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# Journal Pre-proof

Collateral damage: the impact on outcomes from cancer surgery of the COVID-19 pandemic

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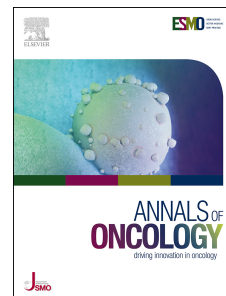
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1 **Collateral damage: the impact on outcomes from cancer surgery of the**  
2 **COVID-19 pandemic**

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38  
39

40 **ABSTRACT**

41

42 **Background:** Cancer diagnostics and surgery have been disrupted by the response of  
43 healthcare services to the COVID-19 pandemic. Progression of cancers during delay will  
44 impact on patient long-term survival.

45 **Methods:** We generated per-day hazard ratios of cancer progression from observational  
46 studies and applied these to age-specific, stage-specific cancer survival for England 2013-  
47 2017. We modelled per-patient delay of three months and six months and periods of  
48 disruption of one year and two years. Using healthcare resource costing, we contextualise  
49 attributable lives saved and life-years gained from cancer surgery to equivalent volumes of  
50 COVID-19 hospitalisations.

51 **Findings:** Per year, 94,912 resections for major cancers result in 80,406 long-term survivors  
52 and 1,717,051 life years gained. Per-patient delay of three/six months would cause  
53 attributable death of 4,755/10,760 of these individuals with loss of 92,214/208,275 life-  
54 years. For cancer surgery, average life-years gained (LYGs) per patient are 18.1 under  
55 standard conditions and 17.1/15.9 with a delay of three/six months (an average loss of  
56 0.97/2.19 LYG per patient). Taking into account units of healthcare resource (HCRU), surgery  
57 results on average per patient in 2.25 resource-adjusted life-years gained (RALYGs) under  
58 standard conditions and 2.12/1.97 RALYGs following delay of three/six months. For 94,912  
59 hospital COVID-19 admissions, there are 482,022 LYGs requiring of 1,052,949 HCRUs.  
60 Hospitalisation of community-acquired COVID-19 patients yields on average per patient 5.08  
61 LYG and 0.46 RALYGs.

62 **Interpretation:** Modest delays in surgery for cancer incur significant impact on survival.  
63 Delay of three/six months in surgery for incident cancers would mitigate 19%/43% of life-  
64 years gained by hospitalisation of an equivalent volume of admissions for community-  
65 acquired COVID-19. This rises to 26%/59% when considering resource-adjusted life-years  
66 gained. To avoid a downstream public health crisis of avoidable cancer deaths, cancer  
67 diagnostic and surgical pathways must be maintained at normal throughput, with rapid  
68 attention to any backlog already accrued.

69

70 **KEY WORDS**71 **Oncology, Survival, Delay, COVID-19, Diagnostics**

72 **INTRODUCTION**

73 Following the first case reports in Hubei province, China in late 2019, a pandemic of COVID-  
74 19 coronavirus was declared by the World Health Organisation in March 2020. Whilst  
75 COVID-19 causes minimal or mild illness in most, a small but appreciable proportion of  
76 individuals require oxygen therapy and often admission to an Intensive Care Unit (ICU). The  
77 ensuing unprecedented pressure on hospital wards and ICUs has necessitated rapid  
78 redeployment of staff and capacity towards the management of COVID-19 cases with  
79 deprioritisation of non-emergency clinical services, including diagnostics and elective  
80 specialist surgery. Concurrently, lockdown of the population has impacted dramatically on  
81 presentation and referral of symptomatic patients from primary into secondary care[1].

82

83 For patients with cancer, delay of surgery has the real potential to increase the likelihood of  
84 metastatic disease, with some patients' tumours progressing from being curable (with near  
85 normal life expectancy) to non-curable (with limited life expectancy)[2]. The situation has  
86 been further exacerbated by recent safety concerns regarding aerosol generation from  
87 endoscopy, cystoscopy and surgery[3, 4].

88

89 Current projections indicate that COVID-19-related disruption may well last for 18 months  
90 or more, until there is either long term effective containment in the population or large-  
91 scale vaccination. To inform healthcare prioritisation and resource allocation, we have  
92 examined the impact on cancer outcomes of different periods of delay of cancer surgery  
93 with disruption extending over variable time periods, comparing resource-weighted  
94 outcomes to hospital management of COVID-19 patients.

