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This is the final peer-reviewed author's accepted manuscript (postprint) of the following publication:

Published Version:

Xu, B., Viswanathan, K., Ahadi, M.S., Ahmadi, S., Alzumaili, B., Bani, M., et al. (2024). Association of the Genomic Profile of Medullary Thyroid Carcinoma with Tumor Characteristics and Clinical Outcomes in an International Multicenter Study. THYROID, 34(2), 167-176 [10.1089/thy.2023.0279].

This version is available at: https://hdl.handle.net/11585/964031 since: 2024-02-29

Published: DOI: http://doi.org/10.1089/thy.2023.0279

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Molecular Profile of Medullary Thyroid Carcinoma (MTC) and Its Impact on Tumor Characteristics and Clinical Outcome: An International Multicentric Study of 290 Patients

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Disclosure Statement: No competing financial interests exist for all contributory authors. Research reported in this publication was supported in part by the Cancer Center Support Grant of the National Institutes of Health/National Cancer Institute under award number P30CA008748 and Italian Government - Ministry of Health Grant No. RF-2011-02350857.

Keywords: Medullary thyroid carcinoma, grade, RET, RAS, prognosis.

Running title: Molecular alterations in medullary thyroid carcinoma.

Abstract

Background: The prognostic impact of germline *RET*, somatic *RET* and *RAS* mutations and their relationship to clinicopathologic parameters and outcomes is poorly defined in medullary thyroid carcinoma (MTC) and needs to be clarified.

Design: Molecular profile, including germline and somatic *RET* and *RAS* mutations were detected using various molecular platforms in 290 primary MTCs from 6 tertiary centers, including 239 from prior an international MTC grading system (IMTCGS) cohort and 51 new cases.

Results: *RET* germline mutations were detected in 40 patients (16.3%). Somatic *RET* and *RAS* mutations were seen in 136 (46.6%) and 57 (19.8%) respectively. *RET* M918T was the most common somatic *RET* mutation (n=75).

Compared with wild type (WT) MTCs, those with *RET* germline mutation were associated with younger age at presentation. MTCs with *RET* somatic mutations were associated with female sex, larger tumor size, advanced AJCC prognostic group, vascular invasion, and high IMTCGS grade. There was no significant difference between MTCs with somatic *RAS* mutations and WT MTCs. When compared with other *RET* somatic mutations, *RET* M918T was more commonly associated with younger age, AJCC prognostic group 4, vascular invasion, extrathyroidal extension, and positive margin.

The presence of any *RET* mutation, *RET* M918T, and *RAS* mutations did not significantly impact overall and disease specific survival (OS and DSS). RET somatic or germline mutations were significant adverse prognostic factor for distant metastasis free survival (DMFS) on univariate survival analysis but lost its significance on multivariate analysis when adjusted for grade and stage. There were no outcome differences between *RET* somatic and *RET* germline mutations, or between *RET* M918T and other *RET* mutations.

Other recurrent molecular alterations included *TP53* (4.2%), *ARID2* (2.9%), *SETD2* (2.9%), *KMT2A* (2.9%), and *KMT2C* (2.9%), *PIK3CA* (1.0%), *ATM* (2.3%), *VHL* (1.0%) and *TERT* promoter mutation (1.0%). Among them, *TP53* mutations were associated with decreased OS, DSS, and LRRFS on univariate survival analysis. There was no association between *TP53* mutations and grade or AJCC prognostic group.

Conclusions: *RET* somatic mutations correlate with high-grade, aggressive primary tumor characteristics, and decreased DMFS. *RET* M918T is associated with aggressive primary tumors but does not impact outcome. We further identified *TP53* mutation as an adverse molecular signature to predict decreased OS, DSS, and LRRFS in MTCs.

Introduction

Medullary thyroid carcinoma (MTC), a neuroendocrine carcinoma derived from parafollicular c-cells, accounts for approximately 2% of all thyroid malignancy and 8% of thyroid cancer-related mortality ¹. It may occur sporadically or in the setting of a germline mutation of *RET* protooncogene (i.e. multiple neuroendocrine neoplasia type 2, MEN2) ²⁻⁵. A large proportion of sporadic MTCs harbor *RET* (particularly *RET* M918T) or *RAS* somatic mutations ⁶⁻¹⁰, making them potential candidates for kinase inhibitor targeted therapies ¹¹.

In 2022, we established an international MTC consortium to develop and validate a powerful prognosticator of MTC, namely the International Medullary Thyroid Carcinoma Grading System (IMTCGS) based on mitotic index, Ki67 proliferation index, and/or tumor necrosis ². Although data for *RET* germline mutations were collected, the underlying molecular profile and its correlation with clinicopathologic features and outcome of MTC remained to be elucidated.

In this study, we investigated the molecular signatures of a large multicentric cohort of 290 patients with primarily resected MTC using RT-PCR based platforms and six different next generation sequencing (NGS) platforms. The aims are two-fold: first to clarify the prognostic significance of somatic *RET* or *RAS* mutations and their correlations with various clinicopathologic parameters, including IMTCGS grade; and second to identify other molecular alterations in MTCs and their prognostic roles.

