Patient-reported outcomes in the ProtecT randomized trial of clinically localized prostate cancer treatments: study design, and baseline urinary, bowel and sexual function and quality of life


*School of Social and Community Medicine, University of Bristol, Bristol, †Bristol Randomised Trials Collaboration, University of Bristol, Bristol, ‡School of Clinical Sciences, University of Bristol, Bristol, §Collaboration for Leadership in Applied Health Research and Care West, United Hospitals Bristol, Bristol, ¶Clinical Practice Research DataLink Group, Medicines and Healthcare Products Regulatory Agency, London, **School of Medicine, Cardiff University, Cardiff, and ††Nuffield Department of Surgery, University of Oxford, Oxford, UK

A.L. and C.M. contributed equally to the paper.
ProtecT Study group members are given in Appendix 1.

Objectives
To present the baseline patient-reported outcome measures (PROMs) in the Prostate Testing for Cancer and Treatment (ProtecT) randomized trial comparing active monitoring, radical prostatectomy and external-beam conformal radiotherapy for localized prostate cancer and to compare results with other populations.

Materials and Methods
A total of 1643 randomized men, aged 50–69 years and diagnosed with clinically localized disease identified by prostate-specific antigen (PSA) testing, in nine UK cities in the period 1999–2009 were included. Validated PROMs for disease-specific (urinary, bowel and sexual function) and condition-specific impact on quality of life (Expanded Prostate Index Composite [EPIC], 2005 onwards; International Consultation on Incontinence Questionnaire-Urinary Incontinence [ICIQ-UI], 2001 onwards; the International Continence Society short-form male survey [ICSmaleSF]; anxiety and depression (Hospital Anxiety and Depression Scale [HADS]), generic mental and physical health (12-item short-form health survey [SF-12]; EuroQol quality-of-life survey, the EQ-5D-3L) were assessed at prostate biopsy clinics before randomization. Descriptive statistics are presented by treatment allocation and by men’s age at biopsy and PSA testing time points for selected measures.

Results
A total of 1438 participants completed biopsy questionnaires (88%) and 77–88% of these were analysed for individual PROMs. Fewer than 1% of participants were using pads daily (5/754). Storage lower urinary tract symptoms were frequent (e.g. nocturia 22%, 312/1423). Bowel symptoms were rare, except for loose stools (16%, 118/754). One third of participants reported erectile dysfunction (241/735) and for 16% (118/731) this was a moderate or large problem. Depression was infrequent (80/1399, 6%) but 20% of participants (278/1403) reported anxiety. Sexual function and bother were markedly worse in older men (65–70 years), whilst urinary bother and physical health were somewhat worse than in younger men (49–54 years, all P < 0.001). Bowel health, urinary function and depression were unaltered by age, whilst mental health and anxiety were better in older men (P < 0.001). Only minor differences existed in mental or physical health, anxiety and depression between PSA testing and biopsy assessments.

Conclusion
The ProtecT trial baseline PROMs response rates were high. Symptom frequencies and generic quality of life were similar to those observed in populations screened for prostate cancer and control subjects without cancer.

Keywords
prostate cancer, treatment, functional status, quality of life, protect trial, ISRCTN 20141297
Introduction

Prostate cancer is the most commonly detected male malignancy in many countries and accounted for 10,837 deaths in the UK in 2012 (Cancer Research UK: http://www.cancerresearchuk.org/health-professional/cancer-statistics/statistics-by-cancer-type/prostate-cancer). Prostate cancer screening with PSA has been shown to identify predominantly clinically localized disease in two randomized trials of screening [1,2]. The main treatments for localized prostate cancer are active surveillance, brachytherapy, external-beam or intensity-modulated radiotherapy and open or robot-assisted radical prostatectomy [3]. The optimum treatment that balances the risks of the intervention and disease control remains uncertain, although two randomized trials of surgery and watchful waiting (supportive care with treatment of symptoms as required) showed a disease-specific survival benefit for radical intervention in the Scandinavian SPCG-4 trial [4], but only for some men in the US PIVOT trial [5].

Clinicians and patients selecting prostate cancer treatments need robust information to enable them to balance symptom and broader impact on quality of life (QoL) against mortality and disease progression outcomes. Validated patient-reported outcome measures (PROMs) [6] are the recommended tools to assess the specific impacts of prostate cancer treatment on incontinence, urinary irritation and obstruction, bowel-related symptoms, sexual function and hormone therapy [7,8]; however, there has been limited use of validated PROMs in localized prostate cancer trials, although recent radiotherapy trials have incorporated PROMs [9]. One of the comparative trials of surgery and watchful waiting for localized disease measured symptoms with a Scandinavian questionnaire [10], whilst the other trial used single items for urinary symptoms, bowel and sexual function [5].

