Molecular testing for miRNA, mRNA and DNA on fine needle aspiration improves the preoperative diagnosis of thyroid nodules with indeterminate cytology

Emmanuel Labourier1*, Alexander Shifrin2, Anne E. Busseniers3, Mark A. Lupo4, Monique L. Manganelli5, Bernard Andruss1, Dennis Wylie1, and Sylvie Beaudenon-Huibregtse1

1Asuragen Inc., Austin, TX; 2Jersey Shore University Medical Center, Center for Thyroid, Parathyroid and Adrenal Diseases, Neptune, NJ; 3Metropolitan Fine Needle Aspiration Service, WA, DC, and Bethesda, MD; 4Thyroid & Endocrine Center of Florida, Sarasota, FL; 5San Diego, CA

Context: Molecular testing for oncogenic mutations or gene expression in fine-needle aspirations (FNAs) from thyroid nodules with indeterminate cytology identifies a subset of benign or malignant lesions with high predictive value.

Objective: Evaluate a novel diagnostic algorithm combining mutation detection and miRNA expression to improve the diagnostic yield of molecular cytology.

Setting: Surgical specimens and preoperative FNAs (n=638) were tested for 17 validated gene alterations using the miRInform Thyroid test and with a 10-miRNA gene expression classifier generating positive (malignant) or negative (benign) results.

Design: Cross-sectional sampling of thyroid nodules with AUS/FLUS or FN/SFN cytology (n=109) was conducted at 12 endocrinology centers across the United States. Qualitative molecular results were compared to surgical histopathology to determine diagnostic performance and model clinical impact.

Results: Mutations were detected in 69% of nodules with malignant outcome. Among mutation-negative specimens, miRNA testing correctly identified 64% of malignant cases and 98% of benign cases. The diagnostic sensitivity and specificity of the combined algorithm was 89% (95% confidence intervals (CI): 73–97%) and 85% (95% CI: 75–92%), respectively. At 32% cancer prevalence, 61% of the molecular results were benign with a negative predictive value of 94% (95% CI: 85–98%). Independently of variations in cancer prevalence, the test increased the yield of true benign results by 65% relative to mRNA-based gene expression classification and decreased the rate of avoidable diagnostic surgeries by 69%.

Conclusions: Multi-platform testing for DNA, mRNA and miRNA can accurately classify benign and malignant thyroid nodules, increase the diagnostic yield of molecular cytology, and further improve the preoperative risk-based management of benign nodules with AUS/FLUS or FN/SFN cytology.

Cytology on ultrasound-guided fine needle aspiration (FNA) biopsies has dramatically improved the clinical management of patients with solid thyroid nodules > 1 cm. In routine clinical practice, this procedure can identify approximately 50% of malignant nodules (50% sensitivity) and 70% of benign nodules (70% specificity)

Abbreviations:
without the need to perform a diagnostic surgery (1–4). Because cytology has both high positive predictive value (PPV) (PPV > 98%) and high negative predictive value (NPV) (NPV > 95%), it allows accurate, preoperative, risk-based classification of thyroid nodules. However, the relatively low diagnostic yield of cytology also results in a large fraction of nodule aspirates, up to 35%, without a definitive benign or malignant diagnosis (Bethesda categories II or VI) (1–4). In addition, the residual risk of thyroid cancer in nodules with indeterminate cytology varies significantly across institutions, in particular for nodules with a preoperative diagnosis of atypia of undetermined significance/follicular lesion of undetermined significance (AUS/FLUS, Bethesda category III) or follicular neoplasm/suspicious for a follicular neoplasm (FN/SFN, Bethesda category IV) (1, 4, 5).

To overcome these limitations, novel diagnostic procedures have been developed and optimized to generate clinically actionable information in FNAs with indeterminate cytology. Qualitative molecular tests for specific gene mutations or RNA fusion transcripts associated with thyroid carcinogenesis can detect malignant nodules with high PPV, from 70 to 100% for different markers (6–11). Alternatively, a microarray-based test combining 7 distinct classifiers and interrogating the expression levels of 167 genes can detect benign nodules with a high NPV of 94% (12). The broad availability of these methods has undoubtedly increased the diagnostic yield of conventional cytopathology and improved the personalized, risk-based management of thyroid cancer patients (13, 14). Yet, they are not without their own limitations. For example, optimized mutation panels can only identify approximately 65% of malignant nodules (65% sensitivity) and the absence of a known oncogenic gene alteration cannot be used to rule out cancer (7, 9). The gene expression classifier only identifies approximately 50% of benign nodules (50% specificity) and most the nodules classified as “suspicious” by the test will go on to be diagnosed as benign (12). Thus, a large number of potentially avoidable diagnostic surgeries are still performed on patients with benign nodules and AUS/FLUS or FN/SFN cytology. Furthermore, when a malignant nodule is diagnosed by surgical histopathology following hemithyroidectomy, a second surgery associated with a higher risk of complication is often required to complete the thyroidectomy (2, 3).

