

Crystal structure of the N-terminal region of human Ash2L shows a winged-helix motif involved in DNA binding

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Ash2L is a core component of the MLL family histone methyltransferases and has an important role in regulating the methylation of histone H3 on lysine 4. Here, we report the crystal structure of the N-terminal domain of Ash2L and reveal a new function of Ash2L. The structure shows that Ash2L contains an atypical PHD finger that does not have histone tail-binding activity. Unexpectedly, the structure shows a previously unrecognized winged-helix motif that directly binds to DNA. The DNA-binding-deficient mutants of Ash2L reduced Ash2L localization to the *HOX* locus. Strikingly, a single mutation in Ash2L_{WH} (K131A) breaks the chromatin domain boundary, suggesting that Ash2L also has a role in chromosome demarcation.

Keywords: Ash2L; histone methyltransferase; MLL; transcription; winged helix motif

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INTRODUCTION

Histone H3 lysine 4 (H3K4) methylation is a conserved epigenetic mark that is correlated with active states for gene expression in eukaryotes (Ruthenburg *et al*, 2007). The multi-subunit Set1 complex is the only histone methyltransferase (HMT) complex that modifies H3K4 in yeast (Briggs *et al*, 2001; Roguev *et al*, 2001). By contrast, mammals contain six Set1-related complexes, which

are collectively referred to as the MLL family HMTs (Malik & Bhaumik, 2010). The catalytic subunits of these complexes, MLL1–4, SET1A and SET1B, share an evolutionarily conserved SET domain, which catalyses the methylation of H3K4 (Ruthenburg *et al*, 2007). In contrast to most other SET-domain-containing HMTs, these MLL family proteins by themselves have low methyltransferase activity (Roguev *et al*, 2001). Their full activities are achieved only in the presence of three common components of the complexes, WDR5, RbBP5 and Ash2L (Dou *et al*, 2006).

Ash2L is a core component of all MLL complexes. Sequence analysis showed two motifs in Ash2L: an amino-terminal plant homeodomain (PHD) finger and a carboxy-terminal SPRY (SPLa/Ryanodine) domain (Fig 1A; Ikegawa *et al*, 1999). Our recent studies show that the SPRY domain of Ash2L mediates the interaction with RbBP5 and has an important role in regulating the methyltransferase activity of MLL complexes (Y. Chen *et al*, unpublished results). PHD finger—found in many eukaryotic chromatin-associated proteins (Bienz, 2006)—is a structural module that recognizes methylated or unmethylated histone tails (Ruthenburg *et al*, 2007).

Here, we present the crystal structure of the N-terminal domain of human Ash2L. The structure shows that Ash2L contains an atypical PHD finger that does not have histone tail-binding activity. Unexpectedly, the structure reveals a previously unrecognized winged-helix motif immediately C-terminal to the PHD finger. The structure, in conjunction with biochemical and cellular analyses, provides new insights into the mechanism by which the winged-helix motif of Ash2L binds to DNA and targets Ash2L to the *HOX* locus.

RESULTS AND DISCUSSION

Overall structure of the Ash2L N-terminal domain

The sequence alignment of many Ash2L proteins showed a conserved region at the amino-terminus, Ash2L_{NTD} (N-terminal domain; residues 1–177), which includes the previously identified

