Review Article

Update on Anaplastic Thyroid Carcinoma: Morphological, Molecular, and Genetic Features of the Most Aggressive Thyroid Cancer

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Anaplastic thyroid carcinoma (ATC) is the most aggressive form of thyroid cancer. It shows a wide spectrum of morphological presentations and the diagnosis could be challenging due to its high degree of dedifferentiation. Molecular and genetic features of ATC are widely heterogeneous as well and many efforts have been made to find a common profile in order to clarify its cancerogenetic process. A comprehensive review of the current literature is here performed, focusing on histopathological and genetic features.

1. Introduction

Anaplastic thyroid carcinoma (ATC) represents the most aggressive extreme of the clinical spectrum of thyroid epithelial neoplasms, being one of the most lethal human tumors.

It constitutes less than 5% of clinically recognized thyroid malignancies but it accounts for more than half of the deaths for thyroid cancer, with a mortality rate that is over 90% and a mean survival of six months after the diagnosis.

It is defined by the WHO as a highly malignant tumor wholly or partially composed of undifferentiated cells that retain features indicative of an epithelial origin, on immunohistochemical or ultrastructural ground [1]. It usually affects elderly people, with a mean age in the mid-60s, and shows a female predominance [1].

In this review we tried to summarize the current knowledge on ATC from both morphological and biological points of view.

2. Morphological Features

Grossly, ATC is well recognized as a large, necrotic, and hemorrhagic mass that is typically widely invasive, often replacing most of the thyroid gland parenchyma with infiltration of the surrounding soft tissue and adjacent structures of the neck (Figures 1(a) and 1(b)).

The morphological spectrum depends on the admixture of three main histological patterns: spindle cell, giant cell, and squamoid [2–4]. These patterns often coexist and are not predictive of patients’ outcome but are historically used to group ATC in major histological categories and to define their main differential diagnoses. The histological categories are sarcomatoid and epithelioid-squamoid.

The small cell category, that was included in older classification of ATC, is no longer considered, as it comprised cases of bona fide lymphomas, medullary carcinomas, and insular carcinomas [2, 3, 5].
Common features to all patterns of ATC are hypercellularity, large foci of necrosis, marked invasiveness, and angiotropism with a tendency to infiltrate medium-sized veins and arteries, replacing their muscular wall [2, 3].

For diagnostic purposes, fine needle aspiration biopsy (FNAB) is an important tool and can provide a correct diagnosis of ATC in up to 84% of cases [6].

FNAB smears are usually composed of a pleomorphic cellular population in a necrotic background (Figures 2(a) and 2(b)). The tumor cells are bizarre, oval to spindle-shaped, dyscohesive elements showing anisocytosis, and irregular sometimes multiple nuclei, perfectly reflecting the sarcomatoid or epithelioid histological morphology.

2.1. Sarcomatoid Category

2.1.1. Histology. Anaplastic thyroid carcinomas with sarcomatoid appearance are characterized by spindle cells and giant cells, the most frequent patterns seen in ATC. In fact, spindle and giant cells have been found, alone or in combination, in at least 50% of cases reported by Carcangiu and colleagues [2].

**Spindle cells** show a fascicular or storiform pattern of growth, indistinguishable from a true sarcoma (Figures 3(a) and 3(b)). These neoplasms are generally well vascularized often resulting in a hemangiopericytoma-like pattern or forming anastomosing channels lined by tumor cells, resembling an angiosarcoma (Figure 3(c)). An odd variation on the theme of the spindle cell form is the paucicellular variant [7, 8]. This infrequent entity was first described by Wan et al. in 1995 as a peculiar subtype of ATC with gross and histological features closely mimicking Riedel's thyroiditis [7]. It is characterized by low cellularity with striking degree of fibrosis and hyalinization, presence of spindle cells resembling fibroblasts or myofibroblasts, absence of obvious nuclear atypia, and sprinkling of lymphocytes. Features allowing a diagnosis of ATC are (1) presence of coagulative necrosis with ghost shadows of preexisting blood vessels, (2) recognition of scattered atypia and mitosis in more cellular areas at the periphery of the fibrosis, (3) detection of blood vessels obliterated by neoplastic spindle cells, and (4) positivity for epithelial markers [7].

