

Departament de Genètica Facultat de Biologia Universitat de Barcelona

TESI DOCTORAL

Ash2 a Drosophila: anàlisi funcional i aproximació al complexe

Mireia Angulo i Parera Barcelona, Juny de 2006



ARTICLE 1

Títol

Activation and repression activities of ash2 in Drosophila wing imaginal disc.

Autors (Any)

Mireia Angulo, Montserrat Corominas i Florenci Serras (2004)

Departament de Genètica, Universitat de Barcelona. Avda Diagonal 645, 08028 Barcelona.

Referència

Development 131, 4943-53

Resum

Els gens del grup Polycomb (PcG) i trithorax (trxG) codifiquen per proteïnes reguladores de la cromatina implicades en el manteniment de les identitats cel·lulars. La seva funció com a reguladors dels gens homeòtics ha estat molt ben documentada però es coneix molt poc sobre els efectes que poden tenir en altres gens diana o processos del desenvolupament. En aquest estudi s'utilitza el patró de venes i intervenes de l'ala com a model per entendre la funció del gen trxG absent, small or homeotic discs 2 (ash2). Es mostra que ash2 es requereix per mantenir l'activació dels gens promotors d'intervena blistered i net, i per reprimir l'expressió de rhomboid, un component de la via de DER, necessària per la formació de venes. Mentre que els fenotips de la pèrdua de funció de la via de DER són rescatats per mutants d'ash2, els fenotips de guany de funció es veuen incrementats. Els nostres resultats també mostren que ash2 actua com a repressor del gen knirps (kni), organitzador de la vena longitudinal L2. En discs mutants d'ash2 l'expressió de Kni es troba estesa per tot el disc. i en clons mutants per ash2. Kni augmenta. La inhibició que ash2 exerceix sobre kni és independent de spalt-major i spalt-related, els quals s'ha postulat que són els reguladors de kni. Tots aquests experiments indiquen que ash2 és essencial per dos processos durant el desenvolupament de l'ala: (1) mantenir el destí cel·lular d'intervena, ja sigui activant els gens promotors d'intervena o inhibint els de vena; i (2) mantenir kni en un estat reprimit en teixits fora de la vena L2.

Research article 4943

Activation and repression activities of ash2 in Drosophila wing imaginal discs

Mireia Angulo, Montserrat Corominas and Florenci Serras*

Departament de Genètica, Universitat de Barcelona, Diagonal 645, Barcelona 08028, Spain *Author for correspondence (e-mail: fserras@ub.edu)

Accepted 23 July 2004

Development 131, 4943-4953 Published by The Company of Biologists 2004 doi:10.1242/dev.01380

Summary

Polycomb (PcG) and trithorax (trxG) group genes are chromatin regulators involved in the maintenance of developmental decisions. Although their function as transcriptional regulators of homeotic genes has been well documented, little is known about their effect on other target genes or their role in other developmental processes. In this study, we have used the patterning of veins and interveins in the wing as a model with which to understand the function of the trxG gene ash2 (absent, small or homeotic discs 2). We show that ash2 is required to sustain the activation of the intervein-promoting genes net and blistered (bs) and to repress rhomboid (rho), a component of the EGF receptor (Egfr) pathway. Moreover, loss-offunction phenotypes of the Egfr pathway are suppressed by ash2 mutants, while gain-of-function phenotypes are enhanced. Our results also show that ash2 acts as a

repressor of the vein L2-organising gene knirps (kni), whose expression is upregulated throughout the whole wing imaginal disc in ash2 mutants and mitotic clones. Furthermore, ash2-mediated inhibition of kni is independent of spalt-major and spalt-related. Together, these experiments indicate that ash2 plays a role in two processes during wing development: (1) maintaining intervein cell fate, either by activation of intervein genes or inhibition of vein differentiation genes; and (2) keeping kni in an off state in tissues beyond the L2 vein. We propose that the Ash2 complex provides a molecular framework for a mechanism required to maintain cellular identities in the wing development.

Key words: ash2, Cellular memory, Imaginal disc, knirps, Drosophila

Introduction

Differential gene expression results in cell diversity, although how different cell identities are established early in development and maintained throughout life is still poorly understood. Most of the transcription factors required for early developmental decisions are expressed transiently, but the gene expression patterns they trigger are maintained during cell division and inherited by daughter cells. Actively dividing cells must preserve individual genes in an on or off expression state after an initial commitment is made, especially given that some regulators disassemble from promoters during DNA replication or mitosis. Thus, developmental decisions may be maintained by the ability to deposit epigenetic marks involving chromatin-modifying complexes to control the cellular memory of gene activity states (Francis and Kingston, 2001; Simon and Tamkun, 2002).

Genes of the Polycomb (PcG) and trithorax group (trxG) encode proteins that are engaged in the regulation of cellular memory (Orlando, 2003). In early *Drosophila* embryonic development, Hox gene expression is controlled by a genetic cascade that includes the segmentation genes (Simon, 1995). When the segmentation proteins decay, Hox expression is maintained in the correct spatiotemporal pattern by the action of PcG and trxG genes, which often act as transcriptional repressor- and activator-chromatin complexes (Francis and Kingston, 2001; Simon and Tamkun, 2002). In addition to Hox

genes, PcG/trxG also act on other target genes (Beltran et al., 2003; Francis and Kingston, 2001). In a genome-wide prediction of PcG/trxG response elements (PRE/TRE) in *Drosophila*, more than 100 elements were identified that mapped to genes involved in development and cell proliferation (Ringrose et al., 2003). However, epigenetic marks are not only restricted to embryonic stages. At later stages, developmental fates are also frozen and inherited by the repressor and activator activities of PcG/trxG complexes. The *Drosophila* wing imaginal disc has proven to be a useful model with which to study how these complexes act to maintain cell identities, as shown for *wingless* (*wg*) and *hedgehog* (*hh*) pathways (Collins and Treisman, 2000; Maurange and Paro, 2002).

The non-neural tissues of the *Drosophila* wing are organised into two types: the A-E intervein regions and the L1-L6 veins (Fig. 1A). The specification of veins in the wing imaginal disc occurs during larval and pupal stages, and is controlled by a network of cell-to-cell interactions, including the *Egfr* signalling pathway (Diaz-Benjumea and Hafen, 1994). The *rho* gene, which encodes a seven-pass transmembrane serine protease, is an activator of *Egfr* (Bier et al., 1990; Sturtevant et al., 1993). *rho* is expressed in rows of cells coinciding with vein primordia and is required for vein formation, as indicated by the observation that the loss-of-function allele *rho*^{ve} displays truncated veins (Diaz-Benjumea and Garcia-Bellido, 1990; Sturtevant et al., 1993). Localised expression of *rho* and *vein* (*vn*), which encodes a diffusible neuregulin class of

ligands, activates the Ras/MAPK signalling cascade necessary for vein differentiation (Sturtevant et al., 1993; Schnepp et al., 1996). By contrast, inhibition of Egfr signalling by the transcription factors blistered (bs) and net is responsible for intervein specification, bs, the Drosophila homologue of the Serum Response Factor, is expressed in a pattern associated with intervein regions and is required for the organisation and differentiation of intervein cells (Fristrom et al., 1994; Montagne et al., 1996). During disc proliferation, bs expression is independent of rho, but during the pupal period bs and rho expression become mutually exclusive (Roch et al., 1998). The net gene, which encodes a basic helix-loop-helix (bHLH) protein, is also expressed in the intervein regions (Brentrup et al., 2000). In contrast to bs, net and rho expression is mutually exclusive in the wing discs of third instar larvae. Lack of net activity causes rho expression to expand, and vice versa. Furthermore, ectopic *rho* expression results in repression of net, thus generating wings with ectopic vein tissue (Brentrup et al., 2000).

