Comprehensive analysis of differential expression profiles of mRNAs and lncRNAs and identification of a 14-lncRNA prognostic signature for patients with colon adenocarcinoma

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Abstract. The objective of this study was to identify potentially significant genes and long non-coding RNAs (lncRNAs) in colon cancer for a panel of lncRNA signatures that could be used as prognostic markers for colon adenocarcinoma (COAD) based on the data from The Cancer Genome Atlas (TCGA). RNA-seq V2 exon data of COAD were downloaded from the TCGA data portal for 285 tumor samples and 41 normal tissue samples adjacent to tumors. Differentially expressed mRNAs and lncRNAs were identified. A functional enrichment analysis of differentially expressed mRNAs was performed, followed by protein-protein interaction (PPI) network construction and significant module selection. Additionally, the regulatory relationships in differentially expressed mRNAs and lncRNAs were assessed, and an lncRNA-lncRNA co-regulation and functional synergistic analysis were performed. Furthermore, the risk score model and Cox regression analysis based on the expression levels of lncRNAs were used to develop a prognostic lncRNA signature. A total of 976 differentially expressed mRNAs and 169 differentially expressed lncRNAs were identified. MDFI and MEOX2 were the PPI network hubs. We found these lncRNAs to be mainly involved in vascular smooth muscle contraction and the cGMP-PKG signaling pathway. Several lncRNA-lncRNA pairs had co-regulatory relationships or functional synergistic effects, including BVES-AS1/MYLK-AS1, ADAMTS9-AS1/MYLK-AS1 and FENDRR/MYLK-AS1. The differential expression profile analysis of four candidate lncRNAs (MYLK-AS1, BVES-AS1, ADAMTS9-AS1, and FENDRR) in COAD tumors were confirmed by reverse transcription-quantitative PCR. Moreover, this study identified a 14-lncRNA signature that could predict the survival for COAD patients.

Introduction

Colon cancer is the third most frequently diagnosed malignancy and one of the leading causes of mortality globally (1). Colon adenocarcinoma (COAD) is by far the most common histologic type of colon cancer (2). Previous findings have shown that the morbidity and mortality rates of colon cancer continue to increase (3). Findings have demonstrated that colon cancer is successfully treated when identified at an early stage (4), and prognosis in colon carcinoma aids in the choice of therapeutic options. However, the underlying molecular mechanism of colon cancer and molecular biomarkers for the survival assessment of this cancer remains poor. Thus, the aim of ongoing research is to identify novel biomarkers, while studying the detailed molecular mechanism remains imperative.

Recent findings have identified multiple genetic alterations that result in tumorigenesis and the progression of colon cancer (5,6). Recently, Kan et al showed that nesfatin-1/nucleobindin-2 (NUCB-2) enhanced the migration, invasion, and mesenchymal phenotype of colon cancer via the LKB1/AMPK/TORC1/ZEB1 pathways (7). The findings of Unger et al demonstrated that stroma-induced insulin-like growth factor 2 (IGF-2) can promote colon cancer progression in a paracrine and an autocrine manner (8). Long non-coding RNAs (lncRNAs) are mostly defined as a class of non-coding RNAs exceeding 200 nucleotides in length (9). Increasing evidence has demonstrated that lncRNAs play key roles in regulating
many crucial biological processes such as genetic imprinting, cell differentiation, apoptosis and cell proliferation, and immune responses. IncRNAs are involved in numerous human diseases including various types of cancer (10,11). There are multiple ways by which IncRNAs can regulate downstream target genes (12). The expression of a c-Myc-activated IncRNA CCAT1 was shown to contribute to colon cancer tumorgenesis, and it also promoted colon cancer cell proliferation and invasion (13). Recently, it was found that, IncRNA-ATB may act on colon tumorigenesis by mediating E-cadherin repression and was also used as a predictor of poor prognosis (14). Additionally, Thorenoor et al reported that IncRNA ZFAS1 may function as an oncogene in colorectal cancer by interacting with cyclin-dependent kinase 1 (CDK1) and this interaction caused indirect destabilization of p53, leading to cell cycle progression and inhibition of apoptosis (15). Nevertheless, current knowledge concerning IncRNA regulation in colon cancer is limited, and more evidence is required to elucidate the gene and global IncRNA expression profiles in colon cancer.

