Review – Urothelial Cancer

Grading of Urothelial Carcinoma and The New “World Health Organisation Classification of Tumours of the Urinary System and Male Genital Organs 2016”

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Abstract

Context: In the management of urothelial carcinoma, determination of the pathological grade aims at stratifying tumours into different prognostic groups to allow evaluation of treatment results, and optimise patient management. This article reviews the principles behind different grading systems for urothelial bladder carcinoma discussing their reproducibility and prognostic value.

Objective: This paper aims to show the evolution of the World Health Organisation (WHO) grading system, discussing their reproducibility and prognostic value, and evaluating which classification system best predicts disease recurrence and progression. The most optimal classification system is robust, reproducible, and transparent with comprehensive data on interobserver and intraobserver variability. The WHO published an updated tumour classification in 2016, which presents a step forward, but its performance will need validation in clinical studies.

Evidence acquisition: Medline and EMBASE were searched using the key terms WHO 1973, WHO/International Society of Urological Pathology 1998, WHO 2004, WHO 2016, histology, reproducibility, and prognostic value, in the time frame 1973 to May 2016. The references list of relevant papers was also consulted, resulting in the selection of 48 papers.

Evidence synthesis: There are still inherent limitations in all available tumour classification systems. The WHO 1973 presents considerable ambiguity for classification of the G2 tumour group and grading of the G1/2 and G2/3 groups. The 2004 WHO classification introduced the concept of low-grade and high-grade tumours, as well as the papillary urothelial neoplasm of low malignant potential category which is retained in the 2016 classification. Furthermore, while molecular markers are available that have been shown to contribute to a more accurate histological grading of urothelial carcinomas,

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1. Introduction

This article reviews the principles behind different grading systems for urothelial bladder carcinoma and discusses their reproducibility and prognostic value. It will not address urinary cytology, as this paper’s focus is on histological criteria. The European Association of Urology (EAU) Nonmuscle Invasive Bladder (NMIBC) Guidelines Panel recommends the use of both the World Health Organisation (WHO) 1973 and WHO 2004/2016 classifications [1]. This article touches on the strengths and weaknesses of the available classifications.

The aim of tumour classification is to be able to identify groups of patients according to different outcomes and manage them accordingly. An ideal classification system will be robust, reproducible, and transparent. It should be relatively easy for pathologists to understand and apply, and for urologists and oncologists to utilise in the treatment decision-making process.

During the last 40 yr, different bladder tumour-grading systems have been introduced, with generally good acceptance. One of the most widely accepted systems has been the WHO 1973 system, which is still used in some parts of Europe [2]. However, the currently recommended system is the new WHO 2016 grading system, which is being proposed for universal use and should be adopted worldwide [3].

1.1. The WHO 1973 classification

The WHO 1973 grading system was proposed by Mostofi et al [2] and differentiates papillary urothelial lesions into three grades: G1, G2, and G3 (Table 1) [4]. Tumours are graded according to the degree of cellular and architectural atypia. The lowest grade (G1) displays nearly no atypia, while the highest grade (G3) displays major atypia with major architectural disorders, such as loss of polarity or pseudostatification. The WHO 1973 is a robust and time-tested grading system. Some centres still prefer it to newer grading systems because it allows a comparison of long-term outcome results between different clinical centres. The major limitation of the WHO 1973 classification is the G2 group of tumours.

1.1.1. G1 tumours

In G1 tumours, the papillary urothelial carcinomas consist of orderly-arranged, nearly-normal, urothelial cells. The tumours display slender papillae with minimal abnormality. Mitosis and necrosis are globally absent. Nuclear pleomorphism is absent, with regular chromatin distribution. If present, nucleoli are small. Cell borders are well defined.

1.1.2. G3 tumours

In G3 tumours, architectural and nucleocytoplasmic heterogeneity is a typical finding. Mitosis can be distributed all over the urothelium, even in superficial layers, and necrosis may be present. There is increased cellularity, nuclear crowding, and a lack of normal differentiation. The nuclei display irregular membranes and shape and sometimes neoplastic giant cells. Chromatin distribution is granular or coarse and prominent nucleoli are common.

