Title page

Title of review
The cytomorphological spectrum of papillary lesions in the breast

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Abstract
The objective of this review is to illustrate the broad spectrum of papillary lesions that can be found in breast fine needle aspirations (FNAC). Papillary tumors of the breast comprise lesions of variable morphology and include entities ranging from benign to high grade malignant. Features of papillary neoplasms invariably describe branching three-dimensional papillary clusters with delicate fibrovascular cords. Cytomorphological criteria for benign and low-grade malignant entities overlap and a definite cytological diagnosis is not always possible. Cellular papillomas may harbor areas of ductal carcinoma in situ (DCIS), atypical ductal hyperplasia (ADH) or lobular neoplasia. DCIS can be both high grade and non-high grade. Immunocytochemistry (ICC) can be helpful, providing there is adequate material for ICC. Relatively high error rates, false negatives as well as false positive diagnoses can be found. In general, papillary lesions may be recognized as such. Reporting strategies will often recommend caution if trying to differentiate benign, cellular papillomas from low-grade carcinomas.

In conclusion, papillary lesions present with a broad spectrum of cytomorphological features that are illustrated in this review.

Introduction
Papillary tumors of the breast comprise lesions of variable morphology and include entities ranging from benign to high grade malignant (table 1). A papillary neoplasm is histologically one that exhibits an arborescent epithelial proliferation with fibrovascular cores and is attached by a stalk to the wall of a dilated duct. They constitute < 2 % of all breast lesions.

Cytomorphological criteria for benign and low-grade malignant entities overlap and a definite cytological diagnosis is not always possible. Features of papillary neoplasms invariably describe branching three-dimensional papillary clusters with delicate fibrovascular cords (1-6). Additional features of papillary carcinomas include moderate to abundant cellular material (7, 8), small papillae arranged in cell balls, tall columnar cells, isolated naked nuclei and hemosiderin laden macrophages (8), irregular groups of predominantly monolayered (two-dimensional) epithelium composed of small, polygonal or cuboidal cells with eosinophilic cytoplasm and rounded, eccentrically placed nuclei (9-24). Features indicating a benign papillary lesion include less cell material, papillae with cohesive stalks surrounded by columnar cells in a honeycomb pattern, apocrine metaplasia, bipolar naked nuclei as well as fewer small papillae and isolated columnar cells (8). Myoepithelial cells within clusters and inconspicuous naked, bipolar nuclei were background also indicate a benign lesion (25). Tse et al report considerable overlaps in the criteria (22) and no demonstrable quantitative differences between papilloma and papillary carcinoma (PC), but with a qualitatively higher degree of atypia in PC and more elaborate and slender papillary fragments. Haji et al found loose cohesive clusters and acinar formation of neoplastic cells to be significant parameters differentiating PC from benign papillomas (26).

Reported accuracy rates in the literature are given as 27 % - 88 % for all papillary lesions (12, 22, 27, 28), 91 % for PC alone (21). Prathiba et al and Jayaram et al reported a sensitivity of 42% and 73%, respectively (12, 19) and a specificity of 88.3 % (12). Relatively high error rates, false negatives (FN) as well as false positive (FP)
diagnoses are reported (15, 19, 21, 22).
Not all lesions diagnosed as papillary on cytology are papillary on histology, the most common error being fibroadenomas and fibrocystic changes (12, 21, 28, 29), but in general, papillary lesions can be diagnosed as such (9, 12, 13, 23, 28).
The known and reported criteria of papillary lesions are almost exclusively based on conventional smears. Many institutions now use liquid based preparations (LBC) alone or as an adjunct to traditional smears. Several features may change and appear different in LBC and could potentially cause both FN and FP diagnoses (29).
The aim of this review is to illustrate the broad spectrum of papillary lesions that can be found in breast FNAC.

