Sex-specific analysis of microRNA profiles in HBV-associated cirrhosis by small RNA-sequencing

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Abstract
Liver cirrhosis represents an advanced stage of chronic liver disease and is associated with significant morbidity, mortality, and risk of cancer development. While sex disparity of liver diseases has been observed, understanding at a genetic level awaits more thorough investigation. In this study, we performed a sex-specific analysis of the microRNA (miR) profiles in hepatitis B virus (HBV)—associated cirrhosis by small RNA-sequencing using clinical tissue samples. Potential associated signaling pathways, downstream gene targets, and upstream regulators were highlighted by computational prediction analyses based on the differentially expressed miRs (DEmiRs). From our results, deregulation of miRs in cirrhosis showed a marked difference between males and females by the degree and pattern. Sixty-five (64 up-regulated, 1 down-regulated) and 12 (6 up-regulated, 6 down-regulated) DEmiRs were found in males and females, respectively, when compared with their respective control group. A number of DEmiRs were only observed in one sex but not the other. In addition, 26 DEmiRs were identified between cirrhosis female and cirrhosis male groups. Fatty acid biosynthesis pathway, extracellular matrix–receptor interaction, p53 signaling, Hippo signaling, tumor necrosis factor signaling, the forkhead box O signaling, as well as gene targets ribosomal protein S27 like, methyl CpG binding protein 2, and estrogen receptor 1, may contribute to the pathogenesis and biological behavior of cirrhosis in a sex-specific manner. Analysis of the Cancer Genome Atlas data set suggested a role of sex-specific DEmiRs in multistep hepatocarcinogenesis. Conclusion: Our findings illustrate that miR profiles in HBV-associated cirrhosis are distinct between the males and females and suggest a potential role of sex-specific biomarkers and molecular mechanisms in disease development and progression.
INTRODUCTION

Cirrhosis represents an advanced stage of chronic liver disease (CLD) and is a major cause per se for liver-related mortality. Chronic hepatitis B (CHB) virus infection is the top-ranking etiology for cirrhosis in Southeast Asia. The annual incidence of CHB to cirrhosis in Asians ranged from 0.1–2.8 per 100 person-years, with a 5-year cumulative incidence of 17%.\(^1\) The annual incidence of hepatic decompensation and liver-related mortality is 2–4 and 2.9 per 100 person-years in the Asian population, respectively.\(^1\) In 2016, cirrhosis was one of the most common indications (51%) for liver transplantation in our center.\(^2\) In addition, cirrhosis is a risk factor for primary liver cancer, with about 80% of hepatocellular carcinoma (HCC) associated with cirrhosis.

Activation of hepatic stellate cells and fibroblasts is a key cellular event during fibrogenesis. Transforming growth factor \(\beta\) (TGF-\(\beta\))/mothers against decapentaplegic homolog (SMAD) and WNT signaling cascades were implicated in liver fibrogenesis.\(^3\) At the genomic level, the mutational landscape has been illustrated using clinical samples in a recent report.\(^6\) In this regard, clues from the clinical and epidemiological perspectives may also provide insights on the dissection of molecular mechanisms in human diseases. Remarkably, sex disparity is characteristically observed in CLD and HCC. For example, autoimmune liver diseases and drug-induced liver injury showed a higher incidence in women than men.\(^5\) Conversely, disease progression of CHB was more rapid in men.\(^6,7\) Worse still, the male sex was identified as an independent risk factor for cirrhosis.\(^1\) Risk of disease-related death was reported to be higher in male than female patients.\(^8–10\) HCC by itself has a higher incidence in males than females with a 5–7:1 ratio.\(^9\) Among patients with cirrhosis, females have better outcome than males in terms of HCC incidence and overall survival.\(^11\) Consistently, the genetic content in hepatitis B virus (HBV)–related HCC was found to be different between the two sexes in terms of copy number variation.\(^12\) Hence, it is plausible that genetic events in CLD could be sex-specific throughout the multistep process.

These pieces of information made us speculate that while cirrhosis appears similar histologically, the molecular signatures may be different between the sexes. This is an important investigative question, as the divergent biological processes may infer a need for sex-specific monitoring and intervention during the multistep clinical course of CHB infection. Thus far, factors accounting for sex disparity are still poorly understood. Sex hormones were one of the first targets investigated for their contribution to the disparity. Male chronic HBV carriers with a higher androgen level progress to HCC at a faster rate.\(^13,14\) Because the HBV genome contains androgen responsive elements, androgen signaling pathway mechanistically promotes hepatocarcinogenesis via inhibition of the tumor suppressor gene.\(^15\)

