Importance
Abiraterone acetate is combined with prednisone, 5 mg, twice daily for metastatic castration-resistant prostate cancer (mCRPC) and with prednisone, 5 mg, once daily for newly diagnosed, high-risk, metastatic castration-sensitive prostate cancer. Understanding the physiological effects of these and other regimens is important.

Objective
To evaluate the safety of abiraterone acetate with 4 glucocorticoid regimens.

Design, Setting, and Participants
Open-label, randomized clinical trial (1:1:1:1) of 164 men with mCRPC from 22 hospitals in 5 countries who were randomly assigned to 1 of 4 intervention groups between June 2013 and October 2014. Analyses were conducted from August 2017 to June 2018.

Interventions
Abiraterone acetate, 1000 mg, once daily with prednisone, 5 mg, twice daily (n = 41), 5 mg once daily (n = 41), 2.5 mg twice daily (n = 40), or dexamethasone, 0.5 mg, once daily (n = 42).

Main Outcomes and Measures
Primary end point was no mineralocorticoid excess (grade ≥1 hypokalemia or grade ≥2 hypertension) through 24 weeks (6 cycles) from treatment.

Results
Of 164 men (median [range] age, 70 [50-90] years) randomized to receive abiraterone acetate, 1000 mg, daily with prednisone, 5 mg, twice daily, once daily, or 2.5 mg twice daily, or dexamethasone, 0.5 mg, once daily, 24 (70.6%) of 34 patients (95% CI, 53.8%-83.2%), 14 (36.8%) of 38 patients (95% CI, 23.4%-52.7%), 21 (60.0%) of 35 patients (95% CI, 43.6%-74.4%), and 26 (70.3%) of 37 patients (95% CI, 54.2%-82.5%), respectively, had no mineralocorticoid excess. Plasma adrenocorticotrophic hormone and urinary mineralocorticoid metabolites after 8 weeks were higher with prednisone, 2.5 mg, twice daily and 5 mg once daily than with 5 mg twice daily or dexamethasone, 0.5 mg, once daily. The level of urinary glucocorticoid metabolites appeared higher in patients who did not meet the primary end point, regardless of glucocorticoid regimen. Total lean body mass decreased in the prednisone groups and total body fat increased in the prednisone, 5 mg, twice daily and dexamethasone groups. In the dexamethasone group, there was an increase in serum insulin and homeostatic model assessment of insulin resistance, while total bone mineral density decreased. In the prednisone, 5 mg, twice daily, 5 mg once daily, 2.5 mg twice daily, and dexamethasone groups, median radiographic progression-free survival was 18.5, 15.3, 12.8, and 26.6 months, respectively.

Conclusions and Relevance
Abiraterone acetate with prednisone, 5 mg, twice daily or dexamethasone, 0.5 mg, once daily met the prespecified threshold for the primary end point (95% CI excluded 50% mineralocorticoid excess); abiraterone acetate with prednisone, 5 mg, once daily or 2.5 mg twice daily did not meet the threshold. Abiraterone acetate in combination with dexamethasone appeared to be particularly active but may be associated with adverse metabolic consequences.

Trial Registration
ClinicalTrials.gov identifier: NCT01867710

JAMA Oncol. doi:10.1001/jamaoncol.2019.1011
Published online June 27, 2019. Corrected on July 11, 2019.
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biraterone acetate, 1000 mg, once daily is approved for use in combination with either prednisone, 5 mg, twice daily or prednisone, 5 mg, once daily based on a phase 3 trials reporting improvements in overall survival for the former in patients with metastatic castration-resistant prostate cancer (mCRPC)\(^1\,^2\) and for the latter in newly-diagnosed, high-risk patients with metastatic castration-sensitive prostate cancer. \(^3\) Prednisone, 5 mg, once daily was also used in phase 3 trials investigating the benefit of abiraterone acetate in hormone-naïve advanced prostate cancer (STAMPEDE [NCT00268476\(^4\) and PEACE-1 [NCT01957436\(^6\)]. When administered without glucocorticoids, abiraterone acetate induces a syndrome of secondary mineralocorticoid excess due to CYP17A1 \(^5\,^6\). In addition to maximizing efficacy for treating prostate cancer, glucocorticoids are combined with abiraterone acetate to prevent this syndrome; however, glucocorticoid exposure may exceed physiological requirements. Different glucocorticoid combination regimens have not been formally tested with abiraterone acetate. Because prolonged exposure to abiraterone acetate plus glucocorticoid is expected from earlier use, there is an urgent need to improve understanding of the physiological effects of these treatments to improve individualized treatment.

