Identification and Characterization of Potential Novel Targets in Thyroid Carcinoma: Evidence of Non-Oncogene Addiction Unveiling Tumor Cell Vulnerabilities

Thesis

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IDENTIFICATION AND CHARACTERIZATION OF POTENTIAL NOVEL TARGETS IN THYROID CARCINOMA: EVIDENCE OF NON-ONCOGENE ADDICTION UNVEILING TUMOR CELL VULNERABILITIES

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The Open University, Milton Keynes (UK)
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Abstract

Thyroid cancer is the most frequent endocrine malignancy, with an incidence constantly increasing. Despite well-differentiated thyroid tumors are generally cured by conventional therapy, a fraction of them relapses and progresses towards undifferentiated forms, characterized by a poor prognosis. Target therapies introduced in clinical testing are often unsuccessful; therefore, novel therapeutic strategies are needed. To identify critical nodal points for therapeutic intervention, our laboratory faced the “non-oncogene addiction” (NOA) paradigm, which asserts that the tumorigenic state relies on the activity of genes that are essential to support the phenotype of cancer cells but not required to the same degree for normal cell viability. By screening a siRNA library on normal and tumor thyroid cell lines, we identified 15 genes whose silencing interfered with the growth of tumor but not normal thyroid cells.

The overall aim of this project was to validate NOA genes to identify thyroid tumor cell vulnerabilities and to explore their role in the regulation of thyroid tumor cell biology. Among the 15 hit genes, we selected MASTL, Cyclin D1 and COPZ1 for validation studies. We confirmed the growth inhibitory effect of their silencing in tumor but not normal thyroid cells and observed that these effects were common to a panel of thyroid tumor cell lines, irrespective of the histotype or genetic lesion. Functional studies on MASTL demonstrated that its depletion in thyroid tumor cells inhibited cell growth, led to mitotic catastrophe and induced DNA damage and cell death. COPZ1 depletion induced abortive autophagy, endoplasmic reticulum stress and apoptotic cell death in thyroid tumor cells, as well as tumor growth inhibition in vivo. Together, our studies identified MASTL and
COPZ1 as vulnerability genes for thyroid tumor cells and provided the rationale for future studies aimed to explore their targeting for potential therapeutic intervention.
1. Introduction

1.1. Thyroid cancer

Thyroid cancer (TC) is the most frequent malignancy of the endocrine system, and notably it is characterized by a global incidence increasing during the last decades. In the United States of America, TC is the fifth most common cancer in women, and accounts for more than 62000 cases occurred in men and women in 2015 (1). Thyroid is a butterfly-shaped organ located on the anterior surface of trachea, at the base of the neck, and it is formed by aggregates of lobules of spherical follicles filled with colloid (2). Epithelial follicular cells mark follicles and are responsible for the production and secretion of thyroid hormones thyroxine (T4) and triiodothyronine (T3), while parafollicular “C” cells, located within follicles, are the neuroendocrine cells responsible for calcitonin secretion (Figure 1.1). Thyroid tumors can originate from both follicular epithelial cells and parafollicular “C” cells. Tumors originating from the follicular cells are distinguished into different subtypes: papillary thyroid carcinoma (PTC, accounting for 80-85% of cases), follicular thyroid carcinoma (FTC, 10-15% of cases), poorly differentiated thyroid carcinoma (PDTC, about 5% of cases) and anaplastic thyroid carcinoma (ATC, 2-3% of cases) (3). Tumors originating from parafollicular C cells are known as medullary thyroid carcinoma (MTC) and account for a small proportion of thyroid malignancies (2-3% of cases).
Explanations for the worldwide increase of thyroid tumor incidence are controversial. Fine-needle aspiration biopsy (FNAB) and histopathological assessment are the cornerstone of diagnostic management of thyroid nodules; however, in the last years a progressive increase in the use of more sensitive techniques, that are high-resolution ultrasonography, computerized tomography, magnetic resonance or positron emission tomography, as diagnostic tools, was observed (4). According to some experts, the increased diagnosis of thyroid tumors is related to the increase in diagnostic sensitivity. Conversely, others attribute the increase of incidence rate to a major exposure to risk factors, like radiation exposure during medical procedures (5), obesity (6), chronic autoimmune thyroiditis (7), or environmental factors (8).

However, in contrast to incidence rates, mortality rates are stable or decreasing both in the United States (9) and in Europe (10), supporting the evidence that probably the increased incidence is mainly related to improvement of surveillance (11). On the other
hand, the problem of probable overdiagnosis of thyroid tumors has emerged as a growing public health concern (12).

1.1.1. Follicular cell-derived thyroid tumors

The majority of thyroid malignancies originates from follicular cells and is distinguished in different subgroups, according to their differentiation status, as well as to some histological and clinical features (Figure 1.2 and Table 1.1). PTC and FTC are classified together as well-differentiated thyroid carcinomas (WDTC) and are generally characterized by a good prognosis, with over 98% five-year survival rate in Europe and North America (13). By contrast, patients with ATC have a poor prognosis, with median survival of 5 months, and less than 20% are alive 1 year after diagnosis (14).

1.1.1.1. Papillary thyroid carcinoma (PTC)

PTC is the most common tumor histotype, occurring more frequently in women than in men, and is also the most common pediatric malignancy. PTC presents as an enlarged solid nodule in more than half of patients and often forms multifocal tumors that can be either unilateral or extended to both thyroid lobes (15). Lymphatic vessels are the main system for PTC diffusion; indeed, cervical lymph node metastases are clinically detectable on initial presentation.

“Classical” PTC is microscopically characterized by a papillary architecture and epithelial cells with “ground glass” nuclei, which include irregularities as grooves and nuclear pseudoinclusions (16). Psammoma bodies, which represent a typical PTC feature, are found in at least 50% of cases, are formed by focal areas of infarction of the tips of papillae and can be found also in tumor stroma and lymphatic spaces. Even if regional
lymph node metastases are common at initial presentation of usual PTC, this feature apparently does not affect long-term prognosis that is generally good for PTC patients.

In addition to classical PTC, which is the most frequent, several histological variants differing in morphological pattern as well as in prognosis, have been described. The most common are papillary thyroid microcarcinoma, follicular, oncocytic, clear cells, diffuse sclerosing, tall cell, columnar cell, solid and cribriform variants (17). These variants have good prognosis, similar to that of classical PTC, with the exception of tall cell, columnar cell, and solid variants, characterized by aggressive behavior, vascular invasion and worse outcome.

1.1.1.2. *Follicular thyroid carcinoma (FTC)*

FTC is characterized by evidence of follicular cell differentiation lacking the diagnostic nuclear features of PTC. It is more frequent in women over 50 years of age (18) and in iodine-deficient areas (19). FTC is an encapsulated tumor and generally presents as unifocal. Unlike PTC, it spreads hematogenously rather than by lymphatic vessels, and distant metastases, primarily to the lung and bone, can be found at the diagnosis (20). FTC can originate from follicular thyroid adenoma (FTA), which is a benign tumor of thyroid gland surrounded by a thin fibrous capsule (21). Indeed, FTAs can acquire oncogenic mutations and progress towards the malignant FTC form.

Hürtle cell carcinoma (HCC), or “oncocytic or oxyphilic”, is considered a rare variant of FTC (and less frequently of PTC) (22). The tumor is composed of at least 75% Hürtle cells, that are large polygonal cells characterized by an abundant eosinophilic granular cytoplasm, due to a high content of mitochondria. As FTCs, HCCs can spread hematogenously to metastasize to bone and lungs, but can also present regional lymph node involvement.
FTC is generally considered a more aggressive disease than PTC, with a prognosis depending on several factors, among which the degree of invasiveness of the tumor: patients with minimally invasive FTC have a survival expectance similar to that of normal population matched for age and sex, while 50% of patients with widely invasive carcinoma die of the disease (18).

1.1.1.3. Poorly Differentiated Thyroid Carcinoma (PDTC)

PDTCs comprise thyroid tumors showing histological and prognostic features that are intermediate between those of differentiated (PTC and FTC) on one hand and anaplastic (ATC) carcinomas on the other. A consensus meeting held in Turin established the diagnostic criteria for PDTC assessment, which include solid/trabecular/insular microscopic growth pattern, lack of well-developed nuclear features of PTCs, convoluted nuclei, tumor necrosis and high mitotic index (23). PDTC can either be the terminal stage of dedifferentiation process of PTC and FTC (Figure 1.2) or grow ex novo (19). It often presents as a unique thyroid mass rapidly growing, and has high rate of metastatic spread to lymph nodes and distant organs, such as lung and bone. Prognosis of PDTC patients is worse than that of PTC, with a 5 year survival rate from 20 to 60% (24).

1.1.1.4. Anaplastic Thyroid Carcinoma (ATC)

ATC is one of the most aggressive and deadly cancers among humans (25). Like PDTCs, it can develop from a pre-existing DTC or can arise ex novo (Figure 1.2). ATC is composed of undifferentiated cells (spindle, epithelioid and pleomorphic giant cells) retaining markers of an epithelial lineage. Histological features include invasiveness, extensive tumor necrosis, pleomorphic nuclei with irregular contours and high mitotic activity with atypical mitotic figures. ATC presents as a large rapidly growing thyroid mass infiltrating surrounding tissues that causes compressive symptoms in patients including dysphagia,
dysphonia, and dyspnea. The peak incidence is in the sixth and seventh decades of life, and the median survival is 6 months from the diagnosis (26). About 50% of patients present distant metastatic disease at the diagnosis, primarily in the lungs but also in bones and brain, and approximately 25% of patients will develop metastases during the clinical course of the disease: it can be so rapid that surgery procedures are often useless.

Figure 1.2 Multistep process of thyroid carcinogenesis
Scheme of the development and dedifferentiation of follicular-cell derived thyroid tumors. Tumor frequencies and molecular alterations that drive the multistep process are indicated. Image adapted from (27).
Table 1.1 Characteristics of thyroid tumors
FTA, follicular thyroid adenoma; PTC, papillary thyroid carcinoma; FTC, follicular thyroid carcinoma; PDTC, poorly differentiated thyroid carcinoma; ATC, anaplastic thyroid carcinoma; *PTC and FTC are usually referred together as well-differentiated carcinomas (WDTC). Table adapted from (3).

<table>
<thead>
<tr>
<th>Tumour type</th>
<th>Cell of origin</th>
<th>Prevalence (% of thyroid cancers)</th>
<th>Standard care and prognosis</th>
<th>Characteristics</th>
</tr>
</thead>
<tbody>
<tr>
<td>FTA</td>
<td>Follicular thyroid cells (which produce thyroid hormone and thyroglobulin)</td>
<td>This is a benign lesion</td>
<td>Conservative monitoring; thyroidectomy if symptomatic</td>
<td>Common benign thyroid tumour; similar architecture to FTC, but typically encapsulated; lacking capsular or vascular invasion; lacking metastasis; lacking nuclear features of PTC</td>
</tr>
<tr>
<td>PTC*</td>
<td>Follicular thyroid cells</td>
<td>80–85</td>
<td>Thyroidectomy and, in selected cases, radioiodine ablation (novel drugs for resistant cases); good overall prognosis</td>
<td>Well differentiated, with papillary architecture and characteristic nuclear features that include enlargement, oval shape, elongation, overlapping and clearing, inclusions and grooves; propensity for lymphatic metastasis; PTC subtypes include conventional PTC (CPTC), follicular-variant PTC (FVPTC), tall-cell PTC (TCP/TCTC) and a few rare variants</td>
</tr>
<tr>
<td>FTC*</td>
<td>Follicular thyroid cells</td>
<td>10–15</td>
<td>Thyroidectomy and radioiodine ablation (novel drugs for resistant cases); good overall prognosis</td>
<td>Well differentiated, hypercellular, microfollicular patterns; lacking nuclear features of PTC; vascular or capsular invasion; propensity for metastasis via the blood stream; Hürthle cell thyroid cancer is a unique subtype of FTC that accounts for 2–3% of thyroid cancers and is characterized by large, mitochondria-rich oncocytes and dense nuclei and nucleoli, as well as a high propensity for metastasis and a poor prognosis</td>
</tr>
<tr>
<td>PDTC</td>
<td>Follicular thyroid cells</td>
<td>5–10</td>
<td>Surgery, radioiodine (in selected cases), chemotherapy, radiotherapy, novel drugs; poor prognosis</td>
<td>Poorly differentiated, often overlapping with PTC and FTC; intermediate aggressiveness between differentiated and undifferentiated thyroid cancers</td>
</tr>
<tr>
<td>ATC</td>
<td>Follicular thyroid cells</td>
<td>2–3</td>
<td>Surgery, chemotherapy, radiotherapy, novel drugs, palliative care; highly and rapidly lethal</td>
<td>Undifferentiated; admixture of spindle, pleomorphic giant and epithelioid cells; extremely invasive and metastatic; highly lethal; may occur de novo or derive from FTC, PTC or PDTC</td>
</tr>
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1.1.2. Genetic alterations in thyroid cancer

In the last decades much effort has been made in studies aimed to identify molecular alterations at the basis of thyroid tumorigenesis (3). Nowadays, it is well known that thyroid cancer originates from various genetic and epigenetic alterations that include: activating/inactivating somatic mutations, alterations in gene expression patterns, microRNA deregulation, aberrant gene methylation; these lesions gradually accumulate to promote tumor progression. The main genetic lesions found in TC consist of point mutation, which is the result of a single nucleotide change within DNA chain, and chromosomal rearrangement, that is a genetic abnormality with breakage and fusion of parts of the same or different chromosomes. Among the different TC histotypes, PTCs have one of the lowest mutation densities of cancers that have been studied by means of whole-exome sequencing (28), while undifferentiated ATCs are characterized by a high
mutation burden (29). To date, it is well known that the main genetic alterations in TC are found in members of the mitogen-activated protein kinase (MAPK) and phosphatidylinositol 3-kinase/protein kinase B (PI3K/AKT) pathways (Figure 1.3).

![Figure 1.3 Main signaling pathways involved in thyroid carcinogenesis](image)

**Figure 1.3 Main signaling pathways involved in thyroid carcinogenesis**
The main genetic alterations that drive thyroid tumor formation and progression are found in members of the MAPK and PI3K/AKT signaling pathways. These signaling cascades have the important role of propagating signals from external stimuli, via cell membrane receptor tyrosine kinases, into the nucleus, to regulate cell proliferation, differentiation and survival processes. Image adapted from (27).

1.1.2.1. **Next-generation sequencing (NGS) revolution**

The introduction of next-generation sequencing (NGS) technologies improved, in the recent years, the understanding of alterations of genes and pathways involved in human carcinogenesis and tumor maintenance (30). NGS is a high-throughput method that allows a massive parallel sequencing of multiple targeted genomic regions, thus being more advantageous than the earlier genome-sequencing methods. It has been recently
applied also on thyroid cancer, with the aim of shedding light on the molecular alterations that drive thyroid tumorigenesis (31,32). In 2014, a study by the TCGA consortium had revolutionized the understanding of PTC genomic landscape (33). A comprehensive multiplatform analysis on a cohort of 496 PTC specimens, which was the largest cohort ever analyzed to date, was performed through next-generation DNA and RNA sequencing, copy-number analysis, DNA methylation and proteomic assays; as result, driving oncogenic lesions already known were confirmed, and novel driver mutations and gene fusions were detected, thus reducing from 25% to 3.5% the so-called “dark matter”, that is the fraction of PTC cases with unknown genetic drivers. The study also proposed a reclassification of PTC into molecular subtypes, as tool for better management of the disease.

These findings paved the way for following NGS studies performed on FTC (34), PDTC (29) and ATC (29,35), with the aim of improving the knowledge of the genetic landscape thyroid tumors.

1.1.2.2. MAPK signaling pathway

The MAPK signaling pathway plays an important role in the regulation of cell growth and survival and also in human tumorigenic processes. In TC, the importance of this pathway has been well established, as many mutations in members belonging to this signaling cascade were identified (3).

- **BRAF**

  *BRAF (v-Raf murine sarcoma viral oncogene homolog B1)* is a proto-oncogene, located on human chromosome 7p24, that encodes a cytosolic serine/threonine protein kinase expressed in most human cells, including thyroid follicular cells (2). BRAF plays a role in the MAPK pathway signal transduction, that is a crucial axis in the regulation of gene expression, cell proliferation and differentiation. Upon RAS activation, wild-type (wt)
BRAF is recruited to the plasma membrane and activated. Activated BRAF triggers sequential phosphorylation of its downstream target, that is mitogen-activated protein kinase kinase (MEK), which, in turn, activates the extracellular signal-regulated kinase (ERK), thus stimulating consequent effectors of the MAPK cascade.

In TC, BRAF can be activated by point mutations, small in-frame deletions or insertions, or chromosomal rearrangements, eventually leading to chronic stimulation of MAPK pathway. The most common mechanism of BRAF activation is a thymidine to adenine substitution at nucleotide position 1799 in exon 15. This nucleotide transversion results in a valine-to-glutamate replacement (V600E), producing the constitutively active BRAFV600E mutant protein by the disruption of hydrophobic interactions between residues in the ATP binding site (36). Other BRAF mutations include adenine to guanine substitution at nucleotide position 1801, which results in a lysine-to-glutamate replacement (K601E), and small in-frame insertions or deletions surrounding codon 600 (37). BRAF rearrangement AKAP9/BRAF was also described and it is the result of paracentric inversion of chromosome 7q that leads to the fusion of BRAF-kinase domain with a portion of AKAP9 gene (38).

Activating BRAF mutations are not specific for TC. On the contrary, BRAF mutations were found in many types of cancer (39), with the highest prevalence (50%) in melanoma (40). BRAF mutations in TC are commonly found in nearly 60% of PTC (33), and are also present in nearly 30% of PDTC and 45% of ATC (29). It was demonstrated that transgenic mice with thyroid-specific expression of BRAFV600E developed PTC closely recapitulating those seen in human tumors (41), thus supporting the role of mutant BRAF in tumor initiation and differentiation. A multicentre study showed that BRAFV600E mutation was associated with aggressive behavior of PTC, including aggressive clinicopathological features, tumor recurrence and treatment failure (42).
**RAS**

*RAS* (*rat sarcoma*) gene family encodes small-GTPases involved in signal transduction from the membrane to the nucleus (43), leading to protein synthesis and regulation of cell survival, proliferation and differentiation. In humans, RAS family consists of three genes: *NRAS*, *KRAS* and *HRAS*.

RAS is a monomeric guanosine-nucleotide binding protein which resides in the inner surface of cell membrane and cycles between an inactive form, when bound to guanosine diphosphate (GDP), and an active form, when bound to guanosine triphosphate (GTP). After binding of the ligand to tyrosine kinase and G-protein coupled receptors, RAS switches to the GTP-bound active conformation and transmits signals on downstream pathways, including MAPK and PI3K/AKT cascades. RAS mutations were found in many tumor types and are usually limited to a small number of amino acids (12, 13, 59 and 61) (44). These single point mutations cause the loss of the GTPase activity of RAS, which is blocked in the constitutively active GTP-bound form and continuously stimulates signal transduction.

*NRAS* and *HRAS* mutations at codon 61 are the most common in TC. In PTC, RAS mutation is the second most frequent genetic alteration identified (13-20%) and is commonly observed in the follicular variant (27,33). Mutations of RAS are found in benign FTAs (3), are common in FTC (49%) (45), and also present in PDTC (45%) and ATC (23%) (46). It is worth to note that, in TC, *BRAF* and RAS mutations are mutually exclusive (33,46). Even though activated mutant RAS is not sufficient to induce tumor formation per se, it may predispose normal cells to acquire further genetic or epigenetic alterations leading to a fully transformed phenotype (47).
• **RET**

**RET (rearranged during transfection)** encodes a transmembrane receptor tyrosine kinase (RTK) that binds, together with its co-receptors of the GDNF family receptor-α (GFRα1-4) family, members of the glial cell line-derived neurotrophic factor (GDNF) family of extracellular signaling molecules (48). Upon ligand binding to the co-receptor, RET dimerizes and autophosphorylates its intracellular tyrosine residues, which serve as docking sites for adaptor proteins that stimulate multiple downstream pathways such as MAPK and PI3K/AKT (49). **RET** is located in the pericentromeric region of chromosome 10q11.2, and is required for the normal development, maturation and maintenance of several tissues and cell types, such as nervous system, kidney and male germ cells (50). Germline loss-of-function mutations of **RET** cause impaired formation of the enteric nervous system and congenital aganglionosis of the colon (Hirschsprung’s disease), while germline activating point mutations are causally related to the hereditary forms of MTC: multiple endocrine neoplasia type 2A and 2B (MEN2A, MEN2B) and familial MTC.

In follicular cell-derived TC, several fusions of **RET** with partner genes, as result of chromosomal rearrangements, were identified (48). **RET** rearrangements, which generate the so-called RET/PTC oncoprotein, occur in 7-20% of PTCs (27,33) and generally result in a juxtaposition of the **RET** 3' sequences, encoding the intracellular tyrosine kinase (TK) domain, with 5' sequences of other partner genes (51). To date, up to 20 different **RET** rearrangements were identified in TC, and all partner genes share the peculiarity of constitutive expression in thyroid follicular cells. The resulting chimeric fusion protein RET/PTC is located in the cytosol, instead of membrane, and is able to form dimers in a ligand-independent manner, leading to chronic stimulation of downstream signaling pathway and tumorigenesis in thyroid cells. The two most common **RET/PTC** rearrangements thus far identified in PTC are **RET/PTC1** and **RET/PTC3**, in which **RET** is
fused to **CCDC6** (also known as **H4**) and **NCOA4** (also known as **ELE1** or **RFG**), respectively. Both rearrangements are intrachromosomal inversions of the long arm of chromosome 10.

Several studies reported an increase in the prevalence of **RET/PTC** rearrangements in PTC patients with a radiation history (52,53). Indeed, ionizing radiations produce double-stranded breaks that can cause the misjoining of chromosome 10 loci, predisposing to intrachromosomal inversion and RET/PTC formation (51).

Apart from PTC, even if rare, **RET/PTC** rearrangements were also found in PDTC and ATC, mainly in tumors associated with a differentiated component (54).

The association of **RET/PTC** rearrangement with clinicopathological features of increased mortality in PTC is still controversial; however, a positive correlation with a more aggressive phenotype and tumor advanced stage was demonstrated in sporadic PTCs with **RET/PTC3** rearrangements, which had both large tumor size and an advanced disease stage at the time of diagnosis; other studies highlighted a role of **RET/PTC3** in metastatic spread (51).

Interestingly, RET rearrangements are not limited to TC, but were identified also other solid tumors such as lung, breast and colon carcinomas (55).

- **TRK**

  **TRK** oncogenes are the product of rearrangements of **NTRK1** (*neurotrophic receptor tyrosin kinase A, or TRKA*) gene on chromosome 1q21-22 (56). **NTRK1** encodes the transmembrane high affinity receptor for the nerve growth factor (NGF), which exerts a critical role in the regulation of development and differentiation of peripheral and central nervous system. Following NGF binding, NTRK1 undergoes dimerization and autophosphorylation on intracellular tyrosine residues; these phosphorylated residues recruit adaptor proteins that activate multiple downstream pathways, such as MAPK and
PI3K/AKT (57). These signals, similar to those triggered by RET, culminate in the induction of genes that regulate cell proliferation and survival.

Deregulation of NTRK1 signaling was found associated with different human diseases: NTRK1 mutations were found in CIPA (Congenital Insensitivity to Pain with Anhidrosis (58), acute myeloid leukemia (59), and deregulated NGF/NTRK1 activity was found associated with prostate (60) and breast (61) cancer development.

Several TRK oncogenes originating from NTRK1 rearrangements were found in TC, particularly in the PTC histotype (56). The most frequent is TRK, that results from the fusion of the NTRK1 sequence, encoding the TK domain, to sequences from of the non-muscle tropomyosin (TMP3) gene on chromosome 1q22-23. Other rearrangements include TRK-T1 and TRK-T2, which derive both from the rearrangement of NTRK1 with translocated promoter region (TPR) gene located on chromosome 1q25, but each displaying a different structure. TRK-T3 oncogene derives from the fusion of portions of NTRK1 and the ubiquitously expressed TRK-fused gene (TFG), located on chromosome 3q11-12 and first identified in this rearranged version (62).

Somatic rearrangements of NTRK1 account for about 1-2% of genetic alterations in PTC. Although occurring less frequently than RET rearrangements, experimental evidence indicate that TRK oncogenes represent an early event in the thyroid carcinogenesis, as transgenic mice carrying TRK-T1 under control of thyroglobulin promoter (Tg-TRK-T1 mice) develop hyperplasia and PTC (63).

As for RET, also NTRK1 rearrangements were identified in solid tumors besides PTC: these consist in colorectal cancer, lung cancer, pediatric brain tumors and glioblastoma, cholangiocarcinoma and sarcoma (55).
1.1.2.3. **PI3K/AKT signaling pathway**

PI3K/AKT signaling pathways plays an important role in the regulation of several cellular functions such as growth, motility and survival (64). The relevance of this pathway in human tumorigenesis and its targeting as therapeutic approach have been widely investigated in the last decade (65,66), also in thyroid tumors (67).

- **PIK3CA**

  *Phosphatidylinositol-4,5-Bisphosphate 3-Kinase Catalytic Subunit α (PIK3CA)* encodes the 110 kDa catalytic subunit of the class I PI3 kinase complex that is activated by RTKs and, when active, phosphorylates phosphatidylinositol-4,5-bisphosphate to produce phosphatidylinositol-3,4,5-trisphosphate (PIP3), the activator of AKT.

  *PIK3CA* involvement in human cancer development is well documented, as somatic activating mutations of this gene were identified in a variety of tumors, such as colorectal, breast and ovarian carcinomas (68). In the context of thyroid, mutations of *PIK3CA* were identified in FTC and ATC histotypes, with a frequency of 10-15%, and in PDTC and ATC either recurrent or with metastatic disease (64). Also *PIK3CA* copy-number gain is reported in TC, mainly in FTC and ATC and uncommonly in PTC. This alteration results in *PIK3CA* amplification and is associated with overexpression of the protein and hyper-phosphorylation of AKT.

- **PTEN**

  *Phosphatase and tensin homolog (PTEN)* encodes the protein phosphatase responsible for the degradation of PIP3, thus antagonizing and terminating the PI3K/AKT signaling.

  *PTEN* has a crucial role as tumor suppressor gene and different mechanisms inhibiting its activity were described for human cancers, including TC (64). These mechanisms include somatic mutations (although with low frequency), loss of heterozygosity or deletion, and promoter hypermethylation. Germline inactivating mutations of *PTEN* cause Cowden’s
syndrome and predispose patients to develop tumors, including FTC. *PTEN* alterations are found in FTC and, more commonly, in ATC (15%) where, together with other genetic alterations of members of the PI3K/AKT pathway, can ideally cooperate and contribute to the aggressiveness of this thyroid tumor (29).

1.1.2.4. Other mutations

- **PAX8/PPARγ**

  The paired box 8/peroxisome proliferator-activated receptor-γ fusion gene (*PAX8/PPARγ*) is another recombinant oncogene in TC, originating from translocation between chromosomal regions 2q13 (*PAX8*) and 3p25 (*PPARγ*) (69). PAX8 is a member of the paired box family of transcription factors required for normal thyroid development; it also induces the expression of thyroid-specific genes in the mature thyroid. PPARγ is a member of the nuclear receptor family of transcription factors and is the master regulator of adipogenesis; it is expressed at very low levels and has no known function in the normal thyroid. The fusion product *PAX8/PPARγ* encodes a nearly full length PPARγ, whose expression is under transcriptional control of PAX8 promoter which is highly active in thyroid follicular cells. Several in vitro experiments provided evidence that *PAX8/PPARG* acts as oncogene in TC (70-72), although the specific mechanism is yet to be defined. *PAX8/PPARγ* rearrangements are prevalent in FTC (30-35%), follicular variant of PTC (13%), and follicular adenomas, while they are absent in undifferentiated PDTC and ATC (73).

- **TP53**

  *TP53* encodes a transcription factor that acts as tumor suppressor, playing a pivotal role in regulation of cell proliferation, DNA damage response and apoptosis (74). In physiological conditions, p53 forms a complex with MDM2, an E3 ubiquitin protein ligase that promotes its degradation by the proteasome. Upon DNA damage or other cellular...
stresses, various pathways activate and culminate in p53 activation by promoting its
dissociation from MDM2. Once activated, p53 binds to DNA and drives the expression of
several genes that lead to cell cycle arrest, apoptosis and/or senescence, contributing to
the maintenance of genome stability. TP53 mutations are found in up to 50% of human
cancers and are generally loss-of-function missense mutations (75). TP53 mutations are
rare in well-differentiated PTC and FTC (76) and also in PDTC (about 8% of frequency),
while they are a common genetic event in the undifferentiated ATCs (up to 73% of cases)
(29). It was demonstrated that TP53 mutations have a role in the progression from well-
differentiated TC to the most aggressive ATC (77), thus supporting the notion that TC
initiation and progression occurs through gradual accumulation of multiple genetic
lesions.

