GRADING OF MEDULLARY THYROID CARCINOMA ON THE BASIS OF TUMOR NECROSIS AND HIGH MITOTIC RATE IS AN INDEPENDENT PREDICTOR OF POOR OUTCOME

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Abstract

Medullary thyroid carcinoma (MTC) is a rare non-follicular cell-derived tumor. A robust grading system may help better stratify patients at risk for recurrence and death from disease. One hundred forty-four MTC between 1988 and 2018 were subjected to a detailed histopathologic evaluation. Clinical and pathologic data were correlated with disease specific survival (DSS), local recurrence free survival (LRFS) and distant metastasis free survival (DMFS). Median age was 53 years (range: 3–88). Median tumor size was 1.8 cm (range: 0.2–11). Lymph node metastases were present in 84 (58%) cases while distant metastases at presentation were found in 9 (6%) patients. Seven (5%) had ≥5 mitoses/10 HPFs. Tumor necrosis was present in 30 cases (20%) while lymphovascular invasion occurred in 41 (28%) of tumors. Extra-thyroidal extension was found in 44 (31%) and positive margins were seen in 19 (14%). There was a strong correlation between increasing tumor size and tumor necrosis (p<0.001). Median follow up was 39 months. In univariate analysis, male gender, higher AJCC stage group, larger tumor size, tumor necrosis, high mitotic index (≥5/10 HPF), nodal status, size of largest nodal metastasis, and elevated post-operative serum calcitonin predicted worse DSS, LRFS and DMFS (p<0.05). Extra-thyroidal extension correlated with DSS and DMFS while positive margins and distant metastasis at presentation imparted worse DSS (p<0.05). In multivariate analysis, tumor necrosis and mitotic activity (5 mitoses/10 HPFs as the cutoff) were the only independent predictors for DSS (p=0.008 and 0.026 respectively). Tumor necrosis was the sole independent prognostic factor for LRFS and
DMFS (p=0.001 and 0.003 respectively). The presence of tumor necrosis and high mitotic rate are powerful independent prognostic factors in MTC and outperform serum calcitonin and stage. We propose a grading system based on tumor necrosis and mitotic activity to better stratify MTC patients for counseling, post resection surveillance and therapy.

Keywords
Medullary thyroid carcinoma; grading; necrosis; mitosis; RET

INTRODUCTION:
Medullary thyroid carcinoma (MTC) is rare accounting for 1–2% of all thyroid malignancies (1). These tumors originate from the neural crest derived parafollicular C cells of the thyroid gland that secrete calcitonin (2, 3). As in many thyroid carcinomas, the clinical course of patients with medullary thyroid carcinoma is variable. Several prognostic factors were shown to confer worse outcome such as age, sex, TNM stage, sporadic versus hereditary disease, distant metastasis, nodal metastatic burden, serum calcitonin, carcinoembryonic antigen (CEA) blood levels, somatic RET mutation status, response to initial therapy risk stratification and extent of thyroidectomy (4–8). However, the predictive value of some of these parameters such as age is controversial (9). Novel and robust prognostic markers of outcome are needed to help guide surgical decision making, systemic therapy and post-resection surveillance strategies.

Despite the initial report by Jacquet of a thyroid tumor with amyloid more than 100 years ago and since the definite histologic description of MTC by Hazard et al in 1959, there has never been a histologic grading system for this entity (10, 11). This is surprising since MTC is a neuroendocrine neoplasm and neuroendocrine tumors in other sites such as lung, pancreas and gastrointestinal tract are usually amenable to grading mainly based on mitotic count, tumor necrosis, and/or Ki-67 proliferation index (12, 13). In order to develop a grading scheme for MTC, we performed a detailed histopathologic review of 144 MTCs and correlated the histologic parameters with outcome and other clinical features. We sought to determine if tumor grade, as measured by mitotic count and/or tumor necrosis would better predict disease-specific outcomes.

MATERIAL and METHODS:
Study population:

The cancer registry at Memorial Sloan Kettering Cancer Center (MSKCC) was searched for all patients with a pathologic diagnosis of MTC, who underwent thyroidectomy between 1986 and 2017. Slides from the primary tumor were available for review by the study pathologists (RAG, BA) on 144 patients. Of note, all the slides were examined by a senior head and neck pathologist with a special interest in thyroid neoplasia (R.A.G). The pathologists were blinded to the patients’ outcome. The study was approved by the Institutional Review Board of MSKCC.
Histopathologic examination:

