Pathogenic mechanisms of deregulated microRNA expression in thyroid carcinomas of follicular origin

Juliane Braun, Stefan Hüttelmaier*

Abstract

Thyroid cancer is one of the most common malignancies of the endocrine system with increasing incidence. The vast majority of thyroid carcinomas derive from thyroid hormone producing follicular cells. Carcinomas of follicular origin are classified as follicular (FTCs), papillary (PTCs), partially differentiated (PDTCs) or anaplastic (ATCs) thyroid carcinomas. While FTCs and PTCs can be managed effectively, ATCs are considered one of the most lethal human cancers. Despite the identification of various genetic alterations, pathogenic mechanisms promoting the progression of thyroid carcinomas are still largely elusive. Over the recent years, aberrant microRNA expression was revealed in all as yet analyzed human cancers, including thyroid carcinomas. In view of the rapidly evolving perception that deregulated microRNA expression serves a pivotal role in tumor progression, microRNAs provide powerful tools for the diagnosis of thyroid carcinomas as well as the identification of potential therapeutic targets. Here, we summarize recent findings on microRNA signatures in thyroid carcinomas of follicular origin and discuss how deregulated microRNA expression could promote cancer progression.

Introduction

Thyroid carcinomas represent the most common cancer of the endocrine system [1]. More than 95% of these carcinomas originate from follicular thyroid cells, whereas only 3% are of C-cell origin, referred to as medullary thyroid carcinomas (MTCs) [2]. The most frequent follicular tumors are benign hyperplastic adenomas (FTAs), whereas papillary thyroid carcinomas (PTCs) are the most frequent thyroid carcinoma (approximately 90%) [3]. FTCs are composed of well-differentiated epithelial cells and can be distinguished by visible changes in nuclear morphology and appearance [4]. Follicular thyroid carcinomas (FTCs) with a prevalence of less than 10% are morphological similar to FTAs but capable of vascular invasion [2]. Although some of these well-differentiated carcinomas behave aggressively, the vast majority of PTCs and FTCs can be managed effectively. In contrast, the rare (2-7%) undifferentiated, anaplastic thyroid carcinomas (ATCs) behave very aggressively, rapidly invade adjacent tissues and are thus considered one of the most lethal human cancers [2]. At present there is no effective treatment of ATCs and death usually occurs within six months after diagnosis [5]. ATCs are characterized by partially or completely undifferentiated cells with a high mitosis rate, necrotic areas, spindle-like cell morphologies as well as giant and occasionally squamous cells [5,6]. Poorly differentiated thyroid carcinomas (PDTCs) present an ‘intermediate’ entity. They appear partially undifferentiated compared to FTCs or PTCs and typically behave more aggressively [5]. Several of these tumors arise de novo, whereas others seem to originate from PTCs or FTCs [7].

Thyroid tumors are supposed to be mainly monoclonal malignancies arising based on somatic mutations of progenitor cells. Putative risk factors for genomic instabilities are radiation exposure, active oxygen-species (H₂O₂ is necessary for thyroid hormone synthesis) and estrogen. Historical events exhibit radiation exposure as the major risk for PTCs, since atomic bomb survivors and Chernobyl victims frequently developed these tumors [8]. One of the major risks for FTCs is dietary iodine deficiency resulting in thyroid proliferation.

* Correspondence: stefan.huettelmaier@medizin.uni-halle.de
Institute of Molecular Medicine, Section for Molecular Cell Biology, Martin Luther University of Halle-Wittenberg, ZAMED Heinrich-Damerow-Str.1, 06120 Halle, Germany
Full list of author information is available at the end of the article
(endemic goiter) as a compensatory mechanism [9]. Genetic predisposition associated thyroid cancers are known for familial polyposis coli (mutations in APC), Cowden disease (mutations in PTEN) and Werner syndrome (mutations in WRN) [2]. Moreover, two sequence variants of loci 9q22.33 and 14q13.3 were found to be associated with a higher risk of FTCs and PTCs [10].