The UK National Institute for Health Research (NIHR) Prostate Testing for Cancer and Treatment (ProtecT) trial compares the effectiveness of active monitoring (a surveillance strategy), radical prostatectomy and external-beam conformal radiotherapy with neoadjuvant androgen suppression for localized prostate cancer. The primary outcome of prostate cancer-specific and intervention-related mortality will be reported at a median 10 years’ follow-up in 2016. The present paper reports the baseline PROMs results to assess their comparability with other studies and to investigate the generalizability of the randomized population, which will assist interpretation of the trial outcome results.

Materials and Methods

Study Design and Participants

The ProtecT trial design, baseline socio-demographic and clinical results have been published elsewhere (Lane et al.; protocol (http://www.nets.nihr.ac.uk/projects/hta/962099)). A feasibility pilot trial (recruitment: 1999–2001) preceded the main trial (recruitment: 2001–2009) [11]. In brief, the trial aimed to establish the effectiveness of radical prostatectomy, external-beam conformal radiotherapy or active monitoring for men with clinically localized prostate cancer. Men aged 50–69 years were invited for PSA testing in primary care, and those with a PSA level ≥3.0 to 19.95 ng/mL proceeded to further diagnostic tests, including prostate biopsies. Men with clinically localized prostate cancer were eligible for randomization if there were no major contra-indications for the radical treatments (exclusions detailed previously) [11]. In total, 2,417 participants were identified with localized disease (of 82,849 men PSA tested) and, of these, 1,643 participants were randomized to three treatments. The primary outcome of definite or probable prostate cancer mortality (including intervention-related-mortality) at a median of 10 years’ follow-up and the secondary clinical, health economic and patient-reported outcomes will be published in 2016 (trial registration: ISRCTN 20141297, http://www.isrctn.com, ClinicalTrials.gov NCT02044172).

Ethics

Approval was obtained from the UK National Health Service Health Research Authority Multicentre Research Ethics Committee (01/04/025).

Patient-Reported Outcomes Measures

Validated PROMs and their specific items (Table 1) were selected using literature reviews and qualitative interviews with participants [12] to capture the major impacts related to prostate cancer treatments: urinary symptoms of incontinence and erectile function after surgery; bowel function after radiotherapy and anxiety or depression from living with untreated cancer during active monitoring [7]. We also aimed to assess the effects of ageing on urinary symptoms over time and wider health issues using generic QoL measures. The trial focused on treatments for localized disease so the Expanded Prostate Index Composite (EPIC) hormonal domain was not used, but this did not impact on the use of other domains, which were scored separately [13]. The European Organisation for Research and Treatment of Cancer Quality of Life Questionnaire C-30 (EORTC-QLQC-30) [14], which has symptom and functional scales, generic QoL and single items, was used at 5 and 10 years’ follow-up (added in 2007).

Data Collection and Analysis

Participants completed paper questionnaires at initial PSA testing (trial recruitment) and at first prostate biopsy clinics, without reminders to non-responders (Fig. 1). The more comprehensive set of assessments completed at first biopsy...
Table 1 ProtecT trial patient-reported outcome measures and completion rates at first prostate biopsy by randomized participants.

<table>
<thead>
<tr>
<th>Outcome</th>
<th>PROM focus (timeframe)*</th>
<th>PROM scales/items presented (range/categories): [domain MID]</th>
<th>Completion by randomised participants (n = 1643) (n completed / n given the PROM)‡</th>
</tr>
</thead>
<tbody>
<tr>
<td>Urinary</td>
<td>ICIQ-UI: incontinence (4 weeks)</td>
<td>Incontinence score (0–21); [1.2] Number of continent men (score of 0, 0–100%) QoL impact (0–10, moderate: 2–6, large: 7–10)</td>
<td>76% (1244/1508)</td>
</tr>
<tr>
<td></td>
<td>EPIC: PCa treatment (4 weeks)</td>
<td>Summary, function and bother scores (0–100): [4.6, 4.2, 5.9] Incontinence and irritative/obstruction scales (0–100): [5.7, 4.6] Pad use (≥1/week)</td>
<td>82% (1244/1508)</td>
</tr>
<tr>
<td></td>
<td>ICSmaleSF: LUTS (4 weeks)</td>
<td>Incontinence score (0–24); [1.0] Voiding score (0–20); [1.5] Daytime frequency (≥2 h/void), nocturia (≥1/night), QoL impact (little, somewhat/a lot)</td>
<td>88% (748/849)</td>
</tr>
<tr>
<td>Bowel</td>
<td>EPIC: PCa treatment (4 weeks)</td>
<td>Summary, function and bother scores (0–100): [4.2, 4.4, 5.0] Bloody stools (ever), loose stools (≥50% of time), fecal incontinence (≥1/week), QoL impact (very small/small, moderate/big)</td>
<td>46% (748/849)</td>
</tr>
<tr>
<td>Sexual</td>
<td>EPIC: PCa treatment (4 weeks)</td>
<td>Summary, function and bother scores (0–100): [11.6, 11.5, 14.8] Erectile function (not firm enough for intercourse), QoL impact (very small/small, moderate/big)</td>
<td>44% (719/849)</td>
</tr>
<tr>
<td>Psychological</td>
<td>HADS: anxiety and depression (1 week)</td>
<td>Anxiety and depression scores (0–21); [1.8, 1.3] Potential clinical cases of anxiety or depression (≥8)</td>
<td>85% (1399/1643)</td>
</tr>
<tr>
<td>Physical</td>
<td>SF-12: overall mental health (4 weeks)</td>
<td>Mental component sub-score (0–100); [3.8]</td>
<td>77% (1260/1643)</td>
</tr>
<tr>
<td>Generic health</td>
<td>SF-12: overall physical health (4 weeks)</td>
<td>Physical component sub-score (0–100); [4.0]</td>
<td>77% (1260/1643)</td>
</tr>
<tr>
<td>Cancer-related Qol.</td>
<td>EQ-5D-3L: health utility (today)</td>
<td>Health utility score (0.594–1.0 scored as perfect health); [0.09]</td>
<td>86% (1413/1643)</td>
</tr>
</tbody>
</table>

