available at www.sciencedirect.com journal homepage: www.europeanurology.com



### <sup>2</sup> Review – Bladder Cancer

- <sup>3</sup> European Association of Urology Guidelines on
- Non–muscle-invasive Bladder Cancer (Ta, T1, and
- <sup>5</sup> Carcinoma in Situ)



### <sup>6</sup> Q3 Marko Babjuk <sup>a,b,\*</sup>, Maximilian Burger <sup>c</sup>, Otakar Capoun <sup>d</sup>, Daniel Cohen <sup>e</sup>, Eva M. Compérat <sup>f</sup>,

Q5

<sup>7</sup> José L. Dominguez Escrig<sup>g</sup>, Paolo Gontero<sup>h</sup>, Fredrik Liedberg<sup>i,j</sup>, Alexandra Masson-Lecomte<sup>k</sup>,

<sup>8</sup> A. Hugh Mostafid<sup>1</sup>, Joan Palou<sup>m</sup>, Bas W.G. van Rhijn<sup>c,n</sup>, Morgan Rouprêt<sup>o</sup>,

<sup>9</sup> Shahrokh F. Shariat<sup>*a,b*</sup>, Thomas Seisen<sup>*o*</sup>, Viktor Soukup<sup>*d*</sup>, Richard J. Sylvester<sup>*p*</sup>

<sup>10</sup> Q4 <sup>a</sup> Department of Urology, Teaching Hospital Motol and 2nd Faculty of Medicine, Charles University Praha, Prague, Czech Republic; <sup>b</sup> Department of Urology, 11 Comprehensive Cancer Center, Medical University Vienna, Vienna General Hospital, Vienna, Austria; <sup>c</sup> Department of Urology, Caritas St. Josef Medical Center, 12 University of Regensburg, Regensburg, Germany; <sup>d</sup> Department of Urology, General Teaching Hospital and 1st Faculty of Medicine, Charles University Praha, 13 Prague, Czech Republic; e Department of Urology, Royal Free London NHS Foundation Trust, Royal Free Hospital, London, UK; f Department of Pathology, 14 Tenon Hospital, AP-HP, Sorbonne University, Paris, France; <sup>g</sup> Department of Urology, Fundación Instituto Valenciano de Oncología, Valencia, Spain; 15 <sup>h</sup> Department of Urology, Città della Salute e della Scienza, University of Torino School of Medicine, Torino, Italy; <sup>i</sup> Department of Translational Medicine, Lund 16 University, Malmö, Sweden; <sup>1</sup> Department of Urology, Skåne University Hospital, Malmö, Sweden; <sup>k</sup> Department of Urology, Université de Paris, APHP, Saint 17 Louis Hospital, Paris, France; <sup>1</sup>Department of Urology, The Stokes Centre for Urology, Royal Surrey Hospital, Guildford, UK; <sup>m</sup> Department of Urology, Fundacio 18 Puigvert, Universitat Autònoma de Barcelona, Barcelona, Spain; <sup>n</sup> Department of Surgical Oncology (Urology), Netherlands Cancer Institute-Antoni van 19 Leeuwenhoek Hospital, Amsterdam, The Netherlands; °GRC 5 Predictive Onco-Uro, Department of Urology, Sorbonne University, AP-HP, Pitié Salpétrière 20 Hospital, Paris, France; P European Association of Urology, Arnhem, The Netherlands

#### Article info

#### Abstract

*Article history:* Accepted August 15, 2021

*Associate Editor:* James Catto

#### Keywords:

Bladder cancer Urothelial carcinoma Bacillus Calmette-Guerin (BCG) Radical cystectomy Cystoscopy Diagnosis Guidelines Follow-up Context: The European Association of Urology (EAU) has released an updated version of the guidelines on non-muscle-invasive bladder cancer (NMIBC). Objective: To present the 2021 EAU guidelines on NMIBC. 05 Evidence acquisition: A broad and comprehensive scoping exercise covering all areas of the NMIBC guidelines since the 2020 version was performed. Databases covered by the search included Medline, EMBASE, and the Cochrane Libraries. Previous guidelines were updated, and the level of evidence and grade of recommendation were assigned. Evidence synthesis: Tumours staged as Ta, T1 and carcinoma in situ (CIS) are grouped under the heading of NMIBC. Diagnosis depends on cystoscopy and histological evaluation of tissue obtained via transurethral resection of the bladder (TURB) for papillary tumours or via multiple bladder biopsies for CIS. For papillary lesions, a complete TURB is essential for the patient's prognosis and correct diagnosis. In cases for which the initial resection is incomplete, there is no muscle in the specimen, or a T1 tumour is detected, a second TURB should be performed within 2–6 wk. The risk of progression may be estimated for individual patients using the 2021 EAU scoring model. On the basis of their individual risk of progression, patients are stratified as having low, intermediate, high, or very high risk, which is pivotal to recommending adjuvant treatment. For patients with tumours presumed to be at low risk and for small papillary recurrences detected more

\* Corresponding author. Department of Urology, 2nd Faculty of Medicine, Charles University, Praha Motol University Hospital, V Úvalu 84, 15006 Praha 5, Czech Republic. Tel.: +42 224 434801; Fax: +42 224 967102.

E-mail address: marek.babjuk@fnmotol.cz (M. Babjuk).

https://doi.org/10.1016/j.eururo.2021.08.010

0302-2838/© 2021 European Association of Urology. Published by Elsevier B.V. All rights reserved.

Intravesical chemotherapy Prognosis Transurethral resection (TUR) **BCG** unresponsive European Association of Urology (EAU)

EUROPEAN UROLOGY XXX (2021) XXX-XXX

than 1 yr after a previous TURB, one immediate chemotherapy instillation is recommended. Patients with an intermediate-risk tumour should receive 1 vr of full-dose intravesical bacillus Calmette-Guérin (BCG) immunotherapy or instillations of chemotherapy for a maximum of 1 yr. For patients with high-risk tumours, full-dose intravesical BCG for 1-3 yr is indicated. For patients at very high risk of tumour progression, immediate radical cystectomy should be considered. Cystectomy is also recommended for BCG-unresponsive tumours. The extended version of the guidelines is available on the EAU website at https://uroweb.org/guideline/non-muscle-invasive-bladder-cancer/. *Conclusions:* These abridged EAU guidelines present updated information on the diagnosis and treatment of NMIBC for incorporation into clinical practice. Patient summary: The European Association of Urology has released updated guidelines on the classification, risk factors, diagnosis, prognostic factors, and treatment of nonmuscle-invasive bladder cancer. The recommendations are based on the literature up to 2020, with emphasis on the highest level of evidence. Classification of patients as having low, intermediate, or and high risk is essential in deciding on suitable treatment. Surgical removal of the bladder should be considered for tumours that do not respond to bacillus Calmette-Guérin (BCG) treatment and tumours with the highest risk of progression. © 2021 European Association of Urology. Published by Elsevier B.V. All rights reserved.

#### 21 Introduction 1.

22 6 This overview represents the updated European Association 23 of Urology (EAU) guidelines for non-muscle-invasive 24 bladder cancer (NMIBC), comprising Ta, T1, and carcinoma 25 in situ (CIS). The information presented is limited to 26 urothelial carcinoma, unless otherwise specified. The aim 27 is to provide practical recommendations for clinical 28 management of NMIBC, with a focus on clinical presenta-29 tion and recommendations.

30 It must be emphasised that clinical guidelines present 31 the best evidence available to the experts, but following 32 guideline recommendations will not necessarily result in 33 the best outcome. Guidelines can never replace clinical 34 expertise when making treatment decisions for individual 35 patients, but rather help to focus decisions that also take the 36 personal values and references/individual circumstances of 37 patients into account. Guidelines are not mandates and do 38 not purport to be a legal standard of care.

#### 39 2. **Evidence acquisition**

40 For the 2021 NMIBC guidelines, new and relevant evidence 41 has been identified, collated, and appraised through a 42 structured assessment of the literature.

43 A broad and comprehensive scoping exercise covering all 44 areas of the NMIBC guidelines since the previous version 45 was published in 2020 was performed. Excluded from the 46 search were basic research studies, case series, reports, and 47 editorial comments. Only articles published in the English 48 language and addressing adults were included. Excluded 49 from the search were basic research studies, case series, 50 reports, and editorial comments. Only articles published in 51 the English language and addressing adults were included. 52 A detailed search strategy is available online at https:// 53 uroweb.org/guideline/non-muscle-invasive-54 bladdercancer/?type=appendices-publications. 55

For sections dealing with staging, diagnosis, and 56 prediction, references cited in this text were assessed 57 according to their level of evidence (LE) according to the 58 2009 Oxford Centre for Evidence-Based Medicine (CEBM)

59 levels of evidence [1]. For sections on disease management 60 and follow-up, a system modified from the 2009 CEBM 61 levels of evidence is used.

62 For each recommendation in the guidelines there is an 63 accompanying online strength rating for which a modified 64 GRADE methodology was used. These key elements are the 65 basis that panels use to define the strength rating of each 66 recommendation. The strength of each recommendation is 67 represented by the word "strong" or "weak" [2].

#### 3. Epidemiology, aetiology, and pathology

#### 3.1. Epidemiology

Bladder cancer (BC) is the tenth most commonly diagnosed 70 cancer worldwide [3]. The age-standardised incidence rate 71 (per 100 000 person-years) is 9.5 for men and 2.4 for women 72 worldwide, and 20 for men and 4.6 for women in the EU [3].

73 Worldwide, the BC age-standardised mortality rate (per 74 100 000 person-years) was 3.3 for men versus 0.86 for 75 women [3]. The incidence and mortality of BC have decreased in some registries, possibly reflecting a decrease 76 in the impact of causative agents [4]. 77

Approximately 75% of patients with BC present with disease confined to the mucosa (stage Ta or CIS) or 80 submucosa (stage T1); for younger patients (<40 yr) this 81 percentage is even higher [5].

#### 3.2. Aetiology

83 Tobacco smoking is the most important risk factor for BC, 84 accounting for slightly less than 50% of cases [6] (LE: 3), 85 followed with occupational exposure to aromatic amines, 86 polycyclic aromatic hydrocarbons, and chlorinated hydro-87 carbons, which are responsible for approximately 10% of all 88 cases [4,7].

89 While family history seems to have little impact [8], 90 genetic predisposition has an influence on the incidence of 91 BC via its impact on susceptibility to other risk factors [9,10].

92 Exposure to arsenic in drinking water increases the risk 93 of BC and chlorination of drinking water and subsequent

Please cite this article in press as: Babjuk M, et al. European Association of Urology Guidelines on Non-muscle-invasive Bladder Cancer (Ta, T1, and Carcinoma in Situ). Eur Urol (2021), https://doi.org/10.1016/j.eururo.2021.08.010

2

68

69

78 79

#### Q1 Table 1 – 2017 TNM classification of urinary bladder cancer

T: Primary tumour	
Tx	Primary tumour cannot be assessed
ТО	No evidence of primary tumour
Та	Noninvasive papillary carcinoma
Tis	Carcinoma in situ: "flat tumour"
T1	Tumour invades subepithelial connective tissue
T2	Tumour invades muscle
T2a	Tumour invades superficial muscle (inner half)
T2b	Tumour invades deep muscle (outer half)
T3	Tumour invades perivesical tissue
T3a	Microscopic invasion
T3b	Macroscopic invasion (extravesical mass)
T4	Tumour invades any of the following: prostate stroma, seminal vesicles, uterus,
	vagina, pelvic wall, abdominal wall
T4a	Tumour invades prostate stroma, seminal vesicles, uterus or vagina
T4b	Tumour invades pelvic wall or abdominal wall
N: Regional lymph nodes	
Nx	Regional lymph nodes cannot be assessed
NO	No regional lymph node metastasis
N1	Metastasis in a single lymph node in the true pelvis (hypogastric, obturator,
	external iliac, or presacral)
N2	Metastasis in multiple regional lymph nodes in the true pelvis (hypogastric,
	obturator, external iliac, or presacral)
N3	Metastasis in common iliac lymph node(s)
M: Distant metastasis	
M0	No distant metastasis
M1a	Nonregional lymph nodes
M1b	Other distant metastases

levels of trihalomethanes are potentially carcinogenic [11]
 (LE: 3). A link between dietary habits and BC risk has been
 suggested [12,13].

Schistosomiasis and exposure to ionising radiation are
 associated with higher BC risk; a weak association was also
 suggested for cyclophosphamide and pioglitazone [11,14]
 (LE: 3).

#### <sup>101</sup> 3.3. Pathology

The information presented in this text is limited to
 urothelial carcinoma, unless otherwise specified.

#### <sup>104</sup> **4. Staging and classification systems**

#### <sup>105</sup> **4.1. Definition of NMIBC**

Papillary tumours confined to the mucosa and invading the 106 lamina propria are classified as stage Ta and T1, respectively, 107 according to the TNM classification system [15]. Flat, high-108 grade tumours confined to the mucosa are classified as CIS 109 (Tis). All of these tumours are grouped under the heading of 110 NMIBC. The term non-muscle-invasive BC, however, represents 111 a group definition; all tumours should be characterised 112 according to their stage, grade, and further pathological 113 characteristics. The term superficial BC should no longer be 114 used as it is incorrect.

#### <sup>115</sup> 4.2. TNM classification

The 2009 TNM classification approved by Union Interna tional Contre le Cancer was updated in 2017 (8th edition;
 Table 1) [15].

#### 4.3. T1 subclassification

Retrospective cohort studies have demonstrated that the<br/>depth and extent of invasion into the lamina propria (T1120substaging) is of prognostic value [16] (LE: 3). Use of T1121substaging is recommended by the 2016 World Health123Organization (WHO) classification [17]. The optimal system<br/>for substaging T1 remains to be defined [17,18].124

#### 4.4. CIS and its classification

CIS is a flat, high-grade, noninvasive urothelial carcinoma. It127can be missed or misinterpreted as an inflammatory lesion128during cystoscopy if not biopsied. CIS is often multifocal and129can occur in the bladder, as well as the upper urinary tract130(UUT), prostatic ducts, and prostatic urethra.131

From a clinical point of view, CIS can be classified as <sup>132</sup> follows: <sup>133</sup>

- Primary: isolated CIS with no previous or concurrent <sup>134</sup> papillary tumours and no previous CIS; <sup>135</sup>
- Secondary: CIS detected during follow-up of patients <sup>136</sup> with a previous tumour that was not CIS; or <sup>137</sup>
- Concurrent: CIS in the presence of any other urothelial <sup>138</sup> tumour in the bladder. <sup>139</sup>

### 4.5. Histological grading of non-muscle-invasive bladder urothelial carcinomas

In 2004 the WHO and the International Society of Urological <sup>142</sup> Pathology (ISUP) published and in 2016 updated a <sup>143</sup> histological classification of urothelial carcinomas that <sup>144</sup> provides a different patient stratification between <sup>145</sup>

Please cite this article in press as: Babjuk M, et al. European Association of Urology Guidelines on Non-muscle-invasive Bladder Cancer (Ta, T1, and Carcinoma in Situ). Eur Urol (2021), https://doi.org/10.1016/j.eururo.2021.08.010

119

126

140

# ARTICLE IN PRESS

### Table 2 – World Health Organization (WHO) classification in 1973 and in 2004/2016 [17]

1973 WHO classification system
Grade 1: well differentiated
Grade 2: moderately differentiated
Grade 3: poorly differentiated
2004/2016 WHO classification system (papillary lesions)
Papillary urothelial neoplasm of low malignant potential (PUNLMP)
Low-grade (LG) papillary urothelial carcinoma
High-grade (HG) papillary urothelial carcinoma

#### Table 3 – World Health Organization 2004 histological classification for flat lesions

Nonmalignant	lesions
--------------	---------

- Urothelial proliferation of uncertain malignant potential (flat lesion without atypia or papillary aspects).
- Reactive atypia (flat lesion with atypia).
- Atypia of unknown significance.
- (Potential) Premalignant lesion
- Urothelial dysplasia.
- Malignant lesion

• Urothelial carcinoma in situ is always high grade.

individual categories compared to the older 1973 WHO
 classification [17] (Tables 2 and 3).

