Integrated analysis of Transcriptomic Data Reveal AP-1 as a Potential Regulation Hub in Inflammation Induced Hyperalgesia Rat

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Research

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Abstract

Background

Inflammation associated chronic pain is a global clinical problem, affecting millions of people worldwide. However, the underlying mechanism of inflammation associated chronic pain remains unclear. Complete Freund's adjuvant (CFA) induced cutaneous inflammation rat has been widely used as an inflammation-induced pain hypersensitivity model.

Methods

Herein, we presented a transcriptomic profile of CFA-induced rat dorsal root ganglion via an approach targeting gene expression, DNA methylation and post-transcriptional regulation.

Results

We identified 418 differentially expressed mRNAs, 120 differentially expressed microRNAs and 2670 differentially methylated regions, which are all highly associated with multiple inflammation related pathways, including NF-κB signaling pathways and IFN signaling pathways. Integrated analysis further demonstrated that AP-1 network, which may work as a regulator of inflammation response, is sophisticatedly regulated at both transcriptomic and epigenetic level.

Conclusions

We believe our data will not only provide drug screening targets for treating chronic pain and inflammation, but also shed a light on the molecular network inflammation induced hyperalgesia.

Introduction

Chronic pain, which affects more than 17% of the all Canadians, as well as approximately 100 million people in the United States [22; 23], has been one of the most common and distressing issues, influencing the society and individuals significantly [5]. Chronic pain is usually accompanied with inflammation, tissue damage, hyperalgesia as a separate condition [18]. Risks factors of chronic pain include socio-demographic, mental, clinical as well as the biological factors [29]. To date, inflammations and tissue damages are widely believed as the leading biological factors to induce chronic pain [7; 24]. However, more studies on the pathogenesis of chronic pain are imperative at molecular level.

Animal inflammation models are widely used to assess the production of inflammatory mediators. Complete Freund's adjuvant (CFA)-induced cutaneous inflammation rat model has been widely characterized in literature and has been used for finding novel anti-inflammatory molecules for years [6]. Swelling, hyperalgesia and allodynia can be used to evaluate the inflammation response in the CFA-induced model [6]. Long-term hyperexcitability of the primary sensory neurons in the dorsal root ganglion (DRG), and the secondary sensory neurons in the spinal cord dorsal horn (SDH), are considered as the
cause of the alteration in pain perception [30]. CFA-induced chronic inflammation causes chronic pain. CFA-induced rat has been used as one of the most common chronic pain animal models [14; 15; 36; 37]. However, the mechanism of how CFA-induced chronic pain remains not fully understood.

Recently, integrated analysis of multi-omics data, which relies on next generation sequencing, has revealed numerous complex molecular networks of basic biological process as well as human diseases [26; 32]. DNA, RNA and protein work synergistically to perform certain biological functions. Focused on chronic pain studies for years, we are extremely interested in understanding the underlying biological network of how CFA induce chronic pain in animals. In fact, there exists a pioneer study published by Tan et al. in 2017. Although only an array-based analysis focused on microRNA (miRNA) and mRNA network was used, they were still able to reveal that several miRNA-mRNA interactions contributed to the inflammatory pain process. For instance, they found that miR-124-3p could attenuate inflammatory pain and decrease IL-6R expression in the spinal cord [17]. Aside from this study, miRNAs have already been widely associated with inflammatory models [1; 12; 27; 33]. Inhibiting some of these miRNAs has been considered as novel treatment for chronic pain [13; 20; 21]. However, only focusing on post-transcriptional regulation is not enough to illustrate how CFA-induced inflammatory process is regulated comprehensively. In fact, our previous work has demonstrated that dysregulation of DNA methylation is involved in the CFA-induced inflammation model [14]. A recent study also described that hypomethylation of nerve growth factors affects Inflammatory hyperalgesia in rats [39]. These studies suggested that epigenetic regulation, especially DNA methylation could contribute to inflammatory pain. In addition, although Tan et al. provided rat SDH tissue expression profiles days after CFA treatment using microarray technology [17], the transcriptomic profile of the dorsal root ganglion (DRG) tissues from the chronic pain animal models is still lacking. This suggested that how the gene expression is affected by CFA treatment in the primary sensory neurons is still not clear.