EPIC, Expanded Prostate cancer Index Composite; EORTC-QLQ-C30, European Organisation for Research and Treatment of Cancer Quality of Life Questionnaire C-30; EQ-5D-3L, EuroQol quality-of-life survey; HADS, Hospital Anxiety and Depression Scale; ICIQ-UI, International Consultation on Incontinence Questionnaire-Urinary Incontinence; ICSmaleSF, International Continence Society short-form male survey; MID, minimal clinically important difference; PROM, patient-reported outcome measure; QoL, quality of life; SF-12, 12-item short-form health survey. *Questionnaire-specific recall period; †MID taken as 0.5 SD of baseline values; ‡excludes men with insufficient items to calculate scales.
were taken as the baseline for subsequent trial outcome analyses. Randomized participants completed postal questionnaires at 6 months and then annually for at least 10 years (ongoing in 2016). Randomized non-responders received reminder letters and questionnaires, followed by telephone calls from trial nurses and then a shortened version of the questionnaire with fewer measures. Some men were not given the International Consultation on Incontinence Questionnaire—Urinary Incontinence (ICIQ-UI; \( n = 135 \)) or EPIC (\( n = 794 \)) measures as they were introduced in the trial during 2001 and 2005, respectively. Data are presented for the return of the PROMs questionnaire booklet and the numbers which could be analysed for each PROM.

The PROMs were analysed as specified by the developers with no additional imputation for missing data. To aid interpretation of the results, the minimal clinically important difference (MID) was calculated as half a standard deviation of baseline values (a commonly used distribution-based method), but unspecified by the PROMs developers because MIDs are a more recently utilized concept (Table 1). Categorical symptom variables were dichotomized to ‘never or low/rarely’ vs ‘ever’ and International Continence Society short-form male survey (ICSmaleSF) items were dichotomized as previously in the ProtecT trial pilot phase [20]. The number of men without incontinence was defined as those with an ICIQ-UI score of 0. Symptom-related QoL items were categorized as none, small and a moderate/severe problem attributable to expected low prevalence of symptoms before treatment. Age was categorized into five-year groups (49–54 years to 65–69 years, with one 49-year-old who was eligible at recruitment). Self-reported ethnicity was categorized using UK Office of National Statistics census groupings (http://webarchive.nationalarchives.gov.uk/20160105160709/http://www.ons.gov.uk/ons/guide-method/measuring-equality/equality/ethnic-nat-identity-religion/ethnic-group/index.html). Socio-economic position was based on men’s current or last paid occupation at recruitment, and was grouped into professional or managerial (e.g. doctor), intermediate (e.g. secretary) and routine or semi-routine
(e.g. postman) using the UK Office of National Statistics categories [21].

Numbers and percentages are presented for categorical data, means and SD values for continuous data, and medians and interquartile ranges for skewed data. Those PROMs with a skewed distribution of results (EPIC, ICIQ-UI) as a result of few participants reporting symptoms are presented as means to aid comparison with other studies. Comparisons across age and occupational social class used Cuzick’s test for trend whilst Wilcoxon’s matched-pairs signed-rank tests (ordinal/continuous data) and McNemar’s test (nominal data) were used for comparisons of recruitment and biopsy data. Variation in selected PROMs completed at first biopsy (EPIC, Hospital Anxiety and Depression Scale [HADS] and a 12-item short-form health survey [SF-12]) was explored according to men’s age at recruitment and occupational social class and whether any differences exceeded the MID. PROMs data collected at PSA testing and biopsy time points were compared, and a sensitivity analysis subsequently assessed the impact of adding PSA testing data if the data from the biopsy time point were missing. All analyses were performed in STATA (version 13).

## Results

### Study and Participant Characteristics and PROMs Completion

The median age of randomized participants was 61 years, the majority were married or co-habiting (84%, 1375) and of white ethnicity (99%, 1606; full baseline demographic and clinical details published elsewhere) [11].

