ORIGINAL RESEARCH Transcriptomic Analysis of Trigeminal Ganglion and Spinal Trigeminal Nucleus Caudalis in Mice with Inflammatory Temporomandibular Joint Pain

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Background: Persistent facial pain heavily impacts the quality of life in patients with temporomandibular joint (TMJ) disorders. Previous studies have demonstrated that long non-coding ribonucleic acid (IncRNA) is an important regulator of pain. In this study, we aimed to analyze lncRNA expression in the whole transcriptome of trigeminal ganglia (TG) and spinal trigeminal nucleus caudalis (Sp5C) in a chronic inflammatory TMJ pain mouse model.

Methods: Chronic inflammatory TMJ pain was induced by intra-TMJ injection of complete Freund's adjuvant (CFA). Mouse TG and Sp5C tissues were harvested on day 4 after CFA injection. The lncRNA expression patterns in the whole transcriptome of TG and Sp5C were profiled with RNA sequencing.

Results: We observed that 38 lncRNAs and 849 mRNAs were differentially expressed after CFA treatment. Gene Ontology and Kyoto Encyclopedia of Genes and Genomes analysis further revealed relationships among those differentially expressed lncRNAs and mRNAs and their potential functions. Specific categories of biological process, cellular processes, and molecular function of the differentially expressed transcripts were ascertained.

Conclusion: Our results suggest that lncRNA expression in the whole transcriptome of trigeminal nociceptive system could contribute to the molecular mechanisms that underlie chronic inflammatory TMJ pain.

Keywords: temporomandibular joint pain, lncRNAs, RNA sequencing, trigeminal ganglia, spinal trigeminal nucleus caudalis

Introduction

Temporomandibular joint disorders (TMDs) affect about 10–15% of the total population.¹ The recent clinical study has shown that 54-68% of the patients with TMDs experience TMJ pain.² However, the molecular mechanisms underlying TMJ pain are not fully understood.

In recent years, long non-coding RNAs (lncRNAs) have been studied in different pain models,^{3–7} but it remains unclear whether lncRNA expression is involved in TMJ pain. It has been reported that lncRNA HOTAIR in chondrocytes contributes to chondrocyte destruction in TMJ osteoarthritis.⁸ In trigeminal neuropathic pain induced by chronic constriction injury of the infraorbital nerve, lncRNA uc.48+ in the trigeminal system plays an important role in pain transmission.9 Our previous study indicates that altered gene expression (such as tumor necrosis factor-alpha) in the trigeminal ganglia (TG) and spinal trigeminal nucleus caudalis (Sp5C) is critical for chronic inflammatory TMJ pain induced by intra-TMJ injection of complete Freund's adjuvant (CFA).¹⁰ Thus, examining the expression of lncRNAs and relevant mRNAs in trigeminal nociceptive system could advance our understanding of the molecular mechanisms for TMJ pain.

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In the present study, we performed RNA sequencing-based lncRNA profiling analysis in TG and Sp5C tissues of mice with CFA-induced chronic inflammatory TMJ pain. We aimed to identify the dysregulated trigeminal lncRNAs and provide new insight into the lncRNA-mediated regulatory network of TMJ pain. The genes we identified in this study may be targeted to develop a novel gene therapy for such pain in patients with TMDs.

Materials and Methods

Animals

C57BL/6 male mice (18–20 g) from Zhengzhou University Laboratory Animal Center (Zhengzhou, Henan, China) were used. The mice were housed under 12 h light–dark condition with water and food available ad libitum. The mice were randomly assigned to CFA and control groups with equal number in each group. We made all efforts to minimize animal suffering and reduce the number of animals we used in this study. All experiments were approved by the Animal Ethics Committee of Zhengzhou University (Animal Protocol Number: 2021328). We carried out these experiments in accordance with the National Institutes of Health Guide for the Care and Use of Laboratory Animals.

