# UNIVERSITY<sup>OF</sup> BIRMINGHAM University of Birmingham Research at Birmingham

# The sex gap in bladder cancer survival

Toren, Paul; Wilkins, Anna; Patel, Keval; Burley, Amy; Gris, Typhaine; Kockelbergh, Roger; Lodhi, Taha; Choudhury, Ananya; Bryan, Richard T

DOI: 10.1038/s41585-023-00806-2

*License:* Other (please specify with Rights Statement)

Document Version Peer reviewed version

Citation for published version (Harvard):

Toren, P, Wilkins, A, Patel, K, Burley, Á, Gris, T, Kockelbergh, R, Lodhi, T, Choudhury, A & Bryan, RT 2023, 'The sex gap in bladder cancer survival: a missing link in bladder cancer care?', *Nature reviews. Urology.* https://doi.org/10.1038/s41585-023-00806-2

Link to publication on Research at Birmingham portal

#### Publisher Rights Statement:

This version of the article has been accepted for publication, after peer review (when applicable) and is subject to Springer Nature's AM terms of use, but is not the Version of Record and does not reflect post-acceptance improvements, or any corrections. The Version of Record is available online at: https://doi.org/10.1038/s41585-023-00806-2

#### **General rights**

Unless a licence is specified above, all rights (including copyright and moral rights) in this document are retained by the authors and/or the copyright holders. The express permission of the copyright holder must be obtained for any use of this material other than for purposes permitted by law.

•Users may freely distribute the URL that is used to identify this publication.

Users may download and/or print one copy of the publication from the University of Birmingham research portal for the purpose of private study or non-commercial research.
User may use extracts from the document in line with the concept of 'fair dealing' under the Copyright, Designs and Patents Act 1988 (?)

•Users may not further distribute the material nor use it for the purposes of commercial gain.

Where a licence is displayed above, please note the terms and conditions of the licence govern your use of this document.

When citing, please reference the published version.

#### Take down policy

While the University of Birmingham exercises care and attention in making items available there are rare occasions when an item has been uploaded in error or has been deemed to be commercially or otherwise sensitive.

If you believe that this is the case for this document, please contact UBIRA@lists.bham.ac.uk providing details and we will remove access to the work immediately and investigate.

# 1 THE SEX GAP IN BLADDER CANCER SURVIVAL - A MISSING LINK IN BLADDER CANCER

## 2 **CARE?**

- 3 Paul Toren<sup>1</sup>, Anna Wilkins<sup>2,3</sup>, Keval Patel<sup>4</sup>, Amy Burley<sup>2</sup>, Typhaine Gris<sup>1</sup>, Roger Kockelbergh<sup>5,6</sup>, Taha
- 4 Lodhi<sup>7</sup>, Ananya Choudhury<sup>7</sup>, Richard T Bryan<sup>6,8</sup>.
- 5

## 6 Affiliations:

- 7 1. CHU de Québec-Université Laval, Quebec City, Canada.
- 8 2. Division of Radiotherapy and Imaging, The Institute of Cancer Research, London, UK.
- 9 3. The Royal Marsden Hospitals NHS Trust, London, UK.
- 10 4. University Hospitals Birmingham NHS Foundation Trust, Birmingham, UK.
- 11 5. University Hospitals of Leicester NHS Trust, Leicester, UK.
- 12 6. Action Bladder Cancer UK, Tetbury, UK.
- Division of Cancer Sciences, University of Manchester and The Christie NHS Foundation Trust,
   Manchester, UK.
- Bladder Cancer Research Centre, Institute of Cancer & Genomic Sciences, University of
   Birmingham, UK.
- 17

## 18 Correspondence:

- 19 Professor Richard T Bryan
- 20 Bladder Cancer Research Centre, Institute of Cancer & Genomic Sciences, University of Birmingham,
- 21 Edgbaston, Birmingham B15 2TT, UK.

## 22 <u>r.t.bryan@bham.ac.uk</u>

23

## 24 ORCIDs

- 25 Paul Toren: 0000-0002-5762-5787
- 26 Anna Wilkins: 0000-0002-0425-584X
- 27 Keval Patel: 0000-0003-1913-0852
- 28 Amy Burley: 0000-0003-3074-6439
- 29 Typhaine Gris: TBC
- 30 Roger Kockelbergh: 0000-0003-2261-3628
- 31 Taha Lodhi: 0000-0002-6891-0866
- 32 Ananya Choudhury: 0000-0002-3561-6580
- 33 Richard Bryan: 0000-0003-2853-4293.

#### 35 ABSTRACT

36 Recent data again bring to light again differences in bladder cancer outcomes between sexes. 37 Uncommon among cancers, bladder cancer outcomes are notably worse for women than for men. 38 Furthermore, bladder cancer is 3-4 times more common amongst men. In this perspective, we review 39 the current understanding of factors which may explain these sex differences. These include understanding the importance of haematuria as a symptom of bladder cancer by both clinicians and 40 41 patients, the resultant delays in diagnosis and referral of female patients with haematuria, and 42 healthcare access. Notably, these factors appear to have geographical variation and are not consistent 43 across all healthcare systems. Likewise, there are inconsistent data relating to sex-specific treatment 44 responses for both NMIBC and MIBC patients. The impact of differences in the microbiome, bladder 45 wall thickness and urine dwell times remain to be elucidated. The interplay of hormone signalling, 46 gene expression, immunology and tumour microenvironment remains complex but likely underpins 47 the sexual dimorphism in disease incidence and stage and histology at presentation. The contribution 48 of these biological phenomena to sex-specific outcome differences is probable, albeit potentially 49 treatment-specific, and further understanding is required. Notwithstanding, we identify opportunities to harness biological differences to improve treatment outcomes, as well as areas of fundamental and 50 51 translational research to pursue. At the level of policy and healthcare delivery, improvements can be 52 made across the domains of patient awareness, clinician education, referral pathways, and guideline-53 based care. Together, we aim to highlight opportunities to close the sex gap in bladder cancer 54 outcomes.

55

#### 57 **INTRODUCTION**

Bladder cancer (BC) is the tenth most common cancer worldwide, responsible for 3% of annual cancer
diagnoses and 2.1% of cancer-related deaths<sup>1</sup>. Most patients (75-80%) present with non-muscleinvasive bladder cancer (NMIBC: stages Ta/T1/Tis) – up to 80% of these patients will experience
recurrence<sup>2</sup>, and up to 44% will progress to muscle-invasive bladder cancer (MIBC: stages T2-4)<sup>2-4</sup>.
Moreover, of the 20-25% of patients initially diagnosed with MIBC, around one-quarter will have
incurable, locally-advanced or metastatic disease<sup>5</sup>. Muscle-invasion thus represents a critical step in
the disease course, carrying a 5-year survival of only 27-50%, despite radical therapies<sup>3</sup>.

Overall, the number of incident BC cases continues to increase, although the age-standardized incidence and mortality appear to be decreasing, paralleled by fewer smoking-related cases<sup>6</sup>. Smoking is the largest modifiable risk factor for BC, with exposure to occupational carcinogens also well documented<sup>7</sup>. However, sex is the largest risk factor for BC, with men 3-4 times more likely to be diagnosed with BC than women<sup>8,9</sup>. Furthermore, there is no clear indication of the differential effects of smoking between men and women<sup>10</sup>. Nonetheless, women tend to have more aggressive tumours at diagnosis and experience worse outcomes thereafter<sup>11-15</sup>.

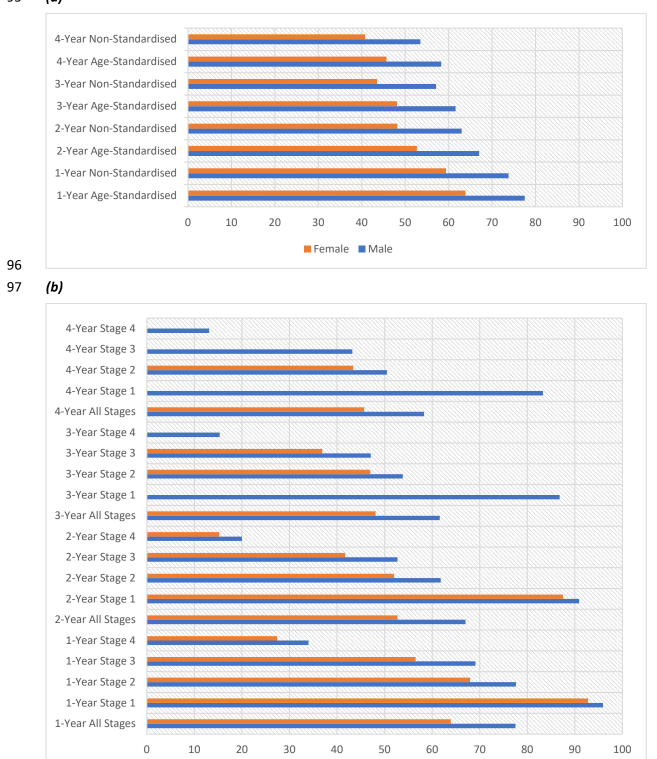
BC cases which are muscle-invasive or metastatic at presentation contribute the most significant morbidity and mortality, while the surveillance of NMIBC drives BC to be one of the most expensive per patient cancers to treat<sup>16</sup>. Therefore, there is a need to identify and fill the sex-based gaps in our understanding and care for all BC patients. In this perspective, we review the differences in outcomes between men and women with BC, discuss our current understanding of these differences, and propose solutions to pursue. These differences are significant given the impact of BC on patients and healthcare systems.

79

#### 80 SEX DIFFERENCES IN OUTCOMES

Recent data regarding cancer survival in England (cancers diagnosed from 2015 to 2019, followed up 81 82 to 2020<sup>17</sup>) appear to lay bare the persistent sex gap in survival for BC patients. Across all stages of BC and for 1-, 2-, 3- and 4-year BC survival, female patients had worse outcomes than male patients 83 84 (Figure 1). With some 3- and 4-year stage-specific survival data missing for female patients (only), a 85 knowledge deficit may already be apparent. Notwithstanding, using an overlapping dataset available for T1–T4 tumours treated from 2013 to 2019, recent analyses of overall survival for urothelial cancer 86 patients in England by Catto et al also illustrate a similar situation<sup>18</sup>. Beyond the UK, such gaps in 87 outcomes continue to be highlighted internationally<sup>19-23</sup>. 88

Figure 1: 1-year, 2-year, 3-year, and 4-year bladder cancer survival (%) for adults (aged 15 to 99 years)
diagnosed in England 2015 to 2019, followed up to 2020<sup>17</sup>. (a) Age-standardised and non-age
standardised net survival (%) by sex. (b) Age-standardised net survival (%) by stage and sex. NB: 3Year Stage 1, 3-Year Stage 4, 4-Year Stage 1, 4-Year Stage 3 and 4-Year Stage 4 data are all missing
for female patients. Age-standardisation represents a weighted-average of mortality rates for each
sex based on the International Classification of Survival Standard.

