1 Role of Chemokine and TNF signaling pathway in oral squamous cell carcinoma:

2 A RNA deep sequencing analysis of oral buccal mucosa squamous carcinoma

- 3 model of Chinese hamster
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16 Abstract

Oral cancer is one of the most common cancers in the world, meanwhile, differentially 17 expressed genes are thought to regulate the development and progression of oral 18 squamous cell carcinomas (OSCC). In this study we screened RNA transcripts from the 19 oral buccal mucosa of healthy male Chinese hamster, divided into 3 groups: a control 20 21 group with no disposal, a solvent control group coated with acetone solvent, and an experimental group coated with 0.5% DMBA acetone solution by high-throughput 22 RNA sequencing. Tophat and Bowtie were used to align the high-quality reads into 23 transcripts, DEseq was used to analysis the expression of differential gene. Then, the 24 25 Gene Ontology (GO) enrichment analysis and Kyoto Encyclopedia of Genes and Genomes (KEGG) pathway enrichment analyses were conducted. The chemokine and 26 27 TNF signaling pathway were differentially expression and the mRNA expression of CXCL1, CXCL2, CXCL3, CCL7, MMP9, monitored by qRT-PCR, increased 28 29 remarkably in the cancer group and coincided with the result of RNA-Sequencing. Meanwhile, the CXCL1, CXCL2, CXCL3, and CCL7 are significantly enriched in the 30 chemokine signaling pathway, and CXCL1, CXCL2, CXCL3, and MMP9 are 31 significantly enriched in the tumor necrosis factor (TNF) signaling pathway. The 32 33 differentially expression of the chemokine and TNF signaling pathway was a response to the invasion of the organism immune system due to oral buccal mucosa squamous 34

carcinoma. All the findings provided novel insights for further molecular researches oforal cancer.

Keywords: oral squamous cell carcinoma (OSCC), Chinese hamster, high-throughput
sequencing, differentially expressed genes (DEGs)

39 Introduction

Oral cancer is one of the most frequent solid cancers worldwide, and oral squamous 40 41 cell carcinoma (OSCC) constitutes around 90% of oral cancers (Siegel et al., 2014). It is highly invasive and metastatic at the advanced stage, and presents a substantial threat 42 to human health. Meanwhile, there are 145000 deaths of OSCC in the world (1.8% of 43 all cancer deaths in the world) annually; including 77% of the burden is in developing 44 countries (Ferlay et al., 2015). Moreover, the incidence rate of OSCC is increasing, 45 especially in younger people. Furthermore, OSCC has a very poor prognosis due to its 46 invasive nature, the survival rate of patients with OSCC has not improved despite the 47 improvements and innovations in diagnostic techniques and treatments (Shlok et al., 48 49 2010). However, clinical samples are pretty difficult to obtain and the less number of clinical samples meeting the experimental requirements is a major problem in oral 50 cancer research, which seriously restricts the development of research on the 51 mechanism of OSCC. Therefore, there is an urgent need for a better animal model of 52 human OSCC lesions to help us better understand the pathogenesis of oral cancer. 53

The hamster cheek pouch is the most relevant known animal system that closely 54 related to the human oral tumor like morphogenesis, phenotype markers and genetic 55 alterations (Raimondi et al., 2005), meanwhile, the hamster buccal pouch mucosa is 56 57 covered by a thin layer of keratinized stratified squamous epithelium that is similar in its thickness to the floor of the mouth and the ventral surface of the tongue in humans 58 (Gimenez-Conti IB and TJ, 1993). Chinese hamster (Cricetulus griseus) also called 59 striped-back hamster, is easy to operate by a single hand, has a buccal pouch on the 60 each side of oral cavity. The establishment of animal models of Chinese hamster oral 61 62 cancer can help us better study the development of oral cancer. In view of the above

advantages, Chinese hamster becomes the ideal animal model of research on OSCCmechanism.

65 Recently, the studies by Shih-Han Lee shows that, in the process of cancer, even if the DNA itself has not changed, the inactivation or alteration of the tumor suppressor 66 gene may also generate. Further studies have shown that alterations in cancer obtained 67 68 during mRNA processing can essentially simulate DNA changes in somatic cells, and patients also have tumor suppressor gene inactivation, which indicate that cancer 69 70 diagnosis of DNA alone may ignore other important molecular information that promotes disease progression (Lee et al., 2018). In common with other cancers, the 71 occurrence and progression of OSCC is a multistep process with the accumulation of 72 genetic and epigenetic changes (Kang and Park, 2001). Evidence from various 73 molecular and genetic studies suggests the association between squamous cell 74 carcinoma initiation and development and the accumulation of genetic alterations at 75 both the DNA and RNA levels (Gibb et al., 2010). Recent studies have indicated that 76 the changes of mRNA expression levels in OSCC are associated with tumor 77 78 development, maintenance, and progression. In addition, mRNAs could be potentially used as biomarkers of OSCC or other oral cancers (Lodi et al., 2010). 79

Compared to traditional techniques, next generation sequencing (NGS) offers 80 greatly improved dynamic ranges and specificity for transcriptome analyses, while 81 sample throughput is continuously increasing and costs are being reduced, and gives a 82 far more precise measurement of transcript expression levels and a far more 83 sophisticated characterization of transcript isoforms, even a far more reliable 84 characterization of allele specific expression patterns (Christopher et al., 2010). 85 86 Although there have been published reports on the successful construction of genomewide mRNA expression profiles in other types of cancer, highlighting the power and 87 capability of high-throughput sequencing techniques, there have been relatively few 88 relating to OSCC (Li et al., 2011; Zhu et al., 2011). Herein, high-throughput RNA 89 sequencing on tumor samples and their matched normal samples combined with qRT-90 PCR analysis were used to investigate the mechanisms of the occurrence and 91 92 development of OSCC. We used the information to explore and predict the molecular

93 functions and biological pathways of the differentially expressed mRNAs. Our results
94 have provided a basis for identifying further molecular markers for the diagnosis and
95 treatment of OSCC.

