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# Erect Wing facilitates context-dependent Wnt/Wingless signaling by recruiting the cell-specific Armadillo-TCF adaptor Earthbound to chromatin

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## SUMMARY

During metazoan development, the Wnt/Wingless signal transduction pathway is activated repetitively to direct cell proliferation, fate specification, differentiation and apoptosis. Distinct outcomes are elicited by Wnt stimulation in different cellular contexts; however, mechanisms that confer context specificity to Wnt signaling responses remain largely unknown. Starting with an unbiased forward genetic screen in *Drosophila*, we recently uncovered a novel mechanism by which the cell-specific co-factor Earthbound 1 (Ebd1), and its human homolog jerky, promote interaction between the Wnt pathway transcriptional co-activators  $\beta$ -catenin/Armadillo and TCF to facilitate context-dependent Wnt signaling responses. Here, through the same genetic screen, we find an unanticipated requirement for Erect Wing (Ewg), the fly homolog of the human sequence-specific DNA-binding transcriptional activator nuclear respiratory factor 1 (NRF1), in promoting contextual regulation of Wingless signaling. Ewg and Ebd1 functionally interact with the Armadillo-TCF complex and mediate the same context-dependent Wingless signaling responses. In addition, Ewg and Ebd1 have similar cell-specific expression profiles, bind to each other directly and also associate with chromatin at shared genomic sites. Furthermore, recruitment of Ebd1 to chromatin is abolished in the absence of Ewg. Our findings provide in vivo evidence that recruitment of a cell-specific co-factor complex to specific chromatin sites, coupled with its ability to facilitate Armadillo-TCF interaction and transcriptional activity, promotes contextual regulation of Wnt/Wingless signaling responses.

**KEY WORDS:**  $\beta$ -Catenin, Context specificity, Earthbound, Erect wing, TCF, Wnt/Wingless, *Drosophila*

## INTRODUCTION

The evolutionarily conserved Wnt/Wingless signal transduction pathway is used repetitively during animal development to regulate cell proliferation, fate determination, differentiation and cell death; deregulation of this pathway is associated with multiple congenital disorders and types of cancer (Clevers, 2006; MacDonald et al., 2009). In response to interaction between Wnt ligands and their transmembrane receptors, the activity of a complex that includes Axin, Adenomatous Polyposis Coli (APC) and glycogen synthase kinase 3 is attenuated, thereby reducing degradation of  $\beta$ -catenin/Armadillo, a key Wnt pathway transcriptional co-activator. Stabilized  $\beta$ -catenin accumulates in cytoplasm and translocates into nucleus, where it associates with DNA-binding transcription factors of the T-cell factor/lymphoid enhancer factor (TCF/LEF) family to activate Wnt pathway target genes (MacDonald et al., 2009).

Diverse target genes are activated upon Wnt stimulation in distinct developmental and cellular contexts, whereas universally induced Wnt target genes are relatively rare (Logan and Nusse, 2004; Clevers, 2006). As the  $\beta$ -catenin-TCF complex is expressed ubiquitously, how does Wnt signaling elicit distinct outcomes in different cells or in the same cell at different stages of differentiation? Extensive efforts during the past three decades have identified core Wnt pathway components that mediate signaling in all cells; yet how cell identity modulates specific Wnt signaling outcomes remains largely unknown.

To identify new factors that promote Wnt signaling, we performed unbiased forward genetic screens in *Drosophila*, anticipating that we might identify core Wnt pathway components as well as cell-specific factors that confer context-specificity to Wnt signaling responses. Indeed, we identified alleles of the well-characterized Wnt pathway transcriptional co-activator Legless/BCL9 (Kramps et al., 2002; Benchabane et al., 2011), as well as alleles of Apc that revealed an unexpected positive regulatory role (Takacs et al., 2008). Furthermore, we recently identified a novel cell-specific factor, Earthbound 1 (Ebd1), that functions as a context-specific regulator of Wingless signaling responses (Benchabane et al., 2011). We found that Ebd1 and its human homolog jerky, which contain centromere binding protein B (CENPB)-type DNA-binding domains, physically associate with and bridge  $\beta$ -catenin/Armadillo and TCF, thereby promoting stability of the  $\beta$ -catenin-TCF complex and recruitment of  $\beta$ -catenin to chromatin (Benchabane et al., 2011). These findings uncovered a novel mechanism by which cell-specific adaptors facilitate interaction between  $\beta$ -catenin/Armadillo and TCF to promote contextual regulation of Wnt signaling responses.

Here, through the same genetic screen, we identify an unanticipated role for the sequence-specific DNA binding transcription factor Erect Wing (Ewg), the fly homolog of human nuclear respiratory factor 1 (NRF1), in the context-specific regulation of Wingless signaling responses. Unexpectedly, Ewg mediates the same subset of Wingless-dependent developmental processes that require Ebd1. Our characterization of Ewg function sheds light on a novel mechanism in which a cell-specific co-factor complex associates with both the core Wingless pathway transcription complex and with distinct chromatin sites to facilitate contextual regulation of Wingless signaling.

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## MATERIALS AND METHODS

### Fly stocks

*ewg<sup>PI</sup>* was isolated in an ethyl methanesulfonate-based mutagenesis screen (Lewis and Bacher, 1968) for dominant suppressors of photoreceptor apoptosis in *Apc1<sup>O8</sup>* mutants. Other lines used were *ewg<sup>1</sup>*, *ewg<sup>2</sup>*, *elav-ewg* (DeSimone and White, 1993), *Apc1<sup>O8</sup>* (Ahmed et al., 1998), *GMR-p35* (Hay et al., 1994), *UAS-TCF<sup>DN</sup>* (van de Wetering et al., 1997) and *ebd1-Gal4* (Benchabane et al., 2011). Canton S flies were used as wild-type controls. Fly crosses were performed at 25°C unless otherwise indicated.

### Mosaic clone analysis

To generate *ewg<sup>2</sup>* germline clones, *ewg<sup>2</sup>* FRT101 was crossed to *ovo<sup>D</sup>* FRT101; *hs-flp* flies. First and second instar larvae were heat shocked at 37°C for 2 hours and female progeny were crossed to *ewg<sup>2</sup>* mutant males carrying duplication *Dp(1;Y)y2611*. *ewg<sup>2</sup>* maternal/zygotic mutant embryos were recognized by lack of Ewg staining.

To generate *ewg*-null mutant eye clones, *ewg<sup>2</sup>* FRT19A flies were crossed to *arm-lacZ* FRT19A; *ey-flp*; retinas from progeny were fixed at 35 to 40 hours APF and stained with Homothorax or cleaved Caspase 3 antibody. *ewg* mutant clones were identified by staining with Ewg antiserum or  $\beta$ -Gal antibody.

### Plasmids

*ewg* cDNA was provided Kalpana White (DeSimone et al., 1996). For bacterial expression, *ewg* cDNA was cloned into the *EcoRI* sites of *pGST-Parallell* (Sheffield et al., 1999) to produce GST-Ewg. For expression in cultured cells, *ewg* cDNA was cloned into the *EcoRI* sites of *pFLAG-CMV-5a* (Sigma). *pProEx-Ebd1* and *pCMV5-Flag-Ebd1* have been described previously (Benchabane et al., 2011). cDNAs for human NRF1 (IMAGE clone 4397528) were obtained from Open Biosystems. For transgenic expression of NRF1 in flies, a V5 epitope tag (GGTAAGCCTATCCCTAACCTCTCCTCGGTCTCGATTCTACG) was added to the N terminus of NRF1, and V5-tagged *NRF1* was cloned into the *pUAST* vector (Brand and Perrimon, 1993) between *XhoI* and *XbaI* sites.

### Immunohistochemistry

Embryos, larval imaginal discs, pupal retinas and pupal flight muscles were fixed and stained as described previously (Rothwell and Sullivan, 2000; Fernandes et al., 2005; Takacs et al., 2008). Primary antibodies used for immunostaining were: guinea pig anti-Ebd1, 1:4000 (Benchabane et al., 2011); rabbit anti-Ewg, 1:1000 (DeSimone and White, 1993); mouse (9F8A9) or rat (7E8A10) anti-Elav, 1:10 (DSHB) (O'Neill et al., 1994); mouse 22C10, 1:50 (DSHB) (Hummel et al., 2000), mouse anti-FasII, 1:20 (DSHB) (Grenningloh et al., 1991), mouse BP102, 1:40 (DSHB) (Seeger et al., 1993); mouse anti-Eve, 1:10 (DSHB); rabbit anti-dMef2, 1:800 (Bour et al., 1995); guinea pig anti-Homothorax, 1:500 (Abu-Shaar et al., 1999); rabbit anti-Homothorax, 1:1000 (Pai et al., 1998); and rabbit anti-cleaved Caspase 3, 1:1000 (Cell Signaling). Secondary antibodies used for immunostaining were goat or donkey Alexa Fluor 488 or 568 conjugates at 1:200 to 1:400 (Molecular Probes), and goat or donkey Cy5 conjugates at 1:200 (Jackson Immunochemicals).

