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Akt methylation by SETDB1 promotes Akt kinase activity and oncogenic functions

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Abstract

Aberrant activation of Akt disturbs proliferation, survival and metabolic homeostasis of various human cancers. Thus, it is critical to understand upstream signaling pathways governing Akt activation. Here, we report that Akt undergoes SETDB1-mediated lysine-methylation to promote its activation, which is antagonized by the Jumonji-family demethylase, KDM4B. Notably, compared with wild-type mice, mice harboring non-methylated mutant *Akt1* not only exhibited reduced body size, but also were less prone to carcinogen-induced skin tumors in part due to reduced Akt activation. Mechanistically, Phosphatidylinositol (3,4,5)-trisphosphate (PIP₃)

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FINANCIAL AND NON-FINANCIAL COMPETING INTERESTS

We have no financial and non-financial competing interests.

interaction with Akt facilitates its interaction with SETDB1 for subsequent Akt methylation, which in turn sustains Akt phosphorylation. Pathologically, genetic alterations including *SETDB1* amplification aberrantly promote Akt methylation to facilitate its activation and oncogenic functions. Thus, Akt methylation is an important step synergizing with PI3K signaling to control Akt activation, suggesting that targeting the SETDB1 signaling could be a potential therapeutic strategy for combatting hyperactive Akt-driven cancers.

INTRODUCTION

Epigenetic regulation, such as DNA methylation and histone modifications, plays important roles in governing gene expression patterns during human development and disease progression ^{1, 2}. Inhibitors targeting epigenetic factors have been explored for cancer therapies and have undergone clinical trials, including DNA methyltransferase 1 (DNMT1) inhibitors, histone deacetylases (HDAC) inhibitors, and histone methyltransferase inhibitors ^{3–5}. Among these epigenetic inhibitors, histone methyltransferase (such as EZH2 and DOT1L) inhibitors display impressive efficacy in cancer patients ^{6, 7}. This robust efficacy may also be attributed to regulating methylation of non-histone proteins such as Rb and p53, in addition to regulating histone methylation ^{8–11}. However, it remains largely unknown whether predominant oncogenic signaling pathways that are frequently activated in human cancers, such as PI3K/Akt signaling pathway, are subjected to methylation-dependent regulation. Thus the identification of the major oncogenic proteins governed by methylation is critical to identify new therapeutic targets.

Hyperactivation of PI3K/Akt signaling is a central module of cell proliferation, survival and metabolic homeostasis in human cancers ^{12, 13}. Physiologically, stimulations derived from various types of growth factors tend to activate Akt, which in turn phosphorylates distinct substrates to perform different biological processes ^{13, 14}. Recently, emerging evidence has demonstrated that distinct signals govern Akt kinase activity e.g. TRAF6/Skp2-mediated positive regulation of Akt in an ubiquitination-dependent manner ¹⁵, CDK2/Cyclin A-mediated positive regulation of Akt in a tail phosphorylation-dependent manner ¹⁶, and pVHL-mediated negative regulation of Akt in a hydroxylation-dependent manner ¹⁷. However, the regulation of Akt as a non-histone substrate by histone methyltransferases is not well defined.

Here, we described that Akt1 methylation in its Linker-region is mediated by the histone methyltransferase SETDB1, which is antagonized by the demethylase KDM4B. Biologically, absence of Akt1 methylation attenuates its kinase activity, represses cell growth, glucose uptake, and tumorigenesis. As a result, deficiency in Akt methylation can physiologically decrease mouse body size and can protect mice from developing carcinogeninduced skin tumors. Thus, our data unravel a profound role for the SETDB1/KDM4B axis in manipulating Akt activity and highlight histone methyltransferase SETDB1 as a potential target for combating hyperactive Akt-driven tumors.

RESULTS

Methylation of Akt enhances its kinase activity

To identify important non-histone proteins regulated in a methylation-dependent manner, we used a specific pan-lysine tri-methylation (K-me3) antibody and performed a mass spectrometry (MS) based screening on cell lysates derived from ovarian cancer cells (Fig. 1a and Supplementary Fig. 1a) ¹⁰. Notably, an Akt1-derived peptide was identified, containing methylated modifications at two nearby evolutionarily conserved lysine residues (K140 and K142) in the Akt1 Linker region (Table S1, Fig. 1b). Furthermore, Akt1 methylation was validated using the K-me3 antibody in cells treated with a global histone methylation inhibitor, 3-Deazaneplanocin A (DZneP) (Supplementary Fig. 1b).

Additional MS analyses on ectopically expressed Akt1 further confirmed the tri-methylation of K140 and K142, where mono- and di-methylation events were also identified (Supplementary Fig. 1c). Moreover, K64, another lysine residue in the PH domain of Akt1, was also identified to be mono-methylated in our system. Interestingly, mutation of both K140 and K142 residues led to a dramatic reduction of Akt1 K-me3 signal in cells (Supplementary Fig. 1d), indicating that K140 and K142 might be the major Akt1 K-me3 residues. In addition, consistent with a previous report ¹⁸, deletion of *Smyd3*, a methyltransferase targeting MAP3K2, did not disturb Akt kinase activity in cells, although the interaction of SMYD3 with Akt1 has been previously reported (Supplementary Fig. 1e-g) ¹⁹. Given that both K140 and K142 sites were identified at endogenous levels through our large-scale non-biased MS approach, we chose to mainly focus on understanding the contribution of K140 and K142 tri-methylation to Akt activity and oncogenic functions in the remainder of this study.

Absence of Akt methylation represses its oncogenic functions in cells and increases resistance to carcinogen-induced skin tumors *in vivo*

In order to reveal the potential biological functions of tri-methylation within its Linker region of Akt1, a methylation-deficient variant of Akt1 (K140R and/or K142R) was ectopically expressed in *AKT1* and *AKT2* double knockout DLD1 cells (DLD1- $AKT1/2^{-/-}$). Compared to Akt1-WT, the Akt1-K140/142R mutant, and to a lesser extent, Akt1-K140R or -K142R, exhibited a significantly reduced activation, as shown by a marked reduction in Akt-pT308 and Akt-pS473, as well as its downstream targets, pGSK3 β and pFOXO3a (Fig. 1c and Supplementary Fig. 1h-j). This observation is coupled with a reduction of Akt1 kinase activity observed *in vitro* (Supplementary Fig. 1k). As a consequence, absence of methylation on K140 and K142 led to reduced cell colony formation, anchorage-independent growth, glucose uptake, and lactate production *in vitro* (Fig. 1d, e and Supplementary Fig. 1l, m) and tumor growth *in vivo* (Fig. 1f-h and Supplementary Fig. 1n, o).

To explore the role of Akt methylation under physiological conditions, we generated Akt1-K140/142R double mutations in HEK293 cells (termed $AKT1^{K140/142R}$) using CRISPR/Cas9 (Supplementary Fig. 1p-r) ²⁰. Notably, K140/142R mutation of endogenous Akt1 decreased Akt phosphorylation in response to various stimuli (Fig. 1i and Supplementary

Fig. 1s-u), and also compromised cell growth, colony formation, glucose uptake, and lactate production (Fig. 1j-m). Next, we generated Akt1-K140/142R double mutation knock-in mice (termed $Akt1^{KUKI}$) (Supplementary Fig. 2a-c). Compared with the WT mice, $Akt1^{KUKI}$ mice exhibited a decrease in body size/weight (Fig. 2a, b and Supplementary Fig. 2d-f), organ size/weight (Fig. 2c and Supplementary Fig. 2g) and Akt kinase activity (Fig. 2d, e and Supplementary Fig. 2h), phenocopying $Akt1^{-/-}$ mice ²¹. To pinpoint whether methylation deficient mutants of Akt1 abrogate tumorigenesis *in vivo*, we utilized a two-step chemical carcinogen (DMBA followed by TPA)-induced skin tumor model ²². In this model, we observed that $Akt1^{KUKI}$ mice displayed a lower incidence of skin tumors and reduced papilloma burden compared with WT mice (Fig. 2f-h), which correlated with decreased Akt signaling (Fig. 2i). These results together support the notion that methylation of Akt in its Linker region likely promotes Akt kinase activity and oncogenic functions both *in vitro* and *in vivo*.

SETDB1 methylates and activates Akt

Next, we set out to identify the physiological upstream methyltransferase(s) for Akt. Consistent with previous reports ^{23–25}, we found that there is a relatively strong physical interaction between Akt1 and two tri-methyltransferases, SETDB1 and EZH2 (Fig. 3a and Supplementary Fig. 2i). Interestingly, genetic alterations of both *SETDB1* and *EZH2* tended to mutually exclusive of alterations in PI3K/Akt pathway-related genes including *PTEN* deficiencies, *EGFR*, *PIK3CA* or *AKT1* amplification/mutations in human melanoma and breast cancer (Supplementary Fig. 2j) ²⁶. Furthermore, SETDB1, but not other methyltransferases including EZH2, could enhance Akt tri-methylation in cells (Supplementary Fig. 2i). This result indicates that unlike EZH2 as a downstream substrate of Akt ²⁵, SETDB1 may be an upstream regulator of Akt methylation (Supplementary Fig. 2k).