96

97 Materials and Methods

98 Establishment of animal model

99 Sixty male Chinese hamsters (Cricetulus griseus, aged 8-10 weeks, 21-25g b.w.) were provided by the Experimental Animal Center of Shanxi Medical University (Taiyuan, 100 China SCXK [Jin] 2015-0001), housed in standard hamster cages, maintained in a 101 temperature-controlled environment with a 12 h light/dark cycle and fed a standard 102 103 hamster diet and water ad libitum(SYXK [Jin] 2015-0001). After one-week acclimation, these hamsters were divided randomly into three groups: treatment group (24 hamsters), 104 which was coated the buccal pouch with 0.5% 7,12-dimethylbenz[a]anthracene 105 (DMBA) acetone solution; solvent control group (12 hamsters), which was only coated 106 107 with acetone solution; control group (24 hamsters), which was subjected to no disposal. The doses were chosen on the basis of the previous studies, and to observe the entire 108 carcinogenesis of oral cancer, animals were treated with DMBA three times a week for 109 15 weeks (Rajasekar et al., 2016; Ramu et al., 2017). All experimental procedures were 110 conducted and performed during the light cycle in accordance with the Animal Care 111 and Use Committee regulations of Shanxi Medical University. 112

113 Histopathological analysis

The tumor tissues were fixed in 4% paraformaldehyde for about 48 hours, transferred to 70% ethanol, then processed in a graded series of ethanol solutions, embedded in paraffin and cut into 4µm thick sections. The sections were stained with hematoxylin and eosin (H & E) for histological examination. The histopathological specimens were observed under light microscope by oral pathologist experts. The pouch pathological changes were determined according to the 12 grade record in the WHO standard (None, 1978).

121 Ultrastructure of oral buccal pouch mucosa observation

The oral buccal mucosa was extracted, washed with physiological saline, cut into 1mm³ pieces, fixed in 2.5% glutaraldehyde for 2h at 4° C dehydrated through a graded series of ethanol, embedded in epoxy resin, trimmed, sectioned, and observed and photographed under a JEM-1011 transmission electron microscope(Tokyo Japan).

126 Enzyme-linked immunosorbent assay (ELISA)

In order to investigate the serum levels of TNF- α , AFP, and SCC-Ag in serum of Chinese hamster with and without OSCC, we collected about 1 ml blood from each animal prior to surgery. Blood samples were allowed to clot for 30 min at room temperature (RT) and serum was obtained by centrifugation at 1300 g for 10 min at 4°C. Serum samples were then aliquoted and stored at – 80°C.

- 132 Serum samples were thawed on ice, the levels of TNF- α (Cat. MM-160001,
- 133 MEIMIAN, China), AFP (Cat. JL-F46753, JiangLai, China), and SCC-Ag (Cat. JL-
- 134 F46768, JiangLai, China) were detectable with the commercially available enzyme-
- linked immunosorbent assay (ELISA) kit according to manufacturer's instructions.
- 136 **RNA Sample Preparation**

All Chinese hamsters were killed by cervical dislocation on the week 15, six pairs of 137 OSCC and normal buccal pouch samples of Chinese hamster, which had been identified 138 by pathologists, were immediately isolated, frozen in liquid nitrogen and stored at -139 80 °C for RNA extraction and gene expression research. Total RNA was extracted from 140 6 pairs of OSCC and normal buccal pouch samples using TRIzol (Invitrogen, USA) 141 142 according to the protocol provided by the manufacturer. The purity and integrity of total RNA were monitored by NanoPhotometer® spectrophotometer (Implen, GER) and the 143 Agilent 2100 Bioanalyzer (Agilent, USA), respectively. 144

145 RNA Library Construction and Deep Sequencing

Briefly, mRNA was isolated from total RNA using the FastTrack MAG mRNA Isolation
Kits according to the manufacturer's instructions (Invitrogen, USA). Before doing any

further steps, Fragmentation Buffer (Agilent, USA) was applied to perform 148 fragmentation on qualified mRNA coming from buccal pouch samples of Chinese 149 hamster. Then, the mRNA was converted into double-stranded cDNA by reverse 150 transcription. Following end repair and A-tailing, paired-end sequencing adaptors 151 complementary to sequencing primers were ligated to the ends of the cDNA fragments. 152 The ligated products were purified on 2% agarose gels, qualified fragments were 153 selected for downstream enrichment by PCR. Each sample was subjected to sequencing 154 155 from both ends by the Illumina Hiseq 2500 sequencing technology.

156 **Data Filtering**

Raw image data received by sequencing were transformed by base calling (CASAVA)
into sequence data (termed Raw Reads). Raw Reads were filtered through data
processing to get Clean Reads. We removed artificial reads, adapter contamination
reads. High-quality reads with a length of >50 bp were reserved.

161 **RNA-seq Reads Alignment**

In our experiment, Tophat v2.0.12 was used to align the reads into transcripts based on
the Cricetulus griseus reference genome in the UCSC (University of California Santa
Cruz) genome database, and Bowtie2 was used to comparison analysis.

165 **Differential Expression Analysis**

166 DEseq was used to analysis the expression of differential gene, compare the expression 167 of the same gene in normal buccal pouch and buccal pouch of OSCC in Chinese 168 hamsters, and select genes with $|\log_2 \text{Ratio}| \ge 1$ and Corrected *P*-value < 0.05 as 169 differentially expressed gene to obtain up-regulated and down-regulated genes.