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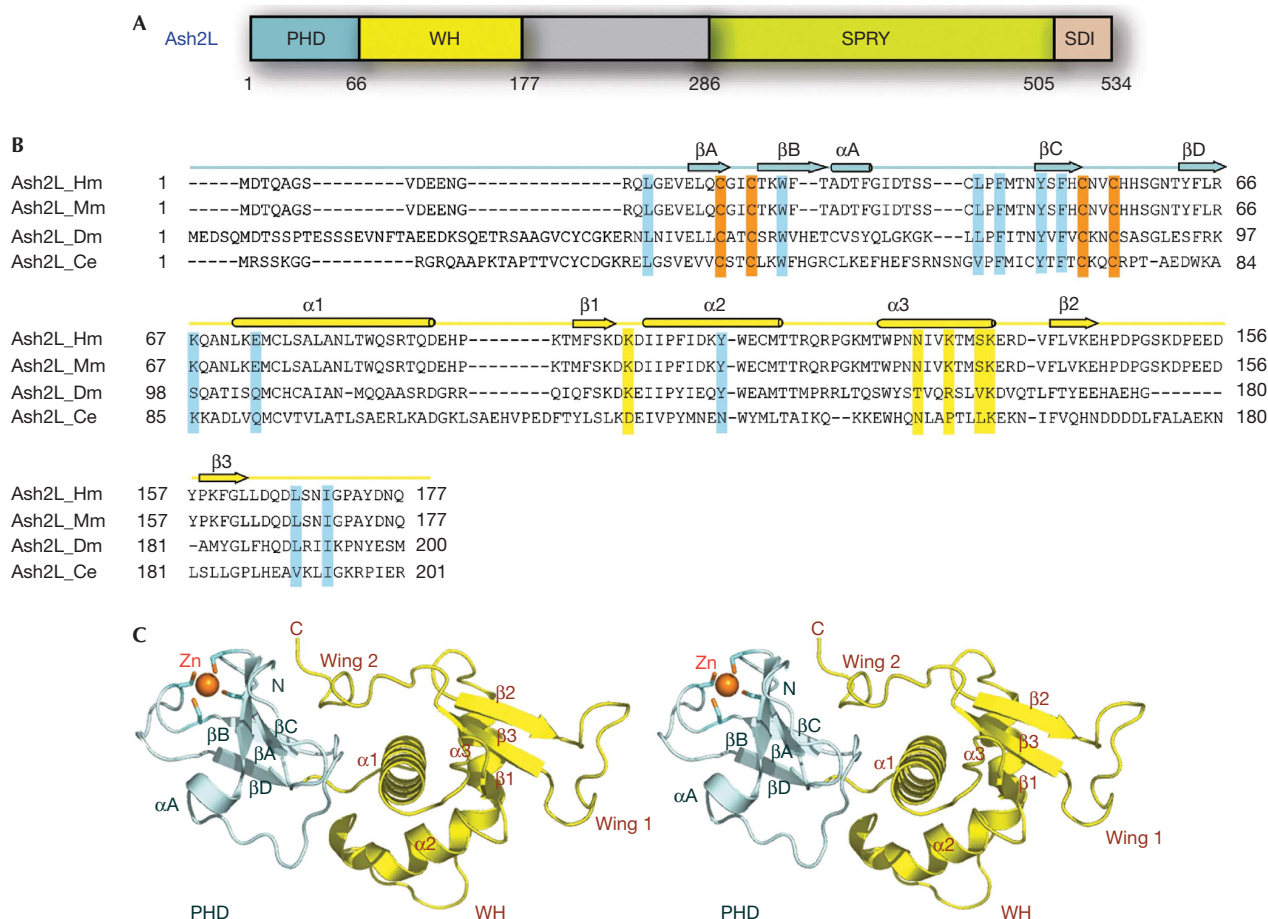


Fig 1 | Crystal structure of the N-terminal domain of Ash2L. (A) Domain organization of Ash2L. The PHD finger is coloured in pale cyan, the winged helix (WH) motif in yellow, the SPRY domain in green and the SDI (Sdc1/DPY30 interaction) motif in pink. (B) Sequence alignment of the N-terminal regions of four Ash2L family members. Secondary structure assignments based on the human Ash2L crystal structure are shown as coloured cylinders (α -helices) and arrows (β -strands) above the sequences. The cysteine residues involved in zinc binding are shown in orange. The putative residues involved in DNA binding are highlighted in yellow, and the residues important for the interaction between the PHD and WH motifs are highlighted in cyan. (C) Stereoview of Ash2L_{NTD}. The PHD finger is coloured in pale cyan and the WH domain in yellow. The zinc ion in sphere representation and the zinc-binding cysteine residues are shown in stick models. Zn, zinc.

PHD finger (Fig 1B; supplementary Fig S1 online). To understand the molecular structure of Ash2L_{NTD}, we determined its crystal structure at 2.1 Å resolution (supplementary Table S1 online). The structure shows that Ash2L_{NTD} has an elongated conformation and is composed of two structurally distinct motifs (Fig 1C). As expected from previous sequence analyses (Ikegawa *et al*, 1999), the N-terminal motif in Ash2L_{NTD} (residues 13–66) is comprised of a PHD finger, consisting of two small anti-parallel β -sheets that are separated by a long, extended loop (Fig 1C). One zinc ion is coordinated by four cysteine residues from the loops between strands in both β -sheets (Fig 1C).

The C-terminal portion of Ash2L_{NTD} (residues 67–174) adopts a compact fold featuring three α -helices and a curved three-stranded β -sheet (Fig 1C). Although sequence analysis failed to identify any known protein motifs in this region, an unbiased search of the database showed structural resemblance of Ash2L_{67–174} with more than 30 winged-helix motifs. Each of

these winged-helix motifs can be superimposed onto Ash2L_{67–174} with a root-mean-square deviation of approximately 3.0 Å in the positions of over 65 equivalent C α atoms. Therefore, we will refer to Ash2L_{67–174} as Ash2L_{WH} (Fig 1A).

The PHD and winged-helix motifs of Ash2L are associated with each other by both van der Waals and electrostatic contacts, enclosing a total surface area of 1,742 Å² (supplementary Fig S2 online). These interactions fix the relative orientation between the PHD and winged-helix motifs and allow Ash2L_{NTD} to adopt a compact structure resembling a single folded unit. Our efforts to prepare Ash2L_{PHD} alone yielded aggregated products (data not shown), suggesting that it requires an interface with the winged-helix motif for stability.