Advances in genome-scale technologies have recently accelerated the discovery of novel biomarker candidates. miRNAs are small, highly conserved RNA molecules that are involved in the pathology of thyroid cancer by regulating key cellular processes such as cell-cycle progression or cell differentiation, proliferation and survival (15, 16). The differential expression of miRNAs in distinct pathological tumor types and at various stages of tumor differentiation or progression has been reported in over 100 publications (17–27). There is currently no validated miRNA test available; however, information derived from miRNA analyses may complement existing molecular diagnostic procedures. Several studies have reported the potential diagnostic utility of miRNAs in preoperative thyroid nodules FNAs (18, 22, 23, 26, 27). Others have shown that miRNA expression changes in papillary thyroid carcinoma are not necessarily correlated with the presence of oncogenic gene mutations (17, 24, 25). In a recent study, we further showed that miRNA expression levels in preoperative FNAs can identify a subset of malignant nodules that are negative for a panel of well-known oncogenic gene alterations (9). In the present work, we took advantage of this characteristic to design a miRNA gene expression classifier that efficiently complements molecular testing for DNA mutations and RNA fusion transcripts. The multianalyte combination test was evaluated in a cohort of thyroid nodules with AUS/FLUS or FN/SFN cytology to establish its diagnostic sensitivity and specificity in preoperative FNAs and model its potential clinical impact on surgery rates. We put forth that this novel strategy can further increase the diagnostic yield of molecular cytology and significantly improve the preoperative risk-based diagnosis of benign thyroid nodules with indeterminate cytology.

Materials and Methods

Algorithm Development

miRNA classifiers were developed using miRNA expression data determined by reverse transcription-quantitative PCR (RT-qPCR), diagonal linear discriminant analysis as training classification models (28), miRNA candidate selection based on effect size, and built-in model normalization enforced by constraints on model coefficients. The case-control training set consisted of 240 surgical specimens collected under a research protocol approved by the University of Michigan institutional review board (IRB). Two independent case-control specimen sets were also used for algorithm optimization: 54 resected tissues acquired from Asterand (Detroit, MI) and 235 remnant nucleic acids samples from preoperative FNAs archived by Asuragen’s CLIA laboratory for research purpose.

Cohort Study

We studied consecutive thyroid nodule FNAs submitted to Asuragen’s CLIA laboratory for evaluation with the miRNAInform Thyroid Test (9, 29) by endocrinology centers across the United States between January 2011 and October 2013 and for whom nucleic acids isolated from the FNAs were available for molecular testing. The study was noninterventional, samples and clinical information were deidentified, and no protected health information or other information identifying study subjects was collected. The protocol was approved by a central investigational
review board, the requirement for informed consent waived according to 45 CFR §46.116(d), and all participating sites signed the study agreement. At the close of the study on January 31, 2014, 282 AUS/FLUS or FN/SFN cytology reports had been received from 20 physicians at 14 sites. Among those 282 aspirations, 113 nodules (40%) had a traceable surgical pathology outcome with a documented histological diagnosis of benign or malignant primary thyroid lesion. Four cases with known history of cancer or previous radioiodine therapy were excluded before performance assessment, resulting in a final set of 109 specimens from 16 physicians at 12 sites for statistical analysis. All specimens and study subjects were distinct from the 529 cases used for development activities.