Approximately two fifths of participants reported a professional or managerial occupation (42%, 684/1643), 16% (259) an intermediate occupation and 42% (678) a routine or semi-routine occupation. Most randomized participants had low-risk disease features at diagnosis, as the median PSA at PSA testing was 4.6 ng/mL and around three quarters had a Gleason score of 6 (77%, 1266) and clinical stage of T1c (76%, 1249).

The majority of participants returned a questionnaire booklet at PSA testing (recruitment, 95%) and first biopsy clinics (88%, Fig. 1). Data that could be analysed for measures completed at first biopsy ranged from 77% for the SF-12 to 88% for EPIC bowel and urinary function (Table 1). As EPIC was introduced during the trial, 52% of randomized participants were asked to complete all PROMs at baseline (849/1643). Response rates for all PROMs were similar by randomized treatment allocation (Tables 2–4).

### Table 2 Urinary function by randomized allocation in ProtecT trial participants

<table>
<thead>
<tr>
<th>PROMs</th>
<th>Summary scores and specific symptoms or QoL Impact</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Active monitoring N = 545</td>
</tr>
<tr>
<td></td>
<td>Radiotherapy N = 545</td>
</tr>
<tr>
<td></td>
<td>Surgery N = 845</td>
</tr>
<tr>
<td>Total N = 1643</td>
<td></td>
</tr>
<tr>
<td>ICQ-UI minimum analysed/asked, n/N (%)</td>
<td>422/500 (84)</td>
</tr>
<tr>
<td>Incontinence mean score (SD)</td>
<td>1.3 (2.5)</td>
</tr>
<tr>
<td>No incontinence, n/N (%)</td>
<td>305 (72)</td>
</tr>
<tr>
<td>Large QoL impact, n/N (%)</td>
<td>3/428 (1)</td>
</tr>
<tr>
<td>EPIC minimum analysed/asked, n/N (%)</td>
<td>244/280 (87)</td>
</tr>
<tr>
<td>Urinary summary (SD)</td>
<td>93.0 (9.6)</td>
</tr>
<tr>
<td>Function score (SD)</td>
<td>95.6 (8.0)</td>
</tr>
<tr>
<td>Bother score (SD)</td>
<td>91.2 (12.3)</td>
</tr>
<tr>
<td>Incontinence score (SD)</td>
<td>93.5 (11.3)</td>
</tr>
<tr>
<td>Irritative/obstructive score (SD)</td>
<td>93.2 (9.5)</td>
</tr>
<tr>
<td>Pad use</td>
<td>1/250 (&lt;1%)</td>
</tr>
<tr>
<td>ICSmaleSF minimum analysed, n/N (%)</td>
<td>471/545 (86)</td>
</tr>
<tr>
<td>Incontinence score (SD)</td>
<td>1.9 (2.1)</td>
</tr>
<tr>
<td>Voiding score (SD)</td>
<td>3.4 (2.9)</td>
</tr>
<tr>
<td>Daytime frequency, n/N (%)</td>
<td>147/470 (31)</td>
</tr>
<tr>
<td>Nocturia, n/N (%)</td>
<td>111/475 (23)</td>
</tr>
<tr>
<td>Little urinary QoL impact, n/N (%)</td>
<td>109/475 (23)</td>
</tr>
<tr>
<td>Somewhat/a lot QoL impact, n/N (%)</td>
<td>20/475 (4)</td>
</tr>
</tbody>
</table>

Urinary, Bowel and Sexual Dysfunction and their QoL Impact

Levels of incontinence were low, indicating few problems, and at baseline <1% of men reported incontinence as a large problem (4/1259 [Table 2]) at diagnosis. Fewer than 1% of men reported using at least one pad per day (5/754). Seventy percent of the participants reported being incontinence-free (873/1244). Urinary function was good (EPIC score 95.1) and bother related to urinary symptoms was low (EPIC score 91.0), with few irritative or obstructive symptoms (EPIC score 93.0), as measured by EPIC. Nocturia was reported by around one fifth of men (312/1423; ICSmaleSF) and around one third also reported a regular daytime frequency (460/1410), although only 3% of men (44/1427) reported these LUTS as being a moderate or severe problem (ICSmaleSF).

Bowel symptom EPIC scores were good (i.e. few problems) and only 3% of men (20/751) reported a moderate or large problem attributable to bowel symptoms (Table 3). The frequency of faecal incontinence or bloody stools was also low, although 16% of participants reported having loose stools at least half of the time (118/754).

Around one third of men (241/735) reported erectile dysfunction and for 16% of participants this was a moderate or large problem (118/731) with the EPIC measure. Sexual function and bother scores were much lower than for urinary and bowel symptoms as assessed by EPIC (Tables 2 and 3).