Genetic alterations in oncogenes that are involved in the activation of cell signalling pathways have been observed in the vast majority of malignant thyroid carcinomas. Mutation of the BRAF gene is most prominent in PTCs with an appearance up to 50% [2,11,12]. One major gain-of-function mutation of BRAF, a substitution of valine to glutamate at position 600 (V600E), results in constitutive activation of the MAPK pathway [13,14]. RAS gene (KRAS, HRAS, NRAS) mutations instead were found in all thyroid cancers: FTAs (frequency: 20-40%), FTCs (frequency: 40-50%), PTDTs (frequency: 20-55%) and ATCs (frequency: 20-60%) [2,11,12]. Activation of the G-protein RAS stimulates MAPK and other signalling pathways like PI3K/AKT. Less frequent are RET rearrangements (20-40% in PTCs, ~10% in PTDTs) due to inter-chromosomal translocations. In ATCs and PTDTs, CTNNB1 (0-25% in PTDTs, 66% in ATCs) and TP53 mutations (20-38% in PTDTs; 67-88% in ATCs) were observed [2,11,12,15,16]. NTRK1 rearrangements were exclusively identified in PTCs (~10%). The proto-oncogene NTRK1 (also known as TRK) encodes a transmembrane tyrosine-kinase receptor for nerve growth factor and activates ERK, PI3K and phospholipase γ (PLCγ) signalling pathways. Notably, all mutations in PTCs are nearly 100% complete. Rearrangements leading to the chimeric protein PAX8-PPARγ were only identified in FTCs (35%) and FTAs (2-10%) [2,3,11]. Additionally, AKT signalling appears most accelerated in ATCs and PTDs due to PTEN mutations and/or PI3KCA amplification [17]. Together the frequency preference of genetic alterations observed in distinct thyroid cancers as well as the observation that well-differentiated thyroid carcinomas precede or co-exist with PTDTs or ATCs supports the view that most undifferentiated thyroid carcinomas evolve by sequential progression (reviewed in: [2]).

In addition to genetic alterations, recent studies indicate that thyroid carcinomas like the majority of as yet analyzed tumors are characterized by aberrant expression of microRNAs (miRNAs). These small, non-coding RNAs of 20-24 nucleotides in length are evolutionarily conserved and control gene expression at the post-transcriptional level [18]. At present, 1366 mature human microRNAs are listed in miR-Base (mirbase.org, release 16). The vast majority of identified and characterized microRNAs target the 3'-untranslated region (3' UTR) of mRNAs and modulate target transcript degradation, translation or both [19,20]. Two classes of microRNAs relevant to cancer are distinguished: ‘onco-miRs’ with tumor promoting effects versus ‘tumor-suppressive’ microRNAs that antagonize cancer progression. In the past years, several studies identified miRNA signatures in thyroid carcinomas aiming to reveal how they modulate thyroid cancer progression and to evaluate their potential for thyroid cancer diagnosis.

**MicroRNA signatures in follicular thyroid carcinomas (FTCs)**

Malignant FTCs and benign FTAs share significant similarities at the morphological and molecular level and thus microRNAs could serve as valuable markers to distinguish these tumors. The first study addressing this aspect on the basis of a limited set of human microRNAs (235 distinct human microRNAs) revealed four microRNAs (miR-346, -328, -192, -197) moderately upregulated by 1.34-1.82 fold in FTCs when compared to FTAs [21]. The authors claimed this moderate upregulation sufficed to distinguish FTAs from FTCs in 74% of analyzed patient samples (23 FTCs, 20 FTAs). In vitro, the upregulation of miR-197 and -346 was associated with elevated proliferation of HEK293T and two FTC-derived cell lines. This pro-proliferative effect was suggested to correlate with miR-controlled expression of ACVR1 (activin A receptor type 1), TSPAN3 (tetraspanin 3), CFLAR (Caspase 8 and FADD-like apoptosis regulator) and EFEMP2 (fibulin 4), although no direct regulation of these targets by miR-197 or -346 was demonstrated [21]. The abundance of these miRNAs was decreased in the majority of analyzed FTCs and was proposed to distinguish FTAs from FTCs [21]. Notably, previous studies suggested activin signalling to act on growth inhibition of FTC-derived cells in vitro, but miRNA-mediated regulation of Activin signalling remains to be shown [22]. TSPAN3 was inversely correlated with the metastatic potential in melanoma and the extracellular matrix (ECM) component EFEMP2 was reported to be over expressed in colon carcinomas but decreased in prostate cancer [23-25].

