Abstract

In the past there has been considerable variation in the way thyroid cytology was reported by pathologists. As the need for improving patient care increased to include more conservative surgical procedures for the management of thyroid lesions, there was a need for a standardized reporting format. This had to be reproducible and universal so that the implications of the reporting categories were clear to all pathologists, surgeons or clinicians. In 2009, The Royal College of Pathologists (RCPath) published ‘Guidance on Reporting of Thyroid Cytology Specimens’ and this guidance has been revised in 2016 [1]. This document details the standardized format for reporting Thyroid FNAC used in the UK today. The classification of thyroid cytology into ‘Thy categories’ is similar to the Bethesda system used in the USA in that it stratifies thyroid FNA’s into various disease categories. The UK system uses five main categories ranging from Thy 1- Thy5 and certain categories are further subdivided into give additional diagnostic information. The Bethesda system uses six categories each with a percentage risk of malignancy. Currently there is little national data available on the percentage of cases that fall into each Thy category. Hence the positive predictive value (PPV) for each Thy category suggested in the RCPath guidance although rudimentary.

This audit was conducted at Wycombe Hospital Cellular pathology department to assess the adherence to RCPath ‘Guidance on reporting of Thyroid Cytology Specimens’. In addition to the aspects of the audit outlined in the guidance, we have also gathered information on the accuracy of local diagnostic opinions compared to those of specialists at the regional MDT to determine if there are large discrepancies between local and specialist practice.

Keywords: Neoplastic; Papillary Carcinoma; Diagnostic; Audit; Cases

Abbreviations: RCPath: Royal College of Pathologists; PPV: Positive Predictive Value

Aims or Objectives

a. To determine in what proportion of thyroid cytology reports a “Thy” category is included as well as a prose explanation of the findings.

b. To determine the percentage of cases that fall into each Thy category.

c. To correlate cytology with subsequent histology in order to determine diagnostic accuracy and the PPV for neoplasia and malignancy in each Thy category. In addition to the audit criteria, the sensitivity, the specificity, false negative rate, false positive rate and overall accuracy should be calculated and compared to those stated in the RCPath guidance.

d. To assess the number of Thy3a-Thy5 cases referred to the regional Thyroid Cancer MDT.

e. To compare the opinion of local pathologists with those of the regional MDT.

Audit Criteria

The agreed criteria range 100%:

Table 1: The agreed criteria range.

<table>
<thead>
<tr>
<th>RCPath Category</th>
<th>% Use of Each Category</th>
<th>PPV for Malignancy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Thy1/1c</td>
<td>18-22</td>
<td></td>
</tr>
<tr>
<td>Thy2/2c</td>
<td>42-51</td>
<td>1.4</td>
</tr>
<tr>
<td>Thy3a</td>
<td>10-May</td>
<td>17</td>
</tr>
<tr>
<td>Thy3f</td>
<td>14-16</td>
<td>Up to 40</td>
</tr>
<tr>
<td>Thy4</td>
<td>4-Feb</td>
<td>Up to 68</td>
</tr>
<tr>
<td>Thy5</td>
<td>10-May</td>
<td>Up to 100</td>
</tr>
</tbody>
</table>

Additional parameters

i. Sensitivity for malignancy between 65-98%

ii. Specificity for malignancy between 76-100%

iii. False negative rate between 0-5%

iv. False positive rate

v. Overall accuracy of 69-97%

No set standard, however there should not be major discrepancies between local and specialist opinions.
Methods

Sample selection

A search for all thyroid FNAs over a set time period (12/05/14 - 12/05/12) was conducted on the local software system (Winpath) using the following search criteria: Series: N, Time Period: 12/05/14 - 12/05/12, T Code: T96000 - Thyroid Gland

Cases where multiple aspirates were received under one lab number were treated as separate cases (i.e. aspirate of left/right sided lesion). The following exclusion criteria were then applied:

a. Cases from April and May 2014 as no subsequent histology was available (shortening the audited time period to 12/05/12-27/03/14).

b. Cases which were erroneously coded as Thyroid FNAs (Lymph node/ salivary gland aspirates).

This yielded a total of 812 cases for review including those which did not have an assigned Thy category. After the results were produced for criteria no. 1, these cases were excluded from the remainder of the audit as criteria 2-4 require a Thy category for analysis.

Data collection

An audit preformed was adapted from the template provided by the RCPath: 'Audit of reporting of thyroid cytology specimens and their correlation with thyroid histology' to record the following data:

1. Was there a prose report included with the Thy category?
2. Was there subsequent histology?
3. If there was subsequent histology, which disease category did the specimen fall into? (non neoplastic, benign neoplasm, papillary thyroid cancer, follicular carcinoma, carcinoma or other)
4. Was the cytology referred to MDT?
5. If referred to MDT, what was the specialist opinion?

