Transcriptional memory from cells-of-origin overrides β-catenin requirement of MLL cancer stem cells

Authors: Teerapong Siriboonpiputtana†♦, Bernd B. Zeisig‡, Magdalena Zarowiecki*, Tsz Kan Fung†, Maria Mallardo†, Chiou-Tsun Tsai†, Priscilla Na Ieng Lau†, Quoc Chinh Hoang†*, Pedro Veiga†, Jo Barnes†, Claire Lynn†, Amanda Wilson†, Boris Lenhard‡§, Chi Wai Eric So†*

Affiliations:

1 Leukemia and Stem Cell Biology Team, Department of Haematological Medicine, Division of Cancer Studies, King’s College London, SE5 9NU, UK

2 Institute of Clinical Sciences, Faculty of Medicine, Imperial College London, London W12 0NN, UK

3 Computational Regulatory Genomics, MRC London Institute of Medical Sciences, London W12 0NN, UK

4 Sars International Centre for Marine Molecular Biology, University of Bergen, N-5008 Bergen, Norway

♦ Current address: Department of Pathology, Faculty of Medicine, Ramathibodi Hospital, Mahidol University, Bangkok, Thailand

*Department of Cancer Research, Vinmec Research Institute for Stem Cells and Gene Technology, 458 MinhKhai, Hà Nội 084, Vietnam

†Equal contribution

*Correspondence to: eric.so@kcl.ac.uk
Abstract: While β-catenin has been demonstrated as an essential molecule and therapeutic target for various cancer stem cells (CSCs) including those driven by MLL-fusions, here we show that transcriptional-memory from cells-of-origin predicts AML patient survival and allows β-catenin independent transformation in MLL-CSCs-derived from hematopoietic stem cells (HSCs)-enriched LSK population but not myeloid-granulocyte progenitors. Mechanistically, β-catenin regulates expression of downstream targets of a key transcriptional-memory gene, Hoxa9 that is highly enriched in LSK-derived MLL-CSCs and helps sustain leukemic self-renewal. Suppression of Hoxa9 sensitizes LSK-derived MLL-CSCs to β-catenin inhibition resulting in abolishment of CSC transcriptional program and transformation ability. In addition, further molecular and functional analyses identified Prmt1 as a key common downstream mediator for β-catenin/Hoxa9 functions in LSK-derived MLL-CSCs. Together, these findings not only uncover an unexpectedly important role of cells-of-origin transcriptional memory in regulating CSC self-renewal, but also reveal a novel molecular network mediated by β-catenin/Hoxa9/Prmt1 in governing leukemic self-renewal.

Keywords: β-catenin, canonical Wnt, MLL leukemia, Hoxa9, Prmt1, AML, cells-of-origin, cancer stem cells
Introduction

Self-renewal is a critical feature of stem cells, but is diminished upon differentiation into their progenitors. During the differentiation process, gene expression programs responsible for self-renewal are downregulated, and frequently replaced by lineage specific transcriptional programs. Increasing evidence suggest that genes involved in promoting normal stem cell self-renewal are commonly hijacked in cancer stem cells (CSCs), which are believed to sustain the disease and be responsible for relapse of various cancers including acute myeloid leukemia (AML)(Fung, Leung et al., 2013, Zeisig, Kulasekararaj et al., 2012). Among them, one of the most striking molecules is β-catenin, which is required for leukemic stem cells (LSCs) driven by MLL-fusion proteins or their downstream targets, Meis1/Hoxa9(Wang, Krivtsov et al., 2010, Yeung, Esposito et al., 2010). Suppression of β-catenin reversed MLL-LSC to pre-LSC stage(Yeung et al., 2010), and its complete inactivation prevented development of leukemia driven by MLL-fusions or Meis1/Hoxa9(Wang et al., 2010, Yeung et al., 2010). While β-catenin is critical for embryonic and a number of somatic stem cells including fetal hematopoietic stem cells (HSCs)(Malhotra & Kincade, 2009, Zhao, Blum et al., 2007), it is largely dispensable for adult HSCs, which can function normally when β-catenin alone or even together with γ-catenin are deleted(Cobas, Wilson et al., 2004, Jeannet, Scheller et al., 2008, Koch, Wilson et al., 2008), highlighting the therapeutic potentials of targeting β-catenin for eradication of LSCs.

In spite of the fundamental difference between stem cells and their progenitors in self-renewal ability, others and we have shown that phenotypically and genetically indistinguishable cancers, including MLL-rearranged leukemia, can arise from not only stem cells, but also their immediate downstream short-lived progenitors with distinctive transcriptional programs(Blanpain, 2013, Cozzio, Passegue et al., 2003, Huntly, Shigematsu et al., 2004, Krivtsov, Twomey et al., 2006, So, Karsunky et al., 2003, Visvader, 2011). Consistently, gene expression signatures associated with stem cells and progenitors correlate with different clinical outcomes in AML(Eppert, Takenaka et al., 2011, Krivtsov, Figueroa et al., 2013). MLL leukemia derived from mouse HSCs enriched Lin−Sca1+ckit+ (LSK) populations can be more aggressive and less responsive to standard chemotherapy than those derived from granulocyte-
myeloid progenitors (GMPs) (Krivtsov et al., 2013). In line with this, a more recent study also reveals that HSC-derived leukemia drives an invasive EMT-related gene expression program, which may partly account for the aggressive nature of the disease (Stavropoulou, Kaspar et al., 2016). In spite of these recent evidence indicating the importance of cancer cells-of-origin in disease pathogenesis, we still do not know if and how they may govern the utilization of molecular pathways for self-renewal, which is a defining feature of CSC and has been a major focus for development of effective cancer therapeutics in the past decade.

Given the important function of β-catenin in CSC biology, we carried out detailed functional biology and molecular studies examining β-catenin requirement in MLL-CSCs originated from different cells-of-origin. Here we report that transcriptional-memory from cells-of-origin that robustly predicts AML patient survival can govern and help to override the β-catenin requirement in MLL-CSCs. Mechanistically, we identify a novel transcriptional network mediated by Hoxa9/Prmt1 in sustaining leukemic self-renewal in the absence of β-catenin in HSCs-derived MLL-CSCs. These findings reveal previously unrecognized functions and molecular networks from cancer cells-of-origin that allow override of β-catenin dependent leukemic self-renewal, adding a new dimension to the ongoing research efforts in developing effective therapeutics for eradication of CSCs.
Results

LSK- but not GMPs-derived MLL-CSCs can override β-catenin requirements for leukemic self-renewal