In order to provide a comprehensive profile of DRG tissues’ transcriptomic response to CFA injection, herein, we used a an approach that target DNA methylation, gene transcription and post-transcriptional regulation, which includes mRNA-seq, small-RNA-seq and targeted bisulfite sequencing, to detect the transcription level alteration. We identified 418 differentially expressed genes, 120 differentially expressed miRNAs and 2670 differentially methylated regions, which are associated with multiple inflammation related pathways. Integrated analysis further indicates that AP-1, a key regulator of inflammation response, is sophisticatedly regulated at both transcriptomic and epigenetic level.

Methods

Animal Model

15 pairs of adult male SD rats in a SPF grade weighing about 200 g ± 5 g were purchased from Experimental Animal Center of Suzhou University. The SD rats maintained at standardized feeding environment including 12-hours shift light-dark cycle and ambient temperature of 24 ± 1°C. The CFA-induced chronic inflammatory pain model was established by unilateral subcutaneous injection of CFA.
0.1 mL at sterile foot skin and was described in our previous research [6] [14]. 10 pairs of the rats (CFA and control) were used for following sequencing experiments, while the rest 5 pairs were used for the validation experiments. All experiments were approved by the Animal Research Ethics Committee from The First People's Hospital of Yancheng, Suzhou Municipal Hospital Affiliated to Nanjing Medical University and Nantong University. Mechanical and Thermal pain threshold measurement was also described in our previous study in detail [14].

**RNA-seq, Small-RNA-seq and Targeted Bisulfide sequencing**

For the NGS-based sequencing, DRG tissues from 10 pairs (Randomly selected) of CFA-treated and untreated rats were extracted. For library construction, RNA-seq library was prepared with NEBNext Ultra RNA with Poly-A selection and was sequenced on an Illumina Hi-Seq 4000. Small RNA-seq library was constructed with NEBNext Multiplex Small RNA Library Prep Kit for Illumina (#7560S). The targeted bisulfide sequencing library is prepared with a DNA library construction TruSeq DNA LT Sample Prep Kit v2 for Illumina. The C-T transition is performed via C-T transition EpiTect Bisulfite kit from Qiagen (Germany).

**RNA sequencing analysis**

After running fastQC (0.11.8) for quality control, Cutadapt (2.1.0) was used for trimming adapters and low-quality sequence (Phred score less than 20) from the raw fastq files. The cleaned fastq files were then mapped to m4 rat genome refence by STAR (2.7.0) with suggested setting. Differential expression analysis was then performed by DEseq2 (1.29.0). Significant differentially expressed genes was defined by P value ≤ 0.05 and log2 fold change ≥ 0.5. Further pathway enrichment analysis was performed with EnrichR. The gene set enrichment analysis (GSEA) was performed by using javaGSEA2-3.0.

The RNA-seq wig file generated by STAR were then used as input for DaPars (0.9.1) to identify alternative polyadenylation events. Significant events were defined by P value less than 0.05 and absolute delta PDUI more than 0.1 [35].

**Small RNA sequencing data analysis**

Small RNA-seq data was analyzed by using sRNAAnalyzer pipeline [34]. Adapter sequences was provided as input to the pipeline for removing adapter contamination. The cleaned reads were then mapped to rat miRNA database to generate count matrix, which was further analyzed by DEseq2 to identify differentially expressed miRNA. Significantly expressed miRNA were defined based on the P value less than 0.05. The miRNA targets were predicted by using ‘Multimir’ (1.10.0) R package. Finally, the pathway enrichment analysis was performed by using EnrichR.

**DNA methylation data analysis**

TrimGalore (0.6.4) was used for sequencing data quality control, adapter removing and trimming low-quality sequence (Phred score less than 20). The cleaned data was then mapped to the m4 rat genome reference by using Bismark (0.22.0). The generated CpG methylation ratios were then used as input for
metilene to call differentially methylated regions (DMRs) [10]. The significant DMRs were defined by P value less than 0.05 and absolute methylation ratio change more than 0.1. Significant DMRs were annotated to nearest genes by using Homer (4.8). Pathway enrichment analysis and motif enrichment analysis were performed by EnrichR and MEME-suite, respectively.

