

A Comparative Analysis of the Influence of Gender, Pathway Delays, and Risk Factor Exposures on the Long-term Outcomes of Bladder Cancer

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1 **A COMPARATIVE ANALYSIS OF GENDER, PATHWAY DELAYS AND RISK FACTOR EXPOSURES**
2 **ON THE LONG-TERM OUTCOMES FROM BLADDER CANCER**

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26

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31

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33 treatment outcome, risk factors.

34

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36

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38

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40 **ABSTRACT**

41 **Background:** The relationship between pathway delays and bladder cancer-specific survival
42 is complex due to the influence of tumour- and patient-specific factors.

43

44 **Objective:** To investigate the influence of tumour factors, patient factors, carcinogen
45 exposure and pathway delays on the long-term outcome of urothelial bladder cancer (UBC).

46

47 **Design, Setting and Participants:** A cohort of 1537 UBC patients were enrolled 1/1/1991-
48 30/6/1992 and followed-up for 17.7years. The period from onset of symptoms to first
49 treatment (TURBT) was divided into 3 components of potential delay.

50

51 **Outcome Measurements and Statistical Analysis:** Associations between patient factors,
52 tumour factors and delay times were analysed using Pearson's chi-squared test and Mann-
53 Whitney U-test. Survival was calculated from date of TURBT to date of death or censor date
54 of 31/12/2010. Competing risks of death were assessed with the cumulative incidence
55 function (CIF); comparisons of CIFs were performed using Gray's test.

56

57 **Results:** At censor, reliable data were available for 1478 patients, of which 75% had died.
58 Females presented more commonly with muscle-invasive bladder cancer (MIBC) (30% vs.
59 26%) and less frequently with pT1 disease (18% vs. 24%) ($p=0.06$), had a longer total delay
60 time (median 120days vs. 106days, $p=0.02$), and those with MIBC had a significantly higher
61 cumulative incidence of death due to UBC (80% vs. 67% at 17years, $p<0.02$). Cox regression

62 identified age, smoking status, and tumour stage, grade, and size as the most significant
63 determinants of poor outcome.

64

65 **Limitations:** We did not capture downstream delays associated with cystectomy or
66 radiotherapy.

67

68 **Conclusions:** Female patients present later than males, and our data suggest that delay in
69 referral may be contributory. The relationship between gender, outcomes, delays and
70 aetiology of UBC is complex.

71

72 **Patient Summary:** We followed a large group of bladder cancer patients for over 17years.
73 The relationship between pathway delays and survival is complex. However, female patients
74 present later than male patients, and our data suggest that delay in referral from general
75 practice may be contributory.

76 **INTRODUCTION**

77 Urothelial bladder cancer (UBC) is the fifth most common cancer in Western societies,
78 accounting for 10,000, 69,000 and 180,000 new cases per year in the UK, USA and EU [1],
79 respectively. The global incidence of the disease is rising reflecting patterns of cigarette
80 smoking and occupational carcinogen exposure [2], the most common aetiological factors
81 [1]. There has been little improvement in the outcome for UBC patients since the 1980s,
82 reflecting complex diagnostic pathways and treatment regimens, and a lack of therapeutic
83 advances [3]. Given these constraints, much attention has been paid to reducing delays in
84 presentation [4], diagnosis and treatment [5].

85

86 For UBC the relationship between time to diagnosis and treatment, and disease-specific
87 survival is complex [6-9]; many tumours are indolent, for which delay in diagnosis does not
88 alter survival [10], and outcomes from aggressive UBCs are multifactorial [6-9]. In addition
89 to delays in healthcare pathways, disease biology (reflected by stage, grade and tumour
90 characteristics [11;12]) and patient-specific factors are important. The latter reflect
91 aetiological agent exposures (e.g. smoking is more common in males) [9;13;14], gender-
92 specific misdiagnoses (e.g. females are more likely to be incorrectly diagnosed with infection
93 [15]) [1;16;17], and potential differences in the molecular pathogenesis of male and female
94 UBC [18].

95

96 To obtain a clearer understanding of factors affecting outcomes in UBC, we have followed a
97 large cohort of prospectively recruited patients since 1991 [9]. This population represents
98 85% of new cases of UBC arising over an 18month period within the West Midlands region

99 of the UK [9]. Here we report long-term outcomes and investigate the influence of gender,
100 carcinogen exposure and pathway delays in this cohort.
101

102 PATIENTS AND METHODS

103 Patients

104 Patients newly-diagnosed with UBC within the West Midlands (UK) were prospectively
105 recruited from 1st January 1991 through 30th June 1992 [9]. Data regarding exposures, dates
106 of onset of symptoms, first referral by GP, first hospital appointment and first treatment
107 (date of TURBT) were collected at recruitment. Data were checked to ensure that TNM
108 classification correlated with histopathology and bimanual examination findings.
109 Discrepancies were resolved by the investigators and the operating Consultant. All patients
110 were notified to the West Midlands' cancer registry, who provided death information at the
111 censor date of 31st December 2010. Ethics committee approval was received prior to study
112 opening. Ex-smoking was defined as abstinence for >12 months. Occupational exposure was
113 identified by 3 assessors (>90% consensus) utilising IARC contemporary evidence to assign
114 no risk, possible risk and definite risk of working in an occupation implicated in the
115 pathogenesis of UBC (see Supplementary Table 1) [19].

116

117 Pathway measures

118 Pathway times were defined as:

- 119 ▪ Time 1: date of onset of patient's symptoms to date of GP's first referral to
120 secondary care;
- 121 ▪ Time 2: date of GP's first referral to secondary care to date of first hospital
122 attendance for urological assessment;
- 123 ▪ Time 3: date of first hospital attendance to date of first treatment by TURBT.

124 Hospital delay included the addition of Times 2 and 3, and Total delay was the summation of
125 all three time periods.