1.1.3. Treatment of thyroid cancer

1.1.3.1. Conventional therapy

The treatment of choice for well-differentiated primary thyroid tumors is surgery, that
consists of partial lobectomy or total thyroidectomy, based on the dimensions and the
clinicopathological features of the thyroid nodule detected (78). Controversy still exists on
the extent of the surgery: partial lobectomy was shown to have the same effects on
patient overall free survival as total thyroidectomy did, but with less frequent adverse
events (such as hypocalcemia and laryngeal nerve paralysis) (79). Nevertheless, with the
exception of unifocal low-risk PTCs smaller than 1 cm, the American Thyroid Association
(ATA) guidelines recommend the complete removal of the thyroid to minimize the risks of
malignancy relapse (80). Together with the thyroid, patients diagnosed with primary PTC
undergo lymph node removal even in the absence of clear nodal metastases, since from
22% to 90% of PTCs spread locally to lymphoid glands. Following surgery, patients are usually treated with radioactive iodine (RAI, $^{131}$I), in order to remove potential occult WDTC foci and decrease the long-term risk of tumor recurrence. In addition to surgery and RAI, thyroid-stimulating hormone (TSH) suppressive treatment is practiced to prevent major adverse clinical events, since WDTCs are usually responsive to TSH stimulation in terms of cell growth. WDTC are generally cured by standard therapy; nevertheless, tumor recurrence can happen, as in up to 20% of patients the tumor relapses and in 10% of patients distant metastases occur (81).

Management of the undifferentiated ATCs is more challenging: surgical removal of the thyroid remains the first choice of therapeutic intervention, but many patients often present at diagnosis with extended local disease invading the surrounding structures (such as trachea and larynx) that makes the tumor mass unresectable (25). Moreover, ATCs are usually refractory to RAI therapy. When surgery is feasible, thyroidectomy is combined with locoregional radiation therapy and chemotherapy with agents such as taxanes or platinum-based antineoplastic drugs, like carboplatin. For patients with unresectable disease, palliative therapy to preserve the airway is mandatory, and a combination of radiations and chemotherapy is applied with the aim of achieving local tumor regression and/or durable local tumor control. However, even if multimodality therapy has shown to prolong patient survival, a successful standardized protocol has not been defined yet and the optimal sequence of multimodality therapy still remains to be determined.

1.1.3.2. **Novel therapies**

In the last decades, the increased knowledge on the molecular events that drive tumorigenesis allowed the development of the so-called precision medicine (82): in cancer, including TC, the improved understanding of the pathogenesis and genetics of
tumors prompted the development of new therapeutic opportunities for patient treatment (Figure 1.4) (83).

**Figure 1.4 Signaling pathways involved in TC pathogenesis and their relevant inhibitors**
Kinase receptors are shown with their corresponding ligands and downstream signaling pathways. Pharmacological inhibitors, which operate at different levels in signaling pathways, are indicated. Image from (83).

A variety of molecular-targeted agents able to inhibit RTKs or downstream effectors is nowadays available. These small molecules are competitive inhibitors of the ATP-binding domains of multiple kinases and acquired great interest as most of their targets belong to the main pathways involved in TC pathogenesis and progression. The potential of this relatively new class of drugs is that their activity may not be restricted to only one target, but that can act on multiple signaling pathways. WDTCs are characterized by increased
peritumoral angiogenesis and high secretion of the vascular endothelial growth factor (VEGF), and the blockade of its receptor (VEGFR) has shown promising results in attenuating tumor cell growth in vivo. Small-molecule tyrosine kinase inhibitors (TKI) often inhibit, in addition to the target kinase, VEGFR and other angiokinase receptors (such as PDGFR, FGFR), thus hindering multiple signaling pathways of potential therapeutic relevance (Table 1.2).

<table>
<thead>
<tr>
<th>DRUG</th>
<th>VEGF-R</th>
<th>c-Kit</th>
<th>RET</th>
<th>PDGF-R</th>
<th>FGF-R</th>
<th>EGF-R</th>
<th>Other targets</th>
<th>Study phase</th>
<th>Approved for thyroid cancer treatment</th>
</tr>
</thead>
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<tr>
<td>Axitinib</td>
<td>Yes</td>
<td>Yes</td>
<td>No</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
<td>dual PI3K/mTOR</td>
<td>II</td>
<td>No</td>
</tr>
<tr>
<td>Bevacizumab</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>dual PI3K/mTOR</td>
<td>II</td>
<td>No</td>
</tr>
<tr>
<td>Cabozantinib</td>
<td>Yes</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>MET, RET-KIF58 rearrangement</td>
<td>III*</td>
<td>Yes</td>
</tr>
<tr>
<td>Imatinib</td>
<td>No</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>Bcr-Abl</td>
<td>II</td>
<td>No</td>
</tr>
<tr>
<td>Lenvatinib</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
<td>RET-KIF58, CCDC63-RET, NeuroRET rearrangement</td>
<td>III*</td>
<td>Yes</td>
</tr>
<tr>
<td>Motesanib</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
<td></td>
<td>II</td>
<td>No</td>
</tr>
<tr>
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<td>No</td>
<td>Yes</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
<td></td>
<td>II</td>
<td>No</td>
</tr>
<tr>
<td>Pazopanib</td>
<td>Yes</td>
<td>No</td>
<td>Yes</td>
<td>No</td>
<td>Yes</td>
<td>No</td>
<td></td>
<td>II</td>
<td>No</td>
</tr>
<tr>
<td>Ponatinib</td>
<td>No</td>
<td>No</td>
<td>Yes</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
<td>Bcr-Abl, FLT3, KIT</td>
<td>II</td>
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<td>Selumetinib</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>MEK</td>
<td>III</td>
<td>No</td>
</tr>
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<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
<td>Raf, FLT3</td>
<td>III</td>
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<tr>
<td>Sunitinib</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
<td>FLT3</td>
<td>II</td>
<td>No</td>
</tr>
<tr>
<td>Vandetanib</td>
<td>Yes</td>
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<td>No</td>
<td>No</td>
<td>No</td>
<td>Yes</td>
<td>RET-KIF58 rearrangement</td>
<td>III*</td>
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</tr>
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<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>BRAF^{V600E, G469V}, CRAF</td>
<td>II</td>
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</tr>
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<td>Everolimus</td>
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<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>mTOR</td>
<td>II</td>
<td>No</td>
</tr>
<tr>
<td>Temsirolimus</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>mTOR</td>
<td>II</td>
<td>No</td>
</tr>
</tbody>
</table>

**Table 1.2 Small-molecule TKIs and their targets tested in clinical trials for TC**
Adapted from (84)

Among the several TKIs tested in phase I, II and III trials for TC treatment, only 4 of them were approved for clinical treatment: sorafenib and lenvatinib for progressive DTC, vandetanib and cabozantinib for MTC (84). Despite the longer patient-free survival duration relative to placebo in randomized clinical trials, these drugs produced several side effect profiles in patients (Table 1.3) and, to date, few data are available about the prolongation of patient overall survival. Of note, treatment of ATC with TKIs has still to be rigorously studied and defined: among the variety of molecules previously listed, only lenvatinib showed promising antitumor activity in ATC patients (85).
As mortality is increased for patients with WDTCs that become refractory to RAI therapy, another strategy that has been developed in the last decades is the development of agents that promote tumor re-differentiation, which can make tumors radioiodine-avid again and therefore treatable with $^{131}$I. Several agents, among which retinoic acid, were tested in RAI-refractory patients but gave no significant benefits (86). A promising candidate for RAI-resensitization is the MEK inhibitor selumetinib, which has been able to increase the uptake of RAI and gave a partial response and decrease in tumor size, especially in patients with NRAS-mutant tumors (87).

Another therapeutic approach for TC cell re-differentiation is the use of epigenetic drugs, specifically histone deacetylase (HDAC) inhibitors. Epigenetic alterations in tumor cells often result in the silencing of genes involved in cell differentiation, given by the hypermethylation of cytosines of the promoter and the subsequent recruitment of HDACs to repress expression of the genes (88). There are various classes of HDACs in cells, thus different compounds were developed to specifically target one class or another.

<table>
<thead>
<tr>
<th>Drug</th>
<th>Vandetanib</th>
<th>Cabozantinib</th>
<th>Sonafenib</th>
<th>Lenvatinib</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name of the study</td>
<td>ZETA</td>
<td>EXAM</td>
<td>DECISION</td>
<td>SELECT</td>
</tr>
<tr>
<td>Tumour</td>
<td>MTC</td>
<td>MTC</td>
<td>ETC</td>
<td>ETC</td>
</tr>
<tr>
<td>Phase</td>
<td>III</td>
<td>III</td>
<td>III</td>
<td>III</td>
</tr>
<tr>
<td>Patients (n)</td>
<td>331</td>
<td>330</td>
<td>417</td>
<td>392</td>
</tr>
<tr>
<td>Partial response (%)</td>
<td>45</td>
<td>28</td>
<td>12.2</td>
<td>64.8</td>
</tr>
<tr>
<td>Stable disease ≥ 6 months (%)</td>
<td>87</td>
<td>NE</td>
<td>42</td>
<td>15.3</td>
</tr>
<tr>
<td>Median progression-free survival (months)</td>
<td>30.5*</td>
<td>11.2</td>
<td>10.6</td>
<td>18.3</td>
</tr>
<tr>
<td>Median overall survival (months)</td>
<td>NE</td>
<td>NE</td>
<td>NE</td>
<td>NE</td>
</tr>
<tr>
<td>Most frequent adverse events (%)</td>
<td>Diarrhoea (56)</td>
<td>Diarrhoea (53)</td>
<td>Diarrhoea (76)</td>
<td>Hypertension (67)</td>
</tr>
<tr>
<td>Any grade</td>
<td>Skin rash (45)</td>
<td>Hand-foot syndrome (50)</td>
<td>Diarrhoea (68)</td>
<td>Diarrhoea (59)</td>
</tr>
<tr>
<td></td>
<td>Nausea (33)</td>
<td>Weight loss (47)</td>
<td>Alopecia (67)</td>
<td>Fatigue or asthenia (59)</td>
</tr>
<tr>
<td></td>
<td>Hypertension (32)</td>
<td>Anorexia (45)</td>
<td>Skin rash (50)</td>
<td>Anorexia (50)</td>
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<tr>
<td></td>
<td></td>
<td>Nausea (43)</td>
<td>Fatigue (49)</td>
<td>Weight loss (46)</td>
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<tr>
<td></td>
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<td>Fatigue (49)</td>
<td>Weight loss (46)</td>
<td>Hypertension (40)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Anorexia (31)</td>
</tr>
<tr>
<td>Molecular targets</td>
<td>VEGF-R, c-Kit, RET, EGFR, RET-ROS rearrangement</td>
<td>VEGF-R, c-Kit, RET, MET, RET-ROS rearrangement</td>
<td>VEGF-R, c-Kit, RET, MET, RET-ROS rearrangement</td>
<td>VEGF-R, c-Kit, RET, PDGF-R, FGFR, RET-ROS, CCDC6-Ret, NcoA4-RET rearrangement</td>
</tr>
</tbody>
</table>

Table 1.3 TKIs approved for the treatment of TC and details of phase III trials
Adapted from (84)
Suberoylanilide hydroxamic acid (SAHA, Vorinostat®) is a pan-HDAC inhibitor that was able to increase Na⁺/I⁻ symporter (NIS) expression and reduce WDTC and ATC cell growth in vitro. Romidepsin (Istodax™) is a class I-HDAC inhibitor able to restore RAI avidity in some TC patients and therefore could potentially increase the response to RAI therapy. Panobinostat (Fraydak®) induced re-expression of NIS and increased response to RAI treatment in vitro, while valproic acid (Depakene®) was able to increase NIS mRNA expression. Nevertheless, none of these compounds produced significant responses in the clinics to date.

1.1.3.3. Conclusions

Despite the improvement of TC therapy, some areas of uncertainty persist (84). Limitations of targeted therapy are mainly made by the toxicity profiles that usually lead to the reduction of therapy doses, the onset of escape mechanisms of resistance to treatment, and notably no data about the long-term effects on patient quality of life is available. ATC still remains one of the deadliest diseases worldwide, as no effective therapy has been identified to date and targeted therapies are now tested in phase II clinical trials.
1.2. Non-Oncogene Addiction

1.2.1. Oncogene and non-oncogene addiction principles

Nowadays it is well known that cancer arises from a mutagenic multistep process during which transforming cells acquire several properties, such as uncontrolled proliferation, resistance to antiproliferative signals and resistance to apoptotic stimuli (89). These properties are shared by all tumor cells and are achieved through random mutations and epigenetic changes in oncogenes and tumor suppressor genes, which eventually lead to the modification of already existing cellular programs normally used during development. The concept of “oncogene addiction” indicates the dependency of tumor cells on the activity of oncogenes and oncogenic pathways that are essential for tumor cell survival (90). This concept put the rationale for therapeutic intervention by targeting the mutated components of key cellular pathways on which tumor cells became dependent. A variety of drugs targeting specific oncogenes in different tumors were developed and tested, such as HER2 in breast cancer (91), BRAF in melanoma (92), EGFR in non-small cell lung cancer (93) and BCR-ABL in chronic myeloid leukemia (94). However, even though remissions were observed in some cases, these were often followed by the recurrence of drug-resistant tumor cells. Moreover, the targeting of oncogenes like RAS or MYC is complex, as the identification of compounds specifically targeting them is difficult. Based on these observations, the “synthetic lethality” concept was exploited. It indicates a phenomenon of lethal combination of mutations/perturbations in two genes, where mutations in only one of those genes have no effect on cell viability (95). The targeting of a synthetic lethal partner of a mutated driver gene in tumor cells may provide two main advantages: the overcoming of the lack of inhibitors for certain oncogenes and the sparing of normal cells during therapeutic
intervention. An example of promising synthetic lethality application in the clinics is the use of poly ADP-ribose polymerase (PARP) inhibitors in advanced ovarian cancer patients with mutations in BRCA1/2, which exhibit specific DNA repair defect and result sensitive to this class of drugs (96).

Beyond the concept of oncogene addiction, it was observed that tumor cells often face increased levels of cellular stresses, caused by the sustained activation of oncogenic pathways, compared to normal cells (97). This phenomenon was referred to as “stress phenotype” of cancer cells (Figure 1.5).

![Image](image.png)

**Figure 1.5 The hallmarks of cancer revisited**

In addition to the six hallmarks of cancer initially proposed by Hanahan and Weinberg (89), additional hallmarks were proposed, in order to highlight the “stress phenotype” of cancer cells. These new hallmarks include DNA damage, metabolic, oxidative, mitotic and proteotoxic stresses and, together with the other cancer hallmarks, promote the tumorigenic state of cancer cells. Image from (98)

Increased cellular stresses associated with the oncogenic activity are tolerated by cancer cells through the activation of stress support pathways, on which they become therefore
addicted. These genes act in oncogenic pathways but are not classical oncogenes themselves, as they are not mutated, aberrantly expressed or responsible for tumorigenesis initiation (98). This phenomenon has been termed “non-oncogene addiction” (NOA) and highlights the potential of non-oncogenes as point of intervention for cancer therapy, since the impairment of their functions would ideally be lethal for tumor cells while sparing normal ones (Figure 1.6).

![Figure 1.6 Schematic representation of cellular non-oncogene addiction effects](image)

Cancer cells, according to NOA paradigm, face elevated cellular stresses produced by collateral events of the tumorigenic process. Interfering with the stress support pathways in tumor cells will lead to their killing, while normal cells are able to tolerate these perturbations. Image adapted from (99).

Tumor cell dependencies on NOA genes provide an attractive target for therapeutic intervention, which could be achieved through two different approaches: stress sensitization, that consists in diminishing the activity of stress support pathways in order to promote cell growth inhibition and cell death, and stress overload, that aims to intensify oncogenic stress to overload the stress support pathways in tumor cells and eventually lead to growth arrest or cell death. Stresses faced by tumor cells are DNA damage, mitotic, proteotoxic, metabolic and oxidative stress (Figure 1.7).
Non-oncogene addiction examples in tumor cells

The tumorigenic state of cancer cells results in several alterations of cellular homeostasis related to the hallmarks of cancer, described in Figure 1.5. These alterations represent potential tumor vulnerabilities, and interfering with regulators of these support pathways could display synthetic lethality with the tumor phenotype. Image from (98)

Because of the high degree of replication and the activity of oncogenes themselves, the rate of spontaneous DNA damage in tumor cells is increased, compared to the normal ones. Indeed, several studies demonstrated that inhibition of members of DNA damage response pathway, such as ATM and CHK1, exerted selective toxicity on tumor cells, as they increased DNA damage stress. Linked to the high degree of replication of tumor cells, there is also mitotic stress, which is the result of increased rates of chromosome segregation errors caused by mutations altering mitosis fidelity. The inhibition of proteins regulating mitosis was found effective in sensitizing tumor cells to stress overload, as it is commonly observed upon treatment with taxol or mitotic kinase inhibitors. Transcriptional alterations and copy-number variations normally happening in tumor cells also result in proteotoxic stress: these changes often produce alterations of protein dosage balance, thus resulting in steady activation of chaperone pathways to maintain normal proteome homeostasis and, in case of unfolded proteins, of the ubiquitin-
proteasome pathway. Indeed, inhibition of the proteasome with bortezomib has proven to be deleterious for tumor cells, and provides an example of proteotoxic stress overload. All these lines of evidence support the concept that NOA genes are attractive targets for therapeutic intervention, since they represent key functional nodes in the oncogenic network and their inhibition would result in system failure. According to the NOA paradigm, several groups already provided promising results in “targeting of non-oncogene addiction”: PARP inhibition in glioblastoma (100), silencing of heat shock factor 1 (HSF1) in multiple cancer cells (101), and HDAC6 inhibition in breast cancer (102) resulted in inhibition of tumor cell growth.

1.2.2. Tools for the identification of cancer cell vulnerabilities

The traditional approach used to identify potential targets for cancer treatment, in the last decades, consisted in the identification of the specific genetic lesions at the basis of tumorigenic process, such as copy-number variations, gene mutations, transcriptional changes and epigenetic modifications (103). Despite being a powerful approach for identifying genetic changes in oncogenes or tumor suppressor genes, some limitations persist: sequencing techniques nowadays mostly limit to coding regions and therefore NOA genes and pathways are not detected, since they are not directly mutated albeit supportive in the tumorigenic process. However, the revolution of RNA interference (RNAi) technologies allowed the development of a functional approach to identify such critical nodal points for cancer therapeutic intervention (104). Functional RNAi screenings are indeed able to identify cancer vulnerabilities, as the investigation is not limited to mutated genes, such as oncogenes, but also includes genes and pathways important for tumor sustainment, that is NOA genes. This strategy is also powerful in terms of target
discovery, as either partial or complete loss of protein function strictly mimics the effects of a pharmacological inhibitor. The wide development of commercially available large-scale RNAi libraries, that can be constructed with different strategies (Figure 1.8), contributed to the improvement of the use of large-scale RNAi-based screenings with the aim of dissecting complex biological processes (105). Such large scale RNAi-based functional screenings performed both *in vitro* (106-108) and also *in vivo* (109,110) were successful in identifying genes essential for the regulation of different types of cancer.
RNAi screenings can be performed using different strategies. Arrayed library screening, using synthetic small interfering RNAs (siRNAs) or short hairpin RNAs (shRNAs), target each gene separately in a multiwell format and various readouts can be used to determine the silencing effect on the phenotype of interest. Arrayed screening using cell microarrays allow to plate cells directly onto microarrays containing the siRNAs/shRNAs, and the target gene is identifiable by the position on the array. Pooled library screens target many genes pools of shRNA expressing vectors are introduced into cells by transfection or infection for pooled library screening; cells are exposed to a selective agent and the shRNA-plasmids present in surviving cells are identified by DNA sequencing or barcode microarray analysis. Image from (111).
Aim of the study

The non-oncogene addiction paradigm highlighted novel tumor cell vulnerabilities that could represent potential points for therapeutic intervention in cancer. Moreover, the improvement of RNAi technologies allowed the development of new tools, that are large-scale RNAi screening libraries, for the investigation of these critical nodal points in the complex cancer network. While NOA was approached for tumors such as myeloma and breast cancer, in the context of thyroid cancer the NOA paradigm is a completely novel concept. Given that current therapies for thyroid tumors are not always effective, particularly for the most aggressive and advanced cases, the identification of TC cell vulnerabilities would be useful for the development of new therapeutic approaches.

Using a siRNA library targeting the whole “druggable” genome, our laboratory has previously performed a loss-of-function based screening on a tumor and a normal thyroid cell lines, in order to identify genes whose silencing interfered with the growth of tumor but not normal cells. This screening allowed us to identify 15 differentially active hits in the tumor versus the normal cell lines.

The overall aim of the project was the identification of thyroid tumor cell vulnerability genes, according to the NOA paradigm, and the dissection of their role in thyroid tumor cell biology, in order to unveil potential novel targets for thyroid cancer treatment. Most of the work was devoted to hit gene validation studies and to studies on MASTL, a vulnerability gene we identified in TC cells. These two sections have been well developed and hereafter presented in two distinct chapters. The thesis includes a third chapter that summarized the studies regarding COPZ1, which is another vulnerability gene identified in TC cells, to which I collaborated.
2. Materials and methods

2.1. Cell Biology

2.1.1. Cell culture

Nthy-ori 3-1 cell line (SV-40 immortalized normal human thyroid follicular cells) was purchased from European Collection of Cell Cultures (ECACC) (Salisbury, UK); tumor BCPAP, TPC-1, WRO82-1, 8505C and KAT-18 cell lines were obtained from Prof. A. Fusco (University of Federico II, Naples, Italy); HTC/C3 cell line was purchased from Riken Cell Bank (Tsukuba, Japan). Information regarding the histotype derivation of cell lines and the mutational status are provided in Table 2.1. HEK 293T cell line was purchased from American Type Culture Collection (ATCC).

Nthy-ori 3-1 cells were cultured in RPMI 1640 medium (Gibco Life Technologies, Carlsbad, CA, USA) supplemented with 10% heat-inactivated fetal bovine serum (FBS) (EuroClone, Pero, Italy). The other cell lines were maintained in DMEM medium (Gibco Life Technologies, Carlsbad, CA, USA) containing 10% FBS and 2mM glutamine. All cell lines were cultured as monolayer at 37°C in a 5% CO₂ humidified atmosphere.

Cell lines were authenticated by short tandem repeat (STR) profiles using Stem Elite ID System (Promega Corporation, Madison, USA) by the Fragment Analysis Facility of Fondazione IRCCS Istituto Nazionale dei Tumori. The profiles obtained matched to their original profiles (112,113). WRO82-1 profile matched to that reported by Xu and colleagues (114) and HTC/C3 to that reported by JCRB Cell Bank (http://cellbank.nibio.go.jp/legacy/celldata/jcrb0164.htm).
Mycoplasma contamination was tested periodically and found negative in all cell lines (PCR Mycoplasma Detection Set, TAKARA Bio Inc).

Table 2.1 List of thyroid cell lines with histological derivation and mutational status

<table>
<thead>
<tr>
<th>Cell lines</th>
<th>Hystological type</th>
<th>Mutational status</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nthy-ori 3-1</td>
<td>normal human thyrocytes</td>
<td></td>
</tr>
<tr>
<td>BCPAP</td>
<td>PTC</td>
<td>BRAF V600E</td>
</tr>
<tr>
<td>TPC-1</td>
<td>PTC</td>
<td>RET/PTC1</td>
</tr>
<tr>
<td>WRO82-1</td>
<td>FTC</td>
<td></td>
</tr>
<tr>
<td>8505C</td>
<td>ATC</td>
<td>BRAF V600E; TP53 R248L</td>
</tr>
<tr>
<td>HTC/C3</td>
<td>ATC</td>
<td>BRAF V600E; TP53 P152L</td>
</tr>
<tr>
<td>KAT-18</td>
<td>ATC</td>
<td></td>
</tr>
</tbody>
</table>

2.1.2. Cell transfection and gene silencing

Transient gene silencing of the hit genes was obtained in thyroid cell lines by the transfection of specific siRNAs, detailed in Table 2.2, using Lipofectamine RNAiMAX reagent (Invitrogen Life Technologies, Carlsbad, CA, USA), according to manufacturer’s protocol. Briefly, cells were seeded at the optimal density and transfected the following day by adding a mixture of Opti-MEM (Reduced Serum Media) (Gibco Life Technologies, Carlsbad, CA, USA) containing 20 nM siRNA and 3 µl/ml Lipofectamine RNAiMAX.
<table>
<thead>
<tr>
<th>siRNA</th>
<th>Sequence/catalog number</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>siNT</td>
<td>ON-TARGET plus Non-targeting Pool siRNA D-0018-10-05</td>
<td>ThermoScientific, Dharmacon Inc., Chicago, IL, USA</td>
</tr>
<tr>
<td>siPSMC3</td>
<td>MISSION esiRNA EHU107331</td>
<td>Sigma-Aldrich, St. Louis, MO, USA</td>
</tr>
<tr>
<td>siMASTL #1</td>
<td>ON-TARGET plus set of 4 siRNAs J-004020-12 5’-CCAUUGAGACGAAAGGUUU-3’</td>
<td>ThermoScientific, Dharmacon Inc., Chicago, IL, USA</td>
</tr>
<tr>
<td>siMASTL #2</td>
<td>ON-TARGET plus set of 4 siRNAs J-004020-11 5’-GCAAAUUGUAUGCAGUAAG-3’</td>
<td>ThermoScientific, Dharmacon Inc., Chicago, IL, USA</td>
</tr>
<tr>
<td>siMASTL #3</td>
<td>ON-TARGET plus set of 4 siRNAs J-004020-10 5’-ACUGGACGCUCUUUGUGUAAG-3’</td>
<td>ThermoScientific, Dharmacon Inc., Chicago, IL, USA</td>
</tr>
<tr>
<td>siMASTL #4</td>
<td>ON-TARGET plus set of 4 siRNAs J-004020-09 5’-GGACAAGUGUUAUGCAGUUA-3’</td>
<td>ThermoScientific, Dharmacon Inc., Chicago, IL, USA</td>
</tr>
<tr>
<td>siMASTL #5</td>
<td>5’-UGUUGAUUUCUGUGGGAUAAdTdT 3’</td>
<td>in house</td>
</tr>
<tr>
<td>siCCND1</td>
<td>MISSION esiRNA EHU153321</td>
<td>Sigma-Aldrich, St. Louis, MO, USA</td>
</tr>
<tr>
<td>siCOPZ1</td>
<td>MISSION esiRNA EHU1040461</td>
<td>Sigma-Aldrich, St. Louis, MO, USA</td>
</tr>
<tr>
<td>siMAP4K5</td>
<td>MISSION esiRNA EHU120861</td>
<td>Sigma-Aldrich, St. Louis, MO, USA</td>
</tr>
<tr>
<td>siCOPE</td>
<td>MISSION esiRNA EHU069701</td>
<td>Sigma-Aldrich, St. Louis, MO, USA</td>
</tr>
<tr>
<td>siPLA2G15 #1</td>
<td>ON-TARGET plus set of 4 siRNAs J-009622-08 5’-GAAGAUGGUCCUGCCUAUGC-3’</td>
<td>ThermoScientific, Dharmacon Inc., Chicago, IL, USA</td>
</tr>
<tr>
<td>siPLA2G15 #2</td>
<td>ON-TARGET plus set of 4 siRNAs J-009622-07 5’-GCAGAGGUACCUGUGAACU-3’</td>
<td>ThermoScientific, Dharmacon Inc., Chicago, IL, USA</td>
</tr>
<tr>
<td>siPLA2G15 #3</td>
<td>ON-TARGET plus set of 4 siRNAs J-009622-06 5’-GGAGAUGGUACCAGCUGUAU-3’</td>
<td>ThermoScientific, Dharmacon Inc., Chicago, IL, USA</td>
</tr>
<tr>
<td>siPLA2G15 #4</td>
<td>ON-TARGET plus set of 4 siRNAs J-009622-05 5’-CAGCGUGGGGUUCCUAAUUC-3’</td>
<td>ThermoScientific, Dharmacon Inc., Chicago, IL, USA</td>
</tr>
</tbody>
</table>
2.1.3. Cell proliferation assays

2.1.3.1. CellTiter-Glo assay

For hit validation studies described in chapter 3, cells (500 cells/well) were reverse transfection in 96-well plate with 20 nM specific siRNAs using the Lipofectamine RNAiMAX reagent and, ten days later, cell viability was assessed by CellTiter-Glo® Luminescent Cell Viability Assay (Promega Corporation, Madison, USA), performed as recommended by the supplier. Luminescence signals were acquired using a microplate reader (TecanUltra, Tecan Trading AG, Switzerland).

For MASTL studies described in chapter 4, HTC/C3 and 8505C cells (1.9 x 10^5) were seeded in 60mm dishes transfected the day later with 20 nM of specific siRNAs using Lipofectamine RNAiMAX. 24 hours after transfection, cells were harvested and plated (2x10^3 cells/well) in 96-well plates; 8505C-TRIPZ-shMASTL #2 (2x10^3) cells were plated in 96-well plates in absence or presence of 2 µg/ml doxycycline. Growth rates were evaluated at the time points indicated in the growth curves after plating by CellTiter-Glo® Luminescent Cell Viability Assay as described above.