To determine the functional requirement of β-catenin in MLL-CSCs derived from different cells-of-origin, we employed the previously described retroviral transduction/transformation assays (RTTA) (Yeung & So, 2009, Zeisig & So, 2009) using HSCs-enriched Lin'Sca-1+c-Kit+ population (LSK), granulocyte/macrophage progenitors (GMPs), and control c-Kit+ cells (mixed population consisting of mostly progenitors) from Ctnnb1fl/fl CreER (Brault, Moore et al., 2001) conditional knockout mice (Figure 1A, Appendix figure S1A, B). Consistent with previous findings (Yeung et al., 2010), β-catenin was not required for MLL-ENL in vitro transformation of c-Kit+ cells (Appendix figure S1C-E), but essential for in vivo development of CSCs (Figure S1F). Similarly, MLL-ENL could transform LSK and GMPs independently of β-catenin in vitro, and formed compact colonies with early myeloid phenotypes (Figure 1B-D, Appendix figure S1G, H). However, while β-catenin deletion in GMP-MLL-ENL abolished its leukemogenic potentials in vivo (Figure 1E), β-catenin deletion had little impact on LSK-MLL-ENL, which could still induce leukemia with indistinguishable phenotypes and largely similar latencies as compared with the wild-type controls (Figure 1F-H). More importantly, LSK-MLL-ENL β-catenin deficient cells could competently induce AML upon secondary transplant (Figure 1F-H, Appendix figure S1I, J), which readout the self-renewal property of CSCs and indicate the largely uncompromised CSC property in the absence of β-catenin in LSK-derived but not GMPs-derived MLL-CSCs. The results could also be reproduced using a different MLL-ENL construct carrying the minimal transformation domain (Slany, Lavau et al., 1998) and MLL-AF9 (Smith, Yeung et al., 2011), and were not due to different expression levels of the MLL fusions in these populations (Appendix figure S1K-M).

To gain further insights into the role of β-catenin in disease development, we followed the in vivo kinetics of the MLL transformed cells derived from different cellular origins with or without β-catenin. The results showed a similar percentage of engraftment across all samples of different cellular origins and genotypes at 16 and 96 hours post transplant (Figure 1I), suggesting that β-catenin deletion did not significantly affect homing and early in vivo proliferation abilities. In contrast to
LSK-derived MLL-CSCs that continued to expand and induced leukemia in the absence of β-catenin, the expansion of GMP-MLL-ENL Ctnnb1^{del/del} cells slowed down at 15 days and were gradually lost in vivo over a 4-months period (Figure 1I), consistent with an impaired self-renewal.

**β-catenin is also not required for leukemia maintenance by LSK-derived MLL-CSCs**

To explore the function of β-catenin in the maintenance of leukemia derived from different origin-specific CSCs, full-blown leukemic cells harvested from primary leukemic mice carrying Ctnnb1-floxed alleles were then treated with either EtOH or tamoxifen prior to transplantation into secondary recipients (Figure 1A). As expected, both EtOH-treated LSK- and GMPs-derived MLL leukemic cells could competently induce leukemia. Inactivation of β-catenin in GMP-MLL-ENL totally abolished their leukemogenic potential (Figure 1J), while β-catenin deficient LSK-MLL-ENL still efficiently induced leukemia and even with a slightly shorter latency in secondary recipients (Figure 1K). In contrast to the absolute requirement of β-catenin for development of various LSCs (Wang et al., 2010, Yeung et al., 2010, Zhao et al., 2007), the finding of largely dispensable function of β-catenin in LSK-derived MLL-CSCs reveals an unexpected and previously unrecognized role of cells-of-origin in governing leukemic self-renewal for both cancer initiation and maintenance.

**Genomic variations do not account for contrasting β-catenin dependence in MLL-CSCs from different cells-of-origin**

To assess if the observed cell-type specific differences could be explained by random genetic changes associated with particular cell types, nucleotide variations were called from all actively transcribed genes using RNA-Seq. Among 23,766,084 high quality base pairs (depth≥10 and quality≥30), the vast majority of sites (23,747,763) were invariant in all our primary cell lines, and identical to the reference genome GRCm38, while a very small proportion, 4663 SNPs (single nucleotide polymorphisms) were invariant and different from GRCm38. In depth variance analysis revealed that the difference between the LSK-MLL-ENL and GMP-MLL-ENL cells was not larger than between normal cells (Figure 2A, Dataset EV1A-C), and there were no fixed differences between the samples that could have caused the
observed phenotypic differences. To further profile the non-coding genomes, whole genome sequence analysis on LSK-MLL-ENL and GMP-MLL-ENL cells covering 918,583,518 high quality base pairs revealed 917,764,811 invariant sites, and a very small number of variants in the samples; 39,846 variants were different between the two biological replicates (mice), and only 17,225 were found different between the two cell types (Figure 2B, Dataset EV1C). Interestingly, the distribution of SNPs differing between cell types or mice was comparable for both comparisons in non-coding and coding regions (Appendix figure S2A). Consistently, SNPs in coding regions occurred in similar proportions in exon, intron and UTR regions in both comparisons (Appendix figure S2B). Moreover, the number of derived SNPs in GMP-MLL-ENL (9309) was higher than those in LSK-MLL-ENL cells (7916) that exhibited β-catenin-independent phenotypes (Dataset EV1C). Additionally we examined and compared the copy number variances (CNVs) between the LSK-MLL-ENL and GMP-MLL-ENL genomes (Figure 2C-D). As a result, we observed very little CNVs in both genomes. There is only a very small genomic region of about 1kb showing CNV between same cell types (i.e., LSK-MLL-ENL vs GMP-MLL-ENL), whereas multiple chromosomal regions of about 50kb exhibiting CNV were detected between samples (i.e., mouse Exp60 vs mouse Expt69 in Dataset EV1D). Importantly, there is also no known coding gene in the 1kb CNV region shared between cell types (Figure 2D), consistently indicating insignificant genomic difference between LSK-MLL-ENL vs GMP-MLL-ENL cells which could account for their contrasting β-catenin dependence. Together, this data reveals relatively few genomic variation in LSK-MLL-ENL compared with GMP-MLL-ENL and the controls, suggesting that non-genomic influence from the cells-of-origin can be a key factor in governing the self-renewal property of genetically and phenotypically indistinguishable cancers.

**Transcriptional memory from cells-of-origin governs self-renewal pathways and predicts AML patient survival**

As self-renewal in normal stem cells is maintained by specific transcriptional programs, we hypothesized that the transcriptional memories from LSK and GMPs would be partially preserved even after transformation, resulting in transcriptional and functional differences observed in the respective CSCs(Zeisig et al., 2012). Thus
RNA-seq analyses of normal LSK, GMPs and their MLL-ENL transformed counterparts were carried out. There were, as expected, large transcriptional differences between normal LSK and GMPs with 4768 significantly differentially expressed genes, including *Hox* genes, *Meis1* and *Evi1* (Figure 2E, Appendix figure S2C, Dataset EV2A-B), while overall gene expression differences between cells of different origin decreased after MLL-ENL transformation (Figure 2E, Appendix figure S2D). Nevertheless, a significantly larger than expected by chance number of genes remained differentially expressed between LSK and GMP even after transformation (Figure 2F, Appendix figure S2C, Dataset EV2C), indicating the presence of “transcriptional-memory” retained from the cells-of-origin. Toppgene functional annotation revealed genes associated with AML are consistently present in both signatures (Appendix figure S2F-I, Dataset EV2D).

