The Role of CpG24 Hyper-Methylation on the Down-Regulation of Rap1Gap in Papillary Thyroid Cancer

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ABSTRACT

Thyroid tumors may emerge following hypermethylation in the administrative area of silencer qualities, for example, Rap1Gap. In this examination, we intended to explore the quality articulation of Rap1Gap and DNA methylation example of CpG24 in Iranian patients with papillary thyroid malignant growth (PTC). Thyroid disease is the most predominant endocrine harm and its rate is expanding around the world. Papillary thyroid disease (PTC) represents around 80% of all thyroid malignant growth. Hereditary and epigenetic alterations are engaged with commencement and movement of thyroid malignant growth, of which transformations prompting the mitogen-activated protein kinase (MAPK) and phosphatidylinositol-4,5-bisphosphate 3-kinase (PI3K) flagging pathways initiation are vital. Other hereditary and epigenetic factors which add to the threatening forecast of PTC are inefficiently characterized. Understanding these atomic modifications and their components may bring about the advancement of novel sub-atomic prognostic and helpful methodologies. DNA hyper-methylation is one of the epigenetic administrative systems which may quiet the objective quality without influencing the DNA grouping.

Rap1Gap quality in the 1p36 locale encodes a kind of GTPase-enacting protein (GAP) that down-directs the movement of Ras-related Rap1 protein. The result of Rap1Gap quality advances the hydrolysis of bound GTP and returns Rap1 to the latent state. RAS superfamily proteins, for example, Rap1, assume key jobs in receptor-connected flagging pathways that control cell development and separation. Rap1 assumes the job in various procedures, for example, cell expansion, separation, and embryogenesis. It is accounted for that Rap1Gap is a tumor silencer quality which is down-directed in different malignancies, for example, squamous cell carcinoma, renal cell carcinoma, melanoma and thyroid disease. Rap1Gap articulation and its action are generally controlled at transcriptional and post-transcriptional levels. Tumors may emerge following hyper-methylation in the administrative locale of Rap1Gap quality. Understanding the example of tumor silencer quality down-guideline and fundamental atomic instruments, in thyroid malignancy, may give a noteworthy clinical effect. In this examination, we intended to research the quality articulation of Rap1Gap and DNA methylation example of CpG24 district in Iranian patients with papillary thyroid disease.

There are a couple of explores done about the job of Rap1Gap quality in deciding thyroid malignant growth forecast. In this examination we found that hyper-methylation of CpG24 is related with tumor size and lymph hub inclusion. In this way, this epigenetic variety can be utilized as a guess biomarker in PTC patients, however progressively extensive examinations are
required. our examination quality incorporates its oddity for Iranian populace. In spite of this quality, there were a few constraints; first, methylation-explicit PCR (MSP) was restricted to give the data at the degree of individual CpG site, in light of the fact that there were different CpG islands in the administrative district of the applicant quality. Second, it was unthinkable for us to discover increasingly thyroid malignant growth cell lines. As per this investigation, we can presume that unusual DNA methylation in the CpG24 locale, autonomous of Rap1Gap quality articulation, could be considered as an epigenetic biomarker for papillary thyroid malignancy.

**Keywords:** Epigenetic; DNA methylation; Papillary thyroid cancer; Tumor suppresser gene; Gene expression