Larval salivary gland polytene chromosomes were prepared as described by Christine James, Paul Badenhurst and Bryan Turner (University of Birmingham, UK) (<http://www.epigenome-noe.net/WWW/researchtools/protocol.php?protid=1>), with the following modifications. Briefly, salivary glands were fixed in 4% formaldehyde, 1% Triton X-100 for 30 seconds, followed by 45% acetic acid, 4% formaldehyde, 2.5% lactic acid for 2 minutes, and spread on glass slides under borosilicate cover slips using the graphite end of a pencil. Slides were blocked with 3% BSA and 0.1% Triton X-100 in PBS prior to staining. Cytological subdivisions in polytene chromosomes were mapped with reference to the DAPI staining pattern (Ashburner, 1972; Sorsa, 1988).

Confocal images shown in Fig. 5D-K, Fig. 6A-C, and supplementary material Fig. S3 and Fig. S7 were obtained with a Nikon A1 confocal microscope. Other fluorescent polytene chromosomes images were obtained on a Zeiss Axioskop 2 microscope. Fluorescent images of other tissues were obtained on a Leica TCS SP UV confocal microscope.

### Histology

Adult eyes and thoraces were fixed as described previously (Cagan and Ready, 1989; Wolff, 2000; Fernandes et al., 2005), embedded in plastic resin (Durcupan, Fluka), sectioned to 1  $\mu$ m and 5  $\mu$ m, respectively, and stained with Toluidine Blue.

### Cell lines and transfection

HEK293T cells were cultured in Dulbecco's modified essential medium, supplemented with 10% fetal bovine serum and transiently transfected using calcium phosphate-DNA precipitation (Hayashi et al., 1997).

### Immunoprecipitation and GST interaction assays

For immunoprecipitations, calcium phosphate-transfected HEK293T cells were lysed 48 hours post-transfection in lysis buffer [50 mM Tris-HCl (pH 7.4), 150 mM NaCl, 1 mM EDTA, 0.5% Triton X-100] containing protease and phosphatase inhibitors cocktail (Thermo Scientific). Cell extracts were immunoprecipitated with anti-FLAG M2 antibody (Sigma) and collected using Protein-A/G PLUS-Agarose beads (Santa Cruz Biotechnology). Immunoprecipitates were separated by SDS-PAGE and immunoblotted with anti-V5 (Invitrogen) and anti-Flag antibodies.

For GST interaction assays, His-Ebd1 was purified from bacterial lysates using TALON metal affinity resin (Clontech). His fusion proteins were eluted off the resin using 150 mM imidazole. GST-Ewg proteins were purified using Glutathione Sepharose 4 Fast Flow beads (GE Healthcare) and incubated with His-Ebd1. Beads were washed in 50 mM Tris-HCl (pH 7.4), 150 mM NaCl, 1 mM EDTA, 0.1% Triton X-100 and associated proteins were detected by immunoblotting with anti-Ebd1 antibody.

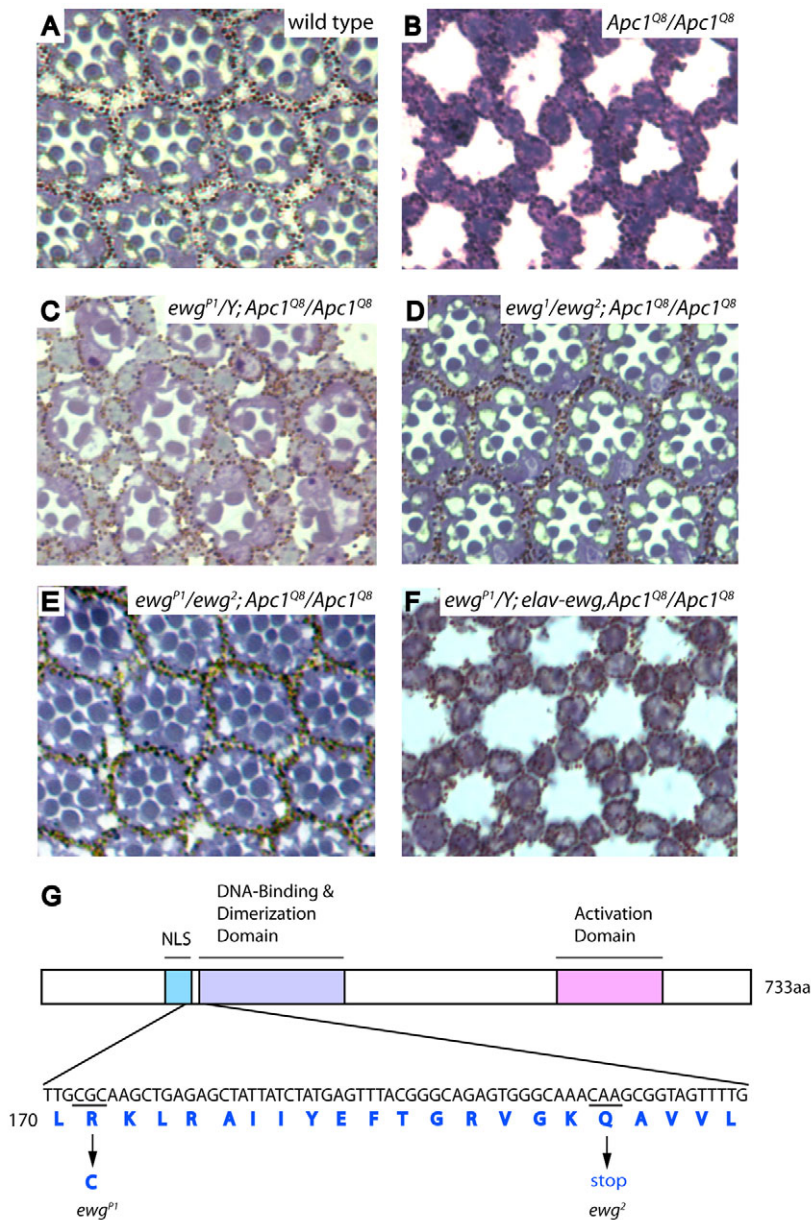
## RESULTS

### Identification of Erect Wing in a forward genetic screen for factors that promote Wingless signaling

During metamorphosis, Wingless signaling induces apoptosis of photoreceptors at the *Drosophila* retinal periphery, thereby selectively eliminating excess neurons (Tomlinson, 2003; Lin et al., 2004). Upon loss of Adenomatous polyposis coli 1 (*Apc1*), constitutive Wingless pathway activation results in apoptosis of all retinal photoreceptors (Ahmed et al., 1998). This *Apc1* mutant phenotype provides a sensitive assay for Wingless pathway activity, as reducing the gene dose of *armadillo*, *TCF* or the Wingless pathway transcriptional co-activator *legless*, by one half is sufficient to suppress the apoptosis (Ahmed et al., 1998; Benchabane et al., 2008). Therefore, to identify components that promote Wingless pathway activation, we performed genetic screens for suppressors of retinal apoptosis in the *Apc1* mutant (Takacs et al., 2008; Benchabane et al., 2011).

As noted above, we recently identified Earthbound 1 (*Ebd1*), a novel member in a protein family containing Centromere Binding Protein B (CENPB)-type DNA binding domains (Masumoto et al., 1989) in this genetic screen (Benchabane et al., 2011). *Ebd1* is expressed primarily in myocytes and neurons, and is required specifically for Wingless-dependent flight muscle development. *Ebd1*, and its human homolog Jerky, function as cell-specific adaptors that interact with both  $\beta$ -catenin/Armadillo and TCF to facilitate formation and promote activity of this core Wnt/Wingless pathway transcription complex.

From the same genetic screen for factors that promote Wingless signaling, we identified an allele on the X chromosome that we termed *PI*. In *Apc1* mutants, partial suppression of photoreceptor apoptosis is observed in hemizygous *PI* mutant males and homozygous *PI* mutant females (Fig. 1C; supplementary material Fig. S1A). Meiotic recombination revealed that *PI* maps to the distal tip of the X chromosome, and is closely linked to *yellow*. Among genes that also map to the X chromosome tip is *erect wing*



**Fig. 1. Ewg is required for photoreceptor apoptosis in response to *Apc1* loss.** (A–F) Cross-sections through adult retinas. (A) Eight photoreceptors are present in each ommatidium with seven photoreceptors visible in this section, arrayed in a trapezoidal pattern. (B) Upon *Apc1* loss, all photoreceptors undergo apoptosis. The pigment cell lattice remains intact. (C–E) Photoreceptor apoptosis is prevented in hemizygous *ewg<sup>P1</sup>* (C), transheterozygous *ewg<sup>1</sup>/ewg<sup>2</sup>* (D) and *ewg<sup>P1</sup>/ewg<sup>2</sup>* mutants (E). (F) Exogenous *ewg* expression driven by the *elav* promoter restores apoptosis in *ewg<sup>P1</sup>; Apc1<sup>Q8</sup>* double mutants. (G) Schematic representation of the Ewg protein. A C-to-T substitution is found in *ewg<sup>P1</sup>*, resulting in an arginine to cysteine substitution in amino acid 171. In *ewg<sup>2</sup>*, an early stop codon at amino acid 178 is caused by substitution of C to T; no mutation was detected in the *ewg*-coding region in the *ewg<sup>1</sup>* allele (G. Gill, personal communication).

