Review of Palliative $^{223}$Ra in Metastatic Castration-Resistant Prostate Cancer: Experience at West Virginia University Cancer Center

Ruta Arays1, Zeeshan Ahmad1, Lorinda Howard2, Kenneth Veselicky3, Joanna Kolodney1, SijinWen4, and Thomas Hogan1

1Department of Medicine, West Virginia University Health Sciences Center, Morgantown, West Virginia; School of Nursing, West Virginia University Health Sciences Center, Morgantown, West Virginia; 2Department of Radiology, West Virginia University Health Sciences Center, Morgantown, West Virginia; and 4Department of Biostatistics, West Virginia University Health Sciences Center, Morgantown, West Virginia

The ALSYMPCA trial of the α-emitter $^{223}$Ra in symptomatic bone-predominant metastatic castration-resistant prostate cancer (mCRPC) reported a median overall survival (OS) of 14.9 mo, versus 11.3 mo for placebo. However, subsequently reported real-world experience with $^{223}$Ra in smaller mCRPC patient cohorts has appeared less successful. We performed a retrospective observational study to review our own $^{223}$Ra experience at West Virginia University (WVU). Methods: Demographic, clinical, laboratory, and imaging data were reviewed in all bone-predominant mCRPC patients treated with $^{223}$Ra at WVU from 2014 to 2019. The number of bone metastases per patient at the start of treatment with $^{223}$Ra was quantified via nuclear bone scans (12 scans, 5 of which also included SPECT/CT), body CT scans (8 scans), and PET/CT scans (4 scans). Standard descriptive statistics were used to study institutional review board–exempted, deidentified patient data. Median survival in ALSYMPCA and WVU patients was compared using a 2-sided, 1-sample log-rank test based on the exponential distributions. The primary endpoint was patient OS after initiating $^{223}$Ra. Results: Twenty-four patients received 98 infusions of $^{223}$Ra; 83% of these patients were referred from outside WVU. Before the first infusion, all 24 had received androgen deprivation therapy. In total, 73 sequential combinations of androgen deprivation therapy were used, 68 of which (93%) preceded the first $^{223}$Ra infusion. Also, before $^{223}$Ra, 19 (79%) patients had received docetaxel and 19 (79%) had received 33 courses of radiation, 24 of which targeted nonprostatic sites. Eleven patients (46%) completed all 6 planned $^{223}$Ra infusions; 13 (54%) stopped early because of clinical deterioration. As of August 2020, only 1 patient remained alive after completing 6 cycles of $^{223}$Ra. Median OS from the first $^{223}$Ra infusion to the last follow-up or death was 8.3 mo (range, 0–44 mo)—nearly 50% less than the ALSYMPCA median survival of 14.9 mo ($P = 0.01$). Compared with ALSYMPCA, more WVU patients received bisphosphonates and docetaxel, more had an Eastern Cooperative Oncology Group performance status of at least 2, more used opiates for pain, more had a greater bone metastasis burden by imaging, and more had lower hemoglobin, albumin, alkaline phosphatase, and prostate-specific antigen levels. Conclusion: Although the science supporting the development and clinical use of $^{223}$Ra is compelling, optimal clinical benefit will likely require earlier referral for $^{223}$Ra, before patients have exhausted most conventional therapies. At WVU, we found that practically all our referred patients had androgen deprivation therapy, radiation, and cytotoxic therapy before starting $^{223}$Ra. We continue to offer $^{223}$Ra therapy to patients with symptomatic bone-predominant mCRPC but are encouraging earlier patient referral.

Key Words: metastatic; castration-resistant; prostate cancer; $^{223}$Ra

DOI: 10.2967/jnmt.120.254474

Prostate cancer is the second commonest cancer in men, with an estimated 174,650 new cases and 31,620 deaths in 2019 in the United States (1). Prostate cancer patients with bone metastasis frequently experience skeleton-related events, including pathologic fractures, hypercalcemia, spinal cord compromise, and pain requiring surgery, radiotherapy, and other interventions (2). A large multicenter, randomized, placebo-controlled phase 3 trial of $^{223}$Ra in metastatic castration-resistant prostate cancer (mCRPC) (ALSYMPCA trial) with positive results led to subsequent approval of $^{223}$Ra by the U.S. Food and Drug Administration to treat symptomatic bone metastasis in patients with mCRPC with no known visceral metastasis (3). The median overall survival (OS) was 14.9 mo in patients who received $^{223}$Ra, versus 11.3 mo in those receiving placebo, a 30% reduction in the risk of death in $^{223}$Ra patients versus placebo. Also, the time to the first symptomatic skeleton-related event was significantly prolonged in patients receiving $^{223}$Ra (3).

