



Whole-exome sequencing reveals genetic underpinnings of tongue carcinoma in Chinese population

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Abstract

Oral tongue squamous cell carcinoma (OTSCC) is a common malignancy, of which the incidence has increased in China in the last decade. Surprisingly, while multiple studies have revealed the mutational features of OTSCC in Western populations, limited data was shown in Asian patients. Herein, we utilized whole-exome sequencing to profile the genetic alterations in 13 Chinese OTSCC and compared them to those from 40 Western patients published in *Cancer Discovery*. In result, some key driver mutations were observed in both Chinese and Western cohorts, such as TP53 (Chinese 60.0% vs Western 60.0%), FAT1 (Chinese 7.7% vs Western 30.0%), CASP8 (Chinese 7.7% vs Western 10.0%) and NOTCH1 (Chinese 15.4% vs Western 10.0%), while mutations in CDKN2A (23.1%) and NTRK3 (23.1%) were only observed in Chinese patients, indicating these two novel mutations might play vital roles in OTSCC tumorigenesis specifically in Asian population. Mutational signatures depicted both common and distinct features across cohorts. In addition, significant copy number loss was found in 7q22.1, 9q13.1, and focal regions spanning CDKN2A and CDKN2B. FOXP1-TEX261 (2p13.3:3p13) fusion, reported in various cancer types, was firstly observed in OTSCC. Also, we identified numerous actionable mutations with FDA approved targeted. Taken together, our study revealed the mutational features of Chinese OTSCC patients, either similar or distinct to those of Caucasian patients. CDKN2A and NTRK3 were observed as two novel drivers that might play essential roles in tumorigenesis in Chinese patients, and were found as two potential therapeutic targets, rendering it promising to develop novel therapies.

Key words Actionable mutation, copy number alteration, oral tongue squamous cell carcinoma, whole-exome sequencing

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Introduction

Oral tongue squamous cell carcinoma (OTSCC) is a common malignancy arising in oral cavity [1], accounting for 354,864 new cancer cases and 177,384 cancer deaths annually worldwide and is particularly prevalent in developing countries [2]. Given the different environment and living lifestyle across the world, different populations might have been exposed to different carcinogens, leading to distinct mutational backgrounds and molecular features underlying tumorigenesis [2]. To date, next-generation sequencing has unearthed key genetic alternations in Caucasian OTSCC population, including somatic mutations in canonical tumor suppressor genes and oncogenes like TP53 and NOTCH1 [3-5]. Nonetheless, limited reports have characterized the genetic underpinnings of OTSCC in Asian population. Herein by utilizing whole-exome sequencing, we scrutinized the mutational features of 13 OTSCC in China, shedding more light on the genetic mechanisms of OTSCC tumor progression in the Chinese population. We further integrated WES data of 40 Caucasian patients published on Cancer Discovery [5], exhibiting both common and unique genetic characteristics between Chinese and Western cohorts. Besides, novel actionable mutations were observed, rendering it promising to develop new targeted therapies for OTSCC patients.

Materials and methods

Patients Enrollment, Sample Collection, Preparation and Ethics

13 patients diagnosed with OTSCC were enrolled from Cancer Hospital Chinese Academy of Medical Sciences from 2012 to 2019. Detailed clinicopathological characteristics of all patients were shown in **Table S1**. All patients underwent resection. Primary tumor tissues together with matched normal specimens were collected surgically and snap frozen in liquid nitrogen and stored at -80. DNA extraction was performed with TIANamp Genomic DNA Kits (Tiangen Biotech, Beijing, China) according to manufacturer's instructions. Written informed consent was collected in sample collection, gene sequencing and data publication. This study was approved by the Institutional Review Board (IRB) of Cancer Hospital Chinese Academy of Medical Sciences and conducted in accordance with the Declaration of Helsinki.

Library Construction and Whole-exome Sequencing

Library construction was performed using a custom 53M length capturing probe, made by Integrated DNA Technologies (IDT, IA, USA). Captured libraries were then pair-end sequenced in 100-bp lengths with Geneplus-2000 sequencing platform (Geneplus, Beijing, China) following the manufacturer's guidance. Reads were further mapped to the reference human genome (hg19) utilizing BWA aligner (version 0.7.10) for mutation calling after filtration.

Mutation Calling

Somatic single nucleotide variants (SNVs), small insertions and deletions (Indels) were detected using MuTect (version 1.1.4) packed in GATK (version 4.0). Variants with high frequencies among unrelated individuals across the world or specifically in east Asia were firstly filtered out as previously reported [6] and were listed in **Table S2**, then further filtered with the method demonstrated in a study [7] in order to screen out the key cancer drivers or cancer associated mutations. In brief, candidate alterations had to be mutations in known cancer associated genes

collected in either Catalogue of Somatic Mutations in Cancer (COSMIC), OncoKB, CIVIC (Clinical Interpretation of Variants in Cancer), CGI (Cancer Genome Interpreter) or IntOGEN (Integrative Onco-Genomics) database (all accessed in July, 2020). Mutations including nonsense, in-frame/frame-shift small insertions and deletions, and mutations in canonical splice-sites were kept. In regard of missense mutations, only those reported in COSMIC with a FATHMM-MKL (Functional Analysis through Hidden Markov Models) score of >0.5 (accessed in August 2020), or those identified in two or more of the following in silico functional analysis algorithms: predication score 0-0.05 in SIFT (Sorting Intolerant from Tolerant), "possibly damaging" or "probably damaging" in Polymorphism Phenotyping-2 (Polyphen2) or predication >0.5 in FATHMM-MKL were kept.