There is a significant shift of patients between the categories of the WHO 1973 and the WHO 2004/2016 systems [19]. The proportion of tumours classified as papillary urothelial neoplasm of low malignant potential (PUNLMP; WHO 2004/2016) has decreased to very low levels in the past decade [20].

#### <sup>154</sup> 4.5.1. Prognostic value of histological grading

155 To compare the prognostic value of both WHO classifica-156 tions, an individual patient data (IPD) analysis of 5145 pri-157 mary Ta/T1 NMIBC tumours from patients at 17 centres was 158 conducted. The WHO 1973 and WHO 2004/2016 systems 159 were both prognostic for progression but not for recurrence. 160 When compared, WHO 1973 was a stronger prognosticator 161 of progression in Ta/T1 NMIBC than WHO 2004/2016. 162 However, a four-tier combination (low-grade [LG]/G1, LG/ 163 G2, HG/G2, and HG/G3) of both classification systems 164 proved to be superior to either classification system alone 165 [21].

In a subgroup of 3311 patients with primary Ta bladder
 tumours, similar prognosis was found for PUNLMP and Ta
 LG carcinomas [22]. Hence, these results do not support the
 continued use of PUNLMP as a separate grade category in
 the WHO 2004/2016 system.

To facilitate clinical utilisation in daily practice, these guidelines provide recommendations for tumours in both classification systems. 171

### 4.6. Inter- and intraobserver variability in staging and grading <sup>174</sup>

There is interobserver variability in the classification of CIS,175with agreement in only 70–78% of cases, in stage T1 versus176Ta tumours, and in tumour grading in both the 1973 and1772004/2016 classifications. The general conformity between178pathologists in staging and grading is 50–60% [23] (LE: 2a).179The WHO 2004/2016 classification provides slightly better180reproducibility than the 1973 classification [19].181

### 4.7. Variants of urothelial carcinoma and lymphovascular invasion

Several variants of urothelial carcinoma have been identified [24,25]. Most of these variants have worse prognosis than pure HG urothelial carcinoma [26] (LE: 3).

182

183

190

198

The presence of lymphovascular invasion (LVI) in TURB187specimens is associated with higher risk of pathological188upstaging and worse prognosis [27] (LE: 3).189

#### 4.8. Molecular classification

Molecular markers, in particular complex approaches such as stratification of patients on the basis of molecular classification, are promising but are not yet suitable for routine application [28]. Guidelines for the classification of BC are presented in Table 4.

5.	Diagnosis	196
----	-----------	-----

5.1. Patient history <sup>197</sup>

A focused patient history is mandatory.

5.2. Signs and symptoms <sup>199</sup>

Haematuria is the most common finding in NMIBC. Visible200haematuria was found to be associated with higher-stage201disease compared to nonvisible haematuria [29]. CIS might202be suspected in patients with lower urinary tract symptoms,<br/>especially irritative voiding.203

5.3. Physical examination 205

A focused urological examination is mandatory, although it does not reveal NMIBC.

#### Table 4 – Guidelines for bladder cancer classification

Recommendation	Strength rating
Use the 2017 TNM system for classification of the depth of tumour invasion (staging).	Strong
Use both the 1973 and 2004/2016 World Health Organization classification systems.	Weak
Do not use the term "superficial" bladder cancer.	Strong

EUROPEAN UROLOGY XXX (2021) XXX-XXX

#### Table 5 – Guidelines for primary assessment of non-muscle-invasive bladder cancer

Recommendation	Strength rating
Take a patient history, focusing on urinary tract symptoms and haematuria.	Strong
Use renal and bladder ultrasound and/or computed tomography (CT) urography during the initial work-up for patients with haematuria.	Strong
Once a bladder tumour has been detected, perform CT urography in selected cases (eg, tumours located in the trigone and multiple or high-risk tumours).	Strong
Perform cystoscopy for patients with symptoms suggestive of bladder cancer or during surveillance. Cystoscopy cannot be replaced by cytology or by any other noninvasive test.	Strong
For men, use a flexible cystoscope, if available.	Strong
Describe all macroscopic features of the tumour (site, size, number, and appearance) and mucosal abnormalities observed during cystoscopy. Use a bladder diagram.	Strong
Use voided urine cytology as an adjunct to cystoscopy to detect high-grade tumours.	Strong
Perform cytology on at least 25 ml of fresh urine or urine with adequate fixation. Morning urine is not suitable because of the frequent presence of cytolysis.	Strong
Use the Paris system for cytology reporting.	Strong

#### <sup>208</sup> **5.4.** *Imaging*

209 Computed tomography (CT) urography is used to detect 210 papillary tumours in the urinary tract, indicated by filling 211 defects and/or hydronephrosis [30]. The necessity to 212 perform baseline CT urography once a bladder tumour 213 has been detected is questionable owing to the low 214 incidence of significant findings obtained [31] (LE: 2b). 215 The incidence of simultaneous upper tract urothelial 216 carcinoma (UTUC) is low (1.8%), but increases to 7.5% for 217 tumours located in the trigone [31] (LE: 2b). The risk of 218 UTUC during follow-up is higher for patients with multiple 219 and high-risk tumours [32] (LE: 2b).

Ultrasound (US) permits characterisation of renal masses, detection of hydronephrosis, and visualisation of intraluminal masses in the bladder, but cannot rule out all potential causes of haematuria [33] (LE: 3). US cannot reliably exclude the presence of UTUC and cannot replace CT urography.

The role of multiparametric magnetic resonance imaging (MRI) in BC diagnosis and staging has not yet been established. A standardised methodology for MRI reporting for patients with BC has been published, but requires validation [34].

#### <sup>227</sup> 5.5. Urinary cytology

Examination of voided urine or bladder-washing specimens
for exfoliated cancer cells has high sensitivity in G3 and
high-grade tumours (84%), but low sensitivity in G1/LG
tumours (16%) [35]. The sensitivity for CIS detection is 28–
100% [36] (LE: 1b).

Cytological interpretation is user-dependent [37]. Evalu ation can be hampered by low cellular yield, urinary tract
 infections, stones, or intravesical instillations; however, in
 experienced hands the specificity exceeds 90% [37] (LE: 2b).

A standardised reporting system redefining urinary
 cytology diagnostic categories was published in 2016 by
 the Paris Working Group [38] and validated in retrospective
 studies [39].

#### <sup>241</sup> **5.6.** Urinary molecular marker tests

Driven by the low sensitivity of urine cytology, numerous
urinary tests have been developed [40]. None of these

markers can replace cystoscopy in routine practice, but the knowledge of positive test results (microsatellite analysis) 245 can improve the quality of follow-up cystoscopy [41] (LE: 246 1b). Promising novel urinary biomarkers assessing multiple 247 targets have been tested in prospective multicentre studies, 248 with a very high negative predictive value [42–44]. 249

#### 5.7. Cystoscopy

The diagnosis of papillary BC ultimately depends on<br/>cystoscopic examination of the bladder and histological251<br/>252evaluation of sampled tissue. CIS is diagnosed by a<br/>combination of cystoscopy, urine cytology, and histological253<br/>254evaluation of multiple bladder biopsies.254

Cystoscopy is initially performed as an outpatient procedure. A flexible instrument with intraurethral lubricant instillation results in better compliance compared to a rigid instrument, especially in men [45] (LE: 1b). Guidelines for the primary assessment of bladder cancer are presented in Table 5. 261

#### 5.8. Transurethral resection of Ta/T1 bladder tumours

The goal of TURB in Ta/T1 BC is to make the correct diagnosis 263 and completely remove all visible lesions. TURB should be 264 performed systematically in individual steps [46] (Table 6). 265

#### 5.8.1. Resection of the tumours

A complete resection, performed using either a fractioned267(separate resection of the exophytic part of the tumour, the268underlying bladder wall and the edges of the resection area)269(LE: 2b) or an en-bloc technique (LE: 1b), is essential to270achieve good prognosis [47,48].271The technique selected depends on the size and location272

The technique selected depends on the size and location of the tumour and experience of the surgeon.

The presence of detrusor muscle in the specimen is 274 considered a surrogate criterion of the resection quality and 275 is required (except for Ta G1/LG tumours). The absence of 276 detrusor muscle is associated with a significantly higher 277 risk of residual disease, early recurrence, and tumour 278 understaging [49] (LE: 1b). 279

In patients with a history of small Ta LG/G1 tumours, <sup>280</sup> fulguration, or laser vaporisation of small papillary <sup>281</sup>

250

262

266

## **ARTICLE IN PRESS**

EUROPEAN UROLOGY XXX (2021) XXX-XXX

### Table 6 – Guidelines for transurethral resection of the bladder, biopsies, and pathology reporting

Recommendation	Strength ratin
n patients suspected of having bladder cancer, perform TURB followed by pathology investigation of the specimen(s) obtained as a liagnostic procedure and initial treatment step.	Strong
Dutpatient fulguration or laser vaporisation of small papillary recurrences can be used in patients with a history of Ta G1/LG tumours.	Weak
Perform TURB systematically in individual steps:	Strong
Bimanual palpation under anaesthesia. This step may be omitted if noninvasive or early treatment for invasive disease is planned;	
Insertion of the resectoscope under visual control, with inspection of the whole urethra;	
Inspection of the whole urothelial lining of the bladder;	
Biopsy from the prostatic urethra (if indicated);	
Cold-cup bladder biopsies (if indicated);	
Resection of the tumour;	
Recording of findings in the surgery report/record;	
Precise description of the specimen for pathology evaluation.	
Performance of individual steps	
Perform en-bloc resection or resection in fractions (exophytic part of the tumour, the underlying bladder wall, and the edges of the esection area).	Strong
woid cauterisation as much as possible during TURB to minimise tissue deterioration.	Strong
ake biopsies from abnormal-looking urothelium. Biopsies from normal-looking mucosa (mapping biopsies from the trigone, bladder lome, and right, left, anterior, and posterior bladder wall) are recommended when cytology is positive, in cases with a history of HG/G3 umours, and for tumours with a nonpapillary appearance. If equipment is available, perform fluorescence-guided (PDD) biopsies.	Strong
ake a biopsy of the prostatic urethra in cases of bladder neck tumour, if bladder CIS is present or suspected, if there is positive cytology vithout evidence of tumour in the bladder, or if abnormalities of the prostatic urethra are visible. If biopsy is not performed during the nitial procedure, it should be completed at the time of the second resection.	Strong
ake a prostatic urethral biopsy from the precollicular area (between the 5 and 7 o'clock positions) using a resection loop. If any biormal-looking areas in the prostatic urethra are observed, these need to be biopsied as well.	Weak
lse methods to improve tumour visualisation (fluorescence cystoscopy, narrow-band imaging) during TURB, if available.	Weak
end the specimens from different biopsies and resection fractions to the pathologist in separately labelled containers.	Weak
he TURB record must describe tumour location, appearance, size and multifocality, all steps of the procedure, and the extent and ompleteness of the resection.	Strong
or patients with positive cytology but negative cystoscopy, exclude UTUC, CIS in the bladder (via mapping biopsies or PDD-guided iopsies), and tumour in the prostatic urethra (via prostatic urethra biopsy).	Strong
erform a second TURB in the following situations:	Strong
After incomplete initial TURB, or in the case of doubt about TURB completeness)	
If there is no detrusor muscle in the specimen after initial resection, with the exception of Ta LG/G1 tumours and primary CIS	
For T1 tumours.	
f indicated, perform a second TURB within 2–6 wk after initial resection. This second TURB should include resection of the primary umour site.	Weak
legister the pathology results of a second TURB, as it reflects the quality of the initial resection.	Weak
form the pathologist of prior treatments (intravesical therapy, radiotherapy, etc).	Strong
he pathology report should specify tumour location, tumour grade and stage, lymphovascular invasion, unusual (variant) histology, ind the presence of CIS and detrusor muscle.	Strong

CIS = carcinoma in situ; HG = high grade; LG = low grade; PDD = photodynamic diagnosis; TURB = transurethral resection of the bladder; UTUC = upper tract urothelial carcinoma.

recurrences on an outpatient basis can reduce the
therapeutic burden [50] (LE: 3).

#### <sup>284</sup> 5.8.2. Bladder biopsies

285 CIS can present as a velvet-like, reddish area that is 286 indistinguishable from inflammation, or it may not be 287 visible at all. For this reason, biopsies from suspicious 288 urothelium should be taken. In addition, for patients with 289 positive urine cytology (see Section 5.5) or with a history of 290 HG/G3 NMIBC and for tumours with a nonpapillary 291 appearance, mapping biopsies from normal-looking muco-292 sa are recommended [51]. If equipment is available, 293 photodynamic diagnosis (PDD) is a useful tool for targeting 294 the biopsy.

#### <sup>295</sup> 5.8.3. Prostatic urethral biopsies

Involvement of the prostatic urethra and ducts in men with
 NMIBC has been reported [52] (LE: 2b). The risk of prostatic
 urethra or duct involvement is higher if the tumour is

located at the trigone or bladder neck, in the presence of299bladder CIS, and in cases with multiple tumours [53] (LE:3003b). On the basis of this observation, a biopsy from the301prostatic urethra is necessary in some cases [52,54].302

303

308

#### 5.9. New methods of tumour visualisation

As a standard procedure, cystoscopy and TURB are performed using white light. However, the use of white light can miss lesions that are present but not visible, which is why new technologies are being developed. 307

#### 5.9.1. PDD (fluorescence cystoscopy)

PDD is performed using violet light after intravesical309instillation of 5-aminolaevulinic acid or hexaminolaevulinic310acid (LE: 1a). In a systematic review and meta-analysis, PDD311had higher sensitivity for detection of tumour lesions than312white light endoscopy at both the patient level (92% vs 71%)313and biopsy level (93% vs 65%) [55]. A prospective314

EUROPEAN UROLOGY XXX (2021) XXX-XXX

368

369

409

410

randomised trial did not confirm a higher detection rate
 among patients with known positive cytology before TURB
 [56].

318 PDD had lower specificity than white-light endoscopy 319 (63% vs 81%) [55]. False positivity can be induced by 320 inflammation or recent TURB and during the first 3 mo after 321 bacillus Calmette-Guérin (BCG) instillation [57,58] (LE: 1a). 322 A systematic review and analysis of 14 randomised 323 controlled trials (RCTs) demonstrated the beneficial effect of 324 fluorescence cystoscopy on the recurrence rate in patients 325 with TURB; however, there were no differences in progres-326 sion and mortality rates [59] (LE: 1a).

### <sup>327</sup> 5.9.2. Narrow-band imaging (NBI)

328 In NBI, the contrast between normal urothelium and 329 hypervascular cancer tissue is enhanced. Improved cancer 330 detection has been observed with NBI flexible cystoscopy 331 and NBI-guided biopsies and resection [60] (LE: 3b). An RCT 332 assessed the reduction in recurrence rates if NBI is used 333 during TURB. Although the overall results of the study were 334 negative, a benefit after 3 and 12 mo was observed for low-335 risk tumours (pTa LG, <30 mm, no CIS) [61] (LE: 1b).

#### <sup>336</sup> 5.10. Second resection

337 A significant risk of residual tumour after initial TURB of Ta/ 338 T1 lesions has been demonstrated [62]. A systematic review 339 demonstrated 51% risk of persistence and 8% risk of 340 understaging for T1 tumours. Most of the residual lesions 341 were detected at the original tumour location [62] (LE: 1a). 342 The prevalence of residual tumours and upstaging to 343 invasive disease after TURB for T1 tumour also remained 344 high in a subgroup with detrusor muscle in the resection 345 specimen [63].

346 A second TURB can increase recurrence-free survival 347 (RFS) [64] (LE: 2a), improve outcomes after BCG treatment 348 [65] (LE: 3), and provide prognostic information [66,67] (LE: 349 3). In a retrospective evaluation of a multi-institutional 350 cohort of 2451 patients with BCG-treated T1 G3/HG 351 tumours, the second resection improved RFS, progres-352 sion-free survival (PFS), and overall survival (OS) only in 353 cases without detrusor muscle in the specimen from the 354 initial resection [68] (LE: 3).

