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DOI: 10.1002/pbc.27509

License: Other (please specify with Rights Statement)

Document Version Peer reviewed version

Citation for published version (Harvard):

Stevens, S, Main, C, Bailey, S, Pizer, B, English, M, Phillips, B, Peet, A, Avula, S, Wilne, S, Wheatley, K, Kearns, P & Wilson, J 2018, 'The utility of routine surveillance screening with magnetic resonance imaging to detect tumour recurrence/progression in children with high-grade central nervous system tumours', *Pediatric Blood & Cancer*. https://doi.org/10.1002/pbc.27509

Link to publication on Research at Birmingham portal

Publisher Rights Statement: Checked for eligibility: 20/11/2018

This is the peer reviewed version of the following article: Stevens SP, Main C, Bailey S, et al. The utility of routine surveillance screening with magnetic resonance imaging to detect tumor recurrence/progression in children with high-grade central nervous system tumors: a systematic review. Pediatr Blood Cancer. 2018;e27509., which has been published in final form at: https://doi.org/10.1002/pbc.27509. This article may be used for non-commercial purposes in accordance with Wiley Terms and Conditions for Use of Self-Archived Versions.

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The Utility of Routine Surveillance Screening with Magnetic Resonance Imaging (MRI) to Detect Tumor Recurrence / Progression in Children with High-Grade Central Nervous System (CNS) Tumors: A Systematic Review

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Text word count: 3,453 Abstract word count: 250 Brief running title: surveillance MRI in paediatric high-grade CNS tumors Key words: Systematic review; Magnetic Resonance Imaging (MRI); Surveillance; High Grade Tumors; Paediatric CNS tumors; Recurrence Tables: 3 Figures: 1

ABSTRACT

The Utility of Routine Surveillance Screening with Magnetic Resonance Imaging (MRI) to Detect Tumor Recurrence / Progression in Children with High-Grade Central Nervous System (CNS) Tumors: A Systematic Review

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Background: Surveillance Magnetic Resonance Imaging (MRI) is routinely used to detect recurrence in children with high-grade central nervous system (CNS) tumors, although no consensus has been reached regarding its effectiveness and whether earlier detection is associated with improved patient outcomes. This review aimed to evaluate this practice and any associated benefits and harms. **Methods**: Systematic searches for relevant studies were undertaken in a number of databases, including MEDLINE and EMBASE, from 1985 to August 2018. Study selection and data extraction was undertaken independently by two reviewers. Due to heterogeneity between studies, no pooling of data was undertaken. Reporting followed PRISMA guidelines. **Results:** No comparative studies were identified. Three retrospective observational studies involving 306 patients were reviewed. All had high risk of bias by virtue of study design. Two studies reported outcomes by symptomatic status both recurrence rates and overall survival for asymptomatic patients were comparable to those for clinically symptomatic patients. No quality of life outcomes were reported. **Conclusion:** There is a paucity of evidence to guide clinical practice as to the effectiveness of MRI surveillance in paediatric patients with high grade CNS tumors. These studies do not clearly demonstrate benefit or harm for the practice. With more research needed, there is a role for researchers to build into future trials data collection on surveillance imaging to give more information for the assessment of imaging frequency and duration in asymptomatic patients. This is an important question, not only to clinicians and patients and their families but also from a health service resource perspective.

1 Introduction

2

Paediatric high-grade central nervous system (CNS) tumors are fast-growing, malignant 3 tumors with metastatic potential and are commonly associated with poor prognosis even after 4 multi-modal treatment. Generally classified by the World Health Organization (WHO) as 5 either grade III or IV tumors, they include glial (anaplastic astrocytoma and glioblastoma 6 multiforme), ependymal (ependymoma, both WHO grade II and III) and embryonal 7 (medulloblastoma and tumors previously known as primitive neuroectodermal tumors 8 (PNET)) tumors, as well as brainstem tumors (diffuse pontine glioma (DIPG)) atypical 9 teratoid/rhabdoid tumor (AT/RT) and pineoblastoma. Many children with high-grade CNS 10 tumors will go on to experience recurrence or progression and the likelihood of this will 11 depend on the histology and location of their first tumor, as well as treatments given.^{1,2} 12 13 In recent years, Magnetic Resonance Imaging (MRI) has become the predominant imaging 14 tool in the management of children with high-grade CNS tumors. The rationale behind 15 routine imaging, or surveillance, is that recurrence or progressive disease detected at an 16 earlier stage may be more responsive to treatment and benefit from a wider range of 17 18 treatment options than disease diagnosed at a later stage from clinical signs and symptoms. However, no consensus has been reached as to whether this leads to improved outcomes for 19 20 patients and their families. 21 The objectives of this review were therefore to: 22 1. assess the diagnostic utility of surveillance MRI in detecting tumor recurrence prior 23

to the emergence of new clinical signs and symptoms compared to the nonroutine use
of MRI upon symptomatic presentation, and assess whether this practice translates to
measurable improvements in clinical outcomes;