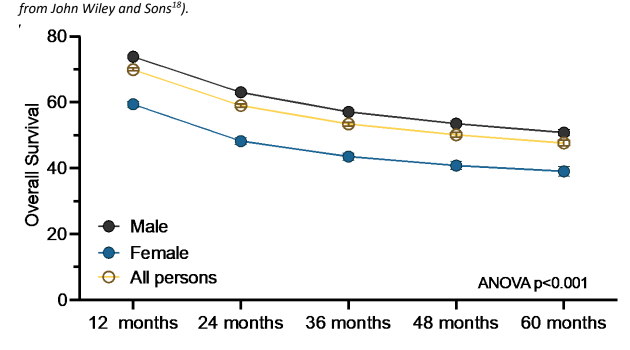


Bladder Female

Bladder Male

95 **(a)** 

Figure 2: Overall survival for patients with T1–T4 bladder cancers treated in England from 2013 to
 2019, plotted using the Kaplan-Meier method with respect to sex (reproduced with kind permission
 from John Wiley and Sons<sup>18</sup>).





102

104 This is neither a new phenomenon, nor an unexplored phenomenon<sup>19,22</sup>, yet it remains unexplained worldwide. Moreover, these findings in BC are the inverse of the situation for most cancer sites where 105 106 cancer outcomes are commonly worse for men<sup>24</sup>. Previous European data from 1999-2007 also reported lower BC survival among women, driven by differences in Northern and Central European 107 countries and the UK<sup>25</sup>. More recent data from the Netherlands also mirrors the UK data<sup>26</sup>. However, 108 other studies have found no differences in treatment or cancer-specific survival<sup>27</sup>. A 2022 systematic 109 review and meta-analysis concluded that female sex was associated with worse cancer-specific 110 111 survival and overall survival in patients with MIBC, but no differences were apparent in the relatively lower number of studies with sex-specific outcomes for NMIBC<sup>28</sup>. Concordant with the recent NHS 112 113 results, this suggests that sex differences in survival are principally driven by patients with non-114 metastatic MIBC at presentation, or those who progress to develop MIBC.

115

## 116 POSSIBLE REASONS FOR DIFFERENCES IN OUTCOMES

The possible explanations are complex – the interplay of the signs and symptoms of the disease, speed of referral into the BC diagnostic pathway, treatment decisions, treatment responses, and biology<sup>29</sup>. It is important to better understand the weight and relevance of each of these factors in order to take concrete steps to further investigate and potentially close the apparent gap in overall outcomes.

#### 121 Recognising the importance of haematuria

Female patients may not recognise the importance of haematuria in a disease where there is a 3:1 male preponderance or may not seek investigation as early<sup>21</sup>. Haematuria is the most common symptom of bladder cancer<sup>2</sup>. However, from a young age, women are much more likely to present with a UTI and have associated haematuria in some instances. This can desensitize both patients and clinicians to not recognize the importance of haematuria, particularly in older women at higher risk of bladder cancer<sup>30</sup>.

128

### 129 Referral for investigations

130 Once haematuria has been identified as a potential symptom of bladder cancer, there is a need for 131 prompt referral according to guidelines. The preponderance of haematuria related to urinary tract 132 infection in women can result in delays in diagnosis – as reported internationally, female patients presenting with haematuria to their primary care provider are not referred for subsequent 133 investigations as rapidly as male patients or are investigated less thoroughly<sup>21,29-34</sup>. Indeed, 134 135 Lyratzopoulos et al. found that women had a significantly higher number of pre-referral consultations 136 than men when presenting with haematuria, with 27% of women requiring 3 or more consultations compared with only 11% of men requiring the same<sup>31</sup>. 137

138

#### 139 Treatment decisions

140 In the setting of NMIBC, sex does not appear to influence the utilisation of adjuvant intravesical therapy<sup>35</sup>, nor the choice of radical treatment for MIBC<sup>36-38</sup> and, in terms of overall treatment 141 142 paradigms, there are currently no differing recommendations between sexes. Although the use of continent urinary diversions appears to be lower in female patients than male patients<sup>39,40</sup>, this does 143 144 not alter survival outcomes. This may reflect higher rates of advanced tumour stage in women, but 145 also potential differences in training and practice patterns, with fewer centres worldwide equally 146 comfortable to perform orthotopic continent diversion in women given anatomical and surgical 147 differences pertinent to differences in tumour stage. However, for female patients undergoing radical cystectomy and bladder substitution, preservation of the uterus and attempted nerve-sparing appear 148 to result in better functional outcomes<sup>41</sup>, yet there remain significant gaps in the adoption of female 149 150 reproductive organ-sparing and nerve-sparing radical cystectomy techniques for patients with organconfined disease<sup>42</sup>. With emerging data from large series confirming the low rate of female 151 reproductive organ involvement at the time of radical cystectomy (4.2-5.7%)<sup>43,44</sup>, there appear to be 152 153 potential sex disparities driven by provider expertise and preference. These disparities may need to 154 be addressed through training and/or the refinement or centralisation of specific provider expertise.

#### 155 Treatment efficacy

156 Evaluating whether female patients derive less benefit from current treatments is more difficult to 157 dissect and may be, in part, driven by biological and anatomical reasons elucidated in the section below. Female patients nonetheless present with worse disease stage<sup>9,33,38,45</sup>, and more often with 158 non-urothelial tumour histology<sup>9,20,46</sup>, thus contextualising subsequent sex-specific treatment 159 160 responses. Studying 24,169 BC patients in The Netherlands, Richters et al. found that, in the first two years after diagnosis, excess mortality rates for women were higher than for men but lower 161 162 thereafter<sup>9</sup>. This applied to both NMIBC and MIBC patients, and baseline differences in age, stage, and 163 histology accounted for only part of the excess mortality gap<sup>9</sup>. Ballas et al. reported similar findings 164 for patients undergoing bladder preservation (trimodality therapy) for T2-T4a N0 M0 MIBC<sup>47</sup>. 165 Marinaro et al. studied 47,229 MIBC patients in the USA and identified increased 90-day mortality 166 following radical cystectomy and worse overall survival in female patients<sup>37</sup>. Others have also highlighted either worse cancer-specific survival in females than males following radical 167 cystectomy<sup>36,45</sup>, or both worse cancer specific and overall survival<sup>11,48</sup>. Notwithstanding, other 168 169 research suggests that higher uptake of neoadjuvant chemotherapy diminishes these sex 170 differences<sup>49</sup>. Such data may indicate that if sex differences in outcomes are attenuated in patients fit 171 enough for chemotherapy then, simply by selection bias, similar outcomes could also be expected in 172 the generally fitter patients enrolled into clinical trials. In the radiotherapy setting, unpublished data 173 from Manchester, UK, regarding 209 MIBC patients treated by radiotherapy with concurrent carbogen and nicotinamide (BCON protocol<sup>50</sup>) show no difference in 5-year cancer-specific survival between 174 175 males and females.

176 In addition to overall survival data, conflicting data report on potential sex differences in response to 177 specific treatments for BC<sup>51</sup>. Evaluating response to intravesical treatment for NMIBC, earlier reports (summarised in a 2018 meta-analysis and systematic review<sup>52</sup>) suggest women have poorer responses 178 179 to BCG. However, there appears to be publication bias in this meta-analysis toward studies reporting 180 a sex-based difference and, thus, there is doubt as to these conclusions suggesting differential responses in NMIBC to BCG<sup>52</sup>. In line with this, two large contemporary cohorts of BCG-treated NMIBC 181 182 patients did not report any sex differences in outcomes of recurrence-free survival or progression-free 183 survival<sup>53,54</sup>. Furthermore, the literature suggests no sex differences in response to intravesical 184 chemotherapy for NMIBC.

For advanced BC, the accumulating evidence does not suggest that sex-based differences exist in disease response to immune checkpoint blockade (ICB); this includes the sex-based analyses presented in recent trials, including KEYNOTE-052<sup>55</sup>, KEYNOTE-361<sup>56</sup>, and IMvigor130<sup>57</sup>. Moreover, the best available data do not suggest sex-specific differences in response to chemotherapy, with Haines *et al.* concluding that "female patients with metastatic urothelial cancer tolerate cisplatin-based chemotherapy similarly to male patients and achieve comparable clinical outcomes"<sup>58</sup>. Biological tumour differences and stage differences in presentation confound these studies, which together suggest that, stage-for-stage, treatment responses remain generally similar between men and women. Hence, there may be regional or treatment-specific circumstances where sex does not appear to be a prognostic factor.

195

#### 196 Biological and anatomical phenomena

Few studies have thoroughly assessed the fundamental sex differences in urothelial transformation and subsequent cancer biology – the interplay of sex hormones, environmental exposures, microenvironment, microbiome, immunology, and genomics are important and highly complex<sup>19,22</sup>. Notwithstanding, the laboratory studies available to date provide some insights into potential drivers of biological differences in bladder tumours which may drive differences in outcomes between men and women.

#### 203 Sex hormones

Knockout studies in mice suggest that the androgen receptor (AR) in the urothelium is important for 204 urothelial carcinogenesis<sup>59,60</sup>. Further, estrogen receptor alpha (ERα) and estrogen receptor beta (ERβ) 205 206 knockout mice experiments suggest that  $ER\alpha$  has a protective role against BC initiation and progression, while ERβ could promote BC<sup>61-63</sup>. These murine results are consistent with 207 208 immunohistochemical studies demonstrating downregulated ERa expression in higher grade and higher stages bladder tumors<sup>64</sup> and upregulated ERβ and aromatase protein expression in higher grade 209 and more aggressive bladder tumors<sup>64-67</sup>. Using the "four cores genotypes" mouse model, which 210 211 decouples chromosomal and gonadal sex, researchers found gonadal sex (i.e. hormonal exposure) had the single largest impact on chemically-induced bladder tumour development<sup>68</sup>; notwithstanding, 212 213 chromosomal sex also independently influenced tumour development<sup>68</sup>.

Sex steroids have direct effects on the activity and function of various subsets of innate and adaptive immune cells and are known to contribute to immunological differences between sexes. The role of systemic hormones in the incidence and progression of bladder tumours remains incompletely defined, particularly in women. Epidemiologic studies of BC among women suggest a later age at menopause, parity (vs nulliparity) and use of hormone replacement therapy (HRT) may be associated with decreased BC incidence<sup>69-73</sup>. However, a meta-analysis suggested that the age of menarche does not affect the risk of BC in women<sup>74</sup>, and so a need for clarification remains.