**Benign or borderline papillary lesions**

**Benign intraductal papillomas** make up approximately 5% of benign breast lesions (30). The mean age at presentation is 48 yrs, but it commonly also presents in the 6th and 7th decades. Most of them are located centrally and they may present with unilateral bloody or serous-bloody nipple discharge. They are often non-palpable with a size range of a few mm up to > 5 cm. Peripheral lesions are often clinically occult, but may cause nipple discharge and eventually a mass as a result of a small cluster of papillomas. They may be mammographically occult and present only as microcalcifications. Retroareolar lesions appear radiologically as a circumscribed benign mass, a solitary dilated duct and rarely as microcalcifications. Ultrasound might reveal a well-defined smooth-walled cystic nodule with solid components. Benign, intraductal papillomas have a varied morphology that is reflected in the cytological specimens (table 2, figure 1a-e). The “plain” papillomas have a moderate cellularity, sheets and groups of benign ductal cells, as well as macrophages and often also apocrine cells. These lesions are cytologically completely benign. Traditionally they are surgically excised, but some argue that it is not mandatory (27, 31). Papeix et al found that 34/36 cases diagnosed as benign with papillary features were indeed benign papillomas and the two remaining cases were stable on radiological follow up. Proliferative lesions tend to be cellular and with a polymorph population consisting of several cell types (benign/usual intraductal hyperplasia = UDH), all with basically benign nuclear criteria (figure 2 a-f). They may harbor areas of ductal carcinoma in situ (DCIS), atypical ductal hyperplasia (ADH) or lobular neoplasia. DCIS can be both high grade and non-high grade. Cells from high grade DCIS are clearly recognized as malignant (32), whereas aggregates and single cells from a non-high grade lesions may not be definitely identified in the smears. In many cases though, the atypical cells may not present in the specimens. Immunocytochemistry can be helpful, providing there is adequate material for ICC (table 3). These cellular and heterogeneous papillary lesions should be diagnosed as equivocal/atypical and surgically excised.

**Florid papillomatosis of the nipple (subareolar papillomatosis)** is a benign epithelial proliferation localized within and around the collecting ducts of the nipple. It comprises < 1% of breast specimens and has an age range of 20-87 yrs (mean 43 yrs). Approximately 2/3 of the women presents with nipple discharge. Some cases may have skin erosion and mimic Paget’s disease of the nipple clinically (30). The cytologic findings are listed in table 4 and they are illustrated in figure 3 a-g. If palpable, the clinical findings are usually very characteristic and help making a confident diagnosis. The FNAC material is moderate to abundant and can be prepared
both as smears, as LBC and as cell block (figure 4 a-e). ICC phenotype (table 2) reveal a benign, polymorph cell composition and supports the diagnosis of a benign lesion.

**Papillary carcinoma variants**

**Intraductal papillary carcinoma in situ** may present with a clear or blood stained nipple discharge. More peripheral lesions may present as a mass. Radiologically it can present both as a single, round and partly cystic lesion as well as micro-calculifications (as part of a DCIS) without a tumor (30). In histology, we find ducts or TDLU (terminal duct-lobular unit) with slender branching fibro-vascular stalks covered by a single cell population of neoplastic cells (Figure 5 a-c). Micropapillary, cribriform and solid growth patterns are also seen. The papillae are lined by neoplastic, columnar cells in one or several layers. The cells are deceptively bland with low - grade atypia (8 figures). The cytological findings mirror the histology and are detailed in table 5.

**Encapsulated papillary carcinoma** (30) is a variant of papillary carcinoma characterized by fine fibrovascular cores covered by neoplastic cells of low or intermediate grade and surrounded by a thick fibrous capsule (figure 6a). In the majority of cases there are no myoepithelial cell layer within the papillae or at the periphery of the lesion. There is a circumscribed round mass on mammography and ultrasound imaging. A frank invasive component is usually IDC and cannot be appreciated in the cytological material. Nipple discharge may be present. Cytologically the features (Figure 6b-g) are the same as the intraductal papillary carcinoma (table 5), and the two entities can only be distinguished histologically.