170 Gene Ontology Functional Analysis

For gene expression profiling analysis, Gene Ontology (GO) enrichment analyses of functional significance were performed by hypergeometric testing to map all differentially expressed genes to GO terms. The calculated *P*-value is corrected and FDR < 0.05 was used as the threshold to judge the differential gene expression of significantly enriched GO terms. The functional classification of the two groups of
genes on the GO (Gene Ontology) was compared and the top ten of the three main
ontology of GO term were selected for GO term enrichment analysis.

178 **KEGG Pathway Analysis**

The KEGG database was used to assign the assembled sequences (http://www.genome. jp/kegg), which can facilitate understanding of the biological functions of genes and identify significantly enriched metabolic pathways or signal transduction pathways in differentially expressed genes. The method of KEGG pathway analysis was the same as the GO analysis, and the significant enrichment pathway was analyzed by KAAS. FDR was used to correct the related parameters, pathways with FDR ≤ 0.05 were considered significantly enriched pathway.

186 Quantitative Real Time PCR

Quantitative real-time PCR (qRT-PCR), the traditional quantification method on gene 187 expression, was adopted to further confirm the findings from the RNA-seq analysis. On 188 the basis of the results, we filtered five significant differentially expressed genes 189 involved in chemokine signaling pathway and TNF signaling pathway. According to 190 Chinese hamster CXCL1 sequence (Accession No: NM_001244044.1), CXCL2 191 sequence (Accession No: XM_003510006.2), CXCL3 sequence (Accession No: 192 NM 001244139.1), CCL7 sequence (Accession No: XM 003495790.2), MMP9 193 sequence (Accession No: XM_007641300.1) and β -actin sequence (Accession No: 194 NM_001244575.1) in GenBank, primers were designed with Primer Premier 5 software, 195 evaluated by Oligo 6 software and subjected to Blast specificity tests at NCBI. β-actin 196 197 was used as the endogenous control and all the primers were synthesized in Huada Gene. Primer sequence information is as Table 1. 198

The qRT-PCR analysis was conducted in a total volume of 20μ L containing 10μ l 2 × SYBR® Premix Ex TaqII (Tli RNaseH Plus) (Takara Bio Inc., China), combined with sense and antisense primers (0.8 μ l, final concentration 0.4 μ M), ddH₂O 6.4 μ l, and 2 μ l diluted cDNA in Optical 8-Tube Strip using the Applied Biosystems 7300

Real-Time PCR Instrument (ABI, USA). The conditions for real-time PCR were as follows: after initial denaturation at 95°C for 30 s, 40 PCR cycles were started with thermo cycling conditions at 95°C for 5 s, 60°C for 30 s, and then a melting curve analysis was performed to verify the specificity of the PCR product. Every sample was analyzed in triplicate. System software and the $2^{-\Delta\Delta Ct}$ method were applied to quantitative calculation.

209 **Data Analysis**

210 Statistical analysis was performed using SPSS v17.0 software and Student's t-test was

used to analyze the OSCC and normal tissue samples. All values in the experiment were

expressed as mean \pm SEM (standard error of the mean) and values of P < 0.05, were

213 expressed statistically significant.

214 **Result**

215 The effect of DMBA-induced oral carcinogenesis

216 To identify cancer tissues of oral buccal pouch mucosa cancer model of Chinese 217 hamster, we examined the histological changes of buccal pouch mucosa by HE stain (Figure 1). It's important to note that in our research no significant differences were 218 observed in solvent control group compared with the control group after 15 weeks 219 acetone solution treatment. Meanwhile, when dissecting, we found that buccal pouch 220 of the cancer group became embrittled and shrinking volume. Furthermore, compared 221 with the control group, we observed the cancer group showed atypical nuclear division 222 with initial keratinization and enlarged nucleoli, cells broke through the basement 223 224 membrane, infiltrating the lamina propria and connective tissue, and tumor islands emerged, was highly differentiated squamous cell and invasive cancer. All of these 225 provided the support for sequence analysis. 226

227 The change of oral buccal pouch mucosa ultrastructure

In order to further understand the alterations in the buccal pouch of the cancer group, we used transmission electron microscope to detect them (Figure 2). Compared with

the control group, in the cancer group, the shape of the cells is irregular, the nucleolus

is concentrated, the nucleus is jagged, the nuclear membrane is invaginated, and thedesmosome is reduced or even disappeared.

233 The levels of the TNF- α, AFP, and SCC-Ag

Measures of TNF- α , AFP, and SCC-Ag in serum of OSCC Chinese hamsters were shown in Figure 3. The level of serum TNF- α in Chinese hamsters with OSCC was significantly lower than healthy control group (P<0.01). While, the level of serum SCC-Ag was higher than healthy control group (P<0.05). Among them, AFP values were not significantly different between control and OSCC.

239 Overview of Sequencing Data from RNA-seq Analysis

In both the cancer and normal groups, about $(32 \sim 44) \times 10^6$ reads $(73 \sim 92\%)$ of the total raw reads) were aligned to *Cricetulus griseus* reference genome sequence among samples, with an average of 38×10^6 reads per sample. The unique alignment sequence of the two groups accounts for $70 \sim 90\%$ of the reference genome. The distribution of the unique alignment sequences in the reference genome was shown in Figure 4.