To study the significantly altered copy number of chromosomal segments, somatic copy number alterations (SCNA) was identified employing GATK (version 4.0) and GISTIC (version 2.0). An in-house algorithm was employed in detecting split reads and discordant read-pairs to identify chromosomal structural variation [6, 8]. Eventually, all candidate somatic variations were manually verified on the integrative genomics reviewer (IGV) browser to filter out the false positive.

To compare the mutational features between Chinese and Caucasian OTSCC populations, mutation data from WES of 40 Caucasian OTSCC patients, published in 2013 on Cancer Discovery [5] were downloaded from cBioPortal for Cancer Genomics database, with detailed clinicopathological information exhibited in **Table S1**. Mutations in genes collected in either COSMIC, OncoKB, CIVIC, CGI or IntOGEN database were kept and subjected to following analyses. To find potential therapeutic targets, we mapped the unfiltered raw SNVs and Indels to OncoKB actionable mutation catalogue (accessed in July 2020), so as to avoid missing any potentially valuable candidates.

Analysis of Mutational Signature and Clonal Architecture

To find the possible mechanisms underlying OTSCC mutagenesis and reason the potential carcinogen-exposure or DNA damage repair deficiency of patients, the trinucleotide altering patterns of unfiltered somatic SNVs and fragment changing patterns of unfiltered Indels were matched to the known signatures described in COSMIC database using YAPSA (version 3.11). Contribution of each signature in samples was quantified and only signatures with contributions of more than 1% of all alterations were illustrated and considered in the subsequent analyses. Clonality of mutational events were inferred by looking at the variant allele frequency (VAF) ratio of mutations, calculated in the way as previously reported [9]. Specifically, VAF ratio was calculated by dividing the VAF of each mutation by the maximum VAF observed in the same sample. A higher VAF ratio suggested the respective event occurred at an earlier stage during tumor progression. Mutations with VAF ratios > 0.75 were determined as clonal mutations while the rest were considered as subclonal mutations.

Statistics

Two-sided Mann-Whitney and Fisher's exact tests were performed on Graphpad Prism (version 7.01) or R (version 3.6.1) to generate the P value. All data processing, analyses and plotting were carried out utilizing R software (version 3.6.1) and/or GraphPad Prism software (version 8.0.2). Statistical significance was defined as P<0.05 for two-tailed student t-tests.

