

Molecular and Functional Mapping of EED Motifs Required for PRC2-Dependent Histone Methylation

Nathan D. Montgomery^{1,2,3}, Della Yee^{1,2},
Stephanie A. Montgomery^{4,5} and Terry Magnuson^{1,2,3*}

¹Department of Genetics,
University of North Carolina at
Chapel Hill, 103 Mason Farm
Road, CB# 7264, Chapel Hill,
NC 27599-7264, USA

²Carolina Center for Genome
Sciences, University of
North Carolina at Chapel Hill,
103 Mason Farm Road,
CB# 7264, Chapel Hill,
NC 27599-7264, USA

³Curriculum in Genetics and
Molecular Biology, University of
North Carolina at Chapel Hill,
103 Mason Farm Road,
CB# 7264, Chapel Hill,
NC 27599-7264, USA

⁴Department of Microbiology
and Immunology, University of
North Carolina at Chapel Hill,
103 Mason Farm Road,
CB# 7264, Chapel Hill,
NC 27599-7264, USA

⁵Carolina Vaccine Institute,
University of North Carolina at
Chapel Hill, 103 Mason Farm
Road, CB# 7264, Chapel Hill,
NC 27599-7264, USA

Received 9 July 2007;
received in revised form
15 October 2007;
accepted 16 October 2007
Available online
22 October 2007

Polycomb group proteins represent a conserved family of developmental regulators that mediate heritable transcriptional silencing by modifying chromatin states. One Polycomb group complex, the PRC2 complex, is composed of several proteins, including the histone H3 lysine 27 (H3K27) methyltransferase enhancer of zeste homolog 2 and the WD-repeat protein embryonic ectoderm development (EED). Histone H3K27 can be mono-methylated (H3K27me1), dimethylated (H3K27me2), or trimethylated (H3K27me3). However, it remains unclear what regulates the number of methyl groups added to H3K27 in a particular nucleosome. In mammalian cells, EED is present as four distinct isoforms, which are believed to be produced by utilizing four distinct, in-frame translation start sites in a common *Eed* mRNA. A mutation that disables all four EED isoforms produces defects in H3K27 methylation [Montgomery, N.D., Yee, D., Chen, A., Kalantry, S., Chamberlain, S.J., Otte, A.P. & Magnuson, T. (2005). The murine polycomb group protein *Eed* is required for global histone H3 lysine-27 methylation. *Curr. Biol.*, **15**, 942–947]. To assess the roles of individual EED isoforms in H3K27 methylation, we first characterized three of the four EED isoform start sites and then demonstrated that individual isoforms are not necessary for H3K27me1, H3K27me2, or H3K27me3. Instead, we show that the core WD-40 motifs and the histone-binding region of EED alone are sufficient for the generation of all three marks, demonstrating that EED isoforms do not control the number of methyl groups added to H3K27.

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Edited by M. Yaniv

Keywords: histone; methylation; Polycomb; WD motif; translational initiation

*Corresponding author. E-mail address: trm4@med.unc.edu.

Abbreviations used: H3K27, H3 lysine 27; EED, embryonic ectoderm development; PcG, Polycomb group; EZH2, enhancer of zeste homolog 2; ES, embryonic stem; MEF, murine embryonic fibroblast; DAPI, 4',6-diamidino-2-phenylindole dihydrochloride; TS, trophoblast stem; YFP, yellow fluorescent protein; PBS, phosphate-buffered saline; NFDm, nonfat dried milk; TBST, Tris-buffered saline Tween-20.

Introduction

Polycomb group (PcG) proteins are a conserved family of development regulators that modify chromatin states in order to mediate heritable transcriptional silencing. PcG-mediated repression is important in diverse biological processes including X-chromosome inactivation, genomic imprinting, and segmental patterning.^{1–9} One PcG complex, the PRC2 complex, is composed of several bona fide PcG proteins, including the histone H3 lysine 27 (H3K27) methyltransferase enhancer of zeste homolog 2 (EZH2) and the WD-repeat protein embryonic ectoderm development (EED).^{9–12}

Histone H3K27 can be monomethylated (H3K27me1), dimethylated (H3K27me2), or trimethylated (H3K27me3).¹³ H3K27me3 is a repressive histone modification that localizes to confirmed targets of PcG silencing, including the inactive X chromosome.^{1,2,14–16} This mark mediates its repressive effect by recruiting to chromatin or at least stabilizing the association of a second PcG complex, PRC1.^{17–20} Conversely, the functions of H3K27me1 and H3K27me2 are not known, although H3K27me1 was recently shown to be present throughout euchromatic regions but absent in the vicinity of the start sites of active genes.²¹

While EZH2's role as the catalytic subunit of PRC2 has been clearly demonstrated, the functions of noncatalytic PRC2 subunits remain poorly defined. Both EED and the Zn finger protein SUZ12 are required to maintain the integrity of PRC2, and mutations in both genes cause EZH2 to become destabilized.^{22,23} Additionally, both *in vivo* and *in vitro*, EED and SUZ12 are required for normal PRC2-mediated H3K27 methylation.^{22–24} However, precise mechanistic details of these functions remain unclear.

Four EED isoforms are found in mammals, and these isoforms are thought to be produced by utilizing four in-frame translation start sites in the *Eed* mRNA. EED isoform usage is regulated developmentally, leading to speculation that EED could also function as a PRC2 regulatory subunit.²⁵ Additionally, EED-2, which has only been observed in undifferentiated stem cells and in tumors, has been proposed to be important in maintaining developmental plasticity.²⁶ However, definitive biochemical functions of the various EED isoforms have not been demonstrated. Previous work using *in vitro* methyltransferase assays postulated that EED isoforms control the substrate specificity of the PRC2 complex. In those initial studies, the largest isoforms, EED-1 and EED-2, appeared to direct EZH2 methyltransferase activity toward histone H1K26, whereas EED-3 and EED-4 appeared to direct EZH2 methyltransferase activity toward H3K27.²⁵ However, a second recent study that also utilized *in vitro* methyltransferase assays failed to detect any difference in the histone substrate preference of PRC2 complexes with or without EED-1 or EED-2.²⁷

Given this discrepancy, in the present study, we aimed to define regions of EED required for each

H3K27 methylation state *in vivo* using mouse embryonic stem (ES) cells that are wild type or mutant for *Eed*. Because all four known EED isoforms associate with EZH2, we examined whether these isoforms might control the number of methyl groups added to H3K27 in a particular nucleosome.^{25,26}

To this end, we definitively characterize three of the four EED isoform start sites and demonstrate that individual isoforms are not necessary for H3K27me1, H3K27me2, or H3K27me3. These results indicate that EED isoforms do not control the enzymatic activity of the PRC2 complex. Instead, we show that EED's core WD-40 motifs and histone-binding region alone are sufficient for the generation of all three marks.

Results

Distinct localization of H3K27 methylation states

Unlike mouse stem cell lines, differentiated mouse cells, such as murine embryonic fibroblasts (MEFs), have a striking nuclear architecture, in which regions of the genome packaged as part of the pericentric heterochromatin are clearly visible by 4',6-diamidino-2-phenylindole dihydrochloride (DAPI) staining as DNA-rich foci. Using this characteristic DNA-staining pattern to provide landmarks, we assessed the localization of H3K27me1, H3K27me2, and H3K27me3 in CD1 MEFs (Fig. 1). For these experiments, we used antibodies specific for each H3K27 methylation state. The specificities of these antibodies have been demonstrated previously by competition assays.¹³ Additionally, positive staining with all three antibodies is lost in *Eed* mutant ES cells and trophoblast stem (TS) cells.²² Because methylation defects at other lysine residues have not been observed in *Eed* mutant ES and TS cells, these results strengthen the conclusion that the antibodies do not cross-react with methyl marks at other residues.

