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## **The association between smoking cessation before and after diagnosis and non-muscle-invasive bladder cancer recurrence: a prospective cohort study**

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## 1 **Abstract**

2

3 **Background:** Smoking is a major risk factor for bladder cancer, but the relationship between  
4 smoking cessation after initial treatment and bladder cancer recurrence has been investigated  
5 less frequently and not prospectively yet.

6 **Methods:** 722 non-muscle-invasive bladder cancer (NMIBC) patients (pTa, pT1 and CIS)  
7 from the prospective Bladder Cancer Prognosis Programme (BCPP) cohort, selected in the  
8 UK between 2005-2011, provided complete data on smoking behaviour before and up to 5  
9 years after diagnosis. The impact of smoking behaviour on NMIBC recurrence was explored  
10 by multivariable Cox regression models investigating time-to-first NMIBC recurrence.

11 **Results:** Over a median follow-up period of 4.21 years, 403 pathologically confirmed  
12 NMIBC recurrences occurred in 210 patients. Only 25 current smokers at diagnosis quit  
13 smoking (14%) during follow-up and smoking cessation after diagnosis did not decrease risk  
14 of recurrence compared to continuing smokers ( $p=0.352$ ).

15 **Conclusions:** Although quitting smoking after diagnosis might reduce the risk of recurrence  
16 based on retrospective evidence, this is not confirmed in this prospective study because the  
17 number of NMIBC patients quitting smoking before their first recurrence was too low.  
18 Nevertheless, this indicates an important role for urologists and other health care  
19 professionals in promoting smoking cessation in NMIBC.

## 20 **Introduction**

21 Bladder cancer (BC) is estimated to be the ninth most frequent cancer worldwide with  
22 approximately 400,000 newly diagnosed cases per year [1]. Compared to other cancers,  
23 mortality rates are generally lower for BC [1] since the majority of BCs diagnosed are non-  
24 muscle-invasive bladder cancers (NMIBC) [2]. However, NMIBC often recurs [3] and has a  
25 risk of progressing to muscle-invasive bladder cancer (MIBC) [4], events which impact on  
26 the quality of life of the patient [5] and generate high disease management costs [6].

27 Although smoking is an established risk factor for BC, its effects has been less  
28 frequently investigated in relation to BC prognosis [7–10]. Although many studies  
29 investigated effectiveness of treatment for NMIBC and MIBC with regard to recurrence,  
30 progression and mortality, most studies did not investigate the effect of smoking or other  
31 factors modifiable by patients on BC prognosis [11]. Nevertheless, the number of studies also  
32 reporting hazard ratios (HRs) for BC recurrence by smoking status at diagnosis has increased  
33 recently and the current body of evidence consistently shows that there is a small association  
34 between smoking and BC recurrence when comparing current smokers to never smokers at  
35 diagnosis [10,12]. However, the impact of smoking cessation after BC diagnosis on  
36 recurrence and mortality has not yet been quantified prospectively [13]. Studies have  
37 investigated the impact of smoking cessation within one year after diagnosis on BC  
38 recurrence, showing a slight decrease in risk of recurrence [14,15], and one study indicating  
39 no effect of quitting after diagnosis on overall or bladder cancer-specific mortality [16].

40 The Bladder Cancer Prognosis Programme (BCPP) followed-up BC patients for five  
41 years post-diagnosis and investigated changes in smoking behaviour in relation to the course  
42 of the disease [17]. The principal aim of this study was to investigate whether smoking  
43 cessation post-diagnosis and smoking behaviour pre-diagnosis influences BC recurrence.

## 44 **Methods**

### 45 **The Bladder Cancer Prognosis programme**

46 This study was conducted within the framework of the West Midlands Bladder Cancer  
47 Prognosis Programme (BCPP), a cohort study in the United Kingdom. Details of the study  
48 are described elsewhere [17]. In brief, individuals were included between December 2005  
49 and October 2011 after referral to participating urology centres due to symptoms suspicious  
50 of BC and followed for a maximum of 5 years from diagnosis. Patients with previous cancer  
51 of the urethra, bladder, ureter, or renal pelvis within the last decade were excluded. The study  
52 was ethically-approved (06/MRE04/65) and all participants gave written informed consent.