In this study, we aimed at unmasking the potential difference in the microRNA (miR) profiles of HBV-associated cirrhosis between males and females. miR is a type of noncoding RNA well known for their epigenetic regulatory functions on gene expression, rendering it a potential useful biomarker. Deregulation of miR was identified from CHB to HCC.\(^16–18\) Moreover, miRs are therapeutic targets. Use of miR-21 inhibitor was reported to be effective in a preclinical model for fibrotic disease in the heart.\(^19\) Alteration of miR profiles in HCC was found to correlate with clinical parameters such as etiology and deregulation specific of signaling pathways.\(^20\) Interestingly, differential miR expression was observed between male and female patients with HCC.\(^21\) As far as cirrhosis is concerned, an enormous effort was put to analyze serum samples obtained from patients with CHB. With clinical tissue samples, a few studies demonstrated the deregulated expression of isolated or a small panel (up to 17) of miRs, and two studies examined the miR profiles by microarray.\(^22,23\) Thus far, deep-sequencing data from HBV-cirrhotic liver tissues are scarce, let alone the potential sex disparity in miR profiles is rarely characterized.

Cirrhosis represents a critical tipping point for investigation. Moreover, the growing body of evidence on regression of liver cirrhosis upon antiviral treatment\(^24\) necessitates a re-examination of the molecular pathogenesis of cirrhosis. Intervention to revert the genetic changes at this stage can also effectively prevent liver-related morbidity and mortality and the development of liver cancer, a deadly cancer despite advances in surgical intervention and targeted therapy. In this study, we carried out miR profiling by small RNA-sequencing (sRNA-seq) using clinical tissue samples from HBV-associated cirrhosis. Our analyses demonstrated that the miR profiles in HBV-associated cirrhosis displayed distinct patterns between males and females, and revealed potential sex-specific biomarkers and molecular mechanisms contributing to fibrogenesis and tumorigenesis.

METHODS

Clinical samples

Formalin-fixed paraffin-embedded (FFPE) clinical tissue samples were retrospectively retrieved from the archive in Department of Pathology, Queen Mary Hospital, Hong Kong. For the test groups, liver explant specimens with a diagnosis of HBV-associated cirrhosis from adult patients were used. Nontumoral liver tissue in liver metastasectomy specimens collected from adult patients with metastatic cancers were used as the control groups. Histological features of the liver
were subsequently selected for sRNA-seq (2 × 151 bp) PCR products ranging between 125 bp and 155 bp according to manufacturer's protocol (Sage Science). Purification and size selection using BluePippin platform allowed for effective separation of PCR products. Following purification, the prepared libraries were subjected to library preparation, which was done using NEBNext Multiplex Small RNA library Prep Set for Illumina (v6.0; New England Biolabs). The adaptor- ligated libraries were enriched by 15 cycles of polymerase chain reaction (PCR) followed by purification and size selection using BluePippin platform according to manufacturer's protocol (Sage Science). PCR products ranging between 125bp and 155bp were subsequently selected for sRNA-seq (2 × 151 bp) with the Illumina NovaSeq 6000.

**Bioinformatics and computational analyses**

Sequencing reads were assigned to each sample using bcl2fastq (Illumina). Number of reads, total throughput, and the quality score, as denoted by Q30, were determined for each sample. Sequencing reads were filtered for adapter sequence. Low-quality sequence, defined as reads with more than 5% unknown bases or more than 50% of bases with quality value less than 10, were excluded. Sequencing reads (≥15bp) were retained. Read alignments were performed using Bowtie2 (v2.2.5). Filtered reads were mapped to human mature miR sequence, and unmapped reads were mapped to hairpin miR sequence (miRBase v21). Reads aligned to reference genome were counted using HTSeq-count (v0.6.0) for each gene-biotype. Alignment and read counts were performed using default parameters with following exceptions: length of the seed substrings to align during multiseed alignment was set to 15; suppression of unpaired alignments as well as discordant alignments for paired reads; and suppression of unpaired reads to align against the reverse-complement (Crick) reference strand. Differential expression analysis of the sRNA-seq was done on allocated group comparison using EBSeq (v1.10.0). The investigators were blinded to the disease background of the group allocation. Differentially expressed miRs (DEmiRs) were defined as false discovery rate (FDR)<0.05 and |fold change (FC)|≥6. Hierarchical clustering of DEmiRs between study groups was generated using Pearson correlation (subtracted from 1), and the algorithm was performed by Clustvis.[25] Predicted miR targets and molecular pathways were generated using DIANA-TarBase v7 and DIANA-miRPath v3.0.[26,27] Transcription factor (TF)–miR regulatory networks were predicted using TFmiR v1.2.[26] Normalized expression of selected mature miRs (Log2[total RPM+1]) in HCC tissues was also obtained from the Cancer Genome Atlas–Liver Hepatocellular Carcinoma (TCGA-LIHC) cohort using the UCSC Xena platform.[29] Data were extracted from paired HCC tissue and the corresponding adjacent nontumoral liver tissue for analysis.