Using the RCPath audit template, the subsequent histology was categorised as follows:

a) Non neoplastic: colloid nodule, hyperplastic nodule, Hashimoto’s thyroiditis, multinodular goitre, normal thyroid tissue.

b) Benign neoplasm: follicular adenoma.

c) Malignant neoplasm: papillary carcinoma, follicular carcinoma, carcinoma, other.

Each case was reviewed from Winpath and the authorised report used to record the relevant data. Subsequent results for the same patient were also checked to enable correlation with histology. If a histology result was present for the same lesion (laterality and site), immediately following the aspirate this was recorded. If however, the patient was investigated with further FNAs; this was recorded as ‘no subsequent histology.’ Each report was then checked for referral to the regional multidisciplinary team (MDT) meeting (as stated in each report) and the subsequent specialist opinion. Only the cases with a supplementary report including the specialist opinion were regarded as referred to MDT. Those cases where histology was reviewed in hindsight due to discrepancy with the initial cytology were not included as a true referral to MDT.

Statistical calculations

In order to calculate the required parameters, Thy 1/1c cases were disregarded to avoid providing false reassurance. Thy2/2c cases were deemed a negative result for malignancy and Thy 3a-Thy5 cases were considered positive. The diagnosis of a follicular adenoma on subsequent histology was considered a positive result. Thus the following definitions were applied:

a) True Negative (FN): Thy 2/2c cytology cases with subsequent 'non neoplastic' histology results.

b) False Negative (FN): Thy 2/2c cytology cases with 'benign i.e follicular adenoma' or 'malignant' history.

c) True Positive (TP): Thy 3a-Thy5 cytology cases with subsequent ‘benign i.e follicular adenoma’ or ‘malignant’ histology.

d) False Positive (FP): Thy 3a-Thy5 cytology cases with subsequent 'non neoplastic' histology results.

Once the data was divided into these four categories, the following standard calculations were used:

- Sensitivity: TP/ (TP + FN)
- Specificity: TN/ (TN+FP)
- Accuracy: (TP +TN)/ (TP +TN+ FP+ FN)
- Positive Predictive Value (PPV): TP/ (TP+FP)
- False Negative Rate: FN/ (TN +FN)
- False Positive Rate: FP/ (TP +FP)

Results and Discussion

Adequacy of report

The majority of cases reviewed (97.7%) were of adequate quality and included both the Thy category and an accompanying prose report as recommended by the RCPath. The remainder of cases did not include the Thy category however a written report was issued from which the diagnosis could be ascertained. These cases were subsequently removed from the sample as the other audit criteria assess the designated Thy category (Table 2).

Table 2: Table showing the quality of reports compared to audit standards.

<table>
<thead>
<tr>
<th>Quality of Report</th>
<th>No. of Cases</th>
<th>Percentage of Total (%)</th>
<th>Audit Standard (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Prose Report and Thy Category</td>
<td>793</td>
<td>97.7</td>
<td>0</td>
</tr>
<tr>
<td>Prose Report Only</td>
<td>19</td>
<td>2.3</td>
<td>0</td>
</tr>
<tr>
<td>Thy Category Only</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

Citation: Kuruppu K, Wathug G (2017) Thyroid Cytology Reporting and Diagnostic Accuracy: A Departmental Audit of the RCPath Guidelines. Int Clin Pathol J 5(4): 00141. DOI: 10.15406/icpjl.2017.05.00141
Distribution of cases into each thy category

The distribution of cases is illustrated. The vast majority of Thyroid FNA’s received by the Wycombe Pathology department fall into the Thy2 category (409 cases, 51.58%). The second largest category is Thy 1 (215 cases, 27.11%), indicating a significant cost in consultant and lab processing time for samples which are insufficient for diagnostic purposes. Very few cases fall into the Thy5 category (10 cases, 1.26 %) and even fewer are categorized as Thy 4 (5 cases, 0.63%). Thy 3 lesions are predominantly designated Thy 3f (50 cases, 6.31 %) with the provision that a follicular lesion cannot be excluded.

This is comparable to a similar audit carried out at NHS Tayside [3]. The author of this Scottish study found the largest proportion of their specimens were Thy2 (44%), followed by Thy3 (26%), Thy 1 (25%), Thy 4 and Thy5 (2.5% each).

In comparison to the percentage usage of each category by the RCPath, our data range extends beyond theirs for Thy1-2c categories and is lower the RCPath for Thy3a-5 cases. The reasons for this may be multifactorial including the skill level of those taking the FNA at different trusts, the availability of expert cytopathologists at tertiary centres and varying degrees of confidence in classifying indeterminate cases.

