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Eminence Grise of the Genome: Long Non-Coding Ribonucleic Acids in Oral Squamous Cell Carcinomas

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Received: 12 December 2019 Accepted: 4 January 2020 Published: 15 January 2020

8 Abstract

Non-coding ribonucleic acids (ncRNAs) are a class of RNA molecules that are transcribed but
 not translated into proteins, but they affect various cellular processes. Around 60

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Index terms— oral squamous cell carcinoma, potentially malignant disorders, non-coding RNAs, long noncoding RNAs, competing endogenous RNA.

14 1 Introduction

ral Squamous cell carcinoma (OSCC) is a heterogenous malignancy which results in decreased survival rates due 15 16 to local recurrence and lymph node metastases [1]. Various other cancers like lymphomas and certain sarcomas possess relatable gene alterations for which effective target drug therapies are developed, but due to complex 17 genomic and epigenomic changes and interactions in OSCC, use of an effective chemotherapeutic agent is still a 18 challenge. Recent research has now focused on epigenomic modifications that effect the gene expression rather 19 than gene mutations to understand these complex mechanisms [2,3,4]. In context to this, non-coding ribonucleic 20 acids (ncRNAs, previously considered as "junk or transcriptional noises") that are transcripts not translated to 21 proteins but are potential effectors of target gene expression have gained additional interest [5,6]. 22 Statistics of Ensemble 1 show that around 34% of the human genome are protein-coding genes. Among which 23

66% of genes encode ribonucleic acids (RNAs) that are not translated into proteins [5,6]. The Encyclopedia of DNA Elements Consortium (ENCODE) revealed that humans have 60554 genes out of which 19815 are proteincoding genes, and the rest represents ncRNAs that regulate gene expression involved in vital physiological and pathological processes [4,7].

They are grouped as house-keeping ncRNAs and regulatory ncRNAs. The house-keeping RNAs include ribosomal [rRNAs], transfer RNAs [tRNAs], small nuclear RNAs [snRNAs] and small nucleolar RNAs **??**snoRNAs]. The regulatory ncRNAs are divided into: a) Short ncRNAs: size < 200 base pairs (bp); b) Long nc 25 RNAs (lncRNA): size > 200 bp; c) Pseudogenes; d) Circular RNAs; e) Intronic RNAs [7,5,6].

The small ncRNAs include small interfering RNAs (siRNAs), PIWI-interacting RNAs (piRNAs), micro RNAs (mi RNAs) and transcription initiator RNAs (tiRNAs) [5,3].

Dr. Reshma Venugopal ?, Dr. Radhika Manoj Bavle ?, Dr. Sudhakara Muniswamappa ? & Dr. Soumya Makarla ? There are about 16,000 lncRNA genes that encode 28,000 lncRNAs. Five types of lncRNA are identified based on the position of DNA protein-coding strands [Figure 1a] from which they are synthesized. Figure ??d: Intergenic lncRNA synthesized from within 1 kb of protein coding on same strand e) Intronic lncRNAs: overlapping either the sense or antisense intronic areas of the protein-coding genes (Figure ??e) [7,8,9]. Figure ??e: Intornic lncRNA synthesized from intronic region of coding gene either same or opposite side

LncRNAs regulate gene expression through epigenetic regulation (chromatin modification & DNA methylation), transcription, and post transcription processing by acting as scaffolds, guides, decoys or repressors, sponges serving as competing endogenous RNAs (ceRNAs) for signaling pathways, and enhancer RNAs. They are involved

43 in pre-mRNA splicing [Table 1] [1,4,6,9,7,10,11,12,13].

44 2 LncRNAs scaffolds

45 Platforms on which multiple enzymatic proteins can be transiently assembled in functional units such as 46 ribonucleoprotein complex (RNP), heterogenous nuclear ribonucleoproteins (hnRNAs) etc. Their interaction 47 is dynamic and exerts regulatory functions during mRNA processing.

48 **3 2.**

49 LncRNAs guides Physically direct the RNAs to specific genomic region by binding to regulatory or enzymatically
 50 active proteins such as transcription factors, chromatin modifiers etc and regulate gene expression either in cis

51 or trans sites.

⁵² **4 3**.

LINCRNA decoys Limit the availability of specific regulatory factors by acting as a molecular sink and sequester RNA-binding proteins, transcription factors, microRNAs, catalytic proteins and subunits of larger modifying complexes. They titrate these factors away from interacting with their native targets, decreasing their bioavailability, inhibiting their normal functions 4.

57 LncRNA sponges Impair miRNAs, explained by competing endogenous RNA (ceRNA) hypothesis.

According to which lncRNAs sequester miRNAs and reduce the availability of miRNA for the target mRNAs.

⁵⁹ The miRNAs play an important role in post transcriptional regulation of protein coding genes and mRNAs. ⁶⁰ LncRNAs actively compete with specific protein-coding mRNA that interact with intracellular pool of miRNAs

LncRNAs actively compete with specific protein-coding mRNA that interact with intracellular pool of miRNAs
 acting as sponges or ceRNA for miRNAs, silence them and reduce the post-transcriptional activity.

62 **5 5**.

63 Signalling lncRNAs Associated with specific signalling pathways or events such as cellular stress leading to 64 transcription activation of specific genes.

