Adding 6 months of androgen deprivation therapy to postoperative radiotherapy for prostate cancer: a comparison of short-course versus no androgen deprivation therapy in the RADICALS-HD randomised controlled trial

Chris C Parker, Noel W Clarke, Adrian D Cook, Howard Kynaston, Charles N Catton, William R Cross, Peter M Petersen, Rajendra A Persad, Fred Saad, Lorna C Bower, John Logue, Heather Payne, Silvia Forcat, Cindy Goldstein, Claire Murphy, Juliette Anderson, Maroie Barkati, David M Bottomley, Jennifer Branagan, Ananya Choudhury, Peter W M Chung, Lyn Cogley, Chee L Goh, Peter Hoskin, Vincent Khoo, Shawn C Malone, Lindsey Masters, Stephen L Morris, Abdenour Nabid, Aldrich D Ong, Rakesh Raman, Kathryn L Tarver, Alison C Tree, Jane Worlding, James P Wylie, Anjali M Zarkar, Wendy R Parulekar*, Mahesh K B Parmar*, Matthew R Sydes*, on behalf of the RADICALS investigators†

Summary

Background Previous evidence indicates that adjuvant, short-course androgen deprivation therapy (ADT) improves metastasis-free survival when given with primary radiotherapy for intermediate-risk and high-risk localised prostate cancer. However, the value of ADT with postoperative radiotherapy after radical prostatectomy is unclear.

Methods RADICALS-HD was an international randomised controlled trial to test the efficacy of ADT used in combination with postoperative radiotherapy for prostate cancer. Key eligibility criteria were indication for radiotherapy after radical prostatectomy for prostate cancer, prostate-specific antigen less than 5 ng/mL, absence of metastatic disease, and written consent. Participants were randomly assigned (1:1) to radiotherapy alone (no ADT) or radiotherapy with 6 months of ADT (short-course ADT), using monthly subcutaneous gonadotropin-releasing hormone analogue injections, daily oral bicalutamide monotherapy 150 mg, or monthly subcutaneous degarelix. Randomisation was done centrally through minimisation with a random element, stratified by Gleason score, positive margins, radiotherapy timing, planned radiotherapy schedule, and planned type of ADT, in a computerised system. The allocated treatment was not masked. The primary outcome measure was metastasis-free survival, defined as distant metastasis arising from prostate cancer or death from any cause. Standard survival analysis methods were used, accounting for randomisation stratification factors. The trial had 80% power with two-sided α of 5% to detect an absolute increase in 10-year metastasis-free survival from 80% to 86% (hazard ratio [HR] 0.67). Analyses followed the intention-to-treat principle. The trial is registered with the ISRCTN registry, ISRCTN40814031, and ClinicalTrials.gov, NCT00541047.

Findings Between Nov 22, 2007, and June 29, 2015, 1480 patients (median age 66 years [IQR 61–69]) were randomly assigned to receive no ADT (n=737) or short-course ADT (n=743) in addition to postoperative radiotherapy at 121 centres in Canada, Denmark, Ireland, and the UK. With a median follow-up of 9.0 years (IQR 7.1-10.1), metastasis-free survival events were reported for 268 participants (142 in the no ADT group and 126 in the short-course ADT group; HR 0.886 [95% CI 0.688-1.140], p=0.35). 10-year metastasis-free survival was 79.2% (95% CI 75.4-82.5) in the no ADT group and 80.4% (76.6-83.6) in the short-course ADT group. Toxicity of grade 3 or higher was reported for 121 (17%) of 737 participants in the no ADT group and 100 (14%) of 743 in the short-course ADT group (p=0.15), with no treatment-related deaths.

Interpretation Metastatic disease is uncommon following postoperative bed radiotherapy after radical prostatectomy. Adding 6 months of ADT to this radiotherapy did not improve metastasis-free survival compared with no ADT. These findings do not support the use of short-course ADT with postoperative radiotherapy in this patient population.

Funding Cancer Research UK, UK Research and Innovation (formerly Medical Research Council), and Canadian Cancer Society.

Copyright © 2024 The Author(s). Published by Elsevier Ltd. This is an Open Access article under the CC BY 4.0 license.

Introduction

In patients receiving radiotherapy as initial treatment for localised prostate cancer, short-course androgen deprivation therapy (ADT) improves metastasis-free survival and overall survival, and is a standard of care for those with intermediate-risk and high-risk clinically localised disease.¹ In patients receiving postoperative radiotherapy after radical prostatectomy, the role of short-course ADT is less certain.

Two randomised controlled trials, GETUG-AFU 16 and RTOG 0534, have previously tested the addition of short-course ADT to salvage radiotherapy to the



oa

Lancet 2024; 403: 2405–15

Published **Online** May 16, 2024 https://doi.org/10.1016/ S0140-6736(24)00548-8

See **Comment** page 2353 See **Articles** page 2416

*Contributed equally

†A full list of investigators is provided in the appendix (pp 9–96)

The Roval Marsden NHS Foundation Trust, London, UK (Prof C C Parker MD. L C Bower PGCert. V Khoo MD(Res). A CTree MD(Res)); The Institute of Cancer Research, London, UK (Prof C C Parker, L C Bower, V Khoo, A C Tree); Department of Urology, The Christie NHS Foundation Trust, Manchester, UK (Prof N W Clarke ChM. I Loque FRCR Prof A Choudhury FRCR, I P Wylie FRCR): Division of Cancer Sciences, University of Manchester, Manchester, UK (Prof N W Clarke, Prof A Choudhury Prof P Hoskin MD); Department of Urology, Salford Royal Hospital Salford UK (Prof N W Clarke); MRC Clinical Trials Unit at UCL. Institute of Clinical Trials and Methodology, University College London, London, UK (A D Cook MSc, S Forcat PhD, C Goldstein MSc, C Murphy BSc, L Masters BSc. Prof M K B Parmar DPhil. Prof M R Sydes MSc); Division of Cancer and Genetics, Cardiff University Medical School, Cardiff, UK (Prof H Kynaston MD): Princess Margaret Cancer Centre. University Health Network, Toronto, ON, Canada

