NSD2 is a conserved driver of metastatic prostate cancer progression

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Deciphering cell-intrinsic mechanisms of metastasis progression in vivo is essential to identify novel therapeutic approaches. Here we elucidate cell-intrinsic drivers of metastatic prostate cancer progression through analyses of genetically engineered mouse models (GEMM) and correlative studies of human prostate cancer. Expression profiling of lineage-marked cells from mouse primary tumors and metastases defines a signature of de novo metastatic progression. Cross-species master regulator analyses comparing this mouse signature with a comparable human signature identifies conserved drivers of metastatic progression with demonstrable clinical and functional relevance. In particular, nuclear receptor binding SET Domain Protein 2 (NSD2) is robustly expressed in lethal prostate cancer in humans, while its silencing inhibits metastasis of mouse allografts in vivo. We propose that cross-species analysis can elucidate mechanisms of metastasis progression, thus providing potential additional therapeutic opportunities for treatment of lethal prostate cancer.

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Metastasis is a complex process that culminates in the progressive accumulation of molecular alterations of cancer cells, which allow them to escape the confines of the tumor, survive during dissemination, and ultimately reside at distant sites, wherein requisite adaptive changes ensue in their new microenvironment. Therefore, it would be most informative to study the biological processes and molecular mechanisms underlying metastatic progression as occur in the context of the whole organism in vivo. However, inherent challenges in accessing primary tumors and their metastases from cancer patients have made it difficult to study de novo metastasis formation. Moreover, most in vivo studies of metastasis have utilized transplantation models wherein cells or tumors are implanted into host organisms, usually immune-deficient ones. While such investigations have advanced our understanding of metastasis mechanisms and have elucidated factors that promote organ tropism, they may not ideally model the cell-intrinsic mechanisms of de novo metastatic progression. Analyses of genetically engineered mouse models (GEMMs) can overcome these obstacles, since they enable access to tumors and their resultant metastases as they arise de novo during cancer progression in the whole organism.

Virtually all prostate cancer deaths are due to metastasis, which arises at advanced disease stages and is often resistant to treatment. Indeed, while patients with locally confined disease have highly favorable outcomes (>95%), the 5-year survival for metastatic prostate cancer is less than 30%. Standard treatment for advanced prostate cancer involves androgen deprivation therapy, which is initially effective but ultimately leads to disease recurrence in the form of castration resistant prostate cancer (CRPC), which is highly aggressive and prone to metastasis. While second generation anti-androgens, such as enzalutamide and abiraterone acetate, are now being used for treatment of CRPC, treatment failure is often associated with progression to even more aggressive subtypes, including neuroendocrine prostate cancer (NEPC). Frequent sites of prostate cancer metastasis are bone and lymph nodes, however, visceral metastases, such as to lungs and liver, are becoming more prevalent in aggressive variants and associated with increased lethality and poor prognosis.

Several recent studies have identified the landscape of recurrent genomic alterations in prostate tumors and metastases. The culmination of these analyses has revealed that metastatic prostate cancer has a significantly higher burden of mutational and somatic copy number alterations compared with primary tumors. These include increased frequency of alterations of key oncogenic and tumor suppressor genes such as AR, PTEN, TP53, and RB1, and aberrant activation of key signaling pathways such as the PI-3 kinase, FGF receptor, and RAS signaling pathways. However, functional analyses of causal drivers of metastatic prostate cancer progression have been hindered by the lack of experimental models that enable biological and molecular investigations of de novo metastasis in context of the whole organism.

In the current study, we have investigated metastatic progression in vivo in a GEMM of prostate cancer. Employing lineage tracing to isolate tumor and metastatic cells, we have defined a molecular signature of metastasis progression. Cross-species computational analyses comparing this mouse signature with a comparable human signature of metastatic prostate cancer progression have identified conserved master regulators of metastasis progression that drive these processes. In particular, we have identified NSD2 as a conserved master regulator of metastatic prostate cancer progression and a robust marker of lethal prostate tumors. Our findings suggest that cross-species investigations based on analyses of de novo metastasis in GEMMs can be broadly used to elucidate mechanisms of metastatic progression and identify potential new therapeutic opportunities for treatment of lethal cancer.

Results

A molecular signature of de novo metastasis progression. To elucidate mechanisms of metastasis progression, we utilized a previously described GEMM of highly penetrant metastatic prostate cancer based on an inducible Cre (CreERT2) expressed under the control of the promoter of the Nkx3.1 homeobox gene. This Nkx3.1CreERT2 allele drives Cre-mediated recombination in an appropriate cell of origin of prostate cancer, while simultaneously resulting in heterozygosity for Nkx3.1, which is prevalent in human prostate cancer. We crossed the Nkx3.1CreERT2 allele with a Pten floxed allele (Ptenfl/fl) and an activatable mutant K-Ras allele (KrasLSL-G12D/+; FlpO/FlpO) to generate NPK mice (for Nkx3.1CreERT2; Ptenfl/fl; KrasLSL-G12D/+). Tumor induction of these NPK mice leads to co-activation of PI3K and RAS signaling, as frequently occurs in lethal prostate cancer in human beings, while these mice develop metastasis with 100% penetrance. These NPK mice also contain a conditionally activatable fluorescent reporter allele, R26RFP, which enables in vivo lineage tracing of primary tumors and their resultant metastases with high efficiency and specificity.

To identify a molecular signature of metastatic progression, we compared expression profiles of primary tumors and metastases from NPK mice. Since these mice exhibit temporal progression from pre-invasive (~1 month), to invasive prostate cancer (~3 months), and ultimately to metastasis (~5 months), we analyzed expression profiles of primary tumors from mice prior to the occurrence of overt metastasis (pre-metastatic, <3 months, n = 8), as well as primary tumors from mice that had developed overt metastases (post-metastatic, ~5 months, n = 8). Further, since these NPK mice metastasize primarily to soft tissues, including lung, liver, and lymph node, we analyzed metastases from these various sites (n = 8, 5, 7, respectively); however, since lung is the most prevalent metastatic site, we focused our molecular analyses primarily on lung. As controls, we performed comparable analyses using non-metastatic primary tumors from NP mice (for Nkx3.1CreERT2; Ptenfl/fl; n = 7).