Results

Mutation Landscape of Somatic Point Mutations

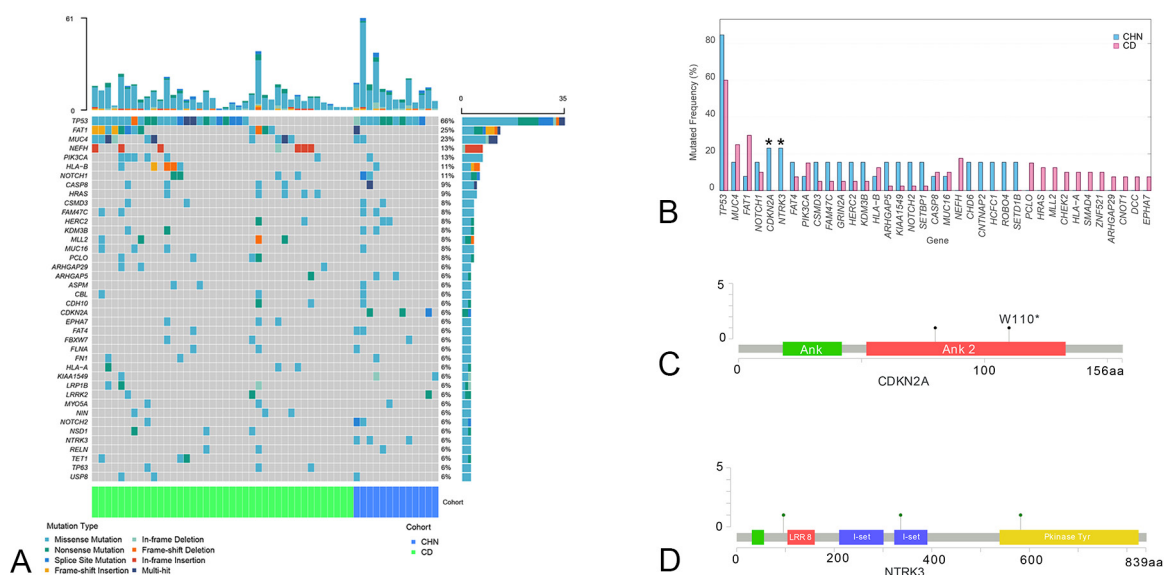


Figure 1. Somatic Single Nucleotide Variations and Small Insertions and Deletions Detected; **A.** Landscape of all mutations identified, panel at the top, mutation counts in each patient, right panel, number of patients harboring mutations in specific genes, panel at the bottom, source of patients, CHN- Chinese patients in our cohort (N = 13); CD-Patients published in Cancer Discovery (N = 40); **B.** Prevalence of Mutations in each cohort, * represents a P value < 0.05; **C.** Locations of sites of CDKN2A somatic SNVs/Indels in our cohort, Ank - ankyrin repeats domain; **D.** Locations of sites of NTRK3 somatic SNVs/Indels in our cohort, LRR8 - Leucine-rich repeat domain; I-set - Immunoglobulin I-set domain; Pkinase Tyr - Tyrosine protein kinases specific active-site signature domain.

Whole-exome sequencing identified a total of 1,456 somatic SNVs and Indels in 13 Chinese OTSCC patients (CHN cohort) before filtration, while 3, 471 mutations in 40 Caucasian patients published in Cancer Discovery (CD cohort) were reported [5]. After filtration, 235 and 536 mutations in known cancer-associated genes were identified in CHN and CD cohorts respectively. In CHN cohort, genes with the highest mutation frequencies were TP53 (11/13, 84.6%), CDKN2A (3/13, 23.1%) and NTRK3 (3/13, 23.1%) (Figure 1A). Other recurrently mutated genes included ARHGAP5, CHD6, CNTNAP2, CSMD3, FAM47C, FAT4, GRIN2A, HCFC1, HERC2, KDM3B, KIAA1549, MUC4, NOTCH1, NOTCH2, ROBO4, SETBP1 and SETD1B, all mutated in 2/13 patients (15.4%) (Figure 1A). Mapping mutations into Reactome (accessed in September 2020) database suggested that mutations were enriched in NOTCH signaling pathway, RUNX3 signaling pathway, VENTX pathway and pathways involved in cellular senescence, which are closely related to cell cycle, apoptosis and stem cell phenotypes, and were shown to be associated with tumorigenesis in various cancer types [10-13]. In regard of CD cohort, recurrently mutated genes included TP53 (24/40, 60%), FAT1 (12/40, 30%), MUC4 (10/40, 25%), NEFH (7/40, 17.5%), PCLO (6/40, 15%), PIK3CA (6/40, 15%), HRAS (5/40, 12.5%), HLA-B (5/40, 12.5%), MLL2 (5/40, 12.5%), CASP8 (4/40, 10%), CHEK2 (4/40, 10%), HLA-A (4/40, 10%), MUC16 (4/40, 10%), NOTCH1 (4/40, 10%), SMAD4 (4/40, 10%) and ZNF521 (4/40, 10%) (Figure 1A).

We then compared the frequencies of mutations in known cancer-associated genes between CHN and CD cohorts (Figure 1B). In result, we observed similar mutational frequencies in some canonical cancer driver genes between two cohorts, such as TP53 (CD 60% vs. CHN 60%), FAT1 (CD 30% vs. CHN 7.7%), CASP8 (CD 10% vs. CHN 7.7%) and NOTCH1 (CD 10% vs. CHN

15.4%), indicating that they might play pivotal roles in OTSCC tumorigenesis. Notably, mutation frequencies of CDKN2A and NTRK3 in our cohort (3/13, 23.0% for both) were observed significantly higher than in CD cohort (0 for both; P = 0.012 for both; Figure 1B), suggesting CDKN2A and NTRK3 might be two novel drivers in OTSCC cancer progression among Chinese population.

To further justify the association of CDKN2A and NTRK3 with OTSCC tumorigenesis, we looked at the locations of their altered nucleotides on the protein-annotated chromosomes accordingly. As shown in (Figure 1C & Figure 1D), 2/3 missense mutations in NTRK3 located in the functional domains, namely immunoglobulin I-set domain and protein tyrosine kinase domain, whereas another mutation located in a non-functional region. As for CDKN2A mutations, 2/3 truncating nonsense mutations located in ankyrin repeats domain while another mutation located at a canonical splice site.

Mutational Signatures of Somatic SNVs and Indels in OTSCC

To gain an insight into OTSCC cancer etiology, we then matched the altering patterns of raw somatic SNVs and Indels to COSMIC single base signatures (SBS) and insertion and deletion signatures (ID) accordingly and quantified the contribution of each signature in both CHN cohort and CD cohort. In result, somatic SNVs were mainly attributed to three SBSs in COSMIC database, SBS39, SBS7b, SBS1 specifically, while Indels were attributed to ID1-14, ID16 and ID17 (Figure 2A).