(*ewg*), the fly homolog of human nuclear respiratory factor 1 (NRF1), a sequence-specific DNA binding transcriptional activator (Fleming et al., 1982; Fazio et al., 2001; Virbasius et al., 1993). We observed that when exposed to carbon dioxide anesthesia, *P1* mutant flies hold their wings in an upright or ‘erect wing’ position, and therefore we sought to determine whether *P1* is an *ewg* mutant allele, and whether two previously isolated mutations in *ewg*, the null allele *ewg<sup>2</sup>* and the hypomorphic allele *ewg<sup>1</sup>* (Fleming et al., 1982), also suppress *Apc1* mutant apoptosis. Transheterozygotes of *ewg<sup>2</sup>* and *ewg<sup>1</sup>* markedly suppress photoreceptor apoptosis in *Apc1* mutants (Fig. 1D), as do transheterozygotes of *ewg<sup>2</sup>* and *P1* (Fig. 1E), suggesting that *P1* is an *ewg* loss-of-function allele. Furthermore, expression of a transgene encoding Ewg, driven by the neuronal-specific *elav* promoter, restores apoptosis in *P1; Apc1* double mutants (Fig. 1F), whereas expression of this *ewg* transgene in wild-type photoreceptors has no effect (supplementary material Fig. S1B), confirming identification of the correct gene. We sequenced the *ewg* coding region in *ewg<sup>P1</sup>* and detected a C-to-T

point mutation that results in a substitution of arginine to cysteine at amino acid 171, a residue in a region that is highly conserved in human NRF1 (Fig. 1G). Together, these data indicate that Ewg is required for Wingless pathway-dependent photoreceptor apoptosis resulting from *Apc1* loss.

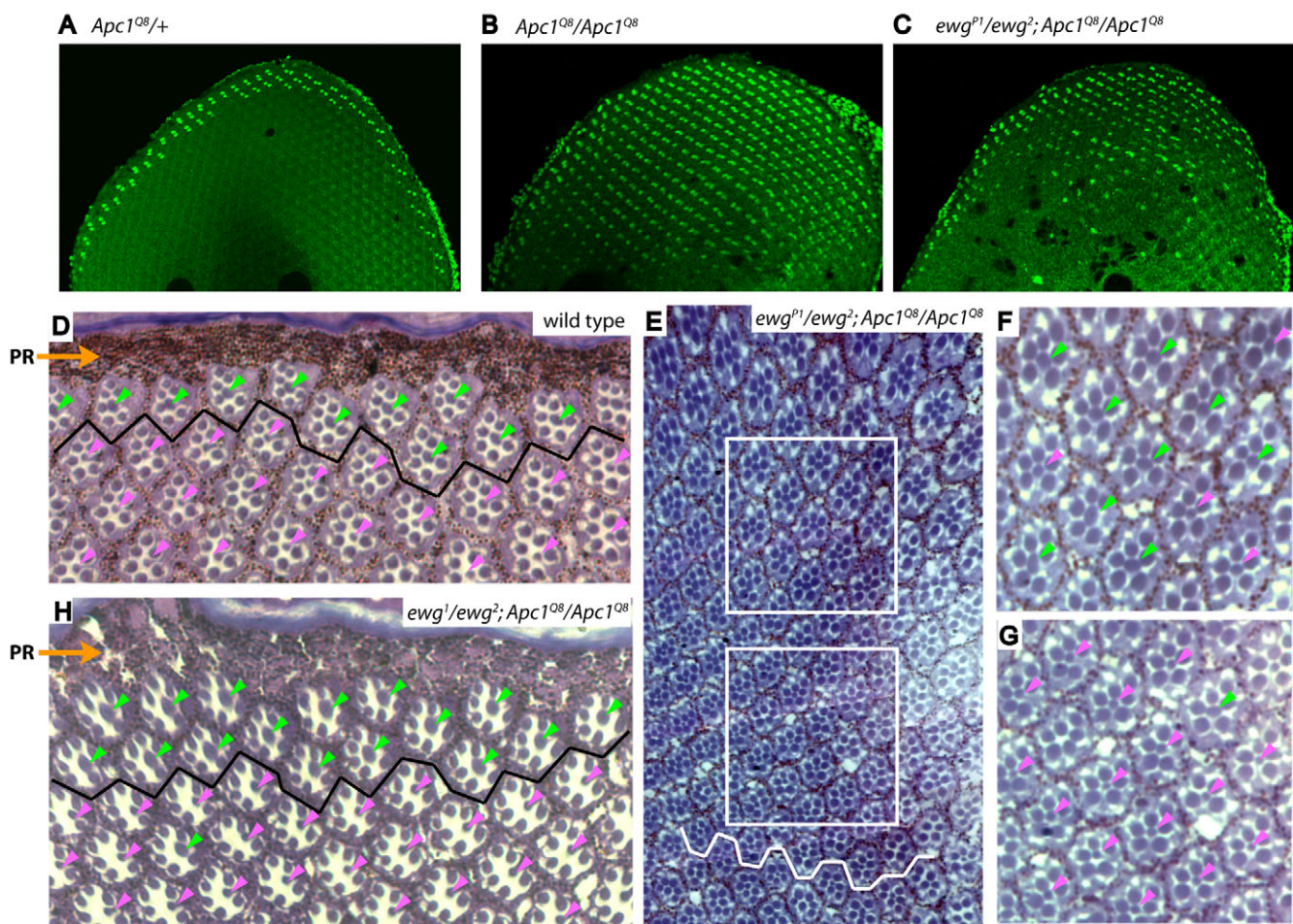
### Ewg promotes Wingless signal transduction

To determine whether Ewg solely facilitates photoreceptor apoptosis, or also promotes other Wingless pathway-dependent processes, we analyzed additional *in vivo* readouts. Prior to their apoptotic death, *Apc1* mutant photoreceptors display two other defects caused by constitutive activation of Wingless signaling: a reduction of photoreceptor length and a misspecification of photoreceptor fate (Ahmed et al., 1998; Benchabane et al., 2008); high level Wingless signaling is required for photoreceptor apoptosis, whereas intermediate level Wingless signaling is sufficient to induce shortened photoreceptor length and aberrant photoreceptor fate specification (Tomlinson, 2003; Benchabane et

al., 2008). Reduction of the Ewg level compromises the level of Wingless signaling in *Apc1* mutant retinas. A partial reduction of Ewg activity, as found in *ewg<sup>1</sup>* homozygotes, reduces Wingless pathway-dependent photoreceptor apoptosis (supplementary material Fig. S2E,F); further reduction of Ewg levels, as found in transheterozygotes of the *ewg<sup>2</sup>* null allele and the *ewg<sup>1</sup>* hypomorphic allele, or in transheterozygotes of *ewg<sup>2</sup>* and *ewg<sup>P1</sup>*, prevents both Wingless pathway-dependent apoptosis and shortening of photoreceptor length (Fig. S2G,H; data not shown). Together, these findings indicate that Ewg facilitates Wingless signaling.

In addition, we examined whether Ewg is also required for mis-specification of photoreceptor fate that results from constitutive Wingless pathway activation upon *Apc1* loss. In wild-type flies, Wingless signaling specifies inner photoreceptors restricted to the dorsal retinal periphery, termed the dorsal rim area (DRA), to adopt

a fate that permits polarized light perception, instead of color light perception (Tomlinson, 2003; Wernet et al., 2003) (Fig. 2A,D). Intermediate level Wingless signaling caused by *Apc1* loss results in aberrant specification of DRA fate in all inner photoreceptors in the dorsal half of the retina, as evidenced by their enlarged rhabdomeres and expression of the transcription factor Homothorax (Hth) (Tomlinson, 2003; Wernet et al., 2003; Benchabane et al., 2008; Takacs et al., 2008) (Fig. 2B). Attenuation of Ewg activity reduces the number of inner photoreceptors adopting a DRA fate in the *Apc1* mutant, as revealed by wild-type rhabdomere size and loss of ectopic Hth expression. We observe partial suppression of DRA fate upon partial reduction of Ewg levels, as found in transheterozygotes of *ewg<sup>P1</sup>* and *ewg<sup>2</sup>* (Fig. 2C,E-G), and nearly complete suppression upon further reduction of Ewg levels, as found in transheterozygotes of *ewg<sup>1</sup>* and *ewg<sup>2</sup>* (Fig. 2H). These findings provide further evidence that Ewg facilitates Wingless