The refined localisation of the L2-L5 veins in the wing depends on positional cues established along the AP axis of the wing imaginal disc (Biehs et al., 1998; de Celis et al., 1996; Sturtevant et al., 1997; Sturtevant and Bier, 1995). The posterior compartment cells express engrailed (en), which activates hh. Hh diffuses through a few cell rows of the anterior compartment, activating vn, knot (kn) and decapentaplegic (dpp), which form the AP organising centre (Bier, 2000). Vn activates Egfr in the borders of the organiser, giving rise to the L3 and L4 vein primordia, while Kn prevents the domain between L3 and L4 responding to vein differentiation, ensuring the intervein fate of this region (Crozatier et al., 2002; Mohler et al., 2000). Dpp diffuses from the AP boundary and activates target genes in a threshold-dependent fashion (Lawrence and Struhl, 1996). The development of L5 is dependent upon the two abutting Dpp target genes, optomotor-blind (omb) and brinker (brk) (Cook et al., 2004). Another Dpp target is the spalt-major (salm)/spalt-related (salr) complex (sal-C) of zinc-finger transcription factors (de Celis et al., 1996; Lecuit et al., 1996; Nellen et al., 1996), which is expressed in the central domain of the wing. The anterior low sal-C-expressing domain promotes the activation of the knirps (kni)/knirpsrelated (knrl) complex (kni-C), which results in L2 specification (Lunde et al., 1998).

Ash2 is a trxG protein that belongs to a 0.5 MDa complex thought to be involved in chromatin remodelling (Papoulas et al., 1998). Ash2 accumulates uniformly in imaginal discs, fat body cells and salivary glands (Adamson and Shearn, 1996). Loss-of-function alleles of this gene cause homeotic transformations (LaJeunesse and Shearn, 1995; Shearn, 1974; Shearn et al., 1987; Shearn, 1989; Shearn et al., 1971) and downregulation of Hox genes (Beltran et al., 2003; LaJeunesse and Shearn, 1995), in addition to severe abnormalities in the wing, such as reduction of intervein and enhancement of vein tissues (Adamson and Shearn, 1996; Amorós et al., 2002). To gain more insight into the function of ash2, we examined whether vein- and intervein-specific genes and vein positioning genes act as putative targets of ash2 function. We found that ash2 is involved in activating intervein-promoting genes and downregulating the Egfr pathway. Moreover, ash2 also acts as a kni repressor independently of sal-C. These results strongly support a role for ash2 in maintaining vein/intervein developmental decisions and vein patterning in the developing wing.

Materials and methods

Drosophila strains

We used the following genetic strains as ash2 alleles: yw; ash2112411/TM6C (Deak et al., 1997; Amorós et al., 2002) and yw; ash2^{II}/TM6C (Amorós et al., 2002; Beltran et al., 2003). Canton S was used as a wild-type strain. To study interactions between ash2 and Egfr signalling pathway elements, we tested the hypomorphic combination $rho^{ve}vn^{l}$, top^{l}/top^{3C8l} (Clifford and Schupbach, 1989; Diaz-Benjumea and Garcia-Bellido, 1990), the gain-of-function allele of Egfr ElpB1/CyO and the rolled gain-of-function mutation D-Raf^{C110}; rl^{Sem} (provided by A. García-Bellido). In order to eliminate the *D-Raf*^{C110} allele, crosses were designed to score the male +/Y; $rl^{Sem/+}$; $ash2^{II/+}$ progeny. The alleles bs^{03267}/CyO (provided by M. Affolter), $E(spl)m\hat{\beta}$ -lacZ (provided by S. Bray) and net^l were used as intervein markers; the stock w; h kni^{ri-1} was used to analyse L2 development. To study the effects of ash2 on kni expression, the minimal L2-enhancer element EX-lacZ, a 1.4 kb fragment that contains an activation and repression domain of the kni gene (Lunde et al., 2003) was provided by E. Bier. For ectopic expression of sal-C, we used the UAS-sal^{64d} transgenic, on the first chromosome, the UAS-salr⁸ transgenic, on the second chromosome, and the nubbin-Gal4 insertion line (provided by J. F. de Celis). We also analysed whether ash2 regulates brk and scalloped (sd) expression by using the brk^{X47} -lacZ transgene (provided by G. Morata) and the sd^{EXT4} stock. The stocks ElpBI/Cyo, net1, w; h kniri-1 and sdEXT4 mentioned above were obtained from the Bloomington Stock Center.

Genetic mosaics

Clones mutant for $ash2^{II}$ were obtained by mitotic recombination using the FLP/FRT technique (Xu and Rubin, 1993). $yw;FRT82Bash2^{II}/TM6C$ flies were crossed with ywhsflp;FRT82BGFP/TM6B and wing imaginal discs from third instar Tubby+ larvae and pupae were dissected. Heat shock was carried out for 30 minutes at $37^{\circ}C$ [52 ± 4 hours after egg laying (AEL)] to induce clone formation. To monitor EX-lacZ (Lunde et al., 2003) expression, a yw; EX-lacZ; $FRT82Bash2^{II}/TM6C$ stock was created and clones were induced as above.

Overexpression of sal-C was obtained by crossing $UAS-Sal^{64d}$; $FRT82Bash2^{II}$ + males with ywhsflp; nubbin-Gal4; FRT82BGFP/TM6B females. Tubby⁺ female larvae were dissected. To monitor brk and sd expression, brk^{X47} -lacZ; $FRT82Bash2^{II}$ /+ males and sd^{EXT4} -lacZ; $FRT82Bash2^{II}$ /+ males were crossed to ywhsflp; FRT82BGFP/TM6B and Tubby⁺ female larvae were dissected. In both cases, only 50% of the progeny contained $ash2^{II}$ clones.

To obtain *Minute*⁺ clones the stock used was *yw; FRT82B arm-lacZ M(3)/TM6C* and heat shock was carried out for 7 minutes at 34°C (110±4 hours AEL). Adult *ash2* mutant *FLP/FRT M*⁺ clones marked with the forked mutation were analysed in males with the following genotype: *ywhsflpf*^{56a}; *FRT82BP[f*⁺]87DM(3)w[124]/*FRT82Bash2*¹¹. The heat shock was carried out for 10 minutes at 37°C (80±12 hours AEL).

Larvae and pupae of the appropriate genotypes were cultured at 25°C and timed in hours AEL or after puparium formation (APF).

Immunohistochemistry

Immunohistochemistry was performed according to standard protocols. Primary antibodies used were: guinea pig anti-Kni (1/50), provided by J. Reinitz; rabbit anti-Salm (1/500), provided by R. Barrio; rat anti-Bs (1/200), provided by M. Affolter; rabbit anti-Plexus (1/1000), provided by H. Matakatsu; rabbit anti-Vestigial (1/20), provided by S. Carroll; mouse anti-En (1/25) from the Developmental

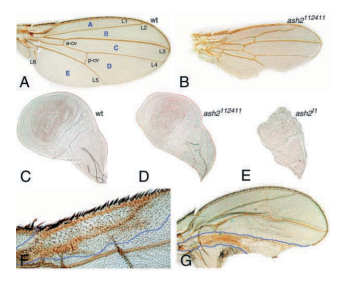


Fig. 1. Wing phenotypes of ash2 alleles. (A) Wild-type (wt) adult wing. Intervein regions (A-E, in blue), longitudinal veins (L1-L6) and anterior (a-cv) and posterior (p-cv) crossveins are marked. (B) Adult wing of ash2¹¹²⁴¹¹ homozygous mutant at the same magnification as A, showing anomalous L2 in which the distal half is crooked, ectopic crossvein tissue between L2-L3 and L4-L5, and a notch in the posterior margin. (C) Wing disc of wild-type third instar. (D) Wing disc of homozygous ash2¹¹²⁴¹¹ third instar. (E) Wing disc of homozygous ash2^{II} third instar. (F,G) Mitotic clones homozygous for ash2^{II} in Minute background. (F) Ventral clone where L2 appears thickened and intervein region A is reduced. (G) Dorsal clone showing ectopic vein tissue between the proximal region of L4-L5.