However, postapproval real-world experience with $^{223}$Ra in mCRPC has often found shorter median survival and
greater complication rates than reported by the ALSYMPCA trial (4–19).

We decided to perform our own retrospective observational study of our clinical experience with $^{223}$Ra in mCRPC patients referred to the WVU Cancer Center.

**MATERIALS AND METHODS**

Demographic, clinical, and laboratory data were collected from electronic medical records (Epic) for every patient with prostate cancer who received $^{223}$Ra during a 5-y period beginning in 2014. No patient who received any $^{223}$Ra at West Virginia University (WVU) during this period was excluded. Because we could not retrieve several outside-laboratory results for one of the patients, laboratory data from that patient were not analyzed. A nuclear radiologist who administered the $^{223}$Ra reviewed nuclear bone scans, nuclear SPECT/CT scans, and PET scans to quantify the number of bone metastases in each patient at the start of $^{223}$Ra. Imaging included 12 nuclear bone scans (5 of which also included SPECT/CT), 8 body CT scans, and 4 PET/CT scans. Patient parameters were compared with those published from the ALSYMPCA trial patients who received $^{223}$Ra (Tables 1–3) (3).

Standard descriptive statistics (mean, median, minimum, maximum, and range) were used to study institutional review board–exempted, deidentified patient data. Median survival in ALSYMPCA and WVU patients was compared using a 2-sided, 1-sample log-rank test based on exponential distributions. The primary endpoint was WVU patient survival from the start of the $^{223}$Ra infusions, calculated using the Kaplan–Meier method (Fig. 1).

**RESULTS**

Over a 5-y period, 24 patients received 98 total monthly infusions of $^{223}$Ra. All patients resided in West Virginia, and 83% were referred from oncology practices outside WVU.

At the initial cancer diagnosis, before $^{223}$Ra, only 4 patients (17%) had disease localized to the prostate and 12 (50%) were TNM classification M1/stage 4. At the time of the first $^{223}$Ra infusion, the median age was 68 y (range, 54–89 y), and the median prostate-specific antigen value was 75.4 ng/mL (range, 1.5–928.1 ng/mL).

The 24 mCRPC patients received, in total, 73 sequential combinations of androgen deprivation therapy, including abiraterone (with prednisone), bicalutamide, degarelix, finasteride, leuprolide, megestrol, enzalutamide, or estrogen patches. All 24 patients received 1 or more trials of various androgen deprivation therapies (100%), 22 received 2 or more (92%), 17 received 3 or more (71%), and 10 received 4 or more (42%). Ninety-three percent of all the different types of androgen deprivation therapy administered preceded the first infusion of $^{223}$Ra.

Eighteen patients (75%) received docetaxel chemotherapy before $^{223}$Ra, and 8 of these had 2 or more chemotherapy agents before $^{223}$Ra. Nineteen patients (79%) received a total of 33 courses of radiation therapy; 9 courses targeted the prostate, and 24 courses targeted other palliative sites. For 18 of the 19 radiation therapy patients (95%), radiation preceded the first infusion of $^{223}$Ra.

All 24 patients (100%) received a first infusion of $^{223}$Ra; 21 (88%), a second; 16 (67%), a third; 13 (54%), a fourth; 13 (54%), a fifth; and 11 (46%), all 6 that were planned. Thus, 13 patients (54%) did not complete all 6 planned infusions of $^{223}$Ra, stopping because of clinical deterioration. The median OS in the 24 patients, from the first $^{223}$Ra

