Genomic alterations in thymoma—molecular pathogenesis?

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Abstract: Thymomas and thymic carcinomas (TCs) are neoplasms of thymic epithelial cells. Thymomas exhibit a low mutational burden and a few recurrently mutated genes. The most frequent missense mutation p.(Leu404His) affects the general transcription factor IIi (GTF2I) and is specific for thymic epithelial tumors (TETs). The clinically indolent types A and AB thymomas express the miRNA cluster C19MC. This miRNA cluster known to be the largest in the human genome, is—with expression otherwise restricted mostly to embryonal tissue—silenced in the more aggressive type B thymomas and TCs. Thymomas associated with the autoimmune disease myasthenia gravis (MG) exhibit more frequent gene copy number changes and an increased expression of proteins homologous to molecules that are targets for autoantibodies. TCs, however, display a higher mutational burden, with frequent mutations of TP53 and epigenetic regulatory genes and loss of CDKN2A. The knowledge of molecular alterations in TETs fosters the understanding of their pathogenesis and provides guidance for further studies that may lead to the development of targeted therapies.

Keywords: General transcription factor IIi (GTF2I); epigenetic; thymic epithelial tumor (TET); thymic carcinoma (TC); myasthenia gravis (MG)

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Introduction

Thymomas and thymic carcinomas (TCs) are rare neoplasms derived from thymic epithelial cells. Thymomas are classified as A, AB, B1, B2 and B3 subtypes dependent on cell morphology and relative proportion of neoplastic epithelial cells and admixed non-neoplastic T-lymphocytes (1). Types A and AB thymomas are indolent tumors. On the other hand, B1, B2 and B3 thymomas exhibit an increasing propensity for intrathoracic spread. However, they usually do not seed hematogenous or lymphatic metastases. TCs, in contrast, are aggressive tumors, mostly squamous cell carcinomas, with a tendency for lymphatic and hematogenous dissemination. Thymic neuroendocrine tumors (NETs) comprise only about 5% of thymic epithelial tumors (TETs). They are morphologically classified similarly to lung NETs into carcinoid (typical and atypical), large cell neuroendocrine carcinoma and small cell carcinoma (1).

The causes for the development of TETs are unknown. There are no identified exogenous factors initiating or promoting the development of TETs. Furthermore, no family clustering has been observed, except for an increased prevalence among Asians/Pacific Islanders (2). The TETs are predominantly a disease of adulthood and exceedingly uncommon in children and adolescents (3). The incidence peaks in the seventh decade of life. This pattern is similar to many other tumors and most likely reflects the accumulation of genetic damage with age.

Thymomas may manifest concurrently with autoimmune diseases, most frequently myasthenia gravis (MG). Ten to 20% of MG patients develop a thymoma and about 30% of thymomas are associated with MG (4,5). Other thymic pathologies associated with MG are follicular thymitis
and, in rare cases, thymolipoma (5). However, the thymus frequently appears histologically normal for age. TCs, in contrast, are not linked with MG or other specific autoimmune diseases, although paraneoplastic autoimmune symptoms may develop on rare occasions.

The genetic alterations present in TETs have been more deeply elucidated within the last 5 years. Similar to other tumor entities this has been feasible largely through the advent of next generation sequencing technologies. The results of these studies show a different pattern of molecular aberrations in thymomas and TCs, with only a few significantly and recurrently mutated genes. The perhaps most striking finding in thymomas is a very frequent point mutation, p.(Leu404His), in the general transcription factor IIi (GTF2I) gene, that so far has not been detected in other tumor entities (6). A further genetic hallmark is the mutation of epigenetic regulatory genes in TCs. The association of thymomas with MG has been linked to an increased gene copy number variation, and gene expression profiling identified overexpressed autoantigens (7). Unfortunately, the molecular profiling efforts have to date added only a few new therapeutic targets for currently available and approved drugs. Nevertheless, the novel insights into the genetic drivers of TETs form a strong basis for further studies on the biology and treatment of these tumors.