Studies Hybridoma Bank of the University of Iowa; and rabbit antiβ-galactosidase (Cappel) (1/1000). Kni, Salm and Vestigial antibodies were pre-absorbed before use.

Secondary antibodies were obtained from Jackson Immuno Research and include: donkey anti-rat-Rhodamine Red (1/200), donkey anti-guinea pig-Cy5 (1/400), goat anti-mouse-FITC (1/200) and donkey anti-rabbit-Rhodamine Red (1/200). Propidium Iodide (Molecular Probes) was used as a nuclear marker after RNAase treatment. Fluorescence was visualised with a Leica TCS confocal microscope.

Whole-mount in situ hybridisation with larval wing discs and pupal wings

In situ hybridisation using digoxigenin-labelled antisense RNA probes was carried out essentially as described previously (Sturtevant et al., 1993; Brentrup et al., 2000).

DIG-labelled riboprobes for net RNA were synthesised using a 2.2 kb insert of netcel Δ 922 (gift of M. Noll) linearised with EcoR1, and for rho from a rho cDNA clone (gift of E. Bier) linearised with HindIII. Sense RNA probes for net and rho did not show detectable signal.

RT-PCR

Total RNA from wild-type and ash211 homozygous larvae was extracted using Trizol (GibcoBRL) and a poly(dT)-24 primer was used for cDNA synthesis. The reaction was carried out in a final volume of 25 µl with five units of avian myeloblastosis virus-RT (Promega) and 200 units of Moloney murine leukaemia virus RT (Gibco). One microlitre of the RT reaction was used for PCR. The specific primers used [5'-cgccgccctgcccttcttc-3' (forward) and 5'gggctgctgctagtcggagtggt 3' (reverse)] were designed to amplify a 369 bp product of the kni gene.

Results

ash2 is required to downregulate Egfr activity

The ash2112411 mutation is a single PlacW insertion (Deak et al., 1997) in the fourth intron of the ash2 gene (Amorós et al., 2002) that causes pharate lethality. Homozygous flies that reach the adult stage (12% at 25°C) are sterile and show reduced wing size, crooked L2 and ectopic vein tissue, mainly extra crossveins and thicker veins (Fig. 1A,B). The ash2¹/₁ allele was generated after an imprecise excision of the ash2112411 insertion and is lethal in late third instar/early pupae. Molecular analysis of the ash2^{II} mutation has shown that it comprises a 2 bp deletion and a 5 bp insertion that result in the absence of the full-length 2 kb transcript (Beltran et al., 2003). Imaginal discs of both alleles are reduced in size, ash2^{II} being smaller than ash2112411 (Fig. 1C-E). The smaller size and abnormal shape suggests that the ash211 mutation alters proliferation and patterning. Clones homozygous for ash2^{II} exhibit impaired proliferation, intervein reduction and extra vein tissue, preferentially close to the normal veins, which appear thickened (Amoros et al., 2002) (Fig. 1F,G). This phenotype is a consequence of intervein cells acquiring morphological features of vein cells, which are typically smaller, more pigmented and with shorter and thicker trichomes than wild-type intervein cells. This extra vein phenotype led us to question whether ash2 functions as a negative regulator of vein differentiation in intervein territories. To test this hypothesis, we assessed genetic interactions with alleles of genes involved in vein/intervein development.

We perturbed the Ras/MAPK signalling pathway in the wing using mutants of genes required for Egfr activation. We first analysed loss-of-function mutants of the pathway. In flies mutant for the hypomorphic viable combination rhove vn¹, activation of the MAPK pathway in presumptive vein cells is prevented and, as a consequence, veins fail to differentiate. By contrast, the triple mutant rhove vnl ash2112411 develops veins (Fig. 2A,B). We observed varying degrees of rescue, ranging from wings that develop only L2 to wings that develop veins almost completely, even with extra crossvein tissue or proximal vein fusions between L2-L3 and L4-L5. Rescue of L2 and L5 is more pronounced than L3 and L4, which are never distally complete, and many wings show notches in the posterior wing margin (77% of cases, n=75 wings; Fig. 2B). In 25% of these cases, wings show a tube-like shape, possibly owing to

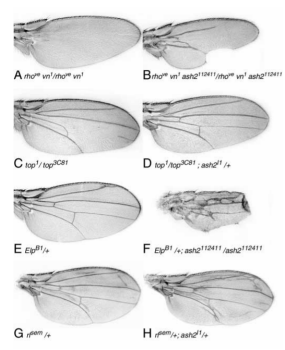


Fig. 2. ash2 downregulates the Egfr pathway during vein development. (A) Lack of wing veins in $rho^{ve} vn^I$ homozygous flies. (B) In the triple mutant $rho^{ve} vn^I ash2^{I124II}$, the loss of veins is rescued. (C) top^I/top^{3C8I} adult wing that lacks the a-cv and the middle part of L4. (D) The combination top^I/top^{3C8I} with one copy of the $ash2^{II}$ allele restores the wild-type vein pattern. (E) The gain-of-function Elp^{BI} results in ectopic vein tissue in distal L2. (F) Elp^{BI} ; $ash2^{I124II}$ individuals show a reduction in wing size, with notches appearing in the posterior margin and extra vein tissue (extra crossveins and a proximal L2-L3 fusion). (G) rl^{Sem} + flies develop wings with ectopic branches of veins. (H) Ectopic vein tissue is enhanced in rl^{Sem} ; $ash2^{II}$ transheterozygous flies. All wings are shown at the same magnification.

detachment of the dorsal and ventral cell layers. This variety of phenotypes is probably due to the variable expressivity found in the $ash2^{11241l}$ allele. The top^1/top^{3C81} allelic combination is a hypomorphic mutation of Egfr (top) in which wings lack the anterior crossvein (a-cv) and a segment of vein L4 (Fig. 2C). Rescue of missing vein tissues is observed in top^1/top^{3C81} ; $ash2^{11/+}$ flies, as shown by the presence of a-cv (58% of the cases, n=93), a complete L4 (4%), or restoration of both a-cv and L4 (28%; Fig. 2D).

We also tested whether ash2 alleles enhance gain-of-function phenotypes of the Egfr pathway. The $Ellipse^{BI}(Elp^{BI})$ allele is an activated form of Egfr. In addition to other phenotypes, Elp^{BI} individuals consistently develop wings with ectopic vein tissue in L2 (Fig. 2E). In Elp^{BI} ; $ash2^{II}/+$ no enhancement was found (data not shown), but $Elp^{BI}/+$; $ash2^{II24II}/ash2^{II24II}$ individuals had reduced wings with ectopic vein tissue (46%, n=50; Fig. 2F), including vein fusion of the proximal regions and extra crossveins between L2 and

L3, and between L4 and L5, but never between L3 and L4. In this case, a graded enhancement was also found, ranging from a wing similar to that of $ash2^{112411}$ to ectopic vein tissue covering most of the wing. Similarly, the MAPK gain-offunction allele, r^{lsem} , generates ectopic vein tissue, a phenotype that is enhanced in $ash2^{11}$ combinations (61%, n=70; Fig. 2G.H).