Ash2L has an atypical PHD finger

PHD finger is a histone mark reader; it uses an aromatic cage formed by three conserved aromatic residues to recognize

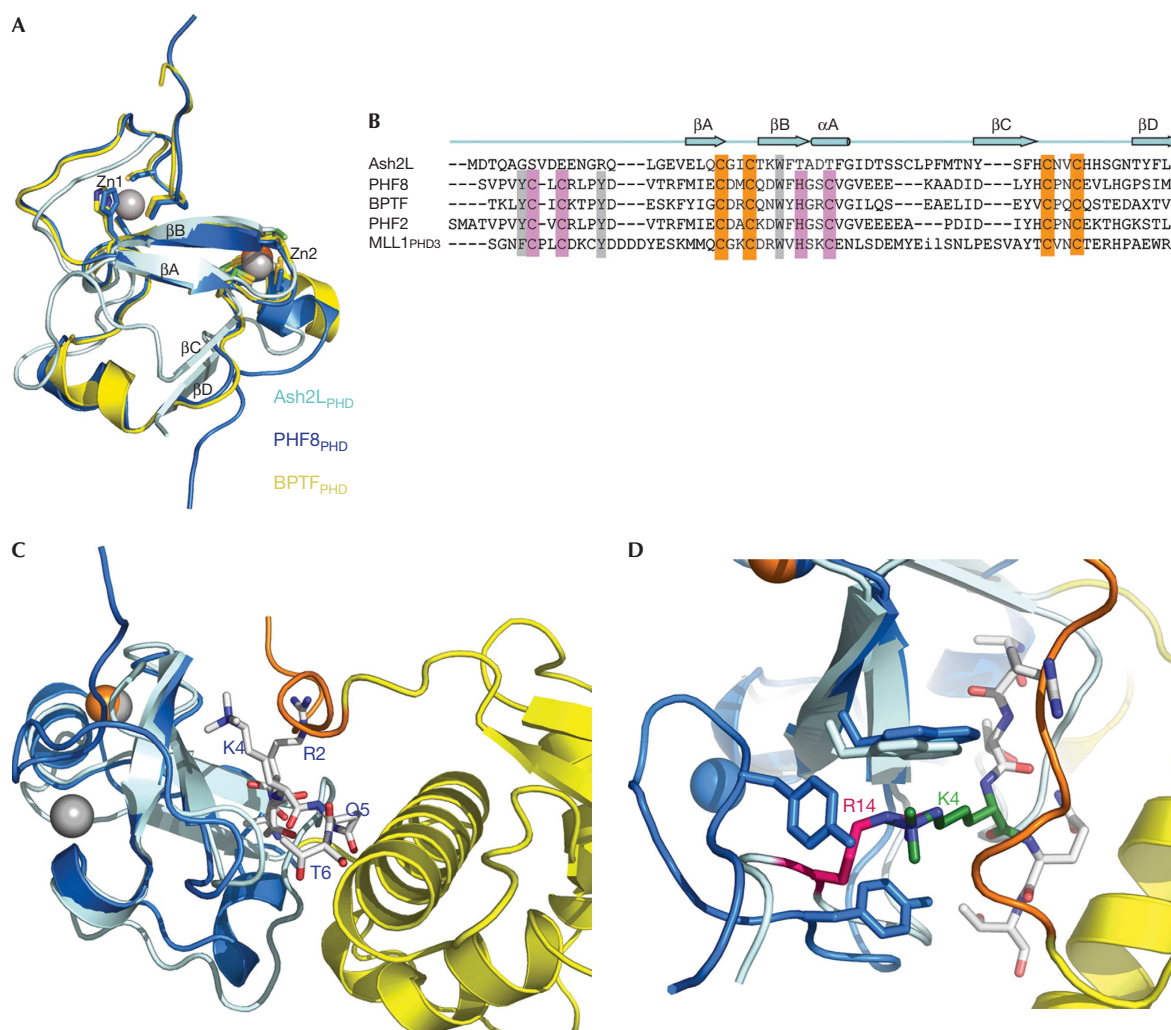


Fig 2 | Ash2L_{PHD} is an atypical PHD finger. (A) Superimposition of Ash2L_{PHD} (cyan), PHF8_{PHD} (marine) and BPTF_{PHD} (yellow). There are two zinc ions in the PHF8_{PHD} structure (grey), but only one zinc ion in Ash2L_{PHD} (orange). (B) Structure-based sequence alignment of Ash2L_{PHD} with four PHD fingers that share high structural similarity. The conserved zinc-binding residues are highlighted in orange, the absent zinc-binding residues in pink and the absent aromatic cage residues in grey. (C) Structural superimposition of Ash2L_{PHD} (cyan) and the PHF8_{PHD}-H3 peptide complex (marine). H3 peptide is shown in sticks (grey). The carboxy-terminal wing 2 from Ash2L_{WH} is shown in orange. (D) The R14 of Ash2L (magenta) and the K4 of H3 (green) occupy the same pocket surrounded by aromatic residues. WH, winged helix; Zn, zinc.

