Bioinformatics evaluation of a novel angiogenesis related genes-based signature for predicting prognosis and therapeutic efficacy in patients with gastric cancer

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Received March 17, 2022; Accepted May 23, 2022; Epub July 15, 2022; Published July 30, 2022

Abstract: Objective: Tumor angiogenesis plays a pivotal role in the development and metastasis of tumors. This study aimed to elucidate the association between angiogenesis-related genes (ARGs) and the prognosis of patients with gastric cancer (GC). Methods: Transcriptomics and clinical data of GC samples were obtained from The Cancer Genome Atlas (TCGA) as the training group and those from Gene Expression Omnibus (GEO, including GSE26253, GSE26091 and GSE66229) as the validation groups. Single-sample gene set enrichment analysis (ssGSEA) was performed for gene set enrichment analysis on the gene set of angiogenesis and divided patients into high- or low-ARG group. Subsequently, to improve the availability of the ARG signature, a ARGs subtype predictor was then constructed by integrating of four machine learning methods, including support vector machine (SVM), least absolute shrinkage and selection operator (LASSO) regression, Random Forest and Boruta (RFB) and extreme gradient boosting (XGBoost). Kaplan-Meier and receiver operating characteristic curves were used to evaluate the performance of prognosis prediction. The EPIC and xCELL method were used to calculate the profile of tumor-infiltrated immune cells. Results: The expression levels of a total of 36 ARGs that correlated with the survival of patients with GC were identified and utilized to establish an ARG-related prognosis signature. The area under the curve for predicting overall survival (OS) in the training group at the 1-, 3- and 5-year was 0.61, 0.64 and 0.76, respectively, and this was further validated using three independent GEO datasets. Moreover, the ARG signatures were significantly correlated with cancer-associated fibroblasts (CAFs), and GC patients that exhibited both high ARG expression level and matrix CAFs level had the most inferior outcomes. The multiple machine learning algorithms were applied to establish a 10-gene ARG subtype predictor, and notably, a high ARG-subtype predictor score was associated with reduced efficacy of immunotherapy, and potential anti-HER2 or FGFR4 therapy, but an increased sensitivity to anti-angiogenesis-related therapy. Conclusion: The novel ARGs-based classification may act as a potential prognostic predictor for GC and be used as a guidance for clinicians in selecting potential responders for immunotherapy and targeted therapy.

Keywords: Angiogenesis, gastric cancer, TCGA, cancer-associated fibroblasts, immune cell infiltration, immunotherapy

Introduction

Gastric cancer (GC) is the fifth most common malignancy and the third most common cause of cancer-related death worldwide [1]. Despite of ongoing advances in treatment strategies, patient outcomes remain poor [2]. GC is a highly heterogeneous disease with differing features, including locations, histological types, molecular classifications and biological behaviors [3]. However, primary tumors with biological heterogeneity are often missed by conventional diagnostic methods, which are often based on TNM staging [4]. Therefore, a novel
risk model for the identification of patients with a high risk of developing GC is required to provide a guideline for personalized medicine in patients with GC.

Angiogenesis is defined as the process in which new capillaries grow from pre-existing vessels, and has been characterized as a vital factor for the proliferation, growth and metastasis of tumor cells in multiple solid tumors [5]. It is considered as one of the hallmarks of cancer, attributing to cancer cell growth by providing oxygen and nutrients, delivering molecules that confer immune and treatment resistance, and facilitating metastasis [6]. Results of previous studies have revealed that a wide variety of genes play crucial roles in angiogenesis, such as vascular endothelial growth factors (VEGFs) and their receptors (VEGFRs), which have been previously established in angiogenic signaling pathways [7].

Results of a previous study revealed that active angiogenesis may result in therapeutic failure and poor outcomes in patients with GC [8]. Meanwhile, Feng et al. demonstrated that angiogenesis signaling was significantly associated with the cytotoxic function in patients with GC with a highly infiltrated immune niche [9]. However, the clinical relevance of angiogenesis-related genes (ARGs) in GC is yet to be fully elucidated.