The diagnosis of MTC was based on the morphologic appearance and/or immunohistochemical profile of the tumor. The MTC was subtyped as classical or other variants on the basis of the most recent WHO classification of endocrine neoplasms (13). The largest dimension of the tumor was based upon a reconciliation of the gross and microscopic findings. The largest size of the largest tumor nodule was used in the analysis. Mitotic rate and tumor necrosis were assessed in the initial thyroidectomy resection and attached lymph nodes. The mitotic rate of the carcinoma was determined by counting 10 contiguous high-power fields (400×) using an Olympus microscope (U-DO model BX-40; Olympus America Inc., Melville, NY). Using that microscope type, 10 high-power fields correspond to 2.4 mm². Mitotic counts were performed in a focused fashion, examining areas that appeared to show greater proliferative activity (so called hot spots). Spontaneous tumor necrosis was classified as absent or present. Tumor necrosis was defined by a “comedo-like” appearance composed of degenerating cytoplasm and punctuate, karyorrhectic nuclear debris (Figures 1 and 2). The presence of fibroblastic stromal reaction, hemorrhage or an identifiable needle track in the necrotic area was attributable to reactive changes induced by prior fine needle aspiration and was therefore not labeled as spontaneous tumor necrosis. Nuclear pleomorphism of the carcinoma cells was characterized as absent/mild, moderate or marked. The presence or absence of amyloid in the tumor was recorded. Infratumoral fibrosis was classified as absent/mild, moderate or prominent. In regard to the status of the tumor capsule, the carcinoma was categorized as completely encapsulated/well circumscribed, partially encapsulated or totally lacking a capsule. Capsular invasion was defined as complete penetration of the capsule by tumor. Only lymphovascular invasion (LVI) of thyroid or extra-thyroid soft tissue vessels was included in the analysis. LVI was diagnosed in accordance with the criteria outlined by the Armed Forces Institute of Pathology fascicle and the WHO classification of endocrine tumors (13, 14). Only when the invasive focus was present in the vascular lumen covered by endothelial cells, or when it was attached to the vessel wall protruding into the lumen of the vessel in a polypoid manner or associated with thrombus formation, it was considered as LVI. For encapsulated tumors, it was defined as invasion of a vessel within or outside the tumor capsule. If the tumor was not encapsulated, any LVI inside or outside the tumor was considered as LVI. The foci of capsular and LVI were counted and subdivided into 2 categories: focal (<4 invasive foci) and extensive (≥4 foci). A tumor was deemed infiltrative if the carcinoma cells invaded in between non-neoplastic thyroid follicles. Extra-thyroidal extension was defined as invasion of peri-thyroid adipose tissue, skeletal muscle or adjacent organs. A surgical margin was considered as positive if tumor was present at the inked resection edge. The number of separate foci of MTC was documented. The number of lymph nodes examined and those with metastasis was recorded along with the size of the largest node positive for tumor, the size of the largest metastatic focus and the presence of extra-nodal extension. The presence of C-cell hyperplasia was documented when feasible.

Clinical and biochemical parameters:

Clinical, biochemical, and follow-up data were obtained by review of the medical records. The following parameters were documented: Age at thyroidectomy, sex, germline RET status, type of thyroid surgery, post-operative serum calcitonin and CEA levels (defined as
the first postoperative level within 30 days of surgery) and distant metastatic status at presentation and on follow up. Locoregional recurrence was defined as structural recurrence in the neck noted on imaging studies while distant recurrence was defined on the basis of biochemical and radiographic evidence of structural disease outside of the neck with or without biopsy. Patients with increased calcitonin without structural correlate were not included in the distant metastasis classification. Locoregional and distant recurrences were recorded independently and the time to recurrence or death was calculated from the date of surgery. The patients’ stage at presentation was assessed using the 8th edition of the American Joint Committee on Cancer (AJCC) staging system (15).

**Statistical analysis:**

All statistical analyses were performed using the SPSS software 24.0 (IBM Corporation, New York, NY, U.S.). Tumor size was correlated with the presence/absence of tumor necrosis using two-tailed Student’s t test. The prognostic significance of each parameter on disease specific survival (DSS), loco-regional recurrence free survival (LRFS) and distant metastasis free survival (DMFS) was calculated using univariate Cox proportional model for log-transformed post-operative CEA and calcitonin levels and log rank tests for all other variables. Factors significant on univariate analysis were subsequently subjected to multivariate analysis using Cox proportional hazards model. Factors that were only applicable to a subset of patients (e.g. the characteristics of LVI and nodal metastasis) were excluded from the multivariate analysis. P values less than 0.05 were considered to be statistically significant.

**RESULTS**

**Clinico-pathologic characteristics of the study cohort**

The clinico-pathologic features of the 144 patients are reported in Table 1. The female to male ratio was approximately 1:1. The median age at diagnosis was 53 years (range: 3–88 years). The median tumor size was 1.8 cm (range: 0.2–11 cm). Four (3%) were treated with lobectomy alone, while the remaining 140 (97%) underwent total thyroidectomy. Mitosis were present in 52 (36%) cases while absent in 92 (64%) of patients. In those 52 mitotically active tumors, the median number of mitoses was 2/10 high power fields (HPFs, range: 1–20) and atypical (abnormal) mitoses were found in 20 (38%) of mitotically active carcinomas. Tumor necrosis was present in 30 (21%) of patients. Patients with tumor necrosis had larger carcinomas (median 3.0 cm) than those without necrosis (median 2.1 cm, p<0.001). The vast majority of cases (87%) displayed none/mild nuclear pleomorphism. A significant (moderate/marked) amount of intratumoral fibrosis was seen in 76% of MTC. A minority of carcinomas (n=26, 18%) were completely encapsulated/well circumscribed. Sixteen (62%) of these 26 totally encapsulated/well circumscribed tumors revealed capsular invasion and 7 (27%) harbored LVI. In eight patients, the dominant tumor was completely encapsulated/well circumscribed and lacked any invasion. These non-invasive tumors comprised 31% of the encapsulated/well circumscribed carcinoma and 6% of the entire study population. Overall, LVI was found in 41 (29%) of individuals. In patients with LVI, extra-thyroidal LVI was seen in 20 (49%) of cases. Extra-thyroidal extension and positive
margins were present in 31% and 14% of cases respectively. In patients with extra-thyroidal extension, invasion was seen in perithyroidal fibroadipose tissue in 39/44 (89%), in skeletal muscle in 4/44 (9%) and in trachea in 1/44 (2%).