27	2.	consider the effect of differing screening intervals on the diagnostic utility of	
28		surveillance MRI and determine the optimal duration of imaging after initial	
29		diagnosis; and	
30	3.	identify any gaps and methodological weaknesses in the current evidence base and	
31		make recommendations to inform the design and analysis of future studies.	
32			
33	The au	thors have also undertaken a systematic review on the effectiveness of surveillance	
34	MRI ir	paediatric low-grade tumors, which forms a companion piece to this review paper. ³	
35			
36	Methods		
37			
38	Standard systematic review methodology was employed and reporting followed the Preferred		
39	Report	ing Items for Systematic Reviews and Meta-Analyses (PRISMA) guidelines. ⁴ A	
40	detaile	d account of the methodology employed in this review can be found in the published	
41	protoco	ol, which is also registered with PROSPERO (CRD42016036802). ⁵ A summary of the	
42	method	ds are described below.	
43			
44	Search	strategy	
45			
46	This re	view formed part of a wider NIHR-funded work programme of systematic reviews	
47	aimed	at assessing the effects of different interventions for the treatment of paediatric CNS	
48	tumors	and therefore searches were not restricted to studies concerned solely with	
49	surveil	lance imaging in children with high-grade tumors. Searches for published studies from	
50	1985 to	O August 2018 were undertaken in several databases, including MEDLINE and	
51	EMBA	SE. (See Supplementary File S1). No language, publication restrictions or study	
52	design	filters were applied.	

53 Study selection 54 55 56 The following inclusion and exclusion criteria were applied: 57 Population: Children and young adults (up to age 25 years) with diagnoses of any type of 58 59 high-grade CNS tumor who were asymptomatic at the time of study recruitment. Given that children undergoing surveillance may have some neurologic sequelae from their tumor and/or 60 its treatment, it would be more accurate to characterise patients as exhibiting no new, stable 61 or improved neurological signs or symptoms. 62 63 Interventions: Routine or surveillance MRI. Studies employing computed tomography (CT) 64 as the sole surveillance imaging modality were excluded. 65 66 67 Outcome Measures: Recurrence rates (by study, tumor type, location and extent of resection), diagnostic yield of imaging, timing of recurrence, change in patient management post-68 recurrence, overall survival (OS), surrogate survival measures (e.g. recurrence-free survival 69 70 (RFS), progression-free survival (PFS)) and quality of survival. Studies reporting outcomes from aggregated CT and MRI scans were excluded. 71 72 Study designs: As randomized controlled trials (RCTs) and nonrandomized comparative 73 studies were initially sought but not identified, the review was extended to include 74 observational studies such as case series. 75 76 Study selection was undertaken by two independent reviewers, with disagreements resolved 77 78 by discussion.

79	
80	Data extraction and risk of bias assessment
81	
82	Data, extracted by one reviewer and checked by a second, were recorded on a standardised
83	proforma developed in Microsoft Word (See Supplementary File S2). Risk of bias was
84	assessed at the study level by two reviewers using a six-point tool devised by the Centre for
85	Reviews and Dissemination (York; CRD) ⁶ designed to assess bias in case series studies.
86	
87	Statistical analysis
88	
89	Due to the design of the included studies and the heterogeneity of outcomes reported, only a
90	descriptive analysis was undertaken.
91	
92	Results
93	
94	Quantity and description of included studies
95	
96	From the electronic database searches, 28 potentially relevant publications were identified
97	with an additional 13 publications identified from citation-checking. On full text
98	examination, 38 were excluded including 11 studies which employed both CT and MRI as
99	surveillance imaging modalities but failed to report results separately for MRI. (See
100	Supplementary File S3). No RCTs or prospective comparative studies were identified. Three
101	retrospective case series studies ⁷⁻⁹ were included in the review. (See Figure 1).
102	

103 The three studies were conducted between 2001 and 2014 and undertaken at single centre

104 institutions. Two studies^{8,9} included patients with high-grade tumors only, with one⁷

including a mix of low- and high-grade tumor patients. (See Table 1).