**Statistical analysis**

Enrichment analysis was detected by Fisher’s Exact test and p-values < 0.05 were considered statistically significant. False Discovery Rate (FDR) correction was used to generate p-adj value in all expression matrix data (RNA-seq, small-RNA-seq and DNA methylation). All data (mean ± SE) were analyzed with a Student’s t test, and a p-value < 0.05 was considered statistically significant.

**Quantitative Real-time PCR assays**

DRG tissues from 4 pairs of the rats (Control and CFA) were separated half portion for qPCR assays. Total RNAs were extracted from cells by using TRIpure reagents (Bioteke Beijing) and then reverse-transcribed by using the first strand cDNA synthesis kit (TSK302S (RT6 cDNA Synthesis Kit Ver 2). Reverse-transcribed products were used as templates for real-time PCR using the 2×T5 Fast qPCR Mix (SYBR Green I). The primers used for different target genes were as follows: Reg3a-F2: CTGTCACCGTGGTAACTGTGG; Reg3a-R2: TGCAGGCTCTACTGCTTGAAC; Reg3b-F: TCAACTGGGAGGAGAACCC; Reg3b-R: TGGCCCTTTGAGATCTGT; Dusp1-F2: AACGAGGCAGTTGACTTTATAGCT; Dusp1-R2: TCTCTGTGTTCTCAG; Egr1-FA: ACCCTACGAGCACCTGACC; Egr1-R: AAGCGGCCAGTATAGGTGATG; rno-miR-384-3p-RT: GTCGTATCGACTGCAGGGTCCGAGGTATTCGCAGTCGATACGACATTGTG; rno-miR-384-3p-F: CGCATTCCTAGAAATTGTT; rno-miR-181b-5p-RT: GTCGTATCGACTGCAGGGTCCGAGGTATTCGCAGTCGATACGACATTGTG; rno-miR-181b-5p-F: GGAACATTCATTGCTTCG; common-R: ACTGCAGGGTCCGAGGTAT; rat-actin-F: AGATCAAGATCTTTAGAGGCTC; rat-actin-R: ACGCAGCTCAGTACAGTCC

**Western blotting**

DRG tissues from three pairs of the mice were collected for the protein extraction (Total n = 6). DRG tissues were washed three times before the tissue preparation. Total lysates were fractionated by SDS/PAGE and then transferred for 1 hour to the nitrocellulose filter (NC) membranes. The membranes were then blocked for 45 min with Tris-buffered saline containing Tween 20 (TBST), which contain 3% (mass/vol) nonfat dry milk. Antibodies for Egr1 were purchased from Invitrogen a-actin were purchased from Cell Signaling. After the incubation with primary antibodies, the membranes were washed and then incubated with HRP-conjugated anti-mouse or anti-rabbit secondary antibodies. The membrane was then followed by detection using ECL Western blotting substrate (Bio-Rad). Quantitative analysis of Western blot results were calculated using Image J software.

**Results**
CFA-induced inflammation rat model

To establish the CFA-induced inflammation rat model, we injected male SD rats (200 ± 5 g of body weight) with 0.1 mL CFA on the plantar skin of the right hind paw. Detail methods were described in the Method section and was published in our previous work. 24 hours after the CFA injection, we measured the rat hind paw withdrawal threshold (PWT) as well as the paw withdrawal latency (PWL) in each group. As expected, we discovered a significant downregulation of both PWT and PWL in the CFA group compared to the saline treated group (Figure 1). These results validated our previous experiments as well [14].

