Review

Impact of Ubiquitination Signaling Pathway Modifications on Oral Carcinoma

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Abstract. Among intra-cellular homeostasis mechanisms, ubiquitination plays a critical role in protein metabolism regulation by degrading proteins via activating a broad spectrum of ubiquitin chains. In fact, ubiquitination and sumoylation signaling pathways are characterized by increased complexity regarding the molecules and their interactions. The Ubiquitin-Proteasome System (Ub-PS) recognizes and targets a broad spectrum of protein substrates. Ubiquitin conjugation modifies each substrate protein determining its biochemical fate (degradation). A major functional activity of Ub-PS is autophagy mechanism regulation. Interestingly, Ub-PS promotes all stages of bulk autophagy (initiation, execution, and termination). Autophagy is a crucial catabolic process that provides protein degradation and for this reason the interaction with Ub-PS is crucial. Furthermore, ubiquitination controls and regulates specific types of selective autophagy. Ub-PS is also involved in oxidative cellular stress and DNA damage response. Additionally, the functional role of Ub-PS in ribosome machinery regulation seems to be crucial. Concerning carcinogenesis, Ub-PS is involved in malignant disease development and progression by negatively affecting the corresponding TGF-B-, MEEK/MAPK/ERK-JNK-dependent signaling pathways. In the current review article, we describe the role of Ub-PS biochemical modifications and alterations in oral squamous cell carcinoma (OSCC).

Ubiquitin is a covalent protein modifier, with major role in protein modification. Ubiquitin pathways comprise a variety of intracellular protein modification systems, which regulate both the final form and the quantity of multiple proteins (1). A growing body of research has enriched our understanding of the role of ubiquitination in a number of physiological processes (2). Existing data indicate that ubiquitination and sumoylation signaling pathways are characterized by increased complexity regarding the participating molecules and their interactions (3-5). The ubiquitin-proteasome system (Ub-PS) recognizes and targets a broad spectrum of protein substrates. Ubiquitin conjugation modifies each substrate protein determining its biochemical fate (degradation). A major functional activity of Ub-PS is autophagy mechanism regulation (6, 7). Interestingly, Ub-PS promotes all stages of bulk autophagy (initiation, execution, and termination). Autophagy is a crucial catabolic process that provides final protein degradation and for this reason the interaction with Ub-PS is crucial (8). Furthermore, ubiquitination controls and regulates specific types of selective autophagy (9-11). Ub-PS is also involved in oxidative cell stress and DNA damage response (12). In conjunction, the functional
role of Ub-PS in ribosome machinery regulation seems to be crucial. Ubiquitination is a significant and major post-translational mechanism for modifying ribosomal proteins. For this reason, the 40S ribosome subunit ubiquitination/deubiquitination balance and the efficient translation eS7 are very important biochemical events (13-15).

Conversely to its physiological activity, abnormal Ub-PS modifications are implicated in non-malignant inflammatory and autoimmune phenotypes including arthritis, atherogenesis, psoriasis, uveitis, etc. (16-19). Furthermore, aberrant ubiquitination appears to be involved in malignant disease development and progression by negatively affecting the corresponding TGF-B-, MEEK/MAPK/ERK-, JNK-dependent signaling pathways (20, 21). Therefore, in the current review article, we describe the role of Ub-PS modifications/alterations in the development of oral squamous cell carcinoma (OSCC) biochemical and molecular substrate.

**Protein Ubiquitination: Molecules and Mechanisms**

Ubiquitin discovery in the 1970s has eventually led to an improved understanding of the nature and interactions of post-translational modifications (22). Ubiquitin is a highly conserved 8.6-kDa (76 amino acids chain) protein, which includes seven lysine residues in its stereo-chemical structure. The ubiquitination procedure represents a major post-translational mechanism that drives protein substrates from synthesis and structural maturation to degradation/breakdown (23, 24). Furthermore, it affects protein-protein interactions, protein localization, motivation, and kinase kinetics into the cell cycle (25, 26). Besides these activities, ubiquitination regulates a variety of intracellular functions including endocytosis, DNA damage repair, and gene transcription and translation (27-29). Ubiquitination also affects inflammatory and autoimmune disease mechanisms by its involvement in the degradation of specific proteins (30).

