Review

Cribriform Patterned Lesions in the Prostate Gland with Emphasis on Differential Diagnosis and Clinical Significance

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Simple Summary: A cribriform structure is defined as a continuous proliferation of cells with intermingled lumina. Various entities may have a cribriform morphology within the prostate gland, ranging from normal, to benign, to borderline and even to malignant lesions. This review summarizes the morphologic features of entities that have a cribriform morphology within the prostate gland, with an emphasis on their differential diagnosis, molecular profile and clinical significance. The basic aim is to assist the pathologist with challenging and controversial cases and inform the clinician on the clinical implications of cribriform morphology.

Abstract: Cribriform glandular formations are characterized by a continuous proliferation of cells with intermingled lumina and can constitute a major or minor part of physiologic (normal central zone glands), benign (clear cell cribriform hyperplasia and basal cell hyperplasia), premalignant (high-grade prostatic intraepithelial neoplasia), borderline (atypical intraductal cribriform proliferation) or clearly malignant (intraductal, acinar, ductal and basal cell carcinoma) lesions. Each displays a different clinical course and variability in clinical management and prognosis. The aim of this review is to summarize the current knowledge regarding the morphological features, differential diagnosis, molecular profile and clinical significance of the cribriform-patterned entities of the prostate gland. Areas of controversy regarding their management, i.e., the grading of Intaductal Carcinoma, will also be discussed. Understanding the distinct nature of each cribriform lesion leads to the correct diagnosis and ensures accuracy in clinical decision-making, prognosis prediction and personalized risk stratification of patients.

Keywords: prostate cancer; intraductal carcinoma; cribriform carcinoma; Gleason Score; prognostic grade group; prognosis

1. Introduction

Glandular structures within the prostate may assume a cribriform morphology as a major or minor part of physiological, non-neoplastic and neoplastic processes. Cribriform is derived from the Latin word *cribrum*, which means sieve and is used to describe something that is pierced with small holes. In histology, cribriform is defined as a continuous proliferation of cells with intermingled lumina. That means that there are cells lying one next to another with round or elongated holes interspaced among them at various intervals. Normal prostate glands and benign (clear cell cribriform hyperplasia and basal cell hyperplasia), premalignant (high-grade prostatic intraepithelial neoplasia), borderline (atypical intraductal cribriform proliferation) or clearly malignant (intraductal, acinar, ductal and basal cell carcinoma) lesions can present with a cribriform morphology. These entities display a different clinical course and variability in clinical management and prognosis. Thus, cribriform morphology can cause a diagnostic challenge for the pathologist. In addition,
even though cribriform morphology in benign entities is probably the result of cells with glandular differentiation pilling up within a pre-existing space (duct), in malignant entities, it may be a purposeful architectural pattern that offers a survival advantage to cancer cells, as it has been associated with adverse prognosis and therapy resistance.

Overall, the aim of this review is to summarize the current knowledge regarding the morphological features of cribriform formations within the prostate gland, their differential diagnosis and the utility of immunohistochemical markers (i.e., PTEN loss and ERG expression) in establishing an accurate diagnosis. Areas of controversy regarding the clinical significance and/or management of cribriform entities will also be discussed. In Table 1, the most important morphologic and molecular features of these entities are summarized.

### Table 1. Summary of differential diagnosis of entities with cribriform morphology within the prostate gland.

| Entity                        | Architecture                                                                 | Cytologic Features                                                                 | Basal Cell Layer | ERG Expression | PTEN Loss |
|-------------------------------|-----------------------------------------------------------------------------|-----------------------------------------------------------------------------------|------------------|----------------|-----------|
| Benign cribriform glands      | Complex epithelium with cribriform pattern and epithelial bridges located in the central zone | High columnar stratified epithelium, granular cytoplasm, small round nuclei without cytologic atypia or prominent nucleoli | Intact           | −             | −         |
| Basal cell hyperplasia        | Nodular lesion, within the transitional zone                                  | Scant cytoplasm, hyperchromatic nuclei without cytologic atypia                    | The lesion involves basal cells | −             | −         |
| Clear cell cribriform hyperplasia | Variant of BPH, medium and large sized acini with cribriform morphology       | Pale to clear cytoplasm, nuclei lack cytologic atypia or prominent nucleoli         | Intact           | −             | −         |
| HGPIN                         | Normal-sized acini with tufting, micropapillary, or flat growth pattern and without expansion of glands | Cytologic atypia, nuclear enlargement and hyperchromasia with prominent nucleoli, no necrosis | Preserved (can be fragmented) | −/+          | −         |
| AIDCP                         | Loose cribriform lumen-spanning architecture                                | Moderate nuclear atypia, absence of necrosis, insufficient to meet the criteria for IDC | Preserved (can be fragmented) | +/−          | Identified |
| Intraductal carcinoma         | Greatly expanded glands with cribriform/solid growth                         | Nuclear atypia (nuclear enlargement, hyperchromatic nuclei) may be present         | Preserved (can be fragmented) | +/−          | Identified |
| Invasive cribriform acinar carcinoma | Continuous prolifération of cells with intermingled lumina                  | Nuclear atypia (prominent nucleoli, hyperchromasia)                               | Absent           | +/−          | Identified |
| Ductal carcinoma              | Papillary, solid and cribriform growth pattern                              | Tall columnar cells, nuclear atypia, mitotic figures                               | Usually absent, may be present        | −/+          | Identified |
| Basal cell carcinoma          | Irregular cribriform formations containing mucin or basement membrane-like material within the lumina, desmoplastic reaction | Hyperchromatic large nuclei with scant cytoplasm                                  | The entire process involves basal cells | −           | Identified |

### 2. Benign Cribriform Formations

The cribriform pattern is commonly observed in normal tissue and benign processes of the prostate gland. Although these entities can become potential mimickers of atypical
and even malignant lesions, under no circumstances must they be confused with cancer, as they have no significant clinical implications. Their features are discussed below.

2. Benign Cribriform Formations

The cribriform pattern is commonly observed in normal tissue and benign processes as they have no significant clinical implications. Their features are discussed below.

2.1. Normal Cribriform Glands

The epithelial tissue of the central zone of the prostate gland near its base can form cribriform glandular structures [1]. These structures are composed of complex papillary epithelium with cribriform morphology, Roman Arches and epithelial bridges [1,2] lined by columnar, pseudostratified epithelial cells with granular eosinophilic cytoplasm and a small round nucleus. Of note, the cytologic atypia or prominent nucleoli are consistently absent. In addition, the presence of an intact and frequently prominent basal cell layer is characteristic and can help to identify the benign nature of the lesion (Figure 1a). This totally normal cribriform morphology has been well understood and is easily distinguishable from other cribriform lesions of the prostate gland—albeit, its differential diagnosis may be more challenging in biopsies.

2.2. Basal Cell Hyperplasia

Basal cell hyperplasia constitutes a focal, nodular process composed of an expansion of basal cells within the acini of prostate gland [3]. Small clusters of proliferating round basal cells that are two or more layers in thickness are formed within a compressed stroma. These cell clusters may grow in a solid, cystically dilated or cribriform pattern [3,4]. The cribriform architectural pattern is composed of irregular round luminal spaces. Cytologically, basal cells have scant cytoplasm with hyperchromatic nuclei and lack cytologic atypia (Figure 1b). Immunohistochemically, the cells in basal cell hyperplasia are positive for basal cell markers (p63, HMWCK) and negative for AMACR [3]. Basal cell hyperplasia is not associated with an adverse prognosis or a higher risk of prostate cancer.

Figure 1. Benign cribriform formations. (a) Glands in the base of the prostate with luminal tufting (upper part) and nuclear pseudostratification. A cribriform architectural pattern is observed, but lack of cytologic atypia and presence of basal cell layer excludes malignancy (×100). (b) Basal cell hyperplasia with a cribriform pattern (×100). (c) Hyperplastic cribriform gland lacking cytologic atypia and displaying abundant pale to clear cytoplasm similar to acinar cells of adjacent prostatic gland (right side) (×200). (d) Clear cell cribriform hyperplasia with bland monotonous nuclei (×200) (scale bar is 50 μm).
2.3. Clear Cell Cribriform Hyperplasia

Clear cell cribriform hyperplasia represents a rare variant in the histologic spectrum of benign nodular prostatic hyperplasia (BPH), is typically located in the transitional zone and is composed of a nodular cluster of medium- and large-sized acini with cribriform morphology. The hyperplastic epithelial cells have a pale-to-clear granular cytoplasm, and their nuclei lack cytologic atypia or prominent nucleoli. A clearly visible basal cell layer is always present (Figure 1c,d). Cribriform hyperplasia does not constitute a risk factor for prostate cancer and the treatment is the same as in BPH.

As far as a differential diagnosis is concerned, the normal and benign cribriform formations described above may be confused with cribriform adenocarcinoma, intraductal carcinoma (IDC) and high-grade prostatic intraepithelial neoplasia (HGPIN), especially when a prostate needle biopsy specimen is evaluated. In this case, the benign cytologic features (absence of prominent nucleoli and lack of cytologic atypia) and the presence of a prominent and continuous basal cell layer support the diagnosis of a benign entity. In addition, the similarity in cellular morphology between the cribriform structure in question and adjacent normal acini is helpful to confirm the non-malignant nature of these cribriform glands.

