A Novel Risk Stratification System for Thyroid Nodules With Indeterminate Cytology—A Pilot Cohort Study

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Background: Thyroid ultrasound (US), fine needle aspiration biopsy (FNAB), and molecular testing have been widely used to stratify the risk of malignancy in thyroid nodules. The goal of this study was to investigate a novel diagnostic approach for cytologically indeterminate thyroid nodules (ITN) based upon a combination of US features and genetic alterations.

Methods: We performed a pilot cohort study of patients with ITN (Bethesda III/IV), who underwent surgical treatment. Based on standardized sonographic patterns established by the American Thyroid Association (ATA), each ITN received an US score (XUS), ranging between 0 and 0.9 according to its risk of thyroid cancer (TC). DNA and RNA were extracted from pathologic material, available for all patients, and subjected to OncomineTM Comprehensive Assay v2 (OCAv2) next-generation sequencing. Each genetic alteration was annotated based on its strength of association with TC and its sum served as the genomic classifier score (XGC). The total risk score (TRS) was the sum of XUS and XGC. ROC curves were generated to assess the diagnostic accuracy of XUS, XGC, and TRS.

Results: The study cohort consisted of 50 patients (39 females and 11 males), aged 47.5 ± 14.8 years. Three patients were excluded due to molecular testing failure. Among the remaining 47 patients, 28 (59.6%) were diagnosed with TC. BRAFV600E was the most common mutation in papillary TC, PAX8-PPARG fusion was present in NIFTP, pathogenic variants of SLX4, ATM, and NRAS were found in Hürthle cell TC and RET mutations in medullary TC. The diagnostic accuracy of XGC and TRS was significantly higher compared with XUS (88 vs. 62.5%, p < 0.001; 85.2 vs. 62.5%, p < 0.001, respectively). However, this increased accuracy was due to significantly better sensitivity (80.7 vs. 34.6%, p < 0.001; 84.6 vs. 34.6%, p < 0.001, respectively) without improved specificity (94.7 vs. 90%, p = 0.55; 85.7 vs. 90%, p = 0.63, respectively).
Molecular testing was not performed or was inconclusive (III and surgical excision for nodules in category IV, in which either surveillance or diagnostic surgery for nodules in category

**INTRODUCTION**

The management of patients with thyroid nodules has evolved with the widespread use of the Bethesda System for Reporting Thyroid Cytopathology (1). Although the approach is standardized, especially for nodules with clearly benign (Bethesda category II) or malignant features (Bethesda category VI), the management of patients with thyroid nodules yielding an indeterminate cytology [Bethesda III—atypia of unknown significance (AUS) or follicular lesion of undetermined significance (FLUS); Bethesda IV—follicular neoplasm (FN) or suspicious for a follicular neoplasm (SFN)], which account for 10–25% of thyroid fine needle aspiration biopsies (FNABs) is still very challenging (1, 2). The current ATA guidelines recommend either surveillance or diagnostic surgery for nodules in category III and surgical excision for nodules in category IV, in which molecular testing was not performed or was inconclusive (3). The standardization of malignancy risk stratification based on ultrasound (US) imaging of the neck has been recently proposed by several organizations. American and European Thyroid Associations have implemented representative pictorial systems based on sonographic patterns of thyroid nodules (3, 4), while the Korean Society of Thyroid Radiology and the American College of Radiologists utilize a scoring system—K-TIRADS and TIRADS (5, 6). These systems are designed to standardize the management strategy and enable easier communication between patients and endocrinologists, radiologists and surgeons (3). Recently established machine learning algorithms, used to characterize the US patterns typical of malignancy, have been also successfully implemented (7–9).

Molecular testing has been evolving as a useful tool in guiding the decision-making regarding surgical vs. conservative management of cytologically indeterminate thyroid nodules (10–13). From single gene mutation testing to next generation sequencing (NGS) multigene panels, as well as gene expression classifiers and microRNA (miRNA) markers, the available diagnostic techniques vary in sensitivity and specificity (14–21).

Despite advances in sonographic systems to classify thyroid nodules according to their risk of malignancy and the added potential utility of molecular diagnosis, overtreatment of thyroid nodules is still commonly observed and associated with substantial side effects and incremental cost (11).

A holistic approach to the patient with an indeterminate thyroid nodule, incorporating clinical, sonographic, cytological, and molecular data is optimal in decision making process. Thus, the goal of this study was to investigate the diagnostic accuracy of a risk stratification system for cytologically indeterminate thyroid nodules based on a combination of US features and genetic abnormalities.