Cancer-associated fibroblasts (CAFs) are the predominant component of the tumor microenvironment (TME), and are involved in critical processes within the TME, including extracellular matrix (ECM) remodeling, reciprocal signaling interactions with cancer cells and crosstalk with infiltrating inflammatory cells [10]. Accumulating evidence indicates that CAFs play crucial roles in the growth, invasion and progression of tumors, and are associated with a poor prognosis in various solid tumors [11-13]. In addition, previous studies have revealed that angiogenesis played a pivotal role in the CAF promotion of invasion and metastasis of tumors [14]. However, the interaction between CAF and angiogenesis, and the outcomes of patients with GC have not been fully elucidated.

Thus, in this study, we sought to evaluate the comprehensive role of ARGs in the prognosis of GC and then develop a robust signature to predict GC patients’ survival based on the ARG related feature using the transcriptomics data from The Cancer Genome Atlas (TCGA) and Gene Expression Omnibus (GEO). Special interests were given to analyze correlation between the ARG signatures and TME in GC. Subsequently, an ARG subtype predictor was established to distinguish the two subtypes in patients with GC, giving stratification not only in survival but also in systemic treatments including immunotherapy.

**Methods and materials**

Aquisition of gastric cancer datasets and selection of angiogenesis-related genes

The RNA-seq (RNA-sequencing) datasets and clinical features of the TCGA-GC cohort from the TCGA database (http://xena.ucsc.edu/) were downloaded as the training group. The following criteria were applied for validation datasets selection in the Gene Expression Omnibus (GEO, https://www.ncbi.nlm.nih.gov/geo/) database: (i) tumor tissue samples were from gastric cancer; (ii) the dataset must have a complete record of the overall survival data; (iii) the expression data must be generated using a genome-wide gene expression method; (iv) the sample size for analysis should be over 100. Then, GSE26253 (containing 432 GC samples), GSE26901 (109 GC samples) and GSE66229 (185 GC samples) were obtained from the GEO as the validation groups.

The Angiogenesis-related genes, a total of 36 genes, were downloaded from Molecular Signatures Database (http://www.gsea-msigdb.org/gsea/msigdb/, MSigDB-Hallmark version 7.4), as the ARGs gene set.

The construction of the ARGs signature based on single-sample gene set enrichment analysis (ssGSEA)

The enrichment score and the absolute enrichment degree of a gene set in each sample were calculated using ssGSEA by an empirical cumulative distribution function according to the given ARGs expression profile. The Gene Set Variation Analysis (GSVA) package was used to calculate the ARGs ssGSEA score of 414 samples from TCGA-STAD. The limma package in R was applied to assess the scores in different tissue types and survival statuses. GSEA (version 4.1.0.24) was performed for gene set enrichment analysis (GSEA) on the gene set (angiogenesis related genes) with 1000 permu-
tations using gene expression profiles of 414 tumor samples. The patients were divided into high and low ARG score (ssGSEA score) groups for further analysis.

**Tumor-infiltrated immune cell analysis**

We firstly evaluated correlation of CYT, ESTIMATE, HLA, ICG, TIL, tumor purity and ARG score by Spearman’s analysis. The EPIC and xCELL methods were used to calculate the infiltration abundance of immune cells in both TCGA and GEO datasets according to the “EPIC and xCELL” package including eight immune cells in EPIC package (version 1.1.5) and thirty-six immune cells in the XCELL package (https://xcell.ucsf.edu/), respectively.

**Construction and validation of the ARGs subtype predictor by multiple machine learning methods**

The 414 GC patients were classified into training (N=331) and testing (N=83) sets based on a ratio of 4:1. First, in the training set, the most important group-relevant features were selected using support vector machine (SVM), least absolute shrinkage and selection operator (LASSO) regression, Random Forest and Boruta (RFB), and extreme gradient boosting (XGBoost) analyses by calculating the importance score for each variable via the glmnet, rms, e1071, caret, randomForest, Boruta, and XGBoost packages in R [15-17]. The expression of the ARGs-related differentially expressed genes (DEGs) were identified using a fold change cutoff value of 2 and a false discovery rate (FDR) <0.05. Then the identified DEGs were selected as the input variable (independent variables), and the status of subtypes was selected as the outcome (binary dependent variables, 0 or 1). The receiver operating characteristic (ROC) curves were used to evaluate the performance of the four machine learning algorithms for feature selection in the training set, and the areas under the ROC curve (AUCs) were subsequently compared. Afterwards, the most critical subtype-related genes were obtained from the intersecting genes among the LASSO, SVM, RFB and XGBoost analyses, and were visualized using a Venn diagram. Finally, the critical genes were analyzed by multivariate logistic regression analysis and used to construct the predictive model, which was termed by ‘Subtype Predictor’. The ROC curve was used to investigate the performance of the subtype predictor which was further used to determine the optimal cutoff values in discriminating different subtypes, as well as the AUC, sensitivity, specificity and accuracy. Finally, the test set was used to validate the predictive performance of the subtype predictor in a similar way.