To focus on cell-intrinsic molecular features of primary tumor and metastatic cells free of the surrounding stromal or other components of the microenvironment, which are likely to differ for each tissue, we isolated YFP-lineage-traced cells from tumors and metastases using fluorescence activated cell sorting (FACS). We then performed RNA sequencing on the purified YFP-labeled cells to generate expression profiles corresponding to pre- or post-metastatic primary tumors (n = 8/group) as well as lung, liver, and lymph node metastases (n = 8, 5, 7, respectively; Supplementary Data 1).

Interestingly, we found that the expression profiles of the pre- and post-metastatic primary tumor cells were highly dissimilar, whereas expression profiles from the post-metastatic primary tumors were very similar to those from lung, liver, and lymph node metastases (Supplementary Data 1; Fig. 2, Supplementary Fig. 1). Specifically, unsupervised principal component analyses (PCA) revealed that expression profiles from the post-metastatic NPK primary tumors clustered more closely with the lung metastases, as well as the metastases to other sites, and further from the pre-metastatic tumors from these NPK mice, whereas the pre-metastatic NPK tumors tended to cluster more closely with non-metastatic NP primary tumors (Fig. 2a; Supplementary Fig. 1a). This relationship was further confirmed by gene set enrichment analyses (GSEA) wherein a differential expression signature comparing post- versus pre-metastatic primary tumors
was significantly enriched with a signature of lung metastases versus pre-metastatic tumors in both the positive (NES = 19.64, \( p < 0.001 \)) and negative (NES = \(-7.52, p < 0.001\)) leading edges (Fig. 2b). Moreover, expression signatures from the other metastatic sites, namely lung, liver and lymph nodes, were also highly enriched relative to each other (Supplementary Fig. 1b–d), providing further evidence of their similarity.

These observations suggest that the most prominent cell-intrinsic molecular changes that occur during metastatic progression in the NPK mice are those that distinguish pre-metastatic from post-metastatic tumors. Hence, taking into consideration: (1) the distinct molecular changes between pre- and post-metastatic NPK tumors; (2) the overall similarity of gene signatures of metastatic cells at the various tissue sites (i.e., lung, liver and lymph node); and (3) that the lung is the major metastatic site in the NPK mice, our subsequent analyses was done using a signature of metastasis progression based on the differentially expressed genes between the pre-metastatic tumors and lung metastases (\( n = 8 \)/group, respectively; two-sample two-tailed Welch t-test; Supplementary Data 1).

Notably, this mouse metastasis progression signature shares molecular features that have been associated with the hallmarks of metastasis progression in other cancer contexts\(^\text{12}\). In particular, differentially expressed genes from the positive leading edge of the GSEA (from Fig. 2b) include those associated with epithelial-mesenchymal transition (e.g., \(Vim\), \(Zeb1\), and \(Twist2\)), cell and focal adhesion (e.g., \(Itga5\), \(Col4a1\), and \(Col4a2\)), membrane type matrix metalloproteinases (e.g., \(Mmp14\)), and developmental pathways (e.g., \(Wnt5A\)) as well as other genes known to promote metastasis of prostate (e.g., \(Etv5\)) or other cancers (e.g., \(Hmga2\)) (Fig. 2c). Similarly, genes from the negative leading edge (from Fig. 2b) include genes associated with the immune response, such as the interferon regulatory factor \(Irf7\) which has been shown to be a critical regulator of immunosurveillance in cancer metastasis\(^\text{36}\) (Fig. 2d).

Furthermore, GSEA of biological pathways comparing the mouse metastasis progression signature with the MSigDB Hallmarks dataset revealed a significant enrichment of pathways that are commonly associated with metastatic progression in other tumor contexts, including epithelial to mesenchymal transition, E2F targets, \(Myc\) targets, TGF beta, and P53 pathway among others (\( p < 0.05; \) Fig. 2e; Supplementary Data 2). Notably, many of the pathways enriched in this metastasis progression signature based on tumor versus lung metastases from the NPK mice were also enriched in analogous signatures based on tumor versus liver or lymph node metastases (\( p < 0.05; \) Supplementary Fig. 1e, f), further emphasizing the overall similarity of cell-intrinsic molecular programs associated with metastasis progression across these various organ sites.

Most notably, the mouse metastasis progression signature was highly conserved with a corresponding signature of human metastatic prostate cancer progression reported by Balk and colleagues, which compares primary tumors with metastatic bone biopsies\(^\text{37}\) (Supplementary Table 1). In particular, GSEA comparing a “humanized” version of the mouse metastasis progression signature with the Balk human prostate cancer metastasis progression signature demonstrated their significant similarity in both the positive (NES = 8.02, \( p < 0.001 \)) and negative (NES = \(-3.38, p = 0.002\)) leading edges (Fig. 2f). Furthermore, GSEA comparing a “humanized” version of the mouse post-metastatic versus pre-metastatic progression signature with this human prostate cancer metastasis progression signature also demonstrated strong enrichment in both the positive (NES = 12.12, \( p < 0.001 \)) and negative (NES = \(-2.67, p = 0.0035\)) leading edges (Supplementary Fig. 1g). Further, the leading edge genes between these GSEA comparisons (i.e., from Fig. 2f and Supplementary Fig. 1g) were highly similar (overlap of the positive leading edges was 90.5% and the negative leading edge was 80%; \(X^2 p < 0.0001\)). Taken together, these molecular analyses define a cell-intrinsic signature of de novo metastasis progression in the NPK mouse model that is highly conserved with metastasis progression of human prostate cancer.

