Integrated bioinformatics analysis and experimental validation reveals fatty acid metabolism-related prognostic signature and immune responses for uterine corpus endometrial carcinoma

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Background: Uterine corpus endometrial carcinoma (UCEC) is the third most common gynecologic malignancy. Fatty acid metabolism (FAM) is an essential metabolic process in the immune microenvironment that occurs reprogramming in the presence of tumor signaling and nutrient competition. This study aimed to identify the fatty acid metabolism-related genes (FAMGs) to develop a risk signature for predicting UCEC.

Methods: The differentially expressed FAMGs between UCEC samples and controls from TCGA database were discovered. A prognostic signature was then constructed by univariate, least absolute shrinkage and selection operator (LASSO) and multivariate Cox regression analyses. Based on the median risk score, UCEC samples were categorized into high- and low-FAMGs groups. Kaplan-Meier (K-M) curve was applied to determine patients’ overall survival (OS). The independent prognostic value was assessed by uni- and multivariate analyses. The associations between the risk score and immune status, immune score, and drug resistance were evaluated. Quantitative Real-time PCR (qRT-PCR) was utilized to confirm FAMGs expression levels in UCEC cells.

Results: We built a 10-FAMGs prognostic signature and examined the gene mutation and copy number variations (CNV). Patients with a high-FAMGs had a worse prognosis compared to low-FAMGs patients in TCGA train and test sets.
We demonstrated that FAMGs-based risk signature was a significant independent prognostic predictor of UCEC. A nomogram was also created incorporating this risk model and clinicopathological features, with high prognostic performance for UCEC. The immune status of each group was varied, and immune score was higher in a low-FAMGs group. HLA-related genes such as DRB1, DMA, DMB, and DQB2 had higher expression levels in the low-FAMGs group. Meanwhile, high-FAMGs patients were likely to response more strongly to the targeted drugs Bortezomib, Foretinib and Ge. The qRT-PCR evidence further verified the significant expression of FAMGs in this signature.

Conclusions: A FAMGs-based risk signature might be considered as an independent prognostic indicator to predict UCEC prognosis, evaluate immune status and provide a new direction for therapeutic strategies.

Keywords: uterine corpus endometrial carcinoma, fatty acid metabolism, risk signature, immune status, prognosis

Introduction

Uterine corpus endometrial carcinoma (UCEC), a frequent gynecological malignant tumor, has been rapidly increasing in recent years. According to the 2022 cancer statistics, there were an estimated 65,950 cases and 12,550 deaths from uterine corpus cancer in United State (1). In China, it is estimated that the number of new cases was 84,520, and deaths were 17,543 of UCEC in young women (2). It, therefore, remains a major public health issue around the world. Despite surgery, chemotherapy, radiotherapy and brachytherapy currently employed to UCEC intervention, there are still considerable numbers of women with more aggressive lesions whose prognosis is dismal (3). A reliable prognosis assessment is the foundation of effective therapy. However, the present predictive system based on clinical, pathological, imaging, and biological features are insufficient to interpret the progressive and prognostic heterogeneity of UCEC (4). Consequently, exploring effective biomarkers to identify individuals with a high risk of recurrence is helpful for precise therapy.

Tumor growth is deeply reliant on the tumor microenvironment (TME), which is characterized by hypoxia, acidity, and nutrition deprivation because tumor cell proliferation is faster than angiogenesis (5). Consequently, tumor cells displayed distinctive metabolic properties from normal cells to handle a variety of adverse situations via a metabolic reprogramming process that supports their growth and survival once the carcinogenic signal is blocked (6). Lipid metabolism is one of three primary energy metabolisms of cells, of which fatty acid metabolism (FAM) is a critical metabolic pathway and plays an important role in cancer pathophysiology (7). Specifically, the fatty acid in particular aids cancer cells in not only maintaining membrane biosynthesis but also supplying a major energy source during metabolic pressure. The FAM pattern tends to be different among various cells and tissues. Previous studies have shown that certain expression patterns of fatty acid metabolism-related genes (FAMGs) were connected with proliferation, prognosis, and immunity of glioma, colorectal cancer, or breast cancer (8–11). However, the pattern of FAMGs in UCEC has not been explored.