In this study, we identified a RNA-seq-based mRNA and IncRNA signature of COAD patients from The Cancer Genome Atlas (TCGA) database using a cohort of >250 cases. Differentially expressed mRNAs and IncRNAs were identified in the tumor samples and normal controls. A functional enrichment analysis of differentially expressed mRNAs was performed, followed by a protein-protein interaction (PPI) network construction and significant module selection. Additionally, regulatory relationships among the differentially expressed mRNAs and IncRNAs were assessed, and IncRNA-IncRNA co-regulation and functional synergistic analyses were carried out. Moreover, the risk score model and Cox regression analysis based on the expression levels of IncRNAs were utilized to develop a prognostic IncRNA signature. The differential expression profile analysis of four candidate IncRNAs in COAD tumors were confirmed by reverse transcription-quantitative PCR (RT-qPCR). The present study successfully identified potentially significant genes and IncRNAs in colon cancer, and provided further insights into the mechanisms and the predictive capacity of IncRNAs underlying colon cancer.

Materials and methods

Public data acquisition and re-annotation. The RNA-seq V2 exon data (level 3, raw count) and clinical information of COAD were downloaded from the TCGA data portal (https://tcga-data.nci.nih.gov/) for a total of 326 samples (285 tumor samples and 41 normal tissue samples adjacent to tumors). Of these, 279 patients had complete prognostic information. Sequence data were generated using the Illumina HiSeq 2000 RNA Sequencing platform. The data were downloaded in March, 2017.

The RNA-seq V2 exon data provided information on the positions of all exons, raw counts, and reads per kilobase per million mapped reads (RPKM). To achieve an accurate and completely annotated dataset, we compared the RNA-seq V2 exon data with the annotation information of the IncRNA chromosomal location of the GENCODE (v25) database (16) (https://www.gencodegenes.org/). If the starting position of an exon was included in the IncRNA or protein-coding RNA in the GENCODE database and the positive and negative strands were consistent, the exon was defined as an IncRNA or protein-coding RNA. The raw data of 3,038 annotated IncRNAs and 19,161 annotated mRNAs were finally analyzed.

Identification of differentially expressed IncRNAs and mRNAs. Findings of a previous study demonstrated that a differential expression analysis based on a variance-stabilizing transformation combined with limma could perform well and were relatively unaffected by outliers (17). In the present study, edgeR+ limma was applied for the differential expression analysis. The raw count data were preprocessed using the R package edgeR (18) (version 3.4, http://www.bioconductor.org/packages/release/bioc/html/edgeR.html). The raw count was normalized to log-counts per million (CPM) values, and the genes whose average expression was lower than the first quartile (Q1) were filtered out. Linear modeling was performed and the mean-variance relationship was adjusted using the precision weights calculated using the voom function (19).

The data were divided into two groups: a COAD group and a normal control group. Using the t-test method provided by the limma package (version 3.10.3, http://www.bioconductor.org/packages/2.9/bioc/html/limma.html) (20), differentially expressed mRNAs and IncRNAs were identified in the COAD group compared with the normal control group. P-values were obtained from the t-test and were adjusted using the Benjamin-Hochberg (BH) (21) multiple testing correction to obtain adj. P-value. Thresholds for the screening of differentially expressed IncRNAs and mRNAs were adj. P-value <0.05 and llog, fold change (FC)>2.

Function, pathway enrichment analysis. The DAVID bioinformatics resource is a web-based application that consists of an integrated biological knowledge base and analytic tools for systematically extracting biological themes behind large gene lists (22). In the present study, functional and pathway enrichment analyses were performed for differentially expressed mRNAs using the DAVID online tools (version 6.8, https://david.ncifcrf.gov/), including Gene Ontology (GO) (23) and Kyoto Encyclopedia of Genes and Genomes (KEGG) pathway enrichment analyses (24). All three GO categories, i.e., biological process (BP), cellular component (CC), and molecular function (MF) were analyzed. GO terms and KEGG pathways with a P-value <0.05 were considered significant.