As previously reported, large foci of H3K27me1 colocalized with the DNA-rich pericentric heterochromatin (Fig. 1a; Ref. 13). Conversely, H3K27me2 and H3K27me3 were specifically excluded from these regions, a pattern opposite to what was observed with H3K27me1 (Fig. 1b and c; Supplementary Fig. 1). HP1- α , an established marker of pericentric heterochromatin,^{28–30} was also enriched in the DNA-rich foci, confirming the identity of these regions (Fig. 1d). Finally, H3K27me2 and H3K27me3, though largely colocalized (Supplementary Fig. 2), were distinguished by the characteristic enrichment of H3K27me3 on the inactive X chromosome in female MEFs (Fig. 1b and c).

In addition to confirming and establishing global localization patterns of H3K27me1, H3K27me2, and H3K27me3, the distinct staining patterns confirm that the H3K27 antibodies are each specific for one H3K27 methylation state and do not cross-react with histones harboring other H3K27 methylation states.

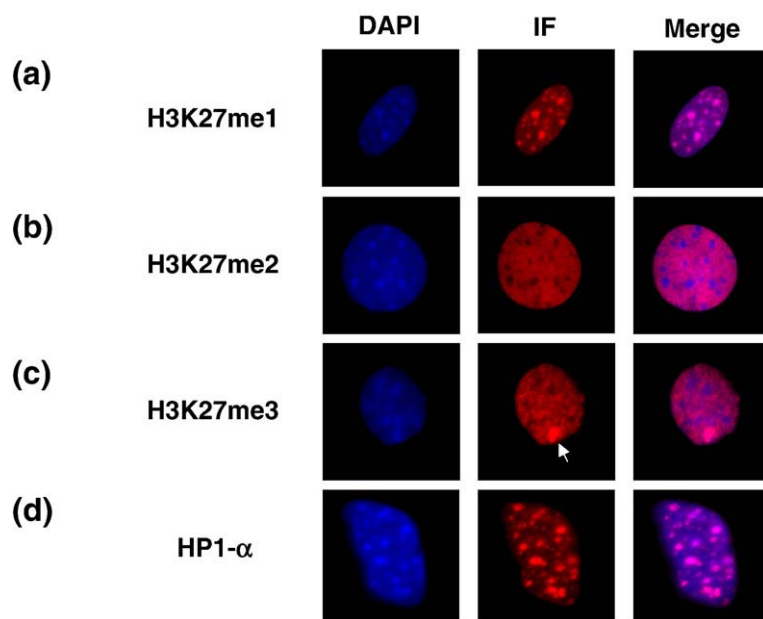


Fig. 1. Localization of histone H3K27 methylation marks. Immunofluorescence analysis of (a) H3K27me1, (b) H3K27me2, (c) H3K27me3, and (d) HP1- α in CD1 MEFs. The inactive X chromosome is indicated by an arrow in (c).

Characterization of EED isoforms expressed in wild-type ES cells

The distinct localization patterns of the three H3K27 methylation states indicate that the number of methyl groups added to H3K27 in a particular nucleosome is a regulated process. Because EED has previously been posited to play a regulatory role in PRC2-mediated histone methylation,²⁵ we aimed to define regions of EED required for H3K27me1, H3K27me2, and H3K27me3. As a first step, we attempted to discern which EED isoforms are required for H3K27 methylation by rescuing *Eed* mutant ES cells, which lack detectable levels of endogenous H3K27me1, H3K27me2, and H3K27me3,²² with constructs expressing individual EED isoforms. However, before a systematic series of rescue experiments was performed, we attempted to confirm the identities of the EED isoforms.

Previous work indicates that all four EED isoforms are present in wild-type, undifferentiated mouse ES cells.²⁶ However, we were consistently only able to resolve three isoforms in multiple, independently derived ES cell lines, as well as in TS cells (Fig. 2 and data not shown). To assess whether our electrophoresis conditions would allow us to resolve all four EED isoforms and, if so, to confirm the identities of the three observed isoforms, we compared the isoforms present in our ES cells to the isoforms present in HeLa cells and in mouse *Wap-T₁₂₁* mammary tumors. Previous work has demonstrated that EED-1, EED-3, and EED-4 are expressed in HeLa cells and that EED-2 is upregulated in many mouse tumors.²⁶ Consistent with those reports, we observed high levels of EED-1, EED-3, and EED-4 and much lower levels of EED-2 in HeLa cells (Fig. 2a). Similarly, EED-2, along with EED-3 and EED-4, was observed in *Wap-T₁₂₁* breast tumors (Fig. 2b). Comparison of the isoforms present in those sources with the isoforms present in our ES cells suggests

that we observe isoforms 1, 3, and 4 but not isoform 2 in our ES cells (Fig. 2).

Kuzmichev *et al.* demonstrated that EED-2 is rapidly downregulated when ES cells are stimulated to differentiate.²⁵ Hence, the most parsimonious explanation for our failure to detect four EED isoforms is that the ES cells employed were beginning to differentiate when they were harvested, causing a downregulation of EED-2. One possible explanation for such differentiation would be the removal of the cells from fibroblast feeders prior to harvesting in order to minimize feeder contamination in our immunoblotting assays. To avoid this complication, we also analyzed EED isoforms in E14 ES cells, which are feeder independent,³¹ but we were still able to detect only three isoforms (Fig. 2).

ES cells, TS cells, and primitive endoderm-derived XEN cells are the only known cell types that can be derived from mouse embryos prior to gastrulation, the embryonic stage when *Eed* mutant embryos arrest.^{32–36} Because only three EED isoforms were resolved in our ES and TS cells and because only trace levels of EED are detected in XEN cells (data not shown), we restricted our analysis to EED isoforms 1, 3, and 4.

Deletion mapping of EED isoform start sites

Although most proteins initiate translation at methionine-encoding AUG codons, an increasing number of proteins are recognized to initiate translation from non-AUG codons. Often, these alternative start sites generate upstream isoforms of proteins also translated from downstream, canonical AUG initiation codons, and typically, the alternative codons differ from the canonical AUG sequence at only one of the three nucleotide positions.^{37,38} For instance, fibroblast growth factor-2 is present as five isoforms in mammalian cells, with

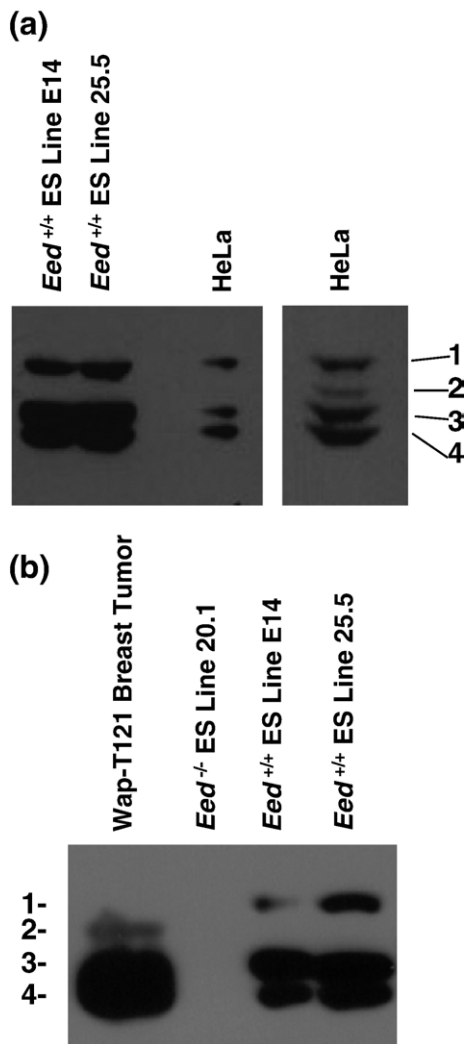


Fig. 2. Confirmation of EED isoform identities. Western blot analysis of EED comparing isoforms observed in ES cells to isoforms observed in (a) HeLa cells or (b) mouse *Wap-T121* breast tumors. On prolonged exposure, EED-2 becomes visible in HeLa lysates (a, right panel). EED isoforms 1–4 are indicated as 1, 2, 3, and 4.

