Research Article

Molecular Subtypes Based on Cuproptosis-Related Genes and Tumor Microenvironment Infiltration Characterization in Colorectal Cancer

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Recent studies have demonstrated the biological significance of cuproptosis modification, a newly discovered programmed cell death, in tumor progression. Nonetheless, the potential role of cuproptosis-related genes (CRGs) in the immune landscape and tumor microenvironment (TME) formation of colorectal cancer (CRC) remains unknown. We comprehensively assessed cuproptosis modification patterns of 1339 CRC samples based on 27 CRGs and systematically analyzed the correlation of these patterns with TME. The CRG-score was constructed to quantify cuproptosis characteristics by LASSO and multivariate Cox regression methods, and its predictive capability was validated in an independent cohort. We identified three distinct cuproptosis modification patterns in CRC. The TME immune cell infiltration demonstrated immune heterogeneity among these three subtypes. Enrichment for multiple metabolism signatures was pronounced in cluster A. Cluster C was significantly correlated with the signaling pathways of immune activation-related, resulting in poor prognoses. Cluster B with mixed features possibly represents a transition phenotype or intratumoral heterogeneity. Then, based on constructed eight-gene CRG-score, we found that the signature could predict the disease-free survival of CRC patients, and the low CRG-score was related to increased neoantigen load, immunity activation, and microsatellite instability-high (MSI-H). Additionally, we observed significant correlations of the CRG-score with the cancer stem cell index and chemotherapeutic drug susceptibility. This study demonstrated that cuproptosis was correlated with tumor progression, prognosis, and TME. Our findings may improve the understanding of CRGs in TME infiltration characterization of CRC patients and contribute to guiding more effective clinical therapeutic strategies.
1. Introduction

Colorectal cancer (CRC) is the third most frequent malignancy and the second biggest cause of cancer-related mortality in the world [1], which has very limited treatment options still, despite the vast amount of research undertaken. Targeted therapies for solid tumors, such as tyrosine kinase inhibitors and immunotherapy, have shown little benefit in CRC [2] (except for BRAF V600E-mutated tumors [3]). The gene expression profiles of CRC have strong clinical relevance. The characteristics of immune cells and stromal cells infiltrating the tumor microenvironment [4, 5], combined with different subtypes of microenvironment-related signals and inherent signals of cancer cells, have always shown predictive value [6, 7]. In recent clinical trials, combining the molecular and histological characteristics of tumors to guide treatment has improved the prognosis of CRC patients [8–10]. The most widely used risk assessment tools are the current molecular subtype of consensus (CMS) and the tumor lymph node metastasis staging system (TNM). However, the current robust classification system has limitations in predicting this highly heterogeneous disease [11–13]. Therefore, more factors related to prognosis need to be considered. And the implementation of molecular classification in the clinical decision-making of CRC is crucial to solving various clinical problems in CRC progression.

Cuproptosis is a novel form of programmed cell death along with copper accumulation, protein lipoylation, and mitochondrial respiration [14]. Cuproptosis differs from other forms of cell death, such as apoptosis, necrosis, autophagy, and ferroptosis, in terms of molecular process. Copper binding causes a hazardous increase of function in lipoylated tricarboxylic acid (TCA) cycle proteins, excess copper increases lipoylated protein aggregation and instability of Fe-S cluster proteins, resulting in proteotoxic stress and cell death. As major regulators of cuproptosis, FDX1 and protein lipoylation play important roles in this process. Copper ionophores are extremely sensitive to cells conducting mitochondrial respiration, which is explained by their high quantities of lipoylated TCA enzymes. CRC tumor cells demonstrate aberrant mitochondrial metabolism as a result of active oncogene and loss of tumor suppressor gene [15], and in the tumor microenvironment (TME), aerobic glycolysis is widely observed in activated immune cells, to support biosynthetic demands [16]. The TME is now well recognized as playing a critical role in carcinogenesis and cancer development. Mitochondria serve as an intriguing signals and inherent signals of cancer cells [17, 18]. Cuproptosis, a kind of programmed cell death that varies from apoptosis, is likely to offer fresh hope for CRC tumor therapy. Despite this, little is known about the involvement of cuproptosis-related genes (CRGs) in prognosis and the tumor microenvironment. As a result, a thorough understanding of the features of TME immune cell infiltration mediated by numerous CRGs should help researchers better understand the underlying mechanism of CRC carcinogenesis and predict immunotherapy response.