SBS39, featured with predominant C > G alterations, was observed with drastically high distribution in head and squamous cell carcinoma and breast cancer and without known proposed etiology [14]. SBS7b, characterized by C > T mutations, was frequently found in skin cancer and was reported to be due to

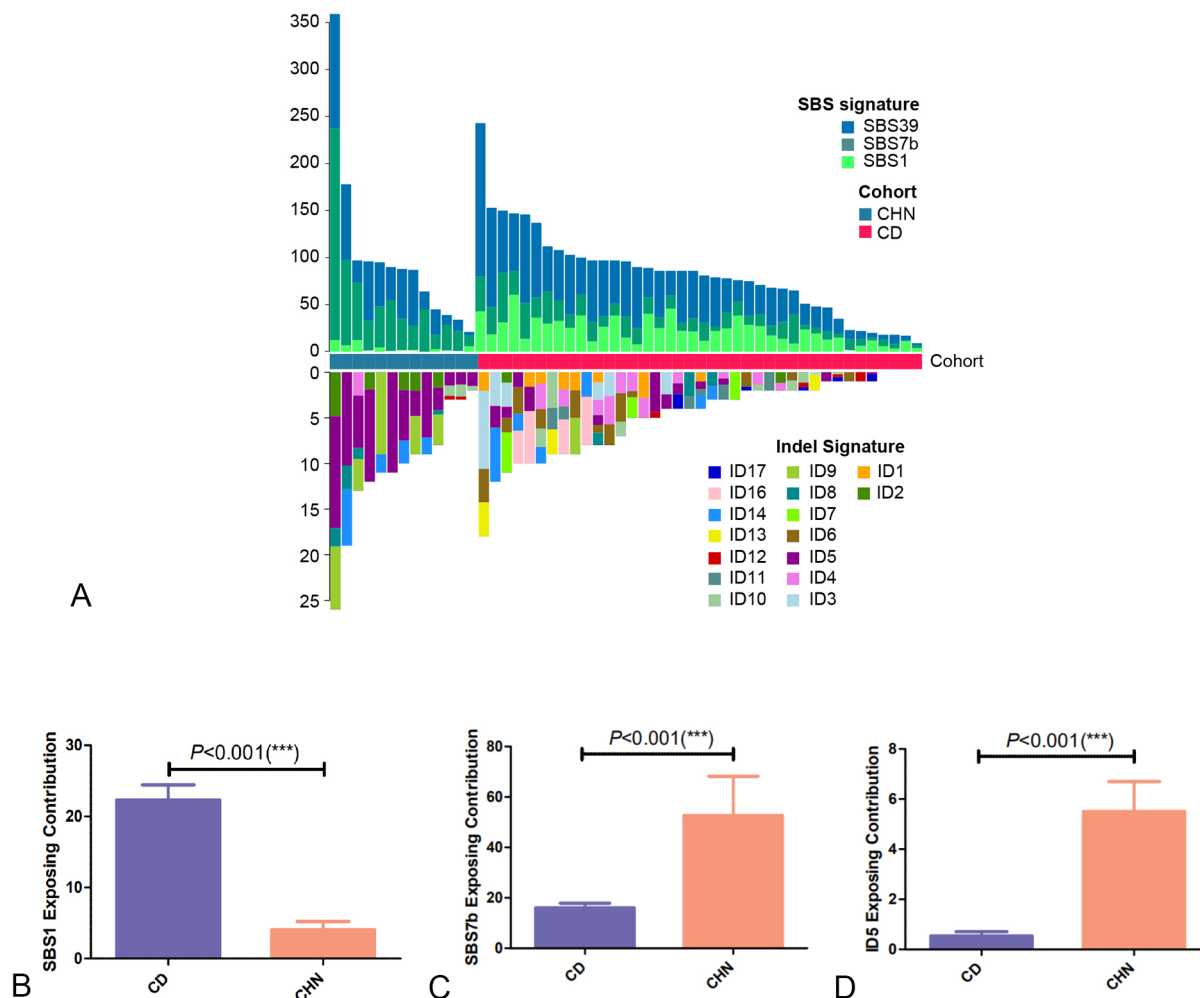


Figure 2. Exposing Contribution of COSMIC signatures in 13 Chinese OTSCC patients; A. panel at the top, exposing contribution of SBS signatures; panel in the middle, source of patients, CHN- Chinese patients in our cohort (N = 13), CD-Patients published in Cancer Discovery (N = 40); panel at the bottom, exposing contribution of Indel signatures; B, C, D. exposing contribution of SBS1, SBS7b, ID5 in CHN vs. CD cohorts accordingly.

exposure to ultraviolet light [14]. SBS1, a clock-like signature with predominant C > T changes, is an endogenous mutational process initiated by spontaneous or enzymatic deamination of 5-methylcytosine to thymine which generates G: T mismatches in double-strand DNA, and was consistent with previous reports that SBS1 was one of the main signatures of OTSCC [15]. In our cohort, ID5 was observed particularly predominant, characterized by predominant single T base deletion, and was reported to be a clock-like signature associated with patients' ages [14].