1.1.3. G2 tumours

In the G2 group of tumours, several different types of noninvasive urothelial carcinomas are grouped broadly together. The architectural and nucleocytoplasmic atypia in G2 cover a broad spectrum of lesions between those seen in grades G1 and G3. The variation in polarity and nuclear aspects can be very important for treatment decision making and even though subclassification of grades was never recommended in the WHO 1973 classification [5], some authors have tried to subclassify G2 tumours into G2a and G2b subgroups on the basis of nuclear polymorphism and mitotic count [6,7].

In addition, the WHO 1973 system has led some pathologists to ambiguously grade tumours that fall between grades 1 and 2 as G1/2, and tumours that fall between grades 2 and 3 as G2/3. Grade G1/2 subgroups contain relatively low-grade (LG) G2 urothelial carcinomas, which thereby improving selection of treatment for a given patient, these are not (yet) part of standard clinical practice.

Conclusions: The prognosis of patients diagnosed with urothelial carcinoma greatly depends on correct histological grading of the tumour. There is still limited data regarding intraobserver and interobserver variability differences between the WHO 1973 and 2004 classification systems. Additionally, reproducibility remains a concern: histological differences between the various types of tumour may be subtle and there is still no consensus amongst pathologists. The recent WHO 2016 classification presents a further improvement on the 2004 classification, but until further data becomes available, the European Association of Urology currently recommends the use of both WHO 1973 and WHO 2004/2016 classifications.

Patient summary: Bladder cancer, when treated in time, has a good prognosis. However, selection of the most optimal treatment is largely dependent on the information your doctor will receive from the pathologist following evaluation of the tissue resected from the bladder. It is therefore important that the classification system that the pathologist uses to grade the tissue is transparent and clear for both urologists and pathologists. A reliable classification system will ensure that aggressive tumours are not misinterpreted, and less aggressive cancer is not overtreated.

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Table 1 – The World Health Organisation 1973 and World Health Organisation 2004/16 classification of urothelial bladder tumours.

| Tumour grade | 1 | 2 | 3 |
|--------------|---|---|---|
| Description 1973 | Slender papillae in an orderly arrangement | Atypia can be more or less marked, increased urothelial layers | Marked atypia, loss of polarity, mitosis |
| Micrograph | ![Image](image1.png) | ![Image](image2.png) | ![Image](image3.png) |
| Biological potential | Papillary neoplasia of low malignant potential | Low | High |
| Description 2004/16 | Slim papillae without atypia, no thickening of the urothelium | Papillae have increased layers, atypia is rare, polarity conserved | Major atypia, major architecture destroyed |
| Micrograph | ![Image](image4.png) | ![Image](image5.png) | ![Image](image6.png) |

are much closer to G1, and relatively high-grade (HG) tumours, which are much closer to G3. This practice creates further ambiguity, complicates the treatment decision-making process and should therefore be avoided [7].

1.2. The WHO/International Society of Urological Pathology 1998 classification system

In 1997, a new multidisciplinary consensus meeting was held to revisit terminology and provide updated recommendations to the WHO on the pathology of urothelial carcinomas.

The WHO/International Society of Urological Pathology (ISUP) classification of 1998 distinguishes papilloma, papillary urothelial neoplasm of low malignant potential (PUNLMP), and LG and HG carcinomas. This classification system was the first to introduce the category of PUNLMP [4]. PUNLMP was subsequently included in the WHO 2004 classification and, although initially controversial [8], is retained in the WHO 2016 classification without any polemic [3].

The distinction made between PUNLMP and LG urothelial carcinoma is based on morphology. While PUNLMP and LG urothelial carcinoma differ on a molecular level, few papers address this aspect. A paper by Barbisan et al [9] described overexpression of FGFR3 and CK20 on a protein level in urothelial neoplasm of low malignant potential, while another paper by Lee et al [10] found lower expression levels of microRNAs-125b in LG noninvasive bladder cancer compared with PUNLMPs. These findings underline why the distinction between both entities is necessary since some pathologists consider PUNLMP to be completely benign, while others consider PUNLMPs to belong to the LG pTa group.