**The evaluation of the efficacy anti-angiogenesis therapy response**

The data from the Genomics of Drug Sensitivity in Cancer (GDSC, https://www.cancerrxgene.org/) were downloaded to explore the response of anti-angiogenesis therapy regimes between high- and low-risk groups. The index of half-maximal inhibitory concentration (IC_{50}) was used for the response evaluation.

**Statistical analysis**

R software (version 3.6.0) was utilized to implement the all-statistical analyses and plots. The X² test was executed to assess the correlation between risk score and clinical characteristics. The significant difference of OS between the low ARGs risk score group and high ARGs risk score group was analyzed by Kaplan-Meier curve and the log-rank test. The relationship between risk scores and OS was evaluated by using the univariate and multivariate Cox proportional hazard regression analyses. The sensitivity and specificity of gene signature risk score in predicting survival were detected using ROC analysis. The prognostic accuracy was estimated by the area under ROC curve (AUC) analysis. In all analyses, P-value <0.05 was set to be statistically significant.

**Results**

**The identification and prognostic analysis of candidate ARGs in the cohort from TCGA**

To investigate ARG variation in GC, a total of 36 ARGs with expression profiles in the cohort obtained from TCGA were selected (Figure 1A). Based on the median value of ARG score (ssGSEA scores), all patients with GC included in this cohort were stratified into high-ARG and low-ARG groups (Figure 1B, the top panel). The middle panel and bottom panel displayed in Figure 1B revealed that the survival rates, including both the OS and DFS rates of patients in the high-risk group differed from those in the
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A

B

C

D

E

F

Am J Transl Res 2022;14(7):4532-4548
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Figure 1. The identification and prognostic analysis of the candidate angiogenesis-related genes in the TCGA cohort. A: The heatmap of the 36 angiogenesis-related genes' expression in gastric cancer tissues. B: The distribution and median value of the ssGSEA scores in the TCGA cohort (up panel). The distribution of OS, OS status, and risk score in the TCGA cohort (middle panel). The distribution of DFS, DFS status, and risk score in the TCGA cohort. C: The Kaplan-Meier curves for the OS of patients in the TCGA cohort, which was divided into high- and low-ARG group. D: The Kaplan-Meier curves for the DFS of patients in the TCGA cohort, which was divided into high-risk group and low-risk group. E: ROC (Receiver operating characteristic) curves showing the 1-, 3-, and 5-year OS and DFS predictive efficiency of the ARGs-signature in the TCGA cohorts. F: The analyses of clinical features between high- and low-ARGs prognosis score groups in GC cohort. ARG: angiogenesis-related genes; OS: overall survival; DFS: disease-free survival; ROC: receiver operating characteristic.

low-risk group. Moreover, compared with those in the low-ARG group, the OS and DFS of patients in the high-risk group were significantly shorter (Figure 1C and 1D). The AUC value predicted that the 1-, 3- and 5-year OS in the training cohort was 0.61, 0.64 and 0.76, respectively, and the AUC value predicted that the 1-, 3- and 5-year DFS was 0.61, 0.68 and 0.65, respectively (Figure 1E). Subsequently, the clinical features of patients with high- or low-ARG prognosis scores were investigated. A high-ARG score was significantly correlated with tumor stage (T) and neoplasm disease stage, as more patients with T4 and/or stage IV disease were presented in the high-ARG group (Figure 1F). We also compared the expression level 36 ARGs between normal and tumor samples in TCGA-GC and other 4 GEO datasets (including GSE13195, GSE13911, GSE27342, GSE63089), and found that the majority of the ARGs were overexpressed in the tumor samples, except SERPINA5 and PGLYRP1 (Figure S1).