(Prof C N Catton FRCPC, PWMChungMBChB): Department of Urology (W R Cross PhD) and Department of Clinical Oncology (J Anderson FRCR, D M Bottomley FRCR), St James's University Hospital, Leeds, UK; Department of Oncology, Rigshospitalet, University of Copenhagen, Copenhagen, Denmark (Peter M Petersen PhD): Department of Urology, Bristol Urological Institute, Bristol, UK (Prof R A Persad FRCS Urol); Department of Urology (Prof F Saad MD) and Department of Radiation Oncology (M Barkati MD), Centre Hospitalier de l'Université de Montréal, Montreal, OC, Canada: Guy's and St Thomas' NHS Foundation Trust, London, UK (L C Bower, S L Morris FRCR); The Prostate Centre, London, UK (Prof H Payne FRCR); Northampton General Hospital, Northampton, UK (J Branagan FRCR); Department of Radiation Oncology, University of Toronto, Toronto, **ON**. Canada (P W M Chung): Derriford Hospital, Plymouth, UK (L Coglev RN): Roval Surrev Hospital, Guildford, UK (C L Goh MD (Res)); Mount Vernon Cancer Centre, Northwood, UK (Prof P Hoskin); The Ottawa Hospital. University of Ottawa, Ottawa, ON, Canada (Prof S C Malone FRCPC); Service de Radio-Oncologie, Centre Hospitalier Universitaire de Sherbrooke, Sherbrooke, QC, Canada (Prof A Nabid FRCPC): Max Rady Faculty of Health Sciences, University of Manitoba, Winnipeg, MB, Canada (A D Ong FRCPC); Kent Oncology Centre, Kent and Canterbury Hospital, Canterbury, UK (R Raman FRCR); Department of Oncology, Queen's Hospital, Romford, UK (K L Tarver FRCR); University Hospitals Coventry and Warwickshire NHS Trust. Coventry, UK (J Worlding FRCR); Department of Oncology, University Hospitals Birmingham, Birmingham, UK (A M Zarkar FRCR); Canadian Cancer Trials Group, Queen's University, Kingston, ON, Canada (Prof W R Parulekar MD)

Research in context

Evidence before this study

In patients receiving postoperative radiotherapy after radical prostatectomy, the role of short-course androgen deprivation therapy (ADT) is uncertain. Two randomised controlled trials, GETUG-AFU 16 and RTOG 0534, have tested the addition of short-course ADT to salvage radiotherapy to the prostate bed. Both found evidence in favour of short-course ADT in terms of freedom from prostate-specific antigen (PSA) progression. However, ADT will inevitably delay PSA progression, and that alone should not be sufficient to change practice. The impact of short-course ADT on longer-term, clinically meaningful outcome measures is not known.

Added value of this study

To our knowledge, RADICALS-HD is the first randomised trial of short-course ADT in men having postoperative radiotherapy that

prostate bed.^{2.3} Both trials found evidence in favour of adding short-course ADT in terms of freedom from progression, their primary outcome measure, although that outcome measure also included prostate-specific antigen (PSA) progression. ADT will inevitably delay PSA progression, and that alone should not be sufficient to change practice. GETUG-AFU 16 also reported a statistically significant improvement in metastasis-free survival (hazard ratio [HR] 0.73, 95% CI 0.54–0.98; p=0.034), whereas RTOG 0534 found no good evidence of an improvement in time to metastasis (0.78, 0.52–1.17; p=0.083). Neither trial has yet reported clear evidence of benefit for adding short-course ADT in terms of cause-specific survival or overall survival.

Given the known morbidity of ADT,4 together with its uncertain long-term benefits in this setting, there has been no consensus on the use of short-course ADT in patients receiving postoperative radiotherapy after previous radical prostatectomy. Previous surveys of UK urological and oncological specialists found variable use of ADT. Of those responding urologists who recommend adjuvant treatment, 72% recommended radiotherapy alone; when ADT was used, 31% recommended shortterm treatment (3-12 months) and 44% recommended either short-term or long-term treatment depending on the patient.56 The relevant clinical guidelines still make only weak recommendations; the ESMO Clinical Practice Guidelines for prostate cancer state that "ADT for 6 months...may be offered to men having salvage RT [radiotherapy]".7 We conducted a large, international, randomised controlled trial to test the efficacy of shortcourse ADT used in combination with postoperative radiotherapy to the prostate bed. Given the limitations of PSA-based outcome measures in any trial of ADT,8 we used metastasis-free survival as the primary outcome measure.

has used a long-term, clinically meaningful, primary outcome measure of metastasis-free survival. The results do not suggest that the impact of short-course ADT on PSA progression will translate into a clinically meaningful benefit in metastasis-free survival or overall survival.

Implications of all the available evidence

These findings are not sufficient to routinely recommend the use of short-course ADT with postoperative radiotherapy for people with prostate cancer. A meta-analysis of all the randomised trials addressing this issue is warranted and will provide the best evidence to guide practice.

Methods

Study design and participants

RADICALS was an international, phase 3, multicentre, open-label, randomised controlled trial in prostate cancer. The protocol addressed questions of the timing of radiotherapy after surgery and the use of ADT with postoperative radiotherapy across separate randomisations with overlapping patient groups.

RADICALS-HD recruited consenting patients with prostate localised cancer due for radiotherapy at any time after previous radical prostatectomy for prostatic adenocarcinoma. The exclusion criteria were previous pelvic radiotherapy, preoperative ADT for longer than 8 months, any ADT within 6 months before surgery, PSA greater than 5 ng/mL, metastatic disease, other active malignancy likely to interfere with protocol treatment or follow-up, or any postoperative hormone therapy. There were no age restrictions. Appropriate ethical review was in place for each participating country (appendix p 8). All participants gave written informed consent. The protocol is available online. This study is registered with the ISRCTN registry, ISRCTN40814031, and with ClinicalTrials.gov, NCT00541047.

Randomisation and masking

Participants in the none-versus-short comparison of RADICALS-HD were randomly allocated to radiotherapy alone (no ADT) or radiotherapy with the addition of 6 months of ADT (short-course ADT). Site staff engaged patients about potential participation in the trial. Those who decided to participate were given the choice, with their clinical team, of being randomly assigned three-way 1:1:1 between no ADT, short-course ADT, and long-course ADT (adding 24 months of ADT), or two-way 1:1 between no ADT and short-course ADT or between short-course ADT and long-course ADT. The large majority of participants chose to be randomly assigned in one of the two-way comparisons, although a small minority of participants chose to be randomly assigned in the three-way comparison. Patients who were allocated to long-course ADT are not included in this analysis because of the small number of participants in the three-way randomisation. Recruitment and random allocation of participants was implemented centrally using minimisation with a random element, stratified by Gleason score, positive margins, radiotherapy timing, planned radiotherapy schedule, and planned type of ADT, in a computerised system. The allocated treatment was open label.

Procedures

If allocated, ADT was to be initiated as soon as possible, and certainly within 2 months after randomisation, and continued for 6 months. Participants were given ADT using gonadotropin-releasing hormone analogue therapy according to site choice using their indicated route, usually subcutaneously, with once-monthly injections recommended. This was supplemented by 3 weeks of an anti-androgen started 1 week before the first gonadotropinreleasing hormone analogue therapy administration. Outside of Canada, daily oral bicalutamide monotherapy 150 mg or monthly subcutaneous degarelix (with nationally approved dosing) were acceptable alternatives. Dose reductions were not possible; treatment could be stopped early if indicated.

Radiotherapy was started approximately 2 months after starting ADT. The intended radiotherapy schedule was prespecified for each participant as $52 \cdot 5$ Gy in 20 fractions over 4 weeks or $66 \cdot 0$ Gy in 33 fractions over $6 \cdot 5$ weeks. The radiotherapy target volume was to include the prostate bed and could also include pelvic lymph nodes. Detailed radiotherapy guidance was given in the protocol.