2.1.3.2. Trypan Blue exclusion test

BCPAP and 8505C (1x10^5), TPC-1 (6 x 10^4), WRO82-1, HTC-C3 and KAT-18 (8 x 10^4) cells were transfection the day after the seeding in 6-well plate with 20 nM of specific siRNAs using Lipofectamine RNAiMAX, following manufacturer’s protocol. At different days after transfection (4 days for TPC-1, 8505C and HTC/C3 cells; 5 days for KAT-18 cells, 6 days for BCPAP cells, 7 days for WRO82-1 cells) the fraction of viable and dead cells was analyzed.

2.1.3.3. Growth inhibition Assay

For experiments with palbociclib, which is a CDK4 and CDK6 inhibitor (115), cells (8x10^4 cells/well) were seeded in black/clear bottom 96-well plates and, the following
day, treated with increasing doses of the palbociclib (PD 0332991) (Selleck Chemicals, Houston, TX, USA, cat. S2768). After 72 hours, cells were fixed with 3.7% paraformaldehyde solution for 30 minutes at room temperature, washed and stained with 1 mg/ml Hoechst 33342. The plates were analyzed with an ArrayScan VTI high-content screening reader (Thermo-Fisher Scientific, Pittsburgh, PA): at least 800 cells were acquired in each well with a 10X objective in one fluorescence channel (XF93 optical filter set). Cell nuclei were automatically recognized and counted in ten fields to estimate the cell number per well as proliferation readout. IC50 values were calculated by sigmoid curve fitting of experimental points.

For MASTL studies, HTC/C3 and 8505C (1.2 \times 10^5) cells were plated in 24 well-plates and transfected the following day with 20 nM of siRNAs using the Lipofectamine RNAiMAX reagent (Invitrogen Life Technologies, Carlsbad, CA, USA). Seven days after transfection, cells were fixed with 3.7% formaldehyde solution (v/v) at room temperature for 15 minutes, washed with PBS and stained for 20 minutes with 0.1% crystal violet (w/v). Crystal violet was then dissolved in 10% acetic acid (v/v) for 30 minutes on a rocker and then diluted 1:4 in ddH₂O. Optical density at \(\lambda=590\) nm was measured by using Ultrospec 2100 Pro spectrophotometer (GE Healthcare Bio-Sciences, Pittsburgh, PA).

### 2.1.4. Immunofluorescence analysis

For hit validation studies described in chapter 3, 8505C cells growing on glass coverslips were transfected with 20 nM control (NT) or MASTL siRNAs and fixed 48 hours later for 10 minutes with 4% paraformaldehyde (Sigma Aldrich, St. Louis, Mo, USA). After permeabilization for 10 minutes with 1% BSA and 0.1% Triton X-100 in PBS, and incubation for 30 minutes with 1X blocking solution (2% BSA in PBS), cells were incubated with anti-β-tubulin (clone TUB 2.1: T4026, 1:400, Sigma Aldrich, St. Louis, MO, USA) or
anti-histone H3 (phospho S10) (E173: ab32107, 1:1000, Abcam, Cambridge, UK) primary antibodies for 1 hour. After washing with PBS, cells were incubated with Alexa Fluor® 546 rabbit (1:500, Invitrogen/Molecular Probes®) and Alexa Fluor 488® mouse (1:500, Invitrogen/Molecular Probes®) secondary antibodies for 1 hour. Slides were then prepared using ProLong Diamond Antifade mountant with DAPI (P36966, Molecular Probes®) and imaged with immunofluorescence microscopy (Eclipse E1000; Nikon Instruments, Inc. NY, USA).

For MASTL studies described in chapter 4, HTC/C3 and 8505C cells growing on glass coverslips were transfected with 20 nM NT or MASTL siRNAs and fixed 120 (for HTC/C3) and 76 (for 8505C) hours later for 10 minutes with 4% paraformaldehyde. After permeabilization performed as described above, slides were prepared using ProLong Diamond Antifade mountant with DAPI (P36966, Molecular Probes®) and imaged with immunofluorescence microscopy. A minimum of 1900-2000 (for HTC/C3) and 3000 (for 8505C) cells was counted for each siRNA treatment, and the number of cells with abnormal nuclei was expressed as percentage of the total count.

### 2.1.5. Annexin V assay

HTC/C3 (1.9 x 10^5) cells were seeded in 60mm dishes and transfected the following day with 20 nM NT and MASTL #5 siRNAs using Lipofectamine RNAiMAX. At 48, 72, 96 and 120 hours after transfection, both adherent and floating cells were harvested and processed for flow cytometry analysis using FITC Annexin V Apoptosis Detection Kit (Becton Dickinson, San Jose, CA, USA): after washing with PBS, cells were resuspended in Annexin V Binding Buffer and incubated with FITC-conjugated Annexin V for 15 minutes, at room temperature in the dark. TO-PRO-3 iodide (1:2000; Invitrogen/Molecular Probes®), instead of propidium iodide, was then added to each sample. Samples were
acquired using BD FACSCanto (Becton Dickinson, San Jose, CA, USA) instrument and analyzed with FlowJo v7.6 software (TreeStar).

2.1.6. Lentiviral-mediated stable MASTL silencing

2.1.6.1. Generation of lentiviral particles

For lentiviral particles generation, TRIPZ-shMASTL #1 (V2THS_177827, ThermoScientific, Dharmacon Inc., Chicago, IL, USA) or TRIPZ-shMASTL #2 (V2THS_261809, ThermoScientific, Dharmacon Inc., Chicago, IL, USA) plasmids were transfected in 293T cells by using the Trans-Lentiviral shRNA Packaging Kit (ThermoScientific, Dharmacon Inc., Chicago, IL, USA) according to manufacturer’s instruction. In details, 293T (5.5x10^6) cells were plated in 100mm dishes and transfected the following day with a mixture of 42 µg of TRIPZ-shMASTL #1 or TRIPZ-shMASTL #2 plasmids, together with 30 µl of Trans-lentiviral packaging mix, by calcium phosphate protocol. After 16 hours, cells were washed and incubated with DMEM, supplemented with 5% FBS, for 48 hours. Supernatant containing lentiviral particles was then collected, centrifuged at 1600 x g at 4°C for 10 minutes, filtered through a sterile 0.45 µM filter (Millipore, Billerica, MA, USA), aliquoted and stored at -80°C. Lentiviral particles were titered according to manufacturer’s instructions.

2.1.6.2. Lentiviral transduction

8505C (6x10^4) cells were seeded in 6-well plates and, the day after, growth medium was replaced with lentiviral particles-containing supernatant (multiplicity of infection, MOI=1), in presence of 6 µl/ml polybrene (Sigma-Aldrich, St. Louis, MO, USA), for 4 hours. After 4 hours incubation, supernatant was replaced with fresh lentiviral supernatant in presence of 6 µl/ml polybrene; 4 hours later supernatant was replaced with complete growth medium. After 48 hours of culture, 1 µg/ml puromycin (Sigma-Aldrich, St. Louis,
MO, USA) was added to medium, in order to select transduced cells. After one-week selection, cells were tested for shRNA production by 1 µg/ml and 2 µg/ml doxycycline (Sigma-Aldrich, St. Louis, MO, USA) administration.

2.2. Biochemical analysis

2.2.1. Protein extraction and sample preparation

Both adherent and floating cells were collected and centrifuged. Proteins were extracted in RIPA modified buffer (20mM Tris-HCl, pH 7.4, 150mM NaCl, 5mM EDTA, 1% NonidetP-40) supplemented with Complete Mini EDTA-free protease Inhibitor Cocktail (Roche, Manheim, Germany), 1mM Na$_3$VO$_4$ and 1mM PMSF. After incubation at 4°C for 20 minutes, samples were centrifuged at 13200 rpm for 30 minutes at 4°C and supernatant with proteins was then recovered. Total amount of protein for each sample was determined by Bradford protein assay (Bio-rad, Hercules, CA, USA) and a total amount of 30-40 µg protein extracts were boiled in NuPAGE LDS sample buffer (Invitrogen, Carlsbad, CA, USA) and then subjected to western blot analysis. Of note, for siCOPZ1-BCPAP and siCOPZ1-TPC-1 cells assayed for trypan blue exclusion test in chapter 3, the entire protein extract was subjected to western blot, as the amount of proteins recovered was less than 20 µg.

2.2.2. Western blot analysis

Protein samples were separated on 4-12% or 10% NuPAGE Novex Gel (Invitrogen, Carlsbad, CA, USA) in MOPS running buffer, under reducing conditions, and resolved proteins were then transferred onto nitrocellulose membranes (GE Healthcare Bio-Sciences, Pittsburgh, PA) using XCell II Blot Module (Invitrogen, Carlsbad, CA, USA). After blocking with 2% bovine serum albumin (BSA) (Sigma-Aldrich, St. Louis, MO, USA) in Tris-
Buffered Saline and Tween 20 (TBS-T [50mM Tris, 150mM NaCl, 0.1% Tween 20]), membranes were incubated overnight at 4°C with primary antibodies, listed in Table 2.3. Antibody binding was assessed by horseradish peroxidase (HRP)-conjugated secondary antibodies (1:5000, GE Healthcare Bio-Sciences, Pittsburgh, PA). Immunoreactive bands were visualized by enhanced chemiluminescence (GE Healthcare Bio-Sciences, Pittsburgh, PA). Densitometric analysis of the bands was performed by using Quantitative One software (Bio-rad, Hercules, CA, USA).

<table>
<thead>
<tr>
<th>Table 2.3 List of antibodies used for western blot analysis</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Actin</strong></td>
</tr>
<tr>
<td>Cdc2</td>
</tr>
<tr>
<td>cleaved PARP (Asp214)</td>
</tr>
<tr>
<td>cleaved-caspase 3 (Asp175)</td>
</tr>
<tr>
<td>COPE</td>
</tr>
<tr>
<td>COPZ1</td>
</tr>
<tr>
<td>Cyclin D1</td>
</tr>
<tr>
<td>gamma-H2AX</td>
</tr>
<tr>
<td>GAPDH</td>
</tr>
<tr>
<td>MAP4K5</td>
</tr>
<tr>
<td>MASTL</td>
</tr>
<tr>
<td>NUDT9</td>
</tr>
<tr>
<td>phospho-cdc2 (Tyr15)</td>
</tr>
<tr>
<td>PLA2G15</td>
</tr>
<tr>
<td>RASD1</td>
</tr>
<tr>
<td>RGS3</td>
</tr>
<tr>
<td>Vinculin</td>
</tr>
</tbody>
</table>
2.3. Molecular biology

2.3.1. RNA extraction, RT-PCR and real time PCR

RNA extraction was performed using NucleoSpin® RNA II extraction kit (Macherey-Nagel GmbH, Düren, Germany). 1 µg of RNA was retro-transcribed using SuperscriptIII (Invitrogen, Carlsbad, CA, USA) with random examers. PCR products were separated on 2% agarose gels, stained with ethidium bromide and visualized under UV transillumination. Primers used for the detection of RGS3, NUDT9, RASD1 and OXTR are listed in Table 2.4. GAPDH was used as housekeeping gene. For Real Time PCR analysis, 20 ng of retrotranscribed RNA were amplified in PCR reactions carried out in triplicate on an ABI PRISM 7900 using TaqMan gene expression assays (Applied Biosystem, Foster City, CA). Hs00212698_m1 was used for COPZ2 expression; human HPRT1 (HPRT-Hs99999909_A1) was used as housekeeping gene for the normalization among samples. Data analysis was performed using the SDS (Sequence Detection System) 2.4 software.

Table 2.4 List of primers used for RGS3, NUDT9, RASD1, OXTR and GAPDH detection by RT-PCR

<table>
<thead>
<tr>
<th>Gene</th>
<th>Primers</th>
<th>Sequence</th>
</tr>
</thead>
<tbody>
<tr>
<td>RGS3</td>
<td>forward</td>
<td>5′-ATGAGAGGGCTGTGGAGCACT-3′</td>
</tr>
<tr>
<td></td>
<td>reverse</td>
<td>5′-TCATGTGTGACTCGAAGGAG-3′</td>
</tr>
<tr>
<td>NUDT9</td>
<td>forward</td>
<td>5′-GACAGATCTAGCTAGCTGAA-3′</td>
</tr>
<tr>
<td></td>
<td>reverse</td>
<td>5′-TATAATGGATCTGACGTG-3′</td>
</tr>
<tr>
<td>RASD1</td>
<td>forward</td>
<td>5′-CGAGGTCTACCAGCTGACAT-3′</td>
</tr>
<tr>
<td></td>
<td>reverse</td>
<td>5′-CACGTCACCAGTTCTCCTG-3′</td>
</tr>
<tr>
<td>OXTR</td>
<td>forward</td>
<td>5′-ATGGAGCGGTCTGGAGATGC-3′</td>
</tr>
<tr>
<td></td>
<td>reverse</td>
<td>5′-GCTCAGGACAAAGGAGGACG-3′</td>
</tr>
<tr>
<td>GAPDH</td>
<td>forward</td>
<td>5′-ACCATCTTCAGGAGCGGACT-3′</td>
</tr>
<tr>
<td></td>
<td>reverse</td>
<td>5′-GGCAGAGATGATGACCTTT-3′</td>
</tr>
</tbody>
</table>
2.4. Gene expression analysis

The thyroid TCGA data set, which contains the molecular data of 496 PTC samples, is the largest data collection so far available (https://tcgadata.nci.nih.gov/tcga/tcgaCancerDetails.jsp?diseaseType=THCA&diseaseName=Thyroid). Illumina HiSeq Level 3 data were downloaded from TCGA Data Portal and RSEM (116) gene normalized expression values of selected genes were derived for 58 PTC patients, in tumor and adjacent normal samples, respectively. The statistical significance of gene expression difference between PTC and normal groups was assessed using Wilcoxon test.
3. Results: siRNA library screening hit validation

3.1. Introduction

The NOA paradigm (section 1.2.1) states that the uncontrolled growth of tumor cells relies on the activity of genes and pathways, not oncogenic themselves but whose activity is essential for tumor cell viability and not for normal ones (98). Genes presenting these features can be identified by loss-of-function RNAi-based genetic approaches that allow the generation of either “synthetic lethal signature”, against specific oncogenes and tumor suppressors, or “cancer lethal signature”, if using both normal and tumor cells for the screening. Several reports suggested that this approach was likely to shed new light on the mechanisms of tumor maintenance: myelomas were found to be addicted to an abnormal regulatory network controlled by IRF4 regardless the oncogenic lesion (117), while genes synthetic lethal with mutant KRAS were identified in several tumors harboring this mutation (118). In order to identify a “cancer lethal signature” in thyroid tumors, our laboratory performed a loss of function siRNA library screening on tumor (BCPAP, PTC-derived, carrying the BRAFV600E oncogene) and normal immortalized (Nthy-ori 3-1) thyroid cell lines (119). Thyroid cell lines were transfected with a siRNA library containing oligos targeting about 9000 potentially druggable genes and colony forming assay (CFA) was performed after 7 (for Nthy-ori 3-1 cells) and 8 (for BCPAP cells) days (Figure 3.1A and 3.1B). Non-targeting siRNA (siNT) and a siRNA targeting the proteasomal subunit PSMC3 (siPSMC3) were transfected as negative and positive controls, respectively. The screening identified genes whose silencing was capable of inhibiting the growth of both cell lines, referred to as “lethal hits”, but also genes whose silencing
selectively inhibited the growth of tumor cells, referred to as “differential hits” (Figure 3.1C and 3.1D).

Figure 3.1 siRNA library screening on normal and tumor thyroid cell lines
A) Schematic representation of siRNA library screening automated procedure; 309 mother 96-well plates, containing 25139 siRNA oligos targeting 9031 genes (three deconvoluted siRNAs for each gene), have been screened in duplicates in BCPAP and NThy-ori 3-1 cell lines. The effect of each siRNAs on the cell viability was analyzed at seven/eight days after transfection by a medium-high throughput CFA. B) Representative colony plates generated from transfection described in A); a indicates siRNA oligos lethal for both cell lines; b indicates siRNA oligos selectively lethal for BCPAP cells; siNT and siPSMC3 represent negative and positive controls, respectively. C) Scatter plot of Colony Formation (CF) values obtained from the screening, reported as % of growth respect to non targeting (siNT) for inter-plate data normalization (dots are the averages of two technical replicates performed in both cell lines). D) Scatter plot of normalized CF values showing the negative controls (siNT, blue), the lethal hits (green) and the differentially active hits (red); insert: distribution of normalized d values after sigma binning. Image adapted from (119).
Technical confirmation of differential hits was then performed by manual transfection of siRNAs on Nthy-ori 3-1 and BCPAP cells, with the same experimental conditions employed in the primary screening. A list of 15 confirmed hit genes, whose inhibition was able to inhibit the growth of tumor but not normal cell lines, was then generated (Table 3.1). Functional annotation analysis revealed that hit genes identified were involved in several biological processes, such as cell cycle and DNA damage checkpoint regulation, vesicular transport and intracellular signal transduction. The relevance and reliability of this screening in identifying TC cell vulnerabilities was given by two lines of evidence. First, among the confirmed hits we found BRAF; this was expected, as BCPAP cells are known to be addicted to BRAFV600E oncogene (120). Second, we identified genes whose silencing was lethal for both normal and tumor cell viability, and most of them were already identified to be essential survival genes by other RNAi screenings (121-123).
### Table 3.1 List of confirmed hits from the screening

The 15 hit genes, whose inhibition resulted to inhibit the growth of thyroid tumor cells, are listed. Representative Gene Ontology-Biological Process (GO-BP) terms are indicated for each of them, by means of DAVID6.7 functional annotation analysis.

<table>
<thead>
<tr>
<th>Gene Symbol</th>
<th>Full Gene Name</th>
<th>Accession</th>
<th>GO-BP Term</th>
</tr>
</thead>
<tbody>
<tr>
<td>MASTL</td>
<td>microtubule associated serine/threonine kinase-like</td>
<td>NM_032844</td>
<td>GO:0006468 protein amino acid phosphorylation&lt;br&gt;GO:0006793 phosphorus metabolic process</td>
</tr>
<tr>
<td>PLA2G15</td>
<td>phospholipase A2, group XV</td>
<td>NM_012320</td>
<td>GO:0006575 cellular amino acid derivative metabolic process&lt;br&gt;GO:0006843 membrane lipid metabolic process</td>
</tr>
<tr>
<td>COPE</td>
<td>coatamer protein complex, subunit epsilon</td>
<td>NM_007263</td>
<td>GO:0006890 retrograde vesicle-mediated transport, Golgi to ER transport&lt;br&gt;GO:0051640 organelle localization</td>
</tr>
<tr>
<td>CCND1</td>
<td>Cyclin D1</td>
<td>NM_053056</td>
<td>GO:0000075 cell cycle checkpoint&lt;br&gt;GO:0000777 DNA damage checkpoint</td>
</tr>
<tr>
<td>RASD1</td>
<td>RAS, dexamethasone-induced 1</td>
<td>NM_016084</td>
<td>GO:0007264 small GTPase mediated signal transduction&lt;br&gt;GO:0016481 negative regulation of transcription</td>
</tr>
<tr>
<td>NUDT9</td>
<td>nudix (nucleoside diphosphate linked moiety X)-type motif 9</td>
<td>NM_024047</td>
<td>GO:0006163 purine nucleoside metabolic process&lt;br&gt;GO:0006811 ion transport</td>
</tr>
<tr>
<td>OXTR</td>
<td>oxytocin receptor</td>
<td>NM_000916</td>
<td>GO:0000165 MAPKK cascade&lt;br&gt;GO:0003012 muscle system process</td>
</tr>
<tr>
<td>BRAF</td>
<td>v-raf murine sarcoma viral oncogene homolog B1</td>
<td>NM_004333</td>
<td>GO:0000165 MAPKK cascade&lt;br&gt;GO:0010941 regulation of cell death</td>
</tr>
<tr>
<td>COPZ1</td>
<td>coatamer protein complex, subunit zeta 1</td>
<td>NM_016057</td>
<td>GO:0006866 intracellular protein transport&lt;br&gt;GO:0006890 retrograde vesicle-mediated transport, Golgi to ER</td>
</tr>
<tr>
<td>MAP4K5</td>
<td>mitogen-activated protein kinase kinase kinase 5</td>
<td>NM_006575</td>
<td>GO:0000165 MAPKK cascade&lt;br&gt;GO:001932 regulation of protein amino acid phosphorylation</td>
</tr>
<tr>
<td>EPHB4</td>
<td>EPH receptor B4</td>
<td>NM_004444</td>
<td>GO:0001769 transmembrane receptor protein tyrosine kinase signaling pathway&lt;br&gt;GO:0045765 regulation of angiogenesis</td>
</tr>
<tr>
<td>DNM3</td>
<td>dynamin 3</td>
<td>NM_015569</td>
<td>GO:0006897 endocytosis&lt;br&gt;GO:0010324 membrane invagination</td>
</tr>
<tr>
<td>REM2</td>
<td>RAS (RAD and GEM)-like GTP binding 2</td>
<td>NM_173527</td>
<td>GO:0006355 regulation of transcription, DNA-dependent&lt;br&gt;GO:0007264 small GTPase mediated signal transduction</td>
</tr>
<tr>
<td>SRPK1</td>
<td>SFRS protein kinase 1</td>
<td>NM_003137</td>
<td>GO:0006396 RNA processing&lt;br&gt;GO:0007243 protein kinase cascade</td>
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</table>
3.2. Aims of the chapter

Functional siRNA library screening identified a number of genes whose silencing interfered selectively with tumor cell viability and, among those, 15 hit genes were confirmed by technical validation. The aims of this chapter are: to identify the molecular profile of selected hits in thyroid cancer; to validate the growth inhibitory effect of these genes by using different siRNA sequences in the two cell models used for the screening; to correlate the growth inhibitory effect with the extent of target gene knockdown; to extend the analysis of the growth inhibitory effect of their silencing on a large panel of thyroid tumor cell lines, representative of the different histotypes.
3.3. Results

3.3.1. Analysis of expression levels of hit genes in normal and PTC samples

In 2014, the TCGA consortium published an important study that improved our understanding of the genomic landscape of PTC (33) (section 1.1.2.1). The study reported analysis of the largest PTC cohort ever analyzed to date, with 58 out of 496 PTC specimens paired with the corresponding normal counterpart. We interrogated the thyroid TCGA dataset with the aim of assessing the expression level and the mutational status of the 15 genes identified by our functional screening. Expression analysis was investigated on both the whole dataset and on the 58 matched tumor and normal samples, and the results obtained were identical. By comparing normal and PTC samples, we classified hit genes in three groups, according to their expression levels (Figure 3.2): i) genes with significant high overexpression in PTC vs normal thyroid, which are CCND1, RGS3, OXTR, RASD1, DNM3 (panel A); ii) genes with equal or slightly different expression in PTC compared to normal thyroid, that are COPE, COPZ1, PLA2G15, SRPK1, REM2, EPHB4, BRAF (panel B); iii) genes that are significantly downregulated in PTC compared to normal thyroid, which are MASTL, MAP4K5, NUDT9 (panel C).
Figure 3.2 Hit gene expression levels in PTC vs normal thyroid specimens
Box plot distribution of RNAseq normalized expression values of the 15 hit genes from the thyroid TCGA dataset analysis in 58 matched normal (N) and PTC samples. A) Genes overexpressed in PTC vs N samples. B) Genes equally expressed in PTC and N samples. C) Genes downregulated in PTC vs N sample. Statistical significance of differences among the two groups of samples was assessed by Wilcoxon test.

For mutational status, we detected BRAF mutations with high frequency, as present in 248 out of 496 samples. With the exception of a silent mutation for COPZ1 in one sample, no mutations affecting the other genes were detected in the whole dataset.

We also retrieved data from the Cancer Cell Line Encyclopedia (CCLE, http://www.broadinstitute.org/ccle/home), which provides public access to genomic data, analysis and visualization for about 1000 cell lines, among which the BCPAP cell line.
We found that all the hit genes are expressed by BCPAP cells, at variable level, ranging from “low expression” to “very high expression” (Table 3.2). *CCND1*, *MASTL*, *MAP4K5*, *SRPK1* and *EPHB4* were found to be at the wild type status, while no data for the mutational status of the other hit genes were available.

### Table 3.2 Expression level of hit genes in BCPAP cells from CCLE

<table>
<thead>
<tr>
<th>Gene</th>
<th>Value</th>
<th>Expression level</th>
</tr>
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<tbody>
<tr>
<td>MASTL</td>
<td>5,2564</td>
<td>low</td>
</tr>
<tr>
<td>PLA2G15</td>
<td>6,7379</td>
<td>moderate</td>
</tr>
<tr>
<td>COPE</td>
<td>10,841</td>
<td>very high</td>
</tr>
<tr>
<td>RGS3</td>
<td>6,5896</td>
<td>moderate</td>
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<tr>
<td>CCND1</td>
<td>9,7431</td>
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</tr>
<tr>
<td>RASD1</td>
<td>6,843</td>
<td>moderate</td>
</tr>
<tr>
<td>NUDT9</td>
<td>8,7534</td>
<td>high</td>
</tr>
<tr>
<td>OXTR</td>
<td>4,5271</td>
<td>low</td>
</tr>
<tr>
<td>BRAF</td>
<td>6,8071</td>
<td>moderate</td>
</tr>
<tr>
<td>COPZ1</td>
<td>10,779</td>
<td>very high</td>
</tr>
<tr>
<td>MAP4K5</td>
<td>8,6024</td>
<td>high</td>
</tr>
<tr>
<td>EPHB4</td>
<td>5,9189</td>
<td>low</td>
</tr>
<tr>
<td>DNM3</td>
<td>4,2194</td>
<td>low</td>
</tr>
<tr>
<td>REM2</td>
<td>6,0285</td>
<td>moderate</td>
</tr>
<tr>
<td>SRPK1</td>
<td>8,5579</td>
<td>high</td>
</tr>
</tbody>
</table>

#### 3.3.2. Analysis of hit gene expression levels in TC cell lines

In order to assess whether these putative “non-oncogenes” are expressed by TC cell lines, we first searched for and purchased commercially available antibodies for western blot analysis for the first 11 hit genes except for *BRAF*. Of note, for some of these genes (*PLA2G15, RGS3, RASD1, NUDT9, OXTR, COPZ1*), no published working antibodies were available at the time of the analysis. We first tested the optimal working conditions for the antibodies purchased by western blot analysis of control cell lines indicated by the antibody datasheets. As reported in Figure 3.3A-F, we were successful in detecting the
expression of MASTL, PLA2G15, COPE, MAP4K5, Cyclin D1 and COPZ1. For RGS3, NUDT9 and RASD1 (Figure 3.3G-I), the antibodies tested did not allow the detection of the corresponding protein.

We investigated the expression of the hit genes in the normal (Nthy-ori 3-1) and tumor (BCPAP) cell lines used for the screening and in a panel of thyroid tumor cell lines, representative of the different histotypes and/or carrying different oncogenes: PTC-
derived (TPC-1, carrying the \textit{RET}/\textit{PTC1} oncogene), FTC-derived (WRO82-1), ATC-derived (8505C and HTC/C3, which carry \textit{BRAFV600E} oncogene, KAT-18). MASTL, Cyclin D1, COPZ1, MAP4K5, PLA2G15 and COPE expression was investigated by western blot (Figure 3.4A). In comparison with Nthy-ori 3-1, MASTL was found to be less expressed in thyroid tumor cell lines; Cyclin D1 expression, except for KAT-18, was higher in the tumor cell lines compared to the normal one; also COPZ1 expression was higher in TC cell lines, compared to Nthy-ori 3-1; PLA2G15 expression was slightly reduced in all the tumor cell lines, except for BCPAP, and strongly reduced in KAT-18; MAP4K5 was found to be expressed at variable levels, with a slight increase in BCPAP, WRO82-1 and 8505C, and a decrease in the remaining; COPE expression in tumor cells was higher compared to normal Nthy-ori 3-1, with the exception of KAT-18. The expression of \textit{RGS3}, \textit{NUDT9}, \textit{RASD1} and \textit{OXTR} was investigated by RT-PCR. All the genes are expressed in the cell lines tested with no appreciable differences (Figure 3.4B).
3.3.3. Analysis of siRNA silencing efficiency for hit validation

Among the list of candidate “non-oncogenes” described above, we selected for validation studies only those genes for which western blot analysis was feasible. We purchased commercially available siRNAs, with oligonucleotide sequences different from those of the library, for MASTL, PLAD2G15, COPZ1, CCND1, COPE and MAP4K5. Four different oligos for MASTL and PLAD2G15, and a smart pool of oligos for CCND1, COPZ1, MAP4K5 and COPE, were used. Nthy-ori 3-1 cells were transiently transfected with the
specific siRNAs and with non-targeting control (siNT) oligos and then subjected to western blot analysis, in order to evaluate silencing efficiency (Figure 3.5). For MASTL, we obtained the highest silencing efficiency with oligos #1 and #4 72 hours after transfection; oligos targeting Cyclin D1, COPZ1 and MAP4K5 completely abrogated protein expression at 72 hours. For PLA2G15 and COPE, siRNA oligos were inefficient or slightly efficient in abrogating protein expression, respectively. Notably, oligos with good silencing efficiency gave no macroscopic effects on the growth and survival of Nthy-ori 3-1 cells, in keeping with what obtained from siRNA library screening.