In comprehensive studies aiming at the identification of microRNA signatures distinguishing thyroid carcinomas from non-transformed thyroid tissue (NT), Nikiforova and colleagues [26] could not confirm the results of Weber et al. [21], except for miR-197. This miRNA showed a higher expression in oncotypic FTCs compared to NTs, whereas expression appeared largely unaffected in conventional FTCs [21,26]. Notably, miR-328 expression, previously suggested being higher in FTCs than in FTAs, was found to be upregulated in FTAs but not in FTCs by Nikiforova et al [26]. The increased expression of miR-221/-222 appears to be a hallmark of...
thyroid cancers of follicular origin, since both miRNAs were found to be expressed at high levels in FTCs, PTCs, PDTCs and ATCs [26-31] (Figure 1). Accordingly, their expression was largely unaffected in FTAs, hyperplastic nodules and medullary thyroid carcinomas (MTCs) [26]. Compared to non-transformed ‘normal’ thyroid (NT) and hyperplastic nodules four additional miRNAs were significantly increased in conventional FTCs (miR-187, -224, -155, -146b) as well as in oncotic FTCs (miR-187, -339, -183, -197). MiR-187 is one of the ten most upregulated microRNAs in PTCs, FTCs and PDTCs but apparently remains unaffected in FTAs. This suggests miR-187 as a useful marker for distinguishing FTCs from FTAs that however is not suitable for discriminating FTCs from other carcinomas of follicular origin. The only few studies addressing miRNA signatures in FTCs focused exclusively on the upregulation of microRNAs. However, by comparing ATPC samples with NTs, FTCs and PTCs, we observed a significant reduction in miR-26a/b and let-7g expression in FTCs. These miRNAs are considered tumor suppressive miRNAs [32-35].

In conclusion, comprehensive miRNA signature analyses in FTCs, in particular with respect to distinct subtypes like oncotic versus conventional FTCs, are still lacking. Thus, it remains largely elusive if FTCs can be distinguished from other thyroid carcinomas or FTAs by altered miRNA expression profiles at this point.

**MicroRNA signatures in papillary thyroid carcinomas (PTCs)**

The upregulation of the miR-221/-222 cluster and miR-181 is the most consistent finding in all studies addressing miRNA signatures in PTCs [26,27,31,36,37]. Moreover, increased expression of miR-146b, -155, -21 and -220 was

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Figure 1 Deregulated miRNA expression in thyroid carcinomas of follicular origin in comparison to normal thyroid tissue. Altered expression of miRNAs observed by indicated studies is represented by color coding. Deregulated expression was classified as up- (red), unaffected (grey) or downregulated (green) in comparison to non-transformed thyroid tissue based on individual studies. Contradictory data are indicated by boxes. The presented data base on different thresholds, were quantified using distinct methods and more importantly distinct miRNA sets. Thus, the presented summary of observed miRNA signatures remains incomplete, preliminary and requires substantial further validation. This is particularly important for FTCs and PDTCs for both of which only a very limited set of microRNAs has been analyzed. FTC: Follicular thyroid carcinoma; PTC: Papillary thyroid carcinoma; PDTC: Poorly differentiated thyroid carcinoma, ATC: Anaplastic thyroid carcinoma.

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[25] Nikolova et al., 2008
[26] Palla et al., 2006
[27] Vonne et al., 2007
[28] Heng et al., 2010
[29] Wei et al., 2009
[30] Schwerin et al., 2009

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reported in at least two individual studies based on comparing signatures between PTCs and NTs (Figure 1). A comparison of PTCs to hyperplastic nodular tissue from formalin fixed paraffin embedded samples further validated the significance of these findings by identifying altered expression of miR-221/-222, -181, -31, and -224 [38]. Upregulation of miR-31 and -224 was also observed by Nikiforova et al. [26]. However, analyses of 40 fine needle aspirate (FNA) specimen and 84 formalin-fixed paraffin-embedded tissues by Chen et al. identified only the upregulation of miR-146b and -222 as potent markers of PTCs [39].