four CUG initiation codons upstream of a canonical AUG start site, and vascular endothelial growth factor is translated from both an upstream CUG and a downstream AUG.^{39–43} Consistent with both of these trends, the upstream EED-1 and EED-2 are postulated to initiate translation from noncanonical GUG codons at positions 169–171 and 274–276 in the *Eed* cDNA, respectively, and the putative start sites for EED-3 and EED-4 are canonical AUG sequences at positions 451–453 and 493–495, respectively (Fig. 3a; Supplementary Fig. 3).^{25,44}

Overall, mouse and human EED are very similar proteins. In fact, the predicted EED-3 and EED-4 isoforms are 100% identical between the two species.⁴⁵ However, the sequences between the putative EED-1 and EED-2 start sites are much more variable.⁴⁵ Given the abrupt boundary between highly variable and nearly identical sequences, we

questioned whether the actual EED-1 start site might be further 3' than previously reported.

To map EED isoform start sites, we generated a series of *Eed* expression constructs progressively truncated at the 5' end of the *Eed* cDNA (Fig. 3a). Individual constructs were then transiently trans-

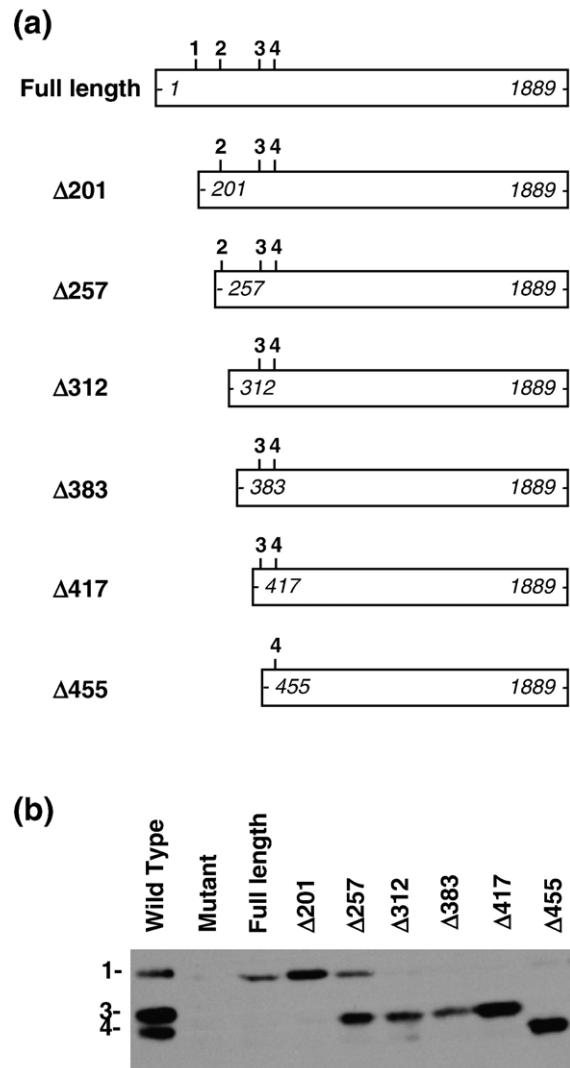


Fig. 3. Deletion mapping of EED translational start sites. (a) Constructs transfected into *Eed* mutant ES cell line 21 (below) are shown. Putative translation start sites for EED-1 (GUG 169–171), EED-2 (GUG 274–276), EED-3 (AUG 451–453), and EED-4 (AUG 493–495) are indicated in the schematic of the endogenous mRNA as 1, 2, 3, and 4 and correspond to codon locations in the mouse *Eed* transcript (accession number: BC012966). The positions of the most 5' and 3' nucleotides of each construct are indicated in italics. Identifiers for each construct are shown on the left and indicate the number of nucleotides removed from the 5' end of the *Eed* cDNA (e.g., Δ210 refers to a construct expressing an *Eed* cDNA lacking the 5' 210 nucleotides of the full-length message). (b) Whole-cell lysates from *Eed*^{+/+} ES cell line 25.5 (wild type), *Eed*^{-/-} ES cell line 21 (mutant), or *Eed*^{-/-} ES cell line 21 transiently transfected with the *Eed* expression constructs shown in (a) were analyzed by Western blotting with an antibody detecting EED.

fected into *Eed* mutant ES cells, which lack detectable endogenous EED, and EED isoform expression was assessed by Western blotting. Because isoform expression could be lost either by deleting past an isoform start site or by deleting an upstream regulatory element required for translation from an intact start site, the absence of a band is uninformative in this assay. However, the continued presence of an isoform after its putative start site has been deleted is strong evidence that the actual start site must be further downstream. Notably, after transfection, the furthest 5' intact translation start site was generally utilized preferentially to downstream translation start sites (Fig. 3). This observation suggests that regulated usage of the various EED translation start sites is not simply a consequence of the interaction between *trans*-acting factors and sequences present in the message but instead may be influenced by upstream regulatory events.

Consistent with the relaxed sequence conservation between mouse and human *Eed* sequences beginning with and immediately downstream from GUG 169–171, EED-1 was expressed from constructs truncated 32 ($\Delta 201$) and even 88 ($\Delta 257$) nucleotides beyond the reported EED-1 start site, suggesting that EED-1 may not initiate translation at GUG 169–171 as previously proposed (Fig. 3b). EED-1 expression was lost only after deleting the 5' 312 nucleotides of the *Eed* cDNA, a deletion extending 143 nucleotides beyond the reported EED-1 start site and 38 nucleotides beyond the published EED-2 start site (Fig. 3b). Consistent with the published identity of the EED-3 start site at AUG 451–453, this isoform was observed after deleting the 5' 417 nucleotides but not after deleting the 5' 455 nucleotides (Fig. 3b). Finally, EED-4 was present even in the largest truncation, which deleted the 5' 455 nucleotides, consistent with the published EED-4 start site residing at AUG 493–495 (Fig. 3b). Notably, our ability to detect EED-1, EED-3, and EED-4 from this series of cDNA expression constructs confirms that alternative splicing of exons within the EED coding sequence is not responsible for the production of these three isoforms.

Confirmation of EED isoform start sites

To verify that GUG 169–171 is upstream of the actual EED-1 start site, we forced expression from this codon by replacing the sequences encoding GUG 169–171 with a canonical translation start site consisting of a consensus Kozak sequence followed by an AUG initiator codon (Fig. 4a). Consistent with the hypothesis that EED-1 translation actually initiates further downstream, the resulting product was substantially larger than EED-1 (Fig. 4b, asterisk). A lower level of an EED-1-sized product, presumably initiating at the actual start site further downstream, was also observed after transfection with this construct (Fig. 4b, arrow).