65 **6 6**.

66 Enchancer IncRNAs Enhance and promote gene activity by altering the 3 dimensional configuration of DNA.

67 LncRNAs act in nucleus or cytoplasm or both, exhibiting three types of interactions: RNA-RNA, RNA-DNA,

and 52 RNA-proteins. Their partners of communication include RNA binding proteins (RBPs), transcription

69 factors, chromatin-modifying complexes, nascent RNA transcripts, mature mRNA, microRNA, DNA, and

⁷⁰ chromatin. [1,7,9,14] In the cytoplasm, they interact with target mRNAs or miRNAs through miRNA response ⁷¹ elements via base paring. They may stabilize or decoy the target transcripts, thus promoting or repressing

⁷² the translation of transcripts to proteins [7]. Cytoplasmic lncRNAs act as sponges and promote micro peptide

73 formation [10,16].

⁷⁴ 7 a) Synthesis of LncRNA b) Functions of LncRNA

LncRNAs affect numerous biological processes such as embryological development, stem-cell biology, development, 75 and differentiation [6]. They are tissue or cell type-specific as indicated by gene expression profiling, possessing 76 a varied expression to different pathophysiological conditions, and tumors. They regulate cell proliferation, 77 survival, apoptosis, invasion, metastases, glycolysis, angiogenesis, growth, tumor-In the nucleus where they are 78 79 mainly localized, they regulate epigenetics of protein-coding genes and alter their expression through chromatin 80 remodeling complexes [such as polycomb repressive complex 2 (PRC 2), H3K9 methyl transferases] and DNA methylation patterns. They control the gene expression at the transcription level, act as decoys, bind to the 81 DNA target sequences, act as alternative splicing regulators (antisense transcripts), are involved in splicing 82 malfunctioning, and act as decoys for splicing [7,15]. The cis-acting lncRNAs are close to transcription site, and 83 trans-acting lncRNAs are on distant genes of chromosomes [1]. 84

LnRNAs originate and are predominantly located in the nucleus. Tissue specific RNA polymerase I, II or 85 III transcribe lncRNAs. They are 5' capped, 3' polyadenylated, have exon/intron length, and undergo splicing 86 of multiple exons through canonical genomic splice motifs. They resemble protein-coding mRNAs, but lack or 87 have a small number of open reading frames. The exon length of lncRNA is the same as proteincoding mRNAs 88 but has fewer exons that are less expressed than protein-coding mRNAs. As the span of lncRNA is more than 89 90 200 bp, they can fold into more complex three dimensional structures unlike, miRNA. The complex structure of 91 lncRNAs determines their specific interaction with transcription factors, histone and chromatin-modifying genes 92 affecting the expression level of a broad spectrum of genes [9,8,10,17]. stroma signaling or genomic stability, thus serving as potential diagnostic, prognostic biomarkers, and therapeutic targets [5,8,10,13,18,19,20]. 93

LncRNAs are dysregulated in several neurological disorders and cancers, demonstrating both oncogenic and tumor-suppressive roles [12,21]. LncRNA Hox antisense intergenic RNA (HOTAIR), for example, functions as an oncogene in breast cancer, colorectal cancer, pancreatic cancer, etc, and increased levels are associated with reduced survival rates [20]. On the other hand, maternally expressed gene 3 (MEG3) is up-regulated in breast, hepatic cancer, and plays an oncogenic role. In contrast, it is downregulated in tongue squamous cell carcinoma (TSCC) and plays a tumor-suppressor role [4]. The present article reviews the expression of lncRNAs in OSCC
 mainly, with a note on its presentation in oral potentially malignant disorders (OPMDs).

Gibb et al. state that in the normal oral mucosa 325 lncRNAs are expressed. In OPMDs, around 164 lncRNAs are aberrantly expressed. Jia et al. studied 3590 differently expressed lncRNAs in TSCC, and found that 1785 were up-regulated, and 1805 were downregulated [3,4,22]. Yu et al. detected 1572 abnormally expressed lncRNAs with 882 up-regulated, and 690 down-regulated lncRNAs [4]. A study on head and neck squamous cell carcinoma (HNSCC) showed that 84 out of 3199 lncRNAs had an impact on survival rates of the patients [1].

106 Studies done by Gao et al. showed six upregulated lncRNAs such as lnc 122-PPP2R4-5, SPRR2D-1, FAM46A-

107 1, BL2-4:1, and MBL2-4:3 (associated with high nodal status) in TSCC and two down-regulated lncRNA viz 108 AL355149.1-1 and STXBP5-1 [17].

¹⁰⁹ 8 II. LNCRNAs Upregulated during OSCC Progression

a) LncRNA H19 First identified long non-coding RNA, coded by gene H19 located in chromosome 11p15.5 in

close association with insulin growth factor (IGF) 2 gene. LncRNA H19 is a transcription factor of the H19/IGF 2 genome blotting cluster, directly activated by c-MYC and down-regulated by p53 contributing to cell growth

113 and proliferation [23].

