The majority of urothelial neoplasms has an exophytic growth pattern, yet some show an inverted architecture [1-3]. A systematic approach to classification of inverted/endophytic urothelial lesions was made in 2012 by the International Consultation on Urologic Disease (ICUD). According to the existing World Health Organization (WHO)/International Society of Urologic Pathology (ISUP) system criteria for exophytic papillary neoplasms and on the basis of the presence and degree of atypia (including assessment of polarity), inverted neoplasms are graded as (1) inverted papilloma (IP), (2) inverted papillary urothelial neoplasm of low malignant potential (PUNLMP), (3) inverted papillary urothelial carcinoma (PUC), (4) low-grade, non-invasive, inverted PUC, (5) high-grade, non-invasive, inverted PUC, or (6) high-grade, invasive [4]. Due to the frequent occurrence of both exophytic and endophytic patterns in the same urothelial lesion, such terminology should apply only to malignant lesions with prominent inverted architecture [5]. However, this approach has been criticized for not considering other architectural and cytological features, namely presence of exophytic papillary structures, type of endophytic pattern (i.e., nests and trabeculae), number of cellular layers, and mitotic index [2,5].

Urothelial carcinoma with an inverted growth pattern (UC-IGP) is a malignant entity within this spectrum of lesions. While efforts have been made over the last few decades to unravel its carcinogenesis and relationship with conventional urothelial carcinoma, the exact classification of inverted urothelial lesions is a matter of debate. The morphological features of UC-IGP pose several issues in differential diagnosis with other mostly benign lesions. Various techniques, including immunohistochemistry, UroVysion, and many molecular methods, have been employed to study the exact nature of this lesion. The aim of this review is to provide a comprehensive overview of the morphological and immunophenotypical aspects of UC-IGP. Moreover, we present and discuss the immunohistochemical and molecular markers involved in diagnosis and prognosis of UC-IGP lesions.

**Key Words:** Urothelial carcinoma; Inverted growth pattern; Immunohistochemistry; Molecular markers

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**Corresponding Author:** Francesca Sanguedolce, MD, PhD, Pathology Unit, University Hospital OO.RR., University of Foggia, Viale Pinto 1, 71122 Foggia, Italy  
Tel: +39-0881736315, Fax: +39-0881736334, E-mail: francesca.sanguedolce@unifg.it
MORPHOLOGICAL FEATURES

According to the current WHO classification, papillary urothelial carcinoma with an inverted growth pattern (PUC-IGP) is a variant of non-invasive PUC and is staged as pTa [6]. Unlike conventional PUC, PUC-IGP shows an endophytic architectural pattern with branching and anastomosing cords of urothelium, some of which have an expansile appearance [7,8]. The stromal-epithelial interface has a smooth profile with delicate vascular architecture. The cytological and architectural features lead similar grading as for conventional UCs, namely low-grade (LG) and high-grade (HG) [8], featuring nuclear atypia, architectural abnormality, and mitotic activity [9,10]. Such changes are present at the surface of the lesion in most cases, further supporting a diagnosis of UC. An exophytic papillary element has been reported in association with the inverted component [1,11], as well as a pseudoexophytic pattern resulting from artifactual fragmentation of the specimen.

The presence of prominent endophytic growth can be misdiagnosed as a pushing border of invasion, yet occasional true lamina propria invasion is supported by a stromal reaction [1,5,7,12] and/or neoplastic cords interweaving with fibers of muscularis mucosae [1,13]. Features such as irregularity of the endophytic nest profile, architectural complexity, and occurrence of single-cell invasion can be useful and should raise suspicion of an invasive lesion [14]. Transurethral resection (TUR)–related artifacts, namely tangential sectioning, cauterization, and crush effect, represent further issues in assessing stromal and/or muscular invasion [1,13].