The WHO/ISUP 1998 grading system also introduced a description and definition of hyperplasia and flat lesions of the bladder.

1.3. The WHO 2004/2016 classification systems

The WHO 2016 system is based on the WHO/ISUP 1998 classification and the WHO 2004 classification (Fig. 1) [8], which refined the criteria of WHO/ISUP 1998 [4]. The division of urothelial tumours into LG and HG tumours, as introduced in the WHO 2004 classification, remains part of the WHO 2016 system.

According to the WHO 2016 system, pTa and pT1 tumours are graded into LG and HG and all detrusor muscle-invasive urothelial carcinomas are considered to be HG tumours. pTa tumours do not invade the lamina propria (no lymphovascular invasion and distant metastasis). However, pT1 tumours do grow under the basement membrane into the lamina propria, and lymphovascular invasion and metastasis can be seen in these cases. In many instances pathologists will identify pT1 tumours as HG tumours, independently of their atypia.

Comparison WHO 1973–2016

![Comparison](image7.png)

Fig. 1 – Comparison of the World Health Organisation (WHO) 1973 and WHO 2004/2016 classification of urothelial carcinoma.

NILGC = noninvasive low-grade papillary urothelial carcinoma;
NIHGC = noninvasive high-grade papillary urothelial carcinoma;
PUNLMP = papillary urothelial neoplasms of low malignant potential.
Strong points of the 2016 classification are clear cut-offs between HG and LG tumours, along with clear, precise descriptions of each grade in order to obtain homogeneous groups of tumours. The system includes three distinct categories: PUNLMP, pTa LG, and pTa HG. It avoids the use of ambiguous grading, G1/2 or G2/3, that fail to place a tumour in a defined grade.

The term noninvasive has also been introduced in the WHO 2016 system to further differentiate noninvasive LG and HG papillary carcinoma from invasive urothelial carcinomas. Consequently, the classification of pTa HG carcinomas in the WHO 2016 system is expansive enough to encompass all HG tumours with similar biological properties. This is a clear advantage of the WHO 2016 compared with the WHO 1973 classification, where these tumours could be randomly distributed between G2 and 3.

1.3.1. WHO 2016 grades and histological groups

1.3.1.1. Papillary urothelial neoplasms of low malignant potential. The PUNLMP category is useful because it does not carry the name cancer. This is important in younger patients, in whom PUNLMP mostly presents, who would otherwise have to carry a diagnosis of cancer with all its psychological and financial consequences. If the criteria for PUNLMP are applied in the very strictest way the classification only pertains to a very restricted group of tumours. The papillae of this exophytic lesion are discreet, slender, and not fused, covered by mostly normal urothelial cells, with absent or minimal cytological atypia. There is a slight increase in the number of atypical cells, polarity is preserved, and nuclei can be increased when compared with normal tissue. Umbrella cells are mostly present. Mitosis, if present, is the exception, and only found in the basal layers. Necrosis is absent. The prognosis of these lesions is very good. Although there is undeniably a risk of recurrence, the risk of progression is low (Table 2).

It should be noted that in 2002 PUNLMP was a new, very controversial entity, and pathologists, as well as urologists, had to get used to it on a diagnostic level. Therefore, as pathologists became more familiar with the entity over time, the intraobserver and interobserver reproducibility improved and cohorts became more robust. As a result, it is likely that the rates found in later studies are more accurate due to an improved understanding of PUNLMP and the recruitment of more consistent cohorts.

Reports of the progression rate of PUNLMP are consistent: it was reported as 8% by Pan et al [11] in 2010, 8% in 2002 by Samaratunga et al [12], and 10% in 2015 by Compérat et al [13]. In Pich et al [14], the median disease-free survival was 76 mo and no patient with PUNLMP showed progression towards a higher-grade tumour. Similar findings were reported by Holmang et al [15], with no progression in his study of 95 patients.

