We studied the expression and activation of the main effector protein kinase of phosphatidylinositol-3-kinase cascade (PI3K) – Akt in conventionally normal tissues, benign and highly differentiated (with and without metastases) human thyroid tumors. There was a difference in the Akt1 amount in tumor tissue compared with normal tissue in papillary carcinomas and tissue of multinodular goiter. Akt expression both in tumor and conventionally normal tissues of follicular adenoma was significantly lower than in follicular carcinoma. The lowest level of Akt expression was observed in tissues of multinodular goiter. Total activity of all three isoforms of Akt1/2/3 was lower in tumors compared to conventionally normal tissue. Thus, Akt activity (according to Thr308 phosphorylation) is not associated with proliferative processes in the tumor tissue of the thyroid. Apoptosis level detected in these tissues was not associated with the protein kinase activity either. Possible mechanisms of signaling cascade PI3K/Akt inhibition in thyroid tumors are discussed.

**Key words:** thyroid gland, benign and malignant tumors, signaling cascade PI3K/Akt.

Proliferative processes in tumor cells are controlled by two main pathways: PI3K/Akt and mitogen-activated protein kinases (MAPK). The latter actually regulates cell division. PI3K/Akt prepares a cell for division.

Akt, also known as protein kinase B (PKB), belongs to a family that combines protein kinases C, cAMP-dependent and cGMP-dependent kinases. The gene encoding Akt was identified as a part of the genome of the murine leukemia transforming virus (v-akt murine thymoma viral oncogene homolog) in 1977 and classified as an oncogene. The man's Akt family includes three evolutionarily conserved isoforms: Akt1 (including 3 splicing variants), Akt2 and Akt3 (2 splicing variants). Akt isoforms play a key role in various cellular processes including apoptosis, growth, proliferation, polarity, migration, DNA repair, glucose transport, metabolism, skeletal muscle and cardiomyocytes contractility, angiogenesis and self-renewal of stem cells. Dysregulation of Akt activity is associated with malignant transformation of cells, cardiovascular disease, type 2 diabetes, muscular hypotrophy and neurodegenerative diseases [1].

Signaling cascade PI3K/PDK/Akt is involved in the regulation of protein synthesis and cell energy supply, i.e. in preparation of a cell for mitosis. Moreover, this signaling cascade inhibits apoptosis, promotes the survival of tumor cells and is activated in many types of cancer [2]. Furthermore, mutations and amplification of individual components of PI3K/Akt pathway causes malignant transformation of cells of different origins, including thyroid [3], while inhibiting the cascade with specific inhibitors enhances the therapeutic effect of anticancer drugs [4]. Akt activity is mainly regulated through phosphorylation of Thr308 and Ser473 amino acid residues by PDK1 and mTORC2 kinases, respectively.

A significant increase of proliferating cell nuclear antigen (PCNA) in thyroid tumors, especially in aggressive tumors with metastases, indicating a substantial increase of proliferative processes was demonstrated previously [5]. The aim of our work was to ascertain the expression and activation peculiarities of Akt as the phosphatidylinositol-3-kinase cascade main effector kinase, and the apoptosis level in human thyroid normal tissues, benign and malignant tumors.
Materials and Methods

The study was conducted on patient’s postoperative material obtained in the surgical department of the Institute. All patients signed informed consent before surgery to the use of the postoperative material for research. As conventionally normal tissue unaltered thyroid tissue was taken, which by morphological criteria did not differ from normal. After removal thyroid tissue was immediately placed on ice and quickly frozen at -80°C.

To determine Akt1 and phospho-Akt1/2/3 (phospho-Thr308) in thyroid homogenates Abcam (UK) ELISA kits ab176658 were used. To determine cleaved PARP ELISA kits ab119690 of the same company were used. Measurements were performed on microplate reader of Bio-tek Instruments (USA) at a wavelength of 450 nm. The study was performed in 3 replications. The number of transformed tissue samples was 3 (follicular carcinoma and multinodular goiter) and 6 (follicular adenoma, encapsulated tumor papillary carcinoma and non-encapsulated tumor papillary carcinoma). Tissue was homogenized in a Retsch TissueLyser II homogenizer (Germany) in a special kit provided buffer that prevents proteins degradation and dephosphorylation. Protein concentration in the lysate was determined with a Novagen (USA) BCA protein assay kit.

The experiments results were presented as $M \pm m$; $n = 3 - 6$. $t$-Student test was used to compare these data groups.