In a recent large series of invasive HG-UC arising in a background of UC-IGP from various sites, Gutierrez et al. [15] reported on bladder tumors presenting at earlier stages (81% pT1) than those involving the upper urinary tract (80% and 43% ≥pT2 in the renal pelvis and ureter, respectively). UC in situ and variant histology were described in approximately 40% and 20% of all cases, respectively [15], the latter being associated with a more aggressive clinical behavior. Conversely, a previous study on 81 non-invasive LG-UC of the bladder, including eight UC-IGP, reported a lower recurrence risk in the inverted group [16]. A first attempt to classify PUC-IGP was conducted in 1997 by Amin et al. [13], who described two main histologic patterns featuring interanastomosing cords and trabeculae (IP-like pattern) and broad bulbous borders (broad-front pattern), respectively [13,17].

LG-PUC-IGP shows mild nuclear atypia in terms of irregular chromatin distribution, enlarged irregular nucleoli, expansile growth with inverted nests and clusters, and increased mitoses [8,18]. HG-PUC-IGP has predominant inverted growth with higher architectural disorder in terms of marked loss of polarity with respect to the basement membrane [4], along with significant nuclear pleomorphism and increased mitotic activity with occasional atypical figures. However, many reported cases of “atypical inverted urothelial papilloma” are described with an exophytic papillary component and significant atypia and/or mitoses, which would best be considered UC with inverted growth [5,19].

IMMUNOHISTOCHEMICAL MARKERS

CD44

CD44 is a stem cell surface marker typically present in the basal layer of normal urothelium; however, UC in situ and the luminal subtype of invasive UC lack CD44 expression [8]. In their recent study on UC-IGPs of various grade, Bang et al. [2] described CD44 expression in two-thirds of their LG cases, while all HG tumors were negative. Further studies are needed to assess the potential of CD44 as a diagnostic and prognostic marker in this setting.

Cytokeratin 20

Cytokeratin 20 (CK20) is a low-molecular weight cytokeratin with diagnostic and prognostic potential in urothelial lesions [20]. CK20 is expressed commonly by superficial cells only in the normal urothelium; therefore, it is a marker of urothelial maturation and differentiation [5]. The immunohistochemical expression of CK20 is of diagnostic value in differentiating IP from UC-IGP [1-3] (Table 1). Moreover, Sun et al. [3] reported that combining Ki67 and CK20 assessment by immunohistochemistry with UroVysion fluorescence in situ hybridization (FISH) showed sensitivity and specificity as high as 89.5% and 100%, respectively.

Cyclin D1

Cyclin D1 is a key regulator of the cell cycle, and its alterations have been implicated in bladder carcinogenesis [21]. Cyclin D1 status has been studied as a prognostic marker in nonmuscle invasive bladder cancer (NMIBC), with conflicting results [21] (Table 1). The LG-IP-like UC reported by Sudo et al. [22] showed cytoplasmic expression of cyclin D1 along with other immunohistochemical markers. Interestingly, Bang et al. [2] found that 28 of 60 (47%) inverted urothelial neoplasms were positive for nuclear cyclin D1, in the absence of significant difference in stain-
ing levels between benign and malignant lesions. Accordingly, cyclin D1 nuclear expression was higher in LG-UC-IGP than in IP (73.7% vs. 69.4%, respectively) in the study by Sun et al. [3], but the difference did not achieve statistical significance (p = .798). Based on these findings, cyclin D1 seems not to be helpful in differential diagnosis of benign and malignant inverted lesions.

**Table 1. Expression of selected immunohistochemical markers in UC-IGP**

| Study                  | Site (No.)                        | CK20 | Cyclin D1 | HER2 | Ki67 | p16 | p53 |
|------------------------|-----------------------------------|------|-----------|------|------|-----|-----|
| Cheon et al. [29]      | Bladder (2)                       | -    | -         | 2/2  | -    | -   | -   |
| Eiber et al. [31]      | Bladder (23)                      | 9/22 | -         | -    | 18/22| -   | -   |
| Jones et al. [1]       | Bladder (29)                      | 17/29| -         | -    | 19/29| -   | -   |
| Terada [32]            | Bladder (3)                       | -    | -         | -    | 3/3  | -   | -   |
| Ehsani et al. [30]     | Renal pelvis (23)                 | -    | -         | -    | 15/23| -   | -   |
| McDaniel et al. [18]   | Bladder (8), renal pelvis (1)     | -    | -         | -    | 3/5  | -   | -   |
| Sun et al. [3]         |                                   | 14/38| 28/38     | -    | -    | -   | -   |
| Bang et al. [2]        | Bladder (19)                      | 4/15 | 3/4       | 1/15 | 2/4  | 4/15| 3/4 |

Values are presented as number (%). UC-IGP, urothelial carcinoma with an inverted growth pattern; CK20, cytokeratin 20; HER2, human epidermal growth factor receptor 2; HG, high-grade; LG, low-grade.