3. Premalignant Cribriform Lesions

High-Grade Prostatic Intraepithelial Neoplasia (HGPIN)

HGPIN is regarded as a precursor lesion to invasive adenocarcinoma. This lesion shares phenotypic and genetic alterations with invasive carcinoma but lacks invasion into the fibromuscular stroma [5]. In HGPIN, luminal epithelial growth takes place within ducts/glands of normal size without expansion of their lumina. This lack of expansion of the involved glandular spaces is contrasted to intraductal carcinoma which typically involves expanded acini and ducts. Cytologic atypia is present and includes nuclear enlargement, hyperchromatic nuclei, prominent nucleoli, slightly amphophilic cytoplasm, and nuclear stratification. Necrosis is absent and the basal cell layer is preserved, although it can be fragmented (Figure 2a). Four main growth patterns were initially described in HGPIN: tufting, micropapillary, cribriform, and flat [6]. However, the cribriform pattern is no longer considered acceptable in HGPIN. Lesions with a loose cribriform pattern and no necrosis or significant pleomorphism are considered atypical cribriform lesions (see below), whereas lesions with dense cribriform pattern and/or necrosis or nuclear enlargement/pleomorphism are diagnosed as IDC.

Figure 2. Preneoplastic and borderline neoplastic cribriform lesions. (a) Multiple foci of high-grade PIN (×40). (b) Atypical intraductal cribriform proliferation in a biopsy. Proliferation is present in <50% of the gland, located at the edge of the core and not accompanied by atypia or necrosis (×100). (c) Atypical intraductal cribriform proliferation. An expanded gland with a loose cribriform architecture is observed. Immunohistochemical expression of the marker ERG is suggestive of the malignant nature of neoplastic cells but is not enough to change the diagnosis to IDC (×100) (scale bar is 50 µm).
The distinction of HGPIN from IDC is particularly important, especially in biopsies, as the two entities have different clinical implications and a different subsequent approach is followed for each of them [7,8]. The distinction is usually straightforward because, as mentioned above, dense cribriform architecture, nuclear pleomorphism and necrosis are not features that are consistent with the diagnosis of HGPIN. Gland expansion, the involvement of >6 glands, presence of an irregular contour and identification of a biphasic cell population favor IDC [8,9]. In cases where it is difficult to distinguish HGPIN from IDC morphologically (i.e., in flat lesions with moderate nucleomegaly), immunohistochemistry can be of use; ERG expression and PTEN loss is observed in IDC, whereas ERG is less frequently expressed and PTEN expression is preserved in HGPIN [7,10–12].

4. Cribriform Lesions with Borderline Clinical Significance

Atypical Intraductal Cribriform Proliferation (AIDCP)

Atypical intraductal cribriform proliferation (AIDCP) is a relatively new term that describes a cribriform lesion with a phenotype more atypical than HGPIN but without meeting all the diagnostic features of IDC. The characteristics of AIDCP on needle biopsies include (a) loose cribriform lumen-spanning architecture beyond that of HGPIN, but lacking significant nuclear pleomorphism or necrosis to meet the criteria for IDC; (b) atypical nuclei with significant pleomorphism but insufficient for a diagnosis of IDC; and/or (c) dense cribriform or solid proliferation of atypical cells partially present in large ducts on the edge of core biopsy specimens (Figure 2b,c) [10]. Basal cells are retained in AIDCP.

The differential diagnosis of AIDCP from HGPIN is usually straightforward, as cribriform architecture is no longer permissible in HGPIN. AIDCP is also easily distinguished from invasive cribriform carcinoma by the presence of basal cells in the case of the former. Immunohistochemistry can help to confirm the presence of basal cells in difficult cases.

The most problematic differential diagnosis is its distinction from IDC. Immunohistochemistry for basal cell markers is not helpful in this setting as both entities show a present, albeit in some cases fragmented, basal cell layer. Similarly, PTEN loss and ERG expression are not helpful as AIDCP and IDC have shown a similar pattern of PTEN and ERG expression [10]. The distinction is, thus, based on morphology and the discriminative features are more a matter of quantity, rather than their presence or absence. When architecture is not dense and nuclear atypia is not high enough to make an IDC diagnosis, then the cribriform lesion is called AIDCP. In our experience, AIDCP commonly coexists with IDC in prostatectomy specimens, meaning that adjacent to ducts with typical features of IDC are ducts with an intraductal proliferation that is very similar to, but falls short of, IDC diagnosis. Whether the criteria for IDC need to be loosened in those cases to include all cribriform intraductal proliferations remains to be determined. This, however, has mainly academic interest, as areas with definite IDC are usually present as are areas with invasive carcinoma and the nature (and nomenclature) of the not-typical-for-IDC cribriform formations will not have any significant implications for the patient’s further management.

From a practical standpoint, the diagnosis of AIDCP has important clinical implications in biopsy specimens as, when isolated, this has correlated with an increased probability of the presence of an invasive carcinoma in repeat biopsy [10,13]. For this reason, the presence of isolated AIDP in biopsies should be treated with close surveillance and re-biopsy and should not be considered as HGPIN [7,11,14]—an entity that does not usually justify a biopsy repeat.

5. Malignant Cribriform Formations

5.1. Intraductal Carcinoma (IDC)

IDC is characterized by a proliferation of malignant epithelial cells within preexisting prostatic ducts and acini with architectural and/or cytological atypia that exceeds that of high-grade prostatic intraepithelial neoplasia [15]. Thus, by definition, IDC retains the basal cell layer, although it is well known that some IDC foci can have dispersed basal cells due to gland distention. Acini and ducts are usually greatly expanded, and the cells often exhibit
remarkable nuclear abnormalities such as enlargement and significant hyperchromasias of the nuclei. Major and minor histological criteria have been used for IDC diagnosis. The major criteria include solid or dense cribriform architecture, significantly enlarged nuclei and non-focal comedonecrosis [16]. The dense cribriform architecture is defined as cells involving >50% of the duct lumen, and the enlarged nuclei were originally defined as ≥6 times the size of the normal cells (Figure 3). However, the later has recently been disputed [17,18] and replaced by marked pleomorphism and nucleomegaly [19]. Minor criteria include irregularly shaped glands with right-angle branching, frequent and easily identifiable mitoses and the existence of two cell populations, one mitotically active at the periphery that exhibits pleomorphism and faint PSA expression, and the other involving quiescent cells in the center with strong PSA expression [20].

Figure 3. Intraductal carcinoma. (a) An expanded gland with cribriform architecture, central necrosis and preservation of basal cells (×100). (b) Confluent foci of expanded glands with dense cribriform architecture (×100). (c) Basal cell layer is fragmented (basal cell marker high molecular cytokeratin 34βE12) (×100) (d) ERG expression is observed in the neoplastic cells (×100) (scale bar is 50 μm).

IDC is usually accompanied by adjacent invasive carcinoma both in biopsies [21,22] and prostatectomy specimens [22], with the invasive carcinoma usually, but not always, being high-grade and high-stage [23,24]. IDC is associated with adverse pathologic parameters [25] (i.e., tumors with advanced Gleason Score [26], extra prostatic extension [27], seminal vesicle invasion [21], large volume [26] and presence of lymph node metastasis [28]). In addition, the presence of IDCp has been independently correlated with a reduction in progression-free survival in hormone-naïve [29,30], treated [31] and CRPC patients [26,32]. Thus, its presence should be noted in the pathology report in both needle biopsies and prostatectomy specimens.

Even when the accompanying invasive carcinoma is low-grade, the finding of IDC in biopsies is associated with a high probability of upgrading in the subsequent prostatectomy and disease progression if an active surveillance protocol is used [23]. Thus, IDC in biopsies is considered an adverse finding and active surveillance is not recommended, even when the invasive carcinoma is low-grade [19,33,34]. Rarely, IDC may be the sole finding either in biopsies [15,35] or prostatectomy specimens [24,32,36]. Of note, in the majority, albeit not all, of cases with isolated IDC (i.e., not accompanied by invasive carcinoma) in biopsies, a high-grade, high-stage invasive carcinoma will be found in the
In addition, in a minority of them, metastatic disease will be observed. It has been stated that the presence of an IDC unaccompanied by invasive carcinoma in a prostate biopsy is usually the result of non-sampling a coexisting high-grade invasive component [37]. Thus, the presence of IDC in a needle biopsy should, at the least, lead to an immediate biopsy to search for unsampled high-grade prostatic carcinoma, or even to definitive therapy [16]. The clinical implications of IDC (and other cribriform-patterned atypical/malignant lesions—see below) are summarized in Table 2.