126

127 Statistical Methods

128 All statistical analyses were performed using Stata 11.2 (*StataCorp LP, College Station,*
129 *Texas, USA*) and R version 2.13.2 (*The R Foundation for Statistical Computing, [http://www.R-](http://www.R-project.org)*
130 *project.org*). Associations between patient and tumour features, with median delay times
131 were analysed using Pearson's chi-squared test for categorical data and the Mann–Whitney
132 U-test for continuous data. Survival was calculated from the date of first TURBT to the date
133 of death or censor date of 31st December 2010, using all-cause mortality. Survival curves for
134 each stage (Ta, T1, T2-4) were constructed using the Kaplan-Meier method and outcomes
135 between groups compared using the log-rank test. We estimated relative survival to
136 calculate the crude probability of death in the general population compared to patients
137 diagnosed with pTa tumours using the user written Stata command *strs* matched for age at
138 diagnosis, sex and year of diagnosis [20]. The calculated probabilities were based upon the
139 Ederer II method. Survival was compared in terms of demographic and tumour
140 characteristics and delay times. A stratified survival analysis was used to test for differences
141 within delay times adjusting for tumour stage and to test for smoking status adjusting for
142 delay times. Cox-proportional hazards models using a complete case approach were applied
143 to investigate the independent effect of age, sex, smoking status, haematuria, tumour stage,
144 grade, type, size and number. We tested the proportional hazards assumption of the models
145 by examining the Schoenfeld and scaled Schoenfeld residuals; in each test the proportional
146 hazards assumption was met. In addition, we evaluated the fit of the models using Cox-Snell

147 residuals which confirmed the models to fit the data well. This formed a base model that
148 was used to adjust the effects of each delay. Hazard ratios with 95% CI and P values are
149 presented.

150 To assess the competing risks of death, we first used a non-parametric test to assess the
151 equality between groups by calculating the cumulative incidence function (CIF) as described
152 by Scrucca et al. [21]. Comparison of specific CIFs was performed using Gray's test [22]. See
153 Supplementary Methods for further details.

154

155 **RESULTS**

156 Cohort description

157 In total, 1537 patients were enrolled into the study and reliable long-term survival data
158 were available on 1478 (96.2%) (**Table 1**). The cohort was typical for UBC, with a male to
159 female ratio of 3:1 and a median age at diagnosis of 69 years for male (IQR 62-76) and 71
160 years for female patients (IQR 64-78). Most patients were current or former cigarette
161 smokers (973, 77%), and 330 (27%) patients were classified as having possible or definite
162 exposure to occupational carcinogens. As detailed previously, patients were treated by
163 contemporaneous standard practice (which did not include re-resection), and surveillance
164 was carried out according to national guidelines [9]. At the censor date, the mean follow-up
165 was 106 months (8.8 years, IQR 22.0-212.8 months) and 1109 patients (75%) had died. The
166 cause of death was known for 983 patients (89% of deaths) (**Table 2**).

167

168 Pathological features

169 There was a significant association between tumour stage and death from UBC ($P < 0.005$,
170 **Figure 1**). Whilst most patients with MIBC died from the disease, the majority of patients
171 with NMIBC died from other causes (**Table 2**); notably, >10% of patients originally
172 presenting with pTa tumours, and >27% of patients originally presenting with pT1 tumours,
173 subsequently died from UBC.

174

175 Gender

176 There was no difference in grade at presentation between the genders ($p = 0.16$). However,
177 females presented more commonly with MIBC (30% vs. 26% for males) and less frequently

178 with pT1 disease (18% vs. 24% for males) ($p=0.06$, **Table 3**). Females had a longer total delay
179 time than males (median 120days vs. 106days, respectively, $p=0.02$) (**Tables 3 and 4**). The
180 majority of this delay arose before hospital referral (**Table 3**); a significantly higher
181 proportion of female patients with visible haematuria encountered a longer delay Time 1
182 than equivalent male patients ($p<0.05$, **Table 4**).

183

184 Female patients with MIBC had a significantly higher cumulative incidence of death from
185 UBC than male patients at 17 years (80% vs. 67%, $p<0.02$). There was no difference in UBC
186 mortality between the genders for pTa and pT1 tumours (14% vs. 15%, $p=0.56$ and 34% vs.
187 38%, $p=0.58$, respectively) (**Figure 2a**), and for other causes of death. Female patients with
188 grade 3 tumours had a significantly higher cumulative incidence of UBC death than males
189 (73% vs. 58%, $p=0.002$), but no difference was seen for grade 1 and 2 (14% vs. 15%, $p=0.65$
190 and 32% vs. 38% $p=0.23$, respectively) (**Figure 2b**).

191

192 Cigarette Smoking

193 At presentation, 77% of patients were current or previous smokers (**Table 1**). As observed in
194 the general population at the time, more men smoked (84%) than women (55%, $p<0.001$,
195 **Table 4**). Based on an age or date of stopping smoking obtained for all previous smokers,
196 the median duration of smoking cessation was 16 years (mean 18.8 years). In univariate
197 analysis there was a trend for cigarette smoking to be associated with increased cumulative
198 incidence of death due to UBC, death from other cancers, and death from other causes, but
199 none reached significance (Supplementary Figure 1a).

200

201 Occupational Carcinogen Exposure

202 We identified that 27% of patients had worked in occupations linked with UBC, with higher
203 exposure in males (31%) than females (14%, $p < 0.0001$, **Table 3**). There was a trend for
204 occupational exposure to be associated with increased cumulative incidence of death due to
205 UBC and death due to other causes, but none reached significance (Supplementary Figure
206 1b).

207

208 Pathway delays

209 The median time from initial onset of symptoms to GP referral was 14days (Time 1, IQR: 0-
210 61), from referral to hospital consultation was 28days (Time 2, IQR: 7-61), and from
211 consultation to first treatment was 20days (Time 3, IQR: 0-50). Patient characteristics by
212 delay times are shown in **Table 4**. Longer delays in Time 3 and Hospital Delay were
213 associated with smaller tumour size ($p < 0.05$ for both). A longer Total Delay was seen for
214 females (120days vs. 106days, $p < 0.05$) and non-smokers (118days vs. 105days, $p < 0.05$)
215 when compared to other patients.

216

217 Analysis of survival by delay stratified for tumour stage demonstrated no impact
218 (Supplementary Table 2), except for patients with MIBC for whom a shorter delay Time 3
219 resulted in worse survival compared to those with longer delay ($p < 0.05$).

220

221 Predictors of Survival

222 Univariate analysis identified histopathological criteria, gender, delays and smoking as
223 factors associated with UBC outcomes. Since these parameters are not necessarily

224 independent, multivariate analysis was used to determine the impact of each feature. Cox
225 regression of delay times adjusted by a base model of independent factors identified that
226 age, smoking status, and tumour stage (MIBC), grade (3), and size (>2cm) were the most
227 significant determinants of poor outcome from UBC (Supplementary Table 3). There was no
228 significant influence of delay, gender or occupational exposure.

229

230

231 **DISCUSSION**

232 Here we report 17-year outcomes from newly-diagnosed cases of UBC within a large
233 geographic region in the UK. We have updated an initial report [9], and now have most
234 cases (75%) followed until death. We are thus able to examine the complex interaction
235 between the tumour, patient gender, carcinogen exposures, pathway delays, and mortality.
236 We identified in univariate and competing risks analysis that many of these factors were
237 associated with disease-specific mortality. However, multivariate analysis identified age,
238 smoking status, and tumour stage, grade, and size as the most significant determinants of
239 poor outcome from UBC. Notably, there was no significant influence of delay, gender or
240 occupational exposure.