#### 222 <u>Genomics</u>

The genomic and molecular understanding of BC has advanced in recent years<sup>75-79</sup>. MIBCs are heterogeneous<sup>80</sup> and characterised by many single nucleotide variants (SNVs) and copy number variants (CNVs)<sup>75,81,82</sup>; loss of multiple tumour suppressors and alteration of multiple pathways are common<sup>77,78</sup>. Six consensus gene expression-based subtypes of MIBC are now recognised and share some characteristics<sup>77</sup>, but which remain heterogeneous with respect to genomic aberrations and behaviour; temporal and spatial plasticity in subtype has also been reported<sup>83</sup>. See **Table 1** below. NMIBC is arguably more complex than MIBC<sup>78,79</sup>, comprising multiple grades of disease<sup>84</sup>.

Within this landscape, BCs (alongside other cancers) demonstrate extensive sex-biased molecular 230 signatures, with sex-biased expressed genes enriched in the sex chromosomes and evidence of sex-231 biased DNA methylation patterns (e.g. TOP2B)<sup>85</sup>. Sex-biased pathways include those related to 232 immune responses, apoptosis and the cell cycle, metabolism, DNA repair and P53 pathways<sup>85</sup>. 233 Furthermore, *KDM6A* alterations are common in BCs (24-33%<sup>75,79</sup>) with the gene functioning as an 234 235 epigenetic regulator of downstream gene expression; importantly, KDM6A escapes X chromosome inactivation<sup>19</sup>. Mouse models appear to demonstrate that loss of *Kdm6a* increases BC risk in female 236 237 mice, and mutations or reduced expression of human KDM6A predicts poor prognosis in female BC patients<sup>68</sup>. Other research has suggested a higher rate of *KDM6A* mutations in NMIBCs from female 238 239 patients<sup>86</sup>.

Class name	Luminal Papillary (LumP)	Luminal Non- Specified (LumNS)	Luminal Unstable (LumU)	Stroma- rich	Basal/Squamous (Ba/Sq)	Neuroendocrino like (NE-like)
% of MIBC	24%	8%	15%	15%	35%	3%
Oncogenic mechanisms	FGFR3+	PPARG+	PPARG+		EGFR+	TP53-
	PPARG+		E2F3+			RB1-
	CDKN2A-		ERBB2+			Cell cycle+
			Genomic instability			
			Cell cycle+			
Mutations	FGFR3 (40%)	<i>ELF3</i> (35%)	TP53 (76%)		<i>TP53</i> (61%)	TP53 (94%)
			ERCC2		<i>RB1</i> (25%)	RB1 (39%)
	<i>KDM6A</i> (38%)		(22%)			
			TMB+ APOBEC+			
Clinical			AFUBLCT			
characteristics	T2 stage +	Older patients +			Women +	
		(80+)			T3/T4 stage +	
Median overall survival (years)	4.0	1.8	2.9	3.8	1.2	1.0

# Table 1: Summary of the main characteristics of the consensus classes of MIBC (adapted from<sup>77</sup>).

subtypes<sup>88</sup>. However, an analysis of NMIBC did not identify differences in molecular subtypes
 according to sex<sup>89</sup>.

#### 255 Tumour immunology & microenvironment

The wide repertoire of immunomodulatory agents used across stages for treatment of BC, including in clinical trials, means that the contribution of sex-specific immunological and microenvironmental factors is likely to be clinically important. Such factors can include genetic, epigenetic and transcriptional effects<sup>85</sup>, which may relate to XX and XY chromosomal differences<sup>90</sup>, as well as hormonal effects<sup>91</sup>. Several pre-clinical and clinical studies highlight these factors and provide some explanation for the divergent observations of increased incidence of BC in male populations yet inferior survival outcomes in women described earlier.

263 Studies of bladder tumour AR expression do not show sex-based differences in expression<sup>92</sup>. However, 264 Kwon et al. have recently described a fascinating androgen-driven mechanism of T cell exhaustion in 265 BC<sup>91</sup>. Their insights provide some explanation for why spontaneous rejection of early immunogenic 266 bladder tumours is less common in males, and hence the male predisposition for the development of 267 BC. The study evaluates three different bladder tumour models, MB49 (transplantable syngeneic 268 tumours), BBN (carcinogen-induced tumours) and BKL171 where BBN-induced tumours develop in a 269 testis-bearing mouse with an XX chromosome to eliminate any immune response to male-specific 270 minor antigens. The authors demonstrate more aggressive tumour growth in male versus female mice with these three models<sup>91</sup>. This effect was eliminated using Tcrb/Tcrd or *RAG2* knockout mice, which 271 272 specifically lack T cells, and reinstated with adoptive transfer of CD8<sup>+</sup> T cells. A two-fold higher 273 frequency of polyfunctional CD8<sup>+</sup> T cells able to produce Interferon gamma (IFN<sub>Y</sub>), Tumour Necrosis 274 Factor Alpha (TNF $\alpha$ ) and Granzyme B (Gzmb) was seen in MB49 tumours of female versus male mice 275 at day 9 of tumour growth.

276 In support of the above observations, single-cell RNA sequencing identified increased effector-like 277 CD8<sup>+</sup> T cells in the female versus male tumour microenvironment (TME). In contrast, the male TME 278 was enriched for progenitor exhausted CD8<sup>+</sup> tumour infiltrating lymphocytes (TILs), as defined by their 279 stem-like genetic profile (i.e. Tcf1/Tcf7<sup>+</sup>). These TILs showed accelerated progression to terminally 280 differentiated Tcf1-Tim3<sup>+</sup> exhausted T cells incapable of restimulation. In keeping with all of the above 281 results, male mice with loss of AR exclusively in CD8<sup>+</sup> T cells were equally protected against cancer as 282 female mice. Finally, the authors observe a negative correlation between type I interferon signalling 283 and AR activity in T cells and they suggest this balance may underlie wider sexual dimorphism in cancer 284 immunity. Insights from the above study are corroborated by observations elsewhere that androgen deprivation therapy can promote responses to ICB<sup>92,93</sup>. The work by Kwon *et al.* is CD8<sup>+</sup> T cell specific; 285 286 CD4<sup>+</sup> T cells are known to be important in BC immunology<sup>94</sup>, and there are likely further sex-specific 287 mechanistic insights to T cell biology to be uncovered in the future.

288 Studies to explore a biologic basis for different responses to BC therapy have yielded interesting 289 results. An analysis of the whole transcriptomes of 460 tumours from the UROMOL cohort, together 290 with multiplex immunofluorescence of tumours from the Kingston Health Services Centre (KHSC) cohort (n=332, 22% female)<sup>95</sup> reported increased expression of the immune checkpoint genes CTLA4, 291 292 PDCD1, LAG3, and ICOS in high-grade tumours from females compared to high-grade tumours from 293 males or low-grade tumours from either sex. In addition, increased expression of CXC ligand 13 294 (CXCL13, an important B-cell–recruiting chemokine) and the B-cell surface-associated molecule CD40 295 were seen more frequently in high-grade tumours from female patients. Intriguingly, the authors also 296 observed increased infiltration of CD163<sup>+</sup> M2-like tumour-associated macrophages (TAMs) in both low- and high-grade NMIBC tumours of female patients versus those of male patients<sup>95</sup>. Of further 297 298 clinical relevance, a higher density of CD163<sup>+</sup> M2-like TAMs and CD79a<sup>+</sup> B cells was independently 299 associated with shorter recurrence-free survival across all high-grade tumours in the KHSC cohort 300 supporting a functional relevance of this sexually dimorphic observation. Together, these findings 301 suggest that the TME of NMIBC from female patients tends towards greater immune exhaustion 302 where immune dysfunction is accentuated by reciprocal communication between increased 303 immunosuppressive macrophage and B cell populations.

304 Elsewhere, research findings indicate that estrogen inhibits interleukin-6 and thus decreases the 305 expression of receptor complexes required for BCG adherence to urothelial cells, such as integrin 306  $\alpha$ 5 $\beta$ 1<sup>96,97</sup>. A therapeutic strategy combining the anti-estrogen therapy ICI 182780 with BCG was shown 307 to improve treatment efficacy in *in vitro* and *in vivo* pre-clinical systems, in part via enhancing TNF $\alpha$ 308 signalling<sup>98</sup>. Furthermore, the immune checkpoint ICOS shows greater upregulation in whole blood 309 following stimulation of healthy female volunteers versus their male counterparts with BCG<sup>99</sup>. 310 However, these findings require careful interpretation in view of the considerable immunological 311 differences between post-menopausal women and younger healthy volunteers. Overall, a current key 312 research gap is to understand the sex-specific longitudinal innate and adaptive immune changes that 313 occur both intra-tumourally and systemically over the months to years following BCG treatment, including how these relate to a differential therapeutic response. 314

Immunological ageing or immunosenesence is known to have sex-specific biological characteristics<sup>100</sup>. A recent study using bulk RNA sequencing showed an enrichment for B cell function-associated pathways in aged healthy female mice bladders versus their aged male counterparts<sup>101</sup>. Multiplex immunofluorescence confirmed a greater number of organised tertiary lymphoid structures (TLS) in the healthy bladders of female mice. Somewhat surprisingly, in murine bladders treated with several weeks of the carcinogen BBN, there was no difference in TLS between male and female mice. Instead, an

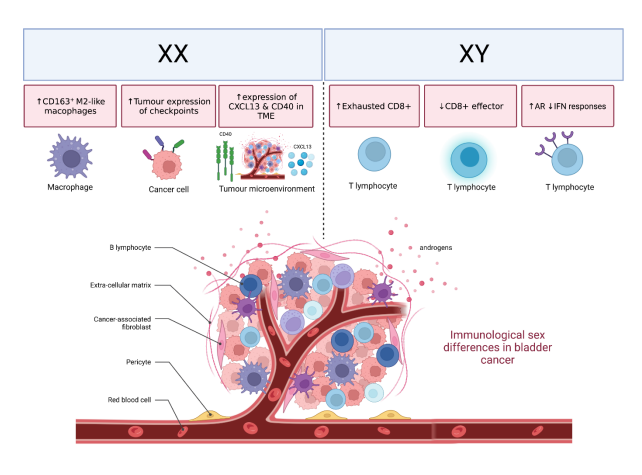
- 321 increase in plasma cells was seen in the lamina propria of female aged mice, and female mice had a more
- 322 immune-infiltrated and oedematous lamina propria across ages.

As outlined earlier, a difference in the response to ICB according to sex has not been clearly established in BC. However, we know that immune cell PD-L1 is associated with inferior survival outcomes across sexes, and that androgen signalling in T cells represses IFNγ to limit ICB responses<sup>93</sup>. Furthermore, estrogen plus a number of X-linked micro-RNAs, including miR-221, miR-222, and miR-106b, can regulate PD-L1 expression<sup>90,102</sup>. Our incomplete understanding of the sex-specific immunogenomic changes underlying differential responses to ICB urgently warrants further research to optimise novel combination strategies across disease stages. See **Figure 3**.