Given that the deletion mapping data indicate that either the true EED-1 start site or some regulatory

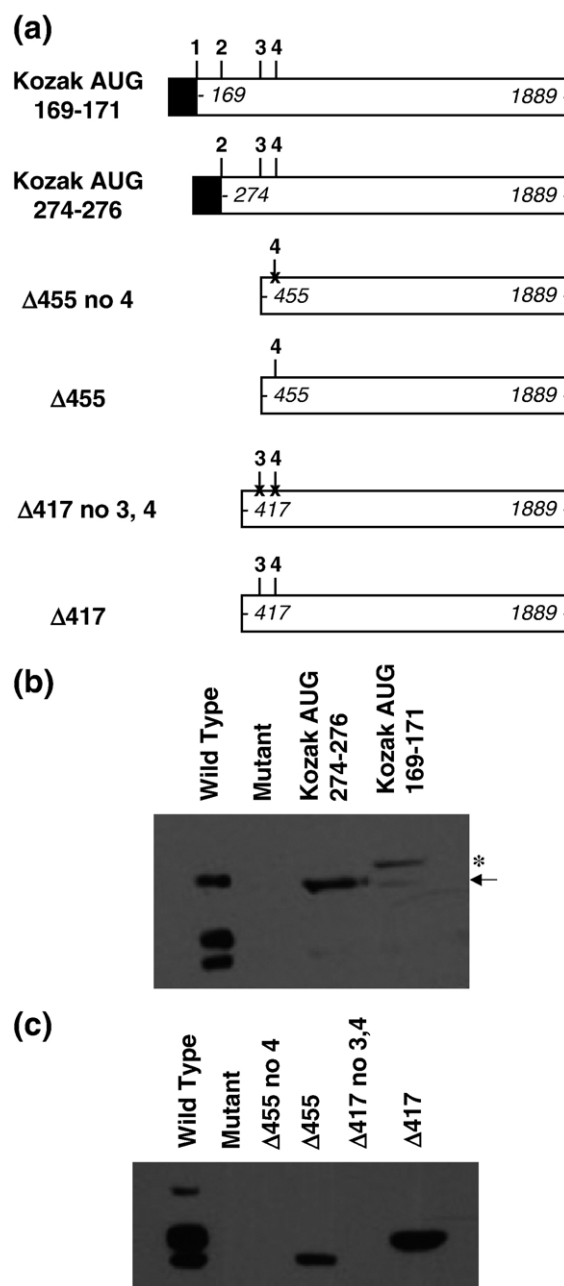


Fig. 4. Verification of EED isoform start sites. (a) Constructs transfected into *Eed* mutant ES cell line 21. The positions of the most 5' and 3' nucleotides of each construct are indicated in italics. Reported EED translational start sites GUG 169–171, GUG 274–276, AUG 451–453, and AUG 493–495 are indicated as 1, 2, 3, and 4, respectively. Black boxes refer to strong Kozak-AUG sequences engineered into the expression construct in order to drive translation from the 169–171 codon (Kozak AUG 169–171) and from the 274–276 codon (Kozak AUG 274–276), respectively. "X" markings through the putative EED-3 and EED-4 start sites in $\Delta 417$ no 3, 4 and in $\Delta 455$ no 4 indicate AUG→AUA mutations intended to disrupt translation initiation. (b and c) Whole-cell lysates from *Eed*^{+/+} ES cell line 25.5 (wild type), *Eed*^{-/-} ES cell line 21 (mutant), or *Eed*^{-/-} ES cell line 21 transiently transfected with the *Eed* expression constructs shown in (a) analyzed by Western blotting with an antibody detecting EED.

element required for translation from that true EED-1 start site must map between nucleotides 257 and 312, we hypothesized that EED-1 may be produced by translation initiating from the previously proposed EED-2 start site at GUG 274–276. To test that possibility, we forced expression of a protein initiating translation at that site by replacing GUG 274–276 with a consensus Kozak followed by an AUG initiator codon (Fig. 4a). Consistent with EED-1 translation initiating at this location, the resulting product was the same size as EED-1 observed in mouse ES cells (Fig. 4b).

Because EED-2 was not expressed in our wild-type ES cells or from any of our truncated expression constructs, we were unable to characterize this isoform. However, driving translation from a candidate start site at GUG 397–399 produced a protein that appears to be smaller than EED-2 (data not shown), suggesting that the EED-2 start site lies between GUG 274–276 and GUG 397–399 or that EED-2 is generated by alternative mechanisms, such as proteolytic processing.

Our deletion mapping data were consistent with presumptive EED-3 and EED-4 start sites residing at AUG 451–453 and AUG 493–495, respectively. To confirm that translation of those isoforms originates at those sites, we engineered AUG→AUA site-directed point mutations into the AUG 451–453 and AUG 493–495 codons (Fig. 4a). Consistent with those codons being the initiation codons for the two smaller isoforms, constructs harboring those mutations failed to express EED-3 and EED-4 (Fig. 4c).

EED isoforms are dispensable for all H3K27 methylation states

To determine whether all three EED isoforms present in our ES cells are required to mediate the three H3K27 methylation states, we monitored H3K27 methylation in *Eed* mutant ES cells transiently transfected with several of the *Eed* cDNA expression constructs. This rescue assay has previously been utilized to demonstrate that protein(s) expressed from a full-length *Eed* cDNA cassette can mediate all three H3K27 methylation states,²² although the role of individual isoforms in that rescue has not been directly assessed. This approach is advantageous relative to an isoform-specific, targeted loss-of-function approach because it avoids the problematic interpretation of negative results in the event that individual isoforms are dispensable for H3K27 methylation.

Constructs harboring site-directed mutations to eliminate EED-3 and EED-4 expression or truncated to eliminate EED-1 expression were able to rescue all three H3K27 methylation states in *Eed* mutant ES cells (Fig. 5d and e). Additionally, a construct retaining only the EED-4 start site also rescued H3K27me1, H3K27me2, and H3K27me3 (Fig. 5f). As previously reported, expression of the *Eed*^{l7Rn5-3354SB} allele, which produces an unstable and nonfunctional protein harboring an L→P substitution, failed to rescue H3K27 methylation (data not shown, but

see Ref. 22). Collectively, these results demonstrate that the three H3K27 methylation states are not dependent on individual EED isoforms.

Disruption of EED WD-40 motifs eliminates H3K27 methylation

Having demonstrated that the sequences distinguishing EED isoforms are not required for H3K27 methylation, we attempted to assess which functional motifs common to all four EED isoforms might be essential for PRC2 activity. EED and its homologs in other organisms are WD-repeat proteins. However, there is disagreement about the number of WD-40 motifs present in EED, with estimates varying between five and seven.^{46–48} In functional studies assessing EED's ability to bind EZH2 or its ability to mediate transcriptional repression when tethered to a GAL4 DNA-binding domain, only five WD-40 motifs have appeared functionally necessary.^{49,50} Those five motifs map to *Eed* cDNA sequences 721–808 (WD-40 motif 1), 1012–1105 (WD-40 motif 2), 1150–1240 (WD-40 motif 3), 1330–1444 (WD-40 motif 4), and 1672–1774 (WD-40 motif 5). To determine whether those same regions are required for EED's ability to mediate H3K27 methylation, we transiently transfected *Eed* mutant ES cells with a series of *Eed* cDNA expression constructs progressively truncated from either the N-terminus or the C-terminus.

A protein lacking the N-terminal 16 amino acids of EED-4 (Δ 5' 541) was able to mediate all three H3K27 methylation states (Fig. 6b), confirming that the N-terminal regions in EED, including those amino acids that distinguish individual isoforms, are not required for the catalytic activity of the PRC2 complex. Conversely, deletion of any of the five putative WD-40 motifs abolished EED's ability to mediate H3K27 methylation (Fig. 6d–g). An intermediate deletion, Δ 5' 697 also left the five putative WD-40 motifs intact. However, protein expressed from this construct retains only 8 amino acids upstream of WD-40 motif 1 and lacks a portion of EED recently shown to be necessary for histone binding.⁵¹ Although the expression of the Δ 5' 697 construct consistently rescued a low level of H3K27me1, little, if any, H3K27me2 and H3K27me3 were observed (Fig. 6c).