Chronic Inflammatory TMJ Pain Model

We injected CFA (10 μ L, 5 mg/mL, Chondrex. Inc.) into the superior joint space of the mouse TMJ bilaterally to induce chronic inflammatory TMJ pain as described in our previous study.¹⁰ Saline was used as the vehicle control.

Orofacial Mechanical Hypersensitivity Test

The calibrated von Frey filaments were used to measure orofacial mechanical pain as described previously.¹⁰ We placed each mouse in a 10-cm-long plexiglass cylinder and the mouse was allowed to poke out their heads but could not turn around in the cylinder. After 60-min acclimation, we applied the von Frey filaments to the trigeminal nerve V3 branch-innervated facial skin area. A positive stimulation-induced response was defined as a sharp withdrawal of the mouse head. Each filament was applied five times for 1-s stimulation, each at 10-s intervals. We started the test with the lowest force of von Frey filament and continued in ascending order. The head withdrawal threshold was determined as the force at which the positive response was produced in three of five stimuli.

Tissue Harvest and RNA Extraction

The mice were sacrificed under isoflurane anesthesia on day 4 after CFA or saline injection, and whole Sp5C and TG tissues were harvested. We selected day 4 post-CFA because the head withdrawal threshold reached the lowest level, which indicates the strongest TMJ pain at that time point. According to the manufacturer's instructions, all RNA content was extracted from the TG and Sp5C tissues using a TRIzol RNA Purification Kit (Invitrogen, Carlsbad, Canada). 1% agarose gel was used to check degradation and contamination in RNA. Nano-Photometer[®] spectrophotometer (Implen, Inc., USA) was utilized to evaluate isolated RNA's purity. A Qubit[®] RNA Assay Kit (Life Technologies, USA) was used for RNA concentration measurement, and quality of isolated RNA was checked with Bio-analyzer 2100 system (Agilent Technologies, USA) with an RNA Integrity Number (RIN) at 8.7.

Library Preparation and RNA Sequencing

For each sample, 3 µg of RNA was used as the input material. Initially, Epi-center Ribo-zero[™] rRNA Removal Kit was used to isolate rRNA, and ethanol precipitation was performed to obtain rRNA-free residues. Afterward, we used NEBNext[®] Ultra[™] Directional RNA Library Prep-Kit for Illumina[®] to prepare genomic libraries with depleted rRNA according to the manufacturer's instructions. NEB-Next First-Strand Synthesis Reaction Buffer (5X) was used for the fragmentation along with divalent cations under elevated temperature range. M-MuLV Reverse Transcriptase, Random hexamer primer, DNA polymerase-I, RNase-H was used to synthesize the first and second cDNA strands. DUTPs replaced all dTTPs in the reaction buffer, and the remaining reads were converted into blunt ends using polymerase/ exonuclease proceedings. Adenylation of 3'-ends of DNA reads was carried out, and then NEBNext-Adaptor ligated to hairpin-loop structures was prepared for hybridization procedure. Using AMPure XP system (Beckman Coulter, Beverly,

USA), we purified the library of fragments to select 150 to 200 bps reads. Afterward, 3 µL of USER Enzyme was used along with adaptor-ligated, optimized cDNA at a temperature of 37°C for about 15 min, followed by 5 min at 95°C. Phusion High-Fidelity DNA polymerase enzyme, Index (X) primers, and Universal PCR primers were used for PCR reaction. Next, gene products were refined, and the quality of the library was evaluated on Agilent Bio-analyzer 2100 apparatus. Novogene Bioinformatics Technology was used for whole transcriptomic libraries, and deep sequencing was performed. Per the manufacturer's instructions, index-coded sample clustering was proceeded on cBot Cluster-Generation System utilizing TruSeq PE Cluster Kit v3-cBot-HS (Illumina, USA). After generating cluster, Illumina HiSeq 4000 system was used for sequencing all libraries, and paired-end 150 bp reads were made. We have deposited our sequencing data in Gene Expression Omnibus (GEO) and the GEO accession number is GSE188922.