We further quantified and compared the contribution of identified SBSs and IDs in CHN and CD cohorts to look at the difference in possible mutagenic exposure and DNA damage repair deficiency between Chinese and Caucasian populations. Interestingly, the exposing contribution of SBS1 was found significantly lower in our cohort compared to CD cohort ($P < 0.001$; **Figure 2B**), whereas

SBS7b and ID5 were both observed with substantially increased contribution in our cohort ($P < 0.001$ for both) (**Figure 2C & Figure 2D**).

Clonal Architecture of Somatic Mutations in OTSCC

To look at the intra-tumor heterogeneity and infer the temporal order of the occurrence of somatic SNVs and Indels in Chinese OTSCC, we gauged the variant allele frequency (VAF) ratio of all alterations in our cohort with the method mentioned in previous literature [9]. Somatic mutations with higher VAF ratios were suggested to occur at earlier stages during tumor progression [9]. Mutations with VAF ratios greater than 0.5 were exhibited in (**Figure 3**), where black dots represent median VAF ratios when multiple mutations in the same genes were identified. Being

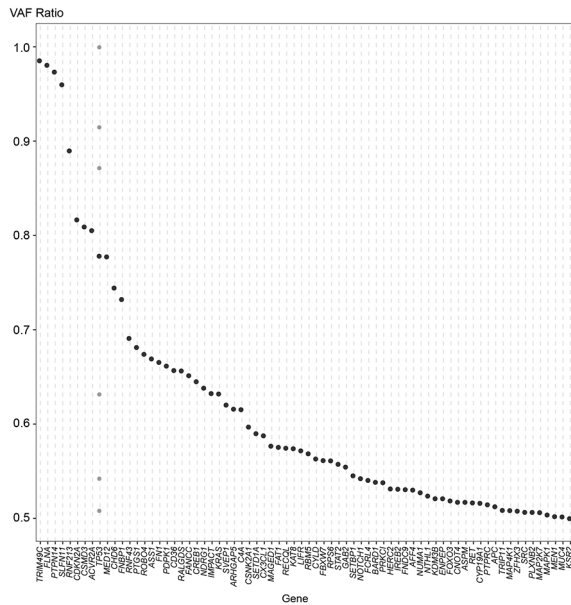


Figure 3. Mutations in genes with VAF ratios equals to or larger than 0.5 in 13 OTSCC patients.

consistent with the findings stated above, mutations in canonical cancer driver genes such as TP53 (0.78), CDKN2A (0.82) and CSMD3 (0.81) were determined as clonal mutations with VAF ratios greater than 0.75, indicating they might occur at an earlier stage compared to other subclonal mutations, and might play pivotal roles in OTSCC tumorigenesis. By contrast, some other

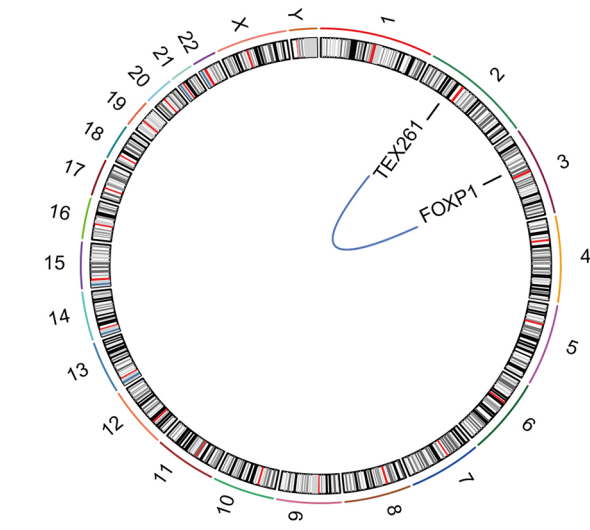


Figure 5. TEX261-FOXP1 fusion identified in 13 OTSCC patients.

canonical cancer drivers like KRAS, FAT1, NOTCH1 and APC were observed with relatively lower VAF ratios, suggesting they might occur at a later stage in tumor progression, exerting functions in tumor development and subclonal evolution. Subclonal mutations with various VAF ratios largely contributed to OTSCC intra-tumor heterogeneity and might be able to prime tumors towards drug resistant, disease relapse and metastasis.

Somatic Copy Number Alteration in OTSCC

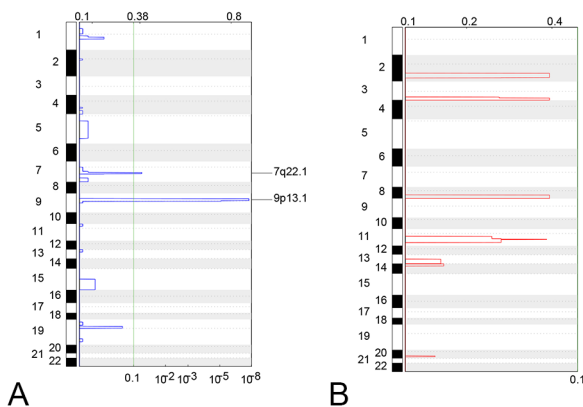


Figure 4. Significant loss (A) or Gain (B) of copy number segments in 13 OTSCC patients. On each figure, the panel in the left represents the chromosome of each CNV segment located on; the numbers on the top represents the G-scores assigned by GISTIC for every cytoband plotted; the numbers on the bottom represents the $-\log_{10}$ transformed q values of each CNV.