EED–EZH2 interaction assessed in living cells by YFP fragment complementation assays

The inability of several truncated *Eed* expression constructs to successfully rescue H3K27 methylation defects in *Eed* mutant cells could reflect an inability of these proteins to associate with EZH2, the catalytic subunit of the PRC2 complex. In order to assess EED–EZH2 interactions, we utilized a yellow fluorescent protein (YFP) fragment complementation assay.⁵² In this assay, test proteins are translationally fused to N- and C-terminal halves of YFP, called Venus 1 and Venus 2, respectively. If the test proteins are able to interact, the two YFP fragments

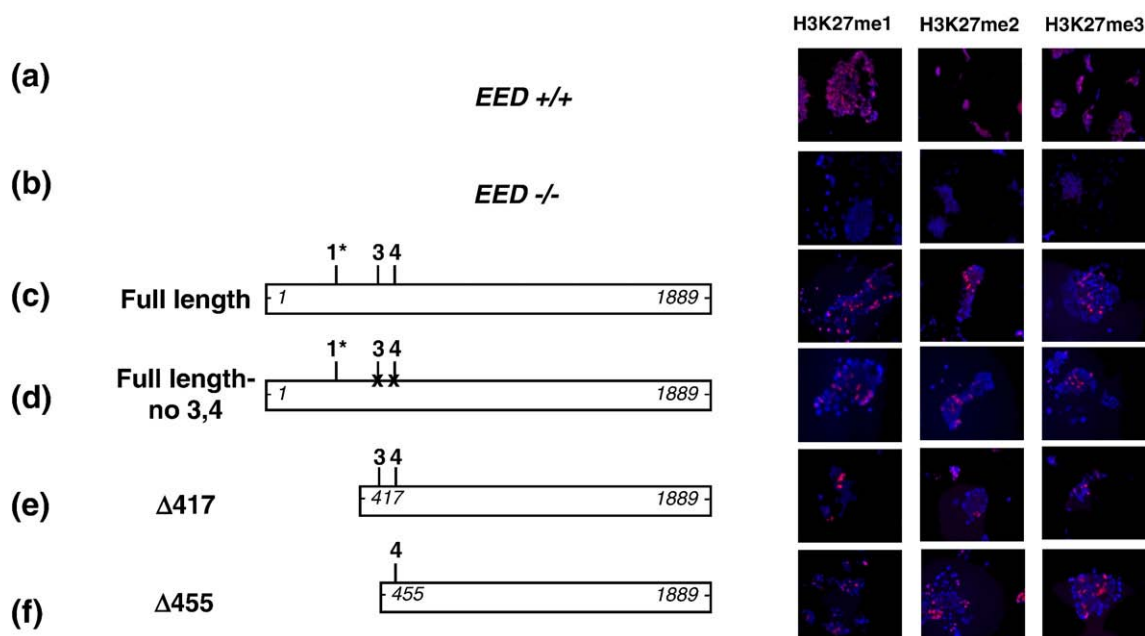


Fig. 5. Histone H3K27 methylation in cells lacking one or more EED isoforms. Immunofluorescence analysis of H3K27me1, H3K27me2, and H3K27me3 in wild-type ES cell line 25.5 (*Eed*^{+/+}) and in *Eed* mutant ES cell line 21 either mock transfected (*Eed*^{-/-}) or transiently transfected with the indicated constructs. *Eed* expression constructs transfected into ES cell line 21 are shown on the left. The positions of the most 5' and 3' nucleotides of each construct are indicated in italics. Isoform start sites at GUG 274–276, AUG 451–453, and AUG 493–495 are shown as 1*, 3, and 4, respectively. 1* discriminates the GUG 274–276 start site for EED-1 reported here from the GUG 169–171 start site reported previously.^{25,44} “X” markings through the putative EED-3 and EED-4 start sites in Full length no 3, 4 and in Δ417 no 3, 4 represent AUG → AUA mutations intended to disrupt translation initiation. EED 1036–1038 L → P refers to *Eed*^{l7Rn5-33545B}, a point mutant protein previously demonstrated to lack H3K27 methyltransferase activity. DAPI-stained DNA is blue, and methylated histones are shown in red. In the transient transfection assay, approximately 10% of the ES cells are successfully transfected, and with constructs expressing functional EED, a similar percentage of cells are rescued.

will be close enough to reconstitute the functional fluorescent complex, which can be assessed by fluorescence microscopy or flow cytometry.

To demonstrate the validity of the YFP fragment complementation assay, we utilized control constructs expressing Venus 1 and Venus 2 fragments translationally fused to a homodimerizing leucine zipper protein, GCN4. When coexpressed in E14 ES cells, robust YFP fluorescence was observed (Fig. 7a). Similarly, Venus 1–EZH2 and EED–Venus 2 were able to interact and reconstitute functional YFP (Fig. 7c; Table 1). This fluorescence was distributed throughout the cell and was specifically dependent on the presence of both constructs, as expression of individual constructs failed to produce YFP activity (Fig. 7h and i).

Next, the truncated *Eed* cDNAs, which were tested previously for their ability to rescue H3K27 methylation defects in *Eed* mutant cells, were fused to Venus 2. Consistent with previous reports,^{49,50} deletion of even a single WD-40 motif from either the N-terminus or the C-terminus of EED appeared to abolish the EED–EZH2 interaction (Fig. 7f and g; Table 1), just as it abolished the ability to rescue H3K27 methylation (Fig. 6d and g).

The *Eed* Δ5' 541 truncation fully rescues H3K27 methylation despite lacking the first 16 amino acids of EED-4 (Fig. 6b). Consistent with that functional readout, in the YFP fragment complementation

assay, this protein was able to robustly interact with Venus 1–EZH2, although at a lower level than the full-length construct (Fig. 7d; Table 1). Expression of the *Eed* Δ5' 697 allele rescued H3K27me1 but not higher H3K27 methylation states. In the YFP fragment complementation assay, no fluorescence over background was observed upon coexpression of Venus 1–EZH2 and EED Δ5' 697–Venus 2, suggesting that this more severe truncation severely limits EED's ability to interact with EZH2.

Truncated EED protein levels

PRC2 subunits are frequently unstable outside of intact protein complexes.^{22,23} As a result, the inability of several truncated proteins to interact with EZH2 raises the possibility that these truncated proteins are not stable. To address this possibility, we assessed levels of FLAG-tagged truncated EED proteins by Western blotting. Consistent with its ability to bind EZH2 and rescue H3K27 methylation, robust levels of EED Δ5' 541 were detected in transfected cells (Fig. 8). Conversely, EED Δ5' 697, EED Δ5' 913, EED Δ3' 1540-End, and Δ3' 1324-End, all of which fail to bind EZH2 in the split YFP assay, were undetectable (Fig. 8). Consequently, there was a strong correlation between H3K27me2/H3K27me3 activity, EZH2 binding, and protein level (Figures 6–8). A striking exception to this