ARG signatures independently predict OS and DFS

In order to validate whether the newly-constructed ARG signature could be utilized to predict the prognosis of patients with GC as an independent risk factor, various clinicopathological parameters were tested between high-risk and low-risk score groups. In the TCGA cohort, the univariate analysis revealed that ARGs (P=0.002), T-stage (P=0.001), N-stage (P<0.001), M-stage (P=0.004), neoplasm-disease-stage (P=0.001) and neoplasm-disease-grade (P=0.021) were markedly correlated with OS. Multivariate Cox regression analysis demonstrated that ARGs, M-stage, and neoplasm-disease-grade were independent risk factors of OS. In addition, univariate and multivariate analyses indicated that ARGs as independent risk factors were correlated with a reduced DFS (Figure 2A).

Subsequently, patients in the training group were stratified into different subgroups according to clinical characteristics, and the correlation between ssGSEA score and OS of patients with GC was further analyzed. The results of the present study demonstrated that following the stratification of different clinical characteristics, the ARG signature score exhibited an effective prognostic effect in patients with high stage (stage III-IV), tumor stage (T3-4) and N stage (N1-3) GC (Figure 2B).

Tumor microenvironment profiles related to ARG score

To further examine the correlation between the ARG score and immune status, ssGSEA was used to calculate the enrichment of different immune cell contents. As demonstrated in Figure 3A, a positive correlation between cytolytic activity (CYT) score, ESTIMATE, human leukocyte antigens (HLA), immune checkpoint gene (ICG), tumor-infiltrated lymphocyte (TIL) and ARG score was identified by Spearman's rank analysis. Further investigations demonstrated a significant negative correlation between tumor purity and ARG score (Figure 3B). Moreover, EPIC analysis of the tumor-infiltrated lymphocyte profiles demonstrated a statistically significant difference between the high- and low-ARG score groups, including CAFs, T cell CD4+, T cell CD8+, macrophages, endothelial cells and NK cells (Figure 3C). The significant difference in CAF content between high- and low-ARG score groups was also determined using XCELL (Figure 3D).

Association between CAF subtype and ARG signature in patients with GC

CAFs are one of the most abundant cell types within the cancer stroma, and these were analyzed in patients with GC with different ARG signature scores. According to the results displayed in Figure 4A-H, only the matrix CAF
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Figure 2. The correlation analysis between ARG score and survival in gastric cancer patients. A: Independent prognostic factors for overall survival (OS) and disease-free survival (DFS) in gastric cancer patients. Univariate and multivariate Cox regression analysis was used to identify the relationship between clinicopathological features (including the risk score) and OS and DFS of patients in the training group. B: Stratified analysis of the correlation between ARG score and survival rate (OS and DFS) of patients with gastric cancer in the training group based on different clinical feature, including neoplasm disease stage (Stage I-II and stage II-III), neoplasm disease grade (G1-2 and G3), tumor stage (T1-2 and T3-4), lymph node metastasis stage (N0 and N1-3), long distant metastasis (M0 and M1).

(mCAF) subtype was significantly correlated with reduced OS and DFS, while the remaining subtypes, including vascular CAF (vCAF) and developmental CAF (dCAF) had no notable...
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Figure 3. Tumor microenvironment related to ARG score in GC samples from the TCGA dataset. (A) Spearman’s rank order correlation between ARG score and ICG, CYT, HLA, ESTIMATE, TIL. (B) The correlation analysis of ARG score and tumor purity. (C and D) Comparison of abundance of immune cells between high- and low-ARG group by EPIC (C) and XCELL (D). CYT: cytolytic activity score; HLA: human leukocyte antigens; ICG: immune checkpoint gene; TIL: tumor-infiltrated lymphocyte.

Figure 3A. Spearman's rank order correlation between ARG score and ICG, CYT, HLA, ESTIMATE, TIL.

Figure 3B. Correlation analysis of ARG score and tumor purity.

Figure 3C. Comparison of abundance of immune cells between high- and low-ARG group by EPIC.

Figure 3D. Comparison of abundance of immune cells between high- and low-ARG group by XCELL.