**Figure 3.5 Evaluation of siRNA silencing efficiency for validation studies**
Western blot analysis of MASTL, COPZ1, PLA2G15, MAP4K5, Cyclin D1 and COPE expression in Nthy-ori 3-1 cells, transiently transfected with the indicated siRNAs and collected at the indicated timepoints. Actin and vinculin represent loading controls.
3.3.4. Effects of MASTL, COPZ1 and Cyclin D1 depletion on BCPAP cell growth

Based on the silencing efficiency of siRNAs previously tested and on background literature data about the hit genes, we selected MASTL, COPZ1 and Cyclin D1 for the validation studies. We transfected siRNAs in the two cell lines used for the screening, Nthy-ori 3-1 and BCPAP. We used the same protocol as for the screening: cells were transfected in 96-well plates with the oligos described in section 3.3.3 and siNT and siPSMC3 controls, and cell growth was evaluated 10 days later by CellTiter-Glo® assay. The luminescence signal, indicative of viable cells, is reported in the scatter plot in Figure 3.6A, normalized with respect to siNT. We observed that siMASTL #1, siMASTL #4, siCOPZ1 and siCCND1 reduced BCPAP cell growth by 60%, 93%, 94% and 86%, respectively; this efficiency is comparable to that of positive control siPSMC3, that is 93% reduction. We did not observe any significant reduction of the growth of BCPAP cells transfected with siMASTL #2 and siMASTL #3; this result is in keeping with the low siRNA silencing efficiency we previously observed in Nthy-ori 3-1 (Figure 3.5). Of note, no significant reduction of Nthy-ori 3-1 cell growth was observed with MASTL-, COPZ1- and CCND1-targeting oligos, while the positive control siPSMC3 was able to reduce cell growth as for BCPAP. We further confirmed the effect of MASTL, COPZ1 and CCND1 silencing with a different experimental approach. BCPAP cells were transfected in 6-well plates with the oligos that previously resulted to be efficient in gene silencing: siMASTL #1 and #4, siCOPZ1 and siCCND1. We also used an additional in-house synthesized siRNA targeting MASTL, that was found to be efficient in MASTL silencing when transfected in Nthy-ori 3-1 (Figure 3.6B). At 6 days after transfection, we noticed an evident effect of MASTL, COPZ1 and CCND1 silencing by microscopy observation: BCPAP cells were less confluent compared to negative control (siNT), with a large number of floating cells (Figure 3.6C). By trypan blue exclusion test, we found a reduction of cell growth, in comparison with
control (siNT), of: 69.5%, 59.7% and 71% for siMASTL #1, #4 and #5, respectively; by 99.6% for siCOPZ1; by 91.5% for siCCND1 (Figure 3.6D). The same cells were tested by western blot at the end of the experiment and complete abrogation of MASTL and Cyclin D1 expression was observed; an insufficient amount of protein extract was recovered from siCOPZ1-transfected cells, due to the massive lethal effect on this cell line, so western blot analysis was not informative (Figure 3.6E).

On the whole, our results confirmed the growth inhibitory effect of MASTL, COPZ1 and Cyclin D1 knockdown on BCPAP cells, thus validating them as vulnerability genes of BCPAP.
Figure 3.6 Effects of MASTL, COPZ1 and Cyclin D1 depletion on BCPAP cell line

A) Scatter plot representing the percentage of cell growth (luminescence values of CellTiter-Glo® assay), normalized to control (siNT), of Nthy-ori 3-1 and BCPAP cells, 10 days after transfection with the indicated siRNAs. One representative out of three experiments is shown. B) Western blot analysis of MASTL expression in Nthy-ori 3-1 cells, transiently transfected with the control (siNT) and the new MASTL (siMASTL #5) siRNAs, and collected at the indicated time points. Actin represents loading control. C) Representative pictures of BCPAP cells 6 days after transfection with the indicated siRNAs. D) BCPAP cell growth evaluated by trypan blue exclusion test 6 days after transfection; cell count of viable cells was normalized to that of control (100% of siNT); data represent the mean +/- sd of two independent experiments; asterisks indicate statistical significant differences by unpaired Student’s t-test (***p<0.001); E) Western blot analysis of MASTL, Cyclin D1 and COPZ1 expression in BCPAP cells, assayed in panel D; actin was used as loading control.
3.3.5. Effects of MASTL, COPZ1 and Cyclin D1 depletion on the growth of several TC cell lines

According to NOA paradigm, tumor cells may share common vulnerabilities based on the stress support pathways needed for sustaining the tumorigenic phenotype, regardless the main oncogenic drivers. We decided to assess whether the survival genes previously confirmed in BCPAP may be involved in such common pathways and therefore represent common vulnerabilities for thyroid tumors. To this aim, we evaluated the functional effects of MASTL, COPZ1 and Cyclin D1 silencing on the panel of TC cell lines in which we have previously determined their expression levels (see Figure 3.4). TC cell lines were transfected in 96-well plates with control (siNT and siPSMC3) siRNAs and siRNAs targeting MASTL, COPZ1 and CCND1, and cell viability was evaluated by CellTiter-Glo® assay 10 days later. We found that silencing of the three genes induced variable extents, ranging from 15% to 95%, of cell growth inhibition (Figure 3.7A) in the majority of cell lines tested, compared to control. Only for CCND1 in 8505C and KAT-18, and for MASTL in KAT-18 cell lines, we did not observe any effect of gene silencing on cell viability. We additionally assessed the effects on cell growth by performing trypan blue exclusion test from 4 to 7 days after cell transfection in 6-well plates, as already performed on BCPAP (detailed previously in 3.3.4). A reduction in the growth of cells upon transfection of siRNAs targeting MASTL, COPZ1 and CCND1, compared to control (siNT) transfected cells, was appreciable by microscopy observation (Figure 3.7B). Cell count revealed that, with the exception of KAT-18 cell lines, silencing of MASTL, COPZ1 and CCND1 significantly reduced the growth of all cell lines tested (Figure 3.7C). Compared to control, inhibition of cell viability was observed by: 30-63% for siMASTL #1, 35-75% for siMASTL #4, 40-75% for siMASTL #5, 75-99% for siCOPZ1 and 70-94% for siCCND1. In KAT-18, we found a decrease of cell growth by 31% for siMASTL #1, 53% for siCOPZ1 and 43% for CCND1; siMASTL #4
and #5 had no significant effect on cell viability; PSMC3 silencing in KAT-18 had a growth inhibitory effect of 53%. We performed western blot analysis on the same samples at the end point of the experiment and observed a high efficiency of gene silencing in all of them (Figure 3.7D). As for BCPAP cells, also in TPC-1 cells we were not able to recover enough protein extract to assess COPZ1 knockdown in siCOPZ1-transfected samples, due to the massive lethal effect of gene silencing on this cell line. Of note, by this alternative approach we were able to notice a growth inhibitory effect of CCND1 silencing in 8505C and KAT-18, and of MASTL silencing in KAT-18 cells, which was not previously observed; this could be related to the variability of silencing efficiency and/or experimental conditions.
Figure 3.7 Effects of MASTL, COPZ1 and Cyclin D1 depletion on a panel of TC cell lines

A) Scatter plot representing the percentage of cell growth (luminescence values of CellTiter-Glo® assay), normalized to control (siNT), of the indicated TC cell lines 10 days after transfection with the indicated siRNAs. One representative out of three experiments is shown. B) Representative images of TPC-1, WRO82-1, 8505C, HTC/C3 and KAT-18 cells, transfected with the indicated siRNAs, before the trypan blue exclusion test. D) Cell proliferation assayed by trypan blue exclusion test 4 days (for TPC-1, 8505C and HTC/C3 cells), 5 days (for KAT-18 cells), 6 days (for BCPAP cells) and 7 days (for WRO82-1 cells) after siRNA transfection; cell count was normalized to that of siNT (100% growth) and expressed as mean +/- sd of two independent experiments; statistically significant differences are indicated by asterisks (ns: not significant; *p<0.05; **p<0.01; ***p<0.001; according to unpaired Student's t-test). E) Western blot analysis of MASTL, COPZ1 and Cyclin D1 expression in TC cells assayed in panel D; actin was used as loading control.

3.3.6. Characterization of TC cell dependency on CCND1, MASTL and COPZ1

In order to explore the potential of MASTL, Cyclin D1 and COPZ1 targeting in TC we performed additional experiments.

Given that Cyclin D1 acts as activator of its dependent kinases CDK4 and CDK6, we tested whether TC cell lines may be susceptible to palbociclib (PD-0332991), a specific inhibitor of CDK4 and 6 (115). The panel of TC cell lines previously tested was treated with palbociclib for 72 hours. The effects of palbociclib administration were appreciable by microscopy observation: after 72 hours, all TC cells treated with palbociclib were less confluent compared to the control (CTRL) (Figure 3.8A). The treatment with increasing doses of palbociclib showed a consistent growth inhibitory effect on all TC cell lines tested (Figure 3.8B). The IC50 values were variable, ranging from <0.003 µM for BCPAP cells to 0.20 µM for HTC/C3 cells (Figure 3.8C).

Since it was reported that MASTL depletion induced multiple mitotic defects (124), we tested whether MASTL-depleted 8505C cells could present the same defects. 48 hours after MASTL silencing in 8505C, immunofluorescence analysis for β-tubulin, phosphorylated histone H3 (Ser10) and DAPI revealed the presence of aberrant mitotic figures (Figure 3.8D): while no anomalies were observed in control siNT (i), MASTL
knockdown was associated with formation of multinuclear cells (ii), lagging chromosomes not aligned to metaphase plate (iii) and chromatin bridges connecting daughter cells (iv).

COPZ1 vulnerability of tumor cells was found to be associated with the downregulation of its paralogous COPZ2 (125). We performed real time PCR in normal (Nthy-ori 3-1) and TC cell lines to assess COPZ2 expression level (Figure 3.8E): we found that all tumor cells, except for WRO82-1, showed a downregulation of COPZ2 mRNA, compared to normal thyrocytes.
Figure 3.8 Characterization of Cyclin D1, MASTL and COPZ1 vulnerabilities of TC cells
A) Representative ArrayScan fields of TC cell lines untreated (CTRL) or treated with 2.2 μM palbociclib for 72 hours and stained with Hoechst (scalebar: 50µm). B) Dose-response curve of TC cell growth treated with palbociclib for 72 hours; nuclei were stained with Hoechst and counted in each well by an ArrayScan high-content screening reader; cell count values derived from nuclear staining (reported as growth parameter normalized with respect to untreated controls) are presented as mean +/- sd of two replicates. C) IC50 values (µM) obtained using a sigmoid function of nonlinear interpolation of experimental points; for KAT-18, IC50 value was derived from graphical analysis. D) Immunofluorescence analysis for β-tubulin (microtubules, green), phosphorylated histone H3 Ser10 (red) and DAPI (nuclei, blue) on 8505C cells 48 hours after transfection with control (siNT) and MASTL (siMASTL) siRNAs; arrows indicate normal mitosis (i), multinuclear cell (ii), lagging chromosomes (iii) and DNA bridge (iv). E) Real time PCR analysis of COPZ2 expression; results are presented as log10-transformed relative quantity (RQ) of COPZ2 mRNA normalized for HPRT housekeeping expression and presented as mean +/- sd of three independent experiments.
3.4. Discussion

Our laboratory performed a loss of function siRNA-based screening on normal and tumor thyroid cell lines, with the aim of identifying potential “non-oncogenes” in thyroid tumor cells. A list of 15 genes, whose silencing was detrimental for tumor but not normal cell viability, was generated. We analyzed their molecular profiles in normal and PTC samples by TCGA dataset interrogation and found, with the exception of BRAF, no mutations of hit genes in PTC samples, while their expression level was equal or differential (both upregulation and downregulation) between normal and tumor specimens. We documented their expression in a panel of thyroid tumor cell lines and selected MASTL, COPZ1 and CCND1 for validation studies. We confirmed their inhibitory effect on BCPAP cell growth, by using siRNA oligos different from those of the library, and verified the abrogation of their expression at protein level in BCPAP cells. Importantly, we observed that their depletion resulted to be detrimental for the growth and survival of several thyroid tumor cell lines, representative of the different histotypes and carrying different oncogenic lesions.

Molecular studies performed in the last years extensively characterized the genomic landscape of PTCs and ATCs. These findings were translated into the use of small-molecule inhibitors, specifically TKIs, in the clinics for the treatment of advanced TC patients who do not respond to conventional therapy (126); even though TKIs showed promising therapeutic strategies, their efficacy gave contrasting evidence in clinical trials, since mechanisms of resistance to TKIs often develop and severe side effects present (127). This evidence underlines the need for alternative approaches aimed to identify novel targets for TC treatment. Large scale RNAi-based functional screenings are a tool nowadays widely used to identify tumor cell vulnerabilities for potential
therapeutic goals (106,107,110). Indeed, these high-throughput screening approaches permit to discover potential target genes to which cancer cells are addicted and that are not necessarily oncogenic themselves. In this chapter, we presented the results of a high-throughput siRNA phenotypic screening on normal and tumor thyroid cells, aimed to identify thyroid tumor cell vulnerabilities (119). For the first time, we identified a panel of genes whose silencing preferentially inhibited the growth of the tumor cell line as compared to that of normal thyrocytes. TCGA dataset analysis revealed no mutations of hit genes (besides BRAF); this suggests that alterations of their function in thyroid tumor cell biology may be excluded, and that they represent “non-oncogenes” in TC. We verified that these genes were actually expressed in the cell models used for the screening and also extended the analysis to a panel of thyroid tumor cell lines; this first approach allowed us to select some of them for validation studies, according to the availability of reagents such as commercial antibodies efficiently working. For hit genes whose expression was not ascertainable by western blot analysis, due to antibodies not efficiently working, we assessed their mRNA expression in TC cell lines by RT-PCR; since western blot was not feasible for them, they were excluded from validation studies. To validate the remaining hits, we first tested siRNA oligos different from those of the library and found that some of them were actually not efficient, thus making further investigation not feasible. We selected MASTL, COPZ1 and CCND1 for the confirmation studies in BCPAP.

Cyclin D1 is a cell cycle regulator that forms a complex with cyclin-dependent kinases (CDK) 4 and 6 to activate them and promote the transition from G1 to S phase of the cell cycle (128). CCND1 is frequently found deregulated in tumors: gene mutations and amplifications, as well as protein overexpression, are events frequently reported. For this reason, Cyclin D1 has been proposed as therapeutic target for cancer treatment. Its
targeting may result difficult since it lacks enzymatic activity; a more feasible approach would be the inhibition of its associated kinases CDK4 and CDK6. Indeed, several CDK4/6 inhibitors are currently under clinical evaluation in different tumors (129). Cyclin D1 was frequently found overexpressed at both protein (130) and mRNA (131,132) levels in TC, but no functional data about its role in thyroid tumorigenesis were reported. We demonstrated that Cyclin D1 inhibition impaired the growth of all TC cell lines tested. Interestingly, the expression level of Cyclin D1 was heterogeneous among them; this finding suggests that TC cell dependency on CCND1 may not require the aberrant expression of the protein. Moreover, we observed that TC cell lines are sensitive to the CDK4/6 inhibitor palbociclib, that is already used for the treatment of ER+/HER2- breast cancer in combination with letrozole (133). Our results prompt the evidence that Cyclin D1 represents a TC cell vulnerability and provide a rationale for further preclinical and clinical studies aimed to develop a therapeutic strategy to inhibit it, directly or indirectly by using CDK4/6 inhibitors, for TC treatment.

MASTL (microtubule associated serine/threonine kinase-like) is a regulator of mitosis as its activity is required for the inhibition of PP2A/B55δ, the main protein phosphatase complex responsible for dephosphorylation of CDK1 substrates. MASTL activity was shown to be necessary to prevent defects in chromosome segregation and avoid prometaphase arrest and mitotic failure (134). Little was known about the role of MASTL in cancer. In lung cancer, it was reported that MASTL inhibition was able to sensitize tumor cells to radiations and to promote cytokinesis defects (135). Another study (136) reported that MASTL has a role in head and neck tumor recurrence and resistance to chemotherapy with cisplatin, thus proposing it as possible therapeutic target. They also found MASTL to be overexpressed in prostate, breast and recurrent head and neck tumors, compared to normal counterpart. In this chapter, we identified MASTL as a
vulnerability gene for thyroid tumors. We observed that its inhibition interfered with the growth of TC cell lines, with variable extents. We also found that MASTL-depleted 8505C exhibited errors in the mitotic process. Moreover, we found no mutations in PTC samples and observed that MASTL is not associated with overexpression in PTC compared to normal samples. Together, these findings highlighted that further investigations about the role of MASTL in TC cell biology were needed, in order to assess whether it could represent a therapeutic target for TC. These studies are presented in the next chapter of the thesis together with a more deepened discussion.

COPZ1 (coatamer protein complex ζ1) is the zeta-subunit of the secretory coatamer protein complex I (COPI), which is involved in vesicular trafficking from Golgi apparatus to endoplasmic reticulum, endosome maturation, autophagy (137,138) and lipid homeostasis (139). Shtutman and colleagues (125) proposed COPZ1 as therapeutic target as they observed that tumor cells become dependent on COPZ1 because of downregulation of COPZ2 and COPZ1 silencing dramatically induced apoptotic cell death. In this chapter, we showed that COPZ1 is a TC vulnerability gene as its silencing led to growth inhibition of all TC cell lines tested. We also found COPZ2 downregulation in the majority of TC cell lines, apart from WRO82-1: even if COPZ2 mRNA level was higher than control, this cell line resulted to be sensitive to COPZ1 depletion; this finding suggests that other mechanisms may be involved in determining COPZ1 sensitivity in TC cells. Further experiments were needed to better understand the mechanisms by which COPZ1 affects TC cell viability, and are presented in an independent chapter of the thesis.

In conclusion, in this chapter we described the identification of a “cancer lethal signature” for thyroid carcinoma, as result of a siRNA-based functional screening on normal and tumor thyroid cell lines. Among the novel potential therapeutic targets that emerged, we functionally characterized Cyclin D1, MASTL and COPZ1 and demonstrated
that their depletion was deleterious for TC cell growth irrespective of tumor histotype and/or genetic lesion. These findings indicate that they represent common examples of “non-oncogene addiction” in thyroid cancer, highlighting them as attractive targets for new therapeutic approaches for thyroid tumors which would spare normal cells, thus having limited side effects.
4. Results: MASTL studies

4.1. Introduction

Microtubule associated serine/threonine kinase-like (MASTL) is the human orthologue of greatwall (GWL), a gene first discovered in Drosophila in 2004, which encodes a kinase involved in the regulation of mitosis (140). The name “greatwall” came from the evidence that mutations in this gene caused defects in chromosome condensation and delay in mitotic progression, thus suggesting that it had a role in protecting chromosome structure. A following study in Xenopus egg extracts revealed that GWL was involved in the complex regulatory feedback loop that drives the phosphorylation of proteins in mitosis progression (141). Central to this complex regulatory loop is cell division cycle protein 2 homolog (cdc2), also known as cyclin-dependent kinase 1 (CDK1): it encodes a serine/threonine kinase that has the essential role of promoting G2/M transition and mitosis entry, once bound to Cyclin B1, thus forming the maturation promoting factor (MPF) complex (142). CDK1 activation is tightly controlled during cell cycle, in order to ensure correct progression through mitosis: during G1, S and G2 phases, MYT1 and WEE1 kinases phosphorylate CDK1 on Thr14 and Tyr15 inhibitory residues, in order to maintain it in an inactive state; at the onset of mitosis, CDC25 phosphatase removes these inhibitory modifications, allowing the activation of CDK1 and, consecutively, of MPF complex (Figure 4.1). Active CDK1 is able to further activate CDC25, as well as to inactivate its inhibitor WEE1. Furthermore, it also phosphorylates polo-like kinase 1 (PLK1), which contributes to a further activation of CDC25. Yu and colleagues showed that, during mitosis, GWL was phosphorylated and activated by CDK1, and GWL depletion
led to accumulation of inhibitory phosphorylations on CDK1 that prevented mitosis progression. It was later demonstrated that GWL contributed to mitosis progression by inhibiting the protein phosphatase 2A (PP2A) associated with the B55δ regulatory subunit (PP2A/B55δ), which is the main phosphatase complex responsible for the dephosphorylation of mitotic substrates of the MPF complex (143). This inhibition was shown to be mediated by two proteins identified as GWL substrates, namely cyclic adenosine monophosphate–regulated phosphoprotein 19 (ARPP19) and α-Endosulfine (ENSA); once activated by GWL, they associated with PP2A/B55δ and inhibited it, thus promoting mitotic entry (144,145). Until recently, no specific regulators of GWL were identified; only the work from Ma and colleagues (146) suggested that protein phosphatase 1 (PP1) could be responsible for GWL inactivation.
Figure 4.1 Regulatory network model of GWL during mitosis entry and exit
At G2/M transition, CDK1/Cyclin B1 complex and GWL are activated. CDK1/CycB1 phosphorylates mitotic substrates and GWL, which contributes to maintenance of mitotic substrate activation by inhibiting PP2A/B55δ complex through ARPP19/ENSA activation. PP2A/B55δ inhibition fully activates mitotic substrates and promotes entry in mitosis. At mitotic exit, Cyclin B1 is ubiquitinated by APC-CDC20 ubiquitin ligase and degraded; GWL is therefore inactivated and PP2A/B55δ consequent activation removes activating phosphorylations from mitotic substrates. Image from (147).

The functional orthologue of GWL in humans is MASTL, a gene that shares 50.2% and 65.7% of sequence homology to Drosophila and Xenopus GWL, respectively, and whose function is well-conserved (148). MASTL is located on chromosome 10 and encodes a protein kinase composed of 879 amino acids and with a molecular mass of about 97 kDa (Figure 4.2). To date, no crystal structures of full-length MASTL are available. Bioinformatic analysis revealed that MASTL has an unusual kinase structure: it contains the DFG and APE motifs, which are commonly shared by protein kinases and delimitate the activation segment (149) but, in between these two motifs, there is a large non-
conserved middle region (NCMR) of about 500 amino acids (150). The structure and the function of NCMR are still ambiguous, but it was demonstrated that this domain prevented MASTL to be expressed in a soluble form in *Escherichia coli*.

![Diagram of MASTL protein structure](image)

**Figure 4.2 MASTL protein structure**
Schematic representation of MASTL kinase structure: between the conserved DFG and APE motifs, it contains a ~500 amino acid region (non-conserved middle region, NCMR) whose function is still not clear; amino acid boundaries of the N- and C-terminals are indicated, as well as the Ser875 phosphorylation site known to be essential for MASTL function. Image from (150).

MASTL has a nuclear localization pattern throughout interphase but, during mitosis, it is exported to the cytoplasm before nuclear envelop breakdown (151). As for *Drosophila* and *Xenopus* GWL, MASTL has the crucial role of inhibiting the phosphatase activity of PP2A/B55δ complex during mitosis, in order to sustain the phosphorylation of CDK1 substrates (124); moreover, its inhibition caused multiple mitotic defects, such as lagging chromosomes, DNA bridges and cytokinesis failure. Also in human cells, MASTL was shown to be inactivated by PP1 phosphatase activity (152).

In addition to the regulation of mitotic progression, GWL was shown to function also as regulator of the DNA damage response (DDR); its depletion was associated with the delay of both checkpoint recovery and progression into cell cycle upon DNA damage induction (153). Human MASTL maintained this function and was also shown to be an essential player for the maintenance of genome integrity: its downregulation delayed the
entry in mitosis of damaged cells, caused a premature activation of the APC/C complex and induced genomic instability associated with genome endoreduplication (154).

In the recent years, evidence of both MASTL involvement in cancer progression and its targeting as promising approach for cancer therapy were provided. MASTL inhibition sensitized recurrent head and neck tumor cells to cisplatin treatment (136) and lung cancer cells to radiation therapy (135). In breast cancer, MASTL upregulation was associated with a more advanced clinical stage (136) and also proposed as indicator of metastatic and disease relapse risk within the ER+ breast tumor subtype (155). Moreover, in both breast and colon cancer cell lines, it was demonstrated that upregulation of MASTL promoted tumor cell growth through the degradation of the phosphatase PHLPP, which usually inhibits AKT activity, thus leading to AKT hyperactivation (156). Based on these recent findings that characterized MASTL as a highly valuable drug target for cancer treatment, a preliminary small-molecule MASTL inhibitor was developed (150); of note, it was created starting from the structure of the GWL minimal kinase domain, since the crystallization of full-length human GWL was not feasible.
4.2. Aims of the chapter

We have previously demonstrated, as described in chapter 3, that *MASTL* is a vulnerability gene for thyroid tumor cells, since its inhibition interfered with the growth of tumor but not of normal thyroid cells. In this chapter we aimed to dissect the molecular mechanisms that are at the basis of TC cell dependency on MASTL function. In particular, we depleted MASTL in TC cells by using RNAi, and investigated the effects of its silencing on cell proliferation, cell division, cell viability and accumulation of DNA damage.
4.3. Results

4.3.1. Effects of MASTL depletion on TC cell growth

We have previously observed that MASTL depletion led to inhibition of the growth of a panel of TC cell lines (section 3.3.5). For the studies hereafter presented, we selected two ATC-derived cell lines, namely HTC/C3 and 8505C; the results obtained would ideally contribute to the design of alternative strategies to ATC treatment, as they currently lack effective therapies (already described in section 1.1.1.4). HTC/C3 and 8505C cells were transfected with control (siNT) and three MASTL (siMASTL #1, #4 and #5) siRNAs and plated 24 hours later for both silencing efficiency and proliferation analyses (Figure 4.3A). By western blot we confirmed the effective silencing of MASTL in both HTC/C3 (Figure 4.3B) and 8505C (Figure 4.3C) cells. Cell proliferation was investigated by CellTiter-Glo assay and we found that MASTL siRNAs reduced the growth of both HTC/C3 (Figure 4.3D) and 8505C cells (Figure 4.3E), although with different extents. The effect of MASTL depletion on cell growth was also tested by colony forming assay as alternative approach: HTC/C3 and 8505C cells were transfected with control and MASTL siRNAs and stained 7 days later with crystal violet (Figure 4.3F-G); even if with different extents, cell growth inhibition was observed for all three MASTL siRNAs used, compared to control. Overall, these results suggest that MASTL is required for the proliferation of TC cells.
Figure 4.3 Effects of MASTL knockdown on TC cell proliferation
A-E) HTC/C3 and 8505C cells were transfected with control (siNT) and MASTL (siMASTL #1, #4 and #5) siRNAs and the following day were plated for silencing efficiency and proliferation ability testing as outlined in (A). Western blot analysis of MASTL expression in HTC/C3 (B) and 8505C (C); actin represents loading control. Growth curves of HTC/C3 (D) and 8505C (E) transfected and analyzed as indicated in A); cell growth was evaluated by CellTiter-Glo assay at the indicated time points after plating, plotted as mean +/- sd and normalized on day 1 values. E-F) Colony forming assay of HTC/C3 (E) and 8505C (F) cells 7 days after transfection with the indicated siRNAs; values represent optical density of Crystal Violet and are indicated as mean +/- sd of two (for HTC/C3) or three (for 8505C) independent experiments. Unpaired Student’s t-test was used to compare control vs MASTL siRNAs (*p<0.05; **p<0.01; ***p<0.001).
4.3.2. Effects of MASTL knockdown on mitosis

MASTL plays a crucial role in the regulation of mitosis progression (124) and we have previously observed the presence of aberrant mitotic figures in TC cells upon its depletion (section 3.3.6). Since it is well established that aberrant mitoses can generate cells with abnormal nuclei (157), such as multiple/lobate nuclei, nuclear blebs and micronuclei, we investigated whether such nuclear aberrations could be produced by MASTL depletion. HTC/C3 cells transfected with control (siNT) and MASTL (siMASTL #1, #4 and #5) siRNAs were scored for the presence of aberrant nuclear structures 120 hours later by immunofluorescence (Figure 4.4A). The nuclear anomalies previously listed were observed in 16.9% of siMASTL #1-cells, 19.3% of siMASTL #4-cells and 21.1% of siMASTL #5-cells, but only 6.1% of control siNT-cells (Figure 4.4B). We performed the same analysis on 8505C cells 76 hours after siRNA transfection (Figure 4.4C). Also in this case, we observed a significant increase of cells displaying aberrant nuclei (Figure 4.4D): 28.1% for siMASTL #1, 36.8% for siMASTL #4 and 41.1% for siMASTL #5, in comparison with 14% for siNT. On the whole, these results suggest that MASTL depletion interferes with TC cell division process originating cells with abnormal nuclei.
Figure 4.4 Analysis of the effects of MASTL knockdown on the structure of TC cell nuclei
A and C) Representative images of HTC/C3 and 8505C cells transfected with control (siNT) and three MASTL siRNAs and stained for nuclei (blue, DAPI) 120 (A) and 76 (C) hours later; cells with aberrant nuclei are indicated by arrows. B and D) Quantification of HTC/C3 and 8505C cells with abnormal nuclei analyzed in A and C); for each set, the percentage of abnormal nuclei (represented as mean +/- sd) was determined by counting a total number of 1900-2000 cells (B) and 3000 cells (D) from 8 and 6 independent experiments, respectively. Unpaired two-tailed Student’s t-test was used to compare control (siNT) vs MASTL (siMASTL #1, #4, #5) depleted cells; asterisks indicate statistical significance (***p<0.001)

4.3.3. Effects of MASTL depletion on TC cell viability

Mitotic aberrations and formation of cells with abnormal nuclei (such as multiple nuclei and micronuclei) are generally described as “mitotic catastrophe”, an event that eventually leads to cell death by apoptosis or necrosis (158). Since we observed both a
reduction of TC cell growth (section 4.3.1) and an increase of TC cells displaying nuclear aberrations upon MASTL depletion (section 4.3.2), we investigated whether MASTL silencing might be associated with occurrence of TC cell death. For the studies hereafter presented, we used siMASTL #5 oligo to deplete MASTL, as we previously observed that it gave the maximum effects on growth inhibition and aberrant nuclei formation. HTC/C3 and 8505C were transfected with siNT and siMASTL #5 and analyzed at different time points. Starting from 72 hours after transfection, we observed morphological changes in HTC/C3 depleted cells, as some appeared to be enlarged in size, together with the presence of floating cells (Figure 4.5A). Western blot analysis showed efficient MASTL silencing, as early as 48 hours after transfection, and an increase of apoptotic markers cleaved Caspase 3 and cleaved PARP, starting from 72 hours after transfection (Figure 4.5B). Moreover, we observed an increase of phosphorylated-CDK1 levels on Tyrosine 15 at the latest time points, thus supporting impairment in cell cycle progression. For 8505C cells, although morphological changes were not marked as for HTC/C3, we observed the presence of floating cells starting from 96 hours after transfection of MASTL siRNA (Figure 4.5C). Similarly to HTC/C3, western blot analysis showed efficient MASTL silencing and increased expression of cleaved Caspase 3, cleaved PARP and phosphorylated-CDK1 in MASTL-depleted cells, compared to control (Figure 4.5D). These findings suggest that MASTL depletion is associated with apoptotic cell death.
A

HTC/C3

48h  72h  96h  120h

siNT

siMASTL #5

B

siNT  siMASTL #5

48 h  72 h  96 h  120 h

+ - + - + - + - + - + - + - + -

MW (KDa)

97  36  89  116

MASTL

P-CDK1 (Y15)

CDK1

Cleaved Caspase 3

GAPDH

C

8505C

48h  72h  96h  120h

siNT

siMASTL #5

D

siNT  siMASTL #5

48 h  72 h  96 h  120 h

+ - + - + - + - + - + - + - + -

MW (KDa)

97  36  89  116

MASTL

P-CDK1 (Y15)

CDK1

Cleaved Caspase 3

GAPDH

Cleaved PARP

GAPDH

88
**Figure 4.5 Effects of MASTL knockdown on TC cell viability**

A and C) Representative images of HTC/C3 (A) and 8505C (C) at the indicated time points after transfection with control (siNT) and MASTL (siMASTL #5) siRNAs. B and D) Western blot analysis of HTC/C3 (B) and 8505C (D) cells for MASTL, phosphorylated CDK1 (Tyr15), total CDK1, cleaved Caspase 3 and cleaved PARP expression; GAPDH was used as loading control. One representative out of two independent experiments is shown for each cell line.