**DNA alterations in TETs**

**GTF2I is a master genetic regulator in thymomas**

The most frequently mutated gene in thymomas is GTF2I. It is mutated in 76–83% of types A and AB thymomas (6-8), less often in types B1, B2 and B3 subtypes and only in 8% of TCs (6,8). TETs with a GTF2I mutation exhibit a less aggressive clinical course than TETs with wild-type GTF2I, with 10 year-survival-rates of 96% and 70%, respectively. This is most likely caused by the higher GTF2I mutation prevalence in the indolent types A and AB thymomas (6).

The GTF2I gene recurrently carries the same point mutation on chromosome 7 at position 74146970T>A in TETs. The mutation leads to an exchange of the amino acid leucine by histidine on the protein level (p.Leu404His). The GTF2I gene codes for the transcription factor TFII-I, which regulates the expression of genes that influence the cell cycle (Figure 1) (9). The activation of TFII-I protein can be triggered by extracellular signals, such as the stimulation of B and T cell receptors and growth factor receptors. In mice the GTF2I gene is indispensable, because a knock-out of the gene causes death at an early embryonic stage (10).

The p.(Leu404His) mutation changes an amino acid in a sequence segment that may function as a protein degradation mark (6). The mutation may thus increase the half-life of TFII-I and thereby support its intracellular accumulation (6,9). This assumption is fostered by the observation that GTF2I mutated TETs exhibit an increase of TFII-I protein in comparison with unmutated TETs, but without concomitant augmentation of mRNA expression (6). The transfection of a p.(Leu404His) mutated GTF2I gene into mouse NIH-3T3 cells increased their proliferation. However, in a soft-agar tumorigenicity assay, no difference between normal or mutated GTF2I transduced cells in their colony forming ability was observed (6). Therefore, mutant GTF2I facilitates cell proliferation but does not exert a strong oncogenic effect.

The GTF2I p.(Leu404His) mutation is characteristic for TETs. For instance, it has not been observed in approximately 10,000 tumors of other entities that have been sequenced by the TCGA (7). However, 6% of angioimmunoblastic T cell lymphomas and less than one percent of other non-thymic tumor entities bear GTF2I missense mutations at other locations (11). Apart from that, a GTF2I-NCOA2 and a GTF2I-RARA gene fusion have been detected in an angiofibroma and in an acute promyelocytic leukemia, respectively (12,13). The GTF2I gene is located on chromosome 7 and the deletion of a region that spans GTF2I is present in patients with Williams-Beuren syndrome and a duplication has been linked with the Somerville-van der Aa syndrome (14). Both diseases exhibit malformations, mental retardation and disturbed behavior. Genome wide DNA sequence polymorphism studies suggested that certain GTF2I sequence variants contribute to the pathogenesis of autoimmune diseases such as rheumatoid arthritis, Sjögren’s syndrome and systemic lupus erythematosus (15-17). However, in thymomas no association of the GTF2I mutation with autoimmune diseases has been
The recognition of the GFT2I p.(Leu404His) mutation as the most prominent genetic alteration in TETs helps to understand their pathogenesis. However, at present the GFT2I mutation cannot be therapeutically targeted. Even so, in clinical practice the identification of a GFT2I mutation may occasionally help pathologists to derive a diagnosis in small biopsies that are difficult to assess.

Irrespective of the GFT2I mutation, overall TETs have a low tumor mutational burden (TMB) with an average of 0.48 mutations per megabase. This is the lowest TMB among 21 other cancers that have been sequenced by The Cancer Genome Atlas (TCGA) (7). In addition to GTF2I only a few other genes were recurrently mutated in TETs at a frequency of at least 3% in a cohort of 155 reported cases (Table 1) (6,7,18,19). These genes were HRAS, TP53, CYLD, PCLO and HDAC4. HRAS mutations affected types A and AB thymomas in ten of eleven mutated TETs. Thus, HRAS is the second most frequently mutated gene in thymomas. TP53 and CYLD mutations do occur in thymomas but are more frequent in TCs (20,21). Mutations in the piccolo presynaptic cytomatrix gene PCLO and the chromatin modifier gene HDAC4 are observed in different thymoma subtypes and TCs with low frequencies and with no significant differences in incidence rates, although the number of published samples is too small to be able to draw firm conclusions.