241

242 Comparison of CIFs demonstrated significant associations between female gender and
243 higher cumulative incidence of death from grade 3 disease and MIBC, concurring with
244 previous reports of worse outcomes for females with UBC [1;16;17;23;24]. Furthermore,
245 there was a trend for female patients to present more commonly with MIBC than male
246 patients. Importantly, female patients experienced a significantly longer Total Delay than
247 male patients. The majority of this delay occurred in Time 1, before referral for investigation
248 in secondary care was implemented by GPs; a significantly higher proportion of female
249 patients with visible haematuria encountered longer delays in Time 1 than equivalent male
250 patients. These data support observations of repeated community-based treatments for
251 suspected urinary infection in symptomatic females [25;26]. As reported by Hollenbeck et
252 al. [7], there were no significant differences in delays between the genders once patients
253 were within secondary care.

254 Female patients with MIBC had a significantly higher cumulative incidence of death from
255 UBC than male patients; it is unlikely that differential utilisation of radiotherapy or
256 cystectomy between the genders would cause this effect, but there is limited evidence to
257 suggest that female patients have worse outcomes from radiotherapy compared to males
258 [27]. However, such effects were not large enough for gender to be an independent
259 prognostic factor in multivariate analysis when adjusted for pathway delays.

260

261 It is a commonly-held belief that more rapid cancer diagnosis and treatment leads to better
262 outcomes, and to suggest otherwise is counterintuitive [5;6]. However, the relationship
263 between delay and survival in UBC is complex [8;9], with no direct linear relationship with
264 any components of delay [6;9]. In the long-term follow-up of this cohort, we have confirmed
265 this complex relationship, as noted by others [6;8;9]: no delay category had a significant
266 influence on survival, except for patients with MIBC for whom a shorter Delay 3 was
267 detrimental. This may represent an anomaly, and there is no clear explanation from our
268 data, but it is feasible that patients with MIBCs with concerning features (e.g. ongoing
269 bleeding) or comorbidities were selected for expedited treatment [9], subsequently
270 succumbing more rapidly as a result of those features or comorbidities. This was also
271 postulated by Liedberg, who demonstrated that a long treatment delay had no influence on
272 survival following cystectomy [8]. Seemingly, once in secondary care, clinicians are good at
273 selecting the highest risk patients and treating them rapidly [9]. Similarly, Nielsen
274 demonstrated that delay from TURBT to radical cystectomy was not independently
275 associated with stage progression or decreased recurrence-free or disease-specific survival
276 [28]. Likewise, for UBC patients treated by radiotherapy, there is no significant influence of

277 treatment delay on survival [29]. However, Hollenbeck investigated delay and survival in
278 29,826 patients with UBC and demonstrated that longer delays from presentation to
279 diagnosis were associated with increased risk of bladder cancer-specific mortality [7], a
280 finding also demonstrated by Gore when assessing the interval between TURBT and
281 cystectomy [30].

282

283 Many patient-related factors analysed here are not independent, e.g. males are more likely
284 to smoke, to have occupational carcinogen exposure, and to be more rapidly referred for
285 the investigation of haematuria [15]. Females are more likely to be non-smokers, are
286 typically exposed to different occupational carcinogens, and are slower to be referred for
287 investigation of haematuria or lower urinary tract symptoms [15;25;26]. In multivariate
288 analysis, we identified that only tumour stage, grade and size, and patient age and smoking
289 exposure were predictors of outcome when adjusted for pathway delays. These reinforce
290 observations from RCTs of bladder cancer treatment, and suggest that gender-related
291 disparities arise at least partly from a disease stage/grade migration due to diagnostic delay.

292

293 A major limitation of this study is a lack of delay data 'downstream' from TURBT. However,
294 the classification and interpretation of such data could be challenging (e.g. classifying the
295 time to definitive treatment of MIBC in the setting of chemoradiotherapy or neoadjuvant
296 chemotherapy/cystectomy), whereas TURBT remains the first intervention for all cases of
297 UBC [11;12]. It could also be suggested that our outcome data are not applicable to modern
298 practice (although outcomes from UBC have remained unchanged for over 30 years [3]), for
299 example, there appear to be high rates of UBC-specific death for patients with Ta and T1

300 tumours; in 1991 these patients may have been understaged and undertreated in an era
301 when re-TUR was rare and the utilisation of intravesical therapies was uncommon.
302 Furthermore, disease surveillance was according to national guidelines and may have
303 limited generalisability for other healthcare systems. Given the nature of multicentre cohort
304 studies, there was also likely to be heterogeneity in both treatment and surveillance
305 strategies between participating units. Finally, the gathering of more comprehensive
306 smoking and occupation data would have been more illuminating than the limited
307 categorical data presented here. The strengths of this study include its prospective nature,
308 its mature and long-term follow-up, and the completeness of data from a large cohort.

309

310

311 **CONCLUSIONS**

312 Our data demonstrate a stage migration to MIBC in female patients at presentation. The
313 relationships between gender, outcomes, delays and aetiology of UBC are complex. Female
314 patients experience a significantly longer Total Delay than male patients, the majority of
315 which results from a delay in referral from general practice to secondary care/urological
316 assessment, and may contribute to stage migration. GPs should be particularly vigilant
317 regarding symptoms that are associated with UBC, and especially in female patients; visible
318 haematuria always requires urgent referral to secondary care for urological assessment.

319

320

321 **LEGENDS FOR TABLES AND FIGURES**

322

323 **Table 1:** Overall patient and tumour characteristics (where recorded).

324

325 **Table 2:** Certified causes of death by tumour stage in the 983 patients where both tumour
326 stage and cause of death were known.

327

328 **Table 3:** Gender-specific characteristics of patients in the cohort.

329

330 **Table 4:** Patient characteristics by delay times (in days), *n* (%). VH=visible haematuria;
331 NVH=non-visible haematuria.

332

333 **Figure 1:** Survival by tumour stage and estimated survival for the general population.

334

335 **Figure 2a:** Cumulative incidence of death due to bladder cancer, other cancer and other
336 causes by gender and tumour stage (solid lines=male patients, dashed lines=female
337 patients).

338

339 **Figure 2b:** Cumulative incidence of death due to bladder cancer, other cancer and other
340 causes by gender and tumour grade (solid lines=male patients, dashed lines=female
341 patients).

342

343

344

345 **Table 1:** Overall patient and tumour characteristics (where recorded).