Ub-PS comprises of three main enzymes: E1 ubiquitin activating enzyme, E2 ubiquitin conjugating enzyme, and E3 ubiquitin ligase enzyme (31). The conjugation of ubiquitin to the lysine residue of other proteins is the result of their sequential action catalyzed by specific deubiquitinating enzymes (32). Biochemically, ubiquitination is based initially on the ubiquitin C-terminal glycine residue activation demanding ATP consumption. Progressively it finally leads to ubiquitin conjugation to the corresponding protein substrates (33). Mono- or multiple ubiquitin chains interact with Lys residues of target proteins. Alternatively, different ubiquitin moieties create new forms of chains by binding to N-terminal methionine residues or to the previously referred lysine residues. Furthermore, other biochemical reactions including acetylation, sumoylation, and phosphorylation provide complexity to the ubiquitin-based modifications. Another important regulatory action of ubiquitin is referred to E2F1 transcription factor regulation (34). Specific linked chains including K11/K48 lead to E2F1 degradation whereas K6, K27, K33, K63 demonstrate a non-degradative influence (35).

Besides, ubiquitination, autophagy plays also major role in intracellular protein degradation. Although these two mechanisms demonstrate different, independent functional pathways, it seems that there is some cross talk and interactions between them. Autophagy represents a degradation mechanism provided by lysosome activation (36). In fact, it acts as a critical catabolic process, as a response to intracellular stress. The formation of specific intracellular domains (autophagosomes) is followed by lysosome-dependent fusion process and molecule degradation (37). Hypoxia, cell organelle dysfunction and damage, oxidative stress and lack of critical ATP-based energy levels are intracellular signals for inducing autophagy motivation (38). At the molecular level, ULK1 serine/threonine kinase represents a major gene activated by stress signals leading to a catarract of molecules phosphorylation (39). The PI3K complex (lipid kinase VPS34, Beclin-1, VPS15, and ATG14) is activated by ULK1 and PI3P/ATG9/ATG2/WIPI interactions leading to autophagosome formation. Interestingly, ATG9-as a transmembrane protein- is responsible for providing membrane substrates to autophagosomes (40).

Concerning the influence of ubiquitination in autophagy regulation, it seems that there is a positive feedback loop. ULK1 and Beclin-1 molecules are targets for ubiquitination under the pressure of E3 ligases and especially the TRAF6, which is activated by the AMBRA1 gene, a PI3K component (41). This gene supports normal function ofULK1 by inducing K63-dependent ubiquitination (42). Similarly, TRAF6 enhances Beclin-1K63-dependent ubiquitination (43). In fact, the two genes Beclin-1 and AMBRA1 regulate not only initiation, but also termination of the autophagy phenomenon. Interestingly, Beclin-1 subsequent degradation is mediated by the influence of HECT-type E3 ligase that provides Lys-11-linked ubiquitination of the gene (44). Additionally, Beclin-1 is indirectly regulated by the E3 ligase PARKIN. Furthermore, AMBRA1 is negatively affected by the expression of phosphorylated mTORC1. Besides the main autophagy procedure, selective autophagy variants are also exposed to the influence of ubiquitination. E3 ligase is involved in aggrephagy, mitophagy, lysophagy, and xenophagy and all of them are stimulated by Lys-63 poly-Ub chains (45).

**Ubiquitination Modifications in Oral Carcinoma**

Oral squamous cell carcinoma (OSCC) represents a major malignancy in Head and Neck Squamous Cell Carcinoma (HNSCC) super-family. OSCCs frequently present an aggressive phenotype, with an increased tendency to present local and distant metastatic lymph nodes, as a result of severe genetic alterations (46). Etiopathogenetic factors that lead to OSCC development and progression include tobacco,
alcohol chronic consumption and also viral-mediated deregulation (47). Concerning viral oncogenic activity, persistent Human Papilloma Virus (HPV) infection is responsible for malignant transformation of the affected oral/oropharyngeal epithelia modifying the host cell genome (48). Concerning ubiquitination modifications in OSCC, a variety of molecules are implicated in them (Figure 1). Ubiquitin D appears to play a pivotal role in malignancy. A study group evaluated its expression by implementing a combined immunohistochemistry (IHC), quantitative real-time polymerase chain (qRT-PCR) reaction and a western blot protocol, in a series of OSCCs (49). They reported that a significant over expression of the molecule correlated with induced proliferation, migration and invasion. They also suggested that ubiquitin D should be potentially considered a prognostic biomarker for monitoring OSCC patients.