methylated or unmethylated lysines (Ruthenburg *et al*, 2007). Ash2L_{PHD} is most similar to the PHD finger in PHF8 (a Jumonji-domain-containing lysine demethylase), which has a root-mean-square deviation value of 1.4 Å (Fig 2A; Horton *et al*, 2010). Despite this overall structural similarity, several structural features of Ash2L_{PHD} are different to PHF8_{PHD}. First, Ash2L_{PHD} lacks half of the conserved C4HC3 signature that mediates the coordination of two zinc ions in PHF8_{PHD} (Fig 2B). Therefore, Ash2L_{PHD} binds to only one zinc ion through the second half of the motif (Figs 1C, 2C). Second, two of the three aromatic residues that form the methyl-lysine recognition cage in PHF8_{PHD} are absent in Ash2L_{PHD} (Fig 2B). When the structure of Ash2L_{PHD} is superimposed with that of PHF8_{PHD} complexed with H3K4me3-containing peptide (Horton *et al*, 2010), the guanidinium group of Ash2L_{PHD} Arg 14 and the trimethyl group of H3K4me3 from the

peptide occupy the same binding site on Ash2L_{PHD} (Fig 2D; supplementary Fig S3A online). Third, Ash2L_{PHD} contains a long, extended loop between strands βB and βC, which precludes Ash2L_{PHD} from binding of any peptide to a site similar to that observed in the PHF8 structure (Fig 2C; supplementary Fig S3B online). These analyses suggest that Ash2L_{PHD} is an atypical PHD finger and by itself lacks the structural features for binding to histone tails. Nevertheless, it is possible that other factors could induce a conformational change in Ash2L to allow histone binding.

Ash2L_{WH} is a DNA-binding motif

The crystal structure of Ash2L_{NTD} shows an unexpected winged-helix motif that could not have been predicted by bioinformatic approaches. The structure of Ash2L_{WH} resembles the winged-helix