The goal of this study is to evaluate the prognostic value and numerous functions of CRGs in the TME in a systematic way. According to The Cancer Genome Atlas (TCGA) and Gene Expression Omnibus (GEO) datasets, 1339 CRC samples were divided into three cuproptosis subtypes based on 27 CRGs, and the differences in survival and immune cell infiltration of these subtypes were explored. Patients were subsequently divided into two gene subtypes based on three cuproptosis subtypes’ differentially expressed genes (DEGs). Additionally, a CRG-score was established to quantify the cuproptosis characteristics of a single tumor. The findings revealed that the CRG-score is a powerful prognostic marker.

2. Materials and Methods

2.1. Data Sources. Figure S1 in the Supplementary Material online depicts the research design used in this investigation. The public databases GEO (https://www.ncbi.nlm.nih.gov/geo/) and TCGA (https://portal.gdc.cancer.gov/) were used to get open CRC gene expression datasets with clinical information and survival outcomes. For the following analyses, a total of six datasets were obtained, containing five datasets from the GEO datasets (GSE17536, GSE39582, GSE17537, GEO161158, and GSE38832) and one from the TCGA CRC dataset (Table 1). Background correction and normalizing of CEL data received from GEO were performed using Robust Multichip Average with quantile normalization. For normalized counts, the FPKM format of RNA-seq data from the TCGA cohort was transformed to log2(TPM+1). GSE17536, GSE17537, GSE39582, GSE38832, and TCGA CRC datasets were pooled, and a batch correction was conducted using the ComBat method. Subsequent analyses involved a total of 1339 CRC patients. Age, sex, TNM stage, BRAF mutation, KRAS mutation, mismatch repair (MMR) status, follow-up period, and survival status were incorporated among the clinical factors. The open-source R/Bioconductor programs were used to handle and analyze all of the data.

2.2. Consensus Clustering and Pathway Enrichment Analysis of CRGs. Twenty-seven CRGs were retrieved from the research of Tsvetkov et al. with a false discovery rate (FDR) < 0.05 as a filter condition [14]. Tables S1 and S2 provide all of the information on these genes. Unsupervised clustering analysis was utilized to categorize CRC patients into different molecular subgroups based on CRG expression using the “ConsensusClusterPlus” package in the R studio program. A consensus clustering approach was performed to estimate the number of clusters [19]. Furthermore, the hallmark gene sets (c2.cp.kegg.v7.4.symbols) were also obtained from the Molecular Signatures Database (MSigDB) database, and the activity of signaling pathways was further accurately assessed by a gene set variation analysis (GSVA) algorithm in each subtype. The possible relationship between subtypes and disease-free survival (DFS) was investigated using Kaplan-Meier survival curves.

2.3. CRC Immune Landscape. CIBERSORT (https://cibersort.stanford.edu/) was used to examine the relative number of
22 immune cells in 1339 CRC samples to better understand their immunological features. We assessed the quantity of distinct immune-infiltrating cell subtypes in each sample based on the gene expression matrix of 1339 samples and the supplied gene expression feature set of 22 immune cell subtypes. The differences in the percentage of immune cells were analyzed using the Wilcoxon rank-sum test in each subtype. We also further observed the differential expression levels of PD-1, PD-L1, and CTLA4 among these subtypes.