330

**Figure 3:** A summary of immunological sex differences in BC: The female TME is characterised by increased infiltration of immunosuppressive CD163+ M2-like macrophages and increased tumour expression of immune checkpoints. In addition, an increase in genes related to B cell recruitment (CXCL13) and function (CD40) has been shown in the female TME. In contrast, the male TME is characterised by increased exhausted CD8+ T cells (both of progenitor and terminally differentiated subtypes), which is driven by AR signalling specifically in CD8+ T cells. **AR**: androgen receptor, **IFN**: interferon, **TME**: tumour microenvironment.

338



339

340

#### 342 <u>Microbiome</u>

Current research seeks to define whether differences in the urinary, tumoral or gut microbiome between men and women may contribute to differential outcomes<sup>103</sup>. While urinary microbiome differences may be associated with the risk of recurrent urinary tract infections (UTIs) contributing to carcinogenesis, the causal implication of potential microbiome differences in tumour progression or treatment response remains to be defined<sup>22</sup>.

#### 348 Anatomy

Anatomically, men have more outlet obstruction related to prostatic enlargement and subsequent detrusor hypertrophy. Women have thinner bladder walls<sup>104</sup>, which may help explain a higher incidence of non-organ-confined disease at diagnosis<sup>45</sup>. Differential urinary dwell times in men and women may contribute to differences in BC development, with men more commonly having higher post-void residuals with age. While conceptually sound, experimental validation of these differences remains lacking and difficult to undertake.

355

#### 356 POSSIBLE STRATEGIES TO MITIGATE THE SEX GAP

It is widely recognised that, despite little difference in treatment patterns or quality measures, female 357 sex is associated with worse overall survival among individuals with MIBC<sup>19</sup>. However, although the 358 359 UK data illustrated above highlight a sex gap in outcomes that is most apparent for stages 2 and above 360 (>pT2), such differences are not uniformly reported internationally. Differences in healthcare system 361 access, cancer registry reporting, and treatment availability may explain the disparity between 362 regions; publication bias in the available literature may also contribute to some discrepancies<sup>105</sup>. Given the differences in immunobiology outlined above, it is perhaps surprising that the reported sex 363 364 differences in outcomes across both NMIBC and MIBC patients remain inconsistent and ambiguous, 365 in contrast to the unequivocal dichotomy in incidence. See **Table 2**. Future initiatives should focus on root causes for sex-specific differences in pathological staging and features at diagnosis<sup>46</sup>, as well as 366 prospectively collecting relevant data<sup>18</sup> – the avoidance of stage migration subsequent to symptom 367 ignorance or delayed referral to secondary care is fundamental<sup>106</sup>. 368

369

## 371 Table 2: Bladder cancer associated phenomena that demonstrate confirmed differences between

## 372 the sexes.

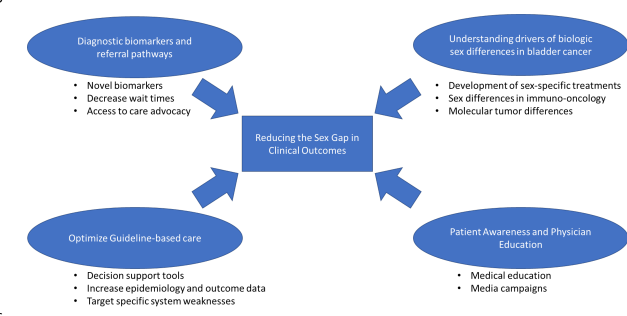
	Male	Female			
Incidence <sup>1</sup>	9.6	2.4			
Age standardized rate (ASR)					
per 100,000 persons per year					
Referral <sup>21,29-34</sup>	Prompt referral	Delayed referral frequent.			
	commonplace.				
Treatment	No differing recommendations between sexes. Differential				
	responses potentially treat	ment- and territory-specific.			
Sex hormones <sup>59,60,64-67,69-74</sup>	Androgens may promote	Differential roles of ERα and			
	carcinogenesis via AR.	ERβ, equivocal role of			
		estrogens.			
Genomics <sup>68,85,86</sup>	Maintained KDM6A expression may be protective (most				
	relevant in females). Methyla	ation differences predominate.			
Histopathology &	For MIBC, urothelial cancer	For MIBC, increased			
molecular subtypes <sup>46,87,88</sup>	predominates, with higher	frequency of basal subtype			
	proportion of luminal	and squamous carcinoma.			
	papillary and neuro-				
	endocrine-like subtypes.				
Immunology &	Evidence of AR-driven T cell	In NMIBC, immune			
microenvironment <sup>91,95</sup>	exhaustion.	exhaustion may result from			
		increased			
		immunosuppressive			
		macrophage & B cell			
		populations.			

373

374

In recent years, increasing adoption of guideline-recommended treatments (e.g. neoadjuvant chemotherapy, trimodality therapy, etc.) may be attenuating such outcome differences between sexes. Notwithstanding, there is a general acceptance across the BC research field that much more work needs to be done to better understand the differences in disease incidence between males and females, and the seemingly worse outcomes for female MIBC patients<sup>19,28</sup>. See **Figure 4**.

- 381 Figure 4: An overview of different approaches to decrease the sex gap in bladder cancer outcomes. Each
- domain and sub-domain likely require adequately-powered prospective studies in order to validate and
   implement strategies that can reduce the sex gap in clinical outcomes.
- 384
- 385



386 387

# 388 Awareness of diagnostic differences between sexes

389 To avoid diagnostic and referral delays, clinicians need to be aware of the differential presentations 390 of BC between sexes. Similar situations are present in other diseases. For example, symptoms experienced during acute coronary syndrome (ACS) differ between the sexes, with a higher proportion 391 392 of women in those patients presenting without typical chest pain<sup>107</sup>. This leads to improved detection 393 in men and worse comparative survival in women presenting with ACS; specific awareness drives have been funded in the UK to reduce this difference. For BC, raising awareness of haematuria as a sign of 394 bladder cancer are also important. Previous work on symptom awareness (such as haematuria<sup>108</sup>) has 395 396 suggested that mass media campaigns combined with targeted higher-intensity community-based 397 programmes for high-risk populations (low socio-economic status, older age, and specific racial groups) can effect change<sup>109</sup>. Campaigns specifically aimed at women may be required. 398

Overall, the importance of haematuria in both sexes should be underlined in primary care where most patients initially present. Primary care physician education drives through presentations at conferences, decision support tools, and practice-specific presentations have been shown to improve referral rates for dementia<sup>110</sup> and their use for early diagnosis of BC should be investigated.

- 403
- 404

#### 405 Addressing clinical & healthcare system factors

406 Addressing health system, access and referral issues leading to stage migration and sex differences in 407 outcomes remains important. Compared to biological differences, problems such as delay in diagnosis 408 and suboptimal treatment are readily modifiable factors to improve outcomes. However, it is 409 important to understand the severity of these problems may vary by jurisdiction. For example, in an 410 American National Cancer Database study, while women with MIBC had poorer overall survival, this difference did not appear to be related to measures of treatment quality, with data suggesting lower 411 rates of treatment delay in women<sup>46</sup>. Notwithstanding, the BC diagnostic and treatment pathway is 412 413 prolonged for all patients and strategies to reduce delays are urgently needed for all patients<sup>111,112</sup>. Accurate diagnostic urinary biomarkers<sup>113,114</sup> may facilitate timely urologic evaluation, and 414 deployment in primary care may be particularly useful for the initial assessment of female haematuria 415 416 patients.

417 Evidently, the relative contribution of delays in diagnosis and evaluation to differences in clinical outcomes is complex and difficult to dissect<sup>115,116</sup>. Although stratified by stage, the data in **Figure 1b** 418 419 do not take account of the difference in outcomes between stages T1a and T1b, T2a and T2b, and T3a 420 and T3b, to which these factors will contribute. Similarly, the extent to which significantly greater sex 421 differences in stage are related to biological differences, access to healthcare, or delays in diagnosis 422 cannot easily be determined with the available data. Socio-economic factors impacting access to care may drive some of the differences<sup>116</sup>. For example, it is striking that differences in the rate of MIBC at 423 424 diagnosis between sexes according to SEER data is greater among African-Americans (30% males vs 425 43% females) versus Caucasians (22% vs 25%). In addition to potential biological drivers of these race 426 differences, access to care is likely a major contributor based on similar data for other cancers<sup>117</sup>. 427 Social barriers which limit access to care and timely referral are thus important to address. BC 428 disproportionately affects patients with lower socio-economic status and research from Canada suggests that this gap has broadened, particularly among women<sup>118</sup>. Similar research elsewhere can 429 430 help to highlight regional and national deficiencies which can stimulate policy and funding changes at 431 a larger level.

As part of healthcare delivery, decision support tools (DSTs) may be useful due to their ability to be embedded within practice electronic medical systems so that they are easily accessible during consultation<sup>119</sup>. They can be automated to draw in background information on smoking status or family history to prompt clinicians to refer for investigation of a certain condition. In an evaluation of a 7-point checklist DST for the assessment of pigmented skin lesions, primary care physicians found such tools easy to use and particularly useful for borderline decision-making<sup>120</sup>. However, widespread use of DSTs relies on the levels of trust placed in the tools, compatibility of the DSTs with specific 439 electronic care systems, and difficulty in usage<sup>119</sup>. Furthermore, the ability of DSTs to affect change in 440 cancer survival is uncertain and requires ongoing investigation. In the context of haematuria, models 441 have been identified that could be used in primary care to guide referrals, with the potential to identify 442 lower-risk patients with visible haematuria and to stratify individuals who present with non-visible haematuria<sup>121</sup>. A recent systematic review identified 13 such models with good discrimination for the 443 444 diagnosis of bladder or kidney cancer (area under the receiver operating curve, AUROC >0.8), although only 8 had been externally validated; all of the studies had either high or unclear risk of bias<sup>121</sup>. The 445 446 authors concluded that external validations in appropriate populations were required before 447 implementation in primary care<sup>121</sup>.

448

### 449 **Optimising guideline-based care**

The establishment and popularisation of guidelines for the referral of women with haematuria are vital to promote standards of care for referral to cystoscopy. Accounting for different age-adjusted cancer risks, some referral guidelines for haematuria differ between men and women<sup>122</sup>. For example, the American Urological Association guidelines give greater weight to non-visible haematuria in men versus women during the 5<sup>th</sup> and 6<sup>th</sup> decade of life<sup>123</sup>. However, the relative paucity of research data to establish these recommendations and the variability of current recommendations between jurisdictions contributes to potential confusion and uncertainty among primary care providers<sup>30</sup>.