106

107 Quality of the research

108

Studies were clinically heterogeneous with study populations varying in terms of both tumor 109 type and disease severity. Study samples were small but patients appeared to be 110 representative of the target population, although it was unclear whether patients were at a 111 similar timepoint in the disease progression. Inclusion and exclusion criteria for each study 112 were explicitly stated. Generally details of previous treatments were not reported. (See 113 Supplementary File S4). There was also variability in terms of reporting and defining of 114 115 outcomes. The terms 'recurrence' and 'progression' were defined in all three studies, although only two reported recurrences as 'symptomatic' and 'asymptomatic' and defined 116 these terms.^{7,9} All three studies reported OS, although only Kornreich⁸ defined the term. (See 117 Table S5). This was also the only study to report PFS. Korones⁷ did not report average 118 duration of follow-up. 119 120

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121 Included studies
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122

123 Korones $(2001)^7$

124

Korones⁷ was a mixed tumor grade study with 112 children at study commencement. Patient
details were provided only for the 46 patients who went on to experience

127 recurrence/progression. Of these, 33 had high-grade tumors. Eight tumor types were

included. The median age of these patients at recurrence was six years (range 0.25 – 21)
although this was not reported by tumor type.

130

All patients underwent surgery as the primary treatment, although this was not further
specified by extent of resection (i.e., gross total resection (GTR) versus sub-total resection
(STR)). At the commencement of surveillance imaging, none of the patients had relapsed
disease.

135

136 With respect to imaging frequency, patients received a median of one scan every 2.5 months

137 (range 1/1 - 1/6.7 months) irrespective of whether they were symptomatic or asymptomatic at

138 recurrence. Frequency of scanning was not reported by tumor type.

139

As only data on recurrent patients was reported, it was not possible to calculate the recurrence
rate for the 33 high-grade tumor patients as a whole, nor by tumor type. The rate of
recurrence/progression by symptomatic status was reported, with 17 patients (52%)
asymptomatic at recurrence. Recurrence by symptomatic status was also reported by tumor
type, with asymptomatic and symptomatic recurrences comparable in number, although the
numbers in each category were very small (ranging from 1-6). (See Table 2). Recurrence by
extent of resection was not reported.

147

The diagnostic yield of imaging for all seventeen asymptomatic patients was 4.4%, i.e., one
asymptomatic recurrence detected every 23 MRI scans. (See Table 2). With respect to CPC,
GCT and AT/RT, there were two asymptomatic recurrences among these tumor types and the

151 diagnostic yield of imaging was 6.5%.

Median time to recurrence from initial diagnosis for all 33 patients was 0.75 years with no 153 significant difference in median time to recurrence between symptomatic and asymptomatic 154 patients at recurrence (0.66 and 0.77 years respectively). Median time to recurrence was not 155 156 reported by individual tumor type, nor by extent of resection. 157 Information regarding local therapy received following recurrence/progression was provided 158 for 26 patients (79%), with eight of 14 asymptomatic patients (57%) undergoing local therapy 159 (surgery with or without stereotactic radiosurgery) compared to only three of 12 symptomatic 160 patients (25%) (p = 0.13). Again, change in patient management was not reported by tumor 161 type. 162 163 Overall survival from recurrence for all 33 patients was reported but only by symptomatic 164 status at recurrence, with median OS for the 17 asymptomatic patients (0.58 years) 165 marginally and nonstatistically significantly greater (p=0.25) than that for the 16 166 symptomatic patients (0.42 years). Median OS was not reported by tumor type. 167 168 Kornreich $(2005)^8$ 169 170 Kornreich⁸ was a retrospective case series study looking at the role of surveillance MRI in the 171 172 management of 15 paediatric patients with DIPG. While the frequency of imaging was not reported, the mean number of MRI scans per patient was six. Thirteen patients (87%) 173 experienced tumor progression while two patients remained stable. Symptomatic status of 174 patients at progression was not reported. 175 176 Median PFS was 0.83 years, ranging from 0 months (in 4 patients who deteriorated 177 immediately from diagnosis without any prior period of stability) to nine years. Treatment 178

(radiotherapy and/or chemotherapy) was planned and not consequent to changes in scans or
recurrence. Median OS was 1.67 years, with three patients (20%) alive at the time of
reporting.

