Sentinel lymph node biopsy in medullary thyroid microcarcinomas

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Abstract. The aim of this prospective study was to analyze accuracy of sentinel lymph node biopsy with methylene blue dye for intraoperative detection of lateral metastases in clinically N0M0 medullary microcarcinomas with calcitonin <1,000 pg/mL and selection of true-positive patients for one-time therapeutic lateral dissection. In addition to total thyroidectomy and central neck dissection, all patients had bilateral sentinel biopsy of jugulo-carotid regions after methylene blue injection to decide upon necessity for lateral dissection. If sentinels were benign on frozen section, additional non-sentinels were extirpated, with no further lateral dissection. If sentinels were malignant, one-time lateral dissection was performed. 20 patients were included in this study. Hereditary disease form was observed in 3/20 (15%) of patients with RET proto-oncogene mutation C634F; remaining 17/20 (85%) were negative for germline mutations. There were no allergic reactions to methylene blue and identification rate of sentinels was 100%. In total, 2/20 (10%) cN0 patients had lymphonodal metastases, thus were reclassified as pN1b. Remaining 18/20 (90%) were classified pN0 based on standard pathohistology. Frozen section findings on sentinels were 100% match with standard pathohistology, and there were no skip metastases in lateral compartments. Sensitivity, specificity and accuracy of sentinel biopsy method with methylene dye and frozen section were 100%. Dzodic’s sentinel lymph node biopsy method can be used for intraoperative assessment of lateral compartments and optimization of initial surgery of medullary microcarcinomas with calcitonin <1,000 pg/mL. This way, cN0 patients with sentinel metastases can receive one-time lateral dissection, and those without benefit from less extensive surgery.

Key words: Medullary thyroid microcarcinoma, Calcitonin below 1,000, Methylene blue dye, Sentinel lymph node biopsy, Lateral neck dissection

MEDULLARY THYROID CARCINOMA (MTC) is a rare thyroid malignancy of C cell origin, characterized by the secretion of a peptide hormone calcitonin (CT). It is accounting for 3% (adults) to 10% (children) of thyroid cancers [1]. MTC appears in two forms: sporadic (75%) and hereditary (25%), with similar gender distribution [1]. MTCs have more aggressive behavior than differentiated thyroid carcinomas, and tend to spread relatively early into regional lymph nodes (LN) [2]. It has been reported that 80% of patients with palpable MTCs have central compartment LN metastases at the time of diagnosis, while 75% and 47% (respectively) have LN metastases in ipsilateral and contralateral jugulo-carotid regions [3].

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Medullary thyroid microcarcinomas (micro-MTC) are defined as tumors with diameter 10 mm or less [4]. Although clinical relevance of sporadic and hereditary micro-MTCs is debatable in literature, there is a significant disease burden associated with this entity [5-11].

Serum CT level is a precise MTC marker, both for preoperative diagnostics, as well as for postoperative assessment of disease relapse. It usually correlates with tumor size and disease extent. However, LN and distant metastases are diagnosed, as well, in patients with lower preoperative CT values [12].

Surgery is the only curative treatment for MTCs. Aim of initial surgery is adequate tumor and LN clearance to achieve good loco-regional disease control [1-3, 13], and, if possible, biochemical cure, commonly used as a surrogate marker for surgical cure [10]. False negative findings on preoperative physical examination and ultrasonography of regional LNs can lead to initial under-treatment of patients with this disease. Due to permanent concerns regarding under- and over-treatment of patients,
surgical recommendations for MTC management are not unanimous, but majority of those agree upon necessity of total thyroidectomy (TT) with prophylactic central lymph node dissection (CLND) [1, 2, 13-18]. The necessity of lateral lymph node dissections (LLND) in clinically evident disease is indisputable; however, its extent in clinically N0 patients is a matter of debate, ranging in the literature from (a) bilateral prophylactic LLND [10, 12, 16, 19], as the most radical approach; or (b) sampling of lateral LNs, as less radical strategy to determine the necessity of complete LLND in histologically proven metastatic disease [20]; to (c) performing ipsilateral LLND only in case of positive central LNs [1, 21-23].