Most interestingly a significant correlation of increased miRNA expression with genetic alterations was observed in PTCs [26]. For instance, miR-221/-222 were more abundant in PTC samples with RAS and BRAF mutations. RAS-mutations moreover correlated with the most severe upregulation of miR-146b. In contrast, RET-mutations were found to be associated with increased expression of miR-155. In vitro analyses using thyroid cancer-derived cells confirmed an ‘oncogene connection’ for upregulated expression of miR-221 and -181b. The abundance of these microRNAs increased upon overexpression of v-raf, v-ras, RET/PTC1, RET/PTC3, E1-Abl, E1a-v-raf, middle T of polyomavirus and v-mos [27]. PTC cell lines harboring RET/PTC1 or BRAF V600E mutations identified additional microRNAs to be severely increased. The most severe upregulation upon BRAF mutation was observed for the miR-200 group [31]. The most significant downregulation was observed for the miR-30 and -200 families. Notably, the comparison of miRNA signatures between ATCs, PTCs and FTCs suggested that downregulation of the miR-30 and -200 families is sufficient to unambiguously distinguish ATCs from NT as well as FTCs and PTCs. As observed for PTCs, the miR-138 was found to be severely reduced in ATC samples as well as in ATC-derived cell lines [30,31,42,43]. Consistent with our studies and the analyses of ATC-derived cells, Nikiforova and colleagues observed a severe upregulation of miR-221/-222 [26,43]. Although they examined the expression of 158 microRNAs and identified 57 down- and 47 upregulated miRNAs (miR-222, -198, -let-7f-1, let-7a-2) by comparing ATCs with non-transformed thyroid tissue [28]. The most significant decrease in expression was determined for miR-30d, -125b-1/2, and -26a which could be confirmed by Schwertheim et al. [36]. In support of these findings, we identified 62 down- and 21 upregulated miRNAs compared to non-transformed thyroid tissue [30]. Strikingly, the decreased miRNAs comprised 12 miRNA-families and 12 clustered miRNA transcription units. The most significant downregulation was observed for tumor suppressive miRNAs of the let-7, miR-26, -30 and -200 families. Notably, the comparison of miRNA signatures between ATCs, PTCs and FTCs suggested that downregulation of the miR-30 and -200 families is sufficient to unambiguously distinguish ATCs from NT as well as FTCs and PTCs. As observed for PTCs, the miR-138 was found to be severely reduced in ATC samples as well as in ATC-derived cell lines [30,31,42,43]. Consistent with our studies and the analyses of ATC-derived cells, Nikiforova and colleagues observed a severe upregulation of miR-221/-222 [26,43]. Although they examined the expression of 158 microRNAs and identified 57 down- and 47 upregulated in the majority of thyroid tumors they only analyzed the ten most upregulated microRNAs in further detail. Finally, the potent onco-miR-21 was found to be highly expressed in ATCs and was observed to promote thyroid tumor growth in mouse [30,44].

In conclusion, ATCs appear to be well-distinguishable from all other thyroid carcinomas by a severe decrease in the expression of various microRNAs. However, future studies have to reveal if altered miRNA profiles suffice to unambiguously distinguish ATCs from PDTCs.

MicroRNA signatures in anaplastic thyroid carcinomas (ATCs)

The most striking difference between ATCs and all other thyroid carcinomas of follicular origin appears to be a severely decreased expression of various miRNAs (Figure 1) [28,30,36]. Visone et al. observed decreased expression of 20 and elevated abundance of only four miRNAs (miR-222, -198, -let-7f-1, let-7a-2) by comparing ATCs with non-transformed thyroid tissue [28]. The most significant decrease in expression was determined for miR-30d, -125b-1/2, and -26a which could be confirmed by Schwertheim et al. [36]. In support of these findings, we identified 62 down- and 21 upregulated miRNAs compared to non-transformed thyroid tissue [30]. Strikingly, the decreased miRNAs comprised 12 miRNA-families and 12 clustered miRNA transcription units. The most significant downregulation was observed for tumor suppressive miRNAs of the let-7, miR-26, -30 and -200 families. Notably, the comparison of miRNA signatures between ATCs, PTCs and FTCs suggested that downregulation of the miR-30 and -200 families is sufficient to unambiguously distinguish ATCs from NT as well as FTCs and PTCs. As observed for PTCs, the miR-138 was found to be severely reduced in ATC samples as well as in ATC-derived cell lines [30,31,42,43]. Consistent with our studies and the analyses of ATC-derived cells, Nikiforova and colleagues observed a severe upregulation of miR-221/-222 [26,43]. Although they examined the expression of 158 microRNAs and identified 57 down- and 47 upregulated in the majority of thyroid tumors they only analyzed the ten most upregulated microRNAs in further detail. Finally, the potent onco-miR-21 was found to be highly expressed in ATCs and was observed to promote thyroid tumor growth in mouse [30,44].