To further investigate the relevance of this cells-of-origin transcriptional-memory gene signature in human leukemia, we employed it to stratify 1290 human AML patients from multiple independent centers (Cancer Genome Atlas Research, 2013, Metzeler, Hummel et al., 2008, Raoni, Harousseau et al., 2007, Raoni, Lancet et al., 2008, Valk, Verhaak et al., 2004, Wouters, Lowenberg et al., 2009) (Dataset EV2E). AML patients with LSK-transcriptional memory signature had much worse prognosis with a median survival 14.5 months as compared to patients with GMP-transcriptional memory signature with median survival 22.7 months (Figure 2G), even though the two groups had similar WBC count (means=40.3, 45.3, t-test p=0.30), age distributions (means= 48.0, 50.0, t-test p=0.07) and cytogenetic risk (cytogenetic risk (1/2/3)=71/199/85, 82/165/66, Chi-square test p=0.16). When compared with the previously identified human HSC signature (Eppert et al., 2011) and MLL leukemic-GMP (LGMP) signatures from different cells-of-origin (Krivtsov et al., 2013), the current transcriptional memory signature represents a stronger predictor to stratify patients into different prognostic subgroups based on both resultant median survivals and p-values (Appendix figure S2J). Moreover, multivariate analyses consistently resulted in significant Cox proportional hazards ratios >1; z-score < 0.1 with both human HSC signature and transcriptional memory signature (Dataset EV2F). Together, these data indicate functional and pathological relevance of the newfound cells-of-origin transcriptional memory in governing human cancer biology beyond known cytogenetic/genetic risk factors.
**Hoxa9 as a key transcriptional memory gene phenocopies β-catenin function in development of origin-specific MLL leukemia**

Given the largely dispensable function of β-catenin in LSK-derived MLL-CSCs, we hypothesize that some self-renewal programs from normal stem cells may persist after transformation, and can sustain self-renewal in the absence of β-catenin. In the transcriptional-memory signature, there were a small number of self-renewal genes such as Hoxa9, Hoxa10 and Meis1 (Figure 2E, F), which are known downstream targets of MLL fusions (Huang, Sitwala et al., 2012, Milne, Briggs et al., 2002, Zeisig, Milne et al., 2004), indicating that their degrees of activation are in part also determined by the cellular origins. Strikingly, RNA-sequencing analysis on MLL-ENL transformed cells upon β-catenin inactivation revealed a specific up-regulation of targets genes suppressed by Hoxa9/Meis1, suggesting a critical function of β-catenin in regulation Hox/Meis1 axis for leukemic self-renewal (Figure 3A, Dataset EV3A-C). Moreover, various stem cell related gene sets were positively enriched in β-catenin deleted LSK-MLL-ENL cells as compared with β-catenin deleted GMP-MLL-ENL (Figure 3B, Appendix figure S3A, Dataset EV3C). β-catenin deleted LSK-MLL-ENL not only expressed higher levels of Hoxa9 (Figure 3C, D) but also showed a negative enrichment for genes repressed by Hoxa9 (Figure 3E, Dataset EV3C). Together, the data consistently suggests a potential Hoxa9 complementation function in replacing β-catenin in LSK-derived MLL-CSCs.

Similar to β-catenin, activation of Hoxa9 enhances self-renewal, while its deletion does not have significant impact on HSCs (Lawrence, Christensen et al., 2005, Smith et al., 2011, So, Karsunky et al., 2004), consistent with the existence of multiple complementary self-renewal pathways in HSCs. We hypothesize if there is indeed a functional complementation between Hoxa9 and β-catenin, Hoxa9 requirement for MLL transformation may also be influenced by cells-of-origin. To address this issue, we used purified hematopoietic populations from Hoxa9 knockout mice for RTTA. While MLL-ENL could competently transform wild-type LSK and GMPs in vitro, only LSK but not GMPs could be transformed by MLL-ENL in the absence of Hoxa9 (Figure 3F, Appendix figure S3B). More importantly, LSK-MLL-ENL Hoxa9−/− similar to wild-type LSK-MLL-ENL could induce serially transplantable leukemia in recipient mice (Figure 3G, Appendix figure S3C), strongly suggesting that Hoxa9 requirement, similar to β-catenin, is also largely determined by
cancer cells-of-origin. These results not only assert the critical function of the newfound cells-of-origin transcription-memory in governing the biology of the resultant disease, but also suggest that LSK-MLL-CSCs may be able to utilize the Hoxa9-mediated self-renewal pathways as a molecular mechanism to overcome targeted disruption of β-catenin function.

**Suppression of Hoxa9 abolishes β-catenin independent transformation in LSK-derived MLL-CSCs**

To gain further molecular insights into the functional interplay between β-catenin and Hoxa9 in mediating self-renewal in origin-specific CSCs, we generated a novel compound Hoxa9<sup>-/-</sup>Ctnnb1<sup>fl/fl</sup> Rosa-CreER Rosa-YFP mouse by crossing Hoxa9<sup>-/-</sup> mice (Smith et al., 2011) with Ctnnb1<sup>fl/fl</sup> Rosa-CreER Rosa-YFP mice for RTTA and RNA-sequencing analysis. While MLL-ENL transduced LSK and GMPs isolated from compound Hoxa9<sup>-/-</sup>Ctnnb1<sup>fl/fl</sup> Rosa-CreER Rosa-YFP mice produced a similar number of first round colonies (Appendix figure S4A), a further β-catenin inactivation significantly compromised their transformation ability resulting in reduced number and size of colonies with early myeloid phenotypes (Figure 4A-B, Appendix figure S4B-D).

LSK-MLL-ENL Hoxa9<sup>-/-</sup>Ctnnb1<sup>-/-</sup> displayed a higher percentage of apoptosis (Figure 4C), and an increase in G2/M arrest at the expense of S-phase (Figure 4D), which might help explain their reduced numbers and colony sizes (Figure 4A). Upon transplantation, both LSK-MLL-ENL Hoxa9<sup>-/-</sup>Ctnnb1<sup>fl/fl</sup> and Hoxa9<sup>-/-</sup>Ctnnb1<sup>del/del</sup> engrafted in comparable levels into the bone marrow and were able to proliferate and transiently expand (Figure 4E). However, inactivation of β-catenin in LSK-MLL-ENL Hoxa9<sup>-/-</sup> led to a gradual loss of their self-renewal property and failed to induce leukemia (Figure 4E-F). Further in vivo limiting dilution analysis revealed similar frequency of CSCs found in wild-type, Hoxa9<sup>-/-</sup>, or Ctnnb1<sup>-/-</sup> LSK-MLL-ENL, ranging from 1/3381 to 1/8625 (Figure 4G and Appendix figure S4E-G), indicating a rather limited impact of single inactivation of Hoxa9 or β-catenin on LSK-derived MLL-CSCs. In contrast, deletion of both proteins in LSK-MLL-ENL resulted in a drastic reduction of CSC frequency (estimated to be below 1/1669041) (Figure 4G and Appendix figure S4H), consistent with a critical functional crosstalk between
Hoxa9 and β-catenin, in which a high level of Hoxa9 expression allows β-catenin independent transformation in LSK-derived MLL-CSCs.