**Human epidermal growth factor receptor 2**

Human epidermal growth factor receptor 2 (HER2) is a tyrosine kinase receptor with oncogenic potential that is overexpressed in 5%–10% of UC, including nonmuscle invasive lesions [23-25]. HER2 expression has prognostic and therapeutic implications, since anti-HER2 targeted drugs are the standard-of-care for patients with HER2+ breast and gastroesophageal cancer [26,27]. However, analytical and pre-analytical issues can affect the reliability of HER2 assessment through current methods, especially with TUR specimens [28]. Cheon et al. [29] found moderate to strong HER2 overexpression (defined as distinct membrane staining) in two malignant bladder lesions with inverted pattern compared to five IP, which were negative for HER2. Similarly, significantly higher levels of HER2 expression (p = .0465) were described in UC-IGP compared with IP and PUNLMP-IGP using a 10% cutoff in a recent study by Bang et al. [2]. In addition, Ehsani and Osunkoya [30] reported a HER2 positivity rate (defined as strong complete membrane staining in > 30% cells) as high as 74% in their series of 46 renal pelvis UC cases; of them, 23 UC-IGP showed HER2 overexpression in 65% of cases, mostly HG-UC. Based on these results (Table 1), HER2 might be useful in differentiating malignant from benign lesions with an inverted growth pattern.

**Ki67**

The proliferative index assessed by scoring Ki67 nuclear antigen has been studied extensively in both NMIBC and muscle invasive bladder cancer (MIBC) [21] and in inverted lesions as well. Overall, it has been reported as diffusely expressed in UC-IGP [1,2,22,31,32] with significantly higher level compared to that in benign inverted lesions [1-3,31]. Consistent with these results, assessment of proliferative activity using proliferating cell nuclear antigen antibody and AgNOR silver colloid staining yielded higher expression rates of both markers in malignant inverted lesions compared to IPs [29]. Eiber et al. [31] suggest that a combined assessment of fibroblast growth factor receptor 3 (FGFR3) mutation status and Ki67 proliferation index can yield a specificity as high as > 90% in differentiating UC-IGP from IP within a consistent histological setting, with a Ki67 labeling index < 5% and wild-type FGFR3 being associated with a benign lesion. It has been suggested that the rate of Ki67 positive cells tends toward constant growth and expansion at the center of the lesion as grade increases [1,33]; therefore, this marker might be an adjunct in disease grading (Table 1).

**p16**

In bladder cancer, p16 has been analyzed either in association with other cell cycle proteins as a prognostic/predictive factor or as an indirect marker of human papillomavirus–induced carcinogenesis [21]. The two studies assessing p16 expression in UC-IGP yielded overlapping results, with higher expression of p16 in malignant compared to benign lesions, without statistical significance (Table 1). Furthermore, in both studies, a more diffuse staining pattern was described in HG-UC versus LG-UC [2,18].

**p53**

The gene encoding tumor suppressor protein p53 is the most common target for mutations in human cancer, and alterations of p53 at a molecular level are found in early bladder carcinogenesis.
events. p53 status seems to have a prognostic role in both NMIBC and MIBC, yet with contradictory results [21,34,35]. Although usually overexpressed in UC-IGP [22,32], no significant difference in p53 expression between benign and malignant inverted lesions was reported using different cutoffs (10%–50%) [2,31]. Conversely, Sun et al. [3] reported significantly higher p53 staining (cutoff, 10%) in a series of LG-UC-IGP compared to IP (p = .001).