Notably, SETDB1 specifically interacted with the PH domain of Akt1, and to a lesser extent, Akt2 and Akt3 (Fig. 3b, Supplementary Fig. 2l, m). Interestingly, mutation of Akt1 methylated residues (K140/142R) could markedly decrease the interaction of Akt1 with SETDB1 (Supplementary Fig. 3a). Importantly, the Tudor domain of SETDB1 was identified to bind Akt1, which was enhanced by enforcing expression of WT, but not the SETDB1-H1224K catalytic inactive mutant (Fig. 3c, Supplementary Fig. 3b). These results indicate that SETDB1 could be a potential reader of the Akt1-K140/142 methylation events through its Tudor domain, and by enhancing its interaction with Akt1, SETDB1 could promote Akt1 methylation on additional residues (such as K64). In addition, SETDB1 promoted Akt1 tri-methylation in an enzymatic activity-dependent manner in cells (Fig. 3d), and mutation of either K140 or/and K142 residue could decrease SETDB1-mediated Akt trimethylation assays also indicate SETDB1 could directly methylate Akt1 at K140 and K142 residues (Fig. 3e). Hence, these results coherently support the notion that SETDB1 serves as a putative Akt1 methyltransferase.

SETDB1 promotes cell growth and glycolysis through methylation of Akt

As histone H3 methyltransferase capable of catalyzing tri-methylation on the lysine 9 of histone H3 (H3K9me3) ²⁷, SETDB1 is amplified in various cancers ^{28–31}. Notably, we

identified that, in addition of being in the nucleus and modifying histone marks, SETDB1 was also observed in the cytoplasm of cells (Supplementary Fig. 3d). To further elucidate the physiological functions attributed to methylated-Akt1, we generated specific antibodies against tri-methylation of Akt1 at K140 (termed K140-me3), which was validated by dot blot and immunoprecipitation assays (Supplementary Fig. 3e, f), and recognized SETDB1-mediated tri-methylation of Akt1 at K140 in cells (Fig. 3f, Supplementary Fig. 3g) and *in vitro* (Fig. 3g, Supplementary Fig. 3h). Additionally, genetic depletion of *SETDB1* markedly reduced Akt1-K140-me3 in cells (Fig. 3h-j and Supplementary Fig. 3i). Compared with Akt1-WT, a cancer patient-derived Akt1-E17K mutant ³² displayed an enhanced interaction with SETDB1, coupled with an increase of its K140 methylation level (Fig. 3k and Supplementary Fig. 3j, k). These findings suggest that Akt1 is a *bona fide* methylation substrate of SETDB1.

Importantly, genetic deletion of *Setdb1* in cells diminished Akt activation (Fig. 3i, j, 1 and Supplementary Fig. 31-o). Furthermore, depletion of endogenous *SETDB1* by shRNA also reduced Akt-pT308 in different cancer cell lines (Supplementary Fig. 4a, b), leading to suppression of colony formation, anchorage-independent growth, and glucose uptake in cells (Supplementary Fig. 4c-f). Conversely, ectopic expression of SETDB1-WT, but not SETDB1-H1224K ²⁷, induced Akt-pT308 in *Setdb1*-deficient MEFs (Fig. 3l). Collectively, these results indicate that the activation of Akt by SETDB1 appears to be methylation-dependent. Of note, consistent with a previous report ²⁸, IHC staining revealed that compared to nevus tissues, SETDB1 is highly expressed in melanoma, especially in metastatic malignancies (Fig. 3m, n), and positively correlates with Akt1 K140-me3 and Akt-pS473 IHC signals (Fig. 3o-q).

SETDB1 is suggested to accelerate melanoma onset in cooperation with BRAF^{V600E 28}. Given that coordinated actions of BRAF^{V600E} with activated Akt are required to promote melanomagenesis ^{33, 34}, we employed a well-established human immortalized melanocyte (HIM) model ³⁵ to examine whether SETDB1 modulates the activation of Akt. To this end, we found that concomitant activation of Akt and ERK signaling by SETDB1 and BRAF^{V600E} led to a marked enhancement of anchorage-independent cell growth, coupled with an increased Akt phosphorylation (Fig. 4a-c). Furthermore, deletion of endogenous *AKT1* in HEK293 cells compromised the oncogenic capability of SETDB1 to promote cell colony formation compared with parental cells (Fig. 4d-f).

More importantly, depletion of *SETDB1* resulted in decreased Akt phosphorylation and cellular malignant phenotypes in Akt1-WT expressing DLD1-*AKT1/2*^{-/-} cells, but did not in Akt1-K140/142R expressing cells that already displayed relatively lower oncogenic capacity than WT cells (Fig. 4g-j). These results indicate that, Akt is likely one of the major downstream pathways through which SETDB1 exerts its oncogenic role. Consistently, re-introduction of constitutively active Akt1 (myr-Akt1) partially restored the colony formation capability of *SETDB1*-depleted A375 cells (Supplementary Fig. 4g, h). Furthermore, SETDB1 was capable of accelerating cell growth and colony formation in *AKT1*^{WT}, but not in the methylation-deficient *AKT1*^{K140/142R} expressing HEK293 cells (Fig. 4k-n). Hence, our results suggest that SETDB1 may exert its oncogenic roles largely through Akt activation in a methylation-dependent manner.

SETDB1-mediated Akt methylation crosstalks with PI3K-mediated Akt phosphorylation

It is well established that Akt activation requires PI3K-dependent generation of PIP₃, which binds with the PH-motif of Akt at the plasma membrane ^{36–38}. We explored the temporal relationship between methylation-dependent and PI3K-dependent activation of Akt in cells. We showed that the interaction between Akt1 and SETDB1, and subsequent methylation of Akt, temporally coincided with Akt-pT308 following insulin or IGF stimulation (Fig. 5a and Supplementary Fig. 4i-k). Interestingly, PI3K inhibitors, but not Akt1 mutated variants (T308A and/or S473A), diminished the interaction of Akt1 with SETDB1, and decreased Akt1-K140 methylation in cells (Fig. 5b and Supplementary Fig. 4l, m), indicating that the PI3K pathway may function upstream of SETDB1-mediated Akt methylation to govern Akt activation. In further support of this notion, the non-PIP₃ binding mutation of Akt1-R25C ³⁹, largely impaired the interaction between Akt and SETDB1, leading to a marked decrease in Akt1-K140 methylation (Fig. 5c, d).

To further pinpoint the potential role of PIP₃ in mediating the interaction between SETDB1 with Akt1-PH motif, we performed *in vitro* binding assays. Results suggest that the interaction of SETDB1 with Akt1-WT, but not Akt1-R25C, could be enhanced by PIP₃ *in vitro* (Fig. 5e and Supplementary Fig. 5a-e), indicating that PIP₃ binding could induce a conformation change of the PH motif, which facilitates Akt1 binding to SETDB1. Importantly, *PTEN* deficiency dramatically increased Akt1-K140 methylation coupled with increased Akt-pT308 (Supplementary Fig. 5f). Together, these data suggest that SETDB1 mediates Akt methylation likely in a PI3K catalytic activity-dependent manner.

Consistent with the finding that methylation of Akt1 could promote Akt-pT308, we observed that WT, but not the catalytically inactive SETDB1-H1224K, could induce the interaction of Akt1 with PDK1 in cells (Fig. 5f and Supplementary Fig. 5g). Moreover, the methylation-deficient Akt1 mutant (K140/142R) or depletion of *SETDB1* displayed an attenuated interaction with PDK1 compared with Akt1-WT (Fig. 5g, h and Supplementary Fig. 5h). In further support of the role for Akt methylation in controlling its activation, reducing Akt methylation by deleting *Setdb1* or introducing the Akt1-K140/142R mutation diminished the interaction of Akt1 with PIP₃ (Fig. 5i, j), and subsequently led to a marked decrease in association of Akt1 with the cell membrane (Fig. 5k, l and Supplementary Fig. 5i-m). These results indicate that methylated Akt1 may have a greater propensity to bind PIP₃ and translocate to the cellular membrane to achieve full activation. Thus, there is likely an intrinsic interplay between PI3K-mediated and SETDB1-mediated pathways to tightly control the spatial and temporal activation of Akt (Supplementary Fig. 5n).