Scheduled follow-up was every 4 months for the first 2 years after randomisation, then every 6 months up to 5 years, and annually thereafter. PSA measurements were taken at every follow-up appointment and as clinically indicated. Imaging was done according to routine clinical practice and was reported locally, without masking of treatment allocation. There was no central review of imaging.

Clinician-reported data were collected at each follow-up visit on diarrhoea, proctitis, cystitis, haematuria, and urethral stricture, graded according to Radiation Therapy Oncology Group toxicity score.⁹ Data for other adverse events were collected, and graded for severity if the event met the criteria to be classified as a serious adverse event.

An algorithm was used to identify deaths with uncertain cause, using the reported primary and contributory causes of death together with disease history during the trial.¹⁰ The process did not refer to allocated treatment. These uncertain causes of death of trial participants were centrally adjudicated by one of three clinical members of the Trial Management Group (CCP, NWC, or CNC). Additionally, for patients in England and Wales, national death registration data were used to confirm deaths and the information contributed to the reviews.

Outcome measures

The primary outcome measure was metastasis-free survival, defined as distant metastasis arising from prostate cancer or death from any cause. Extra-pelvic nodal disease was considered to be metastatic. Secondary outcome measures were freedom from distant metastasis (distant metastasis or death from prostate cancer), overall survival (death from any cause), initiation of non-protocol ADT (ADT other than the policy randomly assigned), clinical progression-free survival (local or nodal progression, metastases, non-protocol ADT, or death from prostate cancer), freedom from treatment failure (PSA progression when on ADT), toxicity, and patientreported outcome measures (PROMs). An additional secondary outcome of treatment failure, defined as PSA progression when on ADT, was not well reported by sites and is not presented. PROMs were collected only in the subset of people also in the RADICALS-RT trial;^{11,12} this included a small number of people, so PROMs are not analysed here.

Safety analyses were done according to allocated treatment, categorised as Radiation Therapy Oncology Group grade 1, 2, or 3 or higher, and analysed with a χ^2 test. Worst grade of adverse events includes routine-assessed toxicities and serious adverse events.

Statistical analysis

This comparison was originally designed as part of a three-way comparison with disease-specific survival as the primary outcome measure and an overall recruitment target of 3053 patients across three arms. However, recruitment was permitted between pairs of arms to facilitate recruitment in the pilot phase and this method was subsequently maintained. Therefore, in 2010, the trial was repowered for separate comparisons of short-course ADT versus long-course ADT (reported separately¹³) and no ADT versus short-course ADT. This was done without any reference to accumulating comparative data within the trial. This separated, pairwise comparison required approximately 1263 patients to observe 128 events.

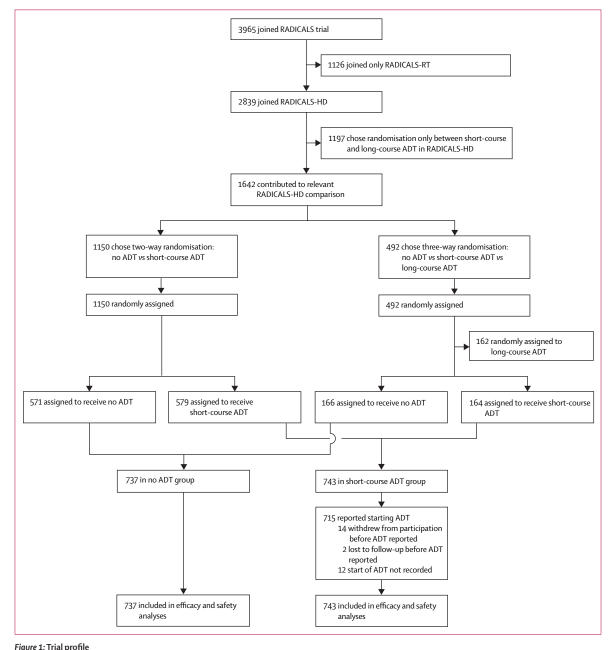
After recruitment and treatment had been completed for all participants, the primary outcome measure was changed in 2019 to metastasis-free survival. This change followed new evidence from the ICECaP study¹⁴ that metastasis-free survival was a robust early surrogate outcome measure for disease-specific survival. This decision did not involve anyone privy to accumulating comparative data from RADICALS-HD.¹⁵ With 200 events from the 1480 participants, this final design had 80% power with two-sided α of 5% to detect an increase in 10-year metastasis-free survival from 80% to 86% (HR=0.67). Based on previous experiences with long-term trials, the calculations conservatively assumed that follow-up might be truncated early

Correspondence to: Prof Matthew R Sydes, MRC Clinical Trials Unit at UCL, Institute of Clinical Trials and Methodology, University College London, London WC1V 9LJ, UK mrcctu.radicals@ucl.ac.uk

See Online for appendix

For the **study protocol** see https://www.mrcctu.ucl.ac.uk/ media/1811/radicals-protocolversion-60-14-dec-2018_signed. pdf between 5 years and 10 years after randomisation for up to 30% of participants.

The full statistical analysis plan is published elsewhere¹⁰ and the details are summarised here. All analyses followed the intention-to-treat principle, with analyses according to allocated group, and were conducted in Stata version 17.0. Follow-up was estimated through reverse censoring on death. For time-to-event outcome measures, the statistical significance of differences between groups was evaluated with the log-rank test, stratified by randomisation stratification factors. Effect estimates were obtained from Cox regression models, also stratified by randomisation stratification factors. The Grambsch–Therneau test was used to test the proportional hazards assumption, with restricted mean survival time becoming the primary estimate of effect if nonproportional hazards were detected, with time restricted (t*) to 10 years. Time-to-event graphs were presented in KMunicate format.¹⁶ When a formal date for stopping ADT was not reported, patients were not included in summaries of time on treatment. Competing risks regression was used for prostate cancer-specific survival



ADT=androgen deprivation therapy.

with other causes of death as competing risks. p<0.05 was considered to indicate statistical significance.

Two prespecified subgroup analyses were planned, by pre-radiotherapy PSA value and by Charlson Comorbidity Index score (excluding age), testing for interaction in the treatment effect." It was hypothesised that patients with higher pre-radiotherapy PSA, and those with less comorbidity, would benefit more from ADT. Exploratory subgroup analysis of all randomisation stratification factors was also planned.

The Independent Data Monitoring Committee (IDMC) met to review data from RADICALS on ten occasions. There were no formal stopping guidelines; the IDMC were asked to give advice on whether the accumulating data from the trial, together with results from other relevant trials, justified continuing recruitment of further patients or further follow-up. The IDMC did not recommend stopping the trial early.

Role of the funding source

The funders of the study had no role in study design (other than organising initial peer review by independent reviewers), data collection, data analysis, data interpretation, or writing of the report. The sponsor took responsibility for these elements, delegated through their staff.