Materials and methods

Study cohort

This retrospective cohort study included 290 resected primary MTCs from six tertiary centers (University of Bologna Medical Center [UB], Bologna, Italy: n=64; Memorial Sloan Kettering Cancer Center [MSKCC], New York, NY, USA: n=54; Institut Gustave Roussy, Villejuif, France: n=45; Brigham and Women's Hospital [BWH], Boston, MA, USA: n=44; Royal North Shore Hospital, Sydney, Australia: n=42, and Emory University Hospital Midtown [EU], Atlanta, GA, USA: n=41). Among them, 239 patients were included in the prior IMTCGS cohort ², whereas the remaining 51 were new patients (UB: n=6; MSKCC: n=4; and EU n=41). One center (BWH) included only patients with somatic MTC, whereas the cases from other centers also contained patients with germline *RET* mutations.

Molecular platforms

Somatic molecular alterations, including *RET* and *RAS* mutations, were detected using either RT-PCR-based platforms targeting only *RET* and/or *RAS* genes (n=99) or six NGS platforms (n=191). Details of the NGS platforms utilized were either commercially available or were described previously ¹²⁻¹⁷ (briefly summarized in **Supplementary Table 1**). Number of cases tested using each NGS platform was as follows: a custom-designed multi-gene panel (n=104), MSK-IMPACT (n=54), ion AmpliseqTM cancer hotspot v. 2 (CHP2, Thermofisher Scientific, n=17), OncoPanel (n=14), Paradigm Cancer Diagnostic (PCDx) platform (Paradigm diagnostics, n=1), and CARIS NGS platform (CARIS life sciences, n=1).

Clinicopathologic review, outcome, and statistical analysis

Clinicopathologic review were performed at each individual participating site. Outcome data, including overall survival (OS), disease specific survival (DSS), distant metastasis free survival (DMFS), and locoregional recurrence free survival (LRRFS), were collected. Statistical analyses were performed using the SPSS software 24.0 (IBM Corporation, Armonk, NY, U.S.). Univariate survival analysis was performed using log rank test for categorical variables and Cox proportional hazards model for continuous variables. Multivariate survival analysis using Cox proportional hazards model adjusted for grade and stage was subsequently conducted. Additionally, comparisons of the clinicopathologic features among each molecular subgroup (*RET* germline mutations, *RET* somatic mutations, *RAS* somatic mutations, and *RET/RAS* wild type) were performed using Chi square test or Fisher's exact test for categorical variables and two-tailed Student's t test for continuous variables.

Results

Clinicopathologic characteristics of the study cohort

This retrospective study cohort had 290 patients with resected primary thyroid MTC. The median age of presentation was 57 years (range: 7 – 88 years, **Table 1**). The male to female ratio was 1:1.1. AJCC prognostic group 4, vascular invasion, extrathyroidal extension, positive margin, and IMTCGS high grade were identified in 39.3%, 42.1%, 28.8%, 18.6%, and 24.8% of the cases respectively. Twenty-one patients (8.2%) had distant metastasis at presentation. External beam radiation therapy and kinase inhibitors were given to 8.2% and 8.7% of patients respectively. The kinase inhibitors used

included selpercatinib (n=7), vandetanib (n=5), cabozantinib (n=3), pralsetinib (n=1), sorafenib (n=1), LOXO-292 (n=1), RAD001 (n=1), and RAD100 (n=1).

RET germline mutations

After excluded the BWH subgroup which contained only somatic MTCs, the frequency of RET germline mutations was 16.3% (40/246, **Figure 1 and Supplementary Table 2**). The most frequently detected germline mutations were C609Y (n=14), C634R (n=8), and G533C (n=3). Among these familial MTCs, *RET* or *RAS* somatic mutations were detected in three cases: one case had germline *RET* Y791F and somatic R*ET* M918T, one had germline *RET* V804Y and somatic *RET* M918T, and the third had *RET* C634R germline mutation and *HRAS* G13R somatic mutation.

RET somatic mutations

RET somatic mutations were detected in 136 cases of MTC (46.6% overall, 53.6% in sporadic MTCs, **Table 1**, **Figure 1**, and **Supplementary Table 1**). Among them, *RET* M918T was the most common somatic mutations, being detected in 75 cases. Other common *RET* somatic mutations affected codons 634 (n=16), 630 (n=11), 620 (n=7), and 883 (n=6).

RAS somatic mutations

RAS somatic mutations were examined in 288 cases, and were identified in 57 MTCs (19.8% in the entire cohort, 22.4% in sporadic MTCs, 44.9% in *RET*-WT sporadic MTCs), including *HRAS* mutations in 37 cases (Q61R n=21, G13R n=10, Q61K n=3, Q61L n=2, and G13V n=1), *KRAS* mutations in 20 cases (G12R n=8, G12V n=3, Q61R n=3, Q61L n=1, Q61K n=1, D54N n=1, A18D n=1, P34L n=1, and C186Mfs*16 n=1), and *NRAS* Q61K mutation in 1 case. One MTC harbored two *RAS* somatic mutations, being *HRAS* G13R and *KRAS* C186Mfs*16. *RAS* and *RET* somatic mutations were mutually exclusive.