**Fig. 2. Ewg is required for aberrant photoreceptor cell fate specification in response to *Apc1* loss.** (A-H) Confocal images of 35–40 hour pupal retinas stained with Homothorax (Hth) antisera (A–C) and cross-sections through adult retinas (D–H). In all panels, dorsal is towards the top. (A,D) In wild-type and *Apc1* heterozygous retinas, inner photoreceptors (PRs) with DRA fate are restricted to one or two rows of ommatidia at dorsal retinal periphery, as indicated by Hth staining in pupae (A) and enlarged rhabdomeres located close to pigment rim (PR) in adults (D, ommatidia above black line, green arrowheads). (B) In *Apc1<sup>Q8</sup>* mutants, constitutive activation of Wingless signaling induces all inner PRs in entire dorsal retinal half to aberrantly express Hth. (C,E–G) In *ewg<sup>P1</sup>/ewg<sup>2</sup>; Apc1<sup>Q8</sup>* double mutants, partial reduction of Ewg activity leads to a reduction in the number of inner PRs expressing Hth and displaying enlarged rhabdomeres, which is most evident near dorsal-ventral equator (C,E). Higher magnifications of the upper and lower boxed areas in E are shown in F and G, respectively. Inner PRs closer to retinal dorsal edge mostly retain DRA fate, as indicated by enlarged rhabdomeres (green arrowheads in F); ommatidia closest to equator (white jagged line in E) mostly adopt a normal cell fate, as indicated by small rhabdomeres (pink arrowheads in G). (E) A montage of photographs compiled in Adobe Photoshop. (H) In *ewg<sup>1</sup>/ewg<sup>2</sup>; Apc1<sup>Q8</sup>* double mutants, inner PRs adopting a DRA fate (green arrowheads) is further reduced to only two ommatidial rows adjacent to the dorsal rim (above solid black line), indicating attenuation of Ewg activity prevents ectopic Wingless signaling in *Apc1* mutants.

signaling. To determine whether Ewg is required for Wingless signaling at the retinal periphery, we generated *ewg*-null mutant clones in otherwise wild-type pupal retinas. We analyzed photoreceptor DRA fate specification as revealed by Homothorax expression, and photoreceptors apoptosis, as revealed by cleaved Caspase 3 (Wernet et al., 2003; Lin et al., 2004; Yu et al., 2002). We detected no defects in DRA fate determination or photoreceptors death upon the loss of Ewg (data not shown), suggesting that Ewg is dispensable for these two Wingless-dependent processes during retinal development.

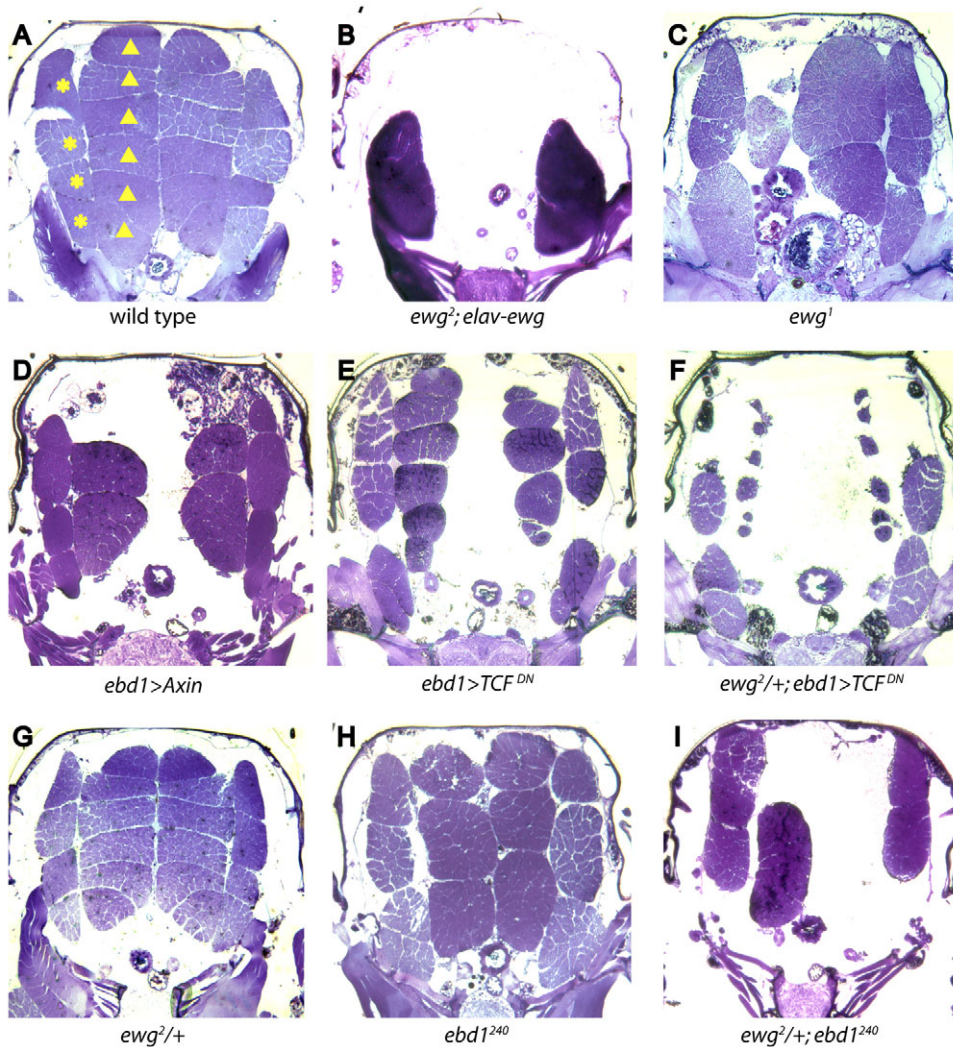
### Ewg facilitates Wingless-dependent flight muscle development

We next sought to identify Wingless-dependent developmental processes that require Ewg activity. Previous studies have revealed that Ewg is important for embryonic neuronal development and adult flight muscle formation (DeSimone and White, 1993). However, in *ewg* mutant embryos, we detect no defects in RP2 motor neuron specification or migration, two well-characterized processes that are dependent on Wingless signaling (Chu-LaGraff and Doe, 1993; Bhat, 2007) (supplementary material Fig. S3, Table S1). In addition, immunostaining of *ewg*-null mutant embryos for the neuronal markers 22C10, FasII and BP102, as well as the muscle marker Mef2, revealed no apparent defects (data not

shown). Furthermore, we detect no defects in Wingless-dependent epithelial cell fate specification (Nusslein-Volhard and Wieschaus, 1980; Wieschaus and Riggleman, 1987; Baker, 1988) (data not shown).

Therefore, we also examined processes that require Ewg activity at later developmental stages. The embryonic lethality resulting from loss of Ewg can be prevented by transgenic expression of Ewg solely in neurons, driven by the *elav* promoter; although these rescued flies are viable, they display loss of indirect flight muscles, resulting from Ewg inactivation in myoblasts (DeSimone et al., 1996) (Fig. 3B). Similarly, hypomorphic *ewg<sup>1</sup>* mutants display either complete loss of dorsal longitudinal muscles or marked reduction in flight muscle number, with enlargement of muscles that remain (Fleming et al., 1982) (Fig. 3C).

Wingless signaling is also required for flight muscle development (Sudarsan et al., 2001; Benchabane et al., 2011). We find that attenuation of Wingless signaling in *ebd1*-expressing cells, caused by overexpression of the negative regulator Axin, results in the reduction of flight muscle number and the enlargement of remaining flight muscles, a phenotype very similar to that observed in *ewg* mutants (Benchabane et al., 2011) (Fig. 3D). The similarity in flight muscle defects found upon attenuation of Ewg or Wingless pathway activity therefore suggested function in the same developmental process.



### Fig. 3. Ewg and Ebd1 are required for adult flight muscle development.

Cross-sections of adult thoraces. (A) In wild type, there are six dorsal longitudinal muscles (DLMs) and seven dorsal ventral muscles (DVMs) on each side of the midline, all DLMs (triangles) and four DVMs (\*) can be seen in this section. (B) DLMs and DVMs of normal number and morphology are lost, and instead are replaced by two large mis-shapen muscles in *ewg<sup>2</sup>; elav-ewg* mutants. (C) The majority of DLMs are lost in *ewg<sup>1</sup>* homozygous mutants. (D) Disrupting Wingless signaling by overexpressing Axin results in partial loss of IFMs. (E) Disrupting Wingless signaling by overexpressing a dominant-negative form of TCF results in reduction of muscle volume. (F) Halving the gene dose of *ewg* enhances muscle defects caused by dominant-negative TCF. (G) Heterozygous *ewg*-null mutants do not have muscle defects. (H) A subset of DLMs is missing in *ebd1<sup>240</sup>* homozygous mutants, and remaining muscles are abnormally enlarged. (I) Halving the gene dose of *ewg* markedly enhances muscle loss in *ebd1<sup>240</sup>* mutants.

To determine whether the requirement of *Ewg* in flight muscle may reflect a role in Wingless-dependent muscle development, we examined genetic interactions between *Ewg* and the Wingless pathway transcriptional co-activator TCF. Attenuation of Wingless signaling, caused by expression of dominant-negative TCF in *ebd1*-expressing cells, results in reduction of flight muscle volume (Benchabane et al., 2011) (Fig. 3E). Halving the *ewg* gene dose markedly enhances severity of flight muscle defects caused by dominant-negative TCF (Fig. 3F), yet causes no muscle defects in an otherwise wild-type background (Fig. 3G), suggesting that *Ewg* may promote TCF activity during flight muscle development. To rule out the possibility that this phenotypic enhancement is due to an increase of Gal4 activity, we examined whether halving *ewg* dose affects the *ebd1-Gal4* driver, but find no significant change in *Gal4* mRNA levels (supplementary material Fig. S4). Thus, the genetic interaction between *Ewg* and TCF (Fig. 3E,F), coupled with the similar mutant phenotypes resulting from attenuation of either *Ewg* or Wnt pathway activity (Fig. 3C,D), support the hypothesis that *Ewg* facilitates Wingless-dependent flight muscle development.