The combination of H19 and enhancer of zest homolog 2 (EZH2) affects signal transduction of bcatenin/glycogen synthase kinase three beta (GSK 3b)/epithelial-mesenchymal transition (EMT) in TSCC, promoting lymph node metastasis and poor prognosis. MiRNA-138 and 630 down-regulate EZH2 and are suppressed by lncRNA H19. Thus H19 and Hox antisense intergenic RNA (HOTAIR) can up-regulate EZH2 decreasing the E-cadherin levels, enhancing the invasive potential of SCC cells with H19 expression being higher in metastatic tumors than nonmetastatic [23].

H19 acts as a ceRNA increasing the level of miRNA lethal(let)-7a targets, a chief regulator of highmobility group AT-hook 2 (HMGA 2) in the process of tumor metastasis. H19 / let-7a/HMGA2 / EMT axis plays a

principal role in the regulation of invasion, metastasis, and is associated with poor prognosis in TSCC [4]. H19

¹²³ over-expression in endothelial cells stimulated angiogenesis. H19 regulates expression of tumor growth factor ¹²⁴ (TGF)-b, which promotes cancer cell migration through enhanced adhesion with extracellular molecules. Notch

and hepatocyte growth factor regulated signaling of H19.

Blocking of Notch and HGF inhibited H19. The inhibition of H19 decreased cell resistance to Fulvestrant and Tamoxifen [10,22,24]. Thus H19 plays a role in reducing the susceptibility of cells to chemotherapeutic drugs.

¹²⁸ 9 b) Hox antisense intergenic RNA-HOTAIR

HOTAIR is highly conserved nuclear lncRNA, a transcript of 2.2 kb, transcribed from the (Homeobox C) HOX
C locus at chromosome 12 but functions at transit close to HOX D locus on chromosome 2 that induces silencing
of transcription [3].

The domain 5' of HOTAIR binds PCR 2, which includes EZH2, SUZ 12, and EED (both polycomb proteins) to the HOX D locus and inhibits its expression. EZH2 is a histone H3 lysine 27 methyl transferase (H3K27me 3) enzyme that catalyzes trimethylation of H3K27, a histone modification associated with long-term transcription repression [23]. EZH2 is a critical epigenetic regulator for various biological processes such as cell proliferation, cell cycle, metastases, and oncogenesis [6,16].

HOTAIR prefers to occupy a guanine-adenine (GA)-rich DNA motif on chromatin, which allows direct
interaction of lncRNA transcript to specific genomic sites (has both cis and trans-regulatory mechanism) [20,23].
Also, the 3' domain of HOTAIR binds the histone demethylase lysine-specific histone demethylase (LSD)1. This
evidence suggests that HOTAIR serves as a platform for two different histone modification complexes [3].

It promotes tumor cell invasion and metastasis by recruiting EZH2 and repressing E-cadherin in OSCC [11]. HOTAIR is differently expressed in the saliva of patients with OSCC with metastases and without metastases [3]. According to Dai et al., HOTAIR 7 is highly expressed in TSCC associated with cell proliferation, apoptosis, metastases, and invasion [4].

High expression of HOTAIR in laryngeal SCC is associated with tumor size greater than 0.9 cm, poor
differentiation, lymph node metastasis, resistance to apoptosis, advanced clinical stages, and also drug resistance
with EZH2 serving as a potential mediator [3,8,20].

HOTAIR is highly expressed in hypoxia and regulates angiogenesis through vascular endothelial growth factor
 A (VEGF A) directly through its promoter sequence or by modifying the levels of glucoseregulated protein-78
 (GRP 78) and angiopoietin 2 (ANG-2) in nasopharyngeal carcinoma [10].

¹⁵¹ 10 c) Ferritin Heavy Chain 1 Pseudogene 3: FTH1P3

FTHIP3 is mapped to chromosome 2p23.3 with a length of 954 nucleotides, it is ferritin pseudogene with a misannotated 3' un-translated region (UTR), which is closely associated with iron-responsive elements (IREs). It affects the post-transcriptional structured cis-acting RNA regulatory elements in the 5' or 3' UTRs of mRNAs. FTH1P3 harbors miR-224-5p cognate site, sponging miRNA 224-5p and consequently modulates the expression of frizzled class receptor five, which acts as an oncogene in OSCC. miRNA 224-5p is a potential tumor suppressor that is suppressed by FTHIP3 in OSCC, contributing to oncogenesis. FTH1P3 is coexpressed with plasminogen activator urokinase (PLAU) and targets OSCC associated genes, including matrix metallopeptidase (MMP) 1, 3, 9, PLAU and interleukin 8 (IL 8) which are essential regulators of tumorigenesis. Ectopic and overexpression of FTH1P3 facilitates cell proliferation, colony formation, tumor progression, metastases, and worsens survival rate in OSCC cases [6,11,22,23,24].