As these results indicate that *ash2* antagonises the Egfr pathway, we tested whether *rho* expression is affected. In situ hybridisation for *rho* mRNA in third instar *ash211* discs showed either no expression at all or expression in only a few scattered cells (Fig. 3A,B), possibly owing to a strong perturbation of patterning in these discs. However, in *ash2112411* homozygous pupal wings, *rho* is expressed and organised in veins, but the domains of *rho* expression are larger (Fig. 3C-F).

ash2 is required for the differentiation of intervein tissue

As several genes that promote intervein specification are antagonists of the Egfr signalling pathway, we tested whether the phenotypes described above are associated with loss of intervein gene activity. During larval stages, net is the key gene involved in intervein development and acts as an antagonist of rho (Brentrup et al., 2000). In third instar wing discs, net transcripts are confined to broad domains corresponding to prospective interveins, in a pattern complementary to rho (Brentrup et al., 2000). net RNA expression is considerably reduced in ash2 mutants, although in ash2112411, some expression is still found in the central domain of the wing pouch (Fig. 4A-C). Likewise, homozygous flies for the net¹/net¹ allele develop extra vein tissue, which is preferentially associated with transverse connections between L2 and L3 and between L4 and L5 (Fig. 4D), resulting in wing blades with an increased number of cells (Diaz-Benjumea and Garcia-Bellido, 1990; Garcia-Bellido and de Celis, 1992). The mutant combination net^{1}/net^{1} ; $ash2^{11}/+$ results in extra vein connections along the proximal and distal L2-L3 region, thickening of veins, blistering and a lanceolated shape to the wing. In addition, the intervein tissue is strongly reduced between veins L2-L3 and L4-L5, and to a lesser extent in A and E intervein regions (Fig. 4E). A more extreme phenotype (75%, n=20 wings; Fig. 4F) was obtained in net^1/net^1 ; $ash2^{112411}/ash2^{112411}$ wings, in which the intervein areas between L2-L3 and L4-L5 are totally missing and result in thicker and fused veins (Fig. 4G). The region between L3 and L4 did not show extra vein tissue and was much less affected.

We also generated double mutant flies to test whether *ash2* interacts with the intervein-promoting gene *bs*. The loss-of-function allele *bs*^{0,3267}, a *PlacZ* insertion within the *bs* gene, is not viable in homozygosis (Karpen and Spradling, 1992). In pharate *bs*^{0,3267} homozygotes, the whole wing blade is transformed into corrugated vein tissue, whilst the hinge region is almost intact. Heterozygous *bs*^{0,3267} flies display small patches of ectopic veins in the proximity of L2 and L5 (Roch et al., 1998) (Fig. 5A). However, flies that are homozygous for *ash2*^{112,411} and heterozygous for *bs*^{0,3267} exhibit blistering (Fig. 5B), reduced wing size associated with localised reduction of the intervein tissue (Fig. 5B,C) and development of extra crossveins (Fig. 5C,D). Veins L2 and L3 are thicker and totally or partially fused in the proximal region. Similarly, the D-intervein region between veins L4 and L5 is reduced and these

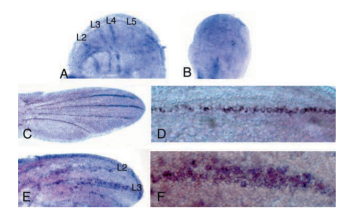


Fig. 3. In situ hybridisation for rho in discs and pupal wings. (A) Wild-type disc. (B) *rho* organisation is lost in *ash2^{II}* homozygous discs. (C) Wild-type pupal wing. (D) High magnification of a wild-type vein. (E) ash2¹¹²⁴¹¹ homozygous pupal wing. Veins L2 and L3 are wider. (F) High magnification of a severe ash2¹¹²⁴¹¹ wing showing vein thickening.

veins are partially fused and thicker. Moreover, in pupal $bs^{03267}/+$; $ash2^{112411}/ash2^{112411}$ wings, rho is expressed in broader domains (Fig. 5E,F). Interestingly, the C-intervein region is much less affected. This region is smaller than the corresponding wild type, probably owing to the overall wing reduction. However, as for net, neither ectopic vein tissue in the C-intervein region nor fusion of L3-L4 was found (Fig. 5C).

To determine whether ash2 function is necessary for transcriptional activation of bs, we used genetic mosaics. In wild-type third instar imaginal wings, bs is expressed in stripes corresponding to the intervein regions (Fig. 5G). When ash211 mutant clones fall within intervein tissue, bs expression is reduced. The most severe cases correspond to those clones located in B or D regions, where Bs expression is completely eliminated. Downregulated expression of bs is weaker in clones located in A, C and E regions (Fig. 5H-J). bs and rho expression is only mutually exclusive during pupal development, when bs promotes intervein differentiation and is expressed in intervein regions with abrupt borders with veins (Roch et al., 1998). In $ash2^{112411}$ pupal wings, bs is also expressed in interveins, although the size of the expression domains is smaller than the corresponding wild type. These bs-expressing domains are separated by wide stripes, corresponding to veins (Fig. 5K-M). The pattern of bs expression in ash2112411 interveins is non-uniform, as indicated by the presence of negatively stained cells spread throughout the wing. Accordingly, pupal ash211 clones result in a clearer downregulation of bs (Fig. 5N-S).

Enhancer of split $[E(spl)m\beta]$, a gene downstream of Notch, is also expressed in the wing pouch in broad domains that correspond to most interveins (de Celis et al., 1997). However, when we generated clones of $ash2^{II}$, we observed no effect on either $E(spl)m\beta$ or plexus (px), another intervein-associated gene (Matakatsu et al., 1999) (data not shown).

As the phenotypes of double mutants in ash2 and either net or bs are reminiscent of mutants with reduced Dpp signalling (de Celis et al., 1996), we investigated whether the expression pattern of the Dpp target gene salm is altered in these flies or in ash2 single mutants. We found that in ash2II/ash2II discs salm is slightly downregulated (Fig. 6A,B). In addition, ash2^{II} clones resulted in weak cell-autonomous downregulation of

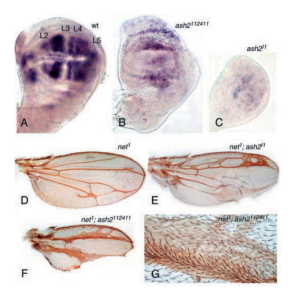


Fig. 4. ash2 is required for net activation. (A-C) Expression of net mRNA in third instar wing discs. (A) Wild-type discs showing net transcripts in intervein regions. The position of veins (lacking net mRNA) is indicated. (B) net expression is reduced in discs ash2¹¹²⁴¹¹/ash2¹¹²⁴¹¹. However, the central domain of the wing pouch shows expression of net. (C) Absence of net mRNA in ash211 wing discs. (D) Wing of net^l homozygous mutant with ectopic vein tissue. (E) net¹/net¹; ash2^{II} heterozygous flies show posterior wing blistering and enhanced ectopic vein connections, especially between L2-L3. (F) Wings of *net*¹; *ash*2¹¹²⁴¹¹ homozygous flies exhibit a total fusion of L2-L3 and L4-L5 with the corresponding loss of intervein regions B and D. The size of the wing is also reduced and the posterior margin is partially lost. (G) High magnification of L2 and L3 fusion, which is mostly associated with L2 thickening, as L5 is for L4-L5 fusion. Note the ventral pattern of corrugation in L2 and the sensillia campaniforme, and dorsal corrugation in L3.