In the context of timely detection and treatment of subclinical, occult, LN disease, while, on the other hand, not over-treating patients without metastases, some authors perform sentinel lymph node biopsy (SLNB), first introduced by Kelemen and co-authors in 1998 [24]. Based on literature data, this method is most commonly used in selection of patients for CLND in differentiated thyroid carcinomas [25-31]. Given that CLND is performed in our center as a standard surgical protocol for thyroid carcinomas, our leading surgeon Prof. Radan Dzodic (last author) introduced SLNB technique for intraoperative staging of lateral LNs in clinically negative (cN0) patients, and published first results in 2006 [32]. Initially, this method was used for differentiated thyroid carcinomas only, but later for MTCs, as well.

The aim of this paper was to present results of the first study that analyzed accuracy of SLNB with methylene blue dye (MBD) for intraoperative staging of lateral LNs in cN0 micro-MTCs with serum CT levels below 1,000 pg/mL, and its usefulness in selection of patients for one-time therapeutic LLND.

Materials and Methods

Patients

This prospective study started in 2007, as a part of the project of the Ministry of Science of Republic of Serbia (project number: 111601/2005; project name: “Early detection of lymphonodal metastases by pathohistological verification of sentinel lymph nodes in malignant epithelial tumors”), with aim to include patients with suspicious or confirmed sporadic or hereditary MTCs meeting following criteria: (a) tumor diameter 10 mm or less, without extrathyroid extension (b) serum CT levels lower than 1,000 pg/mL, (c) clinically negative regional LNs and (d) absence of distant metastases (cM0). It is set in the Institute for Oncology and Radiology of Serbia (IORS).

Patients were either diagnosed primarily in IORS, or they were referred to our center for surgery of suspicious MTC. MTC diagnosis was set based on (a) presence of specific germline mutations or (b) suspicious thyroid tumor or thyroid goiter on physical examination and neck ultrasonography (US) with elevation of basal serum CT levels. If basal CT elevation was inconclusive for MTC diagnosis, calcium-stimulating test was performed for evaluation of stimulated CT increase and decision upon surgery.

Standard preparation for surgery in general anesthesia included: complete thyroid and parathyroid function, complete blood and biochemistry tests, indirect laryngoscopy, chest and tracheal X-ray, abdominal US, internist and anesthesiologist evaluation. In all patients concomitant pheochromocytoma and primary hyperparathyroidism were preoperatively excluded. Preoperative diagnostics was case-oriented in matters of contrast-enhanced computerized tomography (CECT) and fine needle aspiration biopsy (FNAB).

Multidisciplinary IORS Thyroid team decided upon the surgical treatment. Patients were explained in detail the surgical procedure, possible complications and treatment outcomes. Written consent for surgical treatment was obtained from each patient. Surgical protocols for clinically N0 thyroid carcinomas in our center include SLNB with MBD for intraoperative LN staging, and they were approved by IORS Ethical Committee.

Study design

Gender-, age-, disease- and treatment-specific characteristics of the group were analyzed. Given the defined aim, to assess the accuracy of SLNB for intraoperative staging of lateral LNs, we have compared findings of frozen section analysis (FSA) and standard pathohistological analysis (sPH) on sentinel-LNs, as well as evaluated additional lateral LNs on sPH for skip metastases (metastases in non-sentinel-LNs). Sensitivity, specificity, positive and negative predictive values and overall accuracy of the method were calculated.

Surgical treatment

Surgical treatment was uniform for all selected patients, and consisted of TT, prophylactic CLND (level VI), SLNB of both jugulo-carotid regions and FSA of sentinel-LNs. If sentinel-LNs were benign, additional surrounding non-sentinel-LNs from belonging jugulo-carotid region were removed for sPH. Whenever metastases were detected in sentinel-LNs on FSA, a therapeutic LLND was done.

Standard Kocher’s collar incision is made in the natural skin fold, between anterior borders of sternocleidomastoid muscles. After transection of platysma, middle-line dissection and lateralization of infrathyroid muscles, thyroid gland is exposed. Specific surgical technique is
explained in details in next few steps.

