Liquid-Based Cytology in Fine-Needle Aspiration Biopsies of the Thyroid Gland

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**Key Words**
Thyroid nodules · Fine-needle aspiration cytology · Liquid-based cytology

**Abstract**

**Objectives:** Fine-needle aspiration biopsy is regarded as the most important diagnostic tool for thyroid lesions because of its simplicity, safety, and cost-effectiveness. However, its pivotal role in the correct characterization of the majority of nodules is impaired by the difficulties in discriminating benign from malignant follicular-patterned lesions.

**Study Design:** Liquid-based cytology (LBC) is a semiautomated device that has recently become widely available and has gained popularity as a method of collecting and processing both gynecologic and nongynecologic cytologic specimens. It achieves a diagnostic sensitivity as accurate as conventional preparations, especially for its excellent cell preservation and lack of background which decrease the amount of inadequate diagnoses.

**Results:** In many cases the cytologic features are similar in both methods, but the colloid film and the lymphocytic component are more easily evaluated on direct smears whereas nuclear details and colloid globules are better evaluated in LBC slides. The material stored in the preservative solution could be effectively used for the application of immunocytochemical and molecular techniques.

**Conclusions:** LBC-processed biopsies represent a valid alternative to conventional cytology. The possibility of applying additional techniques enhances the efficacy of the cytologic diagnosis of thyroid lesions.

**Introduction**

Fine-needle aspiration biopsy (FNAB) was introduced for the first time in the USA in the 1930s, but it was only until the 1950s that it was recognized in Sweden as an invaluable diagnostic tool. Since then this method has spread worldwide because of its simplicity, safety, and possibility of repetition [1, 2].

The incidence and mortality of thyroid cancer do not qualify this tumor as an important public health problem, but the number of surgical lobectomies performed to assess its presence makes it a disease of economic importance. In this setting FNAB is regarded as the most accurate and cost-effective method for the selection of patients with thyroid nodules [3].

FNAB of a thyroid nodule is preferably carried out under sonographic guidance, and presently the mane-
The thin-layer or LBC technique, originally developed for application in gynecologic cervical smears, has progressively gained consensus after being applied in both nongynecologic and fine-needle aspiration cytology. This method is based on a 2-step procedure: (a) fixation of the totality of the material in an alcohol-based solution (methanol or ethanol, depending on the technique – see below) and (b) automated processing of the material to obtain a thin layer of representative cells. The innovation is a computer-assisted device which allows the transfer of the fixed and partially disaggregated cells onto a single slide. The 2 most common FDA-approved methods for processing the cytologic samples use an alcohol-based fixative solution. In the first method (ThinPrep2000™; Hologic Co., Marlborough, Mass., USA), the cells are aspirated from a methanol-based solution (CytoRich™) and then filtered and transferred onto a positively charged slide with gentle positive pressure. In the latter method the cells are collected in an ethanol-based solution (CytoRich™), centrifuged twice, and then slowly sedimented onto a poly-L-lysinated slide and eventually stained with a specific hematoxylin-eosin stain (SurePath™; TriPath Imaging, Burlington, N.C., USA). The final result for both methods is 1 slide for each lesion where all cells are concentrated in a thin layer on the central area of the slide measuring 20 mm² for ThinPrep and 13 mm² for SurePath [7].

Two methods can be chosen for processing thyroid FNAB with the LBC technique: ‘split sample’ and ‘direct to vial’. The former method requires that the aspirated material be split into 2 equal parts: the first half is smeared onto slides for CS, and the second half is submerged in a preservative solution for LBC. This method is ideal when carried out by a skilled operator; in this setting cytopathologists are generally well trained for an equal subdivision of the material and they can immediately check its adequacy after staining a CS. However, most clinicians and radiologists tend to extrude the majority of the material onto the slides for CS and to save only a small, sometimes nonsignificant portion for LBC. In these cases double sampling of the lesion – the first smeared onto slides for CS and the second ‘direct to vial’ for LBC – is recommended even if the patient may undergo 2 or more biopsies of the same lesion with a higher risk of complications.