As bioinformatic technology develops, numerous approaches have been employed to define meaningful biomarkers (12, 13). In this study, the potential value of FAM in UCEC samples obtained from TCGA database was evaluated by using bioinformatics. We analyzed the differential expression of FAMGs, and selected genes strongly correlated with UCEC prognosis to develop a FAMGs-based risk signature. A scoring system was created to evaluate the probability of survival, as well as immune status and drug sensitivity of UCEC. This risk model offered a novel insight on UCEC prognosis and therapeutic options.

Materials and methods

Data sources

A total of 539 UCEC and 35 normal endometrial cases were retrieved from TCGA database (https://portal.gdc.cancer.gov/). Patients were allocated to TCGA training or test cohort in a 1:1 ratio at random. FAM gene sets, including KEGG FAM pathways, Hallmark FAM genes, and Reactome FAM genes, were acquired from the MSigDB v7.4. A total of 309 FAMGs
were finally ascertained after removing the overlapping genes (14).

Construction and verification of a prognostic signature

The DEGs linked to FAM between UCEC, and normal tissues were screened by R “limma” package (15), with |logFC|>0 and FDR-adjusted $P<0.05$. Upon these FAM-DEGs, the overall survival (OS) related genes with $P<0.05$ were selected by using a univariate cox analysis. Then, candidate FAMGs were confirmed by using the least absolute shrinkage and selection operator (LASSO) and multivariate cox regression analysis via R “glmnet” packages (16).

The risk score is computed by using this formula: $risk \ score = \sum_{i=0}^{n} (coef_i \times Gene_i)$, where $Gene_i$ denotes the expression level of gene $i$ and $coef_i$ denote the regression coefficient of gene $i$. Based on median risk score, UCEC patients were divided into a low- or high-FAMGs group in TCGA training or test set. The log-rank test was utilized to compare the difference in survival status between two groups via R “rms” package. Kaplan-Meier (K-M) analysis of OS or PFS was performed via R “survival” package. To reflect the predictive power of risk model, we plotted the time-dependent receiver operating characteristic (ROC) curve and area under the curve (AUC) for 1-year, 3-year, and 5-year OS via R “timeROC” package (17).

Gene Ontology (GO) (18) and Kyoto Encyclopedia of Genes and Genomes (KEGG) (19) functional enrichment analysis of FAMGs in UCEC were conducted by R “ClusterProfiler” package, where $P<0.05$ indicates a statistical difference.

Establishment of a FAMGs-related nomogram

To verify the independence of the FAMGs-based risk signature, we ran uni- and multivariate cox analyses on risk score and other clinical factors. Then, using the above variables, we established a FAMGs-related clinicopathologic nomogram via R “rms”, “nomogramEx”, and “regplot” packages (20). Then, ROC and calibration curves were utilized to examine the accuracy and discrimination of the nomogram.

Assessment of immune status and drug sensitivity

The association between tumor immune microenvironment (TIME) and this prognostic signature was further assessed. The single-sample gene-set enrichment analysis (ssGSEA) was utilized to quantify the immune activity in high- and low-FAMGs groups. Tumor Immune Dysfunction and Exclusion (TIDE) algorithm was conducted to evaluate the tumor immune escape in two groups based on FAMGs. ESTIMATE algorithm was used to determine the TME score via R software. The expressions of HLA-genes between high- and low-FAMGs group were further compared. To investigate differences in therapeutic effects of small-molecule drugs between two groups, we calculated the half-maximal inhibitory concentration (IC50) values of drugs commonly for UCEC treatment via R “pRRophetic” package [21fron].

Cell culture and qRT-PCR

Human endometriosis cell line hEM15A and UCEC cell ANC3A were cultured in DMEM with 10% FBS in a 5% CO2 incubator at 37°C. Total RNA was extracted by utilizing Trizol reagent (Invitrogen, USA), then cDNAs were generated with a HiScript Synthesis kit (Vazyme, China). Quantitative real-time PCR (qRT-PCR) was completed by utilizing the Fast SYBR Green Master Mix (Roche, USA) on a StepOnePlus Real-Time PCR system (Applied Biosystems, USA). Primer sequences in our work are described in Table S1.

Results

Landscape of FAMGs expression in UCEC

There are 554 UCEC and 35 normal samples retrieved from TCGA dataset. Differentially expressed FAMGs between tumor and normal were presented in a heat map (Figure 1A). The volcano diagram displayed 100 up-regulated FAMGs and 106 down-regulated FAMGs (Figure 1B). Principal component analysis (PCA) was applied to evaluate sample heterogeneity, and the results showed a significant difference (Figure 1C). We also included more normal samples from GTEx database (n=78) to verify this difference (Figure S1). 131 up-regulated FAMGs and 103 down-regulated FAMGs were found. Most of the different FAMGs overlapped. These suggested that FAMGs might have a potential ability to differentiate normal patients from UCEC.