At presentation, 84 (58%) of all patients had nodal disease and 9 (6%) harbored distant metastasis. In patients with nodal metastasis, the median number of positive nodes was 7 (range: 1–60). The median size of the largest nodal metastasis was 1.6 cm (range: 0.01–9.5 cm).

Among the 136 patients with available germline RET mutation analysis, 28 (21%) had familial MTC while 106 (79%) had sporadic tumors. Among the 22 cases with mutation details available, the affected codons were 618 (n=2), 620 (n=2), 634 (n=8), 639 (n=1), 790 (n=2), 791 (n=2, including one Y791F), 804 (n=2), and 918 (n=3). The median post-operative serum calcitonin was 43 pg/ml (range: 0–970000 pg/ml). In regard to post-operative serum CEA, the median was 5.6 ng/ml (range: 0.5–1889 ng/ml).

**Survival analysis**

With a median follow up of 39 months (range 0.3–297 months), the 5-year OS and DSS were 87% and 94% respectively. The 5-year LRFS and DMFS were 77% and 87% respectively. Table 2 shows the p values of univariate survival analysis. Male sex, large tumor size, mitotic index of 5 or more mitoses/10 HPFs, the presence of tumor necrosis, nodal metastasis, AJCC stage grouping, large size of largest lymph node metastatic focus and higher post-operative serum calcitonin levels imparted worse DSS, LRFS and DMFS (Figure 3). An increase in mitotic activity subdivided into 3 categories (<2, 2–10, and >10 mitoses/10 HPFs) was associated with poorer DSS, LRFS and DMFS. Marked nuclear pleomorphism imparted worse DSS, LRFS but not DMFS. Tumor encapsulation improved LRFS but had no effect on DSS and DMFS. Infiltration was associated with shortened LRFS. Neither the number of positive nodes nor extra-nodal extension impacted survival. Patients with extrathyroidal extension had poorer DSS and DMFS with no effect on LRFS. Positive margins correlated with worse DSS only. The presence of LVI imparted worse DSS only while extensive LVI correlated solely with worse DMFS. Distant metastasis at presentation correlated with poorer DSS. Age, familial disease, tumor multifocality, degree of intra-tumoral fibrosis, atypical mitosis, the presence of amyloid and post-operative serum CEA did not significantly impact any outcome measure.

The results of subsequent multivariate survival analysis are shown in Table 3. Factors significant on univariate analysis were entered in a multivariate analysis, with the exception of size of largest nodal metastasis (as it would limit the cases to only those with N1 disease only) and nodal metastasis/tumor size/DM at presentation as they overlapped with the AJCC stage group. Tumor necrosis and mitotic activity (using 5 mitoses/10 HPFs as the cutoff) were the only independent predictors for DSS (p=0.004 and 0.020 respectively). A cutoff of 2 and 10 mitoses per 10 HPFs did not reach significance level on multivariate analysis (p=0.388). Tumor necrosis was the only independent prognostic factor for LRFS and DMFS (p=0.001 and 0.003 respectively). Other factors, including sex, AJCC stage group, infiltration, encapsulation, nuclear pleomorphism, extrathyroidal extension, margin status,
and/or post-operative calcitonin level did not independently predict survival (p>0.05). Mitotic levels (both thresholds) did not independently predict LRFS and DMFS (p>0.05).

Proposed grading system and its prognostic significance

Based on the results of the survival analysis, we herein proposed a two-tiered grading system of MTC based on mitosis and necrosis (Table 4). An MTC could be considered as low-grade if it contained <5 mitosis/10 HPF and no tumor necrosis, and as high grade if it has ≥5 mitosis/10HPF and/or tumor necrosis. Using this grading system, 31 MTCs (22%) were classified as high grade, whereas the remaining 113 (78%) were considered as low grade. Univariate survival analysis using log rank test showed that this grading system predicted DSS (p<0.001, Figure 3C), LRFS (p<0.001), and DMFS (p<0.001). Multivariate analysis using Cox proportional model including the proposed grading scheme and the AJCC stage group showed that both grade and stage were independent prognostic factors for LRFS (stage: p=0.029, hazard ratio=1.548, 95% confidence interval: 1.045–2.293; grade: p<0.001, hazard ratio=5.142, 95% confidence interval 2.367–11.170), and DMFS (stage: p=0.014, hazard ratio=2.174, 95% confidence interval: 1.171–4.037; grade: p<0.001, hazard ratio=10.546, 95% confidence interval 2.782–39.977). We also used the three-tiered grading system utilized in pulmonary neuroendocrine tumors (NET) to grade MTC. This system is based on necrosis and a three-tiered mitotic index (cutoff of 2 and 10 mitosis/2 mm²) (16). Although the pulmonary NET grading scheme was overall significant in univariate and multivariate analysis when adjusted for stage grouping (OS: p<0.001, DSS: p=0.001, LRFS: p<0.001, and DMFS: p<0.001), there were only 2 cases classified as grade 3 (equivalent to pulmonary neuroendocrine carcinomas, >10 mitosis/2 mm²), and their survival did not differ from grade 1, whereas the grade 2 MTC were associated with the worst prognosis (supplementary figure 1).