**METHODS**

We performed a single-institution pilot cohort retrospective study including patients with thyroid nodules who underwent thyroid US and FNAB revealing indeterminate cytology (Bethesda III, IV), and were subsequently subjected to surgical treatment. These patients presented in the Thyroid Outpatient Clinic and the decision to perform FNAB was based on the current ATA guidelines (3). However, there were 2 patients for whom nodules <1 cm were biopsied—one due to cervical lymphadenopathy and one due to family history of papillary thyroid cancer in multiple family members. In the presence of indeterminate cytology, patients were offered either surgery or follow-up based on clinical and ultrasonographic criteria during subsequent visits. Only patients with cytologically indeterminate nodules that had surgery were included in this analysis.

We excluded patients characterized by Bethesda I, II, IV, VI cytology categories and subjects who have not been treated with surgery for indeterminate thyroid nodules.

The study was approved by the NIH Intramural Institutional Review Board. Informed consent has been obtained from each patient after full explanation of the purpose and nature of all procedures performed.

**Evaluation of US Patterns**

Based on American Thyroid Association (ATA) sonographic patterns, each nodule received an annotated ultrasound score ($X_{US}$), according to its risk of malignancy with 0 for benign and very low suspicion pattern (<3% cancer risk per the ATA guidelines), 0.1 for low suspicion (5–10% malignancy risk per the ATA guidelines), 0.2 for intermediate suspicion (10–20% malignancy risk per the ATA guidelines), and 0.9 for high suspicion nodules (>70–90% cancer risk per the ATA guidelines) (3). In other words, numbers of $X_{US}$ refer to the percentages of the upper limit of malignancy per ATA guidelines: 0.1 = 10% (low risk); 0.2 = 20% (intermediate risk); 0.9 = 90% (high risk). Two endocrinologists specialized in thyroid disorders (CG-L and JK-G) reviewed independently the entire registered trailers of thyroid and neck US and were blinded to pathology and molecular test results. The discordance between the evaluation of the nodules was addressed by reviewing the nodules again with a third party—radiologist, until consensus was met.
Evaluation of the Molecular Signature of Thyroid Nodules

DNA and RNA were extracted from formalin-fixed paraffin-embedded tumors at surgery. Next generation sequencing was performed using Oncomine™ Comprehensive Assay v2 (OCAv2) on an Ion Torrent S5 XL sequencer. The Oncomine™ Comprehensive Assay v2 (OCAv2) is a CLIA (Clinical Laboratory Improvement Amendments)-validated commercial pan-cancer targeted NGS panel designed to detect somatic single-nucleotide variants (SNV), insertions and deletions (INDEL), copy number variants (CNV), and gene fusions in 143 genes, including major driver genes in thyroid oncogenesis (Supplemental Figure 1). OCAv2 utilizes Ion AmpliSeq™ chemistry, allowing for DNA and RNA inputs as low as 10 ng of extracted material from formalin-fixed paraffin-embedded (FFPE) tumor samples. The variant calling, annotation, and classification were performed on Ion Reporter software v5.10 (Thermo Fisher). In addition, a droplet digital PCR was performed to assess TERT promoter mutation. TERT promoter c.1-124C>T and c.1-146C>T mutational analysis were performed using the expert design PrimePCR ddPCR TERT C228T_113 Assay and TERT C250T_113 Assay (BIO-RAD, Hercules, CA) on a BIO-RAD QX200 droplet digital PCR (ddPCR) system. The assays were performed in duplicate. The presence of mutation and the fractional abundance of the mutant allele were determined using QuantaSoft v.1.7 (BIO-RAD).

The dataset of ultrasound/FNAB was cross-checked with the dataset of tissue molecular analysis to ensure the identity between biopsied and genotyped nodules. Each nodule received an annotated genomic classifier score (X_GC) based on the probability of a given genomic abnormality being associated with cancer per large molecular databases TCGA and COSMIC v87. Each genomic alteration received a score ranging from 0 to 1, with 0—for no association with cancer and 1—for 100% association with cancer. Genetic abnormalities observed in both benign and malignant lesions received scores between 0 and 1 based on a ratio of their prevalence in malignant and benign lesions. The final genomic classifier (X_GC) score was a sum of all genomic abnormalities given by the formula:

\[
X_{GC} = (X_{SNV/INDEL})n + X_{GF} + X_{CNV}
\]

where X is the annotated score; SNV, single nucleotide variant; INDEL, insertions/deletions; n, number of SNVs/INDELS; GF, gene fusions; CNV, copy number variants (20).