All these data, considered together, show that PUNLMP are not completely benign lesions, may recur and, according to some published data, can show progression, even though some authors dispute the difference between PUNLMP and pTa LG tumours and consider them to belong in the same tumour group (Table 2). The management of PUNLMPs is not yet clear and further research is required to determine follow-up and length of surveillance. Meanwhile, long-term follow-up is recommended, similar to that for pTa tumours.

1.4. Noninvasive papillary LG urothelial carcinoma

LG pTa carcinoma is characterised by orderly-arranged papillae. Although variations in polarity and nuclear size, shape and chromatin distribution are not of primary importance, a specific cytological disorder exists. Mitosis is rare, and, if present, usually occurs in the lower half of the urothelium. The challenges presented by inclusion and tangential cutting may make it difficult to analyse architectural aspects, which can be mistaken for irregular urothelium, while fused glands may be overgraded. However, if different grades are detected in the same lesion, the tumour should be graded according to the highest grade observed. It should be noted that the histological differences between PUNLMP and LG pTa can sometimes be subtle.

1.5. Noninvasive papillary HG urothelial carcinoma

Some HG lesions are characterised by a completely disordered appearance at low magnification due to both cytonuclear and architectural disorganisation. The spectrum of pleomorphism is wide and ranges from moderate to marked, but consensus is to not subdivide this group of lesions into two subgroups. However, the degree of nuclear atypia is important: nuclei have prominent nucleioli and are pleomorphic, with frequent mitosis. Intraepithelial necrosis can be present. The thickness of urothelium can vary considerably; the papillae are fused and display an anarchic growth.

HG tumours must be considered to be aggressive lesions. HG tumours can be pTa, but also pT1–4. Urothelial carcinomas pT2–4 are generally classified as HG tumours according to the WHO 2004/2016 [3,8].

| Reference | Pan et al 2010 [11] | Compérat et al 2015 [12] | Pich et al 2001 [13] | Samaratunga et al 2002 [14] | Holmang et al 2002 [15] |
|-----------|---------------------|--------------------------|----------------------|--------------------------|------------------------|
| n         | 212                 | 42                       | 19                   | 26                       | 68                     |
| Recurrence rate (%) | 18                | 17                       | 47                   | 7                        | 0                      |
| Progression rate (%) | 8%                | 10%                      | Not done             | 8%                       | 14.7%                  |

* International Bladder Cancer Group defines nonmuscle invasive bladder cancer progression as an increase in T stage from carcinoma in situ or T1 (lamina propria invasion), development of T2 or greater or lymph node metastasis.
1.6 Carcinoma in situ

Carcinoma in situ (CIS) is considered to be a HG lesion. It has very variable characteristics. McKenney et al [16] described four different morphological aspects: large-cell pleomorphic type, large-cell nonpleomorphic, small-cell, which does not refer to a neuroendocrine differentiation, but describes cells smaller than in the large-cell group, and clinging denuded form. Also described are three types of cancerisation of the urothelium, where only parts of the urothelium show major atypia. These types are: pagetoid form, with single atypical cancerous cells in the urothelium, the undermining form, in which the basal layer is replaced by highly atypical cells showing all the characteristics of HG lesions, and the overriding form, where the upper layer is replaced [16].

CIS is often observed in the adjacent mucosa of pTa HG lesions. At a molecular level, HG pTa carcinomas resemble invasive urothelial carcinomas. Comparative genomic hybridisation studies have demonstrated deletions at 2q, 5q, 10q, and 18q and gains at 5q and 20q. HG tumours show a high level of progression of 50–65% and a recurrence rate of 14.8–80.7% [17–19].

2 Evidence acquisition

2.1 Intraobserver and interobserver variability and reproducibility

A major challenge for every histological grading system is the degree to which the subjectivity of the pathologist affects the classification. When introducing a new grading system, the comparison is always delicate, as pathologists must move away from the old, familiar, system and try to adopt the new one.