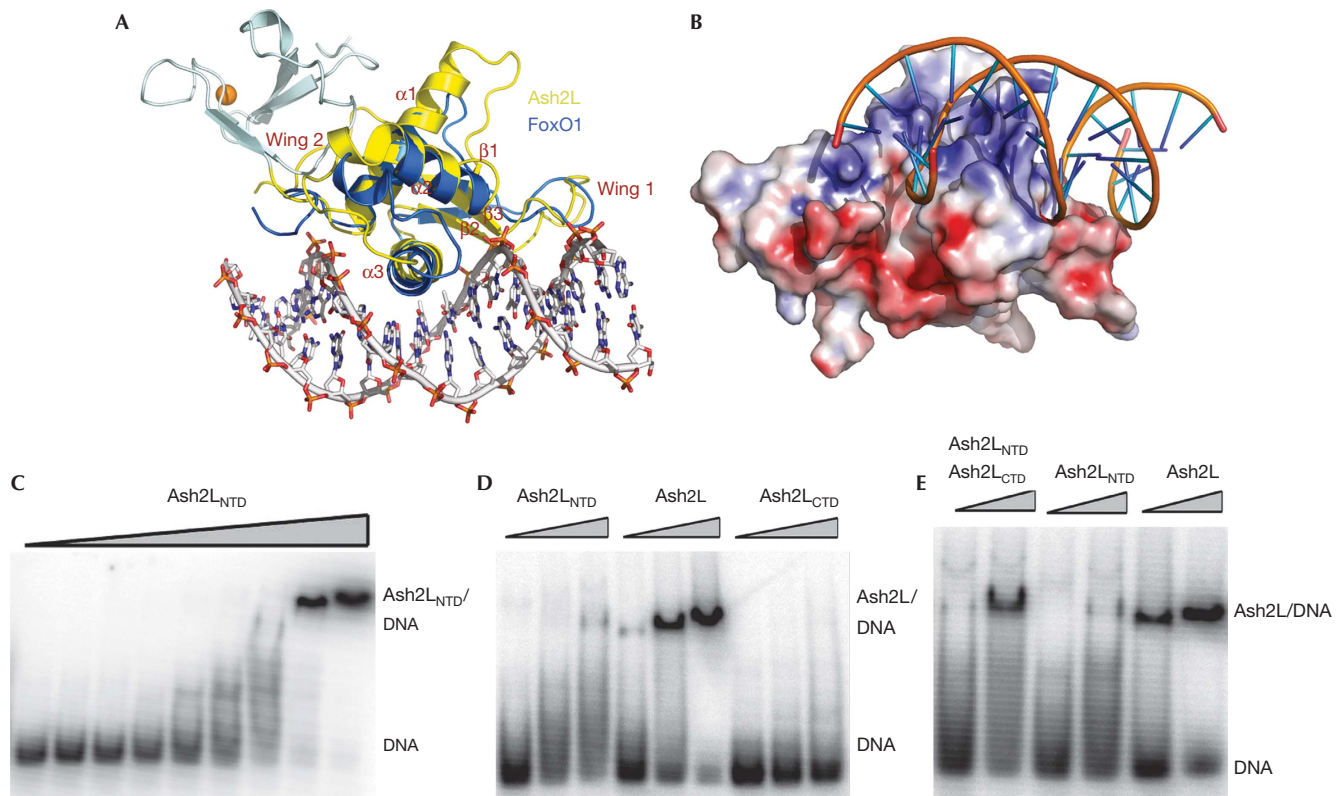


Fig 3 | Ash2L_{WH} is a DNA-binding motif. (A) Overlay of the structures of Ash2L_{NTD} and the FoxO1–DNA complex. Ash2L_{PHD} is shown in cyan, Ash2L_{WH} in yellow and FoxO1 in marine. (B) The surface area of helix $\alpha 3$ of Ash2L_{WH} is highly positively charged and thus compatible with DNA binding. Ash2L_{NTD} is in surface representation and coloured according to its electrostatic potential (positive, blue; negative, red). The modelled DNA structure is from the superimposed FoxO1–DNA complex structure. (C) Ash2L_{NTD} binds to duplex DNA. A concentration of 0.1 μM DNA was incubated with increasing amounts of Ash2L_{NTD} (0.1–25.6 μM). (D) Full-length Ash2L binds with higher affinity to DNA than Ash2L_{NTD}. Three different protein concentrations (1.6, 3.2 and 6.4 μM) were incubated with 0.1 μM DNA. (E) The carboxy-terminal domain of Ash2L enhances Ash2L_{NTD} binding to DNA *in trans*. NTD, N-terminal domain; WH, winged helix.

motif of FoxO1, a forkhead box transcription factor (Brent *et al*, 2008; Fig 3A). The canonical winged-helix motif is a compact α/β -structure consisting of three α -helices, three β -strands and two large loops or wings. Wing 1 connects strands $\beta 2$ and $\beta 3$, whereas wing 2 extends from strand $\beta 3$ to the C-terminus of the winged-helix motif. Similarly to in FoxO1, winged-helix motifs often function as DNA-binding modules and are involved in transcription regulation (Kaestner *et al*, 2000).

The crystal structure of FoxO1 bound to a duplex DNA shows that the DNA recognition helix of the winged-helix motif, $\alpha 3$, is presented to the major groove of the DNA, whereas the two wings also contribute to DNA-binding affinity and specificity (Brent *et al*, 2008). When the structures of Ash2L_{NTD} and the FoxO1–DNA complex are overlaid on the basis of the winged-helix motifs, the $\alpha 3$ helix of Ash2L_{WH} fits into the major groove of the DNA, whereas Ash2L_{PHD} locates at the opposite side of the molecule and presumably does not interfere with the interaction between Ash2L_{WH} and DNA (Fig 3A). The fact that the putative DNA-binding site on helix $\alpha 3$ is highly positively charged, further suggests that Ash2L_{WH} is a DNA-binding module (Fig 3B).

The structural similarities between Ash2L_{WH} and the winged-helix motif of FoxO1 prompted us to ask whether Ash2L_{WH} uses

the same mechanism to bind to DNA. To test this hypothesis, we investigated the DNA-binding activity of Ash2L_{NTD} using electrophoresis mobility shift assay (EMSA). A 60-base-pair oligonucleotide with random sequences in the middle 20 base pairs (N20) was incubated with increasing amounts of Ash2L_{NTD}, and binding was analysed by EMSA. Ash2L_{NTD} bound to the DNA with an equilibrium dissociation constant K_d of 12 μM , which is approximately 1,000 times weaker than the FoxO1–DNA interaction (Fig 3C; Brent *et al*, 2008). This relatively low DNA-binding affinity can be explained by the structural difference between Ash2L and FoxO1. Wing 2 in FoxO1 has an important role in DNA binding, as deletion of this loop abolished the interaction between FoxO1 and DNA (Brent *et al*, 2008). By contrast, wing 2 of Ash2L folds back onto the N-terminal PHD finger and thus does not contribute to DNA binding (Fig 3A). The DNA-binding affinity of Ash2L_{WH} is comparable with that of another forkhead box transcription factor, FoxM1, which also lacks the regular winged loops (Littler *et al*, 2010). Next, we examined whether Ash2L_{WH} binds to DNA in a sequence-specific manner. Several rounds of *in vitro* selection using N20 yielded no obvious consensus DNA-binding sequence (data not shown), suggesting that Ash2L_{WH} probably associates with DNA without strong sequence specificity.