Correlation with histology and PPV

Unfortunately, a significant proportion of the sample did not have subsequent histology (650 cases). This correlates with the substantial number of Thy1 (276 cases) and Thy2 (239 cases) which would not be managed by surgical excision. Excluding Thy1/1c cases, in those which had subsequent histology, 51 were non neoplastic lesions and 35 cases were benign follicular adenomas. 13 cases proved to be papillary carcinomas and 5 were follicular carcinomas. Within the Thy5 category (10 cases), 3 did not have subsequent histology and the remaining 7 cases were all malignant (6 papillary carcinomas, 1 other) which establishes 100% diagnostic accuracy in these cases, as is the set standard.

For the Thy3f category, none of the cases were subsequently diagnosed as follicular carcinomas. There were however four Papillary carcinomas and 1 ‘other’ malignancy. The vast majority of these (22 cases) were benign follicular adenomas. The ‘other’ malignancies encountered included a lymphoma, anaplastic carcinoma and metastasis from a primary elsewhere. The results for each Thy category and the subsequent histology are shown in Table 4.

Table 4: Table of correlation between histology and cytology.

<table>
<thead>
<tr>
<th>Histology</th>
<th>Thy 1</th>
<th>Thy 1c</th>
<th>Thy 2</th>
<th>Thy 2c</th>
<th>Thy 3a</th>
<th>Thy 3f</th>
<th>Thy 4</th>
<th>Thy 5</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>No Histology</td>
<td>188</td>
<td>52</td>
<td>353</td>
<td>27</td>
<td>9</td>
<td>15</td>
<td>3</td>
<td>3</td>
<td>650</td>
</tr>
<tr>
<td>Non Neoplastic</td>
<td>18</td>
<td>7</td>
<td>38</td>
<td>3</td>
<td>1</td>
<td>8</td>
<td>1</td>
<td>0</td>
<td>76</td>
</tr>
<tr>
<td>Benign</td>
<td>8</td>
<td>1</td>
<td>11</td>
<td>0</td>
<td>2</td>
<td>22</td>
<td>0</td>
<td>0</td>
<td>44</td>
</tr>
<tr>
<td>Ptc</td>
<td>1</td>
<td>1</td>
<td>2</td>
<td>0</td>
<td>1</td>
<td>4</td>
<td>0</td>
<td>6</td>
<td>15</td>
</tr>
<tr>
<td>Fc</td>
<td>0</td>
<td>0</td>
<td>5</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>5</td>
</tr>
<tr>
<td>MTC</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Other</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>3</td>
</tr>
<tr>
<td>Total</td>
<td>215</td>
<td>61</td>
<td>409</td>
<td>30</td>
<td>13</td>
<td>50</td>
<td>5</td>
<td>10</td>
<td>793</td>
</tr>
</tbody>
</table>

In this audit, the FNA was viewed as a screening tool for determining which thyroid lesions required surgical excision. Thereby, only Thy2/2c FNA’s were considered ‘negative’. The PPV for both neoplastic lesions (i.e including follicular adenomas and malignancies) and for malignant lesions alone was calculated for Thy3a-5 categories. A PPV for Thy1-2c is not provided as the authors were unable (Table 5) to find evidence of how these were calculated in RCPath guidelines or the definitions used to determine them.

When compared to a large multicentre study done in 2010 [4], which found that the risk of malignancy for ‘indeterminate’ cytology cases was 34% and 98% in malignant cytology cases our PPV for neoplasia in Thy3 cases of 75% is much higher. Our PPV for malignancy in Thy3a cases is also much higher than the RCPath standards. The PPV for Thy4 was lower at 50% however this is likely to be secondary to the small number of cases within this category. We were compliant with the RCPath standard for Thy5 of having a 100% PPV for this category (Table 5).
Table 5: Table of positive predictive values.

<table>
<thead>
<tr>
<th>Category</th>
<th>PPV for Neoplasia (%)</th>
<th>PPV for Malignancy (%)</th>
<th>RCPath PPV for Malignancy (%)</th>
<th>Bethesda - Associated Risk of Malignancy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Thy3a</td>
<td>75</td>
<td>25</td>
<td>17</td>
<td>5-15</td>
</tr>
<tr>
<td>Thy3f</td>
<td>77</td>
<td>14.3</td>
<td>Up to 40</td>
<td>15-30</td>
</tr>
<tr>
<td>Thy4</td>
<td>50</td>
<td>50</td>
<td>Up to 68</td>
<td>60-75</td>
</tr>
<tr>
<td>Thy5</td>
<td>100</td>
<td>100</td>
<td>Up to 100</td>
<td>97-100</td>
</tr>
</tbody>
</table>