RNA-seq profiling of the CFA-induced inflammation model in rat

To investigate the transcriptomic alternation during the CFA-induced inflammation, we performed total RNA-seq with dorsal root ganglion (DRG) from 10 pairs of rats with or without CFA treatment and generated an RNA-seq dataset with high quality in terms of sequencing quality, duplication levels, adapter contamination and mapping rates. Differential expression analysis revealed that 213 and 205 genes were significantly up-regulated or down-regulated, respectively (Figure 2a). Among all the significant differentially expressed genes (DEGs), Reg3b and Reg3a are the top two genes with more than 6-fold increment of expression level (Figure 2b). A quantitative real-time PCR (qRT-PCR) assay was also performed to validate the increment of these mRNA expression (Supplementary Fig. 1). It has been well-known that regenerating gene (Reg) family plays important role in cell proliferation, migration and inflammation response [40]. In order to examine the transcriptomic alteration globally, we also performed pathway enrichment analysis based on all DEGs and observed the enrichment of many inflammation related pathways, such as Interferon alpha/beta signaling pathway, RIG-1-like receptor signaling pathway, TRAF6 mediated NF-κB activation and AP-1 transcription factor network (Figure 2c). Interestingly, The TRAF6 mediated NF-κB has been previously linked to CFA-induced inflammation already [2; 31; 36]. Altogether, these RNA-seq based results not only further validates our success construction of the CFA-induced inflammation rat model but also reveals the specific transcriptomic alternations induced by CFA treatment in rat (Supplementary Table 1).

Alternative polyadenylation (APA) has been shown to regulate gene expression by regulating the interaction between miRNA and RNA molecules [4]. In order to investigate the gene regulation mechanism of the CFA-induced inflammation, we therefore first profiled the APA events between CFA-treated and control rats by DaPars. However, no significant APA events were detected (Supplementary Table 2), indicating that APA is irrelevant to the expression change observed in the CFA-treated rats.

miRNA profiling of the CFA-induced inflammation model in rat

Although we did not observe APA change in the CFA-treated rats, miRNA might regulate the gene expression directly through the miRNA abundance. In fact, there are several miRNAs were reported to be
related to CFA-induced inflammation, including miR-134 in DRG tissue, and miR-124, miR-149, miR-3584 in spinal cord dorsal horn [17; 19]. However, these studies did not include a comprehensive database which should rely on a non-biased next generation sequencing platform. In order to provide a landscape of how miRNA network is regulated right after CFA treatment, we next performed small RNA-seq for the same 10 pairs of rat tissues with or without CFA treatment. As expected, the differential expression analysis showed that 16 miRNAs were up-regulated, such as mo-miR-384-3p, which targets immune related genes like Pik3cd, Nfatc3 and Cblb (Supplementary Table 3). We also identified 104 significantly down-regulated miRNAs, with the top one being mo-miR-181b, which is predicted to target Tnf and Irs2 (Figure. 3a). Heatmap of miRNA expression profile also showed the control and CFA treated rats were well clustered (Figure. 3b). To study the functions of the down-regulated miRNAs unbiasedly, we then performed pathway enrichment based on their target genes. Interestingly, many inflammation related pathways are highly enriched, such as TGF-beta signaling pathway, PDGFB signaling pathway, SHP2 signaling pathway and MAPK signaling pathway (Figure. 3c). These results suggest that, in response to CFA-treatment, miRNA is involved in the regulation of the inflammation response.

**DNA methylation profiling of the CFA-induced inflammation model in rat**

Our previous work demonstrated a promoter demethylation of CXCR4 gene is related to CFA-induced inflammation [14]. To further investigate the gene regulation mechanisms of CFA-induced inflammation response, we profiled the methylation landscape of the CFA-treated rats by performing targeted bisulfite sequencing. In the comparison of control and CFA-treated rats, we totally identified # differential methylated cytosines, suggesting CFA can indeed cause the change of DNA methylation landscape. We further performed differentially methylated regions (DMRs) calling by using the ‘de-novo’ mode of metilene and identified 1401 hypermethylated DMRs and 1269 hypomethylated DMRs (Figure. 4a) (Supplementary Table 4). The majority of DMRs are enriched to the intergenic and intron regions, with only 19 DMRs are located in the promoter regions, suggesting CFA-induced DNA methylation change might not play the role of gene regulation by direct repressing transcription at the gene promoter regions (Figure. 4b). We then assigned DMRs to the nearest genes and performed pathway enrichment. For both hypermethylated and hypomethylated DMRs, we observed the enrichment of inflammation related pathways, such as Interleukin-4 signaling pathway, Insulin signaling pathway, TGF-beta receptor activation of SMADs, Signaling by NGF and Toll receptor cascades (Figure. 4c). Finally, we did motif analysis with the DMRs and found the enrichment of the motifs for Tbp and Mtf1 transcription factors, both of which has been reported to linked to stress response and inflammation response previously [16; 28]. In summary, these results show that DNA methylation is involved in the gene regulation of CFA-induced inflammation response, probably through influencing the binding of inflammation related transcription factors.