**TABLE 1**

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>ALSYMPCA</th>
<th>WVU</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patients (n)</td>
<td>614</td>
<td>24</td>
</tr>
<tr>
<td>Median survival (mo)</td>
<td>14.9</td>
<td>8.3 (P = 0.01)</td>
</tr>
<tr>
<td>Median age (y)</td>
<td>71 (range, 49–90)</td>
<td>68 (range, 54–89)</td>
</tr>
<tr>
<td>&gt;75 y old (n)</td>
<td>171 (28%)</td>
<td>4 (17%)</td>
</tr>
<tr>
<td>Caucasian (n)</td>
<td>575 (94%)</td>
<td>24 (100%)</td>
</tr>
<tr>
<td>Current bisphosphonates (n)</td>
<td>250 (41%)</td>
<td>15 (63%)</td>
</tr>
<tr>
<td>No</td>
<td>364 (59%)</td>
<td>9 (37%)</td>
</tr>
<tr>
<td>Prior docetaxel (n)</td>
<td>352 (57%)</td>
<td>17 (71%)</td>
</tr>
<tr>
<td>Yes</td>
<td>262 (43%)</td>
<td>7 (29%)</td>
</tr>
<tr>
<td>No</td>
<td></td>
<td></td>
</tr>
<tr>
<td>ECOG performance status (n)</td>
<td>0</td>
<td>165 (27%)</td>
</tr>
<tr>
<td>1</td>
<td>371 (60%)</td>
<td>14 (58%)</td>
</tr>
<tr>
<td>2+</td>
<td>77 (13%)</td>
<td>9 (38%)</td>
</tr>
<tr>
<td>WHO ladder for cancer pain (n)</td>
<td>0</td>
<td>3 (13%)</td>
</tr>
<tr>
<td>1</td>
<td>257 (42%)</td>
<td>3 (13%)</td>
</tr>
<tr>
<td>2</td>
<td>151 (25%)</td>
<td>7 (29%)</td>
</tr>
<tr>
<td>3</td>
<td>194 (32%)</td>
<td>11 (48%)</td>
</tr>
</tbody>
</table>

ECOG = Eastern Cooperative Oncology Group; WHO = World Health Organization.
infusion to the last follow-up (August 2020), was 8.3 mo
(range, 0–44 mo; P = 0.01 vs. the ALSYMPCA median
survival) (Tables 1–3). One of the 24 patients currently
remains alive with disease, having completed 6 cycles of
$^{223}\text{Ra}$.

**DISCUSSION**

The science supporting the development and clinical use
of $^{223}\text{Ra}$ is compelling. An $\alpha$-emitter, $^{223}\text{Ra}$ dichloride
($^{223}\text{Ra}$) is bone-seeking and complexes with hydroxyapatite,
releasing short-range (<100 μm), high-energy particles
that exert a selective antitumor effect (20). Areas of
active bone remodeling and bone vascular supply are
the main target sites for $^{223}\text{Ra}$ localization (21). In mouse
models, $^{223}\text{Ra}$ inhibits tumor-induced pathologic bone for-
mation in the tumor microenvironment adjacent to tumor
foci (22).

Preclinical work with mice confirmed that $^{223}\text{Ra}$ selec-
tively accumulates in the bone, with only minute amounts
of daughter radionuclides released from skeletal sites of
$^{223}\text{Ra}$ decay (23). Moreover, dosimetry studies showed that
short-range $\alpha$-radiation from $^{223}\text{Ra}$ substantially spares
bone marrow, with relatively less toxicity than is caused by
$\beta$-emitting $^{89}\text{Sr}$ (23).

A phase 1 clinical trial demonstrated that $^{223}\text{Ra}$ prefer-
etially targets bone metastases rather than diffusely target-
ing healthy bone tissue (24). A randomized, multicenter,
placebo-controlled phase 2 trial assigned patients receiving
external-beam radiation therapy for pain control to receive
either 4 $^{223}\text{Ra}$ injections (50 kBq kg$^{-1}$) at 4-wk intervals or
placebo on the same schedule. The group receiving $^{223}\text{Ra}$
demonstrated a significant decline in alkaline phosphatase
levels, a delay in the time to prostate-specific antigen pro-
gression, and a trend toward reduced skeleton-related
events and improved OS. The safety profile was acceptable,
 hematologic toxicity did not significantly differ between the
2 groups, and no patient discontinued $^{223}\text{Ra}$ because of
treatment-related toxicity (25).

Parker et al. reported the pivotal phase III study of $^{223}\text{Ra}$,
involving 614 $^{223}\text{Ra}$-treated patients versus 307 in a pla-
cebo group, treated at 136 centers in 19 countries (3). At
the time of publication, 532 of 921 patients (58%) had received
all 6 injections of $^{223}\text{Ra}$. The median OS was 14.9 mo in the
$^{223}\text{Ra}$ group and 11.3 mo in the placebo group. $^{223}\text{Ra}$ was
associated with a 30% reduction in the risk of death versus
placebo. The effect of $^{223}\text{Ra}$ on OS was consistent across all
subgroups, and $^{223}\text{Ra}$ was not associated with significantly
more grade 3 or 4 toxic effects than placebo.