**TP53 and epigenetic modifier gene mutations are hallmarks of TCs**

TCs exhibit a higher mutation rate than thymomas (20). The most frequently mutated gene is TP53. It has been altered in 26% of cases by mutation and frequently (32%) by hetero- or homozygous deletion as assessed by FISH (20,21). The presence of mutant TP53 has been associated with poorer overall survival in TC in some studies but this has not been confirmed by others (20-22).

A further genetic hallmark of TCs are recurrent mutations in chromatin modifying and epigenetic regulatory

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**Table 1** Gene alterations in thymic epithelial tumors (TETs) with a frequency >1.5%. The data are retrieved from the cBioPortal for Cancer Genomics (18,19) and encompass the study cohorts of (6,7)

<table>
<thead>
<tr>
<th>Gene</th>
<th>TET (n=155*), %</th>
<th>WHO subtypes, %</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Type A (n=24)</td>
<td>Type AB (n=39)</td>
</tr>
<tr>
<td>GTF2I</td>
<td>43.0</td>
<td>87.5</td>
</tr>
<tr>
<td>HRAS</td>
<td>7.0</td>
<td>29.2</td>
</tr>
<tr>
<td>TP53</td>
<td>5.0</td>
<td>4.2</td>
</tr>
<tr>
<td>CYLD</td>
<td>3.0</td>
<td>–</td>
</tr>
<tr>
<td>PCLO</td>
<td>3.0</td>
<td>4.2</td>
</tr>
<tr>
<td>HDAC4</td>
<td>3.0</td>
<td>4.2</td>
</tr>
<tr>
<td>BCO1</td>
<td>2.6</td>
<td>4.2</td>
</tr>
<tr>
<td>PBRM1</td>
<td>2.6</td>
<td>–</td>
</tr>
<tr>
<td>BRD4</td>
<td>2.6</td>
<td>–</td>
</tr>
<tr>
<td>CSF1R</td>
<td>1.9</td>
<td>4.2</td>
</tr>
<tr>
<td>FGF3</td>
<td>1.9</td>
<td>4.2</td>
</tr>
<tr>
<td>NRAS</td>
<td>1.9</td>
<td>4.2</td>
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<tr>
<td>PAX7</td>
<td>1.9</td>
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</tr>
<tr>
<td>PTTPR7</td>
<td>1.9</td>
<td>4.2</td>
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<tr>
<td>ZMYM3</td>
<td>1.9</td>
<td>–</td>
</tr>
<tr>
<td>NOD1</td>
<td>1.9</td>
<td>4.2</td>
</tr>
</tbody>
</table>

*1 TET sample not otherwise specified; TC, thymic carcinoma.
genes, particularly ASXL1, BAP1, DNMT3, SETD2, SMARCA4, TET2 and WT1 (20). ASXL1 is a polycomb chromatin binding protein, the tumor suppressor BAP1 encodes a histone deubiquitinase, DNMT3A represents a cytosine methyl transferase and SETD2 functions as a H3K36 trimethyl transferase. Furthermore, the SMARCA4 protein represents an ATPase of the SWI-SNF complex, TET2 functions as a methylcytosine dioxygenase and WT1 represents a positive regulator of DNMT3A and SMARCA4. The survival of TC patients with and without epigenetic regulatory gene mutations is similar (20). Another recurrently mutated gene in approximately 11% of TC is CYLD deubiquitinase, which functions as a negative regulator of the NFKB pathway (6,20).

The tyrosine kinase KIT is expressed by 80–86% of thymic squamous carcinomas, but infrequently by non-TCs (23,24). It therefore represents a useful immunohistochemical marker in the histopathologic diagnosis of TC (25). KIT mutations, however, affect only 6–9% of TC, and only in these mutated cases KIT constitutes a potential target for treatment with tyrosine kinase inhibitors (TKIs) (20,21).