Given that TRAF6-mediated Akt ubiquitination has a critical role in regulating Akt membrane translocation ^{15, 40, 41}, we also found that WT, but not SETDB1-H1224K mutant, increased the interaction between Akt1 and TRAF6 (Supplementary Fig. 50), resulting in an increase of Akt ubiquitination (Supplementary Fig. 5p). Furthermore, in comparison with Akt1-WT, Akt1-K140/142R displayed a reduced interaction with TRAF6 (Supplementary Fig. 5q), resulting in decreased Akt ubiquitination (Supplementary Fig. 5r). These findings suggest that SETDB1-induced methylation likely mediates the interaction of Akt1 with TRAF6, and subsequent Akt ubiquitination may promote membrane translocation of methylated Akt (Supplementary Fig. 5n).

KDM4B demethylates and attenuates Akt kinase activity

Importantly, the Jumonji 2 (JMJD2, also termed KDM4) family proteins have been shown to function as specific erasers of SETDB1-mediated H3K9me3 ⁴², therefore, we assessed their ability to demethylate Akt1-K140me3. Notably, among all members of KDM4 family, KDM4A and KDM4B, and to a lesser extent KDM4C, were observed to interact with Akt1 (Fig. 6a and Supplementary Fig. 6a). Furthermore, *in vitro* demethylation assays ⁴² demonstrated that KDM4B-WT, but not the catalytically inactive KDM4B-H189A or other KDM4 members could efficiently erase the tri-methylation of Akt1 at K140 *in vitro* (Fig. 6b and Supplementary Fig. 6b-e) or in cells (Supplementary Fig. 6f). Moreover, the interaction between KDM4B and Akt1 was readily induced by enforcing expression of SETDB1-WT, but not its catalytic mutant (Fig. 6c). We further showed that Akt interacts with KDM4B through its Tudor domain (Supplementary Fig. 6g-j) in a methylation-dependent manner (Fig. 6d, e and Supplementary Fig. 6k, l). Hence, these data indicate that KDM4B might be a *bona fide* demethylase of Akt1.

KDM4B-depletion concomitantly increased Akt-pT308 and K140-me3 in multiple cell lines with Akt1-WT, but not those with *AKT1^{K140/142R}* cells (Fig. 6f-h and Supplementary Fig. 6m-p). Strikingly, depletion of *KDM4A* displayed a reduction in Akt phosphorylation level (Supplementary Fig. 6q, r). In addition, depletion of *KDM4B*, but not *KDM4A*, largely enhanced Akt1 membrane translocation (Fig. 6i and Supplementary Fig. 6s). Furthermore, the interaction between Akt1 and KDM4B primarily occurred in the later phase of insulin stimulation (Supplementary Fig. 6t). This interaction was abolished by PI3K inhibitors (Fig. 6j). Consistent with the finding that KDM4B interacts with methylated-Akt, the patient-derived mutation (Akt1-E17K) increased the interaction between Akt1 and KDM4B, while the non-PIP₃ binding mutation (Akt1-R25C) had the opposite effect (Fig. 6k). Clinically, KDM4B was expressed at relatively high levels in benign nevus compared with malignant melanoma (Fig. 6l, m). This observation correlated with decreased Akt-pS473 and K140-me3 IHC signals (Fig. 6l, n, o). Taken together, these data suggest that KDM4B might function as a negative regulator, antagonizing SETDB1-mediated methylation and activation of Akt.

SETDB1 is a potential therapeutic target of Akt-driven cancers

Given the critical role for the PI3K/Akt pathway in facilitating tumorigenesis, targeting hyperactive PI3K/Akt signaling by PI3K and/or Akt inhibitors have been pursued as promising anti-cancer therapies. However, relatively high-dose of these inhibitors induced cellular toxicity restricting their potential use as a treatment option in cancer patients ^{43, 44}. Thus, our finding that SETDB1 synergizes with PI3K to activate Akt kinase (Supplementary Fig. 7a) indicates that SETDB1 specific inhibitors could benefit cancer patients by targeting both H3K9-mediated epigenetic and PI3K/Akt oncogenic pathways. In support of this hypothesis, we showed that depletion of *SETDB1* significantly decrease A375 tumor growth *in vivo* (Fig. 7a-d), coupled with a reduction in H3K9me3 and Akt phosphorylation (Fig. 7e).

Consistent with a previous report ⁴⁵, we found that Mithramycin A, an anti-neoplastic antibiotic, could markedly decrease SETDB1 expression and H3K9me3 in different cells.

Mithramycin A subsequently led to reduced Akt methylation and phosphorylation (Fig. 7f and Supplementary Fig. 7b) and decreased the interaction of Akt1 and PDK1 (Fig. 7g). Biologically, Mithramycin A attenuated the phosphorylation and colony formation of cells expressing Akt1-WT, but not in *AKT1/2*-depleted DLD1 cells or *AKT1^{K140/142R}*-edited HEK293 cells (Fig. 7h-j and Supplementary Fig. 7c-e). To explore the potential anti-tumor roles of Mithramycin A *in vivo*, we treated nude mice bearing A375 or DLD1 xenografted tumors, and showed that Mithramycin A could significantly decrease tumor growth compared with the vehicle treatment (Fig. 7k-n, and Supplementary Fig. 7f-i). Conceivably, the reduction of SETDB1 expression mediated by Mithramycin A decreased H3K9me3 and Akt phosphorylation (Supplementary Fig. 7j). Taken together, these results suggest that SETDB1 inhibition could possibly benefit cancer patients by repressing Akt oncogenic signaling in addition to its well characterized role in reprogramming the epigenome (Supplementary Fig. 7k).

DISCUSSION

In order to identify novel non-histone methylated proteins involved in oncogenic signaling pathways, we performed a mass spectrometry based high-throughput screen and detected numerous proteins modified by tri-methylation. We found that Akt1 could be methylated in its Linker region, which was enhanced during physiological conditions such as growth factor (Insulin or IGF) stimuli or pathological conditions such as Akt1-E17K mutation or *PTEN* deficiency. Absence of Akt methylation represses its kinase activity, and markedly decreases cell growth, glucose uptake, and tumorigenesis. More strikingly, methylation-deficient *Akt1* knock-in mice exhibit decreased body weight/size, which phenocopies *Akt1* knockout mice ⁴⁶. Importantly, we also observed that methylation-deficient *Akt1* knock-in could attenuate tumorigenesis in a carcinogen-induced skin tumor mouse model. Further investigations will be needed to determine whether knock-in methylation-deficient *Akt1* in genetic alteration models, such as *KRas^{G12D}* mutation and *p53* depletion-mediated lung cancer models (*KP* mice) ⁴⁷, could efficiently compromise the lung tumorigenesis through the decrease of Akt activity.

Among the known upstream signals, Akt1-T308 phosphorylation mediated by PDK1 is pivotal for Akt activation in a PIP₃ dependent fashion ⁴⁸. Here, we observed that SETDB1mediated Akt methylation crosstalks with PI3K-mediated Akt phosphorylation. We suspect that PIP₃ binding to the PH domain of Akt could "unlock" the Akt auto-inhibition confirmation. The conformation change could be necessary for SETDB1 to interact with the Akt PH domain, and to methylate Akt in its exposed Linker region, in turn enhancing the interaction of Akt with its upstream kinase PDK1 (Supplementary Fig. 5n). We also observed that SETDB1 could enhance the interaction of Akt with TRAF6 to facilitate Akt ubiquitination, subsequent PIP₃ binding and membrane translocation (Supplementary Fig. 5n). Our model suggests a loop of Akt activation triggered by PIP₃ accumulation or Akt1-E17K mutation, which could be antagonized by the demethylase KDM4B (Supplementary Fig. 5n). Additionally, the SETDB1-Tudor domain interacts with Akt1 mainly in the context of Akt1 Linker region being methylated. Thus, we speculate that methylation of the Linker region could be the primary modification that enhances SETDB1 interaction with Akt1, and sequentially promotes methylation of Akt on other lysine residues (such as K64 in Akt1 PH

domain). Consistent with our findings, another group also revealed that SETDB1-mediated Akt1 tri-methylation at K64 plays important role for Akt interacting with and being ubiquitinated by TRAF6 to facilitate Akt membrane translocation and kinase activation ⁴⁹. Therefore, the complicated connection and crosstalk of SETDB1-mediate different lysine methylation on Akt including K140/K142 in linker domain and K64 in PH domain warrants further investigation.

Finally, our study reveals that SETDB1 accelerates tumorigenesis largely through Akt activation in a methylation-dependent manner. As such, the function of SETDB1 is not restricted to the nucleus to repress gene transcription, but SETDB1 also methylates non-histone proteins in the cytoplasm such as Akt to activate its oncogenic functions.