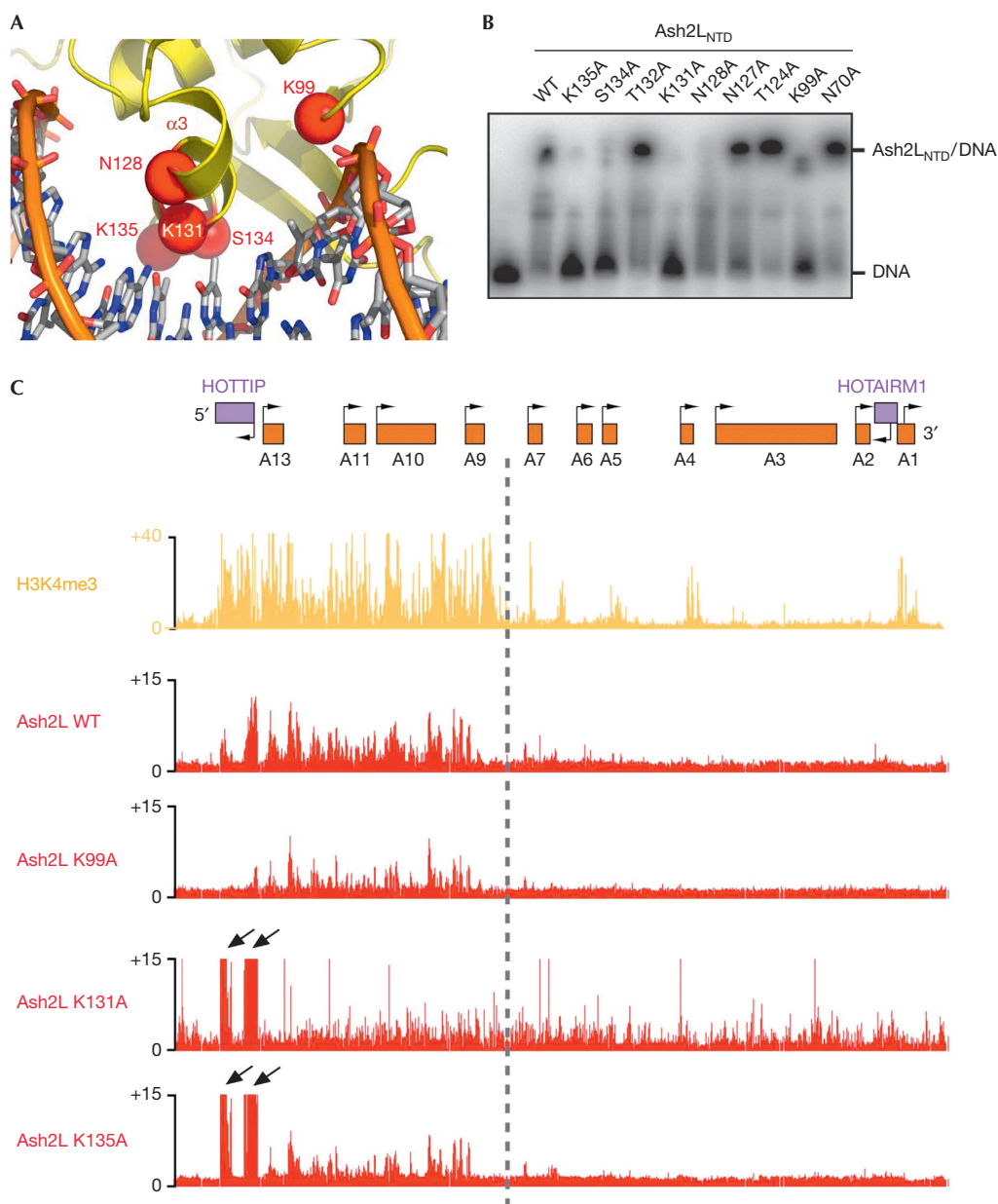


Fig 4 | Mutational analyses of the Ash2L_{WH}-DNA interaction. (A) The modelled interface between helix $\alpha 3$ of Ash2L_{WH} and the DNA major groove. Putative DNA-binding residues in Ash2L_{WH} are labelled. (B) Some point mutations abolished the Ash2L_{NTD}-DNA interaction. A concentration of 0.1 μM of DNA was incubated with 6.4 μM of wild-type (WT) and mutant Ash2L_{NTD} proteins. (C) Chromatin occupancy across approximately 100 kb of *HOXA* is shown. X axis is the genomic coordinate; Y axis shows occupancy of H3K4me3 or the individual Ash2L mutant proteins (chromatin immunoprecipitation/input). Point mutations in Ash2L_{WH} lead to decreased occupancy of these proteins across the 5' *HOXA* locus (K99A, K135A), increased occupancy over HOTTIP (K131A, K135A) and increased occupancy across the entire *HOXA* locus (K131A). Arrows highlight peaks of Ash2L occupancy over HOTTIP on changes in DNA binding. HOTTIP, *HOXA* transcript at the distal tip; WH, winged helix.

Given that the C-terminal SPRY domain of Ash2L mediates the interaction with RbBP5 and has a role in regulating H3K4 methylation by MLL family HMTs (Y. Chen *et al*, unpublished results), we expected that full-length Ash2L would show similar DNA-binding activity as the N-terminal domain. Surprisingly, despite the fact that the C-terminal domain of Ash2L (Ash2L_{CTD}, residues 230–534) alone did not show any detectable DNA-binding activity,

full-length Ash2L bound to DNA with a higher affinity (approximately 3 μM) than Ash2L_{NTD} (Fig 3D). Furthermore, purified Ash2L_{CTD} could still enhance the Ash2L_{NTD}-DNA interaction when added *in trans* (Fig 3E), even though there was no direct interaction between Ash2L_{NTD} and Ash2L_{CTD} in the absence or presence of DNA (supplementary Fig S4 online). Therefore, we propose that Ash2L_{CTD} synergistically cooperates with the winged-helix motif in

DNA binding. Structural information about full-length Ash2L and Ash2_{NTD} complexed with DNA is needed to understand the cooperative DNA binding by the two domains of Ash2L.