457 Should the standard of care for female patients with urothelial carcinoma treatment differ from male 458 patients? This remains a challenging question, particularly as treatment patterns and practice evolves over time. Some historical data suggest that response to treatment may be inferior for women<sup>11,15,52</sup>. 459 460 This includes a meta-analysis suggesting that female sex is associated with poorer cancer-specific 461 survival and inferior responses to BCG. For NMIBC, when evaluated critically, we believe the most reliable and recent data do not support the notion that treatment guidelines should differ between 462 463 men and women. In metastatic urothelial cancer, men and women appear to have similar treatment outcomes across various studies<sup>124</sup>. Therefore, despite the poorer outcomes reported for female 464 patients with advanced BC, it is not clear that alternate standards of care are warranted. 465

466 Overall, accurate and reliable data are essential to identify the weak points in patients' journeys for 467 which improvements in care can translate into better outcomes. This can be applied at both the health 468 system and the hospital level. Equality (and diversity) in recruitment to clinical trials is also essential, 469 with appropriate instigation of meta-analyses where data gaps exist.

470

#### 472 Harnessing biological differences to improve treatment

473 A meaningful assessment of the fundamental sex differences in urothelial transformation and 474 subsequent cancer biology is challenging as it transects the complex interplay of sex hormones, 475 environmental exposures, microenvironment, immunology, genetics and genomics, and the 476 microbiome. Nonetheless, this research is critical to identify specific differences which can be 477 translated to clinical care – a call to action which has recently been emphasised<sup>19</sup>. However, in an era 478 where molecular classification, personalised medicines and targeted therapies are endlessly sought, 479 this 'simple' knowledge may remain years away – with approximately 1 in 4 patients female, many 480 studies are underpowered to answer the question of whether a certain biomarker or classifier or 481 treatment is effective or not in female patients.

482 To address some of these challenges, it is important to design studies accordingly<sup>19</sup>. For example, in pre-clinical studies, attention is needed to account for the sex of origin of cell lines and to include 483 484 studies in both male and female mice. In clinical trials, reporting of results should include sex-based 485 analyses. Furthermore, evaluating prognostic and predictive biomarkers for prognosis and response 486 to therapy according to sex is necessary in correlative analyses of clinical trials. Notably, the 487 recruitment of women into clinical trials is an important pre-requisite for these analyses as this has 488 been a historical challenge. For instance, in a series of well-known NMIBC trials in Spain, only 11% of 489 all recruited patients were women<sup>125</sup>.

490 Biological differences may present unique opportunities to tailor treatment according to sex. 491 Preclinical studies suggest that alterations in sex steroids with AR antagonism may represent a 492 strategy to treat male bladder tumours alone or in combination with immunotherapy. Based on initial 493 pre-clinical studies, AR antagonism may alter immune responsiveness to immunotherapy through alterations in the steroid milieu<sup>92</sup>. Hormonal differences are broadly recognized to impact the immune 494 response<sup>126</sup>, and AR suppressive therapy may improve BC outcomes through a hormonal-mediated 495 496 modulation of the TME<sup>127</sup>. Furthermore, there is now a significant amount of clinical data to suggest 497 that 5-alpha reductase inhibitors (5ARIs) could decrease the incidence and recurrence rates of low-498 grade BC in men. With a 2021 meta-analysis suggesting a HR of 0.46 for recurrence for male NMIBC 499 patients receiving 5ARI therapy, prospective clinical trials are warranted. However, these should be 500 adequately powered to avoid false-negatives from short-term evaluation of a long-acting mechanism 501 of action<sup>128</sup>.

502

503

#### 505 **Future directions**

Research and advocacy are important to address the sex gaps in clinical care. Improved data collection on the natural history and epidemiologic difference in haematuria between sexes can drive the development of best practices and referral pathways<sup>18</sup>. Similarly, understanding the gaps in referral can facilitate targeted campaigns to raise awareness in both the populations at risk and their healthcare providers.

At a more fundamental level, there exist many opportunities to expand our understanding of how sex differences impact the TME of BC and the immune interactions which contribute to anti-cancer activity and response to immune-targeted treatments. Understanding immune differences at both the level of the urothelial and systemic interactions may facilitate more effective and personalised therapies. Clinical research is now emerging to evaluate whether sex-specific treatments are effective, such as

suppression of the androgen axis in combination with existing treatments (e.g. the 'BicaBCa' study,
NCT05327647) in men. Further studies to target sex-specific strategies to decrease recurrences of
NMIBC or progression of MIBC are similarly warranted given existing data<sup>127</sup>, and increased reporting
of sex-based analyses in clinical trials is to be welcomed. Importantly, improved awareness of the sex
gap is also apparent, as illustrated by a number of presentations at this year's American Society of
Clinical Oncology Genitourinary cancers symposium<sup>129-131</sup>.

522

#### 523 CONCLUSIONS

In conclusion, addressing the sex gap in BC outcomes requires coordinated efforts to improve outcomes for women. A greater understanding of the sexual dimorphism of BC biology and immunology may permit personalised, sex-specific biomarkers. There is also a need to develop sexspecific treatments through clinical trials, such as treatments targeting the androgen axis. In parallel, it is important to recognise that implementation of best practices for referral, diagnosis and treatment of BC can provide rapid improvements in outcomes where deficiencies exist. Further research is also needed to identify optimal strategies for the referral and evaluation of haematuria between sexes.

531

533		Reference List
534		
535	1	Bray, F. et al. Global cancer statistics 2018: GLOBOCAN estimates of incidence and mortality
536	-	worldwide for 36 cancers in 185 countries. <i>CA Cancer J Clin</i> <b>68</b> , 394-424,
537		doi:10.3322/caac.21492 (2018).
538	2	Babjuk, M. <i>et al.</i> European Association of Urology Guidelines on Non-muscle-invasive Bladder
539	2	Cancer (Ta, T1, and Carcinoma in Situ). <i>Eur Urol</i> , doi:10.1016/j.eururo.2021.08.010 (2021).
540	3	Witjes, J. A. <i>et al.</i> European Association of Urology Guidelines on Muscle-invasive and
541	5	Metastatic Bladder Cancer: Summary of the 2020 Guidelines. Eur Urol <b>79</b> , 82-104,
542		doi:10.1016/j.eururo.2020.03.055 (2021).
543	4	Sylvester, R. J. et al. European Association of Urology (EAU) Prognostic Factor Risk Groups for
544		Non-muscle-invasive Bladder Cancer (NMIBC) Incorporating the WHO 2004/2016 and WHO
545		1973 Classification Systems for Grade: An Update from the EAU NMIBC Guidelines Panel. Eur
546		<i>Urol</i> <b>79</b> , 480-488, doi:10.1016/j.eururo.2020.12.033 (2021).
547	5	Herbert, A. et al. Population trends in emergency cancer diagnoses: The role of changing
548		patient case-mix. <i>Cancer Epidemiol</i> <b>63</b> , 101574, doi:10.1016/j.canep.2019.101574 (2019).
549	6	Teoh, J. Y. et al. Global Trends of Bladder Cancer Incidence and Mortality, and Their
550		Associations with Tobacco Use and Gross Domestic Product Per Capita. Eur Urol <b>78</b> , 893-906,
551		doi:10.1016/j.eururo.2020.09.006 (2020).
552	7	Cumberbatch, M. G. K. et al. Epidemiology of Bladder Cancer: A Systematic Review and
553		Contemporary Update of Risk Factors in 2018. Eur Urol <b>74</b> , 784-795,
554		doi:10.1016/j.eururo.2018.09.001 (2018).
555	8	Bryan, R. T. <i>et al.</i> A comparison of patient and tumour characteristics in two UK bladder cancer
556		cohorts separated by 20 years. <i>BJU. Int</i> <b>112</b> , 169-175, doi:10.1111/bju.12032 [doi] (2013).
557	9	Richters, A. <i>et al.</i> Bladder cancer survival: Women only fare worse in the first two years after
558		diagnosis. Urol Oncol <b>37</b> , 853-861, doi:10.1016/j.urolonc.2019.08.001 (2019).
559	10	Freedman, N. D., Silverman, D. T., Hollenbeck, A. R., Schatzkin, A. & Abnet, C. C. Association
560		between smoking and risk of bladder cancer among men and women. JAMA <b>306</b> , 737-745,
561		doi:306/7/737 [pii];10.1001/jama.2011.1142 [doi] (2011).
562	11	Uhlig, A. <i>et al.</i> Gender Specific Differences in Disease-Free, Cancer Specific and Overall Survival
563		after Radical Cystectomy for Bladder Cancer: A Systematic Review and Meta-Analysis. J Urol
564		<b>200</b> , 48-60, doi:10.1016/j.juro.2017.11.150 (2018).
565	12	Wang, S. C. <i>et al.</i> The gender difference and mortality-to-incidence ratio relate to health care
566		disparities in bladder cancer: National estimates from 33 countries. Sci Rep 7, 4360,
567		doi:10.1038/s41598-017-04083-z (2017).
568	13	Mungan, N. A. et al. Gender differences in stage-adjusted bladder cancer survival. Urology 55,
569		876-880, doi:S0090429500005239 [pii] (2000).
570	14	Kluth, L. A. <i>et al.</i> Gender-specific differences in clinicopathologic outcomes following radical
571		cystectomy: an international multi-institutional study of more than 8000 patients. Eur Urol 66,
572		913-919, doi:10.1016/j.eururo.2013.11.040 (2014).
573	15	Liu, S. et al. The impact of female gender on bladder cancer-specific death risk after radical
574		cystectomy: a meta-analysis of 27,912 patients. Int Urol Nephrol 47, 951-958,
575		doi:10.1007/s11255-015-0980-6 (2015).
576	16	Mossanen, M. & Gore, J. L. The burden of bladder cancer care: direct and indirect costs. Curr
577		<i>Opin Urol</i> <b>24</b> , 487-491, doi:10.1097/MOU.0000000000000078 (2014).
578	17	Cancer Survival in England, cancers diagnosed 2015 to 2019, followed up to 2020, 2022).
579	18	Catto, J. W. F. et al. Diagnosis, treatment and survival from bladder, upper urinary tract, and
580		urethral cancers: real-world findings from NHS England between 2013 and 2019. BJU Int,
581		doi:10.1111/bju.15970 (2023).
582	19	Theodorescu, D., Li, Z. & Li, X. Sex differences in bladder cancer: emerging data and call to
583		action. Nat Rev Urol 19, 447-449, doi:10.1038/s41585-022-00591-4 (2022).