However, subsequent to $^{223}\text{Ra}$ approval and widespread
use in the United States, multiple reports raised caveats

### TABLE 2

<table>
<thead>
<tr>
<th>Parameter</th>
<th>ALSYMPCA (614 patients)</th>
<th>WVU (23 patients)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total alkaline phosphatase</td>
<td></td>
<td></td>
</tr>
<tr>
<td>≤220 U/L</td>
<td>348 (57%)</td>
<td>18/23 (78%)</td>
</tr>
<tr>
<td>&gt;220 U/L</td>
<td>266 (43%)</td>
<td>5/23 (22%)</td>
</tr>
<tr>
<td>Hemoglobin (g/dL)</td>
<td>12.2 (8.5–15.7)</td>
<td>11.5 (8.6–15.3)</td>
</tr>
<tr>
<td>Albumin (g/L)</td>
<td>40 (24–53)</td>
<td>33.5 (17–39)</td>
</tr>
<tr>
<td>Total alkaline phosphatase (U/L)</td>
<td>211 (32–4631)</td>
<td>115 (68–820)</td>
</tr>
<tr>
<td>Lactate dehydrogenase (U/L)</td>
<td>315 (76–2171)</td>
<td>274 (118–809)</td>
</tr>
<tr>
<td>Prostate-specific antigen (μm/L)</td>
<td>146 (3.8–6026)</td>
<td>75.4 (1.5–928)</td>
</tr>
</tbody>
</table>

*Data are median followed by percentage or range.*

### TABLE 3

<table>
<thead>
<tr>
<th>Parameter</th>
<th>ALSYMPCA</th>
<th>WVU</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bone metastases ($n$)</td>
<td>100 (16%)</td>
<td>3/23 (13%)</td>
</tr>
<tr>
<td>6–20</td>
<td>262 (43%)</td>
<td>5/23 (22%)</td>
</tr>
<tr>
<td>&gt;20</td>
<td>195 (32%)</td>
<td>14/23 (61%)</td>
</tr>
<tr>
<td>Superscan</td>
<td>54 (9%)</td>
<td>1/23 (4%)</td>
</tr>
<tr>
<td>Radiation &lt; 12 wk from screening</td>
<td>Yes</td>
<td>99 (16%)</td>
</tr>
<tr>
<td></td>
<td>No</td>
<td>515 (84%)</td>
</tr>
</tbody>
</table>

*Figures are given as median or percentage.*

**FIGURE 1.** Kaplan–Meier OS of 24 prostate cancer patients at WVU hospital from time of first $^{223}\text{Ra}$ isotope infusion.
concerning patient selection for $^{223}$Ra, finding shorter OS and more adverse events than reported by ALSYMPCA. Most of these have been smaller cohorts in single institutions, and there have also been a few multicenter experiences. Stolten et al. and Vogelzang et al. both looked at outcomes of patients receiving the full 6 cycles versus those who could not complete treatment. The cohorts had 55–184 patients, and OS favored those who completed all 6 cycles of therapy (16 mo vs. 4 mo in the Stolten et al. cohort, and 17 mo in the Vogelzang cohort) (4,5).

Etchebehere et al. reported 41 patients with metastatic prostate cancer treated at M.D. Anderson Cancer Center who had undergone fluoride PET/CT before $^{223}$Ra (26). They defined bone marrow failure as World Health Organization grade 3 hematologic toxicity with no recovery after 6 wk, or recorded death due to bone marrow failure. Bone marrow failure correlated with tumor burden on imaging, which, by multivariable analysis, was the only independent predictor for bone marrow failure. Another report by Etchebehere et al. on 110 $^{223}$Ra-treated patients noted that improved alkaline phosphatase levels and pain scores were associated with improved outcomes (6).

Additionally, there have also been multiple reports showing better outcomes when using $^{223}$Ra earlier in the course of disease. Hague et al., Saad et al., Baldari et al., Wong et al., and Sartor et al. all found this to be the case within their respective studies (8,12,27–29). Specifically, Saad et al. had the largest cohort of these and reported on the early-access program after the ALSYMPCA study and before regulatory approval of $^{223}$Ra in 2013. In total, 839 patients were enrolled from 113 sites in 14 countries; 696 patients received one or more doses of $^{223}$Ra; 403 (58%) had all 6 planned injections (28). Sartor et al. reported the U.S. experience from the expanded-access phase II open-label, single-arm multicenter trial. Those with more advanced disease were less likely to benefit from the isotope. The median OS in those receiving 5–6 injections was not reached, whereas the median survival in those with fewer than 5 injections was 7.5 mo (12).

