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**Abstract:** Epidemiological and biological evidence indicate a causal relationship between the presence of proliferative atrophic lesions (PAH), the development of PIN and prostate cancer. Inflammatory and atrophic lesions of the prostate are widely underestimated and not generally mentioned in pathology reports. We performed a histopathological concordance study among 15 dedicated and non-dedicated genito-urinary pathologists on 116 histological slides containing prostate atrophic lesions, PIN and cancer.
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As stated in the Methods section the cases were selected by two dedicated uropathologists (EB and MF) from two different institutions. Each pathologist was blinded to the other's evaluation and they did not discuss the cases. The gold standard was another GU pathologist (RM) from a third institution who did not participate in case selection. For comparison, we also present data for the overall percentage agreement, which would not be as heavily dependent on the evaluation by a single reviewer.

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Concordance in the histological diagnosis of Focal Prostatic Atrophy Lesions, Acute and Chronic Prostatitis, PIN and Prostate Cancer Among 15 European Pathologists

Francesca Giunchi MD¹, Kristina Jordahl PhD², Enrico Bollito MD³, Maurizio Colecchia MD⁴, Carlo Patriarca MD⁵, Antonietta D’Errico MD¹, Francesco Vasuri MD, PhD¹, Deborah Malvi MD¹, Alessandro Fornari MD³, Luca Reggiani Bonetti MD⁶, Barbara Corti MD¹, Mauro Papotti MD⁷, Paolo DeGiuli MD⁸, Massimo Loda MD⁹, Rodolfo Montironi MD¹⁰, Michelangelo Fiorentino MD, PhD¹ and, Jennifer R. Rider ScD²,¹¹

MF and JR co-shared senior authorship.

¹Pathology Department, S. Orsola-Malpighi Hospital, University of Bologna, Viale Ercolani 4/2, 40138 Bologna, Italy

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³Department of Oncology, Division of Pathology, San Luigi Gonzaga Hospital, Orbassano, Turin area, Regione Gonzole, 10, 10043 Italy.

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11 Department of Epidemiology, Boston University School of Public Health, Boston, 715 Albany Street, T317E, MA 02118, USA.

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Abstract:

Epidemiological and biological evidence indicate a causal relationship between the presence of proliferative atrophic lesions (PAH), the development of PIN and prostate cancer. Inflammatory and atrophic lesions of the prostate are widely underestimated and not generally mentioned in pathology reports.

We performed a histopathological concordance study among 15 dedicated and non-dedicated genito-urinary pathologists on 116 histological slides containing prostate atrophic lesions, PIN and cancer.

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According to the raising relevance of PAH in prostate cancer our results on histopathological concordance support the inclusion of at least PAH in the routine pathology reporting of pathological prostate specimens.

Key words: Atrophic lesions, Inflammation, PAH, Prostate.
**Introduction:**

Long-term chronic inflammation is linked to the development of carcinoma in several organ systems and it is also an important aetiology factor in prostate cancer (PCa). [1] Based on observations in other organs such as the stomach, liver and large bowel in the 1990s, the interest in inflammatory and atrophic lesions in prostate cancer was further cultivated. Inflammatory cells could produce cellular or genomic irreversible damage in prostate cells; cause loss of tolerance to normal prostate antigens; and induce an autoimmune self-perpetuating reaction leading in turn to a “field effect” for the development of PCa [2]. The major events potentially leading to prostate inflammation are infections (virus, fungi, mycobacteria and parasite, and rarely bacteria), hormonal alteration, physical trauma, urine reflux and dietary habits. These exposures can result in injury to the luminal cell layer, which in turn induces reactive (defensive) hyperplasia of basal cells called “proliferative inflammatory atrophy” (PIA), thereby initiating genetic instability. Cytokines released by the inflammatory cells slowly induce epithelial proliferation and angiogenesis with the accumulation of genomic changes, eventually resulting in neoplastic transformation through PIN (prostatic intraepithelial neoplasia). [1] Several studies have referred to a morphological transition between PIA and PIN, as well as PIA and prostate cancer. These observations are further supported by evidence of an elevated proliferative fraction in atrophic areas and the closely proximity of these regions to PIN and PCa. [3, 4] In addition somatic genomic alterations detectable in PIN and PCa have been found in cells in PIA. In particular, these cells show molecular effects of inflammatory stress, such as high levels of glutathione S-transferase P1 (GSTP1), GSTA1 and cyclooxygenase-2 (COX-2). [2]