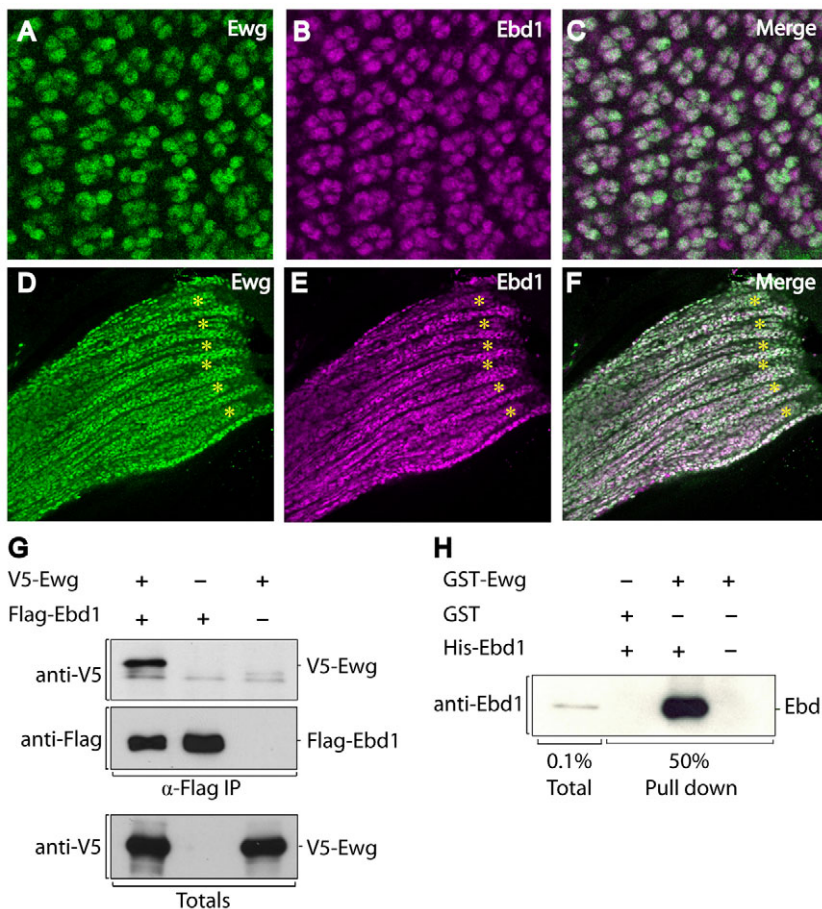
### ***Ewg* acts with *Ebd1* to facilitate Wingless-dependent flight muscle development**

Of note, loss of *Ebd1* activity also results in the reduction of flight muscle number and enlargement of remaining flight muscles, a phenotype similar to that observed in *ewg* mutants and upon disruption of Wingless signal transduction in *ebd1*-expressing cells (Benchabane et al., 2011) (Fig. 3C,D,H). Therefore, to determine whether *Ewg* and *Ebd1* may function in the same developmental

process, we used genetic interaction analyses. Halving *ewg* dose markedly enhances severity of muscle loss defects in *ebd1* mutants, but, as noted above, causes no muscle defects in an otherwise wild-type background (Fig. 3G-I). Their strong genetic interaction with each other and with Armadillo-TCF (Fig. 3E,F) (Benchabane et al., 2011), coupled with their similar mutant muscle phenotype (Fig. 3C,H), support the model that a functional relationship between *Ewg* and *Ebd1* promotes Wingless-dependent muscle development.

### ***Ewg* and *Ebd1* are expressed in neurons and myocytes**

As our functional data suggest that *Ewg* and *Ebd1* act in the same developmental process, we sought to determine whether these two proteins are also expressed in similar sets of cells. Both *Ewg* and *Ebd1* are expressed predominantly in muscles and in the nervous system (DeSimone and White, 1993; DeSimone et al., 1996; Koushika et al., 2000; Benchabane et al., 2011). To determine whether *Ebd1* and *Ewg* are expressed in the same subset of cells during different developmental stages, we performed immunostaining experiments using *Ebd1* and *Ewg* antisera in embryos, larvae and pupae. *Ebd1* and *Ewg* have identical distributions in the nervous system throughout all developmental stages examined (Fig. 4A-F; supplementary material Fig. S5A,B). During embryogenesis, *Ewg* and *Ebd1* are expressed in both the central and peripheral nervous system (supplementary material Fig. S5A-F). In larvae and pupae, *Ewg* and *Ebd1* are expressed in the brain, ventral ganglion, retinal photoreceptors and peripheral neurons (Fig. 4A-C, data not shown). In addition, *Ewg* and *Ebd1* colocalize in myoblasts and



**Fig. 4. *Ewg* and *Ebd1* display similar expression patterns and interact directly.** (A-F) Confocal images of pupal retinas at 24 hours after puparium formation (A-C), pupal myofibers and associated myoblasts (D-F). Tissues are immunostained with *Ewg* antiserum (A,D) and *Ebd1* antiserum (B,E). Merged images are shown in C,F. (A-F) During metamorphosis, *Ewg* and *Ebd1* colocalize in neurons and myocytes, including photoreceptors (A-C), developing myofibers (asterisks) and associated myoblasts (D-F). (G) V5-tagged *Ewg* and Flag-tagged *Ebd1* were co-transfected into HEK 293T cells. Cell lysates were subjected to immunoprecipitation with anti-Flag antibody, followed by immunoblotting with anti-V5 and anti-Flag antibody. Immunoblotting of total lysates is shown in lower panel. (H) GST interaction assay. Bacterially expressed GST-*Ewg* or GST control were purified and incubated with bacterially expressed and purified His-*Ebd1*. Protein bound to GST-*Ewg* or GST control was detected by immunoblotting with *Ebd1* antibody.



developing muscle fibers during metamorphosis (Fig. 4D-F). Although not detected in previous studies, we also observe low levels of Ewg that colocalize with Ebd1 in a segmented pattern outside the nervous system in embryos (supplementary material Fig. S5B-F). Though of low level, this Ewg signal is absent in *ewg*-null mutant embryos, confirming specificity of the signal (supplementary material Fig. S5G-R). The pattern and morphology of these Ewg-expressing cells are consistent with embryonic myoblasts. Available antibodies to myoblast markers are all rabbit derived, as is the Ewg antiserum, precluding direct double-staining experiments. However, these Ewg-expressing cells also express Ebd1, and the vast majority of Ebd1-expressing cells outside the nervous system are myoblasts (Benchabane et al., 2011), suggesting that Ewg is also expressed in embryonic myoblast cells. Their highly similar cell-type-restricted expression patterns, coupled with their similar mutant phenotypes and genetic interactions, are consistent with the model that Ewg and Ebd1 function together in the same cells during development.

### Ewg and Ebd1 associate directly with each other

To test the hypothesis that Ewg and Ebd1 function together as a complex, we examined the physical interaction between these two proteins. We performed co-immunoprecipitation from lysates of HEK293T cells ectopically expressing Ewg and Flag-tagged Ebd1. Immunoprecipitation with Flag antibody followed by immunoblotting with Ewg antiserum revealed that Ebd1 associates with Ewg (Fig. 4G). Furthermore, bacterially expressed GST-Ewg, but not the GST control, associates with bacterially expressed Ebd1, indicating that their interaction is direct (Fig. 4H). Indeed, supporting our conclusion that Ewg and Ebd1 interact directly, Ebd1 was found to interact with Ewg in a genome-wide yeast two-hybrid screen for *Drosophila* protein interactions (Giot et al., 2003). Together with their similar cell-type specific expression patterns, similar loss-of-function phenotypes, and their strong genetic interactions with each other and with Armadillo-TCF, these findings support the model that direct physical interaction between Ewg and Ebd1 underlies their role in mediating context-specific responses to Wingless signal transduction.