Similarly, Jones et al. [1] described a steady increase in p53 expression in UC-IGP versus IP (59% vs. 7%) using a 1% cutoff, suggesting that p53 should be part of a multi-marker panel (along with Ki67 and CK20) to distinguish benign from malignant inverted lesions. Interestingly, there were no statistically significant differences in p53 protein staining between IPs in patients with and without a history of UC [36] and between IPs with and without atypia [37] in two previous studies (Table 1). The use of different cutoffs can impair the reproducibility of immunohistochemical results among studies.

**MOLECULAR FEATURES**

**RAS genes**

HRAS and KRAS are prototype RAS oncoproteins that have been shown to infrequently incur mutations in conventional UC [21,35]. Conversely, HRAS mutations have been reported in inverted tumors [18], along with mutations in other members of the RAS pathway, namely mutations encoding the KRAS G12R and BRAF G469A mutants in LG-UC-IGP and HG-UC-IGP cases, respectively [18]. Moreover, an oncogenic HRAS or KRAS missense mutation was present in nearly all cases of IP and urothelial papilloma according to a recent series [38-40], compared to two of 25 UC-IGP cases (8%). In both cases, further oncogenic mutations in chromatin-modifying genes and/or cell cycle regulators were present. Based on such findings, it has been suggested that an altered RAS pathway supplies the growth and/or progression of inverted urothelial lesions [18].

**Loss of heterozygosity**

Chromosomal aberrations, namely changes of copy numbers of various genetic regions, can occur at several points along the UC pathway that can be detected by cytogenetic studies, including loss of heterozygosity (LOH) analysis [21]. Sung et al. [41] found a very low incidence of LOH at genetic loci, which are frequently lost in both UCs and PNULMPs, by examining four polymorphic microsatellite markers [42] in their series of 39 IPs. LOH analysis performed by Eiber et al. [31] on 62 IPs and 23 UC-IGPs using microsatellite markers at chromosomal loci 9p21, 9q, and 17p13.1 identified statistically significant differences in 9q LOH (13.2% of IPs vs. 36.4% of UC-IGPs, p = .03).

**Telomere shortening**

Telomeres are repetitive DNA sequences that protect chromosome ends. A process known as telomere shortening (TS) occurs with every iteration of DNA replication and cell division. Telomerase is a DNA polymerase that counteracts TS by repairing chromosome ends, and the expression of its catalytic subunit telomerase reverse transcriptase (TERT) is correlated with telomerase activity [35]. TS and telomerase activity are involved in cancer development and progression at different sites, including the bladder [43]. Williamson et al. [44] reported that relative telomere length, as assessed by FISH analysis with a telomere-specific peptide nucleic acid probe, was significantly reduced in UC-IGP compared to IP (p = .001). Interestingly, analysis of relative telomere signal intensity in normal urothelium, non-neoplastic lesions (cystitis glandularis), and IP yielded similar results [44]. Based on these results, the authors argued that IPs do not have premalignant potential, and that benign and malignant inverted lesions develop through different carcinogenetic pathways despite their morphological similarity.

**TERT** promoter mutations have been detected infrequently in IPs [39,45,46]. Similarly, Cheng et al. [10] identified a significantly lower rate of TERT mutations in IP compared with UC-IGP (15% vs. 58%, p = .003). Interestingly, the same C228T mutation was found in inverted lesions as well as in the majority of conventional UC, suggesting that a subset of IP might share a molecular pathway of carcinogenesis with UC-IGP and conventional UC [47]. Such findings suggest the use of TERT mutation analysis in the differential diagnosis of benign and malignant lesions [38,40,47-49].