Retrospective evaluation showed that a second resection performed 14–42 d after the initial resection provides longer RFS and PFS compared to a second resection performed after 43–90 d [69] (LE: 3).

#### <sup>359</sup> 5.11. Pathology report

Pathological investigation of the specimen(s) obtained via 360 TURB and biopsies is an essential step in the decision-making 361 process for BC. Close cooperation between urologists and 362 pathologists is required. To obtain all the relevant informa-363 tion, the specimen collection, handling, and evaluation 364 should follow the recommendations (Table 6) [70]. In difficult 365 cases, an additional review by an experienced genitourinary 366 pathologist can be considered. Guidelines for TURB, biopsies, 367 and pathology report are presented in Table 6.

#### 6. Predicting disease recurrence and progression

#### 6.1. Ta and T1 tumours

Treatment should take into account a patient's prognosis. In370order to predict the risk of disease recurrence and/or371progression, several prognostic models for specified patient372populations have been introduced.373

374 6.1.1. Scoring models using the WHO 1973 classification system 375 6.1.1.1. The 2006 European Organisation for Research and Treatment of 376 Cancer (EORTC) scoring model. The 2006 EORTC scoring model 377 is based on the six most significant clinical and pathological 378 factors for patients mainly treated with intravesical 379 chemotherapy, which are the number of tumours, tumour 380 diameter, prior recurrence rate, category, concurrent CIS, 381 and WHO 1973 tumour grade [71]. Using this model, 382 individual probabilities of recurrence and progression at 383 1 and 5 yr can be calculated.

6.1.1.2. Model for patients with Ta G1/G2 (WHO 1973) tumours treated384with chemotherapy. Patients with Ta G1/G2 tumours receiving385chemotherapy were stratified into three risk groups for366recurrence, taking into account the history of recurrences,387history of intravesical treatment, tumour grade (WHO3881973), number of tumours, and adjuvant chemotherapy380[72].390

391 6.1.1.3. Club Urologico Español de Tratamiento Oncologico (CUETO) 392 scoring model for BCG-treated patients. The CUETO model 393 predicts the risk of recurrence and progression for patients 394 treated with 12 doses of intravesical BCG over a 5- to 6-mo 395 period following TURB. The scoring system is based on 396 evaluation of seven prognostic factors: gender, age, prior 397 recurrence status, number of tumours, T category, associat-398 ed CIS, and WHO 1973 tumour grade.

Using this model, the calculated risk of recurrence is <sup>399</sup> lower than that obtained via the EORTC model. For <sup>400</sup> progression, probability is lower only for high-risk patients <sup>401</sup> [73] (LE: 2a). The lower risks in the CUETO model can be <sup>402</sup> attributed to the use of BCG in this sample. <sup>403</sup>

6.1.1.4. The 2016 EORTC scoring model for patients treated with<br/>maintenance BCG. In patients with intermediate- and high-<br/>risk tumours without CIS treated with 1–3 yr of<br/>maintenance BCG, EORTC risk groups and nomograms for<br/>BCG-treated patients were developed [74] (LE: 2a).404<br/>405

### 6.1.2. Scoring model using the WHO 2004/2016 and WHO 1973 classification systems

411 6.1.2.1. EAU NMIBC 2021 scoring model. To create new prognos-412 tic-factor risk groups using both the WHO 1973 and WHO 413 2004/2016 classification systems, IPD from patients with 414 primary tumours treated with TURB  $\pm$  intravesical chemo-415 therapy were used [22] (see Section 4.5.1). From the 416 multivariate analysis, tumour stage, WHO 1973 grade, 417 WHO 2004/2016 grade, concomitant CIS, number of 418 tumours, tumour size, and age were independent predictors 419 of disease progression [22].

D: 1.

# ARTICLE IN PRESS

Table 7 - Clinical composition of the new European Association ofUrology prognostic-factor risk groups for non-muscle-invasivebladder cancer based on the WHO 2004/2016 or WHO 1973 gradingclassification system [22] a

Risk group	
Low risk	• A primary, single, Ta/T1 LG/G1 tumour <3 cm in diameter without CIS in a patient aged $\leq$ 70 yr
	• A primary Ta LG/G1 tumour without CIS with at most ONE additional clinical risk factors <sup>b</sup>
Intermediate risk	Patients without CIS who are not included in either the low, high, or very high-risk groups
High risk	• All T1 HG/G3 without CIS, EXCEPT those included in the very high-risk group
	• All CIS patients, EXCEPT those included in the very high- risk group
	Stage, grade with additional clinical risk factors: <sup>b</sup>
	• Ta LG/G2 or T1 G1 with CIS and all 3 risk factors
	• Ta HG/G3 or T1 LG with no CIS and at least 2 risk factors
	• T1 G2 with no CIS and at least 1 risk factor
Very high risk	Stage, grade with additional clinical risk factors: $^{ m b}$
	• Ta HG/G3 and CIS with all 3 risk factors
	• T1 G2 and CIS with at least 2 risk factors
	• T1 HG/G3 and CIS with at least 1 risk factor
	• T1 HG/G3 with no CIS and all 3 risk factors

situ; CIS = carcinoma HG = high grade; LG = lowin grade; LVI = lymphovascular invasion; WHO = World Health Organization. Only one of the two classification systems (WHO 1973 or WHO 2004/2016) is required to use this table. If both classification systems are available for an individual patient, the Panel recommends using the risk group calculation based on the WHO 1973 system, as it has better prognostic value. The LG category (WHO 2004/2016) also includes tumours classified as papillary urothelial neoplasm of low malignant potential. The scoring model is based on a meta-analysis of individual patient data, but does not consider patients with primary CIS (high risk) or with recurrent tumours, as well as some pathological parameters such as variant histology (micropapillary, plasmacytoid, sarcomatoid, small-cell, neuroendocrine) and LVI. Nevertheless, on the basis of data from the literature, all patients with CIS in the prostatic urethra, with some variant histology of urothelial carcinoma, or with LVI should be included in the very high-risk group. Patients with recurrent tumours should be included in the intermediate-, high-, or very high-risk groups according to the other prognostic factors they have. <sup>b</sup> Additional risk factors: age >70 yr, multiple papillary tumours, and tumour diameter >3 cm.

This model is used for defining risk groups as this is the
only model in which the WHO 2004/2016 classification
system is included as one of parameters (see Section 6.3).

As the 2021 EAU NMIBC scoring model determines the
 risk of tumour progression, but not recurrence, any of the
 models mentioned in Section 6.1.1 may be used to calculate
 an individual's risk of disease recurrence.

#### <sup>427</sup> 6.1.3. Further prognostic factors

<sup>428</sup> Further prognostic factors have been described in selected
 <sup>429</sup> patient populations:

- For T1 G3 tumours, important prognostic factors were female sex, CIS in the prostatic urethra in men treated with an induction course of BCG, and age, tumour size, and concurrent CIS in BCG-treated patients [52,75] (LE: 2b).
- T1 G3 tumours in bladder (pseudo)diverticulum [76] (LE: 3).
- In patients with T1 tumours, the finding of residual T1 disease at second TURB is an unfavourable prognostic factor [66,67] (LE: 3).

- In patients with T1 G2 tumours treated with TURB, <sup>439</sup> recurrence at 3 mo was the most important predictor of <sup>440</sup> progression [77] (LE: 2b). <sup>441</sup>
- The prognostic value of pathological factors has been discussed elsewhere (see Section 4.6). More research is needed to determine the role of molecular markers in improving the predictive accuracy of currently available risk tables [78]. 446

447

459

486

#### 6.2. Carcinoma in situ

Without any treatment, approximately 54% of patients with448CIS experience progression to muscle-invasive disease [79]449(LE: 3). There are no reliable prognostic factors, but some450studies have reported worse prognosis for concurrent CIS451and T1 tumours compared to primary CIS [80,81], for452extended CIS [81], and for CIS in the prostatic urethra [52]453(LE: 3).454

The response to intravesical treatment with BCG or chemotherapy is an important prognostic factor for subsequent progression and death caused by BC [73,77] 457 (LE: 2a). 458

#### 6.3. Patient stratification into risk groups

460 To be able to facilitate treatment recommendations, the 461 Guidelines Panel recommends the stratification of patients 462 into risk groups according to their probability of progression 463 to muscle-invasive disease (Table 7). The risk group 464 definitions are based on an IPD meta-analysis for primary patients treated with TURB  $\pm$  intravesical chemotherapy 465 and calculation of their progression scores (2021 EAU 466 467 NMIBC scoring model) as presented in Sections 4.5.1 and 468 6.1.2 [22].

For calculation of the risk group for individual patients, 469 either one or both of the WHO 1973 and WHO 2004/2016 470 classification systems may be used. 471

For factors for which IPD were not collected, such as472variant histology, LVI, primary CIS, and CIS in the prostatic473urethra, literature data have been used to classify patients474into risk groups.475

A web-based calculator (www.nmibc.net) and apps (iOS: 476 477 https://apps.apple.com/us/app/eau-nmibc-risk-calculator/ 478 id1578482687 and Android: https://play.google.com/store/ 479 apps/details?id=net.ydeal.nmibc) facilitate determination 480 of a patient's risk group in daily clinical practice. The 481 individual probability of disease progression at 1, 5, and 482 10 yr for the new EAU NMIBC risk groups is presented in 483 Table 8. Guidelines for stratification of patients with NMIBC 484 are presented in Table 9.

#### 7. Disease management 485

#### 7.1. Counselling on smoking cessation

Smoking increases the risk of tumour recurrence and<br/>progression [82] (LE: 3). While it is still controversial<br/>whether smoking cessation in BC will favourably influence487<br/>488<br/>489<br/>489the outcome of BC treatment, patients should be counselled490

Table 8 – Probability of disease progression at 1, 5, and 10 yr for the new European Association of Urology non–muscle-invasive bladder cancer risk groups [22] <sup>a</sup>

New risk groups	Probability of progression, % (95% confidence interval)		
	1 yr	5 yr	10 yr
With WHO 2004/20	16		
Low	0.06 (0.01-0.43)	0.93 ( 0.49-1.7)	3.7 (2.3-5.9)
Intermediate	1.0 (0.50-2.0)	4.9 (3.4-7.0)	8.5 (5.6-13)
High	3.5 (2.4-5.2)	9.6 (7.4-12)	14 (11–18)
Very High	16 (10-26)	40 (29-54)	53 (36-73)
With WHO 1973			
Low	0.12 (0.02-0.82)	0.57 (0.21-1.5)	3.0 (1.5-6.3)
Intermediate	0.65 (0.36-1.2)	3.6 (2.7-4.9)	7.4 (5.5–10)
High	3.8 (2.6-5.7)	11 (8.1-14)	14 (10–19)
Very High	20 (12–32)	44 (30-61)	59 (39–79)

WHO = World Health Organization.

<sup>a</sup> This table does not include patients with variant histologies, lymphovascular invasion, carcinoma in situ in the prostatic urethra, or primary or recurrent carcinoma in situ.

to stop smoking because of the general risks connected to
 tobacco smoking [83] (LE: 3).

#### <sup>493</sup> **7.2.** *Adjuvant treatment*

Although TURB by itself can eradicate a Ta/T1 tumour
 completely, these tumours commonly recur and can
 progress to MIBC. It is therefore necessary to consider
 adjuvant therapy for all patients.

#### <sup>498</sup> 7.2.1. Intravesical chemotherapy

7.2.1.1. A single, immediate, postoperative intravesical instillation of *chemotherapy.* It has been shown that immediate single
instillation (SI) acts by destroying circulating/floating
tumour cells after TURB, as well as via an ablative effect
on residual tumour cells at the resection site and on small
overlooked tumours [84,85] (LE: 3).

Four large meta-analyses have consistently shown that after TURB, SI significantly reduces the recurrence rate compared to TURB alone [86–89] (LE: 1a). In a systematic review and IPD meta-analysis, SI reduced the 5-yr recurrence rate by 14%, although only patients with primary tumours or intermediate-risk recurrent tumours with a prior recurrence rate of one or fewer recurrences per year and those with a 2006 EORTC recurrence score of <5  $^{512}$  benefited [86].  $^{513}$ 

SIs with mitomycin C (MMC), epirubicin, or pirarubicin514have all shown a beneficial effect [86]. SI with gemcitabine515was superior to a placebo control (saline) in an RCT with516remarkably low toxicity rates [90]. The efficacy of continu-517ous saline irrigation in the prevention of early recurrences518has also been suggested [91].519

Prevention of tumour cell implantation should be <sup>520</sup> initiated within the first few hours after TURB [92] (LE: <sup>521</sup> 3). Safety measures should be maintained (Table 10). <sup>522</sup>

523 7.2.1.2. Additional adjuvant intravesical chemotherapy instillations. 524 The need for further adjuvant intravesical therapy depends 525 on prognosis. For patients with low-risk tumours (Table 7), 526 SI reduces the risk of recurrence and is considered to be the 527 standard and complete treatment [86,87] (LE: 1a). For other 528 patients, however, SI remains an incomplete treatment 529 because of the considerable likelihood of recurrence and/or progression (2006 EORTC scoring model and Table 8). 530

Efficacy data for the following comparisons of application schemes have been published. 532

7.2.1.2.1. SI alone versus SI and further repeat instillations. In one<br/>study, further chemotherapy instillations after SI improved533RFS in patients with intermediate-risk tumours [93] (LE:<br/>3355342a).535

7.2.1.2.2. Repeat chemotherapy instillations versus no adjuvant<br/>treatment. Meta-analyses showed an absolute reduction of<br/>13–14% for patients treated with TURB and chemotherapy<br/>instillations over those with TURB alone [94].536<br/>537

7.2.1.2.3. SI and further repeat instillations versus later repeat 539 instillations only. SI might have an impact on recurrence even 540 when further adjuvant instillations are given [95,96]. An 541 RCT comparing SI of MMC with an instillation of MMC 542 delayed until 2 wk after TURB (followed by further repeat 543 instillations in both treatment arms) showed a significant 544 reduction of 9% in the risk of recurrence at 3 yr in favour of SI 545 [95] (LE: 2a). Since the authors' definition of the risk groups 546 differed significantly in the initial publication, they adapted 547 their patient stratification in the second analysis and 548 consistently showed improved efficacy of SI followed by repeat MMC instillations [97]. The results of this study 549

#### Table 9 - Guidelines for stratification of patients with non-muscle-invasive bladder cancer

Recommendation	Strength rating
Stratify patients into four risk groups according to Table 7. A patient's risk group can be determined using the EAU risk group calculator available at www.nmibc.net.	Strong
For information about the risk of disease progression in a patient with primary Ta/T1 tumours, use the data from Table 8.	Strong
Use the 2006 EORTC scoring model to predict the risk of tumour recurrence in individual patients not treated with BCG at www. omnicalculator.com/health/eortc-bladder-cancer.	Strong
Use the 2016 EORTC or the CUETO risk scoring model to predict the risk of tumour recurrence and progression in individual patients treated with BCG intravesical immunotherapy (the 2016 EORTC model is calculated for 1–3 yr of maintenance and the CUETO model for 5–6 m of BCG).	Strong
BCG = bacillus Calmette-Guérin; CUETO = Club Urologico Español de Tratamiento Oncologico; EAU = European Association of Urology Organisation for Reaearch and Treatment of Cancer.	r; EORTC = European

