CABOZANTINIB AND PROSTATE CANCER: INHIBITING SEED AND DISRUPTING SOIL?

Richard J. Lee and Matthew R. Smith

Treatment with cabozantinib, an inhibitor of MET and VEGFR2 signaling, has demonstrated clinical benefit in early trials in men with metastatic prostate cancer. Preclinical evidence suggests that cabozantinib can kill cancer cell seeds while disrupting angiogenesis and stromal cells in the metastatic soil. *Clin Cancer Res;* 20(3); 525–7. ©2013 AACR.

In this issue of *Clinical Cancer Research,* Dai and colleagues provide evidence that the tyrosine kinase inhibitor cabozantinib impacts both cancer cells and microenvironment in bone metastasis models of prostate cancer (1). Cabozantinib is an orally bioavailable inhibitor of MET, VEGFR2, RET, and several other tyrosine kinases. The early clinical experience with cabozantinib in patients with metastatic castration-resistant prostate cancer (mCRPC) demonstrated marked improvements in bone scans of subjects with bone metastases (2). The bone scan improvement was not limited to an improvement in imaging; subjects reported clinical benefits with improved pain perception and decreased narcotic pain medication requirements. Bone scans are an indirect measure of tumor activity because they detect deposition of the technetium tracer by osteoblasts. It is unclear whether cabozantinib leads to clinical benefit by impacting prostate cancer cell viability, osteoblast viability, perfusion of bone tumor deposits, or a combination of effects.

MET and VEGFR2 signaling pathways are active in both bone turnover and prostate cancer progression (Fig. 1; ref. 3). Bone remodeling is a continuous process balancing bone-forming activity by osteoblasts with the resorptive activity of osteoclasts. The balance of bone turnover is mediated by receptor activator of nuclear factor-kappa B (RANK) signaling between osteoblasts and osteoclasts. In addition, MET and VEGFR and their respective ligands, hepatocyte growth factor (HGF) and VEGF, are expressed by both osteoblasts and osteoclasts. HGF/MET and VEGF/VEGFR signaling mediates both autocrine and paracrine roles in the normal activity and survival of osteoblasts and osteoclasts. MET and VEGFR signaling also play important roles in prostate cancer progression and bone metastasis. Increased MET and HGF expression in prostate cancer cells correlate with disease recurrence and metastasis, with highest levels in bone metastases. VEGFR signaling is critical for angiogenesis, a key step in tumor growth. Higher VEGFR2 levels are expressed in high-grade prostate cancers. Prostate cancer cells also express VEGF, unlike their benign counterparts. Higher levels of VEGF are independent predictors of worse overall survival (OS) in men with mCRPC. The activities of MET and VEGFR signaling in bone turnover and metastasis provide a strong rationale for dual inhibition of these pathways as a therapeutic strategy in men with mCRPC.

Inhibition of either MET or VEGF/angiogenesis pathways in men with mCRPC has been evaluated in clinical trials (reviewed in (3)). Four phase III clinical trials targeted angiogenesis with bevacizumab, sunitinib, lenalidomide, or aflibercept; none prolonged OS. A fifth phase III study of the agent tasquinimod is ongoing. A phase II randomized trial evaluating the HGF inhibitor rilotumumab did not improve OS.

Dual inhibition of MET and VEGFR2 with cabozantinib has shown promise in mCRPC and other malignancies (2). Cabozantinib is U.S. Food and Drug Administration–approved for advanced medullary thyroid cancer (MTC), due in part to its inhibition of RET, which is frequently mutated in MTC (4). In a randomized discontinuation study involving subjects with mCRPC, cabozantinib (100 mg daily) was associated with 5% objective response rate at 12 weeks [soft tissue lesions measured by Response Evaluation Criteria in Solid Tumors (RECIST)] and 68% rate of bone scan improvement, including complete resolution in 12%. Bone pain was improved in 67% of evaluable subjects with a decrease in narcotic use in 56% (2). Prostate-specific antigen (PSA) changes were often discordant with bone scan responses. Dose reductions occurred in 62% of subjects due to adverse events. A dose-ranging study of cabozantinib in mCRPC identified a lower dose (40 mg daily) with similar activity but improved tolerability (5). The effects of cabozantinib on OS and pain improvement in men with mCRPC are being evaluated in two current phase III clinical trials (COMET-1 and COMET-2).

In this issue, Dai and colleagues evaluate whether cabozantinib affects prostate cancer cells, bone stroma, or both,
using established cell lines and in vivo models of metastatic progression. Cabozantinib diminished viability of prostate cancer and preosteoblastic cell lines, and inhibited differentiation of preosteoclast cells, consistent with the important roles of MET and VEGFR2 in the normal activity and survival of the three cell types that interact in bone metastases. Cabozantinib affects the intended molecular targets, with decreased phosphorylation of MET and VEGFR2. In mouse models, cabozantinib had differential effects on tumor growth depending on the prostate cancer cell line and the site of tumor growth. Cabozantinib inhibited intratibial metastasis of prostate cancer cell lines that produce osteoblastic or mixed blastic/lytic lesions, but did not affect the intratibial growth of a prostate cancer cell line that produces osteolytic metastases (PC-3). Cabozantinib did inhibit growth of the same PC-3 cell line when implanted subcutaneously, however, suggesting that the drug’s activity is context dependent. Together, these studies suggest that the clinical activity of cabozantinib may result from both inhibiting the seed and disrupting the soil of Paget’s classic “seed and soil” hypothesis of metastasis.