2.4. Construction and Validation of CRG-Score Based on DEGs. With a fold-change > 1.5 and an adjusted P value of 0.05 as filter conditions, the DEGs among the cuproptosis clusters were discovered by the limma R package. Following that, the functional annotation of the DEGs was done with the “clusterProfiler” software, and then Gene Ontology (GO) and Kyoto Encyclopedia of Genes and Genomes (KEGG) enrichment analysis were performed to clarify the biological function of these DEGs. The findings of unsupervised cluster analysis were utilized to identify possible gene categories after detecting prognosis-associated DEGs using univariate Cox regression. Finally, CRC patients were allocated in a 1:1 ratio to the training (n = 670) and testing (n = 669) groups, and a CRG-score related to cuproptosis was created. To reduce the overfitting of recurring characteristics and restrict the range of genes predicting DFS, LASSO Cox regression was used with the R package. The pheatmap and maftools packages in R were used to characterize the DEGs heatmap and mutation landscape, respectively. A P value of less than 0.05 was considered statistically significant for all two-sided statistical P values. R version 4.1.1 was used for all statistical analyses, and no default settings were utilized in any of the investigations.

3. Results

3.1. Genetic Mutation Landscape of CRGs in CRC. Based on the study by Tsvetkov et al., we first identified 27 CRGs through a whole-genome CRISPR-Cas9 positive selection screen using two copper ionophores (Cu-DDC and Elesclomol-copper) in cells (Figure 1(a)). Then, we analyzed the mutational landscape of the CRGs in the TCGA-CRC datasets. As depicted in Figure 1(b) 95 of 535 (17.76%) CRC samples presented genetic mutations among the 27 CRGs. Furthermore, we explored the mutational incidence of copy number variation (CNV), indicating that 27 CRGs demonstrated evident CNV alterations (Figure 1(c)). Among them, SLEN11, SOX2, YEAST2, AHR, MPC1, HAUS5, AFG3L2, COQ7, LIPT1, PDHA1, and GLS had widespread CNV increases, while MUCR1, OXAL1, LIAS, MBTPS1, BRPF1, REXO2, RPL3, CAPRIN1, FDX1, CDKN2A, DLAT, PDHB, and SCAP showed CNV decreases. Figure 1(d) displayed the site of CNV alterations of 27 CRGs on chromosomes. We also analyzed the expression levels of CRGs in CRC and normal tissues and discovered that most CRG

<table>
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<th>Sample type</th>
<th>Number of samples</th>
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<td>HG-U133 plus 2.0</td>
<td>CRC = 191</td>
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where \(\text{Coefi} \times \text{Expri} \) expressed the signature genes and the multivariate Cox regression calculated the Coefi coefficient. The patients were separated into low CRG-score and high CRG-score groups using the median CRG-score as the cut-off value, and then Kaplan–Meier survival analysis was performed.

2.6. Statistical Analyses. The random stratified sampling was carried out using R’s sample function. Meanwhile, the heatmap and maftools packages in R were used to characterize the DEGs heatmap and mutation landscape, respectively. One-way ANOVA and Kruskal–Wallis tests were used to determine the significance of differences between three or more groups. The Wilcoxon test was utilized to compare the differences between the two groups. The correlation coefficients were calculated by the Spearman and distance correlation analysis. The log-rank tests and Kaplan–Meier survival analysis were created using the survminer R program. A P value of less than 0.05 was considered statistically significant for all two-sided statistical P values. R version 4.1.1 was used for all statistical analyses, and no default settings were utilized in any of the investigations.
Altered in 95 (17.76%) of 535 samples.
expression levels were linked to CNV changes (Figure 1(e)), and CRGs with CNV gains, such as YEATS2 and AHR, were significantly elevated in CRC samples.

3.2. Identification of Cuproptosis Subtypes in CRC. This study enlisted 1339 CRC patients from five suitable cohorts (TCGA-CRC, GSE39582, GSE17536, GSE17537, and GSE38832) for
Cuproptosis
Positive correlation with $P < 0.0001$
Risk factors
Negative correlation with $P < 0.0001$
Favorable factors

Cox test, $P$-value
1e−04
0.001
0.01
0.05
1

Figure 2: Continued.
were significantly different.