Results

Between Nov 22, 2007, and June 29, 2015, 1480 patients were randomly assigned to receive no ADT (n=737) or short-course (6 months) ADT (n=743) in addition to postoperative radiotherapy at 121 trial-accredited centres in Canada, Denmark, Ireland, and the UK. (figure 1). Of those, 1150 joined who had been randomly assigned between only these two groups and 330 joined who had been allocated to one of these two groups as part of the three-way randomisation that also included long-course ADT.

The median age of participants was 66 years (IQR 61–69); 1267 (86%) of 1478 had a Gleason score of 7 or higher and 240 (16%) of 1471 had stage T3B disease or higher (table 1). Data on race and ethnicity were not collected. Radiotherapy was initiated in the early salvage setting in 1057 (71%) of 1480 participants. The planned radiotherapy schedule was 66 Gy in 33 fractions for 1021 (69%) of 1480 participants, and the radiotherapy target was the prostate bed only for 1392 (94%) of 1480.

Follow-up at sites for the trial ended on Dec 31, 2021: 1209 of 1480 patients were still in follow-up at that date, 190 were known to have died, and 81 had stopped their participation or been lost to follow-up. Median follow-up was $9 \cdot 0$ years (IQR $7 \cdot 1 - 10 \cdot 1$). Among those still in active follow-up at the end of the trial, minimum follow-up was $4 \cdot 5$ years. The database was locked on May 27, 2022.

In the short-course ADT group, median time from randomisation to starting hormone treatment was 7 days

(IQR 1–14). 28 patients were allocated to short-course ADT with no record of starting treatment, 14 of whom formally withdrew their participation from the trial. A formal date for stopping hormone treatment was not reported for 80 of the patients allocated to short-course ADT.

Metastasis-free survival events were reported for 268 patients, including 142 in the no ADT group and 126 in the short-course ADT group; 78 patients (44 no ADT and 34 short-course ADT) developed metastases but

	No ADT group (n=737)	Short-course ADT All (n=743) (n=1480)			
Age, years	66 (61–69)	66 (61–69)	66 (61-69)		
PSA at randomisation, ng/mL	0.22 (0.12-0.40)	0.20 (0.10-0.40)	0.21 (0.11-0.40)		
<0.3	434 (59%)	471 (63%)	905 (61%)		
0·3 to <0·5	151 (20%)	140 (19%)	291 (20%)		
≥0.5	152 (21%)	132 (18%)	284 (19%)		
Gleason score					
<7	102 (14%)	109 (15%)	211 (14%)		
3+4	345 (47%)	359 (48%)	704 (48%)		
4+3	205 (28%)	189 (25%)	394 (27%)		
>7	83 (11%)	86 (12%)	169 (11%)		
Missing	2	0	2		
T stage					
1 or 2	298 (41%)	305 (41%)	603 (41%)		
3a	325 (44%)	303 (41%)	628 (43%)		
3b or 3c	109 (15%)	122 (17%)	231 (16%)		
4	3 (<1%)	6 (1%)	9 (1%)		
Missing	2	7	9		
Lymph node involvement					
Node negative	418 (57%)	394 (53%)	812 (55%)		
Node positive	25 (3%)	25 (3%)	50 (3%)		
No dissection	292 (40%)	319 (43%)	611 (41%)		
Missing	2	5	7		
Positive margins					
Absent	285 (39%)	271 (36%)	556 (38%)		
Present	452 (61%)	472 (64%)	924 (62%)		
CAPRA-S score		,			
Low (0–2)	111 (15%)	110 (15%)	221 (15%)		
Intermediate (3–5)	386 (53%)	379 (51%)	765 (52%)		
High (≥6)	235 (32%)	249 (34%)	484 (33%)		
Missing	5	5	10		
Charlson Comorbidity Index score	-	-			
0	592 (81%)	602 (82%)	1194 (81%)		
1	106 (14%)	99 (13%)	205 (14%)		
≥2	36 (5%)	37 (5%)	73 (5%)		
Missing	3	5	8		
Country	- -				
UK	584 (79%)	578 (78%)	1162 (79%)		
Canada	145 (20%)	156 (21%)	301 (20%)		
Denmark	6 (1%)	8 (1%)	14 (1%)		
Ireland	2 (<1%)	1 (<1%)	3 (<1%)		
		(Table 1 continues on next page)			

	No ADT group (n=737)	Short-course ADT (n=743)	All (n=1480)
(Continued from previous page)			
Timing of radiotherapy			
Adjuvant	208 (28%)	215 (29%)	423 (29%)
Early salvage	529 (72%)	528 (71%)	1057 (71%)
Planned radiotherapy schedule			
52.5 Gy in 20 fractions	215 (29%)	222 (30%)	437 (30%)
66.0 Gy in 33 fractions	510 (69%)	511 (69%)	1021 (69%)
Other	12 (2%)	10 (1%)	22 (1%)
Planned radiotherapy target			
Prostate bed	700 (95%)	692 (93%)	1392 (94%)
Prostate bed and lymph nodes	37 (5%)	51 (7%)	88 (6%)
Planned hormone therapy			
Gonadotropin-releasing hormone analogue	613 (83%)	624 (84%)	1237 (84%)
Bicalutamide	124 (17%)	119 (16%)	243 (16%)

Data are median (IQR), n (%), or n. Percentages were calculated using the number of participants with available data as the denominator. ADT=androgen deprivation therapy. PSA=prostate-specific antigen. CAPRA-S=Cancer of the Prostate Risk Assessment Post-Surgical.

Table 1: Participant characteristics and pre-randomisation planned treatment

were still alive at the end of the trial, 59 patients (31 no ADT and 28 short-course ADT) had metastases followed by death, and 131 (67 no ADT and 64 short-course ADT) died without having metastases reported. There was no evidence that metastasis-free survival was improved in patients allocated to short-course ADT compared with those allocated to no ADT (HR 0.886 [95% CI 0.688-1.140], p=0.35; table 2, figure 2). There was no evidence of non-proportional hazards in the treatment effect. 10-year metastasis-free survival was 79.2% (95% CI 75.4-82.5) in the no ADT group and 80.4% (76.6-83.6) in the short-course ADT group. The metastasis-free survival treatment effect did not differ meaningfully in either of the two prespecified subgroup analyses, preradiotherapy PSA level (interaction p=0.68) and Charlson Comorbidity Index score (interaction p=0.29; figure 3), nor by any of the exploratory randomisation stratification factors (appendix p 2).