¹⁶² 11 d) Urothelial Cancer-associated 1-UCA1

Located on chromosome 19p13.12, UCA1 regulates the expression of various genes mainly through wingless-163 homeobox gene (WNT)/ b-catenin signaling pathway. The lncRNA is upregulated in OSCC, promoting cellular 164 proliferation and tumor-lymph nodemetastasis (TNM) staging. UCA1 functions as a sponge to miR-184 inhibiting 165 it, miRNA-184, in turn, has an inhibitory effect on phosphatidylinositol 3-kinase (PI3K)/protein kinase B 166 (AKT)/mammalian target of Rapamycin (mTOR) signaling pathway. As a ceRNA, UCA 1 represses the 167 effect of miRNA-184 promoting tumor progression, suppressing the influence of cisplatin-induced apoptosis and 168 chemosensitivity in TSCC cells. UCA1 acts by inhibiting the cisplatinactivated PI3K/Akt signaling pathway, 169 though, in TSCC, UCA1 promotes lymph node metastases more than cell proliferation. 170

In OSCC, it is co-expressed with numerous metabolism-related genes. UCA1 enhances the Warburg effect via mTOR activation followed by activation of signal transducer and activator of transcription 3 (STAT 3) and repression of miR-143. The Warburg effect results in increased hexokinase 2 (HK2) levels and a consequent increase in glycolysis.

Tonghan Zhang et al. discovered the regulation effects of UCA1/miR-124/Jagged 1 (JAG1) axis on tongue cancer which activates Notch pathway [4,5,8,12,13,14,21,25,23].

177 12 e) Metastases associated lung adenocarcinoma transcript 1-178 MALAT1

MALAT1 or nuclear enriched abundant transcript 2 (NEAT2), is a long intergenic non-coding RNA that maps on chromosome 11q13 [23] and is 8.7 kb 203 long. It is very stable due to its triple-helical and nuclear retention element (ENE) like structure located in the nucleus and plays a role in RNA metabolism. Expression of cell cycle genes such as E2F1 transcription factors and the G1/S phase requires MALAT 1. It promotes mitosis through transcription factor B-MYB. MALAT-1 depleted cells are sensitive to p53 levels, indicating that p53 is one of the key molecules involved in the downstream effects of MALAT1.

MALAT 1 acts as a sponge to miRNA-125 b, which upregulates STAT 3 expression promoting OSCC development. It plays a role in EMT in OSCC cells, promoting cell migration and invasion. Its knockdown suppressed N-cadherin and vimentin but induced Ecadherin expression in vitro. In a tongue cancer cell line, MALAT1 targeted miRNA-124 and promoted cancer growth through modulation of JAG1. Studies have shown that upregulated MALAT 1 induced cervical lymph node metastasis in TSCC by increasing BCL2 associated X

190 (BAX) expression.

Upregulated MALAT 1 interacts with EZH2 inducing b-catenin expression activating Wnt/b-catenin signaling pathway, upregulating MMP-7, inducing EMT, enhancing the invasion, and inhibiting apoptosis capacity of TSCC cells. Down-regulation of apoptosisrelated genes BNIP3L (pro-apoptotic BCL-2 family protein) and Neuregulin 1 (NRG1) is noted. Increased levels of MALAT 1 also incite mitogen activated protein kinase (MAPK) and PI3K/Akt. MALAT 1 functions as ceRNA for miRNA-320a, which suppresses forkhead box M1 (FOXMI). FOXM1 is involved in new vessel growth in hypoxic conditions associated with fibroblast growth factor-2 (FGF-2) expression [4,6,14,19,21,25].

Emerges from chromosome 15q15.1 on the strand opposite to OIP5 gene and is involved mainly in the regulation of neurogenesis during development. OIP5-AS1 is an oncogene that serves as a ceRNA by sponging multiple miRNAs such as miR-340-5p, miR-217, miR-200b-3p, miR-223, miR-410, miR-378a, and miR-338-3p. miRNA 338-3p is a tumor suppressor that modulates the expression of neuropilin 1(NRP1). NRP1 is a co-receptor for VEGF and functions as an oncogene in multiple types of cancers. OIP5-AS1 functions as miR-338-3p 'sponge' to trigger NRP1 expression and thus the progression of OSCC. Overexpression of NRP1 promotes EMT by stimulating the nuclear factor-kappa B (NF-kb) pathway.

Knockdown of OIP5-AS1 decreased the levels of NRP1 and significantly inhibited OSCC cell proliferation, migration, invasion, retarded tumor growth, and colony formation [19].

207 **13 3.**

- Long intergenic nonprotein coding RNA 511-LINC00511 ceRNA for miRNA 765 increasing the expression of Laminin subunit gamma 2 (LAMC2), weakening the inhibitory effect of miRNA-765
- High expression: Higher grades of dysplasia in leukoplakia and progression to malignancy. Increases cell
 proliferation and invasion in TSCC. An early biomarker.

²¹² 14 g) Colon cancer-associated transcript-1-CCAT1-S or cancer-²¹³ associated region long non-coding RNA-5-CARLO-5

CCAT1-S is a 2628 nucleotide lncRNA located on chromosome 8q24.21. CCAT1 mediates cell proliferation by inhibiting expression of cyclin-dependent kinase (CDK) inhibitor 1A (CDK N1A) mRNA, the main regulator of G0-G1 phase by increasing the expression of p16, p21, and p27 and thus cell proliferation. Silencing of CCAT1 by siRNA leads to induction of phase 1 cell cycle arrest, increased levels of E-cadherin, and decreased levels of fibronectin and vimentin required for EMT. CCAT1 interacts with transcriptional enhancer c-MYC promoter region through chromatin looping and increases its expression. CCAT1 functions as a ceRNA for miRNA-155-5p, let-7b-5p, and miRNA-490-3p by sequestration and miRNA-218-5p by epigenetic regulation.