9

# EUROPEAN UROLOGY XXX (2021) XXX-XXX

#### Table 10 - Guidelines for adjuvant therapy for Ta/T1 tumours and for carcinoma in situ

General recommendations	Strength ratin
Counsel smokers with confirmed NMIBC to stop smoking.	Strong
The type of further therapy after TURB should be based on the risk groups shown in Section 6.3 and Table 7. For determination of a patient's risk group, use the 2021 EAU risk group calculator available at www.nmibc.net.	Strong
For patients with tumours presumed to be at low risk and those with small papillary recurrences (presumably Ta LG/G1) detected more than 1 yr after previous TURB, offer one immediate chemotherapy instillation.	Strong
For patients with intermediate-risk tumours (with or without immediate instillation), 1-yr full-dose BCG treatment (induction plus 3- weekly instillations at 3, 6, and 12 mo) or instillations of chemotherapy (the optimal schedule is not known) for a maximum of 1 yr is recommended. The final choice should reflect the individual patient's risk of recurrence and progression as well as the efficacy and side effects of each treatment modality.	Strong
For patients with high-risk tumours, full-dose intravesical BCG for 1–3 yr (induction plus 3-weekly instillations at 3, 6, 12, 18, 24, 30, and 36 mo) is indicated. The additional beneficial effect of the second and third years of maintenance should be weighed against added costs, side effects, and problems connected with BCG shortages.	Strong
For patients with very high-risk tumours, discuss immediate RC.	Strong
Offer transurethral resection of the prostate followed by intravesical instillation of BCG to patients with CIS in the epithelial lining of the prostatic urethra.	Weak
The definition of BCG-unresponsive tumours should be respected as it most precisely identifies the patients who are unlikely to respond o further BCG instillations.	Strong
Offer RC to patients with BCG-unresponsive tumours.	Strong
For patients with BCG-unresponsive tumours who are not candidates for RC because of comorbidities, offer preservation strategies (intravesical chemotherapy, chemotherapy and microwave-induced hyperthermia, electromotive administration of chemotherapy, intravesical or systemic immunotherapy; preferably within clinical trials).	Weak
Recommendations: technical aspects for treatment	
ntravesical chemotherapy	
f given, administer a single immediate instillation of chemotherapy within 24 h after TURB.	Weak
Omit a single immediate instillation of chemotherapy in any case of overt or suspected bladder perforation or bleeding requiring bladder irrigation.	Strong
Give clear instructions to the nursing staff to control the free flow of the bladder catheter at the end of the immediate instillation.	Strong
The optimal schedule and duration for further intravesical chemotherapy instillation are not defined; however, the duration should not exceed 1 yr.	Weak
f intravesical chemotherapy is given, use the drug at its optimal pH and maintain the concentration of the drug by reducing fluid intake before and during instillation.	Strong
he length of an individual instillation should be 1–2 h.	Weak
3CG intravesical immunotherapy	
Absolute contraindications to BCG intravesical instillation are:	Strong
During the first 2 wk after TURB;	-
In patients with visible haematuria;	
• After traumatic catheterisation;	
In patients with symptomatic urinary tract infection.	

550 should be considered with caution since some patients did 551 not receive adequate therapy. Another RCT found no impact 552 of SI with epirubicin followed by further chemotherapy or 553 BCG instillations in a cohort of predominantly high-risk BC 554 [98].

7.2.1.2.4. The optimal schedule for intravesical chemotherapy instilla-555 tions. The length and frequency of repeat chemotherapy 556 instillations are still controversial; however, the duration 557 should not exceed 1 yr [96] (LE: 3).

558 7.2.1.3. Options for improving the efficacy of intravesical 559 chemotherapy

7.2.1.3.1. Adjustment of pH, duration of instillation, and drug 560 concentration. One RCT showed that adjusting the urinary 561 pH and decreasing urinary excretion reduced the recurrence 562 rate [99] (LE: 1b). Another trial reported that a duration of 563 1 h for instillation of MMC was more effective than 30-min 564 instillation [100] (LE: 3). Another RCT using epirubicin

documented that concentration is more important than treatment duration [101] (LE: 1b).

#### 7.2.1.3.2. Device-assisted intravesical chemotherapy. Microwaveinduced hyperthermia effect

Promising data have been presented on enhancing the 569 efficacy of MMC using microwave-induced hyperthermia in patients with high-risk tumours [102]. One RCT comparing 570 1 yr of BCG with 1 yr of MMC and microwave-induced 571 572 hyperthermia in patients with intermediate- and high-risk 573 BC revealed greater RFS at 24 mo in the MMC group [103] 574 (LE: 1b).

Hyperthermic intravesical chemotherapy

Different technologies that increase the temperature of instilled MMC are available, but data on their efficacy are still lacking. 578 579

#### Electromotive drug administration

580 The efficacy of MMC using electromotive drug administra-581 tion (EMDA) sequentially combined with BCG in patients with 582 high-risk tumours has been suggested in one small RCT [104].

565 566

567 568

## EUROPEAN UROLOGY XXX (2021) XXX-XXX

583 For application of device-assisted instillations in patients <sup>584</sup> **Q7** with BCG-unresponsive tumours, see Section 7.3.3.

585 7.2.2. Intravesical BCG immunotherapy

586 7.2.2.1. Efficacy of BCG

7.2.2.1.1. Recurrence rate. Five meta-analyses have confirmed 587 that BCG after TURB is superior to TURB alone or 588 TURB + chemotherapy in preventing the recurrence of 589 NMIBC [105-109] (LE: 1a). Three RCTs of intermediate-590 and high-risk tumours compared BCG with epirubicin and 591 interferon (IFN) [110], epirubicin alone [111], or MMC [112] 592 and confirmed the superiority of BCG for prevention of 593 tumour recurrence (LE: 1a). The effect is long-lasting 594 [111,112] and was also observed in a separate analysis of 595 patients with intermediate-risk tumours [111]. An IPD 596 meta-analysis demonstrated a 32% reduction in the risk of 597 recurrence for BCG compared to MMC in trials with BCG 598 maintenance, but a 28% increase for patients treated 599 without BCG maintenance (LE: 1a) [105].

7.2.2.1.2. Progression rate. Two meta-analyses demonstrated 600 that BCG therapy delays and potentially lowers the risk of 601 tumour progression [113,114] (LE: 1a). In a meta-analysis 602 carried out by the EORTC Genito-Urinary Cancers Group 603 (GUCG), tumours progressed in 9.8% of patients treated with 604 BCG compared to 13.8% in the control groups (TURB alone, 605 TURB and intravesical chemotherapy, or TURB with other 606 immunotherapy). The magnitude of the reduction was 607 similar in patients with Ta/T1 papillary tumours and in 608 those with CIS [114]. An RCT with long-term follow-up 609 demonstrated significantly fewer distant metastases and 610 better OS and disease-specific survival for patients treated 611 with BCG when compared to epirubicin [111] (LE: 1b). By 612 contrast, an IPD meta-analysis was not able to confirm any 613 significant difference between MMC and BCG for progres-614 sion, survival, or cause of death [105].

615 The conflicting results in the outcomes of these studies 616 can be explained by differences in patient characteristics, 617 duration of follow-up, methodology, and statistical power. 618 However, most studies showed a reduction in the risk of 619 progression of high-and intermediate-risk tumours if a BCG 620 maintenance schedule was applied.

621 7.2.2.2. BCG strain. A network meta-analysis identified ten 622 different BCG strains used for intravesical treatment, but 623 was not able to confirm the superiority of any BCG strain 624 over another [115]. However, the quality of the source data 625 does not allow definitive conclusions.

626 7.2.2.3. BCG toxicity. BCG intravesical treatment is associated 627 with more side effects than with intravesical chemotherapy 628 [114] (LE: 1a). However, serious side effects are encountered 629 in <5% of patients and can be treated effectively [116] (LE: 630 1b). The incidence of BCG infections after BCG instillations 631 was 1% in a registry-based cohort analysis [117]. It has been 632 shown that a maintenance schedule is not associated with 633 an increase in the risk of side effects when compared to an 634 induction course [116]. Side effects requiring treatment 635 cessation were seen more often in the first year of therapy

636 [118]. Elderly patients do not seem to experience more side 637 effects leading to treatment discontinuation [119] (LE: 2a). 638 No significant difference in toxicity between different BCG 639 strains was demonstrated [120]. Symptoms may be the 640 result of side effects of the BCG treatment or caused by the 641 bladder disease (widespread CIS) itself. Consequently, the 642 burden of symptoms decreases after completion of the 643 treatment in a significant number of patients [121].

644 Major complications can appear after systemic absorp-645 tion of the drug. Thus, contraindications to BCG intravesical 646 instillation should be respected (Table 10). The presence of 647 leukocyturia, nonvisible haematuria, or asymptomatic 648 bacteriuria is not a contraindication to BCG application, 649 and antibiotic prophylaxis is not necessary in these cases 650 [122] (LE: 3).

651 BCG should be used with caution in immunocompro-652 mised patients [123]. The management of side effects after 653 BCG should reflect their type and grade according to the 654 recommendations [124].

655 7.2.2.4. Optimal BCG schedule. Induction BCG instillations are 656 given according to the empirical 6-weekly schedule 657 [125]. For optimal efficacy, the induction course must be 658 followed by maintenance instillations [105,109,113,114] (LE: 659 1a). Many different maintenance schedules have been used. up to a maximum of 27 instillations over 3 yr [126]. 660

7.2.2.4.1. Optimal number of induction instillations and frequency of 661 instillations during maintenance. The optimal number of induc-662 tion instillations and frequency of maintenance instillations 663 were evaluated in the NIMBUS trial. A safety analysis after 345 patients had been randomised demonstrated that a 664 665 lower number of instillations (three instillations for 666 induction and two instillations at 3, 6, and 12 mo) was 667 inferior to the standard schedule (6 instillations for 668 induction and 3 instillations at 3, 6, and 12 mo) regarding 669 the time to first recurrence [127] (LE: 1b). A CUETO RCT 670 showed that for high-risk tumours a maintenance schedule 671 with only one instillation every 3 mo for 3 yr was not 672 superior to induction therapy only, which suggested that 673 one instillation may be suboptimal to three instillations in 674 each maintenance cycle [128] (LE: 1b).

7.2.2.4.2. Optimal length of maintenance. It was demonstrated 675 that at least 1 yr of maintenance BCG is required to obtain 676 superiority of BCG over MMC for prevention of recurrence 677 or progression [113] (LE: 1a).

678 An EORTC RCT showed that when BCG is given at full dose, 3 yr of maintenance (3-weekly instillations 3, 6, 12, 18, 24, 30, and 36 mo) reduces the recurrence rate compared to 1 yr for high-risk but not intermediate-risk tumours. There were no differences in progression or OS [129] (LE: 1b).

679

680

681

682

683 7.2.2.5. Optimal dose of BCG. To reduce BCG toxicity, instillation 684 of a reduced dose has been proposed. However, it has been 685 suggested that a full dose of BCG is more effective for 686 multifocal tumours [130,131] (LE: 1b). The CUETO study 687 compared one-third dose to full-dose BCG and found no 688 overall difference in efficacy. However, a further reduction

### Table 11 – Guidelines for the treatment of Ta/T1 tumours and carcinoma in situ according to risk stratification

Recommendation	Strength rating
EAU low risk group	
Offer one immediate instillation of intravesical chemotherapy after TURB.	Strong
EAU intermediate risk group	
For all patients, either 1-yr full-dose BCG treatment (induction plus 3-weekly instillations at 3, 6, and 12 mo) or instillations of chemotherapy (the optimal schedule is not known) for a maximum of 1 yr is recommended. The final choice should reflect the individual patient's risk of recurrence and progression as well as the efficacy and side effects of each treatment modality. Offer one immediate chemotherapy instillation to patients with small papillary recurrences detected more than 1 yr after previous TURB.	Strong
EAU high risk group	
Offer intravesical full-dose BCG instillations for 1-3 yr or RC.	Strong
EAU very high risk group	
Consider RC and offer intravesical full-dose BCG instillations for 1–3 yr to those who refuse or are unfit for RC.	Strong
BCG = bacillus Calmette-Guérin; EAU = European Association of Urology; RC = radical cystectomy; TURB = transurethral resection of the bladder.	

689 to one-sixth dose resulted in a decrease in efficacy with no 690 decrease in toxicity [132] (LE: 1b). The EORTC did not find 691 any difference in toxicity between one-third and full-dose 692 BCG, but one-third dose BCG was associated with a higher 693 recurrence rate, especially when it was given for only 1 yr 694 [118,129] (LE: 1b). Routine use of one-third dose BCG is 695 complicated by potential technical difficulties in preparing 696 the reduced dose.

#### <sup>697</sup> 7.2.3. Combination therapy

7.2.3.1. Intravesical BCG + chemotherapy versus BCG alone. In one
 RCT, a combination of MMC and BCG was more effective in
 reducing recurrences but more toxic compared to BCG
 monotherapy (LE: 1b). [133]. Improved disease-free survival
 (DFS) but no difference in PFS for patients treated with
 combination treatment comparing to BCG alone were
 observed [134].

705 7.2.3.2. Combination treatment using IFN. In a Cochrane meta-706 analysis of four RCTs, a combination of BCG and IFN-2a did 707 not show a clear difference in recurrence and progression 708 when compared to BCG alone [135]. In one study, weekly 709 MMC followed by monthly BCG alternating with IFN-2 $\alpha$ 710 showed a higher probability of recurrence compared to 711 MMC followed by BCG alone [136]. In addition, an RCT 712 comparing BCG monotherapy with a combination of 713 epirubicin and IFN for up to 2 yr showed that the latter 714 was significantly inferior to BCG monotherapy in preventing 715 recurrence [137] (LE: 1b).

#### <sup>716</sup> 7.2.4. Specific aspects of treatment of CIS

717 7.2.4.1. Treatment strategy. Detection of concurrent CIS
 718 increases the risk of recurrence and progression of Ta/T1
 719 tumours [71,73]. As CIS cannot be cured by an endoscopic
 720 procedure alone, the diagnosis of CIS must be followed by
 721 further treatment using either intravesical BCG instillations
 722 or RC (LE: 4).

723 7.2.4.2. Prospective randomised trials on intravesical BCG or che 724 motherapy. A meta-analysis of clinical trials comparing
 725 intravesical BCG to intravesical chemotherapy in patients
 726 with CIS showed a significantly higher response rate and
 727 lower risk of treatment failure after BCG [138] (LE: 1a).

In an EORTC-GUCG meta-analysis, in a subgroup of 403 patients with CIS, BCG reduced the risk of progression 729 by 35% when compared to intravesical chemotherapy 730 immunotherapy [114] (LE: 1b). The combination of BCG and MMC was not superior to BCG alone [139]. 732

733 7.2.4.3. Treatment of CIS in the prostatic urethra. Patients with CIS 734 are at high risk of extravesical involvement in the UUT and 735 in the prostatic urethra [140]. Patients with extravesical 736 involvement had worse survival than those with bladder CIS 737 alone [140] (LE: 3). Patients with CIS in the epithelial lining 738 of the prostatic urethra can be treated with intravesical instillation of BCG. Transurethral resection of the prostate 739 can improve contact of BCG with the prostatic urethra [141] 740 (LE: 3). 741

For patients with prostatic duct involvement there are<br/>promising results with BCG, but only from small series. The<br/>data are insufficient to provide clear treatment recommen-<br/>dations, and radical surgery should be considered [141] (LE:<br/>3).742<br/>743

The treatment strategy for primary and recurrent 747 tumours after TURB without previous BCG instillations is 748 presented in Table 11. 749

750

#### 7.3. Treatment of failure of intravesical therapy

7.3.1. Recurrence during or after intravesical chemotherapy751Patients with NMIBC recurrence during or after a chemo-<br/>therapy regimen can benefit from BCG instillations.752

Prior intravesical chemotherapy has no impact on the <sup>754</sup> effect of BCG instillations [105] (LE: 1a). <sup>755</sup>

756 7.3.2. Treatment failure after intravesical BCG immunotherapy Several categories of BCG failure, broadly defined as any HG 757 disease occurring during or after BCG therapy, have been 758 759 proposed (Table 12). NMIBC may not respond at all (BCG-760 refractory) or may relapse after an initial response (BCG-761 relapsing). Some evidence suggests that patients with BCG 762 relapse have better outcomes than patients with BCGrefractory disease [142]. 763

To be able to specify the subgroup of patients for whom 764 additional BCG is unlikely to provide benefit, the category of 765 BCG-unresponsive tumour was defined [143], which 766

#### Table 12 – Categories of HG recurrence during or after BCG therapy

#### BCG-refractory tumour

1. If T1 G3/HG tumour is present at 3 mo [144,145] (LE: 3).

2. If TaG3/HG tumour is present after 3 months and/or at 6 mo, after either re-induction or first course of maintenance [146] (LE: 4).