DISCUSSION:

The rarity of MTC and the broad spectrum of its clinical behavior renders prediction of outcome a difficult task (1, 9). A set of clinico-pathologic features have been commonly used to prognosticate these tumors and develop staging systems. By univariate analysis, tumor size, age, male sex, extrathyroidal extension, extent of thyroidectomy, lymph node metastases, distant metastases at presentation, serum calcitonin, CEA blood levels, the presence of familial disease and somatic RET mutation status have been shown to impact survival (1, 4–7, 15, 17–19). However, in multivariate analysis, only age and stage at presentation remain as independent prognostic variables in most publications (1, 4, 18). In congruence with the above studies, we found that male sex, large tumor size, the presence of nodal disease, distant metastases at presentation, stage, elevated post-operative serum calcitonin and extra-thyroidal extension imparted worse survival in univariate analysis. In contrast, older age did not correlate with death or disease recurrence in our cohort. This discrepancy in regard to the prognostic value of age is well known. Indeed, several studies have shown that age does not influence survival in MTC (9, 20, 21). That could be the
reason behind the current American Joint Committee on Cancer (AJCC) system not using age as a staging parameter in MTC.

Many authors have attempted to find histologic markers of outcome in this unpredictable disease. As early as 1966, Williams et al reported an association between a predominant spindle cell pattern, mitosis, necrosis and poorer survival in a series of 67 MTC (22). However, no formal statistical analysis was performed by these authors. Subsequent studies have not shown any prognostic value for cell type, dominant architecture, multicentricity or amyloid deposition (23–25). As shown in Table 2, we confirm the lack of prognostic significance of multicentricity and amyloid deposition.

In 2008, Koperek et al. found a statistically significant correlation between the presence of a desmoplastic stroma and nodal metastasis (26). In contrast, we and others (25, 27, 28) did not find any prognostic value for the presence or the degree of fibrosis in MTC. One possible explanation is the interobserver variability inherent to the evaluation of such a subjective histologic parameter.

In the current series, completely encapsulated and non-infiltrative MTC imparted a statistically better LRFS on univariate analysis. These results are in accordance with the study of Miccoli et al (27). In an analysis of 70 MTC, these authors found that the presence of a complete tumor capsule and lack of infiltration correlate strongly with a high biochemical cure rate and especially an absence of nodal metastasis (27). Therefore, they proposed to use tumor encapsulation and thyroid parenchyma infiltration as tools to stratify patients for cervical lymph node dissection in MTC (27).

The presence of LVI was associated with worse DSS in our patient cohort, and extensive (≥4 foci) LVI with worse DMFS on univariate analysis. This is in congruence with several studies that found a positive correlation between the sole presence of vascular invasion and recurrence and death in MTC (26, 29, 30). While Rios et al. found that the presence of vascular invasion is an independent predictor of recurrence, it lost its predictive value when analyzed in multivariate analysis in the study herein and by others (29). These disparate results could in part be due to differences in the criteria used to diagnose vascular invasion. While we characterized vascular invasion as a tumor thrombus hanging in the lumen of a vessel located in the thyroid or extrathyroidal soft tissue, Pilaete et al describe vascular invasion as “histologic vascular invasion in thyroid specimen or metastatic lymph node” (30). The publication of Rios et al defines vascular invasion as “tumor cells inside or on the vessel walls” without mention of the location of the involved vessels (25). Clearly, additional studies using more uniform criteria for vascular invasion are needed in MTC.

Although nuclear pleomorphism is used to grade many solid tumors, we found very few publications on the subject in MTC. Skopelitou et al. found a correlation between nuclear grade (defined in their article by a combination of nuclear pleomorphism and mitosis) and the proliferative marker Proliferating Cell Nuclear Antigen (PCNA) detected by immunostaining (31). They however did not analyze nuclear pleomorphism on its own and did not assess its effect on survival. In our patient population, nuclear pleomorphism imparted worse DSS and LRFS but not DMFS while Williams et al. did not find such a
correlation (22). There could be many explanations for these disparate findings. The survival impact of nuclear pleomorphism could be confounded by the fact that some MTC display spotty marked nuclear atypia not associated with mitosis (so called endocrine atypia). This phenomenon found in many benign and malignant endocrine and neuroendocrine tumors can be alarming under the microscope but is of no clinical significance (32). Nuclear pleomorphism also suffers from being quite a subjective microscopic feature amenable to significant interobserver variability. For the above reasons, it is our opinion that nuclear pleomorphism is not a robust tool to predict behavior in MTC.