221 27% of oral tumor show overexpression of CCAT1 associated with increased expression of c-MYC; and down-222 regulation of miRNA-155-5p and let 7b-5p. Oral cancer patients with an increased level of CCAT1 are associated 223 with tobacco use, poor prognosis, and aggressive phenotype [6].

²²⁴ 15 h) THOR-CG8846 gene product from transcript

CG8846-RA IGF 2 mRNA binding protein 1 (IGF2BP1), an oncogene belongs to a conserved family of 225 RNA binding proteins present mainly in the cytoplasm. They bind to mRNAs that regulate protein synthesis of 226 k-RAS, MYC gene family, CD44, phosphatase and tensin homolog (PTEN) and IGF 2. It plays a pivotal role in 227 cell proliferation, polarization, metabolism, morphology, differentiation, and migration. IGF2BP1 regulates the 228 radio-/chemo-resistance of cancer cells by increasing the expression of multidrug resistance mutation 1 (MDR1). 229 THOR stabilizes the binding of IGF 2 mRNA with IGF2BP1. It regulates IGF2/mitogen activated 230 kinase/extracellular signal-regulated kinases (IGF2/MEK/ERK) signal pathway in TSCC cells by increasing 231 cell cycle-related proteins cyclin D 1 and E1. THOR functions as a negative prognostic marker by increasing cell 232 proliferation; attenuates cisplatin sensitivity in nasopharyngeal carcinoma; regulates osteosarcoma stemness and 233 mobility [26]. 234

²³⁵ 16 i) Long Intergenic Non-Protein Coding RNA, Regulator

of Reprogramming-IncRNA-ROR Located on chromosome 18q21.31, LncRNA-ROR is 2.6 kb long non-coding 236 RNA and consists of retrotransposons elements such as long interspersed nuclear elements (LINE), short 237 interspersed nuclear elements (SINE) and long terminal repeater (LTR) elements. The location of lncRNA-238 ROR is a binding site for pluripotency transcription factors such as Sox2, Oct4, and Nanog. LncRNA-ROR acts 239 as ceRNA for miRNA 145-5p at post-transcriptional level, modulating the expression of target genes c-MYC, 240 Kl, SOX2, and Oct 4 impacting the differentiation of human embryonic stem cells. It also sponges miRNA-205 241 increasing the half-life of Zinc Finger E-Box Binding Homeobox 2 (ZEB2), thus promoting EMT. LncRNA-ROR 242 is a suppressor of p53 during DNA damage by interacting with hnRNP I, thus directly inhibiting p53 mediated 243 cell cycle arrest and apoptosis [6]. 244

The other lncRNAs associated with OPMDs is presented in table 2 [4,27,28,29]; and OSCC progression in relation to angiogenesis in table 3 [4,6,10,18,21,23,25,29], cell proliferation in table 4 [4,15,19,23,25,27,30,31,32], metastasis in table 5 [1,4,6,8,16,18,22,25,31,33,35], and chemoresistance in table 6 [4,13,29]. Decreased expression of lncRNA PTENP1 is observed in progressive OPMDs [5]. d) Growth Arrest-Specific Transcript 5 antisense 1-GAS 5-AS1 GAS 5 induces apoptosis, is down-regulated in HNSCC, and is associated with poor prognosis. It predicts the response to radical chemotherapy and plays an essential role in the pathogenesis of oral submucous fibrosis (OSF) and its progression to malignancy [4,3,8,22].

²⁵² 17 e) Prostate Androgen Regulated Transcript 1-PART 1

Located on chromosome 5q12, PART 1 functions as a ceRNA for miRNA-301b, which regulates Nuclear Receptor Subfamily 3 Group C Member 2 (NR3C2). Down-regulated NR3C2 promotes cell proliferation, EMT, and metastases. PART1/mir-301b/NR3C2 axis may be associated with TSCC. Androgens regulate PART 1 which is a tumor suppressor. Studies have found less expression of androgen receptor (AR) mRNA in OSCC specimens compared to healthy tissues. ARs might be involved in lessening the progression of OSCC and PART 1 is regulated by androgens, PART 1 study may also be involved in the pathogenesis of OSCC [32,38].

f) Long Intergenic Non-Protein Coding RNA 472-LINC00472 LINC00472 acts as a sponge to miRNA-503
that regulates the expression of Gremlin 2, DAN Family BMP Antagonist (GREM2), which is an antagonist of
bone morphogenetic proteins (BMP). BMP activates the Notch signaling pathway and Wnt/b-catenin signaling.
Higher expression of LINC00472 is associated with a better prognosis [32].
IV.