95

## 96 METHODS

### 97 Data sources

98 Number and age-specific five-year net survival of cancer patients that had potentially  
99 curative surgical resections for non-haematological malignancies between 2013 and 2017  
100 were obtained from Public Health England National Cancer Registration Service (NCRAS)[5].  
101 As well as cancer stage at diagnosis for each cancer type, breast tumour receptor data  
102 allowed subtyping of these cancers as ER+ HER2-, HER+ (any), ER- HER2-, and other.  
103 Estimates for nosocomial infection rates, median duration of hospital stay for each cancer  
104 type, staffing of theatres, ICU and surgical wards were based on information from three  
105 large UK surgical oncology centres. Patterns of administration of adjuvant systemic anti-  
106 cancer therapy (SACT) were based on oncologist-reviewed standard practice guidance[6].  
107 ICU COVID-19 mortality, distribution by age, and duration of stay and proportion referred  
108 into ICU were obtained from ICNARC and data from hospitalised UK cases[7, 8]. Due to lack  
109 of UK data, data from Wuhan was used as the basis for the age distribution of community  
110 infection, age-specific likelihoods of admission from community to hospital, and mortality  
111 rates for non-ICU COVID-19 patients [9, 10] (Supplementary Table 1).

112

### 113 Analysis

#### 114 Impact of COVID-associated delay on cancer outcomes

115 We used published data from studies examining the impact on overall survival from delay in  
116 cancer surgery to estimate per day hazard ratios (HRs) associated with delay for different  
117 cancers (the "Fatality HR") [11-21]. We had sufficient data to generate Fatality HRs for three  
118 tumour types and assigned these to other tumours, based on comparability of 5-year  
119 survival as low (>90%) moderate (50-90%) or high (<50%) progressiveness tumours[5].  
120 Because we were unable to identify any suitable observational data for tumours of high  
121 progressiveness (e.g. oesophageal, gastric), we applied the Fatality HR from tumours of  
122 moderate progressiveness; this is likely to be a conservative assumption (Supplementary  
123 Table 2).

124 By accounting for COVID-related post-surgical mortality and changes in SACT, we adjusted  
125 five-year net survival figures for each cancer for surgical patients under *standard* care to  
126 estimate *current* five-year net survival. To model outcomes of surgery *post-delay*, we apply  
127 to standard five-year net survival, the Fatality HR relating to the specified number of days of

128 delay, again including COVID-related post-surgical mortality. Based on estimates from a UK  
129 surgical oncology centre, supported by the literature, we applied a current per day rate of  
130 nosocomial infection of 5%. Assuming improvement in cold protocols, we modelled  
131 reduction in this rate over time. We estimated COVID-associated surgical mortality based on  
132 per day rate of nosocomial infection, operation-specific duration of post-surgical admission,  
133 and age-specific mortality from infection. We estimated COVID-19 associated mortality for  
134 SACT administration, based on per day rate of nosocomial infection, the frequency of SACT  
135 scheduling, increased risk associated with immunosuppression, and age-specific mortality  
136 from infection. We assumed, where standard-of-care, that SACT offers a uniform survival  
137 benefit (5% in Stage 1, 7.5% in Stage 2 and 10% in Stage 3) and administration would only  
138 continue where this benefit exceeds COVID-related mortality.

139 We used mean life-expectancies per 10-year age-group to calculate life years gained,  
140 averaged per patient. We examined reduction in overall survival and life years gained (LYG),  
141 comparing surgery under standard care, current conditions and post-delay, by cancer type  
142 and by age and stage. Using 2013-2017 surgical workload data, we calculated across all  
143 adult cancers examined, the total number of deaths and life years lost attributable to delay.  
144 To address possible scenarios, we considered per-patient delay of up to six-months, and 1-  
145 and 2-year periods of disruption.

146

#### 147 **COVID-19 outcome**

148 To compare life years associated with timely cancer surgery with that afforded by  
149 hospitalisation of COVID-19 patients, we modelled a volume of community-ascertained  
150 COVID-19 infection resulting in an equivalent volume of hospital admissions to cancer  
151 surgeries (**Supplementary Table 1**).

152

#### 153 **Resource**

154 We analysed healthcare resource units (HCRU) focused specifically on frontline medical and  
155 nursing staff, where one HCRU is one 12-hour shift of direct nursing or medical care. We up-  
156 weighted for shifts from healthcare workers of high-salary (senior doctors) and/or of current  
157 scarcity (anaesthetists, ICU nurses). We calculated HCRUs per patient using estimated  
158 staffing ratios for theatres, ICU and ward care and operation-specific data for theatre hours,  
159 ICU stay and ward days from oncology centres.

160 Details of assumptions and parameter estimates are detailed in **Table 1** and **Supplementary**  
161 **Table 1**. Analyses were performed using STATA (version 15) and transcribed to Excel, to  
162 provide a full visibility of parametrisation, model outputs, and opportunity for the reader to  
163 customise parameters (**Supplementary Materials**).