The secondary outcome measures in which there was clear evidence for a benefit of short-course ADT compared with no ADT were clinical progression-free survival and time to non-protocol (salvage) ADT. For clinical progression-free survival, the HR was 0.544

	No ADT group (n=737)	Short-course ADT group (n=743)	HR (95% CI)*	Log-rank p value*	Proportiona hazards p value†
Metastasis-free survival					
Events	142	126	0.886 (0.688–1.140)	0.35	0.71
Metastases first	75	62			
Prostate cancer death first	4	3			
Death from other causes first	63	61			
RMST (95% CI)‡	9.06 (8.90–9.22)	9.18 (9.03–9.33)			
10-year metastasis-free survival (95% CI)	79·2% (75·4–82·5)	80.4% (76.6–83.6)			
Overall survival					
Events	98	92	0.882 (0.651–1.194)	0.42	0.34
RMST (95% CI)‡	9·45 (9·33–9·57)	9.47 (9.35-9.59)			
10-year overall survival (95% CI)	85.6% (82.2-88.4)	85.3% (81.7-88.3)			
Freedom from distant metastasis					
Events	79	65	0.816 (0.579–1.150)	0.24	0.60
RMST (95% CI)‡	9·39 (9·26–9·53)	9.53 (9.41–9.65)			
10-year freedom from distant metastasis (95% CI)	88.1% (85.0-90.6)	89.9% (87.0-92.2)			
Clinical progression-free survival					
Events	210	131	0.544 (0.433-0.684)	<0.0001	0.071
RMST (95% CI)‡	8.16 (7.94-8.38)	8.96 (8.79–9.14)			
10-year clinical progression-free survival (95% CI)	68·3% (64·3–71·9)	79.4% (75.8–82.6)			
Time to non-protocol ADT					
Events	176	109	0.543 (0.422-0.699)	<0.0001	0.0041
RMST (95% CI)‡	8.45 (8.24-8.66)	9.17 (9.01–9.32)			
10-year freedom from non-protocol ADT (95% CI)	73·3% (69·5–76·7)	82.3% (78.7-85.3)			

ADT=androgen deprivation therapy. HR=hazard ratio. RMST=restricted mean survival time. *Adjusted for randomisation stratification factors. †Grambsch-Therneau test of non-proportional hazards. ‡Restricted to 10 years.

Table 2: Primary and secondary outcome measures

(95% CI 0·433–0·684; log-rank p<0·0001), although with some evidence of non-proportional hazards. For time to non-protocol ADT, the HR was 0·543 (0·422–0·699; log-rank p<0·0001) but with clear evidence of nonproportional hazards, so this is better summarised as improved restricted mean event-free time from 8·45 years (95% CI 8·24–8·66) to 9·17 years (9·01–9·32) with short-course ADT. We found no evidence for a benefit of short-course ADT for overall survival, nor for freedom from distant metastasis, which only included deaths from prostate. In a competing-risks regression model with non-prostate cancer death as the competing risk, the sub-HR was 0·814 (95% CI 0·586–1·130; p=0.22). Causes of death are presented in the appendix (p 3).

During follow-up, Radiation Therapy Oncology Group scale toxicity of grade 3 or higher was reported for 121 (17%) of 737 participants in the no ADT group and 100 (14%) of 743 in the short-course ADT group (p=0.15; table 3). The most commonly reported toxicities of grade 3 or higher were urethral stricture and haematuria. Of the serious adverse events reported, 18 (11 in the no ADT group and seven in the short-course ADT group) were reported as serious adverse reactions and two (both in the short-course ADT group) were reported as suspected unexpected serious adverse reactions (appendix p 4).

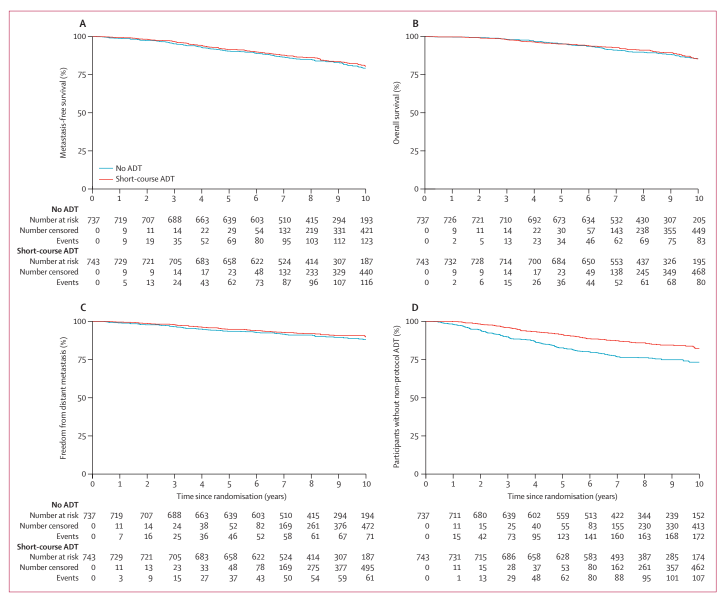


Figure 2: Primary and secondary outcome measures

(A) Metastasis-free survival. (B) Overall survival. (C) Freedom from distant metastasis. (D) Time to non-protocol ADT. Risk tables present the number of participants who, at each timepoint, remain at risk, have been censored, or have had an event. All timepoints add up to the total number of patients. ADT=androgen deprivation therapy.

	No ADT group, n/N	Short-course ADT group, n/N		Hazard ratio (95% CI)	p value
Charlson Comor	bidity Index score				
0	117/592	98/602		0.82 (0.62–1.07)	
1	19/106	19/99	•	1.05 (0.56–1.99)	0.29
≥2	5/36	8/37		1.85 (0.59–5.77)	
Pre-radiotherap	y PSA, ng/mL				
<0.3	76/434	66/471		0.81 (0.59–1.13)	
0.3-0.5	29/151	30/140		1.04 (0.63–1.74)	0.68
>0.5	37/152	30/132		0.96 (0.59–1.55)	
Overall	142/737	126/743	$\langle \rangle$	0.88 (0.69-1.12))
			0.5 1 2		

Figure 3: Pre-planned subgroup analyses

Weighting is by sample size. Age did not contribute to the Charlson Comorbidity Index score. ADT=androgen deprivation therapy. PSA=prostate-specific antigen.

	No ADT group (n=737)			Short-course ADT group (n=743)			p value
	Grade 1–2	Grade 3	Grade 4	Grade 1–2	Grade 3	Grade 4	
Any toxicity	407 (55%)	114 (15%)	7 (1%)	442 (59%)	90 (12%)	10 (1%)	0.17
Diarrhoea	307 (42%)	12 (2%)	1 (<1%)	322 (43%)	13 (2%)	0	0.67
Proctitis	200 (27%)	23 (3%)	0	248 (33%)	20 (3%)	0	0.028
Cystitis	199 (27%)	15 (2%)	2 (<1%)	214 (29%)	17 (2%)	2 (<1%)	0.84
Haematuria	141 (19%)	51 (7%)	2 (<1%)	155 (21%)	28 (4%)	2 (<1%)	0.058
Urethral stricture	74 (10%)	49 (7%)	3 (<1%)	74 (10%)	51 (7%)	7 (1%)	0.65
Data are presented as n (%). Data were missing for six participants in the no ADT group and nine participants in the short-course ADT group. No grade 5 events were recorded. ADT=androgen deprivation therapy.							

Table 3: Maximum toxicity grade reported on Radiation Therapy Oncology Group scales

Five deaths were reported as serious adverse events; none were considered as definitely, probably, or possibly related to trial treatment on clinical review.