2.2 Comparison of reproducibility of WHO 1973 and WHO 2004 classification systems

Pathologists who work together show better reproducibility when grading tumours [20]. Furthermore, reproducibility is much greater when pathologists are using a familiar grading system, rather than a new one. A recent meta-analysis compared both the WHO 1973 and WHO 2004 systems for performance and reproducibility. The authors concluded that the intra- and interobserver variability was slightly less in the 2004/2016 classification, but that it did not show a better prediction of recurrence and progression [21].

The main criticism concerning the reproducibility of the 1973 WHO classification is the poorly-defined morphological criteria. No distinct cut-offs between the different tumour grades were suggested, and there existed a clear lack of reproducibility. The variability of G2 tumours ranged from 13% to 69% and the lack of reproducibility has been a major argument in favour of rethinking the grading system [7,21]. The inclusion of ambiguous grading such as G1/2 and G2/3 gave no clear guidance to clinicians on the course of treatment for such patients.

2.2.1 Recurrence and progression

2.2.1.1 pTa tumours. Several studies have tried to compare the classifications for progression and recurrence with the ultimate aim of finding practice changing results. Cao et al [22] examined 172 pTa carcinomas with good follow-up and demonstrated significant differences between LG and HG papillary carcinomas in recurrence-free survival (p = 0.01). The authors concluded that the 2004 WHO system was superior to the 1973 system for predicting clinical outcomes in patients with urothelial carcinoma, independent of pathologic stage, and underlined the usefulness in pTa tumours. Yin and Leong [23] demonstrated significant differences in recurrence between LGand HG pTa carcinomas (p < 0.5), but none with the WHO 1973 classification. The study of May et al [24] compared the prognostic implications of both WHO systems and showed a significant difference in 5-yr recurrence rate between LG and HG pTa carcinomas, but no significant difference in 5-yr progression-free survival (PFS). In a similar way, G1–3 WHO 1973 pTa tumours had significantly different 5-yr recurrence-free survival (RFS) rates as well as PFS.

Lokeshwar et al [25] reported a significant grade shift in pTa bladder cancers with the use of the WHO 2004 classification, which did not seem to correlate with disease progression. They pointed out the problem of both tumour categories being treated with a range of different treatments, as well as the frequent overtreatment of patients.

2.2.1.2 pT1. Only a few studies comparing the different grading systems in pT1 carcinomas exist. There seems to be some evidence that the 1973 WHO classification might add some benefit to predict survival of patients [22,26,27]. However, the study of Cao et al [22] failed to show a difference for the RFS, PFS, and overall survival between pT1 G2 and G3 carcinomas, while the study of van Rhijn et al [27] showed that G2 and G3 played a significant role as prognosticotor towards muscle invasive carcinomas. The paper of Otto et al [26] showed that the WHO 1973 was a good predictor for disease-specific survival in pT1 bladder carcinomas.

Campbell et al [28] demonstrated moderate interobserver variability (κ = 0.6) in the case of two pathologists when comparing bladder tumours according to the WHO 1998 classification.

In a study of 258 consecutive papillary urothelial carcinomas, Gonül et al [29] showed a correlation between tumour grades in both WHO 1973 and WHO 2004/2016 classification systems (Table 3). Overall agreement between the two pathologists in the study was higher using the WHO 2004/2016 classification than with the WHO 1973 system. Again, PUNLMP showed the lowest degree of agreement. If the diagnosis of PUNLMP was excluded, the kappa (κ) value increased from 0.41 to 0.84 [11].

2.3 Intraobserver variability

Pelucchi et al [30] reported the evaluation of 270 consecutive patients with a first episode of LG pTa tumours. The same pathologist determined the grade for each tumour as
LG pTa and G1 or G2 according to the WHO 1973 and WHO 2004 classification systems, respectively. The 5-yr PFS was 93% for the LG pTa group, 97.6% for the G1 group, and 93.3% for the G2 group, while the 5-yr RFs was 49.4% for the LG group, 62% for the G1 group, and 40% for the G2 group. They concluded that the WHO 1973 system was a better predictor of recurrence than the WHO 2004 system, but that both systems predicted the risk of progression with similar accuracy [13].