Another molecule involved in OSCC progression is the ubiquitin-specific protease 14 (USP14). Implementing experimental colony formation analysis in OSCC cell cultures and tissue specimens, a study group observed a significant over expression of the marker (50). Interestingly, USP14 up-regulation was associated with poor prognosis. Furthermore, USP14 over activation reduced apoptosis and autophagy leading to elevated radio-resistance in them. For these reasons, the authors suggested USP14 as a molecule essential for targeted therapeutic strategies. Similarly, another study focused on the aberrant expression of STUB1 based ubiquitin E3 ligase (51). Implementing a qRT-PCR assay on OSCCs they reported reduced STUB1 expression, associated to significant recurrence rates. Additionally, STUB1 progressive inactivation was correlated with TGM2 up-regulation in these cases, a protein related with malignancy aggressiveness. Another study group evaluated the impact of ubiquitin-specific processing proteases 17 (DUB3) expression on OSCC (52). Applying a qRT-PCR, IHC, western blot and flow cytometry combined assay they observed that DUB3 over expression led to increased cancerous cell proliferation and viability, combined with low apoptotic levels. They also suggested that DUB3 should be a potential target for OSCC therapeutic strategies.

Besides DUB3 over activation, increased expression of ubiquitin-specific protease 22 (USP22)-acting as deubiquitinating hydrolase- is involved in the development and progression of OSSC. A study group co-analyzed USP22, survivin and aurora-B molecules reported high expression levels for all of them. Interestingly, RING finger protein 139 (RNF139) E3 ubiquitin ligase activation reduced cancerous cells proliferation in a molecular study (53). The study group analyzed its expression in SCC9 and SCC25 cell cultures suggesting that the molecule is involved in aggressive phenotypes regarding OSSC (tongue carcinoma).
The role of another molecule – the ubiquitin-specific protease 9 (USP9X) acting as deubiquitinase – in OSCC onset and progression is under investigation. A study group analyzed the co-expression of USP9X and MCL-1, an anti-apoptotic protein, by IHC immunofluorescence, and flow cytometry. In fact, USP9X deubiquinates MCL-1 protein. They also reported high expression levels of both molecules. Interestingly, USP9X/MCL-1 co-over expression was associated to aggressive phenotypes in OSCC combined with poor prognosis. USP9X plays also a significant role by deubiquitinating the critical immune checkpoint protein programmed cell death ligand 1 (PD-L1) in OSCC (54). USP9X stabilizes PD-L1 over expression enhancing also cancerous cell proliferation.

In conclusion, ubiquitination plays a critical role in protein metabolism regulation by degrading proteins, and by also providing intra-cellular homeostasis. Ubiquitination and sumoylation signaling pathways comprise a significant number of molecules. The Ub-PS recognizes and targets a broad spectrum of protein substrates. Ubiquitin conjugation modifies each substrate protein determining its biochemical fate (degradation). A major functional activity of Ub-PS is autophagy mechanism regulation. Interestingly, Ub-PS promotes all the stages of bulk autophagy (initiation, execution, and termination). Autophagy interacts with Ub-PS in the cell micro-environment. Furthermore, ubiquitination controls and regulates specific types of selective. Ub-PS is also involved in oxidative cellular stress and DNA damage response. Interestingly, the functional role of Ub-PS in ribosome machinery regulation seems to be crucial. Concerning carcinogenesis, it is involved in malignant diseases development and progression by negatively affecting the corresponding TGF-B-, MEEK/MAPK/ERK-, JNK-dependent signaling pathways. Continually enriched molecular data reveal new interactions involved in the ubiquitination process that drive cancerous cells to differences and imbalances regarding their metabolic potential providing new knowledge on this topic. For example, specific mutations in signalling pathways -such as NOTCH1C1133Y mutation- strongly interact with E3 ubiquitin ligase (FBXW7) by degrading it (55). Therefore, exploring and understanding the role of ubiquitination modifications in OSCC and neoplasia in general, is a crucial step for identifying patients with specific molecular signatures, potentially eligible for targeted therapeutic strategies.

Conflicts of Interest

The Authors declare no conflicts of interest associated with this article.

Authors’ Contributions

ET, EK, VR, VP, AC researched the literature and drafted the article, with ET a major contributor in writing the article. NM, LM, DR, PP, OD collected the data provided by the corresponding references. All Authors read and approved the final article.

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