We aimed to evaluate administration of abiraterone acetate with prednisone, 5 mg, twice daily, the lower dose of 5 mg once daily used in metastatic castration-sensitive prostate cancer trials, \(^3\,^4\,^7\) and prednisone, 2.5 mg, twice daily. In men with progressive mCRPC, early studies of abiraterone acetate showed tumor response when combined with dexamethasone, 0.5 mg, once daily. \(^5\,^8\) and subsequent reports described tumor responses to abiraterone acetate after a switch from prednisone to dexamethasone. \(^9\) Therefore, our fourth treatment group received dexamethasone, 0.5 mg, once daily. The primary end point was the absence of mineralocorticoid excess in the first 24 weeks of treatment. Additionally, we assessed toxic effects related to glucocorticoid excess or androgen suppression, effects on steroid biosynthesis, and antitumor activity.

Methods

Patients and Study Design

In this open-label, parallel-arm, multicenter, phase 2 study, 164 asymptomatic or minimally symptomatic men with mCRPC received abiraterone acetate 1000 mg once daily and were randomly assigned in a 1:1:1:1 ratio, using an interactive web response system and a computer-generated randomization schedule, to 1 of 4 glucocorticoid regimens: prednisone, 5 mg, twice daily; prednisone, 5 mg, once daily; dexamethasone, 2.5 mg, twice daily; or dexamethasone, 0.5 mg, once daily. The trial protocol is available in Supplement 1. Study treatment was administered in 28-day cycles without interruption. Participants were recruited at 22 centers in 5 countries (eTable 1 in Supplement 2). They were required to have histologically or cytologically confirmed prostate adenocarcinoma with metastatic disease documented by positive bone scan, computed tomography scan, or magnetic resonance imaging and prostate cancer progression documented by prostate-specific antigen (PSA) according to the Prostate Cancer Working Group 2\(^10\) or radiographically according to the modified Response Evaluation Criteria in Solid Tumors (RECIST) version 1.1. \(^11\) Men were surgically or medically castrated, with testosterone levels of less than 50 ng/dL. Men with visceral disease were initially excluded until a protocol amendment in November 2013 (after 61 patients had been accrued), when they were permitted to enroll. Patients with prior cytotoxic chemotherapy, biologic therapy, or inhibition of the androgen receptor with abiraterone acetate plus prednisone for the treatment of mCRPC were excluded.

The planned maximum duration of the main study was 39 cycles (156 weeks) after the first participating patient started study treatment. Patients continued the main study treatment until disease progression, initiation of another prostate cancer treatment, or discontinuation of study treatment for either safety reasons (eg, sustained toxic effects that did not return to grade \(\leq 1\) with appropriate medical management) or patient request. At the end of the main study, men could continue treatment in a separate extension protocol. Patients with grade 3 or 4 hypokalemia and those who required diuretic treatment, a change in glucocorticoid dose, or palliative radiotherapy discontinued the main study treatment but could enroll in the extension protocol and restart the main study treatment after improvement to grade \(\leq 1\) without disease progression. The main study was completed in August 2016, and analyses were conducted from August 2017 to June 2018.

Study visits occurred on days 1 and 15 of cycle 1, day 1 of cycles 2 to 6, day 1 of every 3 cycles thereafter, and 4 weeks after completing the main study. Patients reported any adverse events since the prior visit, and the investigator graded each event using the National Cancer Institute Common Terminology Criteria for Adverse Events (NCI CTCAE) version 4.0. Patients with low potassium or prior history of hypokalemia...
Safety of Glucocorticoid With Abiraterone Acetate for Metastatic Castration-Resistant Prostate Cancer

Original Investigation Research

JAMA Oncology Published online June 27, 2019 E3

could undergo weekly or more frequent laboratory electrolyte evaluation. Additional procedures are described in the eMethods in Supplement 2.