PPI network construction, module extraction and analysis. All the human PPIs present in Mentha (http://mentha.uniroma2.it/about.php), BioGRID (version 3.4; https://wiki.thebiogrid.org/) (25), and HPRD (release 9; http://www.hprd.org/) databases were taken as the background, and the PPIs among the differentially expressed mRNAs identified in the previous step were obtained. The resulting PPIs were then used for the construction of a PPI network using Cytoscape software (26). Additionally, CytoNCA (27) (version 2.1.6; http://apps.cytoscape.org/apps/cytonca), a Cytoscape plugin, was applied for the connectivity degree analysis. The important nodes of the top-ranked connectivity degree were considered as hub proteins (28).

The MCODE plug-in (29) of Cytoscape (26) applies a popular clustering method that can be used to find clusters (highly interconnected regions) in a network. In the present
study, functional modules were extracted from the PPI network using the MCODE plug-in (29). The parameters were set as follows: Included loops, false degree; cut-off, 10; node score cut-off, 0.2; haircut, true; fluff, false; K-core, 2; and max. depth from seed, 100. A KEGG pathway enrichment analysis was performed for the genes in the selected modules using the DAVID tool (22), and the threshold was set at \( p < 0.05 \).

**Analysis of lncRNA-mRNA regulatory relationships.** The Pearson's correlation coefficient \( r \) of each differentially expressed lncRNA and mRNA was calculated based on the expression values of the lncRNA and mRNA of the corresponding samples. In addition, a correlation test was performed to obtain the \( P \)-value. The lncRNA-mRNA pairs with \( |r| > 0.85 \) and \( P < 0.05 \) were identified, and the differentially expressed mRNA of the pair was deemed to be the target gene of the lncRNA. Furthermore, a functional and pathway enrichment analysis of the target genes of these lncRNAs was performed using the R package clusterProfiler (version 3.2.11, http://www.bioconductor.org/packages/release/bioc/html/clusterProfiler.html) (30), of which the enriched functions were considered functional properties of the lncRNAs. The threshold for the enrichment analysis was set at BH-corrected \( P \)-value <0.05.

**lncRNA-lncRNA co-regulation and functional synergistic analysis.** A co-regulatory network among lncRNAs was constructed based on the co-regulated genes targeted by two lncRNAs. If the co-regulated target genes of the two lncRNAs have significant GO BP (level 4) enrichment results, we believe there is a functional synergistic effect between the two lncRNAs. The functional synergistic network was constructed using all of the functional synergistic lncRNAs. The GO BP enrichment analysis was carried out using the R package clusterProfiler (30), with a significant enrichment threshold of BH-corrected \( P \)-value <0.05.

**Prognosis-related lncRNA screening.** The clinical data were analyzed to match the overall survival time (OS) and survival status with samples in the tumor group in the lncRNA matrix. We divided the differentially expressed lncRNAs into two groups according to the median expression value of the tumor group: high and low expression groups. The Kaplan-Meier (K-M) survival curves (31) were plotted and a log-rank test (31) was performed for the two groups. Several survival-related differentially expressed lncRNAs were obtained with \( P < 0.05 \) as the threshold for statistical significance.

In order to further screen for a prognosis-related differentially expressed lncRNA signature, a Random Survival Forests (RSF) (32) was carried out. RSF is a new extension of Random Forests (RF) to survival data. The bootstrap method was used to extract N samples from the original data, and the survival tree model was established to obtain the variable importance measure (VIMP) (33) of each variable, which measures the predictiveness of a variable (a negative value or a value closer to 0 is not predictive). RSF is then used for competing risk data to extract \( N \) samples from the original data, and the survival tree model was established to obtain the variable importance measure (CHF) (22). The differential lncRNA expression data were randomly divided into a test set (75% of the total samples) and a validation set (25% of the total samples), and the R package randomForestSRC (version 2.4.0, https://cran.r-project.org/web/packages/randomForestSRC/index.html) was used for RSF analysis. First, the RSF model was constructed through the test set to obtain the VIMP of each lncRNA in the model, and the VIMP were ranked from high to low. In addition, the lncRNAs were sorted in order and then included into the model to obtain the error rate of the current model; when the error rate was minimum, the current lncRNA combination was regarded as the optimal combination of the RSF models. The RSF model was reconstructed with the optimal combination, and the risk score was obtained by accumulating CHF values at different time points for each patient. The threshold that was used to distinguish high and low risk was set as the median of the risk score.