Results and Discussion

Table 1 shows that there is a fairly high level of Akt expression in all studied tissues, both normal and tumoral. The content of Akt1 in conventionally normal and tumor tissue of follicular carcinoma (FTC) and follicular adenoma (FA) did not differ significantly, in contrast to papillary carcinomas (PTC) and goiter (MNG), where the level of kinase in tumor tissue exceeded the one in conventionally normal tissue almost 2 times (Table 1). It should be noted that Akt expression both in tumor and conventionally normal tissues of FA was significantly lower than in FTC tissues. The lowest level of Akt expression we observed in multinodular goiter (Table 1).

It is known that disruption of the PI3K/Akt pathway as a result of Ras, PTEN, PIK3CA genes mutations and amplifications is one of the reasons for thyroid papillary carcinoma genesis [3], although RET gene rearrangements and mutations in genes encoding the protein kinases of MAPK cascade are more typical of this type of cancer. It is harder to explain the lack of difference in the Akt1 expression in follicular carcinomas compared to the normal tissue, since PI3K catalytic subunit – PIk3cA – gene activating amplification and mutations are found more often in the FTC – 10-15 and 24% of cases, respectively [3].

A different picture was observed regarding Akt1/2/3 activation (Fig. 1). The level of phospho-Akt in conventionally normal tissue of all tumors was considerably higher (almost 8.5 times for the FTC and more than 18 times for nPTC) than in tumor tissue, and was practically absent in encapsulated papillary carcinomas (Fig. 1). Thus, contrary to expectations, the Akt activity in papillary and follicular carcinomas and follicular adenomas was either absent or substantially suppressed, indicating a lack of link between the protein kinase activation by the PI3K/PDK-1 pathway (phosphorylation of Thr308 residue) and enhanced proliferative processes in thyroid tumors, which we observed in the same samples previously [5].

It was established that in tumor tissues Akt suppresses apoptosis by inhibiting caspase-9, pro-apoptotic protein Bad, FKHR and FOHO transcription factors; affects the activity of cell cycle inhibitors p21, p27, Gsk-3β protein kinase and the state of Mdm2 – a tumor suppressor p53 regulator – that generally causes cell cycle dysregulation and un-

<table>
<thead>
<tr>
<th></th>
<th>Conventionally normal tissue (OD)</th>
<th>Transformed tissue (OD)</th>
</tr>
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<tbody>
<tr>
<td>FTC</td>
<td>0.360 ± 0.042</td>
<td>0.335 ± 0.048</td>
</tr>
<tr>
<td>FA</td>
<td>0.221 ± 0.032</td>
<td>0.182 ± 0.073</td>
</tr>
<tr>
<td>iPTC</td>
<td>0.198 ± 0.013</td>
<td>0.327 ± 0.072*</td>
</tr>
<tr>
<td>nPTC</td>
<td>0.228 ± 0.004</td>
<td>0.416 ± 0.146*</td>
</tr>
<tr>
<td>MNG</td>
<td>0.092 ± 0.006</td>
<td>0.159 ± 0.026*</td>
</tr>
</tbody>
</table>

Note. In the table and further figures: FTC – follicular carcinoma, FA – follicular adenoma, iPTC – papillary carcinoma (encapsulated tumors), nPTC – papillary carcinoma (non-encapsulated tumors), MNG – multinodular goiter. OD – optical density at 450 nm per mg of protein, $M \pm m$; $n = 3$ (FTC, MNG) and 6 (FA, iPTC, nPTC); * difference between conventionally normal and tumor tissues is significant, $P < 0.05$.
controlled proliferation; activates IKK and NF-κB-dependent signaling pathway, promoting tumor cell survival, angiogenesis and metastasis formation; increases tumor growth through mTOR activation [2, 6]. Furthermore, excessive Akt activation leads to tumor resistance to radiation and chemotherapy [7, 8]. Therefore, we determined the apoptosis level by cleaved PARP larger fragment (89 kDa) amount, which is characteristic for apoptotic responses. Fig. 2 shows that the PARP fragment amount in papillary carcinomas and goiter doesn’t differ significantly from conventionally normal tissue. In follicular carcinoma samples cleaved PARP amount is higher, and in follicular adenoma samples it is lower compared to conventionally normal thyroid tissue.