3.3. Characteristics of the TME in Different Subtypes. We used GSVA enrichment analysis on the three subtypes to observe whether there were any changes in biological behavior between them (Figure 3(a) and Table S5). Cluster A and cluster B, cluster A and cluster C, and cluster B and cluster C were compared in terms of enrichment analysis, respectively. We observed that enrichment for multiple metabolism signatures was pronounced in cluster A, including pyruvate metabolism, pentose phosphate pathway, and citrate cycle TCA cycle. In contrast, cluster C showed enrichment in terms of pathways associated with immune activation, such as the activated JAK-STAT signaling pathway, TOLL/NOD-like receptor signaling pathways, and antigen processing and presentation. Following TME cell infiltration investigations (Table S6), as shown in Figures 3(b) and 3(a), significant enrichment difference in most immune cells was observed among the three clusters. In cluster A, significant immune-infiltrating cells were plasma cells, T cells CD8, T cells CD4 memory resting and activated, and NK cells resting. Cluster B had an immune cell infiltration level that was halfway between clusters A and C. Similarly, CTLA4, PD-1, and PD-L1 expression levels were found to be higher in cluster C (cluster C vs. cluster A), when three critical immunological checkpoints were examined (Figure 3(c)). We utilized the ESTIMATE method to get TME scores (stromal score, immune score, and estimate score) in the different clusters, and TME scores could evaluate the amount of immunological and stromal components in TME. CRC patients in cluster C had higher TME scores, according to the
Figure 3: Continued.
Figure 3: Correlations of tumor immune microenvironment with three CRG subtypes. (a) GSVA of biological pathways among three distinct subtypes in five datasets, where red and blue represent activated and blue inhibited pathways, respectively. (b) Abundance of 22 infiltrating immune cell types in three CRG subtypes. (c) Expression levels of PD-1, CTLA4, and PD-L1 among three CRG subtypes. (d) Difference of TME score among three CRG subtypes. CRG: cuproptosis-related genes; GSVA: gene set variation analysis; TME: tumor microenvironment.
findings (Figure 3(d)). Up to a point, the higher TME scores in cluster C can indicate tumors in cluster C patients have higher interactions between stromal cells and immune cells.

3.4. Development and Validation of a Gene Signature Based on Cuproptosis-Related Clusters. We identified the DEGs between clusters A and C (P < 0.05) in order to further investigate the probable biological role of the cuproptosis subtypes in CRC, and 702 genes were retrieved. Then, using the “clusterProfile” package, we performed GO and KEGG enrichment analysis and discovered that these DEGs are strongly correlated with immune biological processes (Figures 4(a) and 4(b) and Table S7). Following that, 428 prognosis-related genes were identified using univariate Cox regression analysis for these 702 cuproptosis subtype-related genes. Then, based on the 428 prognosis-related DEGs, a consensus clustering technique was used to divide patients into two genomic subgroups, which were dubbed gene clusters I and II, respectively (Figure 4(c), S3, and Table S8). On the basis of DEG expression levels, we discovered that the two gene subtypes could be distinguished considerably (Figure 4(d)). Furthermore, CRC patients in gene cluster II had a worse DFS than those in gene cluster I, according to the survival analysis (Figure 4(e)). Meanwhile, we found the expression levels of the majority of CRGs (22/27) were significantly different between the two gene subtypes (Figure 4(f)).

To establish a model that could quantify each CRC patient, we assigned randomly CRC patients to a training cohort (n = 670) or a testing cohort (n = 669) at a ratio of 1:1. Then eight of the 428 DEGs were retained by application of LASSO regression and multivariate Cox regression models (Figure S4 and Table S9). We used these eight genes to build a cuproptosis-related signature score which we named the "CRG-score" = (−0.469 × expression level of KLRD1) + (0.281 × expression level of SMARCA1) + (0.185 × expression level of SCG2) + (0.263 × expression level of SERPINE1) + (0.227 × expression level of H3ST2) + (0.230 × expression level of CXCL10) + (0.217 × expression level of SELENBP1) + (0.148 × expression level of MMP12) (Table S10). We divided CRC patients into high and low CRG-score groups with the cut-off value of the median. We further observed molecular subtype distributions with different CRG scores, and DFS outcomes in Figure 5(a), and found that the low- and high-risk CRG-score was significantly differential in the cuproptosis clusters and gene clusters (Figures 5(b) and 5(c)).