**Cell line models and reagents**

Human hepatic stellate cell line LX2 (SCC064) was purchased from MilliporeSigma. Immortalized human hepatocyte cell line MIHA was provided by Dr. Jayanta Roy-Chowdhury, Albert Einstein College of Medicine, New York. LX2 cells were maintained in Dulbecco's modified Eagle's medium (DMEM)–high glucose supplemented with 10% fetal bovine serum (FBS) and 1% penicillin–streptomycin (Gibco by Life Technologies). MIHA cells were maintained in DMEM-high glucose supplemented with 1 mM sodium pyruvate, 10% FBS, and 1% penicillin–streptomycin. All cells were cultured in a humidified incubator at 37°C with 5% CO₂. Cell line was routinely checked and tested negative for mycoplasma contamination before use.

**Cell transfection**

miR mimic for hsa–miR-137-3p (Assay ID: MC10513) and negative control mimic (Cat# 4464058) were purchased from Thermo Fisher Scientific. PcDNA-DEST47-GFP-GFP was a gift from Patrick Van Oostveldt (Addgene plasmid #36139).[30] pDEST-hMeCP2-GFP was a gift from Huda Zoghbi (Addgene plasmid #48078).[31] pCMV-hERalpha was a gift from Patrick Van Oostveldt (Addgene plasmid #36139). [30] pDEST47-GFP-GFP was a gift from Patrick Van Oostveldt (Addgene plasmid #36139). [30] pDEST-hMeCP2-GFP was a gift from Huda Zoghbi (Addgene plasmid #48078). [31] pCMV-hERalpha was a gift from Patrick Van Oostveldt (Addgene plasmid #36139). [30] pDEST47-GFP-GFP was a gift from Patrick Van Oostveldt (Addgene plasmid #36139). [30] pDEST-hMeCP2-GFP was a gift from Huda Zoghbi (Addgene plasmid #48078). [31] pCMV-hERalpha was a gift from Patrick Van Oostveldt (Addgene plasmid #36139). [30]
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Scientific) according to the manufacturer’s instruction. Transfected cells were harvested 24 h following transfection unless otherwise specified.

### RNA extraction from cell line

Total RNA was isolated from cell lines using RNAiso Plus (Takara Bio Inc.) according to the manufacturer’s protocol.

### Reverse-transcription and quantitative PCR

For detection of miR, extracted small RNA was reverse-transcribed using Applied Biosystems Taqman miR reverse-transcription kit in conjunction with a miR-specific stem-loop primer according to the manufacturer’s instructions (Thermo Fisher Scientific). Briefly, 10 ng of RNA was used as template, and primers from TaqMan microRNA assay (Cat# 4427975; Thermo Fisher Scientific) were used for each reverse-transcription reaction using Applied Biosystems ProFlex PCR System (Thermo Fisher Scientific). Complementary DNA samples were stored at −20°C before use. Quantitative PCR was performed using Taqman Universal PCR master mix (without UNG) and TaqMan probe and primers mix according to the manufacturer’s protocol (Thermo Fisher Scientific). Assay was done using Applied Biosystems StepOnePlus real-time PCR system (Thermo Fisher Scientific). TaqMan miRNA assays used were as follows: hsa-miR-200a-5p (ID: 001011), hsa-miR-137-3p (ID: 001129), hsa-miR-34c-5p (ID: 000428), and small nucleolar RNA, C/D Box 44 (RNU44) (ID: 001094).

Expression of miR of interest was normalized to RNU44. For protein coding genes and primary miR, RNA (500 ng) was reverse-transcribed using PrimeScript RT Master Mix (Takara). Quantitative PCR reaction was performed using Applied Biosystems Power SYBR Green PCR master mix (Thermo Fisher Scientific) on StepOne real-time PCR systems (Thermo Fisher Scientific) or the LightCycler 480 system (Roche Diagnostics). Primer sequences used are listed in Table S1. Expression of miRs or genes of interest was normalized to RNU44 or hypoxanthine-guanine phosphoribosyltransferase respectively. Relative expression of miRs or genes of interest was performed using the 2^−ΔΔCT or 2^−ΔΔCt method. Data are presented as mean±SD of values from at least three independent experiments.

### Statistical analysis

Two-tailed unpaired t test, Mann–Whitney U test, or Wilcoxon matched-pairs signed rank test was used to compare differences between two groups whenever appropriate. Kruskal-Wallis test was used for comparison across more than two groups. Statistically compared groups showed similar variance. Diagnostic performance of hsa-miR-200a-5p and hsa-miR-34c-5p was determined by area under receiver operating characteristic curve (AUROC). p <0.05 was considered as statistically significant. Statistical analyses were performed using GraphPad Prism V8.2.1 (GraphPad Software Inc.).