**Table 2.** Clinical implications of malignant cribriform-patterned lesions in pathology reports.

| Cribriform Lesion                                      | Clinical Implication                                                                 |
|--------------------------------------------------------|--------------------------------------------------------------------------------------|
| Atypical intraductal cribriform proliferation (isolated in biopsy) | Increased probability of invasive carcinoma in subsequent biopsy  |
| IDC                                                     | Higher prevalence of homologous DNA repair recombination repair defects Germline testing may be considered |
| IDC with invasive carcinoma PGG1/2 in biopsy            | High probability of upgrading and/or upstaging in subsequent prostatectomy Recommendation for definite treatment (active surveillance not recommended) |
| IDC without invasive carcinoma in biopsy               | High probability of invasive carcinoma of high-grade/-stage in subsequent prostatectomy Recommendation for immediate re-biopsy or definite treatment |
| Cribriform pattern                                      | Higher prevalence of DNA repair mechanisms defects Germline testing may be considered |
| Cribriform pattern in carcinoma PGG2 in biopsy          | Poor prognostic factor Recommendation for definite treatment (active surveillance not recommended) |
| Ductal carcinoma                                        | Higher prevalence of DNA repair mechanisms defects Germline testing may be considered |

Regarding the pathogenesis of IDCp, there are two theories. The first and most prevalent one suggests that IDC results from the spreading of the accompanying invasive high-grade carcinoma into existing non-neoplastic ducts and acini [38]. The adverse pathology and prognosis that IDC is associated with supports the notion that, in the majority of cases, IDC is a late stage of the prostate tumor progression pathway. Indeed, IDC has a genetic profile and genomic instability that is very similar to prostate cancer Gleason 4 and 5 [39] and not lower Gleason Score carcinomas [25]. Loss of \( \text{PTEN} \), CDH1, and BRCA1 and gain of \( \text{MYC} \), features of aggressive invasive carcinomas, are genetic abnormalities shared by IDC too [25,39]. Of note, the presence of IDC is used as an indicator to test patients with PCa for germline mutations of genes of proteins involved in DNA repair through homologous recombination (i.e., \( \text{BRACA2}, \text{BRACA1} \)). The most common chromosomal alteration observed in IDC is the fusion of the \( \text{TMPRSS2–ERG} \) genes, a genetic event that is observed early in the evolution of PCa; when present, it is shared by the adjacent invasive carcinoma [43]. In contrast, when the invasive component is ERG-negative, the associated IDC is also negative. This supports a common clonal origin and indicates that IDC represents an intraductal spread of invasive carcinoma [7,43,44]. The loss of cytoplasmic PTEN expression, a late-onset event occurring in a subclonal population of neoplastic cells during PCa evolution, is another genetic event frequently shared by IDC and the accompanying invasive carcinoma [11,22,25,45,46]. Indeed, previous studies have shown that PTEN-negative HGPIN and IDCp adjacent to invasive cancer are more likely to represent the intraductal spread of malignant cells rather than de novo precancerous lesions [7,43]. The reason why cancer cells make their way inside pre-existing ducts has yet to be explained. Based on the increased prevalence of IDC following therapy and its association with a worse prognosis [31], poor clinical response to therapy [47] and the absence of pathology features of response to therapy in IDC foci [48,49], we may hypothesize that tumor cells find a safe-haven within the pre-existing ducts and away from the tumor stroma; this helps them to better adjust to an ominous microenvironment.
Indeed, experiments with patient-derived xenograft models have shown that cells within IDC are able to withstand castration and regenerate the tumor following testosterone restoration [50]. Additional evidence is needed to support this theory and determine whether this also holds true for IDC in untreated tumors.

The second theory regarding IDC pathogenesis states that, at least in some cases, IDC may represent a precursor lesion. The presence of a few cases of isolated IDC even after full embedding of the prostate supports this theory. IDC may reflect an earlier step in the evolution of the metastatic clone [51] as it has been shown that it is more closely related to the lymph node metastases than the adjacent invasive carcinoma [52], and that IDC and adjacent low-grade carcinoma are not genetically related [24]. The latter is based on the observation that, in 15 cases of IDCp with concurrent low-grade invasive carcinoma analyzed with next-generation sequencing and immunohistochemistry for PTEN and ERG, IDC showed activation in the MAPK/PI3K pathway and a discordant PTEN/ERG status with the adjacent low-grade carcinoma [24]. Thus, in this scenario, IDC may represent a precursor lesion that quickly progresses to a high-grade invasive carcinoma and, as a result, is rarely observed alone.

IDC must be differentiated from high-grade PIN, atypical cribriform lesions and invasive carcinoma. Distinction from HGPIN and AIDCP has been discussed in the previous sections. It is well-established that IDC can be a morphologic mimicker of invasive cribriform carcinoma. The identification of basal cells is the most important feature that distinguishes IDC from invasive PCa. The presence of corpora amylacea and branched architecture supports the diagnosis of IDC and is a helpful clue in cases where basal cells are not readily identified [53]. However, in some cases, a distinction between IDC and invasive cribriform pattern carcinoma cannot be identified with certainty based on H&E alone; therefore, immunohistochemistry with basal cell markers may be used [54]. Of note, the basal cell layer in IDC is often fragmented and may not be present in the plane/level of section. Thus, foci without a basal cell layer are morphologically indistinguishable from adjacent foci that have a basal cell layer, which questions our ability to accurately distinguish IDC from an invasive carcinoma even with the use of immunohistochemistry, especially when performed on a single level [55].

A differential diagnosis of IDC also includes an intraductal spread of the urothelial carcinoma. Neoplastic proliferation is within pre-existing ducts in both cases. The presence of longitudinal nuclear grooves is indicative of the urothelial origin of the cells and may be used as a clue to consider urothelial carcinoma, but is present only in low-grade tumors [56]. A co-existing urothelial carcinoma in situ or papillary urothelial carcinoma in the urethra is also indicative that the neoplastic proliferation within the ducts is of urothelial origin, but these findings may only be appreciated in prostatectomy specimens and, even then, they may not always be present due to epithelial denudation. Immunohistochemistry can help in difficult cases, as IDC expresses PSAP and PSA; whereas urothelial-specific markers (p63, uroplakin, GATA3) are expressed in urothelial carcinomas [8].

Both the International Society of Urological Pathology (ISUP) and the Genitourinary Pathology Society (GUPS), the two international organizations relative to uropathology, agree that IDC is an adverse prognostic factor, that its presence should be noted on the pathology report and that its clinical implications should be commented upon [19,33]. In addition, they both recommend against grading isolated (pure) IDC. However, there are conflicting recommendations regarding the grading of IDC when accompanied by invasive carcinoma and of how to best communicate the adverse clinical implications of IDC to clinicians. ISUP recommends incorporating IDC in Gleason Score/PGG by assigning it a Gleason Grade of 4 (or 5 when comedo necrosis is present) [33], whereas GUPS does not support IDC incorporation in Gleason Score/PGG and only recommends reporting the IDC component as a comment in the pathology report [19]. Table 3 summarizes the current guidelines regarding the grading of IDC and other cribriform-patterned lesions.
Table 3. GUPS and ISUP guidelines regarding the grading of cribriform-patterned lesions.

|                          | GUPS [19]                                      | ISUP [33]                                      |
|--------------------------|------------------------------------------------|------------------------------------------------|
| Isolated IDC             | No grade assigned                              | No grade assigned                              |
| IDC with invasive carcinoma | IDC not included in grading                    | IDC graded as pattern 4 (or 5 when comedonecrosis is present) |
| Comment on IDC clinical significance in pathology report | Recommended                                    | Recommended                                    |
| Cribriform               | Grade as pattern 4 (based on the ISUP 2014 recommendations) [57] | Grade as pattern 4 (based on the ISUP 2005 recommendations) [58] |
| Report the presence of cribriform pattern carcinoma | Yes (biopsies and radical prostatectomies) | Yes (biopsies and radical prostatectomies) |
| Ductal carcinoma         | Grade as pattern 3 if PIN-like ductal [59] and pattern 5 if comedonecrosis present [60] |

There are arguments in favor of both approaches. The distinction of IDC from invasive cribriform carcinoma can be difficult on H&E alone and would require immunohistochemical staining in some cases, creating an extra burden for the pathology laboratory. In addition, as discussed above, the discrimination of IDC from an invasive carcinoma with a cribriform pattern may not be possible even when immunohistochemistry is used. Furthermore, IDC is included in the estimation of tumor extent; thus, it seems logical to include it in PCa grading. Finally, as mentioned above, the presence of IDC is an adverse parameter in patients, even when associated with PGG1 PCa [23]; in addition, excluding IDC from the Gleason Score can underestimate the patient’s risk. In agreement with this, studies have shown that a grading system that incorporates IDC better describes patient prognosis [61,62] and that not only the presence, but also the amount of IDC is associated with adverse prognostic parameters, further supporting its incorporation in grading in a quantitative fashion [54].

On the other hand, IDC is sometimes, albeit rarely, a precursor lesion and has been shown to be a genetically different form of adjacent low-grade carcinoma, indicating that, in these cases, the incorporation of IDC to PGG would, in fact, overestimate the patient’s risk. The distinction of IDC from an invasive carcinoma is not required in all cases; it is only required in cases where a change in PGG will be observed depending on the % of IDC. Studies with routine cases have shown that the inclusion or exclusion of IDC in tumor grading results in a global Grade Group shift in <2% of cases [53,54,63]. However, in a minority of cases, the change in PGG was significant (>2 points) when IDC is incorporated in the Gleason Score [54,63], usually due to the presence of comedonecrosis.

The issue of whether IDC should be included in grading is far from resolved. Even though there are studies that support its incorporation in grading, prospective studies with survival as the endpoint are lacking. In addition, the effect of a grading system where IDC is taken into account may be missed because of the far larger number of cases where the grade is not altered regardless of the presence and amount of IDC. Studies that focus on cases in which the grade would change when the IDC is incorporated and with survival as the endpoint would be informative in terms of which grading system has the most clinical relevance.