²⁶⁴ 18 Applications

Identification of up-regulated or down-regulated lncRNAs in the progression of OSCC is essential as their expression can be altered using appropriate RNA interference machinery such as short hairpin RNAs, miRNAs, siRNAs, oligonucleotides that are complimentary to target lncRNAs, etc. Molecule inhibitors that act by preventing the interaction of lncRNAs with the protein partners, blocking the binding or changing the secondary

structure of the lncRNAs are tried. For instance, silencing of MALAT1 in TSCC, lung adenocarcinoma, cervical 269 cancer, etc. by short hairpin RNA reduced the migration and invasive abilities of cancer cells. Blocking of 270 MALAT1 increased the levels of miRNA 195, which decreased PDL-1 expression in B-cell lymphoma cases, 271 decreasing apoptosis of CD 8+ cells; proliferative and metastatic abilities of the cancerous cells [39]. Down-272 regulation of UCA1, MALAT1, HOTAIR, and FOXCUT by using siRNAs resulted in decreased cell proliferation 273 and increased apoptosis in OSCC [8]. lncRNAs are used as gene therapy drugs to deplete cancer stem cells or 274 reverse their phenotype, thus increasing their sensitivity to radiation and chemotherapy [2]. 275 ν. 276

277 19 Conclusion

LncRNAs function as regulators in the conversion of OPMDs to OSCC, affect angiogenesis, cell proliferation,
 metastases, and predict chemoresistance.

Through reverse-transcriptase polymerase chain reaction, they can be detected in plasma and saliva, and thus serve as biomarkers. Identification of PAC3 in saliva helps in early diagnosis of OSCC, salivary gland tumors, and metastatic disease [8]. Markers such as LINC00974 and NEAT 1 etc. predict the progression of OPMDs. Expression of lncRNAs such as HOTAIR, MALAT 1, etc. found in saliva can help predict metastatic status in OSCC cases. High levels of CILA 1, KCNQ10T1, etc. predict chemoresistance in OSCC cases. Studies have found that specific siRNAs are used to alter the lncRNAs, regulating their expression in OSCC cases that works for a better prognosis. These features facilitate a thought to research these lncRNAs for good treatment options.



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Name of Lnc RNA	Function	Expression in OPMDS/OSCC
Nuclear enriched	Oncogene regulates miR365/ Regulator of G	Over expression: transformation of OPMI
abundant transcript 1-	protein signalling 20 (RGS20) pathway and	OSCC. Elevating proliferation, invasion,
NEAT 1	up regulates cyclin dependent ki- nase (CDK)	nodal metastases and inhibiting apoptosis
	6 through miRNA-107	OSCC. Not found in saliva.
Long intergenic non-	Oncogene-Areca nut constituents have	Higher expiresions
protein coding RNA	shown to activate the TGF-/p-Smad2	fibrinogenesis in OSF and mediates
974-LINC00974	pathway mediated by LINC00974, leading to enhanced myofibroblastic activity	progression of OSF to OSCC.
	Name of Lnc RNA Nuclear enriched abundant transcript 1- NEAT 1 Long intergenic non- protein coding RNA 974-LINC00974	Name of Lnc RNAFunctionNuclearenrichedOncogene regulates miR365/ Regulator of Gabundant transcript 1-protein signalling 20 (RGS20) pathway andNEAT 1up regulates cyclin dependent kinase (CDK) 6 through miRNA-107Long intergenic non-Oncogene-Areca nut constituents haveprotein coding RNAshown to activate the TGF-/p- Smad2974-LINC00974pathway mediated by LINC00974, leading to enhanced myofibroblastic activity

Figure 4: Table 2 :

3

1. Hyaluronan synthase 2 antisense 1-Lnc HAS2- AS1	Marker for hypoxia		Increased HAS HIF-1a increased production of hy thase which intur increases EMT a mour metastases	S2-AS1induces valuronan syn- n and OSCC tu-
2. HIF-1a co- activating	Complexes with HIF-1a, recruit	ment of	Complex is induc stabilizing and	ed by hypoxia,
RNA-lncRNA HIFCAR	HIF1a and p300 that target pro	moters	0	
	in OSCC.			
3. Long inter- genic non-	Acts as a ceRNA for miRNA-29	7 which	Up regulation: A cell proliferation,	ssociated with
coding RNA 668-	inhibits	VE & F	promotiong invasion Knockdown supp	of OSCC. ressed
LINCO 0668	angiogenesis		tumor growth and reduced the expression of proliferation antigen ki-67	
4. FOX C1 up- stream	Influences the expression of mat	rix	Over-expressed in ciated with cell	1 OSCC, asso-
transcript- FOXCUT or	metalloproteins (MMP) 2, 7, 9 a	and	proliferation, colony formation	angiogenesis, and
long intergenic	VEGF A		invasion.	
protein coding RNA 1379- LINC01379				
5. Long non- coding RNA	Hypoxia-inducible lncRNA and	acts as	Enhanced levels with poor clinical	are associated
MIR31	HIF-1?	co-activator	increasingnes and poor oral cancer.	or prognosis in
	angiogenesis			
6. Long non coding RNA	Binds to hnRNP-K complex and	1	Up regulated in (DSCC
p 21-LncRNA- p21	suppresses the expression of p53	1		
-	regulated genes. Induced by hyp inducible factor-1a (HIF 1a) dir during hypoxia, increases levels GLUT-1 and lactate dehydrogen turn increasing glycolysis in can cells.	ooxia ectly of nase in cer		

Figure 5: Table 3 :

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Eminence Grise of the Genome: Long Non-Coding Ribonucleic Acids in Oral Squamous Cell Carcinomas activating HIF1-a resulting in angiogenesis

1. Colon Cancer	Regulates	WNT/b-catenin/	Increased expression: Cell
Associated Tran- script	GSK-3b		proliferation, higher grade of tumour cell and pathological