The optimal combination of prognosis-related differentially expressed lncRNAs was verified by the validation set, and the risk score of each sample was obtained by using the same parameters. The samples were divided into high- and low-risk groups by using the risk score threshold set in the previous step. The log-rank test was conducted on the K-M survival curves of the two groups. Multivariate Cox regression analysis was performed with clinical data to determine whether the identified lncRNA signatures were independent of other clinical variables.

**RNA extraction and RT-qPCR.** COAD tissues and matched normal adjacent tissues of 9 COAD patients were obtained from a residual sample biobank of the First Affiliated Hospital of Harbin Medical University (Harbin, China). The samples used were previously coded and anonymized. Total RNA was extracted using a Total RNA Extraction Kit (Axygen BioScience Inc., Union City, CA, USA; cat. no., AP-MN-MS-RNA-250) following standard protocols. RNA was reversely-transcribed into cDNA using the THUNDERBIRD® SYBR® qPCR Mix (Code no. QPS-201; Toyobo Co. Ltd., Japan). The quantitative detection of lncRNAs was performed using cDNA with ReverTra Ace qPCR RT Master Mix with the gDNA Remover (code no. FSQ-301; Toyobo) and ABI 7500 fast (Applied Biosystems, Thermo Fisher Scientific, Waltham, MA, USA). The reaction conditions used were: heat at 95°C for 1 min; then denaturation at 95°C for 15 sec, annealing at 60°C for 30 sec, and extension at 72°C for 60 sec for 40 cycles. Experiments were performed in triplicate, and the results were normalized to the expression of glyceraldehyde-3-phosphate dehydrogenase (GAPDH). The primers used were: **MYLK-AS1** forward, 5'-AGAGCAGGACAGAGGTTGTTG-3' and reverse, 5'-CCTGGCCTTCCAATCTCAGT-3'; **BVES-ASI** forward, 5'-TTTTACAGTTGTCTCACCATCC-3' and reverse, 5'-TGCACTTCAAGCCACCAT-3'; **ADAMTS9-ASI** forward, 5'-TCCTCATCATGCTTCTC-3' and reverse, 5'-TGCACTTCAAGCCACCAT-3'; **BVES-ASI** forward, 5'-CCTGGCCTTCCAATCTCAGT-3'; **ADAMTS9-ASI** forward, 5'-TCCTCATCATGCTTCTC-3' and reverse, 5'-TGCACTTCAAGCCACCAT-3'; **FENDR** forward, 5'-CCTGGCCTTCCAATCTCAGT-3' and reverse, 5'-CCTGGCCTTCCAATCTCAGT-3'; **GAPDH** forward, 5'-GCTCTGCTCCCTCCCTCGT-3'; **FENDR** forward, 5'-CCTGGCCTTCCAATCTCAGT-3' and reverse, 5'-CCTGGCCTTCCAATCTCAGT-3'; **GAPDH** forward, 5'-GCTCTGCTCCCTCCCTCGT-3'; **FENDR** forward, 5'-CCTGGCCTTCCAATCTCAGT-3'. The comparative cycle threshold (2\(^{-\Delta\Delta C_{T}}\)) method was used to determine the relative quantitative value.

**Statistical analysis.** Data were expressed as mean ± standard deviation (SD), and were analyzed using SPSS 17.0 (SPSS,
Inc., Chicago, IL, USA). The Mann-Whitney test was used to compare differences in lncRNA concentrations between the COAD and control groups. The log-rank test was used to compare differences in survival between the COAD and control groups. The univariate and multivariate Cox regression analyses were used to determine the risk score for different variables. P<0.05 was regarded as statistically significant.

Results

Basic characteristics of the samples. The basic characteristics of 279 patients with COAD are shown in Table I. The mean age of the patients was 64.97±13.3 years. Sixty-nine patients who succumbed to COAD were also included, accounting for 24.73% of the total individuals, and the average survival time was 32.48±29.89 months. Fig. 1 shows the flowchart of the analysis that was utilized in this study.

Screening of differentially expressed lncRNAs and mRNAs. According to the abovementioned method, 14,370 mRNAs and 2,278 lncRNAs were filtered and utilized in the differential expression analysis. Under the criterion for the differential expression analysis, 976 differentially expressed mRNAs were identified, including 304 upregulated and 672 downregulated mRNAs. Additionally, 169 differentially expressed lncRNAs were identified, including 57 upregulated and 112 downregulated lncRNAs.