Step 1: Peritumoral thyroid injection of 0.2–0.5 mL of 1%-MBD solution (using a 27-gauge needle) beneath thyroid capsule, with thermal coagulation of the injection site through surgical tweezers (Fig. 1). In order to avoid MBD leakage, thyroid capsule should not be damaged. Coloring of lymphatic vessels and LNs around thyroid gland is observed immediately (Fig. 2).

Step 2: Removal of the lobe with the suspicious tumor and FSA for pathohistological confirmation of the MTC. Parathyroid glands (PTGs) are not colored with MBD, so their identification and preservation is easier (Fig. 3).

Step 3: Exploration of ipsilateral jugulo-carotid region, focusing on levels II and III, in order to identify colored afferent lymphatic vessels (Fig. 4) and blue-stained, sentinel-LNs that are meticulously removed (Fig. 5) and examined by FSA. If there are no colored LNs, LN belonging to the colored afferent lymphatic is considered to be the sentinel-LN, thus is removed for FSA.

Step 4: Removal of additional surrounding non-sentinel-LNs if sentinel-LNs are benign on FSA; or one-time therapeutic LLND of the side where sentinel-LN is proven to be malignant on FSA.

Step 5: Completion thyroidectomy and level VI clearance, with careful preservation of PTGs on venous-arterial vascular pedicles [33], as well as identification and preservation of both recurrent laryngeal nerves. Due to MBD injection, central LNs are clearly marked with blue, while PTGs remain non-colored, making CLND easier and safer, especially in less experienced hands.

Fig. 1 Injection of 0.2–0.5 mL of 1%-methylene blue dye solution using a 27-gauge needle in thyroid lobe, beneath thyroid capsule, with thermal coagulation of the injection site through surgical tweezers, to avoid the leakage of the tracer

Fig. 2 Methylene blue dye affecting thyroid lobes, lymphatic vessels and lymph nodes of central compartment

Fig. 3 Removal of thyroid lobe, with preservation of parathyroid glands that are not colored with methylene blue dye, while central lymph nodes uptake the vital dye

Fig. 4 Identification of blue-stained afferent lymphatic vessel in the jugulo-carotid region
a PTG is accidentally removed during CLND or devascularized, it should be minced and autotransplanted by Wells technique into the sternocleidomastoid muscle [34].

Step 6: SLNB of contralateral jugulo-carotid region with FSA and further procedure, as described in step 4.

Postoperative course
In the postoperative course, all patients were monitored and managed according to our IORS protocols for post-thyroidectomy patients: measurement of parathyroid hormone, serum calcium and phosphorus levels immediately after surgery and on the first postoperative day, and active evaluation of signs/symptoms of hypocalcaemia, laryngeal nerves injuries or other complications.

After postoperative multidisciplinary team evaluation of each patient, L-thyroxin was initiated. General recommendations for follow-up of MTC patients in our center include: serum CT and CEA measurements to verify biochemical status of the disease, US in all patients, and CECT or $^{99m}$Tc-V-DMSA scans, if indicated, for local or distant relapse assessment. All patients were followed-up in our tertiary center.

Statistical analysis
SPSS (SPSS Inc., IBM, Chicago, USA), version 23, was used for statistical analysis. Descriptive statistical methods (frequencies, percentages, mean, standard deviation (SD) and range) were used to summarize data on patients, disease, treatment and outcomes. MedCalc statistical software for Windows was used for evaluation of diagnostic value of the SLNB method (sensitivity, specificity, positive and negative predictive values and overall accuracy).

Results
In total, from 2007 to 2019, 20 patients met all four criteria and were eligible for inclusion in this study. There was a female predominance in the sample (90%, 18/20 patients). Mean age of patients was 53.90 ± 17.10 (range: 18 to 83) years.

Preoperative findings
Basal CT ranged from 7.2 to 697.0 pg/mL in the group, with a mean value of 234.5 pg/mL and great variations among patients (SD = 217.6).