Molecular Analyses

Surgically resected thyroid lesions and thyroid nodule FNAs were collected, processed and tested for the presence of genetic alterations in the BRAF, RAS, RET or PAX8 genes as previously described (9, 29). miRNA expression in residual total nucleic acids (TNA) samples was measured by RT-qPCR using the miRCURY LNA™ Universal RT microRNA PCR system (Exiqon, Vedbaek, Denmark) and custom-designed Pick-´n-Mix microRNA PCR Panels (Exiqon). Samples were tested in 384-well plates containing primer sets specific for each target miRNA and for a 22-mer, synthetic RNA spiked in every TNA sample as an internal control to assess the quality of the TNA preparation, RT step and qPCR step. During the cohort study, all runs consisted of distinct molecular assays interrogating DNA, RNA or miRNA markers in total nucleic acids samples. The miRNA gene expression classifier reported a qualitative positive or negative result based on the expression levels of 10 miRNA genes, miR-29b-1–5p, miR-31–5p, miR-138–1–3p, miR-139–5p, miR-146b-5p, miR-155, miR-204–5p, miR-222–3p, miR-375, and miR-551b-3p. The other qualitative tests detected the presence of known oncogenic gene alterations, either DNA mutations in the BRAF, HRAS, KRAS and NRAS genes, or the PAX8-PPARG, RET-PTC1 and RET-PTC3 fusion transcripts. Samples positive with either assay were scored as positive and samples negative with all assays were scored as negative. The miRNA classifier was initially trained using well-characterized, surgically-resected, benign or malignant thyroid lesions (n = 240) with the classification thresholds then optimized based on performance evaluated in the original training set alongside independent resected thyroid tissues (n = 54) and preoperative thyroid FNAs (n = 235). Specimens’ characteristics and estimates of MPT performance are summarized in Table 1 and Supplemental Figure 1.

Results

Test Algorithm

The multiplatform mutation and miRNA test (MPT) consisted of distinct molecular assays interrogating DNA, RNA or miRNA markers in total nucleic acids samples. The miRNA gene expression classifier reported a qualitative positive or negative result based on the expression levels of 10 miRNA genes, miR-29b-1–5p, miR-31–5p, miR-138–1–3p, miR-139–5p, miR-146b-5p, miR-155, miR-204–5p, miR-222–3p, miR-375, and miR-551b-3p. The other qualitative tests detected the presence of known oncogenic gene alterations, either DNA mutations in the BRAF, HRAS, KRAS and NRAS genes, or the PAX8-PPARG, RET-PTC1 and RET-PTC3 fusion transcripts.

Data Analyses

Diagnostic sensitivity and specificity were determined using standard 2 by 2 contingency tables comparing qualitative, binary molecular test results (positive or negative) relative to the reference standard diagnoses determined by pathology (benign or malignant). Post-test probability metrics for prevalence values ranging from 0 to 100% were calculated using sensitivity, specificity and Bayes theorem for PPV and NPV. The numbers of true positive, true negative, false positive and false negative molecular results for a given total number of specimens were computed using sensitivity, specificity and prevalence. All calculations assumed conservation of intrinsic test performance in distinct test populations. Unless indicated, 95% confidence intervals (CI) were calculated using the Clopper-Pearson exact method for proportions and p values were calculated using the Fisher exact test for categorical variables. Graphic and statistical analyses were performed in Excel (Microsoft Corp, Redmond, WA) or R version 3.1 (http://www.r-project.org/).

Additional information is available in the Supplemental Materials and Methods.

Cross-Sectional Cohort

To sample a representative cohort of thyroid nodules with indeterminate cytology, consecutive FNAs with known mutation status collected from 12 distinct clinical sites in the United States were tested for miRNA expression. The set consisted of 109 nodules with AUS/FLUS or FN/SFN cytology and a traceable surgical outcome of pri-

| Table 1. Specimens’ characteristics and point estimates of sensitivity and specificity for the multi-platform mutation and miRNA test in 3 independent case-control specimen sets |
|---------------------------------|-----------------|-----------------|
| **Number of cases** | **Training set** | **Fine-needle aspiration** | **Resected tissue** |
| **Female/Male (%)** | 75/25 | 72/28 | 77/23 |
| **Malignant/Benign (%)** | 55/45 | 40/60 | 59/41 |
| **Sensitivity/Specificity (%)** | 84/84 | 84/90 | 84/100 |

The Endocrine Society. Downloaded from press.endocrine.org by [individualUser.displayName] on 04 June 2015 at 13:43. For personal use only. No other uses without permission. All rights reserved.
molecular results in 109 preoperative aspirates. The histopathological reference standard diagnoses were based solely on local pathology expertise and pathologists at each participating site who evaluated the surgically removed thyroid nodules were not aware of the results of molecular testing. There were a total of 74 nodules classified as benign, including 17 Hurthle cell adenomas and 12 follicular adenomas. The 35 malignant nodules consisted of 18 follicular variant of papillary thyroid carcinomas, 10 papillary carcinomas, 5 follicular carcinomas, and 2 Hurthle cell carcinomas (32% thyroid cancer prevalence). All specimens generated valid miRNA gene expression classifier results.