### Ewg and human NRF1 share chromatin association sites in vivo

NRF1, the human homolog of Ewg, contains a basic leucine zipper DNA-binding domain near its N terminus (Virbasius et al., 1993) (Fig. 5A). The NRF1 DNA-binding domain is highly conserved in Ewg, with more than 80% similarity (Efiok et al., 1994). Ewg has been found previously to interact with NRF1 consensus DNA-binding sequences, as revealed by gel mobility shift analysis, and to activate a transcriptional reporter containing multimerized NRF1 DNA-binding sites (Fazio et al., 2001). Together, these observations suggested that, like NRF1, Ewg might also associate with specific chromatin sites. To test this hypothesis in vivo, we took advantage of the presence of endogenous Ewg expression in third instar larval salivary glands (DeSimone and White, 1993), and analyzed the giant salivary gland polyploid chromosomes, which are produced by endoreplication and consist of ~500 closely aligned copies of each homolog that display a highly characteristic chromatin banding pattern. To determine whether Ewg associates with chromatin and, if so, to assess the distribution of Ewg on chromatin sites throughout the genome, we performed immunostaining experiments on squashed polytene chromosomes using Ewg antiserum. As a control for specificity in these studies, we rescued the viability of *ewg*-null mutants with a neuronal-

specific *elav-ewg* transgene that is not expressed in salivary glands (DeSimone and White, 1993). Ewg associates with discrete chromatin sites throughout the genome, and this signal is lost in *ewg*-null mutant chromosomes, confirming antibody staining specificity (Fig. 5B,C).

To determine whether Ewg and human NRF1 share similar chromatin association sites in vivo, we ectopically expressed NRF1 in larval salivary glands. We performed double-label immunostaining studies on polytene chromosomes to simultaneously detect NRF1 and endogenous Ewg. Ewg and NRF1 associate with many of the same chromatin sites throughout the genome (Fig. 5D-G). However, we wanted rule out the possibility that association of NRF1 with chromatin results solely from heterodimerization of NRF1 with Ewg, as NRF1 forms homodimers (Gugneja and Scarpulla, 1997), and the NRF1 dimerization domain is highly conserved in Ewg (Fig. 5A). Therefore, we expressed NRF1 in *ewg*-null salivary gland cells, and found that association of NRF1 with chromatin does not depend on the presence of Ewg (Fig. 5H-K). Thus, the similar chromatin banding patterns of Ewg and NRF1 suggest that Ewg and NRF1 may associate with specific chromatin sites through a similar mechanism.

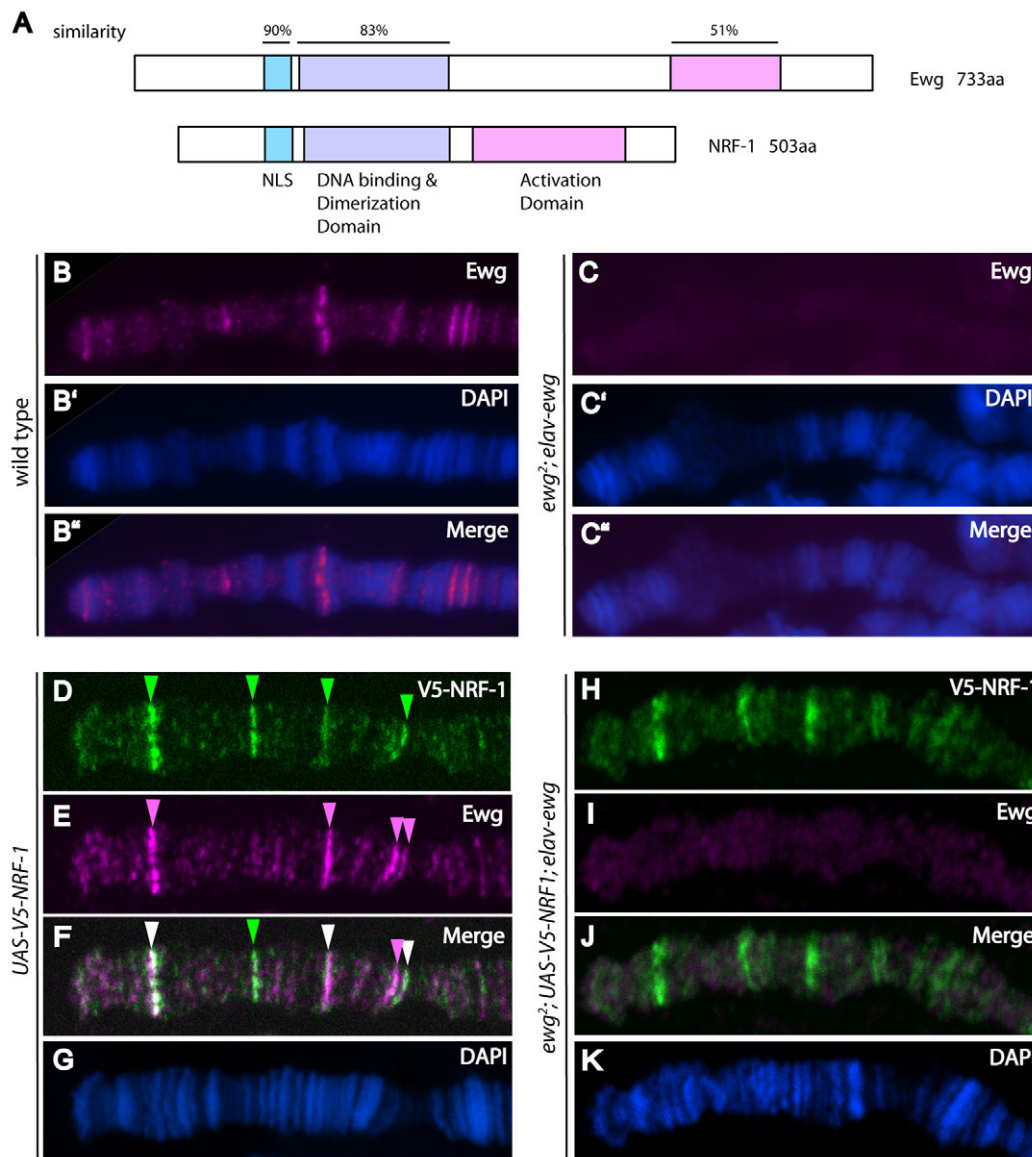
### Ebd1 associates with chromatin in vivo and shares chromatin association sites with Ewg

The physical and genetic interactions between Ewg and Ebd1 raise the possibility that these two factors are present in the same DNA-binding complex. To address this hypothesis, we sought to determine whether Ebd1 also associates with chromatin and, if so, whether Ebd1-associated chromatin sites are shared with Ewg. Immunostaining with Ebd1 antiserum also decorates discrete chromatin sites (supplementary material Fig. S6A), and this signal is lost in *ebd1*-null mutant chromosomes (supplementary material Fig. S6B), confirming Ebd1 antiserum specificity.

To determine whether any chromatin sites at which Ewg and Ebd1 associate are shared, we performed double labeling experiments on polytene squashes with Ebd1 and Ewg antisera. The banding pattern of Ebd1 on polytene chromosomes is highly similar to that of Ewg throughout the genome; many of the polytene bands at which Ebd1 and Ewg associate overlap with each other (Fig. 6A-C). Indeed, Ewg associates with nearly all, if not all, bands at which Ebd1 is found (supplementary material Table S2). However, Ewg associates with some polytene bands at which we detect no Ebd1 (supplementary material Table S2). For example, at the distal end of the X chromosome, Ebd1 associates with 6 of the 10 Ewg-associated bands, whereas Ebd1 is not detected on the other four Ewg-associated bands (Fig. 6F, pink arrowheads; supplementary material Table S2). Together, these findings support the model that Ebd1 and Ewg function as transcription co-factors that act as a complex at discrete promoter/enhancer sites.

### Ewg is required for Ebd1 recruitment to chromatin

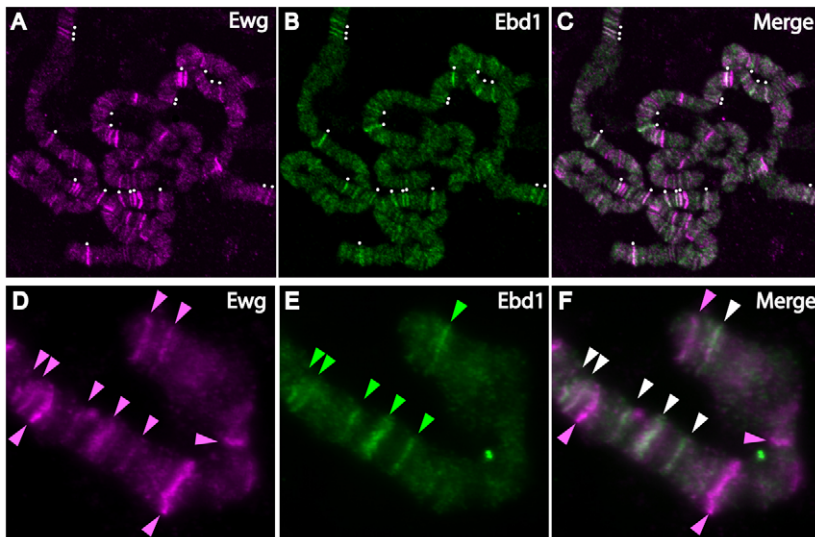
The direct physical interaction between Ebd1 and Ewg and their shared polytene banding patterns suggest that Ewg and Ebd1 associate with chromatin as a complex. To determine whether chromatin recruitment of either Ebd1 or Ewg requires the presence of the other, we analyzed the polytene banding pattern of Ebd1 in the absence of Ewg and, conversely, the banding pattern of Ewg in the absence of Ebd1. Ebd1 is readily detected on wild-type polytenes; by contrast, all association of Ebd1 with chromatin is abolished in the absence of Ewg (Fig. 7). For example, at the distal