**Giant cells** are characterized by deep pleomorphism, having bizarre sometimes multiple hyperchromatic nuclei, abundant eosinophilic cytoplasm, and a plump, oval, or round shape (Figure 4(a)). They are typically interspersed among smaller mononuclear tumor cells with similar cytoplasmic features showing a solid architecture. The formation of alveolar, pseudoglandular, or pseudovascular structures can also be seen, probably due to an artefactual separation of the cells. The cytoplasm of the tumor cells can sometimes assume a clear or granular appearance simulating a clear cell or an oncocytic carcinoma, respectively; the presence of striking pleomorphism, high mitotic activity, and necrosis is strongly suggestive for ATC [2].

**Osteoclast-like multinucleated giant cells** are occasionally present and could be prominent, resembling similar tumors described in breast and pancreas. Osteoclast-like multinucleated giant cells are known to be reactive elements of...
monocytic/histiocytic lineage, immunohistochemically positive for CD68-KP1 (Figure 4(b)) and apparently derived from histiocytoid mononuclear cells via cellular fusion [9]. They give the tumors an appearance reminiscent of giant cell tumor of bone and soft tissue.

**Huge inflammatory infiltrate** is often present, sometimes predominantly neutrophilic in type, giving the tumor an appearance resembling inflammatory variant of malignant fibrous histiocytoma.

**Heterologous elements**, such as bone, cartilage, and skeletal muscle, can also be found. Matrix formation with chondro- and osteosarcomatous differentiation has been reported in up to 5% of anaplastic carcinoma [10]. Rhabdomyosarcomatous appearance has also been described [2, 11]. Carda et al. reported two cases, in which the skeletal muscular differentiation was demonstrated by electron microscopy and immunohistochemistry with positivity for muscle-specific actin, desmin, myogenin, and MyoD1 [11].

2.1.2. **Differential Diagnosis.** Sarcomatoid ATC closely simulates a large variety of soft tissue sarcomas. When a well-differentiated component is lacking and immunohistochemistry fails to demonstrate an epithelial differentiation, this distinction could be really difficult. Two characteristic histological features are helpful to differentiate sarcomatoid anaplastic carcinoma from a true sarcoma: the presence of angulated necrotic foci with neoplastic cells palisading around them as seen in glioblastoma of the central nervous system and the tendency of the spindle neoplastic cells to infiltrate the wall of large-sized veins and arteries [2].

It should be kept in mind however that primary sarcomas of the thyroid are indeed very rare so that it has been suggested that all sarcomatoid tumors of the thyroid gland should be regarded as ATC [12].

Primary sarcomas simulating a sarcomatoid ATC have been reported as case reports: fibrosarcoma [13], leiomyosarcoma [14], chondrosarcoma [15], osteosarcoma [16], and angiosarcoma (including epithelioid variant) [17, 18]. Metastases are possible as well and should be clinically ruled-out [19–21].

In addition, various spindle cell neoplastic and nonneoplastic thyroid lesions could simulate a sarcomatoid pattern and they should be taken into consideration by pathologist during diagnostic process. Differential diagnoses are described in Table 1.

2.2. **Epithelioid-Squamous**

2.2.1. **Histology.** Anaplastic thyroid carcinomas with epithelioid-squamous appearance are histologically less heterogeneous than sarcomatoid tumors. They are characterized by polygonal cells with a clearly epithelial appearance, growing
## Table 1: Differential diagnoses of sarcomatoid category.