53

### 54 **Data collection**

55 At or around time of diagnosis, trained research nurses used semi-structured face-to-face  
56 interviews and questionnaires to collect data on social support, health-related quality of life,  
57 sociodemographics, medical history, and health-related behaviours including smoking  
58 behaviour. Variables on smoking behaviour included current smoking status (never, former,  
59 current), duration (years of smoking), intensity (cigarettes per day), smoking cessation (in  
60 years) and tobacco type (filter, non-filter or rolled cigarettes, cigar or pipe). Monthly smoking  
61 status was also assessed retrospectively by postal questionnaires that were sent out to  
62 participants yearly until the end of follow-up.

63

### 64 **Smoking status at diagnosis and during follow-up**

65 A combined smoking status variable was created indicating continuing smokers, former  
66 smokers who consistently abstained, never smokers, former smokers who started smoking  
67 again, and current smokers who quit smoking post-diagnosis. Patients were considered  
68 quitters when they abstained consistently, so smokers who quit for 3 months and then started

69 again were considered as continuing smokers. Furthermore, for each participant that reported  
70 smoking cessation during follow-up it was confirmed whether this occurred before or after  
71 their first recurrence. If patients quit smoking after their first recurrence, they were  
72 considered as continuing smokers in the time-to-first recurrence analysis.

73

#### 74 **Population at risk**

75 Of the 1,550 cases who agreed to participate, 231 were subsequently identified as not having  
76 BC. Patients who presented with MIBC (n=275) disease at diagnosis were excluded from  
77 analysis because they are fundamentally different from NMIBC with regard to recurrence.  
78 Patients with squamous or adeno-carcinomas of non-urothelial origin or with bladder cancer  
79 as secondary carcinoma were excluded (n=41). In addition to patients presenting with Ta and  
80 T1 tumours, carcinoma in situ (CIS) tumours were included (n=16) since they have an  
81 increased risk of recurrence [18]. In total, 846 (84%) of these patients had provided data on  
82 smoking behaviour at diagnosis and during follow-up and remained under follow-up within  
83 the cohort study. Of the included 846 NMIBC patients, there were 116 patients with  
84 unknown recurrent tumour stage. These 116 unconfirmed events were excluded for other  
85 analyses as well as 8 cases who had radiotherapy (on suspicion of being MIBC cases)  
86 resulting in a NMIBC patient population at risk of recurrence of 722.

87 No systematic guidance or tools were provided to enable patients to quit smoking  
88 after diagnosis, so care as usual was applied by all participating urologists.

89

#### 90 **Statistical analysis**

91 BC recurrence was defined as a new tumour that was the same stage as the primary  
92 tumour (Ta or T1) but also when a primary Ta patient had a T1 recurrence. Patients that  
93 progressed from T1 to T2 disease were not counted as a recurrence but as a progression

94 event. Unfortunately, there were not enough events to also consider biological progression  
95 within this sample of NMIBC patients, as defined in the BCPP cohort [19]. Therefore, this  
96 study only focussed on confirmed recurrence events and patients who experienced a  
97 progression event were censored in the survival analysis when the progression event was  
98 diagnosed.

99         The impact of smoking behaviour on BC recurrence was explored by Cox regression  
100 models—with time since initial transurethral resection of the bladder tumour (TURBT) as the  
101 time-metric—investigating possible differences in likelihood of a first recurrence. We  
102 explored two different Cox regression models: one adjusted for age at diagnosis and sex  
103 (model 1) and one additionally adjusted for BC stage, grade, tumour size and number of  
104 tumours at diagnosis (model 2). This set of confounders was chosen since they are markers of  
105 NMIBC prognosis and are factors that contribute to European Association of Urology (EAU)  
106 risk stratification for clinical decisions[20]. Moreover, they are potentially associated with  
107 smoking behaviour at diagnosis [21]. Consequently, conditional risk set modelling was  
108 applied to investigate time between multiple recurrent events and analysis time was reset at  
109 each event [22]. For this analysis, resection of tumours was added to model 2 as a  
110 confounder. The proportional hazards assumption was checked in all models using  
111 Schoenfeld residuals. Cumulative incidence functions (CIF) corrected for competing risks  
112 (death) were made [23].

113         Furthermore, the differences in mean number of recurrences over 5 years between  
114 never smokers, former smokers and continuing smokers were compared using a multivariable  
115 ANOVA model correcting for pairwise comparisons using Tukey's HSD. There were not  
116 enough BC-related death events (45) or confirmed progression events (19) to allow for  
117 separate analyses. A similarly low number of progression events has been observed in a large  
118 (n=718) NMIBC patient sample before [24].