Risk Stratification Based on Imaging and Molecular Characteristics

A third score, called total risk score (TRS), was calculated as the sum of the ultrasound score (X_US) and the genomic classifier score (X_GC):

\[
TRS = X_{US} + X_{GC}
\]

STATISTICAL ANALYSIS

Receiver operating characteristic (ROC) curves were generated to assess the diagnostic accuracy of X_US, X_GC, and TRS. First, in each scoring system, area under the curve (AUC) values were obtained with different cutoff values for dichotomization between benign and malignant nodules and the best cutoff value with the highest AUC was selected. Second, we compared the three best scoring systems by using AUC, sensitivity, specificity, positive predictive value (PPV) and negative predictive value (NPV) with 95% Wilson confidence intervals (CI). Given that the cancer prevalence may vary in different populations with cytologically indeterminate thyroid nodules, the established sensitivity and specificity were used to calculate PPV and NPV corresponding to the whole spectrum of cancer prevalence utilizing Bayes theorem using R software. All other analyses were based on two-tailed tests using α = 0.05 and conducted using SAS version 9.4 (SAS Institute, Cary, NC, USA).

RESULTS

Among 165 patients with cytologically indeterminate thyroid nodules being currently followed by the National Institutes of Health endocrine service, 96 patients had surgery performed prior to December 2017. We present the results of the first 50 patients (39 females and 11 males) with 87 nodules, for whom a comprehensive molecular analysis of thyroid nodules was performed. Three patients were excluded from further analysis due to insufficient quality of the RNA extracted from the pathology material (Figure 1). The mean age at the time of diagnosis was 47.5 ± 14.8 years (Table 1). The analysis of US patterns of thyroid nodules in the cohort revealed an interobserver agreement rate of 94%. The annotated ultrasound scores (X_US) were 0.9 in 15 nodules, 0.2 in 20 nodules and 0.1 in 15 nodules. Twenty-eight patients (59.6%) were diagnosed with thyroid carcinoma (TC): 15 papillary TC (PTC) (including 4 micro-PTCs identified within biopsied nodule), 6 follicular variant of PTC (PTCFV), 1 non-invasive follicular thyroid neoplasm with papillary-like nuclear features (NIFTP), 2 Hürthle cell TC (HTC), 1 poorly differentiated TC (PDTC), and 3 medullary TC (MTC) (Table 2). Incidentally discovered
TABLE 1 | Baseline characteristics of the study cohort.

| Clinical characteristics                  |       |       |
|------------------------------------------|-------|-------|
| Average ± SD age at diagnosis (years)   | 47.5 ± 14.8 |
| Gender (% female)                        | 37/47 (78.7%) |
| Average ± SD tumor size (cm)             | 2.3 ± 1.7 |
| Surgical procedure                       |       |       |
| Total thyroidectomy                      | 26/47 (55.3%) |
| Lobectomy                                | 21/47 (44.7%) |

TABLE 2 | Histology of 47 cytologically indeterminate thyroid nodules.

|                  | n   | %     |
|------------------|-----|-------|
| Carcinoma        | 28  | 59.6  |
| PTC              | 21  | 75.0  |
| Classic          | 11  |       |
| PTCFV            | 6   |       |
| Microcarcinoma   | 4   |       |
| NIFTP            | 1   | 3.6   |
| HTC              | 2   | 7.1   |
| PDTC             | 1   | 3.6   |
| MTC              | 3   | 10.7  |
| Benign           | 19  | 40.4  |
| Follicular adenoma | 6 | 31.6 |
| Adenomatoid nodule | 13 | 68.4 |

PTC, papillary thyroid carcinoma; PTCFV, papillary thyroid carcinoma follicular variant; NIFTP, non-invasive follicular thyroid neoplasm with papillary-like nuclear features; HTC, Hurthle cell thyroid carcinoma; PDTC, poorly differentiated thyroid carcinoma; MTC, medullary thyroid carcinoma.