<table>
<thead>
<tr>
<th>Thyroid lesions simulating a sarcomatoid pattern</th>
<th>Differential features</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Primary or metastatic sarcoma</strong></td>
<td>It is an exclusion diagnosis:</td>
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<tr>
<td></td>
<td>(i) lack of a well-differentiated component;</td>
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<td></td>
<td>(ii) no epithelial markers;</td>
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<td></td>
<td>(iii) absence of palisading necrosis and neoplastic spindle cells infiltrating the wall of large-sized vessels;</td>
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<td></td>
<td>(iv) presence of extrathyroidal sarcoma clinically detected (in metastatic disease).</td>
</tr>
<tr>
<td><strong>Other malignancies</strong></td>
<td>(i) Adolescent or young adults (mean age 15 years);</td>
</tr>
<tr>
<td><strong>SETTLE</strong> (spindle epithelial tumor with thymus-like elements)</td>
<td>(ii) biphasic pattern of growth with a predominant spindle cell component merging with a mucin-secreting glandular elements; both components have an epithelial phenotype;</td>
</tr>
<tr>
<td><strong>Spindle cell variant of papillary thyroid carcinoma</strong></td>
<td>(iii) generally indolent behavior.</td>
</tr>
<tr>
<td><strong>Spindle cell variant of medullary carcinoma</strong></td>
<td>Metaplastic variant of PTC:</td>
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<tr>
<td></td>
<td>(i) spindle cells retain even if focally nuclear features of PTC;</td>
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<tr>
<td></td>
<td>(ii) consistent immunoreactivity for thyroglobulin.</td>
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<tr>
<td><strong>Solitary fibrous tumor</strong></td>
<td>(i) Low mitotic rate (4 mitoses or fewer per 10 high-power fields);</td>
</tr>
<tr>
<td></td>
<td>(ii) no necrosis or vascular invasion;</td>
</tr>
<tr>
<td><strong>Riedel thyroiditis</strong></td>
<td>(iii) positivity for bcl-2, CD34, CD99, and vimentin and negativity for all epithelial markers.</td>
</tr>
<tr>
<td><strong>Post-fine-needle aspiration Spindle cell nodules of the thyroid</strong></td>
<td>(i) Absence of necrosis;</td>
</tr>
<tr>
<td></td>
<td>(ii) evidence of occlusive phlebitis (no angioinvasion);</td>
</tr>
<tr>
<td></td>
<td>(iii) absence of neoplasm;</td>
</tr>
<tr>
<td><strong>Benign processes</strong></td>
<td>(iv) negativity for epithelial markers;</td>
</tr>
<tr>
<td></td>
<td>(v) generally benign self-limiting disease.</td>
</tr>
<tr>
<td></td>
<td>(i) History of FNA biopsy;</td>
</tr>
<tr>
<td></td>
<td>(ii) size ranging from 3 to 10 mm;</td>
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<tr>
<td></td>
<td>(iii) not encapsulated but relatively circumscribed and located mostly in the center of preexisting thyroid nodules;</td>
</tr>
<tr>
<td></td>
<td>(iv) low mitotic rate;</td>
</tr>
<tr>
<td></td>
<td>(v) immunoreactivity for smooth muscle actin (myofibroblastic origin).</td>
</tr>
</tbody>
</table>

in solid nests, intermingled by desmoplastic stroma (Figures 5(a), 5(b), and 5(c)). Keratinization could be seen even if rarely. Squamoid pattern was present in about 20% of ATCs described in the largest series reported in literature [2, 4] and it is most frequently seen in combination with spindle and/or giant cells patterns.

Two peculiar variants of ATC belong to epithelioid-squamoid category and are described below. **Anaplastic spindle cell squamous carcinoma** is a variant of ATC with both spindle cell elements and squamous islands with focal keratinization, similar to its counterpart described in the breast [22] and oropharynx [23, 24]. It was originally described by Bronner and LiVolsi as a unique subtype of ATC associated with the tall cell variant of papillary thyroid carcinoma (TCV PTC) [25]. In a recent series it was pointed out that this variant of ATC could clinically and histologically mimic a laryngeal squamous cell carcinoma. Therefore, caution is warranted in evaluating laryngeal squamous lesions in patients with known history of TCV PTC and without known risks factors for head and neck carcinogenesis [26].