The data on the predictive value of mitosis and necrosis in MTC is controversial. While most studies performed in the last two decades have found necrosis to be of prognostic value in univariate analysis (24, 33, 34), only Franc et al. showed this feature to be an independent predictor of disease-specific mortality (34). We confirm that tumor necrosis is an independent prognosticator of DSS and expand its independent predictive value to LRFS and DMFS. The reason for these discrepant results could reside in the size of the study cohorts. Indeed, the publications showing independent predictive value have a larger patient population (n=144 for our study and n=109 for the one of Franc et al.) than the one of Dottorini et al. (n=53) which did not find necrosis to be significant in multivariate analysis (10). Another explanation could be the smaller rate of patients with tumor necrosis in the paper of Dottorini et al (7.5%) compared to the study herein (21%) and the one of Franc et al (50%).

In regard to mitosis, the data is scant. There is only one published recent paper that analyzed mitosis using appropriate statistics for survival analysis (33). These authors did not find any survival difference based on mitosis by univariate analysis. In contrast we found mitosis to be an independent predictor of DSS. These divergent results could be due to a difference in cut off for mitotic count. In the study of Rios et al (33), a high mitotic index was described as “mitotic figures predominant in the tumor” while we used a cut off of 5 or more mitosis per 10 HPFs to define high mitotic activity. Whatever the reason for the incongruities in the study of mitosis and necrosis and other histologic variables, this uncertainty led the authors of the most recent AJCC not to incorporate these histologic features in their staging system (15).

To the best of our knowledge, this is the first study that shows that mitosis and tumor necrosis are both independent predictors of poor outcome in MTC. The latter is a particularly strong indicator of worse survival since necrosis was an independent prognostic variable for each of DSS, LRFS and DMFS in this cohort. In our analysis, tumor necrosis was found to be independent from established prognostic factors such as sex, extra-thyroidal extension, AJCC stage and post-operative serum calcitonin making it quite valuable and powerful. In addition, mitosis and tumor necrosis are relatively well-defined and objective histologic features less prone to interobserver variability than for example vascular invasion or cell shape. However, the identification of these parameters still requires the skills of a well-trained pathologist. Indeed, it is important to distinguish tumor necrosis from necrosis related to fine needle aspiration. Tumor necrosis is composed of degenerating cytoplasm and punctuate, karyorrhectic nuclear debris (so called “comedo-like” necrosis) (Figures 1 and 2).
It lacks the fibroblastic stromal reaction, hemorrhage or identifiable needle track found in fine needle aspiration induced necrosis (35).

The fact that mitosis and tumor necrosis are so powerful in predicting outcome in MTC should not come as a surprise. The presence of mitosis and/or tumor necrosis is utilized to grade other neuroendocrine and endocrine tumors such as those from the lung and pancreas (12, 13). Armed with that knowledge, we attempted to use the mitotic rate cut off that stratify neuroendocrine tumors of the lung into three separate prognostically different categories (<2, 2–10, >10 mitoses/10 HPFs, ≈2 mm²) (12). Although this three-tiered mitotic system correlated with survival in univariate analysis, it was not an independent predictor of outcome. We also graded MTC using the three-tiered grading scheme utilized in pulmonary NET (tumor necrosis and the three-tiered mitotic index described above). While the pulmonary grading system was overall significant in univariate and even multivariate analysis when adjusted for stage, there were only 2 cases classified as grade 3 (equivalent to pulmonary neuroendocrine carcinomas), and their survival did not differ from grade 1, whereas the grade 2 MTC were associated with the worse prognosis. This data suggests that the three-tiered grading system of pulmonary NET is impractical for MTC. In view of the latter fact together with the apparent superior predictive value of the two-tiered mitotic index cutoff and its simplicity over a three-tiered mitotic rate grouping, we chose the prognostically significant two tiered cut off at 5 mitoses/10 HPFs to construct our proposed grading system. Interestingly, the same mitotic count cut off can define poorly differentiated thyroid carcinomas of follicular cell origin and separate them from their better differentiated counterparts (36).

Since mitosis seems prognostically important in MTC and neuroendocrine tumors in general, one would assume that the proliferative marker ki-67 will be a helpful tool to predict behavior. For example, the current grading system for pancreatic neuroendocrine neoplasms also include Ki-67 proliferation index as a component of the grading system with a three-tiered cutoff of 2% and 20% (13). In a series of 36 patients with MTC, Tinsel et al. showed a strong correlation between the percentage of Ki-67 immunopositive tumor cells and outcome using a semiautomatic image analysis program (37). However, the ki-67 indices of the primary tumors were overall quite low (median 0.41%, range:0.01–4.5%) making this marker difficult to use in everyday practice and a three-tiered cutoff of 2% and 20% akin to what has been used for pancreatic neuroendocrine neoplasms impractical for MTC (37). There is a clear need to further investigate ki-67 and especially phosphohistone H3 (PHH3) (an immunohistochemical marker specific for mitotically active cells) (38) in MTC.