Conserved master regulators of metastasis progression. We performed cross-species computational analyses to identify
Fig. 2 A molecular signature of de novo metastasis progression. a Principal component analysis (PCA) comparing expression profiles from pre-metastatic (pre-met, blue) or post-metastatic (post-met, red) primary tumors, or lung metastases (lung, black) from NPK mice (n = 8/group). Note that Principal Component 1, capturing 40.6% of gene expression variance, effectively distinguishes pre-metastatic tumors from post-metastatic tumors and lung metastases. b GSEA comparing a reference signature of mouse lung metastases (lung mets) versus pre-metastatic tumors to a query signature of mouse post-metastatic versus pre-metastatic tumors. c, d Heat map representations of differentially expressed genes from the positive and negative leading edges, respectively, from the GSEA in panel b. The color key shows relative expression levels of the differentially expressed genes (red corresponds to overexpressed genes while blue corresponds to underexpressed genes). e Pathway enrichment analysis using the mouse metastasis progression signature defined between lung metastasis versus pre-metastatic tumors, as in panel b, to query the hallmark pathways dataset from the molecular signatures database (MSigDB). Red and blue nodes indicate positive and negative enrichment, respectively (p < 0.05). Thickness of arrows indicate the overlap of genes in the leading edges. The p-values correspond to the GSEA enrichment, and the relative size of the node indicates the relative p-value, as shown. f Cross-species GSEA comparing a reference mouse metastasis progression signature (lung metastasis versus pre-metastatic tumors, as in panel b) with a query gene set from a human metastasis signature defined between bone metastasis biopsies versus primary tumors from Balk et al. (Supplementary Table 1). For GSEA, red vertical bars indicate overexpressed query genes and blue vertical bars indicate underexpressed query genes. GSEA were done using the top 200 differentially expressed genes; p-values were calculated using 1000 gene permutations. ES: enrichment score, NES: normalized enrichment score.
conserved master regulators (MRs) of metastasis progression by interrogating genome-wide regulatory networks, or interactomes, for mouse and human prostate cancer38, using the master regulator inference analysis (MARINA) algorithm39. First, we interrogated the individual mouse and human prostate cancer interactomes with their respective metastatic progression signatures, which defined independent lists of mouse and human MRs of metastatic progression (Fig. 3a). We subsequently integrated these individual mouse and human MR lists using Stouffer integration to define the subset of conserved candidate MRs (n = 485 MRs with Stouffer integrated \( p < 0.0001 \); Supplementary Data 3). Gene ontology analysis of these integrated MRs revealed an over-representation of genes associated with all aspects of epigenetic regulation, including histone modification, DNA methylation, and chromatin remodeling40–42 (n = 136/485 genes, 28%; Supplementary Data 3). Because of the potential significance of perturbations of epigenetic regulation for metastatic progression43,44, particularly in lethal prostate cancer21, and since epigenetic regulators are potential therapeutic targets40–42, we focused on the subset of conserved MRs that are predicted to function as epigenetic regulators.

To further prioritize these candidate MRs, we used a Cox proportional hazard model to assess the association of their MR activity with prostate cancer-specific survival (where activity for a given MR is defined based on the expression levels of its MR transcriptional targets, see Methods). In particular, we used a human prostate cancer cohort described by Sboner et al., which has more than 30 years of clinical follow-up data based on death due to prostate cancer45 (Supplementary Table 1). These analyses identified a subset of 41 MRs whose activities were significantly associated with prostate cancer-specific survival (Wald test \( p < 0.05 \); Fig. 3b). Among these, we focused our subsequent analysis on 8 candidate MRs: (1) that are associated with adverse disease outcome and prostate-cancer specific lethality (Fig. 3b, c); (2) that are broadly activated in multiple metastatic organ sites (Supplementary Fig. 2a); and (3) whose activities are up-regulated (rather than repressed) in metastasis progression (Fig. 3b, c; Supplementary Fig. 2a), and therefore are potentially targets for treatment inhibition. In particular, these 8 candidate MRs are predicted to be highly activated across multiple metastatic sites, namely lung, liver and lymph node (MARINA \( p < 0.001 \); Supplementary Fig. 2a), and are significantly associated with adverse disease outcome (Wald test \( p \leq 10^{-5} \); Fig. 3c).

We further assessed the clinical relevance of these 8 candidate metastasis MRs using several independent cohorts of advanced prostate cancer patients, including those with clinical endpoints of metastasis or lethality due to prostate cancer (Supplementary Table 1). First, we performed hierarchical clustering on the activity levels of the 8 candidate MRs using the Grasso et al. cohort46, which showed that each of these MRs robustly stratify metastases (n = 35) from primary tumors (n = 59) (Fig. 3d). We observed similar findings with a second cohort that included primary tumors from The Cancer Genome Atlas (TCGA; n = 497)27 and metastases from the SU2C cohort (n = 51)21 (Supplementary Fig. 2b).

Additionally, using two independent patient cohorts with clinical follow-up, we found that these 8 candidate MRs predict disease outcome as evidenced by Kaplan–Meier survival analyses based on MR activity levels. In particular, activity of the candidate MRs stratified prostate cancer patients based on their risk of biochemical recurrence using the Glinsky et al. cohort47 (n = 79 primary prostate tumors; log-rank \( p = 0.00605 \); Hazard ratio = 2.452; Fig. 3e). Furthermore, we found that the activities of these candidate MRs also stratified patients based on the risk of death due to prostate cancer in the Sboner et al. cohort45 (n = 281 primary tumors; log-rank \( p = 8.66 \times 10^{-6} \); Hazard ratio = 2.635; Fig. 3f). Notably, the predictive ability of the 8 candidate MRs for adverse disease outcome in both of these cohorts was highly specific when compared to other MRs selected at random (significance for 8 candidate MRs versus randomly selected MRs was \( p\text{-value} = 0.0011 \) for Sboner et al. and \( p\text{-value} = 0.0214 \) for Glinsky et al.; Supplementary Fig. 2c, d).