Optimal Cutoff Value to Discriminate Between Benign and Malignant Lesions Based on the Genomic Classifier Score (XGC)

The XGC cut off value of 0.6 was characterized by the highest diagnostic accuracy of 88% (Supplemental Figure 2B). The sensitivity of XGC of 0.6 was 80.7%, specificity 94.7%, PPV 92.3% (CI 78.2–99.2%), NPV 80% (CI 60.1–91.1%). Based on Bayes' theorem prediction, the NPV and PPV for XGC may range between 83–99% and 43–93%, respectively in different populations (Figure 3).

The diagnostic accuracy of XGC of 0.6 was significantly higher compared with XUS of 0.9 (88 vs. 62.5%, p < 0.001; Figure 4). However, an increased accuracy was due to significantly better sensitivity (80.7 vs. 34.6%, p < 0.001) without further improvement in specificity (94.7 vs. 90%, p = 0.55).

Optimal Cutoff Value to Discriminate Between Benign and Malignant Lesions Based on Total Risk Score (TRS)

The TRS based cutoff of 0.7 was associated with the highest diagnostic accuracy of 85.2% (Supplemental Figure 2C). The sensitivity of TRS 0.7 was 84.6%, specificity 85.7%, PPV 88% (CI 70–95.8%), NPV 81.8% (CI 70.5–92.7%). The NPV and PPV in populations characterized by different cancer prevalence is predicted to range between 82–99% and 27–88%, respectively (Figure 3).

The diagnostic accuracy of TRS of 0.7 was significantly higher compared with the XUS of 0.9 (85.2 vs. 62.5%, p < 0.001; Figure 4). However, the increased accuracy was due to significantly better sensitivity (84.6 vs. 34.6%, p < 0.001) without further improvement in specificity (85.7 vs. 90%, p = 0.63). There was no difference in diagnostic accuracy between TRS and XGC (85.2 vs. 88%, p = 0.46; Figure 4). In addition, in a sub-analysis of our cohort excluding patients with MTC, the sensitivity of XGC was 78.2%, specificity of 95.2% while the sensitivity of TRS was 82.6% and specificity of 85.7% (Supplemental Figures 4–6).

Given a high and non-inferior to molecular testing specificity of US alone for US high-risk nodules, we next performed a subgroup analysis testing the diagnostic accuracy of OCAv2 in group A—US high-risk nodules and group B—US low-to-intermediate risk nodules.

Malignancy Risk Stratification System

Optimal Cut-Off Value Discriminating Between Benign and Malignant Lesions Based on the Ultrasound Score (XUS)

There were no significant differences in diagnostic accuracy determined by AUC between different cutoff points for XUS (Supplemental Figure 2A). However, an XUS cutoff value of 0.9 was selected due to much higher specificity compared to a cutoff value of 0.2 (90 vs. 52.4% p = 0.006). The sensitivity of the US cutoff value of 0.9 was 34.6%, specificity 90%, PPV 81.8% (CI 52.3–97.9%), NPV 52.8% (CI 37–68%). Given a high prevalence of thyroid cancer in our cohort, suggesting a referral and selection bias, Bayes' theorem was used to predict NPV and PPV in populations with different cancer prevalence ranging from 6 to 59% (Figure 2). The NPV and PPV for XUS ranged between 52.8–95% and 18–81.8%, respectively.

Diagnostic Algorithm for Management of Thyroid Nodules

Patients with XUS of 0.9 were characterized by a 5x higher likelihood of cancer than the patients with XUS < 0.9 (OR 5.03, 95%CI 0.95–26.6). The subgroup analysis of 11 patients...
with a $X_{US}$ of 0.9 revealed that implementing $X_{GC}$ or TRS does not lead to improved diagnostic accuracy, as $X_{US}$ alone led to a comparable separation of benign vs. malignant lesions (Supplemental Figure 3A). In contrast, molecular tests significantly improved diagnostic accuracy in patients with $X_{US} < 0.9$ (Supplemental Figure 3B). Diagnostic accuracy of $X_{US}$ of 0.2 was 61%—significantly lower than 82.7% of $X_{GC}$ ($p = 0.04$) and 85.6% of TRS ($p = 0.01$; Supplemental Figure 3B). $X_{GC}$ and TRS significantly improved specificity of cancer diagnosis in this group ($X_{GC}$ vs. $X_{US}$ 94.7 vs. 57.9%, $p = 0.007$ and TRS 94.7 vs. 57.9%, $p = 0.007$), without a significant improvement in sensitivity ($X_{GC}$ vs. $X_{US}$ 70.6 vs. 64.7%, $p = 0.71$ and TRS vs. $X_{US}$ 76.5 vs. 64.7, $p = 0.45$). There was no significant difference in diagnostic accuracy between $X_{GC}$ and TRS (82.7 vs. 85.6%, $p = 0.32$). However, if the analysis had been performed prospectively, TRS < 0.7 could have led to missing diagnosis of cancer in 4 very low risk tumors—two mutation negative FVPTC T1bN0M0, both with $X_{US}$ of 0.2 consistent with intermediate features.
per the US characteristics, one mutation negative classic PTC T1bN0M0 characterized by $X_{US}$ of 0.1 consistent with low risk US features, and one mutation negative micro-PTC T1aN0M0, with $X_{US}$ of 0.1, multifocal, with the maximum diameter of the largest lesion of 0.8 cm—while $X_{GC} < 0.6$ could have led to missing cancer diagnosis not only in the above mentioned 4 very low risk tumors, but also in one PDTC with HRASQ61R mutation, characterized by intermediate risk US features with $X_{US}$ of 0.2. Altogether, this suggests that TRS may have a better clinical performance.