**β-catenin and Hoxa9 co-regulate Prmt1 in LSK-MLL-ENL cells**

To gain further insights into the molecular pathways underlying β-catenin and Hoxa9 mediated leukemic self-renewal, we sought to define common gene sets that were deregulated upon the loss of leukemic self-renewal (i.e., β-catenin inactivation in Hoxa9/−/− LSK-MLL-ENL, Hoxa9 inactivation in Ctnnb1/−/− LSK-MLL-ENL cells). As a result, 38 gene sets were commonly upregulated (Figure 5A) and 33 were commonly downregulated (Figure 5B, Dataset EV4A-B) in compound Hoxa9/−/− Ctnnb1/−/− LSK-MLL-ENL cells. Consistent with the loss of leukemic self-renewal, LSC_maintenance signatures (Somervaille, Matheny et al., 2009) were inversely enriched. Moreover, the Hoxa9/Meis1 targets (Hess, Bittner et al., 2006) were also inversely enriched, supporting the hypothesis that β-catenin and Hoxa9 may co-regulate common sets of genes critical for self-renewal.

In order to specifically identify β-catenin and Hoxa9 co-regulated targets in LSK-MLL-ENL, we performed global quantitative expression analyses using RNA-Seq, and revealed a small fraction of genes ~1% (n=525) showing a synergistic pattern in the double knockout compared to single knockouts alone (Figure 5C, Appendix figure S5, Dataset EV4C). Different protein classes are present in the 321 synergistically up- and 204 synergistically down-regulated gene lists. While hydrolase and cysteine protease inhibitors were enriched in the up-regulated list, methyltransferases amongst others were enriched in the down-regulated list (Figure 5D). These methyltransferases were arginine-specific Prmt1, Prmt5 and Prmt7. Consistently, H4-R3 specific histone methyltransferase activity was amongst the enriched GO: Molecular functions (Figure 5E, Dataset EV4D), suggesting that arginine methylation may be co-regulated by Hoxa9 and Ctnnb1. Indeed, when the targets of Hoxa9/Meis1 gene sets were overlapped with the synergistically up- and down-regulated genes, 19 out of 111 (17%) genes from the Hoxa9/Meis1 targets (Figure 5F, Dataset EV4E), were synergistically regulated by β-catenin and Hoxa9 in LSK-MLL-ENL ($\chi^2$, p=1.4e-22). Moreover, among them is again Prmt1, which had also been independently confirmed by RT-qPCR (Figure 5F). Strikingly, Prmt1 is a key epigenetic modifying enzyme known to be recruited by various fusion proteins
involved in AML pathogenesis (Cheung, Chan et al., 2007, Cheung, Fung et al., 2016, Shia, Okumura et al., 2012), leading us to examine its role in mediating β-catenin/Hoxa9 functions in LSK-MLL-ENL cells.

Prmt1 regulates similar and overlapping transcriptional programs mediated by β-catenin in Hoxa9−/− LSK-MLL-ENL cells

To investigate the transcriptional functions and potential molecular crosstalk between Prmt1 and β-catenin, global transcriptional analyses by RNA sequencing were performed in Hoxa9−/− LSK-MLL-ENL cells in the presence or absence of shRNA-mediated Prmt1 knockdown using previously validated shRNAs (Cheung et al., 2007). As a result, we identified 1416 differentially expressed genes, including 686 differentially up- and 730 differentially down-regulated genes from two biological replicates upon Prmt1 inactivation (Figure 6A, Dataset EV5A). Similar transcriptomic analyses were then performed between Hoxa9−/− LSK-MLL-ENL and Hoxa9−/−Ctnnb1−/− LSK-MLL-ENL, where we identified 342 differentially up-regulated genes and 134 differentially down-regulated genes from two biological replicates (Figure 6A). To assess if Prmt1 and β-catenin may regulate common transcriptional targets, we compared the differentially expressed genes from both analyses. As a result, we revealed similar and highly significant overlapping gene expression signatures associated with the loss of Prmt1 vs β-catenin in Hoxa9−/− LSK-MLL-ENL cells (Figure 6A, Dataset EV5A). While 24 genes were commonly down-regulated by β-catenin and Prmt1 (p<2.4E-7), 68 genes showed the opposite pattern (p<2.7E-22) upon their individual inactivation (Figure 6A). Functional annotation analysis revealed increased myeloid differentiation and apoptosis but reduced histone binding, chromatin silencing and negative regulation of gene expression as dominant GO:Molecular functions and GO:Biological processes upon Prmt1 knockdown and β-catenin knockout (Figure 6B). Strikingly, GSEA revealed that about 75% of the gene sets/pathways affected by Prmt1 knockdown were also regulated by β-catenin knockout in Hoxa9−/− LSK-MLL-ENL cells (Figure 6C, Dataset EV5B) including those involved in leukemic/normal stem cell functions and stemness (Figure 6D), consistent with the hypothesis that Prmt1 mediates β-catenin functions in Hoxa9−/− LSK-MLL-ENL cells. Together, these results strongly suggest a molecular and
functional overlap between Prmt1 and β-catenin in regulating critical transcriptional programs in LSK-MLL-ENL cells.

**Suppression of Prmt1 abolishes Hoxa9 independent transformation in LSK-derived MLL-CSCs**

To finally evaluate the biological function of Prmt1 as a critical mediator for β-catenin/Hoxa9 functions, we first assessed its requirement in mediating Hoxa9−/− LSK-MLL-ENL transformation. Prmt1 expression was independently down-regulated in Hoxa9−/− LSK-MLL-ENL by two different shRNAs, which also resulted in reduction of H4R3 asymmetric dimethylation mark (H4R3me2as) specifically conferred by Prmt1 (Appendix figure S6A,B). Inhibition of Prmt1 expression compromised in vitro MLL-ENL transformation of Hoxa9−/− LSK cells (Figure 7A, Appendix figure S6C) and mimic in vivo inactivation of β-catenin in LSK-MLL-ENL. Similar to Hoxa9−/−Ctnnb1−/− LSK-MLL-ENL cells (Figure 4G), Hoxa9−/−Prmt1 KD LSK-MLL-ENL cells were able to engraft and proliferate short-term, but gradually lost their self-renewal ability (Figure 7B). Crucially, Prmt1 knockdown suppressed oncogenic potential of Hoxa9−/− LSK-MLL-ENL cells, which could otherwise induce leukemia within a month (Figure 7C).

To further determine if Prmt1 as a key mediator for β-catenin/Hoxa9 can also replace the function of Hoxa9, Prmt1 expression was suppressed in Ctnnb1−/− LSK-MLL-ENL (Appendix figure S6A-B). As a result, Prmt1 inhibition mimic Hoxa9 inactivation leading to suppression of colony formation ability of Ctnnb1−/− LSK-MLL-ENL cells (Figure 7D), abolished their self-renewal potentials (Figure 7E) and oncogenic ability in vivo (Figure 7F). Together with the comprehensive global gene expression network analyses, these results consistently indicate Prmt1 as a key player and novel downstream target mediating β-catenin/Hoxa9 functions in LSK-derived MLL-CSCs.