In our cohort, 61 MTCs (21.0%) were devoid of *RET* (somatic or germline) or *RAS* mutations and were grouped as *RET/RAS* wild type (WT).

Correlation of RET/RAS mutations with clinicopathologic parameters

The clinicopathologic characteristics according to RET and RAS mutations status is shown in Table 1.

Compared with WT MTCs, MTC occurring in the familial setting was associated with younger age at presentation (median age: 42 Years in *RET* germline mutation group, 58 years in WT). Other clinicopathologic characteristics did not differ between the two groups.

MTCs with *RET* somatic mutations were associated with aggressive tumor characteristics at presentation compared with WT MTC, including larger tumor size (median size: 2.1 cm in *RET* somatic mutations, 1.9 cm in WT), AJCC prognostic group 4 (56.6% in *RET* somatic mutations, 24.6% in WT), vascular invasion (55.9% in *RET* somatic mutations, 32.8% in WT), and high IMTCGS grade (35.3% in *RET* somatic mutations, 18.0% in WT). Additionally, patients with *RET* somatic mutations were more commonly male (male to female ratio: 1:0.7 in *RET* somatic mutation group, 1:1.4 in WT) and more commonly treated with kinase inhibitor therapy (15.2% in *RET* somatic mutation group, 3.3% in WT group).

Similarly, compared with MTCs with *RET* germline mutations, MTC with *RET* somatic mutations were more aggressive characterized by larger tumor size, AJCC prognostic group 4, vascular invasion, tumor necrosis, high mitotic index, high Ki67 proliferation rate, and distant metastasis at presentation.

The clinicopathologic features did not differ significantly between MTCs with somatic RAS mutations and WT MTCs.

RET M918T somatic mutation was considered a high-risk mutation in this study. When compared with other *RET* somatic mutations, *RET* M918T was more commonly associated with younger age (median age: 52 years in *RET* M918T, 63 years in other *RET* somatic mutations), AJCC prognostic group 4 (69.3% in *RET* M918T, 40.0% in other *RET* somatic mutations), vascular invasion (65.3% in *RET* M918T, 43.3% in other *RET* somatic mutations), extrathyroidal extension (46.7% in *RET* M918T, 20.0% in other *RET* somatic mutations), and positive surgical margin (30.7% in *RET* M918T, 15.0% in other *RET* somatic mutations, **Supplementary Table 3**). Other clinicopathologic parameters did not differ between MTCs with *RET* M918T and those with other *RET* somatic mutations.

The impacts of RET and RAS mutations on clinical outcomes

The prognostic impact of *RET* and *RAS* mutations were determined using univariate survival analysis by log rank test and the results are shown in **Table 2** and **Figure 1**. In brief, *RET* or *RAS* mutation status did not impact OS and DSS.

Compared with WT MTCs, those with *RET* somatic or *RET* germline mutations were associated with shortened DMFS (p<0.05). The 10-year DMFS in MTCs with *RET* somatic mutations, MTCs with *RET* germline mutations, and WT MTCs was 49%, 47%, and 75% respectively. There was also a non-significant trend for *RAS*-mutated MTCs to be associated with improved DMFS (10-year DMFS in *RAS*-mutated MTCs: 88%, p=0.081).

RET or *RAS* mutation profile failed to reach significant levels on multivariate survival analysis using Cox proportional hazards model adjusted for IMTCGS grade and AJCC prognostic group (**Supplementary Table 4**, p>0.05). IMTCGS grade and AJCC prognostic group remained independent adverse prognostic factors (IMTCGS grade: hazard ratio 2.031, 95% confidence interval 1.230 – 3.353, p=0.006; AJCC stage group 4: hazard ratio 1.478, 95% confidence interval 1.221 – 1.789, p<0.001).

Other molecular alterations in MTC

Other mutations were examined in 191 cases using six different NGS platforms ranging from 26 genes to whole exome sequencing. The detailed mutation profile is provided in **Supplementary Table 5**. Recurrent somatic mutations or fusions detected included *TP53* (8/191, 4.2%), *PlK3CA* (2/191, 1.0%), *VHL* (2/191, 1.0%). *TERT* promoter mutation (2/191, 1.0%), *ATM* (2/87, 2.3%), *ARID2* (2/70, 2.9%), *SETD2* (2/70, 2.9%), *KMT2A* (2/70, 2.9%), and *KMT2C* (2/70, 2.9%). The presence of TP53 mutations was associated with decreased OS (log rank test, p=0.027, **Supplementary Figure 1**), DSS (p=0.008), and LRRFS (p=0.018), and showed a non-significant trend towards shortened DMFS (p=0.074). There was no correlation between TP53 mutations and IMTCGS grade or AJCC prognostic group (p=1.000). Other molecular alterations did not significantly correlate with clinicopathologic characteristics or survival.