To further corroborate the occurrence of TC cell death upon MASTL depletion, we performed Annexin V assay on MASTL-depleted and control HTC/C3 cells followed by flow cytometry analysis (Figure 4.6A). We observed that MASTL depletion reduced the number of viable cells in comparison with control (Figure 4.6B): siNT-HTC/C3 viable cells (TO-PRO-3/Annexin V) were 92.7% at 48 hours, 80.2% at 72 hours, 62.2% at 96 hours and 71.0% at 120 hours after transfection; siMASTL-HTC/C3 viable cells were 91.4% at 48 hours, 65.5% at 72 hours, 27.6% at 96 hours and 30.6% at 120 hours after transfection. The percentage of early apoptotic cells (TO-PRO-3/Annexin V) of siNT and siMASTL transfected cells respectively were: 0.9% and 0.6% at 48 hours, 1.9% and 4.5% at 72 hours, 3.4% and 4.0% at 96 hours, 3.6% and 1.3% at 120 hours. Dead cells in apoptosis (TO-PRO-3+/Annexin V+) for control and MASTL-depleted groups respectively were: 4.1% and 3.5% at 48 hours, 13.0% and 22.7% at 72 hours, 28.6% and 42.5% at 96 hours, 18.1% and 31.4% at 120 hours. Cells considered either necrotic or which already ended cell death through apoptosis (TO-PRO-3+/Annexin V-) were respectively for siNT and siMASTL: 2.3% and 4.5% at 48 hours, 5.0% and 7.3% at 72 hours, 5.8% and 26.0% at 96 hours, 7.1% and 36.6% at 120 hours after transfection. The same analysis is currently in progress on 8505C cells. Overall, our findings indicate that MASTL depletion is associated with decreased TC cell viability and increased cell death. This was associated with apoptosis but, given that we also observed an increase of MASTL-depleted cells that are considered either necrotic or apoptotic cells having already lost membrane integrity and Annexin V staining, we could not exclude that also necrotic cell death may occur.
4.3.4. Effects of MASTL depletion on DNA damage

It is well established that chromosome segregation errors are often associated with occurrence of structural chromosomal aberrations and DNA double strand breaks (DSB) (159). Given the effects on mitosis observed upon MASTL depletion, together with
evidence of MASTL involvement in response to DNA damage (136), we investigated whether its depletion may induce DNA DSBs in TC cells. To this aim, we analyzed by western blot the expression of phosphorylated histone H2AX on Serine 139 (referred to as γH2AX), an established marker of DNA DSBs (160), in control (siNT) and MASTL depleted (siMASTL #5) cells. Together with efficient MASTL silencing as early as 48 hours after siRNA transfection, we observed a gradual increase of γH2AX expression levels in both HTC/C3 (Figure 4.7A) and 8505C (Figure 4.7B) MASTL-depleted cells starting from 72 hours after transfection. These results suggest that MASTL depletion may lead to accumulation of DNA damage in TC cells.

**Figure 4.7 Effects of MASTL depletion on occurrence of DNA damage**
Western blot analysis of MASTL and γH2AX expression in HTC/C3 (A) and 8505C (B) cells at the indicated time points after transfection with control (siNT) and MASTL (siMASTL #5) siRNAs. GAPDH was used as loading control. One representative out of two independent experiments is shown. Protein extracts used in panel A for HTC/C3 are the same used in figure 4.4.
**4.3.5. Lentiviral-mediated stable MASTL silencing in TC cells**

We searched for a system to stably inhibit MASTL expression for future studies, in order to obtain a more controlled MASTL knockdown and to explore the long-term effects of its depletion in TC cells. To this aim we purchased an inducible lentiviral shRNA system, namely TRIPZ (Figure 4.8A): the lentiviral vector carried a specific shRNA together with *red fluorescent protein (RFP)* under the control of a tetracycline-inducible promoter, thus allowing the expression of the shRNA only upon doxycycline administration. The plasmid also contained a puromycin-resistance cassette to allow the selection of stably transduced cells. TRIPZ system was already demonstrated to produced efficient gene silencing in other cell models (161,162). We purchased TRIPZ vectors containing two different shRNA sequences targeting MASTL, namely TRIPZ-shMASTL #1 and TRIPZ-shMASTL #2. We transduced 8505C cells with the two vectors and selected transduced cells with puromycin. Cells were then treated with 1 µg/ml and 2 µg/ml doxycycline and RFP expression was monitored: as early as 24 hours after treatment, both 8505C-TRIPZ-shMASTL #1 and 8505C-TRIPZ-shMASTL #2 cells were positive for RFP at fluorescent microscope observation (Figure 4.8B). Western blot analysis of 8505C-TRIPZ-shMASTL #1 and 8505C-TRIPZ-shMASTL #2 cells showed a gradual decrease of MASTL expression upon doxycycline administration, in comparison with untreated cells (Figure 4.8C), thus indicating that the shRNAs targeting MASTL are induced together with RFP after doxycycline administration. These observations support the effectiveness of the system to stably inhibit MASTL expression in 8505C cells.
A

B

8505C-TRIPZ-shMASTL #1

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8505C-TRIPZ-shMASTL #2

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We performed a preliminary experiment to test the growth of 8505C-TRIPZ-shMASTL #2 upon MASTL depletion. We induced the expression of RFP and shMASTL #2 by administration of 2 µg/ml doxycycline and monitored RFP expression, which was clearly detectable even 11 days later in treated cells, and absent in untreated ones (Figure 4.9A).

We observed, by western blot analysis and CellTiter-Glo assay respectively, that doxycycline treatment decreased MASTL expression (Figure 4.9B) and inhibited 8505C cell growth (Figure 4.9C), in comparison with untreated cells. Together, these results further support the effectiveness of lentiviral system and suggest that long-term MASTL depletion impairs the growth of TC cells.
Figure 4.9 Effects of MASTL knockdown on long-term 8505C cell growth
8505C cells transduced with TRIPZ-shMASTL #2 lentiviral vector were tested for RFP expression, MASTL silencing and proliferation ability. A) Representative images of treated and untreated cells at 11 days after doxycycline administration; both visible and RFP fluorescence images are reported. B) Western blot analysis of MASTL expression at the indicated time points after doxycycline treatment in cells represented in A); actin was used as loading control. C) Growth rates of cells, represented in A) and B), at the indicated time points; cell growth was evaluated by CellTiter-Glo assay at the indicated time points after plating, plotted as mean +/- sd and normalized on day 1 values; unpaired two-tailed Student’s t-test was used to compare untreated vs doxycycline-treated cells (*p<0.05; **p<0.01; ***p<0.001).
4.3.6. Set up of IHC analysis of MASTL expression

As preliminary for future studies aimed to assess MASTL expression in thyroid tissues, we purchased MASTL antibody (clone 4F9, Millipore) used for immunohistochemistry (IHC) analysis in the study presented by Wang and colleagues (136) and tested its working conditions on 8505C cells transfected with siNT and siMASTL as positive and negative controls, respectively. By western blot analysis, we first confirmed the reduction of MASTL expression in siMASTL-cells, in comparison with siNT (Figure 4.10A). Then, in collaboration with the immunohistochemistry facility of our institute, we embed 8505C siNT and siMASTL cells in paraffin, in order to mimic conditions of IHC tissue analysis, and performed IHC; nuclear MASTL staining was observed in the positive (siNT) but not negative (siMASTL) control (Figure 4.10B). These observations assert the effectiveness of the antibody for IHC applications and support the feasibility of future studies aimed to assess MASTL expression in thyroid tissues by IHC.

Figure 4.10 IHC analysis of MASTL expression in paraffin embedded 8505C cells
8505C cells were transfected with control and MASTL siRNAs and embedded in paraffin 48 hours later for IHC testing of MASTL antibody. A) Western blot analysis for MASTL expression in siNT and siMASTL 8505C cells; GAPDH represents loading control. B) Representative images of IHC analysis of MASTL expression in 8505C siNT and siMASTL cells; scale bar=50 µm.


4.4. Discussion

In this chapter, we dissected the effects of MASTL depletion on the regulation of thyroid tumor cell biology. We knocked down MASTL in two ATC-derived cell lines by using three different siRNAs and demonstrated a significant reduction of the growth of TC cells, compared to control. We found that MASTL-depleted TC cells displayed increased nuclear aberrations, such as multiple nuclei, micronuclei or lobular nuclei. MASTL depletion also led to increased expression of apoptotic markers, cell death and DNA damage, compared to control.

MASTL recently emerged as promising therapeutic target for cancer treatment: its inhibition was able to reduce the viability of head and neck tumor cells when combined with cisplatin treatment (136) and of non small cell lung cancer cells when combined to radiation therapy (135). Notably, these studies reported that MASTL inhibition alone had a partial (136) or null (135) growth inhibitory effect on tumor cells. To date, MASTL role in the regulation of thyroid tumor cell biology has been never explored. Our studies demonstrated that its inhibition hampered the growth of thyroid tumor cells, thus further pointing out its relevance as possible target for cancer treatment. Of note, we observed that the three different MASTL siRNAs produced different extents of growth inhibition on TC cells. We hypothesized that these differences may be related to the different sequences of siRNAs: as they target different MASTL exons, it may be possible that the secondary structure of the mRNA influences their stability and silencing efficacy during time. Moreover, we decided to set a lentiviral-mediated inducible system to stably silence MASTL expression, in order to obtain a more controlled inhibition of MASTL expression that would last during time; we already tested the system on 8505C cells and observed its
efficacy in depleting MASTL in cells together with inhibition of the growth. This system will be useful for future studies on the role exert by MASTL in thyroid tumor cell biology.

We described in the previous chapter the presence of aberrant mitotic figures in MASTL-depleted TC cells, which were in line with previous observations in other cell models (124,151). The outcome of such aberrant mitoses is indicated as “mitotic catastrophe”, that is an aberrant form of mitosis, whose ultimate step consists in the formation of cells with aberrant nuclear structures, as multiple nuclei, micronuclei and lobular nuclei (158). Mitotic catastrophe is considered as a process preceding cell death, which may occur through apoptotic or necrotic mechanisms either after or during deregulated mitosis. Here, we observed that MASTL silencing led to the increase of TC cells displaying such nuclear anomalies, thus suggesting that MASTL inhibition may cause mitotic catastrophe in TC cells. Moreover, we observed increased cell death upon MASTL depletion. Western blot analysis showed occurrence of apoptosis, but Annexin V assay suggested that also necrotic cell death may occur. Together, these findings agree with the current knowledge about the outcome of cells undergoing mitotic catastrophe (158). The stimulation of mitotic catastrophe emerged as a promising strategy for cancer treatment. As a matter of fact, tumor cells often displayed mitotic catastrophe features upon treatment with ionizing radiations (163) or with drugs that hinder proper cell division, such as agents targeting microtubules, mitotic checkpoint components (164), and mitotic regulators as Aurora kinases, PLK1 and MPS1 (164). The final result of these treatments was tumor cell death (165). With regards to thyroid tumors, the targeting of mitotic regulators has already been explored and promising results were provided: targeting of PLK1 (166) and Aurora kinases (167) through small-molecule inhibitors significantly reduced the growth and induced cell death in ATC-derived cell lines. Our study, by showing unequivocal reduction of cell viability and increase of cell death upon MASTL
depletion, identified a novel mitotic target and strengthened the evidence of mitotic regulators as promising targets for thyroid tumor treatment.

We also found that MASTL silencing led to the accumulation of DNA damage in TC cell; we hypothesized that this increase may be related to different mechanisms. Chromosome segregation errors during mitosis are known to generate structural damage to chromosomes (159) and it is known that mitotic catastrophe cells accumulate chromosomal breaks that eventually trigger DNA damage response and cell death (168). As we observed both aberrant mitotic figures and mitotic catastrophe features in TC cells upon MASTL depletion, we hypothesized that the inhibition of MASTL may lead to structural DNA damage in TC cells. On the other hand, it is known that tumor cells exhibit higher levels of endogenous DNA damage, compared to normal ones, as result of the activity of oncogenes that promote uncontrolled cell growth; therefore, tumor cells become dependent on the activity of pathways that promote damage repair in order to sustain tumorigenic processes (169). Since MASTL is involved in both the regulation of DNA damage response (153) and the maintenance of genome stability during DNA damage response (154), we hypothesized that its inhibition in TC cells may lead to further accumulation of damaged DNA. Whether MASTL-associated DNA damage accumulation is related to one or the other mechanisms, or even related to both, remains to be investigated.

On the whole, our studies supported the notion that MASTL represents a vulnerability for thyroid tumor cells: its inhibition hindered thyroid tumor cell growth and was associated with mitotic catastrophe, cell death and increased DNA damage. Our results, together with recent literature data, propose MASTL as valuable target for cancer therapy, putting the rationale for the development of small-molecule inhibitors to be tested for therapeutic purposes.
5. Results: COPZ1 studies

As previously reported, the aim of our studies was to identify thyroid tumor cell vulnerabilities; as described in chapter 3, among the putative non-oncogenes identified through the functional siRNA screening, we found COPZ1. Even though the majority of my studies were aimed to the dissection of MASTL role in TC cell biology, in our laboratory we also studied the role exerted by COPZ1 in the regulation of TC cell viability, to which I directly collaborated.

COPZ1 encodes the ζ-subunit of coatamer protein complex I (COPI), which is a heptameric vesicular-carrier protein complex (137). COPI most characterized function is the transport of proteins from the Golgi complex back to the endoplasmic reticulum and in between Golgi compartments. In addition to this function, COPI complex was also found to be involved in the maturation of endosomes and regulation of autophagy: depletion of COPI subunits impaired transport of autophagic vacuoles into early endosomes with the consequent accumulation of autophagic vacuoles and inhibition of the autophagic flux (138). Moreover, COPI was identified as regulator of lipid storage and subsequent maintenance lipid homeostasis (139).

COPZ1 recently emerged as target gene for tumor therapy (125). It was demonstrated that tumor cells become dependent on COPZ1 activity because of the downregulation of its isoform COPZ2. The suppression of COPZ1 activity was effective in inducing cell death in both proliferating and non-diving tumor cells, while leaving unaffected normal cells. Also other members of COPI complex were reported to be promising targets for cancer therapy. COPA, which encodes the α-subunit of the complex, emerged from a siRNA screening as vulnerability gene in mesothelioma cells, as its
silencing inhibited both in vitro and in vivo tumor cell growth (170). KRAS mutant tumors showed synthetic lethality with COPB2, that encodes the β'-subunit of COPI (171). The targeting of different members of the COPI complex was associated with abortive autophagy and induction of cell death in different types of tumor cell lines (172) and a recent work identified ARCN1, which encodes the δ-subunit of the complex, as valuable target to block cell growth in different tumors (173). Interestingly, COPZ1 crystal structure was produced, thus laying the foundation for future studies aimed to design specific COPZ1 inhibitors (174).

COPZ1 was among the top-ranked hit genes resulted from the functional screening described in chapter 3; the aim of the studies presented in this chapter was to dissect the mechanisms through which COPZ1 knockdown inhibited the growth of TC cells. The results of this study are hereafter summarized.

We demonstrated that COPZ1 depletion by transient siRNA transfection dramatically impaired the growth of TC cell lines, representative of the different histotypes and carrying different genetic lesions; we also found that its silencing did not affect the growth and viability of normal thyrocytes and human mammary epithelial HME-1 cells. We demonstrated in a PTC-derived (TPC-1) and an ATC-derived (8505C) cell lines that COPZ1 depletion impaired productive autophagy by causing the accumulation of autophagosomes and block of autophagic flux, thus producing an abortive autophagy process. We also observed endoplasmic reticulum stress and unfolded protein response activation upon COPZ1 depletion, that eventually caused apoptotic cell death. In vivo silencing of COPZ1 in 8505C mouse xenografts significantly reduced tumor growth and immunohistochemical analysis on explanted tumors showed decrease of proliferation markers and increase of necrosis. We analyzed the expression levels of COPZ2, that is COPZ1 isoform, in publicly available thyroid data sets, and found that is downregulated in
TC specimens, compared to normal thyroid, thus supporting the hypothesis that COPZ1 vulnerability of TC cells may be related to the absence of its paralogous COPZ2.

Together, our findings demonstrated that COPZ1 represents a vulnerability for TC cells, thus providing a new potential target for the design of therapeutic strategies.

Details and results of the study are presented in the attached reprint of the paper (Appendix B):

6. General discussion and future perspectives

6.1. Discussion and conclusions

The global aim of the thesis was to identify potential novel targets for thyroid cancer treatment. The entire project arose from the evidence that current therapies for thyroid tumors, which are the most common endocrine malignancy, are still not effective in treating the recurrent and advanced forms (84). Most of thyroid tumors are represented by well-differentiated forms, which generally have a good prognosis and standard therapy is generally effective (78). Nevertheless, some tumors relapse and progress to the undifferentiated forms, whose treatment is still a demanding challenge. Despite the recent improvement in both the diagnosis and the knowledge of molecular alterations that drive thyroid tumorigenesis, the lack of effective therapies, especially for the advanced forms, is still present (1). The use of novel therapies such as small-molecule kinase inhibitors improved the survival of patients not responding to conventional treatment, but also produced severe side effects that hampered quality of life of patients (84). To identify nodal points for therapeutic intervention, our laboratory faced the concept of non-oncogene addiction (NOA) in thyroid tumors. NOA concept states that, in response to activation of oncogenes, tumor cells undergo increased cellular stresses and therefore rely on the activity of genes involved in stress support pathways, in order to maintain their tumorigenic capacity (98). Normal cells are not dependent on these genes, as their stress levels are not increased. NOA genes, which are not oncogenes themselves, become therefore attractive targets for therapeutic intervention, as their targeting would ideally be detrimental for tumor cell growth while not affecting normal
cells. To identify potential NOA genes in thyroid tumors, our laboratory performed a loss-of-function screening on tumor and normal thyroid cell lines using a siRNA library targeting about 9000 genes considered “druggable”. RNAi-based loss of function screenings, at variance with the current approaches of NGS technologies that allow the identification of genetic lesions at the basis of tumorigenic processes, demonstrated to be a successful strategy for the identification of “non-oncogenes” essential for tumor growth (106-108,110). Our functional screening identified a list of 15 genes whose inhibition inhibited the growth of tumor but not normal thyroid cells.

The first part of the thesis aimed to validate the 15 hit genes in the original cell models used for the screening and to extend the analysis of the effects of their silencing on the viability of various thyroid cell lines, representative of the different TC histotypes. Thyroid TCGA dataset interrogation showed that hit genes had variable expression levels among PTC samples and the normal counterpart and, notably, were not mutated, thus uncovering their reliability as putative “non-oncogenes”. After selection of efficiently working antibodies and siRNAs, we selected 3 out of the 15 hit genes for validation studies: MASTL (encoding a mitotic kinase), CCND1 (encoding a regulator of CDK4/6 activity) and COPZ1 (encoding a regulator of protein vesicular transport). Functional studies confirmed the growth inhibitory effect of MASTL, CCND1 and COPZ1 silencing in the tumor but not normal cell lines used for the screening and, interestingly, we demonstrated that the growth of other TC cell lines, derived from different tumor histotypes and carrying different genetic lesions, was sensitive to their inhibition. These results demonstrated that MASTL, Cyclin D1, and COPZ1 represent thyroid tumor cell vulnerabilities and are noteworthy for further studies, aimed to explore their targeting as possible strategy for therapeutic intervention. Moreover, this validation study highlighted the effectiveness of large-scale RNAi-based functional screening in identifying tumor cell
vulnerabilities, thus supporting the strength of RNAi-based approaches in the discovery of novel therapeutic targets (106-110).

The studies presented in the second and third part of the thesis aimed to identify the mechanisms through which the inhibition of MASTL and COPZ1, respectively, hindered TC cell growth. We observed that MASTL silencing in two ATC-derived cell lines reduced cell proliferation ability. In parallel, cells exhibited features of mitotic catastrophe, documented by the increase of aberrant nuclear morphologies as micronuclei or multiple nuclei. Moreover, MASTL depletion was associated with TC cell death that was possibly mediated by both apoptotic and necrotic mechanisms, as already discussed in section 4.4. We also found that MASTL silencing led to increased DNA damage levels in TC cells. With regards to COPZ1, we observed that its silencing inhibited the growth of various TC cell lines by inducing autophagosome accumulation and abortive autophagy, endoplasmic reticulum stress and unfolded protein response, which ultimately led to apoptotic cell death. Moreover, COPZ1 silencing inhibited in vivo tumor growth in mouse xenograft and was associated with decreased expression of proliferation markers and increase of necrosis; these findings further supported the reliability of COPZ1 targeting in vivo as potential therapeutic strategy.

Collectively, our studies suggest that MASTL and COPZ1 may effectively represent “non-oncogenes” in thyroid cancer. As previously discussed, thyroid TCGA dataset interrogation revealed that neither MASTL nor COPZ1 were overexpressed in PTC compared to normal counterpart, and, except for one silent mutation of COPZ1, no mutations were reported for both of them in about 500 PTC samples. Thus, both MASTL and COPZ1 properly comply with the concept of NOA (98).

We hypothesized that MASTL may act on mitotic stress support pathway and its depletion may impair stress support and cause tumor cell death (98). Tumor cells often
face with elevated mitotic stress, due to the high number of mutations that alter mitosis fidelity and cause chromosome missegregation; thus, in order to avoid excessive genomic instability, they become dependent on stress support pathways (175). Both sensitization and overload of mitotic stress were exploited as strategies to interfere with tumor cell growth, and are widely exemplified by the targeting of mitotic machinery elements, such as PLK1 and Aurora kinases, as well as by the use of microtubule-targeting agents like chemotherapy drugs (already discussed in section 4.4). Other regulators of mitotic process, such as the spindle assembly checkpoint (SAC) components Mad2 and BubR1, triggered cell death when genetically eliminated in mice and were therefore proposed as targets for inducing further chromosomal instability and consequent cell death in tumor cells (176). Our results suggest that MASTL may effectively represent an example of non-oncogene acting on the mitotic stress support pathway, since we observed that its silencing interfered with TC cell mitosis and was associated with mitotic catastrophe and cell death. On the other hand, also DNA damage and replication stress is often present in tumor cells, as a result of the high degree of replication and the rate of spontaneous DNA damage formation. Targeting of elements that promote DNA repair, such as ATM, was successful in causing stress sensitization and tumor cell death (177). Also radiations and chemotherapy were effective in overloading DNA damage and replication stress and leading to tumor cell death. The increase of DNA damage and cell death observed upon MASTL depletion suggest that MASTL vulnerability of TC cells may be related also to impaired DNA damage stress support pathway.

Regarding COPZ1, our results suggest that its depletion may be associated with proteotoxic stress. COPZ1 depletion in thyroid tumor cells blocked autophagic flux and increased endoplasmic reticulum stress, ultimately leading to apoptosis. It was demonstrated that prolonged endoplasmic reticulum stress results in cell death (178) and
that tumor cells rely on autophagy as mechanism to clear damaged/unfolded proteins and to lower proteotoxic stress (179). We hypothesized that COPZ1 depletion may affect protein balance in cells, leading to an overload of proteotoxic stress that, together with block of productive autophagy, could ultimately drive tumor cell fate towards a programmed form of cell death.

Remarkably, we demonstrated that the growth inhibitory effect of MASTL and COPZ1 targeting was restricted to thyroid tumor cells and did not affect viability of normal cells, as previously reported for other tumor types (125,135,136); these observations support the hypothesis that their targeting in vivo may produce limited side verse effects.

In conclusion, we identified putative non-oncogenes in thyroid cancer and validated MASTL and COPZ1 as “Achille’s heel” of thyroid tumors, thus providing evidence that they represent valuable elements for future studies, aimed to test their inhibition as strategy for thyroid cancer treatment.
6.2. Future perspectives

The studies presented in the thesis highlighted MASTL and COPZ1 as valuable candidates for future studies aimed to validate their targeting as potential therapeutic approach in thyroid cancer management.

MASTL represents an interesting mitotic element that recently emerged as potential target for cancer therapy (135,136) and our studies demonstrated its significance as thyroid tumor cell vulnerability.

To assess its reliability as target for thyroid tumor treatment, the effects of its inhibition in vivo will be investigated using a thyroid tumor mouse xenograft model, already used for COPZ1 studies. MASTL silencing will be performed by in vivo siRNA injection. The set up of the optimal conditions for treatment is in progress.

Deregulated mitoses can cause mitotic cell death and this mechanism often occurs upon mitotic catastrophe (158). Live imaging on TC cells expressing the fluorescent nuclear marker H2B-GFP will allow us to assess the fate of MASTL-depleted TC cells at single cell level, by monitoring cell division and observing nuclear structural changes.

Analysis of the activation status of regulators of DNA damage response pathway will allow us to clarify the effects of MASTL silencing on DNA damage accumulation.