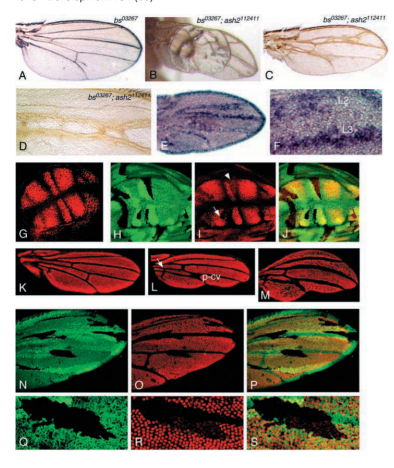


Fig. 5. ash2 is required for activation of bs. (A) Wings of heterozygous bs⁰³²⁶ flies display a small ectopic vein in intervein region D. (B,C) bs⁰³²⁶⁷/+; ash2112411/ash2112411 wings show blisters with extra vein tissue (B) and reduction of interveins B and D alongside partial fusion of L2-L3 and L4-L5 (C). (D) Detail of extra crossveins connecting L2 and L3. (E) *rho* expression in a *bs*⁰³²⁶⁷/+; *ash*2¹¹²⁴¹¹/*ash*2¹¹²⁴¹¹ pupal wing. (F) High magnification of the same image. (G) Bs staining in a wild-type third instar wing disc. The vein territories and the DV boundary are devoid of antibody staining, which is only associated with intervein territories. (H) ash2II clones in a wing imaginal disc (visualised as black spots by the absence of GFP). (I) Bs staining of the same disc. The reduction of Bs staining is more pronounced in the clone located between L2 and L3 (arrow) than in the one between L3 and L4 (arrowhead). (J) Merged green and red channels. (K-M) 24-30 hours APF wings stained with anti-Bs. (K) Wild type. (L) $ash2^{112411}/ash2^{112411}$ where extra vein tissue is restricted to wider p-cv and an extra proximal crossvein (arrow). (M) $ash2^{112411}/ash2^{112411}$ pupal wing of an individual with thicker vein tissues. In both L and M, some interveins are reduced and show some negatively stained cells within the interveins. (N) ash2II mitotic clones in pupal wings. (O) Bs staining of the same wing. (P) Merged green and red channels. (Q-S) High magnification of an ash2II

salm (Fig. 6F-H). However, homozygous $ash2^{112411}$ discs express salm in a central domain (Fig. 6C), as in net^1/net^1 ; $ash2^{112411}/ash2^{112411}$ and $bs0^{3267}/+$; $ash2^{112411}/ash2^{112411}$ discs (Fig. 6D,E). In all these cases, the expression pattern resembles wild type. We also tested for possible alterations of brk, an antagonist of the Dpp signalling pathway that is expressed in peripheral cells of the wing disc in a pattern complementary to the sal-C domain (Campbell and Tomlinson, 1999; Jazwinska et al., 1999). However, $ash2^{11}$ clones in the wing did not show any perturbation of brk expression (Fig. 6I-L).

ash2 inhibits kni expression

We next examined whether the function of ash2 is exclusive to vein/intervein differentiation or whether vein positioning genes are also targeted. As ash2¹¹²⁴¹¹ shows a crooked L2 and the expression of kni-C is required to organise this vein, we studied a possible interaction between the two genes. Analysis with anti-Kni reveals kni expression in L2 and in some cells of the wing pouch margin in wild-type discs (Fig. 7A), while in ash2¹¹ homozygous mutant discs its expression is expanded throughout the whole disc (Fig. 7B). Some expansion of kni

expression is also found in $ash2^{I1241I}$ discs, which in addition show discrete Kni-staining of L2 (Fig. 7C). We also tested the effects in mitotic clones and found that kni is upregulated in $ash2^{II}$ clones (Fig. 7D-L). This effect is cell-autonomous. Activation of kni was observed in twin and $Minute^+$ clones generated in early developing discs, as well as in small clones generated late in development, suggesting that the activation of kni by $ash2^{II}$ can occur at any stage of larval development.

We observed that although *ash2* homozygous cells within the *Minute*⁺ clone show a strong activation of *kni*, the heterozygous cells show residual activity in the whole wing disc (Fig. 7E). To test whether this residual activity is due to the lack of one copy of the *ash2* gene, we generated twin clones (Fig. 7G-I) and found that the residual *kni* expression seen in heterozygous cells was completely missing in cells homozygous for the wild-type allele (Fig. 7J-L). This result supports *kni* de-repression not only in *ash2* homozygous cells but also in *ash2* heterozygous cells. However, whereas Kni in the L2 vein is nuclear, in homozygous *ash2* clones, the localisation is both cytoplasmic and nuclear (Fig. 7M,N), suggesting that although there is an increase in Kni protein, it may not be fully functional. To gain insight into the interaction

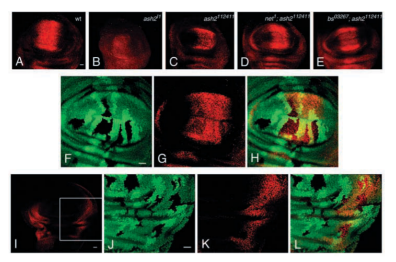


Fig. 6. Patterns of sal (A-H) and brk (I-L) expression in wing discs. (A) Wild type. (B) ash2^{II}/ash2^{II} showing reduced sal expression. (C) ash2¹¹²⁴¹¹/ash2¹¹²⁴¹¹ (D) net^{l}/net^{l} ; $ash2^{l1241l}/ash2^{l1241l}$. (E) $bs^{03267}/+$; $ash2^{112411}/ash2^{112411}$. (F-H) ash211 mutant clones, marked by the absence of GFP. (G) salm expression is reduced in *ash2^{II}* clones. (H) Merged green and red channels. (I) brk expression is visualised by means of the brk-lacZ reporter X47 in a wild-type wing disc. (J) High magnification of the boxed area indicated in I, ash211 clones lack GFP. (K) There is no change in β-galactosidase expression in ash2II mutant cells. (L) Merged images of J and K. Scale bars: 20 µm.

between ash2 and kni we analysed mutant combinations of ash2 and the hypomorphic allele kni^{ri-1}. In kni^{ri-1} wings, the L2 vein fails to develop and the rest of the veins develop normally (Fig. 7O), whereas in kni^{ri-1} ash2¹¹²⁴¹¹ double homozygous mutants, the L2 vein is partially (25%, n=58) or almost completely restored (75%, n=58; Fig. 7P) and rho is expressed in the L2 primordium (Fig. 7Q,R). Further evidence in favour of a role for ash2 as a repressor of kni was gained by semi-quantitative RT-PCR. We found that RNA from ash211 homozygous larvae contains significantly increased levels of kni, compared with wild-type larvae (Fig. 7S).

ash2 regulates kni expression independently of sal-C

We also examined whether repression of kni by ash2 is mediated by regulators of kni expression in the wing. A wellknown regulator of kni in L2 is sal-C. Low levels of sal-C activate kni expression in the presumptive L2 region, whereas higher levels repress kni-C expression (de Celis and Barrio, 2000; Lunde et al., 1998). As we observed some perturbation of salm expression in ash211 tissues, we investigated whether the low levels of salm could be responsible for the ectopic kni expression. To achieve this, we generated ash211 clones in UASsalm or UAS-salr backgrounds, using a nubbin-Gal4 driver. Adult wings overexpressing salm or salr lose L2 and L5 and show severe size reduction (de Celis et al., 1996) (Fig. 8A). Surprisingly, we observed that ash2 clones generated in those flies exhibit de-repression of kni, even when high levels of Salm are maintained in the clone (Fig. 8B-E). Therefore, we conclude that de-repression of kni induced by loss of function of ash2 is independent of sal-C. This interpretation is strengthened by the observation that in mitotic clones and in homozygous ash2^{II} mutants the ectopic expression of kni is not exclusive to the sal-C domain (Fig. 7B,G). In regions outside the sal-C central domain, loss of ash2 also activates kni, in anterior as well as posterior cells.

None of the other possible activators or inhibitors of the L2 enhancer, such as sd, vg or en, showed significant alteration of their expression in ash211 clones (data not shown). Moreover,

using a lacZ reporter construct that includes a minimal L2 enhancer element containing only the activator and repressor sites (EX-lacZ) (Lunde et al., 2003), we observed no βgalactosidase expression in ash211 homozygous discs or in mitotic clones (data not shown). Taken together, these results indicate that the ash2-induced repression of kni is independent of sal-C, and suggest that ash2 could be interacting with a kni enhancer other than L2.