Discussion

In this multicentric retrospective study, we presented a complete dataset including molecular signatures, detailed clinicopathologic features, and outcomes of 290 patients, the largest MTC cohort to date. Our key findings were 1) *RET* somatic mutations were associated with aggressive tumor characteristics, IMTCGS high grade, and decreased DMFS. However, the prognostic values of *RET* somatic mutations were not independent of grade and stage; 2) MTCs with *RET* M918T somatic mutations showed tumor aggressiveness in the primary resection (such as AJCC prognostic group 4, vascular invasion, extrathyroidal extension, and positive margin), but did not impact survival; and 3) *TP53* mutation was identified as a novel adverse molecular signature for survival affecting 4.2% of MTC.

Germline *RET* mutations were detected in 16.3% of the study cohort, lower than the ~25% rate of familial cases reported in the literature ¹⁸. The relatively low frequency of germline *RET* mutations may be explained by the selection bias of our study cohort towards adult patients: only 4 patients (1.4%) of our cohort were age 18 years or under. As the current American Thyroid Association (ATA) guideline recommended prophylactic thyroidectomy at young age for patients with hereditary MTCs, such patients were likely not included in our cohort giving the selection bias towards adult patients ¹⁸. Consistent with what were previously reported ^{5,19-21}, we found that *RET* germline mutations affected codons 533, 609, 611, 618, 634, 791, 804, and 918, with codon 618 being the most prevalent, and that hereditary MTCs occurred at a younger age compared with sporadic MTCs.

RET somatic mutations have been identified in 45% to 70% of sporadic MTCS and have been shown to be associated with larger tumor size ²², high T stage ²³, nodal metastasis ²²⁻²⁴, distant metastasis ²²⁻²⁴, advanced stage ²²⁻²⁴, and decreased survival (overall and disease-free) ²²⁻²⁴ in MTCs. Similarly, we have shown here that MTCs with *RET* somatic mutations had larger tumor size, advance AJCC prognostic group, and decreased DMFS. However, the prognostic values of RET somatic mutations were lost on multivariate survival analysis when adjusted for IMTCGS grade and AJCC prognostic group, while grade and stage remained independent prognostic factors. Together, these data imply that *RET* somatic mutation was associated with more aggressive tumor characteristics and adverse outcome, but its role in MTC may not be independent of grade and stage. In a study of 100 sporadic MTCs, Elisei et al. identified *RET* somatic mutation and advance stage as the only two factors independently correlated with persistent disease ²³. There were several differences between their study and the current study. First, Elisei et al. included only sporadic MTCs, whereas our cohort contained both hereditary and sporadic cases. Second, *RAS* somatic mutation was not studied in Elisei et al. only performed correlation analysis between persistent disease and mutation/stage using multivariable logistic regression model. Multivariant survival analysis was not done. The fact that driver mutations are not independent from stage and other clinico-pathologic factors such as grade is not unique to MTC. Indeed, the same scenario applies to follicular cell derived thyroid carcinomas where *BRAF* V600E mutation was shown to be not independent from clinico-pathologic features such as stage in predicting mortality ²⁵.

Additionally, we have shown for the first time that *RET* somatic mutation correlate with IMTCGS high grade. Najdawi et al. previously reported that there was no correlation between *RET* somatic mutation and IMTGS grade in 44 sporadic MTCs ²⁶, These 44 patients were also included in the current study. We were able to establish an association between IMTCGS high grade and *RET* somatic mutations in the current study, possibly for two reasons: first, we expanded the cohort size to 290 cases; and second, we excluded cases with *RAS* somatic mutations from the control (WT) group.

It is worthwhile to mention that not all IMTCGS high grade MTCs harbored *RET* somatic or germline mutations. Among the 72 cases of high grade MTCs, 55 (76.4%) had *RET* somatic or germline mutations, 9 (12.5%) had *RAS* mutations, whereas the remaining 11 did not harbor *RET* or *RAS* mutations. Therefore, not all patients with IMTCGS high grade MTCs are candidate for kinase inhibitor therapies.

RET M918T somatic mutation, the most prevalent *RET* somatic mutation in MTC, has been implied in several studies to be an adverse molecular signature. Schilling et al. ²⁷ showed that *RET* M918T was associated with decreased OS and increased risk of distant metastasis in 34 patients with sporadic MTCs. However, the authors only examined *RET* M918T mutations in their study, and their control group theoretically included both MTCS with other *RET* somatic mutations and MTCs without *RET* mutations. Therefore, it was impossible to directly compare *RET* M918T somatic mutations with other *RET* somatic mutations in this study. Romei et al. ²⁸ showed that *RET* M918T mutations was associated with larger tumor size and was relatively infrequent in MTC < 1 cm in size. Moura et al. found that MTCs with *RET* somatic mutations involving exons 15 and 16 (including 87% of M918T and 13% of A883F) had higher prevalence of nodal metastasis, stage IV disease, and persistent disease compared with MTCs with other *RET* somatic mutations in 52 sporadic MTCs ²⁹. In contrast, Najdawi et al. ²⁶ showed that *RET* somatic mutations affecting exons 15 and 16 did not impact DSS and progression free survival and did not correlate with IMTCGS grade. In the current study including 75 cases with *RET* M918T somatic mutations, we found that *RET* M918T somatic mutations were associated with younger age, advanced AJCC prognostic group, and other aggressive tumor characteristics (e.g. vascular invasion, extrathyroidal extension, and positive margin) compared with MTCs with other *RET* somatic mutations. However, *RET* M918T somatic mutations lacked any prognostic values to predict OS, DSS, DMFS, and LRRFS in MTCs.