**AP-1 works as a potential regulation hub for CFA-induced inflammation response**
After investigating CFA-induced change of gene expression, miRNA and DNA methylation, we examined their interplay in order to illustrate how these three layers of molecular changes are coordinated. Since majority of miRNAs are downregulated, we then focused on their targets genes which had corresponding increased transcription level (571 genes). These target genes are also enriched for many inflammation related pathways, but with the top one being AP-1 transcription factor network (Figure 5a) (Supplementary Table 5). Interestingly, the DEGs are also enriched in this pathway, further indicating that miRNA mediated repression of AP-1 network had to be mitigated during the CFA-induced inflammation response (Figure 2c and 5b). A validating experiment based on qRT-PCR also showed that Dusp1, one of the two AP-1 related molecules, was significantly highly expressed in the CFA group compared to the control (Supplementary Fig. 2). However, we failed to find the differential expression for Egr1 gene in the qRT-PCR results (Supplementary Fig. 2). In fact, although the RNA-seq result exhibited a 1.5-fold upregulation of Egr1 expression in the CFA group, most of the reads were enriched in the 3’ UTR, indicating a potential post transcriptional regulation event. In order to determine whether Egr1 was regulated by CFA treatment, we detected the protein level of Egr1 and found that Egr1 expression was slightly decreased in the CFA group (n = 3, Supplementary Fig. 3). Compared to the upregulation in mRNA level, this opposite trend of the protein expression may be a consequence of the increased expression of 3’UTR. Furthermore, we also studied the interaction between DNA methylation and gene expression. Since the majority of DMRs are annotated to the intergenic regions, we are not able to observe significantly anti-correlation between differential methylation and differential expression. However, based on the motif analysis, we found that CFA-induced DMRs were surprisingly enriched for the AP-1 binding motif (Figure. 5c) (Supplementary Table 6). Given that AP-1 is a well-known key regulator of inflammation response and has been previously linked to CFA-induced inflammation [3], we speculate that AP-1 may serve an important role as a hub subjected to the regulation of both miRNA and DNA methylation upon CFA treatment.

Discussion

The addition of “omics” to certain molecular term reflects a comprehensive assessment of a set of molecules. Integrated analysis of multiple NGS-based datasets, which highly rely on high-throughput sequencing, has already revolutionized the medical research [8]. Chronic pain is a critical health issue globally, affecting millions of people. However, there exists difficulty for directly obtaining certain tissues directly from patients clinically. Thus, we believed that understanding the molecular networks of chronic pain animal models through a comprehensive approach is an alternative. Herein, we provided an integrated analysis based molecular profile, which focuses on the response of rat DRG tissues to CFA injection, at epigenetic, transcription and post-transcriptional regulation level. Overall, we identified 418 differentially expressed mRNAs, 120 differentially expressed miRNAs and recognized more than 2,500 DMRs in CFA-treated groups. Through gene set enrichment analysis, we validated some of the previously reported CFA-response related signaling pathways that are also highly related to CFA induced inflammation in our dataset as well, such as, NF-κB signaling pathways [36] and IFN signaling pathways [38]. Moreover, we also identified many new genes/pathways that are potentially highly involved in this
pain response model, including Reg family genes (Reg3a, Reg3b) and the AP-1 transcription related pathways. It is interesting that after we adjusted the cell heterogeneity of the methylation sequence data by using CHALM [41], a recent invented software for analyzing methylome, we identified 6832 significant DMRs, suggesting the CFA's impact on rat's methylome is underestimated by the traditional analysis method. Besides inflammation related pathways or terms, the CHALM identified DMRs are also enriched to heart contractions and heart rate, indicating that methylome re-wire is involved in the chronic pain linked heart malfunctions or diseases (Supplementary Table 10). Finally, Based on our multi-omics profiling of the CFA-induced chronic pain model, we selected top 10 differentially expressed genes, miRNAs and differentially methylated regions as the multi-omics signature for our chronic pain model. CFA treatment group and control group can be clearly separated by these signatures (Fig. 6a-c) (Supplementary Table 7–9), and is consistent to the idea that miRNA and DNA methylation regulate mRNA transcription.