Discussion

In this randomised controlled trial, the addition of 6 months of ADT to postoperative prostate bed radiotherapy did not improve metastasis-free survival, but it did delay the time to salvage ADT. These data should inform clinical decision making; 6 months of ADT upfront, at the time of postoperative radiotherapy, improved the 10-year freedom from salvage ADT from $73 \cdot 3\%$ (95% CI 69.5–76.7) to $82 \cdot 3\%$ (78.7–85.3). However, given that it had no meaningful impact on metastasis-free survival, short-course ADT is unlikely to improve overall survival in this setting.

Two other randomised controlled trials have tested the use of short-course ADT with postoperative radiotherapy to the prostate bed. GETUG-AFU 16 was a randomised controlled trial for patients with PSA progression after radical prostatectomy with a primary outcome measure named as progression-free survival.² However, that outcome measure could have been termed biochemical progression-free survival, because the definition of progression included PSA progression. 743 patients were randomly assigned to prostate bed radiotherapy with or without two doses of goserelin 10.8 mg given 3 months

apart. In an updated analysis, at a median follow-up of 9.3 years,² 10-year progression-free survival was 49% for radiotherapy alone and 64% for radiotherapy plus shortcourse ADT (HR 0.54 [95% CI 0.43-0.68]; p<0.0001). Metastasis-free survival was not a prespecified outcome measure of the trial, but 10-year metastasis-free survival was 69% for radiotherapy alone and 75% for radiotherapy plus short-course ADT (HR 0.73 [0.54–0.98]; p=0.039). No difference in overall survival was reported, with 85% and 86% alive at 10 years, respectively (HR 0.93 [0.63-1.39]; p=0.73). Similarly, RTOG 0534 was a trial that enrolled people with PSA progression after radical prostatectomy.3 Its primary outcome measure was freedom from progression and, like GETUG-AFU 16, that definition of progression included increasing PSA as well as clinical progression or death, with PSA driving the event count. 1194 men were randomly assigned to prostate bed radiotherapy with or without 4-6 months of ADT. With a median follow-up of 8.2 years,3 5-year freedom from progression was 71% for radiotherapy alone and 81% for radiotherapy plus short-course ADT (HR 0.64 [0.50-0.82]; p<0.0001). There was no clear evidence of a benefit for adding short-course ADT in terms of distant metastases (0.78 [0.52-1.17]; p=0.083), prostate cancer death (0.79 [0.45-1.38]; p=0.17), or overall survival (0.89 [0.60-1.31]; p=0.245). The results from GETUG-AFU 16 and RTOG 0534 will be combined with those from RADICALS-HD in the DADSPORT meta-analysis (registered PROSPERO. on CRD42022325769), which will provide evidence concerning the effect of short-course ADT on long-term clinical outcomes in this setting. We note that all three trials are consistent with the possibility of an improvement in metastasis-free survival. The DADSPORT meta-analysis will also study the optimum duration of ADT, including the short-course versus longcourse ADT comparison from RADICALS-HD.13

The design of RADICALS-HD differed from GETUG-AFU 16 and RTOG 0534 with regard to the primary outcome measure. Our view was that shortcourse ADT would be expected to delay PSA progression after postoperative radiotherapy, and yet such a positive finding should not be sufficient to justify its use. Rather, the case for or against short-course ADT should rest on more clinically meaningful outcome measures. When RADICALS-HD was first designed, the primary outcome measure was disease-specific survival. This was later amended to metastasis-free survival based on the work of ICECaP.¹⁴ The full details of this change and the broader history of RADICALS are presented elsewhere.¹⁵ The findings from ICECaP14 demonstrated metastasis-free survival to be a useful intermediate outcome measure in trials of non-metastatic prostate cancer: unless a substantial effect is seen on metastasis-free survival, it is unlikely that the intervention will improve overall survival. Therefore, given that these three trials have not demonstrated a substantial impact on metastasis-free

survival for the addition of short-course ADT, we believe that prostate bed radiotherapy alone remains a standard treatment option.

The most promising findings of the RTOG 0534 trial were in the group that received not only the addition of short-course ADT but also pelvic nodal irradiation. In comparison with prostate bed radiotherapy alone, treatment to the prostate bed and the pelvic lymph nodes plus short-course ADT gave the best point estimate for metastasis-free survival benefit (HR 0.79 [95% CI 0.57-1.08], p=0.042) but this did not reach their threshold for statistical significance (p<0.0125). In current clinical practice, we believe that there is a choice to be made between prostate bed radiotherapy alone versus the combination of radiotherapy to the prostate bed and pelvic lymph node plus ADT. Patients prioritising safety might opt for prostate bed radiotherapy alone. Patients willing to accept the increased risk of adverse effects in the hope of greater efficacy might opt for radiotherapy to the prostate bed and the pelvic lymph nodes plus ADT.

The adverse effects of ADT are already well characterised. Therefore, this trial reduced the data collection burden on sites by focusing only on possible radiotherapy side-effects using the Radiation Therapy Oncology Group scale. We believe the focused collection of data and the use of follow-up schedules that reflected normal clinical practice facilitated the conduct of RADICALS-HD. This allowed the trial to recruit sufficient participants to be powered on long-term, clinically meaningful outcome measures.

RADICALS-HD has several limitations. First, the large majority of patients received radiotherapy to the prostate bed alone, whereas results from RTOG 0534 have shown some support for radiotherapy to the pelvic nodes in addition to the prostate bed.3 It remains unclear whether short-course ADT might be beneficial in patients receiving pelvic nodal radiotherapy. Second, RADICALS-HD included patients receiving radiotherapy both in the salvage setting, like in GETUG-AFU 16 and RTOG 0534, and also in the adjuvant setting. Based on the previous results of RADICALS-RT,15 and the ARTISTIC metaanalysis,¹⁸ postoperative radiotherapy is now typically given in the salvage, rather than the adjuvant, setting. Given that some patients receiving adjuvant radiotherapy would have been cured by surgery alone, a lower event rate would be expected than in the salvage radiotherapy setting. In this analysis, we found no evidence of a differential effect for short-course ADT according to the timing of radiotherapy. The DADSPORT meta-analysis (CRD42022325769) of these three trials will include a sensitivity analysis, restricted to those treated in the salvage setting. Third, the good prognosis of this patient group makes clinical trials difficult. Although the trial opened more than 15 years ago, and the comparison accrued around 1500 patients, there are too few events for RADICALS-HD to reliably report on any effect on overall survival or cancer-specific survival. However, given the lack of improvement in metastasis-free survival, it is very unlikely that any benefit will emerge in terms of overall or cause-specific survival. Fourth, although we did a prespecified subgroup analysis by baseline comorbidity score, only a small proportion of the patients had a Charlson Comorbidity Index score of 2 or higher. We cannot be confident that the trial results are generalisable to patients with substantial comorbidity. The trial was in active follow-up during the COVID-19 pandemic from 2020 onwards. Recruitment had been completed many vears previously so neither accrual nor the allocated treatment would have been affected. There is no good reason to think follow-up would have been impacted separately by allocated treatment group during the pandemic. Fifth, data were not collected on ethnicity and race, so we cannot comment on how well the participants reflect the underlying population, especially in light of well known differences in prevalence;19 the trial would not have been powered to look reliably for consistency of effect by ethnicity and race.