Generic Health Status, Anxiety and Depression

Generic physical and mental health sub-scores were similar with UK normative data (SF-12 sub-scores of 50 [Table 4]) [18]. The EuroQoL health utility scale, the EQ-5D-3L, also indicated good overall health (score 0.89). Mean anxiety and depression scores were low, although one fifth of men (278/
had possible clinical levels of anxiety and ~6% of men had depression (80/1399 [Table 4]).

**Symptoms and Quality of Life by Age and Occupational Social Class**

Urinary EPIC summary and bother scores were generally worse in older men than younger men, unlike urinary function, and all bowel scores (where there was no difference with age), but differences in urinary scores did not exceed the MID (Table 5 and Fig. 2). The EPIC sexual summary, function and bother scores were better in the youngest (49–54 years) men compared with the oldest age group (65–69 years), and these differences in summary and functional scores exceeded the MID (Table 5). Anxiety was less frequent in older men, unlike depression which was similar across all age groups. In older men, physical health was worse than in younger men, whilst mental health was better (Table 5) although none of these differences exceeded the MID for HADS and SF-12. There were no strong associations of socio-economic status with quality of life, although there was some evidence of higher levels of depression and reduced bowel function with lower status, but neither exceeded the MID (Table S1). There was no evidence that physical health differed by socio-economic status.

**Generic Physical and Mental Health at PSA Testing and biopsy**

There were no differences in overall mental health (SF-12), and minor differences which did not exceed the MID in physical health (SF-12 < 2 scale points), nor in anxiety (<0.2 points) and depression (0.5 points) at PSA testing and first biopsy clinics (data not shown). The addition of PSA testing PROM data if biopsy PROM data were missing (e.g. an extra 188 participants for the anxiety score) did not alter the biopsy results (data not shown), so this approach was used in the trial outcome analyses (R. Pickard, M. Fabricus & E. McColl, unpublished data).

**Discussion**

There was good completion of a comprehensive range of validated PROMs at diagnostic biopsy before randomization to treatments for localized prostate cancer in the ProtecT trial. Urinary and bowel symptoms were generally infrequent (apart from storage LUTS), whilst sexual dysfunction was reported by around one third of men. Urinary and sexual function were generally worse in older men (65–69 years) than younger men (49–54 years) with clinically important reductions (MID) in sexual function, unlike bowel function where there were no age-related effects. Anxiety and depression were reported in around one fifth and one twentieth of participants, respectively. Overall physical and mental health scores were similar to national normative data. There were no large differences in mental or physical health, anxiety or depression between PSA testing and biopsy assessments. These data, therefore, confirm that the men randomized in the ProtecT study were generally healthy and mostly asymptomatic.

Interpretation of the ProtecT trial clinical results will be enhanced by the use of validated PROMs with pretreatment measurements (recommended by the CONSORT-PRO guidelines) [6] and high response rates. The ProtecT trial results are based on men identified by community-based PSA testing and are hence analogous to screen-detected patients.