**Gene copy number changes in TETs**

Gene copy number aberrations (CNAs) are rare in A and AB thymomas but enriched in B2 and B3 thymomas and TCs (6,7,26). If present, most are large-scale, chromosome arm level CNAs (6,7,26). Deletions involve most frequently the chromosome arms 3p, 6p, 6q, 13q, 16q and 17p (6,7,26,27). Gains of chromosomal material are most common for 1q, 17q and 18 (7,26). In TC the loss of chromosome 16q is most prominent and reported for 67% of cases (7,26). Although no single gene on 16q that is solely crucial for TC pathogenesis has been pinpointed, several candidate tumor suppressor genes are mapped to this chromosome arm, namely CBFB, CDH1, CDH11, CTGF, CYLD and ZFHX3.

A further frequently mutated and/or deleted gene in TC is the tumor suppressor CDKN2A located at 9p21.3. CDKN2A mutations, mostly truncating, have been reported for 11% of TC and a hetero- or homozygous deletion has been observed in 38% of cases in a cohort of 35 TC (21,27). The CDKN2A gene codes for p16\(^{INKA}\) and p14\(^{ARF}\). The two proteins are generated by alternative RNA splicing. The p16\(^{INKA}\) inhibits mitosis by antagonizing cyclin dependent kinases 4 and 6. On the other hand, p14\(^{ARF}\) activates the anti-oncogene TP53. Therefore, a CDKN2A deletion or inactivation by mutation may result in a stimulation of cyclin dependent kinases with subsequent promotion of cell proliferation.

**Genetic aberrations in neuroendocrine thymic tumors**

The WHO classification of thymic NETs is similar to pulmonary NETs and distinguishes carcinoid (typical and atypical), large cell neuroendocrine carcinoma and small cell carcinoma (1,28). So far little is known about the genetic aberrations in thymic NETs, which is likely due to the rarity of these tumors. However, some 25% of thymic carcinoids are associated with multiple endocrine neoplasia-1 (MEN1), a hereditary tumor syndrome that is caused by mutation of the MEN1 gene. In contrast, only 2–8% of patients with MEN1 develop a thymic carcinoid.

A recent study analyzed 63 thymic NETs by low coverage whole genome sequencing to determine gene copy number instabilities (CNI) (29). The report differentiated three groups of thymic NETs with low, intermediate and high CNI. The CNI low and intermediate tumors encompassed in addition to carcinoids also some large cell neuroendocrine carcinomas. On the other hand, the group with high CNI reflected all small cell neuroendocrine carcinomas but contained also a few atypical carcinoids. Therefore, histopathology failed to faithfully predict the CNI grouping. The authors of the study have thus concluded that thymic NETs separate into three genetic clusters which are not satisfactorily recognized by the WHO classification. A morphomolecular grading system for thymic NETs was proposed instead of a mere histologic classification for patient stratification and prognostication (29). However, such a grading system would be difficult to implement in clinical practice, because of the requirement of low coverage whole genome sequencing and assay standardization among diagnostic laboratories.

**The transcriptome of TETs**

**RNA expression profiles mirror the histopathologic classification of TETs**

RNA sequencing (RNA-seq) of 12 thymomas and one TC revealed gene transcription profiles that correlated well with the WHO histopathologic subtyping (30). A similar RNA-seq analysis of a larger cohort of 117 TETs within the TCGA project that integrated RNA-seq, gene copy number
variation, DNA methylation, and protein array findings revealed also a strong correlation of gene expression profiles with histologic subtyping (7). The study differentiated four molecular subgroups. Group 1 were mainly type B thymomas, group 2 TCs, group 3 primarily type AB thymomas and group 4 constituted types A and AB thymomas. RNA-seq detected only a few gene fusions in TETs and none of them was recurrent or involved the GTF2I gene (6,7). Lee et al. evaluated in silico the TET data of TCGA and suggested also the separation of TETs into four groups, although with different molecular features defined by GTF2I mutation, a normal GTF2I phenotype with expression of T cell signaling genes, and TETs with chromosomal stability or chromosomal instability (31). Obviously, the genetic TET grouping by Lee et al. cannot be predicted by histology, reminiscent to molecular subtyping efforts in other tumors, e.g. bladder cancer (31,32).