Effect on the survival of patients with GC. Moreover, comparable results were observed in the three validation sets (GSE26253, GSE66229 and GSE26901, presented in Figure S2A-D). Subsequently, the correlation between ARGs and various CAF subtypes was deter-
Figure 4. Survival analysis of different ARGs and CAFs subgroups. The Kaplan-Meier curves were used for analyzing the OS and DFS of patients in the TCGA cohort, which was divided into high and low CAF density groups (A, B), mCAF (C, D), vCAF (E, F), dCAF (G, H). Spearman’s rank order correlation between ARGs and CAFs including total CAF (I, J, K, L, M, N).
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mined using Spearman’s rank analysis, and results demonstrated that ARGs were positively correlated with various CAFs, including mCAF, vCAF and dCAFs (Figures 4I-L, S2D). Moreover, patients with low-ARG score and mCAF content (ARGs-mCAFs-) exhibited the greatest OS rates. By contrast, patients in the high-ARG score and mCAF content group exhibited the poorest prognosis (ARGs+mCAFs+, Figure 4M, 4N).

Validation of the prognostic ARG signature using external datasets

To further evaluate the robustness and stability of the identified ARG signatures from the training data set, survival and TME analyses were carried out using GSE26253, GSE26091 and GSE66229 datasets. The same formula was used to calculate ARG scores for patients from the validation cohorts, who were divided into low- and high-subgroups according to the median value of ARG score in each cohort. As demonstrated in the validation results, patients with GC in the low-ARG score subgroup exhibited a significantly superior OS and DFS, compared with patients with high-ARG score in all three validation cohorts (GSE26253: OS, P=0.012; GSE26901: OS, P=0.0012 and DFS, P=0.0026; GSE66229: OS, P=0.00011 and DFS, P=0.003; Figure 5). Moreover, the stromal, immune and ESTIMATE scores were markedly increased in the high-ARG score group from the GSE26253 dataset (P<0.001, Figure S3A). The correlation between ssGSEA score, TILs and ICGs (CTLA4, TIM3, LAG3 and IDO1) was analyzed using Spearman’s rank analysis (Figure S3B). A significant correlation was only identified between TIM3 and ARG signature score (R=0.31, P<0.001). Similar results were observed in both GSE26901 and GSE66229 cohorts (Figures S4 and S5).

Cluster analysis based on ARG expression profiles

The consensus clustering analysis was performed to further assess the prognostic implication of ARGs in patients with GC from the training group (TCGA). The 414 patients with GC from the training group were clustered into two subgroups, named cluster 1 and 2. The distinction of gene expression levels between cluster 1 and cluster 2 was further validated using principal component analysis (PCA) (Figure 6A). Moreover, to investigate the overlap among the classification methods of cluster analysis and ARG score, a Venn diagram was used to demonstrate these results (Figure 6B). Results of the survival analysis demonstrated that the OS of patients from cluster 2 was significantly longer, compared with that of patients from cluster 1 (Figure 6C and 6D). In addition, stromal, immune and ESTIMATE scores, as well as TIM level were significantly lower in cluster 2 than those in cluster 1 (Figure 6E, 6F). Cluster 1 exhibited a significantly increased B cell, CAF, CD4 T cell, endothelial cell, macrophage and uncharacterized cell contents (Figure 6G), which was further validated using the XCELL analysis (Figure 6H).

Establishment and validation of the ARG-related prognostic predictor

Within the training set, the most critical subtype-relevant features were identified using four machine learning algorithms, according to the expression levels of DEGs. LASSO, RF, SVM and XGBoost analyses were carried out to identify 33, 114, 80 and 110 genes, respectively. A total of 10 critical genes were identified using the Venn diagram, and the genes were shared by the four featured selection algorithms (Figure 6I). Subsequently, the diagnostic predictive model was constructed using a multivariate lasso-based logistic regression analysis. ARG subtype predictor was calculated using the formula as follows: risk score = 5.869 + 0.852* (expression level of DCLK1) + 0.295* (expression level of PTGIS) + 0.340* (expression level of NUDT10) + 0.598* (expression level of ZFHX4) + 0.290* (expression level of ECRG4). ROC analysis demonstrated an AUC of 0.994 for separating subtypes I and II in the training set (Figure 6J). In addition, the subtype predictor also exhibited a high level
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Figure 5. The prognostic performance of ARG score in the three validation sets. Kaplan-Meier curves of overall survival and disease-free survival of the GSE26901 (A), GSE26253 (B), and GSE66229 (C) cohort.