Although some authors are still skeptical regarding the efficacy of LBC when used alone for diagnosing thyroid lesions [8], good results have been achieved by our (and other) groups, especially in recent years. Since November 2003, the majority of the almost 19,000 FNAB carried out at the 'Agostino Gemelli’ School of Medicine and Hospital of Rome have been processed by LBC alone, adopting the CS for fewer than 2,000 of them performed under manual guidance. This experience has been reported in many studies published from 2005 to date and it emphasizes the efficacy of the LBC technique for the correct preoperative diagnosis of more than 500 malignant neoplasms. In the study published in 2009 by Rossi et al. [9], 3 parameters of efficacy (inadequacy, indeterminacy, and malignancy rates) were evaluated in a series of 10,360 thyroid FNAB. In that investigation the use of ThinPrep alone was as effective as CS in decreasing both inadequate and indeterminate cases whereas the majority of problems occurred when LBC and CS were simultaneously adopted (split sample, see above) [9]. These problems accounted for a high rate of inadequate and false-negative diagnoses and were also reported in many studies published in different countries [10–14]. In series where the direct-to-vial method is used, and all of the cellular material is therefore submitted for cytologic analysis, the average cellularity is generally higher than with the split-sample method and the rates of inadequacy and false positivity decrease significantly [9, 13, 15, 16]. Regarding the results of Geers and Bourgain [17] using the SurePath method, these authors did not achieve good results in terms of the inadequacy rate but there might be some reasons related to the technical differences between ThinPrep and SurePath (see Inadequate Smears). Nevertheless, they claimed a significant reduction of the false-negative diagnoses.

The material remaining in the vial after cytologic diagnosis can be used for the application of ancillary techniques such as immunocytochemistry (ICC), flow cytometry, and molecular biology because the LBC method enables the storage of a variable amount of cells for up to 6 months after the biopsy [18–22].
Inadequate Smears

In the best interest of the patient, the cytopathologist must receive an adequate biopsy specimen to make a meaningful cytopathologic evaluation and a correct diagnosis. The final cytologic specimen must be adequate in terms of cellularity and satisfactory in terms of quality (thickness, fixation, and staining).

An FNAB is defined as unsatisfactory when fixation, smearing, or staining artifacts impair the interpretation of the final slide. A slide is nonrepresentative when the cellularity does not represent the true components of the lesion (e.g., insufficient amount of follicular cells) [23]. In the first instance the inadequacy of the sampling could be attributed to an incorrect technique, and in the latter the characteristics of the lesion do not allow a definitive cytoplasmic diagnosis [24–26]. In both CS and LBC the adequacy criteria are met when at least 6 clusters of 10–20 well-preserved cells are observed [24–28]. When an LBC slide does not contain an adequate number of cells, a second slide can be prepared with the residual cells to meet the adequacy criteria [29]. In the latter scenario up to 18% of cases, diagnosed as inadequate after the first slide, can be reclassified as adequate after the second slide and this result is more significant than the data by Hasteh et al. [30] who dismiss as expensive and useless the preparation of a second ThinPrep slide. The LBC ThinPrep-processed slides may show a few ill-preserved cells at the periphery of the circle because transfer from the preservative solution to the specific slide requires positive pressure which can give rise to cellular artifacts. SurePath, according to Geers and Bourgain [17], yields an inadequacy rate of 25% in thyroid aspiration cytology. This result can probably be attributed to the low sedimentation rate of the colloid droplets which might hinder their inclusion in the final slide [17].

Cystic lesions are most commonly responsible of nonrepresentative cases [31, 32]. A thyroid cyst is a true pseudocystic lesion (without a wall of follicular cells); more frequently it represents cystic or hemorrhagic regression of a nodule which might turn out to be benign or malignant after repeated samplings. LBC slides show many hemosiderin-laden histiocytes with very few clusters of follicular cells and colloid droplets (see below). A conclusive diagnosis is based on the morphologic features of the follicular cells, if their amount meets the adequacy criteria, and on the number of colloid droplets. Otherwise, the final diagnosis is nonrepresentative. In our studies [9, 15] cystic/hemorrhagic lesions accounted for about 7% of all cases.

When an FNAB of a cystic nodule with a solid component is repeatedly nonrepresentative, surgical excision of the nodule may be considered to avoid the possibility of a malignant lesion (between 8 and 19% of cases according to literature) [7, 27, 28, 31, 32].

Benign Diseases of the Thyroid

Goiter

FNAB of nodules in goiters span a wide spectrum of morphologic changes reflecting different stages of the disease: early follicular hyperplasia, cycles of involution/regeneration, and nodule formation. Secondary changes such as oxyphilic metaplasia, recent and old hemorrhage, cystic degeneration, necrosis, granulation tissue, fibrosis, and calcification may occur in all of these stages.