Identification of prognostic FAMGs and genomic variance analysis

Firstly, 206 differentially expressed FAMGs were subjected to univariate Cox analysis, and 28 prognosis-related genes were identified with a $P<0.05$ (Figure 2A). Next, LASSO and multiple Cox analyses were utilized to shrink the range of FAMGs. At last, 10 FAMGs including upregulated PECR, OLAH, ACOT11, ACAT2,
NUDT19, PTGIS, and downregulated GPX1, ADH5, PTGR1, ACADS were identified for prognostic risk model (Figures 2B, C).

The incidence of somatic mutation of FAMGs in UCEC patients was calculated from TCGA cohort. There are 64/518 (12.36%) UCEC patients who experienced mutations of these 10 FAMGs, with a frequency from 0 to 3% (Figure 2D). Among them, ACOT11, ACADS, PECR, PTGR1, PTGIS, OLAH had 3% mutation frequency, followed by ACAT2, ADH5, NUDT19, while GPX1 did not have any mutations. Next, we investigated the CNV of FAMGs and found its prevalence in these 10 FAMGs. The location of CNV changes of the FAMGs on each of their respective chromosomes was shown in Figure 2E. Among them, NUDT19, PTGIS, ACOT11, OLAH, PTGR1, and ADH5 exhibited widespread CNV gain, while GPX1, ACADS ACAT2, and PECR had CNV loss (Figure 2F).

**Construction and validation of FAMGs-based risk signature**

Based on FAMGs median risk score, UCEC patients were categorized into high- and low-FAMGs groups in TCGA training cohort, test cohort and total cohort (Figures 3A). The proportion of alive patients in high-FAMGs group was less than that of low-FAMGs patients among these datasets (Figure 3B). The distribution of these 10 FAMGs expression levels UCEC
Identification of prognostic FAMGs and their characteristic analysis. (A) Forrest plot of 28 FAMGs related with prognosis by univariate regression analysis. (B) LASSO regression analysis. (C) Partial likelihood deviance for LASSO regression. (D) Profiles of genetic mutation in UCEC patients. (E) CNV alteration of FAMGs on chromosomes. (F) Frequencies of CNV gain, loss, and non-CNV among FAMGs. FAMGs: fatty acid metabolism-related genes. LASSO: least absolute shrinkage and selection operator. UCEC: Uterine corpus endometrial carcinoma. CNV: copy number variation.
patients was shown in a heatmap (Figure 3C). K-M curve revealed that low-FAMGs patients’ OS was considerably longer than high-FAMGs patients among these cohorts (Figure 3D), as did their PFS time (Figure S2).

Functional annotation

Further, we identified 835 differentially expressed genes between high- and low-FAMGs groups. To examine the possible biological properties in UCEC, we conducted a functional enrichment analysis of these genes. GO analysis showed that signaling receptor activator activity, tubulin binding, tubulin binding and fatty acid synthase activity were significantly enriched in biological processes (Figure S3A).

KEGG revealed the enrichment in microtubule-based movement, cilium assembly, cilium movement and pattern specification process (Figure S3B).

Independent prognostic value of FAMGs-based risk signature

To elucidate whether FAMGs-based risk signature is an independent prognostic indicator for UCEC, uni- and multivariate Cox analyses were carried out in TCGA total set. As shown in Figure 4A, age ($P = 0.021$), histological type ($P < 0.001$), stage ($P < 0.001$), grade ($P < 0.001$), and risk score ($P = 0.024$) had significant correlation with OS. Multivariate Cox analysis (Figure 4B) showed that age ($P = 0.013$), stage ($P <
0.001), grade (P = 0.007) and menopause status (P = 0.026) and risk score (P = 0.042) were related to OS. We then performed a time-dependent ROC curve to test this risk model’s predicting ability and accuracy (Figure 4C). The result showed that AUC of the 1-, 3-, or 5-year OS was 0.740, 0.761, 0.778, respectively. Importantly, the AUC value for this risk model was 0.740, which was higher than that for the age (0.568), histological type (0.560), and stage (0.719) (Figure 4D). These results indicated that our FAMGs-based risk signature exhibited a great independent predictive value in UCEC.
Establishment of FAMGs-related nomogram