**Fig. 5. Ewg and human nuclear respiratory factor 1 share chromatin association sites in vivo.** (A) Domains conserved between Ewg and mouse NRF-1 (Fazio et al., 2001). Domains in Ewg and NRF1 were aligned individually using ClustalW2. (B-C'') Squashes of larval third instar salivary gland polytene chromosomes, immunostained with Ewg antiserum (B,C). DNA is highlighted by DAPI staining (B',C'). Merged images are shown in B'' and C''. (B-B'') Wild type, distal tip of chromosome X. Ewg is detected on polytene chromosomes at discrete bands (B). (C-C'') Distal tip of chromosome X in *ewg*<sup>2</sup>; *elav-ewg* mutants. Ewg is not detected on polytenes in *ewg*<sup>2</sup>; *elav-ewg* mutant salivary glands (C). (D-K) Squashes of polytene chromosomes from third instar larvae, *UAS-V5-NRF-1* (D-G) and *ewg*<sup>2</sup>/*elav-ewg*/*UAS-V5-NRF-1* (H-K), immunostained with V5 antibody (D,H) and Ewg antiserum (E,I). Merged images are shown in F,J. DNA is highlighted by DAPI staining (G,K). V5-NRF-1 expression results from leaky expression of the *UAS-V5-NRF-1* transgene. Distal tip of chromosome 3L is shown (D-G). NRF1 retains ability to associate with chromatin (H), despite the absence of Ewg (I).

end of the left arm of chromosome 3, between cytological subdivisions 61A and 63C, Ewg and Ebd1 are both detected at the same three sites (Fig. 7A-D, arrowheads). By contrast, in *ewg* null mutant salivary glands, not only Ewg, but also Ebd1 is absent from these sites (Fig. 7E-G). This loss of Ebd1 chromatin association does not result from loss of Ebd1 expression, impaired Ebd1 nuclear localization, or global disruption of chromatin architecture, as Ebd1 is readily detected in nuclei of *ewg*-null mutant salivary gland cells, and the chromatin banding pattern, as revealed by DAPI staining, is intact in *ewg*-null mutants (supplementary material Fig. S7A-I; Fig. 7G). In addition, western blot analysis

indicates that there is no significant reduction in Ebd1 protein level in *ewg*-null mutant glands (supplementary material Fig. S7J). We also sought to determine whether association of Ewg with chromatin is dependent on Ebd1. In the absence of Ebd1, immunostaining of Ewg on polytene squashes reveals distinct bands that are highly similar, if not identical to Ewg-associated chromatin bands detected in wild-type salivary glands (Fig. 7H-J). Together, these findings suggest the existence of an Ewg-Ebd1 complex that associates with distinct chromatin sites, and reveal that recruitment of Ebd1 to chromatin occurs only in the presence of Ewg (Fig. 7K).



**Fig. 6. Ebd1 shares chromatin-association sites with Ewg.** (A-F) Squashes of polytene chromosomes from third instar larvae, immunostained with Ewg antiserum (A,D) and Ebd1 antiserum (B,E). Merged images are shown in C and F. (A-C) Ewg and Ebd1 overlap at many sites throughout the genome. White dots indicate chromatin bands at which Ewg and Ebd1 colocalize. (D-F) A fragment from distal end of chromosome X. Ewg (pink arrowheads) and Ebd1 (green arrowheads) are both detected at discrete bands on polytene chromosomes, and colocalize at six bands (white arrowheads in F). Ewg is associated with four additional bands at which no Ebd1 is detected (pink arrowheads in F).

## DISCUSSION

### An unbiased forward genetic screen in *Drosophila* uncovers a novel mechanism that mediates context-specific Wnt/Wingless signal transduction

Genome-wide expression profiling has revealed that the vast majority of Wnt target genes are context specific, and, by contrast, globally activated target genes are quite rare (Clevers, 2006). Factors that confer context-dependent target gene activation in response to activation of this widely used signal transduction pathway are of fundamental clinical importance, as they provide putative targets for therapeutic intervention; however, the identity of these factors, and their mechanism of action remain poorly understood, and have posed a daunting challenge for the Wnt signaling field (Gordon and Nusse, 2006).

We have taken an unbiased forward genetic approach in *Drosophila* to identify genes that promote Wingless signal transduction. The design of our genetic screen allowed identification of any gene that impacts Wingless signaling downstream, or at the level of the Apc/Axin destruction complex, which in principle could encompass a wide range of functions. Surprisingly, however, among the relatively small number of genetic complementation groups revealed in our screen, we have identified two cell-specific chromatin-associated co-factors that had not been implicated in Wingless signaling previously, but directly associate with each other and probably mediate the same context-specific Wingless-dependent developmental process. The combined activities of the two factors in this complex not only result in its recruitment to discrete chromatin sites, but also promote activity of the Armadillo-TCF transcription complex. Together, these unanticipated findings uncover a novel mechanism that facilitates context-specific regulation of Wingless signaling.

### The Ewg-Ebd1 complex associates with discrete chromatin sites to facilitate context-specific Wingless signaling responses

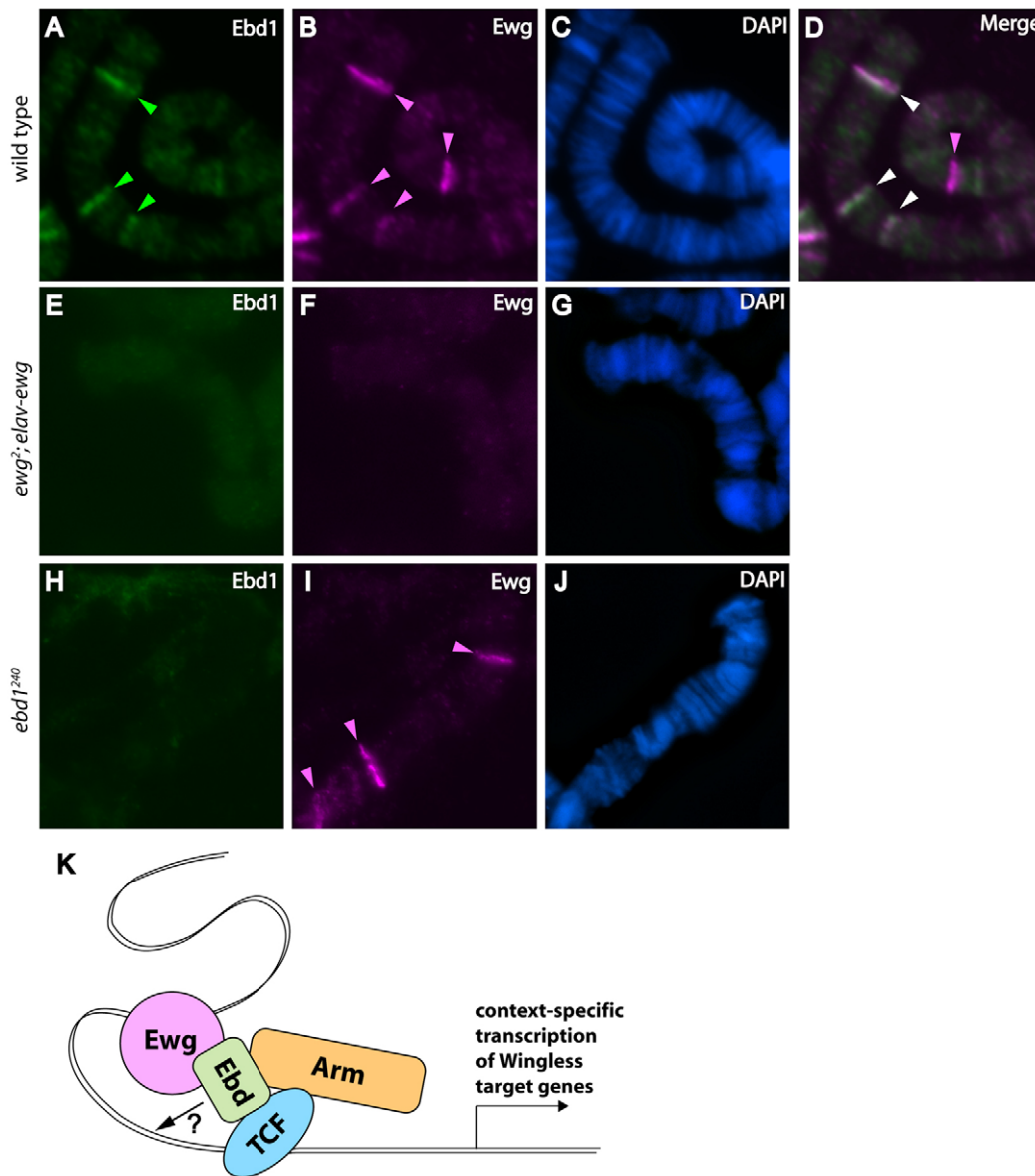
We recently discovered that the novel *Drosophila* CENPB domain protein Ebd1 and its human homolog Jerky contribute to Wnt/Wingless target gene activation by enhancing  $\beta$ -catenin-TCF complex formation and  $\beta$ -catenin recruitment to chromatin (Benchabane et al., 2011). As Ebd1 is expressed in a cell-restricted pattern, these findings provided evidence that context-dependent

responses to Wnt/Wingless stimulation are facilitated by cell-specific co-factors that interact with both  $\beta$ -catenin/Armadillo and TCF to enhance complex formation and activity. However, whether Ebd1 conferred context specificity solely by functioning as a cell-specific Armadillo-TCF adaptor or also by associating with distinct chromatin sites was not known.