The WHO 2004 classification showed better reproducibility than the WHO 1973 system, with the lowest agreement achieved with PUNLMP versus LG pTa. Further training could have a positive impact on the grading of these lesions, especially in the PUNLMP category. HG lesions were the easiest to classify with less intra- and interobserver variability.

2.4. Heterogeneous lesions

Not all urothelial lesions are homogeneous and grade heterogeneity is common. A mixture of two grades in the same tumour have been reported in 3–43% of tumours [31,32]. The WHO/ISUP 1998 consensus meeting suggested reporting the highest grade of a lesion as, at that time, no published evidence existed on this topic [4]. The WHO 2004 system also recommended basing the grade of a heterogeneous tumour on the highest grade observed [8]. Furthermore, the arbitrary criterion to ignore a finding of less than 5% of HG tumour has an unknown impact. Very little research has been carried out to confirm these suggestions for grading [32].

Based on the 1998 WHO/ISUP classification system, Cheng et al [33] suggested a modified grading system and that the volume of a grade should be >5% to be considered for grading [16].

A recent paper from Gofrit et al [34] showed that 5% of resections had mixed grade tumours which he defined as LG tumours with 10% or less of a HG component. As long as the HG component is not higher than 10%, these tumours have an evolution like LG carcinomas.

According to the hypothesis that normal looking urothelium adjacent to CIS and tumours accumulates neoplastic molecular changes [35], the study by Downes et al [35] evaluated those mixed tumours at the morphologic, proteonomic, and molecular levels. The authors demonstrated that LG areas in mixed tumours already exhibited molecular changes associated with disease progression. These findings suggest that molecular changes in HG tumours occur at an early stage of tumour growth, even before the corresponding morphological features become visible, which underlines the current practice of grading papillary urothelial neoplasms based on the highest grade [35].

The distinction between LG and HG tumours is of major importance. In the case of borderline histology, other parameters such as urinary cytology, multifocality, size of the lesion, prior history, and recurrence can play a role in grading. If CIS is present, the tumour should be considered to be a HG lesion. Currently, no authoritative recommendations exist on how to report mixed lesions; neither the WHO 2016, nor the International Collaboration on Cancer Reporting (www.iccr-cancer.org), provide a clear statement on the subject. Using cut-offs (5% according to Cheng [33] 10% according to Gofrit [34]) is arbitrary and not validated; therefore, the pathologist should record a percentage and describe all morphological aspects.

3. Evidence synthesis

3.1. Molecular grading

Urothelial carcinomas are often heterogeneous cell populations, with different morphological aspects and clinical outcomes. Although grading is a major prognostic factor, its usefulness needs to be improved by stronger intra- and interobserver reproducibility in determining tumour grade.

Molecular markers have a role in grading and their use may help to produce a more accurate diagnosis. FGFR-3 and P53 (TP53) mutations have been recognised as key pathways in the genetic development of urothelial neoplasias. In the early 2000s, the first studies combining FGFR-3 and grading were published. They confirmed an association between FGFR-3 mutation and LG tumours (p < 0.0001), with FGFR-3 detected in 84% of G1 tumours [29]. FGFR-3 is one of the most frequently mutated oncogene in urothelial carcinomas. It seems to be associated with LG tumours, noninvasive stage, and low recurrence. In contrast, P53 mutations are linked to a high-tumour grade, advanced stage, and frequent recurrences. Molecular markers can help to refine the accuracy of a histological diagnosis.

3.2. Taxonomy and gene signatures in NMIBC

Since 2012 molecular grading has been refined, Sjödahl et al [36] introduced a five-group system, with the Urobasal group A, which is predominantly made of pTa LG tumours, clearly demonstrating better outcomes. Furthermore, Sjödahl et al [36] also demonstrated that NMIBC still have a urothelial differentiation, which they lose as they become more aggressive. In 2015, Patrchan et al [37] analysed pT1 carcinomas and demonstrated a molecular subtype classification with urobasal (32%), genomically unstable (58%), and squamous cell carcinoma-like (10%) [37]. They also showed that rapidly progressing T1 tumours were of
substrate genotypically unstable or squamous cell carcinoma-like and had high lymphocyte (CD3) levels. Another paper, by Yun et al [38], may show that molecular risk classifiers predict the risk of progression and response to Bacillus Calmette-Guérin and could help to identify the optimal management of HG T1 disease for each patient. Those pT1 carcinomas which progress seem to have a molecular signature close to pT2 carcinomas. Due to the still limited data on NMIBC, no internationally accepted recommendations exist for molecular prognostic or grading markers. Current management will rely on pathologist assessment, based on morphological aspects.