**Solid papillary carcinoma** (30) represents < 1 % of breast carcinomas. It occurs usually in menopausal women with a mean in the seventh decade. About 20-25 % have a bloody discharge. Mammography may show an abnormality or lobular lesion. Ultrasound may also show a lobulated mass that histologically (Figure 7a,b) corresponds to closely apposed, expansile nodules with delicate fibrovascular cores within the nodules. These tumors often reveal neuroendocrine differentiation and/or mucinous components. Conventional invasive growth may be present, often having mucinous or neuroendocrine features (7, 33). The cytological features resemble the two above subtypes of papillary carcinomas (table 5) (Figure 7c-h). The cystic component is virtually absent, and debris and macrophages are not present. The features and findings of the epithelial cells, though, are virtually the same as the two above. Images illustrating the findings are seen in figures z-u. The cytological criteria are those of a low - grade carcinoma (7). The presence of an eventual additionally invasive component cannot be evaluated in these three lesions. Primary surgery should then be complete excision, but should not include removal of the sentinel node (SN) or extensive surgery.

An overview and comparison of cytological features and immunocytochemical findings in benign and malignant papillary lesions is given in tables 3 and 6.

**Invasive micropapillary carcinomas** (30) are composed of small, hollow and morula-like clusters of cancer cells and surrounded by clear stromal places (figure 8a). There is usually a reversed polarity, an “in side out” growth pattern, whereby the apical pole of the cells faces the stroma and not the luminal surface. They make up 0.9-2 % of invasive breast cancers (4), but up to 7.4 % may show invasive breast cancers may have partial micropapillary growth pattern (5). The mean age is as in IDC, 75 % are grade 2 and 3, estrogen and progesterone receptors (ER/PgR) are usually positive, whereas HER-2 is variable, but often positive. They usually present
as a palpable mass. Cytologically these present as high grade carcinoma cells in well
demarcated, angulated micropapillary groups lacking a fibrovascular core with
irregular crowded nuclei (Figure 8b-d) and are clearly malignant (34-36).
According to the WHO (world health organization) definition (30), **invasive
carcinoma** is a tumor with predominantly papillary growth pattern (> 90
%) in the invasive component. It is a rare tumor with no known clinical characteristics
or known epidemiological data. The tumor resembles papillary carcinomas from other
organ sites, in particular ovary, tube and lung that are the main differential diagnoses.
The cytological features have not been described.

There are no conflicts of interest.

Table 1. Papillary lesions of the breast
- Intraductal papilloma
  - Intraductal papilloma NOS (not otherwise specified) with or without
    benign hyperplasia
  - Intraductal papilloma with atypical ductal hyperplasia
  - Intraductal papilloma with ductal carcinoma in situ
  - Intraductal papilloma with lobular neoplasia
- (florid) (subareolar) papillomatosis of the nipple
- Intraductal papillary carcinoma in situ
- Encapsulated papillary carcinoma
- Solid, papillary carcinoma
- Invasive papillary carcinoma
- Invasive micropapillary carcinoma

Table 2. Cytological findings in (benign) intraductal papillomas:
- Variable cellularity with a basic benign pattern
- Epithelial components:
  - Cohesive papillary clusters
  - (Micro-) papillary clusters, bipolar cells
  - Complex, folded, monolayer or three dimensional epithelial aggregates
  - Small groups and monolayer sheets
  - Benign nuclei or nuclei showing moderate to distinct cellular/nuclear
    pleomorphism but with a fine chromatin pattern
  - Nucleoli may be distinct
  - Hyperplasia with and without atypia showing a mixed cell population,
    both benign and irregular/atypical with threedimensional aggregates
    that may resemble ADH/low grade DCIS (solid and/or cribriform)
  - Usually the epithelial fragments are rather cohesive
  - a (usually small) population of single cells is not uncommon
  - Myoepithelial cells/nuclei
  - Apocrine cells
- Stromal fragments, partly papillary with vascular core
- Debris and macrophages
Table 3. Overview of papillary lesions and their immunophenotypes