Here, we provide multiple lines of evidence consistent with the model that the putative sequence-specific DNA-binding transcriptional activator Ewg is a physical and functional partner of Ebd1 in regulating context-specific Wingless signaling, and is required for chromatin recruitment of the Ewg-Ebd1 complex. Together, our studies suggest that Ebd1 and Ewg act together in the same context-specific Wingless-dependent developmental process, interact functionally with Armadillo-TCF, are expressed in similar cell-restricted patterns, associate physically with each other and are recruited to similar chromatin sites throughout the genome. Furthermore, all recruitment of Ebd1 to chromatin is abolished in the absence of Ewg. Based on these findings, we propose a model in which context specificity in Wingless signaling is achieved in part through association of Ewg with distinct enhancer sites, and recruitment of Ebd1 to these enhancers (Fig. 7K). In addition, our previous work suggested that by binding both Armadillo and TCF, Ebd1 stabilizes the Armadillo-TCF complex (Benchabane et al., 2011). Together, our findings support the model that Ebd1 promotes recruitment of the Armadillo-TCF complex to enhancers containing Ewg-binding sites, thereby facilitating context specificity in Wingless signal transduction. Alternatively, Ebd1 might also function in an Ewg-independent manner by solely providing an Armadillo-TCF adaptor function that stabilizes the Armadillo-TCF complex and thereby facilitates transcriptional activation of a subset of Wingless target genes requiring relatively high Armadillo-TCF levels.

### Ewg, a putative sequence-specific DNA-binding protein, promotes recruitment of Ebd1, a cell-specific Armadillo-TCF adaptor, to discrete chromatin sites

Our ability to visualize endogenous Ebd1 and Ewg on larval salivary gland polytene chromosomes reveals that Ewg is essential for association of Ebd1 with chromatin, and that nearly all, if not all, chromatin sites with which Ebd1 associates are also bound by Ewg. By contrast, the association of Ewg with chromatin is not



**Fig. 7. Ewg is required to recruit Ebd1 to chromatin.** (A–J) Squashes of polytene chromosomes from third instar larvae, immunostained with Ebd1 (A,E,H) and Ewg antiserum (B,F,I). DNA is highlighted by DAPI staining (C,G,J). (A–D) Ewg and Ebd1 overlap at the majority of their binding sites in wild-type polytenes. (E–G) Ebd1 is not detectable on chromatin in the absence of Ewg. (H–J) Ewg association with chromatin is maintained in absence of Ebd1. (K) A model depicting function of Ewg and Ebd1 in context-specific Wingless signaling. Ewg associates with specific chromatin sites and recruits Ebd1. Ebd1 bridges Arm and TCF, stabilizing this complex, facilitating its recruitment to chromatin, and thereby promoting context-specific activation of a subset of Wingless target genes.

dependent on Ebd1; furthermore, not all Ewg-associated chromatin sites are bound by Ebd1, which probably reflects the broader role for Ewg in muscle and neuronal development. Together, these findings suggest that Ewg is necessary but not sufficient for recruitment of Ebd1 to chromatin, and either additional factors or specific DNA sequences are important for association of Ebd1 with chromatin. In particular, the presence of CENPB-type DNA-binding domains in Ebd1 indicates that Ebd1 may bind DNA directly, and therefore the association of Ebd1 with chromatin may require not only Ewg, but also sequence-specific Ebd1 DNA-binding sites. If so, Ewg might bind and remodel specific chromatin sites, thereby allowing Ebd1 to access nearby cognate binding sites or, alternatively, physical association with Ewg may

result in a conformational change in Ebd1 that enables DNA binding. Similarly, association of Ebd1 with the Armadillo-TCF complex may facilitate association of Ebd1 and/or Armadillo-TCF with distinct chromatin sites; currently, we are unable to test this hypothesis in vivo, as our fixation and immunostaining conditions do not detect the indirect association of endogenous Armadillo with polytene chromosomes. Nonetheless, the extensive physical and genetic interactions between Ewg, Ebd1 and Wingless pathway transcriptional components, coupled with association of Ewg-Ebd1 with distinct chromatin sites support the model that context-specific activation of Wingless target genes is facilitated by interaction of Armadillo-TCF with cell-specific co-activator complexes at specific sites in chromatin.

### The Ebd1-Ewg complex promotes context-specific Wingless-dependent adult muscle development

Our findings also suggest that Ebd1-Ewg complex function is required for only a context-specific subset of the Wingless-dependent processes that direct IFM development (Sudarsan et al., 2001; Benchabane et al., 2011). For example, Wingless secreted from larval wing imaginal disc epithelia specifies the fate of myoblasts associated with the wing disc that are important for IFM development (Sudarsan et al., 2001). In addition, Wingless signaling regulates the specification of tendon cells and the position of flight muscle attachment sites (Ghazi et al., 2003). Further evidence for Wingless signaling in IFM development was revealed by hypomorphic *wingless* mutants, some of which display a morphologically normal external appearance, but are nonetheless flightless (Sharma and Chopra, 1976; Deák, 1978). Indeed, our analysis of these hypomorphic *wingless* mutants reveals defects in late steps of muscle differentiation that result in partial loss of IFMs (Benchabane et al., 2011). In addition, we find strong genetic interactions between Ewg and the Armadillo-TCF complex (Fig. 3E,F). Together with the genetic and physical interactions between Ewg and Ebd1, as well as our previous findings that Ebd1 facilitates Wingless signaling in adult muscle development (Benchabane et al., 2011), the genetic interaction between Ewg and TCF provides further evidence supporting the model that Ewg and Ebd1 facilitate Wingless-dependent adult flight muscle development. However, as documented previously (DeSimone et al., 1996), we do not detect Ewg expression in IFM precursors until pupation, suggesting that Ewg does not participate in early steps of Wingless-dependent specification of IFM myoblasts or tendon cells, but instead promotes later steps in IFM growth and differentiation. Previous studies revealed that context specificity in Wnt signaling responses is facilitated by either crosstalk between signal transduction pathways or replacement of TCF with tissue-specific DNA-binding transcription co-factors that associate with  $\beta$ -catenin (Halfon et al., 2000; Guss et al., 2001; Kioussi et al., 2002; Notani et al., 2010; Olson et al., 2006) (see also MacDonald et al., 2009; McNeill and Woodgett, 2010). Thus, our findings suggest a novel mechanism in which a sequence-specific DNA-binding transcription factor recruits a cell-specific Armadillo-TCF adaptor to modulate context-specific Wingless signaling responses.

### Wnt signaling and NRF-1 in vertebrate brain and retinal development

Ewg activity in neuronal differentiation can be functionally replaced by its human homolog nuclear respiratory factor 1 (NRF1) (Haussmann et al., 2008), a sequence-specific DNA-binding transcription factor important for activation of genes required in mitochondrial biogenesis and respiration, and for regulation of histone gene expression (Calzone et al., 1991; Virbasius et al., 1993; Gomez-Cuadrado et al., 1995; Dhar et al., 2008). Functional analysis of vertebrate NRF1 during development has been hampered by early lethality, as mouse *Nrf1*-null embryos die near the time of uterine implantation (Huo and Scarpulla, 2001). Fortuitously, however, inactivation of the zebrafish *NRF1* homolog, *not really finished* (*nrf*), has improved our understanding of NRF1 function during vertebrate development; fish *nrf*-null mutants survive up to 14 days of development with the primary developmental defect restricted to markedly smaller brains and retinas (Becker et al., 1998). The tissue-restricted nature of the fish *nrf*-null mutant phenotype was unexpected, given the broader

expression of the fish *nrf* gene (Becker et al., 1998) and the global roles for NRF1 in mitochondrial and histone function revealed by in vitro studies. However, these in vivo analyses revealed that fish *Nrf*, like fly Ewg, acts in specific tissue-restricted developmental processes. Of note, Wnt signaling is required for patterning of the mouse central nervous system (Thomas and Capocchi, 1990; Moon et al., 2004; Ciani and Salinas, 2005). Similarly, Wnt signaling is also crucial for brain and eye development in zebrafish (Lekven et al., 2001; Lekven et al., 2003; Dorsky et al., 2003; Nyholm et al., 2007), thus raising the possibility that, like Ewg, vertebrate *Nrf* may also facilitate a context-specific subset of Wnt-dependent developmental processes.

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### Competing interests statement

The authors declare no competing financial interests.

### Supplementary material

Supplementary material available online at <http://dev.biologists.org/lookup/suppl/doi:10.1242/dev.068890/-/DC1>

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