- 58420Sadighian, M. & Porten, S. Gender differences in oncologic and functional outcomes in<br/>patients with bladder cancer undergoing radical cystectomy with urinary diversion. *Curr Opin*<br/>Urol **29**, 542-547, doi:10.1097/MOU.0000000000660 (2019).
- 58721Burge, F. & Kockelbergh, R. Closing the Gender Gap: Can We Improve Bladder Cancer Survival588in Women? A Systematic Review of Diagnosis, Treatment and Outcomes. Urol Int **97**, 373-589379, doi:10.1159/000449256 (2016).
- 59022Koti, M. et al. Sex Differences in Bladder Cancer Immunobiology and Outcomes: A591Collaborative Review with Implications for Treatment. Eur Urol Oncol 3, 622-630,592doi:10.1016/j.euo.2020.08.013 (2020).
- 59323Shu, T. D. *et al.* Disparities in cause-specific mortality by race and sex among bladder cancer594patients from the SEER database. *Cancer Causes Control*, doi:10.1007/s10552-023-01679-x595(2023).
- 59624Dong, M. et al. Sex Differences in Cancer Incidence and Survival: A Pan-Cancer Analysis. Cancer597Epidemiol Biomarkers Prev 29, 1389-1397, doi:10.1158/1055-9965.EPI-20-0036 (2020).
- 59825Marcos-Gragera, R. *et al.* Urinary tract cancer survival in Europe 1999-2007: Results of the599population-based study EUROCARE-5. *Eur. J. Cancer*, doi:S0959-8049(15)00708-X600[pii];10.1016/j.ejca.2015.07.028 [doi] (2015).
- Richters, A., Leliveld, A. M., Goossens-Laan, C. A., Aben, K. K. H. & Ozdemir, B. C. Sex
  differences in treatment patterns for non-advanced muscle-invasive bladder cancer: a
  descriptive analysis of 3484 patients of the Netherlands Cancer Registry. *World J Urol* 40,
  2275-2281, doi:10.1007/s00345-022-04080-6 (2022).
- 60527Blindheim, A. *et al.* T1 bladder cancer in Norway: treatment and survival. *Scand J Urol* 54, 370-606375, doi:10.1080/21681805.2020.1803401 (2020).
- 60728Mori, K. *et al.* Impact of sex on outcomes after surgery for non-muscle-invasive and muscle-608invasive bladder urothelial carcinoma: a systematic review and meta-analysis. World J Urol,609doi:10.1007/s00345-022-04116-x (2022).
- 61029Dobruch, J. *et al.* Gender and Bladder Cancer: A Collaborative Review of Etiology, Biology, and611Outcomes. *Eur Urol* **69**, 300-310, doi:10.1016/j.eururo.2015.08.037 (2016).
- Solution Structure
  Solution Structure</l
- Lyratzopoulos, G., Abel, G. A., McPhail, S., Neal, R. D. & Rubin, G. P. Gender inequalities in the
  promptness of diagnosis of bladder and renal cancer after symptomatic presentation:
  evidence from secondary analysis of an English primary care audit survey. *BMJ Open* 3,
  doi:bmjopen-2013-002861 [pii];10.1136/bmjopen-2013-002861 [doi] (2013).
- Ngo, B., Perera, M., Papa, N., Bolton, D. & Sengupta, S. Factors affecting the timeliness and adequacy of haematuria assessment in bladder cancer: a systematic review. *BJU Int* 119 Suppl 5, 10-18, doi:10.1111/bju.13821 (2017).
- Bryan, R. T. *et al.* A Comparative Analysis of the Influence of Gender, Pathway Delays, and Risk
  Factor Exposures on the Long-term Outcomes of Bladder Cancer. *Eur. Urol. Focus* 1, 82-89,
  doi:S2405-4569(15)00007-3 [pii];10.1016/j.euf.2015.01.001 [doi] (2015).
- 34 Zhou, Y. *et al.* Prolonged Diagnostic Intervals as Marker of Missed Diagnostic Opportunities in
  Bladder and Kidney Cancer Patients with Alarm Features: A Longitudinal Linked Data Study. *Cancers (Basel)* 13, doi:10.3390/cancers13010156 (2021).
- Banforth, K. N. *et al.* Care Quality and Variability in the Use of Intravesical Therapy for Initial
  Treatment of Nonmuscle Invasive Bladder Cancer Within a Large, Diverse Integrated Delivery
  System. *Urology* 131, 93-103, doi:10.1016/j.urology.2019.03.035 (2019).
- 63136Grajales, V. et al. Associations Between Female Sex and Treatment Patterns and Outcomes for632Muscle-invasive Bladder Cancer. Urology 151, 169-175, doi:10.1016/j.urology.2020.06.058633(2021).

- 63437Marinaro, J. *et al.* Sex and Racial Disparities in the Treatment and Outcomes of Muscle-635invasive Bladder Cancer. Urology 151, 154-162, doi:10.1016/j.urology.2020.06.087 (2021).
- 636 38 Heberling, U. *et al.* Gender and Mortality after Radical Cystectomy: Competing Risk Analysis.
  637 Urol Int **101**, 293-299, doi:10.1159/000487445 (2018).
- 63839Farber, N. J. *et al.* Disparities in the Use of Continent Urinary Diversions after Radical639Cystectomy for Bladder Cancer. *Bladder Cancer* 4, 113-120, doi:10.3233/BLC-170162 (2018).
- Bachour, K. *et al.* Trends in urinary diversion after radical cystectomy for urothelial carcinoma.
  World J Urol **36**, 409-416, doi:10.1007/s00345-017-2169-3 (2018).
- 642 41 Gross, T., Meierhans Ruf, S. D., Meissner, C., Ochsner, K. & Studer, U. E. Orthotopic ileal
  643 bladder substitution in women: factors influencing urinary incontinence and hypercontinence.
  644 *Eur Urol* 68, 664-671, doi:10.1016/j.eururo.2015.05.015 (2015).
- Gupta, N. *et al.* Practice Patterns Regarding Female Reproductive Organ-Sparing and NerveSparing Radical Cystectomy Among Urologic Oncologists in the United States. *Clin Genitourin Cancer*, doi:10.1016/j.clgc.2023.01.010 (2023).
- 648 43 Bree, K. K. et al. Contemporary Rates of Gynecologic Organ Involvement in Females with 649 Muscle Invasive Bladder Cancer: A Retrospective Review of Women Undergoing Radical 650 Cystectomy following Neoadjuvant Chemotherapy. J Urol 206, 577-585, 651 doi:10.1097/JU.000000000001784 (2021).
- 65244Lobo, N. *et al.* Is It Safe To Spare Gynaecological Organs in Female Patients Undergoing Radical653Cystectomy? A Multi-Institutional Study of Three Tertiary Pelvic Cancer Centres. Journal of654Clinical Urology 15, 94, doi:10.1177/20514158221077479 (2022).
- 65545Rosiello, G. *et al.* The effect of sex on disease stage and survival after radical cystectomy: a656population-based analysis.*UrolOncol***39**, 236e231-236e237,657doi:10.1016/j.urolonc.2020.09.004 (2021).
- Krimphove, M. J. *et al.* Sex-specific Differences in the Quality of Treatment of Muscle-invasive
  Bladder Cancer Do Not Explain the Overall Survival Discrepancy. *Eur Urol Focus* 7, 124-131,
  doi:10.1016/j.euf.2019.06.001 (2021).
- 661 47 Ballas, L. K. *et al.* Disparities in male versus female oncologic outcomes following bladder
  662 preservation: A population-based cohort study. *Cancer Med* 10, 3004-3012,
  663 doi:10.1002/cam4.3835 (2021).
- Williams, S. B. *et al.* Survival differences among patients with bladder cancer according to sex:
  Critical evaluation of radical cystectomy use and delay to treatment. *Urol Oncol* 35, 602 e601602 e609, doi:10.1016/j.urolonc.2017.05.022 (2017).
- 667 49 Venkat, S. et al. Does neoadjuvant chemotherapy diminish the sex disparity in bladder cancer 668 survival after radical cystectomy? Urol Oncol 40, 106 e121-106 e129, doi:10.1016/j.urolonc.2021.09.003 (2022). 669
- Hoskin, P. J., Rojas, A. M., Bentzen, S. M. & Saunders, M. I. Radiotherapy with concurrent
  carbogen and nicotinamide in bladder carcinoma. *J Clin Oncol* 28, 4912-4918,
  doi:10.1200/JCO.2010.28.4950 (2010).
- bacillus Calmette-Guerin immunotherapy for T1G3/HG bladder cancer. *World J Urol* 39, 33373344, doi:10.1007/s00345-021-03653-1 (2021).
- 676 52 Uhlig, A. *et al.* Gender-specific Differences in Recurrence of Non-muscle-invasive Bladder
  677 Cancer: A Systematic Review and Meta-analysis. *Eur Urol Focus* 4, 924-936,
  678 doi:10.1016/j.euf.2017.08.007 (2018).
- 67953Bree, K. K. et al. Impact of sex on response to BCG in non-muscle invasive bladder cancer680patients: a contemporary review from a tertiary care center. World J Urol **39**, 4143-4149,681doi:10.1007/s00345-021-03755-w (2021).
- 68254Fadel, J. *et al.* Analysis of sex-based differences to Bacillus Calmette-Guerin for non-muscle683invasive bladder cancer. *Urol Oncol*, doi:10.1016/j.urolonc.2022.09.024 (2022).