In conclusion, ATCs appear to be well-distinguishable from all other thyroid carcinomas by a severe decrease in the expression of various microRNAs. However, future studies have to reveal if altered miRNA profiles suffice to unambiguously distinguish ATCs from PDTCs.

MicroRNA signatures in poorly differentiated thyroid carcinomas (PDTCs)

PDTCs are poorly defined and show morphological characteristics of both, differentiated and anaplastic thyroid carcinomas. Aiming to distinguish PDTCs from PTCs and ATCs on the molecular level, Schwertheim et al. investigated the expression of two different sets of microRNAs (‘set 1’: miRNA-146b, -181b, -21, -221 and -222, all upregulated in PTCs; ‘set 2’: miRNA-30d, -125b, -26a, -30a-5p and let7d, all downregulated in ATC). Abundance of these microRNAs was analyzed in comparison to four NTs in 15 PDTCs, nine PTCs and
nine ATCs [36]. ‘Set 1’ microRNA expression was slightly increased in PDTCs but did not significantly differ from NTs. Since PTCs and ATCs instead showed a more robust upregulation of these microRNAs, the authors suggested ‘set 1’ microRNAs as a promising diagnostic tool to distinguish PDTCs from PTCs or ATCs. MicroRNAs of ‘set 2’ instead appear useful to discriminate PDTCs from PTCs, since they were expressed at low levels in PDTCs and ATCs but significantly upregulated in PTCs. Nikiforova et al. investigated the microRNA expression of four PDTCs but did not aim to distinguish PDTCs from PTCs or ATCs [26]. In contrast to Schwertheim et al., Nikiforova and colleagues found miR-181b, -221, -222 and -146b to be upregulated in all four analyzed PDTCs when compared to NTs [26].

Although preliminary evidence indicates that miRNA profiles could be useful molecular markers to identify PDTCs, additional and more comprehensive studies are required.

**Onco-miRs in Thyroid cancer**

Onco-miRs are typically classified by an upregulation in cancer and the targeting of transcripts encoding effectors antagonizing tumor progression (Figure 2, red). In the following we discuss validated and suggested phenotypic consequences of aberrant onco-miR expression observed in thyroid cancer.

**miR-221/-222**

The overexpression of miR-221/-222 is a hallmark of thyroid malignancies [26-28,30,31]. Moreover, upregulation of both miRNAs was also shown in hepatocellular carcinomas (HCC), glioblastoma and gastric cancer [45-47]. Antagonizing the function of miRs-221/222 in PTC-derived cells interfered with cell proliferation and
growth [27]. In subsequent studies it was demonstrated that both miRNAs negatively regulate expression of the cyclin-dependent kinase (CDK) inhibitor p27Kip1[29]. This inhibitory role most likely involves miR-221/-222 directed control of FOXO3, a key transcriptional activator of p27Kip1[48,49]. Notably, FOXO3a miRNA was significantly downregulated in PTCs and FOXO3a represses onco-miR-21 in lung cancer cells [49,50]. An inhibitory role of both miRNAs was also observed for p57Kip2 expression [47,51]. Hence, elevated expression of miR-221/-222 presumably indicates aberrant proliferation and cell growth due to their role in antagonizing cell cycle arrest by targeting crucial cell cycle gatekeepers like CDK-inhibitors. Supporting this view, reduced expression of p27Kip1 was observed in many primary thyroid tumors as well as thyroid carcinoma-derived cell lines [52].