Two critical points differentiate LBC from CS. First, the cells in each slide are a monolayered representative sample of all of the material collected in the vial. The whole cellularity is seldom collected in a single LBC slide; thus, a variable number of them is stored in the preservative solution to be used for additional investigations. Second, the automated process causes some changes in both cellular and background morphology (table 1). One of the most important changes occurring in LBC compared to CS slides is the appearance of the colloid which is fragmented during the filtering procedure. Therefore, this substance is observed as small droplets in the background of benign nodules: the higher the number of these colloid droplets, the higher the likelihood of benignancy of the nodule [15, 20, 34]. Thus, the LBC evaluation of the amount of colloid is quantitative whereas in CS the presence of colloid in the background usually does not require quantitation (fig. 1). The lack of watery colloid, which has been considered a major problem by some authors [8, 12–16], can be considered a minor diagnostic detail provided this substance is accurately sought instead of being only roughly evaluated at a lower magnification. Colloid droplets can sometimes be detected mixed with hemorrhagic debris and hemosiderin-laden histiocytes within a cystic lesion (see above). Unlike CS, the LBC picture may be safely interpreted, in the presence of a small amount of typical follicular cells, as goiter with hemorrhagic changes [33, 35].

Thyroiditis

A cytologic diagnosis of thyroiditis can be very puzzling at times, and the clinical and immunologic findings must be considered [36, 37]. A clinical picture of a tender...
<table>
<thead>
<tr>
<th>Diagnosis</th>
<th>CS</th>
<th>LBC</th>
<th>Histologic correspondence</th>
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<td><strong>Nonneoplastic lesions</strong></td>
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<td>Colloid nodule</td>
<td>Abundant and clumped colloid; large sheets of small thyrocytes with ‘hyalinized stroma’; foamy ‘colloidophagic’ histiocytes</td>
<td>Clusters of small monomorphic thyrocytes with clear, sometimes ‘granular’ cytoplasm; small clumps of dense colloid (colloid globules); foamy ‘colloidophagic’ histiocytes</td>
<td>Diffuse or nodular goiter; macrofollicular adenomatous nodule in a goiter</td>
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<td>Thyroiditis</td>
<td>Inflammatory cells (mostly mature lymphocytes) in the background; fragments of reticular tissue with epithelioid histiocytes; small clusters of thyrocytes sometimes with oxyphilic metaplasia; scant colloid; plurinucleated histiocytes</td>
<td>Small clusters of thyrocytes or oxyphilic cells mixed with lymphohistiocytes (lymphoepithelial clusters); the same cells are present in the background; few droplets of colloid; large plurinucleated histiocytes</td>
<td>Granulomatous De Quervain’s thyroiditis; lymphocytic ‘Hashimoto type’ thyroiditis</td>
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<td><strong>Follicular lesions</strong></td>
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<td>FN</td>
<td>Scant colloid; microfollicles or small clusters of medium-sized thyrocytes, sometimes with slight nuclear pleomorphism (hyperfunction) and rounded nuclei; fibrovascular tissue; hemorrhage with hemosiderin-laden histiocytes</td>
<td>Small clusters of medium-sized thyrocytes with pleomorphic nuclei, generally with regular outlines; fibrin flakes; scant or no colloid droplets; hemosiderin-laden histiocytes</td>
<td>Adenomatous nodule; follicular adenoma; minimally invasive follicular carcinoma; FVPC</td>
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<td>OFN</td>
<td>Scant colloid; sheets or clusters of oxyphilic cells; hemorrhage; scattered inflammatory cells and reticular tissue in the background may be seen in thyroiditis</td>
<td>Small aggregates of oxyphilic cells with brilliant cytoplasmic granules and large hyperchromatic pleomorphic nuclei; fibrous tissue; scant colloid globules (with inflammatory cells; see Thyroiditis)</td>
<td>Oxyphilic adenomatous nodule; oxyphilic neoplasm; oxyphilic hyperplastic nodule in thyroiditis</td>
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<td><strong>Suspicious for carcinoma</strong></td>
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<td>Follicular lesion suspicious for PC</td>
<td>Absent colloid; small clusters or microfollicles of medium-to-large-sized thyrocytes with moderate nuclear pleomorphism, irregular nuclear membrane, clearing, and grooves (no pseudo-inclusions or papillae), and often oxyphilic cytoplasms; hemorrhage; fibrovascular tissue</td>
<td>Small clusters of medium-sized thyrocytes (see FN) mixed with scattered aggregates of large cells with pleomorphic nuclei, clear chromatin, and irregular nuclear outlines; no papillae or nuclear pseudo-inclusions; fragments of fibrous tissue; if prominent nucleioli: hyperplastic nodule (Graves’ disease)</td>
<td>FVPC; ‘toxic’ or hyperplastic adenoma; follicular carcinoma. The frozen-section examination is suggested</td>
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<td><strong>Malignant neoplasms</strong></td>
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<td>PC</td>
<td>Absent colloid; large irregular sheets or papillae lined by large thyrocytes with severe nuclear pleomorphism and irregularities, with pseudo-inclusions and grooves; hemorrhage; fibrovascular tissue</td>
<td>Aggregates of thyrocytes with large and elongated nuclei with grooves and focal pseudo-inclusions; plurinucleated giant cells; fibrin filaments; iron-laden histiocytes; small and thin papillae (uncommon)</td>
<td>PC</td>
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<td>Medullary carcinoma</td>
<td>Small clusters of cells with round or ovoid slightly pleomorphic nuclei eccentrically disposed (‘plasmacytoid cells’); fragments of hyaline material; scant colloid; calcitonin and CEA positivity</td>
<td>Isolated cells with eccentrically disposed (‘plasmacytoid cells’) or cylinder-shaped round or ovoid slightly pleomorphic nuclei; calcitonin and CEA positivity</td>
<td>Medullary carcinoma</td>
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OFN = Oxyphilic follicular neoplasm; FVPC = follicular variant of papillary carcinoma.
gland with small inconspicuous nodules whose cytology shows mature lymphocytes, plasma cells, and scattered giant cells is likely to be a granulomatous thyroiditis (De Quervain’s).