- Balar, A. V. *et al.* Efficacy and safety of pembrolizumab in metastatic urothelial carcinoma:
  results from KEYNOTE-045 and KEYNOTE-052 after up to 5 years of follow-up. *Ann Oncol* 34,
  289-299, doi:10.1016/j.annonc.2022.11.012 (2023).
- 68756Powles, T. *et al.* Pembrolizumab alone or combined with chemotherapy versus chemotherapy688as first-line therapy for advanced urothelial carcinoma (KEYNOTE-361): a randomised, open-689label, phase 3 trial. *Lancet Oncol* **22**, 931-945, doi:10.1016/S1470-2045(21)00152-2 (2021).
- 690 57 Galsky, M. D. *et al.* Atezolizumab with or without chemotherapy in metastatic urothelial
  691 cancer (IMvigor130): a multicentre, randomised, placebo-controlled phase 3 trial. *Lancet* 395,
  692 1547-1557, doi:10.1016/S0140-6736(20)30230-0 (2020).
- 69358Haines, L. et al. The impact of gender on outcomes in patients with metastatic urothelial694carcinoma. Clin Genitourin Cancer 11, 346-352, doi:10.1016/j.clgc.2013.04.010 (2013).
- Hsu, J. W. *et al.* Decreased tumorigenesis and mortality from bladder cancer in mice lacking
  urothelial androgen receptor. *Am J Pathol* 182, 1811-1820, doi:10.1016/j.ajpath.2013.01.018
  (2013).
- 69860Miyamoto, H. *et al.* Promotion of bladder cancer development and progression by androgen699receptor signals. J. Natl. Cancer Inst **99**, 558-568, doi:99/7/558 [pii];10.1093/jnci/djk113 [doi]700(2007).
- Hsu, I., Vitkus, S., Da, J. & Yeh, S. Role of oestrogen receptors in bladder cancer development.
   *Nat Rev Urol* 10, 317-326, doi:10.1038/nrurol.2013.53 (2013).
- 70362Hsu, I. *et al.* Estrogen receptor alpha prevents bladder cancer via INPP4B inhibited akt pathway704in vitro and in vivo. Oncotarget 5, 7917-7935, doi:10.18632/oncotarget.1421 (2014).
- 70563Hsu, I. *et al.* Suppression of ERbeta signaling via ERbeta knockout or antagonist protects706against bladder cancer development. *Carcinogenesis* **35**, 651-661, doi:10.1093/carcin/bgt348707(2014).
- 70864Miyamoto, H. *et al.* Expression of androgen and oestrogen receptors and its prognostic709significance in urothelial neoplasm of the urinary bladder. *BJU Int* **109**, 1716-1726,710doi:10.1111/j.1464-410X.2011.10706.x (2012).
- Nguyen, D. P. *et al.* Association of Aromatase With Bladder Cancer Stage and Long-Term
  Survival: New Insights Into the Hormonal Paradigm in Bladder Cancer. *Clin Genitourin Cancer* **15**, 256-262 e251, doi:10.1016/j.clgc.2016.05.017 (2017).
- 71466Shen, S. S. *et al.* Expression of estrogen receptors-alpha and -beta in bladder cancer cell lines715and human bladder tumor tissue. *Cancer* **106**, 2610-2616, doi:10.1002/cncr.21945 (2006).
- 716 67 Tuygun, C. *et al.* Sex-specific hormone receptors in urothelial carcinomas of the human urinary
  717 bladder: a comparative analysis of clinicopathological features and survival outcomes
  718 according to receptor expression. *Urol Oncol* 29, 43-51, doi:10.1016/j.urolonc.2009.01.033
  719 (2011).
- 72068Kaneko, S. & Li, X. X chromosome protects against bladder cancer in females via a KDM6A-721dependent epigenetic mechanism. Sci Adv 4, eaar5598, doi:10.1126/sciadv.aar5598 (2018).
- 72269Huang, A. T. *et al.* Bladder cancer and reproductive factors among women in Spain. *Cancer*723*Causes Control* **20**, 1907-1913, doi:10.1007/s10552-009-9384-1 (2009).
- 70 Bai, Y. *et al.* Parity and bladder cancer risk: a dose-response meta-analysis. *BMC Cancer* 17, 31, doi:10.1186/s12885-016-3023-5 (2017).
- 726 71 Cantwell, M. M., Lacey, J. V., Jr., Schairer, C., Schatzkin, A. & Michaud, D. S. Reproductive factors, exogenous hormone use and bladder cancer risk in a prospective study. *Int J Cancer* 728 **119**, 2398-2401, doi:10.1002/ijc.22175 (2006).
- 72972Daugherty, S. E. *et al.* Reproductive factors and menopausal hormone therapy and bladder730cancer risk in the NIH-AARP Diet and Health Study. *Int J Cancer* **133**, 462-472,731doi:10.1002/ijc.28022 (2013).
- 73 73 McGrath, M., Michaud, D. S. & De Vivo, I. Hormonal and reproductive factors and the risk of
  733 bladder cancer in women. *Am J Epidemiol* 163, 236-244, doi:10.1093/aje/kwj028 (2006).

- 74 Li, Y. D., Gao, L., Gou, Y. Q., Tan, W. & Liu, C. Age of menarche and primary bladder cancer
  risk: A meta-analysis and systematic review. *Urol Oncol* 40, 346 e317-346 e326,
  doi:10.1016/j.urolonc.2022.022 (2022).
- 737 75 Comprehensive molecular characterization of urothelial bladder carcinoma. *Nature* 507, 315738 322, doi:nature12965 [pii];10.1038/nature12965 [doi] (2014).
- 739 76 Van, B. J. *et al.* Bladder cancers arise from distinct urothelial sub-populations. *Nat. Cell Biol*740 16, 982-991, doi:ncb3038 [pii];10.1038/ncb3038 [doi] (2014).
- 741 77 Kamoun, A. *et al.* A Consensus Molecular Classification of Muscle-invasive Bladder Cancer. *Eur* 742 *Urol* 77, 420-433, doi:10.1016/j.eururo.2019.09.006 (2020).
- 743 78 Lindskrog, S. V. *et al.* An integrated multi-omics analysis identifies prognostic molecular
  744 subtypes of non-muscle-invasive bladder cancer. *Nat Commun* 12, 2301, doi:10.1038/s41467745 021-22465-w (2021).
- 746 79 Goel, A. *et al.* Combined exome and transcriptome sequencing of non-muscle-invasive bladder
   747 cancer: associations between genomic changes, expression subtypes, and clinical outcomes.
   748 *Genome Med* 14, 59, doi:10.1186/s13073-022-01056-4 (2022).
- 74980Meeks, J. J. *et al.* Genomic heterogeneity in bladder cancer: challenges and possible solutions750to improve outcomes. Nat Rev Urol **17**, 259-270, doi:10.1038/s41585-020-0304-1 (2020).
- Kiu, X. *et al.* Highly prevalent TERT promoter mutations in bladder cancer and glioblastoma.
   *Cell Cycle* 12, 1637-1638 (2013).
- 75382Robertson, A. *et al.* Comprehensive Molecular Characterization of Muscle-Invasive Bladder754Cancer. *Cell* **171**, 540-556 (2017).
- 75583Sfakianos, J. *et al.* Epithelial plasticity can generate multi-lineage phenotypes in human and756murine bladder cancers. *Nat Commun.* **11**, 2540 doi: 2510.1038/s41467-41020-16162 (2020).
- Soukup, V. *et al.* Prognostic Performance and Reproducibility of the 1973 and 2004/2016
  World Health Organization Grading Classification Systems in Non-muscle-invasive Bladder
  Cancer: A European Association of Urology Non-muscle Invasive Bladder Cancer Guidelines
  Panel Systematic Review. *Eur Urol* 72, 801-813, doi:10.1016/j.eururo.2017.04.015 (2017).
- 76185Yuan, Y. *et al.* Comprehensive Characterization of Molecular Differences in Cancer between762Male and Female Patients. *Cancer Cell* **29**, 711-722, doi:10.1016/j.ccell.2016.04.001 (2016).
- Response of Non-invasive Bladder Cancer with Distinct Metabolic
  Profile and Female Gender Bias in KDM6A Mutation Frequency. *Cancer Cell* 32, 701-715 e707,
  doi:10.1016/j.ccell.2017.08.005 (2017).
- Ball, K. S., Hounsome, L., Verne, J. & Kockelbergh, R. Non-transitional cell carcinoma only 766 87 767 partly explains adverse survival outcomes in females with T1–T4 bladder cancer: A summary 768 of UK epidemiological data. Journal of Clinical Urology 10, 14-18, 769 doi:10.1177/2051415816679529 (2017).
- de Jong, J. J. *et al.* Distribution of Molecular Subtypes in Muscle-invasive Bladder Cancer Is
  Driven by Sex-specific Differences. *Eur Urol Oncol* 3, 420-423, doi:10.1016/j.euo.2020.02.010
  (2020).
- 773
   89
   Hedegaard, J. et al. Comprehensive Transcriptional Analysis of Early-Stage Urothelial

   774
   Carcinoma.
   Cancer
   Cell
   **30**,
   27-42,
   doi:S1535-6108(16)30209-4

   775
   [pii];10.1016/j.ccell.2016.05.004 [doi] (2016).
- Care, A. *et al.* Sex disparity in cancer: roles of microRNAs and related functional players. *Cell Death Differ* 25, 477-485, doi:10.1038/s41418-017-0051-x (2018).
- 77891Kwon, H. *et al.* Androgen conspires with the CD8(+) T cell exhaustion program and contributes779to sex bias in cancer. *Sci Immunol* **7**, eabq2630, doi:10.1126/sciimmunol.abq2630 (2022).
- 780 92 Toren, P. *et al.* Androgen receptor and immune cell PD-L1 expression in bladder tumors
  781 predicts disease recurrence and survival. *World J Urol* **39**, 1549-1558, doi:10.1007/s00345782 020-03358-x (2021).
- Guan, X. *et al.* Androgen receptor activity in T cells limits checkpoint blockade efficacy. *Nature* **606**, 791-796, doi:10.1038/s41586-022-04522-6 (2022).

- 785
   94
   Oh, D. Y. *et al.* Intratumoral CD4(+) T Cells Mediate Anti-tumor Cytotoxicity in Human Bladder

   786
   Cancer. *Cell* **181**, 1612-1625 e1613, doi:10.1016/j.cell.2020.05.017 (2020).
- 787 95 Chenard, S. *et al.* Sexual Dimorphism in Outcomes of Non-muscle-invasive Bladder Cancer: A
  788 Role of CD163+ Macrophages, B cells, and PD-L1 Immune Checkpoint. *Eur Urol Open Sci* 29,
  789 50-58, doi:10.1016/j.euros.2021.05.002 (2021).
- 790 96 Zhang, G. J. *et al.* Autocrine IL-6 production by human transitional carcinoma cells upregulates
   791 expression of the alpha5beta1 firbonectin receptor. *J Urol* 163, 1553-1559 (2000).
- Guise, A. I., Chen, F., Zhang, G. & See, W. The effects of physiological estrogen concentration
  on the immune response of urothelial carcinoma cells to bacillus Calmette-Guerin. *J Urol* 185,
  298-304, doi:10.1016/j.juro.2010.09.004 (2011).
- 79598Shang, Z. et al. Targeting estrogen/estrogen receptor alpha enhances Bacillus Calmette-796Guerin efficacy in bladder cancer. Oncotarget 7, 27325-27335, doi:10.18632/oncotarget.8756797(2016).
- Piasecka, B. *et al.* Distinctive roles of age, sex, and genetics in shaping transcriptional variation
  of human immune responses to microbial challenges. *Proc Natl Acad Sci U S A* 115, E488-E497,
  doi:10.1073/pnas.1714765115 (2018).
- 801 100 Marquez, E. J. *et al.* Sexual-dimorphism in human immune system aging. *Nat Commun* **11**, 751,
   802 doi:10.1038/s41467-020-14396-9 (2020).
- Hamade, A. *et al.* Sex differences in the aging murine urinary bladder and influence on the tumor immune microenvironment of a carcinogen-induced model of bladder cancer. *Biol Sex Differ* 13, 19, doi:10.1186/s13293-022-00428-0 (2022).
- Yang, L. *et al.* Posttranscriptional Control of PD-L1 Expression by 17beta-Estradiol via PI3K/Akt
  Signaling Pathway in ERalpha-Positive Cancer Cell Lines. *Int J Gynecol Cancer* 27, 196-205,
  doi:10.1097/IGC.00000000000875 (2017).
- 809 103 Pederzoli, F. et al. Sex-specific Alterations in the Urinary and Tissue Microbiome in Therapy-810 Urothelial Bladder Cancer Patients. Eur Urol Oncol 3, 784-788, naive 811 doi:10.1016/j.euo.2020.04.002 (2020).
- 812104Anzia, L. E. *et al.* Comprehensive non-invasive analysis of lower urinary tract anatomy using813MRI. Abdom Radiol (NY) 46, 1670-1676, doi:10.1007/s00261-020-02808-9 (2021).
- Sarrio-Sanz, P. *et al.* Mortality prediction models after radical cystectomy for bladder tumour:
  A systematic review and critical appraisal. *Eur J Clin Invest* 52, e13822, doi:10.1111/eci.13822
  (2022).
- 817106Zhou, Y., Funston, G., Lyratzopoulos, G. & Walter, F. M. Improving the Timely Detection of818Bladder and Kidney Cancer in Primary Care. Adv Ther **36**, 1778-1785, doi:10.1007/s12325-019-81900966-x (2019).
- 820107Haider, A. *et al.* Sex and gender in cardiovascular medicine: presentation and outcomes of821acute coronary syndrome. *Eur Heart J* **41**, 1328-1336, doi:10.1093/eurheartj/ehz898 (2020).
- 822108Merriel, S. W. D. *et al.* A prospective evaluation of the fourth national Be Clear on Cancer823'Blood in Pee' campaign in England. *Eur J Cancer Care (Engl)* **31**, e13606,824doi:10.1111/ecc.13606 (2022).
- 825
   109
   Wakefield, M. A., Loken, B. & Hornik, R. C. Use of mass media campaigns to change health