119 NMIBC patients who died before the end of follow-up (n=157) were censored at time  
120 of death and patients who underwent cystectomy (n=15) were censored at the date of  
121 cystectomy (13). Other patients were considered lost to follow-up when the date on which  
122 patients were last seen in the hospital for bladder cancer-related therapy or the date on which  
123 they filled in their last follow-up questionnaire was before the end of follow-up (5 years).



## 124 **Results**

### 125 **Number of recurrences and characteristics of population at risk**

126 All 722 patients at risk of recurrence were followed over a median period of 4.21 years (IQR  
127 = 2.64-5.00 years). The majority of patients (506, 70%) were followed for at least 3 years.  
128 Over this period of follow-up, 210 NMIBC patients experienced at least one confirmed  
129 recurrence event. These 210 NMIBC patients accumulated a total of 403 confirmed  
130 recurrence events in the cohort.

131 Most cases were male (79%) and around the age of 70 (Table 1). Furthermore,  
132 continuing smokers seemed to be underrepresented in the low EAU risk group (12%), those  
133 who quit smoking seemed more likely to be younger and female, and continuing smokers  
134 seemed more likely to present with multiple tumours at diagnosis (Table 1). In the  
135 multivariate models, 26 patients were not included in the analysis due to missing data on age  
136 (n=7), number of tumours at diagnosis (n=15) and tumour size (n=4). Because participants  
137 were recruited from multiple centers, patients were treated by multiple urologists with  
138 different individual thresholds to perform certain therapies. Therefore, not all patients were  
139 treated exactly according to the EAU guidelines [20], which is often the case in actual  
140 clinical practice [25].

141 **Table 1. Patient characteristics at diagnosis & number of recurrences over 5 years for**  
 142 **722 NMIBC patients treated with transurethral resection by smoking category.**

	Overall (n=722)	Combined smoking status					p- value*
		Never smoker (n=103)	Former smoker (n=266)	Continuing Smoker (n=186)	Former smoker who started again (n=150)	Quitters after diagnosis (n=17)	
<b>Age in years</b>							<0.001
Median (25th-75th percentile)	71 (63-77)	72 (61-79)	72 (67-79)	67 (57-74)	72 (64-77)	62 (56-67)	
<b>Sex</b>							<0.001
Male	573 (79%)	63 (61%)	231 (87%)	139 (75%)	129 (86%)	11 (65%)	
Female	149 (21%)	40 (39%)	35 (13%)	47 (25%)	21 (14%)	6 (35%)	
<b>EAU risk group</b>							<0.001
Low	128 (18%)	28 (27%)	71 (27%)	23 (12%)	4 (3%)	2 (12%)	
Intermediate	383(53%)	50 (49%)	131 (49%)	97 (52%)	91 (61%)	14 (82%)	
High	211 (29%)	25 (24%)	64 (24%)	66 (36%)	55 (37%)	1 (6%)	
<b>Number of tumours</b>							<0.001
1	429 (61%)	70 (70%)	179 (69%)	100 (55%)	69 (46%)	11 (65%)	
2-7	258 (36%)	27 (27%)	74 (28%)	76 (42%)	75 (50%)	6 (35%)	
>=8	22 (3%)	3 (3%)	8 (3%)	6 (3%)	5 (3%)	0 (-)	
<b>Tumour size</b>							0.068
<3cm	445 (63%)	68 (68%)	174 (67%)	105 (58%)	85 (57%)	13 (76%)	
>=3cm	260 (37%)	32 (32%)	84 (33%)	77 (42%)	63 (43%)	4 (24%)	
<b>Grade</b>							0.001
1	212 (30%)	34 (34%)	99 (38%)	51 (28%)	26 (17%)	2 (13%)	
2	257 (36%)	34 (34%)	75 (28%)	73 (40%)	66 (44%)	9 (56%)	
3	245 (34%)	33 (33%)	90 (34%)	60 (32%)	57 (38%)	5 (31%)	
<b>Stage</b>							0.590
pTa	476 (66%)	68 (66%)	184 (69%)	115 (62%)	95 (63%)	14 (82%)	
pT1	239 (33%)	35 (34%)	79 (30%)	69 (37%)	53 (35%)	3 (18%)	
pCis	7 (1%)	0 (-)	3 (1%)	2 (1%)	2 (1%)	0 (-)	
<b>No of recurrences</b>							0.337
1	108 (51%)	18 (62%)	28 (46%)	33 (53%)	27 (52%)	2 (33%)	
2	46 (22%)	6 (21%)	16 (26%)	16 (26%)	6 (11%)	2 (33%)	
>3	56 (27%)	5 (17%)	17 (28%)	13 (21%)	19 (37%)	2 (33%)	
<b>Smoking intensity</b>							0.076
1-9 cigarettes	128 (29%)	NA	55 (30%)	23 (21%)	42 (34%)	8 (50%)	
10-19 cigarettes	140 (32%)	NA	53 (28%)	42 (38%)	42 (34%)	3 (19%)	
>20 cigarettes	167 (38%)	NA	78 (42%)	45 (41%)	39 (32%)	5 (31%)	
<b>Smoking duration</b>							<0.001
1-9 years	45 (10%)	NA	26 (14%)	2 (2%)	16(14%)	1 (6%)	
10-19 years	83 (19%)	NA	43 (23%)	10 (9%)	29 (25%)	1 (6%)	
20-29 years	87 (20%)	NA	46 (25%)	12 (11%)	27 (23%)	2 (13%)	
30-39 years	88 (21%)	NA	37 (20%)	28 (25%)	19 (16%)	4 (25%)	
>40 years	127 (30%)	NA	32 (17%)	60 (54%)	27 (23%)	8 (50%)	
<b>Smoking cessation</b>							0.051
<20 years	48 (12%)	NA	23 (9%)	NA	25 (17%)	NA	
21-40 years	208 (51%)	NA	134 (51%)	NA	74 (49%)	NA	
>40 years	155 (38%)	NA	104 (40%)	NA	51 (34%)	NA	