Hereditary form of disease was observed in three patients (15%) with RET proto-oncogene mutation C634F. Patient 1 was a male, aged 46, with basal CT of 221.6 pg/mL, a suspicious tumor in the left lobe and negative LNs on US and physical examination. Patient 2 was a female, aged 18, with basal CT of 42.0 pg/mL, a suspicious tumor in the right lobe and clinically negative LNs. Patient 3 was a female, aged 72, with basal CT of 42.6 pg/mL, bilateral thyroid tumors on US and physical examination, and negative LNs. Patient 1 and 2 had no clinical or biochemical signs of adrenal or PTG tumors. Patient 3 had prior surgery for pheochromocytoma and PTG adenoma. Remaining 17 patients (85%) were negative for germline mutations based on standard genetic testing.

FNAB was done by endocrinologist in 4/20 (20%) patients, suggesting of colloid goiter in half and normal thyroid tissue in other 50% of patients.

Surgical treatment
All patients had TT, prophylactic CLND (level VI), bilateral SLNB of jugulo-carotid regions and FSA of sentinel-LNs. None of the patients had allergic reactions to MBD. In all patients blue-stained LNs were identified in lateral compartments (100% identification rate).

In 90% (18/20) of patients examined sentinel-LNs were benign on FSA (Table 1). Here, additional surrounding non-sentinel-LNs were removed from the respective jugulo-carotid regions for sPH analysis. In 10% (2/20) of patients, both with hereditary MTC (patient 1 and 2), LN metastases were detected in sentinel-LNs on FSA, thus therapeutic LLNDs of respective jugulo-carotid compartments were done. Patient 1 had bilateral LLND of regions II-V, since sentinel-LN metastases were found in both jugulo-carotid regions, while LLND was performed on the right side in patient 2.

Pathohistological reports
Unilateral unifocal tumors were found in 90% (18/20) of patients, presenting as sporadic MTC in 16 and hereditary form in 2 patients. One patient (5%) with sporadic disease form had unilateral multifocal MTC,
and one patient (5%) with hereditary MTC had bilateral microcarcinomas (Fig. 6). As shown in Table 2, mean tumor size in the group was 8.05 ± 2.04 (range: 3 to 10) mm. Capsular and lymphovascular invasion was not observed in this group. Associated thyroid disease was present in 50% (10/20) of patients, which is given in details in Table 3.

The average number of dissected central LNs was 3.10 ± 1.48 (range: 2–7), and total number of sentinel-LN metastases in two patients selected on FSA was 1/1 LN per both lateral compartments in patient 1 (Fig. 7) and 1/4 LNs in ipsilateral compartment of patient 2.

A mean of 11.15 ± 6.33 (range: 6–28) non-sentinel-LNs were removed from jugulo-carotid regions (LLND included). In 2 patients that were FSA-selected for one-time LLND, number of additional lateral LN metastases was 2/11 and 2/12 in bilateral LLND in patient 1, and 1/15 in unilateral LLND in patient 2.

Average number of dissected central LNs was 9.61 ± 2.54 (range: 5–15). Central LN metastases were verified in those 2 patients with sentinel-LN metastases on FSA. In patient 1, 3/12 central LNs were affected, while patient 2 had metastasis in 1/9 dissected central LNs.

### SLNB diagnostic value

Definite pathohistological reports on sentinel-LNs were 100% match with FSA findings. In total, 10% (2/20) of patients were reclassified from cN0 to pN1b due to SLNB with MBD and FSA. Remaining 90% (18/20) of patients were confirmed to be pN0 based on sPH analysis of central LNs, sentinel-LNs and non-sentinel-LNs of lateral compartments.

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**Table 1** Pathohistological characteristics of sentinel lymph nodes

| SLN characteristics | % (n) of patients |
|---------------------|------------------|
| Identification rate | 100% (20/20)     |
| FSA findings        |                  |
| benign              | 90% (18/20)      |
| malignant           | 10% (2/20)       |
| sPH findings        |                  |
| benign              | 90% (18/20)      |
| malignant           | 10% (2/20)       |
| FSA and sPH match   | 100% (20/20)     |
| TOTAL               | 100% (20)        |

Legend: SLN, sentinel lymph node; FSA, frozen section analysis; sPH, standard pathohistology