of performance in distinguishing the subtypes as evaluated in the test set, with an AUC value of 0.979 (Figure 6K). Kaplan-Meier survival analysis revealed that GC patients with a high-risk score presented with significantly inferior OS and PFS in the GSE26901 cohort (Figure 6L, 6M). The AUC of predicting 1-, 3- and 5-year OS in the GSE26901 cohort were 0.7, 0.65 and 0.64, respectively, and the AUC of predicting 1-, 3- and 5-year OS in the GSE26901 cohort were 0.66, 0.62 and 0.59, respectively (Figure 6N, 6O). The association between risk-score and patient survival was also identified in the GSE66229 cohort, as patients with a high risk score exhibited a significantly reduced DFS and OS (Figure 6O, 6P). The prediction efficacy of the constructed ARG signature was stable and adequate in the GSE66229 cohort (the AUC of predicting 1-, 3- and 5-year OS was 0.59, 0.63 and 0.64, respectively; the AUC of predicting 1-, 3- and 5-year DFS was 0.57, 0.64 and 0.65, respectively, Figure 6R, 6S).

Identification of the molecular characteristics, and the efficacy of immunotherapy and targeted therapy

As demonstrated in Figure 7, the percentage of patients with recurrent GC in the GSE26901 cohort was significantly different, and patients that suffered with disease recurrence exhibited significantly higher risk scores (Figure 7A). Moreover, the EMT subtype of GC was significantly higher than the other three GC subtypes, including MSI, TP53 positive and TP53 negative in GSE66229 (Figure 7B). The higher predicted capacity was concentrated in the GS subtype, compared with the CIN, POLE, MSI and EBV infection subtypes (Figure 7C). The low-risk score group presented an increased tumor mutational burden compared with the high score group (Figure 7D, 7E), and the Kaplan-Meier curves demonstrated that patients with the low-risk score exhibited an improved prognosis compared with those with the high-score in the IMvigor210 dataset (Figure 7F). Patients who responded to anti-PD-1/L1 immunotherapy exhibited lower low-risk scores compared with those without response (P=0.051, Figure 7G). In addition, the mutation counts, FGFR4 and HER2 were significantly negatively correlated with the risk score (Figure 7H). Notably, in accordance with angiogenesis, patients with a high ARG subtype score exhibited a significantly lower IC50 level of specific VEGF and PDGF receptor inhibitors, which may inhibit angiogenesis and the development of tumors. These included suni-
Figure 6. Construction and validation of a ARGs subtype predictor. (A) Principal component analysis (PCA) of the two clusters (cluster 1 and 2) in the TCGA cohort. (B) A Venn diagram of different expression genes of different clusters. The Kaplan-Meier curves for the OS (C) and DFS (D) of patients in the TCGA cohort, which was divided into cluster 1 and cluster 2 group. (E) Comparison of the immune score, stromal score, and ESTIMATE score between cluster
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1 and cluster 2. (F) The expression level of TIM3 in the two clusters. Comparison of the contents of infiltrating immune cells between cluster 1 and cluster 2 was conducted by ESTIMATE (G) and XCELL (H). (I) A 10 most critical subtype specific genes were identified by Venn diagram, which were shared by four feature selection algorithms. ROC curves of the subtype predictor in distinguishing two subtypes in the train set (J) and test set (K). Kaplan-Meier survival analysis also suggested that patients’ OS and PFS were significantly different between Subtype I and II both in GSE26901 (L and M) cohort. ROC curves showing the 1-, 3-, and 5-year OS (N) and DFS (O) predictive efficiency of the ARGs-signature in the GSE26901 cohort. Kaplan-Meier survival analysis also suggested that OS and PFS of patients were markedly different between Subtype I and II both in GSE66229 cohort (P, Q). (J, K) ROC curves showing the 1-, 3-, and 5-year OS (R) and DFS (S) predictive efficiency of the ARGs-signature in the GSE66229 cohort. ROC: Receiver operating characteristic; OS: overall survival; DFS: disease-free survival.

tinib, sorafenib, pazopanib and axitinib. Although these drugs have not been approved for GC treatment to date, patients with GC with a high-risk signature score may experience promising benefits from anti-angiogenesis therapy (Figure 7I).