Quality Control, Reference Genome Mapping, Transcriptome Assembly, and Transcripts Predicted with Coding Potential

Perl in-house scripts were compiled for raw reads data of fastq format. Clean reads were acquired after eliminating lower quality reads, which are adapters and contain ploy-N. After that, Q20 value, Q30 value, and the clean data were evaluated. All downstream analyses were based upon refined data having higher quality. Referred species genomic file (.fna) and genetic model annotation file (.gtf) were retrieved directly by the ftp genome web-server. Bowtie2 version 2.2.8 software was used for the development of referred genome index against target genes, and refined reads having paired-end were assembled against the downloaded reference genome using HISAT2 (version 2.0.4).¹¹ HISAT2 was executed along "RNA-stranded-ness RF" default parameters. Next, these mapped-reads for all samples were aligned using String-Tie tool (version 1.3.1)¹² under a referenced dependent method. String-Tie tool utilizes an advanced networking flow algorithm along with an additional de-novo assembly step to align and quantify fuller-length transcripts that represent extra spliced variants for all gene loci. Transcripts were predicted to have coding putative by the following procedures: 1) Coding/Non-Coding-Index (CNCI, version 2)¹³ with default parameters was used; 2) Coding Putative Calculator (CPC, 0.9-r2),¹⁴ NCBI eukaryotes protein database were used with a pre-set e-value of "1e-10" in our analysis; 3) Pfam Scan (version 1.3), default parameters of -E 0.001 –domE 0.001¹⁵ were used; 4) Phylogenetic codon substitution frequency (PhyloCSF, version 20121028),¹⁶ a multi-species genome sequence alignment set, was built and executed, and genes that do not have the coding potential were chosen as our query cluster of lncRNAs.

Quantification and Analysis of Differential Expression of Genes

Fragments-per-kilo-base of exon-per-million fragments mapped (FPKM) is computed based upon the length of fragments and mapped reads count against each fragment. For the evaluation of FPKM values for lncRNAs along with putative coding genes, a tool Cuffdiff (version: 2.1.1) is utilized for all samples.¹⁷ FPKM values for all genes were executed by the collective sum of all FPKM values for all gene group transcripts. All transcripts are referred to as differentially expressed (DE) genes that have an adjusted *P* value of less than 0.05.



Figure I Intra-TMJ injection of CFA induces inflammatory TMJ pain in mice. CFA injection significantly decreased the head withdrawal threshold in trigeminal nerve V3 branch-innervated skin areas compared to the saline control group (*p < 0.05, ***p < 0.001 vs the saline control group at corresponding time points). The decreased head withdrawal threshold started from day I post-CFA injection and lasted at least for 10 days, and the lowest head withdrawal threshold was shown on day 4 after CFA injection. n = 6 for each group.

Analyses of Gene Ontology (GO) and Kyoto-Encyclopedia of Genes and Genomes (KEGG)

GO analysis of lncRNA query genes or DE genes was conducted with GO sequencing. The R-based package, which is used for gene length biases, was verified and corrected.¹⁸ GO referred to as corrected *P* value <0.05 was assumed to be potentially enriched from DE genes. KEGG is a database resource that can be used to understand different biological



Figure 2 General information regarding the quality of and correlations of the data. (A) Clean reads as a percentage of raw reads for each sample. (B) Pearson correlation analysis results.

systems and pathways.¹⁹ The KOBAS tools were utilized for the statistical enrichment test of lncRNA or DE genes and target genes in KEGG pathways.²⁰

IncRNA Target Gene Prediction

PhyloFit tool was used for the computation of all phylogeny models for conserved or non-conserved reads in different organisms of species. After that, we evaluated a conservative scoring set in coding genes and lncRNAs. Functional annotation of reference mRNAs in *cis/trans* was used to predict query genes of lncRNAs. We scanned 100 kb upstream and downstream of lncRNA for putative *cis* mRNAs. Co-expressed mRNA sequences of lncRNAs along Pearson correlation-coefficients under/above 0.95 and above 100 kilo-base in genome distances from the lncRNA and situated upon alternative chromosomes were referred to as genes that are *trans*-regulated.