The study protocol and amendments were approved by the institutional review board at each participating site, and the study was conducted according to the provisions of the Declaration of Helsinki and the International Conference on Harmonization of Good Clinical Practice Guidelines. All patients provided written informed consent before participating.

Outcomes

The primary end point of no mineralocorticoid excess was defined as experiencing neither of 2 mineralocorticoid excess toxic effects, grade ≥1 hypokalemia or grade ≥2 hypertension, during the first 24 weeks of treatment (ie, 6 cycles). This end point was derived from treatment emergent adverse event data, defined using the Medical Dictionary for Regulatory Activities (MedDRA)3 and graded according to NCI CTCAE v4.0. Secondary end points were global safety profile by NCI CTCAE v4.0; changes in plasma ACTH and urinary metabolites (eMethods and eTable 2 in Supplement 2); incidence of exogenous glucocorticoid adverse effects, namely changes in fasting serum insulin, homeostatic model assessment of insulin resistance (HOMAIR; calculated as insulin × glucose ÷ 22.5), total lean body mass, total body fat, and bone mineral density; and clinical benefit, including antitumor activity from serum PSA values, radiographic progression-free survival, and influence on quality of life (based on the EuroQol 5-dimensional questionnaire and Functional Assessment of Cancer Therapy-Prostate Cancer questionnaire). Total lean body mass and total body fat were assessed using bone density scan data.13 To avoid artifacts in interpretation due to bone metastases, we only report body fat changes involving the arms and total body. An exploratory secondary objective to evaluate associations with outcome and plasma androgen receptor aberrations14 will be reported separately.

Trial Size Calculation and Statistical Analyses

The assumption was that for a glucocorticoid regimen to be considered of interest, 75% of patients in a treatment group would need to meet the primary end point, based on previously reported rates in a regulatory phase 3 trial.2 In each treatment group, an exact binomial test with a 5% 1-sided significance level had 89% power to detect the difference between 50% and 75% with a sample size of at least 30 patients; a treatment group would be rejected if 50% of patients or fewer met the primary end point. Analyses were conducted separately within each group, and the study was neither designed nor powered to compare results across groups.

Changes in steroid metabolites (after 2 cycles) and exogenous glucocorticoid adverse effects (at cycle 12 and at the end of the main study) were examined with the Wilcoxon signed rank test. Values for urinary metabolites after 2 cycles were compared among patients who met the primary end point and those who did not at 24 weeks using the Wilcoxon 2-sample test. Statistical significance was defined as \( P < .05 \), with no correction for multiple testing. Missing values for the end-of-main-study visit were imputed using the last-observation-carried-forward method; for other time points, only patients with data were included. Patient-reported quality of life was summarized descriptively at baseline and at cycles 6 and 18. Radiographic progression-free survival was analyzed with the Kaplan-Meier product-limit method. The trial was not designed to compare antitumor activity between the 4 treatment groups.

Results

Patient Characteristics

Of 204 men assessed for study eligibility, 164 (median [range] age, 70 [50-90] years) were randomly assigned between June 2013 and October 2014 to 1 of 4 glucocorticoid regimens (intention-to-treat population) and 163 received at least 1 dose (safety population) (Figure 1). Patient disposition details are provided in eFigure 1 in Supplement 2. Most baseline characteristics were balanced across treatment groups, although incidences of elevated blood pressure at baseline were 46.3% for prednisone, 5 mg, once daily and ranged from 26.8% to 33.4% in the other groups. The mean time from diagnosis to randomization was 101.4 months in the dexamethasone, 0.5 mg, once daily group and ranged from 59.6 to 69.2 months in the other groups (eTable 3 in Supplement 2). Between 39.0% and 56.1% of patients (16 to 23 patients) in each treatment group were taking any antihypertensive medication at baseline. Median treatment exposure was 12.9 months overall (12.9, 10.6, 8.4, and 18.4 months in the prednisone, 5 mg, twice daily, once daily, and 2.5 mg twice daily, and dexamethasone groups, respectively), with a maximum follow-up time on treatment of 35.6 months.