The trial was run when bone scan and CT scan were regarded as the standard imaging modalities. In recent years, new imaging techniques, such as prostate-specific membrane antigen (PSMA) PET, have become available, and it is interesting to consider the impact that this might have had on interpreting the trial results. PSMA PET is more sensitive than conventional imaging techniques and this would tend to bring forward the detection of metastatic disease during follow-up and increase the event rate. However, patients with detectable metastatic disease at baseline were excluded from the trial, and so the use of PSMA PET would lead to a lowerrisk trial population and a lower event rate. We do not envisage that the use of new imaging techniques would have had a differential impact between the two trial groups.

In conclusion, RADICALS-HD found that the addition of 6 months of ADT to postoperative prostate bed radiotherapy did not improve metastasis-free survival, although it did delay the time to salvage ADT. This reduction in salvage ADT could be considered insufficient to justify 6 months of ADT at the time of radiotherapy. However, it is unclear whether the reduction in salvage ADT means that some patients will avoid the need for salvage ADT entirely, or whether the start of salvage ADT is merely being delayed. In our view, these findings are not sufficient to routinely recommend the use of shortcourse ADT with postoperative radiotherapy.

Contributors

CCP was the chief investigator. CCP, MRS, NWC, HK, CNC, and MKBP were responsible for trial design. CCP, MRS, NWC, HK, and MKBP were grant holders in the UK and CNC and WRP were grant holders in Canada. CCP, NWC, ADC, HK, PMP, CNC, WRC, JL, WRP, LCB, SF, CM, HP, RAP, FS, MKBP, and MRS were members of the Trial Management Group. SF and CM were responsible for trial operations. CCP, ADC, MRS, and MKBP were responsible for the analysis plan. ADC and MRS did the final analyses, and checked and verified the data. CCP, ADC, and MRS wrote the first draft of the manuscript. All authors had full access to data, collated data, interpreted data, edited, reviewed, and approved the final manuscript, and had final responsibility for the decision to submit for publication. All authors affirm that the manuscript is an honest, accurate, and transparent account of the study being reported; that no important aspects of the study have been omitted; and that any discrepancies from the study as planned have been explained.

Declaration of interests AN reports honoraria for lectures, presentations, speakers bureaus,

manuscript writing, or educational events from Merck. ADC reports research grants for the STAMPEDE trial from Janssen, Astellas, Novartis, Sanofi, and Clovis. ACT reports research grants to their institution from Elekta, Varian, and Accuracy; honoraria for talks from Elekta, Accuracy, and Janssen; travel assistance for radiotherapy conferences from Elekta; serving on the data safety board for two academic trials; and roles as the Chair of the MR-Linac Consortium and the genitourinary lead editor for the International Journal of Radiation Oncology, Biology, Physics. AC reports receiving grants or contracts from the National Institute for Health and Care Research, Manchester Biomedical Research Centre, Cancer Research UK, Medical Research Council (MRC) UK, Prostate Cancer UK, Bayer UK, and Elekta; and payment or honoraria for lectures, presentations, speakers bureaus, manuscript writing, or educational events from the American Society for Radiation Oncology, the American Society of Clinical Oncology, Cancer Research UK, Roche, AstraZeneca, and Bristol Myers Squibb (BMS). CNC reports support for the present manuscript from the Canadian Cancer Trials Group; and payment or honoraria for lectures, presentations, speakers bureaus, manuscript writing, or educational events from Bayer Corp, Knight Therapeutics, and AbbVie. CLG reports participating in the independent data monitoring and steering committee for the BARCODE study at the Institute of Cancer Research in London (principal investigator Prof Ros Eeles). CCP reports consulting fees from AAA, ITM Radiopharma, Myovant, and Clarity Pharmaceuticals to his institution; and payment or honoraria for lectures, presentations, speakers bureaus, manuscript writing, or educational events from Janssen and Bayer to his institution. CM reports that they are employed by University College London (UCL) as Trial Manager for RADICALS. FS reports grants or contracts from Janssen, Bayer, Merck, Pfizer, Astellas, BMS, Novartis, Sanofi, and AstraZeneca to his institution; consulting fees from Janssen, Bayer, Astellas, BMS, Novartis, Sanofi, AstraZeneca, Merck, and Pfizer to him; and payment or honoraria for lectures, presentations, speakers bureaus, manuscript writing, or educational events from Janssen, Bayer, Myovant, Astellas, BMS, Novartis, Sanofi, AstraZeneca, Merck, and Pfizer to him. LCB reports their previous Institute of Cancer role was funded in part by a Biomedical Research Centre grant which was paid to the institution; the role was not connected to the RADICALS trial. MRS reports research grants and biomarker testing costs, all to their institution and all active in the past 36 months but on research outside of this research, from Astellas, Clovis Oncology, Janssen, Novartis, and Sanofi-Aventis; consulting fees from Eli Lilly; speaker fees at a clinical trial statistics training meeting for clinicians (no discussion of particular drugs) from Lilly Oncology, Janssen, and Eisai; and is an independent member of many independent data monitoring committees but all for academic sponsors and unpaid. NWC reports honoraria for lectures, advisory boards, and symposia from AstraZeneca, Janssen, Bayer, and Pfizer; support for travel to and attendance at a European meeting from Bayer; participation on the independent data monitoring committee for the Probio trial (Karolinska), and the trial steering committee for the Capi 28 trial (AstraZeneca) and the STAMPEDE trial (MRC); and is the Joint National Clinical Lead for the National Prostate Cancer Audit.

PWMC reports grants or contracts from the Canadian Association of

Radiation Oncology ACURA program and the Canadian Institute of

Health Research; payment or honoraria for lectures, presentations,

board for AAA Nordic, MSD, and Pfizer Denmark. SCM reports

speakers bureaus, manuscript writing, or educational events from Pfizer

and EMD Serono; and participation on a data safety monitoring board or

advisory board from TerSera, AbbVie, Tolmar, and Knight Therapeutics.

PMP reports participation on a data safety monitoring board or advisory

For more on the MRC Clinical Trials Unit's approach to data sharing see http://www.ctu.mrc. ac.uk/our_research/datasharing

For data sharing requests see http://www.ctu.mrc.ac.uk/our_ research/datasharing/ application_process honoraria from Janssen, Astellas, Knight Therapeutics, AMGEN Pharmaceutical, AstraZeneca, AbbVie, and Bayer; and support for attending meetings or travel from TerSera and Sanofi. VK reports a Cancer Research UK grant for the CORE trial to his institution; honoraria from Accuracy, Astellas, Bayer, and Boston Scientific; meeting attendance support from Accuray, Astellas, AstraZeneca, Bayer, BMS, Boston Scientific, Janssen, Merck Sharp & Dohme, and Norvatis; is an unpaid member of the independent data monitoring committee for the EPIC trial and the advisory board for Bayer and Janssen; and is the unpaid Chair of National Cancer Research Institute Penile subgroup of the Bladder and Renal Clinical Studies Group. WRC reports consulting fees from Bayer and Janssen; payment or honoraria for lectures, presentations, speakers bureaus, manuscript writing, or educational events from Astellas, Bayer, Janssen, AAA Novartis, and Myriad Genetics; and support for attending meetings or travel from Janssen, AAA Novartis, and Bayer. All other authors declare no competing interests.