**Discussion**

Self-renewal as a defining feature of normal and cancer stem cells is tightly regulated by complex transcriptional networks. Most of the current targeted therapies and their intended clinical utility are developed without considering the cancer cells-of-origin, which can have distinctive self-renewal and transcriptional properties. This
traditional view has been challenged by the identification of phenotypically indistinguishable leukemia from different cells-of-origin (Cozzo et al., 2003, Huntly et al., 2004, So et al., 2003), which exhibit different responses to standard chemotherapy treatment (George, Uyar et al., 2016, Krivtsov et al., 2013, Stavropoulou et al., 2016). However until now, very little is known about impact of cells-of-origin on cancer self-renewal and the molecular pathways underpinning this defining feature of CSCs. By performing comprehensive genomic and transcriptomic analyses on origin-specific CSCs in combination with various in vitro and in vivo functional genomic assays, here we provide the experimental evidence for the presence of cells-of-origin transcriptional memory governing molecular pathways available for CSC self-renewal, urging that both genetic mutations and transcriptional memory inherited from cells-of-origin determine the resultant CSC biology and heterogeneous responses to treatment. Key components of the canonical Wnt/β-
catenin signaling pathway are recurrently deregulated in various human cancers, and a number of inhibitor are in early phase clinical trials (Anastas & Moon, 2013). This is particularly relevant to leukemia as normal HSCs remain largely intact upon a complete inactivation of β-catenin and targeting β-catenin represents a promising venue for eradication of LSCs (Fung et al., 2013). However, our results reveal an added dimension of cancer heterogeneity conferred by cells-of-origin transcriptional memory, and predict that pharmacological targeting of β-catenin is unlikely to be effective in MLL leukemia originated from HSCs. We would like to point out that all the published literatures including our studies related to cancer cells-of-origin were performed using mouse cells, which could have different features from the human counterparts. Given the recent evidences from mouse models indicate the importance of the cells-of-origin in governing the resultant cancer biology, future relevant studies in human cell systems are paramount to give necessary insights and to improve our understanding of the cancer cell biology and designing effective therapeutics in the human diseases.

In contrast to its essential function in embryonic and other somatic stem cells, β-catenin is dispensable for adult HSCs, suggesting the presence of residual canonical Wnt signaling for normal HSC function (Malhotra & Kincaide, 2009) or an alternative molecule/pathway compensated for β-catenin in adult hematopoiesis. Interestingly, most of the known molecules/pathways involved in canonical Wnt signaling predominately identified in epithelial cells or ES cells are not significantly affected
upon β-catenin deletion in MLL leukemia in regardless their cells-of-origin (Dataset EV2A, EV3C). In contrast, we identified a number of novel β-catenin targets including those downstream of Meis1/Hoxa9 that are critical for HSC self-renewal, consistently indicating co-regulation of common self-renewal pathways by β-catenin and Hoxa9 in hematopoietic cells. Hoxa9 recently proposed as a key component of human LSC signature in AML (Jung, Dai et al., 2015) can mediate Bmi-1 independent leukemic self-renewal (Smith et al., 2011) and resistance to PARPi treatment in AML (Esposito, Zhao et al., 2015). In line with this finding, others and we have also reported the ability of β-catenin or Hoxa9 in promoting HSC self-renewal (Argiropoulos & Humphries, 2007, Malhotra & Kincade, 2009, Zeisig et al., 2012), but deletion of either one of them yields only mild hematopoietic phenotypes (Cobas et al., 2004, Jeannet et al., 2008, Koch et al., 2008, Lawrence et al., 2005, Smith et al., 2011, So et al., 2004), suggesting their overlapping function in HSC self-renewal. This is further supported by identification of Prmt1 as a key common downstream target that mediates their transcriptional and self-renewal functions. However, the lack of good quality ChIP-grade antibodies against Hoxa9 and β-catenin makes it unfeasible to reliably determine if they might directly bind to Prmt1 regulatory regions. While the lack of known consensus binding sites of Hox or β-catenin/Tcf by in silico analysis (data not shown) suggests an intermediate instead of direct regulation, future investigation is critical to gain further insights into any potential direct functional interactions and regulations. Nevertheless, our comprehensive transcriptomic analyses in combined with functional genomic studies have revealed the novel molecular networks mediated by β-catenin/Hoxa9/Prmt1 in regulating leukemic self-renewal in LSK-derived MLL-CSCs, and exemplify the intricate diversity in molecular pathways utilized by cancer cells to evade therapies, and underscores that simultaneous targeting of multiple self-renewal pathways may be required for successful elimination of certain CSCs.

Intriguingly, activation of canonical Wnt/β-catenin in MLL-CSCs has also recently been identified as a major mechanism for development of resistance to pharmacological inhibition of BET (Fong, Gilan et al., 2015, Rathert, Roth et al., 2015), a targeted therapy at its early clinical phase for MLL leukemia, further highlighting the importance of adequately targeting β-catenin in the context of cells-of-origin for future targeted cancer therapies. Given the challenges involved in developing clinically effective inhibitors to β-catenin (Fung et al., 2013), the
identification of Prmt1 as a nexus for mediating leukemic self-renewal in LSK-MLL-ENL transformed cells not only provides novel mechanistic insights into the downstream targets and molecular networks regulated by β-catenin, but also suggest an alternative avenue for targeting β-catenin in MLL-CSCs.

**Material and Methods**

**Animals and transplantation studies**

All experimental procedures were approved by King’s College London ethics committees and conform to the UK Home Office regulations. For all in vivo experiments, mice were distributed into their respective groups randomly. Investigators were not blinded to the sample identity. Mice were considered leukemic when an engraftment of donor cell (>30%) was detected in the bone marrow. Ctnnb1^{fl/fl} mice (Brault et al., 2001) were crossed with Rosa26-CreER Rosa26-YFP mice to generate Ctnnb1^{fl/fl} RosaCre-ER RosaYFP mice. These mice were crossed with Hoxa9^{-/-} Ctnnb1^{fl/fl} RosaCre-ER RosaYFP mice. Compound homozygous animals were used for experiments. C57BL/6 or SJL mice were given 11Gy total body γ-irradiation and injected via tail vein with test cells mixed with C57BL/6 or SJL bone marrow nuclear cells. Mice were culled when sign of sickness appeared. Survival curves were produced using GraphPad Prism software, and survival differences tested with the log-rank test. For leukemia development experiments, up to 500,000 test cells mixed with 200,000 C57BL/6 or SJL bone marrow nuclear cells were transplanted. For primary transplants n=5-18 mice/cohorts and for secondary transplants n=3-10 mice/cohorts were used. For in vivo limiting dilution assays, varying numbers of cells from indicated populations were transplanted into n=5 sublethally irradiated syngeneic mice/cohorts/mice and monitored for disease development. ELDA was used for statistical analysis (Hu & Smyth, 2009). For in vivo homing experiments, 5 million test cells were transplanted into n≥4 (when studying genetic ablation) and n=2 (when studying shPRMT1 knockdown) sublethally irradiated syngeneic mice/mice/cohorts/time point. Homing differences were statistically tested using a two-tailed t-test in Excel.

**Hematopoietic stem and progenitor purification**

Mouse femur and tibias were prepared and c-kit+ (CD117) cells isolated (Zeisig & So, 2009) using MACS (Miltenyi Biotech Technology, Germany). LSK and GMP populations were isolated as previously described (Yeung & So, 2009). Briefly, LSK (lin-, Sca-1+, c-Kit+) and GMP (lin-, Sca-1-, c-Kit+, CD34+, CD16/32lo) were isolated from lineage negative cells after lineage (Gr-1, Mac-1, B220, Ter119, CD3e, CD4, CD8) (Biolegend/eBiosciences) depletion using a BD FACS ARIA cell sorter. Post-sort purity of greater than 97% was routinely achieved. For functional analysis, sorted LSK and GMP were plated in Methocult M3434 (Stem Cell Technologies, Canada) and after 7-10 days of incubation different colony types were scored.