To evaluate the functional relevance of the 8 candidate MRs for tumor growth and metastasis progression in vivo, we used an allograft cell model derived from the NPK mice, which recapitulates the pattern of NPK primary tumor growth and metastasis when engrafted into host mice3. In particular, we performed shRNA-mediated silencing of each of the candidate MRs using a minimum of 2 shRNAs for each gene (Supplementary Fig. 3; Supplementary Table 3). Analysis in vitro revealed that NPK cells having individually silenced MRs displayed reduced colony formation and reduced invasive potential compared with the control cells, albeit to varying extents for each MR (\( p < 0.05 \), two-tailed Student’s t-test; Supplementary Fig. 3b–d). Furthermore, when engrafted in vivo, these MR-silenced NPK cells displayed reduced tumor growth (\( p < 0.05 \), two-way ANOVA) and/or reduced incidence of metastasis compared with the control cells, also to varying degrees for each MR (\( p < 0.05 \), two-tailed Student’s t-test; Supplementary Fig. 3e–g). Taken together, these cross-species computational systems analyses have identified conserved master regulators of metastasis progression that are associated with adverse disease outcome and functionally relevant for prostate cancer progression.

**NSD2 is a driver of metastatic prostate cancer progression.** Among the candidate MRs, the highest level of MR activity (Fig. 3c) as well as experimentally determined functional activity (Supplementary Fig. 3) were observed for the histone methyltransferase, Nuclear receptor binding SET Domain protein 2 (NSD2). Notably, NSD2 is a putative cofactor of androgen receptor48 that has been previously implicated in advanced prostate cancer49–51, and has been shown to collaborate with RAS signaling in other tumor contexts52. Therefore, we examined the expression of NSD2 at the mRNA and protein levels in non-metastatic and metastatic contexts in both mouse and human prostate cancer (Fig. 4). In the mouse prostate, we found that Nsd2 protein is expressed at low levels in non-metastatic tumors and is strongly expressed in nuclei of lethal castration-resistant adenocarcinomas (CRPC-Adeno; Fig. 4b). In human prostate cancer, we found that NSD2 expression is increased during cancer progression at both the mRNA and protein levels (Fig. 4b–e). In particular, expression of NSD2 mRNA levels were significantly higher in more advanced (Gleason \( \geq 4 + 4 \); n = 104) versus earlier stage (Gleason <4+4; n = 173) prostate primary tumors reported in TCGA27 (\( p = 2.65 \times 10^{-9} \) two-sample two-tailed Welch t-test; Fig. 4b). Further, NSD2 expression was significantly higher in prostate cancer metastases reported in the SU2C cohort21 (n = 51) as compared with primary tumors from TCGA (n = 333; \( p = 1.64 \times 10^{-10} \) two-sample two-tailed Welch t-test; Fig. 4c).

To evaluate expression of NSD2 protein in human prostate cancer, we performed immunohistochemistry on a human prostate cancer tissue microarray (n = 100) comprised of benign tumors (n = 26), non-lethal prostate adenocarcinoma (n = 25), lethal castration-resistance adenocarcinomas (CRPC-Adeno;
**Fig. 3** Conserved master regulators of metastasis progression. **a** Strategy: mouse and human metastasis progression signatures (as in Fig. 2f) were used to interrogate mouse and human prostate cancer interactomes, respectively, using the MARiNa algorithm. Independent lists of mouse and human master regulators (MRs) were integrated to identify conserved MRs, which were prioritized by clinical and functional validation. **b** Scatter plot showing the association of the 136 conserved MRs (Supplementary Data 3) to clinical outcome using the Sboner et al. dataset, which reports prostate cancer-specific survival as the clinical endpoint (Supplementary Table 1). The Y axis represents the Cox proportional hazard p-value and the X axis represents the fold change based on MR activity. MRs that are inactive (blue) relative to primary tumors have negative fold change values and those that are active (red) have positive fold change values. **c** Summary of the 8 candidate MRs depicting their positive (activated; red bars) and negative (repressed; blue bars) targets. Shaded boxes show the ranks of differential activity and differential expression (darker is higher and lighter is lower); the numbers indicate their rank in the differential expression signature (gray indicates that a specific gene is not present on the utilized gene expression platform, yet its targets are present). **d** Kaplan–Meier survival analysis based on the activity levels of the 8 candidate MRs in: **e** Glinsky et al. (n = 79), with biochemical recurrence as the clinical end-point; and **f** Sboner et al. (n = 281), with prostate cancer-specific survival as the clinical endpoint (Supplementary Table 1). P-values were estimated using a log-rank test to determine the difference in outcomes between patients with higher MR activity levels (red) versus those with lower/no MR activity (blue).
While NSD2 was either not expressed or expressed at low levels in the non-lethal tumors, its expression increased dramatically in advanced disease stages and was particularly robust in the most aggressive phenotypes, namely CRPC adenocarcinomas and NEPC (p < 0.01, two-tailed Fisher’s exact test; Fig. 4e). To further evaluate the relationship of NSD2 expression with progression to lethal prostate cancer, we
examined matched sets of primary tumors and metastases from the same patient (n = 3). Whereas expression in the primary tumors was scattered and focal, NSD2 was robustly expressed in metastases from these patients (Fig. 4f). These findings extend previous studies showing increased expression of NSD2 in advanced prostate cancer.

To evaluate the functional consequences of NSD2 for disease progression and metastasis, we used the mouse NPK metastatic allograft model, as described above, as well as human DU145 prostate cancer cells, which model aggressive disease. In particular, we used lentiviral gene delivery to introduce a minimum of two independent shRNAs to silence NSD2 specifically. We used lentiviral gene delivery to introduce a minimum of two independent shRNAs to silence NSD2. In particular, we used lentiviral gene delivery to introduce a minimum of two independent shRNAs to silence NSD2. In particular, we used lentiviral gene delivery to introduce a minimum of two independent shRNAs to silence NSD2.