Primary End Point

Of 163 patients who received any treatment, 19 discontinued before week 24 without an adverse event of either grade 2 or higher hypertension or grade 1 or higher hypokalemia; therefore, 144 patients were evaluable for the primary end point (Figure 1). Of these patients, 85 (59.0%) experienced neither grade 2 or higher hypertension nor grade 1 or higher hypokalemia during the first 24 weeks of treatment and met the primary end point. The breakdown by treatment group was 24 of 34 patients (71%; 95% CI, 54%-83%) treated with prednisone, 5 mg, twice daily, 14 of 38 patients (37%; 95% CI, 23%-53%) treated with prednisone, 5 mg, once daily, 21 of 35 patients (60%; 95% CI, 44%-74%) treated with prednisone, 2.5 mg, twice daily, and 26 of 37 patients (70%; 95% CI, 54%-83%) treated with dexamethasone (Table). The lower boundary of the 95% CI was higher than 50% (the protocol-defined criterion) with prednisone, 5 mg, twice daily and dexamethasone but less than 50% with prednisone, 5 mg, once daily and 2.5 mg twice daily. Overall, grade 2 or higher hypertension occurred in the first 24 weeks in 49 (34.0%) of 144 evaluable patients, and grade 1 or higher hypokalemia occurred in 18 (12.5%) of 144 patients (Table), including 2 patients with grade 3 hypokalemia in the prednisone, 5 mg, once daily group.

Treatment Effect on Steroid Biosynthesis

In keeping with the prednisone, 2.5 mg, twice daily and 5 mg once daily groups not meeting the primary end point for ab-
sence of mineralocorticoid excess toxic effects, plasma ACTH increased significantly during treatment from baseline to cycle 3 with abiraterone plus prednisone, 5 mg, once daily (median, 8.95 pmol/L; interquartile range [IQR], 4.39-12.63) (P < .001) and abiraterone plus prednisone, 2.5 mg, twice daily (median, 3.97 pmol/L; IQR, 0.42-9.24) (P < .001). Plasma ACTH did not change significantly with prednisone, 5 mg, twice daily (median, −1.07 pmol/L; IQR, −3.08 to 0.30) (P = .16) and decreased significantly with dexamethasone (median, −1.82 pmol/L; IQR, −3.39 to −0.10) (P = .02) (Figure 2A and eTable 4 in Supplement 2).

To further evaluate upregulation of steroids upstream of CYP17A1 inhibition, urinary steroid metabolites were evaluated. From baseline to cycle 3, deoxycorticosterone metabolites significantly increased (all P < .001) with prednisone, 5 mg, twice daily (absolute median, 11.4 μg/24 h; IQR, −1.9 to 60.9 and median percentage change, 149%; IQR, −15% to 324%); 5 mg once daily (absolute median, 75.9 μg/24 h; IQR, 46.7 to 176.2 and median percentage change, 862%; IQR, 351%-1491%); and 2.5 mg twice daily (absolute median, 100.5 μg/24 h; IQR, 36.9 to 158.6 and median percentage change, 716%; IQR, 402%-1391%) but not with dexamethasone (Figure 2B and eTable 4 in Supplement 2). Corticosterone metabolites significantly increased (each P < .001) with prednisone, 5 mg, once daily (absolute median, 4452.6 μg/24 h; IQR, 2610.2-6573.0 and median percentage change, 984% IQR, 622%-1851%) and prednisone, 2.5 mg, twice daily (absolute median, 4913.5 μg/24 h; IQR, 1650.6 to 7787.9 and median percentage change, 1016%; IQR, 471%-1846%) (eFigure 2A and eTable 4 in Supplement 2). At first assessment, at cycle 3, median suppression