**Retroviral transduction and transformation (RTTA) assays**
RTTA were performed as previously described with some modifications (Zeisig & So, 2009). The MSCV-MLL-ENL construct has been described previously. The two independent shRNAs against murine Prmt1 have been reported previously (Cheung et al., 2007). Briefly, isolated c-Kit+, LSK or GMP cells were cultured overnight in RPMI+10%FBS supplemented with 20ng/ml SCF, 10ng/ml IL3 and 10ng/ml IL6 prior to viral transduction with virus particles carrying MLL-ENL by centrifugation at 800xg at 32°C for 2hr. Cells were plated in M3234 Methylcellulose medium supplemented with 20ng/ml SCF, 10ng/ml of each IL3, IL6 and GM-CSF and appropriate antibiotic selection on the following day. Colonies were scored after 7 days and replated every 7 days. To induce the deletion of Ctnnb1, 20nM tamoxifen (Sigma, US) was added to the Methylcellulose medium in the second round of plating and YFP positive were sorted after the second round of plating using a BD FACS ARIA and plated into the 3rd round. After the 4th round of plating, cells were cultured in R20/20 to establish cell lines as previously described (Yeung et al., 2010). Differences in colony numbers were statistically tested using a two-tailed t-test in Excel.

**Phenotypic analysis**

Immunophenotypic analysis was performed by FACS using fluorochrome-conjugated monoclonal antibodies to murine c-Kit (2B8 clone), Mac-1 (M1/70) and Gr-1 (RB6-8C5) (eBiosciences). Staining was generally performed on ice for 15mins and washed once before analysis using a BD LSR II system. Differences in surface marker expression were statistically tested using a two-tailed t-test in Excel.

**Western blot**

Cell lysates from primary transformed cells or sorted leukemic cells were isolated and subjected to western blot as described (Yeung et al., 2010).

**Histone extraction and detection**

Histone proteins were prepared by acid extraction (Abcam protocol, http://www.abcam.com/protocols/histone-extraction-protocol-for-western-blot). Briefly, cells were lysed for 10 minutes on ice in TEB buffer (PBS/0.5%Triton X 100/protease inhibitors) at a cell density of 10^7 per ml. The nuclei were harvested by centrifugation, and washed in half the volume of TEB. Pellets were resuspended in 0.2N HCl at a density of 4x10^7 nuclei per ml and histones were extracted overnight at 4°C. Samples were centrifuged to pellet debris and the supernatant containing the histones were transferred into a new Eppendorf tube. 10ul histones were mixed with 40ul PBS and 50ul 2xSDS loading buffer and incubated at 95°C for 8minutes. 25ul of the denatured histones were subjected to western blot using a 10% Next gel (AMRESCO LLC). Antibodies against H4R3me2a (Active Motif) and Histone H3 (Abcam) were used.

**Genotyping PCR**

Genomic DNA was isolated and β-catenin genotyping PCR was performed as described previously (Brault et al., 2001). Primer sequences for Hoxa9 are available on request. Differences in gene expression were statistically tested using a two-tailed t-test in Microsoft Excel.
**q-RT-PCR**

q-RT-PCR was performed on StepOne qPCR machine (Applied Biosystems) using Taqman or SYBR green chemistries. See Appendix Table S1 for the primer sequences used throughout the paper for validation.

**Apoptosis and cell cycle analysis**

MLL-ENL transduced Hoxa9^{-/-}Ctnnb1^{fl/fl} cells were treated for 72 hours with or without tamoxifen and stained with AnnexinV for apoptosis. Cell cycle was analyzed using the BRDU Flow kit (BD Pharmingen) according to manufacturer’s instructions. Cells were analyzed using a BD LSR II system (BD, USA) and differences were statistically tested using a two-tailed t-test in Excel.

**RNA sequencing**

300ng to 1ug of mirVANA (Ambion) isolated total RNA was used for RNA-Seq library preparation using TruSeq Stranded Total RNA kit (Illumina) and sequenced on Hiseq2000 platform (Illumina) as per manufacturers recommendations. All samples are listed in Dataset EV1A.

**Whole genome sequencing**

Genomic DNA was isolated with the QIAamp DNA Micro kit (Qiagen) according to the manufacturer’s instructions, and 30ng of genomic DNA was used as input for each library preparation. Whole-genome sequencing libraries were generated using the tagmentation method as previously described (Wang, Gu et al., 2013), but without bisulfite treatment and with minor modifications. Briefly, genomic DNA was subjected to tagmentation with a hyperactive Tn5 transposase (Epizyme), which fragmented the DNA and appended sequencing adaptors in a single step. After PCR amplification of libraries, DNA fragments of 200-800bp were double-side selected using SPRI Ampure XP beads, with left-right ratios of 1.5-0.55. Purified libraries were subjected to 125-bp paired-end sequencing on an Illumina HiSeq2500 machine.

**Bioinformatic analyses**

**Mapping and read counts**

The FASTQ files were de-tagged, and the quality of the FASTQ files inspected using FastQC v.0.11.2 (Andrews, 2015). Remaining adapters were trimmed using TRIMGalore! v.0.3.7 (Kreuger, 2015). Whole genome sequencing reads were mapped to the Ensembl mouse genome GRCm38 (Aken, Ayling et al., 2016) using Bowtie2 v.2.2.5 (Langmead & Salzberg, 2012). RNA-Seq reads were mapped to the Ensembl mouse genome using TopHat2 v2.0.13 (Kim, Pertea et al., 2013), reads were filtered for quality and counted using samtools v.0.1.18 and bedtools v2.23.0-10-g447cb97 (Li, Handsaker et al., 2009, Quinlan & Hall, 2010) (Dataset EV1B).
SNP calling
SNP calling (for both RNA-Seq and WGS) was performed using samtools v.0.1.18 mpileup and bcftools v.1.2 (using htslib v.1.2.1) (Li, 2011), SNPs were leniently filtered only for quality $\geq$Q30, and custom scripts used to produce summary stats (Dataset EV1C). To summarize the regions in which SNPs fell, variant call files and Ensembl GTF annotation (Mus_musculus.GRCm38.82.gtf) were converted to bed format using custom scripts and compared using bedtools intersect v2.23.0. Bedtools complement v2.23.0 (Quinlan & Hall, 2010) was used to extract intronic and intergenic coordinates.