### Table 5

**Symptoms and quality of life by age group in men randomized in the ProtecT trial.**

<table>
<thead>
<tr>
<th>Age group</th>
<th>49-54 years</th>
<th>55-59 years</th>
<th>60-64 years</th>
<th>65-70 years</th>
<th>P*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Symptoms (MID)‡</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Minimum analysed/asked, n/N (%)</td>
<td>76/87 (87)</td>
<td>200/225 (89)</td>
<td>220/270 (81)</td>
<td>223/267 (84)</td>
<td></td>
</tr>
<tr>
<td>Urinary summary (4.6)</td>
<td>93.1 (9.9)</td>
<td>94.0 (7.2)</td>
<td>92.2 (9.5)</td>
<td>92.0 (9.9)</td>
<td>0.022</td>
</tr>
<tr>
<td>Urinary function (4.2)</td>
<td>92.9 (12.8)</td>
<td>95.7 (7.0)</td>
<td>94.9 (8.0)</td>
<td>95.3 (8.1)</td>
<td>0.627</td>
</tr>
<tr>
<td>Urinary bother (5.9)</td>
<td>93.3 (10.3)</td>
<td>92.7 (9.2)</td>
<td>90.2 (12.4)</td>
<td>89.6 (13.1)</td>
<td>~0.001</td>
</tr>
<tr>
<td>Bowel summary (4.2)</td>
<td>93.7 (10.3)</td>
<td>93.1 (8.6)</td>
<td>93.1 (8.2)</td>
<td>94.4 (7.6)</td>
<td>0.357</td>
</tr>
<tr>
<td>Bowel function (4.4)</td>
<td>91.7 (11.1)</td>
<td>91.3 (9.0)</td>
<td>91.6 (8.7)</td>
<td>93.0 (7.9)</td>
<td>0.170</td>
</tr>
<tr>
<td>Bowel bother (5.0)</td>
<td>95.4 (11.6)</td>
<td>94.9 (9.8)</td>
<td>94.7 (10.0)</td>
<td>95.7 (9.7)</td>
<td>0.755</td>
</tr>
<tr>
<td>Sexual summary score (11.6)</td>
<td>72.9 (20.6)</td>
<td>65.8 (21.0)</td>
<td>60.0 (23.2)</td>
<td>56.2 (23.8)</td>
<td>~0.001</td>
</tr>
<tr>
<td>Sexual function score (11.5)</td>
<td>66.1 (21.2)</td>
<td>58.8 (20.3)</td>
<td>53.0 (22.4)</td>
<td>48.3 (24.0)</td>
<td>~0.001</td>
</tr>
<tr>
<td>Sexual bother score (14.8)</td>
<td>87.9 (24.1)</td>
<td>81.1 (28.0)</td>
<td>75.9 (30.8)</td>
<td>74.0 (30.4)</td>
<td>~0.001</td>
</tr>
<tr>
<td>Mental/physical health†(MID), n/N (%)</td>
<td>142/189 (75)</td>
<td>346/418 (83)</td>
<td>403/532 (76)</td>
<td>369/504 (73)</td>
<td></td>
</tr>
<tr>
<td>Anxiety score (1.8)§</td>
<td>5.5 (3.5)</td>
<td>5.2 (3.5)</td>
<td>4.8 (3.6)</td>
<td>4.5 (3.3)</td>
<td>~0.001</td>
</tr>
<tr>
<td>Depression score (1.3)§</td>
<td>2.5 (2.8)</td>
<td>2.4 (2.7)</td>
<td>2.5 (2.5)</td>
<td>2.4 (2.3)</td>
<td>0.123</td>
</tr>
<tr>
<td>Mental health score (3.8)§</td>
<td>51.9 (9.2)</td>
<td>53.3 (7.3)</td>
<td>53.9 (7.7)</td>
<td>55.3 (6.5)</td>
<td>~0.001</td>
</tr>
<tr>
<td>Physical health score (4.0)§</td>
<td>52.9 (7.6)</td>
<td>51.6 (7.8)</td>
<td>50.8 (8.0)</td>
<td>50.3 (7.9)</td>
<td>~0.001</td>
</tr>
</tbody>
</table>

HADS, Hospital Anxiety and Depression Scale; MID, minimal clinically important difference; PROM, patient-reported outcome measure; SF-12, 12-item short-form health survey.

Data are EPIC summary scores (so), unless otherwise indicated. *Cuzick’s test; MID 0.5 so of baseline values with domains in bold where exceeded between youngest and oldest age groups. ‡Details of PROMs and their administration in Table 1 and Fig. 1. §HADS; SF-12; © 2016 The Authors
with largely low-risk disease, which reflects contemporary practice in many countries, unlike the earlier SPCG-4 trial [4]. The comprehensive range of PROMs encompasses the domains identified to assess treatments for prostate cancer by the National Cancer Institute prostate cancer working group [7], the International Consortium for Health Outcome Measurements for prostate cancer [8] and the ongoing core outcome set for trials of interventions for localized prostate cancer, unpublished data). The EPIC-26 PROM (a shorter version of EPIC) is now recommended for assessing prostate cancer treatments [8] and EPIC was added to the trial in 2005, so the majority of participants will have outcome assessments using this measure. Overall QoL and psychological PROMs also assessed the broader impacts of prostate cancer treatment, unlike in many previous studies [22–24].

There are some potential limitations to these baseline results in that the characteristics of non-responders to questionnaires were not compared with those of responders, although major differences are unlikely given the high response rates. Less severe psychological impacts may have been overlooked as the HADS was developed to detect clinical depression and anxiety [17]. Psychological distress was reported by around one fifth of ProtecT trial participants (33/171) completing the Impact of Events Scale at PSA testing and biopsies in a substudy [25], which was similar to the frequency of anxiety reported with HADS in the present study by randomized participants. Baseline PROMs assessments occurred during the diagnostic process, which may have influenced participants’ perceptions of symptoms but were needed before randomization and there were also only minor differences between PSA and biopsy assessments. Furthermore, we present MIDs to aid interpretation and comparison with other findings but these PROMs were developed before the MID concept was widespread and this had not been validated nor was it in the prespecified trial analysis plan.

The applicability of the present results to men of non-white ethnicity is unclear, as ProtecT trial participants were mostly of white ethnicity. The percentage of white participants in the trial was broadly similar to the older male population at seven of the cities in the ProtecT trial (UK census data), whilst the percentage of black participants was lower than in the population of one city and Asian men in two cities. Participants in screening trials are often healthier than the general population and, in the European Randomised Screening for Prostate Cancer (ERSPC) trial screened arm, cardiovascular mortality was lower than in the unscreened arm [26]. The ProtecT trial baseline overall physical and mental health and HADS scores in randomized participants were similar to UK population data [18,27], as was broadly the dietary energy intakes of ProtecT participants with prostate cancer [28]. The socio-economic distribution of randomized participants in the ProtecT trial is similar to national data (e.g. 39% of men aged 55–64 years with a professional occupation in census data compared with 42% in ProtecT trial participants) [21].