Protein–Protein Interaction Network Analysis

Depending upon the screening of mRNAs-lncRNAs gene pairs, a co-expression network was developed, Pearson'scoefficient of correlation (PCC) greater than 0.95 or *P* value <0.05. For the prediction of protein/protein interaction (PPI) for DE genes, STRING database was referred. BLASTx (version 2.2.28) was utilized to retrieve query sequences of those genes that were not retrieved by developed networks, and for all chosen referred organisms, networks were built as per known interactions. The Cytoscape tool was utilized for the visualization of PPIs, which is developed by the Institute for Systems Biology (Seattle, Washington, USA).

Statistical Analysis

Data are expressed as the mean \pm SEM. Two-way ANOVA with repeated measures was performed for behavioral data analysis. For all comparisons, the statistical significance standard was set at p < 0.05.

Results

Intra-TMJ Injection of CFA Induces Chronic Inflammatory TMJ Pain

We observed that intra-TMJ injection of CFA significantly decreased the head withdrawal threshold in trigeminal nerve V3 branch-innervated facial skin areas compared to the saline group (Figure 1). The CFA-induced TMJ pain started from day 1 post-CFA injection and lasted for at least for 10 days. On day 4 post-CFA injection, the head withdrawal threshold reached the lowest value, indicating the strongest TMJ pain at that time point.

Quality Control and Data Correlations

RNA-seq data quality was examined by analyzing the ratio of clean data to all data (Figure 2A). The ratio of clean reads to all raw reads was more than 95% for all samples, which suggests that the quality of RNA-seq data was high and they can be used for subsequent analysis. We used Pearson correlation coefficient to analyze correlations as calculated in R (Figure 2B). The R² values in each group were more extensive than 0.90, indicating that the samples' biological replicates in each group met analysis requirements.

Genes	mRNA (C-TG vs S-TG)	mRNA (C-Sp5C vs S-Sp5C)	IncRNA (C-TG vs S-TG)	IncRNA (C-Sp5C vs S-Sp5C)	TUCP-RNA (C-TG vs S-TG)	TUCP-RNA (C-Sp5C vs S-Sp5C)
Up- regulated	229	145	2	6	5	5
Down- regulated	228	247	28	2	6	2
Total	457	392	30	8	11	7

 Table I Differentially Expressed mRNAs and IncRNAs in the TG and Sp5C After TMJ Pain Induction

Abbreviations: C-TG, TG in the CFA group; S-TG, TG in the saline group; C-Sp5C, Sp5C in the CFA group; S-Sp5C, Sp5C in the saline group; TUCP, transcripts of uncertain coding potential.







Figure 3 Volcano plots, venn diagrams, and heat maps of DE mRNAs. (A and B) Volcano plots illustrating comparisons of DE mRNAs between CFA group and saline control group in the TG and Sp5C. (C–E) Venn diagrams showing overlap of DE mRNAs between CFA group and saline control group in the TG and/or Sp5C: total DE mRNAs; up-regulated DE mRNAs; down-regulated DE mRNAs. (F) Heat map displaying the hierarchical clustering of DE mRNAs in each group (red color indicates up-regulation and blue color indicates down-regulation). C_TG, TG from CFA group; S_TG, TG from saline control group; C_Sp5C, Sp5C from CFA group; S_Sp5C, Sp5C from saline control group.