NSD2 has been reported to function as a histone methyltransferase that targets the histone H3 di-methyl mark on lysine 36 (H3K36me2) in mouse tumors. Accordingly, we found that silencing of NSD2 in either human or mouse cells resulted in a modest but reproducible reduction of the H3K36me2 mark, while not altering the mono-methyl marks on lysine 36 (H3K36me1) or other histone marks such as tri-methyl lysine 27 (H3K27me3) or lysine 9 (H3K9me3) (Fig. 5c). Furthermore, NSD2 silencing in either mouse NPK cells or human DU145 cells in vitro resulted in a 5–10 fold inhibition of colony formation (p < 0.0001, two-tailed Student’s t-test), as well as significantly decreased invasion (p < 0.01, two-tailed Student’s t-test; Fig. 5d–g).

Moreover, analyses of NPK metastatic allografts in vivo revealed that Nsd2 silencing resulted in increased overall survival (n = 10/group; p = 0.0005, log-rank; Fig. 6a) as well as a significant reduction of metastatic burden while not affecting primary tumor growth (n = 9/group; p < 0.03, Mann–Whitney U test; Fig. 6b–d). Notably, these Nsd2-silenced tumors had profoundly reduced expression of Nsd2 protein compared with the control tumors, as well as reduced expression of the corresponding H3K36me2 mark, but not other histone marks, such as H3K9me3 or H3K27me3 (Fig. 6e, f). Taken together, these observations demonstrate that increased expression of NSD2 is associated with lethal and metastatic prostate cancer, and establish the functional relevance of NSD2 for metastatic prostate cancer progression.

To consider whether it might be feasible to pharmacologically target NSD2 activity to inhibit prostate cancer progression and tumor growth, we used a small molecule inhibitor of NSD2 called MCTP-39 (3-hydrazinoquinoxaline-2-thiol), which has been reported to be a lysine-HMTase inhibitor that is a competitor of the SAM (Sterile Alpha Motif) domain. We found that MCTP-39 inhibited the H3K36me2 mark, while not affecting other histone marks such as H3K9me3 or H3K27me3 (Fig. 7a); notably, the degree of reduction the H3K36me2 mark following treatment with MCTP-39 in vitro was comparable to the degree that the H3K36me2 mark was reduced following silencing of NSD2 in vitro (Fig. 5c).

Treatment with MCTP-39 in human DU145 cells in vitro resulted in a significant dose-dependent reduction in colony formation (>10 fold; p < 0.01 two-tailed Student’s t-test; Fig. 7b). Since we found that this inhibitor was well tolerated in vivo (Supplementary Fig. 4), we evaluated the effect of MCTP-39 on tumor growth of human prostate cells in vivo by establishing DU145 xenografts (Fig. 7c–f). We found that DU145 xenografts treated with MCTP-39 had a significant decrease in tumor volume (n = 14 vehicle-treated and n = 15 MCTP-39-treated; p < 0.001, two-way ANOVA; Fig. 7c, d). The resulting MCTP-39 treated tumors had reduced expression of the H3K36me2 mark, but not other histone marks such as H3K9me3 or H3K27me3 (Fig. 7e, f). Together with the results of silencing NSD2 in vivo, these findings regarding MCTP-39 treatment suggest that NSD2 may be a target for intervention in advanced prostate cancer.

Discussion

Our study demonstrates the value of cross-species integration of molecular data from genetically engineered mouse models (GEMM) and human cancer to elucidate cell-intrinsic mechanisms of de novo metastasis progression. Notably, our current study, which identifies conserved drivers of metastatic progression by isolation of lineage-marked cells directly from tumors and metastases from a prostate cancer GEMM, complements and extends previous work that identified mechanisms of lung cancer metastasis using cell lines generated from tumors and metastases of lung cancer GEMM. Among its advantages for our current investigations, the NPK mouse model displays a highly penetrant metastatic phenotype with consistent temporal progression, and it incorporates in vivo lineage tracing of the primary tumors and metastases. Thus, by comparing metastasis progression signatures from NPK mice with comparable signatures from human prostate cancer, we have identified conserved master regulators (MRs) of metastasis progression that are associated with adverse disease outcome. We propose that the general strategy of integrating molecular analyses of tumors and metastases from relevant GEMMs with cross-species computational analyses of human cancer can be broadly adopted to identify new targets for prevention, detection, and potentially treatment of metastasis progression for other cancer types.

The capability of generating transcriptomic data from lineage-marked tumor and metastatic cells from different organ sites that are free from other stromal and tissue-specific cells, allowed us to elucidate cell-intrinsic gene expression changes that occur during cancer progression. Surprisingly, we found that the predominant gene expression differences that occur during metastasis progression arise in the transition from pre-metastatic to metastatic tumors and are shared among metastases from various organs. This is similar to findings of a recent study of metastasis progression in mouse model of pancreatic cancer.

Notably, our analyses of purified tumor and metastatic cells free of other tissue components reveals an overall similarity of cell-intrinsic metastasis progression across the various metastatic sites, thus supporting the concept that organ-site tropic factors may be contributed by the tumor microenvironment at the metastatic site. We speculate that organ-site specific factors act in collaboration with cell-intrinsic drivers of metastasis progression, such as those identified herein. Furthermore, our findings, which suggest that there are common cell-intrinsic drivers of metastasis progression across organ sites in the mouse model, is consistent with a study of human prostate cancer, which reported the inherent similarity of tumors and metastases from the same patient, and therefore support the feasibility of investigating agents that target metastatic progression in advanced prostate cancer patients.

Interestingly, we find that conserved master regulators of metastasis progression are highly enriched for genes that are predicted to function as regulators of the epigenome, including those that modify DNA and histones, or remodel chromatin architecture. Consistent with our findings, genomic sequencing of prostate tumors has identified several mutations in epigenetic genes particularly in advanced prostate cancer. Furthermore, dysregulation of the epigenome is associated with metastatic progression of human prostate cancer.