245 Analysis of Differentially Expressed Genes

Analysis of the data indicated that there were 194 significantly differentially expressed 246 genes in the treated samples, of which 66 (34.02%) were down-regulated and 128 247 (65.97%) were up-regulated (Figure 5). The top five up-regulated and down-regulated 248 genes with the significant changes in expression were shown in the Figure 6A, and the 249 top thirty of the altered genes was listed in Table 2. The hierarchical cluster analysis of 250 251 differentially expressed genes in Cancer and Normal groups revealed that multiple gene modules were formed by many genes with similar expression trends, and these genes 252 may be involved in the occurrence, metastasis, and invasion of oral squamous cell 253 carcinoma (Figure 6). 254

255 Differential Expression GO Analysis

In the GO (Gene Ontology) statistics of differentially expressed genes, 184 genes were 256 classified according to the GO classification method, and the amount of up-regulated 257 and down-regulated differentially expressed genes in each subclass was calculated 258 (Figure 7). The GO enrichment analysis of differentially expressed genes in cancer 259 samples identified 120 biological processes, 18 cellular components, and 2 molecular 260 261 functions that are closely related to OSCC and the top ten of them are listed in the Table 3. 262

263 **KEGG Pathway Enrichment Analysis**

Based on the analysis of KEGG pathway enrichment, we found that differentially 264 expressed genes in the cancer group were mainly enriched in 8 signaling pathways 265 266 including tumor necrosis factor pathway (TNF), chemokine pathway, cytokine interaction pathway, extracellular matrix receptor interaction pathway, protein digestion 267 pathway and absorption pathway; and the number of enriched gene in the cytokine 268 interaction pathway and chemokine pathway is the highest, suggesting that these 269 270 pathways may play an crucial role in tumorigenesis, invasion and metastasis (Table 4).

We screened five differentially expressed genes related to OSCC, the Table 5 clearly 271 reflects that CXCL1, CXCL2, CXCL3, CCL7, and MMP9 are enriched in multiple GO 272 terms and KEGG pathways. As is depicted in the Figure 8(Kanehisa et al., 2017), 273 CXCL1, CXCL2, CXCL3, and CCL7 are significantly enriched in the chemokine 274 signaling pathway (Figure 8A), whereas CXCL1, CXCL2, CXCL3, and MMP9 are 275 significantly enriched in the tumor necrosis factor signaling pathway (Figure 8B), 276 suggesting that CXCL1, CXCL2, CXCL3, CCL7, and MMP9 may participate in the 277 278 development of OSCC by molecular dialogues and interactions in metabolic pathways.

279

Expression Analysis by qRT-PCR

Based on the RNA deep sequencing analysis of the expression, five genes, including 280 CXCL1, CXCL2, CXCL3, CCL7 and MMP9, were selected for confirmation as well 281 as to monitor their expression with qRT-PCR, and the data was statistically analyzed as 282 follows (figure 9). 283

Compared with normal group, in the cancer group the mRNA expression level of 284 CXCL1, CXCL2, CXCL3, CCL7 and MMP9 increased remarkably. Meanwhile the 285 286 transcriptome sequencing revealed that the differentially expressed genes: CXCL1, CXCL2, CXCL3, CCL7 and MMP9 were also significantly up-regulated in the cancer 287 group compared to the normal group. The relative expression of mRNA detected by 288 qRT-PCR and the transcriptome sequencing data (RPKM) were highly consistent, 289 indicating that the sequencing data was reliable, suggesting that CXCL1, CXCL2, 290 291 CXCL3, CCL7 and MMP9 may promote the development, invasion and metastasis of OSCC. 292

293 Discussion

Oral cancer is the most common malignancy of head and neck cancer in the world and 294 it is a leading cause of cancer-related mortality. Epidemiological data have shown that 295 high risk factors, such as tobacco, alcohol and Human papillomavirus infection are 296 297 closely related to OSCC (Chi et al., 2015). Recently, the incidence of oral cancer is rising with ages every year and more seriously, the patients are mostly diagnosed at a 298 299 relatively late stage which lead to increase in the cost of treatment and decrease in treatment outcomes(Oh et al.). In the present biological era, the prognosis of this deadly 300 disease has improved to some extent because of the enhancement of technologies 301 (Frédérique et al., 2012), but diagnosing the tumor at its initial stages relapse, and 302 metastasis are the major challenge to improve the scope of patient survival (Dahiya and 303 Dhankhar, 2016). Therefore, it is urgent and imperative to delve into the molecular 304 mechanisms of the development and progression of oral cancer to guide clinical 305 306 treatment and prognosis.

Since Sally first established the DMBA-induced oral carcinogenesis model in the cheek pouch of hamster in 1954, it has become a classic animal model of OSCC (Salley, 1954). In present study, we successfully constructed the animal model of OSCC using tri-weekly applications of a 5% solution of DMBA in acetone onto the cheek pouch of Chinese hamsters over about a 15-week period. For models like the hamster model for

OSCC, high-throughput sequencing provides a powerful tool for analyzing both mRNA 312 expression patterns and quantitative expression levels, as it profiles thousands of genes 313 simultaneously. New high-throughput sequencing technologies have enabled the 314 detection of novel transcripts through increased sensitivity. These recent advances have 315 facilitated more comprehensive and more thorough research into the effects of 316 transcription and translation (Zeng JH et al., 2019). This technology is much more 317 efficient than the now outmoded and time-consuming methods used in earlier work, 318 and is becoming the broadest transcriptome research tool available. 319