3. If CIS (without concomitant papillary tumour) is present at 3 mo and persists at 6 mo after either reinduction or a first course of maintenance. For patients with CIS present at 3 mo, an additional BCG course can achieve a complete response in >50% of cases [146] (LE: 1b).

4. If HG tumour appears during BCG maintenance therapy.

#### **BCG-relapsing tumour**

Recurrence of G3/HG (WHO 1973/2004) tumour after completion of BCG maintenance, despite an initial response.

BCG-unresponsive tumour

BCG-unresponsive tumours include all BCG refractory tumours and those with T1/Ta HG recurrence within 6 mo of completion of adequate BCG exposure <sup>b</sup> or CIS within 12 mo of completion of adequate BCG exposure [143] (LE: 4).

#### **BCG intolerance**

Severe side effects that prevent further BCG instillation before completing treatment [124].

BCG = bacillus Calmette-Guérin; CIS = carcinoma in situ; HG = high grade; LE = level of evidence; LG = low grade; WHO = World Health Organization.

<sup>a</sup> LG recurrence during or after BCG treatment is not considered to be a BCG failure.

<sup>b</sup> Adequate BCG therapy is defined as completion of at least five of six doses of an initial induction course plus at least two of six doses of a second induction course or two of three doses of maintenance therapy.

767	comprises BCG-refractory [144–146] and some BCG-relaps-
768	ing tumours (Table 12).

769	7.3.3. Treatment of BCG-unresponsive tumours, late BCG-relapsing
770	tumours, LG recurrences after BCG treatment, and patients with BCG
771	intolerance

- Patients with BCG-unresponsive disease are unlikely to
  respond to further BCG therapy; RC is therefore the standard
  and preferred option. Several bladder preservation strategies are currently being investigated, including cytotoxic
  intravesical therapies [147], device-assisted instillations
  [148,149], intravesical immunotherapy [150], systemic
  immunotherapy [151], and gene therapy [152].
- 779 An RCT including patients with predominantly high-risk 780 NMIBC failing at least one previous BCG induction course 781 demonstrated that MMC combined with microwave-in-782 duced hyperthermia provided 35% overall DFS at 2 yr as 783 compared to 41% in the control arm (treated with either 784 BCG, MMC, or MMC and electromotive drug administration 785 at the discretion of the investigator) [149]. The systemic 786 immunotherapy drug pembrolizumab was recently granted 787 US Food and Drug Administration approval on the basis of a 788 phase 2 study showing a 40% complete response rate in 789 BCG-unresponsive CIS [151]. Promising data from a phase 790 3 multicentre trial with intravesical nadofaragene firade-791 novec were published, showing a complete response in 792 53.4% of patients with BCG-unresponsive CIS [152].

Repeat BCG therapy may be appropriate for non-HG and <sup>793</sup> even for some HG recurrent tumours, namely those <sup>794</sup> relapsing beyond 1 yr after BCG exposure [153] (LE: 3). <sup>795</sup>

Treatment decisions in LG recurrences after BCG should be individualised according to the tumour characteristics. <sup>797</sup> Little is known about the optimal treatment for patients <sup>798</sup> with high-risk tumours who could not complete BCG <sup>799</sup> instillations because of intolerance. Treatment options for the various categories of BCG failure are presented in Table 13. <sup>802</sup>

#### 7.4. Radical cystectomy for NMIBC

803

There are several reasons to consider immediate RC for selected patients with NMIBC: 804

- The staging accuracy for T1 tumours via TURB is low, with 27–51% of patients upstaged to muscle-invasive tumour at RC [154,155] (LE: 3). 808
- Some patients with NMIBC experience disease progression to muscle-invasive disease (Table 8).
   810
- Patients who experience disease progression to the muscle-invasive stage have worse prognosis than those who present with primary muscle-invasive disease [156].

The potential benefit of RC must be weighed against its <sup>814</sup> risks, morbidity, and impact on quality of life, and should be <sup>815</sup>

Category	Treatment options
BCG-unresponsive	1. RC.
	2. Enrolment in clinical trials assessing new treatment strategies.
	3. Bladder-preserving strategies for patients unsuitable for or refusing RC.
Late BCG-relapsing	1. RC or a repeat BCG course according to the individual situation.
T1/Ta HG recurrence	2. Bladder-preserving strategies.
>6 mo or carcinoma in situ	
>12 mo since last BCG exposure	
recurrence after BCG for primary intermediate-risk tumour	1. Repeat BCG or intravesical chemotherapy.
	2. RC.

#### Table 13 - Treatment options for the various categories of BCG failure

# ARTICLE IN PRESS

Table 14 – Guidelines for follow-up of patients after transurethral resection of the bladder for non-muscle-invasive bladder cancer

Recommendation	Strength rating
Base follow-up of Ta/T1 tumours and carcinoma in situ on regular cystoscopy.	Strong
Patients with low-risk Ta tumours should undergo cystoscopy at 3 mo. If negative, subsequent cystoscopy is advised 9 mo later, and then yearly for 5 yr.	Weak
Patients with high-risk and those with very high-risk tumours treated conservatively should undergo cystoscopy and urinary cytology at 3 mo. If negative, subsequent cystoscopy and cytology should be repeated every 3 mo for a period of 2 yr, every 6 mo thereafter up to 5 yr, and then yearly.	Weak
Patients with intermediate-risk Ta tumours should have an in-between (individualised) follow-up scheme using cystoscopy.	Weak
Regular (yearly) upper tract imaging (CT-IVU or IVU) is recommended for high-risk and very high-risk tumours.	Weak
Endoscopy under anaesthesia and bladder biopsies should be performed when office cystoscopy shows suspicious findings or if urinary cytology is positive.	Strong
During follow-up for patients with positive cytology and no visible tumour in the bladder, mapping biopsies or PDD-guided biopsies (if equipment is available) and investigation of extravesical locations (CT urography, prostatic urethra biopsy) are recommended.	Strong
For patients initially diagnosed with Ta LG/G1-2 bladder cancer, use ultrasound of the bladder during surveillance if cystoscopy is not possible or is refused by the patient.	Weak

discussed with patients. It is reasonable to propose
immediate RC for patients with NMIBC who are at very
high risk of disease progression (see Sections 6.3 and
Table 7) [52,71,73,157] (LE: 3).

Early RC is strongly recommended for patients with BCG unresponsive tumours and should be considered for late
 BCG-relapsing HG tumours (Tables 10 and 13). A delay in RC
 may lead to shorter disease-specific survival [158] (LE: 3).

### 824 8. Follow-up of patients with NMIBC

Owing to the risk of recurrence and progression, patients with NMIBC need surveillance following therapy. The frequency and duration of cystoscopy and imaging follow-up should reflect the individual patient's degree of risk (see the guidelines in Table 14).

When planning the follow-up schedule and methods, the following points should be considered:

- Prompt detection of muscle-invasive and HG/G3 nonmuscle-invasive recurrence is crucial because a delay in diagnosis and therapy can be life-threatening. Therefore, the best surveillance strategy for these patients will continue to include frequent cystoscopy and cytology.
- 837 Tumour recurrence in the low-risk group is nearly always 838 of low stage and LG/G1. Small Ta G1/LG papillary 839 recurrence does not present an immediate danger to 840 the patient and early detection is not essential for 841 successful therapy [159] (LE: 2b). Fulguration of small 842 papillary recurrences on an outpatient basis could be safe 843 [160] (LE: 3). Multiple authors have suggested active 844 surveillance in selected cases [161] (LE: 3/2a).
- The first cystoscopy after TURB at 3 mo is an important prognostic indicator for recurrence and progression [77,162–164] (LE: 1a). Therefore, the first cystoscopy should always be performed 3 mo after TURB in all patients with Ta or T1 tumours or CIS.
- For low-risk tumours, the risk of recurrence after 5 yr of recurrence-free status is low [163] (LE: 3). Therefore, for

low-risk tumours, discontinuation of cystoscopy or replacement with less invasive methods can be considered after 5 yr of follow-up [164].

- For tumours originally classified as intermediate, high, or very high risk and treated conservatively, recurrences after 10 yr of tumour-free status are not unusual [165]
   (LE: 3). Therefore, life-long follow-up is recommended [164].
- The follow-up strategy must reflect the risk of extravesical recurrence (prostatic urethra in men and UUT in both genders). 860
- The risk of UUT recurrence is higher for patients with multiple and high-risk tumours [32] (LE: 3). 864
- 865 • Research has been carried out into the usefulness of 866 urinary cytology versus urinary markers as an adjunct to 867 cystoscopy in NMIBC follow-up [42,43,166]. One prospec-868 tive randomised study found that knowledge of positive 869 test results (microsatellite analysis) can improve the 870 quality of follow-up cystoscopy [41] (LE: 1b), supporting 871 the adjunctive role of a noninvasive urine test performed **Q8**<sup>872</sup> before follow-up cystoscopy [41] (see Section 5.7.3).
- For patients initially diagnosed with Ta G1–2/LG BC, US of the bladder or a urinary marker may be used for surveillance if cystoscopy is not possible or is refused by the patient [167].
- According to current knowledge, no urinary marker can replace cystoscopy during follow-up or reduce the cystoscopy frequency on a routine basis.

Author contributions:Marko Babjuk had full access to all the data in the880study and takes responsibility for the integrity of the data and the881accuracy of the data analysis.882

Study concept and design: Babjuk.

 Acquisition of data: Babjuk, Burger, Capoun, Cohen, Compérat, Dom 884

 inguez Escrig, Gontero, Liedberg, Masson-Lecomte, Mostafid, Palou, van
 885

 Rhijn, Rouprêt, Shariat, Seisen, Soukup, Sylvester.
 886

 Aragueir, and interpretation of data; Babiuk, Burger, Capoun, Cohen
 887

Analysis and interpretation of data:Babjuk, Burger, Capoun, Cohen,887Compérat, Dominguez Escrig, Gontero, Liedberg, Masson-Lecomte,888Mostafid, Palou, van Rhijn, Rouprêt, Shariat, Seisen, Soukup, Sylvester.889

883

- <sup>890</sup> Drafting of the manuscript: Babjuk.
- <sup>891</sup> Critical revision of the manuscript for important intellectual content:
- Babjuk, Burger, Capoun, Cohen, Compérat, Dominguez Escrig, Gontero,
   Jiddharg, Massan Lecomto, Mastafid, Palay, yan Philip, Boyurêt, Sharist
- <sup>893</sup> Liedberg, Masson-Lecomte, Mostafid, Palou, van Rhijn, Rouprêt, Shariat,
- <sup>894</sup> Seisen, Soukup, Sylvester.
- <sup>895</sup> Statistical analysis: None.
- <sup>896</sup> *Obtaining funding*: None.
- <sup>897</sup> Administrative, technical, or material support: Babjuk.
- <sup>898</sup> Supervision: Babjuk.
- <sup>899</sup> Other: None.

900 Financial disclosures: Marko Babjuk certifies that all conflicts of interest, 901 including specific financial interests and relationships and affiliations 902 relevant to the subject matter or materials discussed in the manuscript 903 (eg, employment/affiliation, grants or funding, consultancies, honoraria, 904 stock ownership or options, expert testimony, royalties, or patents filed, 905 received, or pending), are the following: Marko Babjuk is a company 906 consultant for Astellas and Ipsen Pharma s.r.o.; holds an advisory board 907 position for Ferring; receives company speaker honoraria from Janssen, 908 Olympus, Astellas, and Ipsen; and participates in trials run by Hamlet 909 Pharma, Ferring, and Sotio. Maximilian Burger is a company consultant 910 and receives speaker honoraria from Medac GmbH, Janssen-Cilag, Bayer 911 HealthCare AG, Merck Sharp & Dohme GmbH, Ipsen, Photocure, Pfizer, 912 and Bristol-Myers Squibb. Otakar Capoun has received consultation fees 913 from Janssen; has received company speaker honoraria from Janssen, 914 Ipsen, Astellas, and Bayer; has received fellowship/travel grants from 915 Janssen, Ipsen, and Astellas; and participates in trials by Janssen, Aragon 916 Pharmaceuticals, and Bayer s.r.o. José L. Dominguez Escrig has 917 participated in clinical trials by COMBAT BRS, BTS, Presurgy, Ipsen, 918 STORZ, Arquer, and Angiodynamics; is the national coordinator and 919 responsible for the design of the CUETO Physion-Arquer Trial; and is a 920 proctor for Angiodynamics. Paolo Gontero is a company consultant for 921 Arquer, Ferring, Ismar Healthcare, Lightpoint, and Photocure; has 922 received research grants from AB Medica, Astellas, Coloplast, Ipsen, 923 Janssen, and Storz; and has received lecture grants from Cepheid and 924 Medacs. Alexandra Masson-Lecomte has received research support from 925 the European Urological Scholarship Program and Ipsen Pharma; has 926 received consultancy fees from Ipsen Pharma, Astra Zeneca, Ambu, 927 Ferring, BMS, and Janssen Cilag; has received company speaker 928 honoraria from Astellas, Ferring, Janssen, and Ipsen Pharma; and 929 participates in studies by Janssen Cilag and Roche. A. Hugh Mostafid 930 received speaker honoraria from Medac and Bristol-Myers Squibb and 931 participates in trials by AstraZeneca PLC, Merck, and Cepheid UK. Joan 932 Palou is a company consultant for Arguer Diagnostics; receives honoraria 933 or consultation fees from Combat BRS, Olympus, Sanofi Pasteur, and 934 Cepheid, and participates in trials by Ipsen, COMBAT BRS, Presurgy, 935 STORZ, Archer, Arquer Diagnostics, IDL Biotech AB, and Palex Medical SA. 936 Bas W.G. van Rhijn is a company consultant for AstraZeneca, Ferring, and 937 QED Therapeutics. Morgan Rouprêt has received research support from 938 GSK, Pfizer, and Roche; has received consultancy fees from Lilly, GSK, 939 Ipsen, Astellas, Takeda, Sanofi Pasteur, Medac, Ferring, and Janssen Cilag; 940 has received company speaker honoraria from Roche, Zambon, Janssen, 941 Astellas, Ipsen Pharma, and Bayer S.A.S; and participates in studies by 942 Pfizer and Roche. Shahrokh F. Shariat is a company consultant for 943 Olympus and Jansen; receives company speaker honoraria from Astellas, 944 AstraZeneca, Bayer, BMS, Cepheid, Ferring, Ipsen, Janssen, and Lilly; 945 participates in company-sponsored speaker bureaus for BMS, MSD, 946 Roche, Ipsen, and Olympus; participates in trials by Roche, MSD, and 947 BMS; and owns patents for a method to determine prognosis after 948 therapy for prostate cancer, methods to determine prognosis after 949 therapy for bladder cancer, prognostic methods for patients with 950 prostatic disease, and a soluble Fas urinary marker for detection of 951 bladder transitional cell carcinoma. Richard J. Sylvester receives consultation fees from Arquer Diagnostics; is a company consultant952for Medac GmbH and Arquer Diagnostics; and receives research support953from Ferring International Center SA. The remaining authors have954nothing to disclose.955