Then, in the training and testing sets, we sought to establish the possible predictive value of each CRC patient's CRG-score (Figures S5 and S6). The Kaplan–Meier survival curves in two datasets demonstrated that CRC patients with low risk had a considerably better disease-free survival than those with high risk (log-rank test, P < 0.001; Figure 5(d)). Furthermore, for 1-, 3-, and 5-year survival rates, the AUC values of CRG-score were 0.701, 0.724, and 0.687, respectively (Figure 5(e)). To validate the applicability and generalizability of the CRG-score, we further verified this score in an independent cohort with a total of 191 CRC patients (GSE161158) and obtained the same results (Figure S7). We further included CRG-score as an effective indicator and other clinical characteristics in univariate and multivariate Cox regression analysis and found that CRG-score could be recognized as an independent factor that predicts the prognosis of CRC patients (Table 2). Additionally, we also created a predictive nomogram based on the CRG-score and clinicopathological parameters to predict the 1-, 3-, and 5-year DFS status of each patient in the training and testing sets, and calibration plots revealed that the constructed nomogram performed similarly to an ideal model (Figures S8a, S8b, S8d, and S8e). Meanwhile, in the training and testing sets, we also evaluated the prediction accuracy of the CRG-score with clinicopathological markers, and the findings revealed that our signature's AUC was better than that of other clinical criteria (AUC = 0.723) (Figures S8(c) and S8(f)).

3.5. Correlation of CRG-Score with TME, TMB, MSI, and CSC Index and Drug Susceptibility. We discovered that CRG-score might play an essential role in clinical prediction based on these findings. Then we observed whether the CRG-score may be used as a guide for treatment, particularly immunotherapy. With different CRG scores, we looked at TME cell infiltration. First, when the CRG-score grew, the immuneScore from ESTIMATE analysis decreased, but the stromalScore had the reverse impact (Figure 5(f)). Furthermore, immunological activation-related cells such as T cells CD8, NK cells, and macrophages M1 had significant negative correlations with CRG-score (P < 0.001), but M2 macrophages had a positive connection with CRG-score owing to polarization (P < 0.001) (Figure 5(g)). Furthermore, we discovered that the majority of immune invading cells were highly associated with the eight CRG-score genes (Figure 5(h)).

TMB and MSI were important predictors of tumor immune response and played an indispensable role in carcinogenesis and progression. As a result, we investigated the function of MSI and TMB in the CRG-score in greater depth. Low CRG-score was connected with MSI-H status, whereas the high CRG-score was associated with microsatellite stable (MSS) status, according to correlation studies (Figures 6(a) and 6(b)). The Spearman correlation analysis revealed a weak correlation between CRG-score and TMB (R = −0.15) (Figure 6(c)), and no significant difference in TMB between the low-risk and high-risk groups (Figure 6(d)). As a result, we looked at the changes in somatic mutation distribution in the CRG-score signature. We observed the top 20 somatic mutation genes with high mutational frequencies in TCGA CRC cohort and found no significant changes in TMB or mutation types between the high and low CRG-score groups (Figures 6(e) and 6(f)).

In addition, CRG-score was shown to be inversely linked with the CSC index in Figure 6(g), showing that CRC cells with a lower CRG-score exhibited more unique stem cell features and a lower degree of cell differentiation (R = −0.38, P < 0.001). Then, we chose chemotherapeutic drugs commonly used in CRC treatment to observe how sensitive individuals in low- and high-risk categories were to them. Interestingly, CRC patients with a high CRG-score had lower IC50 values for shikonin, whereas
Figure 4: Continued.
chemotherapeutics including sorafenib, ABT.888, and gefitinib had considerably lower IC50 values in the low CRG-score group (Figure 6(h), S9).

4. Discussion

Despite the introduction of the term cuproptosis, the cellular processes, overall impact, and TME infiltration features mediated by the combined actions of numerous CRGs have yet to be thoroughly understood in CRC. The 27 CRGs were studied in terms of prognostic value, functions in the immune microenvironment, and putative regulatory mechanisms in CRC in this work. Despite the low mutational intensity of CRGs, the majority of CRGs (22/27) were found to be differently expressed between normal and malignant tissues, with a total of 22 genes having substantial prognostic significance. The finding of differences in expression levels of CRGs indicated the latent function of CRGs in CRC tumorigenesis.