Different from earlier studies, our study considered the development and progression 320 of OSCC holistically, including a variety of pathways and genes, rather than just a single 321 factor. Using high-throughput sequencing analysis, we evaluated mRNA expression 322 profiles of OSCC and normal cheek pouch mucosa tissues, and identified 128 mRNAs 323 324 that were up-regulated and 66 down-regulated in cancer tissues compared with normal tissues. What's more, because genes often cooperate with each other to perform their 325 biological functions, pathway analyze facilitate further understanding genes and their 326 327 roles. By the KEGG enrichment analysis, we found that differentially expressed genes were enriched in 317 pathways, of which 8 pathways were significantly enriched. In 328 329 addition, we used GO analysis to screen genes which were significantly enriched in pathways, and selected five genes(CXCL1, CXCL2, CXCL3, CCL7, and MMP9) that 330 were highly expressed in OSCC, may be closely related to the development of oral 331 cancer and significantly enriched in chemokine signaling pathway and TNF signaling 332 pathway for qRT-PCR to validate transcriptome sequencing. The GO functional 333 enrichment analysis showed that the differentially expressed mRNAs in OSCC were 334 335 mainly enriched in cellular components, including extracellular matrix, extracellular space and extracellular region, etc; involved molecular functions that included cytokine 336 activity and extracellular matrix structural constituent; involved biological processes 337 that included granulocyte chemotaxis, granulocyte migration, cell chemotaxis, cell 338 migration, cell motility and localization of cell, etc. RT-qPCR confirmed that the 339 expression of CXCL1, CXCL2, CXCL3, CCL7 and MMP9 were consistent with RNA-340 seq. Meanwhile, qRT-PCR confirmed that CXCL2 had the largest differential 341

expression folds in OSCC tissues, revealing that CXCL2 may play a central role in the 342 molecular mechanism of OSCC. The high expression of CXCL2 suggests that it can be 343 344 used as a new target for diagnosis, treatment and prognosis of OSCC, and it also provides novel ideas and important theoretical basis for the search for potential 345 molecular markers of oral cancer. These findings suggest that there is still an 346 inflammatory response during OSCC and the previous differential expression analysis 347 also reflected that chemokine is still highly expressed in OSCC. In summary, the 348 inflammatory response is in an uncontrollable state and it is difficult to restore the 349 body's homeostasis, eventually resulting in the production of tumors. 350

Chemokines are a group of small proteins(8–12 KDs), responsible for transmitting 351 signals for cell migration, inflammation regulation, and angiogenesis, classifed into 352 four subgroups referred to as CXC/ α , CC/ β , CX3C/ δ or C/ δ families (Baggiolini, 1998; 353 Zlotnik and Yoshie, 2000). They are mostly known for their role in immunesurveillance 354 and infammatory responses, but they have been also implicated in many pathological 355 processes of malignant tumor (Mahboobeh et al., 2014). Recently, the research of 356 chemokines in the oncology has drawn extensive attention. Meanwhile, related studies 357 have demonstrated the involvement of chemokines and their receptors in various tumors 358 such as liver cancer, nasopharyngeal cancer, and breast cancer (Jing et al., 2016; Shah 359 et al., 2015; Weitzenfeld et al., 2016). Abnormal function of chemokines in cancer 360 promotes cell survival, facilitated proliferation, angiogenesis, and metastasis in 361 multiple types of tumors (Paola et al., 2011). The research of Vera Levina indicate that 362 chemokines and growth factors produced by tumors by binding to the cognate receptors 363 on tumor and stroma cells could provide proliferative and anti-apoptotic signals helping 364 365 tumors to escape drug-mediated destruction (Vera et al., 2010). Furthermore, it is believed that chronic inflammatory conditions facilitate oral carcinogenesis, and 366 functions of cytokine-dependent and chemokine-dependent immunoregulatory 367 368 pathways are apparent in oral carcinoma.

Among them, Jung(Da-Woon et al., 2010) confirmed that *CCL7* may participate in the invasion and metastasis of OSCC through a molecular dialogue with the *CCR1* and *CCR3*. In this study, the chemokine *CXCL2*, *CXCL3*, and *CXCL1* were all highly

expressed in OSCC, and Peng (Peng et al., 2015)also detected significant upregulation 372 of CXCL2 and CXCL3 by microarray analysis of OSCC in rats, indicating abnormal 373 expression of CXCL1, CXCL2 and CXCL3 is an important factor leading to pathological 374 changes in the oral cancer model. Furthermore, OSCC microarray analysis also 375 reflected abnormal expression of CXCL1, CXCL2 and CXCL3 in tumor tissues 376 (Sanjukta et al., 2015), while clinical and follow-up basic studies on CXCL1, CXCL2, 377 and CXCL3 are rare in oral cancer and most of them are found in liver cancer, prostate 378 379 cancer and breast cancer (Gui et al., 2016; Li et al., 2015; See et al., 2014). Thence, the further study of these significantly different chemokines in the pathogenesis has 380 important implications for the diagnosis and treatment of oral cancer. 381

Matrixmetallo-proteinases (MMPs) are major proteolytic enzymes that are involved 382 in the normal extracellular matrix (ECM) turnover. Its main function is to degrade and 383 remodel the ECM, maintain the dynamic balance of ECM, and participate in many 384 pathological and physiological processes in the body. Many of clinical studies have 385 confirmed the high expression of MMP9 and MMP13 in OSCC (Jingqiu et al., 2015; 386 387 Monteiro et al., 2016; Nanda et al., 2014), which is consistent with our research. Monteiro (Monteiro et al., 2016) applied immunohistochemistry to detect the 388 expression of MMP9 protein in 60 cases of OSCC and found that MMP9 was strongly 389 expressed in cytoplasm of tumor cell in 83.7% of patients, suggesting that MMP9 is a 390 potential tumor marker of oral cancer. Yu (Yu et al., 2011) knocked out the chemokine 391 receptor CXCR4 gene in Tca8113 cells, as a result, the expression levels of MMP9 and 392 MMP13 were significantly reduced, and the invasion and metastasis of cells were 393 weakened, suggesting that MMP9 and MMP13 play a pivotal role in the invasion and 394 395 metastasis of OSCC. The matrix metalloproteinase we screened is MMP9, which is enriched with CXCL1, CXCL2, and CXCL3 in TNF signaling pathway, suggesting that 396 MMP9 may promote tumorigenesis through the interaction with chemokines. Gao et al. 397 (Gao et al., 2015) said that CXCL5/CXCR2 axis activates the PI3K/AKT signaling 398 pathway of the tumor signaling pathway, leading to the up-regulation of MMP2 and 399 MMP9, which promotes the metastasis of bladder cancer, confirming the interaction 400 between MMP9 and chemokines. However, the specific interaction between MMP9 and 401

402 *CXCL1, CXCL2, CXCL3* in OSCC still needs further study. Judging from these figures, 403 we can draw the conclusion that the expression trend of the five genes in qRT-PCR was 404 consistent with the transcriptome sequencing, which validated the reliability of RNA-405 seq data and further demonstrated that the chemokine signaling pathway and TNF 406 signaling pathway may closely related to the occurrence and development of OSCC.