**Table 2** Pathohistological characteristics of medullary thyroid microcarcinomas

| Tumor characteristics |
|------------------------|
| % (n) of patients      |
| Tumor size             | 8.05 ± 2.04 mm |
| range                  | 3–10 mm       |
| Capsular invasion      |                 |
| no                     | 100% (20/20)  |
| yes                    | 0% (0/20)     |
| Lymphovascular invasion|               |
| no                     | 100% (20/20)  |
| yes                    | 0% (0/20)     |
| Tumor pattern          |                 |
| unilateral unifocal    | 90% (18/20)   |
| sporadic              | 88.9% (16/18) |
| hereditary             | 11.1% (2/18)  |
| unilateral multifocal* | 5% (1/20)     |
| bilateral, multicentric** | 5% (1/20) |
| TOTAL                  | 100% (20)     |

Legend: * sporadic MTC; ** hereditary MTC

**Table 3** Pathohistological reports on associated thyroid diseases

| Associated thyroid disease                                      |
|----------------------------------------------------------------|
| % (n) of patients                                              |
| None                                                           | 50% (10/20) |
| Hashimoto’s thyroiditis                                        | 15% (3/20)  |
| Follicular adenoma                                             | 15% (3/20)  |
| C-cell hyperplasia                                             | 5% (1/20)   |
| Micro-PTC                                                      | 10% (2/20)  |
| Micro-PTC + Hurthle cell tumor of uncertain malignant potential| 5% (1/20)   |
| TOTAL                                                          | 100% (20)   |

Legend: PTC, papillary thyroid carcinoma
In 18 patients that had negative sentinel-LNs on FSA and sPH there were no skip metastases in additional LNs that were extirpated for analysis. In 2 patients that were FSA-selected for one-time LLND of affected compartments, sPH examination revealed additional LN metastases in dissected compartments, meaning positive sentinel-LNs on FSA were 100% predictive for presence of additional lateral LN metastases.

Sensitivity, specificity, positive and negative predictive value of SLNB method with MBD and FSA were 100%. The overall accuracy of the method was 100% (Table 4).

Postoperative follow-up data

In the immediate postoperative course, all patients were carefully evaluated for complications. There were neither postoperative bleedings, nor permanent hypothyroidism in the group. One patient (5%) was with mild hoarseness, that resolved spontaneously (thus was attributed to neuropraxia), but he also had a prolonged lymphatic leakage that could not have been resolved with conservative approach, so a re-do surgery was performed with suture of injured lymphatic vessel on the right side and full recovery after being discharged from hospital.

Patients were followed-up in average 79.75 ± 36.40 (range: 1–145.75) months. During this time, serum CT and CEA were within normal ranges in all patients, suggesting complete biochemical cure.

Discussion

Although micro-MTCs have earlier been considered harmless, studies that analyzed clinical behavior of this entity have shown a significant disease burden associated with micro-MTCs [5-11].

Role of CT screening for active detection of MTCs, especially in nodular goiter patients, remains debatable in the literature [10, 19, 35]; however, this is the routine clinical practice in our center.

Every increase in basal CT levels should raise suspicion of MTC and further diagnostics should be performed. Still, lower CT values might not always be of diagnostic significance, due to potential confounding effects of comorbidities (e.g. sporadic C-cell hyperplasia, hypercalcaemia, neuroendocrine tumors, etc.), drugs (e.g. proton pump inhibitors), smoking, or due to assay related factors, gender specific differences, and other [1]. Stimulated CT should help in differential diagnosis, although test is not available in all countries.

Preoperative CT usually correlates to tumor size and disease extent; however, it can lead to inadequate disease staging and under-treatment, given that patients may have LN metastases in lower CT levels [12]. In our study, two patients with LN metastases had basal CT levels of 221.6 and 42.0 pg/mL, while those with higher values, over 500 pg/mL, for example, were staged pN0. Physical examination and US evaluation are used to guide the extent of LN surgery, but are not always reliable [1, 15]. Intraoperative assessment of LNs by a surgeon was also shown not to be accurate enough, with sensitivity of 64% [36].