We then divided CRC patients into three cuproptosis subgroups (Clusters A, B, and C) using the unsupervised clustering approach. Cuproptosis clusters were connected to distinct cellular metabolism-related signaling pathways and immunological infiltrations, and were shown to be independent of clinicopathological characteristics. As a result, we...
Figure 5: Continued.
found two gene clusters with diverse immunological activities and roles based on DEGs connected to the subgroups' signature. By LASSO and multivariate Cox regression models, CRG-score was also constructed to quantify the cuproptosis gene subgroups. In the TME, MSI, CSC index, and drug susceptibility, CRC patients with low- and high-risk scores had significantly different prognoses. Specifically, when compared to standard clinical markers like the TNM...
Figure 6: Continued.
stage, the CRG-score outperformed them in predicting patient prognosis.

Cuproptosis is a kind of cell death that is dependent on copper and protein lipoylation [14]. It differs from other types of cell death. Copper levels are very high in CRC patients' serum and tissue [20–23]. Furthermore, cancer cells have elevated copper concentrations, which are thought to be critical for angiogenesis and metastasis [24]. Cuproptosis is therefore anticipated to play a key role in the establishment and progression of neoplasms. Our findings consistently showed that differences in mRNA transcriptomes between different cuproptosis subtypes were significantly related to metabolism and immune-related biological pathways, such as patients with cluster A had more metabolism reprogramming features and better DFS, and patients with cluster C had higher TME scores and showed enrichment in pathways associated with immune activation. The cuproptosis-related prognostic signature was found to be a good predictor of CRC patients' prognosis. These results indicate that the cuproptosis could lead to different biological behaviors in CRC patients, confirming the existing conclusions.

According to mounting evidence, the TME of CRC plays a critical role in tumor progression, invasiveness, metastasis, drug resistance, and even the maintenance of the cancer stem-like phenotype [25], which is primarily mediated by immune cells (macrophages, T cells, B cells, and so on), cytokines, chemokines, and exosomes [26]. Throughout cancer development, the TME constantly alters with great complexity. Immune interaction is a fundamental characteristic of CRC and a potential treatment target. The key components of TME are stromal cells and immune cells, and immune and stromal scores are linked to colorectal cancer components of TME are stromal cells and immune cells, and immune checkpoints (CTLA4, PD-1, and PD-L1), with a worse prognosis. Patients with a high CRG-score, as well as increased PD-1, PD-L1, and CTLA-4 expression, were shown to be more likely to react to immune checkpoint blocking. The exact mechanism linking cuproptosis with immunity, however, remains unknown. These findings might help us learn more about the association between cuproptosis and CRC TME infiltration cells.
Tumors have metabolic processes that have been altered to support cancer development [35]. The variability and flexibility of metabolism across malignancies and TME have been highlighted in recent research [36]. Metabolic heterogeneity is caused by a variety of mechanisms in the TME, including interactions between cancer cells and other TME components, such as immune cells, and extracellular matrix. In our study, we found that enrichment for multiple metabolism signatures was pronounced in cluster A, such as citrate cycle TCA cycle and pyruvate metabolism. Metabolic heterogeneity is significant because it has an impact on treatment vulnerabilities and may predict clinical outcomes. Copper-based compounds have been proven in several studies to be powerful cytotoxic agents that may specifically target cancer cells in vitro and in vivo [37–39]. Moreover, copper affects the proliferation of cancer cells, which can modulate tumor tissue homeostasis [40, 41]. The growing interest in the design of copper-based compounds as anticancer drugs stems. But, the exploitation of copper toxicity has been less successful in cancer. One of the main factors is that copper ionophore-induced cell death is mediated by the pathways of mitochondrial respiration. Copper ionophores are almost 1000-fold more sensitive in mitochondrial respiration-dependent cells than in glycolysis-dependent cells [14]. The metabolic activity of cancer cells is aberrant; the Warburg Effect is a frequent metabolic trait of cancer cells that entails a preference for aerobic glycolysis over oxidative phosphorylation to create ATP and biosynthetic building blocks [17]. It is not clear whether metabolic changes in CRC patients with cluster A are driven or mediated by cuproptosis-related mechanisms, or whether these metabolic changes are caused by the tumor itself, resulting in different cuproptosis-related changes. However, the molecular subtypes based on CRGs may certainly have some guiding significance for the treatment of copper-based compounds in CRC.