164

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## 166 RESULTS

### 167 Impact of surgical delay on survival for different cancers

168 The greatest rates of deaths arise following even modest delays to surgery in aggressive  
169 cancers, with over 30% reduction in survival at six months and over 17% reduction in  
170 survival at three months for patients with stage 2 or 3 cancers of the bladder, lung,  
171 oesophagus, ovary, liver, pancreas and stomach (**Table 2, Supplementary Table 3,**  
172 **Supplementary Materials**). Accounting for nosocomial COVID-19 infection, for cancers with  
173 a relatively good overall prognosis, delay of surgery by three months had a minimal impact  
174 on survival: <1% for all Stage 1 ER+ and HER2+ breast cancers, for example. In older patients  
175 (>70 years), for early stage colorectal, kidney and ER+ breast cancers, the current impact on  
176 survival of COVID-related mortality exceeded the impact of three or even six months delay  
177 (**Table 2, Supplementary Table 3**).

178

179 For a high proportion of solid cancers, survival at five years is generally considered to be  
180 equivalent to cure. Predicated on this assertion, we considered life-years gained adjusting  
181 for resource (resource adjusted life years (RALYGs)). Perhaps unsurprisingly, most benefit is  
182 afforded in younger age groups for operations that are shorter with no associated ICU  
183 requirement. For example, trans-urethral resection of stage 1 bladder cancers affords on  
184 average 23.4 RALYG per patient age 30-39, whereas cystectomy for stage 2 bladder cancer is  
185 only associated with 1.2 RALYGs in that age group (**Supplementary Table 4**). In the context  
186 of prioritisation, avoidance of a six-month delay restitutes on average 4.1 RALYGs in the  
187 former group, compared to 0.7 in the latter (**Table 3, Supplementary Table 5**). Wide local  
188 excision for breast cancer has low resource requirement and therefore confers substantial  
189 RALYGs, even in good prognosis subtypes.

190

### 191 Impact of surgical delay on cancer survival combined across cancer types

192 Each year, 94,912 surgical resections for common invasive adult cancer types are performed  
193 in England, with 80,406 of those patients surviving their cancer at five years. A surgical delay  
194 of three months across all incident solid tumours over one year would incur 4,755 excess  
195 deaths, escalating to 10,760 excess deaths for a six-month delay. This includes at six  
196 months, attributable deaths of 2,980 for colorectal cancer 1,439 for lung cancer and 804 for  
197 breast cancer (**Figure 1**).

198

199 For a high proportion of solid cancers, five-year survival is generally considered to be  
200 equivalent to cure. Predicated on this assertion, across all cancers a delay of three months  
201 in treatment would lead to a reduction of 92,214 life-years and for six months' reduction of  
202 208,275 life years (**Table 3**). Prior to the COVID-19 crisis, each year cancer surgery was  
203 directly responsible for 1,717,051 LYGs. This represents on average 18.1 LYG per patient,  
204 which markedly reduces to 17.1 with three months' delay and to 15.9 with six months'  
205 delay. Cancer surgery per year requires 764,765 units of healthcare resource. Assuming this  
206 to be unchanged by delay, this affords on average 2.25 RALYG per patient under standard  
207 conditions, reducing to 2.12 with three months' delay and 1.97 with six months of delay, an  
208 average loss of 0.12 and 0.27 RALYGs, respectively, per patient.

209

### 210 **Resource comparison for outcomes afforded by cancer surgery and COVID-19** 211 **management**

212 For contextualisation, we compare the impact of cancer surgery delay to hospital care for  
213 patients with community-acquired COVID-19 infection. COVID-19 ICU admission for those  
214 aged 40-49 yielded on average 27.5 LYG and 0.8 RALYG. Those aged >80 years admitted to  
215 ICU benefit by on average 2.1 LYG and 0.06 RALYG. For non-ICU admission, average benefit  
216 is 9.3 LYG and 1.5 RALYG for those aged 40-49 and 1.4 LYG and 0.2 RALYG for those aged  
217 >80 years (**Supplementary Materials**). These estimates are inherently conservative as they  
218 do not take into account the impact on life expectancy of the excess comorbidities  
219 associated with many hospitalised COVID-19 cases.

220

221 COVID-19 community-acquired infection of 683,083 individuals would result in 94,912  
222 hospital admissions (*i.e.* the equivalent number to number of annual admissions for cancer  
223 surgery). For these 94,912 admissions, 16,135 will require ICU (critical cases) and 78,777 will  
224 not require ICU (severe cases). 1,052,949 units of healthcare resource are required in total  
225 and there are 15,587 deaths, 25,752 attributable lives saved, and 482,022 attributable LYGs  
226 (8,241 deaths/7,894 attributable lives saved/223,227 LYGs for ICU admissions, 7,346/  
227 17,858/ 258,795 for non-ICU). This represents on average 5.08 LYG and 0.46 RALYG per  
228 hospitalised COVID-19 patient.