The C19MC miRNA cluster characterizes A and AB thymomas

The C19MC miRNA cluster is strongly expressed in types A and AB thymomas, but virtually absent in type B thymomas and TCs (21,30). It is a large imprinted miRNA cluster on chromosome 19q13.42 that is normally expressed only during embryonic development and is present in the placenta but silenced in adult tissues. An exception are medullary and cortical thymic epithelial cells which do express C19MC (33). Therefore, the expression of C19MC in A and AB thymomas may reflect a physiological state, and expression is likely silenced in type B thymomas and TCs by a mechanism that is currently not known, but might involve promoter methylation (34). In types A and AB thymomas the miRNAs of the C19MC cluster may activate the PI3K/AKT pathway as proposed by Radovich et al. (30).

The C19MC miRNA cluster is expressed also in a few non-thymic malignancies. In the rare and aggressive brain tumors ETANTRs (embryonal tumors with abundant neuropil and true rosettes) and ETMRs (embryonal tumors with multilayered rosettes) the C19MC cluster is amplified or fused with the TTYH1 gene promoter, which results in overexpression (35,36). Additionally, the C19MC cluster is expressed in adenomas of the thyroid and parathyroid, tamoxifen non-responsive breast tumors and liver cell carcinomas and mesenchymal hamartomas. The mechanism of overexpression in these tumors are variable and encompass gene amplification, chromosomal translocation, and hypomethylation (37-40).

The expression of a further miRNA cluster, called C14MC and located on chromosome 14q32, is also reduced in TCs, albeit not all miRNAs of the cluster are silenced (21). The C14MC cluster has been regarded to possess a tumor suppressive function in GIST and glioma (41,42). The downregulation of anti-oncogenic C14MC miRNAs might thus similarly promote the genesis of TCs.

The reports on non-clustered miRNA expression in TETs are more heterogenous and therefore difficult to integrate into a common pattern (21,43,44). This may be caused by the use of different techniques (microarrays, next generation sequencing/NGS) and varied TET subtype composition and size of the study cohorts. Enkner et al. observed in association with NGS the upregulation of mainly oncogenic miR-21, miR-9-3 and miR-375 and the reduction of putative tumor suppressive miR-34b, miR-34c, miR-130a and miR-195 in TCs as compared to type A thymomas (21). Notably, in non-small cell lung cancer TP53 downregulates PD-L1 via miR-34, and functional loss of TP53 and consecutive silencing of miR-34 increases PD-L1 and thereby supports immune evasion (45). TCs express PD-L1 more frequently than type A thymomas, and this might likewise be caused by a decrease of miR-34.

Gaeci et al. compared the miRNA expression of 54 TETs with normal thymic tissue by microarray hybridization and identified 87 differently expressed miRNAs (43). Similar to the observation by Enkner et al. an upregulation of oncogenic miR-21-5p was recognized among the most significantly deregulated miRNAs. In addition, the silencing of the tumor suppressive miR-145-5p was found to be significant, and an epigenetic regulation of miR-145-5p expression, previously demonstrated in other tumor entities, was revealed in TETs as well (43,44).