\*Kruskal-Wallis test for continuous and chi-square test for categorical variables

**143 Associations between smoking behaviour pre and post-diagnosis and BC recurrence**

144 Although HR estimates for smoking cessation pre-diagnosis indicated a protective  
145 association with BC recurrence, the p for linear trend was not statistically significant  
146 ( $p_{\text{trend}}=0.126$ ) and therefore the association cannot be considered as strong (Table 2). No  
147 association between smoking status and risk of recurrence was observed in the multivariable  
148 model (Table 2). Interestingly, when compared to continuing smokers (HR=1.04, 95%  
149 CI=0.65-1.66) HRs were similar for those who quit smoking ( $p=0.352$ ) and former smokers  
150 who started again post-diagnosis ( $p=0.431$ ) (Table 2). Additionally, the cumulative incidence  
151 function shows that cumulative incidence of BC recurrence was lowest for former smokers  
152 and never smokers (Figure 1).

153

**154 Insert Figure 1 here**

155

**156 Figure 1. Cumulative incidence functions with correction for competing risk (death)**  
**157 indicating cumulative incidence of first recurrence per category of smoking**  
**158 status in NMIBC patients treated with TURBT.**

159

160 Only 25 smokers (14%) of the 174 current smokers originally recorded at diagnosis  
161 quit smoking at any point during follow-up. Three quitters were excluded for full analysis for  
162 not having information on their date last seen and another five had missing data regarding the  
163 invasiveness of their recurrent events. Of the 480 former smokers at diagnosis, 172 (36%)  
164 started smoking (any form of tobacco) again post-diagnosis in all included 846 NMIBC  
165 patients.

166            Exposure to environmental tobacco smoke during childhood (HR=1.17, 95%CI=0.81-  
167 1.68) or adulthood (HR=1.02, 95%CI=0.76-1.36) did not seem to have any impact on time to  
168 first recurrence (Table 2).

169 **Table 2. Cox regression analysis investigating the association between combined**  
 170 **smoking status, smoking cessation before diagnosis and passive smoking and time-to-**  
 171 **first recurrence in NMIBC patients treated with TURBT.**