5.2. Invasive Acinar Carcinoma with Cribriform Pattern

The acinar adenocarcinoma is the main histologic type of prostate carcinoma, accounting for the majority of all primary carcinomas of the prostate [15]. Prostate cancer grading was originally proposed in 1974 by Donald Gleason [64] and, with important refinements across the years [57,58] and the recent adoption of the Prognostic Grade Grouping nomenclature [65], it remains one of the most important prognostic parameters in PCa. Among the Gleason Grades (GG), GG4 is the most heterogeneous pattern both morphologically and clinically. Architecturally, four growth patterns are recognized within GG4: ill-formed glands, fused glands, and glomeruloid and cribriform patterns. Clinically, numerous studies reported a correlation between the cribriform pattern with adverse pathologic features
and adverse outcomes compared to the other GG4 patterns [66]. In particular, tumors with a cribriform pattern have adverse pathologic parameters more frequently than tumors without this pattern [39,54,67,68] and the patients show earlier PSA recurrence [69–72], more frequent metastasis [72,73] and shorter disease-specific survival [73]. Its clinical significance is best appreciated in GS 7 tumors. Patients with a biopsy of GS 7 PCa without the cribriform pattern have a similar prognosis to those with a GS 6 tumor [39]. In line with the clinical and pathologic observations, unfavorable molecular characteristics are observed in tumors with a cribriform pattern (and intraductal carcinoma) which is similar to the alterations observed in aggressive prostate cancer, including increased genomic instability [74,75], oncogene amplifications, and tumor suppressor genes copy number alterations and point mutations [74]. Thus, the presence of a cribriform pattern in a GS7 tumor on the biopsy is considered a contraindication for active surveillance. Both GUPS and ISUP agree that the cribriform invasive carcinoma should be assigned a Grade 4 (or 5 when comedo necrosis is present) [57]. In addition to being factored in grade, a comment regarding the presence and clinical significance of the cribriform pattern should be included in the report in both biopsies and prostatectomy specimens [19].

Cribriform acinar carcinoma includes both small, rounded cribriform glands characterized by a continuous proliferation of cells with intermingled lumina [57,76,77] and larger irregular cribriform glands (>3 times the size of normal glands). Cytologic features of atypia typical of PCa such as nuclear enlargement, anaplasia and prominent nucleoli are also observed (Figure 4). A basal cell layer is typically lacking. It is also worth mentioning that the cribriform pattern shows a higher inter-observer reproducibility among pathologists in comparison with the other Gleason pattern 4 variants [78].

![Figure 4](image_url)

Figure 4. Invasive cribriform acinar carcinoma. (a) Merging of cribriform structure to clearly invasive carcinoma (×100). (b) Complex cribriform structure invading in the periprostatic adipose tissue verifies the invasive nature of the structure (×100). (c) Absence of staining for basal cell marker verifies the invasive nature of these cribriform glandular structure (Gleason pattern 4) (Immunohistochemistry with the basal cell marker high molecular cytokeratin 34βE12) (×100) (scale bar is 50 μm).

Mimickers of invasive cribriform carcinoma include IDC and ductal carcinoma. The distinction of cribriform pattern carcinoma from IDC is analyzed above. The distinction from ductal carcinoma is reported in the next section. Finally, less frequently, invasive cribriform carcinomas need to be distinguished from urothelial carcinoma involving the prostate, as gland-like lumina may be observed in the nests of urothelial carcinoma. In some instances, the cytologic characteristics of the cells (round cells with nucleoli and limited pleomorphism versus elongated nuclei with grooves) or the presence of an in situ/papillary component versus other patterns of PCa may be indicative of urothelial or prostate carcinoma, respectively. In difficult cases, the immunohistochemistry for prostate and urothelial-specific markers may be performed as mentioned in the IDC section.

The pathogenesis of cribriform pattern carcinoma remains obscure. A molecular profile unique to cribriform patterned invasive carcinoma has been shown by numerous studies (reviewed in [79]). These molecular alterations frequently involve pathways associated...
with aggressive tumor characteristics (i.e., MYC, MAPK, DNA repair, integrin signaling) and may account for the aggressive behavior of the cribriform pattern. However, these alterations may neither be the result nor the consequence of cribriform pattern, as they are not cribriform-specific. One of the most characteristic histologic features of the cribriform pattern in invasive carcinomas is that the majority of the intraglandular cells lack contact with the stroma [80,81]. In contrast, in Gleason Grade 3 and other patterns of Gleason Grade 4 (fused glands, poorly formed glands), most of the tumor cells are in contact with the stroma [81]. This feature leads us to hypothesize that, similar to IDC, tumor cells find a safe haven within cribriform structures, away from the potentially harmful effects of the stroma. In addition, hypoxia that is higher within cribriform formations [75], probably as a result of the limited contact of the cells to the vessel-containing stroma [81], and the effects of a specific type of tumor-associated fibroblast found around cribriform structures that is distinct from fibroblasts associated with other morphologic patterns [82], may account for the aggressive behavior of cribriform-patterned carcinoma.

Numerous studies have shown that tumors with a cribriform (and intraductal carcinoma) pattern have distinct genetic and epigenetic signatures [41,71,79,83] compared to tumors without these aggressive patterns. However, head-to-head comparisons of gene expression between cribriform and non-cribriform areas within the same tumor are very sparse and mostly performed using immunohistochemistry. Proteins reported to be up-regulated in cribriform areas (compared to other GG4 patterns) include EGFR [84] and the cell proliferation marker ki67 [85]. In addition, CD44 downregulation [84] and PTEN [86] loss are more common in cribriform compared to non-cribriform areas. Integrins have also shown distinct patterns of expression in cribriform and non-cribriform areas [87], which is not surprising given that these molecules are the cells’ sensor of the extracellular environment [88]. It has been hypothesized that a prostate cancer ‘nimbus’ (gathering of stormy clouds in Latin) [75], characterized by the gathering of multiple unfavorable events (i.e., hypoxia [75] genetic instability [74,75] and various adverse molecular alterations [75,79]), happens in cribriform carcinoma and probably accounts for its aggressiveness.

5.3. Ductal Carcinoma

Ductal carcinoma is an uncommon and aggressive morphologic type of prostate cancer. It is found in 1–6% of PCa cases, usually mixed with high-grade acinar adenocarcinoma [89] and represents only 1% of prostate carcinomas in its pure form [15,45]. It typically presents as a mass within the prostatic urethra and large periurethral prostatic ducts, although tumors morphologically similar to ductal carcinoma may be observed in the peripheral zone too [90]. When located in the periurethral area, it presents with hematuria and obstructive symptoms.

Clinically, ductal carcinoma is associated with worse pathologic parameters [89] and is an independent indicator of worse biochemical-free [90,91], metastasis-free [92] and overall survival [93] compared to acinar carcinoma. Morphologically, ductal carcinoma may have various patterns, usually more than one within the same case, the most characteristic being a papillary pattern with true fibrovascular cores lined by tall pseudostratified columnar cells (Figure 5) [94]. A glandular, cribriform and solid pattern may also be observed, but in these cases, a distinction from acinar carcinoma is more difficult and relies on the identification of the characteristic cellular features [94]. The neoplastic cells are tall, columnar and pseudostratified, resembling endometrial adenocarcinoma, hence why it was originally thought to be of mullerian origin [95]. Significant nuclear atypia and high mitotic activity are also frequently observed. The cytoplasm is typically amphophilic, although it can be clear. Immunohistochemical staining for basal cell markers may be focally positive [96], as ductal carcinoma is well known for its ability to involve and expand pre-existing glands.
Despite the morphologic and phenotypic differences between ductal and acinar carcinoma, few molecular differences have been depicted between the two entities [97] and are better appreciated when pure forms of ductal adenocarcinoma are examined. Genetic alterations in genes associated with chromatin modification, the PI3K and APC pathway, cell cycle/apoptosis, and transcription regulation, similar to advanced and metastatic acinar prostate cancer, have been shown in ductal carcinoma [98]. In addition, an intrinsic upregulation of androgen-resistance pathways [93] and frequent DNA repair gene aberrations [99,100] have been described in ductal adenocarcinoma and may account for its aggressive clinical behavior.

To account for its aggressive behavior, ductal carcinoma is assigned a Gleason Grade 4 [58]. Similar to acinar carcinoma, a Gleason Grade 5 is given when comedo necrosis is present [45]. Due to its more favorable clinical behavior [59], PIN-like ductal adenocarcinoma should be assigned a Gleason Grade 3.

The cribriform pattern of ductal carcinoma needs to be distinguished from intraductal, and invasive acinar cribriform carcinoma. The typical cellular features, i.e., tall columnar pseudostratified cells may be difficult to appreciate in this pattern, making the distinction more difficult, especially in a biopsy specimen [94]. The presence of true papillae, slit-like lumina and elongated cells are in favor of ductal carcinoma, whereas punched-out lumens and rounded nuclei are typically observed in invasive acinar and intraductal carcinoma [94,101]. Immunohistochemistry is not particularly helpful. PTEN loss and ERG expression have been described less frequently in ductal carcinoma compared to acinar carcinoma [102]; however, both markers may be observed in ductal carcinoma; hence, they cannot be used as a reliable factor for differential diagnosis between the two entities. Similarly, a lack of basal cell markers is in favor of ductal carcinoma versus intraductal carcinoma; however, the opposite is not true, as ductal carcinomas may spread within prostatic ducts [96].