229

230 It is therefore noteworthy, that a delay of surgery by six months results in 208,275 lost life-  
231 years for an annual quota of surgical patients: this equates to 43% of the total 482,022 life-  
232 years gained from hospitalisation of an equivalent number of community-acquired COVID-  
233 19 cases. This rises to 59% when adjusted for differences in resource (RALYGs).

234

### 235 **Sensitivity Analysis**

236 The outcomes from the model were mostly sensitive to changes in the Fatality HR for the  
237 per-day delay: varying this by  $\pm 8\%$  (1SD) caused the average LYG with a six-month delay to  
238 range from 15.7-16.1, and attributable LY lost by 2.00-2.39. Sensitivity analysis for other  
239 parameters is shown in **Supplementary Table 2**.

240

241

**242 DISCUSSION**

243 We provide estimates derived from reported surgical outcomes to quantify the impact on  
244 survival of delay of cancer treatment, within the parameters of the assumptions of the  
245 model.

246

**247 Implications for healthcare planning**

248 For aggressive cancers, our analysis demonstrates that even short delays (three months)  
249 have a significant impact on patient survival. However, even for cancers of comparatively  
250 favourable prognosis, a delay of six months will result in significant summed attributable  
251 deaths as many of these cancers are common. Delay will also result in tumours being more  
252 advanced, meaning not only is survival poorer, but that the upstaged cancers will be more  
253 costly to treat both in terms of surgery and/or chemotherapy. Furthermore, resource  
254 requirements (for example, ICU stay) are dramatically higher for the many who will  
255 inevitably present as emergencies such as with obstruction, perforation or acute bleeding of  
256 the gastrointestinal tract[22].

257

258 Critical to mitigating cancer deaths is recognition that delay or bottleneck may arise at any  
259 point in the linear patient journey from (i) self-presentation of the symptomatic patient to  
260 primary care, (ii) primary care review and referral into secondary care (iii) diagnostic  
261 investigation, and (iv) surgery (or radiotherapy) with curative intent. Alongside any 'bulge' in  
262 accumulated cases will be the normal stream of incident cancer presentations. In the face of  
263 prolonged stress, it will be challenging to provide extra capacity to address these bulges  
264 alongside standard demands. In the short term, to avoid knock-on delays, immediate  
265 diversion of supra-normal resource volumes are required to process the backlog of cases  
266 that will have accrued in the initial months of the pandemic, in which referrals,  
267 investigations, and surgeries have been reduced by up to 80%[1]. In the medium-long term  
268 (over the next 3-24 months), avoidance of delay to cancer surgery should be of the highest  
269 priority: urgent attention is required to ensure sufficient resourcing for standard capacity of  
270 all pathway elements in primary care, cancer diagnostic, and surgical.

271

272 Delay in cancer surgery will have a highly deleterious health and economic impact. For the  
273 most part, the surgery will still be required (and may be more complex and costly) but

274 results in rapid diminution resultant life-years gained and resource-adjusted life-years.  
275 Comparing equivalent-sized hospital populations adjusted for resource, the health impact of  
276 delaying cancer surgery for six months will approximate 60% of health gains of  
277 hospitalizations for community acquired COVID-19 infection. We need to consider  
278 resourcing in the likely event of sizeable requirement for COVID-19 management for a  
279 sustained period of time, potentially up to two years. Although large facilities may be  
280 built/repurposed for COVID-19 management, these facilities are competing for the same  
281 fixed pool of healthcare workers that provide care for treating non-COVID-19 disease.

282

283 Currently, where the rate of nosocomial infection is high, for older groups in particular,  
284 surgery and/or SACT may in the short-term offer more risk than benefit (see Supplementary  
285 Materials). Active focus is required to establish 'cold' sections of the healthcare system,  
286 with rigorous protocols for staff screening and shielding protocols. This will serve to  
287 minimise nosocomial acquisition and mortality from COVID-19, to protect staff, and also to  
288 provide reassurance to the public regarding uptake of diagnostics and surgery for cancer.

289

290 Urgent review by professional bodies is required regarding best protection of their staffing  
291 groups, and guidance on surgical and diagnostic practice commensurate with the true  
292 risks[3].

293

#### 294 **Implications for prioritisation amongst cancer patients**

295 Given an accrued backlog of cases and ongoing tight competition for resources, decisions  
296 regarding surgical prioritisation may be required for a number of years, with capacity  
297 varying geographically and temporally. Recognising its limitations regarding assumptions  
298 and parameters, we propose a model that provides a rational approach by which to  
299 evaluate across patients of different ages, tumour types, and stages, the benefit and  
300 resource implications of their cancer surgery. We highlight in our model those age-stage  
301 groups for which COVID-related mortality currently exceeds survival benefit for surgery  
302 and/or SACT. Whilst these and other groups for whom benefit is marginal will be the most  
303 rationale to delay, they will nevertheless require monitoring and surgery downstream.  
304 Longitudinal planning, monitoring of progression, dynamic re-prioritisation, and capacity-  
305 planning will inevitably be highly challenging.