Prostatic inflammation and atrophy are not routinely collected as part of a standard histopathologic review, yet there is considerable interest in the potential role of atrophy and inflammation in the development and progression of prostate cancer. Recent studies have related the presence of prostatic atrophic lesions and inflammation to prostate cancer risk [5, 6, 7, 8] and survival [9], with mixed results. One possible explanation for the inconsistent findings is misclassification of the various lesion types due to subjectivity in grading.
The increasing correlation among inflammation, atrophic lesions, PIN and PCa led De Marzo et al. to propose a histological classification of the different atrophic lesions in 4 morphological patterns: simple atrophy (SA), partial atrophy (PA), post atrophic hyperplasia (PAH) and simple athropy with cyst formation (SACF). PAH can be considered a surrogate of PIA. While other studies have shown that PIN is frequently overdiagnosed in pathological specimens [10], the reliability of the assessment of inflammation and atrophic changes is unclear. A concordance study found that 83% of atrophic lesions were classified as the correct subtype, but there was substantial variability in accuracy across specific lesion types [11]. To specifically address this gap in the literature, we undertook a histopathological concordance study that focused on inflammation, atrophy, PIN and cancer among 15 European pathologists, including both those dedicated and non-dedicated to genitourinary pathology.

Materials and Methods:

Two dedicated uropathologists (MF, EB) selected 61 slides of prostate tissue and identified 121 areas of interests (ROIs) with inflammation, atrophic lesions, PIN and PCa on the basis of the original pathology report from biopsies, radical prostatectomy and TURP specimens. The slides were selected from the Pathology archives at the S. Orsola-Malpighi Hospital in Bologna and the S. Luigi Gonzaga Hospital in Turin. None of the included patients was treated with hormone ablation or radiation therapy. Specifically, the two pathologists who selected the cases asked for the re-cutting of three copies of each selected block. Then they drew circles around the area of interests with a bullpen. The three sets of slides were anonymized and circulated among the 15 pathologists who accepted to join the study for histological revision of prostate atrophic lesions according to the De Marzo et al. classification, [11]. Twenty-one of the 61 slides included at least a single focus of PCa together with atrophic lesions. Of the 30 pathologists originally invited to the study, 15 professionals from 9 centers accepted to review the slides and completed the evaluation form. Eight of these 15 pathologists were dedicated experienced uropathologists while 7 were general pathologists. Participating pathologists were asked to independently record the presence of the following histological features: prostate cancer, PIN, SA, SACF, PAH/PIA, PA, acute prostatitis, and chronic prostatitis. Two of the uropathologists did not review all of slides for all lesion types, leaving 116 slides for analysis of
prostate cancer, 115 for most types of inflammation and atrophy, and 114 for simple atrophy with cyst formation. The entire review process took about two years.

Statistical Methods

We used two methods to evaluate the interpreter reliability of each histological feature. First we considered the overall percentage agreement, which is the average percent agreement across all possible rater pairs. We also estimated kappa statistics, which range from -1 to 1, where 1 indicates perfect agreement, 0 represents the agreement expected by chance, and values <0 indicate less agreement than is expected by chance. A suggested interpretation of kappa is that 0.21-0.40 is fair agreement, 0.41-0.60 is moderate agreement, and 0.61-0.99 is almost perfect agreement. The exact kappa coefficient was developed because it reduces to Cohen’s un-weighted Kappa for two raters. Second, we used the rating of one particularly experienced dedicated genitourinary pathologist (RM) as the “gold standard” by which to compare all other raters. We calculated the mean percentage agreement and mean Cohen’s Kappa across the 15 pairs. Comparisons were made among all 15 pathologists, and the subset of dedicated GU pathologists (N=8).

Results:

Table 1 shows that the overall percent agreement between all possible pairs of reviewers was 80.2% for prostate cancer, 67% for PIN, and 48.7% for any atrophic changes. While the kappa statistics for prostate cancer indicated nearly perfect agreement, agreement for PIN and atrophic changes were in the fair to moderate range. When specific types of focal prostatic atrophy were considered, the overall percentage agreement ranged from 5.2% for SA to 43% for SACF, all with kappa statistics indicating modest precision. In particular the agreement for PAH and partial atrophy was similar and low: 26.1% and 22.6% respectively. The average agreement between pairs of raters was 20% for chronic inflammation and 53% for acute inflammation.