<table>
<thead>
<tr>
<th>Tumor type</th>
<th>Benign/ malignant</th>
<th>p63</th>
<th>High molecular weight (HMW) cytokeratins CK5/6, CK14</th>
<th>ER/PgR</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intraductal papilloma</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1. NOS</td>
<td>benign</td>
<td>positive</td>
<td>positive in myoepithelial cells</td>
<td>patchy positive</td>
</tr>
<tr>
<td>2. with benign hyperplasia (UDH= usual ductal hyperplasia))</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3. with atypical ductal hyperplasia (ADH)</td>
<td>borderline</td>
<td>mixed</td>
<td>mixed</td>
<td>Mixed patchy and uniform</td>
</tr>
<tr>
<td>4. with lobular neoplasia (ALH (atypical lobular hyperplasia) + LCIS (lobular carcinoma in situ))</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>5. with ductal carcinoma in situ (DCIS)</td>
<td>in situ</td>
<td>negative</td>
<td>negative</td>
<td>uniform and 100 % positive</td>
</tr>
<tr>
<td>Subareolar papillomatose</td>
<td>benign</td>
<td>positive</td>
<td>positive in myoepithelial cells</td>
<td>patchy positivity</td>
</tr>
<tr>
<td>Intraductal papillary carcinoma</td>
<td>in situ; usually low-grade with occasionally high grade lesions</td>
<td>negative in papillary areas; positive in duct periphery</td>
<td>negative</td>
<td>uniform and 100 % positive</td>
</tr>
<tr>
<td>“encapsulated”/ intracystic papillary carcinoma</td>
<td>In situ or invasive, low grade</td>
<td>negative</td>
<td>negative</td>
<td>uniform and 100 % positive</td>
</tr>
<tr>
<td>Solid papillary carcinoma</td>
<td>Invasive or invasive, low grade</td>
<td>negative</td>
<td>negative</td>
<td>uniform and 100 %</td>
</tr>
<tr>
<td>invasive papillary carcinoma</td>
<td>invasive</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>invasive micropapillary carcinoma</td>
<td>invasive, high grade</td>
<td>negative</td>
<td>negative</td>
<td>usually positive</td>
</tr>
</tbody>
</table>

Table 4. Cytologic findings in subareolar papillomatosis
- Moderate or high cellularity with a basic benign pattern
- Small amount of debris, inflammatory cells and siderophages may be found
- Aggregates and smaller epithelial groups
- Basically cohesive, irregular aggregates
- Irregular shapes
- Micropapillary, macrophages, naked nuclei
- Uniform nuclei with finely distributed chromatin and small nucleoli
- Little anisonucleosis, occasional hyperchromatic nuclei possible
- Occasional) dispersed epithelial cells
- Apocrine cells may be present

Table 5. Cytological features of papillary carcinomas (intraductal, encapsulated and solid)
- May be cystic on aspiration
- Cell material is usually abundant
- Epithelial cells are monotonous and appear “monoclonal” showing low grade nuclear/cell atypia (usually G1)
- Anisonucleosis, hyperchromasia, coarse chromatin and prominent nucleoli are uncommon (occasionally G2-G3).
- Benign bipolar cells are absent from the background, myoepithelial cells are not seen within the groups or are very scarce
- Large papillary cell clusters forming arborizing arrays bearing overlapping, palisaded cells on a fibrovascular core may be present (as in papilloma)
- Cells may be dispersed and the fibrovascular cores denuded
- (pseudo)-papillary arrangement of cells (in the solid-papillary type)
- micropapillary as well as irregular small epithelial cell groups
- The epithelial cells are often distinctly columnar in appearance
- Evaluation of invasive component not possible
- Microcalcifications are common findings
- Intracytoplasmic vacuoles are not rare
- neuroendocrine differentiation common in solid-papillary type
- Often extensive dissociation in single cells, plasmacytoid
Table 6. Morphologic - diagnostic considerations cellular (benign) papillary lesions versus papillary carcinoma