In July 2018, the safety committee of the European Medicines Agency—the Pharmacovigilance Risk Assessment Committee—recommended restricting the use of $^{223}$Ra-dichloride to patients who had only 2 previous treatments for metastatic prostate cancer or cannot receive other treatments (13,14). The committee recommended that $^{223}$Ra not be used with abiraterone plus corticosteroid, with other systemic cancer therapies except for hormonal therapy, and in asymptomatic patients or those with few osteoblastic bone metastases. The committee further suggested that patients be carefully assessed for fracture risk and that use of bisphosphonates or RANKL inhibitors be considered to increase bone strength before $^{223}$Ra. Although O’Sullivan et al. made a case against these restrictive recommendations, the European Medicines Agency countered that the recommendations were not only because of detrimental effects on bone fractures when used with abiraterone but also because of risk of lymph node or visceral progression during $^{223}$Ra bone-focused treatment (30,31).

Of interest, one study has looked at possible predictive measures of survival in patients receiving $^{223}$Ra. Dittman et al. used quantitative bone SPECT/CT before $^{223}$Ra in 60 patients with mCRPC and found that isotope uptake in the central skeleton varied from 11% to 56% of the injected dose. After $^{223}$Ra, median OS in all 60 patients was 15.2 mo. When skeletal uptake was 26% or more, median OS was 7.3 mo; when skeletal uptake was less than 26%, median OS was 30.8 mo (17). The authors concluded that initial quantitative SPECT/CT bone scanning could predict subsequent patient survival after treatment with $^{223}$Ra. This finding may be of future interest once more data become available.

After reviewing our own patient experience, we identified certain factors likely contributing to our patients’ shorter-than-expected median OS. Practically all androgen deprivation, radiation, and cytotoxic therapy was given before referral to WVU for $^{223}$Ra. Although most referred patients initially seemed eligible for $^{223}$Ra on the basis of Eastern Cooperative Oncology Group performance status and the usual laboratory parameters, many appeared to receive only minor benefit from $^{223}$Ra as administered. When compared with the original ALSYMPCA prostate cancer patients, more patients at WVU received bisphosphonates and docetaxel, had a performance status of more than 2, took more opiates for cancer pain, had a greater bone metastasis burden, and had lower hemoglobin, albumin, alkaline phosphatase, and prostate-specific antigen levels (Tables 1–3).

Potential limitations of our study are that it was observational and retrospective, and our Appalachian referral-patient results may be hard to extrapolate to other practice settings. The ALSYMPCA trial was multicenter, multinational, and prospective, whereas our WVU data were from a single referral center. The number of $^{223}$Ra-treated patients studied in the 2 trials differed by over 25-fold, as we had a relatively small number of patients in our dataset (the last patient reported here started $^{223}$Ra in October 2018). Also, the data-collection time frames of the 2 studies differed by more than 5 y. Imaging in our patients included 12 nuclear bone scans (5 of which also included SPECT/CT), 8 body CT scans, and 4 PET/CT scans. Thus, only 7 patients had a nuclear bone scan, and use of higher-resolution imaging in the other 17 might have increased our detection of the number of bone lesions versus earlier studies using only conventional nuclear bone scans (17). Despite these issues, our report adds to the growing amount of real-world data concerning use of $^{223}$Ra, which will hopefully guide the future clinical application of this promising isotope.

CONCLUSION

We think that our initial referral pattern resulted in more heavily pretreated cancer patients receiving $^{223}$Ra, with suboptimal results versus ALSYMPCA. Hopefully, the referral pattern will improve as patients become more aware of $^{223}$Ra availability and use, with earlier patient demand...
and earlier referral for $^{223}$Ra. Currently, $^{223}$Ra appears to be an evolving and promising treatment for bone-predominant metastatic prostate cancer and potentially for other osteoblastic tumors, including metastatic breast cancer, differentiated thyroid cancer, and renal cell carcinoma (32). However, treatment benefit will likely be optimal when $^{223}$Ra is given earlier, targeted to patients who have not already exhausted most conventional therapies. Going forward, we continue to offer $^{223}$Ra isotope therapy for patients with symptomatic bone-predominant mCRPC but are emphasizing earlier referral to our center.

DISCLOSURE

No potential conflict of interest relevant to this article was reported.

REFERENCES