It was expected that increased Akt activity/content would lead to apoptosis intensity reduction and vice versa. However, increased amount of cleaved PARP against reduced kinase activity was observed only in follicular adenoma tissue (Fig. 2). Thus, Akt content and activity are not related to the apoptosis level in the studied tissues.

The fact of considerable kinase activity inhibition in thyroid carcinomas deserves special attention. A possible explanation for this are data indicating that Akt might participate in the replicative senescence of normal and tumor cells [9-12], a phenomenon, that along with apoptosis inhibits tumor growth. In addition, Akt can stimulate apoptosis [13] and inhibit the breast cancer cells migration [14].

Cell cycle regulator (inhibitor of cyclin-dependent kinases) p21WAF1 is phosphorylated only by Akt1, that negatively regulates the cell cycle and proliferation [14]. Thus, under certain conditions Akt may show anti-proliferative and cancerostatic properties. This phenomenon was first demonstrated regarding MAPK and was named oncogene toxicity. It was shown that although Ras and Raf oncogenes are often involved in malignant transformation of thyroid cells, constitutive activation of this cascade in tumor tissues leads to growth arrest and senescence in many cases [15-18]. For example activated Ras or c-Raf-1 cause cell growth arrest by producing and secreting autocrine/paracrine factor LIF (leukemia inhibiting factor) in human medullary thyroid carcinoma cells [15]. Sustained activation of Raf/MEK/ERK signaling pathway causes growth arrest, accompanied by corresponding cell cycle regulators states changes (pRB dephosphorylation, E2F1 down-regulation and p21WAF1 up-regulation), specific changes in cells morphology and c-Myc or RET expression in LNCaP, U251, and TT human tumor lines (the latter - medullary thyroid carcinoma) [17].

Cancer cells induce special protective mechanisms, such as the heat shock protein mortalin synthesis [19], which inhibits MAPK expression and activation and thus protects cells from aging, growth arrest and apoptosis. Therefore, it is possible that, as is the case with MAPK [5, 16, 17, 19], thyroid tumor cells initiate special defense mechanisms that inhibit Akt activation and thus protect themselves against oncogene toxicity – senescence and cell cycle arrest.
Another question that arises in the analysis of the obtained data is how a tumor cell replaces inactive Akt – one of the major protein kinases that control growth, energetics and cell division. The likely answer is contained in the works that suggest Akt replacement by other protein kinases in PI3K signaling pathway, including Sggk3 (serum/glucocorticoid regulated kinase) [20].

There is also the possibility of alternative Akt activation through Ser473 residue phosphorylation of mTORC2 and DNA-PK protein kinases complexes. However, it is known that such phosphorylation stimulates full Akt activity and, consequently, suppresses apoptosis by inhibiting FOXO proapoptotic protein family [21], which we have not observed in most thyroid tumors (Fig. 2). In addition, a number of protein kinases that are involved in oncogenesis, activate Akt, phosphorylating other amino acid residues of the kinase. Thus, Ack1 (TNK2) phosphorylates Tyr176 residue; Src and RTK6 – Tyr215, Tyr315 and Tyr326; TBK1 (TANK-binding kinase 1) – Thr195, Ser378, Ser473 [22].

Thus, our findings indicate that Akt activity (by Thr308 phosphorylation) is not associated with thyroid tumor tissue proliferative processes. Apoptosis level, which was determined in the same tissues, does not correlate with the protein kinase activity either.

Ключевые слова: щитоподобная зазолаза, доброкакачественные и злонакачественные узлы, механизм угнетения PI3K/Akt сигнальной каскады.

АКТИВНОСТЬ ПРОТЕИНКИНАЗЫ Akt В ОПУХОЛЯХ ЩИТОПОДОБНОЙ ЖЕЛЗЫ ЧЕЛОВЕКА

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Выводы и рекомендации. Уровень экспрессии и активации главной каскадной протеинкиназы PI3K/Akt в опухолях щитовидной железы человека был достоверно выше, чем в нормальных тканях. Экспрессия Akt в нормальных тканях щитовидной железы была достоверно выше, чем в опухолевых тканях. Уровень апоптоза,
который определялся в этих же тканях, также не коррелирует с активностью протеинкиназы. Обсуждаются возможные механизмы подавления активности сигнального каскада РІ3К/Akt в опухолях щитовидной железы.

Ключевые слова: щитовидная железа, доброкачественные и злокачественные опухоли, сигнальный каскад РІ3К/Akt.

Reference


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