 $2\text{-}\mathrm{CCAT2}$

affect metastases. stage (stage II/III) of OSCC, but does not

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2. Long intergenic	Targets miR-328-5p and miR-	Up regulation enhances cell
non-	939-5p	proliferation
protein coding	Regulation of p-AKT, p-	and inhibits cell apoptosis in
RNA 939 or	mTOR, P62 and BCL-2	TSCC. Over expression: In-
lncRNA RP5-	Oncogene: activates the	hibition of apoptosis, pro-
916L7.2 3. Cancer	Wnt/?-catenin signalling	motes cell proliferation, in-
Susceptibility	pathway Up regulates miR-	creases local recurrence in
9-CA SC 9 4.	185-5p target gene cyclin D2	OSCC cases. High expres-
MYC-induced	(CCND2) by competitively	sion: promotes cell prolifera-
long non-coding	sponging miR-185-5p and	tion Up-regulation results in
RNA-MINCR	then activating CCND2	OSCC cell proliferation, tu-
5. Long non-	signalling pathways involved	mor growth, increased Ki-
coding RNA	in cell cycle progression from	67 index and decreased the
Protein Disulfide	G1 to S phase	survival rates. Prognostic
Isomerase Family		biomarker to distinguish pa-
A Member 3		tients with higher risks of
Pseudogene 1-		OSCC progression. and reg-
LncRNA PDIA3P		ulates $G0/G1$ stage in OSCC.
а т		
6. Long non-	Induced by higher levels of	Upregulated TUG increased
6. Long non- coding RNA	Induced by higher levels of notch 1. Acts a	Upregulated TUG increased cell proliferation
6. Long non- coding RNA Taurine	Induced by higher levels of notch 1. Acts a sponge of miR-145 that regu-	Upregulated TUG increased cell proliferation and decreased apoptosis.
6. Long non- coding RNA Taurine upregulated	Induced by higher levels of notch 1. Acts a sponge of miR-145 that regu- lates Wnt/b-	Upregulated TUG increased cell proliferation and decreased apoptosis.
6. Long non- coding RNA Taurine upregulated gene 1-LncRNA	Induced by higher levels of notch 1. Acts a sponge of miR-145 that regu- lates Wnt/b- catenin signalling pathway.	Upregulated TUG increased cell proliferation and decreased apoptosis.
6. Long non- coding RNA Taurine upregulated gene 1-LncRNA TUG 1	Induced by higher levels of notch 1. Acts a sponge of miR-145 that regu- lates Wnt/b- catenin signalling pathway. TUG 338	Upregulated TUG increased cell proliferation and decreased apoptosis.
6. Long non- coding RNA Taurine upregulated gene 1-LncRNA TUG 1 & 338	Induced by higher levels of notch 1. Acts a sponge of miR-145 that regu- lates Wnt/b- catenin signalling pathway. TUG 338 dysregulated down stream	Upregulated TUG increased cell proliferation and decreased apoptosis.
6. Long non- coding RNA Taurine upregulated gene 1-LncRNA TUG 1 & 338	Induced by higher levels of notch 1. Acts a sponge of miR-145 that regu- lates Wnt/b- catenin signalling pathway. TUG 338 dysregulated down stream factors STAT 1,	Upregulated TUG increased cell proliferation and decreased apoptosis.
6. Long non- coding RNA Taurine upregulated gene 1-LncRNA TUG 1 & 338	Induced by higher levels of notch 1. Acts a sponge of miR-145 that regu- lates Wnt/b- catenin signalling pathway. TUG 338 dysregulated down stream factors STAT 1, ATK and caspase 3.	Upregulated TUG increased cell proliferation and decreased apoptosis.
 Long non- coding RNA Taurine upregulated gene 1-LncRNA TUG 1 & 338 7 Long Intergenic 	Induced by higher levels of notch 1. Acts a sponge of miR-145 that regu- lates Wnt/b- catenin signalling pathway. TUG 338 dysregulated down stream factors STAT 1, ATK and caspase 3. Regulates expression of Ec-	Upregulated TUG increased cell proliferation and decreased apoptosis. Up-regulation inhibits apop-
 6. Long non- coding RNA Taurine upregulated gene 1-LncRNA TUG 1 & 338 7 Long Intergenic non- 	Induced by higher levels of notch 1. Acts a sponge of miR-145 that regu- lates Wnt/b- catenin signalling pathway. TUG 338 dysregulated down stream factors STAT 1, ATK and caspase 3. Regulates expression of Ec- tonucleotide	Upregulated TUG increased cell proliferation and decreased apoptosis. Up-regulation inhibits apop- tosis. Decreased
 6. Long non- coding RNA Taurine upregulated gene 1-LncRNA TUG 1 & 338 7 Long Intergenic non- coding RNA 261- 	Induced by higher levels of notch 1. Acts a sponge of miR-145 that regu- lates Wnt/b- catenin signalling pathway. TUG 338 dysregulated down stream factors STAT 1, ATK and caspase 3. Regulates expression of Ec- tonucleotide Pyrophosphatase/ Phospho-	Upregulated TUG increased cell proliferation and decreased apoptosis. Up-regulation inhibits apop- tosis. Decreased expression shows better prog-
 6. Long non- coding RNA Taurine upregulated gene 1-LncRNA TUG 1 & 338 7 Long Intergenic non- coding RNA 261- 	Induced by higher levels of notch 1. Acts a sponge of miR-145 that regu- lates Wnt/b- catenin signalling pathway. TUG 338 dysregulated down stream factors STAT 1, ATK and caspase 3. Regulates expression of Ec- tonucleotide Pyrophosphatase/ Phospho- diesterase	Upregulated TUG increased cell proliferation and decreased apoptosis. Up-regulation inhibits apop- tosis. Decreased expression shows better prog- nosis in TSCC
 6. Long non- coding RNA Taurine upregulated gene 1-LncRNA TUG 1 & 338 7 Long Intergenic non- coding RNA 261- LINC00261 	Induced by higher levels of notch 1. Acts a sponge of miR-145 that regu- lates Wnt/b- catenin signalling pathway. TUG 338 dysregulated down stream factors STAT 1, ATK and caspase 3. Regulates expression of Ec- tonucleotide Pyrophosphatase/ Phospho- diesterase (ENPP) 4 and ENPP5 that	Upregulated TUG increased cell proliferation and decreased apoptosis. Up-regulation inhibits apop- tosis. Decreased expression shows better prog- nosis in TSCC
 6. Long non- coding RNA Taurine upregulated gene 1-LncRNA TUG 1 & 338 7 Long Intergenic non- coding RNA 261- LINC00261 	Induced by higher levels of notch 1. Acts a sponge of miR-145 that regu- lates Wnt/b- catenin signalling pathway. TUG 338 dysregulated down stream factors STAT 1, ATK and caspase 3. Regulates expression of Ec- tonucleotide Pyrophosphatase/ Phospho- diesterase (ENPP) 4 and ENPP5 that are involved in	Upregulated TUG increased cell proliferation and decreased apoptosis. Up-regulation inhibits apop- tosis. Decreased expression shows better prog- nosis in TSCC
 6. Long non- coding RNA Taurine upregulated gene 1-LncRNA TUG 1 & 338 7 Long Intergenic non- coding RNA 261- LINC00261 	Induced by higher levels of notch 1. Acts a sponge of miR-145 that regu- lates Wnt/b- catenin signalling pathway. TUG 338 dysregulated down stream factors STAT 1, ATK and caspase 3. Regulates expression of Ec- tonucleotide Pyrophosphatase/ Phospho- diesterase (ENPP) 4 and ENPP5 that are involved in tumor development. Low ex-	Upregulated TUG increased cell proliferation and decreased apoptosis. Up-regulation inhibits apop- tosis. Decreased expression shows better prog- nosis in TSCC
 6. Long non- coding RNA Taurine upregulated gene 1-LncRNA TUG 1 & 338 7 Long Intergenic non- coding RNA 261- LINC00261 	Induced by higher levels of notch 1. Acts a sponge of miR-145 that regu- lates Wnt/b- catenin signalling pathway. TUG 338 dysregulated down stream factors STAT 1, ATK and caspase 3. Regulates expression of Ec- tonucleotide Pyrophosphatase/ Phospho- diesterase (ENPP) 4 and ENPP5 that are involved in tumor development. Low ex- pression of	Upregulated TUG increased cell proliferation and decreased apoptosis. Up-regulation inhibits apop- tosis. Decreased expression shows better prog- nosis in TSCC