Discussion

In the present study, we developed a risk stratification system that predicts GC patients’ survival from ARG feature, and verified its accuracy on independent validation datasets. The ARG score and related risk model could robustly and stably predict patient outcomes, having correlation with TME and immune feature. Furthermore, we also found the application value of the established risk score in treatment efficacy.

Angiogenesis plays a crucial role in tissue repair and regeneration in physiological and pathological processes, and is one of the hallmark events in cancer development [18]. Under normal physiological conditions, endothelial cells are sensitive to angiogenic signals and involved in angiogenesis by maintaining a high degree of plasticity. In contrast, in many disease states, including cancer, rheumatoid arthritis and atherosclerosis, aberrant angiogenesis will further accelerate the progression of these diseases, and is considered a hallmark of these disease states [19]. Previous evidence revealed that identified ARG may be used as a prognostic marker for the survival of patients with GC [20]. Results of the present study demonstrated that the survivals of patients with GC were significantly decreased in the high-ARG score group compared with those in the low-ARG score group, which is in consistent with previous research [21]. These results further supported that the ARG signature may act as an important prognostic prediction marker for GC patients.

Angiogenesis interacts with TME in cancer, and cell contents in TME interact with surrounding cells via the circulatory and lymphatic systems to further impact the development and progression of cancer [22]. The TME is mainly composed of three types of cell components, endothelial cells, immune cells (granulocytes, lymphocytes and macrophages) and fibroblasts [23]. GC patients with high ARGs score had significantly higher abundance of CAFs, which are a major component of the tumor stroma and crucial in facilitating crosstalk between cancer cells and the TME. Previous studies have demonstrated that CAF and endothelial cells play important roles in tumor cell proliferation, tumor angiogenesis, escaping from immune surveillance and metastasis [24]. Meanwhile, CAFs could promote breast cancer cell invasion by inducing the upregulation of genes involved in angiogenesis [25]. In addition, results of numerous previous studies revealed that CAFs increased angiogenesis by inducing the aberrant expression of key angiogenic factors. We also found that GC patients with both high abundance of mCAF and high ARG score had the most inferior survival.

We identified 10 key genes related to the ARG subtype, and the majority of them have been suggested as oncogenes that promote tumor development, invasion and metastasis in GC, including DCLK1 [26], NUDT10 [27], PTGIS [28], CHRLD1 [29], AGTR1 [30] and CNTN1 [31]. Previous studies suggested that most of these oncogenes could promote tumor by regulation TME, inducing EMT and angiogenesis in tumor [32]. In contrast, PCDH9 [33] and ECRG4 [34] were regarded as tumor suppressers in GC. Further in vitro and or in vivo studies are merited to give deep insights on how these genes regulating angiogenesis and tumor development in GC, and whether they could be therapeutically targeted to improve patients’ survival.

To date, chemotherapy remains the cornerstone of the systemic treatment for patients with unresectable advanced or recurrent GC,
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but with only limit benefits in improving both the PFS and OS [34]. The development of precision medicine, including targeted and immune therapies, are of urgent needs to overcome the treatment dilemma of GC. Notably, the finally established ARG subtype predictor risk score could not only robustly and stably predict patients’ prognosis but also provide indications for therapeutic efficacy. Anti-angiogenesis therapy, especially those targeted at VEGF and/or its receptor, has shown promising antitumor efficacy in multiple types of cancers, including gastric cancer. Antibody blockade-based (bevacizumab [35] and ramucirumab [36] and tyrosine kinase inhibitor (TKI)-based strategies anlotinib and apatinib [37]) have been approved to use as monotherapy or combination with chemotherapy for gastric cancer by the USA and Chinese administrations. Some other TKIs, including sorafenib, sunitinib, pazopanib and axitinib have shown clinical benefits by combing with chemotherapy in patients with GC [38]. In this study, we found albeit high-risk patients had worse survival, and they were more sensitive to these anti-angiogenesis TKIs, which is in consistent with the mechanism how the predictor constructed. Another emerging novel therapy for GC is the immune checkpoint inhibitors. However, they were only efficacious to small part of patients. MSI and TMB are the FDA-approved pan-cancer biomarkers associated with better clinical outcomes treated with pembrolizumab, an anti-PD-1 blockade, and have been proved in the GC [39]. Interestingly, we found that the risk score was significantly negatively correlated with the TMB, and patients in the MSI subtype had the lowest risk score. In concordance with these results, high risk patients had inferior survivals and lower response rate to atezolizumab based on analysis of the Imvigor210. Furthermore, we found the risk score was also negatively correlated with ERBB2 and FGFR4 levels. ERBB2, also known as HER2, is overexpressed in 10-30% of GC and is one of the most important actionable biomarkers in GC. Multiple targeted strategies, including antibodies, antibody drug conjugate (ADC) and TKIs had proved promising anti-tumor effects in GC [40]. In addition, recent clinical trials, such as KEYNOTE-811, further provided clinical rationales for the combination of anti-HER2 and anti-PD-1/L1 for treating HER2-positive GC patients [41]. In short, these results provided potential treatment stratification for GC patients, and patients with higher risk score may have better clinical benefits from anti-angiogenesis therapy but lower possibility to response to anti-HER2 and/or immune checkpoint blockades.