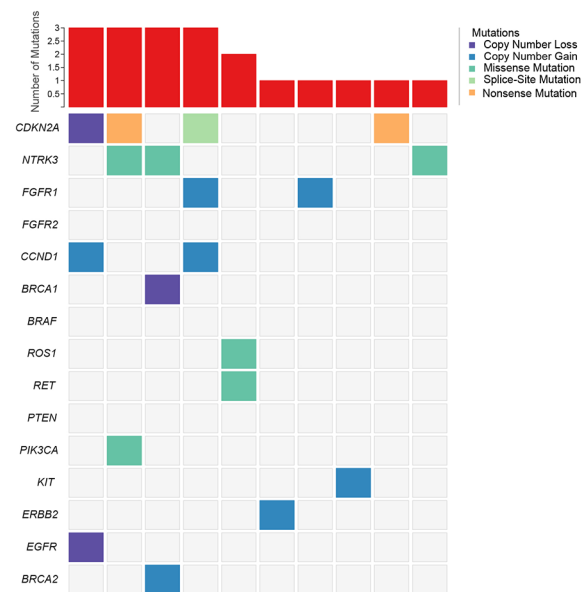


Figure 6. Mutations in Actionable Mutations by Searching OncoKB actionable genes in raw mutations of 13 Chinese OTSCC patients.

To identify the recurrent focal somatic copy number alterations (SCNA), the GISTIC algorithm and the 2Mb windows setting were applied to our cohort. Among population, two significant regions of deletion, including 7q22.1 and 9q13.1, were detected (**Figure 4A**), while no significant amplified large segment was identified (**Figure 4B**).

We also manually checked the smaller focal regions of amplification and deletion of individual patients in order to characterize the driver genes significantly implicated in copy number changes. As a result, canonical oncogenes such as MYC, CCND1, FGFR1, ERBB2, PDGFRA, and KIT were gained or amplified whereas tumor suppressor genes such as CDKN2A, CDKN2B, and BRCA1 were lost or deleted (**Table S3**).

Chromosomal Variation in OTSCC

A total of 28 structural changes were identified in our cohort (10/13, 76.9%). Of particular interest, among 10 OTSCC patients harboring chromosomal changes, one case was paid special attention to and discussed in detail due to the detection of cancer associated FOXP1-TEX261 (2p13.3:3p13) fusion (**Figure 5**), which might exert potential functions in enticing tumorigenesis and were previously reported in other cancer types encompassing aldosterone-producing adenoma [16], prostate cancer acute lymphoplasmic leukemia and etc [17-19].

Actionable Mutations of OTSCC in Chinese Population

In order to develop novel therapeutic interventions for OTSCC, we looked at the actionable mutations in our cohort by mapping unfiltered raw mutations into OncoKB actionable mutation catalogue (**Figure 6**), so that all potential candidate targets could be kept. Multiple actionable genes were observed with recurrent mutations, including CDKN2A, NTRK3, FGFR1, FGFR2, CCND1, BRCA1 and etc, functioning as potential candidates for developing novel targeted therapies. Of particular interest, CDKN2A and NTRK3 were observed mutated with the highest prevalence in Chinese cohort whereas no mutation in these genes was found in Caucasian cohort, suggesting they might play essential roles in OTSCC tumorigenesis and could be considered as two promising therapeutic targets, specifically in Asian population.

Discussion

While incidence of head and neck cancer (HNC) has been gradually decreased in the last decades, several subtypes of HNC showed an increasing incidence, particularly for those occurred in oral cavity [20]. In this study, we illustrated the genomic landscape of OTSCC to dissect its genetic underpinnings. Albeit with a modest sample size, we integrated mutation data of 40 Caucasian OTSCC from a published dataset and validated the key findings of previous report as well as unearthed several novel drivers that might essentially contribute to OTSCC tumor progression. Lastly, we have shown that CDKN2A and NTRK3 together with some other actionable genes might be novel potential therapeutic targets, rendering it promising to develop new targeted therapies to prolong patients' survival.

Overall, compared to Western patients, the mutation frequencies of some known cancer associated genes in Chinese patients were quite close [5]. To this end, some key mutations in oncogenes and tumor suppressor genes such as TP53, PIK3CA, CASP8, NOTCH1, MUC4, FAT1 and etc were identified in both our cohort and CD cohort, proving their roles in OTSCC tumorigenesis as previously reported [5]. Also, our results suggested that the

pathways these genes enriched in, NOTCH signaling pathway and CASP8 cellular necrosis pathway for instance, might be engaged in OTSCC cancer progression.