RAS mutations are the predominant driver mutations in *RET*-WT sporadic MTC, being detected in 11% (range: 9% to 20%) of MTC overall, 13% (range: 0% to 43%) of sporadic MTCs, and 61% (range: 0% to 81%) of *RET*-WT sporadic MTCs ^{7,9,10,30-33}. Similarly, we detected *RAS* mutations in 19.8% of all MTCs, 22.4% of sporadic MTCs, and 44.9% of *RET*-WT sporadic MTCs. Although we showed a non-significant trend of *RAS*-mutated MTCs with improved DMFS, overall RAS mutations did not impact tumor characteristics or outcomes in MTC as shown previously ²⁴.

It appears from the above data that additional molecular events other than *RET* or *RAS* mutations are needed for the aggressive behavior seen in high grade MTC. In that regard, we herein reported *TP53* mutation as a novel prognostic molecular alteration in MTCs. *TP53* mutations was uncommon in MTCs, being reported in 9% (8/88 cases) in the literature ³³ and 4.2% in our cohort. It may co-exist with *RET* or *RAS* driver mutations as shown in our cohort and in a previous publication ³³. The current study was the first to show that *TP53* mutation was associated with decreased OS, DSS, and LRRFS in MTC.

This international multicentric MTC consortium allowed us to provide detailed clinical, pathologic, and molecular characteristics of MTC. We have previously established and validated a prognostically relevant grading scheme ². We herein presented evidence that *RET* somatic mutations and *TP53* also carried prognostic values in MTC, although (possibly) not independent of grade and AJCC prognostic group. The next step is to establish a nomogram incorporating these prognostic factors, in order to better risk stratify patients with MTCs.

_	All cases (N=290)	RET germline mutations (n=40)	RET somatic mutations (n=133)	RAS somatic mutations (n=56)	RET/RAS wild type (WT, n=61)	P values (<i>RET</i> somatic vs. WT)	P values (<i>RET</i> germline vs. WT)	P values (<i>RAS</i> vs. WT)	P values (<i>RET</i> somatic vs. RET germline)
Male: female ratio	138:152 (1:1.1)	18:22 (1:1.2)	77:56 (1:0.7)	18:38 (1:2.1)	35:36 (1:1.4)	0.031	NS	NS	NS
Age (years)	57 (7-88)	42 (7-66)	59 (22-88)	57 (7-79)	58 (25-84)	NS	<0.001	NS	<0.001
Tumor size (cm)	2.0 (0.1- 11.0)	1.2 (0.1-6.0)	2.1 (0.4-8.0)	2.0 (0.3- 11.0)	1.9 (0.3-6.0)	0.036	NS	NS	0.006
AJCC prognostic group						<0.001	NS	NS	0.006
1	92 (31.7%)	13 (32.5%)	29 (21.3%)	24 (42.9%)	26 (42.6%)				
2	43 (14.8%)	4 (10.0%)	13 (9.6%)	12 (21.4%)	14 (23.0%)				
3	41 (14.1%)	11 (27.5%)	17 (12.5%)	7 (12.5%)	6 (9.8%)				
4	114 (39.3%)	12 (30.0%)	77 (56.6%)	13 (23.2%)	15 (24.6%)				
Vascular invasion	122 (42.1%)	12 (30.0%)	76 (55.9%)	17 (30.4%)	20 (32.8%)	0.005	NS	NS	0.007
Extrathyroidal extension	83 (28.8%)	9 (22.5%)	47 (34.6%)	14 (25.0%)	15 (25.4%)	NS	NS	NS	NS
Positive margin	54 (18.6%)	5 (12.5%)	32 (23.5%)	11 (19.6%)	8 (13.1%)	NS	NS	NS	NS
First post-op CEA	5 (0-38335)	2 (0-600)	7 (0-26300)	5 (1-539)	3 (0-38335)	NS	NS	NS	NS
First post-op calcitonin	13 (1- 970000)	48 (1-7979)	24 (1- 970000)	5 (1-68791)	4 (1-16000)	NS	NS	NS	NS
Calcitonin doubling time									
Never doubled	105/163 (64.4%)	23/32 (71.9%)	40/75 (53.3%)	19/27 (70.4%)	23/31 (74.2%)	NS	NS	NS	NS
Months if doubled	17 (3-139)	13 (3-139)	15 (3-84)	27 (8-35)	15 (3-99)	NS	NS	NS	NS
IMTCGS grade						0.027	NS	NS	NS
Low grade	218 (75.2%)	33 (82.5%)	88 (64.7%)	47 (83.9%)	50 (82.0%)				
High grade	72 (24.8%)	7 (17.5%)	48 (35.3%)	9 (16.1%)	11 (18.0%)				
Necrosis	43 (14.8%)	1 (2.5%)	30 (22.1%)	4 (7.1%)	8 (13.1%)	NS	NS	NS	0.002
Mitotic index (/2mm ²)	1 (0-29)	0 (0-7)	1 (0-18)	1 (0-9)	1 (0-7)	0.028	NS	NS	0.049
KI67 (%)	2 (0-58)	1 (0-15)	2 (0-30)	2 (0-58)	2 (0-30)	NS	NS	NS	0.016
DM at presentation	25 (8.7%)	1 (2.5%)	19 (14.4%)	1 (1.8%)	4 (6.7%)	NS	NS	NS	0.047
Radiation therapy	21 (8.2%)	4 (10.3%)	14 (12.2%)	1 (2.1%)	2 (3.7%)	NS	NS	NS	NS
ТКІ	25 (8.7%)	2 (5.0%)	20 (15.2%)	1 (1.8%)	2 (3.3%)	0.015	NS	NS	NS