When a single gold standard genitourinary pathologist was designated, the mean percent agreement increased: 96.6% for prostate cancer (kappa = 0.88) and 91.7% for PIN (kappa=0.55). There was
agreement with the gold standard by 92.5% of the raters regarding the presence of any atrophic changes, but the kappa statistic indicated that this agreement was often due to chance (kappa=0.47). Simple atrophy had the lowest agreement (65.2%), followed by PAH (71.9%), partial atrophy (84.7%), and simple atrophy with cyst formation (88.4%). Pathologists agreed with the gold standard rater for chronic and acute inflammation in 71.9% and 69.4% of ratings, respectively. The dedicated genitourinary pathologists consistently had higher agreement for all lesion types, which was especially pronounced for atrophic changes and specific types of focal atrophy. The agreement for atrophic lesions (92.2%) was stratified as follows: 86.9% for SACF, 84.6% for PA, 77.4% for PAH and 64.9% for SA. The agreement among non-genitourinary pathologists with the gold standard was similar to the results for dedicated genitourinary pathologists.

**Discussion:**

Our current understanding of PIA has raised questions about the role of this lesion in the development of prostate cancer. Moreover, the association between chronic inflammation/PAH and prostate cancer appears to be stronger compared to the other atrophic lesions such as SA. Nevertheless few studies have tested the diagnostic agreement among pathologists for atrophic lesions, which will impact the results of studies of this topic and inform the need to conduct centralized pathological reviews. In this study we evaluated the ability of general and genitourinary-dedicated pathologists to recognize atrophic lesions according to the classification proposed by De Marzo in order to determine whether there is utility in introducing the characterization of these lesions into standard pathology reports. Our results highlighted a moderate agreement for atrophic lesions among both genitourinary and general pathologists, but with patterns that varied by atrophy type. The agreement was favorable for PAH and simple atrophy with cyst formation among all the pathologists. However, for PA and SA the concordance was suboptimal. Further training on recognizing these lesions types may be warranted. With respect to the diagnosis of prostate cancer and PIN, our results confirmed several previous reports that found excellent agreement for cancer and very good agreement for PIN. Our concordance data support the standard reporting of PIN.
Our study is affected by several limitations. The number of raters and reviewed cases were both modest. However, we estimated that the 121 slides included in the study would allow us to identify differences in lesion attribution among the 15 participating pathologists. Other limitations derive from the binary categorization of the variables as presence or absence of each histological feature. Had we included in the review the extent of inflammation, the concordance findings may have changed. Similarly, we provided the reviewers with H&E slides that were already circled for areas of interest to be scored. If areas of interest were not indicated, as in real-world clinical and research settings, non GU-dedicated pathologists may have underestimated the atrophic lesions. Despite the reference pathologist being the most experienced, the reference gold standard was likely imperfect. However, if we selected as the gold standard, for instance, the majority opinion of the raters we may have introduced more substantial bias.

This study did not confirm results from other pathology review studies where dedicated genitourinary pathologists displayed higher diagnostic concordance for prostate cancer, PIN and atrophic lesions compared to general pathologists [12]. In our data, differences in the agreement with the gold standard for dedicated and general European pathologist were minimal.

Given the growing interest in inflammation in prostate cancer risk assessment, our results support the inclusion of at least PAH/PIA in the standard pathology reporting of all pathological prostate specimens, but that some specific types of atrophy and for inflammation, additional training may improve concordance.

Compliance with Ethical Standards:

Funding

No external funds were obtained for this work.

Conflict of Interest

The authors declare no conflicts of interest.
References


Table 1. Concordance among 15 pathologists in histological assessment of inflammation, atrophy, PIN and cancer

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<th>Subjects</th>
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Figure Legend: Figure A) Morphological patterns of the focal prostate atrophic lesions: partial atrophy (PA), simple atrophy with cyst formation (SACF), post-atrophic hyperplasia (PAH) and simple atrophy (SA) (H&E, 100x magnification). B) Morphological pathway of progression of prostate cancer development through simple atrophy, PAH and PIN (H&E, 100x magnification).

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