General recommendation for treatment of MTC is TT with prophylactic CLND, even in patients with micro-MTCs [1, 2, 13-18], and this surgical strategy is used in

Table 4  Diagnostic values of sentinel lymph node biopsy method using methylene blue dye and frozen section analysis

| SLN status  | Lateral lymph nodes status | TOTAL (n) | Diagnostic SLN biopsy value |
|------------|---------------------------|----------|---------------------------|
| Positive (n) | Positive (n) | Negative (n) | 2 | PPV = 100% (2/2) |
| Negative (n) | 0 | 18 | 18 | NPV = 100% (18/18) |
| TOTAL       | 2 | 18 | 20 | |

SN = 100% (2/2)  SP = 100% (18/18)  FNR = 0% (0/2)  ACC = 100%

Legend: n, number of patients; SLN, sentinel lymph node; SN, sensitivity; SP, specificity; FNR, false negative rate; PPV, positive predictive value; NPV, negative predictive value; ACC, accuracy
our center, as well. However, the necessity and the extent of LLND are debatable in the referent literature. Undoubtedly, in clinically evident lateral LN metastases (macroscopically or by US), therapeutic LLND is required. However, recommendations are not unanimous regarding the indications for LLND in the clinical absence of metastases.

Many studies upon clinical behavior of MTCs acknowledged that LN metastases occur early in the course of MTC, regardless the tumor size [2, 37]. For example, at the time of diagnosis, over 70% of patients with palpable MTC [36] and over 35% of micro-MTCs [9] have LN metastases. In addition, the risk of lateral LN involvement in patients with central LN metastases is at least 70% [23], while over a third of patients with central also have contralateral lateral LN metastases [38].

Due to these data reports on high rate of LN metastases in MTCs and many reports on prognostic significance of initial surgery for patients’ survival [39], significant number of surgeons have approached to MTC treatment more extensively, even in the absence of lateral LN involvement. Namely, majority of surgical teams [2, 10, 12, 16, 19, 36, 37] advocate bilateral prophylactic LLND in all MTC patients. The aim is to achieve higher biochemical cure rate (postoperative CT normalization), lower rate of reoperations and better survival rates. Some data suggest that contralateral LLND can be omitted in sporadic MTCs, if there is no evidence of central and ipsilateral lateral LN involvement, but it is still necessary to perform routine bilateral LLNDs in hereditary forms [1, 2].

Less radical surgical approach for clinically N0 patients implies performing ipsilateral LLND only in case of positive central LNs [1, 21-23], or if sampling of lateral LNs harbor metastatic disease [20].

Break-through in minimally invasive approach to LN management in thyroid carcinomas has to be contributed to Kelemen [24], who used isosulfan blue dye (IBD) to mark LNs of the central and lateral neck regions, based on well-established, widely used model of SLNB in breast carcinoma and melanoma. Main goals of SLNB are: to accurately detect LN metastases, to optimize the extent of LN surgery (patients with metastases are selected for complete dissection, others not), to minimize morbidity from unnecessary surgery.

There are various tracers available for LN mapping, like: vital blue dyes (MBD, IBD and Patent Blue V dye), fluorescent dyes (indocyanine green), radionuclide (Tc99m-labeled radiocolloid), Carbon nanoparticles or magnetic particles [15]. Majority of tracers require some additional equipment for imaging, like: (a) gamma camera and hand-held gamma probe or single-photon emission computed tomography for radionuclide tracers [40], (b) SPY Elite system and Hamamatsu PDE-Neo probe for indocyanine green fluorescence [41], or (c) magnetic particle imaging scanner and handheld magnetic probe if magnetic particles are used [42]. Unavailability of these tracers due to health care regulations and the costs and complexity of imaging equipment can be challenging, especially for hospitals in developing countries. Certain tracers are also associated with complications like skin tattooing, especially carbon black, reported by Zhang et al. to leave a black staining for over a year after breast surgery in 42% of patients [43]. In thyroid surgery, injection is made into the thyroid gland; however, the skin staining might occur due to tracer leakage. Anaphylactic reactions associated with vital dyes have also been reported in the literature; however, all are contributed to IBD and patent blue, none to MBD injection [44]. Indocyanine green can also lead to allergic reactions in patients with iodine or shellfish allergies, since it contains sodium iodide [41].