407 To our knowledge, our study presents the first genomewide profiling of mRNAs of squamous cell carcinoma in oral buccal pouch of Chinese hamster by high-throughput 408 409 RNA-Seq. In addition, our findings support previous studies reporting that the progression of OSCC was influenced by chemokines, suggesting that chemokine 410 signaling pathway, Cytokine-cytokine receptor interaction signaling pathway, and TNF 411 signaling pathway may play a central role in the invasion and metastasis of OSCC 412 (Farkas et al., 2013; Toung et al., 2011). Peeping a spot to see overall picture: local 413 delicate change was packed with the complex issues of the whole organism. But for the 414 further verification and exploration, researches on the cellular level and the significant 415 expression of proteins during the development and progression of OSCC should be 416 carried out. 417

418 Conclusions

All the findings, including the chemokine signaling pathway, TNF signaling pathway 419 and classic genes provided an extensive bioinformatics analysis of DEGs and revealed 420 a series of targets and pathways, which may affect the carcinogenesis and progression 421 of OSCC, for future investigation. These findings add to significant insights into the 422 diagnosis and treatment of this disease. Meanwhile, it should be noted that this study 423 424 examined the DEGs in oral cancer Chinese hamster animal model by qRT-PCR and bioinformatics analysis; further research needs to be done to explore more specific 425 mechanisms. Notwithstanding its limitation, these findings significantly improved the 426 understanding of underlying molecular mechanisms in OSCC, and the key genes and 427 pathways could be used as diagnostic and therapeutic targets and diagnostic biomarkers. 428

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433 **Competing interests**

The authors declare no competing or financial interests.

435 Author contributions

G.H.S. designed the study and contributed funding. G.Q.X., J.N.W., and B.HF.
established animal model, completed RNA sequencing and statistical analyze. L.F.X,
X.T.W., J.P.G. AND R.J.X. collected samples and processed samples. Z.Y.C. provided
the necessary tools and instruments for the experiments. G.Q.X. and J.P.G. contributed
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565 Figure legend

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581

Figure 3. The levels of the AFP, SCC-Ag, and TNF- α in oral buccal pouch mucosa between control and cancer group. TNF- α in Chinese hamsters with OSCC was significantly lower than healthy control group (P<0.01). While, the level of serum SCC-Ag was higher than healthy control group (P<0.05). Among them, AFP values were not significantly different between control and OSCC.

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Figure 4. Sequence quality and unique alignment sequence distribution of cancer and
normal groups. A: Sequencing quality distribution of the samples. B: The distribution of the
unique alignment sequence in each region of genes in the reference genome.

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592 Figure 5. Histograms and volcano plot differentially expressed gene in cancer and normal 593 groups. A: Histogram, Y-axis is the amount of differentially expressed genes, X-axis is the 594 classification of differentially expressed genes, red indicates up-regulated genes, green indicates 595 down-regulated genes, and blue indicates total differentially expressed genes. B: volcano plot, Yaxis is -log10 (P-value) of differentially expressed genes, X axis is the log2 FoldChange of the 596 597 differentially expressed genes, red indicates up-regulated genes and green indicates down-598 regulated genes. There were 66 (34.02%) were down-regulated and 128 (65.97%) were upregulated. 599

600

Figure 6. Hierarchical clustering of differentially expressed genes in oral squamous cell carcinoma and normal tissues. A: Hierarchical clustering graph, each rectangle represents the expression value of a certain gene (row) in a certain sample (column), and the gene expression changes from blue (low expression) to red (high expression). B: The top five up-regulated and downregulated genes.

606

Figure 7. GO classification statistics histogram of differentially expressed genes between
 cancer and normal groups. Red indicates up-regulated genes, green indicates down-regulated
 genes, and all differentially-expressed genes are classified into 67 different subclasses (third-level
 entry) according to GO classification.

- 611
- **Figure 8.** A: chemokine signaling pathway. B: tumor necrosis factor signaling pathway.
- All signaling pathways are cited from www.kegg.jp/kegg/kegg1.html.
- 614

Figure 9. qRT-PCR validation transcriptome sequencing results. The first line is the relative
expression level of differentially expressed genes in oral buccal mucosa tissues of Chinese hamster
detected by qRT-PCR (n=3). The second line is the expression level of differentially expressed genes
in each tissue detected by transcriptome sequencing (RPKM, n=3).

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Table:

Gene	Sequence	Lengtl
CXCL1	5'-GTGTCAACCACTGTAAGAGAAGCA-3'	24
CACLI	5'-ACACATTCCCTCACCCTAATACAAA-3'	25
CXCL2	5'-CCAGACAGAAGTCATTGCCACTC-3'	23
CACL2	5'-GCCTTGCCTTTGTTCAGTATCTTT-3'	24
CVCL 2	5'-TGAGGCAGGAAAGGAGGAAG-3'	20
CXCL3	5'-TGTTCAAAGCAAACAGGAGAGG-3'	22
CCL7	5'-CCCTGGGAAGCTGTGATCT-3'	19
	5'-GGCTTTGGAGTTTGGGTTTTC-3'	21
MMP9	5'-CCTTGTCACTTTCCCTTCACCTT-3'	23
	5'-ATTTGCGGTCGGTGTCGT-3'	18
β-Actin	5'-GTGCTGTCCCTGTATGCCTCT-3'	21
	5'-GTCACGCACAATTTCCCTCTC-3'	21

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Table2. Top thirty genes altered in cancer group.