3.3. Markers

3.3.1. Tumour recurrence

A study by Compérat et al [39] compared the different grades of pTa tumours and the expression of MiB-1 and aurora kinase A, which are markers for early mitosis. The study found that aurora kinase A and MiB-1 combined with the WHO 2004 grading system formed a powerful predictor of tumour recurrence.

Another study found an FGFR-3 mutation in all G1 and G2 tumours and also showed that patients with the mutation developed fewer recurrences (p = 0.004). Van Rhijn et al [40] suggested a molecular grading system in tumours with an FGFR-3 mutation in combination with MiB-1 expression. The study showed that FGFR-3 mutation was present in 88% of G1 tumours, but in only 16% of G3 tumours. The authors concluded that molecular grading was a relatively simple and reproducible method.

3.3.2. Tumour progression

A study by van Oers et al [41] used tissue microassays to investigate the expression of Ki-67, p53, and CK20, as markers for disease progression, and SNaPshot analysis to detect FGFR-3 mutation. Abnormal CK20 staining was strongly associated with higher tumour grades. In a group of pTaG1 tumours, 59% presented with abnormal CK20 staining, whereas 82% carried the FGFR-3 mutation. In a group of bladder tumours with a normal CK20 pattern, the FGFR-3 gene was mutated in 89%, whereas FGFR-3 mutations were only found in 37% of cases with abnormal CK20 staining. All markers were strong predictors for survival in univariate studies. In multivariate analysis, they were not independent from classical pathological parameters. None of the molecular markers was significantly associated with tumour recurrence [28].

The same research group also performed a prospective study linking the WHO 1973 and 2004 grading systems to Ki-67 and CK20 expression and to FGFR-3 mutation [42]. The study showed that both grading systems remained statistically significant independent predictors of progression (p = 0.005). However, the combination of WHO 2004 grade and FGFR-3 mutation status allowed improved risk stratification for patients with HG nonmuscle-invasive tumours [43].

Van Rhijn et al [44] proposed an alternative grading to validate the European Organisation for Research and Treatment of Cancer risk score, taking into account both MiB-1 expression and FGFR-3 mutation status. FGFR-3 mutations were significantly related to favourable disease. However, MiB-1 overexpression was seen in pT1 HG tumours with high European Organisation for Research and Treatment of Cancer risk scores [44].

It is clear that molecular markers can add benefit to a molecular grading system and research is ongoing. Until such data are available, no authoritative recommendations can be made. At the moment these, and other, markers are under investigation and only few specialised centres use them in daily practice [45–47]. None is recommended in the EAU guidelines [1].

The latest research in molecular classification suggests that markers might eventually become more important for determining prognosis and treatment than tumour grading. Sjödahl et al [48] has used molecular classification to identify five subtypes of tumours, while Choi et al [49] has identified three subgroups, which will impact treatment decisions. It should be noted that more data are available for muscle-invasive bladder cancer.

3.4. New concepts in pT1 HG tumours

The WHO 2016 classification system divides pT1 tumours into LG and HG tumours, but it is increasingly recognised that most pT1 tumours are probably HG [3]. The decision to stratify pT1 carcinoma as LG was taken because while they often recur they show progression in only 5% of cases [4].

However, tumours that invade the lamina propria have an aggressive potential and can develop metastasis. In addition, it is difficult to avoid either up-staging or down-staging these lesions, as shown in a recent study by Compérat et al [20], which highlighted the difficulty of a group of experienced genitourinary pathologists in detecting an invasive tumour. Kappa (κ) score for interobserver reproducibility ranged from 0.42 to 0.6 (mean: 0.49), demonstrating the difficulty of staging correctly.