In particular, we have demonstrated that the nuclear receptor binding SET Domain Protein 2 (NSD2) is a robust marker of lethal metastatic prostate cancer and a key driver of prostate cancer metastasis, extending previous studies that have reported...
the relevance of NSD2 in prostate cancer. NSD2 was discovered as the overexpressed product of the t(4;14)(p16.3;q32.3) translocation in multiple myeloma, and alternatively named Multiple Myeloma SET domain containing protein (MMSET), and was identified as a target gene on the 4p16 deletion for the Wolf-Hirschhorn Syndrome, and alternatively called Wolf-Hirschhorn Syndrome Candidate 1 (WHSC1). Previous studies have shown that genomic alterations occur in other cancer types in addition to multiple myeloma including pediatric leukemia and laryngeal tumors. In prostate cancer, NSD2 has been shown to be up-
regulated in advanced tumors coordinating with the activation of PI-3 kinase signaling\textsuperscript{49}, and to be a cofactor of androgen receptor\textsuperscript{48}.

Notably, the role of NSD2 in cancer has been shown to be dependent on its activity as a histone methyltransferase for the histone H3 di-methyl K36 (H3K36me2)\textsuperscript{50,61,62}. In the current study, we show that MCTP-39, a putative inhibitor of NSD2\textsuperscript{56}, inhibits prostate tumor growth in vivo. However, several caveats preclude us from drawing the direct conclusion that MCTP-39 is acting to inhibit NSD2 activity in this context, including the potential activity of unknown metabolites and the potential lack of specificity of MCTP-39 given its relatively simple chemical structure\textsuperscript{56}.

Nonetheless, our study demonstrates that NSD2 is a functional driver of prostate cancer metastasis and suggests that it may be target for treatment of advanced prostate cancer. Notably, the activity of NSD2 as a histone methyltransferase has been shown to be coordinately regulated by EZH2\textsuperscript{51}, a major component of the histone methyltransferase polycomb repressive complex 2 (PRC2), which is also dysregulated in prostate cancer. Additionally, in multiple myeloma, NSD2 has been shown to be a regulator of DNA damage response that impacts resistance to chemotherapy\textsuperscript{52}. These previous studies suggest that combination therapy targeting NSD2 together with inhibition of PI-3 Kinase, AR, EZH2, and/or DNA repair mechanisms, all of which are themselves targetable and highly relevant for prostate cancer, may prove to be efficacious for treatment of metastatic prostate cancer. We further proposed that these combination treatments can be evaluated in co-clinical assays using the NPK mouse model described herein.

**Fig. 5** Silencing of NSD2 abrogates tumorigenicity in vitro. Panels a–g show in vitro analyses of NSD2 silencing in a mouse metastatic cell line (NPK cells) and a human advanced prostate cancer cell line (DU145 cells). Cells were infected with control shRNA or two independent shRNAs for mouse or human NSD2, respectively. a, b Validation of NSD2 silencing in NPK and DU145 cells, as indicated, using quantitative real-time PCR (qPCR). c Western blot analyses of NSD2-silenced or control NPK and DU145 cells, as indicated, showing reduced expression of NSD2, which is accompanied by reduction of the H3K36me2 mark, but not the H3K36m1 or the other histone marks shown. The position of a molecular marker is shown; uncropped images are provided in Supplementary Figure 5. d, e Colony formation assays in NPK and DU145 cells, as indicated showing quantification (left) representative images (right). f, g Invasion assays in NPK and DU145 cells, as indicated showing quantification (left) and representative images (right). Experiments were done in three independent biological replicates each in triplicate; p-values were calculated using a two-tailed Student’s t-test. Error bars represent the standard deviation (s.d.) from the mean.

**Fig. 6** Silencing of Nsd2 abrogates metastasis in vivo. Panels a–f show in vivo analyses of Nsd2 silencing in a mouse metastatic cell line (NPK cells). Cells (1 x 10\textsuperscript{6} cells) were engrafted subcutaneously into the flank of nude mice and the mice were monitored for up to 40 days. Studies were done using 2 independent shRNA for Nsd2; representative data for shRNA#1 is shown. a Survival analyses with the endpoint being tumor volume of 1.5 cm\textsuperscript{3} (n = 10/group). The p-value was calculated using a log-rank test. b Analyses of tumor weights (in grams) at the time of killing (total n = 9/group). c Number of lung metastases per mouse (total n = 9/group). d Number of lung metastases per mouse (total n = 9/group). Scale bars represent 100 microns. e Representative immunostaining of shControl and shNsd2 tumors using the indicated antibodies (n = 4/group). Scale bars represent 50 μ. f Western blot analysis showing representative cases from the shControl (lanes 1, 2) and shNsd2 (lanes 3, 4) tumors using the indicated antibodies (total n = 4/group). The position of a molecular marker is shown; uncropped images are provided in Supplementary Figure 5.
Methods

Expression profiling of lineage-marked cells. All experiments using animals were performed according to protocols approved by and following all ethical guidelines required by the Institutional Animal Care and Use Committee (IACUC) at Columbia University Irving Medical Center or the Ethics Committee for Animal Research (CEIC) at Bellvitge Biomedical Research Institute. For molecular profiling analyses, lineage-marked cells from primary tumors and/or metastases were collected from Nes;loxP/loxP, R26R^{YFP} (NP) and Nes;loxP/loxP, R26R^{flox/flox}, Kras^{G12D/2D}; R26R^{flox/flox} (NPK) mice, which have been previously published. Note that inclusion of the lox-stop-lox R26R^{flox/flox} allows for lineage tracing specifically in prostate cells at the time of tumor induction. These NP, and NPK mice have been maintained in our laboratory on a predominantly C57BL/6 background. All studies were done using littermates that were genotyped prior to enrollment; mice were randomly enrolled to treatment or control groups and only male mice were used because of the focus on prostate.