Data sharing

The RADICALS trial data are held at the MRC Clinical Trials Unit at UCL, which encourages optimal use of data by using a controlled access approach to data sharing. Requests for data can be made at any time and can be initiated by email to mrcctu.datareleaserequest@ucl.ac.uk or via our website. There is a formal application process, whereby the request will undergo review by the trial team, as well as independent researchers, to ensure that the proposed research is both ethical and has a strong scientific rationale. Data will not be released if this would compromise the ongoing research. The specific data and associated documents to be shared will be dependent on the nature of the individual request and this will be documented in a formal data sharing agreement.

Acknowledgments

We recognise the efforts of all trial team members at the trial units and hospitals who have supported and engaged with RADICALS. A list of investigators and oversight committee members across the comparisons of the RADICALS protocol is given in the appendix (pp 9–96). We particularly thank Peter Neville and Alan Stirling, the current patient and public involvement members of the Trial Management Group. We recognise and thank all of the participants of the trial and the families and friends who supported them. Clinical trials only happen because people choose to join them. Grant funding in the UK was provided by the Clinical Trials Advisory Award Committee on behalf of Cancer Research UK (UK/C7829/A6381). Funding in Canada was provided by the Canadian Cancer Society (704970). The trial was further supported at the MRC Clinical Trials Unit at UCL by a core grant from the MRC, now part of UK Research and Innovation (MC_UU_12023/28). UK sites were part of the Health Research Clinical Research Network. AC and PH are supported by the National Institute for Health Research (NIHR) Manchester Biomedical Research Centre. This Article represents independent research funded in part by the NIHR Biomedical Research Centre at the Royal Marsden NHS Foundation Trust and the Institute of Cancer Research. The views expressed are those of the authors and not necessarily those of the NIHR or the Department of Health and Social Care.

References

- Kishan AU, Sun Y, Hartman H, et al. Androgen deprivation therapy use and duration with definitive radiotherapy for localised prostate cancer: an individual patient data meta-analysis. *Lancet Oncol* 2022; 23: 304–16.
- 2 Carrie C, Magné N, Burban-Provost P, et al. Short-term androgen deprivation therapy combined with radiotherapy as salvage treatment after radical prostatectomy for prostate cancer (GETUG-AFU 16): a 112-month follow-up of a phase 3, randomised trial. *Lancet Oncol* 2019; 20: 1740–49.
- 3 Pollack A, Karrison TG, Balogh AG, et al. The addition of androgen deprivation therapy and pelvic lymph node treatment to prostate bed salvage radiotherapy (NRG Oncology/RTOG 0534 SPPORT): an international, multicentre, randomised phase 3 trial. *Lancet* 2022; 399: 1886–901.
- 4 Taylor LG, Canfield SE, Du XL. Review of major adverse effects of androgen-deprivation therapy in men with prostate cancer. *Cancer* 2009; 115: 2388–99.

- 5 Lee LW, Clarke NW, Ramani VA, Cowan RA, Wylie JP, Logue JP. Adjuvant and salvage treatment after radical prostatectomy: current practice in the UK. *Prostate Cancer Prostatic Dis* 2005; 8: 229–34.
- 6 Morris SL, Parker C, Huddart R, Horwich A, Dearnaley D. Current opinion on adjuvant and salvage treatment after radical prostatectomy. *Clin Oncol (R Coll Radiol)* 2004; 16: 277–82.
- 7 Parker C, Castro E, Fizazi K, et al. Prostate cancer: ESMO Clinical Practice Guidelines for diagnosis, treatment and follow-up. Ann Oncol 2020; 31: 1119–34.
- 8 Spratt DE. Evidence-based risk stratification to guide hormone therapy use with salvage radiation therapy for prostate cancer. *Int J Radiat Oncol Biol Phys* 2018; **102**: 556–60.
- 9 Cox JD, Stetz J, Pajak TF. Toxicity criteria of the Radiation Therapy Oncology Group (RTOG) and the European Organization for Research and Treatment of Cancer (EORTC). Int J Radiat Oncol Biol Phys 1995; 31: 1341–46.
- 10 Parker CC, Parmar MKB, Sydes MR, Cook A. RADICALS trial statistical analysis plan. May 27, 2022. https://doi.org/10.5281/ zenodo.6586525 (accessed May 1, 2024).
- 11 Parker CC, Petersen PM, Cook AD, et al. Timing of radiotherapy (RT) after radical prostatectomy (RP): long-term outcomes in the RADICALS-RT trial [NCT00541047]. Ann Oncol 2024; published online May 16. https://doi.org/10.1016/j.annonc.2024.03.010.
- 12 Parker CC, Clarke NW, Cook AD, et al. Timing of radiotherapy after radical prostatectomy (RADICALS-RT): a randomised, controlled phase 3 trial. *Lancet* 2020; 396: 1413–21.

- 13 Parker CC, Kynaston H, Cook AD, et al. Duration of androgen deprivation therapy with postoperative radiotherapy for prostate cancer: a comparison of long-course versus short-course androgen deprivation therapy in the RADICALS-HD randomised trial. *Lancet* 2024; published online May 16. https://doi.org/ S0140-6736(24)00549-X.
- 14 Xie W, Regan MM, Buyse M, et al. Metastasis-free survival is a strong surrogate of overall survival in localized prostate cancer. *J Clin Oncol* 2017; **35**: 3097–104.
- 15 Parker CC, Clarke NW, Catton C, et al. RADICALS-HD: reflections before the results are known. *Clin Oncol (R Coll Radiol)* 2022; 34: 593–97.
- 16 Morris TP, Jarvis CI, Cragg W, Phillips PPJ, Choodari-Oskooei B, Sydes MR. Proposals on Kaplan–Meier plots in medical research and a survey of stakeholder views: KMunicate. *BMJ Open* 2019; 9: e030215.
- 17 Charlson ME, Pompei P, Ales KL, MacKenzie CR. A new method of classifying prognostic comorbidity in longitudinal studies: development and validation. *J Chronic Dis* 1987; 40: 373–83.
- 18 Vale CL, Fisher D, Kneebone A, et al. Adjuvant or early salvage radiotherapy for the treatment of localised and locally advanced prostate cancer: a prospectively planned systematic review and meta-analysis of aggregate data. *Lancet* 2020; **396**: 1422–31.
- 19 James ND, Tannock I, N'Dow J, et al. The *Lancet* Commission on prostate cancer: planning for the surge in cases. *Lancet* 2024; 403: 1683–722.