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1 **NHTM_40**

2 **URINARY BIOMARKERS FOR THE DIAGNOSIS OF UROTHELIAL BLADDER CANCER**

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18 **Keywords:**

19 Bladder cancer; urinary biomarkers; diagnosis.

20 **ABSTRACT**

21 Urothelial bladder cancer is a common cancer associated with considerable burden for both patients
22 and healthcare providers alike. The majority of patients present with non-muscle-invasive bladder
23 cancer (NMIBC) which, although not immediately life-threatening, requires appropriate initial
24 management and long-term surveillance which is both invasive and costly. Accurate diagnostic
25 urinary biomarkers could be transformational in this setting, yet have proved to be a significant
26 challenge to bladder cancer scientists over the last two decades. Such biomarkers would need to
27 represent a range of tumour grades and stages, encompass inter- and intra-tumour heterogeneity,
28 and compete with the current diagnostic gold standard of cystoscopy with a sensitivity and
29 specificity of 85% and 87%, respectively. For the field to move forward in this current exciting era of
30 high-throughput proteomics and genomics, bladder cancer scientists need to find a consensus on
31 the optimal urinary substrate (DNA, RNA, protein, etc) and deliver robust well-designed studies in
32 the correct populations with appropriate statistical input. Issues relating to tumour heterogeneity
33 and anticipatory diagnosis also require considerable thought. The challenge remains unchanged.

34 **FOCAL POINTS**

- 35 • Accurate urinary biomarkers for the diagnosis of urothelial bladder cancer could be
36 transformational for patients and healthcare providers alike.
- 37 • To date, despite several FDA approvals, no such markers are routinely used in the clinical
38 setting.
- 39 • Poor study design and non-representative study populations are major contributory factors
40 and recent systematic reviews have highlighted such weaknesses.
- 41 • Low grade and low stage tumours are common, yet the most difficult to diagnose non-
42 invasively; however, they should be incorporated into study populations in proportions
43 representative of the incident and/or recurrent disease patient setting to avoid bias.
- 44 • Promising urinary biomarker substrates include proteins and nucleic acids, with inherent
45 strengths and weaknesses.
- 46 • Enduring challenges remain inter- and intra-tumour heterogeneity, and anticipatory
47 diagnosis.

48

49

50 **COMMENTARY**

51 Urothelial bladder cancer (UBC) is the seventh commonest cancer in Western societies [1], resulting
52 in 69,000 and 180,000 new cases per year in the USA and EU, respectively. The vast majority of new
53 cases are diagnosed following single or repeated episodes of haematuria (blood in the urine) which
54 is investigated by cystoscopy (inserting a “telescope” via the urethra into the bladder) and around
55 10% of patients investigated for haematuria will be diagnosed with UBC [2]. Following initial
56 treatment by transurethral resection of bladder tumour, 75-85% of these patients will be diagnosed
57 with non-muscle-invasive tumours (NMIBC, stages pTa/pT1/pTis), and the remainder muscle-
58 invasive tumours (MIBC, stages pT2-4) [3]. Thereafter, treatment strategies differ markedly: patients
59 with MIBC are likely to undergo more radical therapy with combinations of chemotherapy and
60 radiotherapy or cystectomy (removal of the bladder) [4], whereas those with NMIBC will be treated
61 with intravesical therapy (therapies delivered into the bladder) followed by cystoscopic surveillance
62 (regular inspection of the bladder) [5]. Schedules of cystoscopic surveillance (and the nature of
63 intravesical therapy) are determined by the risk category of NMIBC (low-, intermediate- or high-risk)
64 [5]. With disease recurrence a lifetime risk across all NMIBC categories (up to 80% [6]), and
65 progression to MIBC an important consideration for high-risk NMIBC patients (up to 45% [6;7]),
66 cystoscopic surveillance represents the mainstay of longer term management for all NMIBC patients.
67 Urine cytology is often used as an adjunct to cystoscopy: the microscopic detection of cancer cells in
68 the urine is a very specific indicator of UBC but has poor sensitivity for low-grade UBC, resulting in
69 low overall sensitivity [8].

70 Cystoscopy is invasive and burdensome for patients and expensive for healthcare providers [9;10],
71 such that from diagnosis to death on a per patient basis UBC is one of the most expensive
72 malignancies to manage [11]. Therefore, non-invasive or urinary biomarkers for the accurate and
73 reliable detection of urothelial bladder cancer (UBC) have the potential to be transformational for
74 both UBC patients and healthcare providers by reducing reliance on cystoscopy for diagnosis and
75 surveillance. Furthermore, this setting is fertile yet challenging ground for translational medicine.

76 Since UBCs are in direct contact with urine, urine is considered to be a promising biospecimen for
77 developing non-invasive tests to detect and characterise bladder tumours. However, UBCs are highly
78 heterogeneous with high mutational burden and variable copy number aberrations and gene
79 expression profiles [12;13]; thus, different tumours may release different biomarkers (necessitating
80 multimarker tests), and early-stage and low-grade tumours may only release very small amounts of
81 such markers, potentially impairing test sensitivity [14]. Markers must also be highly tumour-specific
82 so that haematuria itself and other non-malignant conditions do not generate false positives [14;15].
83 In the search for better urinary biomarkers genomic, proteomic and metabolomic approaches have
84 all yielded promising results [14;16-19]. Despite such work over several decades [20], a 2015
85 WHO/ICUD consensus stated that [8]:

- 86 • *Despite considerable advances in recent years, the authors feel that at this stage the added value*
87 *of molecular markers for the diagnosis of urothelial tumors has not yet been identified.*
- 88 • *Current data suggest that some of these markers may have the potential to play a role in*
89 *screening and surveillance of bladder cancer.*
- 90 • *Well-designed protocols and prospective, controlled trials will be needed to provide the basis to*
91 *determine whether integration of molecular markers into clinical decision-making will be of value*
92 *in the future.*

93 We recently undertook a systematic review of diagnostic and prognostic urinary protein biomarkers
94 and formed similar conclusions [20], principally that:

- 95 • *The majority of urine biomarker studies contain bias or are insufficiently reported.*
- 96 • *The urinary concentrations of a large number of proteins are increased by the presence of*
97 *bladder cancer, but most proteins are not increased in all cases and are not specific to bladder*
98 *cancer.*
- 99 • *NMP22, BTA, UBC and Cyfra 21-1 are the only well-validated urinary protein biomarkers and*
100 *their sensitivities and specificities are well below those of cystoscopy.*

101 We considered our approach to this systematic review to be stringent yet pragmatic [20], such that
102 it would provide a useful resource for workers in the field. We applied a number of criteria to define
103 whether individual studies provided “equivocal” or “unequivocal data” regarding a particular
104 biomarker(s) [20]. Unequivocal data were generated by studies which comprised of ≥ 20 cancer
105 patients and ≥ 20 controls; sensitivity and specificity had to be reported. Importantly, we also
106 required unequivocal studies to comprise $\geq 25\%$ stage pTa tumours (generally, smaller tumours and
107 more difficult to detect non-invasively, and whose incidence is c.50% [3;21]) and $\geq 15\%$ grade 1
108 tumours (the least cellularly and molecularly abnormal tumours [13] so also difficult to detect, and
109 whose incidence is c.25% [21]). These parameters ensured that the selected unequivocal studies had
110 to possess an element of statistical relevance, and also be representative of a normal UBC patient
111 population. Furthermore, if unequivocal data were generated from ≥ 3 studies, then we considered
112 the biomarker data to be validated.

113 We also classified the identified proteins as either “possible” or “unlikely” biomarkers dependent
114 upon whether the combined sensitivity and specificity was $\geq 80\%$ or $< 80\%$, respectively. White light
115 cystoscopy is currently the gold standard detection method for UBC, the reported sensitivity and
116 specificity of which vary greatly but a 2012 meta-analysis arrived at values of 85 and 87%,
117 respectively [22]; any urinary biomarker would need to match or improve upon cystoscopy to be
118 acceptable to patients and urologists. Hence, we were permissive in our definition of a possible
119 biomarker. Yet, as described, very few studies could be considered as unequivocal, although these
120 studies did report several possible biomarkers: fibronectin, clusterin, CEACAM1, apolipoprotein A4,
121 calprotectin, CD147, coronin-1A, DJ-1, reg-1, stathmin-1, and γ -synuclein [20].

122 We specifically limited our review to soluble urinary proteins as historically this has been the main
123 focus of UBC urinary biomarker research. Additionally, with the technology currently available, they
124 are the easiest class of biomolecule to use for point-of-care testing or to combine in an economical
125 single multiplex assay for the detection of UBC (should a suitable biomarker panel be determined).

126 We also envisage that measuring volatile metabolites [23], or advances in DNA sequencing may
127 allow point-of-care testing in the not too distant future. In fact, recent publications make a strong
128 case for DNA-based biomarkers being the frontrunners in the race to reduce reliance on cystoscopy
129 [24-28]. Although the amount of DNA that can be extracted from urine is low and variable, PCR and
130 advanced analysis techniques such as next generation sequencing allow identification of tiny
131 amounts of tumour DNA in the majority of urine samples, even in the presence of an excess of non-
132 tumour DNA [27]. Genome wide copy number changes in urinary DNA, microsatellite analysis,
133 methylation and mutations have all been used for the purpose [24-28]. Studies of urinary DNA have
134 focussed almost exclusively on DNA extracted from the urinary cell-pellets obtained by centrifuging
135 urine; however, we and others have found that cell-free DNA (cfDNA) in the urine supernatant
136 contains a higher fraction of tumour DNA than cell pellet DNA, and we are optimistic that urinary
137 cfDNA could underlie a clinically useful test for UBC detection [24;29]. As with protein biomarkers,
138 the performance of DNA biomarkers requires thorough evaluation prior to clinical uptake,
139 particularly in the disease surveillance setting.

140 Whatever the biomarker substrate (proteins, nucleic acids, etc) or source (urine supernatant, cell
141 pellet, etc), the field now needs to concentrate on designing and delivering the right studies in the
142 right patient populations and with due statistical consideration so that evidence synthesis is robust,
143 results are reproducible, and product marketing is not premature. Issues such as inter- and intra-
144 tumour heterogeneity should also be addressed, which may require the utilisation of biomarker
145 panels comprised of 10s or 100s of individual markers [19;24]. And the conundrum of “anticipatory”
146 or “pre-emptive” diagnosis requires clarification - the scenario whereby a patient is urinary
147 biomarker positive and cystoscopy negative, yet who develops recurrence within the following 12-24
148 months. Should such patients be treated as false positive, be placed under closer surveillance, be the
149 subject of personalised biomarkers based upon the tumour’s biomarker expression, or even be
150 treated pre-emptively with intravesical therapies? If the biomarker is highly specific, then the latter
151 three options could all be appropriate. The future is exciting and challenging.

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Figure 1: A suggested urothelial cancer biomarker research pipeline