Distribution of Molecular Results by Surgical Outcome

Among the 35 nodules with a confirmed malignant outcome, 24 were positive by mutation testing and 20 were positive by miRNA classifier (Table 2). Seven (7) out of the 11 mutation-negative cases were positive by miRNA classifier resulting in an overall cancer detection rate of 89% (31/35) for the MPT. The 4 malignant cases negative by mutation testing and by miRNA gene expression classifier were 3 follicular variants of papillary thyroid carcinoma, 0.8 to 3.0 cm in size, and a 3.0 cm follicular thyroid carcinoma (Supplemental Table 1). Among the 74 nodules with confirmed benign outcome, 10 were positive by mutation testing and 6 were positive by miRNA classifier. Five (5) out of the 6 miRNA-positive cases were also positive for a RAS mutation resulting in an overall benign detection rate of 85% (63/74) for the MPT. The single benign nodule negative by mutation testing but positive by miRNA gene expression was from a study subject who had undergone total thyroidectomy and had a 0.9 cm follicular variant of papillary thyroid carcinoma in the opposite thyroid lobe (Supplemental Table 1).

Diagnostic Performance

The performance characteristics of the MPT and corresponding 95% CI obtained in the cross-sectional cohort study are summarized in Table 3. The MPT accurately classified both the malignant nodules and the benign nodules. Among the cases with a positive MPT result, the post-test probability of a malignant nodule (PPV) was 74% (31/42) and among the MPT-negative cases, the post-test probability of a benign nodule (NPV) was 94% (63/67). The overall odds of a correct molecular result were 44 times higher than the odds of an incorrect result (P < .01). Importantly, the rate of negative calls generated by the MPT was high (67/109 or 61% of all nodules evaluated), ie, the MPT identified 67 nodules with a low residual risk of thyroid cancer (4/67 or 6%). In the clinical setting, those nodules would be candidates for active surveillance without surgery. At 32% prevalence, the MPT had a true negative call rate of 58% (63/109) and would efficiently reduce the number of surgical procedures from 74 diagnostic surgeries in the absence of molecular testing (68% of all nodules evaluated) to 11 surgeries after reclassification by the MPT (10% of all nodules evaluated).

Clinical Relevance at Different Prevalence

The predictive value of any diagnostic procedure, whether based on pathology review or molecular testing, varies according to the pretest probability of the condition of interest in the evaluated population. As the prevalence of thyroid cancer in nodules with AUS/FLUS or FN/SFN cytology can be different at various institutions and pathology practices (1, 4), we next calculated the PPV and NPV of the MPT in populations with different pretest cancer probability using the Bayes’ theorem and the sensitivity and specificity observed in the present study. The PPV was predicted to be > 50% for any prevalence > 15% and the NPV would range from 93 to 98% for cancer probability > 15% and < 35% (Figures 1A and 1B). Because the rate of true benign calls generated by the MPT

| Table 2. | Cohort characteristics and summary of molecular results in 109 preoperative aspirates |
|---|---|---|
| Surgical histology | Malignant | Benign |
| Number of cases | 35 | 74 |
| Age range [average] | 24−79 [52] | 21−89 [58] |
| Female/Male (%) | 79/21 | 72/28 |
| Mutation positive | 24 | 10 |
| miRNA positive | 20 | 6 |
| Positive by either test | 31 | 11 |
| Negative by both tests | 4 | 63 |

| Table 3. | Performance of the multi-platform miRNA and mutation test* |
|---|---|---|
| | Cohort | AUS/FLUS | FN/SFN |
| Number of cases | 109 | 58 | 51 |
| Sensitivity (%) | 89 [73–97] | 94 [73–100] | 82 [57–96] |
| Specificity (%) | 85 [75–92] | 80 [64–91] | 91 [76–98] |
| PPV (%) | 74 [58–86] | 68 [46–85] | 82 [57–96] |
| NPV (%) | 94 [85–98] | 97 [84–100] | 91 [76–98] |

PPV, positive predictive value; NPV, negative predictive value.