Discussion

An important step towards understanding how cell determination is maintained is the identification of targets of the PcG and trxG genes. Although ash2 has been considered to be a member of the trxG (Adamson and Shearn, 1996), its biological and molecular function is still unknown. In this work, we show evidence that ash2 is required for normal vein/intervein patterning, and demonstrate that it plays a role in two major biological events – determination of intervein identity and maintenance of kni repression beyond L2.

Identifying intervein and vein target genes of ash2

Loss of ash2 function causes differentiation of ectopic vein tissue, indicating that ash2 is required for intervein development, where it functions as an activator of the intervein-promoting genes net and bs, restricting rho expression to vein regions. In addition, the loss-of-function phenotypes of Egfr alleles are rescued in ash2 mutants, while the gain-of-function phenotypes are enhanced. Furthermore, rho mRNA exhibits an expanded expression pattern in ash2 mutant tissues. Thus, ash2 promotes the maintenance of intervein fate, either by activation of *net* and *bs* or by repression of the Egfr pathway. As rho and bs/net expression is mutually exclusive, we cannot determine whether the Ash2 complex interacts directly with one or all of them. However, as bs expression is inhibited by the loss-of-function of ash2 during larval and pupal stages, we can propose that ash2 acts as a long-term chromatin imprint of bs that is stable throughout development.

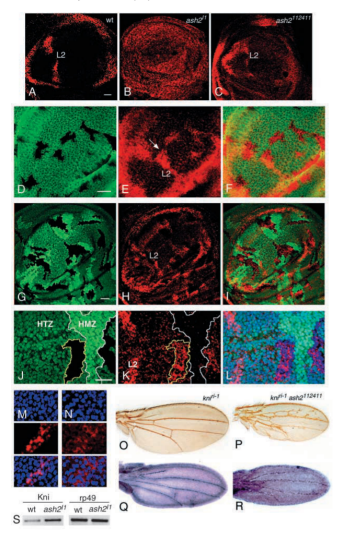


Fig. 7. ash2 regulates kni expression. kni in wild-type (A). $ash2^{II}/ash2^{II}$ (B) and $ash2^{I1241I}/ash2^{I1241I}$ (C) wing discs. (D) Minute+ clones homozygous for ash211 induced at 110±4 hours AEL are marked by the absence of βgalactosidase. (E) Expression of kni is cell-autonomously upregulated by the loss of ash2 function (arrow). There is residual kni expression in heterozygous tissue. (F) Merged images of C and D. (G) Clones homozygous for ash2^{II} induced at 52±4 hours AEL are marked as black spots by the absence of GFP and twin clones of wild-type cells appear as intense green spots. (H) kni expression is upregulated in homozygous ash211 mutant cells and residual kni disappears in cells homozygous for GFP. (I) Merged green and red channels of F and G. (J) Detail of a clone. Homozygous GFP cells are outlined in white (HMZ) and homozygous $ash2^{II}$ cells are outlined in yellow. HTZ. heterozygous tissue. (K) Anti-Kni staining of the same disc. One mutant copy of ash2 is enough to de-repress kni expression. (L) Merged images of I and J, and an additional channel with nuclear labelling (propidium iodide, shown in blue). (M) Detail of endogenous Kni (red) showing nuclear localisation (blue) in the L2 domain (pink in lower panel corresponds to nuclear labelling). (N) Detail of an ash2II clone; Kni is mainly cytoplasmic, although nuclei also contain some protein. (O) Adult kni^{ri-1}/kni^{ri-1} wing. (P) kni^{ri-1}/kni^{ri-1} ash2¹¹²⁴¹¹/ash2¹¹²⁴¹¹ wing showing rescue of L2 vein. (Q) rho expression in pupal kni^{ri-1}/kni^{ri-1} wing. (R) rho expression in pupal kniri-1/kniri-1 ash2112411/ash2112411 wing. There is distal expression in L2. (S) RT-PCR showing increased kni mRNA in ash211 homozygous larvae compared with wild type. Rp49 is shown as a control. Scale bars: 20 µm.

Our results in adult clones and from analysis of genetic interactions suggest that ash2 acts principally by maintaining B and D intervein regions, as the C intervein remains unaltered in ash2 mutants. This region is under the control of organising genes that respond to the Hh signal (Tabata and Kornberg, 1994; Zecca et al., 1995). One of these genes is kn, which prevents vein differentiation in the C intervein (Crozatier et al., 2002; Mohler et al., 2000) and is required for the expression of bs in this domain (Vervoort et al., 1999). bs expression is regulated by two enhancer elements: the boundary enhancer, which is dependent on hh and controls bs expression in the C intervein region through kn; and another enhancer dependent on Dpp activity, which controls bs expression in B and D intervein domains (Nussbaumer et al., 2000). Thus, the role of ash2 as a positive regulator of bs is mainly restricted to regions beyond the kn domain where the Dpp dependent bs enhancer is active. It has been found that some combinations of *dpp* alleles and mosaic clones of *sal-C* result in elimination of B and D intervein regions, along with fusion of their flanking veins (de Celis et al., 1996). Although the genetic interactions between *ash2* and either *bs* or *net* could be the result of a synergistic failure to activate genes downstream of Dpp, our results indicate that this may not be the case because *salm* is expressed in the central domain of the wing pouch of those mutant combinations.

It has been recently shown that another trxG complex, the Brm complex, is involved in regulating wing vein development (Marenda et al., 2004). The authors found that components of that complex interact genetically with *net* and *bs* at pupal stages to regulate the expression of *rho*, and that the complex is specifically required in cells within and bordering L5 to mediate proper signalling. There are some key differences

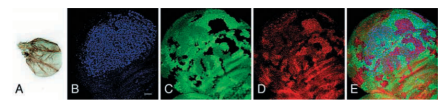


Fig. 8. kni upregulation by ash2^{II} is independent of sal-C downregulation. (A) Overexpression of sal-C from UAS-sal^{64d} using a nubbin-Gal4 driver results in loss of L2 and L5 and severe reduction of wing size. (B) Wing imaginal disc showing ectopic expression of salm throughout the wing pouch. (C) ash2^{II} homozygous clones are devoid of GFP. (D) kni expression is also upregulated in ash2^{II} mutant clones even, when salm is overexpressed. (E) Merged images. Scale bars: 20 µm.

between the Brm complex and Ash2: (1) Ash2 maintains bs expression from the third instar stage; (2) the Ash2 complex is mainly required for interveins B and D; and (3) the enhancement or suppression phenotypes of the genetic interactions with Egfr and intervein-promoting alleles are much stronger for ash2 than for the Brm complex. Taken together, these results suggest that ash2 plays a crucial role in intervein identity and that each trxG complex acts in a specific spatiotemporal program to maintain organ identity.

Ash2 complex maintains kni in an off state

The positioning of vein tissues depends on the sal-C patterning dictated by the Dpp signalling pathway (Sturtevant et al., 1997; Sturtevant and Bier, 1995). Low levels of sal-C in the anterior compartment are required for the expression of kni-C, which triggers the differentiation of L2 (de Celis and Barrio, 2000; Lunde et al., 1998). We have shown that lack of ash2 activity results in downregulation of salm and upregulation of kni. Thus, it is possible that within the sal-C domain, the ectopic expression of kni is a result of low levels of salm. However, when high levels of salm or salr are maintained by ectopic activation, lack of ash2 nevertheless results in de-repression of kni. Moreover, kni is also cell-autonomously de-repressed by loss-of-function of ash2 in cells outside of the sal-C expression domain. Thus, the repression state in the whole wing must be maintained by factors other than sal-C. The kni/knirl L2enhancer is subdivided into activation binding sites for Brk, En and the Sd/Vg complex, and repression binding sites for Sd/Vg, En, Salr and Brk (Lunde et al., 2003). We did not observe changes either in β -gal expression from the EX-lacZ enhancer or in sd, vg, brk or en expression in clones lacking ash2. Therefore the de-repression of kni in ash2 mutant cells must be accounted for by a mechanism entirely different from that of the signal-dependent induction of L2, perhaps through another enhancer more global than that of L2.