Mice were induced to form tumors at 2–3 months of age by administration of tamoxifen (Sigma-Aldrich, Allentown, PA) using 100 mg/kg once daily for 4 consecutive days and monitored for 1 to 9 months, during which time the NP mice develop prostate adenocarcinoma that progresses to overt metastasis. At the time of killing, prostate tumors from NP or NPK mice, as well as tissues with overt metastases from the NPK mice, as detected by ex-vivo fluorescence, were collected in ice cold phosphate-buffered saline (PBS). Tissues were digested in one part of collagenase/hyaluronidase (Stem Cell Technologies, Cambridge, MA) and nine parts of DMEM-F12 and 10% fetal bovine serum (FBS) at 37 °C for 3 h. Samples were pelleted at 350G in an Eppendorf 5810 R tabletop centrifuge for 5 min at 4 °C, re-suspended in 0.25% trypsin/EDTA (Stem Cell Technologies, Cambridge, MA), and incubated for 1 h on ice. Cells were collected by centrifugation as above, and incubated in a cocktail of pre-warmed dispase (5 mg/ml) plus DNase1 (1 mg/ml) (Stem Cell Technologies, Cambridge, MA) for 10 min at 37 °C; after which cells were filtered through a 40 μm cell strainer, pelleted and re-suspended in 1% PBS/FBS and proceed to the sorter. The YFP-lineage marked cells were purified using a BD FACs Aria II sorter and the YFP + population isolated using the PE/FTC (R-Phycocerythrin/Fluorescein isothiocyanate) channels to gate the YFP + population. Cell pellets were resuspended in 10 μl of Trizol and flash-frozen in liquid nitrogen. RNA was prepared using a MagMAX-96 total RNA isolation kit (Life Technologies). Total RNA was enriched for mRNA using poly-A pull-down; only samples having between 200 ng and 1 μg with an RNA integrity number (RIN) > 8 were used. Library preparation was done using an Illumina TruSeq RNA prep-kit, and the libraries were sequenced using an Illumina HiSeq2500 by multiplexing samples in each lane, which yields targeted number of single-end/paired-end 100 bp reads for each sample, as a fraction of 180 million reads for the whole lane. Raw counts were normalized and the variance was stabilized using DESeq2 package (Bioconductor) in R-studio. R v3.3.0 (The R Foundation for Statistical Computing, ISBN 3-900051-07-0). A complete list of differentially expressed genes is provided in Supplementary Data 1.

Cross species computational analyses. Differential gene expression signatures were defined as a list of genes ranked by their differential expression between two different sets of samples (e.g., local cells vs primary tumors) estimated using a two-sample two-tailed Welch t-test. For cross species analyses, the human gene expression signatures were defined based on published prostate cancer cohorts (Supplementary Table 1) and the mouse gene expression signatures were generated from the RNA sequencing analyses as described above (Supplementary Data 1). For the mouse signatures, a minimum of 5 samples were used for each group as necessary to estimate statistical significance in the two-sample two-tailed Welch t-test and GSEA.

For comparison with human genes, mouse genes were mapped to their corresponding human orthologs based on the homoloGene database (NCBI) so that mouse-human comparisons were done using the “humanized” mouse signatures. For gene set enrichment analysis (GSEA) normalized enrichment score (NES) and p-values were estimated using 1,000 gene permutations. Pathway enrichment analysis was done using GSEA to query the Hallmark Pathways dataset from the MSigDB (i.e., Molecular Signatures Database) and the Broad Institute, where NES and p-value were estimated using 1,000 gene permutations (Supplementary Data 2). Master regulator (MR) analysis was performed using the MASTER Regulator Inference algorithm (MARINA) to query the mouse and human prostate cancer interactomes, respectively, as published previously (Supplementary Data 3).

Master regulator activity analyses. Transcriptional activity of master regulators (MRs) was estimated using expression levels of their transcriptional targets and reflects their enrichment in the signature being queried. In particular, targets of a particular MR are used as a query gene set to estimate their enrichment in the reference signature of interest (e.g., metastatic progression signature). If positive targets are overexpressed (i.e., enriched in the overexpressed tail of the reference signature) and/or negative targets are underexpressed (i.e., enriched in the underexpressed tail of the reference signature), such MR is considered active (i.e., its transcriptional activity is positive). If the converse is the case, the MR is repressed (i.e., its transcriptional activity is negative).

The relationship of MR activity levels for clinical outcome was assessed using four independent datasets (Supplementary Table 1); Sboner et al., which reports
death due to prostate cancer as the clinical end-point; Grasso et al., which reports metastases versus primary tumors as a binary outcome; Grasso et al. and combined SU2C and TCGA cohort were utilized to evaluate the two independent experiments each done using 10 mice/group. Tumors were monitored by caliper measurement twice weekly for approximately 4 weeks and tumor volumes were calculated using the formula \[\text{Volume} = (\text{width})^2 \times \text{length}/2\]. At the time of euthanasia, tumors were harvested and weighed, and the number of lung metastases was documented ex vivo by visualizing their fluorescence using an Olympus SZX16 microscope equipped with epifluorescence capabilities. The total number of lung metastatic nodules (MR filteration model) was counted for each mouse and the p-value was calculated by comparing the control and each shRNA silences using a two-tailed Student’s t-test or the Mann–Whitney U test.

For analyses of NSD2 inhibition, MCTP-39 (3-hydrazinylquinoxaline-2-thiol) was purchased from Enamine Ltd. (Ukraine). For studies performed in mice, the MCTP-39 was further purified by dynamic lipid synthesis on organic synthesis core facility (ORCF) at Memorial Sloan Kettering Cancer Center. The MCTP-39 was subjected to silica gel chromatography, microfiltrated on a 0.2 μm Teflon membrane and lyophilized. The resulting solid was analyzed by UPLC (ultra-high pressure liquid chromatography) on reverse phase C18 silica gel in addition to 1H and 13C NMR (Bruker 600 MHz machine).

In two in vitro, 3 × 10^5 DU145 cells were seeded in triplicate in three independent experiments (aggregate total n = 9) in a 100 mm dish. 24 h following plating, the compound MCTP-39 (5 mM in DMSO) was added to a final concentration of 2.5 μM or 5 μM and incubated for 72 h; DMSO was used as a vehicle control. Cells were collected and lysed for western blot analysis or were seeded for colony formation assays as described above.