*Numbers between brackets indicate the 95% confidence intervals.
would be high in the relevant range of 15 to 35% prevalence (72 to 55%; Figure 1C), the MPT was predicted to significantly reduce the number of avoidable surgeries (Figure 1D). Independently of potential variations in thyroid cancer prevalence in the clinical setting, the use of the MPT would result in a constant 6.7-fold or 85% decrease in the number of diagnostic surgeries that may be performed in the absence of molecular testing (Figure 1D, dash line).

**Discussion**

In the present study, we showed that a molecular test combining miRNA expression and gene mutation detection can increase the diagnostic yield of molecular cytology by accurately classifying thyroid nodules with AUS/FLUS or FN/SFN cytology into benign or malignant categories. Our results underscore the value of this novel diagnostic algorithm and highlight key performance metrics that are important to improve the management of patients with thyroid nodules.

Cytopathology and molecular testing on indeterminate nodules are routinely combined in the clinical setting to identify different subsets of benign or malignant nodules prior to thyroid surgery. Molecular testing itself is often a combination of multiple assays with distinct analytical and/or clinical performance characteristics (6–9). For example, mutation testing includes \textit{BRAF} c.1799T>A, a mutation exclusively associated with thyroid carcinomas (100% specificity), as well as other oncogenic alterations, such as RAS mutations or PAX8-PPARG fusion transcripts, to identify different subsets of BRAF-negative carcinomas (10, 11). Similarly, the miRNA classifier interrogates well-known miRNAs associated with thyroid carcinogenesis (17–22, 24–26) to identify a subset of malignant nodules that are negative by mutation testing. One key feature of the miRNA classifier, however, is that it was designed and optimized to have high specificity in order to increase the sensitivity of the MPT without impacting significantly its specificity. In our cohort study, >90% (10/11) of the misclassified benign specimens were positive for a gene alteration and the single apparent false-positive by miRNA expression alone was from the left thyroid lobe of a study subject with multiple bilateral circumscribed follicular epithelial cell lesions and a papillary carcinoma in the right lobe. Among the mutation-negative cases, the specificity of the miRNA classifier was extremely high, 98% in both the cohort study (63/64) and the 3 case-control specimen sets (240/246).

The relatively high specificity of the MPT is a critical characteristic not only because it enables high and actionable PPV (less false positive) but also because it increases the benign call rate (more true negative). In a simplified clinical management algorithm in which all patients with AUS/FLUS or FN/SFN nodules would be sent to diagnostic surgery, the proportion of patients with a benign nodule who would undergo a potentially avoidable surgery is [1-prevalence], ie, 68% for the 32% thyroid cancer prevalence observed in our cohort study (Figure 2A). Following molecular reclassification with the MPT, 61% of the patients would have a negative MPT result with a low residual risk of malignancy (ROM) of 6% and only 39% of the patients might be referred to surgery because of a positive MPT result. With a PPV of 74%, only 10% of all the patients evaluated would have undergone surgery to remove a nodule later diagnosed as benign by histopathology (0.39x[1–0.74]=0.1). The relative decrease in the rate of avoidable surgery from 68% to 10% is the specificity of the MPT ([0.68–0.1]/0.68 = 0.85; see also algebraic demonstration in Supplemental Materials and Methods). Thus, use of the MPT in the clinical setting could potentially result in a 6.7-fold reduction in the number of unnecessary diagnostic surgeries, independently of variations in cancer prevalence.