The low levels of salm expression associated with ash211 clones may also be explained by de-repression of kni. In dorsal tracheal cells, kni/knrl activity represses salm transcription, and this repression is essential for branch formation. Similarly the establishment of the border between cells acquiring dorsal branch and dorsal trunk identity entails a direct interaction of Knirps with a salm cis-regulatory element (Chen et al., 1998). Also in the wing, kni and knrl are likely to refine the L2 position by positive auto-regulation of their own expression and by providing negative feedback to repress salm expression (Lunde et al., 1998).

It is possible that the de-repression of kni, intervein

inhibition and appearance of extra vein tissues are linked events. The kni-C complex organises the development of the L2 vein by activating rho and inhibiting bs (Lunde et al., 1998; Montagne et al., 1996). Thus, kni-C participates in L2 morphogenesis by functioning downstream of salm and upstream of vein-intervein genes. The ectopic activation of kni by lack of ash2 could trigger intervein repression and vein activation. Indeed, ectopic activation of UAS-kni results in broad expression of *rho* and elimination of Bs expression in pupal wings, leading to the production of solid vein material (Lunde et al., 1998). However, in adult clones not all ash2 mutant cells develop vein tissue. This raises the possibility that de-repressed kni may not be fully functional, as ectopic kni is often localised to the cytoplasm rather than the nucleus. Alternatively, ash2 could have independent functions in the wing, maintenance of the repressed state of kni alongside maintenance of the intervein condition, by acting on different

The ash2¹¹²⁴¹¹ mutation can partially rescue the loss of L2 in *kni^{ri-1}* mutants. This is in contrast to our observation that the L2 enhancer appears not to mediate the effect of ash2. The kni^{ri-1} allele is a 252 bp deletion in the enhancer of L2 (Lunde et al., 2003) that results in lack of kni expression in L2 (Lunde et al., 1998). It has been shown, however, that it is possible to rescue the vein-loss phenotype of kni^{ri-1} by expressing a *UAS*rho transgene in L2 (Lunde et al., 2003). In addition, double mutant flies for kniri-1 and net partially rescue L2 (Diaz-Benjumea and Garcia-Bellido, 1990). It is therefore likely that the antagonistic effect of ash2 on rho could account for the partial rescue of L2 in kniri-1 ash2112411 wings, as rho mRNA is expressed in the rescued L2.

Some PcG genes are known to be required for the maintenance of kni expression domains in the embryo (McKeon et al., 1994; Pelegri and Lehmann, 1994; Saget et al., 1998). It is also likely that some trxG genes or other complexes of trxG proteins, such as the Ash2 complex (Papoulas et al., 1998), may interact with repressor sequences necessary to keep kni expression in an off state beyond L2. Moreover, in a genome wide prediction screen it has been shown that kni contains PRE/TREs (Ringrose et al., 2003). Thus, we propose here that ash2 acts as regulator of kni expression in the wing through an epigenetic mechanism of cellular memory similar to the trx-G regulation of homeotic genes, albeit that it remains to be seen whether kni is a direct or indirect target of ash2.

Cellular memory and morphogen gradients

A well-studied mechanism through which to induce and

preserve cell identities in wing imaginal discs is the response to gradients of the morphogen Dpp. This raises questions about the extent to which the response to Dpp occurs through concentration-dependent mechanisms or cellular memory. There is compelling evidence in favour of the existence of Dpp gradients that organise the pattern and growth of the wing imaginal disc (Podos and Ferguson, 1999; Strigini and Cohen, 1999). Dpp signalling causes a graded transcriptional regulation of brk by an interaction between the Dpp transducers and a brk morphogen-regulated silencer (Muller et al., 2003). Thus, brk appears to respond to direct morphogenetic signalling rather than remembering the inputs of previous developmental events. However, whereas activation of salm requires continuous signalling through the Dpp pathway (Lecuit et al., 1996; Nellen et al., 1996), other targets of Dpp, such as *omb*, remember exposure to the signal (Lecuit et al., 1996). We have shown here that stable regulation of other genes involved in wing development, such as kni repression, and net and bs activation, would also respond to the cellular memory conferred by epigenetic marks of the Ash2 complex. Thus, both mechanisms - morphogen-dependant, which will be required for growth and patterning, and epigenetic, which will keep specific genes in an off or on state - are likely to act simultaneously to maintain cellular identities within the wing.

Because many developmental regulators are only expressed transiently during development, the function of epigenetic complexes is likely to be very dynamic. The developmental events required for the construction of the wing, as with many other morphogenetic events, cannot only rely on an on or off state of gene expression. Instead, morphogenesis is rather malleable and epigenetic marks could act as a means to facilitate, rather that fix, the preservation of developmental fates. It may well be that the epigenetic marks of the Ash2 complex allow changes in chromatin structure to assist the access of proteins that activate or repress gene expression. From an epigenetic point of view, the ultimate refinement of morphogenesis and maintenance of cellular memory will depend upon the interaction of these chromatin remodelling complexes with the factors that trigger or inhibit transcription.

We are grateful to J. F. de Celis, F. Roch and E. Moreno for advice and constructive criticism. We thank A. García Bellido, J. F. de Celis, M. Affolter, E. Bier, E. Moreno and G. Morata for providing stocks; J. Reinitz, M. Affolter, H. Matakatsu, S. Carroll and R. Barrio for providing antibodies; and M. Noll, E. Bier and J. F. de Celis for the constructs used to make riboprobes. We also acknowledge the Bloomington Stocks Center and the Developmental Studies Hybridoma Bank for sending all the stocks and antibodies requested. We especially thank E. Moreno and S. Beltran for helpful comments on the manuscript, and M. Milán and M. Marsal for advice on pupal wings and in situ hybridisation. M.A. is the recipient of a fellowship from Universitat de Barcelona (UB). This work was supported by the Ministerio de Ciencía y Tecnologia (Spain), grants BMC2000-0766, BMC2003-05018 and HI2001-0009.

References

- Adamson, A. L. and Shearn, A. (1996). Molecular genetic analysis of Drosophila ash2, a member of the trithorax group required for imaginal disc pattern formation. Genetics 144, 621-633.
- Amoros, M., Corominas, M., Deak, P. and Serras, F. (2002). The ash2 gene is involved in *Drosophila* wing development. *Int. J. Dev. Biol.* 46, 321-324.