5.4. Basal Cell Carcinoma

Basal cell carcinoma is an unusual type of prostate cancer composed of basal cells [15]. Morphologically, it shows an adenoid cystic carcinoma-like pattern, with irregular cribriform formations containing mucin or basement membrane-like material within the lumina. The stroma is usually desmoplastic. Cytologically, the nuclei are large and hyperchromatic with scant cytoplasm. Nucleoli may or may not be prominent. Basal cell carcinoma has been considered an indolent tumor; however, almost half of the cases are associated with high-risk features and local recurrence [103]. Immunohistochemically, the tumor is positive for basal cell makers p63 and 34bE12 and negative for luminal markers PSA and PAP [104]. CK7 can be positive and AMACR staining is weak to negative [15]. At the molecular level, basal cell carcinoma shows a basal cell gene signature [105] with genetic aberrations that are distinct from acinar adenocarcinoma, i.e., frequent EGFR overexpression and absence.
of ERG rearrangements [103]. Differential diagnoses from other cribriform lesions of the prostate are based on the characteristic cytologic and architectural features mentioned above, as well as the distinct immunohistochemical profile. The most important distinction is from basal cell hyperplasia, a benign condition with excellent prognosis. The presence of an adenoid cystic carcinoma-like pattern and anastomosing, variably sized and irregularly shaped nests and tubules are in favor of basal cell carcinoma [15]. A widely invasive pattern in between normal prostatic acini and the presence of extraprostatic extension are virtually diagnostic of basal cell carcinoma (versus basal cell hyperplasia). Bcl-2 expression and high ki67 are more commonly observed in basal cell carcinoma compared to basal cell hyperplasia [106] and can be of use in difficult cases.

6. Conclusions

Overall, understanding the distinct nature of each cribriform lesion is fundamental for rendering the correct diagnosis and ensuring accuracy in clinical decision-making, prognosis prediction and personalized risk stratification of patients. In this review, we summarized the recent literature on the differential diagnosis of all neoplastic and non-neoplastic cribriform formations and explained how cribriform architecture could alter management decisions for prostate cancer patients. All the data and information collected in the review aim to help the pathologist deal with challenging and controversial cases of cribriform prostate patterns in the daily laboratory work practice. A graphical illustration of the various cribriform patterned lesions is shown in Figure 6. The goal is to make the correct diagnosis so as the patient can receive the appropriate clinical management and follow-up.

Figure 6. Graphical illustration of the various entities with cribriform morphology. Clear cell hyperplasia is characterized by a loose cribriform proliferation of clear cells without atypia. In atypical intraductal cribriform proliferation (AIDCP), atypical cells proliferate within a duct lumen (note the myoepithelial cells at the periphery); however, the architecture is loose, and the cells take up <50% of the surface of the lumen. In intraductal carcinomas (IDC), three patterns may be observed (alone or in combination): dense cribriform (>50% of the surface) or even solid (not shown) proliferation of atypical cells within the duct lumen (note the presence of myoepithelial cells, albeit they may be sparse in some cases) and loose architecture, but with either nuclear atypia or comedonecrosis. In cribriform carcinoma myoepithelial cells are missing (cribriform carcinoma is assigned a Gleason Grade 4). However, a Gleason Grade 5 is assigned if comedonecrosis is present (note again the lack of myoepithelial cells, consistent with invasive carcinoma). In ductal carcinoma, lumina are slit-like (instead of round) and cells are columnar. Myoepithelial cells are usually absent (although intraductal spread may be observed, not shown here).
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References

1. Srodon, M.; Epstein, J.I. Central zone histology of the prostate: A mimicker of high-grade prostatic intraepithelial neoplasia. Hum. Pathol. 2002, 33, 518–523. [CrossRef] [PubMed]
2. McNeal, J.E. Prostate. In Histology for Histopathologists; Mills, S.E., Ed.; Lippincott Williams & Wilkins: Philadelphia, PA, USA, 2007.
3. Hosler, G.A.; Epstein, J.I. Basal cell hyperplasia: An unusual diagnostic dilemma on prostate needle biopsies. Hum. Pathol. 2005, 36, 480–485. [CrossRef]
4. Rioux-Leclercq, N.C.; Epstein, J.I. Unusual Morphologic Patterns of Basal Cell Hyperplasia of the Prostate. Am. J. Surg. Pathol. 2002, 26, 237–243. [CrossRef]
5. Bostwick, D.; Montironi, R.; Sesterhenn, I.A. Diagnosis of Prostatic Intraepithelial Neoplasia: Prostate Working Group 1 Consensus Report. Scand. J. Urol. Nephrol. 2000, 34, 3–10. [CrossRef] [PubMed]
6. Montironi, R.; Mazzucchelli, R.; Lopez-Beltran, A.; ScarPELLi, M.; Cheng, L. Prostatic intraepithelial neoplasia: Its morphological and molecular diagnosis and clinical significance. BJU Int. 2011, 108, 1394–1401. [CrossRef] [PubMed]
7. Hickman, R.A.; Yu, H.; Li, J.; Kong, M.; Shah, R.B.; Zhou, M.; Melamed, J.; Deng, F.M. Atypical intraductal cribriform proliferations of the prostate exhibit similar molecular and clinicopathologic characteristics as intraductal carcinoma of the prostate. Am. J. Surg. Pathol. 2017, 41, 550–556. [CrossRef] [PubMed]
8. Magers, M.; Kunju, L.P.; Wu, A. Intraductal carcinoma of the prostate morphologic features, differential diagnoses, significance, and reporting practices. Arch. Pathol. Lab. Med. 2015, 139, 1234–1241. [CrossRef] [PubMed]
9. Cohen, R.J.; Wheeler, T.M.; BonkhoFF, H.; RubiN, M.A. A proposal on the identification, histologic reporting, and implications of intraductal prostatic carcinoma. Arch. Pathol. Lab. Med. 2007, 131, 1103–1109. [CrossRef]
10. Morais, C.L.; Han, J.S.; Gordetsky, J.; Nagar, M.S.; Anderson, A.E.; Lee, S.; Hicks, J.L.; Zhou, M.; Magi-Galluzzo, C.; Shah, R.B.; et al. Utility of PTEN and ERG Immunostaining for Distinguishing High-grade PIN From Intraductal Carcinoma of the Prostate on Needle Biopsy. Am. J. Surg. Pathol. 2015, 39, 169–178. [CrossRef] [PubMed]
11. Lotan, T.L.; Gumuskaya, B.; Rahimi, H.; Hicks, J.L.; Iwata, T.; Robinson, B.D.; Epstein, J.I.; De Marzo, A.M. Cytoplasmic PTEN protein loss distinguishes intraductal carcinoma of the prostate from high-grade prostatic intraepithelial neoplasia. Mod. Pathol. 2013, 26, 587–603. [CrossRef] [PubMed]
12. Shah, R.B.; Yoon, J.; Liu, G.; Tian, W. Atypical intraductal proliferation and intraductal carcinoma of the prostate on core needle biopsy: A comparative clinicopathological and molecular study with a proposal to expand the morphological spectrum of intraductal carcinoma. Histopathology 2017, 70, 693–702. [CrossRef]
13. Jiang, H.; Zhou, Z.; Jin, S.; Xu, K.; Zhang, H.; Xu, J.; Sun, Q.; Wang, J.; Xu, J. PRMT9 promotes hepatocellular carcinoma invasion and metastasis via activating PI3K/Akt/GSK-3β/Snail signaling. Cancer Sci. 2018, 109, 1414–1427. [CrossRef] [PubMed]
14. Divatia, M.K.; Ro, J.Y. Ym1-57–1054. Oncosei Med. J. 2016, 57, 1054–1062. [CrossRef] [PubMed]
15. Humphrey, P.A.; Moch, H.; Cubilla, A.L.; Ulbright, T.M.; Reuter, V.E. The 2016 WHO Classification of Tumours of the Urinary System and Male Genital Organs-Part B: Prostate and Bladder Tumours. Eur. Urol. 2016, 70, 106–119. [CrossRef] [PubMed]
16. Guo, C.C.; Epstein, J.I. Intraductal carcinoma of the prostate on needle biopsy: Histologic features and clinical significance. Mod. Pathol. 2006, 19, 1528–1535. [CrossRef] [PubMed]
17. Varma, M.; Delahunt, B.; Egevad, L.; Samarautunga, H.; Kristiansen, G. Intraductal carcinoma of the prostate: A critical re-appraisal. Virchows Arch. 2019, 474, 525–534. [CrossRef] [PubMed]
18. Samarautunga, H.; Delahunt, B.; Yaxley, J.W.; Johannsen, S.; Egevad, L. Intraductal Carcinoma of the Prostate: Extreme Nuclear Size Is Not a Diagnostic Parameter. Am. J. Surg. Pathol. 2021, 45, 1527–1533. [CrossRef]
19. Epstein, J.I.; Amin, M.B.; Fine, S.W.; Algbaga, F.; Aron, M.; Baydar, D.E.; Beltran, A.L.; Brimo, F.; Cheville, J.C.; Colecchia, M.; et al. The 2019 Genitourinary Pathology Society (GUPS) White Paper on Contemporary Grading of Prostate Cancer. Arch. Pathol. Lab. Med. 2021, 145, 461–493. [CrossRef]
20. Cohen, R.J.; McNeal, J.E.; Baillie, T. Patterns of differentiation and proliferation in intraductal carcinoma of the prostate: Significance for cancer progression. Prostate 2000, 43, 11–19. [CrossRef]
21. Watts, K.; Li, J.; Magi-Galluzzo, C.; Zhou, M. Incidence and clinicopathological characteristics of intraductal carcinoma detected in prostate biopsies: A prospective cohort study. Histopathology 2013, 63, 574–579. [CrossRef]
22. Humphrey, P.A. Histopathology of prostate cancer. Cold Spring Harb. Perspect. Med. 2017, 7, a030411. [CrossRef]
23. Khani, F.; Epstein, J.I. Prostate Biopsy Specimens With Gleason 3 + 3 = 6 and Intraductal Carcinoma. Am. J. Surg. Pathol. 2015, 39, 1383–1389. [CrossRef] [PubMed]
24. Khani, F.; Webker, S.E.; Hicks, J.L.; Robinson, B.D.; Barbieri, C.E.; De Marzo, A.M.; Epstein, J.I.; Pritchard, C.C.; Lotan, T.L. Intraductal carcinoma of the prostate in the absence of high-grade invasive carcinoma represents a molecularly distinct type of in situ carcinoma enriched with oncogenic driver mutations. *J. Pathol.* 2019, 249, 79–89. [CrossRef] [PubMed]