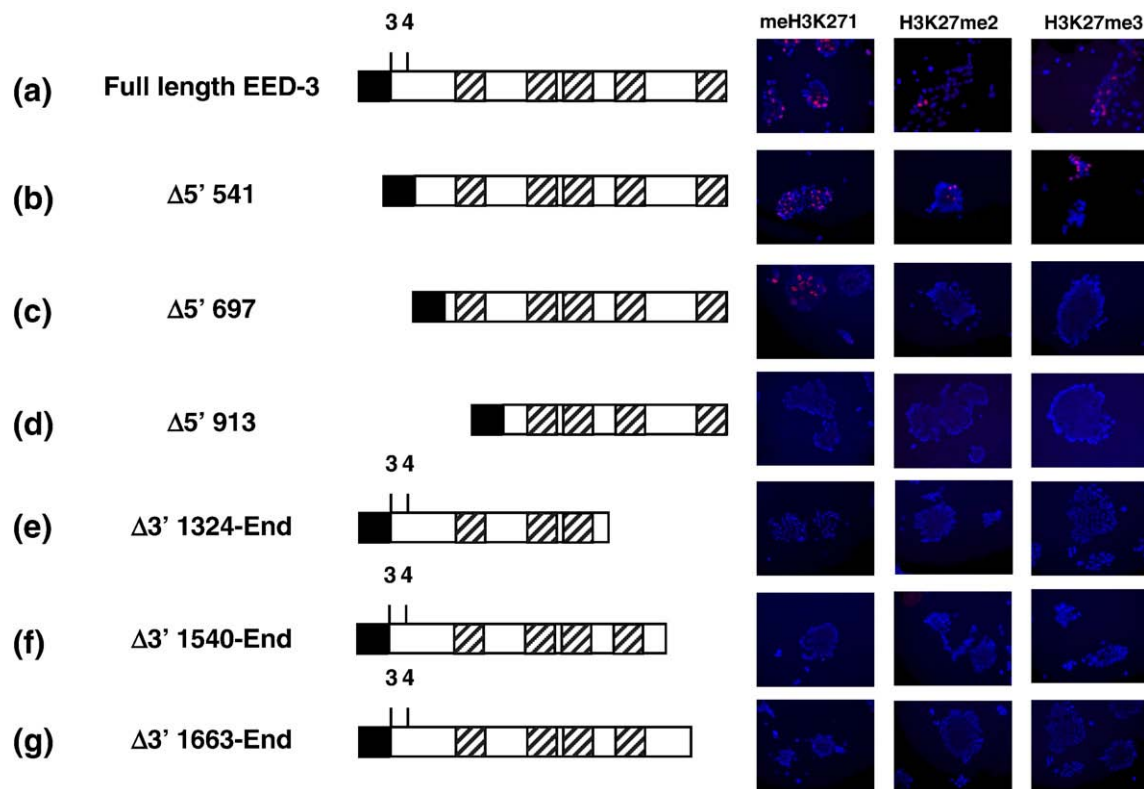


Fig. 6. Functional mapping of required WD-40 motifs in EED. Immunofluorescence analysis of H3K27me3 in *Eed* mutant ES cell line 21 either mock transfected (*Eed*^{-/-}) or transiently transfected with the indicated constructs. Below each schematic, nucleotides encoding for the N- and C-terminal amino acids of each protein are indicated in italics. Above each schematic, 3 and 4 refer to translation start sites for EED isoforms 3 and 4. Diagonally lined boxes refer to putative WD-40 motifs encoded by cDNA sequences 721–808 (WD-40 motif 1), 1012–1105 (WD-40 motif 2), 1150–1240 (WD-40 motif 3), 1330–1444 (WD-40 motif 4), and 1672–1774 (WD-40 motif 5).⁴⁹ Filled boxes refer to consensus Kozak+ATG initiator sequences engineered into the construct. DAPI-stained DNA is blue, and methylated histones are shown in red.

trend was observed with the *Eed* Δ3' 1663-End allele. Despite failing to bind EZH2 robustly in the YFP complementation assay (Fig. 7), this protein was stably detected in transfected cell populations (Fig. 8).

Discussion

EED isoforms are not required for any H3K27 methylation state

Although the biological functions of H3K27me1 and H3K27me2 have not been defined, it is clear from localization studies that the three H3K27 methylation states are partitioned to nucleosomes associated with distinct regions of the genome (Fig. 1; Refs. 13,21). Interestingly, whereas we and others have demonstrated by immunofluorescence that foci of H3K27me1 staining colocalize with the pericentric heterochromatin, a recent study utilizing chromatin immunoprecipitation demonstrated that H3K27me1 is found throughout chromatin (Fig. 1; Refs. 13,21). Likely, the apparent pericentric enrichment of H3K27me1 observed by immunofluorescence simply reflects the local enrichment of his-

tones in the compacted pericentric heterochromatin, as opposed to any specific enrichment of this mark as had previously been suggested.¹³

Nevertheless, H3K27me1, H3K27me2, and H3K27me3 do appear to localize to nucleosomes associated with distinct regions of the genome (Fig. 1). This specificity, in turn, implies that the number of methyl groups added to a particular nucleosome is a regulated process. In the present study, we have defined those regions of EED required for H3K27 methylation. Initially, we hypothesized that EED isoforms could function as regulatory switches controlling H3K27 methylation states. However, none of the EED isoforms were specifically required in order to generate H3K27me1, H3K27me2, or H3K27me3 (Fig. 5). These results demonstrate that the N-terminal extensions discriminating EED isoforms are not required for the enzymatic activity of the PRC2 complex.

EED WD-40 motifs and histone-binding regions are required for H3K27 methylation

The ability of *Eed* expression constructs to rescue H3K27 methylation was dependent upon the presence of all five EED WD-40 motifs (Fig. 6). Truncated proteins lacking the N- or C-terminal WD-40