MASTL overexpression was found in head and neck, prostate and advanced breast carcinomas (136), while for other tumors, including thyroid cancer, its expression level has been never explored. MASTL expression at protein level will be assessed by IHC analysis of normal and tumor thyroid tissues, and correlated with gene expression data derived from analysis of publicly available thyroid dataset. Together, these results will allow the characterization of MASTL expression status in thyroid cancer.
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List of abbreviations

AKT  Protein kinase B
ATC  Anaplastic thyroid carcinoma
BRAF  v-Raf murine sarcoma viral oncogene homolog B1
CCND1  Cyclin D1
CDK1  Cyclin dependent kinase 1
CDK4  Cyclin dependent kinase 4
CDK6  Cyclin dependent kinase 6
COPE  coatomer protein complex , subunit epsilon
COPZ1  Coatomer protein complex, subunit zeta-1
DNM3  Dynamin 3
EPHB4  EPH receptor B4
FNAB  Fine-needle aspiration bioppsy
FTA  Follicular thyroid adenoma
FTC  Follicular thyroid carcinoma
HCC  Hürtele cell carcinoma
HDAC  Histone deacetylase
MAP4K5  Mitogen-activated protein kinase kinase kinase kinase 5
MAPK  Mitogen-activated protein kinase
MASTL  Microtubule associated serine/threonine kinase-like
MTC  Medullary thyroid carcinoma
NGS  Next-generation sequencing
NOA  Non-oncogene addiction
NTRK1  Neurotrophic receptor tyrosin kinase A
NUDT9  Nudix (nucleoside diphosphate linked mojety X)-type motif 9
OXTR  Oxytocin receptor
PAX8  Paired box 8
PDTC  Poorly differentiated thyroid carcinoma
PI3K  Phosphatidylinositol 3-kinase
PIK3CA  Phosphatidylinositol-4,5-biphosphate 3-kinase catalytitc subunit α
PLA2G15  Phospholipase A2, group XV
PP2A  Protein phosphatase 2A
PPARγ  Peroxisome proliferator-activated receptor γ
PTC  Papillary thyroid carcinoma
<table>
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<th>Abbreviation</th>
<th>Full Form</th>
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<tr>
<td>PTEN</td>
<td>Phosphatase and tensin homolog</td>
</tr>
<tr>
<td>RAI</td>
<td>Radioactive iodine</td>
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<tr>
<td>RAS</td>
<td>Rat sarcoma</td>
</tr>
<tr>
<td>RASD1</td>
<td>RAS, dexamethasone-induced 1</td>
</tr>
<tr>
<td>REM2</td>
<td>RAS (RAD and GEM)-like GTP binding 2</td>
</tr>
<tr>
<td>RET</td>
<td>Rearranged during transfection</td>
</tr>
<tr>
<td>RGS3</td>
<td>Regulator of G-protein signaling 3</td>
</tr>
<tr>
<td>RNAi</td>
<td>RNA interference</td>
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<tr>
<td>RTK</td>
<td>Receptor tyrosine kinase</td>
</tr>
<tr>
<td>SRPK1</td>
<td>SFRS protein kinase 1</td>
</tr>
<tr>
<td>TC</td>
<td>Thyroid cancer</td>
</tr>
<tr>
<td>TKI</td>
<td>Tyrosine kinase inhibitor</td>
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<tr>
<td>WDTC</td>
<td>Well-differentiated thyroid carcinoma</td>
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Publications


Appendixes

Appendix A


Supplementary Table S1, Table S2 and Table S3 were not included in Appendix A due to their size; they can be downloaded at the following link:

http://www.impactjournals.com/oncotarget/index.php?journal=oncotarget&page=rt&op=suppFiles&path[]=5282&path[]=0

Appendix B

Appendix A
Identification of thyroid tumor cell vulnerabilities through a siRNA-based functional screening

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Keywords: thyroid cancer, non-oncogene addiction, MASTL, Cyclin D1, COPZ1

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ABSTRACT

The incidence of thyroid carcinoma is rapidly increasing. Although generally associated with good prognosis, a fraction of thyroid tumors are not cured by standard therapy and progress to aggressive forms for which no effective treatments are currently available. In order to identify novel therapeutic targets for thyroid carcinoma, we focused on the discovery of genes essential for sustaining the oncogenic phenotype of thyroid tumor cells, but not required to the same degree for the viability of normal cells (non-oncogene addiction paradigm). We screened a siRNA oligonucleotide library targeting the human druggable genome in thyroid cancer BCPAP cell line in comparison with immortalized normal human thyrocytes (Nthy-ori 3–1). We identified a panel of hit genes whose silencing interferes with the growth of tumor cells, while sparing that of normal ones. Further analysis of three selected hit genes, namely Cyclin D1, MASTL and COPZ1, showed that they represent common vulnerabilities for thyroid tumor cells, as their inhibition reduced the viability of several thyroid tumor cell lines, regardless the histotype or oncogenic lesion. This work identified non-oncogenes essential for sustaining the phenotype of thyroid tumor cells, but not of normal cells, thus suggesting that they might represent promising targets for new therapeutic strategies.

INTRODUCTION

The incidence of thyroid cancer (TC), the most common endocrine malignancy, has increased significantly over the last few decades [1]. All TCs, except medullary thyroid carcinoma, are derived from thyroid follicular cells. Differentiated tumors, namely papillary and follicular carcinoma (PTC and FTC, 80% and 10% of cases, respectively) are generally cured by current treatment involving surgery, thyroid hormone and radioactive iodine therapy; however, they occasionally progress into undifferentiated, more aggressive and lethal carcinomas. Poorly differentiated (PDTC) and anaplastic (ATC) carcinoma, although rare, are the most clinically aggressive TCs; for ATC patient the survival is reduced to few months [2]. Genetic studies performed in the last 25 years have dissected the molecular bases of TC. PTC is associated with a high frequency of deregulation of RTK/RAS/RAF/MAPK pathway [3]. This includes oncogenic
rearrangements of the RET and NTRK1 receptors, and point mutations in the BRAF and RAS genes, with the BRAFV600E as the most frequent PTC alteration. The genetic landscape of PTC has been very recently expanded by integrated genomic characterization studies which identified several novel driver alterations [4]. FTC is associated with RAS mutations and PAX8/PPARG rearrangements. RAS mutations are common in PDTC. ATC is associated with mutations of BRAF, RAS, TP53, PTEN, and PIK3CA. The identification and functional characterization of differentially expressed genes and miRNAs have also contributed to the understanding of thyroid tumor biology, and provided important diagnostic/prognostic tools [5–8].

Molecular findings have been translated into the clinical setting, leading to clinical experimentation of agents targeting thyroid oncoproteins or downstream pathways [9]. Several kinase inhibitors, primarily targeting BRAF and angiogenic kinases, are under evaluation in clinical trials [10]. Nevertheless, no effective targeted therapies are currently available for aggressive TC that are not curable by the standard therapeutic approaches.

The traditional approach for cancer target discovery is based on the identification of driver oncogenes through the characterization of the molecular alterations of primary tumors. However, according to the concept of “non-oncogene addiction” (NOA), the uncontrolled growth of cancer cells often relies on normal genes, that are not themselves oncogenes or otherwise mutated, whose activity is essential for tumor cells but not required to the same extent by normal cells [11]. NOA genes cannot be identified by genetic approaches aimed at identifying gene mutation/aberrancy, but rather by large-scale siRNA-based functional screening, a strategy which is nowadays widely used for the identification of tumor cell vulnerabilities to be explored for therapeutic purposes [12–15].

In order to identify novel, unforeseen molecular targets for thyroid carcinoma, in this study we have undertaken the screening of a siRNA oligonucleotide library in tumor (BCPAP) and normal (Nthy-ori 3–1) thyroid cell lines, and we have identified a set of genes whose silencing reduces the growth of tumor cells, while sparing growth of non-transformed cells of the same tissue origin. Among the top ranked genes, we selected Cyclin D1 (CCND1), MASTL and COPZI. All of them passed a secondary confirmation study, which unequivocally demonstrated the dependency of BCPAP cells upon their activity. Interestingly, we found that silencing of Cyclin D1, MASTL and COPZI inhibits the growth of several further thyroid tumor cell lines.

RESULTS

Druggable genome siRNA screening

To identify genes affecting growth of thyroid tumor cells, we conducted an RNAi-based phenotypic screening, examining effects on cell growth. The papillary thyroid carcinoma BCPAP cell line, carrying the BRAFV600E mutation, and the immortalized normal human thyrocyte Nthy-ori 3–1 cell line were transfected with a siRNA library containing 25139 siRNA oligos targeting about 9000 potentially “druggable” genes (3 duplexes/gene, on average), and with a non-targeting siRNA (siNT) and a siRNA targeting the proteasomal subunit PSMD3 as negative and positive controls, respectively. Cells were transfected at low density in 96-well plates and colony formation assay (CFA) was performed after 7 (Nthy-ori 3–1) or 8 (BCPAP) days. Images of a representative plate for each of these lines are shown in Figure 1A. We preferred CFA to short-term (48–72 hours) proliferation assay, since it allows the detection of long-term consequences of “weak” phenotypes (our unpublished results). The screening results are shown in Figure 1B: scatter plots represent the fluorescence signal, derived from CFA acquisition, normalized with respect to siNT (% siNT) of Nthy-ori 3–1 and BCPAP cells transfected in duplicate with the library siRNA oligos (the complete list is reported in Table S1). Of note, the uneven distribution of data across graph diagonal denotes slightly higher transfection efficiency for Nthy-ori 3–1 than for BCPAP. Genes essential for cell viability of BCPAP, but not Nthy-ori 3–1 cells, were identified through the “d parameter” (defined in Materials and Methods). d values close to 0 denote preferential inhibition of BCPAP cell proliferation with respect to Nthy-ori 3–1. Based on data distribution, a threshold of -3σ (corresponding to d = 47.2) was applied to define differentially active hits: 398 siRNA oligos (1.58%), targeting 386 genes, were found to be below this threshold and thus were defined as “differential hits” (Figure 1C; hit list is reported in Table S2). A significant preferential activity towards BCPAP cells was observed for 12 genes with 2 oligos out of 3, and for the remaining 374 genes with 1 oligo out of 3; the latter include BRAF, consistent with the notion that BCPAP cells are addicted to BRAFV600E oncogene [16]. No genes emerged with 3/3 oligos among hits. Functional annotation clustering analysis was performed on the 386 gene list (382 DAVID IDs), using Gene Ontology–Biological Process (GO-BP) and Gene Ontology–Molecular Function (GO-MF) annotation terms and medium classification stringency. A significant Enrichment score (>1.3) was found in 15 out of the 117 annotation clusters that were globally identified. The top ranked GO-terms, representative for the 15 significant clusters, have been reported in Figure S1A.

By setting an arbitrary threshold of 20% colony growth with respect to siNT controls, we identified 1695 siRNA oligonucleotides (6.74%) capable of inhibiting cell growth both in BCPAP and Nthy-ori 3–1, therefore defined as “lethal hits” (Figure 1C). Two hundred and seventeen genes emerged as indiscriminately lethal hits with 2/3 (163) or 3/3 (54) oligonucleotides (Table S3). Most of them encode proteins involved in fundamental processes,
and some, such as the kinases PLK1, WEE1, AURKB and several proteasome subunits, have previously been shown to be essential for cell survival, emerging as top-ranking lethal hits in RNAi-mediated phenotypic proliferation screens in different tumor cell lines [14, 17, 18].

Confirmation of differentially active hits

Eighty-four siRNA oligonucleotides, targeting 28 genes, were selected for confirmatory studies. Hits were prioritized for technical confirmation based on $d$ values of individual oligos and on specific interest of identifying candidate druggable target genes. siRNAs were picked from library plates and transfected manually in triplicate on Nthy-ori 3–1 and BCPAP cells using the same experimental conditions as employed in primary screening. In these experiments a threshold of $-2\sigma$ (corresponding to $d = 49.8$) was applied to define oligos with significantly low $d$ values, i.e. displaying differential activity. We considered equally confirmed any siRNA oligos that in confirmation experiments gave the same phenotype of the screening, including the ones inactive or lethal for both cell lines. Figure 2A shows the comparison between primary screening and confirmation results. Eighteen out of 35 siRNA oligos (targeting 15 genes), which had shown differential activity ($d < -3\sigma$) in the primary screening were confirmed by manual transfections ($d < -2\sigma$). All of 49 oligos which were not found to be differentially active in the screening run ($d > -2\sigma$) were confirmed as such by manual transfections ($d > -2\sigma$). Overall, the phenotype of 67 out of 84 oligos was confirmed. Representative images of outgrown colonies from the confirmatory study are shown in Figure 2B and the corresponding quantifications are reported in Figure 2C. The list of confirmed hits is shown in Table 1: 3 genes were confirmed with 2 oligos (MASTL,
PLA2G15 and COPE) and 12 genes were confirmed with 1 oligo (RGS3, CCND1, RASD1, NUDT9, OXTR, BRAF, COPZ1, MAP4K5, EPHB4, DNM3, REM2, SRPK1). Functional annotation analysis indicates that these genes are involved in several biological processes, such as cell cycle control, DNA damage and cell death, vesicular transport and endocytosis, cellular metabolic processes and intracellular signal transduction (Table 1). Network analysis performed by Ingenuity Pathway Analysis (IPA) identified two major connection networks as reported in Figure S1B.

Molecular profiling of confirmed hit genes

The thyroid TCGA data set [4], including 496 PTC samples, 58 of which matched with normal thyroid, was interrogated for assessing the mutational status and the expression level of hit genes in PTC. As expected, oncogenic BRAF mutations were detected with high frequency, being present in 248 samples. No mutations affecting the other genes were detected, with the exception of a silent mutation in COPZ1 gene in one case (data not shown). Expression analysis of hit genes was investigated on the whole data set (data not shown), as well as on the 58 matched tumor and normal samples (Figure 3); in both cases identical results were obtained. According to their expression levels, hit genes can be classified in three groups: 1) genes with significantly high overexpression in PTC vs normal thyroid: CCND1, RGS3, OXTR, RASD1, DNM3; 2) genes with equal or slightly different expression in PTC and normal thyroid: COPE, COPZ1, PLA2G15, SRPK1, REM2, EPHB4, BRAF; 3) genes that are significantly downregulated in PTC vs normal thyroid: MAP4K5, NUDT9, MASTL. No differences among different PTC variants (classical, follicular, tall cell) or tumor stage were observed (data not shown).

Data retrieved from the Cancer Cell Line Encyclopedia (http://www.broadinstitute.org/ccle/home) showed that all the hit genes are expressed in BCPAP cells, with levels ranging from “low” to “very high” (data not shown).
Table 1. List of the confirmed hits after the primary screening.

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<tr>
<th>Gene Symbol</th>
<th>Full Gene Name</th>
<th>Accession</th>
<th>GO-BP Term</th>
<th>Confirmed active oligos</th>
<th>d (active oligos)</th>
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<td><strong>MASTL</strong></td>
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shown). Mutation analysis data indicated a low number (50/1651) of mutated genes in BCPAP cells. Cyclin D1, MASTL, MAP4K5, SRPK1 and EPHB4 genes were found to be of wild-type status, whereas no information was available for the remaining hit genes. These findings suggest that the addiction of BCPAP cells to the activity of the hit genes is unlikely related to alterations of the cognate proteins caused by mutations.

### Hit gene expression in thyroid cell lines

We investigated the expression of 10 genes (the top 11 hits except BRAF, see Table 1), in the normal (Nthy-ori 3–1) and tumor (BCPAP) cell lines used for the siRNA library screening, as well as in a panel of thyroid tumor cell lines, representative of different tumor histotypes, and/or carrying different oncogenes: PTC-derived (TPC-1, carrying the RET/PTC1 oncogene); FTC-derived (WRO82–1); ATC-derived (8505C and HTC/C3, carrying the BRAFV600E oncogene; KAT-18). The expression of MASTL, PLA2G15, COPE, Cyclin D1, COPZ1 and MAP4K5 was investigated by western blot analysis (Figure 4A). Cyclin D1 was found to be 1.3 to 2.7 fold overexpressed in 6 out of 7 tumor cell lines with respect to Nthy-ori 3–1, with the exception of KAT-18, in which it was found less expressed. For COPE a slight increase in the protein expression levels in tumor cells, except KAT-18, with respect to Nthy-ori 3–1 was observed. MASTL was found frequently less expressed in tumor cells, in keeping with what was observed by gene expression analysis in tumor samples (Figure 3). COPZ1 protein expression was slightly increased in all the tumor cell lines, and greater in 8505C. PLA2G15 expression was slightly reduced in all the tumor cell lines, except BCPAP, and strongly reduced in KAT-18. For MAP4K5, a slight increase in BCPAP, WRO82–1 and 8505C, and a decrease in the remaining cell lines were observed.

The expression of RASD1, NUDT9, RGS3 and OXTR genes was investigated by RT-PCR, as we failed to unequivocally detect the specific protein by western blot analysis (data not shown). As shown in Figure 4B, the genes are expressed in all the cell lines analyzed, and no differences can be appreciated.

### Hit validation in BCPAP cell line

Among the candidate BCPAP vulnerability genes identified by the screening we selected CCND1, MASTL and COPZ1 for a validation study using commercially available siRNAs, different from those of the library screening. Four different oligos for MASTL, and a smart pool of oligos for CCND1 and COPZ1 were used (details are reported in Table S4). Firstly, the efficiency of gene knock-down was evaluated by investigating the expression of corresponding proteins in Nthy-ori 3–1 cells transiently transfected with specific siRNAs and with non-targeting oligos (siNT) as control (Figure 5A). For MASTL the highest silencing efficiency was obtained with oligos 1 and 4 (89 and 95% of inhibition at 72 hours, respectively). siRNA targeting COPZ1 and CCND1 completely abrogated protein expression. Of note, in keeping with the siRNA library screening results, no macroscopic effects on Nthy-ori 3–1 cell growth/survival were observed (data not shown).

To investigate the effect of MASTL, CCND1 and COPZ1 gene silencing on BCPAP and Nthy-ori 3–1 cell viability, cells were transfected in 96 well plates with the different oligos above described, or with NT and PSMC3 siRNAs as controls, and analyzed by CellTiter-Glo assay 10 days later. Scatter plots in Figure 5B represent the luminescence signal (average of two replicates, normalized with respect to siNT) of cells transfected with each siRNA oligo. siMASTL#1, siMASTL#4, siCCND1 and siCOPZ1 reduced cell viability by 60%, 93%, 94% and 86%, respectively, an efficiency comparable to that of the control siRNA PMSC3 (93%). No effects were observed for siMASTL#2 and siMASTL#3, in keeping with their low silencing efficiency (Figure 5A).
The effect of MASTL, CCND1, COPZ1 gene silencing in BCPAP cells was further confirmed by a different experimental approach. Cells were transfected in 6-well plates with the oligos which showed efficient silencing in the previous experiment (siCCND1 and siCOPZ1 smart pools, siMASTL #1 and #4 oligos), and the newly in house synthesized MASTL oligo #5 (see Figure S2 for silencing efficiency). Six days later the effect of MASTL, COPZ1 and CCND1 silencing on both cell growth and cell viability was evident by microscope observation: silenced BCPAP cultures were less confluent than control (siNT); in addition, they presented numerous floating (most likely dead) cells (Figure 5C). Trypan blue exclusion assay showed a reduction of cell growth by 69.5%, 59.7% and 71% for siMASTL#1, #4 and #5, respectively; by 99.6% for siCOPZ1, by 91.5% for siCCND1 in comparison with the control (siNT). Western blot analysis performed at the end point of the experiment (Figure 5E) showed the complete abrogation of MASTL and Cyclin D1 protein expression. The same analysis was not feasible for COPZ1: due to the massive lethal effect of siCOPZ1, an insufficient amount of protein extract was recovered (data not shown).

Overall these results confirmed the growth inhibitory effect of Cyclin D1, MASTL, and COPZ1 gene silencing, thus providing evidence that BCPAP cells are addicted to the activity of these genes.

**Figure 3: Hit gene expression levels in PTC.** Box plot distribution of RNAseq normalized expression values from the thyroid TCGA dataset of 15 selected genes in 58 matched normal (N) and PTC samples; A, B and C. represent genes overexpressed, equally or downregulated in PTC vs normal samples, respectively. The statistical significance of differences among the groups was assessed using Wilcoxon test.

Irrespective of the specific oncogenic driving lesion and histotypes, thyroid tumors may share common effector
pathways for sustaining their transformed phenotype. To assess whether the survival genes confirmed in BCPAP cells are involved in such common pathways, we investigated the effect of \textit{CCND1}, \textit{MASTL} and \textit{COPZ1} gene silencing in a panel of thyroid tumor-derived cell lines representative of different tumor histotypes.

Cell lines were transfected in 96 well plates with oligos targeting \textit{MASTL}, \textit{CCND1} and \textit{COPZ1} genes and analyzed by CellTiter-Glo assay 10 days later. As reported in Figure 6A, gene silencing induced variable extents (15–95%) of cell growth inhibition in the majority of cell lines tested; no effect of \textit{CCND1} silencing in 8505C and KAT-18 cells, and of \textit{MASTL} silencing in KAT-18 was observed. We also assessed viability of cells transfected in 6-well plates by trypan blue exclusion assay performed four-seven days after siRNAs transfection. As reported in Figure 6B, in all the cell lines analyzed (except KAT-18), \textit{MASTL}, \textit{COPZ1} and \textit{CCND1} gene silencing led to a significant decrease of cell viability: by 30–63% for siMASTL#1, by 35–75% for siMASTL#4, by 40–75% for siMASTL#5, by 75–99% for COPZ1 and by 70–94% for siCCND1. KAT-18 cells were found to be less sensitive to all the siRNA oligos: we found a decrease of cell growth by 31% for siMASTL#1, by 53% for siCOPZ1 and by 43% for CCND1, and by 52.3% for the siPSMC3 control, and no effect for siMASTL#4 and #5. The reduction of cell growth in all the cell lines was also documented by images taken before cell count: silenced cells appeared less confluent in comparison with the control, and presented variable fractions of detached and floating cells (Figure S3). Concomitantly, western blot in Figure 6C shows the high efficiency of \textit{MASTL}, \textit{CCND1} and \textit{COPZ1} silencing in the same samples. The same analysis was not feasible for TPC1 cells transfected with siCOPZ1, due to massive lethal effect (data not shown). Importantly, this experiment revealed a growth inhibitory effect of \textit{CCND1} silencing in 8505C and KAT-18, and of \textit{MASTL} silencing in KAT18 cells, which was not previously observed. This could be ascribed to variability in the experimental conditions and/or silencing efficiency.

Figure 4: Hit gene expression in different thyroid tumor cells. A. Western blot analysis of MASTL, Cyclin D1, COPZ1, MAP4K5, PLA2G15 and COPE protein expression in a panel of thyroid cell lines; values represent band intensity, assessed by densitometry analysis, reported as the ratio to actin (loading control) and normalized for the Nthy-ori 3–1 value. B. RGS3, RASD1, NUDT9 and OXTR mRNA expression by RT-PCR in a panel of thyroid cell lines; GAPDH was used as housekeeping control. One out of three representative experiments is shown.
We next performed some experiments to further corroborate the dependency of thyroid cancer cells upon Cyclin D1, MASTL and COPZ1 activity.

We tested thyroid tumor cell lines for susceptibility to palbociclib (PD-0332991), a specific inhibitor of Cyclin D1 associated kinases CDK4/6 [19]. Treatment with increasing doses of palbociclib showed a consistent antiproliferative effect in thyroid cancer cells, with an IC$_{50}$ value ranging from $<$0.003 μM for BCPAP cells to 0.20 μM for HTC/C3 cells (Figure 7A and 7B). Representative images of thyroid cancer cell lines untreated or treated with 2.2 μM palbociclib for 72 h are reported in Figure 7C.

It has been reported that inhibition of MASTL induces multiple mitotic defects [20]. To investigate this issue in MASTL-depleted thyroid tumor cells, we performed the analysis reported in Figure 8. Asynchronous 8505C cells

Figure 5: Effect of MASTL, CCND1 and COPZ1 silencing on BCPAP cells. A. Western blot analysis of MASTL, Cyclin D1 and COPZ1 protein expression in Nthy-ori 3–1 cells, transiently transfected with siRNAs; actin was used as loading control. B. Scatter plot graph representing the percentage of cell growth (luminescence signal by CellTiter-Glo assay), normalized to that of Non Targeting oligo (siNT) 100%, 10 days after siRNA transfection in Nthy-ori 3–1 and BCPAP cells; siPSMC3 was used as positive control; one out of three representative experiments is shown. C. Representative pictures at 10X magnification of BCPAP cells 6 days after siRNA transfection. D. Cell proliferation assay of BCPAP cells 6 days after siRNAs transfection; the growth rate was determined by the trypan blue exclusion assay. Cell count of viable cells was normalized to that of siNT (100%); data represent the mean +/- sd of two independent experiments; the asterisks indicate differences significant by the unpaired Student’s t-test (***p < 0.01, ****p < 0.001). E. Western blot analysis of MASTL and Cyclin D1 expression performed in the same samples of panel D; actin was used as loading control for cell extracts.
were transfected with the siMASTL or NT siRNA oligos and, 48 hours later, analyzed by immunofluorescence for β-tubulin, phospho-histone H3, and DNA for the presence of mitosis and nuclei aberrations. Several abnormalities associated with MASTL knockdown [20] were observed in MASTL-depleted 8505C cells. These include:

Figure 6: Effect of CCND1, MASTL and COPZ1 silencing on different thyroid tumor cell lines. A. Scatter plot graph representing the percentage of cell growth (luminescence signal by CellTiter-Glo assay), normalized to that of Non Targeting oligo (siNT) 100%, 10 days after siRNA transfection in a panel of thyroid cell lines; siPSMC3 was used as positive control. One out of three representative experiment is shown. B. Cell proliferation assay 4 days (for TPC1, 8505C and HTC/C3 cells), 5 days (for KAT-18 cells), 6 days (for BCPAP cells), 7 days (for WRO82–1 cells) after siRNAs transfection; the growth rate was determined by the trypan blue exclusion assay. Cell count of viable cells was normalized to that of siNT (100%); data represent the mean +/− sd of two independent experiments; the asterisks indicate differences significant by the unpaired Student’s t-test (*P < 0.05, **P < 0.01, ***p < 0.001). C. Western blot analysis of MASTL, Cyclin D1 and COPZ1 expression performed in the same samples of panel B; actin was used as loading control for cell extracts.
multinuclear cells (panel B); abnormal mitotic figures, documented by the presence of condensed chromosomes that did not correctly align to metaphase plate (panel C); and chromatin bridges connecting daughter cells during anaphase (panel D). None of these abnormalities was observed in the control siNT cells (panel A). These results suggest that the growth inhibitory effect of MASTL silencing in thyroid tumor cells may be the consequence of failure of correct cell division process.

Shtutman et al [21] have recently reported that tumor cell COPZ1 vulnerability is related to downregulation of the paralogous COPZ2 gene. To assess whether this occurs also in thyroid tumor cells we investigated the expression of COPZ2 by Real Time PCR (Figure 9). With respect to control Nthy-ori 3–1 all the tumor cell lines analyzed, but WRO 82–1, showed downregulation of COPZ2 mRNA. This result indicates that for the majority of thyroid tumor cell lines the COPZ1 dependence may result from downregulation of COPZ2.

Collectively, our study identified Cyclin D1, MASTL and COPZ1 as thyroid cancer cell specific vulnerabilities, suggesting that they could provide a common therapeutic target in thyroid cancer treatment.

**DISCUSSION**

Despite much progress having been made in the genetic and molecular characterization of thyroid carcinoma, few or no therapeutic options are currently available for patients with aggressive and iodine-refractory thyroid tumors. Kinase inhibitors (KIs) in particular are being intensively studied in clinical trials in thyroid cancer: of note, various BRAF inhibitors for tumors that bear mutation of this gene [22–24], as well as a series of multi-kinase inhibitors, are emerging as potentially effective options in the treatment of advanced TC that are not responsive to traditional therapies [10]. However, the efficacy of KIs in patients with differentiated thyroid
carcinoma has given contrasting evidence in the clinical trials, probably due to the drug resistance (as in other tumor types). Furthermore, KIs might cause significant side effects. This underlines the need for further efforts aimed at the identification of novel therapeutic approaches for thyroid tumors [10].

Large-scale siRNA-based functional screening on cancer cell lines are nowadays widely used for the identification of tumor cells vulnerabilities to be explored for therapeutic purposes [12]. Interestingly, such approaches allow the discovery of target genes that are required by tumor cells in specific contexts, thus providing the premise for the identification of therapeutic approaches with no or limited effect on normal cells.

In this paper, we described, for the first time, a high-throughput siRNA phenotypic screening campaign for the identification of thyroid tumor cell vulnerabilities. By screening a “druggable genome” siRNA library, we identified 398 siRNA oligos (targeting 386 genes) which preferentially inhibited the growth of the BCPAP papillary thyroid cancer cell line as compared to that of non-transformed thyrocytes (Nthy-ori 3–1). Fifteen out of 28 hit genes were technically confirmed. The relevance of this screening study and its ability to identify bona fide cell vulnerabilities is supported by two lines of evidence. First, BRAF was detected among the top BCPAP vulnerability genes. This was expected, as the dependency of BCPAP cells upon BRAFV600E oncogene has been previously documented [16]. Second, the screen efficiently detected genes that were indiscriminately lethal for both the Nthy-ori 3–1 and BCPAP cell lines: in many cases, these were found to be genes that encode proteins of essential function and have been identified as survival genes in previous functional screenings [14, 18, 25].

Figure 8: MASTL silencing causes abnormal mitotic figures. 8505C cells transfected with the indicated siRNAs were analyzed by immunofluorescence for β-tubulin (green), phospho-histone H3 (red) and DAPI (DNA, blue). Arrows indicate normal mitosis (A), multinuclear cell (B), condensed chromosomes not correctly congressed to metaphase plate (C), DNA bridge (D) (Magnification 40x).
Analyzing the thyroid TCGA data set, we found no mutations affecting the confirmed hit genes, besides BRAF, suggesting that functional alterations in PTC may be excluded. With respect to gene expression the hit genes were found to be either equally or differentially (both upregulated and downregulated) expressed in PTC in comparison with normal samples.