The uptake among tracers differs, as well as identification rate of sentinel-LNs. For example, Tc99m uptakes more rapidly than indocyanine green [41], while carbon nanoparticles that are with very small diameter have even faster uptake [43]. MBD has a fast uptake and high identification rate of sentinel-LNs, ranging from 84% in our previously published study on micro-PTCs [45] to 100% in this study on micro-MTCs. The MB uptake can be interfered by severe thyroiditis or large goiters [45]. The accuracy of SLNB method with MBD and FSA of sentinel-LNs in reported study with micro-PTCs [45] was 97.25%, while in this series of patients it reaches 100%.

Given the availability, low price, feasibility of injection, speed of uptake, identification rate, accuracy and absence of anaphylactic reactions, MBD could be considered as the most appropriate tracer for SLNB in thyroid carcinomas [32, 45-47].

However, this method is not commonly used worldwide. Based on 4 meta-analyses [27, 29-31], it is mainly used for detection of central LNs metastases in differentiated thyroid carcinomas. To our knowledge, none of the SLNB studies published so far has the same surgical technique as here described [32]. Differences between original Dzodic’s SLNB method, used in our center [32], and other published SLNB techniques are following: (1) cancer type: used for papillary and medullary carcinomas, (2) tracer: MBD as a single tracer, (3) neck compartment (lymphatic basin): jugulo-carotid regions, (4) pathohistological analysis: intraoperative, FSA, (5) aim of the method: immediate decision on one-time LLND. Ikeda [48] and Lee [49] also perform staging of lateral LNs; however, they use indocyanine green and Tc99m (respectively), not MBD, they do not perform FSA on sentinel-LNs, and they use it for papillary carcinomas,
not MTC. There are only two publications on SLNB for detection of lateral LN metastases in micro-MTC after Tc99m tumor injection: one case report [50] and one pilot study [51], both published in 2014, showing that this type of LN staging is useful. However, SLNB with MBD can provide reliable results, with fewer costs and technical problems.

Although recommendations for the necessity and the extent of LLND are based on serum CT levels and preoperative US staging of LNs, there is no consensus on the most appropriate CT level-based risk stratification throughout available guidelines [1, 16-19]. A recent publication from 2019 by Norwegian authors [52] showed that basal CT cannot be used to predict the need for prophylactic LLND in patients with MTC due to many inconsistencies with CT level and presence of lateral LN metastases. In their series, lateral LN metastases were found in 16% of patients with CT ≤500 pmol/L and 50% of those with CT 501–1,000 pmol/L. On the other hand, 19% of N0 patients had CT over 500 pmol/L and 17% of N1b patients had CT ≤500 pmol/L. Similarly, out of 18 US-cN0 patients in our study that were also pathohistologically “cleared” for LN metastases (confirmed pN0), 16 of them with basal CT above 20 pg/mL were candidates for elective ipsilateral LLND, based on some recommendations [13], while 8 out of 18 with basal CT above 200 pg/mL were eligible for elective contralateral LLND. Authors strongly suggest that a benefit for patients can be observed if SLNB with MBD is performed for intraoperative LN staging and decision upon necessity for LLND, regardless the status of central LNs. On the other hand, two patients with C634F mutation were identified to harbor metastases in central and lateral compartments, although metastases were not detected by preoperative US investigation and CT values were 42.0 and 221.6 pg/mL.

In our study of 20 cN0 patients with micro-MTCs and CT levels below 1,000 pg/mL, which was expected to be a good prognostic group, we have showed that: (1) 2/20 patients actually had central and lateral LN metastases; (2) additional lateral LN metastases were found in patients who had LLND, meaning positive sentinel-LNs were predictive for metastases in other lateral LNs; and (3) patients that were “cleared” by sentinel-LN FSA to be negative did not have “skip” metastases in non-sentinel-LNs.

The authors find Dzodic’s SLNB method for LN staging useful for intraoperative assessment of lateral LNs in cN0 patients and optimization of initial surgery of micro-MTCs with CT below 1,000 pg/mL. This way, patients with FSA proven sentinel-LN metastases can receive one-time LLND, while those with negative sentinel-LNs benefit from less extensive surgery and potentially decreased complication rate.

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Disclosure

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