This study has some limitations. First, all analyses are based on several public databases, and the selection bias may affect their robustness, the stability of the type required multi-angle and multidatabase validation, and the link between cuproptosis and TME requires additional experimental verification. Furthermore, when assessing the prognosis of cuproptosis status, some treatment modalities such as surgery, radiotherapy, and chemotherapy were not included in the analysis of survival models, which may cause certain accuracy problems. In addition, tumor metabolic heterogeneity is an issue that must be addressed, and with the advancement of cuproptosis-related research, more focused solutions for different types of tumors may be provided.

5. Conclusion

In conclusion, this research discovered three subtypes of cuproptosis-related molecules in CRC, as well as their biological function, demonstrating the heterogeneity of the TME in CRC. Finally, we created a CRG-score signature to measure cuproptosis traits statistically. Furthermore, the CRG-score signature might be a useful prediction tool for predicting CRC patients’ prognosis and guiding appropriate therapeutic therapy. In summary, this research revealed new information on the involvement of cuproptosis and TME in boosting clinical outcomes and allowing tailored immunotherapy choices for CRC patients.

Data Availability

The data used to support the results of this study are available from The Cancer Genome Atlas (TCGA) database (https://portal.gdc.cancer.gov/repository) and the Gene Expression Omnibus (GEO) database (https://www.ncbi.nlm.nih.gov/geo/).

Conflicts of Interest

The authors declare that they have no conflicts of interest regarding the publication of this study.

Authors’ Contributions

F.W and F.H designed and supervised the research project and revised the manuscript until submitting it. H.H and Z.L performed research and drafted the manuscript. Y.X, P.Q, and X.Z performed the experiment and analyzed data. Y.X, X.L, and L.K collected the data and analyzed the data. Y.Z, L.Y, W.M, and Y.L reanalysed the results and interpretation. D.H, XX, and X.Z performed the figures and edited the data. D.H, X.S, and M.Z. revised the manuscript. All authors read and approved the final manuscript. All authors reviewed and approved the final version of the manuscript prior to submission. Hao Huang, Zhiping Long, and Yilin Xie contributed equally to this work and are the co-first authors.

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Supplementary Materials

Supplementary 1. Table S1-S10. Supplementary Table S1: Whole-genome CRISPR screen for the Elesclomol-Cu and Cupric-DDC in cells. Table S2: Summary of 27 cuproptosis-related genes. Table S3: The prognostic value of 27 cuproptosis-related genes in CRC patients. Table S4: The subtype of 27 cuproptosis-related genes in CRC patients. Table S5: The differential states of biological pathways in distinct cuproptosis subtypes by GSVA enrichment analysis. Table S6: Relative fractions of tumor-infiltrating immune cells of CRC patients by CIBERSORT’ algorithm. Table S7: Functional analysis of the 702 differentially expressed genes between the cuproptosis subtypes. Table S8: The gene subtype of cuproptosis subgroups related genes in CRC patients. Supplementary Table S9 The prognostic values of 428 cuproptosis subtype related genes in
CRC patients. Table S10: Multivariate Cox regression analysis of 8 CRGs associated with DFS in CRC patients.

Supplementary 2. Figure S1-S9. Figure S1: Flowchart of the research. Figure S2: Unsupervised clustering of cuproptosis cluster and consensus matrix heatmaps for k = 2, 4-9. Figure S3: Unsupervised clustering of cuproptosis-related gene cluster and Consensus matrix heatmaps for k = 3 – 9. Figure S4: The identification of candidate prognostic genes by LASSO regression analysis. Figure S5: Construction of CRG_score model in the training dataset. Figure S6: Validation of CRG_score model in the testing dataset. Figure S7: Validation of CRG_score model in the GSE161158 dataset. Figure S8: Development and evaluation of a prognostic nomogram in the training and testing sets. Figure S9: Relationships between CRG_score and chemotherapeutic sensitivity.

References