A nomogram integrating the risk model, age, grade, stage, histological type, and menopause status was created to predict UCEC patients’ OS (Figure 4E). By summing the point for each prognostic factor, a total point was generated for each patient, and higher total points meant a worse outcome. The calibration plot showed a close agreement with the ideal model, demonstrating the nomogram’s perfect stability and discrimination (Figure 4F). ROC curve showed that this nomogram (AUC = 0.854) had a superior predictive ability than a separate parameter, such as age (AUC = 0.600), stage (AUC = 0.747), or risk model (AUC = 0.779) (Figure 4G).

Prognostic power of FAMGs in UCEC patients

We examined the prognostic power of FAMGs-based risk signature for UCEC patients under different clinicopathological factors, including age, grade, histological type, menopause status and stage. K-M analysis for each subgroup revealed that low-FAMGs patients have longer OS than high-FAMGs patients, no matter their age, grade, and stage, as is the same situation in patients with EAC or post-menopause (Figure 5). These suggested that FAMGs-based risk signature has strong predictive power in most populations with different clinical features.

Analysis of immune microenvironment

We further evaluate the relationship between TIME and this prognostic signature. Immune status of low- and high-FAMGs patients revealed some degree of heterogeneity (Figure 6A). In addition, high-FAMGs group had a lower TIDE score compared with low-FAMGs group (Figure 6B). Regarding the TME score, high-FAMGs patients had lower immune scores, and ESTIMATE score than low-FAMGs patients, but no significant difference in Stromal score between two groups (Figure 6C). Furthermore, we observed that high-TMB (tumor mutational burden) was linked to a better OS ($P<0.001$, Figure 6D). We then combined FAMGs with TMB to divide patients into high-TMB/low-risk, low-TMB/low-risk, high-TMB/high-risk, and low-TMB/high-risk groups.

FIGURE 5
Prognostic power in different clinical subgroups. K-M curve analyses for patient subgroups, including (A) age, (B) grade, (C) histological type, (D) menopause, (E) stage. K-M: Kaplan-Meier.
As seen in Figure 6E, a significant difference among all groups was identified (P < 0.001), with high-TMB/low-risk group showing the highest OS. Furthermore, we also found that most of HLA-related genes, such as DRB1, DMA, DQB2, DMB were expressed significantly higher in low-FAMGs group (Figure 6F).

Response to therapeutic drugs

Given that the risk score is linked to a poor prognosis, it is necessary to investigate the impact of FAMGs-based risk signature on the drug resistance for UCEC. IC50 was designed to predict the therapeutic response to common targeted drugs. As shown in Figures 7A-C, low-FAMGs samples had greater IC50 values of Bortezomib, Foretinib and Gefitinib compared with high-FAMGs patients. Moreover, the risk score is inversely correlated with drug sensitivity (Figures 7D-F). These results suggested that high-FAMGs patients were more responsive to Bortezomib, Foretinib and Gefitinib.

Validation of FAMGs expression

To further validate the expression of FAMGs in this signature, we performed a qRT-PCR experiment to detect the
The difference between UCEC cells and normal endometrial cells. As expected, the expressions of PECR, OLAH, ACOT11, ACAT2, NUDT19 and PTGIS were upregulated, whereas GPX1, ADH5, PTGR1 and ACADS were downregulated (P<0.05) in UCEC cells compared to normal cells (Figure 8). These were consistent with the above bioinformatic results.

**Discussion**

UCEC is one of the most prevalently diagnosed gynecologic malignancy and ranks sixth among female tumors worldwide (22). In recent years, UCEC incidence and mortality have been increased with a younger trend. Early-stage UCEC could be
surgically removed followed by chemoradiotherapy, with a 5-year survival rate of up to 90% (23). Nevertheless, advanced-stage patients at a high risk of recurrence had a worse prognosis, some prospective trials have attempted to identify these patients to develop effective adjuvant therapy (24–26), but to date, no interventions have been proven to improve OS. Traditional clinicopathological parameters are insufficient for precisely forecasting the outcome of UCEC, as patients in same clinical stage may exhibit distinct clinical features (27). As a result, the discovery of novel biomarkers for the prognosis and therapy of UCEC has become an urgent clinical issue to be resolved.