Notably, mutations in CDKN2A and NTRK3, both occurred in 23.1% patients in our cohort, were instead undetected in CD cohort, and both were categorized in OncoKB database as actionable genes, potentially offering novel targets for therapeutic development. Previously, CDKN2A and NTRK3 were reported to play key roles in various cancer types in the Chinese population including non-small cell lung cancer [21]. CDKN2A is a canonical tumor suppressor gene encoding p16INK4a protein, and was demonstrated to be a key driver triggering tumorigenesis in various cancer types [8, 22, 23]. The encoded p16INK4a protein exerts functions to orchestrate multiple signaling pathways in modulating cell cycle and senescence, through regulating the cyclin-dependent kinase (CDK) 4/6 and cyclin D complexes [8, 22, 23]. Moreover, genetic and epigenetic alterations in CDKN2A were exhibited to be indicative of disease relapse and dismal prognosis in cancer patients [8, 22, 23]. In our cohort, 2 truncating nonsense mutations and 1 mutation in canonical splice site were identified in 13 patients, and both 2 nonsense mutations located in the functional ankyrin repeats domain, which was shown to be involved in protein-protein interaction of p16INK4a to constrain cells from passing through cell cycle, and was suggested to play key roles in p16INK4a function [24, 25]. Our result indicated that mutations in CDKN2A might lead to the loss of p16INK4a regulatory capacity in modulating cell cycle arrest, and further result in OTSCC tumorigenesis. In addition, clonal CDKN2A mutation with high VAF ratio detected in our cohort suggested that CDKN2A might be altered at an early stage compared to other genes with subclonal mutations. Significant copy number loss of CDKN2A together with CDKN2B were also found recurrent in our cohort, which potentially resulted in a decreased expression level of p16 proteins, reducing its ability to maintain normal cell cycle arrest. Similarly, a recent study showed that CDKN2A copy number loss was observed highly prevalent (146/401, 36.4%) in HPV-negative head and neck squamous cell carcinoma, stringently associated with patients' dismal prognosis [26]. A germline polymorphism CDKN2B-AS1 (rs2151280 T>C) was also shown by another study to be associated with a higher risk of lung cancer [27]. Besides, differential expression in tumor tissues was found in various cancer types including hepatoblastoma, colorectal cancer and pituitary adenomas etc [28-31]. The down-regulation of CDKN2A and CDKN2B were found to associate with the recurrence of disease in oral cancer patients [32]. In 2003, Yakushiji et al showed that by comparing 25 pairs of primary oral squamous cell carcinoma (OSCC) and matched normal oral mucosa tissues, hypermethylation and down-regulation in both mRNA and protein expression levels were found in 48% samples, demonstrating a mechanism of CDKN2A undergoing hypermethylation, down-regulation to eventually trigger OSCC tumorigenesis [33]. In-vitro knock-in and knock-down assays of CDKN2A exhibited its basic functions in triggering G1 phase arrest during cell cycle in cancer cell-lines, where CDKN2A down-regulation was shown to be associated with enhanced cell proliferation [34, 35]. Besides, a CRISPR-Cas9 engineered CDKN2A knock-out model demonstrated depletion of CDKN2A facilitated melanocyte in-vitro motility and in-vivo invasiveness, indicating the function of CDKN2A to inhibit E2F1-mediated transcriptional activation of BRN2, which in turn to regulate down-stream signaling cascades to trigger melanoma metastasis [36]. Taken together, these results further solidified the conjecture that CDKN2A might be a novel cancer driver gene in OTSCC. The discrepancy of CDKN2A mutation in our cohort compared to CD cohort might be due to modest sample sizes of these two studies, while another possible explanation is somatic CDKN2A mutations might be specific to

Asian OTSCC population whereas they occur less frequently in Caucasian population, awaiting more substantiation.

NTRK3 is another cancer suppressor gene regulating cell survival and apoptosis, encoding neurotrophin-3 receptor, which functions as a tyrosine kinase receptor to orchestrate multiple signal transduction cascades involved in cellular proliferation and was demonstrated to play important roles in various cancer types [37, 38]. Similarly, NTRK3 was found differentially expressed in various cancer types. Methylation and down-regulation were observed in tumors where reconstitution induced cell apoptosis, and a decreased expression of NTRK3 was reported to be associated with neoplastic transformation, cell migration and invasion in-vitro and in-vivo [39-42]. Aberrantly methylated NTRK3 was shown to be a biomarker of colorectal cancer [43]. Down-regulation of NTRK3 was suggested to increase cancer cell invasion, migration and proliferation [42, 44]. Moreover, ETV6-NTRK3 fusion was reported to be driver alteration triggering tumorigenesis of multiple cancer types, including secretory breast cancer, colorectal cancer and Spitz tumor [37, 45-48]. Recently, NTRK somatic mutations and ETV6-NTRK3 fusion were also reported in head and neck squamous cell cancers with a prevalence of 7.9% and 2.4% in Chinese patients (3/127; 10/127) respectively, and one of the ETV6-NTRK3 positive patient benefited from crizotinib treatment, offering an alternative treatment strategy [49]. Similar to CDKN2A, NTRK3 was observed recurrently mutated in our cohort, while no mutation was detected in CD cohort. 2 missense mutations of NTRK3 were found located in the immunoglobulin I-set domain and protein tyrosine kinase domain, leading to NTRK3 loss of function. Therefore, NTRK3 might also be another novel driver facilitating tumorigenesis and tumor development in OTSCC, and possibly specific to Asian population. However, ETV6-NTRK3 fusion was not detected in our cohort, possibly due to a limited sample size and/or a different sequencing strategy utilized in our study.