DE mRNAs and IncRNAs in TMJ Pain

To identify previously unrecognized molecular features of TMJ pain, high-throughput RNA-seq was carried out. TG and Sp5C tissues were harvested from mice on day 4 after intra-TMJ injection of CFA or saline. We extracted and sequenced the total TG and Sp5C RNA from these groups. DE mRNAs and lncRNAs in the comparison between CFA (C) and Saline (S) groups are listed in Table 1. There were 457 DE mRNAs in the TG (229 upregulated and 228 downregulated); There were 392 DE mRNAs in the Sp5C (145 upregulated and 247 downregulated); There were 30 DE lncRNAs in the TG (2 upregulated and 28 downregulated); There were 8 DE lncRNAs in the Sp5C (6 upregulated and 2 downregulated); There were 11 DE transcripts of uncertain coding potential RNAs (TUCP-RNAs) in the TG (5 upregulated and 6 downregulated); There were 7 DE TUCP-RNAs in the Sp5C (5 upregulated and 2 downregulated).

DE mRNAs and lncRNAs in each group were also displayed with volcano plots, Venn diagrams, and heat maps. We generated volcano plots to show the DE mRNAs profiles with the statistical significance threshold set at P < 0.05 (Figure 3A and B). To determine the similarity between TG and Sp5C comparisons, Venn diagrams were drawn (Figure 3C–E). A heat map of the DE mRNAs is displayed in Figure 3F. The heat map revealed the variations in DE mRNAs among groups (Figure 3F). Representatives of differentially expressed mRNAs in the TG and Sp5C after TMJ pain induction are shown in Table 2. Similarly, we drew volcano plots to show the DE lncRNAs profiles with the statistical significance threshold set at P < 0.05 (Figure 4A and B). To determine the similarity between TG and Sp5C comparisons, Venn diagrams were drawn (Figure 4C–E). We then screened the candidate lncRNAs. Four tools (CPC, CNCI, PFAM, and PhyloCSF) were used to analyze the coding potential of lncRNAs. The candidate lncRNAs were designated by four analytical tools and used in subsequent analyses (Figure 4F). A heat map showed the variations in DE lncRNAs among groups (Figure 4G). Moreover, the class distribution of DE lncRNAs was assessed, including lncRNA, antisense, and intronic transcripts. Detailed biotypes of these lncRNAs are displayed in Figure 4H, which indicates that most lncRNAs are in the intergenic lncRNA category. Representatives of differentially expressed lncRNAs in the TG and Sp5C after TMJ pain induction are shown in Table 3.

Genomic Features of IncRNAs

It has been reported that lncRNAs are shorter in length and less conserved than protein-coding transcripts.¹⁷ Consistently, our results showed that the predicated lncRNAs tended to contain fewer exons (Figure 5A), and their genes were shorter in length than protein-coding transcripts (Figure 5B). Furthermore, we identified that lncRNAs tended to be shorter in Open Reading Frame (Orf) length than protein-coding genes (Figure 5C). We observed that lncRNAs tended to be expressed at lower levels than mRNAs (Figure 5D). We also observed that the predicted lncRNAs were less conserved than protein-coding transcripts (Figure 5E), which is consistent with a previous report.²¹

mRNA	Gene_Id	Gene_Name	S_FPKM	C_FPKM	Fold-Change	P-value
mRNA (TG)	ENSMUSG0000009350	Мро	0.16	1.72	1.56	0.04
mRNA (TG)	ENSMUSG0000042312	S100a13	34.75	39.19	4.43	0.04
mRNA (TG)	ENSMUSG0000021025	Nfkbia	14.95	20.63	5.69	0.04
mRNA (TG)	ENSMUSG0000028931	Kcnab2	60.26	21.07	-39.19	0.04
mRNA (TG)	ENSMUSG0000054934	Kcnmb4	11.47	9.28	-2.19	0.02
mRNA (TG)	ENSMUSG0000054477	Kcnn2	3.21	0.92	-2.29	< 0.01
mRNA (Sp5C)	ENSMUSG0000025958	Crebl	1.43	4.99	3.56	0.02
mRNA (Sp5C)	ENSMUSG0000004221	lkbkg	3.83	7.11	3.28	< 0.01
mRNA (Sp5C)	ENSMUSG0000020366	Mapk9	2.39	5.46	3.06	0.04
mRNA (Sp5C)	ENSMUSG0000021508	Cxcl14	11.51	23.23	11.72	< 0.01
mRNA (Sp5C)	ENSMUSG0000034656	Cacnala	9.41	11.26	1.85	0.03
mRNA (Sp5C)	ENSMUSG0000038077	Kcna6	20.41	17.48	-2.93	< 0.01
mRNA (Sp5C)	ENSMUSG0000041329	Atp1b2	2.12	0.64	-1.48	0.04
mRNA (Sp5C)	ENSMUSG0000039904	Gpr37	37.49	30.15	-7.33	0.02