Table 1. Clinicopathologic characteristics according to RET and RAS mutation status.

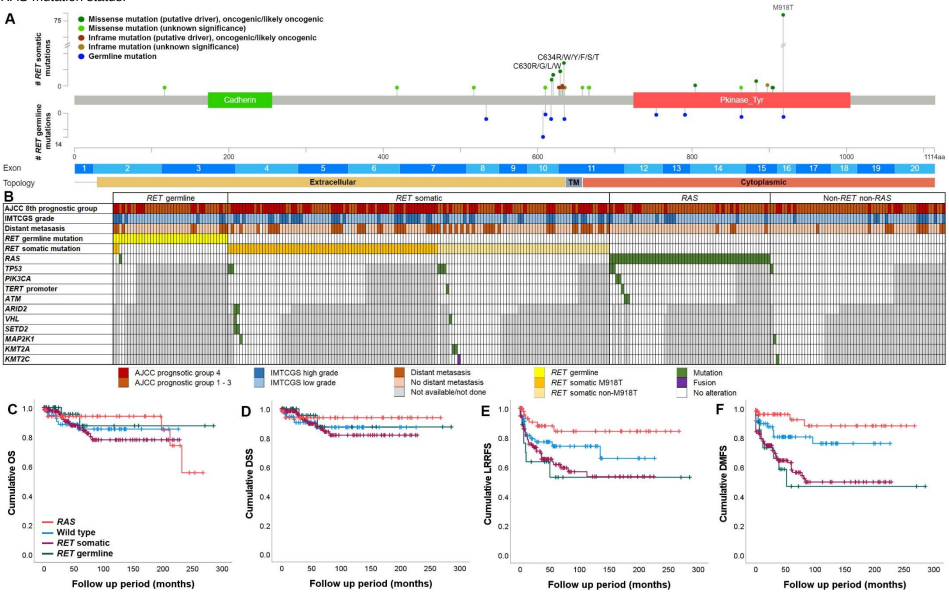
* The three cases with both *RET* germline mutations and somatic *RET* or *RAS* mutations are classified as *RET* germline mutations in this table. Values are expressed as n (column %) for categorical variables, or median (range) for continuous variables. NS: not significant. DM: distant metastasis,

 Table 2. Impact of RET and RAS mutation on prognosis in medullary thyroid carcinoma.
 Values in the table are p values.
 Bold values: significant p values.

 OS: overall survival, DSS: disease specific survival, DMFS: distant metastasis free survival, LRRFS: locoregional recurrence free survival, WT: wild type.

	OS	DSS	DMFS	LRRFS
RET somatic vs. RET germline	0.387	0.556	0.745	0.630
RET somatic vs. RET/RAS WT	0.748	0.854	0.010	0.118
RET germline vs. RET/RAS WT	0.422	0.557	0.045	0.156
RAS mutated vs. RET/RAS WT	0.406	0.407	0.081	0.193
RET M918T somatic vs. other RET somatic	0.202	0.328	0.375	0.531

Figure 1. *RET* germline and somatic mutations in medullary thyroid carcinoma (MTC) and their impact on clinical outcome. (A) lollipop plot. Cadherin: Cadherin domain. Pkinase_Tyr: protein tyrosine kinase. TM: Transmembrane. (B) Oncoprint showing the clinicopathologic features and mutation profile of MTCs. (C-F) Kaplan Meier curves for overall survival (OS, C), disease specific survival (DSS, D), locoregional recurrence free survival (LRRFS, E), and distant metastasis free survival (DMFS, F). *RET* mutations, somatic or germline, are associated with decreased LRRFS and DMFS. OS and DSS do not differ according to *RET* or *RAS* mutation status.