METHODS

Cell culture, transfection and cell fractionations.

Cell line sources are detailed in the Reporting Summary. All cell lines were free of mycoplasma contamination (tested by the vendor). HEK293, HEK293T, DLD1, OVCAR5 and A375 cells were cultured in DMEM medium supplemented with 10% FBS, 100 units of penicillin and 100 µg/ml streptomycin. HCT116 PTEN^{+/+} and PTEN^{-/-} cells (as gifts from Dr. Todd Waldman in School of Medicine, Georgetown University). Setdb1^{f/f}-ER-Cre mouse embryonic fibroblasts (MEFs) were obtained as gifts from Drs. Yoich Shinakai and Matthew C. Lorincz ⁵⁰. DLD1-AKT1^{-/-}AKT2^{-/-} (termed AKT1/2^{-/-}) and counterpart cells were kindly provided by Dr. Bert Vogelstein (Johns Hopkins University School of Medicine), and these cells were also maintained in DMEM medium supplemented with 10% FBS. The human immortalized melanocyte HIM was cultured in keratinocyte serum-free medium (K-SFM, Gibco Life Technologies), supplemented with epidermal growth factor (EGF; 5 ng/ml) and bovine pituitary extract (40 pg/ml). Cell transfection was performed using Lipofectamine and Plus reagents, as described previously ¹⁷. Packaging of lentiviral shRNA or cDNA expressing viruses and retroviral cDNA expressing viruses, as well as subsequent infection of various cell lines were performed according to the protocols described previously ¹⁷. Following viral infection, cells were maintained in the presence of hygromycin (200 µg/ml) or puromycin (1 µg/ml), depending on the viral vectors used to infect cells.

To prepare primary MEFs, *Jmjd2b^{flox/flox}* mice were crossed and fibroblasts were established from E13.5 embryos according to standard procedures. In brief, the embryos were dissociated and then trypsinized to produce single-cell suspensions. The cells were maintained in DMEM supplemented with 10% fetal calf serum (FCS), L-glutamine, and antibiotics. The cells were infected with phage-Cre lentiviruses for 2 days to deletion *Jmjd2b. Kras;p53* and *Kras;p53;sMYD3–/–* cells derived from mouse lung and pancreas were obtained from Pawel K. Mazur at Stanford University School of Medicine ¹⁸.

Cell fractionations were performed with Cell Fractionation Kit (CST9038). Kinase inhibitors Wortmannin (Selleck S2758), BKM120 (Selleck S2247) and LY294002 (Selleck S1105) were used at the indicated doses. Mithramycin A (Sigma M6891), growth factors including EGF (Sigma E9644), insulin (Invitrogen 41400–045) and IGF (Sigma SRP3069), were used

at the indicated doses. PIP₃ beads (P-B00Ss) and free PIP₃ (P-3908) were purchased from Echelon Biosciences.

Plasmid construction.

Constructs of pcDNA3-HA-Akt1, pcDNA3-HA-Akt2, pcDNA3-HA-Akt3 and pcDNA3-HA-myr-Akt1, pcDNA3-HA-Akt1-E17K, pET-GSK3β, pCMV-Flag-PDK1 were previously described ¹⁷. pCMV-flag-EZH2, pCMV-flag-SET7, pCMV-flag-SET8 and pCMV-flag-MMSET were bought from Addgene. pCDNA-flag-SETDB1 and pCDNA-flag-SETDB1-H1224K were as gift from Dr. Gerd P. Pfeifer ⁵¹. pCMV-Flag-KDM4A, 4B, 4C and 4D were obtained from Dr. Yang Shi (Harvard Medical School). pCMV-GST-PDK1, pCMV-GST-SETDB1, pCMV-GST-Akt1, pCMV-GST-Akt1-PH (aa1–108), pCMV-GST-Akt1-Linker (aa109–150), pCMV-GST-Akt1-KD (aa151–408), pCMV-GST-Akt1-HM (aa409–481), pCMV-GST-SETDB1-Tudor (aa257–318), pCMV-GST-KDM4A, pCMV-GST-KDM4C, pCMV-GST-KDM4D, pCMV-GST-KDM4B, pCMV-GST-KDM4B-jmj (aa1–180), pCMV-GST-KDM4B-Linker (aa181–425) and pCMV-GST-KDM4B-PH-Tudor (aa181–425) were cloned into mammalian expression GST-fusion vectors. Details of plasmid constructions are available upon request.

Various Akt1 and KDM4B mutants were generated using the QuikChange XL Site-Directed Mutagenesis Kit (Stratagene) according to the manufacturer's instruction. All mutants were generated using mutagenesis PCR and the sequences were verified by DNA sequencing. sh-Akt1-resistant mutants were generated with the specific primers as previously described ¹⁶.

Antibodies.

All antibodies were used at a 1:1000 dilution in TBST buffer with 5% non-fat milk for western blot. Anti-phospho-Ser473-Akt antibody (4060), anti-phospho-Thr308-Akt antibody (2965), anti-Akt1 antibody (2938), anti-Akt total antibody (4691), anti-PDK1 antibody (13037), anti-SMYD3 antibody (12859), anti-H3K9me3 antibody (13969), anti-Pan-Kme3 antibody (14680), anti-AIF antibody (5318), anti-Histone H3 antibody (4499), anti-H3K4me2 antibody (9725), anti-phospho-Ser9-GSK3β antibody (5558), anti-GSK3β antibody (12456), anti-phospho-FOXO1 (Thr24)/FOXO3A (Thr32) antibody (9464), anti-FOXO3A antibody (2497), anti-GST antibody (2625), anti-pS6K1 (Thr389) antibody (9205), anti-S6K1 antibody (2708) and anti-pS240/244-S6 antibody (5364) were obtained from Cell Signaling Technology. Anti-SETDB1 antibody (11231) was obtained from Proteintech. Anti-KDM4A (A300-860A) and anti-KDM4B (A301-478A) antibodies were bought from Bethyl. Anti-Akt1 agarose beads (sc-5298) and polyclonal anti-HA antibody (sc-805) were obtained from Santa Cruz. Polyclonal anti-Flag antibody (F-2425), monoclonal anti-Flag antibody (F-3165, clone M2), anti-Tubulin antibody (T-5168), anti-Flag agarose beads (A-2220), anti-HA agarose beads (A-2095), peroxidase-conjugated antimouse secondary antibody (A-4416) and peroxidase-conjugated anti-rabbit secondary antibody (A-4914) were obtained from Sigma. Monoclonal anti-HA antibody (MMS-101P) was obtained from Covance.

The polyclonal Akt1-K140-me3 antibodies generated by Cell Signaling Technology (CTS) were derived from rabbit, with each hydroxylation residue produced four clones. The

antigen sequence used for immunization was Akt1 aa130–152 (GAEEMEVSLAKPKHRVTMNEFEY). K stands for tri-methylation residue in this synthetic peptide. The antibodies were affinity purified using the antigen peptide column, but they were not counter selected on unmodified antigen.

shRNAs, sgRNAs and CRISPR/CAS9-mediated knock-in and knockout assay.

shRNAs against *SETDB1* as gift obtained from Dr. H. Phillip Koeffler ⁵², and shRNAs against *KDM4A* as gift obtained from Dr. Cun-yu Wang ⁵³. shRNAs against *KDM4B* were reported previously ⁵⁴.

The lentiviruses for CRISPR/Cas9-based editing of *AKT1* were generated by cloning the annealed short guide RNAs (sgRNAs) into BsmBI-digested lentiCRISPR V2 vector, which encodes both Cas9 and an sgRNA of interest, as previously described ²⁰. The sgRNAs were designed by CRISPR Design tool (crispr.mit.edu) as listed below:

sghAKT1#1 Forward: 5'-CACCGTGAAGAGATGGAGGTGTCCC-3';

Reverse: 5'-AAACGGGACACCTCCATCTCTCAG-3';

sghAKT1#2 Forward: 5'-CACCGTGGGGACAGGCCTCACCACG-3';

Reverse: 5'-AAACCGTGGTGAGGCCTGTCCCCAC-3';

For knock-in experiments, the lentiCRISPR V2 constructs containing sgRNA (as mentioned above) were co-transfected with template DNA into HEK293 cells. After selected with puromycin for two days, resulting cells were seeded into 96 well plates with 1 cell/well. After culture around 2 weeks, the single clone was expanded to 12 well-plate, at the same time, the genomic DNA of each clone was harvested by using DNA Extract Solution (Epicentre), and PCR was performed with the primers covered the guild RNA sequence (around 600bp). The PCR products were digested with SacII, which can digest the knock-in (KI) genome but not the wild-type genome *AKT1*. Following that, the genomic DNAs of digestible clone were subjected to sequence to confirm the knock-in sequence. The examined clones were expanded and prepared for further studies.