In conclusion, this study confirmed the prognostic role of ARGs and its correlation between ARGs and TME, especially CAFs in GC. In addition, a 10-gene based risk model was constructed, providing not only prognosis but also potential treatments stratification. However, this study has serval limitations. First, though the performance of the established ARG and related prognostic signature was stable and robust, the construction and validation of them were all based on the public data. So, a confirmatory test using local tumor tissue samples from patients with gastric cancer needs to be carry out in future. Second, the mechanisms of the identified 10 genes involved in ARG subtype predictor in regulating tumor angiogenesis and development are still unknown. Further in vitro or in vivo study is merited to clarify the biology function.
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Acknowledgements

This work was supported in part by a grant from the National Natural Science Foundation of China (NSFC No. 82002122) and Changning district medical and health research projects of Shanghai (CNKW2018Y35) to Ning Ma, National Natural Science Foundation of China (NSFC No. 81472479) to Bin Wang and Foundation of Naval Medical University (2021QN26) to Jie Li.

Disclosure of conflict of interest

None.

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Figure S1. Comparison of the expression level 36 ARGs between normal and tumor samples in TCGA-GC, GSE13195, GSE13911, GSE27342 and GSE63089 datasets. *P<0.05; **P<0.01; ***P<0.001; ****P<0.0001; ns: not significant.
Figure S2. Association between CAFs subtypes and ARG signature in GC patients in three validation cohorts. The difference in the survival between high- and low-ARG risk score in GSE26901 (A), GSE66229 (B) and GSE26253 (C). (D) The correlation between ARGs and matrix cancer-associated fibroblasts (mCAFs). OS: overall survival; DFS: disease-free survival; RFS: recurrence.
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Figure S3. ARGs signature associated tumor microenvironment analysis in GSE26253. (A) The difference in Stromal, Immune and ESTIMATE score between high- and low-ARGs score samples in GSE26253. (B) Correlations between ARG score and TILs, ICGs (CTLA4, TIM3, LAG3 and IDO1) were analyzed by Spearman analysis. Tumor-infiltrated immune cells were analyzed by XCELL (C) and EPIC (D). *P<0.05; **P<0.01; ***P<0.001; ****P<0.0001; ns: not significant.
Figure S4. ARGs signature associated tumor microenvironment analysis in GSE26901. (A) The difference in Stromal, Immune and ESTIMATE score between high- and low-ARGs score samples in GSE26901. (B) Correlations between ARG score and TILs, ICGs (CTLA4, TIM3, LAG3 and IDO1) were analyzed by Spearman analysis. Tumor-infiltrated immune cells were analyzed by XCELL (C) and EPIC (D). *P<0.05; **P<0.01; ***P<0.001; ****P<0.0001; ns: not significant.
Figure S5. ARGs signature associated tumor microenvironment analysis in GSE66229. (A) The difference in Stromal, Immune and ESTIMATE score between high- and low-ARGs score samples in GSE66229. (B) Correlations between ARG score and TILs, ICGs (CTLA4, TIM3, LAG3 and IDO1) were analyzed by Spearman analysis. Tumor-infiltrated immune cells were analyzed by XCELL (C) and EPIC (D). *P<0.05; **P<0.01; ***P<0.001; ****P<0.0001; ns: not significant.