306

307

**308 Broader and International relevance**

309 While we have used data for England, cancer survival is broadly similar across most  
310 economically developed countries, so the impact of delay per tumour is broadly applicable  
311 across Europe. However, variation in incidence of cancer, life expectancy and population  
312 age structure mean that predictions regarding total case numbers and life-years gained and  
313 lost are more difficult to extrapolate, even when scaling for relative size of reference  
314 population.

315 Whilst customised for surgical delay due to the COVID-19 pandemic, this model could  
316 readily be adapted to quantify the impact of surgical delay due to other causes.

317

**318 Limitations**

319 As with any model-based analysis, our predictions are predicated on the validity of  
320 assumptions and estimates used for parameterisation. While we have made use of  
321 observational data, our approach simplifies the complexity of cancer progression and is  
322 solely survival-focused. For healthcare planning, a more elaborate model capturing stage-  
323 shifting may offer additional utility. We base our analysis on survival data from 2013-17; for  
324 some tumour types, standard-of-care and survival has evolved since this time. Our  
325 modelling of the benefit of SACT is simplistic as the scheduling, benefits and  
326 immunosuppressive consequences vary by chemotherapy regimen. Whilst we have included  
327 in our model the impact withholding of SACT if nosocomial infection risk is high, we have  
328 not modelled additional reduction in survival from delays in administration of adjuvant  
329 therapy. Mortality from nosocomial COVID-19 infection during surgical admission or  
330 attendance for chemotherapy is based on a uniform per-day risk of infection: these may  
331 vary between institutions. While our resourcing analysis deliberately focuses on the  
332 requirement for the direct medical and nursing staff who most limit healthcare provision,  
333 we acknowledge it does not capture other 'costs' incurred in hospital care, primary care,  
334 and social care.

335 Our model of COVID-19 admissions is limited by availability of detailed individual-level UK  
336 data, in particular for non-CCU hospital admissions; this model is also conservative in regard  
337 of disregarding impact of co-morbidities on life expectancy.

338

**339 Further research**

340 Within our current approach, we only estimate the effects of a specified period of per-  
341 patient delay. Contemporaneous data for NHS activity offers the prospect of developing  
342 dynamic models to predict the impact of (i) differential prioritisation of patient groups, (ii)  
343 different patterns of re-presentation of ‘accumulated’ cases alongside incident cases, and  
344 (iii) varying release of bottlenecks in primary care, diagnostics, and surgery. Evaluation is  
345 also important for the alternative management approaches being adopted, such as  
346 radiotherapy with curative intent where surgery is gold-standard or *a priori* hormonal  
347 treatment for prostate and ER-positive breast cancers. For any strategies involving  
348 deliberate delay to surgery, models for re-staging and dynamic re-prioritisation are  
349 essential. We have focused on the impact to surgery with curative intent; analyses are also  
350 required to quantify the impact on mortality of changes to life-extending chemo- and radio-  
351 therapy for patients with Stage 4 disease.

352

**353 CONCLUSION**

354

355 Compared to COVID-19 management, cancer surgery is highly impactful in regard to life-  
356 years gained per resource expended. Delay in diagnosis and surgery cause exponential  
357 burden of attributable mortality. The COVID-19 pandemic has placed unprecedented strain  
358 on health care provision. It is highly plausible that surges of population infection, lock-  
359 downs, resource competition, bottlenecks, and back-logs could recur over the next two  
360 years. Supra-normal capacity is required to manage backlogs of accumulated cancer cases  
361 alongside ongoing incident cases. To avoid a deferred public health crisis of unnecessary  
362 cancer deaths, urgent ringfencing of substantial resources is required.

363

364

**365 LEGENDS FOR FIGURES**

366

367 **Figure 1:** Impact from 6-months delay lasting one year for all solid cancers analysed and six  
368 common cancer types in England expressed in **a:** Attributable deaths **b:** Life years Lost

369

**370 Author contributions**

371 C.T., M.E.J., A.S. and R.S.H. designed the model. M.E.J. provided cancer progression models.  
372 J.B. generated and quality-assured the NCRAS datasets applied to the model. M.E.J., J.B.  
373 C.T., R.S.H., A.S., C.A., G.L., M.W., and P.D.P.P provided epidemiological expertise in  
374 parameterisation of the model. F.G provided microbiology expertise in estimation of  
375 nosocomial infection rates. S.A.B, S.J., D.L.N, P.W., J.L., J.M.H, N.Y. and Y-E.S provided details  
376 of clinical pathways and estimates of clinical resourcing. B.T., A.G. and C.L. quality assured  
377 and user-tested the model. B.T. and C.L. assembled figures for presentation. C.T drafted the  
378 manuscript, with substantial contribution from A.S., R.S.H., M.E.J., G.L., M.W. and C.S.. All  
379 authors contributed to the final manuscript.