AKT-K140/142R guild DNA sequence:

GGAGATGGACTTCCGGTCGGGCTCACCCAGTGACAACTCAGGGGCTGAAGAGAT GGAGGTGTCCCTCGCC<u>AGGCCGCGGC</u>ATCGCGTGGTGAGGCCTGTCCCCACTTC TGCCTGTGCCTGGGGCTGCCTTGGACTGTGGAGGGCTGGGTG (the bold indicated mutant nucleotides to restitute K140/142 to R140/142, and at the same time to generate a new Sac II digest site. The shadow nucleotides indicated the destruction of sgRNArecognized site.).

Generation and validation of Akt1^{K140/142R} knock-in mice.

sgRNAs were synthesized by using *in vitro* transcription (IVT) method (PNA Bio Company) with the sequence: TGAAGAGATGGAGGTGTCCC and TGTGGTGAGTCTGGGCTCTG. Cas9 enzyme protein was obtained from PNA Bio. Mutated mouse genomic single strand

template DNA (ssODN) was synthesized from IDT with the sequence: GGACTCAAGAGGCAGGAAGAAGAAGAAGACGATGGACTTCCGATCAGGCTCACCCAGT GACAACTCAGGGGCTGAAGAGATGGAGGTGTCCCTCGCC<u>AGGCCGCGG</u>CATCG TGTGGTGAGTCTGGGCTCTGCTTCTGCTGGGGGCTGCCAGGGGTTGTGGAGGGAC CTGGTAGGTCCTGGTATTCCTGCCTGGCTTCCTGTATGG (the bold indicated mutant nucleotides to restitute K140/142 to R140/142, and at the same time to generate a new Sac II digest site. The shadow nucleotides indicated the destruction of sgRNA-recognized site.). The mixture of Cas9 protein (200 ng/µl), sgRNA(s) (100 ng/µl) and ssODN DNA (10–70 ng/ µl) in 50 µl was utilized for generating knock-in mice by BIDMC transgenic facility.

The genomic DNAs derived from the tail of edited mice were extracted with QuickExtract DNA extraction solution (Epicentre, QE09050), and the PCR was performed with primers: mAkt1-geno-5' 5-AGGCCCAGGATCTGAGTGG and mAkt1-geno-3' 5-TCAGCGGGCATCTTCATATTAC. The PCR products were digested with Sac II for 1 hr or subjected for Sanger sequencing with the primer mAkt1-seq-3' 5-GACAAGCACTCTGCCAACTG. The validated heterogeneous knock-in mice (termed founder mice, P1) were selected and crossed to each other, and got the WT, heterogeneous and homogenous Akt1K140/142R knock-in mice (termed P2). Furthermore, we crossed homogenous knock-in mice with wild-type C57BL6 mice (Jackson Lab), and got the heterogeneous Akt1K140/142R knock-in mice (termed P3). Then we crossed these heterogeneous knock-in mice to each other and got the WT, heterogeneous and homogenous knock-in mice (termed P4). In this time point, we monitored the mice body size, organ size as well as Akt signaling among WT, heterogeneous and homogeneous Akt1K140/142R knockin mice. To further diminish the nonspecific effect of genomic editing, we would cross back the homogenous knock-in mice (P4) with wild-type C57BL6 mice again to generate heterogeneous knock-in mice (P5) for further crossing and phenotype monitoring.

All experimental procedures were approved by the Institutional Animal Care & Use Committee (IACUC, RN150D) at Beth Israel Deaconess Medical Center with protocol #043–2015. The research projects that are approved by the IACUC are operated according the applicable Institutional regulations. The Institute is committed to the highest ethical standards of care for animals used for the purpose of continued progress in the field of human cancer research.

IAP-LC-MS/MS screening.

The experiments were performed as previously reported ¹⁰. Briefly, OVCAR5 cell lysates were proteolytically digested, then purified over a Sep-Pak C18 cartridge. After dissolving in IAP buffer, the recovered peptides were purified with tri-methylation-specific antibodies. Finally, the enriched peptides were subjected to LC-MS/MS to identify the tri-methylation sites in corresponding proteins.

Two-step chemical carcinogen-induced skin tumors.

The method of two-step chemical carcinogen inducing skin tumors were described previously 22,56 . In brief, the shaved naked back of 4–6 week WT or $Akt1^{K140/142R}$ knock-in mice were painted with 150 µg DMBA (Sigma D3254) resolved in 200 µl acetone. After

recovered two weeks, the mice were treated with 20 μ g TPA (LC laboratories, P1680) in 200 μ l acetone twice/week for 15 weeks. The neoplasms were monitored and then dissected after mice euthanized, and subjected to HE and IHC staining.

Immunoblot (IB) and immunoprecipitation (IP) analyses.

Cells were lysed in EBC buffer (50 mM Tris pH 7.5, 120 mM NaCl, 0.5% NP-40) supplemented with protease inhibitors (Complete Mini, Roche) and phosphatase inhibitors (phosphatase inhibitor cocktail set I and II, Calbiochem). The protein concentrations of whole cell lysates were measured by the Beckman Coulter DU-800 spectrophotometer using the Bio-Rad protein assay reagent. Equal amounts of whole cell lysates were resolved by SDS-PAGE and immunoblotted with indicated antibodies. For immunoprecipitation analysis, 1000 μ g lysates were incubated with the indicated antibody (1–2 μ g) for 3–4 hrs at 4°C followed by 1 hr incubation with Protein A/G sepharose beads (GE Healthcare). The recovered immuno-complexes were washed five times with NETN buffer (20 mM Tris, pH 8.0, 150 mM NaCl, 1 mM EDTA and 0.5% NP-40) before being resolved by SDS-PAGE and immunoblotted with indicated antibodies. Quantification of the immunoblot band intensity was performed with Image J software.

Melanoma and Nevus TMA and IHC staining.

Tissue microarray (ME1004f) containing 18 nevi, 33 metastatic melanomas and 49 malignant melanomas were obtained from Biomax.

Immunohistochemistry was performed on four micron-thick, FFPE sections. FFPE sections were deparaffinized using xylene and rehydrated in graded ethanol. Sections were heated with a pressure cooker to 125°C for 30 seconds and 90°C for 10 seconds in citrate buffer (pH 6.0) for antigen retrieval. All sections were incubated with peroxidase (Dako #S2003) and protein blocking reagents (Dako #X0909) for 5 minutes each. Sections were then incubated with anti-pS473-Akt (CST #4060, 1:50), anti-K140-me3-Akt (1:200), anti-pS420/424-S6 (CST #5364, 1:1500) and anti-SETDB1 (CST #93212, 1:100) antibody diluted in Dako diluent with background reducing components (Dako #S3022) for 1 hr at room temperature. Following primary antibody incubation, sections were incubated with monoclonal mouse anti-rabbit immunoglobulins (Dako #M0737) for 30 minutes at room temperature. Afterwards, sections were incubated with Envision+ System-HRP Labeled Polymer Anti-Rabbit (Dako #K4003) for 30 minutes. All sections were developed using the DAB chromogen kit (Dako #K3468) and lightly counterstained with hematoxylin. The score of the IHC signals was judged by two independent pathologists blindly.

Purification of GST- and His-tagged proteins from bacteria.

Recombinant GST-conjugated KDM4B catalytic domain and His-conjugated GSK3β were generated by transforming the BL21 (DE3) *E. coli* strain with pGEX-KDM4B-Cata and pET-28a-GSK3β, respectively. Starter cultures (5 ml) grown overnight at 37°C were inoculated (1%) into larger volumes (500 ml). Cultures were grown at 37°C until an O.D. of 0.8, following which protein expression was induced for 12–16 hrs using 0.1 mM IPTG at 16°C with vigorous shaking. Recombinant proteins were purified from harvested pellets. Pellets were re-suspended in 5 ml EBC buffer and sonicated (5 cycles of 10 sec each at 50%

output). Insoluble proteins and cell debris were discarded following centrifugation in a tabletop centrifuge (13000 rpm/4°C/15 min). Each 1 ml supernatant was incubated with 50 μ l of 50% Glutathione-sepharose slurry (Pierce) or Ni-beads (Qiagen) for 3 hrs at 4°C. The Glutathione beads were washed 3 times with PBS buffer (1 ml per wash) and stored at 4°C in PBS buffer containing 10% glycerol or eluted by elution buffer. The Ni-beads were washed 3 times with 50 mM Tris-Cl (pH 8.0) containing 20 mM Imidazole, and eluted by Tris-buffers containing 100 mM Imidazole. Recovery and yield of the desired proteins (or complexes) was confirmed by analyzing 10 μ l of beads by Coomassie blue staining, and quantified against BSA standards.