Variable	Number (%)
Gender (1478 responses, 100%)	
Male	1097 (74)
Female	381 (26)
Haematuria at presentation (1171 responses, 79%)	
Visible	1021 (87)
Non-visible	67 (6)
None	83 (7)
Age, years (1478 responses, 100%)	
<60	315 (21)
61-70	478 (32)
71-80	495 (33)
>80	190 (13)
Smoking history (1260 responses, 85%)	
Current smoker	330 (26)
Previous smoker	643 (51)
Never smoked	287 (23)
Occupational exposure (1240 responses, 84%)	
Known or suspected increased relative risk	330 (27)
No increased relative risk	910 (73)
Tumour type (1404 responses, 95%)	
Papillary	903 (64)
Solid	246 (18)
Mixed	255 (18)
Tumour number (1392 responses, 94%)	
Single	1042 (75)
2 or more	350 (25)
Tumour size, cm (1366 responses, 92%)	
≤2	552 (40)
>2	814 (60)
Tumour stage (1300 responses, 88%)	
pTa	658 (51)
pT1	291 (22)
T2-T4	351 (27)
Grade (1347 responses, 91%)	
Well (G1)	475 (35)
Moderate (G2)	513 (38)
Poor and anaplastic (G3)	359 (27)

346 **Table 2:** Certified causes of death by tumour stage in the 983 patients where both tumour stage and cause of death were known.

Cause	pTa (%)*					pT1 (%)*					T2-T4 (%)*	Total
	Well (G1)	Moderate (G2)	Poor and anaplastic (G3)	Unknown	All grades	Well (G1)	Moderate (G2)	Poor and anaplastic (G3)	Unknown	All grades	All grades	
Bladder Cancer	31 (12)	30 (19)	5 (36)	1 (09)	67 (15)	11 (38)	39 (34)	24 (35)	5 (42)	79 (35)	219 (69)	365
Other Cancer	65 (25)	31 (20)	2 (14)	2 (18)	100 (23)	5 (17)	12 (11)	10 (14)	2 (17)	29 (13)	24 (8)	153
Other Causes	165 (63)	94 (61)	7 (50)	8 (73)	274 (62)	13 (45)	63 (55)	35 (51)	5 (42)	116 (52)	75 (24)	465
Total	261	155	14	11	441	29	114	69	12	224	318	983

*Rounded proportions may sum to more than 100%

347

Table 3: Gender-specific characteristics of patients in the cohort.

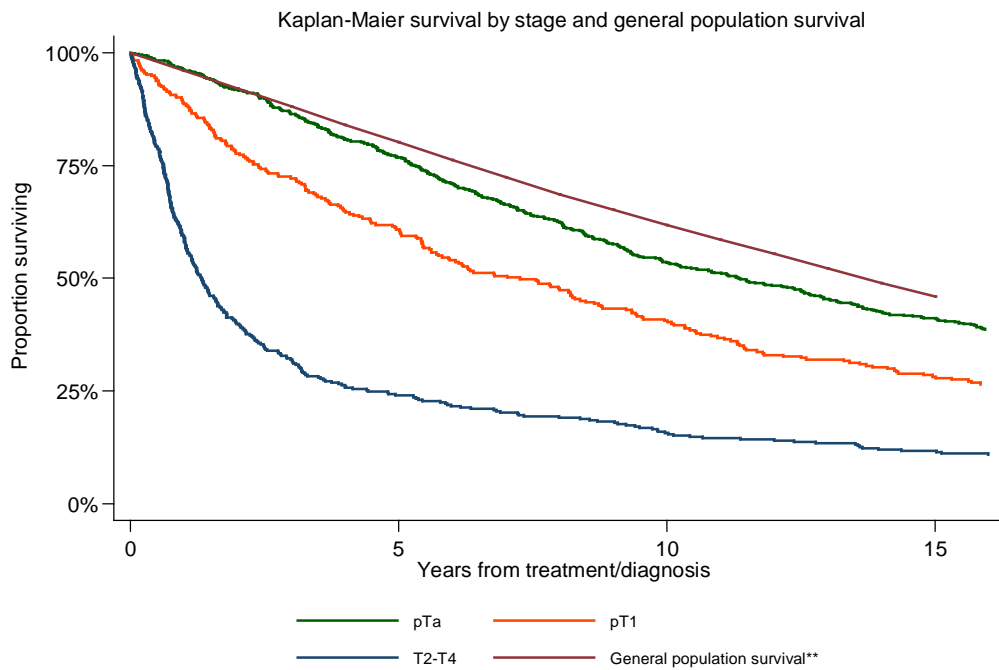
	Males	Females	p
Proportions	74%	26%	<0.001
Haematuria at presentation			
Visible	87%	89%	0.673
Non-visible	6%	5%	
None	7%	6%	
Grade			
G1	34%	39%	0.157
G2	39%	35%	
G3	27%	26%	
Stage			
Ta	50%	52%	0.060
T1	24%	18%	
T2-4	26%	30%	
Median Delay Time 1			
Delay 1 ≤14 days	51%	46%	0.101
Delay 1 >14 days	49%	54%	
Median Delay Time 2			
Delay 2 ≤28 days	27	29	0.498
Delay 2 >28 days	52%	50%	
Median Delay Time 3			
Delay 3 ≤20 days	22	18	0.152
Delay 3 >20 days	48%	52%	
Median Hospital Delay			
Hospital Delay ≤68 days	68	68	0.848
Hospital Delay >68 days	51%	51%	
Median Total Delay			
Total Delay ≤110 days	106	120	0.024
Total Delay >110 days	52%	45%	
History of smoking			
	48%	55%	<0.001
History of occupational exposure			
	31%	14%	<0.001

Table 4: Patient characteristics by delay times (in days), *n* (%).