Mutational analysis of the Ash2L_{WH}–DNA interaction

To provide further support for the hypothesis that Ash2L_{WH} is a DNA-binding motif, we identified several putative DNA-binding residues in Ash2L_{WH} on the basis of the structural superposition of the winged-helix motifs of Ash2L and FoxO1 (Fig 4A). Ash2L_{NTD} proteins with single mutations of these residues were checked for their DNA-binding activity. All the mutant proteins behaved well in solution, indicating that these mutations did not affect protein folding and stability (supplementary Fig S5A online). Notably, alanine substitution of a cluster of residues on helix α 3 (Asn 128, Lys 131, Ser 134 and Lys 135) at the predicted Ash2L_{WH}–DNA interface either completely abrogated or weakened the DNA binding by Ash2L_{NTD} (Fig 4B). In addition, mutation of Lys 99 from loop L₁₂ between helices α 1 and α 2 also impaired the stability of the Ash2L_{NTD}–DNA complex (Fig 4B). By contrast, mutations of four residues—Asn 70, Thr 124, Asn 127 and Thr 132, which are at the peripheral region of the interface and presumably make less or no contribution to the interaction—had little effect on DNA binding (Fig 4B). Taken together, these results support the notion that Ash2L_{WH} is a DNA-binding motif of Ash2L.

Ash2L DNA-binding drives its chromosome localization

To investigate the significance of Ash2L_{WH} *in vivo*, we determined whether the DNA-binding activity of this motif has a role in targeting Ash2L to the *HOXA* locus (Wang et al, 2011). Our previous studies showed that core components of the MLL complex, WDR5, RbBP5 and Ash2L, are required for *HOX* gene expression (Dou et al, 2006). We expressed Myc-tagged wild-type and three DNA-binding-deficient mutants (K99A, K131A and K135A) of Ash2L in anatomically distal cells (primary human foreskin fibroblasts) and examined their localization across *HOXA* by loci-wide chromatin immunoprecipitation (ChIP), followed by tiling array analysis (ChIP–chip). The wild-type Ash2L showed a sharply demarcated localization; it only occupied the 5' *HOXA* locus from *HOXA9* to *HOXA13*, which corresponds to the transcriptionally active chromatin domain marked by H3K4me3 (Fig 4C). The broad distribution of Ash2L across the 5' *HOXA* locus also suggests that Ash2L probably does not bind to DNA with strong sequence specificity, consistent with the *in vitro* selection result.

In contrast to the wild-type Ash2L, all three DNA-binding-deficient mutants showed less occupancy on *HOXA9*–*HOXA13*. The K99A mutation led to less binding of Ash2L across the 5' *HOXA* locus overall (Fig 4C). Both Ash2L^{K131A} and Ash2L^{K135A} showed more-severe defects than Ash2L^{K99A}. In addition to the lower occupancy on 5' *HOXA* overall, Ash2L^{K131A} and Ash2L^{K135A} also showed increased occupancy at the *HOTTIP* (*HOXA* transcript at the distal tip) DNA (Fig 4C). Our recent studies showed that the MLL complex initially occupies the *HOTTIP* element and subsequently spreads to *HOXA9*–*HOXA13* by the *HOTTIP* RNA, a long intergenic non-coding RNA that coordinates the activation of several 5' *HOXA* genes (Wang et al, 2011). Thus, the accumulation of Ash2L^{K131A} and Ash2L^{K135A} at *HOTTIP* suggested that the sequential transfer of Ash2L from *HOTTIP* to the rest of the locus is blocked. Dissimilarly to Ash2L^{K99A} and

Ash2L^{K135A}, which were still confined within the 5' *HOXA* locus, Ash2L^{K131A} spread across the boundary between *HOXA9* and *HOXA7* into the 3' *HOXA* genes (Fig 4C). This indicates that Ash2L^{K131A} is unable to obey the chromatin domain boundary set up by chromosomal looping and existing histone marks.

Next, we examined whether the DNA-binding activity influences Ash2L recruitment to the promoter of *HOXC8*, another *HOX* gene, the expression of which is regulated by Ash2L (Dou et al, 2006). Wild-type Ash2L preferentially localized to the *HOXC8* locus (supplementary Fig S5B online). By contrast, when the K131A and K135A mutants were expressed in cells, the *HOXC8*-associated Ash2L showed an approximately 50% reduction compared to wild-type Ash2L-expressing cells (supplementary Fig S5B online). We conclude that the DNA-binding activity of Ash2L has an important role in targeting Ash2L to the *HOX* loci for gene expression regulation.