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### Table. Primary End Point: Mineralocorticoid Excess Adverse Events During the First 24 Weeks

<table>
<thead>
<tr>
<th>Variable</th>
<th>Treatment Group</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Prednisone, 5 mg, Twice Daily (n = 34)</td>
</tr>
<tr>
<td>Met the primary end point²</td>
<td>No. (%)</td>
</tr>
<tr>
<td>95% CI ¹</td>
<td>(54-83)</td>
</tr>
<tr>
<td>Investigator reported an adverse event</td>
<td></td>
</tr>
<tr>
<td>Grade ≥ 2 hypertension and grade ≥ 1 hypokalemia, No. (%)</td>
<td>2 (6)</td>
</tr>
<tr>
<td>Grade ≥ 2 hypertension alone, No. (%)</td>
<td>7 (21)</td>
</tr>
<tr>
<td>Grade ≥ 1 hypokalemia alone, No. (%)</td>
<td>1 (3)</td>
</tr>
</tbody>
</table>

² Patients in all treatment groups received abiraterone acetate.
³ Patients completed the 24 weeks of treatment (6 cycles) or discontinued treatment early and experienced either hypertension (National Cancer Institute Common Terminology Criteria for Adverse Events grade ≥ 2) or hypokalemia.
⁴ Investigator reported neither hypertension (National Cancer Institute Common Terminology Criteria for Adverse Events grade ≥ 2) nor hypokalemia during the first 24 weeks (6 cycles).
of urinary excretion of major androgen metabolites androstosterone (eFigure 2B in Supplement 2) and dehydroepiandrosterone (eFigure 2C in Supplement 2) was greater than 90% (eTable 4 in Supplement 2). Androgen precursor metabolites were also significantly suppressed (both P < .001) with prednisone, 5 mg, twice daily (absolute median, −330.8 μg/24 h; IQR, −555.5 to −165.8; median percentage change, −81%; IQR, −89% to −63%) and dexamethasone (absolute median, −262.1 μg/24 h; IQR, −369.0 to −89.5; median percentage change, −88%; IQR, −93% to −56%). However, androgen precursor metabolites were not significantly suppressed with prednisone, 5 mg, once daily or prednisone, 2.5 mg, twice daily (Figure 2C and eTable 4 in Supplement 2).

In exploratory analyses, 24-hour urinary excretion of cortisol (Figure 2D) and cortisone (eFigure 3B in Supplement 2) were significantly higher in patients in any treatment group who experienced grade 2 or higher hypertension or grade 1 or higher hypokalemia in the first 24 weeks of treatment, as determined by the Wilcoxon 2-sample test.

**Adverse Events**

Over the main study, grade 3 adverse events of hypertension were reported in 7% (n = 3), 22% (n = 9), 13% (n = 5), and 7% (n = 3) of patients in the prednisone, 5 mg, twice daily, prednisone, 5 mg, once daily, prednisone, 2.5 mg, twice daily, and dexamethasone groups, respectively. Grade 3 hypokalemia was reported in 7.3% (n = 3) of patients in the prednisone, 5 mg, once daily group and in no other patients; all other events of hypokalemia were grade 1 or grade 2 (eTable 6 in Supplement 2). Neither grade 4 hypokalemia nor grade 4 hypertension was reported in any treatment group.
sion were reported, and no patient discontinued study treat-
ment for persistent hypertension. One patient in the
prednisone, 5 mg, once daily group with an unrelated gastro-
intestinal illness had a prolonged hospitalization for a serious
adverse event of hypokalemia. No patient had a serious ad-
verse event of hypertension. Cumulative incidences of hyper-
tension and hypokalemia over time are shown in Figure 4 in
Supplement 2. Other adverse events of interest such as pe-
ripheral edema, cardiac effects, and liver function are sum-
marized in Table 7 in Supplement 2.