1	HOXA transcript at the distal tip- HOTTIP		
2	A disinte- grin and metalloproteinase with a throm- bospondin type 1 motif 9 antisense 2- ADAMTS9- AS2		
3	Krueppel- like factor 8 (KLF 8) regulated Long non coding		
Year4. 2020	AC132217.4 Long Intergenic Non- Protein Coding RNA 958- LINC00958		
Volu5me 6. XX 7. 8.	Long intergenic		
Is-	non-		
sue V	protein-		
Ver-	RNA 673-		
sion	LINC00673		
Ι	Long intergenic		
	non-		
	coding BNA		
	00152-		
	LINC00152		
	Small nucleolar		
	RNA host	10	
	gene 20- LncRNA		

 $\mathbf{5}$

1. Chemotherapy-induced	Activates Wnt/b catenin pathway
lncRNA 1-CILA1	
2. KCNQ1 overlapping	Transcriptionally silences KCNQ1
transcript 1-KCNQ1OT1	locus by regu- lating histone methylation, ceRNA to miRNA 211-5p
	that regulates Ezrin (also known as cytovillin or villin-2) /focal adhesion kinase (Fak)/Src (non receptor tyrosine kinase) signalling
3. Long non coding RNA-	Possess complementary sequence to
lnc-p23154	miR-378a-3p promoter region. miRNA
	378-3p targets 3?UTR of Glut1,
	inhibiting its expression & in turn glycolysis. Lnc-p23154 interacts with miR-378a-3p promoter to repress its transcription, and then increases Glut1 expression.
III. LNCRNAs Down-Regulated d OSCC Progressiona) Maternally expressed gene 3-M	uring 11 EG3

Mapped to chromosome 14q32.2, MEG 3 is expressed in normal human tissues and posses tumor suppressor

19 CONCLUSION

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