25. Montironi, R.; Zhou, M.; Magi-Galluzzi, C.; Epstein, J.I. Features and Prognostic Significance of Intraductal Carcinoma of the Prostate. *Eur. Urol. Oncol.* 2018, 1, 21–28. [CrossRef] [PubMed]

26. Miyai, K.; Divatia, M.K.; Shen, S.S.; Miles, B.J.; Ayala, A.G.; Ro, J.Y. Clinicopathological analysis of intraductal proliferative lesions of prostate: Intraductal carcinoma of prostate, high-grade prostatic intraepithelial neoplasia, and atypical cribriform lesion. *Hum. Pathol.* 2014, 45, 1572–1581. [CrossRef]

27. McNeal, J.E.; Yemoto, C.E. Spread of adenocarcinoma within prostatic ducts and acini. Morphologic and clinical correlations. *Am. J. Surg. Pathol.* 1996, 20, 802–814. [CrossRef]

28. Kryvenko, O.N.; Gupta, N.S.; Virani, N.; Schultz, D.; Gomez, J.; Amin, A.; Lane, Z.; Epstein, J.I. Gleason score 7 adenocarcinoma of the prostate with lymph node metastases: Analysis of 184 radical prostatectomy specimens. *Arch. Pathol. Lab. Med.* 2013, 137, 610–617. [CrossRef]

29. Van Der Kwast, T.; Al Daoud, N.; Collette, L.; Sykes, J.; Thoms, J.; Milosevic, M.; Bristow, R.G.; Van Tienhoven, G.; Warde, P.; Mirimanoff, R.O.; et al. Biopsy diagnosis of intraductal carcinoma is prognostic in intermediate and high risk prostate cancer patients treated by radiotherapy. *Eur. J. Cancer* 2012, 48, 1318–1325. [CrossRef]

30. Kimura, K.; Tsuzuki, T.; Kato, M.; Saito, A.M.; Sassa, N.; Ishida, R.; Hirabayashi, H.; Yoshino, Y.; Hattori, R.; Gotoh, M. Prognostic value of intraductal carcinoma of the prostate in radical prostatectomy specimens. *Prostate* 2014, 74, 680–687. [CrossRef]

31. Efstathiou, E.; Abrahams, N.A.; Tibbs, R.F.; Wang, X.; Pettaway, C.A.; Pisters, L.; Mathew, P.F.; Do, K.; Logothetis, C.J.; Troncoso, P. Morphologic Characterization of Preoperatively Treated Prostate Cancer: Toward a Post-Therapy Histologic Classification. *Eur. Urol.* 2010, 57, 1030–1038. [CrossRef]

32. Miyai, K.; Divatia, M.K.; Shen, S.S.; Miles, B.J.; Ayala, A.G.; Ro, J.Y. Heterogeneous clinicopathological features of intraductal carcinoma of the prostate: A comparison between “precursor-like” and “regular type” lesions. *Int. J. Clin. Exp. Pathol.* 2014, 7, 2518–2526. [PubMed]

33. Van Leenders, G.J.L.H.; van der Kwast, T.H.; Grignon, D.J.; Evans, A.J.; Kristiansen, G.; Kweldam, C.F.; Litjens, G.; McKenney, J.K.; Melamed, J.; Mottet, N.; et al. The 2019 International Society of Urological Pathology (ISUP) Consensus Conference on Grading of Prostatic Carcinoma. *Am. J. Surg. Pathol.* 2020, 44, e87–e99. [CrossRef] [PubMed]

34. Gandhi, J.S.; Smith, S.C.; Paner, G.P.; McKenney, J.K.; Sekhri, R.; Osunkoya, A.O.; Baras, A.S.; DeMarzo, A.M.; Cheville, J.C.; Jimenez, R.E.; et al. Reporting Practices and Resource Utilization in the Era of Intraductal Carcinoma of the Prostate. *Am. J. Surg. Pathol.* 2020, 44, 673–680. [CrossRef] [PubMed]

35. Robinson, B.D.; Epstein, J.I. Intraductal carcinoma of the prostate without invasive carcinoma on needle biopsy: Emphasis on radical prostatectomy findings. *J. Urol.* 2010, 184, 1328–1333. [CrossRef] [PubMed]

36. Grypari, I.M.; Logotheti, S.; Lazaris, A.C.; Kallidonis, P.; Fokaefs, E.; Melachrinou, M.; Zolota, V.; Tzelepi, V. Isolated Intraductal Carcinoma of the Prostate. *Eur. Urol. Oncol.* 2021, 4, 207–215. [CrossRef]

37. Schneider, T.M.; Osunkoya, A.O. ERG expression in intraductal carcinoma of the prostate: Comparison with adjacent invasive prostatic adenocarcinoma. *Mod. Pathol.* 2014, 27, 1174–1178. [CrossRef] [PubMed]

38. Robinson, B.; Magi-Galluzzi, C.; Zhou, M. Intraductal carcinoma of the prostate. *WHO Classification of Tumours of the Urinary System and Male Genital Organs* 4th ed.; Holger Moch Peter, A.; Humphrey Thomas, M.; Ulbright, V.E.R. (Eds.) International Agency for Research on Cancer: Lyon, France, 2016.

39. Kweldam, C.F.; van der Kwast, T.; van Leenders, G.J. On cribriform prostate cancer. *Transl. Androl. Urol.* 2018, 7, 145–154. [CrossRef]

40. Risbridger, G.P.; Taylor, R.A.; Clouston, D.; Sliwinski, A.; Thorne, H.; Hunter, S.; Li, J.; Mitchell, G.; Murphy, D.; Frydenberg, M.; et al. Patient-derived Xenografts Reveal That Intraductal Carcinoma of the Prostate Is a Prominent Pathology in BRCA2 Mutation Carriers with Prostate Cancer and Correlates with Poor Prognosis. *Eur. Urol.* 2015, 67, 496–503. [CrossRef]

41. Giri, V.N.; Knudsen, K.E.; Kelly, W.K.; Cheng, H.H.; Cooney, K.A.; Cookson, M.S.; Dahut, W.; Weissman, S.; Soule, H.R.; Petrylak, D.P.; et al. Implementation of Germline Testing for Prostate Cancer: Philadelphia Prostate Cancer Consensus Conference 2019. *J. Clin. Oncol.* 2020, 38, 2798–2811. [CrossRef]

42. Iczkowski, K.A.; Egved, L.; Ma, J.; Harding-Jackson, N.; Alenga, F.; Billis, A.; Camparo, P.; Cheng, L.; Clouston, D.; Comperat, E.M.; et al. Intraductal carcinoma of the prostate: Interobserver reproducibility survey of 39 urologic pathologists. *Ann. Diag. Pathol.* 2014, 18, 333–342. [CrossRef] [PubMed]

43. Schneider, T.M.; Osunkoya, A.O. ERG expression in intraductal carcinoma of the prostate: Comparison with adjacent invasive prostatic adenocarcinoma. *Mod. Pathol.* 2014, 27, 1174–1178. [CrossRef] [PubMed]

44. Zhou, M. High-grade prostatic intraepithelial neoplasia, PIN-like carcinoma, ductal carcinoma, and intraductal carcinoma of the prostate. *Mod. Pathol.* 2018, 31, S71–S79. [CrossRef] [PubMed]
46. Haffner, M.C.; Weier, C.; Xu, M.M.; Vaghasia, A.; Gürel, B.; Gümüşkaya, B.; Esopi, D.M.; Fedor, H.; Tan, H.L.; Kulac, I.; et al. Molecular evidence that invasive adenocarcinoma can mimic prostatic intraepithelial neoplasia (PIN) and intraductal carcinoma through retrograde glandular colonization. *J. Pathol.* 2016, 238, 31–41. [CrossRef] [PubMed]

47. Wilkinson, S.; Ye, H.; Karzai, F.; Harmon, S.A.; Terrigino, N.T.; VanderWeele, D.J.; Bright, J.R.; Atway, R.; Trostel, S.Y.; Carrabba, N.V.; et al. Nascent Prostate Cancer Heterogeneity Drives Evolution and Resistance to Intense Hormonal Therapy. *Eur. Urol.* 2021, 80, 746–757. [CrossRef]

48. McKay, R.R.; Xie, W.; Ye, H.; Fennessy, F.M.; Zhang, Z.; Lis, R.; Calagiu, C.; Rathkopf, D.; Laudone, V.P.; Bubley, G.J.; et al. Results of a Randomized Phase II Trial of Intense Androgen Deprivation Therapy prior to Radical Prostatectomy in Men with High-Risk Localized Prostate Cancer. *J. Urol.* 2021, 206, 80–87. [CrossRef]