Pancreatitis-associated proteins, which are from Reg families, have been previously linked to modulation of spinal sensory pathways in pathological pain states [9]. Our results demonstrated that Reg3b and Reg3a were significantly upregulated for 12-fold and 6-fold, respectively, in CFA-treated groups, indicating that the Reg family genes may play a crucial role in regulating inflammatory pain response in DRG tissues. AP-1 transcription factor is composed of dimerization of a bZIP (basic region leucine zipper) domain via the Fos and Jun subunits. AP-1 regulation network was previously linked to chronic pain response [3]. However, it was never recognized as a central pathway in pain response before. In our dataset, AP-1 network is considered as the regulation central hub. Not only the CFA-induced miRNA and mRNA interactions are highly enriched for AP-1 networks, but also the CFA-induced DMR motifs are enriched for the AP-1 transcription binding activities. Therefore, we hypothesized that inhibiting certain AP-1 network genes, such as Egr1, which is recognized both as a pleiotropic inflammatory trans-activator [25] and a chronic pain contributor [11], could have a chance to alleviate chronic pain. Interestingly, most of the differentially expressed miRNAs (104/120) were downregulated after CFA treatment. The global downregulation of miRNA itself is an interesting phenomenon, which was missed in the previous array-based study [17]. Notably, in these downregulated miRNAs, many of them are targeting AP-1 network genes. Thus, we speculate adding back of these miRNAs could inhibit the overactivation of AP-1 network, thus alleviate pain. Overall, we believe AP-1 network plays a central role in regulating inflammatory pain responses through a Methylation-transcription-posttranscription regulation axis.

In conclusion, this study provides a comprehensive transcriptomic profile of the CFA-induced inflammatory pain rat model via an approach that target DNA methylation, gene expression as well as post-transcriptional regulation. Our study has certain limitations. In the first place, although we included 10 pairs of rats for the CFA treatment, we only have a single time point post CFA injection, which is 24 hours. Future studies should focus on the 48hours, 72hours or even 7 days post treatment, as CFA could induce a chronic inflammation as well. In addition, although we demonstrated that AP-1 is likely to work as a regulation hub for CFA-induced inflammation response potentially, we did not include the biochemistry assays to further investigate the alteration of the AP-1 signal regulation. Future studies could focus on the effect of individual molecule or gene of AP-1 network on regulating chronic pain.
Conclusion

Our study provides the first comprehensive molecular profile of the CFA-induced inflammatory pain model using the integrated omics approach. Our results revealed AP-1 signal may play a central role in regulating pain-related signaling.

Abbreviations

CFA = Complete Freund's adjuvant
miRNA = microRNA
DRG = dorsal root ganglion
SDH = spinal cord dorsal horn
SD rat = Sprague Dawley Rat
FDR = False Discovery Rate
PWT = paw withdrawal threshold
PWL = paw withdrawal latency
DEGs = differentially expressed genes
APA = Alternative polyadenylation
DMR = differentially methylated regions

Declarations

Ethics approval and consent to participate: Approved by Nantong University.

Consent for publication: All authors consent to the publication of the manuscript.

Availability of data and material: Data are available from the corresponding author on request.

Competing interests: The authors do not have any competing interest.

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Authors’ Contribution

Lei Wei designed research and analyzed data. Feng Li, Miqun Wang, Huibin Su, Xuedong Wu, Haiyan Qiu, Wang Zhou, Chunli shan, Yuan Zhou, Gang Chen performed research and analyzed data; Xiang Zhu, Lei Wei wrote the paper.

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**Supplementary Figure Legend**
1. qRT-PCR experiments validated that both Reg3a and Reg3b gene expression were highly increased in the CFA treatment group compared to the control. P-value is calculated from Student’s t-test.

2. qRT-PCR experiments demonstrated that the two AP-1 related genes, Dusp1 was significantly increased in the CFA treated group. P-value is calculated from Student’s t-test. Egr1 was not significantly increased.

3. Western blotting assays showed a slightly decrease of Egr1 protein expression in the CFA group (p = 0.0001) by Student’s t-test.