Functional enrichment analysis of differentially expressed mRNAs. The GO and KEGG pathway enrichment analysis was performed on the upregulated and downregulated mRNAs, respectively. The top 10 BP terms, top 5 CC terms, top 5 MF terms, and top 10 pathways enriched by separate upregulated and downregulated mRNAs are shown in Fig. 2. The results showed that the upregulated genes were significantly enriched in the GO terms associated with embryonic development, cell proliferation, and pathways, such as the Wnt signaling pathway and the cytokine-cytokine receptor interaction pathway. Additionally, the downregulated genes were significantly involved in the GO terms associated with ion homeostasis and drug metabolism-related pathways.

PPI network construction and module selection. We identified 339 PPIs formed by 317 differentially expressed mRNAs in the PPI network, of which 166 genes were upregulated and 211 genes were downregulated. The top 10 nodes with higher degrees in the PPI network are shown in Table II, including the MyoD family inhibitor (MDFI; degree=20), mesenchyme homeobox 2 (MEOX2; degree=13), thyroid hormone receptor interactor 13 (TRIP13; degree=11), and C-X-C motif chemokine receptor 10 (CXCL10). Additionally, further sub-network module mining was performed in the network, and two significant modules (clusters) were identified, of which the larger module was comprised of 84 mRNAs and 113 PPIs (Fig. 3A). To infer the biological function of these mRNAs in the larger module, they were subjected to a KEGG pathway analysis. The results showed that these genes were significantly enriched in the Ras signaling pathway. The other module contained three mRNAs, i.e., C-C motif chemokine receptor 10 (CCR10), C-C motif chemokine ligand 19 (CCL19), and C-X-C motif chemokine ligand 13 (CXCL13).

Analysis of co-expression of lncRNAs and mRNAs. We identified 612 lncRNA-mRNA pairs that were significantly and differentially co-expressed, which consisted of 48 differential lncRNAs and 169 differentially expressed genes with co-regulatory relationships. The mRNAs with co-regulatory relationships were regarded as target genes for lncRNAs. To improve the understanding of the function of the differentially expressed lncRNAs that targeted genes, KEGG pathway enrichment analysis was carried out for the top 10 lncRNAs with the most number of target genes (n≥18). Nine pathways were identified, and we found that these lncRNAs were mainly involved in vascular smooth muscle contraction, the cGMP-PKG signaling pathway, and the oxytocin signaling pathway (Fig. 3B).

IncRNA-lncRNA co-regulation and functional synergistic analysis. An lncRNA-lncRNA co-regulation network was constructed based on the co-regulated target genes of two lncRNAs (Fig. 3C). This co-regulation network included 30 nodes (lncRNAs) and 161 edges (interactions). All of the lncRNAs included in this network were downregulated. In addition, we found that several lncRNAs had more co-regulatory relationships with other lncRNAs, such as MYLK-AS1 (n=19),...

<table>
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<th>Value</th>
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<tr>
<td>Age (mean ± SD)</td>
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</tr>
<tr>
<td>Sex (male/female)</td>
<td>154/125</td>
</tr>
<tr>
<td>Pathologic_M (M1/M0/NA)</td>
<td>39/189/51</td>
</tr>
<tr>
<td>Pathologic_N (N2/N1/N0)</td>
<td>46/70/163</td>
</tr>
<tr>
<td>Pathologic_T (T4/T3/T2/T1/NA)</td>
<td>38/191/43/6/1</td>
</tr>
<tr>
<td>Stage (IV/III/II/I/NA)</td>
<td>39/79/108/44/9</td>
</tr>
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<td>191/54/11/23</td>
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<tr>
<td>Survival status (dead/alive)</td>
<td>69/210</td>
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Table II. The top 10 nodes with a higher degree in the PPI network.