182

183 Perreault $(2014)^9$

184

Perreault⁹ was a retrospective case series study which sought to assess the benefits of
surveillance MRI in a cohort of 258 high-grade tumor patients. There were seven tumor types
included. (See Table 1). All patients underwent surgery as the primary treatment although
this was not further specified by extent of resection. At commencement of surveillance
imaging, none of the patients had relapsed disease.

190

While frequency of scanning was not reported, the median number of MRI scans per patient
across all tumor types was 13, ten of the brain and three spinal. (See Table 3). The interval
since last MRI for symptomatic patients was not longer for symptomatic compared to
asymptomatic patients (mean 3.9 versus 4.8 months).

195

196 Rates of recurrence/progression were also reported by symptomatic status. (See Table 3). With respect to first recurrences (n=113), there was a slight predominance of asymptomatic 197 198 (46%) compared to symptomatic recurrences (42%), whereas for subsequent recurrences (n=125) the converse was the case (29% versus 58%). Recurrences (both first and 199 subsequent) by symptomatic status were also reported by tumor type where, in the case of 200 medulloblastoma and ependymoma, this trend continued with the majority of first recurrences 201 202 asymptomatic and second symptomatic. Conversely, for sPNET, the majority of first recurrences were symptomatic and second asymptomatic. For HGG, the majority of both first 203 and second recurrences were symptomatic. For the remaining tumor types (GCT, AT/RT and 204

pineoblastoma), the number of recurrences was so small that caution should be exercised
when comparing recurrences by symptomatic status (most notably AT/RT, with 100% of first
recurrences asymptomatic based on only four patients). Recurrences among glioma patients
were more frequently symptomatic compared to those patients with other tumor types (68
versus 38 % respectively; p=0.003). The rate of recurrence by extent of resection was not
reported.

211

A breakdown of MRI scans by both tumor type and site of imaging was reported, with diagnostic yield across all tumor types of 8.3% for brain recurrence only (range 2.1% to 21.6%), 3.8% for combined brain-spine recurrence (range 1.6% to 19.7%) and 0.9% for spine recurrence only (range 0.7% to 4.9%). (See Table 3).

216

Median time to recurrence from initial diagnosis was 1 year, although it is unclear whether this relates to first or all recurrences. Median time to recurrence by tumor type was reported but, again, it is unclear if this relates to first or all recurrences. (See Table 3). No significant difference in median time to recurrence was reported between symptomatic and asymptomatic patients at recurrence (1.0 and 0.92 years respectively; p>0.8). The time by which greater than 90% of recurrences had occurred for each individual tumor type was also reported. (See Table 3). Median time to recurrence by extent of resection was not reported.

Change in patient management following first recurrence was reported for 93% of patients,
with 59% of patients undergoing new treatments, 11% continuing with existing treatment,
16% scheduled for palliative care and 7% undergoing closer interval surveillance MRI. New
treatments consisted of chemotherapy (22% standard dose and 4% high dose with stem cell
support), radiotherapy (6%), radiosurgery (2%), surgery (5%) and unspecified multi-modal

therapy (20%). Change in patient management post-recurrence by tumor type was notreported.

232

There was no significant difference (p > 0.3) in median OS from recurrence between
symptomatic and asymptomatic patients (1.92 years and 2.25 years respectively). Median OS
by tumor type was not reported.

236

237 Discussion

238

This systematic review is one of a series evaluating treatments for children with CNS tumors. 239 Underpinning the reviews was consultation with clinical experts and a Patient and Public 240 Involvement (PPI) group, consisting of mothers of children with CNS tumors. The PPI group 241 in particular expressed concerns about over-scanning, especially in situations where scanning 242 is no longer able to influence prognosis as in the case of patients for which nothing further 243 can be clinically done. As well as the unknown risks associated with repeated administration 244 of contrast materials such as Gadolinium,¹⁰ anaesthesia and sedatives, the PPI group spoke of 245 what has come to be termed 'scanxiety', i.e. an overwhelming feeling of stress experienced 246 247 by both patient and family around the time of scanning. As one parent put it "At times, it seems like life and all its decisions revolve around scanning, which serves as a constant 248 249 reminder of the cancer and acts as an obstacle to resuming normal behaviour."