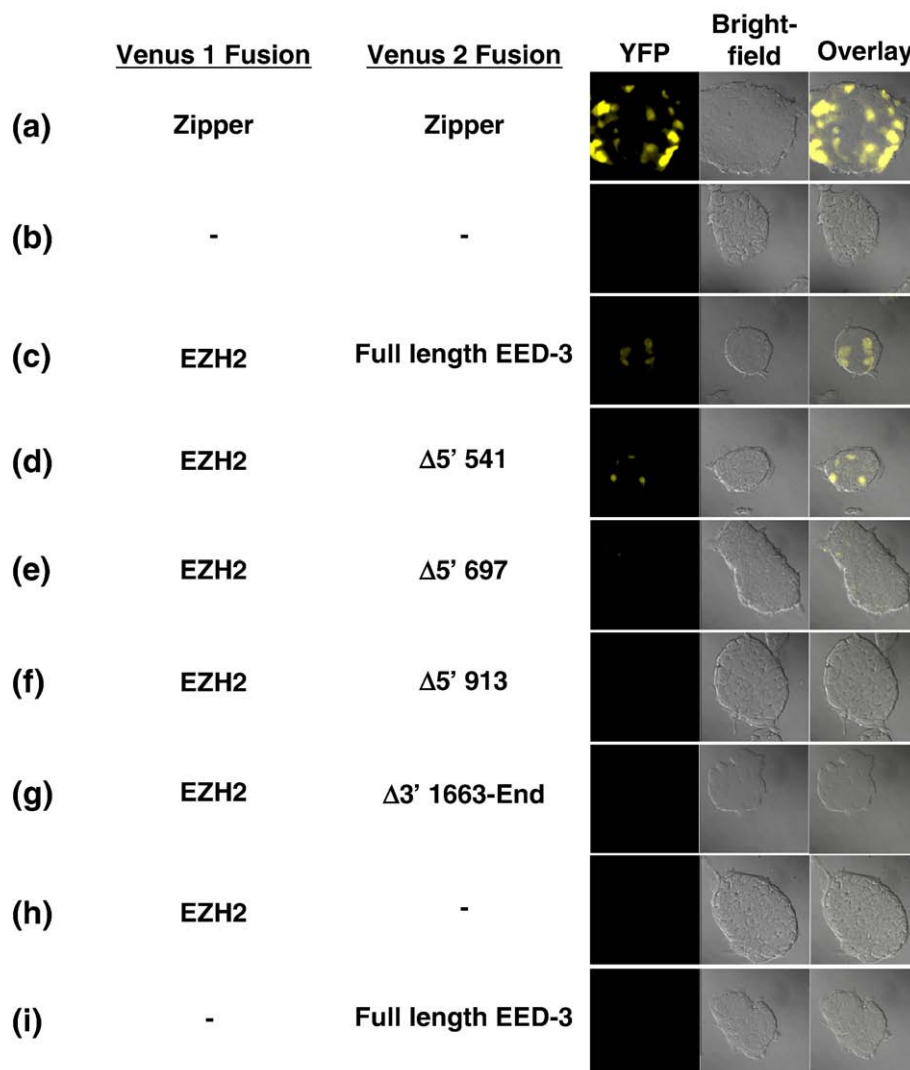


Fig. 7. EED–EZH2 interaction assessed by YFP fragment complementation assays. YFP fragment complementation assays in E14 ES cells transiently transfected with constructs expressing the following Venus 1 and Venus 2 fusions: (a) Venus 1–GCN4 and GCN4–Venus 2, (b) no DNA, (c) Venus 1–EZH2 and Full-length EED-3–Venus 2, (d) Venus 1–EZH2 and EED Δ5' 541–Venus 2, (e) Venus 1–EZH2 and EED Δ5' 697–Venus 2, (f) Venus 1–EZH2 and EED Δ5' 913–Venus 2, (g) Venus 1–EZH2 and EED Δ3' 1663-End–Venus 2, (h) Venus 1–EZH2 and no Venus 2, and (i) no Venus 1 and Full-length EED-3–Venus 2.

motif were also unable to bind to EZH2 in a YFP fragment complementation assay (Fig. 7e and f), confirming that EED–EZH2 association is likely

Table 1. Split YFP complementation by EED–EZH2 interaction

	Percentage of positive cells (mean±SD)
Full length	11.27±0.33
Δ5' 541	6.21±0.52
Δ5' 697	0.18±0.04
Δ5' 913	0.12±0.02
Δ3' 1663-End	0.28±0.05
No transfect	0±0

E14 ES cells were cotransfected with Venus 1–EZH2 and EED–Venus 2 expression constructs. Triplicate samples were analyzed using fluorescence-activated cell sorting. The mean percentage of YFP-positive cells and corresponding standard deviation from a representative experiment are shown.

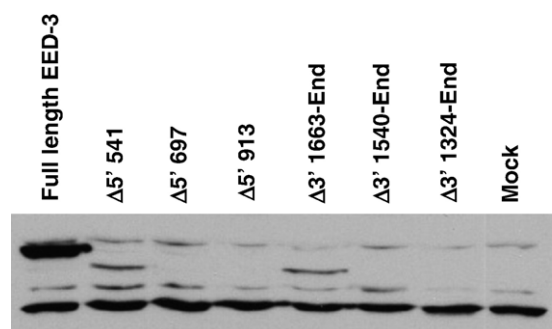


Fig. 8. Levels of truncated EED proteins. The stability of truncated, tagged EED proteins was assessed by Western blotting lysates from transfected E14 ES cells with an anti-FLAG antibody. Three background bands observed with the FLAG antibody serve as loading controls and are indicated by asterisks.

necessary for normal H3K27 methylation. These observations are also consistent with earlier studies indicating that all five WD-40 motifs are required for EZH2 binding and EED-mediated transcriptional repression.^{49,50}

Interestingly, whereas the $\Delta 5'$ 541 construct was able to rescue all three H3K27 methylation states, the protein expressed from the $\Delta 5'$ 697 construct, which also retains all five WD-40 motifs, was able to mediate a low level of H3K27me1 but no H3K27me2 or H3K27me3 (Fig. 6b and c). Recent work suggests that the N-termini of EED and of its fly homolog ESC bind to histone H3 and that the histone-binding N-terminus of ESC is required for H3K27me3.⁵¹ The $\Delta 5'$ 697 construct deletes the region of EED required for histone H3 binding. Accordingly, the reduced levels of H3K27 methylation mediated by this construct suggest that, as in *Drosophila*, the histone H3-binding N-terminus of EED may be required for full activity of the PRC2 complex. However, unlike the studies with ESC, binding between EED and EZH2 also appears to be compromised when the histone-binding domain is deleted (Fig. 7e). Furthermore, $\Delta 5'$ 697 protein levels were dramatically reduced, suggesting that the inability to bind EZH2 may destabilize this protein (Fig. 8). Given that interpretation, it is unclear whether the reduced activity of the protein expressed from the $\Delta 5'$ 697 construct is a consequence of its inability to bind histones or its reduced ability to bind EZH2 and resultant instability. Finally, as a third explanation, it is possible that reduced $\Delta 5'$ 697 protein levels could be due to reduced expression. While formally possible, this explanation is less likely, given that the $\Delta 5'$ 541 and $\Delta 5'$ 697 constructs have the same promoters, enhancers, 5' and 3' untranslated regions, and canonical Kozak sequences.

EED-1, EED-3, and EED-4 translation start sites map to GUG 274–276, AUG 451–453, and AUG 493–495

In mapping regions and motifs in EED required for H3K27 methylation, we have also defined *in vivo* the translation start sites for three of the four EED isoforms. Previous work has suggested that the four EED isoforms are generated by translation initiating at GUG 169–171, GUG 274–276, AUG 451–453, and AUG 493–495, respectively.^{25,44} This interpretation followed from early studies by Denisenko and Bomsztyk, who assessed EED proteins translated in a cell-free system.⁴⁴ Subsequently, Kuzmichev *et al.* generated an isoform-restricted EED antibody, called α NT, which was raised to peptides encoded by *Eed* cDNA sequences from positions 258–453 (M26). Because α NT recognized EED-1 and EED-2 but not EED-3 and EED-4, those results demonstrated that EED-1 and EED-2 must include the α NT epitope and must initiate translation upstream of M451–453, as predicted by Denisenko and Bomsztyk.⁴⁴

Here, we have directly assessed the identity of three EED translation start sites by a combination

approach involving deletion mapping, forced translation from reported start sites, and site-directed mutagenesis of candidate initiation codons. These experiments definitively map EED-1, EED-3, and EED-4 start sites to GUG 274–276, AUG 451–453, and AUG 493–495, respectively. Importantly, because the informative α NT antibody utilized by Kuzmichev *et al.* recognizes amino acids that would be present not only in a hypothetical protein initiating at GUG 169–171 but also in a protein initiating at GUG 274–276, the EED-1 initiation site reported here, the immunoblotting data from Kuzmichev *et al.* are fully consistent with the results presented in the present work, although our conclusions disagree with the conclusions made by those authors.²⁵

On the other hand, these results do directly contradict the earlier work of Denisenko and Bomsztyk.⁴⁴ Denisenko and Bomsztyk assessed EED proteins expressed from truncated and engineered *Eed* cDNAs comparable to those utilized here. However, whereas we assessed translation from those messages in living cells, Denisenko and Bomsztyk utilized a rabbit reticulocyte cell-free system. In their assay, a truncation that would have deleted the 5' 222 nucleotides in our constructs eliminated EED-1 expression, although a truncation extending 35 nucleotides further ($\Delta 257$) did not disrupt EED-1 expression in our assay (Fig. 3b). Additionally, Denisenko and Bomsztyk observed apparent upregulation of EED-1 after mutating GUG 169–171 and flanking sequences to more closely resemble a canonical initiation sequence, whereas a similar experiment in our assays produced a protein larger than EED-1 (Fig. 4b).

The differences in our respective methodologies may account for the discrepancy in our results. The major advantage to the transient transfection assay employed in this report is that it allows direct comparison of EED expressed from expression constructs to endogenous EED expressed in the same cell type. Moreover, a protein initiating at GUG 169–171 is predicted to be less than 4 kDa larger than a protein initiating at GUG 274–276. In our hands, resolving proteins initiating at those sites required large gels run at low voltage. Accordingly, differences in electrophoresis conditions could also explain discrepancies in our results.