<table>
<thead>
<tr>
<th>(Benign) cellular papillary lesion</th>
<th>Papillary carcinoma (in situ and/or invasive)</th>
</tr>
</thead>
<tbody>
<tr>
<td>heterogeneous (&quot;polyclonal&quot;) cell population</td>
<td>monomorphous (&quot;monoclonal&quot;) cell population</td>
</tr>
<tr>
<td>basic benign pattern, but may have a population of cells showing low grade nuclear atypia/anisonucleosis</td>
<td>low grade nuclear atypia in the vast majority of cases</td>
</tr>
<tr>
<td>straight or curved tubular structures representing adenosis in papilloma</td>
<td></td>
</tr>
<tr>
<td>mostly cohesive, but with single cells</td>
<td>more discohesive, often extensive</td>
</tr>
<tr>
<td>threedimensional cells aggregates representing/resembling low grade DCIS, solid and/or cribriform</td>
<td>threedimensional cells aggregates representing/resembling low grade DCIS, solid and/or cribriform</td>
</tr>
<tr>
<td>papillary and micropapillary groups</td>
<td>papillary and micropapillary groups</td>
</tr>
<tr>
<td>fibrovascular stalks</td>
<td>fibrovascular stalks</td>
</tr>
<tr>
<td>debris and macrophages</td>
<td>debris and macrophages</td>
</tr>
</tbody>
</table>

References


Figure legends

Figure 1. Benign papilloma

a. Magnification x 50, PAP (Papanicolaou). Large and partly folded sheets of benign ductal cells in benign papilloma
b. Magnification x 200, MGG (May-Grunwald Giemsa). Monolayer sheet benign ductal cells. In the background some cystic debris and macrophages
c. Magnification x 200, PAP, SurePath® (Becton and Dickinson). Smaller groups and a short strip, debris, macrophages.
d. Magnification x 400, PAP, SurePath. Micropapillary group with benign nuclear characteristics
e. Magnification x 400,PAP, ThinPrep® (Hologic). A distinct single cell population, but benign nuclear features, especially the fine, uniform chromatin structure.

Figure 2. Cellular, benign papilloma
a. Magnification x 100, MGG. Cellular material with stromal fragment, irregular sheets, groups and aggregates of ductal cells as well as a single cell population.

b. Magnification x 200, MGG. Smaller groups and sheets, mixed ductal and apocrine as well as numerous macrophages. In addition a minor single cell population.

c. Magnification x 200, MGG. Cellular papilloma with three-dimensional epithelial clusters and metachromatic basement membrane-like globules

d. Magnification x 200, MGG. Complex epithelial sheets with transition from ductal to apocrine cell type.

e. Magnification x 400, MGG. Cellular polymorphism in a papilloma with extensive, but histologically benign hyperplasia. Nuclear irregularities, but uniform chromatin structure.

f. Magnification x 200, HE (Hematoxylin Eosin). Proliferative papilloma with adenosis; same lesion as 2e.

Figure 3. Subareolar papillomatosis

a. Magnification x 50; HE, cell block. FNAC material from a cellular papillary lesion located in the nipple.

b. Magnification x 50, MGG. Abundant cellularity with large, complex and folding epithelial aggregates

c. Magnification x 100, MGG. Complex cohesive aggregates

d. Magnification x 100, MGG. Complex aggregate with part of the stromal papillary stalk and a minor component of single cells

e. Magnification x 100, MGG. Irregular, medium sized aggregates, both ductal and apocrine.

f. Magnification x 200, MGG. Micropapillary structures

g. Magnification x 400, MGG. Very regular, benign nuclei with uniform chromatin structure

Figure 4. Subareolar papillomatosis immunophenotype

a. Magnification x 100, cell block stained with p63

b. Magnification x 100, cell block stained with CK 5/6

c. Magnification x 100, LBC (ThinPrep) with cytospin preparation stained with p63

d. Magnification x 100, LBC (ThinPrep) with cytospin preparation stained with CK 5/6