From our result, 76.9% OTSCC harbored mutations sensitive to off-the-shelf targeted therapeutic drugs, rendering targeted therapy a highly promising means of intervention to this malignancy. We manually checked the FDA approved medications in OncoKB database (Memorial Sloan Kettering Cancer Center) for these candidate genes. Among which, gefitinib, afatinib, osimertinib and dacomitinib were approved for EGFR+ non-small cell lung cancer, rucaparib was approved for BRCA1+ ovarian cancer, sunitinib, imatinib and regorafenib were approved for KIT+ gastrointestinal stromal tumor, and fulvestrant with alpelisib was approved for PIK3CA+ breast cancer etc. It's worth noting that even though these drugs were approved in numerous cancer types, limited numbers of drugs were developed or approved to be applied to OTSCC to date, therefore developing new therapeutic strategies for this cancer type might be of great significance to improve patients' treatment opportunities.

The mutational signatures of oral cavity cancer appear to vary depending on the exposure of environmental carcinogens [50], as habitual consumption of tobacco, alcohol, and betel quid (BQ) were recognized as the major etiologic factors for this type of malignancies. Previous literature has depicted the distinctive mutational features in tumors associated with cigarette smoking [51], alcohol drinking [52], and BQ chewing [53, 54]. In present study, we conducted a quantitative measurement of mutational patterns for our cohort and CD cohort. In result, though consistent signatures were identified between our cohort with previous report, significant difference was observed in contribution of mutational signatures between two cohorts. The discrepancy in signature distribution is considered reasonable given that the living environment and lifestyle largely vary across the populations.

Copy number alteration analysis revealed the deleterious somatic SCNA in hallmark genes of tongue cancer, such as amplification

of MYC, CCND1 and FGFR1, and were also identified in previous reports [16, 55, 56]. In addition, high expression of ERBB2 was reported to exhibit important roles in the development of tongue cancer in patients with smoking history and/or alcohol consumption [57], and PDGFRA expression was significantly higher in oral cancer cohort with or without the establishment of tobacco risk factor [58]. Also as stated above, the down-regulation of CDKN2A and CDKN2B were found to associate with the recurrence of disease in oral cancer patients [32]. In conclusion, our SCNA findings were consistent with previous literature and have vital implications in clinically relevant applications.

Despite no overall structural variation pattern scrutinized, multiple chromosomal changes occurred in individuals might potentially contribute to cancer progression. In one patient, a FOXP1-TEX261 (2p13.3:3p13) fusion was observed (**Figure 6**). FOXP1 (Forkhead Box P1) encodes transcription factor Foxp1 expressed across the whole system, and was previously reported to exert functions in regulating immune response, cancer progression, organ development, and showed oncogene properties in acute lymphoblastic leukemia [19, 59-62]. As chromosomal changes involving FOXP1 was frequently found in various malignancies, FOXP1 served as a highly potential gene target for developing therapeutic interventions [62].

Conclusion

We reported the mutational analyses of 13 Chinese OTSCC patients. Apart from recently reported well-characterized alterations like TP53 mutation and CASP8 mutation in OTSCC, we also demonstrated several undescribed genetic changes such as CDKN2A, NTRK3 and FOXP1-TEX261 fusion. Also, several genes were identified as novel therapeutic targets, laying a solid foundation for developing new effective interventions.

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Author contributions

NL designed and supervised the study, RXH collect and preserve the patient samples. SHW, JN, ZY and YF analyzed the data and composed the article. SX collected clinical data and samples. YY, QF, CZ, ZXT, JL verified the data and methods. All the authors have read the article and were involved in the review and revision.

Competing interests

The authors declare that they have no conflicts of interest in this study.

Ethical statement

The study was approved by the Ethic Committee of Cancer Hospital Chinese Academy of Medical Sciences (ID: NCC-002760).

Additional files

[Additional file 1:](#) Table S1. Clinicopathological Characteristics of Patients Enrolled and in External Datasets.

[Additional file 2:](#) Table S2. Raw Mutations Identified from 13

Chinese OTSCC patients.

[Additional file 3](#): Table S3. Significantly Implicated Driver Genes in Focal CNV Segments.

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