It is widely recognized that a major cause of the development of UCEC is obesity (28), which is linked to fatty acid oxidation, synthesis, accumulation and metabolic regulation. Additionally, a majority of FAMGs are known to be tightly related to malignancy and prognosis of cancers. The evidence supports a potent prognostic value of FAMGs for UCEC patients. The advent of bioinformatics has allowed us to examine the specific pattern of FAMGs for UCEC. In our study, a prognostic signature with 10 FAMGs was designed to predict UCEC patients’ survival time based on TCGA cohort. Among them, the expression levels of six genes (PECR, OLAH, ACOT11, ACAT2, NUDT19, PTGIS) were significantly upregulated and 4 genes (GPX1, ADH5, PTGRI, ACADS) were downregulated in UCEC samples compared to normal tissues, which have been validated by PCR experiment. Substantial evidence revealed that most of these FAMGs functioned as oncogenes or tumor-suppressor genes in various cancers (29–36). We also determined the prevalence of genetic mutation in UCEC samples, as well as CNV alterations. These confirmed the significant effect of fatty acid metabolism on UCEC malignant progression. Furthermore, a scoring system was created to identify and verify the prognostic value of FAMGs-based risk signature, allowing for the effective risk categorization of UCEC. High-FAMGs UCEC patients had a worse OS or PFS compared with low-FAMGs group. Importantly, FAMGs-based risk signature exhibited great accuracy and independence in predicting UCEC prognosis than other factors. In addition, a nomogram integrating our signature and patient characteristics was created as a superior tool to predict the survival of UCEC. We also discovered that this signature offered predictive value for different subgroup of patients with specific clinical characteristics. These findings indicated that FAMGs-based risk signature may be a reliable predictor of UCEC.

Tumor metabolism not only modulates the signaling for tumorigenesis and survival, but also an antitumor immune response by releasing intermediate metabolites to affect the expression of immune molecules (38). FAM is a vital metabolic route involved in the immune response, which contains anabolic and catabolic activities for energy homeostasis, as well as metabolites generation that keeps cell membrane structure and function, stores energy and allows cross-talk between tumor and immune cells (39).

In our study, the effect of FAMGs in the risk signature on UCEC TME was further investigated. Patients with low- and high-FAMGs displayed remarkably different immune status. Following processing by ESTIMATE algorithm, it was discovered that low-FAMGs group had a higher estimate score relative to high-FAMGs group, indicating this risk signature may be able to function as a new immune indicator in UCEC. Then, TIDE technique was applied to anticipate clinical response to immune checkpoints and suggested that high-FAMGs patients were more effective to immunotherapy. We continue to investigate the TMB of UCEC patients. The results showed that patients with higher TMB possessed a better prognosis than those with lower TMB. Similar findings have been observed in other cancers (40–42), highlighting the possibility that TMB might serve as a prognostic marker for guiding more efficient immunotherapeutic approaches (43). By combing with FAMGs risk score, patients with low-TMB/high-FAMGs had the lowest survival probability than other groups. In addition, given the importance of HLA-related genes to immune system (44), we compared their expression levels in high- and low-FAMGs group, and found that most of these genes were generally increased in low-FAMGs group. Moreover, the resistance and sensitivity of common targeted drugs were measured to evaluate the prognostic ability of this risk model for therapeutic outcomes. These findings proved that FAMGs-based risk signature was associated with immune status and tumor treatment of UCEC.

Nevertheless, our study also has several limitations. First, the research was completely performed based on TCGA database, but lacked external cohorts for validation. Second, the regulatory mechanism of fatty acid metabolism in UCEC TME warrants further investigation. Third, the value of FAMGs-based risk signature for clinical application requires multi-center, large sample trials to be processed.

In conclusion, we identified significant FAMGs in UCEC and constructed a FAMGs-based risk signature to predict the patients’ prognosis using systematic bioinformatic analyses. Patients with high-FAMGs had a worse prognosis than low-FAMGs patients. This signature might be regarded as an independent indicator to estimate the survival time, immune response and treatment effect. This study provided us a new understanding and direction on the evolution of FAMGs in UCEC.
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.

Author contributions

CG, YH, LC, and YL designed this work. YW, YB, and NZ analyzed the data, FJ and HZ wrote this manuscript. LZ edited and revised the manuscript. All authors contributed to the article and approved the submitted version.

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Conflict of interest

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

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Supplementary material

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fonc.2022.1030246/full#supplementary-material

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