Three hit genes, namely Cyclin D1, MASTL and COPZ1, were selected for further confirmation. Their vulnerability for BCPAP cells was confirmed by using siRNA oligos different from the screening. Interestingly, we showed that the silencing of Cyclin D1, MASTL and COPZ1 reduces the growth and survival of several different TC cell lines, regardless the histotype and the oncogenic lesion.

Cyclin D1 is an important cell cycle regulator that promotes G1/S phase transition by activation of CDK4 and CDK6 kinases [26]. Cyclin D1 is frequently overexpressed in human cancer, and it has been proposed as a therapeutic target. Several approaches aimed at its inhibition are presently ongoing. As for all cyclins, while the lack of enzymatic activity makes direct targeting of Cyclin D1 difficult, a much more amenable approach is to target cyclin functions by inhibition of their associated kinases. Indeed, several CDK4/6 inhibitors have shown promising antitumor activity in experimental systems, and are presently under clinical evaluation for different tumor types [26]. Alternative approaches are also currently under investigation, including the targeting of factors which regulate Cyclin D1 protein turnover, such as the USP2 deubiquitinase [27]. In thyroid carcinoma overexpression of Cyclin D1 at mRNA and protein level has been documented, and has been suggested to contribute to tumor progression [28–30]. However, no functional data dissecting the role of Cyclin D1 in thyroid tumorigenesis are available. We found that Cyclin D1 inhibition led to a consistent reduction of cell growth in all the thyroid tumor cell lines analyzed. This was not associated with Cyclin D1 expression level, suggesting that the dependency of thyroid tumor cells upon Cyclin D1 activity may not require aberrant protein expression. We also found that all the thyroid tumor cell lines are sensitive to palbociclib (PD-0332991), a selective CDK4/6 inhibitor approved for the treatment of ER+/HER2- breast cancer in combination with letrozole [31]. Whether the antiproliferative effect after Cyclin D1 silencing or palbociclib treatment in thyroid tumor cells, but not in normal ones, is related to the presence of functional pRB, as reported for other tumor types [19], remains to be investigated. Our studies identified Cyclin D1 as a therapeutic target for thyroid tumors, and provide a rationale for preclinical and clinical studies aimed to assess the efficacy of inhibitors of Cyclin D1 function in thyroid cancer.

MASTL (Microtubule associated serine/threonine kinase-like) regulates mitosis progression by inhibiting PP2A/B55δ, the principal protein phosphatase complex that dephosphorylates CDK1 substrates. MASTL activity is essential for prevention of defects in chromosome condensation and segregation, prometaphase arrest and mitotic collapse during cell division [32]. Furthermore, it modulates DNA damage response by promoting checkpoint recovery and cell cycle progression [33]. Little is known about MASTL in cancer. Nagel et al. [34] reported that the knockdown of MASTL, through promoting defects in in cytokinesis and a shortened G2/M phase arrest, is capable of sensitizing lung cancer cells to radiations. Notably, MASTL inhibition affects tumor cell but not normal primary fibroblast radiosensitivity, thus suggesting that MASTL could represent a druggable target to combine with radiotherapy. In head and neck carcinoma MASTL was found involved in progression and recurrence and proposed as therapeutic target [35].

![Figure 9: Expression of COPZ2 in thyroid tumor cell lines.](image_url) Real-time PCR analysis of COPZ2 gene expression; results are presented as log10-transformed relative quantity (RQ) of COPZ2 mRNA normalized for HPRT1 housekeeping gene expression. Data represent the mean +/- sd of three independent experiments.

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have identified MASTL as vulnerability gene for thyroid carcinoma. In fact, MASTL silencing interfered with the growth of all thyroid cell lines, although with variable extents. Moreover, evidence of failure of correct cell division process was found in MASTL-depleted cells. Of note, our study identified MASTL as a novel mitotic division process in thyroid tumor, in addition to PLK1 and AURKs, both overexpressed and proposed as therapeutic targets in ATC [36, 37]. Remarkably, we found that MASTL vulnerability in thyroid tumor cell lines is not associated with overexpression; on the contrary, MASTL is slightly downregulated in PTC samples and in thyroid tumor cells. Overall these findings highlights the need of further investigation in order to assess whether MASTL may represent a useful target in different tumor types, as well as the search/design of strategies to block the activity of MASTL, for which no specific inhibitors have been so far identified.

COPZ1 (coatamer protein complex ζ1) is a subunit of coatamer protein complex I (COPI), a secretory vesicle coat protein complex involved in Golgi apparatus and endoplasmic reticulum traffic, endosome maturation, autophagy [38, 39] and lipid homeostasis [40]. COPZ1 has been proposed as a tumor-specific gene target. It has been shown that tumor cells become dependent on COPZ1, as a result of a tumor-specific downregulation of its isoform COPZ2 [21]. Interestingly, COPZ1 knockdown kills both proliferating and non-dividing tumor cells, but does not affect normal cells, thus suggesting that COPZ1-targeting therapies have the potential for eradicating cancer cells, independently of whether they are actively proliferating. Our screening and validation identified COPZ1 as a thyroid tumor cell specific survival gene. Interestingly, we also found a significant downregulation of COPZ2 mRNA expression in the majority of thyroid tumor cell lines. An exception is represented by WRO82–1 which, despite expressing COPZ2 mRNA levels higher than control, is sensitive to COPZ1 silencing. This raises the possibility that other mechanisms, in addition to COPZ2 downregulation, may be involved in determining tumor cell COPZ1 dependency. Whether the thyroid tumor cell dependency upon COPZ1 is related to downregulation of COPZ2 isoform, or to deregulation of other COPI components, as well as the modalities through which COPZ1 inhibition leads to growth suppression of thyroid tumor cells, remains to be investigated.

In conclusion, by a siRNA-based functional screening we have identified a signature of vulnerability for thyroid carcinoma cells, thus identifying novel potential therapeutic targets in this setting. Further characterization of three hit genes, namely Cyclin D1, MASTL and COPZ1, demonstrates that their abrogation selectively impairs the growth of several thyroid tumor cell lines, irrespective of the histotype or driving genetic lesion, thus indicating that they represent common examples of “non-oncogene addiction” in thyroid cancer. Variable extents of cell growth inhibition among the cell lines upon silencing of these target genes were observed. Whether these differences are related to genetic background and/or intrinsic sensitivity to cell death remains to be investigated. The dependency upon Cyclin D1, MASTL and COPZ1 activity is restricted to tumor cells and not normal thyroid cells, as previously reported for other tumor types [21, 27, 34, 35]. Thus we envisage that Cyclin D1, MASTL and COPZ1 are attractive targets for new therapeutic approaches for thyroid cancer which spare normal cells and which would thus have limited side effects.

MATERIALS AND METHODS

Cell lines

Nthy-ori 3–1 cell line (SV-40 immortalized normal human thyroid follicular cells) was purchased from European Collection of Cell Cultures (ECACC) (Salisbury, UK); BCPAP, TPC-1, WRO82–1, 8505C, KAT-18 were obtained from Prof. A. Fusco (University Federico II, Naples, IT); HTC/C3 was purchased from Riken Cell Bank (Tsukuba, Japan).

Nthy-ori 3–1 and the PTC-derived BCPAP cell lines were cultured in RPMI 1640 medium (Gibco Life Technologies, Carlsbad, CA, USA) supplemented with 10% (v/v) heat-inactivated fetal bovine serum (FBS) (EuroClone, Pero, Italy), penicillin (100 U/ml) and streptomycin (100 mg/ml). The other cell lines PTC-derived TPC1; FTC-derived WRO82-1; ATC-derived 8505C, KAT18 and HTC/C3 were maintained in DMEM (Gibco, Life Technologies, Carlsbad, CA, USA) medium containing 10% FBS and 2 mM glutamine. All cell lines were cultured as monolayer at 37°C in a 5% CO₂ humidified atmosphere.

Cell lines were genotyped at the Fragment Analysis Facility of Fondazione IRCCS Istituto Nazionale dei Tumori, using Stem Elite ID System (Promega Corporation, Madison, USA) according to the manufacturer’s instructions and ATCC guidelines. The profiles obtained matched to their original profiles [41, 42]. WRO82–1 profile matched to that reported by Xu et al. [43] and HTC/C3 to that reported by JCRB Cell Bank (http://cellbank.nibio.go.jp/legacy/celldata/jcrb0164.htm).

Mycoplasma contamination was tested periodically and found negative in all cell lines (PCR Mycoplasma Detection Set, TAKARA Bio Inc).

siRNA transfection optimization

Optimal conditions for reverse siRNA transfection of Nthy-ori 3–1 and BCPAP cells were determined by comparing cell viability after transfecting a positive control siRNA targeting the proteasome subunit PSMC3, and a negative control non-targeting oligo (siNT), in the presence of different cell densities and Lipofectamine
RNAiMAX reagent (Invitrogen Life Technologies Carlsbad, CA, USA) concentrations. The best reagent and transfection conditions were those that produced the least (<10%) cell loss with siNT, and the greatest (>90%) cells decrease with the lethal PMSC3 oligo.

**High throughput siRNA screening**

The human Silencer Select Druggable Genome siRNA Library V4 (Ambion Life Technologies, Carlsbad, CA, USA) is composed by 309 96-well plates containing 25139 unique lyophilized siRNA oligos targeting 9031 human genes (three oligos per gene, on average). Stock plates were obtained by dissolving oligonucleotides with 100 μl of DEPC-treated water (Ambion Life Technologies, Carlsbad, CA, USA) (final concentration 2.5 μM) and by adding NT and PSMC3 siRNA oligos (2.5 μM) in alternate position in the empty column 12 of each plate (see Figure 1A). Library oligonucleotides were further diluted to 800 nM with DEPC-treated water (mother plates). Daughter plates were prepared by transferring 17 μl/well of mother plate oligonucleotide solution in plates prefilled with 119 μl of Opti-MEM® I medium (Gibco Life Technologies, Carlsbad, CA, USA) containing 4 μl/ml of lipofectamine® RNAiMAX. The solution was gently mixed by pipetting and incubated for 20 minutes room temperature, and then 30 μl/well were split into four transfection plates, to which 120 μl/well of cell suspension (1250 cells/ml) were added. Two plates for each cell line (Nthy-ori 3–1 and BCPAP) were used. Each well contained 150 cells, 20 nM oligo, RNAiMAX 0.1% vol/vol. After transfection, plates were kept at room temperature for 15–20 minutes to minimize uneven distribution of the cells in each well [44], and then incubated at 37°C for 7–8 days by stacking in alternate orientations to limit uneven medium evaporation and to facilitate gas exchange. As a result of this procedure, no edge effects or evidences of asymmetric cell growth emerged from screening data analysis, and visual inspection of plate scans confirmed homogeneous cell distribution in all plates. The following custom-barcoded 96-well microplates purchased from Greiner Bio-One (Frickenhausen, Germany) were used: U-bottom microplate for mother and daughter plates; CELLSTAR®, Black /μClear® plate for transfection plates. All liquid handling and plate tracking were performed with a Freedom EVO® robotic platform (Tecan, Männedorf, Switzerland); disposable MCA384 Tips and reservoirs were purchased from Tecan.

**siRNAs**

Information on siRNAs used in the transfection setup, as control in the screening, or in confirmation and validation studies is summarized in the Table S4. Oligos produced in house were designed and synthesized with stabilizing modifications (i.e.: phosphorylation in 5’ position of the antisense strand and 2’-O-methyl ribosyl substitution at position 2 of the sense strand) known to reduce off-target effect [45].

**Colony formation assay (CFA)**

Seven-eight days after transfection, cells were fixed with formaldehyde 3.7% v/v solution at room temperature for 30 minutes, washed with PBS and stained for 20 minutes with a PBS solution containing 0.05% v/v Triton® X-100 (Sigma-Aldrich, St. Louis, MO, USA) and TO-PRO®-3 iodide (Invitrogen Life Technologies (Carlsbad, CA, USA) diluted 1:1000. After a second wash, 50 μl PBS were left in each well. Plates were sealed and analyzed with an Odyssey® infrared scanner (LI-COR Biosciences, Lincoln, NE, USA) which provides whole-well fluorescence quantification of cells stained with TO-PRO®-3 dye. The fluorescence intensity (FI) was considered as readout, which is proportional to both colony number and size.

**Screening data analysis**

FI values were registered in a proprietary Oracle database through Symyx Assay Explorer® and matched with the library information. The FI of each sample was divided by the average FI of the three non-targeting oligos within the same plate (set as 100%) for interplate normalization. TIBCO SpotFire (Boston, MA, USA) and Microsoft Excel (Redmond, WA, USA) were used for statistical analysis and visualization of screening data distributions. The means and standard deviations of FI (% siNT) values and z*-scores [46] were calculated from the two technical replicates for each cell line and reported in Table S1 (Excel file with screening results).

To identify genes whose RNAi-mediated inhibition affects the growth of BCPAP but not Nthy-ori 3–1 cells, a distance factor (d) expressed as:

\[
    d = \sqrt{(100 - %NT_{\text{Nthy-ori } 3-1})^2 + (%NT_{\text{BCPAP}})^2}
\]

was introduced. “d” represents the distance between the %NT value of any given siRNA oligo and the point of maximal lethality towards BCPAP without effects on Nthy-ori 3–1 (coordinate %NT_{\text{Nthy-ori } 3-1} = 100, %NT_{\text{BCPAP}} = 0). Screening hits were selected among siRNA oligos with significantly low d values, whereas oligos either equally inactive on both cell lines, or equally lethal on both cell lines, or preferentially lethal on Nthy-ori 3–1, are excluded. Based on the distribution analysis of d scores, a -3σ threshold (corresponding to d = 47.2) in the primary screening, and a -2σ threshold (corresponding to d = 49.8) in the confirmation study, were applied to define hits.

**Cell proliferation assay**

Cells (500 cells/well) were reverse transfected in 96-well plate with 20 nM specific siRNAs using
the Lipofectamine RNAiMAX reagent, according to manufacturer’s instruction. Ten days later, cell viability was assessed by CellTiter-Glo Luminescent Cell Viability Assay (Promega Corporation, Madison, USA), performed as recommended by the supplier. Luminescence signals were acquired using a microplate reader (TecanUltra, Tecan Trading AG, Switzerland).

For the trypan blue exclusion assay, BCPAP and 8505C (1 x 10^6), TPC-1 (6 x 10^5), WRO82–1, HTC/C3 and KAT-18 (8 x 10^5) cells were transfected the day later in 6-well plate with 20 nM of specific siRNAs using Lipofectamine RNAiMAX, according to manufacturer’s instruction. At different days after transfection (4 days for TPC1, 8505C and HTC/C3 cells; 5 days for KAT-18 cells, 6 days for BCPAP cells, 7 days for WRO82–1 cells) the fraction of viable and dead cells was analyzed. For palbociclib experiments, cells (8000 cells/well) were seeded in black/blank bottom 96-well plates and, the following day, treated with increasing doses of the palbociclib (PD 0332991) inhibitor (Selleck Chemicals, Houston, TX, USA, cat. S2768). After 72 hours, cells were fixed with 3.7% paraformaldehyde solution for 30 minutes at RT, washed and stained with 1 mg/ml Hoechst 33342. The plates were analyzed with an ArrayScan VTI high-content screening reader (Thermo-Fisher Scientific, Pittsburgh, PA): at least 800 cells were acquired in each well with a 10X objective in one fluorescence channel (XF93 optical filter set). Cell nuclei were automatically recognized and counted in ten fields to estimate the cell number per well as proliferation readout. IC50 values were calculated by sigmoid curve fitting of experimental points.

**Gene Ontology and gene expression analyses**

The functional annotation analysis on the identified gene lists was performed by means of the Database for Annotation, Visualization and Integrated Discovery (DAVID) tool v6.7 (https://david.ncifcrf.gov/). Gene Ontology (GO) Biological Process (BP) and Molecular Function (MF) terms were used as annotation categories and medium classification stringency was set for the functional annotation clustering analysis. The annotation clusters with an Enrichment score (ES) >1.3 were selected and representative GO-terms were indicated for each significant cluster. To assess the interactions between the validated fifteen genes, network analysis was carried out using the Ingenuity Pathway Analysis Tool (Ingenuity® Systems, http://www.ingenuity.com). The Core Analysis function was used to compare genes pooled from literature and other molecules in IPA’s database.

The thyroid TCGA data set, containing the molecular data on nearly 500 PTC, is the largest data collection so far available (https://tcgadata.nci.nih.gov/tcga/tcgA). Illumina HiSeq Level 3 data were downloaded from TCGA Data Portal and RSEM [47] gene normalized expression values of selected genes were derived for 58 PTC patients, in tumor and adjacent normal samples, respectively. The statistical significance of gene expression difference between PTC and normal groups was assessed using Wilcoxon test.

**RNA extraction, RT-PCR and Real time PCR**

RNA extraction and RT-PCR were performed as previously described [48], using the following primer: 5′-ATGAGAGGCTCTTGAGACACT-3′ (forward), 5′-TCATGTCATGACTTGACAGGAG-3′ (reverse) for RGS3 fragment amplification; 5′-GCAGATCTTACGTACGTAG-3′ (forward), 5′-TATAATGGATCTACAGGGTG-3′ (reverse) for RASD1; 5′-ATGTTGACCGTCTCAGGAT-3′ (forward), reverse 5′-CAGTGCCACGTCTCTGGT-3′ (reverse) for OXTR [49].

For Real Time PCR analysis 20 ng of retrotranscribed RNA were amplified in PCR reactions carried out in triplicate on an ABI PRISM 7900 using TaqMan gene expression assays (Applied Biosystem, Foster City, CA). Hs00212698_m1 was used for COPZ2 expression; human HPRT1 (HPRT-Hs99999909_A1) was used as housekeeping gene for the normalization among samples. Data analysis was performed using the SDS (Sequence Detection System) 2.4 software.

**Western blot analysis**

Western blot analysis was performed as previously described [48], using the following antibodies: anti-Cyclin D1 (DCS-6: sc-20044, 1:500), anti-COPE, (E-20: sc-12104, 1:500), anti-PLA2G15 (LYPLA3, H-167: sc-135297, 1:500), anti-MAP4K5 (KHS, N-19: sc-6429, 1:1000), anti-COPZ1 (D-20: sc-13349, 1:750), anti-GAPDH (6C5, sc-3223: 1:2000) purchased from Santa Cruz, CA, USA; anti-MASTL (ab86387, 1:7000) from Abcam, Cambridge, UK; anti-β-actin (A2066, 1:5000) and anti-vinculin (V9131, 1:1000) from Sigma-Aldrich, St Louis, Mo, USA.

**Immunofluorescence analysis**

Cells growing on glass coverslips were transfected with 20 nM NT or MASTL siRNAs and, 48 hours later, fixed for 10 minutes with 4% paraformaldehyde (Sigma Aldrich, St Louis, Mo, USA). After permeabilization for 10 minutes with 1% BSA and 0.1% Triton X-100 in PBS, and incubation for 30 minutes with 1X blocking solution (2% BSA in PBS), cells were incubated with anti-β-tubulin (clone TUB 2.1: T4026, 1:400, Sigma Aldrich, St Louis, Mo, USA) or anti-histone H3 (phospho S10) (E173: ab32107, 1:1000, Abcam, Cambridge, UK) primary antibodies for 1 hour. After washing with PBS, cells were incubated with Alexa Fluor® 546 rabbit
(1:500, Invitrogen/Molecular Probes®) and Alexa Fluor 488® mouse (1:500, Invitrogen/Molecular Probes®) secondary antibodies for 1 hour. Slides were then prepared using ProLong Diamond Antifade mountant with DAPI (P36966, Molecular Probes®) and imaged with immunofluorescence microscopy (Eclipse E1000; Nikon Instruments, Inc. NY, USA).

ACKNOWLEDGMENTS AND FUNDING

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CONFLICTS OF INTEREST

The authors declare no conflict of interest.

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**Supplementary Figure S1:** A. Pie chart of functional annotation clustering analysis on 386 genes by DAVID 6.7 tool. Only 15 annotation clusters with an Enrichment Score (ES) > 1.3 are reported. A representative Biological Process (BP) or Molecular Function (MF) GO-term and the percentage of the genes on the 382 DAVID IDs globally identified are indicated for each significant annotation cluster; for each selected cluster the ES is also indicated. B. Visual representation of the two principal networks generated by Ingenuity Pathway Analysis (IPA) including the fifteen validated hits. Solid lines represent a direct interaction between the two gene products and dotted lines mean an indirect interaction.
Supplementary Figure S2: western blot analysis of MASTL protein expression in Nthy-ori 3–1 cells, transiently transfected with MASTL#5 siRNA; actin was used as loading control.
Supplementary Figure S3: representative pictures at 10X magnification of TPC-1, WRO82–1, 8505C, HTC/C3 and KAT-18 cells, taken before the trypan blue exclusion assay.
**Supplementary Table S1:** List of 25139 siRNA oligonucleotides of the human Silencer Select Druggable Genome siRNA Library V4 (Ambion Life Technologies, Carlsbad, CA, USA) and screening results. Average and standard deviation (StDev) of the fluorescence intensity (FI) normalized on siNT (%NT), and average and standard deviation (StDev) of z* scores for BCPAP and Nthy-ori 3–1, distance factor (\(d\)) and binned \(d\) (based on \(\sigma\) distribution) are shown. The gene symbols, full gene names, Refseq (http://www.ncbi.nlm.nih.gov/refseq/about/) accession numbers, siRNA batches and their corresponding plate blocks were provided by Ambion.
Supplementary Table S2: List of 398 siRNA oligonucleotides resulted to preferentially inhibit the proliferation of BCPAP cells with respect to Nthy-ori 3–1 (d < -3σ) and thus defined as “differential hits”.
Supplementary Table S3: List of 1695 siRNA oligonucleotides resulted to inhibit the proliferation of both BCPAP and Nthy-ori 3–1 (signal intensity < 20% siNT), therefore defined as “lethal hits”.
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<td>in house</td>
<td>transfection setup, screening, confirmation</td>
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<tr>
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<td>5′-CGGCUGAAGUGGCAUAAAtt-3′</td>
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<td>validation</td>
</tr>
</tbody>
</table>

List of siRNA oligos used in the transfection setup, as control in the screening, or in confirmation and validation studies
Appendix B
Targeting COPZ1 non-oncogene addiction counteracts the viability of thyroid tumor cells

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**Targeting COPZ1 non-oncogene addiction counteracts the viability of thyroid tumor cells**

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**Running title:** Targeting COPZ1 in thyroid tumors

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Abstract

Thyroid carcinoma is generally associated with good prognosis, but no effective treatments are currently available for aggressive forms not cured by standard therapy. To find novel therapeutic targets for this tumor type, we had previously performed a siRNA-based functional screening to identify genes essential for sustaining the oncogenic phenotype of thyroid tumor cells, but not required to the same extent for the viability of normal cells (non-oncogene addiction paradigm). Among those, we found the coatamer protein complex ζ1 (COPZ1) gene, which is involved in intracellular traffic, autophagy and lipid homeostasis. In this paper, we investigated the mechanisms through which COPZ1 depletion leads to thyroid tumor cell death. We showed that siRNA-mediated COPZ1 depletion causes abortive autophagy, endoplasmic reticulum stress, unfolded protein response and apoptosis. Interestingly, we observed that mouse tumor xenografts, locally treated with siRNA targeting COPZ1, showed a significant reduction of tumor growth.

On the whole, we demonstrated for the first time the crucial role of COPZ1 in the viability of thyroid tumor cells, suggesting that it may be considered an attractive target for novel therapeutic approaches for thyroid cancer.

Keywords: thyroid carcinoma, COPZ1, non-oncogene addiction, cell death
1. Introduction

Cancer cells often depend on non mutated genes and pathways to support their unchecked growth. The activity of these genes is essential for tumor cells, but not required to the same extent by normal cells; this concept is known as "non-oncogene addiction" (NOA) paradigm [1]. NOA genes represent tumor vulnerabilities and their inhibition results in the failure of the oncogenic phenotype. Nowadays, large-scale siRNA-based functional screening of cancer cell lines represent a powerful strategy for the identification of NOA genes that can be investigated as therapeutic targets for many types of tumors [2]. We used this approach to unveil nodal points for therapeutic intervention for thyroid carcinoma (TC), which represents the most frequent endocrine cancer, with a rapidly increasing incidence [3].

The majority of TC originates from epithelial cells and includes well-differentiated papillary (PTC) and follicular (FTC) carcinomas, poorly differentiated (PDTC) and undifferentiated anaplastic carcinomas (ATC). Most TC are effectively treated by standard therapy, involving surgery, thyroid stimulating hormone and radioiodine. Nevertheless, a fraction of patients cannot be cured due to local recurrence (in up to 20% of patients), distant metastasis (in approximately 10% at 10 years [4]) and/or radioresistant disease. Furthermore, patients with PDTC and ATC have a very poor prognosis, with survival of few months in the case of ATC [5;6]. Even though recent molecular findings allowed the development of several therapies designed to specifically target thyroid oncoproteins or their downstream pathways, this strategy showed modest results and only partial response [7;8]. Few or no therapeutic options are currently available for patients with aggressive and iodine-refractory thyroid tumors. Thus, there is a need to better understand the mechanisms of thyroid carcinogenesis and to improve the treatment of the most aggressive tumor forms not curable by standard therapy.

We have recently discovered several thyroid tumor cell vulnerabilities. By screening a siRNA library, we identified a set of genes whose silencing inhibited the growth of a panel of thyroid tumor cells, but not of normal immortalized thyrocytes. The COPZ1 gene was found among the top ranked
genes [9]. COPZ1 (coatamer protein complex ζ1) belongs to the coatamer protein complex I (COPI), an heptameric complex which is involved in: assembly of coated vesicles on Golgi membranes, retrograde transport of luminal and membrane proteins in the ER-Golgi secretory pathway, endosome maturation, autophagy [10;11] and lipid homeostasis [12]. It has been demonstrated that COPZ1 knockdown causes cell death in both proliferating and nondividing tumor cells. Unlike tumor cells, normal cells are not sensitive to COPZ1 inhibition [13]. We identified COPZ1 as a thyroid tumor cell specific survival gene, as its inhibition induced cell death in tumor cell lines but not in immortalized thyrocytes. This evidence, together with a not significant variation of expression and absence of mutations in PTC, allowed us to classify COPZ1 as an example of "non-oncogene addiction” for thyroid cancer cells [9].

Here we investigated the mechanisms through which COPZ1 depletion leads to thyroid tumor cell death and explored its potential use as therapeutic target for thyroid carcinoma. We found that cell death induced by COPZ1 depletion is associated with abortive autophagy, ER stress, UPR and apoptosis. Interestingly, local treatment with siCOPZ1 oligos reduced tumor growth in an in vivo model of thyroid carcinogenesis.
2. Materials and Methods

2.1 Cell lines
Thyroid tumor cell lines, primary thyrocytes and mammary epithelial h-TERT-HME1 cells were cultured as previously described [9;14;15]. Source and culture conditions are reported in Supplementary. GFP-LC3 stable cell lines were generated by transfecting 2µg of pEGFP-LC3 plasmid (from Tamotsu Yoshimori’s laboratory [16], using the Lipofectamine 3000 reagent (Invitrogen Life Technologies, Carlsbad, CA, USA) for TPC-1, and Lipofectamine LTX (Invitrogen Life Technologies) for 8505C cells, according to manufacturer’s instruction. Transfected cells were selected and propagated in medium containing G418 (500µg/ml, Invitrogen Life Technologies).

2.2 Detection of LC3 puncta
Cells growing on glass coverslips were transfected as described below and 24h, 48h, 72h later fixed for 20 minutes with 4% paraformaldehyde. After washing with PBS, cells were treated with the ProLong Diamond Antifade mountant with DAPI (P36966, Molecular Probes, Invitrogen Life Technologies) and imaged with immunofluorescence microscopy (Eclipse E1000; Nikon Instruments, Inc. NY, USA). The percentage was calculated as the ratio between cells displaying LC3 puncta and total cells (at least 150), considering six randomly selected 60X fields.

2.3 siRNA reagents and transfection
siRNA transfection was performed using 20 nM of siRNA oligos (see list in Supplementary) and the Lipofectamine RNAiMAX reagent (Invitrogen Life Technologies), according to manufacturer’s instruction.

2.4 Colony formation assay
Cells were plated in 100 mm dishes and transfected the following day with 20 nM of siRNAs. One day later, cells were collected and plated as follows: 2x10^4 cells in 24 well-plates for colony
formation assay; 1.5x10^5 cells in 60 mm dishes for western blot analysis. For the colony formation assay, cells were fixed five days later with formaldehyde 3.7% v/v solution for 30 minutes, washed with PBS and stained for 20 minutes with 0.1% crystal violet (w/v). Pictures were taken using an Epson image scanner.

2.5 Western blot analysis

WB analysis was performed as previously described [17]. The list of antibodies is provided in Supplementary.

2.6 RNA extraction and Real time PCR

RNA extraction and Real time PCR were performed as previously described [17]. The following TaqMan gene expression assays (Applied Biosystem, Foster City, CA) were used: Hs01023197_m1 for COPZ1 and Hs00358796_g1 for CHOP expression; Hs02800695_m1 for HPRT1, used as housekeeping gene for normalization among samples.