This study has several limitations. First, it is retrospective in nature. This is however almost always the rule in a rare disease such as MTC. Secondly, we did not evaluate the effect of somatic RET or RAS mutation on prognosis since some of our patients were operated in outside institutions making the retrieval of their tissue difficult. In that regard some authors have found that somatic RET mutation status correlated with worse outcome (7, 39) while others did not find such a relationship (40, 41). In a study of 51 sporadic MTC cases, Moura et al. divided their patients into three groups: Those with RET mutations in exons 15 and 16, cases with other RET mutations and patients who were wild type for RET. The group with
RET mutations in exons 15 and 16 had a higher prevalence and number of nodal metastasis than those with other RET mutations (40). There were however no statistically significant survival differences between the three groups of patients (40). Ciampi et al. found a correlation between RAS mutation and better outcome in MTC but it did not reach statistical significance (42). Although the data in the literature is overall encouraging (43), larger studies using multivariate analysis are needed to assess the additive prognostic value of molecular profiling in sporadic MTC.

Despite these limitations, this study has many advantages. It was performed on a large series of patients (n =144) followed in the same institution. It consisted of a very meticulous and especially detailed histopathologic analysis. Recurrence was defined as structural recurrence and therefore as a clinically relevant outcome measure. It demonstrated that mitosis and necrosis are very powerful indicators of poor survival in MTC independent from well-established prognostic factors such as AJCC stage at presentation and post-operative serum calcitonin. In the clinical setting, the American Thyroid Association (ATA) guidelines state that “TNM classification and other factors, such as the postoperative calcitonin level and the calcitonin and CEA doubling times, should be used to predict outcome and to help plan long-term follow-up of patients with MTC” (1). In the opinion of the ATA panel, stage lacks important prognostic factors such as calcitonin and CEA doubling times (1). However, these latter parameters change over time, being longer when the disease is in its early stages and shorter in later stages, when there is disease progression (42, 44). The grading of MTC based on mitosis and necrosis will hopefully pave the way to a customized patient follow-up and therapy from diagnosis. Herein, we propose to grade MTC into low grade (<5 mitosis/10 HPF and no tumor necrosis) and high grade (≥5 mitosis/10HPF and/or tumor necrosis, Table 4). This grading system is independent from stage in predicting LRFS and DMFS and the sole independent prognostic factor for DSS in our cohort. This will enable the treating clinician to better counsel the patient, design the optimal follow up and better select those individuals who may benefit from systemic therapy.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Figure 1.
66-year-old male with a medullary thyroid carcinoma (MTC) containing tumor necrosis in the primary treated initially by thyroidectomy and lymph node dissection. Patient was M0 at presentation with post-operative serum calcitonin and CEA of 898 pg/ml and 2.4 ng/ml respectively. The patient died of disease 2 years and 4 months after initial surgery. A: Medium power view of an H&E section showing a classical MTC (m) growing in solid nests infiltrating adjacent non-neoplastic thyroid (thy) (100x). B: Immunostain for calcitonin on the same tissue section present in A confirming the MTC diagnosis (100X). Tumor is immunopositive for calcitonin (m) while the non-neoplastic thyroid is negative (thy) (100x). C: H&E from another area of the primary tumor showing fresh tumor necrosis (n) surrounded by viable tumor (m) (200x). D: High power view of an H&E from another focus of tumor necrosis (n) showing nuclear debris (arrow). This appearance of the necrosis is referred to as “comedo-like”. The viable tumor cells (m) display no significant nuclear pleomorphism (200x).
Figure 2.
53-year-old male with a medullary thyroid carcinoma (MTC) containing tumor necrosis in the regional nodes at presentation treated initially by thyroidectomy and lymph node dissection. Patient was M0 at presentation with post-operative calcitonin and CEA of 124 pg/ml and 20.4 ng/ml respectively. The patient died of disease 4 years and 4 months after initial surgery. A: Medium power view of an H&E section from the primary showing a classical MTC (m) growing in solid nests adjacent to non-neoplastic thyroid (thy) (100x). B. Medium power view of an H&E section from a regional node with metastatic tumor at presentation. There is fresh tumor necrosis (n) surrounded by viable tumor (m) (200x). C. Higher power view of the necrotic area in B showing tumor necrosis (n) containing nuclear debris (arrow). The viable tumor cells (m) do not show significant nuclear atypia (200x). D. Immunostain for calcitonin on the same tissue section present in B confirming the MTC diagnosis. The viable tumor surrounding the necrotic focus (n) is immunopositive for calcitonin (m) (100X).
Figure 3.
Kaplan-Meier plots demonstrating the impacts of mitotic index (A), tumor necrosis (B), and the proposed grading system (C) on disease specific survival.
### Table 1:
Clinico-pathologic characteristics of 144 patients with medullary thyroid carcinoma (MTC) cohort