Factor	Grouping	N	Time 1: Initial symptom to GP referral		Time 2: GP referral to first consultation		Time 3: Consultation to first Treatment		Hospital Delay		Total Delay	
			≤14	>14	≤28	>28	≤20	>20	≤68	>68	≤110	>110
Median age, years			70	69	69	70	70	69	70	69	69	70
IQR			62-77	61-76	61-76	62-77	62-77	62-76	62-76	62-76	62-76	62-76
Sex	Male	1097	548 (76)	523 (72)	553 (75)	518 (73)	511 (73)	560 (76)	549 (74)	525 (74)	558 (77)*	513 (72)*
	Female	381	171 (24)	199 (28)	184 (25)	187 (27)	193 (27)	178 (24)	188 (26)	184 (26)	168 (23)	203 (28)
Tumour stage	pTa	658	340 (54)	314 (48)	345 (52)	309 (50)	312 (51)	342 (51)	321 (49)	333 (52)	320 (50)	334 (52)
	pT1	291	140 (22)	149 (23)	140 (21)	149 (24)	131 (21)	158 (24)	148 (23)	142 (22)	146 (23)	143 (22)
	T2-T4	351	154 (24)	186 (29)	176 (27)	164 (26)	168 (27)	172 (26)	181 (28)	161 (25)	179 (28)	161 (25)
Tumour size, cm	≤2	552	286 (42)	259 (38)	272 (39)	274 (41)	244 (37)	302 (43)	247 (36)	301 (45)	259 (38)	287 (42)
	>2	814	388 (58)	419 (62)	417 (61)	390 (59)	412 (63)*	395 (57)*	439 (64)*	370 (55)*	417 (62)	390 (58)
Presenting with haematuria	VH (M)	755	301 (78)*	445 (72)*	452 (75)	149 (73)	259 (73)	487 (75)	405 (73)	343 (75)	390 (76)	356 (72)
	VH (F)	266	86 (22)	174 (28)	149 (25)	111 (27)	98 (27)	162 (25)	148 (27)	113 (25)	124 (24)	136 (28)
	NVH (M)	52	26 (79)	25 (76)	22 (76)	29 (78)	16 (76)	35 (78)	26 (76)	25 (78)	23 (74)	28 (80)
	NVH (F)	15	7 (21)	8 (24)	7 (24)	8 (22)	5 (24)	10 (22)	8 (24)	7 (22)	8 (26)	7 (20)
	None (M)	64	28 (82)	35 (73)	34 (69)	29 (88)	28 (65)	35 (90)	33 (73)	31 (82)	29 (74)	34 (79)
	None (F)	19	6 (18)	13 (27)	15 (31)	4 (12)	15 (35)*	4 (10)*	12 (27)	7 (18)	10 (26)	9 (21)
Smoking	Never	287	133 (21)	147 (24)	142 (22)	138 (23)	140 (23)	140 (22)	138 (22)	142 (23)	126 (20)	154 (25)
	Ever	973	486 (79)	471 (76)	490 (78)	467 (77)	459 (77)	498 (78)	492 (78)	467 (77)	500 (80)*	457 (75)*

* $P < 0.05$

Figure 1: Survival by tumour stage and estimated survival for the general population.



** General population survival for pTa tumours, matched by year of diagnosis, sex and age

Figure 2a: Cumulative incidence of death due to bladder cancer, other cancer and other causes by gender and tumour stage (solid lines=male patients, dashed lines=female patients).

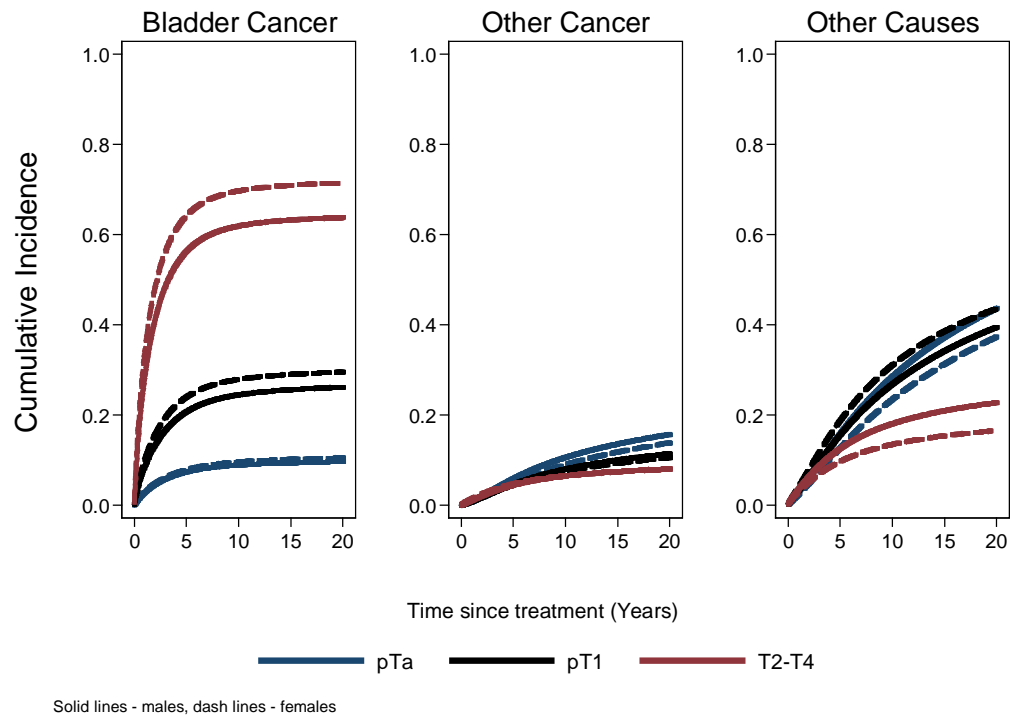
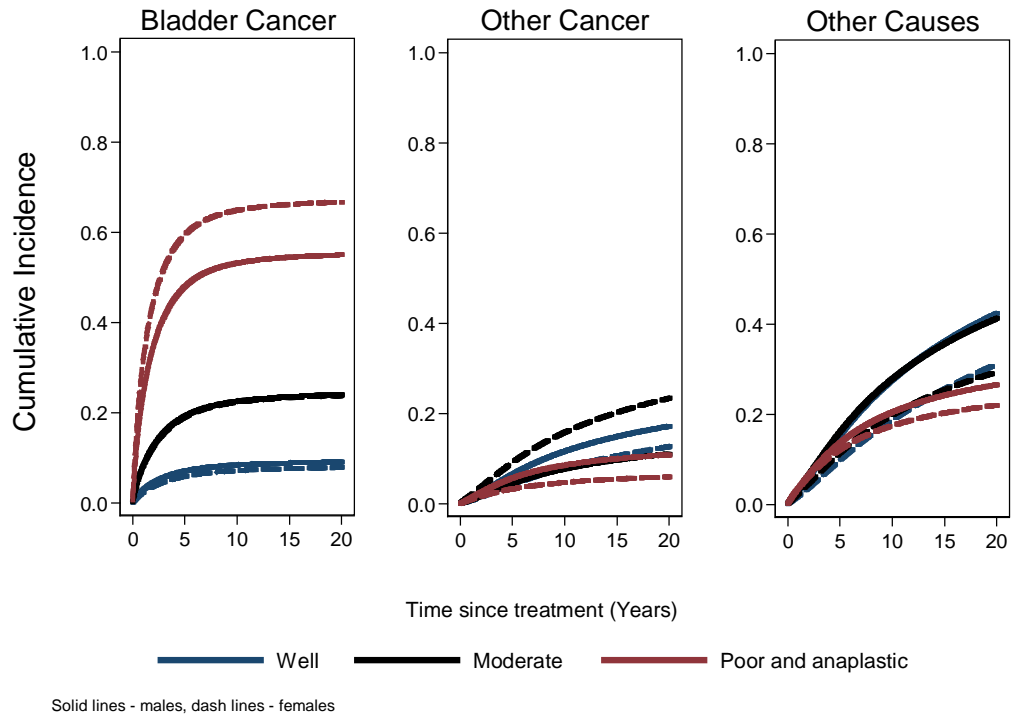


Figure 2b: Cumulative incidence of death due to bladder cancer, other cancer and other causes by gender and tumour grade (solid lines=male patients, dashed lines=female patients).