Molecular reclassification of benign thyroid nodules to reduce the number of surgeries and associated healthcare costs has gained considerable attention in the past few years with the commercial availability of the Afirma gene expression classifier (AGEC). Although our study was not...
designed to compare different molecular methods, post-test probability metrics such as predictive values or surgery rates can easily be modeled using estimates of diagnostic sensitivity and specificity determined in representative clinical populations. In a recent multicenter cross-sectional cohort study, Alexander and colleagues reported a sensitivity of 90% and a specificity of 52% for the AGEC in thyroid nodules with AUS/FLUS or FN/SFN cytology (12). With a specificity of 52%, the AGEC would indeed reduce by 52% the rate of avoidable surgeries, from 76% of all nodules evaluated without molecular testing (24% cancer prevalence observed) to 37% after AGEC testing. This rate would be significantly improved to 11% with the MPT (P < .01), in part because of the higher PPV and mainly because of the higher benign call rate (Figure 2B). At 24% prevalence, 41% of the calls generated by the AGEC were “benign” with a 6% ROM (39% true benign call rate) while 67% of the MPT calls would be “benign” with a 4% ROM (65% true benign call rate). The relative increase in true benign call rate from 39% to 65% (1.65-fold or 65%) and the relative decrease in the rate of avoidable surgeries from 37% to 11% (3.3-fold or 69%) are both directly proportional to the specificity of the 2 molecular methods and are therefore independent of thyroid cancer prevalence (see algebraic demonstration in Supplemental Materials and Methods).

The relatively high sensitivity of the MPT (89%) is also an important characteristic as it contributes to PPV (more true positive) and is critical to reach actionable NPV (less false negative). The NPV observed in our study (94% at 32% prevalence) was not statistically different from the NPV of the AGEC (P = 1.0) while the large differences in PPV and true benign call rate between the 2 methods were significant (P < .01). Notably, the widths of the 95% CI for NPV and true benign call rate were similar with both methods, indicating similar precision in the estimation of performance (Supplemental Figures 2A to 2C). Statistical analyses performed either on nodules with AUS/FLUS cytology or on nodules with FN/SFN cytology showed similar patterns for NPV, PPV and true benign call rate, but with broader 95% CI as the number of computed features was smaller for each independent cytology category (Table 3 and Supplemental Figures 2D to 2I).

Recent advances in the understanding of the molecular pathogenesis of thyroid cancer and in next generation sequencing (NGS) technologies have exponentially increased the number of genetic alterations that can be interrogated in a single nodule aspirate. Nikiforov and colleagues recently reported a high sensitivity (90%) and specificity (93%) for the ThyroSeq v2 NGS panel in nodules with FN/SFN cytology at a single institution (30). Thus, NGS with extended mutation panels could further increase the sensitivity of the MPT by detecting additional malignant cases. However, NGS could also decrease specificity (and therefore the benign call rate) by detecting germline or low-level somatic mutations of unknown clinical significance in nodules classified as benign by surgical pathology. The clinical utility of this novel technology will require careful consideration of study design- and institution-specific parameters that can dramatically impact test performance such as FNA collection methods and sites, pathology review procedures, or histopathology features and cutoffs used to establish the final benign/malignant classification.

Cross-sectional cohort studies are designed to sample consecutive specimens in the source population and then retrospectively assess the history of exposures and outcomes that are available at the end of the study (7, 9, 12–14, 30). Estimation of diagnostic performance is therefore restricted to thyroid nodules with available surgical histology. Variations in local clinical practices and pathology classification may affect the calculation and extrap-
olation of performance at different cancer prevalence and
limit the comparison across studies performed in distinct
populations. In addition, our study was designed to eval-
uate clinical cases that had been submitted to molecular
testing as part of their diagnostic work-up at diverse en-
docrinology centers. Although this design guaranteed that
only FNAs truly representative of the target clinical pop-
ulation were interrogated, we do acknowledge that it may
have caused a study bias by increasing the number of mu-
tation-positive cases with available surgical outcome. This
in turn may have artificially decreased the specificity of
mutation testing and increased its sensitivity (69% ob-
served). Regardless, the miRNA classifier correctly iden-
tified 64% of the mutation-negative malignant cases. Even
with a sensitivity of 55 to 60% for mutation testing alone,
ie., lower than the range of 61 to 75% reported in the
literature (6–9), the sensitivity of the MPT would be 83 to
86%. With specificity at 85%, the NPV of the MPT would
still be high (94 to 95% NPV at 24% prevalence) and
similar to other methods routinely used in the clinical
setting.