#### Funding/Support and role of the sponsor: None.

#### References

- Phillips B. Oxford Centre for Evidence-based Medicine levels of Q9 evidence. Updated by Jeremy Howick, March 2009. Oxford, UK: <sup>957</sup> CEBM; 2009.
- [2] Guyatt GH, Oxman AD, Kunz R, et al. Going from evidence to recommendations. BMJ 2008;336:1049–51.
- [3] International Agency for Research on Cancer. Estimated number of new cases in 2020, worldwide, both sexes, all ages. Geneva, <sup>959</sup> Switzerland: World Health Organization; 2021.
- [4] Chavan S, Bray F, Lortet-Tieulent J, Goodman M, Jemal A. International variations in bladder cancer incidence and mortality. Eur Urol 2014;66:59–73.
- [5] Comperat E, Larre S, Roupret M, et al. Clinicopathological characteristics of urothelial bladder cancer in patients less than 40 years old. Virch Arch 2015;466:589–94.
   962
- [6] van Osch FH, Jochems SH, van Schooten FJ, Bryan RT, Zeegers MP.
   Quantified relations between exposure to tobacco smoking and bladder cancer risk: a meta-analysis of 89 observational studies.
   <sup>963</sup> 964
   Int J Epidemiol 2016;45:857–70.
- [7] Pesch B, Taeger D, Johnen G, et al. Screening for bladder cancer with urinary tumor markers in chemical workers with exposure to aromatic amines. Int Arch Occup Environ Health 2014;87:715–24.
   966
- [8] Egbers L, Grotenhuis AJ, Aben KK, Witjes JA, Kiemeney LA, Vermeulen SH. The prognostic value of family history among patients with urinary bladder cancer. Int J Cancer 2015;136:1117–24.
   967
- [9] Zhong JH, Zhao Z, Liu J, Yu HL, Zhou JY, Shi R. Association between APE1 Asp148Glu polymorphism and the risk of urinary cancers: a meta-analysis of 18 case-control studies. OncoTargets Ther 2016;9:1499–510.
   970
- [10] Martin C, Leiser CL, O'Neil B, et al. Familial cancer clustering in urothelial cancer: a population-based case-control study. J Natl Cancer Inst 2018;110:527–33.
- [11] Steinmaus C, Ferreccio C, Acevedo J, et al. Increased lung and bladder cancer incidence in adults after in utero and early-life arsenic exposure. Cancer Epidemiol Biomarkers Prev 2014;23:1529–38.
- [12] Witlox WJA, van Osch FHM, Brinkman M, et al. An inverse association between the Mediterranean diet and bladder cancer risk: a pooled analysis of 13 cohort studies. Eur J Nutr 2020;59:287–96.
- [13] Jochems SHJ, Reulen RC, van Osch FHM, et al. Fruit consumption and the risk of bladder cancer: a pooled analysis by the Bladder Cancer Epidemiology and Nutritional Determinants Study. Int J Cancer 2020;147:2091–100.
- [14] Tuccori M, Filion KB, Yin H, Yu OH, Platt RW, Azoulay L. Pioglitazone use and risk of bladder cancer: population based cohort study. BMJ 2016;352:i1541.
- [15] Brierley JD, Gospodarowicz MK, Wittekind C, editors. International Union Against Cancer TNM classification of malignant tumors. ed.
   <sup>981</sup> 8. New York, NY: Wiley-Blackwell; 2017. p. 263.
- [16] van Rhijn BW, van der Kwast TH, Alkhateeb SS, et al. A new and highly prognostic system to discern T1 bladder cancer substage.
   <sup>982</sup> Eur Urol 2012;61:378–84.
- [17] Moch H, Humphrey P, Ulbright T, Reuter VE. WHO classification of tumours of the urinary system and male genital organs. ed. 4 Lyon, France: International Agency for Research on Cancer; 2016.
- [18] Colombo R, Hurle R, Moschini M, et al. Feasibility and clinical roles of different substaging systems at first and second transurethral
   984

15

## ICLE IN PF

EUROPEAN UROLOGY XXX (2021) XXX-XXX

985

1014

1015

1018

1019

1020

1021

1022

1025

1026

1027

resection in patients with T1 high-grade bladder cancer. Eur Urol
Focus 2018;4:87–93.

- [19] Soukup V, Capoun O, Cohen D, et al. Prognostic performance and 986 reproducibility of the 1973 and 2004/2016 World Health Organi-987 zation grading classification systems in non-muscle-invasive blad-988 der cancer: a European Association of Urology Non-muscle 989 Invasive Bladder Cancer Guidelines Panel Systematic Review. Eur Urol 2017;72:801-13.
- [20] Hentschel AE, van Rhijn BWG, Bründl J, et al. Papillary urothelial 990 neoplasm of low malignant potential (PUN-LMP): still a meaning-991 ful histo-pathological grade category for Ta, noninvasive bladder 992 tumors in 2019? Urol Oncol 2020;38:440-8.
- [21] van Rhijn BWG, Hentschel AE, Bründl J, et al. Prognostic Value of 993 the WHO1973 and WHO2004/2016 Classification Systems for 994 grade in primary Ta/T1 non-muscle-invasive bladder cancer: a 995 multicenter European Association of Urology Non-muscle-inva-996 sive Bladder Cancer Guidelines Panel study. Eur Urol Oncol 997 2021:4:182-91.
- [22] Sylvester RJ, Rodríguez O, Hernández V, et al. European Association 998 of Urology (EAU) prognostic factor risk groups for non-muscle-999 invasive bladder cancer (NMIBC) incorporating the WHO 2004/ 1000 2016 and WHO 1973 classification systems for grade: an 1001 update from the EAU NMIBC Guidelines Panel. Eur Urol 1002 2021:79:480-8.
- [23] Mangrud OM, Waalen R, Gudlaugsson E, et al. Reproducibility and 1003 prognostic value of WHO1973 and WHO2004 grading systems in 1004 TaT1 urothelial carcinoma of the urinary bladder. PLoS One 1005 2014:9:e83192.
- [24] Veskimae E, Espinos EL, Bruins HM, et al. What is the prognostic 1006 and clinical importance of urothelial and nonurothelial histologi-1007 cal variants of bladder cancer in predicting oncological outcomes 1008 in patients with muscle-invasive and metastatic bladder cancer? A 1009 European Association of Urology Muscle Invasive and Metastatic 1010 Bladder Cancer Guidelines Panel Systematic review. Eur Urol Oncol 1011 2019;2:625-42.
- [25] Comperat EM, Burger M, Gontero P, et al. Grading of urothelial 1012 carcinoma and the new "World Health Organisation classification 1013 of tumours of the urinary system and male genital organs 2016". Eur Urol Focus 2019;5:457-66.
  - [26] Seisen T, Comperat E, Leon P, Roupret M. Impact of histological variants on the outcomes of nonmuscle invasive bladder cancer after transurethral resection. Curr Opin Urol 2014;24:524-31.
- [27] Mari A, Kimura S, Foerster B, et al. A systematic review and meta-1016 analysis of the impact of lymphovascular invasion in bladder 1017 cancer transurethral resection specimens. BJU Int 2019;123:11-21.
  - [28] Marzouka NA, Eriksson P, Rovira C, Liedberg F, Sjodahl G, Hoglund M. A validation and extended description of the Lund taxonomy for urothelial carcinoma using the TCGA cohort. Sci Rep 2018;8:3737.
  - [29] Ramirez D, Gupta A, Canter D, et al. Microscopic haematuria at time of diagnosis is associated with lower disease stage in patients with newly diagnosed bladder cancer. BJU Int 2016;117:783-6.
  - [30] Trinh TW, Glazer DI, Sadow CA, Sahni VA, Geller NL, Silverman SG. Bladder cancer diagnosis with CT urography: test characteristics and reasons for false-positive and false-negative results. Abdom Radiol 2018;43:663-71.
- [31] Palou J, Rodriguez-Rubio F, Huguet J, et al. Multivariate analysis of 1023 clinical parameters of synchronous primary superficial bladder 1024 cancer and upper urinary tract tumor. J Urol 2005;174:859-61.
  - [32] Millan-Rodriguez F, Chechile-Toniolo G, Salvador-Bayarri J, Huguet-Perez J, Vicente-Rodriguez J. Upper urinary tract tumors after primary superficial bladder tumors: prognostic factors and risk groups. J Urol 2000;164:1183-7.
  - [33] Hilton S, Jones LP. Recent advances in imaging cancer of the kidney and urinary tract. Surg Oncol Clin North Am 2014;23:863-910.

- [34] Panebianco V, Narumi Y, Altun E, et al. Multiparametric magnetic 1028 resonance imaging for bladder cancer: development of VI-RADS 1029 (Vesical Imaging-Reporting and Data System). Eur Urol 1030 2018;74:294-306.
- [35] Yafi FA, Brimo F, Steinberg J, Aprikian AG, Tanguay S, Kassouf W. 1031 Prospective analysis of sensitivity and specificity of urinary cytol-1032 ogy and other urinary biomarkers for bladder cancer. Urol Oncol 1033 2015;33:66.e25-3.
- [36] Tetu B. Diagnosis of urothelial carcinoma from urine. Mod Pathol 2009;22(Suppl 2):S53-9.

1034

1049

1055

- [37] Raitanen MP, Aine R, Rintala E, et al. Differences between local and 1035 review urinary cytology in diagnosis of bladder cancer. An inter-1036 observer multicenter analysis. Eur Urol 2002;41:284-9.
- [38] Rosenthal D, Wojcik E, Kurtycz D. The Paris system for reporting 1037 urinary cytology. Cham, Switzerland: Springer International Publishing: 2016.
- [39] Meilleroux J, Daniel G, Aziza J, et al. One year of experience using 1038 the Paris system for reporting urinary cytology. Cancer Cytopathol 1039 2018:126:430-6.
- [40] Soria F, Droller MJ, Lotan Y, et al. An up-to-date catalog of available 1040 urinary biomarkers for the surveillance of non-muscle invasive 1041 bladder cancer. World J Urol 2018;36:1981-95.
- [41] van der Aa MN, Steyerberg EW, Bangma C, van Rhijn BW, Zwarthoff EC, van der Kwast TH. Cystoscopy revisited as the gold standard for 1042 detecting bladder cancer recurrence: diagnostic review bias in the 1043 randomized, prospective CEFUB trial. J Urol 2010;183:76-80.
- [42] Valenberg F, Hiar AM, Wallace E, et al. Prospective validation of an 1044 mRNA-based urine test for surveillance of patients with bladder 1045 cancer. Eur Urol 2019;75:853-60.
- [43] D'Andrea D, Soria F, Zehetmayer S, et al. Diagnostic accuracy, 1046 clinical utility and influence on decision-making of a methylation 1047 urine biomarker test in the surveillance of non-muscle-invasive 1048 bladder cancer. BJU Int 2019;123:959-67.
- [44] Konety B. Evaluation of Cxbladder and adjudication of atypical cytology and equivocal cystoscopy. Eur Urol 2019;76:238-43.
- [45] Krajewski W, Koscielska-Kasprzak K, Rymaszewska J, Zdrojowy R. 1050 How different cystoscopy methods influence patient sexual satis-1051 faction, anxiety, and depression levels: a randomized prospective 1052 trial. Qual Life Res 2017;26:625-34.
- [46] Suarez-Ibarrola R, Soria F, Abufaraj M, et al. Surgical checklist 1053 impact on recurrence-free survival of patients with non-mus-1054 cle-invasive bladder cancer undergoing transurethral resection of bladder tumour. BJU Int 2019;123:646-50.
- [47] Teoh JY, MacLennan S, Chan VW, et al. An international collabora-1056 tive consensus statement on en bloc resection of bladder tumour 1057 incorporating two systematic reviews, a two-round Delphi survey, 1058 and a consensus meeting. Eur Urol 2020;78:546-69.
- [48] Richterstetter M, Wullich B, Amann K, et al. The value of extended 1059 transurethral resection of bladder tumour (TURBT) in the treat-1060 ment of bladder cancer. BJU Int 2012;110:E76-9.
- [49] Mariappan P, Zachou A, Grigor KM. Detrusor muscle in the first, 1061 apparently complete transurethral resection of bladder tumour 1062 specimen is a surrogate marker of resection quality, predicts risk of 1063 early recurrence, and is dependent on operator experience. Eur Urol 2010;57:843-9.
- [50] Planelles Gomez J, Olmos Sanchez L, Cardosa Benet JJ, Martinez Lopez E, Vidal Moreno JF. Holmium YAG photocoagulation: safe 1064 and economical alternative to transurethral resection in small 1065 nonmuscle-invasive bladder tumors. J Endourol 2017;31:674-8.
- [51] van der Meijden A, Oosterlinck W, Brausi M, Kurth KH, Sylvester R, de Balincourt C. Significance of bladder biopsies in Ta,T1 bladder 1066 tumors: a report from the EORTC Genito-Urinary Tract Cancer 1067 Cooperative Group. EORTC-GU Group Superficial Bladder Commit-1068 tee. Eur Urol 1999:35:267-71.

1076

1077

1078

1079

1083

1084

1085

1086

1099

1100

1101

1102

1103

1104

1105

### ARTICLE IN PRESS

- [52] Palou J, Sylvester RJ, Faba OR, et al. Female gender and carcinoma in situ in the prostatic urethra are prognostic factors for recurrence, progression, and disease-specific mortality in T1G3 bladder cancer patients treated with bacillus Calmette-Guerin. Eur Urol 2012;62:118–25.
- [53] Mungan MU, Canda AE, Tuzel E, Yorukoglu K, Kirkali Z. Risk factors
   for mucosal prostatic urethral involvement in superficial transitional cell carcinoma of the bladder. Eur Urol 2005;48:760–3.
  - [54] Brant A, Daniels M, Chappidi MR, et al. Prognostic implications of prostatic urethral involvement in non-muscle-invasive bladder cancer. World J Urol 2019;37:2683–9.
  - [55] Mowatt G, N'Dow J, Vale L, et al. Photodynamic diagnosis of bladder cancer compared with white light cystoscopy: systematic review and meta-analysis. Int J Technol Assess Health Care 2011;27:3–10.
- [56] Neuzillet Y, Methorst C, Schneider M, et al. Assessment of diagnostic gain with hexaminolevulinate (HAL) in the setting of newly diagnosed non-muscle-invasive bladder cancer with positive results on urine cytology. Urol Oncol 2014;32:1135–40.
  - [57] Draga RO, Grimbergen MC, Kok ET, Jonges TN, van Swol CF, Bosch JL. Photodynamic diagnosis (5-aminolevulinic acid) of transitional cell carcinoma after bacillus Calmette-Guerin immunotherapy and mitomycin C intravesical therapy. Eur Urol 2010;57:655–60.
  - [58] Ray ER, Chatterton K, Khan MS, et al. Hexylaminolaevulinate fluorescence cystoscopy in patients previously treated with intravesical bacille Calmette-Guerin. BJU Int 2010;105:789–94.
- [59] Chou R, Selph S, Buckley DI, et al. Comparative effectiveness of fluorescent versus white light cystoscopy for initial diagnosis or surveillance of bladder cancer on clinical outcomes: systematic review and meta-analysis. J Urol 2017;197:548–58.
- [60] Kim SB, Yoon SG, Tae J, et al. Detection and recurrence rate of
   transurethral resection of bladder tumors by narrow-band imag ing: prospective, randomized comparison with white light cystos copy. Invest Clin Urol 2018;59:98–105.
- [61] Naito S, Algaba F, Babjuk M, et al. The Clinical Research Office of the Endourological Society (CROES) multicentre randomised trial of narrow band imaging-assisted transurethral resection of bladder tumour (TURBT) versus conventional white light imaging-assisted TURBT in primary non-muscle-invasive bladder cancer patients: trial protocol and 1-year results. Eur Urol 2016;70:506–15.
  - [62] Cumberbatch MGK, Foerster B, Catto JWF, et al. Repeat transurethral resection in non-muscle-invasive bladder cancer: a systematic review. Eur Urol 2018;73:925–33.
    - [63] Naselli A, Hurle R, Paparella S, et al. Role of restaging transurethral resection for T1 non-muscle invasive bladder cancer: a systematic review and meta-analysis. Eur Urol Focus 2018;4:558–67.
  - [64] Eroglu A, Ekin RG, Koc G, Divrik RT. The prognostic value of routine second transurethral resection in patients with newly diagnosed stage pT1 non-muscle-invasive bladder cancer: results from randomized 10-year extension trial. Int J Clin Oncol 2020;25:698– 704.
- [65] Gordon PC, Thomas F, Noon AP, Rosario DJ, Catto JWF. Long-term
   outcomes from re-resection for high-risk non-muscle-invasive
   bladder cancer: a potential to rationalize use. Eur Urol Focus
   2019;5:650–7.
- [66] Bishr M, Lattouf JB, Latour M, Saad F. Tumour stage on re-staging
   transurethral resection predicts recurrence and progression-free
   survival of patients with high-risk non-muscle invasive bladder
   cancer. Can Urol Assoc J 2014;8:E306–10.
- [67] Palou J, Pisano F, Sylvester R, et al. Recurrence, progression and cancer-specific mortality according to stage at re-TUR in T1G3 bladder cancer patients treated with BCG: not as bad as previously thought. World J Urol 2018;36:1621–7.