250

Although the use of surveillance MRI is standard practice throughout the developed world in the management of children with high-grade CNS tumors, this systematic review did not identify any RCTs evaluating this intervention. After excluding 11 high-grade tumor surveillance imaging studies which employed both CT and MRI but did not report results separately by imaging modality,¹¹⁻²¹ the review included three retrospective, single arm

studies (n=306 patients) with MRI employed as the sole imaging modality. It could be argued
that in excluding studies employing CT imaging, the review has lost valuable data on
surveillance. However, the reason for focussing on MRI, other than its superior sensitivity, is
that MRI studies are more recent than CT studies and therefore encompass an era of
improved survival and greater salvageability of patients due to improved treatments.

261

The findings of the review were mixed. Korones⁷ concluded that "asymptomatic recurrences 262 were detected in only a small proportion of surveillance scans and had no impact on survival 263 in children with high-grade tumors." Kornreich⁸ reported on 15 patients with DIPG and 264 compared the findings of 51 surveillance scans with those from clinical examination and 265 reported a high degree of concordance (87%), suggesting that for DIPG, surveillance MRI is 266 providing little information over and above that conveyed by clinical symptoms and signs 267 and therefore its utility may be limited. Ultimately, surveillance imaging did not affect the 268 treatment given, nor the outcome. Based on this evidence, it could be argued, albeit 269 tentatively, that certain tumor types may be more amenable to surveillance MRI than others 270 and that for aggressive tumors such as DIPG, where often any period of clinical stability is 271 extremely limited, there is a very short window of opportunity for surveillance imaging to 272 exploit. In support of this, Kornreich⁸ reported four patients with zero time to progression. 273 However, with other, less aggressive high-grade tumor types, the use of MRI surveillance 274 may be of value. For example, with Perreault⁹ asymptomatic recurrence rates were higher for 275 ependymoma and medulloblastoma compared to other tumor types, suggesting that 276 surveillance might potentially be beneficial to these patients, although in this study 277 asymptomatic patients across all tumor types did not benefit from improved overall survival 278 279 compared to symptomatic patients. Unfortunately, the potential for bias within case series is considerable and therefore conclusions from this review are tentative and should be viewed 280 with extreme caution. 281

283	There were several reporting problems that made comparison across the studies problematic.
284	Korones failed to report frequency of MRI imaging by tumor grade or type thereby rendering
285	a cross-study comparison of the effect of differing imaging schedules on the rate of
286	asymptomatic recurrence for different tumor types impossible. ⁷ Similarly, Kornreich ⁸ did not
287	report patients by symptomatic status at time of progression. Only Perreault ⁹ reported patients
288	and recurrences by tumor type and symptomatic status, enabling observations to be drawn
289	which could potentially inform the design of future trials. However, it is important to
290	appreciate that the data analysed in these studies were acquired for clinical purposes for
291	which assessment of surveillance imaging protocols was not an objective.
292	
293	The initial aim of this review was to assess the effectiveness of surveillance MRI. RCTs were
294	required to do this but as none were found, focus was switched to finding studies that were
295	specifically conducted to describe surveillance scanning. With just three studies meeting the
296	inclusion criteria, one criticism of this review which emerged from the peer review process
297	was that the co-operative trials should have been hand-searched for information on
298	surveillance. This does raise an interesting point about the best way to systematically review
299	paediatric oncology trials. Systematic reviewing (especially employing Cochrane
300	methodology) was developed with single question trials involving more common diseases in
301	mind, i.e. A versus B, whereas paediatric oncology trials tend to be co-operative, multi-modal
302	trials that attempt to answer a variety of questions within a single trial due to the rarity of the
303	diseases. In response to the peer review feedback, a search of co-operative trials in
304	medulloblastoma was undertaken to determine whether there was data within these trials to
305	inform the review question. Of 27 trials, surveillance MRI scanning intervals appeared to be
306	arbitrary and variable, with few reasons given for the surveillance schedules. (See
307	Supplementary File S6). Only one study, not identified in our systematic review searches