Purification of His-Akt1 protein from insect S9 cells.

Akt1 cDNA was cloned into an insect cell expression vector, pFastBac-HT A (Invitrogen), in frame with a Histidine (His) tag at the N-terminus. Constructs were transformed into DH10Bac bacteria (Invitrogen) to generate bacmid DNAs. Akt baculoviruses were produced in Sf9 insect cells according to the manufacturer's specification. His-tagged proteins were purified on HisPur Cobalt resin (Thermo Scientific) and eluted by Imidazole (Sigma). In brief, Sf9 insect cells expressing His-tagged Akt proteins were collected and washed once with ice-cold Phosphate Buffered Saline (PBS). Cell pellet was then re-suspended for 30 minutes at 4°C in lysis buffer containing 10 mM HEPES pH 7.6, 3 mM MgCl₂, 300 mM KCl, 5% glycerol, 0.5% NP-40, 10 mM Imidazole, 1 mM Na₂VO₄, 20 mM NaF, 1 mM sodium pyrophosphate, 25 mM β-glycerophosphate and complete EDTA-free protease inhibitors (Roche). The cell lysate was cleared by centrifugation for 20 min at $21,000 \times g$ and 4 °C. While centrifuging the supernatant a second time in the same condition, the HisPur Cobalt resin (Thermo Scientific) was equilibrated in the lysis buffer. The cell lysate was incubated with the resin for at least one hour and afterward extensively washed with the lysis buffer. His-tagged Akt proteins were then eluted from the resin in several fractions using a buffer containing 10 mM HEPES (pH 7.6), 3 mM MgCl₂, 300 mM KCl, 10% glycerol and 250 mM imidazole.

Peptide synthesis and dot immunoblot assays.

N-terminal biotinylated peptides used for dot blot assays were synthesized at Tufts Medical School in 0.1 mmol scale. The sequences were listed below:

Akt1-K140-WT (aa131–151): AEEMEVSLAKPKHRVTMNEFE;

Akt1-K140-me3: AEEMEVSLA*KPKHRVTMNEFE;

Akt1-K140/142-me3: AEEMEVSLA*KP*KHRVTMNEFE (*Denoted tri-methylation).

Peptides were diluted into 1 μ M for further biochemical assays. For dot blot assays, peptides were spotted onto nitrocellulose membrane allowing solution to penetrate (usually 3–4 mm diameter) by applying it slowly as a volume of 1 μ l. The membrane was dried, and blocked in non-specific sites by soaking in TBST buffer with 5% non-fat milk for immunoblot analysis as described previously ¹⁷.

Akt in vitro kinase assays.

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Akt *in vitro* kinase assays were adapted from a protocol described previously ¹⁶. Briefly, 1 μ g of the bacterially purified His-GSK3 β fusion proteins were incubated with immunoprecipitated Akt1 from HEK293 cells transfected with various mutant Akt1 in the presence of 100 ng [γ S]ATP (abcam 138911) in the kinase reaction buffer (50 mM Tris pH 7.5, 1 μ M MnCl₂, 2 mM DTT) for 30 minutes at 30°C. The reaction was subsequently stopped by adding in 0.1 mM EDTA, and further react for another 1 hour by adding in the pNitrobenzyl mesylate (abam 138910) to alkylate the thiophosphorylation site on the substrates. The reaction was stopped by adding 3 x SDS loading buffer and resolved by SDS-PAGE. Phosphorylation of His-GSK3 β was detected by specific antibody to recognize thiophosphate ester as previously reported ⁵⁷.

In vitro methylation and demethylation assays.

The *in vitro* methylation assays were performed as described ⁵⁸. In briefly, Flag-tagged SETDB1 were expressed and purified from HEK293T cells serving as the source of methyltransferase. The Akt1 proteins including GST-tagged Akt1 expressed and purified from HEK293T cells, recombinant his-Akt1 obtained from Thermofisher Scientific, or the Akt1 peptides synthesis from Tufts were used as substrates. The reactions were performed in the condition of 50 mM Tris-HCl (pH 8.5), 5 mM MgCl₂, 4 mM DTT and 10 μ M S-adenosyl-methionine (SAM) at 37°C for 6 hrs.

Demethylation assays were performed as reported previously ⁵⁹. In briefly, the bacterially purified KDM4B catalytic domain (1–400aa) were used as the source of demethylase, and the synthetic Akt1-K140-me3 peptides were used as substrates. The reactions were performed in the condition of 50 mM Tris-HCl (pH 8.2), 2% glycerol, 0.1% MgCl₂, 0.1% KCl and 0.1 mM PMSF at 37°C for 12 hrs.

In vivo ubiquitination assays.

Akt ubiquitination assays were performed as reported previously ⁴¹. Briefly, His-ubiquitin along with different variants of Akt1 were transfected into cells, and resulting cells were lysed in buffer A (6 M guanidine-HCl, 0.1 M Na₂HPO4/NaH₂PO₄, and 10 mM imidazole [pH 8.0]) and subjected to sonication. After high-speed centrifuged, the supernatants were incubated with nickel-beads (Ni-NTA) (QIAGEN) for 3 hrs at room temperature. The products were washed twice with buffer A, twice with buffer A/TI (1 volume buffer A and 3 volumes buffer TI), and one time with buffer TI (25 mM Tris-HCl and 20 mM imidazole [pH 6. 8]). The pull-down proteins were resolved in 8% SDS-PAGE for immunoblot analysis.

Mass spectrometry analyses.

For mass spectrometry analysis, anti-HA immunoprecipitations (IP) were performed with the whole cell lysates derived from three 10 cm dishes of HEK293 cells co-transfected with Flag-SETDB1 and HA-Akt1. The IP proteins were resolved by SDS-PAGE, and identified by Coomassie staining. The band containing Akt1 was reduced with 10 mM DTT for 30 minutes, alkylated with 55 mM iodoacetamide for 45 minutes, and in-gel-digested with trypsin enzymes. The resulting peptides were extracted from the gel and analyzed by

microcapillary reversed-phase (C_{18}) liquid chromatography-tandem mass spectrometry (LC-MS/MS), using a high resolution QExactive HF Orbitrap (Thermo Fisher Scientific) in positive ion DDA mode (Top 8) via higher energy collisional dissociation (HCD) coupled to a Proxeon EASY-nLc II nano-HPLC ⁶⁰. MS/MS data were searched against the Uniprot Human protein database (version 20151209 containing 21,024 entries) using Mascot 2.5.1 (Matrix Science) and data analysis was performed using the Scaffold 4.4.8 software (Proteome Software). Peptides and modified peptides were accepted if they passed a 1% FDR threshold.

Colony formation assays.

Cells were seeded into 6-well plates (300 or 600 cells/well) and left for 8–12 days until formation of visible colonies. Colonies were washed with PBS and fixed with 10% acetic acid/10% methanol for 20 minutes, then stained with 0.4% crystal violet in 20% ethanol for 20 minutes. After staining, the plates were washed and air-dried, and colony numbers were counted.

Soft agar assays.

The anchorage-independent cell growth assays were performed as described previously ¹⁶. Briefly, the assays were preformed using 6-well plates where the solid medium consists of two layers. The bottom layer contains 0.8% noble agar and the top layer contains 0.4% agar suspended with 1×10^4 or 3×10^4 cells. 500 µl complete DMEM medium was added every 7 days to keep the top layer moisture and 4 weeks later the cells were stained with iodonitrotetrazolium chloride (1 mg/ml) (sigma I10406) for colony visualization and counting.

Glucose uptake and lactate production assays.

For glucose uptake assays, cells were cultured in 60 cm plates, and were starved for 24 hrs with serum-free and glucose-free DMEM, then subjected to 20 μ M 2-NBDG (Sigma 72987) containing glucose-free DMEM for different time points. Last, the cellular glucose uptake was quantified by FACS analysis using BD FACSDiva 6.1.3 software.

For lactate production assays, cells were planted into 60 cm plates, and were cultured for 24 hrs, then the cultured medium was collected, and lactate concentration was measured with lactate test strips and Accutrend Lactate analyzer (Accutrend Lactate, Nova Biomedical). At the same time, the viable cell numbers were calculated under the microscope. Finally, the relative lactate production was calculated with the formula (relative lactate production = lactate concentration/cell numbers) and normalized with the ratio of control cells.

Mouse xenograft assays.