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A COMPARATIVE ANALYSIS OF GENDER, PATHWAY DELAYS AND RISK FACTOR EXPOSURES

ON THE LONG-TERM OUTCOMES FROM BLADDER CANCER

RT Bryan*, T Evans*, J Dunn, G Iqbal, S Bathers, SI Collins, ND James,

JWF Catto⁺, DMA Wallace

SUPPLEMENTARY METHODS & DATA

SUPPLEMENTARY METHODS

To assess the competing risks of death in our cohort, we first used a non-parametric test to assess the equality between groups by calculating the cumulative incidence function (CIF) as described by Scrucca et al. [1]. Comparison of specific CIFs was performed using Gray's test [2]. We then extended our analysis to investigate the effects of other covariates (stage at diagnosis, tumour grade, gender, smoking status, occupational exposure risk and age group at diagnosis), present in our data on the CIF. We constructed flexible parametric models using the user written Stata command `stpm2` in order to calculate the cause-specific hazard for each cause and for each covariate of interest [3]. Gender and cause of death were modeled as time-varying covariates. We used information from the Akaike Information Criterion (AIC) and Bayesian Information Criterion (BIC) for model selection. Post-estimation we applied the user written `stpm2cif` command [4], so that the cumulative incidence function for each model of interest could be derived and graphed.

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LEGENDS

Supplementary Table 1: Occupations with known or suspected exposure to urothelial carcinogens, and chemicals implicated in urothelial carcinogenesis. These data were utilised by the assessors to assign risk of occupational exposure.

Supplementary Table 2: Survival by delay times stratified for tumour stage.

Supplementary Table 3: Cox regression of delay times adjusted by a base model of independent factors.

Supplementary Figure 1a: Cumulative incidence of death due to bladder cancer, other cancer and other causes by smoking status (blue line=non-smoker, red line=current or ex-smoker).

Supplementary Figure 1b: Cumulative incidence of death due to bladder cancer, other cancer and other causes by risk of occupational exposure to bladder carcinogens (blue line=known or suspected risk of occupational exposure, red line=no increased risk of occupational exposure).

Supplementary Table 1: Occupations and carcinogens associated with urothelial carcinogenesis.

Occupations with exposure to urothelial carcinogens	Occupations with suspicion of an excess of bladder cancer	Implicated chemicals
Manufacture of rubber & rubber products	Leather working	1-Naphthylamine
Cable manufacturing industry	Manufacture and use of paint	2-Naphthylamine
Manufacture of dyestuffs	Plastics industry	3,3'-Dichlorobenzidine
Manufacture of organic chemicals	Medical and nursing	3,3'-Dichlorobenzidine hydrochloride
Gasworks, coke oven and iron foundry working	Textile printing and dyeing	3,3'-Dimethoxybenzidine (o-Dianisidine)
Rodent extermination	Hairdressing	3,3'-Dimethylbenzidine (o-Tolidine)
Sewage works	Aluminium refining and smelting	4,4'-Methylene bis (2-Chloroaniline)
Manufacture of firelighters/patent fuels	Security printing	4,4'-Methylenedianiline (MDA)
Laboratory work	Mechanics and land transport working	4-Aminobiphenyl
	Machine turning	4-Chloro-o-toluidine (4-COT)
		Aniline
		Auramine
		Benzidine
		Benzidine dihydrochloride
		Benzidine hydrochloride
		Benzidine sulphate
		Magenta
		Phenyl 1-naphthylamine
		Phenyl b-naphthylamine

Supplementary Table 2: Survival by delay times stratified for tumour stage.

DELAY, N DAYS	N	Dead (N)	%Alive	O/E	Median [95% CI] survival, years	Surviving at years				
						1	3	5	10	
Time 1 (P=0.09)										
≤14	719	524	27.1%	0.97	7.8 [6.7,8.7]	86%	70%	61%	42%	
>14	722	554	23.3%	1.03	6.4 [5.6,7.5]	83%	67%	58%	39%	
Time 1 by tumour stage (P=<0.001)										
pTa										
≤14	340	220	35.3%	0.86	11.8 [9.5,13.4]	97%	87%	78%	55%	
>14	314	212	32.5%	0.90	10.8 [9.1,12.6]	96%	86%	75%	52%	
pT1										
≤14	140	102	27.1%	0.97	8.0 [6.1,10.4]	90%	76%	65%	42%	
>14	149	120	19.5%	1.08	5.7 [4.8,9.3]	88%	69%	58%	40%	
T2-T4										
≤14	154	140	9.1%	1.21	1.3 [1.0,1.7]	58%	31%	23%	15%	
>14	186	166	10.8%	1.19	1.3 [1.0,1.6]	59%	33%	25%	17%	
Time 2 (P=0.09)										
≤28	737	565	23.3%	1.03	6.2 [5.4,7.4]	83%	66%	57%	38%	
>28	705	513	27.2%	0.97	8.2 [7.0,9.0]	86%	71%	62%	43%	
Time 2 by tumour stage (P=<0.001)										
pTa										
≤28	345	238	31.0%	0.92	9.6 [8.6,11.5]	97%	85%	75%	49%	
>28	309	194	37.2%	0.84	12.9 [11.1,15.1]	96%	88%	79%	59%	
pT1										
≤28	140	109	22.1%	1.04	6.2 [4.4,10.0]	87%	70%	57%	42%	
>28	149	113	24.2%	1.01	7.8 [5.8,9.3]	91%	75%	65%	40%	