The LBC picture of lymphocytic thyroiditis is similar to CS with one exception: in a CAT the amount of lymphocytes in the background of the slide can be higher than normal because of the spinning of the material before the automated process. When a CAT is suspected, the detection of lymphoepithelial clusters on an inflammatory background is the pivotal clue for the diagnosis (fig. 2) [16]. This setting represents the opposite of goiter: LBC diagnosis of thyroiditis requires identification of the lymphoepithelial clusters, and the simple detection of lymphohistiocytic cells in the background of the slide does not warrant a diagnosis of thyroiditis. According to the literature, the LBC diagnosis of thyroiditis is reliable [8]. Oxyphilic hyperplastic nodules in a CAT should not be surgically removed because they represent the functional replacement of parenchyma infiltrated by the inflammatory cells. The occurrence of either an oxyphilic carcinoma or a non-Hodgkin primary lymphoma within a CAT is exceedingly uncommon compared to the frequency of papillary carcinoma (PC) which can easily be detected by cytology. Thus, identification of lymphoepithelial clusters, mostly when made up of oxyphilic cells, virtually rules out an oxyphilic neoplasm and warrants a simple follow-up for the patient.

**Toxic Goiter**

FNAB is not usually the first choice in cases of toxic goiter unless a ‘cold’ lesion is detected during a scan. However, some cases in which the cytopathologist faces this diagnosis may occur: (a) a preclinical hyperfunctioning nodule, where a decrease of the circulating TSH is the only clinical sign; (b) a ‘cold’ nodule within a toxic goiter, where the sampled lesion can be a hyperplastic nodule inhibited by the surrounding toxic goiter; (c) a rapidly growing toxic nodule during suppressive therapy, and (d) a hyperplastic nodule in a CAT. The cytologic picture of a toxic nodule is that of a follicular neoplasm (FN) showing microfollicles layered by medium-sized thyrocytes with distinctive vacuolated cytoplasms (‘fire-flare’ or ‘flame cells’; fig. 3) [38, 39]. The colloid in the background is scant and focal nuclear pleomorphism may be detected in a longstanding goiter. Scattered lymphoid cells may be present in a smear of a toxic goiter and, on the other hand, cells with features of hyperfunction may appear in an otherwise obvious colloid nodule or in thyroiditis.