- [68] Gontero P, Sylvester R, Pisano F, et al. The impact of re-transurethral resection on clinical outcomes in a large multicentre cohort of patients with T1 high-grade/grade 3 bladder cancer treated with bacille Calmette-Guerin. BJU Int 2016;118:44–52.
- [69] Baltaci S, Bozlu M, Yildirim A, et al. Significance of the interval between first and second transurethral resection on recurrence and progression rates in patients with high-risk non-muscle-invasive bladder cancer treated with maintenance intravesical bacillus Calmette-Guerin. BJU Int 2015;116:721–6.
- [70] Grignon D, Brimo F, Comperat E, et al. Urinary tract carcinomas biopsy and transurethral resection specimen. Sydney, Australia: 1122 International Collaboration on Cancer Reporting; 2019.
- [71] Sylvester RJ, van der Meijden AP, Oosterlinck W, et al. Predicting recurrence and progression in individual patients with stage Ta T1
   bladder cancer using EORTC risk tables: a combined analysis of 2596 patients from seven EORTC trials. Eur Urol 2006;49, 466–5.
- [72] Lammers RJ, Hendriks JC, Rodriguez Faba OR, Witjes WP, Palou J, Witjes JA. Prediction model for recurrence probabilities after intravesical chemotherapy in patients with intermediate-risk non-muscle-invasive bladder cancer, including external validation. World J Urol 2016;34:173–80.
- [73] Fernandez-Gomez J, Madero R, Solsona E, et al. Predicting non-<br/>muscle invasive bladder cancer recurrence and progression in<br/>patients treated with bacillus Calmette-Guerin: the CUETO scoring<br/>model. J Urol 2009;182:2195–203.1129<br/>1130
- [74] Cambier S, Sylvester RJ, Collette L, et al. EORTC nomograms and risk groups for predicting recurrence, progression, and diseasespecific and overall survival in non-muscle-invasive stage Ta-T1 urothelial bladder cancer patients treated with 1–3 years of maintenance bacillus Calmette-Guerin. Eur Urol 2016;69:60–9.
- [75] Gontero P, Sylvester R, Pisano F, et al. Prognostic factors and risk groups in T1G3 non-muscle-invasive bladder cancer patients initially treated with bacillus Calmette-Guerin: results of a retrospective multicenter study of 2451 patients. Eur Urol 2015;67:74–82.
- [76] Voskuilen CS, Seiler R, Rink M, et al. Urothelial carcinoma in bladder diverticula: a multicenter analysis of characteristics and clinical outcomes. Eur Urol Focus 2020;6:1226–32.
- [77] Palou J, Rodriguez-Rubio F, Millan F, et al. Recurrence at three months and high-grade recurrence as prognostic factor of progression in multivariate analysis of T1G2 bladder tumors. Urology 2009;73:1313–7.
- [78] Alkhateeb SS, Neill M, Bar-Moshe S, et al. Long-term prognostic value of the combination of EORTC risk group calculator and molecular markers in non-muscle-invasive bladder cancer patients treated with intravesical bacille Calmette-Guerin. Urol Ann 2011;3:119–26.
- [79] Lamm DL. Carcinoma in situ. Urol Clin North Am 1992;19:499– 508.
- [80] Losa A, Hurle R, Lembo A. Low dose bacillus Calmette-Guerin for carcinoma in situ of the bladder: long-term results. J Urol 2000;163:68–71.
- [81] Griffiths TR, Charlton M, Neal DE, Powell PH. Treatment of carcinoma in situ with intravesical bacillus Calmette-Guerin without maintenance. J Urol 2002;167:2408–12.
- [82] Rink M, Xylinas E, Babjuk M, et al. Smoking reduces the efficacy of intravesical bacillus Calmette-Guerin immunotherapy in nonmuscle-invasive bladder cancer. Eur Urol 2012;62:1204–6.
- [83] Crivelli JJ, Xylinas E, Kluth LA, Rieken M, Rink M, Shariat SF. Effect of smoking on outcomes of urothelial carcinoma: a systematic review of the literature. Eur Urol 2014;65:742–54.
- [84] Brocks CP, Buttner H, Bohle A. Inhibition of tumor implantation by intravesical gemcitabine in a murine model of superficial bladder cancer. J Urol 2005;174:1115–8.

EUROPEAN UROLOGY XXX (2021) XXX-XXX

- [85] Oosterlinck W, Kurth KH, Schroder F, Bultinck J, Hammond B, Sylvester R. A prospective European Organization for Research and Treatment of Cancer Genitourinary Group randomized trial comparing transurethral resection followed by a single intravesical instillation of epirubicin or water in single stage Ta, T1 papillary
- carcinoma of the bladder. J Urol 1993;149:749–52.
- [86] Sylvester Rj, Oosterlinck W, Holmang S, et al. Systematic review and individual patient data meta-analysis of randomized trials comparing a single immediate instillation of chemotherapy after transurethral resection with transurethral resection alone in patients with stage pTa-pT1 urothelial carcinoma of the bladder: which patients benefit from the instillation? Eur Urol 2016;69:231-44.
- [87] Sylvester RJ, Oosterlinck W, van der Meijden AP. A single immediate postoperative instillation of chemotherapy decreases the risk of recurrence in patients with stage Ta T1 bladder cancer: a metaanalysis of published results of randomized clinical trials. J Urol 2004;171:2186–90.
- [88] Abern MR, Owusu RA, Anderson MR, Rampersaud EN, Inman BA.
   Perioperative intravesical chemotherapy in non-muscle-invasive
   bladder cancer: a systematic review and meta-analysis. J Natl Compr Cancer Netw 2013;11:477–84.
- [89] Perlis N, Zlotta AR, Beyene J, Finelli A, Fleshner NE, Kulkarni GS.
   Immediate post-transurethral resection of bladder tumor intravesical chemotherapy prevents non-muscle-invasive bladder cancer recurrences: an updated meta-analysis on 2548 patients and quality-of-evidence review. Eur Urol 2013;64:421–30.
- [90] Messing EM, Tangen CM, Lerner SP, et al. Effect of intravesical instillation of gemcitabine vs saline immediately following resection of suspected low-grade non-muscle-invasive bladder cancer on tumor recurrence: SWOG S0337 randomized clinical trial. JAMA 2018;319:1880–8.
- [91] Zhou Z, Zhao S, Lu Y, et al. Meta-analysis of efficacy and safety of continuous saline bladder irrigation compared with intravesical chemotherapy after transurethral resection of bladder tumors. World J Urol 2019;37:1075–84.
- [92] Bohle A, Jurczok A, Ardelt P, et al. Inhibition of bladder carcinoma cell adhesion by oligopeptide combinations in vitro and in vivo. J Urol 2002;167:357–63.
- [93] Tolley DA, Parmar MK, Grigor KM, et al. The effect of intravesical mitomycin C on recurrence of newly diagnosed superficial bladder
   cancer: a further report with 7 years of follow up. J Urol 1996;155:1233-8.
- [94] Huncharek M, McGarry R, Kupelnick B. Impact of intravesical chemotherapy on recurrence rate of recurrent superficial transitional cell carcinoma of the bladder: results of a meta-analysis. Anticancer Res 2001;21:765–9.
- [95] Bosschieter J, Nieuwenhuijzen JA, van Ginkel T, et al. Value of an immediate intravesical instillation of mitomycin C in patients with non-muscle-invasive bladder cancer: a prospective multicentre randomised study in 2243 patients. Eur Urol 2018;73:226–32.
- [96] Sylvester RJ, Oosterlinck W, Witjes JA. The schedule and duration
   of intravesical chemotherapy in patients with non-muscle-inva sive bladder cancer: a systematic review of the published results of
   randomized clinical trials. Eur Urol 2008;53:709–19.
- [97] Bosschieter J, Nieuwenhuijzen JA, Vis AN, et al. An immediate, single intravesical instillation of mitomycin C is of benefit in patients with non-muscle-invasive bladder cancer irrespective of prognostic risk groups. Urol Oncol 2018;36:400.e7–400.e14.
- [98] Elsawy AA, El-Assmy AM, Bazeed MA, Ali-El-Dein B. The value of
   immediate postoperative intravesical epirubicin instillation as an
   adjunct to standard adjuvant treatment in intermediate and high risk non-muscle-invasive bladder cancer: a preliminary results of
   randomized controlled trial. Urol Oncol 2019;37:179.e9–179.e18.

[99] Au JL, Badalament RA, Wientjes MG, et al. Methods to improve efficacy of intravesical mitomycin C: results of a randomized phase III trial. J Natl Cancer Inst 2001;93:597–604.

1201

1202

1203

1204

1205

1216

1217

1218

1219

1220

1221

1222

1223

- [100] Giesbers AA, Van Helsdingen PJ, Kramer AE. Recurrence of superficial bladder carcinoma after intravesical instillation of mitomycin-C. Comparison of exposure times. Br J Urol 1989;63:176–9.
- [102] Arends TJ, van der Heijden AG, Witjes JA. Combined chemohyperthermia: 10-year single center experience in 160 patients with nonmuscle invasive bladder cancer. J Urol 2014;192:708–13.
   [102] Arende TJ, Nativ O, Maffaarjini M, et al. Berulte of a randomized
   [215] Arende TJ, Nativ O, Maffaarjini M, et al. Berulte of a randomized
- [103] Arends TJ, Nativ O, Maffezzini M, et al. Results of a randomised controlled trial comparing intravesical chemohyperthermia with mitomycin C versus bacillus Calmette-Guerin for adjuvant treatment of patients with intermediate- and high-risk non-muscleinvasive bladder cancer. Eur Urol 2016;69:1046–52.
- [104] Di Stasi SM, Giannantoni A, Giurioli A, et al. Sequential BCG and electromotive mitomycin versus BCG alone for high-risk superficial bladder cancer: a randomised controlled trial. Lancet Oncol 2006;7:43–51.
- [105] Malmstrom Pu, Sylvester Rj, Crawford De, et al. An individual<br/>patient data meta-analysis of the long-term outcome of random-<br/>ised studies comparing intravesical mitomycin C versus bacillus<br/>Calmette-Guerin for non-muscle-invasive bladder cancer. Eur Urol<br/>2009;56:247–56.1224<br/>1225
- [106] Shelley MD, Kynaston H, Court J, et al. A systematic review of intravesical bacillus Calmette-Guerin plus transurethral resection vs transurethral resection alone in Ta and T1 bladder cancer. BJU
   Int 2001;88:209–16.
- [107] Han RF, Pan JG. Can intravesical bacillus Calmette-Guerin reduce recurrence in patients with superficial bladder cancer? A meta-analysis of randomized trials. Urology 2006;67:1216–23.
   [1232] 1233
- [108] Shelley MD, Wilt TJ, Court J, Coles B, Kynaston H, Mason MD.
   Intravesical bacillus Calmette-Guerin is superior to mitomycin C in reducing tumour recurrence in high-risk superficial bladder cancer: a meta-analysis of randomized trials. BJU Int 2004;93:485–90.
   <sup>1235</sup> 1236
   1237
   1238
   1238
   1239
- [109] Bohle A, Jocham D, Bock PR. Intravesical bacillus Calmette-Guerin1240versus mitomycin C for superficial bladder cancer: a formal meta-<br/>analysis of comparative studies on recurrence and toxicity. J Urol12412003;169:90–5.1243
- [110] Duchek M, Johansson R, Jahnson S, et al. Bacillus Calmette-Guerin is superior to a combination of epirubicin and interferon-alpha2b in the intravesical treatment of patients with stage T1 urinary bladder cancer. A prospective, randomized, Nordic study. Eur Urol 2010;57:25–31.
   1244
- [111] Sylvester RJ, Brausi MA, Kirkels WJ, et al. Long-term efficacy results of EORTC Genito-Urinary Group randomized phase 3 study 30911 comparing intravesical instillations of epirubicin, bacillus Calmette-Guerin, and bacillus Calmette-Guerin plus isoniazid in patients with intermediate- and high-risk stage Ta T1 urothelial carcinoma of the bladder. Eur Urol 2010;57:766–73.
   1249

   1249
   1250
   1251
   1251
   1251
   1252
   1253
   1254
   1253
   1254
- [112] Jarvinen R, Kaasinen E, Sankila A, Rintala E. Long-term efficacy of<br/>maintenance bacillus Calmette-Guerin versus maintenance mito-<br/>mycin C instillation therapy in frequently recurrent TaT1 tumours<br/>without carcinoma in situ: a subgroup analysis of the prospective,<br/>randomised FinnBladder I study with a 20-year follow-up. Eur Urol<br/>2009;56:260–5.1255<br/>1258<br/>1259
- [113] Bohle A, Bock PR. Intravesical bacille Calmette-Guerin versus nitomycin C in superficial bladder cancer: formal meta-analysis 1261

Please cite this article in press as: Babjuk M, et al. European Association of Urology Guidelines on Non-muscle-invasive Bladder Cancer (Ta, T1, and Carcinoma in Situ). Eur Urol (2021), https://doi.org/10.1016/j.eururo.2021.08.010

18

1157

1158

1337

1338

1339

1340

1263of comparative studies on tumor progression. Urology12642004;63:682-6.