### RESULTS

#### Clinicopathological characteristics of the study cohort

Liver tissues from 22 patients with HBV-associated cirrhosis (11 males and 11 females) were studied. Representative microscopic images of cirrhotic liver tissues are shown in Figure 1A. The clinicopathological parameters of the four groups, namely cirrhosis from males (cirrhosis-M), cirrhosis from females (cirrhosis-F), control from males (ctrl-M) and control from females (ctrl-F), are provided in Table S2. Raw data obtained from sRNA-seq underwent standard pipeline for quantification of mature miRs expression in each sample followed by identification of DEmiRs. We made use of the pathway enrichment analyses, downstream gene targets prediction, and TF-miR regulatory network to investigate the biological significance and their associated molecular mechanisms in HBV-associated cirrhotic liver. A simplified workflow from sample preparation to sRNA-seq is shown in Figure 1B.

#### miR expression profiles in HBV-associated cirrhosis display distinct patterns between males and females

Mean total throughput from the 32 samples was 7.2 Gb (range: 5.9–12.5 Gb). Filtered read pairs aligned to reference genome were counted, and the numbers of filtered read pairs mapped to mature miR were comparable between groups (Figure 2A). Mean percentage of filtered reads of all 32 samples was 87.9% (range: 72.3%–96.0%), whereas the mean percentage of filtered read pairs mapped to mature miR was 14.9% (range: 2.9%–29.7%). Percentage of filtered read pairs mapped to mature miR, hairpin miR, piwi-interacting RNA, ribosomal RNA, and unmapped read pairs of each sample were listed in Table S3. Total number of mature miR identified from the cirrhosis-M, cirrhosis-F, ctrl-M, and ctrl-F groups were 1836, 1830, 1479, and 1451, respectively. DEmiRs were defined as FDR <0.05 and |FC|≥6. Between the ctrl-M and ctrl-F groups, no DEmiRs were identified. Between the cirrhosis-F and ctrl-F groups, 12 miRs (6 down-regulated, 6 up-regulated) were differentially expressed. Between the cirrhosis-M and ctrl-M
groups, 65 miRs (1 down-regulated, 64 up-regulated) were identified as DEmiRs. Twenty-six miRs were differentially expressed (23 down-regulated, 3 up-regulated in cirrhosis-F) between the cirrhosis-F and cirrhosis-M groups. By comparing all groups with cirrhosis with control samples, 45 DEmiRs (4 down-regulated, 41
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(A) Number of filtered miRNA

(B) Heatmap comparison of miRNA profiles

(C) Heatmaps of miRNA expression in different groups

(D) Venn diagram of miRNA overlap

(E) Relative expression of selected miRNAs

(F) ROC curves for miRNA detection
up-regulated) were identified (Figure 2B). Heatmaps with hierarchical clustering were constructed based on DEmiRs identified between specific groups (cirrhosis-F and ctrl-F, cirrhosis-M and ctrl-M, and cirrhosis-F and cirrhosis-M) (Figure 2C). The top 10 DEmiRs found in the cirrhotic liver group when compared with the respective control of each sex are summarized in Table 1. To understand whether cirrhotic livers from both sexes share a list of similar DEmiRs, we compared the lists of DEmiRs and found that only four miRs (miR-24-2-5p, miR-34c-5p, miR-200a-5p, and miR-934) were commonly deregulated in both groups with cirrhosis. These DEmiRs were up-regulated in the cirrhotic liver groups when compared with their respective control groups. To further highlight, most of the DEmiRs were specifically deregulated in one sex only (Figure 2D). The numbers of DEmiRs and degree of miR alteration in terms of FC in the male group appeared to be much higher than that in the female group. A cross comparison between the lists of DEmiRs identified in cirrhosis-M versus ctrl-M, cirrhosis-F versus ctrl-F, and cirrhosis (M+F) versus ctrl (M+F) was also performed. Analyses

| miR ID     | miR name         | Fold change | Regulation type | FDR   |
|------------|------------------|-------------|-----------------|-------|
| MIMAT0000686 | hsa-miR-34c-5pa  | 22.327      | Up              | 0.000 |
| MIMAT0019970 | hsa-miR-4796-5p  | 11.334      | Down            | 0.041 |
| MIMAT004977  | hsa-miR-934a     | 10.879      | Up              | 0.005 |
| MIMAT0027661 | hsa-miR-6880-3p  | 8.491       | Down            | 0.000 |
| MIMAT0004496 | hsa-miR-23a-5p   | 8.075       | Up              | 0.011 |
| MIMAT0028118 | hsa-miR-7110-3p  | 7.265       | Down            | 0.000 |
| MIMAT0027591 | hsa-miR-6845-3p  | 7.239       | Down            | 0.000 |
| MIMAT0027450 | hsa-miR-6775-5p  | 7.095       | Down            | 0.001 |
| MIMAT0001620 | hsa-miR-200a-5pa | 6.565       | Up              | 0.021 |
| MIMAT0004497 | hsa-miR-24-2-5pa | 6.495       | Up              | 0.047 |