The hyperfunctioning features described above are important in the decision for follow-up rather than surgery in a single follicular-structured nodule [38]. Since the malignant risk in a toxic nodule is low the detection of hyperfunctioning thyrocytes rules out this possibility, avoiding an unnecessary thyroidectomy.
Follicular Lesions

There are very few differences in the cytologic pictures of FN in LBC compared to CS. This diagnosis is based on the identification of microfollicles made up of medium-sized follicular cells in a background with scant colloid. The amount and morphology of the follicular cells allow the inclusion of an individual lesion in one of the categories that have been recently devised in Europe and in the USA. There are 3 possible scenarios:

1. The lesion is highly cellulated but the cells are monomorphous with occasional enlarged nuclei. This lesion may correspond to the cellular adenomatous nodule of DeMay and is usually included in the nonneoplastic category of European classifications (Thy 2 of the BTA classification and TIR 2 of the Italian classification) [31, 40]. On the other hand, the Bethesda classification has established a different category for this picture which is defined as ‘a follicular lesion of undetermined significance’ or ‘atypical cells of undetermined significance’. The difference between these two categories resides in the different risk of malignant occurrence; in the American classification it is in the range of 5–15% (mostly follicular carcinoma) whereas in the European systems it is closer to nonneoplastic lesions.

2. The lesion is mostly follicular structured and is composed of follicular cells with elongated and clear nuclei (sometimes with grooves and peripheral nucleoli) (fig. 5a). Papillae, psammomatous bodies, or nuclear pseudoinclusions are not identified. These cases are included in the category of suspicious for PC which bears a risk of malignancy ranging between 50 and 70%. This category warrants surgical removal of the nodule as a follicular variant of PC is very likely to be found upon histologic examination (more than 90% of cases).

LBC pictures of these microfollicular-patterned lesions allow a more thorough evaluation of the nuclear details of the follicular cells because the background is devoid of blood clots and fibrin.

Another advantage of the LBC technique over CS in the diagnosis of follicular lesions is the possibility of ap-
plying additional investigation methods (ICC, flow cytometry, and molecular biology). These techniques, especially ICC and molecular biology, are particularly helpful in refining the diagnosis of follicular lesion and can be applied to the cells left in the vial which are safely stored at room temperature for up to 6 months after the sampling of the lesion (see above).

**Oxyphilic (Hürthle Cell) Lesions**

The FNAB of an oxyphilic lesion is categorized as an ‘oxyphilic or Hürthle cell neoplasm’ when follicles made up of more than 80% oxyphilic cells are identified. The colloid amount may be scant (but is sometimes abundant), and features of old hemorrhage (hemosiderin-laden histiocytes) may coexist. Both fire-flare cells (detected in hyperfunctioning lesions or in juvenile thyroiditis – see above) and small thyrocytes (more than 20% of the cellular component) suggest a benign lesion with an oxyphilic component. Unlike their follicular counterpart, oxyphilic cells may feature nuclear enlargement and pleomorphism either in benign neoplasms or even in hyperplastic lesions. The most paradigmatic case is represented by hyperplastic nodules in CAT which may show a striking nuclear pleomorphism of the oxyphilic component [1, 2, 41–49]. Some authors have attempted to correlate the atypia of oxyphilic cells (and some other features such as transgressing vessels) with the risk of malignancy, but their results are still debatable [47, 49]. Thus, a lesion made up of oxyphilic cells should be included in the FN category (Thy 3 of the BTA classification and TIR 3 of the Italian classification) if more than 80% of the cells are oxyphilic. Otherwise, it should be included in the nonneoplastic category as it might be either a hyperplastic nodule in a CAT or a oxyphilic component of a goiter.

**Thyroid Malignant Neoplasms**

The cytologic diagnosis of thyroid malignancy does not differ substantially in LBC preparations with respect to CS. As mentioned before, the clear background facilitates the identification and characterization of the cellular details distinctive of each neoplasm.

The most important malignant tumor which should be appropriately identified is papillary carcinoma. LBC diagnosis of PC is straightforward when the nuclear pseudoinclusions (major criterion) are detected even within tridimensional clusters of cells with nuclear elongation and clearing (minor criterion – see previous paragraph; fig. 5b) [50, 51]. Papillary structures and psammoma bodies are seldom identified. The lack of fibrin clots and red blood cells in the background enables the identification of the distinctive PC cells within a hemorrhagic or cystic degeneration of the tumor. In earlier re-

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**Fig. 5.** a Group of thyrocytes showing nuclear elongation and clearing and irregularity of the nuclear membrane outline, suspicious for carcinoma (Papanicolaou, ThinPrep, ×1,000). b Clear pseudoinclusion (‘Orphan Annie’s eye’) centered in the nucleus for the diagnosis of PC (Papanicolaou, ThinPrep, ×1,000).
ports, the difficulty in detecting the distinctive nuclear features of PC was one of the most important objections against adoption of the LBC technique in thyroid cytology. However, the most recent investigations have somewhat corrected those data and emphasized that accurate evaluation of both the major and the minor diagnostic criteria of PC plays a pivotal role in the preoperative characterization of PC. This diagnostic efficacy includes the diffuse sclerosing and tall cell variants of PC which warrant an aggressive surgical approach.