Conclusions

The biological functions of the four reported EED isoforms remain unclear. Both *in vivo* and *in vitro*, all isoforms analyzed appear capable of facilitating PRC2-dependent histone methylation (Fig. 5; Ref. 27). Accordingly, it is uncertain whether the four isoforms are even functionally necessary in living cells. However, if isoform-specific functions do exist, the *in vivo* characterization of EED isoform start sites reported here is an important first step in defining those functions. One critical function common to the EED isoforms is the stabilization of EZH2 in PRC2

complexes. Here, we have demonstrated that the five WD-40 motifs in EED are critical to this stabilization and to PRC2-mediated H3K27 methylation.

Materials and Methods

Cell lines and culture

CD1 MEFs were plated on gelatin-coated coverslips and grown to subconfluency. Wild-type ES cell line E14, wild-type ES cell line 25.5, and *Eed*^{17Rn5-3354SB/17Rn5-3354SB} (herein referred to as *Eed* mutant or *Eed*^{-/-}) ES cell line 21 were grown first on irradiated fibroblast feeders before being plated on coverslips without feeders and grown to subconfluency.^{31,53} In transfection experiments, ES cells plated on coverslips were transfected using Lipofectamine 2000™ (Invitrogen) and harvested 24–48 h later.

Immunofluorescence

Cells on coverslips were permeabilized with CSK buffer (100 mM NaCl, 300 mM sucrose, 3 mM MgCl₂, 10 mM Pipes, pH 6.8) and fixed in 4% paraformaldehyde that had been diluted in phosphate-buffered saline (PBS). To stain, we washed cells in PBS and incubated them with blocking buffer (PBS, 5% goat serum, 0.2% Tween-20, 0.2% fish skin gelatin). After blocking, samples were incubated with primary antibody [anti-H3K27me1 (Upstate), anti-H3K27me2,¹³ anti-H3K27me3,¹³ or anti-HP1- α (Upstate)], which had been diluted 1:250 in blocking buffer. Subsequently, cells were washed in PBS/0.2% Tween-20 and then incubated with secondary antibody [Goat Anti-Rabbit Alexa 594 (Molecular Probes)] diluted 1:250 in blocking buffer. Blocking and antibody incubations were always performed in a humid chamber at 37 °C. Subsequently, cells were washed again in PBS/0.2% Tween-20 and mounted with Vectashield™ containing DAPI (Vector Laboratories). Stained slides were visualized by fluorescence microscopy; then, black-and-white images were captured with a Spot CCD digital camera before being pseudocolored and merged with Spot software V3.5.9 (Diagnostic Instruments Inc.).

Plasmid construction

For mapping and rescue experiment, constructs expressing *Eed* cDNAs were cloned into the shuttle TA-cloning vector pGEM®-T Easy (Promega) and then subcloned into EcoRI-digested pTarget™ (Promega) by conventional molecular biology techniques. The orientation and identity of all inserted sequences were confirmed by fully sequencing the cDNA and the cloning junctions.

Eed cDNAs were truncated by PCR, utilizing forward primers that annealed within the *Eed* cDNA and reverse primers anchored in pTarget (primer sequences available upon request). Site-directed point mutations were generated by standard methods. Briefly, primers spanning EED-3 and EED-4 start sites but harboring ATG → ATA mutations were used as forward primers with a reverse primer anchored in pTarget. Finally, in order to engineer strong translation start sites, we included in-frame consensus Kozak(GCCACC)-ATG 5' extensions on forward primers.

For YFP fragment complementation assays, PRC-amplified *Eed* and *Ezh2* fragments were cloned into split

YFP expression vectors.⁵² Briefly, *Eed* PCR products were TA cloned into pGEM®-T Easy (Promega) and then subcloned into the NotI and ClaI sites at the 5' end of Venus 2. Similarly, PCR-amplified *Ezh2* fragments were TA cloned before subcloning into the BspEI and XbaI sites at the 3' end of Venus 1.

Western blotting

Wild-type ES cell line 25.5, wild-type ES cell line E14, mock-transfected *Eed* mutant ES cell line 21, and *Eed* mutant ES cell line 21 transfected with various expression constructs were harvested in urea lysis buffer (7.5 M urea, 0.01 M Tris, pH 8.0, 0.1 M NaH₂PO₄) 48 h after transfection. Urea lysates were also harvested from mouse *Wap-T₁₂₁* mammary tumor tissue, which is generated by tissue-specific expression of T₁₂₁, a fragment of SV40 T antigen that interferes with the function of retinoblastoma family proteins.⁵⁴ Proteins were separated on 10% SDS-PAGE gels in Tris-glycine running buffer and transferred to Immobilon-PVDF membranes (Bio-Rad) in Tris-glycine methanol transfer buffer. Membranes were blocked in 5% nonfat dried milk (NFDM; Food Lion)/Tris-buffered saline Tween-20 (TBST; 50 mM Tris-HCl, pH 7.4, 150 mM NaCl, 0.1% Tween-20) and then incubated overnight at 4 °C with a mouse monoclonal anti-EED antibody⁵⁵ diluted 1:400 in 3% NFDM/TBST or Sigma M2 anti-FLAG antibody diluted 1:1000 in 3% NFDM/TBST. Membranes were vigorously washed in TBST and then incubated overnight at 4 °C with horseradish-peroxidase-conjugated goat anti-rabbit antibody (Pierce) diluted 1:3000 in 5% NFDM/TBST. Membranes were then vigorously washed in TBST and TBS (50 mM Tris-HCl, pH 7.4, 150 mM NaCl) before adding Super-Signal West Dura Extended Duration Substrate (Pierce) developing reagents. Finally, blots were exposed to film and developed.

YFP fragment complementation assays

E14 ES cells grown on coverslips were transiently transfected with YFP fragment complementation plasmids expressing GCN4, EZH2, or EED. Twenty-four hours after transfection, the cells were washed in 1× PBS, fixed for 20 min in 4% paraformaldehyde, and then washed thoroughly in 1× PBS. Fixed cells were mounted in a polyvinyl alcohol/glycerol mounting medium with 2% N-propyl gallate and 0.25% 1,4-diazabicyclo[2,2,2]octane. Subsequently, YFP was visualized with a Carl Zeiss LSM5 fluorescence microscope.

For quantification of YFP fluorescence, cells were harvested 24 h posttransfection, trypsinized, washed in 1× PBS, and then fixed in 1% formaldehyde/PBS. Relative levels of fluorescence were determined using a CyAn ADP flow cytometer (DakoCytomation, Inc.). For each sample, at least 200,000 total events were analyzed. Percentage of YFP-positive cells and YFP fluorescence intensity were determined using DakoCytomation Summit software version 4.3.

Acknowledgements

We thank A. Otte (EED antibody), T. Jenuwein (H3K27me2 and H3K27me3 antibodies), S. Bultman

(*Wap-T₁₂₁* tumor tissue), and S. Michnick (YFP fragment complementation assay vectors) for generously sharing reagents. Additionally, we are grateful to B. Bagnell of the University of North Carolina Microscopy Services Laboratory, D. Ciavatta, and the University of North Carolina Flow Cytometry Core Facility for technical assistance. Finally, we thank D. Ciavatta and S. Chamberlain for helpful comments on the manuscript. This research was supported by a National Institutes of Health grant to T.M. and a Howard Hughes Medical Institute predoctoral fellowship to N.D.M.

Supplementary Data

Supplementary data associated with this article can be found, in the online version, at [doi:10.1016/j.jmb.2007.10.040](https://doi.org/10.1016/j.jmb.2007.10.040)

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