| miR ID     | miR name         | Fold change | Regulation type | FDR   |
|------------|------------------|-------------|-----------------|-------|
| MIMAT0000429 | hsa-miR-137     | 55.552      | Up              | 7.91E-05 |
| MIMAT0019873 | hsa-miR-4742-3p | 38.201      | Up              | 5.21E-03 |
| MIMAT0001620 | hsa-miR-200a-5pa| 32.236      | Up              | 1.27E-08 |
| MIMAT000281  | hsa-miR-224-5p   | 28.706      | Up              | 2.54E-09 |
| MIMAT0000686 | hsa-miR-34c-5pa  | 23.782      | Up              | 1.62E-06 |
| MIMAT0004428 | hsa-miR-135a-5p | 23.676      | Up              | 1.76E-05 |
| MIMAT004977  | hsa-miR-934a    | 23.390      | Up              | 1.06E-04 |
| MIMAT0000416 | hsa-miR-1-3p    | 17.146      | Up              | 4.04E-04 |
| MIMAT0000454 | hsa-miR-184     | 16.407      | Up              | 3.27E-03 |
| MIMAT0020924 | hsa-miR-642a-3p | 16.258      | Up              | 2.20E-04 |

*aCommonly deregulated miR identified in males and female.*
showed that 27 (of 64 DEmiRs from male-only comparison and of 41 DEmiRs from M+F comparison) and 5 (of 6 DEmiRs from female-only comparison and of 41 DEmiRs from M+F comparison) were commonly up-regulated. None (of 1 DEmiRs from male-only comparison and of 4 DEmiRs from M+F comparison) and 1 (of 6 DEmiRs from female-only comparison and of 4 DEmiRs from M+F comparison) were commonly down-regulated (Figure 2D). These findings suggest that HBV-associated cirrhosis bears distinct deregulated miR expression patterns in males and females. Expression of miR-200a-5p and miR-34c-5p in the same cohort was evaluated by quantitative PCR (Figure 2E). With AUROC analysis, both miR-200a-5p and miR-34c-5p appeared to be a good marker (AUROC = 0.832 and 0.718, respectively) to differentiate HBV-associated cirrhosis from controls (Figure 2F).

**Prediction analyses highlight potential sex-specific molecular mechanisms and regulators in HBV-associated cirrhosis**

First, Kyoto Encyclopedia of Genes and Genomes pathway enrichment analyses were performed based on the 65 DEmiRs and 12 DEmiRs identified in cirrhosis-M and cirrhosis-F, respectively (cirrhosis vs. control groups in each sex). Gene targets of these DEmiRs were derived from DIANA-Tarbase v7.0, which gene targets and miRs interactions were experimentally validated, and subsequent pathway enrichment analyses were conducted using DIANA-miPath v3 (Figure 3A). Although certain pathways are commonly deregulated in both males and females, including proteoglycans in cancer, cell cycle, viral carcinogenesis, adherens junction, and TGF-β signaling pathway, some human pathological processes related pathways were only highlighted in one sex. For example, fatty acid biosynthesis was highlighted in females, which was not identified in males. Contrarily, extracellular matrix–receptor interaction, p53 signaling pathway, Hippo signaling pathway, tumor necrosis factor signaling pathway, and the forkhead box O signaling pathway, for example, were specifically enriched in males. Apart from revealing the “sex-specific” DEmiR-associated molecular pathways enriched in the groups with cirrhosis when compared with their respective control groups, DEmiRs and the biological significance of these DEmiRs in the cirrhotic liver between females and males (cirrhosis-F vs. cirrhosis-M) were also determined. This comparison was made to evaluate the molecular differences between females and males with the same disease. The top five DEmiRs (miR-486-3p, miR-216a-5p, miR-144-5p, miR-2681-3p, and miR-137) were all down-regulated in the cirrhotic-F group (Table 2). We further identified downstream effectors of these five DEmiRs using DIANA-Tarbase v7. Ribosomal protein S27 like (RPS27L) was the sole target of three DEmiRs, namely miR-137, miR-144-5p, and miR-486-3p (Figure 3B). Liver disease–associated TF-miR network was constructed to determine the upstream regulators of identified DEmiRs (cirrhosis-F vs. cirrhosis-M) using TFmiR database. Methyl CpG binding protein 2 (MECP2) and estrogen receptor 1 (ESR1) were highlighted as potential transcription regulators of the DEmiRs (MECP2 on miR-184 and miR-137; ESR1 on miR-18a, miR-106a, miR-18b, and miR-20b). Hepatic stellate cells are recognized to be a major cell type implicated in liver fibrogenesis with their role in extracellular matrix production during liver injury and regeneration. In an attempt to verify the prediction, MECP2 and ESR1 were each over-expressed in LX2 cells, and expression of the respective primary miRs were examined. Up-regulation of primary miR-137 and miR-184 was observed following overexpression of MECP2 (at 24 h after transfection), whereas overexpression of ESR1 did not alter all predicted miRs (Figure 3C). To evaluate the involvement of normal hepatocytes in this regulation, MIHA cells were also adopted to examine the expression of primary miR-137 and miR-184. The results showed that expression of these miRs was not significantly altered in MECP2-overexpressing MIHA cells (Figure S1).