| Characteristics          | Number of patients (%) | Characteristic          | Number of patients (%) |
|--------------------------|------------------------|-------------------------|------------------------|
| Age, years               |                        | Encapsulation (n=143)   |                        |
| Median (years)           |                        | Completely encapsulated | 26 (18%)               |
| <55                      |                        | Partially encapsulated  | 57 (40%)               |
| ≥55                      |                        | None                    | 60 (42%)               |
| Sex                      |                        | Lymphovascular invasion (LVI)|                   |
| Male                     | 73 (51%)               | Absent                  | 103 (72%)              |
| Female                   | 71 (49%)               | Present                 | 41 (28%)               |
| Tumor size (n=141)       |                        | Extent of LVI (n=40)    |                        |
| ≤2cm                     | 80 (57%)               | Focal <4 foci           | 31 (77.5%)             |
| 2.1–4cm                  | 42 (30%)               | Extensive (≥4 foci)     | 9 (22.5%)              |
| >4cm                     | 19 (13%)               | Extrathyroidal LVI (n=41)|                      |
| Mitotic index a          |                        | Absent                  | 21 (51%)               |
| <2/10 HPFs               | 117 (82%)              | Present                 | 20 (49%)               |
| 2–10/10 HPFs             | 25 (17%)               | Extrathyroidal extension (n=143)|  |
| > 10/10 HPFs             | 2 (1%)                 | Absent                  | 99 (69%)               |
| Mitotic index b          |                        | Present                 | 44 (31%)               |
| <5/10 HPFs               | 137 (95%)              | Fibrosis (n=143)        |                        |
| ≥5/10 HPFs               | 7 (5%)                 | No fibrosis             | 6 (4%)                 |
| Atypical mitosis (n=52)  |                        | Mild fibrosis           | 28 (20%)               |
| Absent                   | 32 (62%)               | Moderate fibrosis       | 46 (32%)               |
| Present                  | 20 (38%)               | Prominent fibrosis      | 63 (44%)               |
| Tumor necrosis           |                        | Separate focus of MTC (n=129)|                   |
| Absent                   | 114 (79%)              | Absent                  | 98 (76%)               |
| Present                  | 30 (21%)               | Present                 | 31 (24%)               |
| Nuclear pleomorphism     |                        | C cell hyperplasia (n=103)|                   |
| Mild                     | 125 (86%)              | Absent                  | 85 (83%)               |
| Moderate                 | 18 (13%)               | Present                 | 18 (17%)               |
| Marked                   | 1 (1%)                 | Lymph node status       |                        |
| Variant                  |                        | N0/Nx                   | 60 (42%)               |
| classical                | 142 (98%)              | <5 positive LN          | 29 (20%)               |
| follicular variant       | 1 (1%)                 | ≥5 positive LN          | 55 (38%)               |
| paraganglioma like       | 1 (1%)                 | Size of largest nodal metastasis (n=81)|              |
| Amyloid (n=142)          |                        | <1 cm                   | 26 (32%)               |
| Absent                   | 59 (42%)               | ≥1 cm                   | 55 (68%)               |
| Present                  | 83 (58%)               | Extranodal extension (n=82)|                   |
| Infiltration (n=143)     |                        | Absent                  | 19 (23%)               |
| Absent                   | 19 (13%)               | Present                 | 63 (77%)               |
| Present                  | 124 (87%)              | Distant metastasis at presentation|               |
| Margin (n=135)           |                        | M0                      | 135 (94%)              |

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| Characteristics                        | Number of patients (%) | Characteristic | Number of patients (%) |
|---------------------------------------|------------------------|----------------|------------------------|
| Negative                              | 116 (86%)              | M1             | 9 (6%)                 |
| Positive                              | 19 (14%)               |                |                        |
| RET germline mutation status (n=134)  |                        |                |                        |
| AJCC 8th prognostic stage groups      |                        |                |                        |
| I                                     | 43 (30%)               | Wild type      | 106 (79%)              |
| II                                    | 16 (11%)               | Mutated        | 28 (21%)               |
| III                                   | 22 (15%)               | Proposed grading |                       |
| IV                                    | 43 (62%)               | Low grade      | 113 (78%)              |
|                                       |                        | High grade     | 31 (22%)               |

a Mitotic rate used in classification of neuroendocrine tumors of lung

b Mitotic rate used in Memorial Sloan Kettering Cancer Center (MSKCC) definition of poorly differentiated follicular cell-derived thyroid carcinoma

HPFs: high power fields, LN: lymph node

AJCC: American Joint Committee on Cancer

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Table 2:
Correlation between clinicopathologic variables and outcomes