Gene	Log ₂ Ratio	Corrected <i>P</i> -value [*]	Gene	Log ₂ Ratio	Corrected <i>P</i> -value [*]
CXCL2	26.69	5.54E-15	CCL7	5.88	2.13E-05
MMP13	11.69	1.18E-14	IL1B	11.53	2.13E-05
CCL3	26.24	4.13E-14	PTX3	7.31	2.38E-05
LCN1	25.19	1.27E-10	CAIV	5.85	2.38E-05
MMP9	7.42	4.35E-09	CXCL7	22.94	4.54E-05
CXCL3	6.52	1.42E-07	OLR1	22.97	4.61E-05
STRA6	7.15	1.68E-07	RN45s	-8.19	5.53E-05
LOC100754872	23.63	6.98E-07	THBS1	4.72	6.72E-05
Serpine1	6.05	8.31E-07	IL-36 y	6.67	9.12E-05
TNN	8.20	1.23E-06	CXCL1	5.77	0.0001
CTHRC1	8.46	1.35E-06	LOC100766767	-7.59	0.0001
GPx6	-24.40	4.67E-06	MS4A7	5.99	0.0001
CTRP6	5.52	6.88E-06	SFRP2	5.02	0.0001
LOC100764819	-8.33	1.40E-05	POSTN	4.81	0.000115
SLN	-6.62	1.49E-05	MMP12	5.23	0.000121

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Table3. GO enrichment analysis of differentially expressed genes in cancer and normal groups.

635	(FDR<0.05) (BP: Biological Process; CC: Cellular Component; MF: Molecular	r Function)
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GO name	GO	GO classification	Number	<i>P</i> -value	FDR
GO hame	category	number	of genes	1 vulue	TDR
granulocyte chemotaxis	BP	GO:0071621	11	4.40E-12	5.60E-08
granulocyte migration	BP	GO:0097530	11	1.00E-11	6.36E-08
cell chemotaxis	BP	GO:0060326	15	2.20E-11	8.27E-08
leukocyte chemotaxis	BP	GO:0030595	13	2.60E-11	8.27E-08
myeloid leukocyte migration	BP	GO:0097529	12	6.10E-11	1.55E-07
cell migration	BP	GO:0016477	31	1.10E-10	2.33E-07
cell motility	BP	GO:0048870	32	1.70E-10	2.70E-07
localization of cell	BP	GO:0051674	32	1.70E-10	2.70E-07
leukocyte migration	BP	GO:0050900	15	2.30E-10	3.25E-07
neutrophil chemotaxis	BP	GO:0030593	9	4.80E-10	6.11E-07
extracellular matrix	CC	GO:0031012	29	2.10E-19	3.04E-16
extracellular space	CC	GO:0005615	42	1.30E-18	9.40E-16
extracellular region	CC	GO:0005576	85	3.00E-18	1.45E-15
proteinaceous extracellular matrix	CC	GO:0005578	25	3.60E-17	1.30E-14
extracellular region part	CC	GO:0044421	73	8.20E-17	2.37E-14
extracellular matrix component	CC	GO:0044420	11	8.10E-09	1.95E-06
sarcoplasmic reticulum	CC	GO:0016529	8	1.70E-08	3.51E-06
sarcoplasm	CC	GO:0016528	8	4.20E-08	7.59E-06
collagen trimer	CC	GO:0005581	9	7.80E-08	1.25E-05
striated muscle thin filament	CC	GO:0005865	4	1.50E-06	2.17E-04
cytokine activity	MF	GO:0005125	7	6.70E-06	0.026
extracellular matrix structural constituent	MF	GO:0005201	5	2.60E-05	0.049

KEGG signal pathway	Map	Count	Q-value	Genes
Malaria	map05144	5	0.002	CSF3;IL1B;THBS2;SELP;CCL2
ECM-receptor interaction	map04512	6	0.005	LOC103164586;TNC; THBS2;TNN;SPP1;COL1A1
Salmonella infection	map05132	6	0.005	CXCL1;CXCL3;CXCL2;IL1B; CCL4; CCL3
Chemokine signaling pathway	map04062	8	0.005	CCL7;CXCL1;CXCL3;CXCL2; CCL4;CCL3;CCL2;CXCL7;
Protein digestion and absorption	map04974	6	0.005	LOC10316458; OL18A;COL7A; COL12A;COL1A1; SLC1A1;
Cytokine-cytokine receptor interaction	map04060	8	0.006	CSF3R;CCL7;CSF3;IL1B; CCL4;CCL3;CCL2;CXCL7
TNF signaling pathway	map04668	6	0.008	CXCL1;CXCL3;CXCL2; MMP9;IL1B;CCL2;
Toll-like receptor signaling pathway	map04620	5	0.047	IL1B;CCL4;CCL3;SPP1;CTSK;

6	4	2	

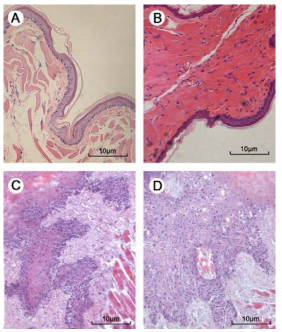
643 Table5. GO and KEGG enrichment pathways of 5 screened genes TOP 3 GO-BP Gene P-value GO-BP KEGG enrichment pathway enrichment pathway CXCL1 0.0001 13 Salmonella infection cell chemotaxis cell migration Chemokine signaling pathway cell motility TNF signaling pathway CXCL2 41 granulocyte chemotaxis Salmonella infection 5.54E-15 granulocyte migration Chemokine signaling pathway cell chemotaxis TNF signaling pathway CXCL3 1.42E-07 33 granulocyte chemotaxis Salmonella infection granulocyte migration Chemokine signaling pathway cell chemotaxis TNF signaling pathway CCL7 101 2.13E-05 granulocyte chemotaxis Chemokine signaling pathway granulocyte migration Cytokine-cytokine receptor interaction cell chemotaxis MMP9 4.35E-09 60 cell migration cell motility TNF signaling pathway localization of cell