The expression of the c-kit tyrosine kinase receptor is suppressed or reduced in most tested FTCs and PTCs [53]. Indicating a role of miR-221/-222 in this regulation, it was demonstrated that both miRNAs target c-kit in melanoma-derived cells [54]. Interestingly, 70% of cutaneous melanomas are characterized by BRAF mutations which was also shown to be a precaution of miR-221/-222 upregulation in PTC [26]. Various studies showed that miR-221/-222 downregulate expression of the tumor-suppressive phosphatase PTEN leading to an upregulation of PI3K/AKT signaling [55,56]. This pathway blocks apoptosis and promotes invasion by modulating focal adhesion kinase (FAK) phosphorylation and matrix metalloprotease (MMP) expression levels [57]. PI3KCA amplifications as well as PTEN mutations were reported for many FTCs, PTCs and ATCs suggesting that AKT signaling is sustained by genomic alterations and modified by post-transcriptional control in thyroid carcinomas [17,58,59]. Another pro-apoptotic factor and inhibitor of MMP expression regulated by miR-221/-222 is TIMP3. Reduced expression of PTEN and TIMP3 facilitated by this miRNA-cluster induces TRAIL (tumor necrosis factor (TNF)-related apoptosis inducing ligand) resistance and upregulation of MMPs [55]. In vivo evidence for the growth promoting role of elevated miR-221/-222 expression was provided by Xenograft studies in nude mice. The constitutive overexpression of this miRNA cluster resulted in an increased gain of Xenograft weight and volume upon subcutaneous injection of tumor-derived cell lines [47]. Together these findings indicate the overexpression of miR-221/-222 as a key event in thyroid tumor progression. Upregulation of both miRNAs presumably promotes uncontrolled growth and could as well modulate the invasive potential of tumor cell.

miR-21
Upregulated expression of miR-21 is observed in multiple types of cancers, such as breast, liver, brain, prostate and myometrial cancers but also in PTCs and ATCs [44,60,61]. Most recently it was shown that oncogenic Ras induces miR-21 expression via activating Raf/MAPK and AKT signaling in rat thyroid cells and in a mouse model of lung tumorigenesis [44]. MiR-21 predominantly acts in an anti-apoptotic manner by interfering with the expression of PTEN and PDCD4 [62-64]. Thus, this miRNA promotes AKT signaling similar to the miR-221/-222 cluster and thereby stimulates its own expression. Elevated AKT signaling and the direct regulation of tropomyosin 1 (TPM1) and maspin (PI5) affects cell morphology, motility and enhances cell invasion [65,66]. By the direct targeting of RECK and the indirect effect on TIMP3 expression, upregulation of miR-21 promotes cell migration and invasiveness but also blocks apoptosis [67]. Hence, the overexpression of miR-21 in thyroid cancers is likely to promote uncontrolled growth and potentially upregulated invasiveness of tumor cells.

miR-155
MiR-155 overexpression was observed in FTCs, PTCs and ATCs [26,31]. In leukemia, miR-155 together with miR-21 stimulates PI3K/AKT signaling by blocking expression of SHIP1, another AKT inhibitory phosphatase [68,69]. In addition miR-155 interferes with the expression of RhoA and serves a role in modulating tumor cell invasion as well as epithelial-to-mesenchymal-transition (EMT) [70]. TGFβ induces miR-155 expression and thereby promotes TGFB-induced EMT and tight junction dissolution involving the depletion of RhoA. Like miR-221/-222, miR-155 also directly targets FOXO3a [71]. Hence, upregulation of miR-155 in thyroid cancers is presumed to promote cell survival, invasiveness and resistance to chemotherapeutics, as for instance demonstrated in breast cancer cells [71].

Tumor-suppressive miRNAs in thyroid cancer
In contrast to onco-miRs, tumor-suppressive micro-RNAs are typically downregulated during tumor progression and target transcripts encoding oncogenic factors. The regulatory role of key tumor-suppressive miRNAs in thyroid cancer is discussed in the following (Figure 2, green).