49. McKay, R.R.; Berchuck, J.; Kwak, L.; Xie, W.; Silver, R.; Bubley, G.J.; Chang, P.K.; Wagner, A.; Zhang, Z.; Kibel, A.S.; et al. Outcomes of Post-Neoadjuvant Intense Hormone Therapy and Surgery for High Risk Localized Prostate Cancer: Results of a Pooled Analysis of Contemporary Clinical Trials. *J. Urol.* 2021, 205, 1689–1697. [CrossRef]

50. Porter, L.H.; Hashimoto, K.; Lawrence, M.G.; Pezaro, C.; Clouston, D.; Wang, H.; Papargiris, M.; Thorne, H.; Li, J.; Ryan, A.; et al. Intraductal carcinoma of the prostate can evade androgen deprivation, with emergence of castrate-tolerant cells. *BJUI Int.* 2018, 121, 971–978. [CrossRef]

51. Chua, M.L.K.; van der Kwast, T.H.; Bristow, R.G. Intraductal Carcinoma of the Prostate: Anonymous to Ominous. *Eur. Urol.* 2017, 72, 496–498. [CrossRef]

52. Egevad, L.; Delahunt, B.; Kristiansen, G.; Samaratinga, H.; Varma, M. Contemporary prognostic indicators for prostate cancer incorporating International Society of Urological Pathology recommendations. *Pathology* 2018, 50, 60–73. [CrossRef]

53. Rijstenberg, L.L.; Hansum, T.; Hollemans, E.; Kümmerlin, I.P.; Bangma, C.H.; Kwast, T.H.; Roobol, M.J.; Leenders, G.J.L.H. Intraductal carcinoma has a minimal impact on Grade Group assignment in prostate cancer biopsy and radical prostatectomy specimens. *Histopathology* 2020, 77, 742–748. [CrossRef] [PubMed]

54. Tzelepi, V.; Grypari, I.M.; Logotheti, S.; Stavros Kontogiannis, P.K.; Melachrinou, M.; Zolota, V. Contemporary Grading of Prostate Cancer: The Impact of Grading Criteria and the Significance of the Amount of Intraductal Carcinoma. *Cancers* 2021, 13, 5454. [CrossRef] [PubMed]

55. Varma, M.; Epstein, J.I. Head to head: Should the intraductal component of invasive prostate cancer be graded? *Histopathology* 2021, 78, 231–239. [CrossRef]

56. Amin, M.B. Histological variants of urothelial carcinoma: Diagnostic, therapeutic and prognostic implications. *Mod. Pathol.* 2009, 22, 596–5118. [CrossRef] [PubMed]

57. Epstein, J.I.; Egevad, L.; Amin, M.B.; Delahunt, B.; Srigley, J.R.; Humphrey, P.A. The 2014 International Society of Urological Pathology (ISUP) Consensus Conference on Gleason Grading of Prostatic Carcinoma: Definition of Grading Patterns and Proposal for a New Grading System. *Am. J. Surg. Pathol.* 2016, 40, 244–252. [CrossRef]

58. Epstein, J.I.; Allsbrook, W.C.J.; Amin, M.B.; Egevad, L.L. The 2005 International Society of Urological Pathology (ISUP) Consensus Conference on Gleason Grading of Prostatic Carcinoma. *Am. J. Surg. Pathol.* 2005, 29, 1228–1242. [CrossRef] [PubMed]

59. Paulk, A.; Giannico, G.; Epstein, J.I. PIN-like (Ductal) Adenocarcinoma of the Prostate. *Am. J. Surg. Pathol.* 2018, 42, 1693–1700. [CrossRef]

60. Epstein, J.I. Prostatic Ductal Adenocarcinoma: A Mini Review. *Med. Princ. Pract.* 2010, 19, 82–85. [CrossRef]

61. Van Leenders, G.J.L.H.; Kweldam, C.F.; Hollemans, E.; Kümmerlin, I.P.; Nieboer, D.; Verhoeof, E.I.; Remmers, S.; Incrocci, L.; Bangma, C.H.; van der Kwast, T.H. Improved Prostate Cancer Biopsy Grading by Incorporation of Invasive Cribriform and Intraductal Carcinoma in the 2014 Grade Groups. *Eur. Urol.* 2020, 77, 191–198. [CrossRef]

62. Kato, M.; Hirakawa, A.; Kobayashi, Y.; Yamamoto, A.; Ishida, R.; Sano, T.; Kimura, T.; Majima, T.; Ishida, S.; Funahashi, Y.; et al. The influence of the presence of intraductal carcinoma of the prostate on the grade group system’s prognostic performance. *Prostate* 2019, 79, 1065–1070. [CrossRef]

63. Chen-Maxwell, D.; Prendeville, S. Grading of prostate cancer: The impact of including intraductal carcinoma on the overall Grade Group assigned in diagnostic biopsies. *Histopathology* 2020, 77, 503–507. [CrossRef] [PubMed]

64. Gleason, D.F.; Mellinger, G.T. Prediction of Prognosis for Prostatic Adenocarcinoma by Combined Histological Grading and Clinical Staging. *J. Urol.* 1974, 111, 58–64. [CrossRef]

65. Matsuo, A.; Epstein, J.I. Grading of Prostate Cancer: Past, Present, and Future. *Curr. Urol. Rep.* 2016, 17, 25. [CrossRef] [PubMed]

66. Dong, F.; Yang, P.; Wang, C.; Wu, S.; Xiao, Y.; McDougal, W.S.; Young, R.H.; Wu, C.-L. Architectural heterogeneity and cribriform pattern predict adverse clinical outcome for Gleason grade 4 prostatic adenocarcinoma. *Am. J. Surg. Pathol.* 2013, 37, 1855–1861. [CrossRef] [PubMed]

67. Rubin, M.A.; de La Taille, A.; Bagiella, E.; Olsson, C.A.; O’Toole, K.M. Cribriform carcinoma of the prostate and cribriform prostatic intraepithelial neoplasia: Incidence and clinical implications. *Am. J. Surg. Pathol.* 1998, 22, 840–848. [CrossRef]

68. Keefe, D.T.; Schieda, N.; El Hallani, S.; Breau, R.H.; Morash, C.; Robertson, S.J.; Mai, K.T.; Belanger, E.C.; Flood, T.A. Cribriform morphology predicts upstaging after radical prostatectomy in patients with Gleason score 3 + 4 = 7 prostate cancer at transrectal ultrasound (TRUS)-guided needle biopsy. *Virchows Arch.* 2015, 467, 437–442. [CrossRef]

69. O’Brien, C.; True, L.D.; Higano, C.S.; Rademacher, B.L.S.; Garzotto, M.; Beer, T.M. Histologic changes associated with neoadjuvant chemotherapy are predictive of nodal metastases in patients with high-risk prostate cancer. *Am. J. Clin. Pathol.* 2010, 133, 654–661. [CrossRef]
70. Iczkowski, K.A.; Torkko, K.C.; Kotnis, G.R.; Wilson, R.S.; Huang, W.; Wheeler, T.M.; Abeyta, A.M.; La Rosa, E.G.; Cook, S.; Werahera, P.N.; et al. Digital quantification of five high-grade prostate cancer patterns, including the cribriform pattern, and their association with adverse outcome. *Am. J. Clin. Pathol.* 2011, 136, 98–107. [CrossRef]

71. McKenney, J.K.; Wei, W.; Hawley, S.; Auman, H.; Newcomb, L.F.; Boyer, H.D.; Fazli, L.; Simko, J.; Hurtado-Coll, A.; Troyer, D.A.; et al. Histologic Grading of Prostatic Adenocarcinoma Can Be Further Optimized: Analysis of the Relative Prognostic Strength of Individual Architectural Patterns in 1275 Patients From the Canary Retrospective Cohort. *Am. J. Surg. Pathol.* 2016, 40, 1439–1456. [CrossRef]

72. Hollemans, E.; Verhoef, E.I.; Bangma, C.H.; Rietbergen, J.; Osanto, S.; Pelger, R.C.M.; van Wezel, T.; van der Poel, H.; Bekers, E.; Hollemans, J.; et al. Cribriform architecture in radical prostatectomies predicts oncological outcome in Gleason score 8 prostate cancer patients. *Mod. Pathol.* 2021, 34, 184–193. [CrossRef]

73. Kweldam, C.F.; Wildhagen, M.F.; Steyerberg, E.W.; Bangma, C.H.; van der Kwaast, T.H.; van Leenders, G.J.L. Cribriform growth is highly predictive for postoperative metastasis and disease-specific death in Gleason score 7 prostate cancer. *Mod. Pathol.* 2015, 28, 457–464. [CrossRef]

74. Böttcher, R.; Kweldam, C.F.; Livingstone, J.; Lalonde, E.; Yamaguchi, T.N.; Huang, V.; Yousif, F.; Fraser, M.; Bristow, R.G.; van der Kwaast, T.; et al. Cribriform and intraductal prostate cancer are associated with increased genomic instability and distinct genomic alterations. *BMC Cancer* 2018, 18, 8. [CrossRef] [PubMed]