likely due to indexing, evaluated the number of patients who had relapse detected through 308 surveillance MRI compared to symptom-based relapse.²² (This study reported that 45 relapses 309 were detected on surveillance MRI, with 20 detected from symptoms alone. Of these, patients 310 detected from symptoms had a significantly shorter survival post-relapse than those detected 311 by surveillance MRI (p<0.01), although OS post-primary diagnosis was not statistically 312 significantly different. This could be due to lead time bias or that patients in the symptomatic 313 relapse group possibly have more aggressive tumors). Finding the evidence in a systematic 314 way, from identifying the relevant publications to finding the information within the trial 315 publications (often results are written into the discussions) can be challenging in these large 316 co-operative trials. In future, we recommend that systematic reviewers consider hand-317 searching relevant co-operative trials, whilst bearing in mind that the main aim of these trials 318 might differ from that of the systematic review. We also urge authors of co-operative trials to 319 improve the transparency of their publications, especially with respect to database indexing 320 as well as signposting and organization of information within the papers. 321

322

The paucity of data evidenced in this review may be due to the complexity of surveillance in 323 these patients, with frequency of monitoring depending on tumor type, disease status (newly 324 325 diagnosed, resistant or relapsed), extent of metastatic spread and previous treatments. Other factors such as pseudo-progression and radiation necrosis can also complicate the 326 327 interpretation of scans, making it a difficult area to investigate. However, there is a need to examine this question further in order to guide clinicians in developing optimal evidence-328 based surveillance strategies, to help parents and children understand the need for 329 surveillance and to optimise the use of health service resources. There is a role for researchers 330 to build into future, large co-operative trials methodology that investigates the role of 331 332 surveillance MRI or at the very minimum, collects and reports data on the trial surveillance

333 MRI practice, as well as incorporating quality of life data collection, particularly regarding

anxiety around surveillance and the reassurance that it may also afford.

335

336 Conclusion

338	Only three retrospective observational studies with a high risk of bias were identified to guide
339	clinical practice of surveillance MRI for children with high-grade CNS tumors. ⁷⁻⁹ These
340	studies do not clearly demonstrate benefit or harm for this practice, nor do they define
341	methods or intervals for maximal effectiveness. To resolve this, more research is needed with
342	the ultimate endpoints of surveillance relating to survival and quality of life, as opposed to
343	surrogate outcomes such as the detection of tumor growth. As most of the patients within this
344	group are treated within the context of a co-operative clinical trial, this research could be built
345	into trial protocols for very little extra investment. It is an important question, not only to
346	clinicians and patients and their families but also as a health service resource question.
347	
348	
349	Figure 1: PRISMA diagram of flow of studies through the selection process
350	Supplementary file S1: Search strategy
351	Supplementary file S2: Data extraction and quality assessment proforma
352	Supplementary file S3: List of excluded studies
353	Supplementary file S4: Quality assessment of included studies
354	Supplementary file S5: Definitions of recurrence / progression and symptomatic /
355	asymptomatic provided by study authors
356	Supplementary file S6: MRI imaging schedules in co-operative trials in paediatric
357	medulloblastoma

Compliance with ethical standards

Disclosure of potential conflicts of interest: The authors declare that they have no conflict of interest.

Informed consent: Not required for this type of study (i.e. systematic review).

Ethical approval: Not required for this type of study (i.e. systematic review).

Research involving Human Participants and/or animals: This article does not contain any studies with human participants or animals performed by any of the authors.

Authors' contributions

SPS conceived and designed the study and wrote the article. CM conceived and designed the study and read and commented on the article. SB, BP, ME, RP conceived the study concept, provided clinical input and read and commented on the article. AP, SA, SW and PRK provided clinical input and read and commented on the article. KW conceived and designed the study, provided methodological and statistical input and read and commented on the article and commented on the article. JSW conceived and designed the study, revised the article and is the guarantor of the review. All authors read and approved the final manuscript.

Acknowledgments

We would like to acknowledge the input of the Patient and Public Involvement (PPI) group and the wider clinical team who helped to frame the review question and contribute to the direction of the paper.

Funding

This paper presents independent research funded by the National Institute for Health Research (NIHR) under its Research for Patient Benefit (RfPB) Programme (Grant Reference Number PB-PG-1112-29122). PK, KW and JW received funding from Cancer Research UK. AP is funded by an NIHR Research Professorship (13-0053). The views expressed are those of the author(s) and not necessarily those of the NHS, the NIHR, the Department of Health and Social Care or Cancer Research UK.

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