	Age & sex adjusted			Multivariable model*		
	HR	95% CI	number of events / patients at risk	HR	95% CI	number of events / patients at risk
<b>Combined smoking status</b>						
Never smoker	1.00	ref	29/103	1.00	ref	28/99
Former smoker	0.79	0.51-1.24	61/266	0.78	0.48-1.24	59/254
Continuing smoker	1.17	0.75-1.83	62/186	1.04	0.65-1.66	61/180
Former smoker who started again**	1.04	0.65-1.64	51/150	0.87	0.53-1.41	49/146
Current smoker who quit smoking***	1.25	0.52-3.00	6/17	1.47	0.63-3.41	6/17
<b>Smoking cessation (in years) ****</b>						
<20 years	0.81	0.46-1.43	15/48	0.82	0.46-1.46	15/47
21-40 years	0.76	0.53-1.08	57/208	0.74	0.51-1.08	54/200
>40 years	0.67	0.44-1.02	39/155	0.71	0.46-1.09	38/148
<b>p for trend</b>	0.070			0.126		
<b>Exposed to passive smoking during childhood?</b>						
No	1.00	ref	36/142	1.00	ref	35/138
Yes	1.23	0.86-1.75	173/576	1.17	0.81-1.68	168/554
<b>Exposed to passive smoking during adulthood?</b>						
No	1.00	ref	74/261	1.00	ref	74/261
Yes	1.03	0.77-1.38	135/454	1.02	0.76-1.36	135/454

\* All estimates adjusted for age, sex, stage, grade, tumour size and number of tumours

\*\* Former smoker who started again and current smoker who quit smoking not included in former smokers at diagnosis

\*\*\* Smokers who quit after their first event are considered as current smokers

\*\*\*\* Reference category = current smokers at diagnosis, estimates also include former smokers who started again after diagnosis

173            Table 3 shows HRs for time to first recurrence by smoking intensity, duration and  
174 pack-years. No linear trends were observed although the highest categories showed the  
175 highest point estimates for both smoking intensity and pack years. For smoking duration the  
176 HRs were divergent and did not indicate any trend ( $p_{\text{trend}}=0.729$ ) at all.

177 **Table 3. Multivariable Cox regression analysis concerning the association between**  
 178 **smoking pack-years, intensity and duration (recorded at diagnosis) with time to first**  
 179 **recurrence in NMIBC patients treated with TURBT.**  
 180

	Age & sex adjusted			Multivariable model*		
	HR	95% CI	number of events / patients at risk	HR	95% CI	number of events / patients at risk
<b>Never smoker</b>	1.00	ref	29/103	1.00	ref	28/99
<b>Pack-years</b>						
1-9 packyears	0.86	0.53-1.42	36/141	0.81	0.48-1.37	34/134
10-19 packyears	0.95	0.54-1.67	22/81	0.92	0.51-1.65	22/80
20-29 packyears	0.93	0.49-1.77	15/58	0.81	0.42-1.60	15/57
30-39 packyears	0.70	0.35-1.43	11/55	0.60	0.30-1.22	11/53
>40 packyears	1.28	0.76-2.14	30/86	1.14	0.66-1.97	29/83
p for trend	0.365			0.688		
<b>Smoking intensity (cigarettes/day)</b>						
1-9 cigarettes	0.83	0.50-1.38	32/128	0.81	0.47-1.38	30/122
10-19 cigarettes	0.75	0.45-1.28	31/140	0.61	0.35-1.07	31/138
20+ cigarettes	1.24	0.79-1.96	55/167	1.16	0.72-1.85	54/160
p for trend	0.112			0.198		
<b>Smoking duration (in years)</b>						
1-9 years	1.03	0.52-2.05	12/45	0.97	0.48-1.95	12/43
10-19 years	0.94	0.54-1.62	22/83	0.85	0.48-1.50	21/78
20-29 years	0.79	0.45-1.39	21/87	0.79	0.44-1.44	20/85
30-39 years	1.08	0.61-1.89	26/88	0.93	0.52-1.66	25/85
40+ years	1.00	0.60-1.64	36/127	0.88	0.52-1.49	36/124
p for trend	0.917			0.729		

\* All estimates adjusted for age, sex, stage, grade, tumour size and number of tumours at diagnosis

182           When considering multiple events that have occurred in patients (Table 4) the HRs  
 183 are similar to the time to first recurrence analysis (HR for continuing vs never smokers is  
 184 1.10, 95%CI=0.72-1.69). However, continuing smokers seemed to have experienced more  
 185 recurrences than never smokers on average over 5 years on average, however not  
 186 significantly (0.64 vs 0.45, p=0.308).