The molecular pathogenesis of MG

About 30% of thymoma patients exhibit an autoimmune disorder, in particular MG. The autoantibodies are directed against muscle specific antigens, mainly the acetylcholine receptor (AChR) in the postsynaptic muscle membrane (4,46). Localized or general muscle weakness is the predominant symptom. Although the disease-inducing antibodies have been well characterized, the pathogenetic mechanisms triggering the autoimmune disease are less understood. However, similar to other autoimmune diseases an association of MG with certain major histocompatibility complex (MHC)—classes has been observed (4).
The recent genomic characterization of thymomas by the TCGA have revealed that thymomas with MG exhibit a higher prevalence of gene copy aberrations than thymomas in the absence of MG (7). However, which of the deletions and gains of chromosomal material are crucial for the development of MG could not be clarified, because none of the gene CNAs was significantly accumulated in the thymomas with MG. Furthermore, MG did not segregate with any gene mutation and DNA methylation or gene expression pattern (7). Surprisingly, thymomas associated with MG do not express complete prototypical target molecules of autoantibodies such as AChR, titin, and muscle ryanodine proteins 1 and 2. Expression of the AChR alpha subunit was only 3.0-fold higher in 32 thymomas with MG than in 72 tumors without MG. By contrast, expression of NEFM (neurofilament medium) which shares sequence similarities with AChR, titin, and ryanodine receptors was 23.8-fold higher in MG positive compared with MG negative thymomas (7). Furthermore, the neuronal ryanodine receptor 3, which shares homology with muscle ryanodine receptors 1 and 2, was 5.5-fold upregulated in MG positive thymomas. In summary, MG in thymoma concurs with an increased intratumoral expression of genes that share sequence homologies with autoantibody targets.

Targeted therapies in TETs

Surgical resection is the primary therapy for TETs. In addition, for unresectable or incompletely resected tumors, chemo- and radiotherapy are used (47). At present no targeted therapy for TETs is firmly established. This is brought about by the low number of recurrent and actionable mutations and the rarity of TETs, which makes clinical trials difficult. Nevertheless, activating mutations in the KIT oncogene, which are present in approximately 5% of TCs, have been explored as a therapeutic target for TKIs with encouraging results in a few patients (48,49). However, more informative clinical studies with a larger number of KIT mutated patients are needed. Approximately 80% to 86% of TCs express KIT protein, but this does not correlate with their response to TKIs (25,47). Therefore, only TCs with an activating KIT mutation are potential candidates for a TKI therapy.

The genomic profiling of TETs revealed additional candidate molecular targets (6,7,20,21). Amongst them the miRNA cluster C19MC, which is a unique feature of types A and AB thymomas (21,30). The cluster activates the PI3K pathway which might be effectively blocked in the rare incidences of recurrent types A and AB thymomas. The common mutation of genes that modify the epigenetic signatures of DNA and chromatin in TCs could offer an additional chance for therapeutic intervention (20).

Immune checkpoint inhibitors may also evolve as a therapeutic option in clinically advanced TETs. PD-L1 is frequently expressed by tumor cells, particularly in B3 thymomas and TCs (21,50-53). In 40 patients with recurrent TCs who had progressed after at least one line of chemotherapy the immune-checkpoint inhibitor pembrolizumab achieved complete or partial response in 22.5% and stable disease in 53% of patients (54). Another phase II clinical trial with pembrolizumab enrolled 26 TC and seven thymoma patients (55). Five (19.2%) of the TCs exhibited a partial response and 14 (53.8%) had stable disease. Of the seven thymoma patients, two achieved partial response and five stable disease. In both studies high PD-L1 protein expression was associated with better response to pembrolizumab. Although the observed responses of TETs to immune checkpoint inhibitor therapy are promising, the high incidence rate of severe autoimmune adverse advents that has been observed is of major concern, in particular in the thymoma patients (54,55).

Conclusions

In summary, exemplary prominent insights derived from molecular profiling are the identification of GTF2I as an important gene for the development of TETs, the discovery of frequent epigenetic regulatory gene mutations in TCs and the recognition of the large miRNA cluster C19MC in types A and AB thymomas. The translation of the molecular insights into a clinical benefit for TET patients will need the development of novel targeted medicines that address different molecules and the organization of multi-center trials, because of the diversity and infrequency of recurrent genetic alterations and the rareness of TETs.

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Footnote

Conflicts of Interest: The authors have no conflicts of interest to declare.
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