The ProtecT trial baseline PROM results could be compared either with men undergoing prostate cancer screening prior to prostate cancer treatment or men with benign urological conditions or community-based populations. ERSPC participants without prostate cancer reported good urinary and bowel function, low bother (EPIC score) and reduced sexual function, which worsened in older men (Dutch questionnaire) [29] in a similar manner to randomized participants in the ProtecT trial. There were few changes in quality of life and anxiety during ERSPC screening [30,31]
and their SF-36 scores were consistent with the ProtecT trial SF-12 scores at PSA testing and biopsy (scoring equivalency exists between questionnaires [32]). Men with high levels of anxiety during ERSPC screening were predicted to have high anxiety at recruitment [33], which was also observed in the wider ProtecT trial cohort [25]. Non-cancer controls in the American Prostate Lung and Colorectal Cancer screening trial showed similar patterns of bowel, sexual and urinary symptoms (EPIC score) to those of men in ProtecT trial at baseline [34].

None of the major localized prostate cancer trials [4,5,35] comparing treatments include baseline assessments made before treatment, unlike several prostate cancer cohorts. Rates of erectile function, pad use (1%) and problematic urinary symptoms (2%) in US patients before treatment, measured with EPIC-26 [36], were similar to baseline ProtecT trial findings. Dutch patients subsequently undergoing surgery or radiotherapy also had similar levels of pad use, a slightly lower frequency of loose stools (10%) and greater erectile dysfunction (~50%) [37] than results presented here.

American prostate cancer cohorts that recruited either in 1994 to 1995 (Prostate Cancer Outcomes Study [PCOS]) or 2011 to 2012 (Comparative Effectiveness Analysis of Surgery and Radiation [CEASAR]), with baseline assessments made by recalling symptoms before diagnosis with the University of California Los Angeles Prostate Cancer Index (UCLA-PCI, a precursor to EPIC) and EPIC-26, respectively, were compared for patterns over time [38]. Incontinence was relatively frequent but was higher in the newer cohort (CEASAR: 27%, PCOS 19%), whilst pad use was higher in both cohorts than reported here (CEASAR: 7%, PCOS 3%). Erectile dysfunction was very prevalent in the recent CESAR cohort (45%, PCOS: 24%) and the ProtecT trial baseline results at around one third of men was broadly similar to both cohorts. Bowel function was good in the CESAR cohort [39] but was not compared with PCOS because the UCLA-PCI did not include all the bowel questions included in the EPIC score.

An Australian cohort of men without prostate cancer (matched to men with the disease) [40] showed similar levels of incontinence (1% pad use), a slightly higher percentage of major bowel problems (6%) and similar erectile dysfunction (22%), assessed using the UCLA-PCI, to those seen in the participants of the ProtecT trial men at baseline. The prevalence of LUTS increased with age and was up to 30% in men aged ≥80 years, based on UK general practice data [41]. Some symptom PROMs used in the ProtecT trial were validated with clinical or community populations, e.g. ICIQ-UI where the community sample incontinence score (1.6) was similar to the ProtecT trial results in the present study, and was lower than for urology clinic attendees (2.4) [15]. Men listed for surgery for LUTS had a mean incontinence score of 4.0 and a voiding score of 8.9 (ICSmaleSF) [16], which indicated worse symptoms than for participants in the ProtecT trial before treatment. The UCLA-PCI results for US men without cancer were very similar to ProtecT trial baseline results [42], except that bowel function and bother were worse in older men, possibly because the mean age of older men completing UCLA-PCI was 79 years, which exceeded the upper age limit for ProtecT trial men (69 years). The EPIC sexual function scores of older men in the ProtecT trial were below a clinical threshold for potency (EPIC sexual domain score of <60) [43].

In conclusion, there were generally low levels of urological and bowel symptoms but more erectile dysfunction in participants before randomization and treatment for localized prostate cancer in the ProtecT trial. Symptom frequencies and trends with age were similar to those found in other cohorts. These results help characterize the trial population and will facilitate the robust evaluation of treatment impacts when the ProtecT trial outcomes are published in 2016.

Acknowledgements

The ProtecT trial is funded by the UK National Institute for Health Research Health Technology Assessment Programme (projects 96/20/06, 96/20/99, http://www.nets.nihr.ac.uk/projects/hta/962099) with the University of Oxford as sponsor. Department of Health disclaimer: The views and opinions expressed therein are those of the authors and do not necessarily reflect those of the Department of Health.

Conflict of Interest

Chris Metcalfe reports grants from UK NIHR Health Technology Assessment and Health Services Research Programme and Clinical Trials Unit Support Funding, outside the submitted work. Jane Blazeby, Jenny L. Donovan, Tim J. Peters and Athene Lane reports grants from National Institute for Health Research and Clinical Trials Unit Support Funding during the conduct of the study. Jenny L. Donovan, Freddie C. Hamdy, David E. Neal and Tim J. Peters are NIHR senior investigators. Some of Jenny L. Donovan’s time is supported by the Collaboration for Leadership in Applied Health Research and Care West at University Hospitals Bristol NHS Foundation Trust. The remaining authors have no conflict of interest to declare.