187

188 **Table 4. Conditional risk set model investigating time between multiple recurrence**  
 189 **events in NMIBC patients treated with TURBT by smoking status at diagnosis and**  
 190 **after diagnosis.**

	HR*	95% CI	number of events / patients at risk	Mean number of recurrences over 5 years (95% CI)
<b>Smoking status</b>				
Never smoker	1.00	ref	43/99	0.45 (0.28-0.63)
Former smoker	0.71	0.47-1.08	108/254	0.45 (0.33-0.57)
Continuing smoker	1.10	0.72-1.69	116/180	0.64 (0.47-0.81)
Former smoker who started again	0.89	0.56-1.43	108/146	0.82 (0.57-1.06)
Current smoker who quit smoking**	0.85	0.35-2.04	18/19	0.84 (0.10-1.58)

\* All estimates adjusted for age, sex, stage, grade, tumour size, number of tumours and resection of recurrent tumour

\*\* Smokers who have quit after their first event (n=2) are also included

191



## 192 **Discussion**

### 193 **Smoking cessation post-diagnosis and BC recurrence & clinical implications**

194 The reported HRs give reason to believe that quitting smoking does not influence the  
195 likelihood of NMIBC recurrence over 5 years when compared to continuing smokers in our  
196 sample. However, the number of quitters in our prospective sample was small which  
197 complicates drawing conclusions for this group. Another (retrospective) patient cohort study  
198 which assessed smoking cessation post-diagnosis concluded that quitting smoking  
199 significantly reduced risk of recurrence (HR=0.45, 95% CI=0.25-0.83, comparing quitters to  
200 continuing smokers), however the proportion of quitters (~43% of current smokers at  
201 diagnosis) was also considerably larger [14]. In another retrospective cohort study, Fleshner  
202 et al concluded that it remained unclear whether smoking cessation at time of diagnosis is  
203 beneficial with regard to BC recurrence [15] although Aveyard et al. estimated that the  
204 Fleshner study shows a HR of 0.71 (95% CI=0.48-1.05) when comparing quitters to  
205 continuing smokers[26], which is similar to the estimate observed in the study by Chen et al.  
206 Taken together, the limited evidence at this point seems to indicate that quitting smoking at  
207 or closely after diagnosis could reduce risk of recurrence. However, even across several  
208 smoking-related cancer sites such as lung cancer where this association is stronger, evidence  
209 to imply a strong, causal relationship between smoking behaviour after diagnosis and  
210 recurrence is still limited [27] so more prospective research is needed.

211         Considering the prolonged latency period for the development of BC after exposures  
212 [2], it is credible that the association between altering smoking behaviour post-diagnosis and  
213 likelihood of a first recurrence or multiple recurrences over 5 years is not as strong as the  
214 association between smoking and carcinogenesis. Similarly, epidemiological evidence  
215 suggests that pre-diagnostic smoking cessation does not immediately lower the risk of BC  
216 [28], also indicating a longer latency period than 5 years. Furthermore, it is considered that a

217 first BC recurrence is often the result of incomplete resection and/or tumour cell re-  
218 implantation, and that genuine new tumour formation only plays a more important role in  
219 later recurrences [29]. It is therefore reasonable to suggest that, because of the DNA-  
220 damaging effects of cigarette smoke [30], modifying smoking behaviour may only influence  
221 later recurrences and possibly those that may occur beyond the follow-up period of 5 years  
222 reported here.

223 Notwithstanding the results from our study, when considering the impact of  
224 comorbidities on overall survival in BC patients [31] which include several smoking-related  
225 diseases [32] and other evidence indicating beneficial and significant results of post-  
226 diagnostic smoking cessation in retrospective studies [14,15], it is evident that smoking  
227 cessation should be encouraged for NMIBC patients at diagnosis.

228 It is striking that only 14% of current smokers at diagnosis in our sample quit  
229 smoking post-diagnosis. There are examples of successful smoking cessation interventions in  
230 urology [33], and several studies found that when patients were diagnosed with BC they were  
231 more likely to quit smoking [34,35]. Therefore, urologists should continue to improve  
232 smoking cessation counselling in newly diagnosed NMIBC patients and to be current on the  
233 available tools to improve smoking cessation figures. Moreover, more intervention clinical  
234 research investigating smoking cessation programmes in NMIBC patients is warranted.

235

### 236 **Smoking behaviour pre-diagnosis & exposure to environmental tobacco smoke**

237 Smoking cessation was most beneficial, with regard to reducing the risk of recurrence, the  
238 longer before diagnosis it happened compared to continuing smokers. This was the strongest  
239 association observed in our study and has been observed in other studies as well, although not  
240 consistently [12]. Other results were in line with earlier studies investigating smoking status

241 at diagnosis and BC recurrence as well, by indicating a slightly increased risk of recurrence  
242 in NMIBC patients for current smokers compared to never smokers in a meta-analysis [10].