Gene differential expression
Differential expression was determined either using DESeq2 v.1.10.1 (Love, Huber et al., 2014), negative binomial GLM fitting and Wald statistics; or with limma v.3.26.9 (Ritchie, Phipson et al., 2015) using voom to normalize read counts, and eBayes to determine differential expression (as detailed in Expanded View Datasets). Functional enrichment analysis was conducted using the GSEA software, using various Molecular Signature Databases (c2 set version 5.0, version 6.0 and all_gene_sets) (MSigDB) (Subramanian, Tamayo et al., 2005) appended with a custom made gene set for LSC stem cell maintenance (Somervaille et al., 2009) (Dataset EV3B), comparing log2-fold changes in gene expression as the ranking metric. For the GSEA analysis, human-mouse gene orthologues were identified using MGI list of orthologous genes (Blake, Bult et al., 2014), and Ensembl bioMart used to transfer MGI IDs to Ensembl gene IDs (Kinsella, Kahari et al., 2011). In order to find genes which showed a synergistic effect of simultaneous knockout of Cbnt1 and Hoxa9, differential expression was tested with DESeq2 Negative Binomial GLM fitting and Wald statistics. The data was subjected to five DE analyses, which served to categorize the genes into 78 different classes. These comparisons are 1. LSK WT to Ctnnb1 KO, 2. LSK WT to HoxA9 KO, 3. Additive effect LSK WT to Ctnnb1 KO and Hoxa9 KO, 4. LSK Ctnnb1 KO to Ctnnb1:HoxA9 interaction, 5. LSK HoxA9 KO to Ctnnb1:HoxA9 interaction. For overlapping significantly differentially expressed genes, heatmaps were plotted using the function heatmap.2 from the R-package gplots v3.0.1 (Warnes, Bolker et al., 2016) with Z-score scaling of rows and/or columns.

Survival analysis
Statistical analysis and data visualization was performed using R (R Core Team, 2014). Survival and Cox proportional hazards ratio analysis was conducted using R-packages survival v.2.39.5 (Therneau, 2015) and survcomp v.1.20.0 (Schoeder, Culhane et al., 2011) on all patients which had complete survival and expression data from the following datasets; GSE1159 (n=293), GSE12417; GPL570 (n=79), GPL96 (n=163), GSE14468 + GSE6891 (n=602), GSE5122 (n=58), GSE8970 (n=34) available from the Gene Expression Omnibus database ncbi.nlm.nih.gov/geo/ and The Cancer Genome Atlas AML dataset (n=183) (Cancer Genome Atlas Research, 2013). Summary data for patient cohorts are presented in Dataset EV2D. Samples were normalized using the GENENORM algorithm from the R-package inSilicoMerging v.1.15.0 (Taminau, Meganck et al., 2012) with prior DESeq2 Vst transformation of RNA-Seq samples used in this study to derive the cell-of-origin specific signature (Love et al., 2014). All probe intensities were translated into human genes and averaged prior to merging. MLL patients in the datasets were identified from associated karyotyping and PCR diagnosis.
Data availability

All WGS has been submitted to the ENA study PRJEB14461. RNA-Seq is available on ArrayExpress E-MTAB-3647.

Statistical analysis

All the experimental results were analysed using two-tailed Student’s t-test, χ² test, hypergeometric test, as indicated in the figure legends and Expanded View Datasets. Groups that were statistically compared shared a similar variance, as shown in the figures. P<0.05 was considered as statistically significant. For GSEA analysis, FDR-q values <0.25 were considered as statistically significant (Subramanian et al., 2005). The log-rank test was used to compare survival curves.

Acknowledgements

We would like to thank Sam Tung, Jayant Rane for assistance on bioinformatics analyses; Erica Tse for graphical assistance; Peter Parker and Ryan Driskell for critical advice on the manuscript. This work was supported mainly by a Cancer Research UK (CRUK) program grant A16753 and in part by a Bloodwise program grant to CWE So.

Author Contributions

TS, BBZ, TKF, MM, CTT, PL, PV, JB and AW performed experiments, BBZ contributed to writing the manuscript, MZ, QCH, CL and BL contributed the bioinformatics analyses and editing the manuscript, CWES contributed to overall research design, direction and writing of the manuscript.

Conflict of interest

The authors have declared that no conflict of interest exists.

References


myeloid leukemia stem cells employs a transcriptional program shared with embryonic rather than adult stem cells. Cell Stem Cell 4: 129-140


Therneau T (2015) A Package for Survival Analysis in S. In http://cran.r-project.org/package=survival


Figure 1. MLL-ENL leukemic stem cells derived from LSK or GMP populations have contrasting functional requirements of β-catenin for their initiation and maintenance of disease. (A) Schematic overview of the experimental procedures. Keys and colour codes in the legend box indicate the cells-of-origin and the β-catenin status of MLL-ENL transduced cells in the following experiments (B-K). (B) Colony numbers in serial replating assay of the different MLL-ENL transduced cells. Data are represented as mean ± SD (C, G) PCR validation of Ctnnb1 deletion on genomic DNA isolated from the indicated MLL-ENL transduced GMP and LSK cells (C) or leukemic cells (G). L, 100bp ladder; W, wild type control, F, Ctnnb1 floxed allele, D,
Ctnnb1 deleted allele, N, negative control. (D, H) Cell lysates from indicated MLL-ENL transduced GMP and LSK cells after the fourth round of plating (D) or from indicated leukemic mice (H) were blotted with anti-β-catenin (top) and anti-actin (bottom) antibodies. (E, F) Kaplan-Meier survival curves of indicated MLL-ENL transduced cells transplanted into primary recipient (solid lines) and secondary recipient mice (dotted lines). N=10 mice per group were used in primary transplants (solid lines) in (E). N=15 mice were used for Ctnnb1fl/fl (blue solid line) and n=10 mice were used for Ctnnb1del/del (green solid line) in (F). For all secondary transplants (dotted lines) n=5 mice were used per group. (I) Percentage of CD45.2+ donor cells in the bone marrow of recipient mice at the indicated time points post-transplantation of the indicated GMP-MLL-ENL and LSK-MLL-ENL transduced cells. Data are represented as mean ± SD. N=4, two-tailed t-test was performed. (J-K) Kaplan-Meier survival curves of secondary transplanted GMP-MLL-ENL (J) and LSK-MLL-ENL (K) primary leukemia bone marrow cells treated with DMSO as floxed controls or tamoxifen for β-catenin deletion prior to secondary transplantation (n=3 per group). See also Appendix figure S1.
Figure 2. Cells-of-origin transcriptional-memory predicts survival in AML patients. (A-B) Number of identified genomic variants in indicated MLL-ENL transformed cells using RNA-Seq (A) and genomic sequencing (B). A two-tailed t-test was performed in (A). LSK, wildtype LSK cells; GMP, wildtype GMP cells. (C, D) Manhattan plots indicating estimated length of CNVs (C) or number of genes in 28

Average gene expression (reads/gene, log-scale), all genes n=43,632

<table>
<thead>
<tr>
<th>LSK vs GMP</th>
<th>LSK vs GMP (MLL-ENL)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>UP</strong></td>
<td>1525</td>
</tr>
<tr>
<td><strong>Down</strong></td>
<td>1111</td>
</tr>
<tr>
<td>p&lt;8e-16</td>
<td>1223</td>
</tr>
<tr>
<td>p&lt;3e-13</td>
<td>1254</td>
</tr>
</tbody>
</table>

Survival %

n=1290  
GMP-like  
LSK-like  
p<0.0014  
Overall survival time (months)