Mouse xenograft assays were performed as described previously ⁴. Briefly, 2×10^6 DLD1-*AKT1/2^{-/-}* cells stably expressing WT or K140/142R mutant form of Akt1, or 1×10^6 A375 cells knocked down *SETDB1* were injected into the flank of 8 female nude mice (NCRNU-M-M from Taconic, 4–5 weeks of age). Tumor size was measured every three days with a caliper, and the tumor volume was determined with the formula: LxW²×0.5, where L is the longest diameter and W is the shortest diameter. In the drug treatment group, when the

tumor reached 100 mm³, the mice were i.p. injected with Mithramycin A (0.2 mg/kg) or PBS (as a negative control) in the other day. After around 18 days, mice were sacrificed and xenografted solid tumors were dissected, then tumor weights were measured and recorded post-necropsy.

Statistics and reproducibility

Statistical analysis (two-tailed Student's *t* test) was performed using Prism 8 (GraphPad Software) for Figs. 1h; 2b, 2c and 7n and Supplementary Fig. 7i in this manuscript. To calculate the *P* value between groups in Figs. 1f, 1j; 2h, 2l; 7b, 7l and Supplementary Figs. 7g, two-way analysis of variance (ANOVA) analysis was performed with Prism 8. *Chi-square* test was also performed to analyze the *P* value for Figs. 3j, 3l, 3m and Figs. 6m, 6n, 6o by using Prism 8. All the *P* values calculated above are indicated in the corresponding figures, and derived from three independent experiments with 3 technical replicates for each experiment. MS screens were performed twice successfully. All the western blots were performed twice independently with similar results. *In vitro* methylation assays were performed once in Fig. 3e, and performed twice in Fig. 3g and supplementary Fig. 3h independently with similar results. *In vitro* demethylation , ubiquitination, soft agar, colony formation, glucose uptake and lactate production assays were performed twice independently with similar results. To knock out or knock down genes of interest, at least two independent sgRNAs or shRNAs were employed to generate cell lines, and similar results were found in all cell lines.

Reporting Summary

Further information on experimental design is available in the Nature Research Reporting Summary linked to this article.

Data availability

The mass spectrometry-based screening data generated in this study have been deposited in ProteomeXchange under the accession code PXD011657. The SETDB1, EZH2, PTEN, EGFR, PIK3CA and AKT1 genetic alterations in TCGA datasets was integrated from cBioPortal database (www.cbioportal.org), with the query of gene "ESET", "EZH2", "PTEN", "EGFR", "PIK3CA" and "AKT1" for both mutation and copy number alterations (CNA) in different cancer types, such as BRCA and Melanoma. Following are the information of each cancer studies. One dataset for BRCA: TCGA, provisional. One dataset for Melanoma: TCGA, provisional.

Source data for Fig. 1e-f, 1h, 1j-m; 2b-c, 2g-h; 3n, 3p-q; 4c, 4f, 4h, 4j, 4l, 4n; 6m-o; 7b, 7d, 7j, 7l, 7n and Supplementary Fig. 1l-m; 2e-f; 3o; 4c-f, 4h; 7e, 7g, 7i have been provided as Supplementary Table 2. All other data supporting the findings of this study are available from the corresponding author upon reasonable request.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Fig. 1. Akt methylation promotes its activity and oncogenic functions

a, A schematic workflow of IAP-LC-mass spectrometry (MS)/MS experiments. OVCAR5 cell lysates were proteolytically digested to perform IAP-LC-MS/MS assays. b, Alignment of MS-characterized Akt1 putative methylation residues among different species, Akt2 and Akt3. c, Immunoblot (IB) analysis of whole cell lysates (WCL) derived from DLD1- $AKT1/2^{-/-}$ cells infected with indicated Akt1 encoding virus and selected with hygromycin (200 µg/ml) for 72 hrs before harvesting. Data shown represent two independent experiments. d-e, cells generated in c were subjected to colony formation and soft agar assays. The experiment was performed twice independently with three repeats, and exhibited similar results (d). Representative images were shown in d and relative colony numbers derived from two independent experiments were plotted in e. f-h, Cells generated in (c) were subjected to mouse xenograft assays. Tumor sizes were monitored (f), and dissected tumors were weighed (g, h). Error bars in f and h are mean \pm s.e.m, n = 7 mice. i-k, K140R and K142R mutations of Akt1 were genetically engineered in HEK293 cells by the CRISPR/ CAS9-based technique. Resulting cells were serum-starved for 12 hrs, and harvested for IB analysis at different time points after stimulation with insulin (100 nM) (i). The experiment was performed twice independently with similar results (i). Cells generated in (i) were assessed for proliferation (j) assays. The experiment in j was performed three times independently and exhibited similar results. Error bars in **j** are mean \pm s.e.m, n = 3 independent experiments. k,l,m, Cells generated in i were subjected to colony formation (k), glucose uptake (I) and lactate production (m) assays. The experiment was performed twice independently with three repeats, and exhibited similar results (k,l,m). Relative colony

numbers, glucose and lactate levels derived from two independent experiments were plotted in $(\mathbf{k,l,m})$. Two-way ANOVA analysis was performed in $(\mathbf{f,j})$ to calculate the *P* value. Detailed statistical tests are described in the Methods. Source data for \mathbf{e} , \mathbf{f} , \mathbf{h} and $\mathbf{j-m}$ are shown in Supplementary Table 2. Scanned images of unprocessed blots are shown in Supplementary Fig. 8.



Fig. 2. Methylation deficient *Akt1* knock-in mice display reduced body size/weight and resist to chemical carcinogen-induced skin tumorigenesis *in vivo*

a, Mice derived from the same litter were imaged at age of 4 weeks old. **b**, The mice derived from the age of 4 weeks were weighed (including 15 male mice with 5 WT, 5 heterogeneous, and 5 homogeneous- $Akt1^{K140/142R}$ genetic background status). Error bars are mean \pm s.e.m, n = 5 mice. *P* values were calculated using two-tailed unpaired Student's *t* test. **c**, The mice in (**b**) were euthanized and their organs were dissected and weighed. Error bars are mean \pm s.e.m, n = 5 mice. *P* values were calculated using two-tailed unpaired Student's *t* test. **d**, IB analysis of WCL derived from livers or hearts of WT (+/+), $Akt1^{K140/142R}$ -knock-in heterogeneous (+/KI) or homogeneous (KI/KI) mice from the same litter at age of 4 weeks. The experiment was performed twice, independently, with similar results. **e**, Graphic representation of H&E and IHC staining of heart and liver tissues derived

from WT or $Akt1^{K140/142R}$ -KI mice. Scale bar, 50 µm. This experiment was performed twice, independently, with similar results. **f**, The side view of 12-weeks old mice derived from WT or $Akt1^{K140/142R}$ knock-in mice were treated with chemical carcinogen (DMBA following with TPA) (n = 16 for WT mice; n = 11 for Akt1-K140/142-KI mice). The neoplasm lesions were arrowed. The tumor incidence (**g**) and lesion numbers (**h**) of the mice described in (**f**) were calculated and plotted. Error bars are mean ± s.e.m (for WT n = 16 mice; for Akt1-K140/142-KI n=11 mice). After treatment 12 weeks with DMBA/TPA, the mice were euthanized and the H&E and IHC staining were performed (**i**). Scale bar, 1 mm. The experiment in (**i**) was performed twice, independently, with similar results. Statistical source data for **b**,**c**,**g** and **h** are shown in Supplementary Table 2. Scanned images of unprocessed blots are shown in Supplementary Fig. 8.



Fig. 3. SETDB1 methylates Akt on K140 and K142 to promote its kinase activity

a-d,f,h, IB analysis of immunoprecipitates (IP), GST pulldown products and WCL derived from A375 and OVCAR5 (**a**) or HEK293 cells transfected with indicated constructs (**b-d, f**) or *AKT1^{K140/142R}*-KI cells stably expressed WT or catalytically inactive SETDB1-H1224K (**d**). IgG was used as a negative control. **e**, *In vitro* methylation assays were performed with recombinant His-Akt1 proteins purified from insect cells as substrates and purified Flag-SETDB1 from HEK293T cells as the source of methyltransferase in the presence/absence of ³H-SAM. **g**, *In vitro* methylation assays were performed with IP Flag-SETDB1 derived from HEK293T cells as the source of methyltransferase, and the synthetic Akt1 peptides containing K140 and K142 as the substrate. Akt1-K140-me3 peptides were used as a positive control. **h**, IB analysis of IP products and WCL derived from A375 cells lentivirally transfected with shRNAs against *SETDB1*. Resulting cells were selected with puromycin for

72 hrs before harvesting. **j**, *Setdb1* conditional knockout MEFs were treated with or without 4-OHT (500 nM) for 48 hrs to deplete endogenous *Setdb1*, then resulting cells were serumstarved for another 20 hrs and stimulated with insulin (100 nM) for 15 min before being harvested and subjected to IP and IB analysis. **k**, IB analysis of IP products and WCL derived from HEK293 cells transfected with indicated constructs. **l**, *Setdb1* conditional knockout MEFs were infected with WT or H1224K-SETDB1 encoding virus and selected with puromycin for 72 hrs, then treated with or without 4-OHT (500 nM) for another 48 hrs before harvesting for IB analysis. **m-q**, Image illustration of the immunohistochemistry (IHC) staining for SETDB1, pS473-Akt1 and K140-me3-Akt1 in melanoma TMA (**m,o**). Scale bar, 50 µm. The distribution of SETDB1 staining was plotted in (**n**) (n = 97 tissue specimens). The correlation of pS473-Akt or K140-me3-Akt with SETDB1 were plotted in (**p,q**) (n = 95 tissue specimens). Western-blots in (**a-1**) were performed twice, independently, with similar results. *P* values were calculated using *Chi-Square* test in (**n-q**). Statistical source data for **n-q** are shown in Supplementary Table 2. Scanned images of unprocessed blots are shown in Supplementary Fig. 8.