380

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403 The authors have no relevant disclosures to declare.

404

#### 405 **Highlights**

- 406 • Lockdown and re-deployment due to the COVID-19 pandemic is causing significant  
407 disruption to cancer diagnosis and management.
- 408 • 3-month delay to surgery across all Stage 1-3 cancers is estimated to cause >4,700  
409 attributable deaths per year in England.
- 410 • The impact on life years lost of 3-6 month to surgery for Stage 1-3 disease varies  
411 widely between tumour types.
- 412 • Strategic prioritisation of patients for diagnostics and surgery has potential to  
413 mitigate deaths attributable to delays.
- 414 • The resource-adjusted benefit in avoiding delay in cancer management compares  
415 favourably to admission for COVID-19 infection.

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COMPONENT OF MODEL	ELEMENTS	DATA SOURCE	COMMENT	Reference/specific values	
Life years lost due to delay in surgery	Proportion of patients surviving after surgery	5 year survival rates for cancer surgery in England	Age, site, and stage-specific 5-year cancer survival in individuals in whom major resection was performed	PHE NCRAS[4]	
	Decrease in survival due to delay in treatment	Observational studies of increased death rate due to delay in treatment	Hazard ratio for increase in death rate for each day delay in treatment based on estimates from literature, applied to standard survival rates. Applied to tumours depending on tumour aggressiveness	<b>Cancer progressiveness based on 5y survival:</b> Low: >90%, Moderate: 50-90% High: <50%  <b>Per day Hazard ratio for fatality [10-20]:</b> Low: 0.0030, Mod: 0.0056 High: 0.0056	
	COVID-related post-surgical mortality. SACT-related mortality	Nosocomial infection rate		Based on literature, estimate from clinical site data	5 % per day[29]
		Mortality from COVID-infection		Age-specific data from international series	<b>0-39 y</b> 0.2% <b>30-39 y</b> 0.2% <b>40-49 y</b> 0.4% <b>50-59 y</b> 1.3% <b>60-69 y</b> 3.6% <b>70-79 y</b> 8.0% <b>80+ y</b> 14.8%
		Survival benefit from SACT		Expert clinical interpretation of literature	<b>Stage 1:</b> 5% <b>Stage 2:</b> 7.5% <b>Stage 3:</b> 10% [30]
		Increase in COVID-related mortality due to SACT		Based on UK and international literature	2-fold [7, 8]
		Life-expectancy after survival	General population mean life-expectancies per 10 year age-band	Expected remaining life years in treated group based on proportion who survive after treatment (with and without delay)	ONS Life Tables[31]
	Healthcare resourcing	Duration of operation, ICU and inpatient ward stay	Data from UK surgical oncology centres	Calculated as Healthcare Resource Unit (HCRUs) of direct clinical care. 1 HCRU= one 12 hour medical/nursing shift	
Staffing ratios in theatre, wards, ICU					

**Table 1:** Summary of sources for parameters estimates for cancer surgical model (see Supplementary Table 1 for full description)