CONCLUSIONS

Mammalian MLL family complexes regulate a range of gene expressions, and the recruitment of these MLL complexes to their target genes involves a combination of mechanisms (Ruthenburg et al, 2007). Recent studies suggested that MLL interactions with menin and LEDGF (lens epithelium-derived growth factor) have a key role in targeting MLL complexes to the *HOX* locus (Yokoyama & Cleary, 2008). By contrast, the CXXC motif in MLL is dispensable for this localization. Instead, MLL_{CXXC} binds to non-methylated CpG DNA sites that are essential for maintenance of appropriate epigenetic marks and continued transcription (Cierpicki et al, 2009). As Ash2L is a common core component of all MLL complexes, we propose that Ash2L provides a nonspecific DNA-binding activity that collaborates with other gene-specific mechanisms to stabilize the association of MLL complexes with active chromatin domains. We demonstrate that a single mutation in the winged-helix motif of Ash2L (K131A) overcomes the chromatin domain boundary set up by chromosomal looping and histone marks, and allows Ash2L to spread into otherwise silent chromatin domains. It is possible that Ash2L either directly regulates chromatin domain demarcation or has an indirect role by interacting with other protein factors that define chromatin domain boundaries. Revealing the molecular mechanism by which Ash2L is involved in chromatin demarcation is an interesting direction for future investigations.

METHODS

Protein preparation. Ash2L_{NTD} was cloned into a pET28b-based vector with a 6xHis-SUMO tag fused at the N-terminus. Se-Met protein was expressed in *Escherichia coli* B834 (DE3) supplemented with 10 μ M ZnSO₄ in the M9 medium. The protein was purified by Ni-NTA affinity resin and on-bead digestion using Ulp1 protease, followed by gel filtration chromatography on Hiloal Superdex 75 equilibrated with 25 mM Tris–HCl, pH 8.0, and 150 mM NaCl. The purified protein was concentrated to 20 mg/ml and stored at –80 °C for crystallization.

Crystallization and structure determination. The crystals were grown by hanging-drop vapour diffusion at 4 °C. The precipitant/well solution contained 100 mM Tris–HCl, pH 8.5, and 20% PEG8000. Crystals were gradually transferred to a harvest solution containing 100 mM Tris–HCl, pH 8.5, 25% PEG8000 and 25% glycerol before flash-freezing in liquid nitrogen. SAD data was

collected at Advanced Photon Source beamline 21ID-G. Six Se sites were located and refined, and the SAD (single-wavelength anomalous dispersion) phases were calculated using SHARP (Vonrhein *et al*, 2007). A model was built into the experimental electron density. The model was then iteratively refined in CNS (crystallography and NMR system; Brunger, 2007).

Electrophoresis mobility shift assay. The sequence of DNA template used for the assay was 5'-CGCTCGAGGGATCCG AATTC(N₂₀)TCTAGAAAGCTTGTCGACGC-3'. The randomized double-stranded DNA was obtained by annealing N20 template with 3' primer (5'-GCGTTCGACAAGCTTTCTAGA-3'), followed by filling of 5' overhangs to form blunt ends using Klenow enzyme. The oligonucleotides were radiolabelled with [γ -³²P]ATP using T4 polynucleotide kinase. Radiolabelled oligonucleotides were separated from free [γ -³²P]ATP on G25 spin column. Ash2L in binding buffer (25 mM Tris-HCl, pH 8.0, 150 mM NaCl, 2 mM dithiothreitol and 10% glycerol) was mixed with 10 nM ³²P-labelled single-stranded DNA or double-stranded DNA in a total volume of 15 μ l. The reaction mixtures were incubated at 4 °C for 30 min before being loaded onto a 4–20% non-denaturing polyacrylamide gel. The gels were then dried and visualized using a PhosphorImager.

ChIP–chip. Complementary DNAs for each of the Ash2L mutants were cloned with a Myc epitope tag into lentiviral constructs under the control of the tetracycline operator. Replication incompetent, VSVg-coated lentiviral particles were packaged in 293T cells and used to infect primary human foreskin fibroblasts. The degree of protein induction was determined by titrating in doxycycline (Sigma) to reach an expression level equivalent to that of endogenous Ash2L. Cells were collected 72 h postinduction for ChIP–chip experiments. ChIP–chip was performed using H3K4me3 (Abcam, Cambridge, MA) and Myc (Abcam) antibodies, as described previously (Rinn *et al*, 2007). Retrieved DNA and input chromatin were competitively hybridized to custom tiling arrays interrogating human HOXA loci at 5 base-pair resolution, as described previously (Rinn *et al*, 2007).

ChIP and quantitative real-time-PCR. ChIP analyses were performed using the Chromatin Immunoprecipitation Assay Kit (Millipore) and the protocols recommended by the manufacturer. Real-time PCR quantification of ChIP was performed using Taqman probes and an ABI Prism 7500 (Applied Biosystems), and both the relative quantification and percent input methods as described previously (Dou *et al*, 2006).

The structure was deposited in the Protein Data Bank, with the accession code 3RSN.

Supplementary information is available at EMBO reports online (<http://www.emboreports.org>).

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CONFLICT OF INTEREST

The authors declare that they have no conflict of interest.

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