Based on this analysis, we propose the following algorithm for the management of thyroid nodules with indeterminate cytology (Figure 5).

1) $X_{US}$ high risk = 0.9—no added benefit in specificity of molecular testing—consider referral for surgical treatment
2) $X_{US} < 0.9$ (low to intermediate risk nodules)—diagnostic accuracy improves with molecular testing—consider surgery if $TRS \geq 0.7$; if $TRS < 0.7$, a watchful waiting strategy with observation of tumor behavior over time might be a reasonable option as all cancers that could have been missed in this strategy were very low risk micro-PTCs.

**DISCUSSION**

We propose a novel diagnostic algorithm for patients with Bethesda III/IV cytology diagnosis, based on a combination of US patterns and the molecular signature of thyroid nodules. We show that OCAv2 molecular profiling is not associated with a significant improvement in specificity of cancer diagnosis in cytologically indeterminate thyroid nodules characterized by high-risk US features. However, OCAv2 molecular profiling is useful in improving diagnostic accuracy of cytologically indeterminate thyroid nodules characterized by low-to-intermediate sonographic features. Therefore, we propose a sequential approach for patients with AUS/FLUS/FN/SFN cytology diagnosis that consists of:

1) Reviewing patients’ US images and considering surgical treatment for those with ATA high-risk US features nodules, without further molecular profiling, as the US alone-based specificity of cancer diagnosis in these nodules is non-inferior to molecular profiling;
2) performing molecular profiling for US low-to-intermediate risk nodules and considering surgical treatment for patients characterized by molecular score TRS of ≥0.7 and watchful waiting strategy with an observation of tumor growth over time as a reasonable option for molecular score TRS < 0.7.

This approach should be associated with an improved cost to benefit ratio, as routine implementation of molecular diagnostics is expensive. Costs of molecular testing vary according to insurance coverage but reported costs for the most comprehensive tests range between $3,000–$4,800 per nodule (12, 13). That being said, some centers may utilize high-risk molecular signature (e.g., concomitant BRAFV600E and TERT promoter mutation, EIF1AX mutations and THADA fusions) to guide the extend of surgery, in which case analysis of molecular signature of all nodules is warranted.

Several studies have suggested that sonographic patterns can effectively stratify the prevalence of malignancy in indeterminate thyroid nodules (22–25). A retrospective study of 173 indeterminate thyroid nodules reported a cancer rate of 75% in high suspicion nodules compared to 16.8% in very low, low and intermediate suspicion nodules, per ATA US patterns (24). Another retrospective study, also using ATA US patterns, including 463 indeterminate thyroid nodules found 5 times higher likelihood of malignancy in high risk patterns compared with low-to-intermediate risk patterns (25).

However, the appropriate risk stratification of thyroid nodules based on US patterns depends on the experience of the ultrasonographer and the characteristics of the equipment. Choi et al. compared interobserver and intraobserver variations in ultrasound assessment of thyroid nodules and concluded that the final assessments of experienced radiologists were highly accurate (26). These results were corroborated by another study that demonstrated that trainees receiving one-on-one instruction from experienced radiologists improved their diagnostic performance for evaluating thyroid nodules with ultrasonography (27).