T2-T4										
≤28	176	160	9.1%	1.21	1.1	[0.9,1.5]	55%	30%	21%	14%
>28	164	146	11.0%	1.19	1.4	[1.1,2.1]	63%	34%	27%	18%
Time 3 (P=0.45)										
≤20	704	520	26.1%	0.99	6.8	[5.5,8.1]	82%	66%	56%	39%
>20	738	558	24.4%	1.01	7.5	[6.4,8.4]	87%	71%	62%	42%
Time 3 by tumour stage (P=<0.001)										
pTa										
≤20	312	208	33.3%	0.89	10.1	[9.0,12.8]	95%	84%	73%	51%
>20	342	224	34.5%	0.88	12.1	[10.0,13.4]	98%	89%	80%	56%
pT1										
≤20	131	90	31.3%	0.92	8.8	[5.8,11.4]	90%	76%	63%	47%
>20	158	132	16.5%	1.12	6.2	[5.0,8.2]	88%	70%	59%	35%
T2-T4										
≤20	168	155	7.7%	1.23	1.0	[0.8,1.2]	53%	26%	18%	12%
>20	172	151	12.2%	1.17	1.8	[1.3,2.4]	65%	38%	30%	20%
Hospital Delay (P=0.30)										
≤68	737	560	24.0%	1.02	6.1	[5.4,7.4]	82%	65%	56%	37%
>68	709	522	26.4%	0.98	8.1	[6.8,8.9]	87%	72%	62%	43%
Hospital Delay by tumour stage (P=<0.001)										
pTa										
≤68	321	221	31.2%	0.92	9.6	[8.4,11.6]	96%	84%	74%	49%
>68	333	211	36.6%	0.85	12.9	[11.1,14.1]	97%	89%	79%	58%
pT1										
≤68	148	112	24.3%	1.01	7.8	[5.0,10.0]	87%	72%	59%	43%
>68	142	111	21.8%	1.04	7.0	[5.6,8.6]	90%	73%	63%	39%
T2-T4										
≤68	181	160	11.6%	1.18	1.1	[0.9,1.3]	55%	30%	23%	17%

>68	161	148	8.1%	1.23	1.7	[1.2,2.1]	64%	34%	26%	16%
Total Delay (P=0.26)										
≤110	726	552	24.0%	1.02	6.9	[5.6,7.9]	83%	66%	58%	39%
>110	716	526	26.5%	0.98	7.4	[6.4,8.6]	86%	71%	60%	41%
Total Delay by tumour stage (P=<0.001)										
pTa										
≤110	320	217	32.2%	0.91	10.5	[9.0,12.3]	96%	86%	76%	52%
>110	334	215	35.6%	0.86	12.5	[9.9,14.4]	96%	87%	78%	55%
pT1										
≤110	146	112	23.3%	1.03	8.4	[6.2,10.5]	88%	74%	66%	46%
>110	143	110	23.1%	1.03	5.7	[4.4,8.2]	90%	71%	57%	36%
T2-T4										
≤110	179	160	10.6%	1.19	1.1	[0.8,1.4]	54%	30%	22%	16%
>110	161	146	9.3%	1.21	1.6	[1.2,2.1]	64%	34%	26%	16%

Supplementary Table 3: Cox regression of delay times adjusted by a base model of independent factors.

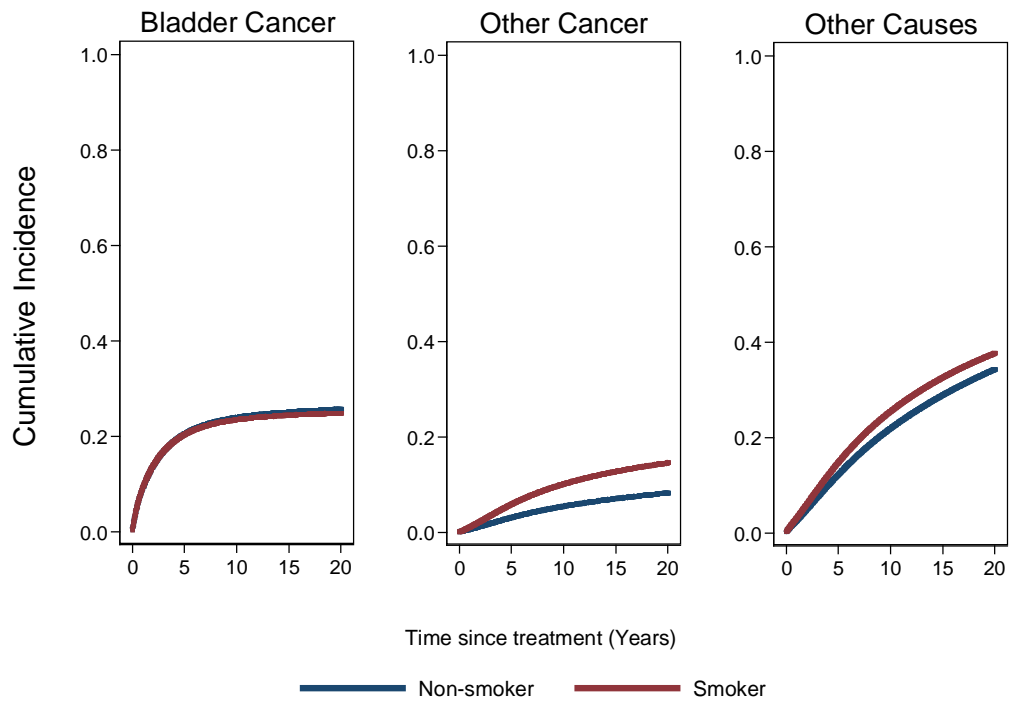
Factor	Grouping	Coefficient	z	P	Hazard ratio (95% CI)
Time 1 adjusted by base model (n=1001)					
Time 1	(≤14, >14 days)	0.10	1.34	0.18	1.11 (0.95,1.28)
Age	(Continuous)	0.06	13.98	<0.001	1.06 (1.05,1.07)
Tumour type					
	(Papillary)				1 (1,1)
	(Mixed)	0.24	1.67	0.09	1.27 (0.96,1.67)
	(Solid)	0.13	1.11	0.27	1.14 (0.91,1.43)
Tumour stage					
	(pTa)				1 (1,1)
	(pT1)	0.10	0.95	0.34	1.11 (0.90,1.36)
	(T2-T4)	0.50	3.43	<0.001	1.65 (1.24,2.20)
Smoking	(Never, ever)	0.36	3.72	<0.001	1.44 (1.19,1.74)
Occupational exposure	(No increased risk, known or suspect risk)	0.06	0.70	0.48	1.06 (0.90,1.26)
Sex	(Female, Male)	0.02	0.16	0.87	1.02 (0.84,1.22)
Tumour grade					
	(Well differentiated)				1 (1,1)
	(Moderately differentiated)	0.02	0.23	0.82	1.02 (0.85,1.23)
	(Poorly differentiated)	0.31	2.26	0.02	1.36 (1.04,1.77)
Tumour size (cm)	(≤2, >2)	0.15	1.90	0.06	1.17 (0.99,1.36)
Time 2 adjusted by base model (n=1001)					
Time 2	(≤28, >28 days)	-0.11	-1.44	0.15	0.90 (0.78,1.04)
Age	(Continuous)	0.06	13.89	<0.001	1.06 (1.05,1.07)
Tumour type					
	(Papillary)				1 (1,1)
	(Mixed)	0.24	1.72	0.09	1.27 (0.97,1.68)
	(Solid)	0.13	1.11	0.27	1.14 (0.90,1.43)