Medullary thyroid carcinoma (MTC) is a difficult diagnosis in thyroid cytology. It usually relies on the identification of a double population of both plasmacytoid and spindle cells with variable nuclear pleomorphisms (fig. 6a). The amounts of these components can be variable in a way that an MTC may alternatively look like a plasmacytoma or a mesenchymal neoplasm. As the cytologic diagnosis of MTC is very important in order to establish the most appropriate surgical strategy, the LBC technique offers the opportunity to identify the presence of calcitonin in neoplastic cells (fig. 6b). To avoid false-positive results, the negativity for thyroglobulin of the neoplastic parafollicular cells needs to be confirmed [55].

Anaplastic thyroid carcinoma (ATC) is an uncommon finding in thyroid cytology as this tumor presents a rapid growth which frustrates attempts to plan a surgical strategy. The LBC picture of ATC usually shows a background of necrotic debris and fibrovascular fragments with small clusters of large round or spindle cells with pleomorphic nuclei and prominent nucleoli. Aggregation of these cells and their large cytoplasms helps to distinguish this neoplasm from a large cell non-Hodgkin malignant lymphoma, which represents, from a clinical and morphologic viewpoint, the most important differential diagnosis. In many cases the cellularity may be exceedingly scant so that the neoplastic cells can be misdiagnosed as histiocytes. Immunostains for cytokeratins may be useful to confirm the epithelial origin of these cells which usually do not express thyroglobulin or TTF-1.

LBC diagnosis of the large cell variant of malignant non-Hodgkin lymphoma, which must be distinguished from ATC (see above), is usually simple. The correct characterization of this tumor, which prevents a surgical approach, relies also on the immunocytochemical expression of LCA, CD20, bcl-6, and other lymphocytic antigens. Insular thyroid carcinoma is a very rare occurrence and shows a solid pattern made up of small thyrocytes with dark nuclei. Small flecks of necrotic debris may sometimes be seen in the background. Metastatic carcinoma to the thyroid gland may occasionally present as a single nodule mimicking a primary tumor. In a background of necrotic debris or hemorrhagic material, clusters of neoplastic cells with features of adenocarcinoma or squamous cell carcinoma are detected. The lung, breast, kidney, large bowel, and larynx are the most common primary sites [56].
**Special Techniques**

**Immunocytochemistry**

Immunohistochemistry was introduced in the early 1970s for specific use directed toward the definition of the nature of lesions and then became an invaluable diagnostic tool in both surgical pathology and cytopathology. The antibodies utilized include those directed at cytoplasmic and membrane-bound antigens as well as nuclear and matrix antigenic sites.

ICC plays an important role in the differential diagnosis between follicular and C cell-derived neoplasms and in the identification of primary or metastatic thyroid neoplasms (e.g., malignant lymphoma).

One of the most interesting insights into the role of ICC in thyroid neoplasms is the evaluation of the markers of malignancy which may distinguish malignant from benign lesions regardless of the presence of capsular or vascular invasion.

The popularity and usefulness of the technique is reflected by the large number of papers over the last 15 years. The majority of this information is synthesized in a recent paper by Colasacco et al. [57] in which the authors analyzed a total of 100 journal articles dealing with immunocytochemical stains on diagnostic cytology.

Immunocytochemical staining may be carried out on cells stored in preservative solution by preparing an appropriate number of additional slides. The use of ICC on LBC yields excellent results with most immunoreagents in terms of the staining pattern, intensity of the reaction, and the smaller amount of reagent due to the clear background and smaller size of the LBC slide [58].

The quality of the immunocytochemical reaction on thin-layer preparations, as far as morphologic details and purity of background are concerned, is much better than CS.