<table>
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<tr>
<th>mRNA</th>
<th>Upregulated/downregulated</th>
<th>Degree</th>
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<tbody>
<tr>
<td>MDFI</td>
<td>Up</td>
<td>20</td>
</tr>
<tr>
<td>MEOX2</td>
<td>Down</td>
<td>13</td>
</tr>
<tr>
<td>TRIP13</td>
<td>Up</td>
<td>11</td>
</tr>
<tr>
<td>MMP3</td>
<td>Up</td>
<td>10</td>
</tr>
<tr>
<td>NR3C1</td>
<td>Down</td>
<td>9</td>
</tr>
<tr>
<td>TFAP2A</td>
<td>Up</td>
<td>9</td>
</tr>
<tr>
<td>ADRB2</td>
<td>Down</td>
<td>8</td>
</tr>
<tr>
<td>COL1A1</td>
<td>Up</td>
<td>8</td>
</tr>
<tr>
<td>CRYAB</td>
<td>Down</td>
<td>8</td>
</tr>
</tbody>
</table>
ADAMTS9-AS2 (n=19), and FENDRR (n=19), where n represents the number of lncRNAs that had co-regulatory relationships with an lncRNA. Moreover, several lncRNA/lncRNA pairs had a larger number of co-regulated target genes, such as MYLK-AS1/RP11-1336O20.2 (n=57), ADAMTS9-AS1/MYLK-AS1 (n=53), ADAMTS9-AS1/RP11-1336O20.2 (n=53), and BVES-AS1/MYLK-AS1 (n=51), where n represents the number of co-regulated target genes of two lncRNAs.
On the other hand, functional synergistic network was constructed, as shown in Fig. 3D, consisting of 14 lncRNAs and 44 functional synergistic relationships. Among those functional synergistic lncRNA/lncRNA pairs, several lncRNA-lncRNA pairs had a strong functional synergistic effect, including FENDRR/RP11-1336020.2 (n=35), ADAMTS9-AS2/RP11-1336020.2 (n=32), MYLK-AS1/RP11-1336020.2 (n=28) and FENDRR/MYLK-AS1 (n=28), where n represents the number of GO terms in which the two lncRNAs were commonly involved.

**Prognosis-related lncRNA screening.** A K-M survival analysis of the differential expressed lncRNAs revealed 17 lncRNAs that were significantly associated with survival. The differential lncRNA expression data were randomly divided into a test set (75% of the total samples) and a validation set (25% of the total samples), and the R package randomForestSRC (version 2.4.0, https://cran.r-project.org/web/packages/randomForestSRC/index.html) was used for the RSF analysis. First, the RSF model was constructed through the test set to obtain the VIMP of each lncRNA in the model, and then the VIMPs were ranked from high to low. Furthermore, the lncRNAs sorted in order were included into the model to obtain the error rate of the current model; when the error rate was minimal, the corresponding lncRNA combination was regarded as the optimal combination of RSF models. The RSF model was reconstructed with the optimal combination, and the risk score was obtained by accumulating CHF values at different time points for each patient, and the threshold that was used to distinguish high and low risk was set as the median of the risk score.

An RF model was constructed in the test set (n=209) to obtain the VIMP of each lncRNA. According to the abovementioned method, when the top 14 variables with VIMP ranked in order were included into the model, the smallest error rate was achieved (Fig. 4A). We reconstructed the RSF model for the top 14 lncRNAs (AP000525.9, LL22NC03-N64E9.1, RP11-115D19.1, MAMDC2-AS1, DRAIC, BVES-AS1, PGM5-AS1, RP11-353N14.4, RP11-542B15.1, BMS1P17, LINC01234, etc.).
RP11-10A14.5, LINC01354, and BMS1P18), and obtained the risk score (range: 0.21-28.89) for each patient based on the CHFs of the above 14 prognosis-related lncRNAs. The samples were divided into two groups according to the median of risk score: a high-risk group (risk score ≥4.12, n=105) and a low-risk group (risk score <4.12, n=104). Additionally, the log-rank test demonstrated that the survival rate of the high-risk group was significantly lower than that of the low-risk group (P<0.0001).