For in vivo studies, 5 × 10^5 DU145 cells in 50% matrigel were injected subcutaneously into the flanks of nude mice. When tumors reached 200 mm^3, mice were randomized to the treatment (1% Carboxymethylcellulose, 0.1% Tween80, 5% DMSO) or vehicle (MCTP-39 (10 μg/mg) in vehicle) groups using cage-based randomization, which were administered by oral gavage daily for 3 months. Tumor volume was measured by calipers twice weekly, and estimated by the formula \[\text{Volume} = \text{(width)}^2 \times \text{length}/2\]. Total mice analyzed were: vehicle = 14; MCTP-39 (2015); and 35 (2016) independent experiments. Two-way analysis of variance (ANOVA) was used to calculate the significance (p-value) of the difference between the vehicle and treatment groups.

**Statistical analyses.** The Cox model and Kaplan–Meier analysis were used during the surv and coxph functions from the survcomp package (Bioconductor). Statistical significance was estimated with Wald test and log-rank test, respectively. For Kaplan–Meier survival analysis, k-means clustering was done on the activity levels of the MRs to cluster patients into two groups: one group with increased activity of the candidate MRs and one group with decreased MR activity. To compare the predictive ability of candidate MRs to results at random, we selected a random (equally sized, n = 8) group of MRs and utilized their activity levels for Kaplan–Meier survival analysis. This procedure was repeated 1000 times and log-rank p-values from all iterations were used to build a Null model. The empirical p-value was then estimated as a number of times log-rank p-values for a random group of 8 MRs reached or outperformed our original log-rank value was then estimated as a number of times log-rank p-values from the corresponding authors upon reasonable request.

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**In vivo studies.** Cells were lysed with 1X radioimmunoprecipitation assay (RIPA) buffer (0.1% SDS, 1% deoxycholate sodium salt, 1.0% Triton-X 100, 0.15 mM NaCl, 10 mM Tris–HCl (pH 7.5), 1 mM EDTA supplemented with protease inhibitor cocktail (Roche, 11 836 153 001), 1% phosphatase inhibitor cocktail 3 (Sigma, P8340), and 0.5% PMFS (Sigma). Protein lysates (5 μg per lane) were resolved by SDS-PAGE and transferred onto a PVDF membrane (GE Healthcare, Amersham), then blocked with PBS-T (phosphate-buffered saline and 0.1 Tween-20) containing 5% non-fat dry milk. Incubation with primary antibody was done at 4°C overnight, followed by incubation with secondary antibody for 1 h at room temperature. Detection was performed using the ECL Plus Western Blotting Detection system (Leica Microsystems, IL, USA). Slides were de-paraffinized and heat-mediated antigen retrieval was performed using the Bond Epitope Retrieval 2 solution at pH 9 and incubated with the anti-NSD2 primary antibody (Supplementary Table 2). Nuclear staining intensity was evaluated by a pathologist and scoring was done blinded and defined as negative (or present in <5% of nuclei), weak, moderate, or strong. Immunostaining of mouse prostate tissues and metastases was done as described previously. Briefly, 3 μm paraffin sections were deparaffinized in xylene, followed by antigen retrieval in antigen retrieval solution (Vector Labs, BA-8000, CA). Slides were washed twice in 10% normal goat serum, then incubated with primary antibody overnight at 4°C, followed by incubation with secondary antibodies for 1 h. For immunostaining, the signal was enhanced using the Vectastain ABC system and visualized with NovaRed Substrate Kit (Vector Labs). All antibodies used in this study, as well as antibody dilutions, are described in Supplementary Table 2.

**Functional validation studies.** Mouse cell lines were isolated from lung metastases from Nkx2-1 animals and their genotype and authentication as described previously. Human cell lines were purchased from and authenticated by ATCC (American Type Culture Collection). Cells were grown in RPMI media supplemented with 10% FBS (ThermoFisher, Bridgewater, NJ). Only early passage cells were used for all studies herein. Cells were routinely tested to ensure that they are free of mycoplasma using the MycoAlert Mycoplasma Detection Kit (Invitrogen, Life Technologies, CA). For RNA-mediated silencing, a minimum of two independent shRNA clones were used for each gene using the pLKO.1 lentiviral vector system following manufacturer’s instructions (Sigma-Aldrich, Allentown, PA). The sequences for all mouse and human shRNA used in this study are provided in Supplementary Table 3. As a control, we used a pLKO.1 lentiviral vector with shRNA targeting the Luciferase gene (SHC007, Sigma-Aldrich).

Analysis of RNA expression was done by quantitative real time PCR using the Quantitee SYBR Green PCR kit (Qiagen, Germantown, MD) using mouse or human GADPH as the control. Sequences of all primers are provided in Supplementary Table 5. For in vivo experiments, 3 × 10^5 DU145 cells were seeded in triplicate in three independent experiments (aggregate total n = 9) in 60-mm plates and grown for 10 days in RPMI-1640 media (Gibco, Bridgewater, NJ) supplemented with 10% FBS (BenchMark Gemini Bio-Products, Sacramento, CA). Colonies were visualized by staining with crystal violet and quantified using ImageJ software (National Institute of Health) and ImageJ. For in vivo assays, 2.5 × 10^6 cells were seeded in three independent experiments (aggregate total n = 9) in BD FluoroBlok inserts (BD Biosciences, Billerica, MA) in FBS-free media and supplemented with 10% FBS was used in the lower chamber as a chemotraherrent.

For in vivo assays, 1 × 10^5 cells were injected subcutaneously into the flank of immunodeficient NCR nude mice (male, Taconic, Rensselaer, NY); we performed...
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**Author contributions**
A.A., A.M., and A.G. designed and performed experiments and analyzed data. K.R., J.C., J.A., L.P., and S.F.-M. performed experiments and analyzed data. M.A.R. and M.M.S. supervised the study. A.C., A.A., and C.A.S. designed experiments and supervised the study. C.A.S., A.A., A.G., and A.M. wrote the manuscript; all authors provided comments.

**Additional information**
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