Tumour stage					
	<i>(pTa)</i>				1 (1,1)
	<i>(pT1)</i>	0.11	1.03	0.31	1.11 (0.91,1.37)
	<i>(T2-T4)</i>	0.52	3.58	<0.001	1.69 (1.27,2.24)
Smoking	<i>(Never, ever)</i>	0.36	3.69	<0.001	1.44 (1.19,1.74)
Occupational exposure	<i>(No increased risk, known or suspect risk)</i>	0.05	0.58	0.56	1.05 (0.89,1.24)
Sex	<i>(Female, Male)</i>	0.01	0.10	0.92	1.01 (0.84,1.21)
Tumour grade					
	<i>(Well differentiated)</i>				1 (1,1)
	<i>(Moderately differentiated)</i>	0.02	0.22	0.82	1.02 (0.85,1.23)
	<i>(Poorly differentiated)</i>	0.30	2.24	0.03	1.35 (1.04,1.76)
Tumour size (cm)	<i>(≤2, >2)</i>	0.15	1.89	0.06	1.16 (0.99,1.36)
Time 3 adjusted by base model (n=1001)					
Time 3	<i>(≤20, >20 days)</i>	-0.09	-1.15	0.25	0.92 (0.79,1.06)
Age	<i>(Continuous)</i>	0.06	13.91	<0.001	1.06 (1.05,1.07)
Tumour type					
	<i>(Papillary)</i>				1 (1,1)
	<i>(Mixed)</i>	0.24	1.71	0.09	1.27 (0.96,1.68)
	<i>(Solid)</i>	0.13	1.11	0.27	1.14 (0.91,1.43)
Tumour stage					
	<i>(pTa)</i>				1 (1,1)
	<i>(pT1)</i>	0.10	1.00	0.32	1.11 (0.90,1.36)
	<i>(T2-T4)</i>	0.51	3.50	<0.001	1.66 (1.25,2.21)
Smoking	<i>(Never, ever)</i>	0.37	3.73	<0.001	1.44 (1.19,1.75)
Occupational exposure	<i>(No increased risk, known or suspect risk)</i>	0.06	0.69	0.49	1.06 (0.90,1.25)
Sex	<i>(Female, Male)</i>	0.01	0.09	0.93	1.01 (0.84,1.21)
Tumour grade					
	<i>(Well differentiated)</i>				1 (1,1)
	<i>(Moderately differentiated)</i>	0.01	0.13	0.89	1.01 (0.84,1.22)

	<i>(Poorly differentiated)</i>	0.32	2.35	0.02	1.37 (1.05,1.79)
Tumour size (cm)	(≤2, >2)	0.15	1.87	0.06	1.16 (0.99,1.36)
<i>Hospital delay adjusted by base model (n=1003)</i>					
Hospital delay	(≤68, >68 days)	-0.06	-0.84	0.40	0.94 (0.81,1.09)
Age	(Continuous)	0.06	13.95	<0.001	1.06 (1.05,1.07)
Tumour type					
	<i>(Papillary)</i>				1 (1,1)
	<i>(Mixed)</i>	0.24	1.68	0.09	1.27 (0.96,1.67)
	<i>(Solid)</i>	0.13	1.09	0.28	1.14 (0.90,1.43)
Tumour stage					
	<i>(pTa)</i>				1 (1,1)
	<i>(pT1)</i>	0.11	1.04	0.3	1.12 (0.91,1.37)
	<i>(T2-T4)</i>	0.51	3.50	<0.001	1.66 (1.25,2.21)
Smoking	<i>(Never, ever)</i>	0.37	3.73	<0.001	1.44 (1.19,1.75)
Occupational exposure	<i>(No increased risk, known or suspect risk)</i>	0.06	0.66	0.51	1.06 (0.89,1.25)
Sex	<i>(Female, Male)</i>	0.01	0.14	0.89	1.01 (0.84,1.22)
Tumour grade					
	<i>(Well differentiated)</i>				1 (1,1)
	<i>(Moderately differentiated)</i>	0.02	0.21	0.83	1.02 (0.85,1.23)
	<i>(Poorly differentiated)</i>	0.32	2.35	0.02	1.37 (1.05,1.79)
Tumour size (cm)	(≤2, >2)	0.14	1.77	0.08	1.15 (0.98,1.35)
<i>Total delay adjusted by base model (n=1001)</i>					
Total delay	(≤110, >110 days)	0.01	0.18	0.86	1.01 (0.88,1.17)
Age	(Continuous)	0.06	13.92	<0.001	1.06 (1.05,1.07)
Tumour type					
	<i>(Papillary)</i>				1 (1,1)
	<i>(Mixed)</i>	0.24	1.70	0.09	1.27 (0.96,1.68)
	<i>(Solid)</i>	0.13	1.09	0.28	1.14 (0.90,1.43)
Tumour stage					

	<i>(pT_a)</i>			<0.001	1 (1,1)
	<i>(pT₁)</i>	0.11	1.02	0.31	1.11 (0.91,1.37)
	<i>(T₂-T₄)</i>	0.51	3.46	<0.001	1.66 (1.25,2.21)
Smoking	<i>(Never, ever)</i>	0.37	3.73	<0.001	1.44 (1.19,1.75)
Occupational exposure	<i>(No increased risk, known or suspect risk)</i>	0.06	0.70	0.48	1.06 (0.90,1.26)
Sex	<i>(Female, Male)</i>	0.01	0.12	0.91	1.01 (0.84,1.22)
Tumour grade					
	<i>(Well differentiated)</i>				1 (1,1)
	<i>(Moderately differentiated)</i>	0.02	0.19	0.85	1.02 (0.85,1.23)
	<i>(Poorly differentiated)</i>	0.31	2.29	0.02	1.36 (1.05,1.78)
Tumour size (cm)	<i>(≤2, >2)</i>	0.16	1.92	0.05	1.17 (1.00,1.37)

Supplementary Figure 1a: Cumulative incidence of death due to bladder cancer, other cancer and other causes by smoking status (blue line=non-smoker, red line=current or ex-smoker).



Supplementary Figure 1b: Cumulative incidence of death due to bladder cancer, other cancer and other causes by risk of occupational exposure to bladder carcinogens (blue line=known or suspected risk of occupational exposure, red line=no increased risk of occupational exposure).