In routine practice, only the concordance of a panel should be considered in the diagnosis of thyroid nodules, especially in cases of follicular lesions. The use of more than one immunomarker is a further guarantee of a correct diagnostic approach, especially when a concordant result is expressed by the cells [15].

There have been only few experiences reported in the literature dealing with ICC applied to LBC. Leung et al. [59] reported satisfactory results with ICC applied in LBC slides from different body sites, except from lymphoma markers. Dabbs et al. [60] showed that ICC in LBC from a variety of neoplastic and nonneoplastic samples resulted in greater intensity and distribution of proper staining compared with conventional cytology.

The studies of our group have demonstrated the possibility of applying ICC to diagnostic thyroid cytology with excellent results not only for the mutual distinction of malignant neoplasms [15, 56] but also in the differential diagnosis between benign and malignant FN [20].

However, based on the data in the literature, no 'magic single marker' may unmistakably distinguish benign from malignant follicular-patterned neoplasms [61].

In thyroid literature, HBME-1, Galectin-3, and RET proto-oncogene have shown the best specificity and sensitivity in discriminating benign differentiated tumors from malignant ones. These data emerged from one of the papers of our group in which the combination of nuclear pleomorphism and positivity of the antibody panel resulted in a slight increase in the specificity and diagnostic accuracy of FNAB (75% and 89% respectively), but this improvement became quite significant when oxyphilic neoplasms, showing a distinctive cytologic picture, were excluded.

**Molecular Techniques**

To also contribute to this purpose, recent advances in the molecular genetics of thyroid cancer are being applied for the development of new diagnostic markers for FNA samples in an attempt to differentiate benign thyroid nodules from malignant ones [62–64]. PC, the most common thyroid malignancy, may carry BRAF, RET/PTC, or RAS mutations [65–67]. These mutually exclusive somatic mutations are found in more than 70% of PC, and some of them are associated with more aggressive tumor behavior. Several studies have demonstrated the feasibility of detecting BRAF, RET/PTC, or RAS mutations in thyroid FNA samples and have shown that this may improve the cytologic diagnosis [67–69].

For this reason, a great number of studies have addressed the possible use of molecular tests for FNA samples in order to improve the accuracy of the cytologic diagnosis of thyroid nodules. Some groups have explored the diagnostic role of BRAF mutations with prospective and retrospective studies which evidenced that all BRAF-positive FNA samples yielded PC at histology [70].

A recent paper by Nikiforov et al. [65] emphasized the diagnostic utility of molecular testing for a panel of molecular mutations consisting of BRAF, RAS, RET/PTC, and PAX-PPARɣ in 480 FNA samples from thyroid nodules which were prospectively tested and yielded 32 mutations with 31 malignant surgical diagnoses and only 1 case of follicular adenoma [66].

The possible diagnostic usefulness of molecular markers is reflected in the last guidelines published by the...
American Thyroid Association. These guidelines indicate that the use of molecular markers such as BRAF, RAS, RET/PTC, and PAX8-PPARγ may be considered (although with a low recommendation rate) for patients with indeterminate FNA cytology to help guide their clinical management. In fact, BRAF mutations correlate with aggressive tumor characteristics such as extrathyroidal extension, advanced tumor stage at presentation, and lymph node or distant metastases as in the review by Xing [63, 71–75].

LBC can offer the possibility of molecular testing for common somatic mutations in thyroid FNAB as the nucleic acids are stable in the preservative solution for up to 6 months after sampling. In some cases the material can also be used for the immunocytochemical evaluation of HBME-1 and Galectin-3 expression which represents a helpful investigation in discriminating low and high malignant FN. In this setting, the future possibility of a guideline encompassing the combined use of ICC and molecular tests for supplementing the morphologic diagnosis could be the starting point for a complete preoperative assessment of thyroid lesions.

As shown in our preliminary study the evaluation of BRAF positivity could be of prognostic value for the prediction of nodal metastases in the central neck compartment [76].

Conclusions

The LBC method applied to the diagnosis of thyroid lesions represents, when the cytopathologist is skilled in thyroid cytology and is aware of the morphologic changes in respect to CS, a powerful diagnostic tool. Its use reduces the number of nondiagnostic (both inadequate and artifactual) and indeterminate cases without impairing the ability to detect the distinctive features of carcinoma. Storage of a variable amount of well-preserved cells allows the application of immunocytochemical and molecular techniques which dramatically improve the efficacy of the morphologic diagnosis.

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