	Stage	30-39 y	40-49 y	50-59 y	60-69 y	70-79 y	80+ y
Bladder	1	15.8%	15.8%*	26.3%	18.4%	21.9%	23.8%
	2	36.0%	35.9%	32.7%	31.9%	29.0%	28.6%
	3	35.9%	35.8%*	34.8%	34.1%	32.4%	29.3%
Breast (ER+, HER2-)	1	1.5%	0.6%	-0.3%	-1.5%	-3.2%	-3.1%
	2	5.9%	2.8%	2.4%	0.7%	-1.3%	-5.6%
	3	13.4%	8.2%	9.2%	9.2%	9.1%	2.5%
Breast (ER-, HER2-)	1	6.2%	4.3%	5.4%	2.3%	0.5%	4.1%
	2	13%	12.2%	11.3%	10.0%	12.7%	14.0%
	3	18.2%	19.8%	19.4%	18.5%	18.2%	16.0%
Breast (HER2+)	1	0.4%	0.9%	1.0%	0.5%	-1.7%	3.5%
	2	4.2%	3.1%	3.4%	3.0%	3.3%	6.5%
	3	11.3%	7.0%	9.6%	8.8%	13.9%	15.0%
Colon and rectosigmoid junction	1	2.1%	4.9%	4.5%	3.0%	-1.5%	-2.8%
	2	16.7%	15.9%	14.0%	14.7%	15.0%	4.8%
	3	29.9%	29.1%	29.2%	28.5%	30.2%	28.8%
Kidney	1	2.1%	2.6%	6.0%	5.1%	0.5%	-2.5%
	2	13.2%	17.0%	11.5%	16.1%	13.8%	26.4%
	3	19.8%	23.5%	25.7%	24.9%	23.5%	22.2%
Larynx	1	11.5%	16.3%	19.0%	16.9%	11.2%	20.1%
	2	29.5%	29.5%*	20.5%	31.7%	32.3%	32.5%
	3	33.9%	33.8%*	35.4%	34.2%	32.8%	20.7%
Lung (non-small cell)	1	5.4%	14.3%	25.4%	27.5%	29.6%	24.0%
	2	31.6%	34.2%	34.8%	34.5%	32.3%	29.6%
	3	35.7%	35.7%	34.1%	29.6%	27.9%	19.6%
Melanoma of skin	1	1.1%	2.5%	0.4%	1.2%	0.2%	2.8%
	2	19.9%	22.5%	24%	28.2%	27.1%	34.4%
	3	29.0%	30.8%	31.4%	33.5%	31.4%	31.5%
Oesophagus	1	31.6%	31.5%	29.8%	29.4%	24.7%	29.9%
	2	35.9%	35.8%*	35.4%	34.3%	32.2%	28.3%
	3	35.8%	34.2%	30.4%	31.9%	27.0%	25.3%
Ovary	1	4.6%	7.1%	10.8%	10.4%	11.3%	-1.1%
	2	16.9%	26.2%	28.9%	29.6%	31.9%	35.3%
	3	31.5%	35.9%	33.8%	31.5%	28.6%	21.0%
Pancreas	1	1.0%*	9.6%*	12.7%	15.4%	20.2%*	28.1%
	2	23.8%	35.9%*	27%	23.6%	21.4%	25.9%
	3	24.8%	24.7%*	32.3%	33.2%*	31.4%*	24.1%
Prostate	1	1.4%*	1.4%	-0.3%	-0.7%	1.6%	15.4%
	2	0.0%*	-0.1%	-0.3%	-0.7%	-1.5%	16.9%
	3	0.0%*	-0.1%	-0.3%	-0.7%	-1.5%	17.8%
Stomach	1	12.2%	18.6%*	29.3%	21.4%	11.1%	-6.5%
	2	35.0%	27.9%*	35.2%	34.4%	32.2%	18.0%
	3	35.0%	32.3%	33.2%	32.3%	28.9%	26.8%
Uterus	1	3.3%	5.6%	6.1%	9.5%	12.6%	6.0%
	2	13.2%	18.4%	18.9%	26.5%	32.6%	33.0%
	3	10.2%	31.1%	33.4%	35.8%	33.1%	33.6%

**Table 2:** Reduction in five-year net survival as a consequence of six-month delay to surgery for 13 cancer types, by tumour stage and age of diagnosis.

Reduction in survival above the median is represented in red, at the median in yellow and below the median in green. Survival analysis is based on per-day hazard ratios for disease fatality. \* indicates strata estimates of lower confidence whereby crude rather than net survival estimates were applied.

	Stage	30-39	40-49	50-59	60-69	70-79	80+
Bladder	1	4.1*	3.3*	4.1	2.0	1.5	0.8
	2	0.7*	0.6	0.4	0.3	0.1	0.1
	3	0.7*	0.6*	0.4	0.3	0.2	0.1
Breast (ER+, HER2-)	1	0.3	0.1	0.0	-0.1	-0.2	-0.1
	2	1.2	0.5	0.3	0.1	-0.1	-0.2
	3	2.8	1.4	1.2	0.8	0.5	0.1
Breast (ER-, HER2-)	1	1.3	0.7	0.7	0.2	0.0	0.1
	2	2.7	2.0	1.4	0.9	0.7	0.4
	3	3.8*	3.3	2.4	1.6	1.0	0.5*
Breast (HER2+)	1	0.1	0.2	0.1	0.0	-0.1	0.1
	2	0.9	0.5	0.4	0.3	0.2	0.2
	3	2.4	1.2	1.2	0.8	0.8	0.4
Colon and rectosigmoid junction	1	0.1	0.1	0.1	0.0	0.0	0.0
	2	0.6	0.4	0.3	0.2	0.1	0.0
	3	1.0	0.8	0.6	0.4	0.3	0.1
Kidney	1	0.1	0.1	0.2	0.1	0.0	0.0
	2	0.5*	0.5	0.2	0.2	0.1	0.1
	3	0.7*	0.7	0.6	0.4	0.2	0.1
Larynx	1	0.4*	0.4	0.4	0.2	0.1	0.1
	2	0.9*	0.7*	0.4*	0.4	0.3	0.1*
	3	1.0*	0.8*	0.6	0.4	0.3	0.1*
Lung (non-small cell)	1	0.2	0.3	0.5	0.3	0.2	0.1
	2	0.9*	0.8	0.6	0.4	0.2	0.1
	3	1.1*	0.8	0.6	0.4	0.2	0.1
Melanoma of skin	1	0.4	0.7	0.1	0.2	0.0	0.1
	2	2.1	1.9	1.5	1.2	0.7	0.5
	3	3.0	2.6	2.0	1.5	0.9	0.4
Oesophagus	1	0.6*	0.4	0.3	0.2	0.1	0.1*
	2	0.6*	0.5*	0.4	0.3	0.1	0.1*
	3	0.6*	0.5	0.3	0.2	0.1	0.1*
Ovary	1	0.5	0.6	0.7	0.5	0.3	0.0
	2	1.8*	2.2	1.8	1.3	0.9	0.5
	3	0.8	0.8	0.5	0.4	0.2	0.1
Pancreas	1	0.0*	0.1*	0.1*	0.1	0.1*	0.1*
	2	0.4*	0.5*	0.3	0.2	0.1	0.1*
	3	0.4*	0.4*	0.4*	0.3*	0.1*	0.1*
Prostate	1	0.0*	0.0	0.0	0.0	0.0	0.1
	2	0.0*	0.0	0.0	0.0	0.0	0.1*
	3	0.0*	0.0	0.0	0.0	0.0	0.1*
Stomach	1	0.3*	0.3*	0.4	0.2	0.1	0.0
	2	0.7*	0.4*	0.4	0.3	0.2	0.0
	3	0.7*	0.5	0.4	0.3	0.1	0.1
Uterus	1	0.3	0.4	0.3	0.4	0.3	0.1
	2	1.1*	1.3	1.0	1.0	0.7	0.4
	3	0.9*	2.2	1.8	1.3	0.8	0.4