- 1265 [114] Sylvester RJ, van der MA, Lamm DL. Intravesical bacillus Calmette 1266 Guerin reduces the risk of progression in patients with superficial
   1267 bladder cancer: a meta-analysis of the published results of ran 1268 domized clinical trials. J Urol 2002;168:1964–70.
- [115] Boehm BE, Cornell JE, Wang H, Mukherjee N, Oppenheimer JS, Svatek RS. Efficacy of bacillus Calmette-Guerin strains for treatment of nonmuscle invasive bladder cancer: a systematic review and network meta-analysis. J Urol 2017;198:503–10.
- 1272 [116] van der Meijden AP, Sylvester RJ, Oosterlinck W, Hoeltl W, Bono AV.
   1273 Maintenance bacillus Calmette-Guerin for Ta T1 bladder tumors is 1274 not associated with increased toxicity: results from a European 1275 Organisation for Research and Treatment of Cancer Genito-Urinary 1276 Group phase III trial. Eur Urol 2003;44:429–34.
- [117] Larsen ES, Nordholm AC, Lillebaek T, Holden IK, Johansen IS. The
   epidemiology of bacille Calmette-Guerin infections after bladder
   instillation from 2002 through 2017: a nationwide retrospective
   cohort study. BJU Int 2019;124:910–6.
- [118] Brausi M, Oddens J, Sylvester R, et al. Side effects of bacillus
  [1282 Calmette-Guerin (BCG) in the treatment of intermediate- and
  high-risk Ta, T1 papillary carcinoma of the bladder: results of
  the EORTC Genito-Urinary Cancers Group randomised phase
  3 study comparing one-third dose with full dose and 1 year with
  3 years of maintenance BCG. Eur Urol 2014;65:69–76.
- [119] Oddens JR, Sylvester RJ, Brausi MA, et al. Increasing age is not associated with toxicity leading to discontinuation of treatment in patients with urothelial non-muscle-invasive bladder cancer randomised to receive 3 years of maintenance bacille Calmette-Guerin: results from European Organisation for Research and Treatment of Cancer Genito-Urinary Group study 30911. BJU Int 2016;118:423–8.
- [120] Unda-Urzaiz M, Cozar-Olmos JM, Minana-Lopez B, et al. Safety and efficacy of various strains of bacille Calmette-Guerin in the treatment of bladder tumours in standard clinical practice. Actas Urol Esp 2018;42:238–48.
- [121] Danielsson G, Malmstrom PU, Jahnson S, Wijkstrom H, Nyberg T, Thulin H. Bladder health in patients treated with BCG instillations
   for T1G2-G3 bladder cancer – a follow-up five years after the start
   of treatment. Scand J Urol 2018;52:377–84.
- 1300 [122] Herr HW. Outpatient urological procedures in antibiotic-naive patients with bladder cancer with asymptomatic bacteriuria. BJU Int 2012;110:E658–60.
- Roumeguere T, Broeders N, Jayaswal A, et al. Bacillus Calmette Guerin therapy in non-muscle-invasive bladder carcinoma after
   renal transplantation for end-stage aristolochic acid nephropathy. Transplant Int 2015;28:199–205.
- <sup>1305</sup>Q10 [124] Witjes JA, Palou J, Soloway M, et al. Clinical practice recommenda <sup>1306</sup>tions for the prevention and management of intravesical therapy <sup>1307</sup>associated adverse events. Eur Urol Suppl 2008;7:667–74.
- Interview (125) Morales A, Eidinger D, Bruce AW. Intracavitary bacillus Calmette Guerin in the treatment of superficial bladder tumors. J Urol
   1976;116:180–3.
- [126] Lamm DL, Blumenstein BA, Crissman JD, et al. Maintenance bacillus Calmette-Guerin immunotherapy for recurrent TA, T1 and carcinoma in situ transitional cell carcinoma of the bladder: a randomized Southwest Oncology Group Study. J Urol 2000;163:1124–9.
- I27] Grimm MO, van der Heijden AG, Colombel M, et al. Treatment of high-grade non-muscle-invasive bladder carcinoma by standard number and dose of BCG instillations versus reduced number and standard dose of BCG instillations: results of the European Association of Urology Research Foundation randomised phase III
   clinical trial "NIMBUS". Eur Urol 2020;78:690–8.

- [128] Martinez-Pineiro L, Portillo JA, Fernandez JM, et al. Maintenance therapy with 3-monthly bacillus Calmette-Guerin for 3 years is not superior to standard induction therapy in high-risk nonmuscle-invasive urothelial bladder carcinoma: final results of randomised CUETO study 98013. Eur Urol 2015;68:256–62.
   <sup>1322</sup>
- [129] Oddens J, Brausi M, Sylvester R, et al. Final results of an EORTC-GU
   [127] Oddens J, Brausi M, Sylvester R, et al. Final results of an EORTC-GU
   [1327] Cancers Group randomized study of maintenance bacillus Calmette-Guerin in intermediate- and high-risk Ta, T1 papillary carcinoma of the urinary bladder: one-third dose versus full dose and 1 year versus 3 years of maintenance. Eur Urol 2013;63:462–72.
   [1327] Isolard M, Sylvester R, et al. Final results of an EORTC-GU
   [1328] Isolard M, Sylvester R, et al. Final results of an EORTC-GU
   [1329] Isolard M, Sylvester R, et al. Final results of an EORTC-GU
   [1329] Isolard M, Sylvester R, et al. Final results of an EORTC-GU
   [1329] Isolard M, Sylvester R, et al. Final results of an EORTC-GU
   [1329] Isolard M, Sylvester R, et al. Final results of an EORTC-GU
   [1329] Isolard M, Sylvester R, et al. Final results of an EORTC-GU
   [1329] Isolard M, Sylvester R, et al. Final results of an EORTC-GU
   [1329] Isolard M, Sylvester R, et al. Final results of an EORTC-GU
   [1329] Isolard M, Sylvester R, et al. Final results of an EORTC-GU
   [1320] Isolard M, Sylvester R, et al. Final results of an EORTC-GU
   [1320] Isolard M, Sylvester R, et al. Final results of an EORTC-GU
   [1320] Isolard M, Sylvester R, et al. Final results of an EORTC-GU
   [1320] Isolard M, Sylvester R, et al. Final results of an EORTC-GU
   [1320] Isolard M, Sylvester R, et al. Final results of an EORTC-GU
   [1320] Isolard M, Sylvester R, et al. Final results of an EORTC-GU
   [1320] Isolard M, Sylvester R, et al. Final results of an EORTC-GU
   [1320] Isolard M, Sylvester R, et al. Final results of an EORTC-GU
   [1320] Isolard M, Sylvester R, et al. Final results of an EORTC-GU
   [1320] Isolard M, Sylvester R, et al. Final results of an EORTC-G
- [130] Martinez-Pineiro JA, Flores N, Isorna S, et al. Long-term follow-up of a randomized prospective trial comparing a standard 81 mg dose of intravesical bacille Calmette-Guerin with a reduced dose of 1334
   [131] Martinez Diracire JA, Martinez Diracire L, Scheme E, et al. Has a 2
- [131] Martinez-Pineiro JA, Martinez-Pineiro L, Solsona E, et al. Has a 3fold decreased dose of bacillus Calmette-Guerin the same efficacy against recurrences and progression of T1G3 and Tis bladder tumors than the standard dose? Results of a prospective randomized trial. J Urol 2005;174:1242–7.
- [132] Ojea A, Nogueira JL, Solsona E, et al. A multicentre, randomised prospective trial comparing three intravesical adjuvant therapies for intermediate-risk superficial bladder cancer: low-dose bacillus Calmette-Guerin (27 mg) versus very low-dose bacillus Calmette-Guerin (13.5 mg) versus mitomycin C. Eur Urol 2007;52:1398–406.
   [1341] 1342
- [133] Solsona E, Madero R, Chantada V, et al. Sequential combination of mitomycin C plus bacillus Calmette-Guerin (BCG) is more effective but more toxic than BCG alone in patients with non-muscleinvasive bladder cancer in intermediate- and high-risk patients: final outcome of CUETO 93009, a randomized prospective trial. Eur Urol 2015;67:508–16.
  [1346] 1346
- [134] Huang D, Jin YH, Weng H, Huang Q, Zeng XT, Wang XH. Combination of intravesical bacille Calmette-Guerin and chemotherapy vs. bacille Calmette-Guerin alone in non-muscle invasive bladder cancer: a meta-analysis. Front Oncol 2019;9:121.
   [135] 1351
- [135] Shepherd AR, Shepherd E, Brook NR. Intravesical bacillus Calmette-Guerin with interferon-alpha versus intravesical bacillus
   Calmette-Guerin for treating non-muscle-invasive bladder cancer.
   Cochrane Database Syst Rev 2017;2017:CD012112.
- [136] Jarvinen R, Marttila T, Kaasinen E, et al. Long-term outcome of patients with frequently recurrent non-muscle-invasive bladder carcinoma treated with one perioperative plus four weekly instillations of mitomycin C followed by monthly bacillus Calmette-Guerin (BCG) or alternating BCG and interferon-alpha2b instillations: prospective randomised FinnBladder-4 study. Eur Urol 2015;68:611-7.
  [136] Jarvinen R, Marttila T, Kaasinen E, et al. Long-term outcome of patients with frequently recurrent non-muscle-invasive bladder 1359
  [136] Jarvinen R, Marttila T, Kaasinen E, et al. Long-term outcome of patients with frequently recurrent non-muscle-invasive bladder 1369
  [136] Jarvinen R, Marttila T, Kaasinen E, et al. Long-term outcome of patients with frequently recurrent non-muscle-invasive bladder 1360
  [136] Jarvinen R, Marttila T, Kaasinen E, et al. Long-term outcome of patients with one perioperative plus four weekly instillations of mitomycin C followed by monthly bacillus Calmette-Guerin (BCG) or alternating BCG and interferon-alpha2b instillations: prospective randomised FinnBladder-4 study. Eur Urol 2015;68:611-7.
- [137] Marttila T, Jarvinen R, Liukkonen T, et al. Intravesical bacillus Calmette-Guerin versus combination of epirubicin and interferon-alpha2a in reducing recurrence of non-muscle-invasive bladder carcinoma: FinnBladder-6 study. Eur Urol 2016;70:341–7.
   [136] 1365
   [136] 1365
   [136] 1365
   [136] 1365
   [136] 1365
   [136] 1365
   [136] 1365
   [136] 1365
   [136] 1365
   [136] 1365
   [136] 1365
   [136] 1365
   [136] 1365
   [136] 1365
   [136] 1365
   [136] 1365
   [136] 1365
   [136] 1365
   [136] 1365
   [136] 1365
   [136] 1365
   [136] 1365
   [136] 1365
   [136] 1365
   [136] 1365
   [136] 1365
   [136] 1365
   [136] 1365
   [136] 1365
   [136] 1365
- [138] Sylvester RJ, van der Meijden AP, Witjes JA, Kurth K. Bacillus
   Calmette-Guerin versus chemotherapy for the intravesical treatment of patients with carcinoma in situ of the bladder: a metaanalysis of the published results of randomized clinical trials. J
   Urol 2005;174:86–91.
- [139] Kaasinen E, Wijkstrom H, Rintala E, Mestad O, Jahnson S, Malmstrom PU. Seventeen-year follow-up of the prospective randomized Nordic CIS study: BCG monotherapy versus alternating therapy with mitomycin C and BCG in patients with carcinoma in situ of the urinary bladder. Scand J Urol 2016;50:360–8.
   <sup>1373</sup>
- [140] Solsona E, Iborra I, Ricos JV, Monros JL, Dumont R, Almenar S.
   Extravesical involvement in patients with bladder carcinoma in situ: biological and therapy implications. J Urol 1996;155:895–9.
   [141] Palevi L Parial L Klat L et al. Userbalia carcinome of the prostate.
- [141] Palou J, Baniel J, Klotz L, et al. Urothelial carcinoma of the prostate.
   <sup>13</sup>Urology 2007;69:50–61.

## **ARTICLE IN PRESS**

EUROPEAN UROLOGY XXX (2021) XXX-XXX

- 1381 [142] Herr HW, Milan TN, Dalbagni G. BCG-refractory vs. BCG-relapsing 1382 non-muscle-invasive bladder cancer: a prospective cohort outcomes study. Urol Oncol 2015;33:108.e1-e.
- Kamat AM, Sylvester RJ, Bohle A, et al. Definitions, end points, and clinical trial designs for non-muscle-invasive bladder cancer: recommendations from the International Bladder Cancer Group. J Clin Oncol 2016;34:1935–44.
- In 1387
   Issa
   Issa
- <sup>1391</sup> [145] Herr HW, Dalbagni G. Defining bacillus Calmette-Guerin refractory
   <sup>1392</sup> superficial bladder tumors. J Urol 2003;169:1706–8.
- <sup>1393</sup> [146] Sylvester RJ, van der Meijden A, Witjes JA, et al. High-grade Ta
   <sup>1394</sup> urothelial carcinoma and carcinoma in situ of the bladder. Urology
   <sup>1395</sup> 2005;66:90–107.
- Intravesical gemcitabine for non-muscle invasive bladder cancer. Cochrane Database Syst Rev 2012;2012:CD009294.
- Racioppi M, Di Gianfrancesco L, Ragonese M, Palermo G, Sacco E, Bassi PF. Electromotive drug administration (EMDA) of mitomycin
   C as first-line salvage therapy in high risk "BCG failure" non muscle invasive bladder cancer: 3 years follow-up outcomes. BMC Cancer
   2018;18:1224.
- 1402 [149] Tan WS, Panchal A, Buckley L, et al. Radiofrequency-induced thermo 1403 chemotherapy effect versus a second course of bacillus Calmette 1404 Guerin or institutional standard in patients with recurrence of non 1405 muscle-invasive bladder cancer following induction or maintenance
   1406 bacillus Calmette-Guerin Therapy (HYMN): a phase III, open-label,
   1407 randomised controlled trial. Eur Urol 2019;75:63–71.
- In the second sec
- <sup>1412</sup> [151] Wright KM. FDA approves pembrolizumab for BCG-unresponsive
   <sup>1413</sup> NMIBC. Oncology 2020;34:44.
- Id14 [152] Boorjian SA, Alemozaffar M, Konety BR, et al. Intravesical nadofaragene firadenovec gene therapy for BCG-unresponsive nonmuscle-invasive bladder cancer: a single-arm, open-label, repeat-dose clinical trial. Lancet Oncol 2021;22:107–17.
- <sup>1418</sup> [153] Gallagher BL, Joudi FN, Maymi JL, O'Donnell MA. Impact of previous bacille Calmette-Guerin failure pattern on subsequent response to bacille Calmette-Guerin plus interferon intravesical therapy. Urology 2008;71:297–301.

- [154] Fritsche HM, Burger M, Svatek RS, et al. Characteristics and outcomes of patients with clinical T1 grade 3 urothelial carcinoma treated with radical cystectomy: results from an international cohort. Eur Urol 2010;57:300–9.
   [155] Turker D. Rostrom DI. Waredawski ML et al. Unstaging of urothelial
- [155] Turker P, Bostrom PJ, Wroclawski ML, et al. Upstaging of urothelial cancer at the time of radical cystectomy: factors associated with upstaging and its effect on outcome. BJU Int 2012;110:804–11.
   <sup>1426</sup>
- [156] Moschini M, Sharma V, Dell'oglio P, et al. Comparing longterm outcomes of primary and progressive carcinoma invading bladder muscle after radical cystectomy. BJU Int 2016;117:604–10.
   [157] Willie D, Formander MI, Diekstein PL et al. Clinical outcomes of [1433]
- [157] Willis DL, Fernandez MI, Dickstein RJ, et al. Clinical outcomes of cT1 micropapillary bladder cancer. J Urol 2015;193:1129–34.

1434

1435

1436

1437

1438

1439

1440

1441

- [158] Raj GV, Herr H, Serio AM, et al. Treatment paradigm shift may improve survival of patients with high risk superficial bladder cancer. J Urol 2007;177:1283–6.
- [159] Gofrit ON, Pode D, Lazar A, Katz R, Shapiro A. Watchful waiting policy in recurrent Ta G1 bladder tumors. Eur Urol 2006;49:303–6.
- [160] Herr HW, Donat SM, Reuter VE. Management of low grade papillary bladder tumors. J Urol 2007;178:1201–5.
- [161] Hurle R, Lazzeri M, Vanni E, et al. Active surveillance for low risk nonmuscle invasive bladder cancer: a confirmatory and resource consumption study from the BIAS project. J Urol 2018;199:401–6.
   [142] 1443
- [162] Takenaka A, Yamada Y, Miyake H, Hara I, Fujisawa M. Clinical
   outcomes of bacillus Calmette-Guerin instillation therapy
   for carcinoma in situ of urinary bladder. Int J Urol 2008;15:309–13.
- [163] Mariappan P, Smith G. A surveillance schedule for G1Ta bladder
   cancer allowing efficient use of check cystoscopy and safe discharge at 5 years based on a 25-year prospective database. J Urol 2005;173:1108–11.
- [164] Soukup V, Babjuk M, Bellmunt J, et al. Follow-up after surgical treatment of bladder cancer: a critical analysis of the literature.
   Eur Urol 2012;62:290–302.
- [165] Holmang S, Strock V. Should follow-up cystoscopy in bacillus
   Calmette-Guerin-treated patients continue after five tumour-free years? Eur Urol 2012;61:503–7.
   <sup>1456</sup> 1457
- [166] Kavalieris L, O'Sullivan P, Frampton C, et al. Performance characteristics of a multigene urine biomarker test for monitoring for<br/>recurrent urothelial carcinoma in a multicenter study. J Urol<br/>2017;197:1419–26.1458<br/>1459<br/>1460
- [167] Niwa N, Matsumoto K, Hayakawa N, et al. Comparison of outcomes1462between ultrasonography and cystoscopy in the surveillance of1463patients with initially diagnosed TaG1-2 bladder cancers: a1464matched-pair analysis. Urol Oncol 2015;33:386.e15-e.1465