References

Lane et al.


11 Lane JA, Donovan JL, Davis M et al. Active monitoring, radical prostatectomy, or radiotherapy for localised prostate cancer: study design and diagnostic and baseline results of the ProteCt randomised phase 3 trial. *Lancet Oncol* 2014; 15: 1109–18


20 Brindle LA, Oliver SE, Dedman D et al. Measuring the psychosocial impact of population-based prostate-specific antigen testing for prostate cancer in the UK. *BJU Int* 2006; 98: 777–82


42 Litwin MS. Health related quality of life in older men without prostate cancer. *J Urol* 1999; 161: 1180–4


© 2016 The Authors

BJU International published by John Wiley & Sons Ltd on behalf of BJU International
Correspondence: Athene Lane, School of Social and Community Medicine, Canynge Hall, University of Bristol, 39 Whatley Road, Bristol BS8 2PS, UK.
e-mail: epjaxl@bristol.ac.uk

Abbreviations: PROM, patient-reported outcome measure; ProtecT trial, Prostate Testing for Cancer and Treatment trial; NIHR, National Institute for Health Research; QoL, quality of life; EPIC, Expanded Prostate Index Composite; ICIQ-UI, International Consultation on Incontinence Questionnaire-Urinary Incontinence; MID, minimal clinically important difference; HADS, Hospital Anxiety and Depression Scale; ERSPC, European Randomised Screening for Prostate Cancer; PCOS, Prostate Cancer Outcomes Study; CEASAR, Comparative Effectiveness Analysis of Surgery and Radiation; UCLA-PCI, University of California Los Angeles Prostate Cancer Index.

Appendix 1


Radiotherapy: Helen Appleby, Dominic Ash, Dean Aston, Steven Bolton, Graham Chalmers, John Conway, Nick Early, Tony Geater, Lynda Goddall, Claire Heymann, Deborah Hicks, Liza Jones, Susan Lamb, Geoff Lambert, Gill Lawrence, Geraint Lewis, John Lilley, Aileen MacLeod, Pauline Massey, Alison McQueen, Rollo Moore, Lynda Penketh, Janet Poterton, Neil Roberts, Helen Showler, Stephen Slade, Alasdair Steele, James Swinsoe, Marie Tiffany, John Townley, Jo Treeby, Joyce Wilkinson, Lorraine Williams, Lucy Wills, Owain Woodley, Sue Yarrow Histopathologists: Selina Bhattacharai, Neeta Deshmukh, John Dormer, Malee Fernando, John Goepel, David Griffiths, Ken Grigor, Nick Mayer, Jon Oxley, Mary Robinson, Murali Varma, Anne Warren. Data Management, Research and Statistics: Lucy Brindle, Michael Davis, Dan Dedman, Liz Down, Hanan Khazragui, Chris Metcalfe, Sian Noble, Tim Peters, Hilary Taylor, Marta Tazewell, Emma Turner, Julia Wade, Eleanor Walsh, Grace Young; Administration: Susan Baker, Elizabeth Bellis-Sheldon, Chantal Bougard, Joanne Bowtell, Catherine Brewer, Chris Burton, Jennie Carlton, Nicholas Christoforou, Rebecca Clark, Susan Coull, Christine Croker, Rosemary Currer, Claire Daisey, Gill Delaney, Rose Donohue, Jane Drew, Rebecca Farmer, Susan Fry, Jean Haddow, Alex Hale, Susan Halpin, Belle Harris, Barbara Hattrick, Sharon Holmes, Helen Hunt, Vicky Jackson, Donna Johnson, Mandy Le Butt, Jo Leworthy, Tanya Liddiatt, Alex Martin, Janee Mauree, Susan Moore, Gill Moulam, Jackie Mitch, Alena Nash, Kathleen Parker, Christopher Pawsey, Michelle Purdie, Teresa Robson, Lynne Smith, Jo Snoeck, Carole Stenton, Tom Steuart-Feilding, Chris Sully, Caroline Sutton, Carol Torrington, Zoe Wilkins, Sharon Williams, Andrea Wilson and others. Data Monitoring Committee: Chairs: Adrian Grant and Ian Roberts; Deborah Asby, Richard Cowan, Peter Fayers, Killian Mellon, James N’Dow, Tim O’Brien, Michael Sokhal. Trial Steering Committee: Chair: Michael Baumb; Jan Adolphson, Peter Albertsen, David Dearaley, Fritz Schroeder, Tracy Roberts, Anthony Zietman.

Supporting Information

Additional Supporting Information may be found in the online version of this article:
Table S1 Symptoms and quality of life by occupational social class in men randomised in the ProtecT trial.