Figure 5. Intraductal papillary carcinoma (papillary DCIS)

a. Magnification x 100, HE. Slender papillary structures lined by one layer of epithelial cells with low-grade nuclear atypia and delicate fibrovascular stalks

b. Magnification x 50, MGG. Overview of smear with large papillary aggregate and numerous cell groups of varying sizes

c. Magnification x 100, MGG. On larger magnification the branching capillaries are seen
d. Magnification x 200, MGG. A smaller papillary fragment with central capillary surrounded by low-grade atypical cells. There is mild to moderate nuclear pleomorphism and scattered single cells.
e. Magnification x 400, MGG. On larger magnification a fine to somewhat coarse chromatin is seen and inconspicuous nucleoli.
f. Magnification x 400, MGG. Micropapillary epithelial cell group with low-grade nuclear atypia.
g. Magnification x 400, MGG. Smaller epithelial cell groups with low-grade nuclear atypia.
h. Magnification x 200, HE. Micropapillary epithelial proliferations with varying nuclear atypia, up to high grade (G1-3).
i. Magnification x 200, HE. Papillary epithelial proliferation with high grade atypical nuclei (G3).
j. Magnification x 400, MGG. Micropapillary DCIS G1-3 with proteineous debris, macrophages and micropapillary epithelial cell groups. One of the micropapillary groups shows a low-grade nuclear atypia whereas the rest are high grade with prominent pleomorphism.
k. Magnification x 400, PAP. Loosely cohesive sheet with high grade malignant cells: coarse irregularities in the nuclear contour, irregular peripheral chromatin brim, coarse nuclear chromatin with clearing and prominent nucleoli. Cytoplasmic vacuolization is seen in some of the cells.

Figure 6. Encapsulated, papillary carcinoma
a. Magnification x 10, HE. Well demarcated papillary carcinoma (the two central nodules). In addition an ordinary invasive ductal carcinoma to the right and above.
b. Magnification x 50, MGG. Overview with large, branching papillary aggregate as well as smaller groups.
c. Magnification x 100, MGG. Monolayer (two-dimensional) sheets as well as a complex partly three-dimensional and partly monolayer aggregate. In addition we can see numerous single cells and some “strips”
d. Magnification x 100, MGG. Numerous micropapillary cell groups.
e. Magnification x 200, MGG. The cells groups are loosely cohesive with numerous single cells. There is a mild to moderate pleomorphism.
f. Magnification x 400, MGG. A loosely cohesive sheet with nuclei in the size range of 2-3 X RBC.
g. Magnification x 630, PAP, ThinPrep. Small epithelial groups with low-grade nuclear atypia. The nuclear chromatin is finely granular with accentuated peripheral chromatin brim and there are small to medium sized nucleoli.

Figure 7. Solid-papillary carcinoma
a. Magnification x 50, HE. Solid growing carcinoma with a well demarcated border.
b. Magnification x 200, HE. Low-grade nuclear atypia and intratumoral branching vessels.
c. Magnification x 100, MGG. Abundant cell material with large, pseudopapillary and three-dimensional epithelial aggregates, smaller groups and a large single cell population.
d. Magnification x 100, MGG. Loosely cohesive cells with abundant single cells

e. Magnification x 100, PAP. Large, three-dimensional aggregates, loosely cohesive cells

f. Magnification x 400, MGG. Loosely to non-cohesive, low-grade carcinoma cells with plasmacytoid appearance. Microcalcification (arrow).

g. Magnification x 400, PAP, SurePath. Mainly single cell population of plasmacytoid cells with delicate, finely granular chromatin and small, but visible nucleoli

h. Magnification x 400, PAP. Abundant, loosely cohesive and plasmacytoid cells. The chromatin if fine granular, but more distinct than in the SurePath preparation. Nucleoli are small, but distinct.

Figure 8. Invasive, micropapillary carcinoma

a. Magnification x 400, HE. Micropapillary carcinoma groups with high grade nuclear atypia (G3)

b. Magnification x 100, MGG. Numerous micropapillary epithelial cell groups

c. Magnification x 100, PAP. Numerous micropapillary epithelial cell groups

d. Magnification x 400, PAP. High grade nuclear atypia with coeaearse chromatin, distinct nuclear size variation and in conspicuous nucleoli

e. Magnification x 400, MGG. High grade atypical cells in micropapillary arrangement; clearly irregular nuclear contour and irregular chromatin. Nuclear size is 4-5x RBC.