Supplementary table	1. Next generation sequencing molecular platforms and genes tested with each platform.	
Platforms	Gene tested	Ī

Platforms	Gene tested
A custom-designed multi-gene panel	26 oncogenes or tumor suppressor genes as described previously ¹⁴ , including BRAF, c-KIT, CTNNB1, DICER1, DPYD, EGFR, EIF1AX, GNA11, GNAQ, GNAS, H3F3A, HRAS, IDH1, IDH2, KRAS, MED12, MET, NRAS, PDGFRa, PIK3CA, PTEN, RET, RNF43, SMAD4, TERT promoter, TP53, TSHR, and VHL.
MSK-IMPACT	505 cancer-related genes as described previously ^{13,15} .
Ion AmpliSeq Cancer Hotspot Panel v2 (CHP2)	Commercially available platform of 50 oncogenes and tumor suppressor genes and four fusions, including ABL1, AKT1, ALK, APC, ATM, BRAF, CDH1, CDKN2A, CSF1R, CTNNB1, EGFR, ERBB2, ERBB4, EZH2, FBXW7, FGFR1, FGFR2, FGFR3, FLT3, GNA11, GNAQ, GNAS, HNF1A, HRAS, IDH1, IDH2, JAK2, JAK3, KDR, KIT, KRAS, MET, MLH1, MPL, NOTCH1, NPM1, NRAS, NTRK1, PDGFRA, PIK3CA, PTEN, PTPN11, RB1, RET, ROS1, SMAD4, SMARCB1, SM0, SRC, STK11, TP53, and VHL.
OncoPanel	298 to ~500 genes as described previously ^{12,17} .
Paradigm Cancer Genetics Platform	A commercially available platform of 230 genes as described previously ¹⁶ .
CARIS NGS platform	A commercially available whole exome sequencing platform.

Supplementary Table 2. RET germline and somatic mutations in medullary thyroid carcinoma. ATA: American Thyroid Association.

<i>RET</i> mutations	n	ATA risk (for germline mutation) or OncoKB		
		annotation (for somatic mutation)		
Germline mutations (40	/246, 16.			
M918T	2	ATA highest		
V804L	2	ATA modest		
V804M	1	ATA modest		
Y791F	1	ATA modest		
C634R	8	ATA high		
C634F	2	ATA high		
C634Y	1	ATA high		
C618R	1	ATA modest		
C618Y	1	ATA modest		
G611Y	1	ATA modest		
C611S	1	ATA modest		
C609Y	14	ATA modest		
G533C	3	ATA modest		
Exon 11	1	N/A		
Somatic mutations (13	5/290, 46.	6%)		
M918T	73	Oncogenic		
M918T, G601R	1	Oncogenic, unknown		
M918T, P516L	1	Oncogenic, unknown		
S904A, D898_E901del	1	Likely oncogenic, unknown		
S904P, D898_E901del	1	Likely oncogenic, unknown		
D898_E901del	1	Unknown		
A883F	6	Oncogenic		
G861Q	1	Unknown		
V804M	3	Oncogenic		
K666E	1	Unknown		
K666N	1	Unknown		
C634R	8	Oncogenic		
C634W	3	Likely oncogenic		
C634Y	1	Likely oncogenic		
C634F	1	Likely oncogenic		
C634S	1	Likely oncogenic		
C634T	1	Likely oncogenic		
C634_I638del	1	Unknown		
E632_C634del	1	Likely oncogenic		
E632_L633del, H658Y	1	Likely oncogenic, unknown		
D631_C634del	1	Likely oncogenic		
D631_L633delinsE	1	Likely oncogenic		
C630R	5	Oncogenic		
C630G	3	Likely oncogenic		

C630L	1	Likely oncogenic
C630_D631del	1	Likely oncogenic
C630W	1	Likely oncogenic
L629_L633del	1	Likely oncogenic
C620R	3	Oncogenic
C620S	3	Likely oncogenic
C620Y	1	Likely oncogenic
C618A	1	Likely oncogenic
C618G	1	Likely oncogenic
C618R	1	Oncogenic
C611S	1	Likely oncogenic
R417P	1	Unknown
R112C	1	Unknown

Supplementary table 3. Clinicopathologic features of medullary thyroid carcinoma with RET M918T somatic mutations and those with other RET somatic mutations. NS: not significant