**Lymphoepithelioma-like ATC** is a subtype of epithelioid-squamoid ATC characterized by histologic features similar to those of lymphoepithelioma of the nasopharynx and lymphoepithelioma-like carcinoma (LELC) of other sites [27]. It is composed of sheaths of epithelial cells in a rich inflammatory background including lymphocytes and some plasma cells. Tumor cells are immunoreactive for epithelial membrane antigen and keratin but are negative for thyroglobulin. Notably there is not association with EBV infection, as in LELC of organs that are not embryologically derived from primitive pharynx or foregut such as skin [28], urinary bladder [29], and uterine cervix [30].

### 2.2.2. Differential Diagnosis.

Pure squamous cells carcinoma of the thyroid is exceedingly rare and is listed as a separate entity in the WHO [1]. Its clinical presentation and behavior are the same of ATC. It is by definition not associated with other types of thyroid carcinoma.
When a thyroid tumor is almost composed of squamoid elements it could be also necessary to rule out a direct invasion from an upper airway primary or a metastatic process firstly from the lung. Careful clinical examination is the most important clue, particularly to exclude metastases. The search of a well-differentiated component, by extensive sampling of the surgical specimen, is also a helpful feature in identifying the tumor origin (Figure 6) and it is mandatory in these cases. Notably Toner et al. reported some cases of ATC with endotracheal presentation, showing metaplasia or atypical, probably regenerative, epithelial changes in the adjacent airway epithelium that could be easily misinterpreted as an in situ component [31].

In thyroid, squamous differentiation may be seen in other neoplastic settings, without the meaning of anaplastic transformation. Squamous differentiation can be present as a result of a metaplastic process in papillary carcinoma, most commonly in the diffuse sclerosing variant [32], in medullary carcinoma, in mucoepidermoid carcinoma, and in sclerosing mucoepidermoid carcinoma with eosinophilia [1]. Squamous differentiation is also present in most cases of “carcinoma showing thymus-like differentiation” (CASTLE), which is thought to arise either from ectopic thymus or remnants of branchial pouches [33, 34].

On the other hand, nonneoplastic squamous cells can be present as embryonic remnants in the thyroglossal duct or structures derived from the branchial pouch (e.g., thymic epithelium) and as squamous metaplasia in thyroiditis or as a reparative phenomenon following FNAB [35] and postradiation therapy.

Differential diagnoses are summarized in Table 2.

2.3. Immunohistochemical Features. ATCs show a variable immunophenotype. Immunoreactivity for cytokeratin is present in 40% to 100% of cases according to the different series [2, 36–38]. Vimentin is consistently present in the spindle cell component, whereas EMA and CEA are particularly expressed in the squamoid cells [2].
Table 2: Differential diagnoses of squamoid category.

<table>
<thead>
<tr>
<th>Thyroid lesion simulating a squamoid pattern</th>
<th>Differential features</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pure squamous cells carcinoma</td>
<td>(i) Entirely composed of squamous cells; (ii) no evidence of another type of thyroid carcinoma in close proximity.</td>
</tr>
<tr>
<td>Primary head and neck squamous cell carcinoma</td>
<td>(i) Presence of in situ component in head and neck structures; (ii) features of “ab extrinseco” involvement of thyroid parenchyma; (iii) PAX8 is consistently negative.</td>
</tr>
<tr>
<td>Metastatic squamous cell carcinoma of the lung</td>
<td>(i) Presence of a lung nodule clinically detected; (ii) PAX8 is consistently negative.</td>
</tr>
<tr>
<td>Diffuse sclerosing variant of papillary thyroid carcinoma</td>
<td>(i) Abundant psammoma bodies; (ii) neoplastic cells retain nuclear features of PTC; (iii) immunoreactivity for thyroglobulin.</td>
</tr>
<tr>
<td>Mucoepidermoid carcinoma</td>
<td>(i) Combination of squamoid and mucinous components; (ii) thought to represent papillary carcinoma with extreme degree of squamous and mucinous metaplasia; (iii) low grade thyroid neoplasm.</td>
</tr>
<tr>
<td>Sclerosing mucoepidermoid carcinoma with eosinophilia</td>
<td>(i) Fibrohyalin stroma; (ii) striking infiltration of eosinophils; (iii) mucin secretion is often present; (iv) typically arising in Hashimoto thyroiditis (thought to derive from metastatic squamous nests associated with inflammatory infiltrate).</td>
</tr>
<tr>
<td>CASTLE (carcinoma with thymus-like elements)</td>
<td>(i) Lymphoepithelioma-like carcinoma with foci of squamous differentiation; (ii) pushing margins; (iii) immunoreactivity for CD5, bcl-2, high molecular weight keratin, mcl-1 (thought to be ectopic thymic carcinoma arising from remnants of the branchial pouch); (iv) usually indolent behavior with tendency to late recurrences.</td>
</tr>
</tbody>
</table>

