancer patients in GSE14333.
The 41-GPS Reflects the Drug Resistance of Cell Lines

We compared the chemo radiation resistance of nine colorectal cancer cell lines, of which SW403 and SW837 were the most resistant (Figure 6A). To test whether 41-GPS prediction would reflect the resistance of cell lines, we divided the patients again into sensitive and resistant groups based on the median resistance to chemo radiation and compared the number of response gene pairs between the two groups. We then used the 41-GPS to predict each cell line and calculated the proportion of response gene pairs. The results showed that the average response ratio of the resistant group was 0.459, which was lower than that of the sensitive group (0.569; T-test, \( p < 0.05 \); Figure 6B). These findings confirm the validity of our signature.

DISCUSSION

The identification of transcriptional predictive signatures of LARC response to nCRT would provide an additional reference for selecting patients most likely to benefit from therapy. In this study, we identified a predictive signature for the LARC response to nCRT based on REO and random forest algorithms. In previous studies, the REO algorithm showed resistance to experimental batch effects (7). Based on this, a more efficient and robust strategy (random forest) was used to identify our signature, which solved the limitations of our previous method regarding the selection of starting features (17). This signature performed well in predicting the pathological response and long-term survival prognosis of LARC patients receiving nCRT, and its prediction effect was also highly correlated with the resistance of colorectal cancer cell lines. Our experimental results showed the superiority of the random forest algorithm in omics data and the robustness of the 41-GPS. We believe that the 41-GPS could...
strongly predict nCRT responders prior to treatment. It is worth mentioning that cohort GSE87211 in our study contained both classification labels and survival data. This is direct evidence of the effectiveness of the 41-GPS. To the best of our knowledge, this is the first study to apply such data to the verification of rectal cancer markers.

This study, however, is subject to limitations. When the measurement scale or the number of categories of the predictor variables are different, the results of the random forest will be biased; however, this is inevitable (32). In the near future, more robust random forest models should be explored in large cohorts to define the importance metric influencing accuracy.

In conclusion, this study identified a signature of 41 gene pairs that can predict the nCRT response of LARC. This signature may be useful for individual clinical applications. It may help clinicians avoid risks for patients who will not benefit from F5-FU-based nCRT 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 in the article/Supplementary Material.

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**AUTHOR CONTRIBUTIONS**

ZX and YG: conceptualization. YG: funding acquisition and project administration. ZX, SY, and YG: methodology. YL, MH, YD, LL, and WP: supervision. ZX: writing—original draft. ZX, SY, HC, GH, and YG: writing—review and editing. All authors contributed to the article and approved the submitted version.

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**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmed.2021.744295/full#supplementary-material
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