**Sex-specific DEmiRs in HBV-associated cirrhosis carry potential functional significance**

As some DEmiRs were only identified in the male but not the female group (Table 1), we questioned what
functions these sex-specific DEmiRs may exert in the liver. Hepatic stellate cells are a major cell type implicated in liver fibrogenesis; thus, miR-137-3p mimic was used in LX2 cells to study its functional effects in vitro. Transfection efficiency was first confirmed by quantitative PCR (Figure 4A). We quantified the expression of myofibroblast marker smooth muscle actin and fibrogenic markers collagen type I (COL1A1), COL1A2, TGF-β1, TIMP1, and TIMP2 by quantitative

**TABLE 2** Top 5 DEmiRs identified in cirrhosis female group compared with cirrhosis male group

| miR ID | miR name       | Fold change | Regulation type | FDR      |
|--------|----------------|-------------|-----------------|----------|
| MIMAT0004762 | hsa-miR-486-3p | 24.251      | Down            | 1.39E-09 |
| MIMAT0000273 | hsa-miR-216a-5p | 19.331      | Down            | 1.62E-08 |
| MIMAT0004600 | hsa-miR-144-5p | 17.777      | Down            | 7.16E-08 |
| MIMAT0013516 | hsa-miR-2681-3p | 16.279      | Down            | 0.006    |
| MIMAT0000429 | hsa-miR-137   | 14.855      | Down            | 1.93E-05 |

Note: In the present study, we demonstrated the distinct miR expression patterns in HBV-associated cirrhosis between sexes using sRNA-seq. Prediction of upstream and downstream regulators based on the DEmiRs suggests sex-specific molecular mechanisms implicated in the pathogenesis and biological behavior of liver cirrhosis.

**FIGURE 4** Functional significance of sex-specific DEmiRs was explored. (A) Validation of miR-137-3p expression in miR-137-3p mimic (1 nm) transfected LX2 cells. Relative expression data are normalized to RNU44 (n = 3); two-tailed unpaired t test. Data are presented as mean ± SD. (B) mRNA expression of fibrogenic genes smooth muscle actin alpha 2, collagen type I alpha 1, COL1A2, TGF-β1, TIMP1, and TIMP2 following transfecting miR-137-3p mimic (1 nm) in LX2 cells (n = 3). Relative expression data are normalized to HPRT; two-tailed unpaired t test. Data are presented as mean ± SD. (C) Expression of female-specific DEmiRs (miR-23a-5p [n = 19], miR-4796-5p [n = 17], and miR-7110-3p [n = 6]) in female paired hepatocellular carcinoma (HCC) tissues and nontumoral liver tissues obtained from The Cancer Genome Atlas–Liver Hepatocellular Carcinoma (TCGA-LIHC) data set; Wilcoxon matched-pairs signed-rank test. Data are presented as median with interquartile range (IQR). (D) Expression of male-specific DEmiRs (miR-224-5p [n = 26], miR-135a-5p [n = 16], miR-184 [n = 16], miR-1-3p [n = 25], and miR-4742-3p [n = 19]) in male paired HCC tissues and nontumoral liver tissues obtained from the TCGA-LIHC data set; Wilcoxon matched-pairs signed-rank test. Data are presented as median with IQR; *p < 0.05, **p < 0.01, and ****p < 0.0001. Abbreviations: NT, nontumoral liver tissues; T, HCC tumor tissues
PCR upon transfecting miR-137-3p mimic in LX2 cells (Figure 4B). From the results, the fibrogenic genes were not significantly altered in the miR-137-3p mimic–transfected cells. With further experiments, we found that expressions of fibrogenic genes smooth muscle actin alpha 2, COL1A1, TGF-β1, and TIMP2 were up-regulated at 48 h (but not at 24 h) following overexpression of MECP2, whereas expressions of miR-137 and miR-184 were not significantly altered at this time point (Figure S2). These findings suggest that miR-137-3p per se may not be sufficient to mediate the effect of MECP2 on hepatic stellate cells. To further explore the potential significance of the sex-specific DEmiRs in multistep hepatocarcinogenesis, we analyzed the expression of the top-listed DEmiRs identified in the cirrhosis-M versus control-M, as well as cirrhosis-F versus control-F comparison, from TCGA-LIHC data set. Some analyses were not feasible due to the small number of paired samples (which include miR-6880-3p, miR-6845-3p, miR-6775-5p, miR-137, and miR-642a-3p). Among the three female-specific DEmiRs, miR-23a-5p, miR-4796-5p and miR-7110-3p showed no significant difference in female HCC tumor tissues when compared with nontumoral liver tissues (Figure 4C). Among the five male-specific DEmiRs, miR-224-5p, miR-135a-5p and miR-184 (all up-regulated in cirrhosis vs. control) were up-regulated in male HCC tissues compared with nontumoral liver tissues, whereas miR-1-3p and miR-4742-3p showed no significant difference between groups (Figure 4D).