2.7 Apoptosis assay

Cells were transfected with 20 nM of siRNA oligos in the presence of 2 µM of the fluorogenic substrate for activated caspase-3/7 (CellEvent™ Caspase-3/7 Green Detection Reagent, Invitrogen Life Technologies). Live-cell fluorescence images were taken 72h later with microscope (Eclipse TE2000-S; Nikon Instruments).

2.8 In vivo studies

Female CD-1 nu/nu mice (5-weeks old) (Charles River, Calco, Italy) were injected subcutaneously into the left flank with 8505C cells (10x10^6). When xenograft tumors reached the mean volume of 0.085 mm^3 (range 0.040-0.144), they were randomly divided into three groups and then locally treated with 20 µg of siCOPZ1 oligos (4457308, Ambion® ID s22427) or non-targeting oligos
(4457289, Ambion® \textit{In Vivo} Negative Control #1 siRNA) mixed in 50 µl of MaxSuppressor\textsuperscript{TM} \textit{In Vivo} RNA-LANCEr II reagent (Bioo Scientific, Austin, TX, USA), or with MaxSuppressor\textsuperscript{TM} only. Treatments were carried out at day 17, 20, 23, 27, 30 after cell injections. Tumor growth was followed for 34 days and assessed by monitoring tumour weight (TW), as previously described \cite{18}. Animal studies were reviewed and approved by the Ethics Committee for Animal Experimentation of the hosting institution and are in accordance with the guidelines of the UK Coordinating Committee for Cancer Research \cite{19}.

2.9 \textit{Immunohistochemistry}

Serial sections from paraffin-embedded tumor xenografts (2-µm thick) were stained with hematoxylin and eosin and evaluated under a light microscope to assess the pattern of tumor growth. Antigen retrieval was performed using 1 mM citrate buffer (pH 6), then sections were immunostained with a primary rat polyclonal antibody anti-human COPZ1 (1:400 SAB4500896 Sigma-Aldrich St Louis, Mo, USA), anti-human Ki-67 (Mib1) (1:200 M7240 Dako, Agilent Technologies, CA, USA). Immunostains were performed using standard immunoperoxidase protocol followed by diaminobenzidine chromogen reaction (Dako \textsuperscript{TM} REAL\textsuperscript{TM} EnVision\textsuperscript{TM} Detection System, K5007 Dako). The intensity of staining for COPZ1 was scored as low (1+), medium (2+) or high (3+). The percentage of cells immunostained for Ki67 and necrotic cells was estimated counting one 10X field, randomly selected. One sample of each group of tumor explants was excluded from the IHC analysis due to technical problems.

2.10 \textit{Gene/miRNA expression analysis}

Gene expression data of \textit{COPZ2} gene (log scale) from 58 normal-thyroid tissue samples compared to 31 ATC, 72 PTC and 17 PDTC samples, were obtained combining GSE3467 \cite{20}, GSE6004 \cite{21}, GSE33630 \cite{22;23} and GSE76039 \cite{24} publicly available datasets. Raw intensity expression values on U133 Plus 2.0 array (Affymetrix, Santa Clara, CA) were processed by Robust Multi-array
Average procedure [25], with the re-annotated Chip Definition Files from BrainArray libraries version 20.0.0 [26], available at http://brainarray.mbni.med.umich.edu. Batch effects were removed by using sva R/Bioconductor package [27]. The analysis of the thyroid TCGA dataset from 58 paired normal/PTC samples was performed as previously described [9]. Normalized reads per million miRNA mapped (RPM) data were obtained for the same 58 tumor samples from Illumina HiSeq Level 3 isoform quantification files (TCGA Data Portal website), by summing up the reads aligned to each 3p or 5p miRBase v16 mature strands [28;29]. Pearson’s correlation method was applied to evaluate the correlation between RNAseq normalized gene and miRNA read counts (log scale) in 58 tumor samples.

3. Results

3.1 COPZ1 depletion affects thyroid tumor cell viability

The issue that COPZ1 represents vulnerability for thyroid tumor cell lines but not for normal thyrocytes [9] is further documented by the colony forming assay in Figure 1A. Upon transfection of siRNA oligos targeting COPZ1 (siCOPZ1), a drastic growth inhibition was observed in thyroid tumor cell lines (PTC-derived: BCPAP and TPC-1; FTC-derived: WRO82-1; ATC-derived: 8505C, KAT-18, and HTC/C3) in comparison to non-targeting (siNT) transfected cells, and to COPZ1-depleted Nthy-ori 3-1 cells, normal immortalized thyrocytes. Western blot (WB) analysis performed in parallel documented the efficient COPZ1 silencing in all the cell lines (bottom panel). SiRNA transfected TPC-1, 8505C, HTC/C3, and Nthy-ori 3-1 cell lines were also analyzed by time-lapse phase-contrast video microscopy (Supplementary Videos). Unlike siNT-transfected cells, COPZ1-depleted tumor cells showed morphological changes, such as shrinking and rounding up 30 hours (h) (for TPC-1 cells) and 48h (for 8505C and HTC/C3 cells) after silencing. Then, cells detached from dishes and reached massive cell death at the end of the experiment (68h for TPC-1 cells, 89h for 8505C and HTC/C3 cells). None of these effects was observed in COPZ1-depleted NThy-ori 3-1 cells, which proliferated as the control (siNT) and reached confluence at the end of the experiment (68h). In Figure 1B, representative images of each cell lines at the end of experiment are reported.
Further images documenting cell morphological changes after COPZ1 depletion are shown in Supplementary Figure S1. We also showed the lack of effect of COPZ1 silencing on primary human thyrocytes and human mammary epithelial HME-1 (Figure 1C): COPZ1 depletion leaves unaffected the growth of these non malignant cells, similarly to the negative control siNT; on the contrary, a massive cell death was observed in cells transfected with siRNA oligos targeting Ubiquitin B (siUBB), used as control. WB analysis 72h after siRNA transfection documented the complete abrogation of COPZ1 protein expression.

3.2 COPZ1 depletion affects autophagy

COPI complex members are involved in autophagy [11] and are required for completion of the autophagic process. Moreover, the depletion of COPI members impairs productive autophagy, as demonstrated in prostate, breast and ovarian carcinoma cells [13;30]. TPC-1 and 8505C cell lines, stably expressing the autophagosome marker GFP-LC3 (hereafter referred as TPC1-GL3 and 8505-GL3) were transfected with siCOPZ1 and siNT oligos, and analyzed at different time points by fluorescence microscopy for the presence of the typical autophagy-associated LC3 puncta (Figure 2A). The number of siNT-transfected cells displaying LC3 puncta remained low over the time. On the contrary, the number of COPZ1-depleted cells presenting puncta significantly increased over the time. In TPC1-GL3 cells puncta were detected in 5.7% and 38% at 24h and 48h, respectively; the analysis at 72h was not feasible, as cells detached from glass coverslip due to the lethal effect of COPZ1 silencing. In 8505-GL3 cells, puncta were present in 6%, 14% and 34%, at 24h, 48h and 72h, respectively. In Figure 2B, representative images of COPZ1-depleted cells displaying LC3 puncta and relative controls are shown. Autophagosome formation upon COPZ1 silencing was also demonstrated by assessing the conversion of the endogenous LC3-I to LC3-II by WB. Accumulation of LC3-II protein was observed in COPZ1-depleted naive TPC-1 and 8505C cells. In both cell lines, the endogenous LC3-II was barely detectable at 48 h and progressively increased at the latest time points (Figure 2C). Since an increased steady-state expression of LC3-II could
suggest persistent accumulation of autophagosomes, impaired recycling and consequently block of the autophagic flux, we investigated whether COPZ1 depletion causes abortive autophagy. We monitored by WB whether LC3-II accumulation was associated with increased expression of SQSTM1/p62, an LC3-binding protein mainly degraded through autophagy catabolism and whose increase is a marker of abortive autophagy [31]. We observed an increase of SQSTM1/p62 in both COPZ1-depleted cell lines in comparison to controls (Figure 2D). Such increase was 1.4 and 1.6 fold in TPC-1 cells at 72h and 96h, and 2.9 and 2.5 fold in 8505C cells. The persistent accumulation, rather than turnover/degradation of autophagosomes, and the concomitant increase of SQSTM1/p62 protein expression suggest that COPZ1 depletion is associated with abortive autophagy.

3.3 COPZ1 depletion induces endoplasmic reticulum stress and unfolded protein response

As COPZ1 regulates vesicular transport, its depletion could be associated with protein accumulation in the endoplasmic reticulum (ER) and with a deficit in proteins coming back from the Golgi. Since these effects could be linked to the activation of ER stress and of unfolded protein response (UPR), we investigated the expression of some components of these pathways. COPZ1-depleted TPC-1 and 8505C cells showed an increased expression of the chaperone GRP78/BiP, which, upon ER stress, is the major responsible for the activation of downstream UPR signals [32;33]. In TPC-1 cells, the level of GRP78/BiP increases by 1.5, 2.8 and 11 fold at 48h, 72h and 96h after siRNA transfection, respectively, in comparison to the control; whereas, in 8505C cells the increase was by 1.8, 3.9 and 3.2 fold (Figure 3A). Concomitantly, we observed an increase of another marker of ER stress, the ER lumen chaperone calnexin (1.2 at 48h and 1.6 from 72h fold increase in TPC-1; 1.4 and 2.8 fold increase in 8505C cells) (Figure 3A). CHOP, also known as GADD153, is transcriptionally upregulated during ER stress and UPR [34;35]. We analyzed the expression of CHOP and COPZ1 by Real Time PCR in TPC-1 and 8505C cells at different time points after COPZ1 silencing. As shown in Figure 3B, COPZ1 depletion led to a drastic increase of CHOP expression in comparison
to the control. In conclusion, our data indicate that COPZ1 depletion causes the onset of ER stress and UPR.

3.4 COPZ1 depletion induces apoptotic cell death

Prolonged ER stress or failure of adaptive response can result in cell death [36]. Moreover, CHOP activation triggers apoptotic death [34;35]. TPC-1 and 8505C cells were transfected with siCOPZ1 or siNT oligos in the presence of the fluorescent substrate for activated caspase-3/7. Live-cell fluorescence microscopy performed 72h after transfection showed a significant increase of Caspase-3/7 activity in COPZ1-depleted cells in comparison to the siNT control (Figure 4A). The induction of apoptosis by siCOPZ1 is also documented by the WB analysis in Figure 4B: COPZ1 depletion increases cleaved PARP and cleaved caspase-3 levels, which were detectable at 48h and 72h after siRNA transfection in TPC-1 and 8505C cells, respectively. Overall, these results suggest that the depletion of COPZ1 induces caspase-dependent apoptosis in thyroid tumor cells.

3.5 In vivo effects of COPZ1 depletion

We used 8505C cell mouse xenografts as in vivo model of thyroid carcinogenesis and, in a preliminary experiment, we tested the in vivo efficacy of siRNA targeting COPZ1. Nude mice were xenografted with 8505C cells and then locally treated with siCOPZ1, explanted and analyzed by WB three days later. As shown in Figure 5A, a consistent reduction (by 40%) of COPZ1 protein in siCOPZ1-treated tumor in comparison to vehicle- or siNT- treated control was observed. The expression of COPZ1 protein in tumor explants was also analyzed by immunohistochemistry (IHC). The intensity of staining for COPZ1 in siCOPZ1 explant was reduced in comparison to siNT explant: score 1+ vs 3+. This was associated with a reduction of the rate of proliferating cells, determined by Ki67 staining: 15% in siCOPZ1 explant vs 50% in siNT explant (Figure 5B). After proving the efficacy of siCOPZ1 in an in vivo setting, we evaluated the effect of COPZ1 silencing on in vivo tumor growth. Nude mice were xenografted with 8505C cells and, at day 17 after, they
were locally treated twice a week with siCOPZ1, or with siNT, or with the vehicle only. At day 34, tumors were excised and weighted. As shown in Figure 5C, treatment with siCOPZ1 significantly reduced tumor growth compared to siNT or vehicle treatments (controls). Then, tumor explants were analyzed by IHC for the expression of COPZ1 protein. COPZ1 immunostaining was high in two siNT samples and ranged from medium to high in two other samples. Conversely, in COPZ1-depleted explants it was very low (two samples), low with medium areas (two samples) or medium (one sample). In line with this, the mean rate of proliferating cells (Ki67) was 53.7% for siNT and 31% for siCOPZ1 explants. Concomitantly, when we evaluated the neoplastic cellularity, we observed an increase of necrosis (20%) in siCOPZ1 explants in comparison with the controls (11.2%) (Figure 5D). Representative pictures of selected explants are shown in Figure 5E: the siNT-9 sample showed high COPZ1 and Ki67 (50%) expression and low necrotic cells (5%), whereas the siCOPZ1-19 sample displayed an opposite state, being the COPZ1 staining low, Ki67 20% and necrotic cells 25%. Overall, our results demonstrated that the local treatment with siCOPZ1 productively reduced tumor growth in our model of thyroid carcinoma in nude mice.

3.6 COPZ2 is downregulated in thyroid cancers

COPZ1 vulnerability is related to downregulation of the paralogous COPZ2 gene in tumor cells. While normal cells tolerate COPZ1 inhibition because COPZ2 can replaces its activity in the COPI complex, the absence of COPZ2 in tumor cells confers sensitivity to COPZ1 depletion. Interestingly, COPZ2 was found downregulated in several primary tumors compared to their normal counterparts [13]. We have previously provided evidence that dependence of thyroid tumor cell lines on COPZ1 may result from downregulation of COPZ2, since COPZ2 mRNA expression is downregulated in the majority of analyzed thyroid tumor cell lines [9]. In order to assess whether also thyroid tumors belonging to different histotypes may be sensitive to COPZ1 targeting, we interrogated several publicly available datasets for the expression status of COPZ2. The first analysis combined data from different datasets, including 58 normal thyroid, 72 PTC, 17 PDTC and
31 ATC samples [20-24]: COPZ2 was found downregulated in all the tumor histotypes (Figure 6A). Furthermore, COPZ2 gene was significantly downregulated in the 58 PTC samples from the thyroid TCGA dataset [28], compared to the paired normal counterpart (Figure 6B, left). Therefore, the frequent downregulation of COPZ2 in thyroid tumors allow predicting their sensitivity to COPZ1 inhibition.

The first intron of the COPZ2 gene encodes the precursor of miR-152 [37], which exerts a tumor suppressor role and is frequently downregulated in several tumors. miR-152 and its hosting gene COPZ2 are transcribed from the same promoter and are concurrently silenced in tumor cells [13]. To assess mirR-152 expression in thyroid tumors, data were retrieved from TCGA and correlated with COPZ2 expression. The analysis shows a notable positive correlation between COPZ2 and hsa-miR-152 RNAseq expression levels (Figure 6B, right). This result indicates that miR-152, concurrently to COPZ2, is downregulated in thyroid cancer, and it may exert an oncosuppressor role also in this tumor type.

4. Discussion

By performing a siRNA-based functional screening, we have recently discovered several thyroid tumor cell vulnerabilities, consisting in a set of genes, defined non-oncogenes, whose silencing inhibited the growth of a panel of thyroid tumor cells, but not that of normal immortalized thyrocytes. The COPZ1 gene was found among the top significant genes, exerting a role in sustaining the phenotype of thyroid tumor cells. In this study, we dissected the mechanisms leading to cell lethality upon COPZ1 depletion, and explored its employability as therapeutic target for thyroid carcinoma.

First, we confirmed that COPZ1 represents vulnerability for several thyroid tumor cell lines since its silencing drastically impairs their viability, and demonstrated the lack of effect also in normal primary thyrocytes, and mammary epithelial cells.

Our study, aimed at better characterizing the mechanisms leading to cell death upon COPZ1 depletion, showed that, similarly to other members of the coatamer complex [11;30], COPZ1 was a
critical player of autophagy. Indeed, COPZ1 silencing initiated autophagy, which became abortive due to an increase rather than a turnover of autophagosomes. Concomitantly, COPZ1-depleted cells displayed the activation of markers of ER stress and UPR. Whether the impairment of autophagy is the cause or the consequence of ER stress remains to be investigated. We then demonstrated that the onset of a prolonged ER stress exacerbates cell death, since COPZ1-depleted cells rapidly underwent caspase-dependent apoptosis.

Interestingly, in support to what observed by functional studies, a preliminary gene expression profiling of COPZ1-depleted 8505C cells indicated that the main upregulated canonical pathways consist of the unfolded protein and ER stress response, lipid metabolism and cell death (apoptosis). In parallel, proteomic profiling of COPZ1-depleted 8505C cells by nano-scale liquid chromatographic tandem mass spectrometry, indicated that the main involved pathways deal with vesicle trafficking, autophagy and caspase cascade in apoptosis (our unpublished results), in keeping with what observed by gene expression analysis.

Our results suggest that thyroid tumor cells dependence upon the activity of COPZ1 may be related to its pivotal involvement on the regulation of autophagy, UPR and ER stress, processes generally known as cytoprotective and adaptive mechanisms for cancer cell survival. Thus, COPZ1 inhibition may represent a strategy to target these processes, and convert their effect into a lethal mechanism. Interestingly, normal cells, which are probably not strictly dependent on these mechanisms, are preserved from cell death. However, apart from the well known role of COPZ1 in vesicle trafficking, the possible direct involvement of COPZ1 in cancer cell survival is still unclear. How COPZ1 depletion modulates the activation of intracellular multiple signalling pathways, executors of the final lethal effect, remains to be better investigated. This will allow identifying other critical nodes in the oncogenic network whose inhibition will result in cell collapse.

In our study, the most striking effect of COPZ1 silencing was observed in mouse tumor 8505C xenografts: in fact, siCOPZ1-treated tumors showed a consistent reduction of tumor growth.
Moreover, IHC analysis of tumor explants showed that COPZ1 silencing was associated with the decrease of the proliferative marker Ki67 and with an increase of necrosis. Further studies are needed to identify the mediators mainly involved in this process. An attractive hypothesis is that COPZ1 depletion could establish an inflammatory microenvironment through the release of cytokines/chemokines; this is supported by gene expression profile of COPZ1-depleted cells, showing upregulation of genes coding for cytokines involved in mediating communication between immune cells. More information on mediators released after cell death will be provided by the characterization of the cellular secretome content after COPZ1 depletion.

Tumor cell sensitivity to COPZ1 depletion is related to downregulation of its paralogous COPZ2 gene. COPZ1 inhibition is tolerated by normal cells because COPZ2 substitutes for it in the COPI complex, whereas tumor cells, characterized by the absence of COPZ2, display sensitivity. This “synthetic lethality” renders COPZ1 depletion an attractive strategy to kill tumor cells. By interrogating publicly available data sets including a large collection of thyroid tumors we found that COPZ2 was frequently downregulated in PTC, PDTC and ATC samples, thus indicating that the majority of thyroid tumors may be eligible for COPZ1 targeting. As reported in other tumor types [13], we observed a concomitant low expression of miR-152, encoded within the COPZ2 gene, which exerts a tumor suppressor role and is downregulated in different tumors, including thyroid ones [38]. Whether the downregulation of COPZ2 and miR-152 in thyroid tumors is due to promoter hypermethylation, as reported for other tumor types [39], remains to be investigated. Moreover, we cannot exclude that COPZ1 dependency of thyroid tumors may also rely on other mechanisms or on other components of the COPI complex. Indeed, we had previously observed that the FTC-derived WRO82-1 cell line is sensitive to COPZ1 depletion, even though it expresses high level of COPZ2 mRNA [9].

Recent reports indicate that, as for COPZ1, targeting other COPI complex members may represent a therapeutic approach for cancer. A siRNA-based functional screening proposed COPA gene, encoding the α subunit of COPI complex, as a potential therapeutic target for mesothelioma [40].
Claerhout et al have shown that targeting the COPI complex members is a potent approach to block productive autophagy and stimulate cell death in cancer cells [30]. Recently, ARCN1, the δ-subunit of the COPI complex, has been proposed as a new therapeutic target [41]. COPB2 was found as synthetic lethal partner of mutated KRAS [42]. The co-occurrence of mutations in KRAS and LKB1, present in a small fraction of lung adenocarcinomas, drives addiction to the COPI complex members [43]. Of note, COPE gene, encoding the ε subunit of COPI complex, emerged from our screening in the list of genes specifically inhibiting thyroid tumor cell growth [9]. All this evidence suggests that targeting the coatamer complex, and consequently causing abortive autophagy and ER stress, may represent an optimal strategy to kill cancer cells, especially those resistant to conventional therapy.

In conclusion, we demonstrated that COPZ1 represents vulnerability for thyroid tumor cells, thus providing the first evidence of non-oncogene addiction for this tumor type. This adds a new element to the list of potential therapeutic targets, which includes, in addition to the driver oncogenic proteins and downstream pathways, several genes and miRNAs differentially expressed in thyroid tumors, some of which identified by our group [17;18;44-47], and whose role in promoting or suppressing the transformation process has been functionally demonstrated. On the whole, our results pave the way for designing novel therapeutic approaches based on COPZ1 inhibition useful for thyroid tumor patients who do not respond to standard therapies and rapidly progress to malignant forms. Notably, being effective on tumor but not on normal cells, these novel therapeutic approaches may have limited side effects.

5. Conflict of interest

None

6. Acknowledgements
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7. Author contribution

AMC conceived the study, designed and performed the experiments, interpreted data and wrote the manuscript; CE, TK, GA, MG, DMT, CL and PS performed the experiments; LD performed the experiments and provided the editing of the manuscript; BB, TC, MG and NA supervisioned the study and revised the manuscript; GA conceived and supervisioned the study, wrote the manuscript.

8. References

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9. Figure legends

Figure 1. Effect of COPZ1 silencing on cell viability. (A) Colony forming assay of thyroid cell lines, 5 days after siRNA transfection; (B-C) Representative pictures at 10X magnification of Nthy-ori 3-1, TPC-1, 8505C, HTC/C3 cells, primary normal thyrocytes and human mammary epithelial cells (HME-1) at the indicated time points after siRNA transfection. In (A) and (C), cells were assessed by WB for protein levels of COPZ1 72h after siRNA transfection; β-actin was used for normalization of gel loading. siNT: Non-Targeting siRNAs; siCOPZ1: siRNAs targeting COPZ1; siUBB: siRNA targeting Ubiquitin B.

Figure 2. Effect of COPZ1 silencing on autophagy. (A) Percentage of TPC1- and 8505C-GL3 cells displaying LC3 puncta at different time points after siCOPZ1 and siNT oligos transfection; the asterisks indicate differences significant by the unpaired Student’s t-test (*P<0.05, **P<0.01). (B) Representative pictures of TPC1- 8505C-GL3 cells, 48h and 72h after siRNA transfection,
respectively; cells were analyzed by fluorescence microscopy for the presence of LC3 puncta (green; arrows); nuclear DNA was stained with DAPI. (C-D) TPC-1 and 8505C cells were analyzed by WB for the expression of LC3-I/II, SQSTM1/p62 and COPZ1 proteins, at the indicated time points after siRNA transfection; β-actin was used for normalization of gel loading. In 8505C cells (D), p62 and COPZ1 expression in the same cell extracts was assessed in two separate gels.

**Figure 3.** Effect of COPZ1 silencing on ER stress and UPR. (A) WB analysis of calnexin and BiP protein expression in TPC-1 and 8505C cells, at the indicated time points after siRNA transfection; β-actin was used for normalization of gel loading. In TPC-1 cells, calnexin, COPZ1 and BiP expression in the same cell extracts was assessed in two separate gels; in 8505C cells, calnexin and BiP expression was assessed in the same gel of Figure 2D. (B) Real-time PCR analysis of COPZ1 and CHOP gene expression in TPC-1 and 8505C cells at the indicated time points after siRNA transfection; results are presented as log10-transformed relative quantity (RQ) of COPZ1/CHOP mRNA normalized for HPRT1 housekeeping gene expression. Data represent the mean +/- sd of two independent experiments.

**Figure 4.** Effect of COPZ1 silencing on cell death. (A) TPC-1 and 8505C cells were transfected with siCOPZ1 or siNT oligos in the presence of 2 µM of the fluorogenic substrate for activated caspase-3/7; live-cell fluorescence images (10X magnification) were taken 72h later with microscope. (B) WB analysis of cleaved PARP (arrowhead), cleaved caspase 3 and COPZ1 protein expression, at the indicated time points after siRNA transfection; β-actin was used for normalization of gel loading. Cleaved PARP expression in the same cell extracts was assessed in two separate gels.

**Figure 5.** Effect of COPZ1 silencing on 8505C in vivo tumor growth. (A) WB analysis of COPZ1 expression in tumour xenografts, excised three days after siRNA treatment; actin was used as
loading control for cell extracts. The graph represents the densitometric analysis of the bands; data are reported as ratio of COPZ1 to actin and normalized for the siNT sample value. (B) IHC analysis of COPZ1 and Ki67 expression in siNT and siCOPZ1 tumor explants. (C) *In vivo* tumorigenicity assay: the line within each box represents the median tumor weight (g) value of 10 controls (five vehicle and five siNT) and six siCOPZ1 treated explants at the end of the experiment (34 days); upper and lower edges of each box represent the 75th and 25th percentile, respectively; upper and lower bars indicate the highest and lowest values less than one interquartile range from the extremes of the box. *P* = 0.026, controls vs. siCOPZ1-treated mice, Kruskal-Wallis Test. (D) Mean percentage of Ki67 positive and necrotic cells in tumor explants (four siNT and five siCOPZ1 samples); the asterisk indicates differences significant by the unpaired Student’s t-test (*P* < 0.05). (E) IHC analysis of COPZ1 and Ki67 expression and H&E staining of two representative tumor explants (siNT-9, siCOPZ1-19); black arrow indicates areas of necrosis.

**Figure 6.** Expression of *COPZ2* and mir-152 in thyroid carcinomas. (A) Scatter plot distribution of *COPZ2* gene expression levels (log$_2$ scale) in 58 normal thyroid-tissues, 72 PTC, 17 PDTC and 31 ATC primary tumor samples; in the graph the median is recorded. Nonparametric pairwise multiple comparisons between all analyzed groups were performed by Dunn’s test and Benjamini-Hochberg correction, following a significant Kruskal-Wallis test (*p* = 5.733 e$^{-11}$). (B) (left) Scatter plot distribution of RNAseq normalized expression values (natural scale) from the thyroid TCGA dataset of *COPZ2* in 58 matched normal (N) and PTC samples. Illumina HiSeq Level 3 data were downloaded from TCGA Data Portal website and RSEM gene normalized expression values were derived for 58 PTC patients, in tumor and adjacent normal samples, respectively; in the graph the median is recorded. Differential gene expression levels between tumor and normal groups were evaluated by Wilcoxon test (p-value= 2.442x10$^{-10}$); (right) dispersion plot of *COPZ2* (x-axis) and hsa-miR-152 (y-axis) RNAseq normalized expression levels (log$_2$ scale) in the 58 PTC samples. Normalized reads per million miRNA mapped (RPM) data were obtained for the same 58 tumor
samples from Illumina HiSeq Level 3 isoform quantification files (TCGA Data Portal website), by summing up the reads aligned to each 3p or 5p miRBase v16 mature strands. Pearson’s correlation method was applied to evaluate the correlation between RNAseq normalized gene and miRNA read counts in 58 tumor samples ($r = 0.67$, p-value $= 8.41 \times 10^{-9}$).
Figure 1

A

B

C
Figure 2

(A) Bar graphs showing the percentage of cells with LC3 puncta in TPC1-GL3 and 8505-GL3 cells treated with siNT or siCOPZ1 after 24, 48, and 72 hours.

(B) Fluorescence images of TPC1-GL3 and 8505-GL3 cells treated with siNT or siCOPZ1 showing the effect on LC3 puncta.

(C) Western blots for TPC-1 and LC3-I, LC3-II, and β-actin in TPC-1 and 8505C cells treated with siNT or siCOPZ1 after 48, 72, and 96 hours.

(D) Western blots for p62 and COPZ1 in TPC-1 and 8505C cells treated with siNT or siCOPZ1 showing the effect on p62 and COPZ1 expression over time.
Figure 3
Figure 4

A

Bright field  Caspase 3/7  Bright field  Caspase 3/7

D

TPC-1  8505C
dPARP

β-actin

dCASP-3

COPZ1

β-actin
Figure 5

A

COPZ1

\[ \text{β-actin} \]

vehicle siNT siCOPZ1

\[ \text{COPZ1/β-actin} \]

vehicle siNT siCOPZ1

B

\[ \text{siNT} \]

COPZ1 20X K67 20X

C

\[ \text{Tumor weight (g)} \]

controls siCOPZ1

D

\[ \text{Explants} \]

siNT siCOPZ1 siNT siCOPZ1

E

\[ \text{siNT-5} \]

COPZ1 20X K67 20X

\[ \text{siCOPZ1-19} \]

COPZ1 20X K67 20X H&E 20X
Highlights

- COPZ1 represents an example of non-oncogene addiction in thyroid tumor cells
- COPZ1 depletion affects thyroid tumor cell viability
- Cell death is associated with abortive autophagy, ER stress, UPR and apoptosis
- *In vivo* COPZ1 depletion reduces the growth of thyroid tumor mouse xenografts
- COPZ1 represents an attractive therapeutic target for thyroid cancer