243 Another recent study not included in the aforementioned meta-analysis shows similar  
244 HRs (HR=1.49, 95% C.I.=0.95-2.33) for current smokers at diagnosis [8]. However, when  
245 including this study and our study (data from continuing smokers) in the meta-analysis the  
246 pooled HR barely changes from 1.27 (95% CI=1.09-1.46) to 1.26 (95% CI= 1.12-1.40) [10],  
247 indicating a significantly increased risk of recurrence for current smokers at diagnosis  
248 compared to never smokers. Possibly, the lack of association for continuing smokers in this  
249 study can be explained through multiple synchronous tumours being present at diagnosis in  
250 epithelial tumours. This theory of “field cancerization” proposes that (pre-)malignant  
251 transformation of cells has already occurred at different sites across the urothelium,  
252 explaining why (changing) smoking exposure will not have a large impact on disease  
253 prognosis [36].

254 Additionally, given that recent reviews indicate no considerable heterogeneity between  
255 studies that do not show an association between environmental tobacco smoke and risk of  
256 BC, it is unlikely that we would have shown any substantial association with BC recurrence  
257 either [37,38].

258 Because no substantial association between smoking status pre-diagnosis and BC  
259 recurrence was observed in adjusted models it is possible that the tumour characteristics  
260 associated with BC recurrence (stage, grade, tumour size, number of tumours) included as  
261 confounders in these models overshadow the effects of smoking behaviour in determining  
262 risk of BC recurrence [21] and possible also mortality since no association between quitting  
263 smoking after diagnosis and all-cause or bladder-cancer-specific mortality was observed in a  
264 large retrospective cohort study[16]. Moreover, since current smokers at diagnosis in our  
265 cohort have been associated with having a higher stage, higher grade and larger tumour size

266 compared to never smokers [39], smoking behaviour might play a more crucial role in  
267 determining risk of recurrence already before diagnosis through promoting unfavourable  
268 tumour characteristics associated with BC recurrence at diagnosis, although in a Dutch cohort  
269 of 323 UBC patients there was only a weak association between smoking intensity and  
270 increased risk of a more aggressive tumour type [40].

271

## 272 **Strengths and weaknesses**

273 Despite the prospective nature of our study there were some limitations restricting the  
274 analyses. Due to the relatively short follow-up of this study, long term effects of smoking  
275 cessation post-diagnosis could not be assessed and the number of deaths due to BC in the  
276 NMIBC patients within our cohort was too low for Cox regression analysis. Also, it was not  
277 possible to obtain detailed information on adjuvant therapy for all patients, so differences in  
278 adjuvant therapy could not be considered in the statistical analysis. Additionally, we did not  
279 correct for biomarkers of BC recurrence such as mutations in the *FGFR3* or *TP53* genes [41],  
280 although they might work together with smoking intensity in predicting BC outcome [42].

281 Furthermore, one of the caveats of using only self-reported questionnaire data to  
282 assess smoking exposure was likely demonstrated in our sample of NMIBC patients. The  
283 large proportion (about 1 in 3) of former smokers pre-diagnosis who reported to have started  
284 smoking again post-diagnosis is implausible and is probably observed due to  
285 misclassification of either the questionnaire at baseline or during follow-up. A high  
286 misclassification rate (47%) when comparing self-reported data on smoking behaviour to  
287 cotinine values in blood was also shown in another sample of bladder cancer patients  
288 undergoing surveillance [43]. Preferably, future studies should consider more reliable ways  
289 of verifying smoking exposure through biochemical analysis.

290           Unfortunately, at the start of the study we did not anticipate this small proportion of  
291   quitters after diagnosis which is why the analysis concerning quitters is underpowered.

## 292 **Conclusion**

293 Although quitting smoking after diagnosis might reduce probability of recurrence based on  
294 retrospective evidence, the number of NMIBC patients quitting smoking in our prospective  
295 study was low. This indicates an important role for urologists and other health care  
296 professionals in promoting smoking cessation in NMIBC. Based on the current evidence,  
297 smoking cessation pre-diagnosis seems to have the largest impact on reducing risk of  
298 recurrence after NMIBC diagnosis.

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