Long-Term Metabolic and Musculoskeletal Effects

Insulin resistance was evaluated by recording changes in fast-
ing serum insulin or HOMA-IR. The mean change from base-
line for fasting serum insulin in all patients was 9.2 pmol/L (95% CI, −14.5 to 33.0) at cycle 12 and 17.7 pmol/L (95% CI, 1.0-34.3) at the end of the main study. Significant increases in fasting serum insulin and HOMA-IR were observed in the dexameth-
asone group only, both at cycle 12 (fasting serum insulin, 53.7 pmol/L; 95% CI, 16.6-90.8; \( P < .001 \) and HOMA-IR, 2.15; 95% CI, 0.24-4.07; \( P = .004 \) (Figure 3A, B)) and at the end of the main study (fasting serum insulin, 47.5 pmol/L; 95% CI, 18.8-76.2; \( P = .002 \) and HOMA-IR, 1.99; 95% CI, 0.59-3.38; \( P = .01 \)) (eFigure 5 in Supplement 2). No patient required insulin. Two pa-
tients started medications for non–insulin-dependent diabe-
etes (both were in the dexamethasone group).

Body composition showed a significant decrease in total
lean body mass from baseline to cycle 12 in the prednisone, 5
mg, twice daily (−5.8%; 95% CI, −8.6% to −3.0%) (\( P < .001 \)),
prednisone, 5 mg, once daily (−2.5%; 95% CI, −4.9% to −0.0%) (\( P = .02 \)), and prednisone, 2.5 mg, twice daily groups (−5.6%; 95% CI, −8.5% to −2.6%; \( P < .001 \)) (Figure 3C) and in all groups at the end of the main study (eFigure 5 in Supplement 2). In keeping with higher glucocorticoid exposure, a significant in-
crease in total body fat in the prednisone, 5 mg, twice daily and dexamethasone groups was observed when comparing base-
line to cycle 12 (12.3%; \( P = .001 \) and 19.2%; \( P < .001 \), respectively) (Figure 3D) or to the end of the main study (eFigure 5 in Supplement 2) but was not observed with the lower glucocorticoid regimens. A small but significant decrease in total body bone mineral density from baseline to the end of the main study was seen in the dexamethasone group (−2.0%; 95% CI, −3.4% to −0.6%); no other change in the bone mineral density of the total body or the arms was observed at cycle 12 or at the end of the main study (eTable 8 in Supplement 2).

**Clinical Benefit**

Among the 164 randomized patients evaluated, a PSA decrease of greater than or equal to 50% was observed in 26 patients (63.4%; 95% CI, 48.1%-76.4%) treated with prednisone, 5 mg, twice daily (n = 41), prednisone, 5 mg, once daily (n = 40), prednisone, 2.5 mg, twice daily (n = 38), or dexamethasone, 0.5 mg, once daily (n = 41). Dotted lines represent the PSA responders (50% reduction and 90% reduction in PSA).

Prostate-specific antigen plots for prednisone, 5 mg, once daily, prednisone, 2.5 mg, twice daily, and dexamethasone, 0.5 mg, once daily exclude patients who were missing either baseline or postbaseline PSA values. Increases greater than 25% were cut off at 25%.

**Discussion**

Abiraterone acetate plus prednisone is being used earlier in the treatment paradigm for prostate cancer and because patients benefit for a longer period, treatment exposure is increased.
We therefore designed a randomized clinical trial to support physician decision making in selecting a glucocorticoid regimen that balances controlling manifestations of endogenous mineralocorticoid excess with the adverse consequences of nonphysiological exposure to exogenous glucocorticoids. Manifestations of endogenous mineralocorticoid excess often present within 24 weeks of treatment, whereas the adverse consequences associated with exogenous glucocorticoids are usually associated with cumulative exposure. Prednisone, 5 mg, twice daily and dexamethasone, 0.5 mg, once daily met the prespecified threshold for the primary end point (the 95% CI excluded 50% mineralocorticoid excess), but prednisone, 5 mg, once daily and 2.5 mg twice daily did not meet the threshold.