| Characteristics                                      | DSS   | LRFS  | DMFS  |
|------------------------------------------------------|-------|-------|-------|
| Age                                                  | 0.742 | 0.227 | 0.641 |
| Sex                                                  | 0.008 | 0.050 | 0.018 |
| AJCC prognostic stage group                          | 0.014 | 0.003 | 0.012 |
| Tumor size                                           | 0.001 | 0.042 | 0.013 |
| Mitotic index (cut off 2 and 10/10 HPFs)             | 0.025 | 0.029 | <0.001|
| Mitotic index (cut off 5/10 HPFs)                    | <0.001| 0.004 | 0.001 |
| Atypical mitosis                                     | 0.431 | 0.441 | 0.435 |
| Tumor necrosis                                       | <0.001| <0.001| <0.001|
| Nuclear pleomorphism                                 | <0.001| 0.049 | 0.072 |
| Amyloid                                              | 0.611 | 0.972 | 0.538 |
| Fibrosis                                             | 0.501 | 0.060 | 0.269 |
| Infiltration                                         | 0.201 | 0.033 | 0.331 |
| Encapsulation                                        | 0.614 | 0.010 | 0.704 |
| Lymphovascular invasion (LVI)                        | 0.041 | 0.090 | 0.408 |
| Extent of VI                                         | 0.526 | 0.609 | 0.009 |
| Extrathyroidal VI                                    | 0.145 | 0.287 | 0.188 |
| Extrathyroidal extension                             | 0.013 | 0.165 | <0.001|
| Margin                                               | 0.013 | 0.623 | 0.051 |
| Separate focus of MTC                                 | 0.679 | 0.539 | 0.689 |
| Nodal status                                         | 0.042 | 0.001 | 0.005 |
| Number of metastatic LN (<5 vs. ≥5)                  | 0.324 | 0.959 | 0.859 |
| Size of largest nodal metastasis                     | 0.022 | 0.003 | 0.012 |
| Extranodal extension                                 | 0.215 | 0.642 | 0.096 |
| Post-operative serum calcitonin (all cases)          | 0.001 | 0.001 | 0.004 |
| Post-operative serum calcitonin (excluding those with DM at presentation) | 0.003 | <0.001 | 0.004 |
| Post-operative serum CEA                             | 0.216 | 0.392 | 0.188 |
| DM at presentation                                   | 0.046 | 0.504 | NA    |
| Familial MTC                                         | 0.492 | 0.724 | 0.844 |

\(a\) Mitotic rate used in classification of neuroendocrine tumors of lung

\(b\) Mitotic rate used in MSKCC definition of poorly differentiated follicular cell-derived thyroid carcinoma

DSS: Disease specific survival, LRFS: Loco-regional free survival, DMFS: Distant metastasis free survival, MTC: Medullary thyroid carcinoma, CEA: carcinoembryonic antigen, DM: Distant metastasis, NA: not available.

CEA and calcitonin are log-transformed.

Values in red: significant p values.

AJCC: American Joint Committee on Cancer
Table 3:
Multivariate analysis of clinico-pathologic parameters impacting survival

|                | P values | Hazard ratio | 95% CI       |
|----------------|----------|--------------|--------------|
| **DSS**        |          |              |              |
| Sex            | 0.287    | 0.257        | 0.021–3.132  |
| AJCC stage group | 0.405   | 2.255        | 0.332–15.317 |
| Mitotic index (<5 vs. ≥5/10 HPFs) | 0.026 | 58.302       | 1.621–2096.993 |
| Tumor necrosis  | 0.008    | 20–497       | 2.179–192.788 |
| Nuclear pleomorphism | 0.314 | 3.479        | 0.308–39.286  |
| Lymphovascular invasion | 0.451 | 1.942        | 0.346–10.913  |
| Extrathyroidal extension | 0.929 | 0.881        | 0.054–14.299  |
| Margin         | 0.979    | 1.024        | 0.168–6.235  |
| Post-op calcitonin level | 0.062 | 2.433        | 0.955–6.198   |
| **LRFS**       |          |              |              |
| Sex            | 0.595    | 0.777        | 0.307–1.969  |
| AJCC stage group | 0.195   | 1.369        | 0.851–2.201  |
| Mitotic index (<5 vs. ≥5/10 HPFs) | 0.177 | 3.085        | 0.601–15.841  |
| Tumor necrosis  | 0.001    | 4.110        | 1.770–9.541  |
| Nuclear pleomorphism | 0.144 | 2.476        | 0.733–8.367  |
| Infiltration    | 0.976    | 603263.045   | 0-indefinite |
| Encapsulation   | 0.173    | 0.594        | 0.280–1.257  |
| Post-op calcitonin | 0.161 | 1.271        | 0.909–1.778  |
| **DMFS**       |          |              |              |
| Sex            | 0.323    | 0.497        | 0.124–1.986  |
| AJCC stage group | 0.136   | 1.837        | 0.827–4.085  |
| Mitotic index (<5 vs. ≥5/10 HPFs) | 0.209 | 4.264        | 0.444–40.942  |
| Tumor necrosis  | 0.003    | 5.166        | 1.748–15.271 |
| Extrathyroidal extension | 0.820 | 0.861        | 0.238–3.116  |
| Post-op calcitonin | 0.217 | 1.453        | 0.803–2.632  |

HPF: High power fields, 400x, CI: confidence interval.

Values in red: significant p values.

AJCC: American Joint Committee on Cancer
Table 4:
Proposed grading system for medullary thyroid carcinoma.

| Grade   | Mitoses/10 HPFs | Tumor Necrosis |
|---------|-----------------|----------------|
| Low     | <5              | no             |
| High    | >=5             | and/or         |

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