Fig. 4. Oncogenic function of SETDB1 depends on the activation of Akt

a-c, Human immortalized melanocytes (HIM) were lentivirally infected with indicated constructs, and selected with puromycin and hygromycin for 72 hrs before harvesting for IB analysis (a). Resulting cells were subjected to soft agar assays (b). d, IB analysis of CRISPR/CAS9-mediated AKT1 knockout and parental HEK293 cells which were lentivirally infected with the constructs encoding SETDB1. e,f, Cells described in d were subjected to colony formation assays (e). g-j, DLD1-AKT1/ $2^{-/-}$ cells were infected with virus encoding WT or mutated Akt1, and selected with hygromycin for 72 hrs. Resulting cells were lentivirally infected with shRNA against SETDB1 (with shCtr as a negative control) and selected with puromycin for 72 hrs, and were harvested for IB analysis (g), cell proliferation (h), colony formation and soft agar (i) assays. k-n, AKT1 K140/142R-edited and parental HEK293 cells were lentivirally infected with SETDB1-WT or SETDB1-H1224K encoding constructs and selected with puromycin for 72 hrs before harvesting for IB analysis (k). Resulting cells were subjected to proliferation (l) and colony formation (m) assays. The experiments in (b,e,i,m) were performed twice independently with three repeats, and exhibited similar results. Relative colony numbers derived from two independent experiments were plotted in (c,f,j,n). The cell proliferation assays in (h,l) were performed three times independently, and cell numbers were quantified in (h,l). Error bars are mean \pm s.e.m, n = 3 independent experiments. Two-way ANOVA analysis was performed in (h,l) to calculate the Pvalue. Detailed statistical tests are described in the Methods. Source data for c,f,h,j,l and n are shown in Supplementary Table 2. Western-blots in (a,g,k) were performed twice, independently, with similar results. Scanned images of unprocessed blots are shown in Supplementary Fig. 8.



Fig. 5. SETDB1-medidated methylation of Akt synergizes with PI3K to activate Akt

a,b, A375 cells were serum-starved for 20 hrs, then stimulated with insulin (50 nM) at different time points (a) or post-treatment of various PI3K inhibitors (b) before harvesting for IP and IB analysis. c,d, IB analysis of IP products and WCL derived from HEK293 cells transfected with indicated constructs stimulated without (c) or with IGF (100 ng/ml) (d) before harvesting. e, In vitro binding assays were performed with recombinant GST-Akt1 protein purified from mammalian cells, and flag beads bound SETDB1. The binding was performed in 4° C for 4 hrs incubated with or without PIP₃ (20 μ M) and subjected to IB analysis. f-h, IB analysis of Akt1-IP and WCL derived from HEK293 cells infected with indicated SETDB1 encoding constructs (f), AKT1K140/142R and its parental HEK293 cells (g) and SETDB1 depleted A375 cells (h). i,j, IB analysis of PIP₃ pull-down products and WCL derived from Setdb1 conditional knockout MEFs treated with or without 4-OHT (500 nM) for 48 hrs (i) or from HEK293 cells transfected with indicated constructs (j). Where indicated, empty beads (Ctr) serve as a negative control. k,l, IB analysis of cell fractionations separated from Setdb1 conditional knockout MEFs treated with or without 4-OHT (500 nM) for 48 hrs (k) or from AKT1K140/142R-edited and parental HEK293 cells (l). All Western-blots above were performed twice, independently, with similar results. Scanned images of unprocessed blots are shown in Supplementary Fig. 8.



Fig. 6. KMD4B demethylates Akt to inhibit Akt kinase activity

a,c-e, IB analysis of Akt1-IP products and WCL derived from A375, OVACR5 cells (**a**), SETDB1 expressing HEK293 cells (**c**), *AKT1^{K140/142R}*-edited and parental HEK293 cells (**d**) and *SETDB1*-depleted A375 cells (**e**). IgG was used as a negative control. **b**, IB analysis of *in vitro* de-methylation assays performed with synthetic Akt1-K140-me3 peptides as substrate, and bacterially purified catalytic domain of KDM4B as the source of demethylase. **f**, IB analysis of WCL derived from primary *Kdm4b* conditional knockout MEFs infected with or without phage-Cre for 48 hrs before harvesting. **g**, A375 cells were lentivirally infected with shRNA against *KDM4B*. Resulting cells were serum starved for 20 hrs, then stimulated with IGF (100 ng/ml) before harvesting for IP and IB analysis. **h**, IB analysis of IP products and WCL derived from A375 cells infected with lentivirus against *KDM4B*. **i**, IB analysis of cell fractionations separated from A375 cells lentivirally infected with shRNA

against *KDM4B* or *KDM4A*. **j**,**k**, HEK293 cells were transfected with indicated constructs and treated with different PI3K inhibitors for 1 hr (**j**) before subjected to GST pull-down assay and IB analysis. **l-o**, IHC staining of KDM4B, pS473-Akt1 and K140-me3-Akt1 in melanoma TMA (**l**). Scale bar, 50 µm. The distribution of KDM4B staining was plotted in (**m**, n = 97 tissue specimens). The correlations of pS473-Akt or K140-me3-Akt1 with KDM4B were plotted in (**n**, n = 96 tissue specimens; **o**, n = 95 tissue specimens). All *P* values were calculated using *Chi-Square* test. Detailed statistical tests were described in Methods. Western-blots were performed twice, independently, with similar results. Statistical source data for **m-n** are shown in Supplementary Table 2. Scanned images of unprocessed blots are shown in Supplementary Fig. 8.



Fig. 7. Deficiency of SETDB1 inhibits Akt kinase activity and oncogenic function

a-d, *SETDB1*-depleted A375 and control cells were subjected to mouse xenograft assays. Tumor sizes were monitored (**a**,**b**). Tumors were dissected (**c**) and tumor mass were weighed (**d**). Error bars are mean \pm s.e.m, n = 7 mice. *P* values were calculated by using two-way ANOVA analysis (**b**) and two-tailed unpaired Student's *t* test (**d**). **e**, The phosphorylation status of Akt1 (pT308-Akt) and methylation of H3K9 (H3K9me3) were detected by IB analysis with WCL derived from recovered xenografted tumors. **f**, IB analysis of IP products and WCL derived from A375 cells treated with different doses of Mithramycin A for 72 hrs before harvesting. **g**, HEK293 cells were transfected with indicated constructs and treated with different doses of Mithramycin A for 72 hrs before harvesting for GST pull-down

assays and IB analysis. **h-j**, $AKT1^{K140/142R}$ -edited and parental HEK293 cells were treated with different doses of Mithramycin A for 72 hrs and subjected to IB analysis (**h**). Meantime, resulting cells were subjected to colony formation (**i**) assay. The experiment was performed twice independently with three repeats, and exhibited similar results (**i**). Representative images were shown in (**i**) and relative colony numbers derived from two independent experiments were plotted in (**j**). **k-n**, Mithramycin A treatment reduced *in vivo* tumorigenesis of xenografted A375 cells. When the tumors of xenografted A375 cells reached 100 mm³, the mice were treated with Mithramycin A (0.2 mg/kg) or PBS (as a negative control). Tumor sizes were monitored in (**k**,**l**) and tumor mass were weighed and presented in (**m**,**n**). Error bars are mean \pm s.e.m, n = 8 mice. *P* values were calculated by using two-way ANOVA analysis (**l**) and two-tailed unpaired Student's *t* test (**n**). Detailed statistical tests are described in the Methods. Source data for **b**, **d**, **j**, **l** and **n** are shown in Supplementary Table 2. Scanned images of unprocessed blots are shown in Supplementary Fig. 8.