We selected hypertension and hypokalemia as indicators of clinically relevant mineralocorticoid excess because other clinical features, including lower limb edema and general fluid overload, are less feasible to standardize in a multicenter trial. To support our clinical end point, we confirmed a significant rise in plasma ACTH and a subsequent rise in mineralocorticoid production with the lower prednisone doses. We hypothesize that monitoring of urinary glucocorticoid metabolite levels could identify patients at risk of mineralocorticoid excess and allow tailoring of the glucocorticoid dose.

A strength of the trial was monitoring for long-term adverse effects of glucocorticoid excess. Total lean body mass decreased significantly in all 3 prednisone groups and total body fat increased significantly in the prednisone, 5 mg, twice daily and dexamethasone groups. Significant increases in insulin resistance and decreases in total bone mineral density were observed only with dexamethasone. Androgen precursors were significantly suppressed with dexamethasone, 0.5 mg, once daily and prednisone, 5 mg, twice daily. The antitumor activity associated with abiraterone acetate plus glucocorticoids in the present study was similar to previous reports. In this disease setting; dexamethasone appeared particularly active but may be associated with adverse metabolic consequences.

Limitations
A limitation of the trial is that the size calculation was based on the primary end point related to the safety of individual regimens; therefore, the number of patients accrued does not allow for direct comparisons among the 4 treatment groups. Second, although patients were randomly assigned to treatment, differences at baseline could contribute to differences in outcomes. Third, testing for physiological effects and their association with clinical manifestation of mineralocorticoid excess were exploratory analyses, and we did not correct for multiple testing. Overall, the observations of our phase 2 trial require further validation.

Conclusions
This trial provides results consistent with the approved use of abiraterone acetate with prednisone, 5 mg, twice daily for the treatment of mCRPC. Long-term adverse metabolic and musculoskeletal changes are small and do not appear to have a detrimental effect on patient-reported quality of life. Different glucocorticoid regimens make distinct compromises on control of mineralocorticoid excess, changes in body composition, and development of insulin resistance. With careful monitoring, the risk of hypokalemia with lower glucocorticoid doses can be mitigated, as demonstrated in the LATITUDE and STAMPEDE trials where no major safety concerns were raised. A lower dose of prednisone appears to reduce long-term risks of insulin resistance, increased body fat, and loss of bone mineral density.
National Institute for Health Research (NIHR) Birmingham Biomedical Research Centre at the University Hospitals NHS Foundation Trust & University of Birmingham (grant reference number BRC-1215-20009). The views expressed are those of the authors and not necessarily those of the NIHR or the Department of Health and Social Care. Dr Sterneberg has received honoraria from Janssen, Astellas, Clovis, and AstaZeneca and has consulted for Bayer, Ferring, Sanofi, and Pﬁzer. Dr Feyera$mb; end has received research funding from Janssen, Bayer, Astellas, Ipsen, Roche, and MDX Health and has received honoraria from Janssen, Ipsen, Sanofi, and Astellas. Dr Berruti has consulted for and received honoraria from Janssen. Dr Joniau reports grants and personal fees from Janssen, Astellas, Ferring, Roche, and Bayer; personal fees from Ipsen; and grants from MDX Health outside the submitted work. Dr Lefresne, Ms Lahaye, Dr Shelby, and Ms Pissart are employees of Janssen. Ms Lahaye and Ms Pissart are shareholders in Johnson & Johnson. Dr Jones has received honoraria from Janssen and Astellas, and research funding from Astellas. Dr Tombal has received research funding from Astellas and Ferring, and honoraria from Janssen, Amgen, Astellas, Ferring, Bayer, and Sanofi. No other disclosures were reported.

Funding/Support: The study was funded by Janssen EMEA. Dr Attard is a Cancer Research UK Advanced Clinician Scientist fellow supported by grant No. A22744.

Role of the Funder/Sponsor: Grant No. A22744.

Additional Contributions: The authors thank the study participants and study centers (eTable 1 in the Supplement) for their contributions. The study participants did not receive compensation. The study centers received compensation for the costs of conducting the study. Jonathan Latham (PharmaScribe, LLC) provided medical writing assistance and PAREXEL International provided graphics assistance with financial support from Janssen Pharmaceutica NV.

Data Sharing Statement: See Supplement 3.

REFERENCES