**UroVysion**

UroVysion is a multicolor FISH-based urine assay with higher sensitivity than urine cytology in the setting of UC screening and follow-up [21]. This assay assesses amplification of chromosomes 3, 7, and 17 along with deletion of 9p21. Such alterations have been detected in up to 79% of UC-IGPs through different studies [1,5,7,33]. In their multimethod study on a series of 15 IPs and 29 UC-IGPs, Jones at al. [1] yielded normal results for all cases of IP, while UC-IGP demonstrated chromosomal abnormalities typical of conventional UC, including gains of chromosomes 3 and 7. Accordingly, UroVysion positivity, defined as a gain of at least two of chromosomes 3, 7, or 17 or a homozygous loss of 9p21, yielded increased sensitivity and specificity rates
Microsatellite instability

Microsatellite instability (MSI) resulting from errors in DNA replication is a distinctive feature of several tumors, including BC, where it has been reported to be associated with increased grade and stage [21] and to have a predictive role, especially in NMIBC [25]. MSI status can be assessed either directly by microsatellite analysis, a PCR-based technique, or indirectly by assessing the MSI factors MutL homologue 1 (MLH1), MutS homologue 2 (MSH2), and MutS homologue 6 (MSH6).

A frequent association between MSI positive status and inverted growth pattern in tumors of the upper urinary tract has been described [50,51], mostly highlighted by MSH2 and/or MSH6 protein loss. Eiber et al. [31] investigated MSI in their series of IPs and UC-IGPs through both microsatellite analysis and MLH1, MSH2, and MSH6 immunostaining; however, they failed to find a significant correlation between MSI status and diagnosis [31]. Based on these findings, they argued that microsatellite unstable inverted tumors of the upper urinary tract represent a distinct subgroup of inverted urothelial tumors.

FGFR3

FGFR3 is a member of the family of fibroblast growth factor tyrosine kinase receptors that is involved in urothelial carcinogenesis through a papillary pathway associated with low cellular grade and lack of invasion [52]. Alterations of the FGFR3 gene have been described in UC-IGP, namely activating mutations encoding S249C, R248C, and G370C [18]. Since such alterations have been detected in both LG-UC-IGP and HG-UC-IGP, a possible explanation could be the non-invasive nature of inverted lesions overall, despite their grade of differentiation [18].

Distinct molecular alterations in inverted urothelial lesions

Evidence from molecular genetic studies has shown that IPs have additional distinct molecular features compared to their malignant counterparts, such as low tumor mutational burden, mutations in the mitogen-activated protein kinase/ERK pathway, along with a lack of the prevalent APOBEC mutation signature [1,7,31,36,38-41]. On the other hand, both conventional UC and UC-IGP carry overlapping genetic alterations [53], namely mutations in FGFR3, TP53, CDKN1A, PIK3CA, FBXW7, ERBB2, and NOTCH1 [18,40]. Interestingly, the specific point mutations at FBXW7 R505, ERBB2 V842, and NOTCH1 R1594 have not been reported in conventional urothelial cancers [54]. UC-IGP arising in the upper urinary tract has been described in association with hereditary non-polyposis colorectal cancer syndrome/Lynch syndrome, with DNA mismatch repair gene abnormalities and MSI [55], and the latter are more frequent in tumors of the renal pelvis and ureter than in bladder primaries and can be used as prognostic markers [25].

Differential diagnosis

Several authors have highlighted the high potential for misinterpretation of UC-IGP as IP due to overlapping morphological features [1,7,11-15,22,44]; however, a combination of morphologic, immunohistochemical, and molecular genetic assessments can be helpful in achieving a correct diagnosis [1]. This task is particularly challenging when the cystoscopy-obtained biopsy tissue is limited, extensively fragmented, heavily inflamed, and/or obscured by crush or cautery artifacts [15]. IPs are relatively less frequent than conventional urothelial benign papillary neoplasms, can be encountered anywhere throughout the urinary tract, and do not undergo malignant transformation [56].

Amin et al. [13] analyzed 18 UC-IGPs and established several morphologic criteria to distinguish them from IPs. UC-IGP tends to have an exophytic papillary surface, thick irregular cords or trabeculae, grade-dependent cytological atypia, and decrease to lack of maturation, spindling, or peripheral palisading [2,3,5,10,18,32]. Cytologic atypia presents in the form of nuclear pleomorphism, irregular chromatic structure, and/or enlarged uneven nucleoli [57]. Furthermore, UC-IGPs tend to have greater mitotic activity above the basal layer with occasionally atypical mitotic figures [3,57]. The presence of UC in situ in the surface urothelium is a further hint to diagnosis of UC-IGP [7].