In summary, we have demonstrated that a diagnostic
algorithm combining miRNA expression and gene muta-
tion detection yields clinically actionable molecular in-
formation in thyroid nodules with AUS/FLUS or FN/SFN
cytology. Based on the high PPV and NPV of the MPT, it
is reasonable to propose that patients with positive (ma-
lignant) MPT results may be sent to surgery while patients
with negative (benign) MPT results may benefit from a
more conservative management, ie, active follow-up with-
out surgery (Figure 2). On one hand, knowledge of the
mutational status would provide valuable diagnostic,
prognostic and theranostic information for the selection of
optimal and personalized therapeutic strategies. On the
other hand, the MPT would significantly increase the be-
nign call rate and decrease the rate of unnecessary di-
agnostic surgeries, thus further reducing the number of
2-step total thyroidectomies and the total number of sur-
greries performed. Our data and predictive models under-
score substantial improvements to thyroid cancer patient
management and collateral cost saving opportunities.

Acknowledgments
The authors would like to thank Prof. Thomas J. Giordano from
the Department of Pathology at the University of Michigan Med-
ical School for his expert pathology review of the surgical cases
used for miRNA classifier training as well as the personnel from
Asuragen’s molecular laboratory and from every participating
clinical site for their respective contribution.

Address all correspondence and requests for reprints to: Em-
manuel Laborier, PhD, Asuragen Inc., 2150 Woodward Street,
Suite 100, Austin, TX 78744, USA, Email: manulabourier@gmail.com.

This work was supported by.

Disclosure Summary: BA is an employee of Asuragen Inc. EL,
DW and SBH were employees of Asuragen Inc. at the time of the
study. EL is a consultant for PDI Inc. AS, AEB, MAL and MLM
have no conflicts of interest to declare.

References
1. Bongiovanni M, Spitalle A, Faquin WC, Mazzucchelli L, Baloch ZW.
The Bethesda System for Reporting Thyroid Cytopathology: a meta-
2. Cooper DS, Doherty GM, Haugen BR, Kloos RT, Lee SL, Mandel
SJ, Mazzaferri EL, Melcer B, Pacini F, Schlumberger M, Sherman SI,
Steward DL, Tuttle RM. Revised American Thyroid Association
management guidelines for patients with thyroid nodules and dif-
Vitti P, Nodules AAETFoT. American Association of Clinical En-
docrinologists, Associazione Medici Endocrinologi, and European-
Thyroid Association Medical Guidelines for Clinical Practice for
the Diagnosis and Management of Thyroid Nodules. Endocr Pract.
2010;16:S1–43.
DL, Zeiger MA, Westra WH, Wang Y, Khanafshar E, Felleagar G,
Rosai J, Livolsi V, Lamman RB. A large multicenter correlation study
5. Banks ND, Kowalski J, Tsai HL, Somervell H, Tufano R, Dackiw
APB, Marohin MR, Clark DP, Umbricht CB, Zeiger MA. A Diagno-
sic Predictor Model for Indeterminate or Suspicious Thyroid
F. Impact of proto-oncogene mutation detection in cytological spec-
imens from thyroid nodules improves the diagnostic accuracy of
7. Nikiforov YE, Ohori NP, Hodak SP, Carty SE, LeBoau SO, Ferris
RL, Yip L, Seethala RR, Tublin ME, Stang MT, Coyne C, Johnson
JT, Stewart AF, Nikiforov MN. Impact of mutational testing on the
diagnosis and management of patients with cytologically indeter-
minate thyroid nodules: a prospective analysis of 1056 FNA sam-
8. Nikiforov YE, Steward DL, Robinson-Smith TM, Haugen BR,
Molecular testing for mutations in improving the fine-needle aspira-
tion diagnosis of thyroid nodules. J Clin Endocrinol Metab. 2009;
94:2092–2098.
JM, Babu V, Blevins TC, Moore P, Andruss B, Labourier E. Cen-
tralized molecular testing for oncogenic gene mutations comple-
ments the local cytopathologic diagnosis of thyroid nodules. Thy-
10. Nikiforov MN, Kimura ET, Gandhi M, Biddinger PW, Knauja KA,
Basolo F, Zhu Z, Giannini R, Salvatore G, Fusco A, Santoro M,
Fagim JA, Nikiforov YE. BRAF mutations in thyroid tumors are
restricted to papillary carcinomas and anaplastic or poorly differ-
entiated carcinomas arising from papillary carcinomas. J Clin En-
docrinol Metab. 2003;88:5399–5404.
11. Nikiforov MN, Lynch RA, Biddinger PW, Alexander EK, Dorn
GW, 2nd, Tallini G, Kroll TG, Nikiforov YE. RAS point mutations and
PAK8-PPAR gamma rearrangement in thyroid tumors: evi-