Table 2 Representative Differentially Expressed mRNAs in the TG and Sp5C After TMJ Pain Induction

Abbreviations: S_FPKM, fragments-per-kilo-base of exon-per-million fragments mapped in the saline group; C_FPKM, fragments-per-kilo-base of exon-per-million fragments mapped in the CFA group.



Figure 4 Volcano plots, venn diagrams, and heat maps of DE IncRNAs. (A and B) Volcano plots illustrating comparisons of DE IncRNAs between CFA group and saline control group in the TG and Sp5C. (C-E) Venn diagrams showing overlap of DE IncRNAs between CFA group and saline control group in the TG and Sp5C. (C-E) Venn diagrams showing overlap of DE IncRNAs between CFA group and saline control group in the TG and/or Sp5C: total DE IncRNAs; up-regulated DE IncRNAs; down-regulated DE IncRNAs. (F) Venn diagram showing candidate IncRNAs selected by four tools (CPC, CNCI, PFAM, and phyloCSF) for subsequent analyses. (G) Heat map displaying the hierarchical clustering of DE IncRNAs in each group (red color indicates up-regulation and blue color indicates down-regulation). (H) A pie chart showing three categories of the altered IncRNAs. C_TG, TG from CFA group; S_TG, TG from saline control group; C_Sp5C, Sp5C from CFA group; S_Sp5C, Sp5C from saline control group.

IncRNA	Gene_ld	Gene_Name	S_FPKM	C_FPKM	Fold-Change	P-value
IncRNA (TG)	ENSMUSG0000086844.1	AC163434.5	11.94	7.68	-4.26	< 0.01
IncRNA (TG)	ENSMUSG0000074415.13	RP23-442M18.1	9.74	7.54	-2.20	0.04
IncRNA (Sp5C)	ENSMUSG0000092274.2	Neat I	1.98	3.73	1.75	0.04
IncRNA (Sp5C)	ENSMUSG00000105843.1	RP23-240E15.3	2.02	4.44	2.42	0.01
IncRNA (Sp5C)	ENSMUSG0000086606.1	RP23-23015.2	7.78	5.24	-2.54	< 0.01

Table 3 Representative Differentially Expressed IncRNAs in the TG and Sp5C After TMJ Pain Induction

Abbreviations: S_FPKM, fragments-per-kilo-base of exon-per-million fragments mapped in the saline group; C_FPKM, fragments-per-kilo-base of exon-per-million fragments mapped in the CFA group.

Functional Analysis of DE mRNAs and IncRNAs

The pathway patterns predicted by KEGG could provide information for future studies on mRNAs' roles in TMJ pain mechanisms. Using KEGG pathway enrichment analysis of DE mRNAs, we identified top 20 dysregulated pathways in the TG (Figure 6A) and Sp5C (Figure 6B). Moreover, GO analysis of the DE mRNAs was conducted to ascertain specific categories of biological process (BP), cellular processes (CC), and molecular function (MF) of the DE transcripts. For the DE mRNAs in the TG, a histogram displaying the number of genes for each GO term was shown in Figure 6C. For the DE mRNAs in the Sp5C, a histogram displaying the number of genes for each GO term was shown in Figure 6D.