**DISCUSSION**

In this study, by using next generation sequencing, we performed a sex-specific analysis of the miR profiles with the liver tissue samples from HBV cirrhosis. Our findings showed that the deregulated miR profiles in male and female patients with cirrhosis are relatively distinct. The degree of deregulation was higher in males versus females as from the number of differentially expressed genes and number of predicted signaling pathways. Most of the DEmiRs in cirrhotic male samples were up-regulated, whereas half in the female counterpart were down-regulated. Common DEmiRs in both sexes constitute a minority. Our further computational and bioinformatics analyses also highlighted several signaling pathways, downstream effectors, and upstream regulators that potentially play a key role in the pathogenesis of liver cirrhosis in each sex. Analysis of the TCGA-LIHC data set suggested a role of sex-specific DEmiRs in multistep hepatocarcinogenesis.

We compared our findings with reports from expression analysis using clinical samples collected from patients with HBV. Among the DEmiRs in cirrhosis-M versus ctrl-M, cirrhosis-F versus ctrl-F, as well as cirrhosis-F versus cirrhosis-M groups, only a small proportion have been reported in the studies on expression analysis, especially in the cirrhosis-F versus cirrhosis-M groups. With reference to two previous reports on miR profiling of HBV liver tissues by microarray, some of the DEmiRs were also identified in our current study, which mostly belong to the cirrhosis-M versus ctrl-M groups (for example, miR-150-5p, miR-155-5p, miR-200a-3p, miR-200b-3p, miR-338-3p, and miR-642a-3p as revealed by Chen et al. in the S4 versus S0 group; and miR-96-5p, miR-1-3p, miR-10b-5p, miR-218-5p, and miR-23a-3p as shown by Singh et al. in their analysis between advanced fibrosis and early fibrosis groups). On comparing the DEmiRs identified in different group comparisons, the pattern of DEmiRs identified in cirrhosis-F versus cirrhosis-M was not readily discernible in the cirrhosis-M or cirrhosis-F versus the respective control group in the same sex. This was likely due to the different reference groups used in the comparisons (cirrhotic-M and ctrl-M/ctrl-F, respectively).

Some of the top-listed DEmiRs identified in our current study were reported to show profibrotic functions using cell line models (e.g., miR-34c[33,34]). The top-ranking male-specific miR identified in our study, miR-137, was not reported to exert any pro-fibrogenic functions thus far. Consistently, expression of fibrogenic genes was not altered in LX2 cells following miR-137 transfection from our in vitro experiments. Likewise, miR-486-3p and miR-216a-5p, identified from the cirrhosis-F versus cirrhosis-M analysis, were not reported to result in liver fibrosis from functional studies. Of note, while miR-200a was up-regulated in LX2 cells following miR-137 transfection and the functional phenotype may rely on an interplay between multiple markers.

Fatty acid biosynthesis pathway was identified as a deregulated signaling pathway in cirrhosis-F group (vs. ctrl-F) but not in the male counterpart, suggesting that fatty acid synthesis may still play an important role in liver fibrogenesis among female patients with primary etiology other than fatty liver disease. Apart from nonneoplastic conditions, fatty acid biosynthesis pathway has been found to facilitate tumorigenesis via multiple mechanisms. Moreover, target prediction analysis showed that RPS27L is a common target for the DEmiRs being down-regulated in cirrhotic-F when compared with the cirrhotic-M group. RPS27L encodes...
a ribosomal protein and was reported to suppress tumorigenesis by directly regulating p53.\textsuperscript{[41]} These findings suggest that RPS27L may be a factor accounting for lower malignant transformation potential in female cirrhosis. The identified DEmiRs by comparing cirrhosis-F versus cirrhosis-M highlighted ESR1 and MeCP2 as potential upstream master regulators. From our experiments, we could validate the regulation of DEmiRs following overexpressing MECP2 but not ESR1 in LX2 cells. Of note, MeCP2 was reported as a regulator of miRs/genes involved in liver fibrosis.\textsuperscript{[42–44]} Regarding results on ESR1, we speculate that its effect on miR regulation may also depend on estrogen level in the cells. Because LX2 is a male cell line, the regulation of ESR1 on sex-specific DEmiRs may not be fully characterized by manipulation of gene targets alone.