miR-200
The miR-200 family (comprising miR-141, -200a, -200b, -200c and -429) was identified as a potent suppressor of EMT [72-75]. Reduced expression of this miRNA-family was previously reported for stomach, breast and
ovarian cancers [76]. Surprisingly, moderately upregulated expression of some miR-200 family members was observed in PTCs and FTCs, whereas a severe downregulation of this microRNA family was found in all ATCs analyzed [30]. This observation supports the view that the miR-200 family serves a key role in preserving an epithelial phenotype or morphology, respectively. Elevated levels of the miR-200 family interfere with the expression of EMT-promoting factors like ZEB1, ZEB2, SNAI2, SMAD2, TGFβR1 and TGFβ2. This antagonizes transcriptional repression of E-cadherin and the miR-200 genomic clusters (miR-141/-200c and miR-200a/-200b/-429) [30,72-74]. In agreement, severely reduced E-cadherin levels have been described as a common characteristic of primary ATCs as well as ATC-derived cells [30,77]. Moreover, elevated expression of TGFβ2 was observed in ATC and PTC compared to FTC and PTC samples [78]. Beyond the antagonistic role in TGFβ-induced EMT, the miR-200 family presumably also modulates actin dynamics by interfering with the expression of WAVE3, an actin cytoskeleton remodeling protein [79]. Taken together, this provides strong evidence that the loss of miR-200 expression presents a hallmark in the progression of thyroid carcinomas culminating in TGFβ-dependent EMT and elevated invasiveness, as observed for ATCs [30]. The loss of miR-200 expression is probably potentiated by TGFβ-mediated upregulation of miR-155 expression and enhancement of miR-21-maturation, for both of which a pro-metastatic function was proposed [70,80]. More recently, regulation of LIN28B by miR-200 members was identified in prostate cancer-derived cells [81]. LIN28 is a stem cell factor and a powerful inducer of pluripotency [82]. It also represses maturation of the tumor-suppressive let-7 family by binding to the loop region of let-7 precursors [83-85]. Additionally, the miR-200 family apparently promotes apoptosis, as demonstrated in colorectal cancer-derived HCT116 cells. In these cells the repression of FAP1 (Fas-associated phosphatase 1) by the miR-200 family was correlated with elevated susceptibility to TNF-receptor CD95-dependent apoptosis [86]. In summary, these findings indicate reduced expression of the miR-200 family as a key trigger for dedifferentiation and potentiated aggressiveness observed in ATCs.

let-7
Downregulation of the let-7 family is observed in all thyroid carcinomas of follicular origin [27,28,30]. Let-7 was initially identified as a factor promoting differentiation in C. elegans and was since then validated as a key regulator of gene expression in various organisms [87,88]. The let-7 family is ubiquitously expressed in adult mouse tissue and its reduced expression is considered to be a hallmark in cancer progression [89]. In accord with their tumor-suppressive role, members of the let-7 family target cell cycle regulators and oncogenes like RAS, HMGA1/2, MYC, IGF2BP1 and LIN28 [34,89-93]. Thus it appears likely that the capability to antagonize activating RAS mutations and thus uncontrolled proliferation is severely compromised by reduced levels of the let-7 family. Like activating RAS mutations, the overexpression of MYC is associated with enhanced cell growth and reduced serum dependency in various malignancies [94]. Accordingly, the frequently observed increase of MYC levels in ATCs is likely to correlate with downregulation of the let-7 family [78,95,96]. Notably, MYC is a key regulator of miRNA transcription and negatively controls expression of various tumor-suppressive miRNA clusters including once again the let-7 family but in addition the miR-30, -26, -34 and -29 families [97,98]. Another layer of post-transcriptional regulation is provided by the control of MYC mRNA degradation via the RNA binding protein IGF2BP1 (Insulin-like growth factor 2 mRNA binding protein 1) [99]. IGF2BP1 shows an oncofetal expression pattern and becomes de novo synthesized in various cancers [100]. This is in agreement with a reduction of let-7 expression, since members of this miRNA family interfere with IGF2BP1 expression. Thus, MYC-directed repression of the let-7 family is likely to promote the expression of IGF2BP1 that in turn sustains MYC expression [97]. HMGA1 and HMGA2, two other key factors in tumor progression that are controlled by the let-7 family, support the RAS/MEK-facilitated induction of EMT by enhancing SNAI1 expression [101-103]. Moreover, HMGA proteins promote cell growth and HMGA1 depletion induces programmed cells death in ATC-derived cells [104]. In conclusion, loss of the let-7 family is likely to be an early event in the progression of thyroid carcinomas. The severe reduction of this tumor-suppressive miRNA family promotes uncontrolled tumor growth and presumably also affects the invasive potential of tumor cells at later stages.