Table 3: Immunohistochemical features of ATCs.

<table>
<thead>
<tr>
<th>Immunostain</th>
<th>Percentage of positive cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cytokeratin</td>
<td>40–100%</td>
</tr>
<tr>
<td>Vimentin</td>
<td>100% (in spindle cells)</td>
</tr>
<tr>
<td>EMA</td>
<td>30–50% (in squamous cells)</td>
</tr>
<tr>
<td>CEA</td>
<td>Rarely (in squamous cells)</td>
</tr>
<tr>
<td>Thyroglobulin</td>
<td>0% (false positivity due to nonneoplastic thyroid follicular cells entrapped in the tumor or diffusion from destroyed normal follicles)</td>
</tr>
<tr>
<td>TTF-1</td>
<td>0%</td>
</tr>
<tr>
<td>RET/PTC oncoprotein</td>
<td>0%</td>
</tr>
<tr>
<td>Calcitonin</td>
<td>0%</td>
</tr>
<tr>
<td>PAX8</td>
<td>0–79% of ATCs (probably depending on antibody used); 92% of ATCs with squamoid features</td>
</tr>
</tbody>
</table>

PAX8 staining of ATCs have had widely disparate results [40, 43, 44]. Nevertheless PAX8 seems to have a useful diagnostic role in specific settings, having been found in 79% of ATCs and in up to 92% of ATCs showing squamoid features, whereas it is negative in head and neck squamous carcinoma and lung carcinoma [40, 45]. Immunohistochemical features are summarized in Table 3.

3. Genetic Features

Even though ATC is a rare disease, a consistent amount of information is currently available on the genetic alterations that are most frequently associated with this tumor [46, 47] (Figure 8).

3.1. Somatic Gene Mutations. Mutations in the components of the principal oncogenic pathways (MAPK, PI3K, Wnt, etc.) have been described to occur with high frequency in ATC. It is known that more than 90% of thyroid cancer harbor mutations in the MAPK pathway [48]. RAS mutations that occur both in benign and malignant thyroid cancers are detected also in ATCs, with variable frequency ranging from 6 to 50% of cases depending on series [49–53]. By contrast, RET/PTC rearrangements, which account for about 15–20% of PTCs, are rarely found in ATCs [47, 52]. Mutations in the BRAF gene, which occur in more than 50% of well-differentiated PTCs [54–58], are only detected in 25% of ATC cases [59, 60]. This lower frequency is in apparent contrast...
Figure 7: An ATC made up of pleomorphic epithelioid cells arranged in loosely cohesive nests; focally, intracytoplasmic vacuoles are present ((a), (b)). Immunohistochemical stains with keratin 7 (c) and TTF1 (d) highlight entrapped thyroid follicles.

with the role of this mutation in driving aggressiveness of thyroid tumors, which has been proposed and largely debated in the past decade [61–63].