75. Chua, M.L.K.; Lo, W.; Pintilie, M.; Muric, J.; Lalonde, E.; Bhandari, V.; Mahamud, O.; Gopalan, A.; Kweldam, C.F.; van Leenders, G.J.L.H.; et al. A Prostate Cancer “Nimbusos”: Genomic Instability and SchLAP1 Dysregulation Underpin Aggression of Intraductal and Cribriform Subpathologies. *Eur. Urol.* 2017, 72, 665–674. [CrossRef]

76. Kweldam, C.F.; van der Leenders, G.J.; van der Kwaast, T. Grading of prostate cancer: A work in progress. *Histopathology* 2019, 74, 146–160. [CrossRef] [PubMed]

77. Montironi, R.; Cimadamore, A.; Cheng, L.; Lopez-Beltran, A.; Scarpelli, M. Prostate cancer grading in 2018: Limitations, implementations, cribriform morphology, and biological markers. *Int. J. Biol. Markers* 2018, 33, 331–334. [CrossRef] [PubMed]

78. Kweldam, C.F.; Nieboer, D.; Algaba, F.; Auman, H.; Newcomb, L.F.; Boyer, H.D.; Fazli, L.; Simko, J.; Hurtado-Coll, A.; Troyer, D.A.; et al. Histologic Grading of Prostatic Adenocarcinoma Can Be Further Optimized: Analysis of the Relative Prognostic Strength of Individual Architectural Patterns in 1275 Patients From the Canary Retrospective Cohort. *Am. J. Surg. Pathol.* 2016, 40, 1439–1456. [CrossRef]

79. Hesterberg, A.B.; Gordetsky, J.B.; Hurley, P.J. Cribriform Prostate Cancer: Clinical Pathologic and Molecular Considerations. *Urology* 2021, 155, 47–54. [CrossRef] [PubMed]

80. Shah, R.B.; Cai, Q.; Aron, M.; Berney, D.M.; Cheville, J.C.; Deng, F.-M.; Epstein, J.; Fine, S.W.; Genega, E.M.; Hirsch, M.S.; et al. Diagnosis of “cribriform” prostatic adenocarcinoma: An interobserver reproducibility study among urologic pathologists with recommendations. *Am. J. Cancer Res.* 2021, 11, 3990–4001. [PubMed]

81. Verhoef, E.I.; van Cappellen, W.A.; Slotman, J.A.; Kremers, G.-J.; Ewing-Graham, P.C.; Houtsmuller, A.B.; van Royen, M.E.; van Helleman, J.; et al. Digital quantification of five high-grade prostate cancer patterns, including the cribriform pattern, and their association with adverse outcome. *Virchows Arch.* 2018, 474, 146–160. [CrossRef] [PubMed]

82. Hesterberg, A.B.; Rios, B.L.; Wolf, E.M.; Tubbs, C.; Wong, H.Y.; Schaffer, K.R.; Lotan, T.L.; Giannico, G.A.; Gordetsky, J.B.; et al. Gleason grade 4 prostate adenocarcinoma patterns: An interobserver agreement study among genitourinary pathologists. *Histopathology* 2016, 69, 441–449. [CrossRef] [PubMed]

83. Montironi, R.; Cimadamore, A.; Cheng, L.; Lopez-Beltran, A.; Scarpelli, M. Prostate cancer grading in 2018: Limitations, implementations, cribriform morphology, and biological markers. *Int. J. Biol. Markers* 2018, 33, 331–334. [CrossRef] [PubMed]

84. Xiao, G.-Q.; Nguyen, E.; Unger, P.D.; Sherrod, A.E. Comparative expression of immunohistochemical biomarkers in cribriform and pattern 4 non-cribriform prostatic adenocarcinoma. *Exp. Mol. Pathol.* 2019, 103, 271–286. [CrossRef] [PubMed]

85. Olkhov-Mitsel, E.; Siadat, F.; Kron, K.; Liu, L.; Savio, A.J.; Trachtenberg, J.; Fleschner, N.; van der Kwaast, T.; Bapat, B. Distinct DNA methylation alterations are associated with cribriform architecture and intraductal carcinoma in Gleason pattern 4 prostate tumors. *OncoLett.* 2017, 14, 390–396. [CrossRef]

86. Ronen, S.; Abbott, D.W.; Kravtsov, O.; Abdelkader, A.; Xu, Y.; Banerjee, A.; Iczkowski, K.A. PTEN loss and p27 loss differ among morphologic patterns of prostate cancer, including cribriform. *Hum. Pathol.* 2017, 45, 85–91. [CrossRef] [PubMed]

87. Connell, B.; Kopach, P.; Ren, W.; Joshi, R.; Naber, S.; Zhou, M.; Mathew, P. Aberrant integrin αv and α5 expression in prostate adenocarcinomas and bone-metastases is consistent with a bone-colonizing phenotype. *Transl. Androl. Urol.* 2020, 9, 1630–1638. [CrossRef] [PubMed]

88. Goel, H.L.; Li, J.; Kogan, S.; Languino, L.R. Integrins in prostate cancer progression. *Endocr. Relat. Cancer* 2008, 15, 657–664. [CrossRef]

89. Ranasinha, N.; Omer, A.; Philippou, Y.; Harriss, E.; Davies, L.; Chow, K.; Chetta, P.M.; Erickson, A.; Rajakumar, T.; Mills, I.G.; et al. Ductal adenocarcinoma of the prostate: A systematic review and meta-analysis of incidence, presentation, prognosis, and management. *BJU Int. Compas* 2021, 2, 13–23. [CrossRef] [PubMed]

90. Seipel, A.H.; Wiklund, F.; Wiklund, N.P.; Egevad, L. Histopathological features of ductal adenocarcinoma of the prostate in 1,051 radical prostatectomy specimens. *Virchows Arch.* 2013, 462, 429–436. [CrossRef] [PubMed]

91. Tan, Y.G.; Khalid, F.; Huang, H.H.; Chen, K.; Tay, K.J.; Lau, W.K.O.; Cheng, C.W.S.; Ngo, N.T.; Yuen, J.S.P. Prostatic ductal adenocarcinoma variant predicts worse pathological and oncological outcomes: Insight from over 1000 consecutive patients from a large prospective uro-oncology registry. *Prostate* 2021, 81, 242–251. [CrossRef] [PubMed]
92. Chow, K.; Bedő, J.; Ryan, A.; Agarwal, D.; Bolton, D.; Chan, Y.; Dundee, P.; Frydenberg, M.; Furrer, M.A.; Goad, J.; et al. Ductal variant prostate carcinoma is associated with a significantly shorter metastasis-free survival. *Eur. J. Cancer* 2021, 148, 440–450. [CrossRef]

93. Ranasinghe, W.; Shapiro, D.D.; Hwang, H.; Wang, X.; Reichard, C.A.; Elshehtawi, M.; Achim, M.F.; Bathala, T.; Tang, C.; Aparicio, A.; et al. Ductal Prostate Cancers Demonstrate Poor Outcomes with Conventional Therapies. *Eur. Urol.* 2021, 79, 298–306. [CrossRef] [PubMed]

94. Seipel, A.H.; Delahunt, B.; Samaratunga, H.; Amin, M.; Barton, J.; Berney, D.M.; Billis, A.; Cheng, L.; Comperat, E.; Evans, A.; et al. Diagnostic criteria for ductal adenocarcinoma of the prostate: Interobserver variability among 20 expert uropathologists. *Histopathology* 2014, 65, 216–227. [CrossRef] [PubMed]

95. Melicow, M.M.; Epstein, J.I. Immunohistochemical Antibody Cocktail Staining (p63/HMWCK/AMACR) of Ductal Adenocarcinoma and Gleason Pattern 4 Cribriform and Noncribriform Acinar Adenocarcinomas of the Prostate. *Am. J. Surg. Pathol.* 2007, 31, 889–894. [CrossRef] [PubMed]

96. Seipel, A.H.; Delahunt, B.; Samaratunga, H.; Egevad, L. Ductal adenocarcinoma of the prostate: Histogenesis, biology and clinicopathological features. *Pathology* 2016, 48, 398–405. [CrossRef] [PubMed]

97. Simper, N.B.; Jones, C.L.; MacLennan, G.T.; Montironi, R.; Williamson, S.R.; Osunkoya, A.O.; Wang, M.; Zhang, S.; Grignon, D.J.; Eble, J.N.; et al. Basal cell carcinoma of the prostate is an aggressive tumor with frequent loss of PTEN expression and overexpression of EGFR. *Hum. Pathol.* 2015, 46, 805–812. [CrossRef] [PubMed]

98. Hennes, D.; Dragovic, A.; Sewell, J.; Hoh, M.Y.; Grills, R. Primary basal cell carcinoma of the prostate with concurrent adenocarcinoma. *IJI Case Reports* 2020, 3, 57–60. [CrossRef] [PubMed]

99. Su, X.; Long, Q.; Bo, J.; Shi, Y.; Zhao, L.; Lin, Y.; Luo, Q.; Ghazanfar, S.; Zhang, C.; Liu, Q.; et al. Mutational and transcriptomic landscapes of a rare human prostate basal cell carcinoma. *Prostate* 2020, 80, 508–517. [CrossRef] [PubMed]

100. Yang, X.J.; McEntee, M.; Epstein, J.I. Distinction of basaloid carcinoma of the prostate from benign basal cell lesions by using immunohistochemistry for bcl-2 and Ki-67. *Hum. Pathol.* 1998, 29, 1447–1450. [CrossRef]