28
CNV areas (D) on the y-axis in the respective chromosomal positions shown in the x-axis. (E) MA-plots showing the log2-fold gene expression changes in the normal (left panel) and MLL-ENL transformed (right panel) cells as indicated. (F) Transcriptional memory signature; the overlap of differentially expressed genes in GMP-LSK in normal vs MLL-ENL transformed cells is significantly enriched using a hypergeometric test. (G) Survival differences between patients clustered using transcriptional memory signatures with a log-rank test. See also Appendix figure S2.
Figure 3. Key transcriptional-memory gene *Hoxa9* may help to overcome β-catenin dependent transformation in LSK-derived MLL-CSCs. (A) Gene set enrichment analysis (GSEA) shows “Targets of Hoxa9/Meis1, down” (Hess et al., 2006) for the indicated comparison. (B) Log10-fold FDR q-values of the indicated gene sets positively enriched in β-catenin deleted LSK-MLL-ENL compared to β-catenin deleted GMP-MLL-ENL. (C-E) RNA-seq log2-fold change of key self-renewal genes (C), RT-qPCR validation of *Hox/Meis1* expression represented as mean ± SD of 3 independent experiments. Two-tailed t-test was performed (D), and GSEA showing “Hoxa9_dn.v1_up” (Faber, Krivtsov et al., 2009) for the β-catenin deleted LSK-MLL-ENL to β-catenin deleted GMP-MLL-ENL comparison (E). (F) Colony numbers in serial replating assay of indicated MLL-ENL transduced cells. Data are represented as mean ± SD. (G) Kaplan-Meier survival curve of mice transplanted with Hoxa9-/- (n=18) or WT LSK-MLL-ENL (n=13) transformed cells (solid lines) and secondary recipient mice (n=5 for Hoxa9-/- and n=4 for wt, dotted lines) as indicated. Comparisons between Hoxa9 WT and Hoxa9/-/- were not significantly different (ns). See also Appendix figure S3.
Figure 4. Deletion of both β-catenin and Hoxa9 abrogates leukemia development in LSK-MLL-ENL cells. Keys and colour code in the left top corner indicate the origin and the β-catenin and Hoxa9 status of MLL-ENL transduced cells. (A) Colony numbers in serial replating assay of MLL-ENL transduced cells. Images of typical 4th round colonies shown above. Data are represented as mean ± SD. (B) Summary of immunophenotypic analysis (Figure S4D) of LSK-MLL-ENL Hoxa9-/-Ctnnb1fl/fl with or without 72h Tamoxifen treatment. Data are represented as mean ± SD. N=3, two-tailed t-test was performed. (C-E) % of apoptotic cells (AnnexinV assay) (E), cells in the indicated cell cycle phases (BrdU assay) (F) and CD45.2+ donor cells in the bone marrow of recipient mice at the indicated time points post-transplantation (G) of LSK-MLL-ENL transduced cells. Data are represented as mean ± SD. N=4, two-tailed t-test was performed. (F) Kaplan-Meier survival curve of mice transplanted with LSK-MLL-ENL carrying different Hoxa9 and Ctnnb1 genotypes. N=5 per genotype was used. (G) Summary of the LSC frequencies obtained from the in vivo limiting dilution experiments (Figure S4E-H) using the LSK-MLL-ENL leukemic cells with different genotypes. See also Appendix figure S4.
Figure 5. β-catenin and Hoxa9 co-regulate Prmt1 in LSK-MLL-ENL cells. (A, B) Venn diagram showing the overlap of 33 commonly enriched upregulated (A) and 38 commonly enriched downregulated (B) highly significant (FDR<0.05) gene sets between indicated comparisons (i.e., β-catenin inactivation in Hoxa9⁻/⁻ LSK-MLL-ENL, Hoxa9 inactivation in Ctnnb1⁻/⁻ LSK-MLL-ENL cells), including “Hoxa9/Meis1_DN” (Hess et al., 2006), “LSC_maintenance_down” (Somervaille et al., 2009), “Hoxa9/Meis1_UP” (Hess et al., 2006) and “LSC_maintenance_up” (Somervaille et al., 2009) as indicated. (C) A small set of genes is synergistically regulated by β-catenin and Hoxa9 (the effect of knockout of both genes is smaller or larger than could have been predicted from their single knockouts). Example sets of genes synergistic up-regulated (top), down-regulated (bottom) and non-significantly regulated (bottom right) are shown (all sets are in Figure S5). Keys and colour code in the top corner indicate the origin and the β-catenin and Hoxa9 status of MLL-ENL transduced cells. (D) Pantherdb protein classes (http://pantherdb.org/) are shown for the 321 synergistic up and 204 synergistic down genes as indicated. P-value was obtained after Bonferroni correction. (E) The FDR qvalues of the top5 Toppgene GO:Molecular function (https://toppgene.cchmc.org/enrichment.jsp) are shown for the synergistic up and downregulated genes as indicated. (F) Statistically significant directionality in the overlap of the synergistic genes with the Hox/Meis1_DN gene set (Dataset EV5B) including Prmt1, which has also been independently validated by RT-qPCR (right panel). See also Appendix figure S5.
Figure 6: Prmt1 regulates similar and overlapping transcriptional programs mediated by β-catenin in LSK-MLL-ENL cells. (A) Heatmap analysis and Venn diagram showing the commonly regulated genes after the loss of function of Prmt1 and β-catenin in Hoxa9−/− LSK-MLL-ENL cells. (B) The FDR q-values of the top GO:Molecular function (top panel) and GO:Biological process (bottom panel) (https://toppgene.cchmc.org/enrichment.jsp) for the indicated up- and down-regulated genes is shown. (C) Venn diagrams showing the total, positively and negatively regulated genesets identified by GSEA upon the loss of function of Prmt1 and β-catenin in Hoxa9−/− LSK-MLL-ENL cells. (D) Examples of stem cell and stemness signatures which were commonly higher expressed in the single knockout Hoxa9−/− LSK-MLL-ENL cells compared to Prmt1 down-regulated or β-catenin inactivated Hoxa9−/− LSK-MLL-ENL cells.
Figure 7. Suppression of Prmt1 abolishes β-catenin or Hoxa9 independent transformation in HSC-derived MLL-CSCs. (A) The relative number of colonies from Hoxa9^-/- LSK-MLL-ENL leukemic cells with empty vector or shPRMT1. Data are represented as mean ± SD. (B) The % of CD45.2+ donor cells in the bone marrow of recipient mice at the indicated time points post-transplantation of Hoxa9^-/- LSK-MLL-ENL leukemic cells with empty vector or shPrmt1. Data are represented as mean ± SD. (C) Kaplan-Meier survival curve of secondary recipient mice transplanted with Hoxa9^-/- LSK-MLL-ENL leukemic cells with vector control or shPrmt1. (D) The relative number of colonies from Ctnnb1^-/- LSK-MLL-ENL leukemic cells with empty vector or shPRMT1. Data are represented as mean ± SD. (E) The % of CD45.2+ donor cells in the bone marrow of recipient mice at the indicated time points post-transplantation of Ctnnb1^-/- LSK-MLL-ENL leukemic cells with empty vector or shPrmt1. Data are represented as mean ± SD. (F) Kaplan-Meier survival curve of secondary recipient mice transplanted with Ctnnb1^-/- LSK-MLL-ENL leukemic cells with vector control or shPrmt1. See also Appendix figure S6.