miR-30
Significantly reduced expression of the miR-30 family was identified in ATCs. However, moderately reduced expression is also observed in FTCs, PTCs and PDTCs, with the exception of miR-30b in FTCs [28,30]. The first hint of a potential tumor-suppressive role of this miRNA family was revealed by the identification of UBC9 as a direct target [105]. Downregulation of this E2-conjugating enzyme for sumoylation interferes with cell growth and cancer progression. In bladder cancer, reduced expression of the miR-30 family was associated with upregulation of the tumor marker cytokeratin 7, although a functional role of this potential regulation
remains elusive [106]. We identified the miR-30 family as an antagonist of TGFβ-induced EMT and in vitro invasiveness of ATC-derived cells [30]. This role apparently involves the direct targeting of SMAD2 and ZEB2. In agreement, the forced expression of miR-30 family members interfered with in vitro invasiveness of ATC-derived cells and correlated with a downregulation of the mesenchymal marker vimentin. However, in contrast to the miR-200 family, expression of the miR-30 family is apparently controlled in a ZEB-independent manner [30]. Intriguingly, the miR-30 family also targets the stem cell factor LIN28, as previously demonstrated for other key tumor-suppressive miRNA families like miR-200 and let-7 [107]. Hence, the identification of deregulated expression of the miR-30 family in ATCs identified yet another tumor-suppressive miRNA cluster modulating EMT and invasiveness of tumor cells.

**Conclusions**

The comprehensive view of distinct miRNA signatures in thyroid carcinomas of follicular origin provides novel insights in the molecular pathologies of these malignancies. However, knowledge of target miRNAs controlled by deregulated miRNAs in thyroid cancers is still sparse at present. To reveal how altered expression of micro-RNAs promotes or antagonizes thyroid tumor progression it is thus required to identify novel miRNA targets in future studies.

In thyroid carcinomas the most striking observation is that ‘less’ aggressive FTCs and PTCs (compared to ATCs) are apparently characterized by an upregulation of ‘oncogenic’ miRNAs (e.g. miR-221/-222 or miR-146b). These miRNAs mainly act in a pro-proliferative and anti-apoptotic manner. In accord with the clonal progression hypothesis, ATCs reveal a severe reduction of tumor-suppressive miRNAs (Figure 1). This decrease presumably promotes dedifferentiation, which morphologically manifests as an epithelial-to-mesenchymal-transition (EMT) driven by reduced expression of the miR-200 and -30 families (see Fig. 1 in [30]). Notably, reduction of these miRNA families is likely to provide a valuable diagnostic tool for distinguishing ATCs from FTC or PTCs. Unresolved aspects of immediate diagnostic and potentially therapeutic importance address the question if miRNA signatures are suitable to distinguish FTAs from FTCs. The presented studies seem to be insufficient to define a subset of microRNAs that unambiguously discriminates these thyroid cancers. Thus, it is required to re-evaluate miRNA signatures in thyroid cancers of follicular origin in a comprehensive manner based on pathologically unambiguously classified primary samples. These studies need to be based on the same evaluation method, for instance microarray-based microarrays, to generate data sets allowing an unbiased direct comparison of miRNA signatures.

Despite their potential as diagnostic tools, miRNAs could also provide promising therapeutic targets based on onco-miR inhibition or restoring levels of tumor suppressive miRNAs. The stage for such approaches was set in rodents already in 2005 by demonstrating that the role of miR-16 as well as -122 could be antagonized by the intravenous injection of anti-miRs [108]. Vice versa, the adenoviral delivery of the tumor-suppressive miR-26 was shown to suppress mouse liver tumorigenesis [33]. However, in clinical practice the specificity of miRNA or anti-miR delivery remains an essential limitation due to off-site effects in non-neoplastic organs.

A third strategy could be envisioned on the level of epigenetic silencing of tumor-suppressive miRNAs. In breast and prostate cancer cell lines methylation of the miR-200 cluster was observed and inhibition of methyltransferases released this transcription block [109]. Moreover, epigenetic silencing might explain the observation that complete miRNA transcription units are downregulated in ATCs (Table I in [30]). Hence, spatially restricted and cluster specific release of epigenetic silencing could provide a valuable tool in the treatment of cancer but as yet such approaches are not available.

In conclusion the here reviewed findings reveal the potency and current limitations of miRNAs in diagnosis, prognosis and potentially therapeutic strategies for the treatment of thyroid cancer. However, substantial additional work will be required to establish miRNAs in clinical practice and reveal the molecular networks via which altered miRNA expression promotes thyroid cancer progression.

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**Authors’ contribution**

Both authors contributed equally

**Competing interests**

The authors declare that they have no competing interests.

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