3.1.1. PIK3CA. Gain of function mutations in the PIK3CA (phosphatidylinositol-4,5-bisphosphate 3-kinase, catalytic subunit alpha) gene is found in 25–40% of cancers, with alterations mainly clustering in two hotspots within the helical (exon 9) and catalytic (exon 20) domains [64]. In thyroid cancer, PIK3CA mutations are rare in PTCs (0–5% depending on series) but more frequent in poorly differentiated and anaplastic thyroid cancer (from 11 to 23%). As well, amplification of the PIK3CA genomic locus in 3q26.3 is found in about 40% of ATC suggesting that alteration of the PI3K depending pathway plays a pivotal role in the pathogenesis of ATCs [50, 51, 53, 65].

3.1.2. TERT. Somatic mutations in the promoter of the TERT (Telomerase Reverse Transcriptase) gene have been described as highly recurrent in different types of cancer including thyroid cancer [66–69]. Up to 50% of ATCs (33–50%) have been shown to carry these mutations. Intriguingly, TERT promoter mutations seem to occur prevalently in those tumors harboring mutated BRAF or RAS, suggesting that TERT alteration is acquired later during tumor development and may provide a functional advance to BRAF or RAS-driven tumors by enabling acquisition of additional genetic defects leading to disease progression.

3.1.3. CTNNBI. Wnt pathway appears also to play a relevant function in ATC development. Mutations in CTNNBI (β-Catenin) gene leading to a constitutively active Wnt-signaling have been reported in 25–60% of ATCs [70]. CTNNBI is a major component of the E-cadherin cell-cell adhesion complexes and a role of this protein in the epithelial-mesenchymal transition process has been demonstrated [71]. Intriguingly, the transdifferentiation of well-differentiated thyroid tumor cells toward a nondifferentiated status has been proposed as one of the major processes in the pathogenesis of ATCs.

3.1.4. p53 and PTEN. Besides gain of function alterations in key oncogenes, tumor development and progression rely significantly on the inactivation of tumor suppressor genes. p53 and PTEN genes are involved in the negative regulation of cell proliferation and in promoting apoptosis and are frequently impaired during tumor progression. More than 50% of ATCs have been reported to carry loss of function mutations in the p53 gene. As well, the overexpression of p53, which may reflect altered function of the protein in the absence of mutation, has been frequently observed in ATC. Loss of function alterations in the PTEN gene, which inhibit the activation of the PI3K pathway, has been reported to occur in 4 to 16% of ATC [50, 51, 53].

3.2. Somatic Chromosomal Aberration. It is well established that the accumulation of genetic alterations is a driving mechanism of tumor growth and spread to distant sites.
Several studies have investigated genomic instability and DNA copy number variations in ATC with the intent to understand the impact of genomic damage on the genesis and progression of this tumor. Liu and colleagues used real-time PCR analysis to investigate the copy number of a panel of genes involved in MAPK and PI3K pathway in thyroid cancer including a series of 51 ATCs. They observed that genes coding for tyrosine kinase receptors (RTK) like EGFR, PDGFR, VEGFR, KIT, and MET are frequently amplified in thyroid cancer and in particular in the ATC histotype [51]. Wreesman and colleagues used CGH technique to investigate the molecular-cytogenic profile of different histotypes of thyroid cancer to define chromosomal regions that could be specifically associated with the development of ATC [72]. These authors observed several chromosomal abnormalities that were common to both well-differentiated and nondifferentiated thyroid cancer (like gain of 5p15, 5q11–13, 19p, and 19q and loss of 8p) and that could represent early event in the genesis of these tumors. Furthermore, they found alterations like gain in 3p13-14, loss of 5q11–31, and gain in 11q13 that were exclusive of the genome of the 15 ATCs analyzed and that may represent late genetic events driving the transformation of a preexisting thyroid cancer into the aggressive ATC histotype. Using the same approach, Rodrigues et al. investigated the chromosomal profiles of 7 ATCs, showing that chromosomal imbalances affect the genome of all cases analyzed [73]. Intriguingly, the chromosomal regions affected by the alterations were extremely heterogeneous, suggesting the existence of a high-grade genetic interneoplastic diversity in ATCs. Besides
chromosomal imbalances, Miura and colleagues reported that 6 out of 10 ATCs showed aneuploidy [74].