Our study could provide some insights on sex-specific biomarkers to monitor the progression of liver fibrosis (e.g., miR-137, miR-4742-3p, and miR-224-5p for male patients and miR-4796-5p, miR-6880-3p, and miR-23a-5p for female patients). The great magnitude of miRs and signaling pathway derangement may explain the high risk of liver fibrosis progression in males. In addition, the differential predicted upstream regulators and downstream effectors between the two sexes, such as fatty acid biosynthesis pathway, RPS27L, ESR1 and MeCP2, may offer insights in explaining the sex difference in disease course and in devising antifibrotic therapeutics for different patient groups. By referring to the expressions of some DEmiRs from the TCGA-LIHC data set, we might be able to gather some insight on the functions of sex-specific DEmiRs. For the female-specific DEmiRs, none (of the 3) female-specific DEmiRs was deregulated in female HCC, whereas 3 (of the 5) up-regulated male-specific DEmiRs identified in our study were also up-regulated in male HCC. While miR-224-5p and miR-135a-5p were also up-regulated in female HCC versus nontumor tissue, concordant changes were not observed in the cirrhosis versus normal comparison. These findings suggest that male-specific DEmiRs may play a functional role in the multistep events of tumor development. One merit of our study is the use of next generation sequencing in the identification of miRs from clinical samples. In a previous report, miR changes were studied with next generation sequencing of hepatitis C liver tissues.\textsuperscript{[45]} For HBV, miR profiling was mostly performed with microarray method in previous studies.\textsuperscript{[22,23]} A limitation of our study is the cohort size, which was largely due to the number of female cirrhosis cases falling into the inclusion criteria. Clinicopathological parameters (e.g., age, albumin, bilirubin levels) may constitute confounding effects in the analysis of DEmiRs. Expansion of the study cohort or inclusion of samples with other primary liver disease etiologies for group comparison could allow a more comprehensive analysis. Another limitation is the lack of validation of target gene predictions by real-time quantitative PCR. The samples were extracted from FFPE tissues and contained fragmented RNAs with a major peak below 200 nucleotides (range: 40%–96% of total). The fragmented size rendered gene-expression quantification suboptimal from a technical point of view.

Sex disparity has been observed in some human diseases, both neoplastic and nonneoplastic. The disparity includes incidence, pathogenic mechanisms, and biological behavior. It is therefore an important aspect to be addressed in the study of human diseases. Results from our study elaborated the commonly deregulated miRs and sex-specific DEmiRs in HBV-associated cirrhosis. The former potentially represent some critical genetic events in cirrhosis, while the latter likely explain the sex disparity in the susceptibility and biological cause of CLD. Identification of these miRs together with their molecular mechanisms will help enrich our understanding of the genetic complexity of HCC. Based on our results, sex-specific serum miR markers could potentially be identified and validated for disease monitoring and risk stratification in patients with HBV. Despite the disparities between hepatic and blood miR expressions from some studies, some miRs were identified to show concordant differential expression patterns in tissue and blood samples from patients with liver fibrosis. To highlight, a positive correlation was observed between the expression of hepatic miR-122, miR-222, miR-33a, and miR-571 with corresponding serum levels.\textsuperscript{[46–50]} Hence, evaluation of a tissue-serum expression correlation for specific miRs is indicated. Moreover, sex-specific molecular targets could be of value for further studies to delineate the potential points of intervention more precisely, which hopefully will contribute to improve the prognosis of CLD.

**AUTHOR CONTRIBUTIONS**

Data acquisition: Kristy Kwan-Shuen Chan, Kwan-Yung Au, Wai-Ching Fung, Cheuk-Yan Wong, Albert Chi-Yan Chan, and Regina Cheuk-Lam Lo. Data analysis: Kristy Kwan-Shuen Chan, Kwan-Yung Au, and Regina Cheuk-Lam Lo. Data interpretation: Kristy Kwan-Shuen Chan, Kwan-Yung Au, and Regina Cheuk-Lam Lo. Manuscript draft: Kristy Kwan-Shuen Chan and Regina Cheuk-Lam Lo. Statistical analysis: Kristy Kwan-Shuen Chan. Study concept, design, and supervision: Regina Cheuk-Lam Lo.

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CONFLICT OF INTEREST
Nothing to report.

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Supporting Information
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