These data add to our understanding of metastasis and raise important questions about the role of cabozantinib in the management of mCRPC. First, context of tumor type and metastatic niche are critical. The same cell line (PC-3) grew tumors in bone and soft tissue, but cabozantinib only decreased tumor-associated vasculature and inhibited growth in the subcutaneous tumors. Because bone is a prevascular tissue, bone metastases may be less dependent on angiogenesis for nutrient supply, compared with subcutaneous tumors. Thus, some subcutaneous tumors may be more susceptible to antiangiogenic therapies. These studies highlight the need for accurate modeling of the heterotypic interactions between tumor cells and stroma.

Second, not all bone metastases are created equal. Prostate cancer can produce osteoblastic, osteolytic, or mixed blastic/lytic lesions. Could the nature of the bone metastases predict susceptibility to dual VEGFR/MET inhibition? Cell lines that create predominantly osteolytic metastases (like PC-3) may release more growth factors from the bony matrix and thereby have less dependence on angiogenesis for establishment of viable tumors. The therapeutic implication is that such tumors may be less susceptible to antiangiogenic therapy and by extension, less sensitive to dual pathway inhibition of therapies like cabozantinib. Because we know from large clinical trials that inhibition of just one pathway does not improve OS, perhaps we can use these observations to identify predictive biomarkers (i.e., denoting predominance of osteoblastic lesions) that can improve patient selection for cabozantinib therapy.

Third, inhibition of two important targets may be insufficient to diminish overall tumor burden in patients. Much
like PC-3 response to cabozantinib is dependent on whether the tumor is in bone or soft tissue, men with CRPC metastases to both soft tissue and bone may require coadministration of multiple targeted therapies that can disrupt different kinds of soil.

Finally, the promise of cabozantinib may lie in its ability to target both seed and soil. By inhibiting prostate cancer cell growth and inducing apoptosis, cabozantinib may be tumoricidal. By disrupting vasculature or affecting viability and activity of osteoblasts and osteoclasts, cabozantinib may be tumoristatic. The clinical evidence of rapid improvement in bone scans but discordant PSA response in some subjects with mCRPC suggests that the initial effect may relate to altered perfusion of bone metastases. The discordance between PSA and bone scan response in some patients suggests that cabozantinib may be tumoricidal in some tumors but tumoristatic in others. In conclusion, the compelling preclinical observations by Dai and colleagues provide mechanism(s) for the clinical activity of cabozantinib in mCPRC. Their observations may also inform future development of cabozantinib including the rational selection of combination therapies.

**Disclosure of Potential Conflicts of Interest**

R.J. Lee received research support from Exelixis. M.R. Smith was a consultant/advisory board member of Exelixis.

**Authors' Contributions**

Conception and design: R.J. Lee, M.R. Smith

Analysis and interpretation of data (e.g., statistical analysis, biostatistics, computational analysis): R.J. Lee, M.R. Smith

Writing, review, and/or revision of the manuscript: R.J. Lee, M.R. Smith

Administrative, technical, or material support (i.e., reporting or organizing data, constructing databases): R.J. Lee

**Grant Support**

This study was supported by the Department of Defense Prostate Cancer Research Program under Award W81XWH-09-1-0471 and a Conquer Cancer Foundation Career Development Award (R.J. Lee), by NIH Midcareer Investigator Award No. 5K24CA121990 (M.R. Smith); and competitive research awards from the Prostate Cancer Foundation.

Received October 18, 2013; accepted October 29, 2013; published OnlineFirst November 27, 2013.

**References**


Cabozantinib and Prostate Cancer: Inhibiting Seed and Disrupting Soil?

Richard J. Lee and Matthew R. Smith


Updated version
Access the most recent version of this article at:
doi:10.1158/1078-0432.CCR-13-2636

Supplementary Material
Access the most recent supplemental material at:
http://clincancerres.aacrjournals.org/content/suppl/2014/02/04/1078-0432.CCR-13-2636.DC1

Cited articles
This article cites 5 articles, 4 of which you can access for free at:
http://clincancerres.aacrjournals.org/content/20/3/525.full.html#ref-list-1

E-mail alerts
Sign up to receive free email-alerts related to this article or journal.

Reprints and Subscriptions
To order reprints of this article or to subscribe to the journal, contact the AACR Publications Department at pubs@aacr.org.

Permissions
To request permission to re-use all or part of this article, contact the AACR Publications Department at permissions@aacr.org.