The use of molecular markers for cytologically indeterminate thyroid nodules has been evolving over the past two decades (11–13). We used the Oncomine® Comprehensive Assay v2 (OCAv2), a pan-cancer 143-gene panel focused on potentially actionable oncogenes relevant in precision medicine. We created a scoring system based on the strength of association with cancer ranging from 0 (no association with cancer) to 1 (100% association with cancer). Similarly, ThyroSeq v3, a 112-gene panel, uses a genomic classifier score in which each detected association with cancer). Similarly, ThyroSeq v3, a 112-gene panel, uses a genomic classifier score in which each detected association with cancer). Similarly, ThyroSeq v3, a 112-gene panel, uses a genomic classifier score in which each detected association with cancer). Similarly, ThyroSeq v3, a 112-gene panel, uses a genomic classifier score in which each detected association with cancer. Similarly, ThyroSeq v3, a 112-gene panel, uses a genomic classifier score in which each detected association with cancer. Similarly, ThyroSeq v3, a 112-gene panel, uses a genomic classifier score in which each detected association with cancer. Similarly, ThyroSeq v3, a 112-gene panel, uses a genomic classifier score in which each detected association with cancer. Similarly, ThyroSeq v3, a 112-gene panel, uses a genomic classifier score in which each detected association with cancer. Therefore, while with proposed by us approach there might be a higher risk of missing thyroid cancer amongst indeterminate thyroid nodules due to lower sensitivity compared with Thyroseq v3, its high specificity may help with avoiding unnecessary surgeries.

Another widely utilized diagnostic approach is the Afirma assay by Veracyte, Inc. It is based on the use of messenger-RNA (mRNA) expression and a proprietary machine learning algorithm to classify the risk of malignancy of a given nodule into benign or suspicious. The Afirma Gene Expression Classifier (GEC) was validated in a large cohort of patients in 2012 as a relatively good rule-out test with high sensitivity and a NPV from 75 to 100% (14). However, post-validation studies have shown that Afirma GEC did not perform as expected on Hurthle cell-rich lesions and had a lower than anticipated malignancy rate within GEC-suspicious nodules (28–30). An updated version of Afirma called Genomic Sequencing Classifier (GSC) was validated in 2018 with a reported sensitivity of 91% and a specificity of 68% (31). An independent comparison between Afirma GEC and GSC showed that the latter version has a higher benign call rate compared to the former, predominantly for Hurthle cell cytology (32). These findings were recently corroborated by another study, from a single academic tertiary center, that also reported an improvement in specificity and PPV of Afirma GSC, while maintaining high sensitivity and NPV (33). Compared with Afirma, OCAv2 is characterized by lower sensitivity of 84.6%, but higher specificity of 85.7%.

ThyGeNEXT/ThyraMIR, previously known as ThyGenX/ThyraMIR, is a next generation sequencing test for mutations (10 genes) and fusions (6 genes) implicated in thyroid tumorigenesis complemented with expression analysis of 10 microRNAs (miRNA). It uses a proprietary algorithm to classify each nodule as having a high risk or low risk miRNA profile (12). This combined algorithm achieved a sensitivity of 89% and a specificity of 85% among cytologically indeterminate nodules in a cross-sectional study (18). No post-validation studies have been reported. Compared with ThyGeNEXT/ThyraMIR, OCAv2 is characterized by similar sensitivity and specificity. While sensitivity and specificity of a diagnostic test depend on test performance, negative predictive value (NPV) and positive predictive value (PPV) depend on the prevalence of disease in the population. Our study was most likely associated with a referral and selection bias, as the prevalence of cancer in our sample of 50 patients was as high as 59%. Analysis of all 96 patients who underwent surgery revealed cancer prevalence of 45.8% (44/96) and assuming that non-operated patients were characterized by benign disease, cancer prevalence would have been 26.6% (44/165) (Figure 1). The analysis of two recent reports testing performance of Afirma GSC amongst patients with indeterminate thyroid nodules, who underwent surgery, revealed a similar cancer prevalence of 50–55% (32, 33). The TRS-based NPV of 82% is significantly lower than reported in these studies NPV of 100%, while TRS-based PPV of 88% is significantly better than reported in these studies PPV of 50–60% (32, 33). Compared with Afirma GSC, in populations with cancer prevalence 50–59%, TRS may perform better in avoiding surgery for benign nodules but might be associated with higher risk of missing cancer. That being said, implementing our approach
performed a Bayes’ theorem-based simulation, documenting that NPV and PPV of TRS in populations with cancer prevalence of 6–59% would range from 82 to 99% and 27–88%, respectively. In particular, comparing with Thyroseq v3 tested on a population with 28% cancer prevalence (21), TRS may perform similarly—as Thyroseq v3 with NPV of 97% performs slightly better than predicted TRS-based NPV of 93.4% while Thyroseq v3 PPV of 66% is slightly worse than predicted TRS-based PPV of 69.7% (Figure 3). The performance of ThyGeNEXT/ThyraMir in a population with cancer prevalence of 32% is very similar to predicted TRS performance—NPV of 94 vs. 93% and PPV of 74 vs. 72%, respectively [Figure 3; (18)]. Obviously, only head to head comparisons in well-designed non-inferiority trials would enable reaching any conclusions about the accuracy of the above-mentioned tests, as the comparison performed above is based on simulation rather than actual hard data.