When the additional parameters were calculated we achieved the targets set by the RCPath guidance for specificity and overall accuracy only. The other parameters fell outside the RCPath targets. The overall sensitivity of 64% fell short of the target range. The false negative rate of 31% was much higher than the target of 0-5% and the false positive rate was higher than recommended at 23%. Compared to a similar study in 2008 [6] which found a sensitivity of 92.6%, a specificity of 91.6% and a false negative rate of 2.3%, our local department rates need significant improvement, particularly for sensitivity and false positive/negative rates. A specificity of 80% suggests that local pathologists are good at identifying true negative cases. Although the sensitivity rate is lower than the target range, it is only so by 1%, suggesting that positive cases are in fact being picked up well and a larger sample size might yield higher results. The high false negative/positive rates may be explained by the inclusion of benign follicular adenomas as positive results especially as 31% of all follicular adenomas in our sample came from a Thy2 (negative) FNA. The overall accuracy of 72% falls within the required range and thus the department is performing adequately, but results could be markedly better. These results are limited by the small number of cases used (143 cases) to calculate these statistics as all those without histology (650 cases) were not included. It is also important to note that the target ranges provided by the RCPath guidance is likely to evolve as more data becomes available in the future (Table 6).

Table 6: Table of additional parameters compared to target range.

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Percentage (%)</th>
<th>RCPath targets (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sensitivity</td>
<td>64</td>
<td>65-98</td>
</tr>
<tr>
<td>Specificity</td>
<td>80</td>
<td>76-100</td>
</tr>
<tr>
<td>False Negative Rate</td>
<td>31</td>
<td>0-5</td>
</tr>
<tr>
<td>False Positive Rate</td>
<td>23</td>
<td>0-5.7</td>
</tr>
<tr>
<td>Overall Accuracy</td>
<td>72</td>
<td>69-97</td>
</tr>
</tbody>
</table>

Correlation with specialist opinion

In addition to the set standards, we compared the initial diagnostic opinion of the local pathologists with the final diagnostic opinion of the specialists at the regional MDT to determine if there were major discrepancies. When cases were initially given a Thy5, there was agreement with the specialists for 90% of these cases. Only one case was downgraded to Thy4. Of the four Thy4 cases which were sent to MDT, two cases were agreed as Thy4, one case was upgraded to Thy5 and one case was downgraded to Thy3a. This result is difficult to interpret accurately as there are so few numbers of cases in this category. However, downgrading to Thy3a will confer a difference in management for the patient who may subsequently undergo a hemithyroidectomy instead of a total thyroidectomy [5-7].

Figure 1: Graph of all cases referred to MDT.

Figure 2: Graph of all cases which should be referred to MDT.

Referral to regional MDT

The standards set by the RCPath suggest that 100% of Thy 4 and Thy5 cases should be referred to MDT. As per local protocols, Thy3a and Thy3f should also be referred. The results show that 92% of Thy3a, 90% of Thy3f, 80% of Thy 4 and 100% of Thy 5 cases were referred to MDT and a subsequent specialist opinion was available (Figure 1 & 2).
After 12 Thy3a cases were submitted for specialist opinion, three were upgraded and two were downgraded. This means a 41.6% diagnostic resolution of these indeterminate cases. There were a total of 45 Thy3f cases referred to MDT. 57% were agreed as Thy3f, 27% were downgraded and 16% were upgraded. This infers a 43% diagnostic resolution. These statistics compare with a published literature review which found a 42.5% diagnostic resolution on ‘indeterminate’ cases sent for second opinion.

It was outside the scope of this audit to compare the subsequent histology in cases where there was discrepancy between local and regional opinion (Figure 3).

**Conclusion**

This audit shows that the department is 97.7% compliant with audit criterion 1 which requires a prose report to accompany the Thy category. The percentage usage of each category is vastly different at our trust to those stated by the RCPPath standards. In addition, we have achieved the targets for specificity (80%) and overall accuracy (72%) however the sensitivity (64%), the false negative rate (31%) and the false positive rate (23%) fall far outside the recommended range. We achieved 100% compliance with Criterion 4, but only for Thy 5 cases. All other categories (Thy 3a, Thy3f and Thy 4) included some cases which should have been referred to MDT but for unknown reasons were not sent. The value of referral to regional experts was highlighted by a 41.6% and 43% diagnostic resolution of Thy3a and Thy3f cases respectively.

**Recommendations**

a. Continue to collect data on the proportion of cases which fall into each Thy category to be contributed to national data.

b. To improve the PPV of Thy3a-Thy4 cases, consider encouraging attendance at regional MDT’s, teaching sessions and courses to train staff to increase diagnostic accuracy.

c. Inform all medical staff and administrative staff which cases should be referred to MDT. Develop a Performa which would trigger an automatic referral to MDT for the relevant cases.

d. Re-audit in 12 months.

**Acknowledgment**

None.

**Conflict of Interest**

The authors declare they have no actual or potential competing interests.

**References**