Conversely, IPs have a smooth, dome-shaped surface due to the endophytic growth of uniform cords and trabeculae, usually lack an exophytic element, are more circumscribed, feature palisading at the periphery and spindling or streaming in the center of the trabeculae, and cytologic atypia is weak to absent [3,31]. It is clear that IPs lack stromal invasion; however, IPs with foamy or vacuolated cytoplasm have been described occasionally [19]. At cystoscopy, IP usually appears as a single peduncle mass with smooth surface, while UC-IGP presents as wide-based, cauliflower-like multiple masses with an uneven surface [5] (Fig. 1).

It has been suggested that most lesions diagnosed in the past as IP with concurrent UC were actually UC-IGPs [58], and some IPs were labeled as LG-UC-IGPs [59], resulting in confusion regarding the actual incidence of each disease. Moreover, a cate-
gory of “IP with atypia” or “atypical IP” was introduced by some authors, referring to a subset of IP with malignant potential [36,37]. Compared to UC-IGP, IP with atypia has lower mitotic index and proliferative activity. Interestingly, Brimo et al. reported on a series of 12 UC-IGPs encompassing areas within the tumor whose morphological features were identical to those of IP in each case [17]. Immunohistochemically, higher rates of p53 and CK20 expression and increased Ki67 proliferative index were seen in UC-IGP compared to IP [1,7,14,44]. Sun et al. [3] described higher Ki67 and CK20 expression in LG-UC-IGP than IP. Interestingly, Broussard et al. [37] found a higher incidence of Ki67 and p53 in IPs with atypia compared to conventional IPs. Moreover, 24 of 38 (63.2%) LG-UC-IGP cases were positive for UroVysion FISH, whereas all IPs showed no gains of chromosomes 3, 7, or 17 and absence of 9p21 loss, suggesting that IP arises from pathogenetic mechanisms that differ from those that produce UC (Fig. 1).

As a lesion with an endophytic growth pattern, a large-nested variant of invasive UC enters the differential diagnosis with UC-IGP. However, involvement of the detrusor muscle as well as the high variability of size and shape, irregular profile, and infiltrative architecture of large-nested UC allow distinction between the two [60].

**CONCLUSION**

In conclusion, UC-IGP and IP are distinct entities with peculiar biologic behaviors and clinical outcomes that can be difficult to distinguish due to their morphological commonalities. However, misdiagnosis should be avoided since IP is a benign disease, while UC-IGP can warrant further treatment or surveillance depending on grade [2,7,56]. Data from the literature assess the role of ancillary techniques, namely immunohistochemistry and FISH, in supporting a proper diagnosis [3]. However, studies on large case series are warranted to further elucidate the molecular mechanisms and diagnostic and prognostic markers of UC-IGP.

**Ethics Statement**
Not applicable.

**Availability of Data and Material**
Data sharing not applicable to this article as no datasets were generated or analyzed during the study.

**Code Availability**
Not applicable.

**ORCID**
Francesca Sanguedolce https://orcid.org/0000-0001-7459-3521
Beppe Calò https://orcid.org/0000-0001-9285-9426
Marco Chirico https://orcid.org/0000-0002-0637-1481
Ugo Falagario https://orcid.org/0000-0002-1152-3005
Gian Maria Busetto https://orcid.org/0000-0002-7291-0316
Magda Zanelli https://orcid.org/0000-0002-8733-9933
Alessandra Bisagni https://orcid.org/0000-0001-8484-7871
Maurizio Zizzo https://orcid.org/0000-0002-4516-9040
Giuseppe Carrieri https://orcid.org/0000-0002-6609-6434
Luigi Cormio https://orcid.org/0000-0002-1126-5368

**Author Contributions**
Conceptualization: FS. Project administration: FS, MZ (Magda Zanelli), LC. Supervision: GC, LC. Writing—original draft: FS, BC, MC, UF, GMB, MZ (Magda Zanelli), AB, MZ (Maurizio Zizzo), SA. Writing—review & editing: FS, SA, GC, LC. Approval of final manuscript: all authors.

**Conflicts of Interest**
The authors declare that they have no potential conflicts of interest.

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