Using the same model and parameters, we used a risk score of 4.12 as the threshold in the validation set and divided the samples into a high-risk group (n=52) and a low-risk...
The Cox model was constructed with the other known clinical indicators, including age, pathologic stage, pathology_T stage, pathology_N stage, and sex, and the risk score was based on the 14 lncRNAs. As shown in Table III, the univariate Cox regression analysis revealed that the risk score, age, pathologic stage, pathology_T stage, pathology_N stage, and pathologic_M stage were all significantly associated with OS in COAD patients. Moreover, multivariate Cox regression analysis revealed that the risk score was still significantly associated with OS even after adjustment with other clinical factors (P<0.001) (Table III). The results showed that the prognostic power of the 14-lncRNA signature was independent of other clinical variables for predicting the survival of patients with COAD. The hazard ratio (HR) of each variable is shown in Fig. 4C.

**Validation of several lncRNAs using RT-qPCR.** We evaluated the expression levels of four candidate lncRNAs by RT-qPCR analysis between the COAD tissues and paired adjacent normal tissues of 9 patients from a residual sample biobank of our hospital. The present study was approved by the Ethics Committee of the First Affiliated Hospital of Harbin Medical University (Harbin, China) and written informed consent was obtained from all the subjects. The tissues were collected between April, 2017 and September, 2017. The mean age of the patients was 68.3 years (range, 57-82 years) and the ratio of male to female was 7:2. As shown in Fig. 5, MYLK-AS1, BVES-AS1, ADAMTS9-AS1, and FENDRR were all down-regulated in COAD samples compared to those in control samples (P<0.05), which was consistent with the lncRNA expression profiles determined by the above mentioned bioinformatics analysis.

**Discussion**

In this study, we performed a bioinformatics analysis of the data from COAD and control samples downloaded from the TCGA database to investigate the global expression profile of lncRNAs and mRNAs in colon cancer and to identify diagnostic biomarkers for colon cancer. A total of 976 differentially expressed mRNAs and 169 differentially expressed lncRNAs were identified. MDFI and MEOX2 were the PPI network hubs. By analysis of target genes of differentially expressed lncRNAs, we found these lncRNAs were primarily involved in vascular smooth muscle contraction and the cGMP-PKG signaling pathway. Several lncRNA-lncRNA pairs had co-regulatory relationships or functional synergistic effects, including BVES-AS1/MYLK-AS1, ADAMTS9-AS1/MYLK-AS1, and FENDRR/MYLK-AS1. Moreover, this study identified a 14-lncRNA signature that could be used to predict the survival times for COAD patients.

MDFI is a known inhibitor of myogenic differentiation (35). MDFI regulates the Wnt signaling pathway (36), which plays a significant role in cancer development and progression (37). A recent study has demonstrated that MDFI was significantly methylated in colorectal cancer tissues (38). In the present study, MDFI was identified as a hub protein in the PPI network. These findings suggested that MDFI plays a critical role in the progression of COAD, which requires further validation. Moreover, MEOX2 was also identified as a hub protein in the

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</tr>
<tr>
<td>Sex (male/female)</td>
<td>0.62 (0.33-1.19)</td>
<td>0.66 (0.34-1.31)</td>
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</table>
| HR, hazard ratio; CI, confidence interval.
PPI network. Chen et al demonstrated that MEOX2 regulated nuclear factor-xB (NF-xB) activity in vascular endothelial cells and suggested MEOX2 as a possible molecular target for the anti-angiogenic therapy such as cancer treatment (39). This finding again provided evidence that MEOX2 may play a role in COAD carcinogenesis, while further experimental evidence is required for validation.

Vascular smooth muscle cells (VSMCs) are the main cell type of the vascular wall and have critical functions in vascular diseases (40). Vasculogenesis involves the de novo formation of blood vessels and occurs with the recruitment of VSMCs (41). Increasing evidence has demonstrated that vasculogenesis is critical to tumor growth and metastasis (42). On the other hand, it has been shown that nitric oxide increased the migration and invasion of colon cancer cells by upregulating matrix metalloproteinases (MMP)-2/9 via the cGMP-PKG-ERK signaling pathways (43). In addition, Li et al demonstrated that sulindac sulfide can selectively inhibit colon tumor cell growth by increasing intracellular cGMP levels and activating cGMP-PKG signaling (44). In this study, by analyzing the differential expression of IncRNA target genes, we found that these IncRNAs were mainly associated with vascular smooth muscle contraction and cGMP-PKG signaling pathways. The results of the present were consistent with those of previous studies (42-44), further suggesting the potentially significant roles of identified IncRNAs in the progression of COAD.