The strengths of our study rely on broad clinical data and pathologic diagnosis available for all patients. All ultrasound studies were performed with the same equipment and reported by board-certified radiologists and independently reviewed by two endocrinologists, blinded to histological diagnosis. All cytological and histological diagnosis were made by board-certified and experienced pathologists. Moreover, Oncomine is a widely available assay used by many molecular diagnostics laboratories. In addition, the algorithm proposed, utilizing a combination of US features and molecular diagnostics in malignancy risk stratification of thyroid nodules, might be applicable to any other molecular test available worldwide in different institutions.

We do acknowledge a significant referral and selection bias in our cohort as a potential limitation of this study. Some patients were referred to our institution with the intent of having surgery. Moreover, as a retrospective study, we could not control for factors that prompted certain patients for surgery instead of conservative management. Yet, we do provide a simulation of the performance of XUS, XGC, and TRS in cohorts with different cancer prevalence according to the Bayes theorem, with ranges of NPV and PPV for each score (Figure 3).

Moreover, this pilot study is limited by the small sample size and reduced number of Hürthle cell carcinomas, NIFTP, and follicular architecture tumors. The nomenclature revision of encapsulated follicular variant of PTC (EFVPTC) to non-invasive follicular thyroid neoplasm with papillary-like nuclear features (NIFTP) (34) may represent the acceptance of borderline/precursor lesions in the thyroid (35); additional information is warranted about the molecular signature of these tumors. The relative high prevalence of classic PTC in our cohort (11/21) is different from the literature (35–37) and may be responsible for the enrichment in BRAF-like mutations in our cohort as compared to RAS-like mutations, thus increasing the specificity of our diagnostic approach. It will be important to test the performance of our algorithm in cohorts characterized by higher prevalence of RAS-like tumors. We are currently conducting a prospective study to obtain validation of this pilot study in an independent cohort with analysis of the assay performance on cytologic specimens, using a new version of Oncomine (OCAv3).

**CONCLUSION**

We propose a diagnostic algorithm utilizing a combination of US features and next generation sequencing that appears to provide a cost-effective diagnostic tool to guide the management strategy of indeterminate thyroid nodules. Our data suggest that molecular testing could be avoided in US high-risk nodules diagnosed in centers with experienced endocrinologists/radiologists evaluating US images, as the specificity of cancer diagnosis in such scenarios is non-inferior to molecular testing. Molecular testing might be beneficial in low-to-intermediate risk sonographic patterns of thyroid nodules as evaluation of genetic landscape in such lesions increases the specificity of cancer diagnosis, and as such, may lead to the avoidance of unnecessary surgeries in these patients.

**DATA AVAILABILITY STATEMENT**

The datasets generated for this study can be found in the NCBI BioProject repository, in the following link: https://www.ncbi.nlm.nih.gov/bioproject/PRJNA600873/.

**ETHICS STATEMENT**

The studies involving human participants were reviewed and approved by NIH Intramural Institutional Review Board. The patients/participants provided their written informed consent to participate in this study.

**AUTHOR CONTRIBUTIONS**

J-KG: conception and design of the study, revision of the final version of the manuscript. CG-L: organization of the database and data management. J-LM: patients recruitment and patients care coordination. AF, MR, SP, and LX: review of pathology material and performance of the molecular testing. LW and KB: revision of the final version of the manuscript. CG-L: organization of the database and data management. J-LM: patients recruitment and patients care coordination. AF, MR, SP, and LX: review of pathology material and performance of the molecular testing. LW and KB: revision of the final version of the manuscript. All authors contributed to manuscript revision, read, and approved the submitted version.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fendo.2020.00053/full#supplementary-material

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**Conflict of Interest:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.