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647 **Figure:**

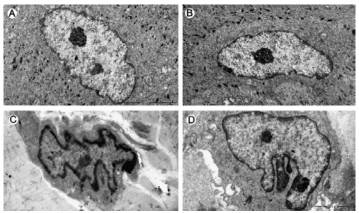
648 Figure 1



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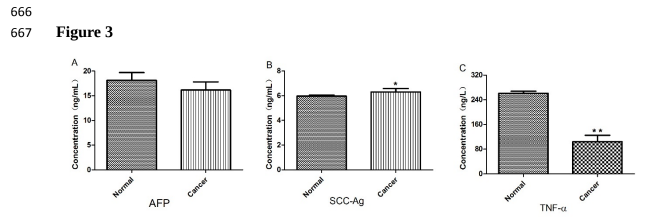
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the muscular layer, and the basal cells are arranged in a neatly arranged shape. Meanwhile, in C &
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657 Figure 2



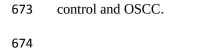
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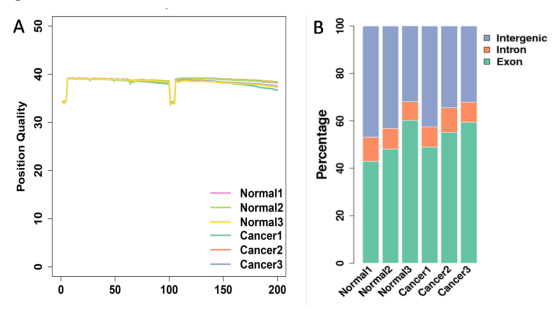




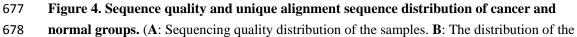
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675 **Figure 4**

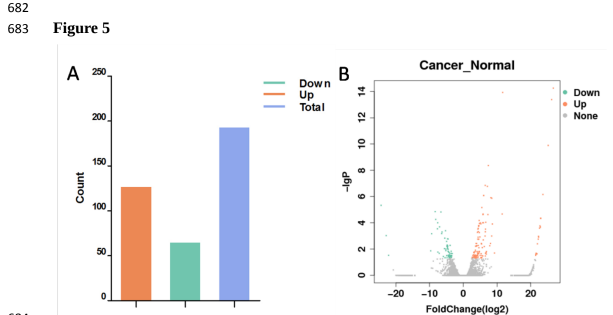


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unique alignment sequence in each region of genes in the reference genome.)

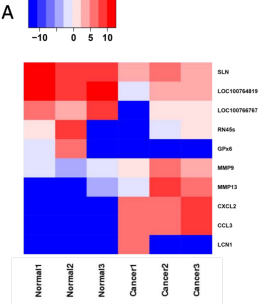
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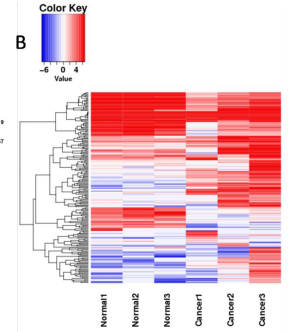




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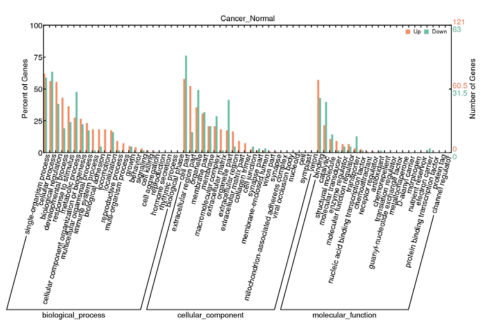




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- 701
- 702 Figure 7



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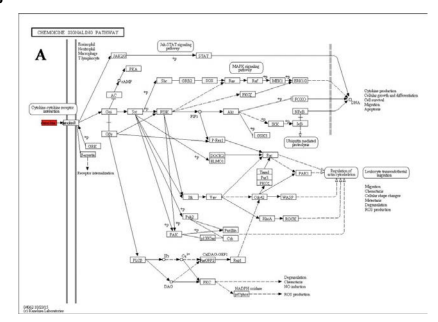
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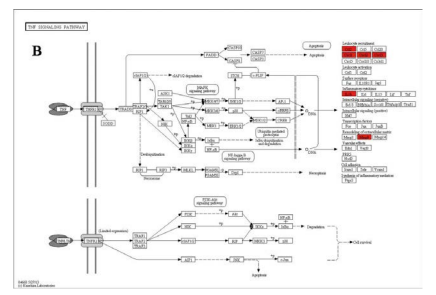
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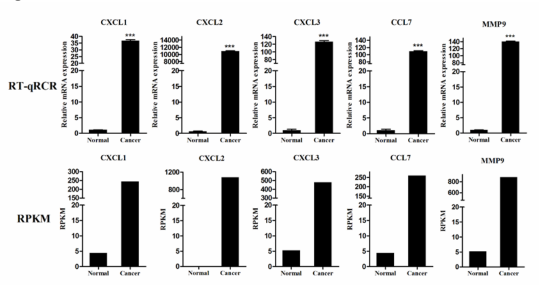


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