   826
   behaviour. Lancet **376**, 1261-1271, doi:10.1016/S0140-6736(10)60809-4 (2010).
- 827110Arsenault-Lapierre, G. *et al.* Improving dementia care: insights from audit and feedback in828interdisciplinary primary care sites. *BMC Health Serv Res* 22, 353, doi:10.1186/s12913-022-82907672-5 (2022).
- 830111Russell, B. *et al.* A Systematic Review and Meta-analysis of Delay in Radical Cystectomy and831the Effect on Survival in Bladder Cancer Patients. *Eur Urol Oncol* **3**, 239-249,832doi:10.1016/j.euo.2019.09.008 (2020).
- Bryan, R. T. *et al.* Comparing an Imaging-guided Pathway with the Standard Pathway for
  Staging Muscle-invasive Bladder Cancer: Preliminary Data from the BladderPath Study. *Eur Urol*, doi:10.1016/j.eururo.2021.02.021 (2021).

- Ward, D. G. *et al.* Highly Sensitive and Specific Detection of Bladder Cancer via Targeted Ultradeep Sequencing of Urinary DNA. *Eur Urol Oncol*, doi:10.1016/j.euo.2022.03.005 (2022).
- Humayun-Zakaria, N., Ward, D. G., Arnold, R. & Bryan, R. T. Trends in urine biomarker
  discovery for urothelial bladder cancer: DNA, RNA, or protein? *Translational Andrology and Urology* 10, 2787-2808 (2021).
- 841 115 Wallace, D. M., Bryan, R. T., Dunn, J. A., Begum, G. & Bathers, S. Delay and survival in bladder
  842 cancer. *BJU. Int* 89, 868-878 (2002).
- Russell, B. *et al.* Systematic review of the association between socioeconomic status and bladder cancer survival with hospital type, comorbidities, and treatment delay as mediators. *BJUI Compass* 2, 140-158, doi:10.1002/bco2.65 (2021).
- Esnaola, N. F. & Ford, M. E. Racial differences and disparities in cancer care and outcomes:
  where's the rub? *Surg Oncol Clin N Am* **21**, 417-437, viii, doi:10.1016/j.soc.2012.03.012 (2012).
- 848118Densmore, R., Hajizadeh, M. & Hu, M. Trends in socio-economic inequalities in bladder cancer849incidence in Canada: 1992-2010. Can J Public Health 110, 722-731, doi:10.17269/s41997-019-85000227-y (2019).
- 851119Sutton, R. T. *et al.* An overview of clinical decision support systems: benefits, risks, and852strategies for success. NPJ Digit Med **3**, 17, doi:10.1038/s41746-020-0221-y (2020).
- 853120Walter, F. M. *et al.* Using the 7-point checklist as a diagnostic aid for pigmented skin lesions in854general practice: a diagnostic validation study. *Br J Gen Pract* **63**, e345-353,855doi:10.3399/bjgp13X667213 (2013).
- 856121Harrison, H. *et al.* Risk prediction models for symptomatic patients with bladder and kidney857cancer: a systematic review. *Br J Gen Pract* **72**, e11-e18, doi:10.3399/BJGP.2021.0319 (2022).
- Linder, B. J., Bass, E. J., Mostafid, H. & Boorjian, S. A. Guideline of guidelines: asymptomatic microscopic haematuria. *BJU Int* **121**, 176-183, doi:10.1111/bju.14016 (2018).
- 860
   123
   Barocas, D. A. et al. Microhematuria: AUA/SUFU Guideline. J Urol 204, 778-786,

   861
   doi:10.1097/JU.0000000001297 (2020).
- 862124Sorce, G. *et al.* Survival trends in chemotherapy exposed metastatic bladder cancer patients863and chemotherapy effect across different age, sex, and race/ethnicity. Urol Oncol 40, 380864e319-380 e327, doi:10.1016/j.urolonc.2022.03.014 (2022).
- 865125Fernandez-Gomez, J. et al. Prognostic factors in patients with non-muscle-invasive bladder866cancer treated with bacillus Calmette-Guerin: multivariate analysis of data from four867randomized CUETO trials. Eur Urol 53, 992-1001, doi:10.1016/j.eururo.2007.10.006 (2008).
- 868126Schafer, J. M. *et al.* Sex-biased adaptive immune regulation in cancer development and869therapy. *iScience* **25**, 104717, doi:10.1016/j.isci.2022.104717 (2022).
- 870127Kourbanhoussen, K. *et al.* Switching Cancers: A Systematic Review Assessing the Role of871Androgen Suppressive Therapy in Bladder Cancer. *Eur Urol Focus* 7, 1044-1051,872doi:10.1016/j.euf.2020.10.002 (2021).
- 873128Dekalo, S. *et al.* 5alpha-reductase inhibitors and the risk of bladder cancer in a large,874population-based cohort. *Urol Oncol*, doi:10.1016/j.urolonc.2022.09.004 (2022).
- Niegisch, G. *et al.* Treatment patterns, indicators of receiving systemic treatment, and clinical outcomes in metastatic urothelial carcinoma: A retrospective analysis of real-world data in Germany. *J Clin Oncol* **41** (2023).
- Westhofen, T. *et al.* Gender specific differences in health-related quality of life for patients
  with bladder cancer following radical cystectomy. *J Clin Oncol* **41** (2023).
- 131 Ibilibor, C. Eliminating Differences in Outcomes by Race and Gender in Urothelial Carcinoma.
   *J Clin Oncol* **41** (2023).
- 882

#### 884 COMPETING INTERESTS

885 PT reports research funding from AstraZeneca, as well as personal fees as a consultant from Abbvie, 886 Bayer, Knight Pharmaceuticals, Tolmar and TerSera. AW and AB disclose funding received from 887 AstraZeneca for a PhD studentship for AB supervised by AW. AW discloses funding from imCORE for RE-ARM trial analysis. AW acknowledges funding from the RMH/ICR Cancer Research UK RadNet 888 889 Centre. RTB discloses research funding from Janssen, QED Therapeutics and UroGen Pharma, and 890 consultancy for Nonacus Limited and Cystotech ApS. AC discloses funding from Cancer Research UK, 891 National Institute of Health Research, Prostate Cancer UK, UK Research & Innovation, Elekta AB, 892 honoraria from Bayer PLC, Janssen, AZ, ASTRO, ASCO, Roche, Merck and is Editor in Chief of BMJ 893 Oncology. AC is supported by the NIHR Manchester Biomedical Research Centre.

894

### 895 KEY POINTS

- Bladder cancer is a common malignancy with a 3-4 to 1 male preponderance, suggesting
   relative 'resistance' to the development of the disease in females compared to males.
   Environmental exposures, hormone signalling, gene expression, immunology and tumour
   microenvironment likely coalesce to underpin this sexual dimorphism.
- Although bladder cancer outcomes are considered to be notably worse for females than
   males, this finding is not consistent across territories and treatment modalities, suggesting
   the considerable influence of healthcare system factors on outcomes. Such factors may
   include diagnostic delays and discrepancies in the appropriate and timely use of guideline based care.
- Addressing healthcare system factors by the implementation of best practices for referral,
   diagnosis and treatment could provide rapid improvements in outcomes where deficiencies
   exist.
- Nevertheless, the biological phenomena driving the sexual dimorphism in disease incidence are likely to also influence treatment responses, and better understanding of these mechanisms through carefully-designed fundamental research, and preclinical and clinical studies, may reveal sex-specific biomarkers or treatment approaches to benefit all bladder cancer patients.
- 913
- 914

#### 915 TABLE & FIGURE LEGENDS

**Table 1:** Summary of the main characteristics of the consensus classes of muscle invasive bladder
 cancer (MIBC) (adapted from<sup>77</sup>).

918 **Table 2:** Bladder cancer associated phenomena that demonstrate confirmed differences between the919 sexes.

Figure 1: 1-year, 2-year, 3-year, and 4-year bladder cancer survival (%) for adults (aged 15 to 99 years)
diagnosed in England 2015 to 2019, followed up to 2020<sup>17</sup>. (a) Age-standardised and non-age
standardised net survival (%) by sex. (b) Age-standardised net survival (%) by stage and sex. NB: 3Year Stage 1, 3-Year Stage 4, 4-Year Stage 1, 4-Year Stage 3 and 4-Year Stage 4 data are all missing for
female patients. Age-standardisation represents a weighted-average of mortality rates for each sex
based on the International Classification of Survival Standard.

Figure 2: Overall survival for patients with T1–T4 bladder cancers treated in England from 2013 to
2019, plotted using the Kaplan-Meier method with respect to sex (reproduced with kind permission
from John Wiley and Sons<sup>18</sup>).

**Figure 3:** A summary of immunological sex differences in BC: The female TM) is characterised by increased infiltration of immunosuppressive CD163+ M2-like macrophages and increased tumour expression of immune checkpoints. In addition, an increase in genes related to B cell recruitment (CXCL13) and function (CD40) has been shown in the female TME. In contrast, the male TME is characterised by increased exhausted CD8+ T cells (both of progenitor and terminally differentiated subtypes), which is driven by AR signalling specifically in CD8+ T cells. **AR**: androgen receptor, **IFN**: interferon, **TME**: tumour microenvironment.

Figure 4: An overview of different approaches to decrease the sex gap in bladder cancer outcomes.
Each domain and sub-domain likely require adequately-powered prospective studies in order to
validate and implement strategies that can reduce the sex gap in clinical outcomes.

939

## 940 SHORT SUMMARY

Bladder cancer outcomes are considered worse for women than for men, while incidence is 3-4 times
higher in men. Understanding biological phenomena and health system factors driving these
differences is essential to improve outcomes and develop novel treatment approaches.