The WHO 2016 includes substaging for pT1 tumours, which is a better prognostic factor than grade. The International Collaboration on Cancer Reporting committee recommends the provision of some measurement of volume, or depth of invasion (depth in mm), and/or total maximum dimension of invasive tumour (mm) and/or invasion superficial to, or involving, the muscularis mucosa, and/or deep to muscularis mucosae (the last item is equivalent to pTa1/b).

At a molecular level, it has been proven that these lesions are mostly aneuploid, although diploid tumours may exist. FGFR-3 mutations are common. Recurrence is relatively frequent, and about 10% of patients will show 50% progression towards a higher grade of tumour or stage of disease. Disease-specific mortality between 0.5% and 4% has been reported.

In the WHO 1973 system, invasive urothelial carcinomas can theoretically be considered as G1–3 [4]. It is agreed that no invasive carcinoma can be G1. Since these neoplasms are very heterogeneous, it is easily possible for
sampling to miss the presence of higher-grade carcinoma during biopsy.

Finally, it should be noted that muscle-invasive bladder cancers should always be considered HG.

3.4.1. Histological subtypes with deceptively-benign aspects

These types of urothelial carcinomas are sometimes difficult to stage, because they display a bland invasive front. It is important to recognise them, but also to be aware of the problems in confirming an eventual invasion, especially in case of superficial, detrusor lacking biopsies. Some of these tumour variants have very bland aspects, although they are aggressive tumours, which is probably one of the reasons why, historically, some invasive urothelial carcinomas were considered LG.

3.4.1.1. Nested carcinoma. Nested carcinoma was first described in 1979 by Stern [50], who did not realise it was malignant. It took 10 yr for nested carcinoma to be recognised as malignant.

Many pathologists considered the nested variant to be a LG tumour because of its low mitotic activity and mild pleomorphic aspects. In case of only a few tumour nests, which are not invading the detrusor muscle, this may be the case. However, this lesion has been reported to have an aggressive behaviour, and should be considered HG according to the WHO 2004 classification [8,20,51].

3.4.1.2. Urothelial carcinoma with small tubules and microcystic form. A similar problem exists for urothelial carcinoma with small tubules and the microcystic form. Although only a few cases have been reported, and it is therefore difficult to make recommendations, these carcinomas should be considered aggressive as soon as they have invaded the lamina propria [52].

3.4.1.3. Inverted variant of urothelial carcinoma. Urothelial carcinoma with inverted papilloma-like endophytic growth also belong to the group of benign-looking carcinomas. Borders are regular and pathologists may have the impression of tangential cutting. Furthermore, solid pushing borders can mask the invasiveness of the tumour, even while it is destroying the bladder wall, and even muscle-invasive tumours can be difficult to recognise. Nevertheless, these carcinomas are aggressive as soon as they invade the lamina propria and must be considered as HG [53].

One of these pushing inverted entities, known as large-nested urothelial carcinoma, has been described by Cox and Epstein [54]. Atypia in this entity are nearly completely absent. However, this tumour can invade the detrusor muscle and perivesical fat and is considered to be a HG tumour (Figs. 2 and 3) [55].

4. Conclusions

Bladder cancer grading classifications are not perfect. The EAU NMIBC Guidelines recommend that both the WHO 1973 and WHO 2004/2016 classifications should be used [1], until further validation of the 2004/2016 grading classification. The WHO 1973 and the 2004/2016 systems each have their advantages and disadvantages. A recent systematic review by the NMIBC Guidelines Panel confirmed these findings [21].

A major advantage of the 2004/2016 classification system is that it avoids ambiguous grades and the groups are clearly classified by histological descriptions. This may be why one can expect a higher intraobserver reproducibility in due time.

The clinical behaviour of NMIBC is closely related to the tumour differentiation/grade. The addition of molecular markers will be a step forward on the path to more personalised medicine. It is important that urologists understand the strengths and weaknesses of the classification used to grade their specimens. Pathologists and urologists are partners in providing the best treatment for patients, and are required to work closely together by sharing experience and research.
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