Summarizing the data currently available, two major considerations emerge: (1) the type of chromosomal alterations that characterize ATCs is widely heterogeneous and up to now it is not possible to define a "common" profile of alterations that is specific of this tumor type. This observation implies that different kinds of genetic damage may contribute to the genesis of this tumor type. (2) ATCs are characterized by a higher number of chromosomal alterations with respect to well-differentiated and poorly differentiated thyroid cancer. However, several studies reported that the amount of genetic damage does not directly correlate with the grade of aggressiveness or the outcome of ATCs. Based on these considerations, we may hypothesize that the high-grade genomic instability observed in ATCs is a side effect of the loss of restraining mechanisms of cell proliferation rather than being the cause of tumor progression. Indeed, a number of mitotic proteins involved in cell cycle check points or engaged in chromosome assembly and segregation have been shown to be deranged in ATCs [75, 76]. These include the three members of the Aurora kinase family. Aurora kinases are implicated in several aspects of chromosome segregation and cytokinesis. Expression of all Aurora kinases and in particular of Aurora A is strongly induced in ATC cells [75, 76] and overexpression of Aurora A has been shown to induce centrosome amplification and to potentiate the oncogenic function of Ras [77, 78].

Evidences exist of a negative cross-talk between Aurora A kinase and p53 [79, 80]. Considering the fact that p53 is mutated or aberrantly expressed in a wide proportion of ATCs, it is likely to suppose that these alterations may affect the balance between p53 and Aurora A with relevant consequences on chromosome stability. The possibility to counteract the misfunctioning mitotic proteins has been considered a potential therapeutic strategy for cancer with high grade genetic damage. Indeed, inhibitors of Aurora kinases alone or in combination with other drugs, including microtubule inhibitors, showed an important anticancer effect in preclinical models of ATCs indicating this approach as a possible therapeutic strategy for ATCs treatment [75, 81].

4. Histogenesis

In literature there are indirect, although convincing, evidences that ATC represents a terminal dedifferentiation of preexisting well-differentiated thyroid carcinoma (WDTC) in most, if not all, cases. A large portion of ATC develops in longstanding goiters or in the context of preexisting, incompletely treated papillary or follicular thyroid cancers. Likewise, careful examination of primary ATC tumors reveals coexisting areas of WDTC in 80% to 90% of cases [82, 83]. This better differentiated tumor is usually a papillary carcinoma or one of its variants (particularly Warthin-like and tall cell variant), but it may also be a follicular carcinoma, as well
as an oncocytic carcinoma, or an insular carcinoma [4, 26, 84, 85]. It has been suggested that if an extensive sampling is performed, foci of WDTC are eventually found in every specimen of ATC [83]. Furthermore, it has been postulated that the sharply outlined sclerohyaline nodules sometimes present within undifferentiated carcinoma represent the burn-out residue of such well-differentiated components [86].

Anaplastic transformation may also take place in a metastatic focus, (Figures 9(a), 9(b), and 9(c)) thus supporting the idea that these lesions originate through the dedifferentiation of preexisting well-differentiated cancer [87–89]. Nevertheless, according to the current genetic data it is conceivable that not all the ATCs arise as temporal aggressive evolution of a preexisting WDTC. If the ATC phenotype was always the temporal aggressive evolution of a preexisting WDTC then common founding alterations between the ATCs and the WDTC subgroups should be identified. Whole genome studies showed that the chromosomal asset of ATCs and WDPTC is widely different [51, 72–74] supporting the hypothesis that not all thyroid cancers start as indolent lesions but some of them may originate as already aggressive nondifferentiated cancer.

Conflict of Interests
The authors declare that there is no conflict of interests regarding the publication of this paper.

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