somatic mutations. No. not significan	RET M918T somatic mutation (n=75)	<i>RET</i> non-M918T somatic mutation (n=60)	P values
Male: female ratio	44:31 (1.4:1)	34:26 (1.3:1)	NS
Age (years)	52 (22-84)	63 (27-88)	<0.001
Tumor size (cm)	2.1 (0.4-8.0)	2.0 (0.5-9.0)	NS
AJCC prognostic group			<0.001
1	10 (13.3%)	19 (31.7%)	
2	6 (8.0%)	7 (11.7%)	
3	7 (9.3%)	10 (16.7%)	
4	52 (69.3%)	24 (40.0%)	
Vascular invasion	49 (65.3%)	26 (43.3%)	0.015
Extrathyroidal extension	35 (46.7%)	12 (20.0%)	0.002
Positive margin	23 (30.7%)	9 (15.0%)	0.042
First post-op CEA	15 (0-26300)	4 (1-1152)	NS
First post-op calcitonin	83 (1-970000)	6 (1-4481)	NS
Calcitonin doubling time			
Never doubled	20/41 (48.8%)	20/33 (60.6%)	NS
Months if doubled	18 (3-84)	10 (4-57)	NS
Grade			NS
Low grade	46 (61.3%)	42 (70.0%)	
High grade	29 (38.7%)	18 (30.0%)	
Necrosis	16 (21.3%)	14 (23.3%)	NS
Mitotic index (/2mm ²)	1 (0-18)	1 (0-15)	NS
KI67 (%)	3 (0-30)	2 (0-25)	NS
Distant metastasis at presentation	8 (10.8%)	11 (18.3%)	NS
Radiation therapy	6 (9%)	9 (17.6%)	NS
Kinase inhibitor therapy	15 (20.3%)	6 (10%)	NS

Supplementary table 4. Multivariate survival analysis for distant metastasis free survival using Cox proportional hazards model. Bold p values: significant p values.

	P values	Hazard ratio (95% confidence interval)
RET or RAS mutations		
RET and RAS wild type		Reference
RET germline mutations	0.154	1.945 (0.779-4.856)
RET somatic mutations	0.217	1.525 (0.780-2.979)
RAS somatic mutation	0.147	0.428 (0.136-1.349)
IMTCGS grade	0.006	2.031 (1.230-3.353)
AJCC stage group 4	<0.001	1.478 (1.221-1.789)

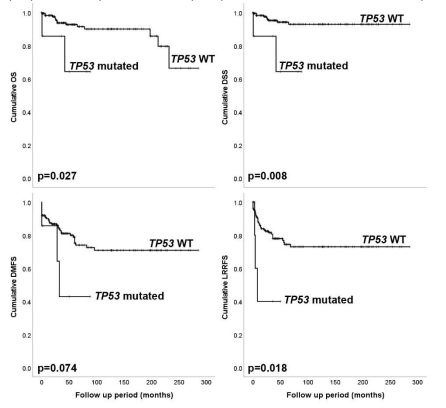
Supplementary Table 5: Other molecular alterations in medullary thyroid carcinoma.

RET or RAS somatic mutations	Other molecular alterations		
<i>RET</i> M918T	<i>TP53</i> G360W		
<i>RET</i> M918T	<i>TP</i> 53 V197E		
<i>RET</i> M918T	MGA W2174*, DIS3 R689Q		
<i>RET</i> M918T	FOXA1 P71Dfs*130, FANCA S858R		
<i>RET</i> M918T	ATM R3008H, SETD2 Y1666C, PIK3CB R321Q		
<i>RET</i> M918T	ARID2 Q1062*, SETD2 R1625C		
<i>RET</i> M918T, <i>RET</i> G601R	<i>MAPK1</i> E322K		
<i>RET</i> S904P <i>, RET</i> D898_E901del	RPS6KA4 S236L		
RET D898_E901del	KMT2C::HILPDA fusion		
<i>RET</i> A883F	VHL W117dup		
RET V804M	<i>TP53</i> R156C		
RET C634R	<i>KMT</i> 2A M1585Ifs*7		
RET C634R	ARID2 F233Lfs*59, MUTYH R95W		
RET C634R	<i>RB1</i> H339_D340fs		
RET C634F	TP63 R379C, EP300 D1399N, CHEK2 R346C		
RET C634_I638del	TGFBR1 E125Dfs*19		
RET L629_L633del	ARID1A S2096*		
RET C620Y	APC R854*		
RET C618R	TP53 V172D		
RET C618G	KMT2A K461Nfs*106		
NRAS Q61K	<i>TERT</i> c124 C>T		
KRAS G12V	<i>PIK3CA</i> R108H		
KRAS D54N	<i>TERT</i> c124 C>T		
HRAS Q61R	TP53 Q317*		
HRAS Q61R	SMAD4 L200Pfs*6		
HRAS Q61R	PIK3CA D1017N		
HRAS Q61K	TP53 R248W, KMT2C R5432W, SMAD2 R321Q		
HRAS G12V	EGFR T790M, VHL R161Q, RAD50 D69N, SPEN M612Cfs*2, MAX R60Q		
-	MAP2K1 K57N		
-	MAP2K1 E102_I103del		
-	KMT2C A1685S		
-	GNAS R201C		

-	<i>ATM</i> R337C
-	<i>TP53</i> T256I
-	ALK R395H, MST1 G673S, MST1 R651*, FAT1 R1205*, LZTR1 R340*
-	IDH2 R140Q, FOXF1 G162S

-: negative for RET and RAS mutations.

Supplementary Figure 1. *TP53* mutation is associated with decreased survival in medullary thyroid carcinoma. Kaplan Meier curves for overall survival (OS), disease specific survival (DSS), distant metastasis free survival (DMFS), and locoregional recurrence free survival (LRRFS). WT: wild type.



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