**Table 3:** Estimated average life years gained per unit of healthcare resource for cancer surgery for 13 cancer types, by tumour stage and age of diagnosis comparing current surgery to surgery after six months delay based on 5-year net survival.

\* indicates strata estimates of lower confidence whereby crude rather than net survival estimates were applied. Values for LYG per HCRU above the median are represented in blue, at the median in white and below the median in red.

CANCER SURGERY					
Reference time period (months)		12		24	
Per patient delay (months)		3	6	3	6
Per day rate of nosocomial infection (current)		5%			
STANDARD CONDITIONS	Major resections for cancer-	94,912		189,823	
	HCRUs-total	764,765		1,529,529	
	LY gained-total	1,717,051		3,434,102	
	Lives saved-total	80,406		160,812	
	LY gained from cancer	18.1			
	LY gained from cancer	2.2			
IMPACT of DELAY	Deaths attributable to delay-total	4,755	10,760	9,511	21,521
	LY lost attributable to delay-total	92,214	208,275	184,428	416,549
	LY gained from cancer treatment post-delay- average per patient	17.1	15.9	17.1	15.9
	LY lost attributable to delay-average per patient	0.97	2.19	0.97	2.19
	LY gained per HCRU from cancer treatment post-delay-average per patient	2.12	1.97	2.12	1.97
	LY lost per HCRU attributable to delay-average per patient	0.12	0.27	0.12	0.27
HOSPITALISATION OF COMMUNITY-ACQUIRED COVID INFECTION					
Reference time period (months)		12		24	
Community infections		683,083		1,366,167	
Hospital Admissions	Total admissions	94,912		189,823	
	ICU admissions	16,135		32,270	
	non-ICU admissions	78,777		157,553	
Health care resource units (HCRUs)	Total	1,052,949		2,105,899	
	ICU	556,657		1,113,313	
	non-ICU	496,293		992,586	
Deaths	Total	15,587		31,173	
	ICU	8,241		16,481	
	non-ICU	7,346		14,692	
Total lives saved -attributable to hospital admission	All	25,752		51,504	
	ICU	7,894		15,789	
	non-ICU	17,858		35,715	
Total LY gained -attributable to hospital admission	All	482,022		964,044	
	ICU	223,227		446,454	
	non-ICU	258,795		517,591	
LY gained -average per patient	All	5.08			
	ICU	13.83			
	non-ICU	3.29			
LY gained per HCRU -average per patient	All	0.46			
	ICU	0.40			
	non-ICU	0.52			
Comparison	LY lost through <u>delay</u> in cancer treatment as a proportion of LY gaineds from hospitalisation from COVID-19	19%	43%	19%	43%
	RALY lost through <u>delay</u> in cancer treatment as a proportion of RALY gaineds from hospitalisation from COVID-19	26%	59%	26%	59%

**Table 4:** Summary outcomes from delays in cancer surgery, with comparison to an equivalent number of admissions for community-acquired COVID-19 infection. Only major resections for common adult cancers included. Reference population: England. LY: life years. RALY: resource adjusted life years. HCRU: healthcare resource units