Many studies have been focused on the roles of IncRNAs in cancer initiation and progression (45,46). In the present study, through the IncRNA-IncRNA co-regulation and functional synergistic analysis, we identified several pairs that had significant co-regulatory relationships or functional synergistic effects, such as BVES-AS1/MYLK-AS1, ADAMTS9-AS1/MKYK-AS1, and FENDRR/MYLK-AS1. A previous study has reported that a decreased expression of FENDRR was able to predict poor prognosis in gastric cancer, and FENDRR was able to regulate gastric cancer cell metastasis by affecting fibronectin expression (47). FENDRR was also downregulated in the current study. At present, few studies have investigated the roles of ADAMTS9-AS1, BVES-AS1, and MYLK-AS1. In the present study, these three IncRNAs were all downregulated in COAD. Moreover, the downregulation of MYLK-AS1, BVES-AS1, ADAMTS9-AS1, and FENDRR in COAD tumors were further validated by RT-qPCR. Our results suggested that these differentially expressed IncRNAs may play critical roles in the development and progression of COAD. However, since studies on the interactions and biological functions of these IncRNAs are still lacking in patients with colon cancer, many issues need to be addressed in the future.

Gene expression profile-based prognostic IncRNA signatures for prognosis prediction in patients with cancer have been previously investigated (48-50). In the present study, we reported that the expression of 14 IncRNAs can be used to predict the clinical outcome of COAD. A K-M survival analysis of the identified differentially expressed IncRNAs and the risk score method were performed, resulting in 14 prognostic IncRNA markers, including BVES-AS1. Further survival analysis demonstrated a clear separation in the survival curves between the patient groups with high- or low-risk scores in the training or testing datasets, indicating the predictive power of the 14-IncRNA signature. The relationship between differentially expressed IncRNAs and the survival of colorectal cancer patients has been investigated in small samples using distinct approaches (51). For instance, Li et al analyzed the prognostic value of 21 IncRNAs in 30 patients with colorectal cancer using a PCR array (51). The current study used data based on RNA-seq technology using a larger cohort from the TCGA database. In addition, when taking other clinical factors into account, such as age, pathologic stage, pathology_T_stage, pathology_N_stage, pathology_M_stage, and sex, the multivariate Cox regression analysis revealed that the 14-IncRNA signature was independent of the conventional clinicopathological factors, and can be used as a risk factor for the prognosis of colon cancer (P<0.001). However, the functions of only a few IncRNAs have been indicated. A previous study reported that LL22NC03-N64E9.1 conferred an oncogenic function in human colorectal cancer via partially repressing KLF2 transcription (52). Sakurai et al showed that IncRNA DRAIC may have a tumor suppressive role (53). Thus, no thorough functional annotation data are available for the 14 prognostic IncRNAs in the current study. However, we validated the differential pattern of one of these 14 IncRNAs, BVES-AS1, in COAD patients compared with controls. Further functional annotation of these prognostic IncRNAs may increase our understanding of their biological implications in determining COAD prognosis.

In conclusion, our study identified several potentially significant genes (MDFI and MEOX2) and IncRNAs (BVES-AS1, MYLK-AS1, ADAMTS9-AS1, and FENDRR) in the progression of COAD. Moreover, a 14-IncRNA signature was identified that could be used to predict the survival times for patients with COAD. However, the data from the TCGA database were based on RNA-seq data, and other in vitro and in vivo experiments are needed to verify the current findings. These findings may lead to novel insights pertaining to patient prognosis and contribute to the development of novel therapeutic targets against colon cancer in future.

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Availability of data and materials

The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request.

Authors' contributions

DP conceived and designed the research, YX and ZZ acquired the data. YX and ZZ were involved in the analysis and interpretation of data. YZ performed the statistical analysis. YX was involved in drafting and revising the manuscript for important intellectual content. LZ and AZ were involved in the analysis and interpretation of data and performed the statistical analysis. All authors read and approved the final manuscript.
Ethics approval and consent to participate

The study was approved by the Ethics Committee of the First Affiliated Hospital of Harbin Medical University (Harbin, China). Informed consent was signed by the patients and/or guardians.

Consent for publication

Not applicable.

Conflict of interest

The authors declare that they have no conflict of interest.

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