To identify potential target genes of the DE lncRNAs, we set a threshold for collocation as 100 kb upstream and 100 kb downstream of each lncRNA. Using the KEGG pathway enrichment analysis of the DE lncRNAs, we revealed top 20 significantly enriched pathways in the TG (Figure 7A) and Sp5C (Figure 7B). Moreover, GO analysis of the DE lncRNAs was conducted to ascertain specific categories of the DE transcripts. For the DE lncRNAs in the TG, a histogram displaying the number of genes for each GO term is shown in Figure 7D.

Protein–Protein Interaction (PPI) Networks Under TMJ Pain

To explore the lncRNA-mediated molecular mechanisms for TMJ pain, we further conduct PPI network analysis. We identified that the PPI network consisted of 2063 nodes and 4535 edges in the TG (Figure 8A) and the PPI network consisted of 87 nodes and 91 edges in the Sp5C (Figure 8B). Genes were displayed as nodes and PPIs were displayed as edges. The node size is proportional to the degree of the node. Interesting genes were enlarged and labeled with red color. The PPI networks indicated the reliability of our sequencing data and provided a lot of information on downstream targets of the DE lncRNAs.

Discussion

Previous studies have shown that the expression of protein-coding genes is altered in the TG neurons following CFA- or capsaicin-induced orofacial pain.^{22,23} In recent years, several lncRNAs have been reported to regulate different types of pain.^{3,4,24–26} However, the potential functions of lncRNAs in orofacial pain are not fully understood. Xiong et al demonstrate that lncRNA uc.48+ is upregulated in the TG and plays an essential role in pain transmission of trigeminal neuralgia.⁹ In the present study, we analyze the transcripts of both TG and Sp5C in a CFA-induced inflammatory TMJ pain mouse model. Using the sequencing technique, we carried out a comprehensive genome-wide transcriptional screening of lncRNAs and mRNAs during CFA-induced TMJ pain. Our results indicate that on day 4 post-CFA when maximal TMJ pain is induced, the expression patterns of many mRNAs and lncRNAs in both TG and Sp5C are significantly altered.

The DE mRNAs and lncRNAs identified in this study are associated with each other and contribute to pain regulation. It has been reported that decreased potassium channel gene expression can reduce voltage-gated potassium currents in ganglion neurons for pain regulation.²⁷ In this study, we observed that after TMJ pain induction, Kcnab2, Kcnn2 and Kcnmb4 in the TG and Kcna6 in the Sp5C were decreased (see Table 2). Meanwhile, we also observed that TMJ pain induction decreased AC163434.5 (predicted target mRNA: Kcnab2) in the TG (see Table 3). These results suggest that potassium channel gene regulation in the trigeminal nociceptive system contributes to the pathogenesis of TMJ pain. The



Figure 5 Comparison of the identified IncRNAs and mRNAs. (A) Distribution of the number of exons in the identified mRNAs and IncRNAs. (B) Distribution of transcript lengths in the identified mRNAs and IncRNAs. (C) Distribution of number of open reading frame (Orf) in the identified mRNAs and IncRNAs. (D) Expression level analysis in the identified mRNAs and IncRNAs. (E) Conservative analysis of sequences in the identified mRNAs and IncRNAs.



Figure 6 KEGG and GO analyses of the DE mRNAs. (A and B) KEGG analysis of the DE mRNAs showing top 20 dysregulated pathways in the TG and Sp5C comparing CFA group with saline control group. (C and D) GO analysis of the DE mRNAs showing the number of genes for each GO term in the TG and Sp5C comparing CFA group with saline control group. C_TG, TG from CFA group; S_TG, TG from saline control group; C_Sp5C, Sp5C from CFA group; S_Sp5C, Sp5C from saline control group.



Figure 7 KEGG and GO analyses of the DE lncRNAs. (A and B) KEGG analysis of the DE lncRNAs showing top 20 dysregulated pathways in the TG and Sp5C comparing CFA group with saline control group. (C and D) GO analysis of the DE lncRNAs showing the number of genes for each GO term in the TG and Sp5C comparing CFA group with saline control group. C_TG, TG from CFA group; S_TG, TG from saline control group; C_Sp5C, Sp5C from CFA group; S_Sp5C, Sp5C from saline control group.



Figure 8 PPI networks under TMJ pain in the TG and Sp5C. (A) PPI networks constructed by DE IncRNAs target mRNAs in the TG comparing CFA group with saline control group. (B) PPI networks constructed by DE IncRNAs target mRNAs in the Sp5C comparing CFA group with saline control group. Genes were displayed as nodes and PPIs were displayed as edges. The node size is proportional to the degree of the node. Interesting genes were enlarged and labeled with red color.

DE mRNAs and lncRNAs we identified provide consistent information for specific gene regulation in TMJ pain. Moreover, previous studies have demonstrated that the Mitogen-activated protein kinase (MAPK) signaling pathway is involved in pain sensitization²⁸ and TMJ pain.²⁹ In the present study, we identified increased Mapk9 in the Sp5C after TMJ pain induction (see Table 2), which indicates a potential role of MAPK signaling in TMJ pain. Additionally, lncRNA RP23-240E15.3 and its predicted target mRNA Cxcl14 as well as lncRNA RP23-230I5.2 and its predicted target mRNA Gpr37 are altered after TMJ pain induction (see Tables 2 and 3). The change in the expression of these genes may also contribute to TMJ pain. Therefore, our results in this study provide potential targets for us to further investigate the underlying molecular mechanisms for TMJ pain in the future.

We next conduct a KEGG pathway analysis of the identified DE mRNAs and lncRNAs in the TG and Sp5C. The top 20 dysregulated pathways for each condition are summarized. We further conduct GO analysis of the identified DE mRNAs and lncRNAs to ascertain specific categories of the DE transcripts. For instance, the cAMP signaling pathway we identified is involved in inflammatory TMJ hypersensitivity³⁰ and neuropathic pain,³¹ the MAPK signaling pathway we identified can enhance TMJ nociceptive processing,²⁹ and the Ras signaling pathway we identified has been reported to contribute to immune processes in the pathophysiology of TMD pain.³² Furthermore, PPI network analysis of the identified DE lncRNAs shows interesting genes that are mostly relevant to pain regulation. Therefore, transcriptomic analysis data we report here not only demonstrate several common signaling pathways in pain mechanisms but also indicate some potential genes, including potassium channel genes, as well as those genes in cAMP, MAPK and Ras signaling pathways, may be specifically involved in TMJ pain. However, the identified genes have not been validated to confirm their roles in TMJ pain. We will conduct an experimental validation of the identified genes in our future studies.

Conclusion

Our results in this study highlight important transcriptional changes induced by intra-TMJ injection of CFA into the TG and Sp5C. We identify DE mRNAs and lncRNAs in the trigeminal nociceptive system of mice with inflammatory TMJ pain. Based on bioinformatic analysis, we reveal that the dysregulated mRNAs and lncRNAs are mostly involved in pain regulation. Our study advances our understanding of different gene expressions in TMJ pain. Additional functional analyses are needed to further unravel the potential roles of the identified transcriptional modification in the molecular mechanisms underlying TMD and TMJ pain.

Data Sharing Statement

The datasets generated during and/or analysed during the current study are available in the GEO repository, and the GEO accession number is GSE188922.

Ethics Approval

This study was approved by the Animal Ethics Committee of Zhengzhou University (Animal Protocol Number: 2021328).

Author Contributions

All authors made substantial contributions to conception and design, acquisition of data, or analysis and interpretation of data; took part in drafting the article or revising it critically for important intellectual content; agreed on the journal to which the article will be submitted; gave final approval of the version to be published; and agreed to be accountable for all aspects of the work.

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Disclosure

The authors declare that there are no conflicts of interest.

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