- Beltran, S., Blanco, E., Serras, F., Perez-Villamil, B., Guigo, R., Artavanis-Tsakonas, S. and Corominas, M. (2003). Transcriptional network controlled by the trithorax-group gene ash2 in *Drosophila melanogaster*. Proc. Natl. Acad. Sci. USA 100, 3293-3298.
- Biehs, B., Sturtevant, M. A. and Bier, E. (1998). Boundaries in the Drosophila wing imaginal disc organize vein-specific genetic programs. Development 125. 4245-4257.
- Bier, E. (2000). Drawing lines in the *Drosophila* wing: initiation of wing vein development. *Curr. Opin. Genet. Dev.* **10**, 393-398.
- Bier, E., Jan, L. Y. and Jan, Y. N. (1990). rhomboid, a gene required for dorsoventral axis establishment and peripheral nervous system development in *Drosophila melanogaster*. Genes Dev. 4, 190-203.
- Brentrup, D., Lerch, H., Jackle, H. and Noll, M. (2000). Regulation of *Drosophila* wing vein patterning: net encodes a bHLH protein repressing rhomboid and is repressed by rhomboid-dependent Egfr signalling. *Development* 127, 4729-4741.
- Campbell, G. and Tomlinson, A. (1999). Transducing the Dpp morphogen gradient in the wing of *Drosophila*: regulation of Dpp targets by brinker. Cell 96, 553-562.
- Chen, C. K., Kuhnlein, R. P., Eulenberg, K. G., Vincent, S., Affolter, M. and Schuh, R. (1998). The transcription factors KNIRPS and KNIRPS RELATED control cell migration and branch morphogenesis during *Drosophila* tracheal development. *Development* 125, 4959-4968.
- Clifford, R. J. and Schupbach, T. (1989). Coordinately and differentially mutable activities of torpedo, the *Drosophila melanogaster* homolog of the vertebrate EGF receptor gene. *Genetics* 123, 771-787.
- Collins, R. T. and Treisman, J. E. (2000). Osa-containing Brahma chromatin remodeling complexes are required for the repression of wingless target genes. *Genes Dev.* 14, 3140-3152.
- Cook, O., Biehs, B. and Bier, E. (2004). brinker and optomotor-blind act coordinately to initiate development of the L5 wing vein primordium in *Drosophila*. *Development* 131, 2113-2124.
- Crozatier, M., Glise, B. and Vincent, A. (2002). Connecting Hh, Dpp and EGF signalling in patterning of the *Drosophila* wing; the pivotal role of collier/knot in the AP organiser. *Development* 129, 4261-4269.
- de Celis, J. F. and Barrio, R. (2000). Function of the spalt/spalt-related gene complex in positioning the veins in the *Drosophila* wing. *Mech. Dev.* 91, 31-41
- de Celis, J. F., Barrio, R. and Kafatos, F. C. (1996). A gene complex acting downstream of dpp in *Drosophila* wing morphogenesis. *Nature* 381, 421-424
- de Celis, J. F., Bray, S. and Garcia-Bellido, A. (1997). Notch signalling regulates veinlet expression and establishes boundaries between veins and interveins in the *Drosophila* wing. *Development* 124, 1919-1928.
- Deak, P., Omar, M. M., Saunders, R. D., Pal, M., Komonyi, O., Szidonya, J., Maroy, P., Zhang, Y., Ashburner, M., Benos, P. et al. (1997). P-element insertion alleles of essential genes on the third chromosome of *Drosophila melanogaster*: correlation of physical and cytogenetic maps in chromosomal region 86E-87F. Genetics 147, 1697-1722.
- Diaz-Benjumea, F. J. and Garcia-Bellido, A. (1990). Behaviour of cells mutant for an EGF receptor homologue of *Drosophila* in genetic mosaics. *Proc. R. Soc. Lond., B, Biol. Sci.* 242, 36-44.
- Diaz-Benjumea, F. J. and Hafen, E. (1994). The sevenless signalling cassette mediates *Drosophila* EGF receptor function during epidermal development. *Development* 120, 569-578.
- Francis, N. J. and Kingston, R. E. (2001). Mechanisms of transcriptional memory. Nat. Rev. Mol. Cell Biol. 2, 409-421.
- Fristrom, D., Gotwals, P., Eaton, S., Kornberg, T. B., Sturtevant, M., Bier, E. and Fristrom, J. W. (1994). Blistered: a gene required for vein/intervein formation in wings of *Drosophila*. *Development* 120, 2661-2671.
- Garcia-Bellido, A. and de Celis, J. F. (1992). Developmental genetics of the venation pattern of *Drosophila. Annu. Rev. Genet.* 26, 277-304.
- Jazwinska, A., Kirov, N., Wieschaus, E., Roth, S. and Rushlow, C. (1999). The *Drosophila* gene brinker reveals a novel mechanism of Dpp target gene regulation. *Cell* 96, 563-573.
- Karpen, G. H. and Spradling, A. C. (1992). Analysis of subtelomeric heterochromatin in the *Drosophila* minichromosome Dp1187 by single P element insertional mutagenesis. *Genetics* 132, 737-753.
- LaJeunesse, D. and Shearn, A. (1995). Trans-regulation of thoracic homeotic selector genes of the Antennapedia and bithorax complexes by the trithorax group genes: absent, small, and homeotic discs 1 and 2. Mech. Dev. 53, 123-130.
- Lawrence, P. A. and Struhl, G. (1996). Morphogens, compartments, and pattern: lessons from drosophila? Cell 85, 951-961.

- Lecuit, T., Brook, W. J., Ng, M., Calleja, M., Sun, H. and Cohen, S. M. (1996). Two distinct mechanisms for long-range patterning by Decapentaplegic in the Drosophila wing. Nature 381, 387-393.
- Lunde, K., Biehs, B., Nauber, U. and Bier, E. (1998). The knirps and knirpsrelated genes organize development of the second wing vein in Drosophila. Development 125, 4145-4154.
- Lunde, K., Trimble, J. L., Guichard, A., Guss, K. A., Nauber, U. and Bier, E. (2003). Activation of the knirps locus links patterning to morphogenesis of the second wing vein in Drosophila. Development 130, 235-248.
- Marenda, D. R., Zraly, C. B. and Dingwall, A. K. (2004). The Drosophila Brahma (SWI/SNF) chromatin remodeling complex exhibits cell-type specific activation and repression functions. Dev. Biol. 267, 279-293.
- Matakatsu, H., Tadokoro, R., Gamo, S. and Hayashi, S. (1999). Repression of the wing vein development in Drosophila by the nuclear matrix protein plexus. Development 126, 5207-5216.
- Maurange, C. and Paro, R. (2002). A cellular memory module conveys epigenetic inheritance of hedgehog expression during Drosophila wing imaginal disc development. Genes Dev. 16, 2672-2683.
- McKeon, J., Slade, E., Sinclair, D. A., Cheng, N., Couling, M. and Brock, H. W. (1994). Mutations in some Polycomb group genes of *Drosophila* interfere with regulation of segmentation genes. Mol. Gen. Genet. 244, 474-483.
- Mohler, J., Seecoomar, M., Agarwal, S., Bier, E. and Hsai, J. (2000). Activation of knot (kn) specifies the 3-4 intervein region in the Drosophila wing. Development 127, 55-63.
- Montagne, J., Groppe, J., Guillemin, K., Krasnow, M. A., Gehring, W. J. and Affolter, M. (1996). The Drosophila Serum Response Factor gene is required for the formation of intervein tissue of the wing and is allelic to blistered. Development 122, 2589-2597.
- Muller, B., Hartmann, B., Pyrowolakis, G., Affolter, M. and Basler, K. (2003). Conversion of an extracellular Dpp/BMP morphogen gradient into an inverse transcriptional gradient, Cell 113, 221-233.
- Nellen, D., Burke, R., Struhl, G. and Basler, K. (1996). Direct and longrange action of a DPP morphogen gradient. Cell 85, 357-368.
- Nussbaumer, U., Halder, G., Groppe, J., Affolter, M. and Montagne, J. (2000). Expression of the blistered/DSRF gene is controlled by different morphogens during Drosophila trachea and wing development. Mech. Dev. **96**, 27-36.
- Orlando, V. (2003). Polycomb, epigenomes, and control of cell identity. Cell 112, 599-606
- Papoulas, O., Beek, S. J., Moseley, S. L., McCallum, C. M., Sarte, M., Shearn, A. and Tamkun, J. W. (1998). The Drosophila trithorax group proteins BRM, ASH1 and ASH2 are subunits of distinct protein complexes. Development 125, 3955-3966.
- Pelegri, F. and Lehmann, R. (1994). A role of polycomb group genes in the regulation of gap gene expression in Drosophila. Genetics 136, 1341-1353.
- Podos, S. D. and Ferguson, E. L. (1999). Morphogen gradients: new insights from DPP. Trends Genet. 15, 396-402.
- Ringrose, L., Rehmsmeier, M., Dura, J. M. and Paro, R. (2003). Genome-

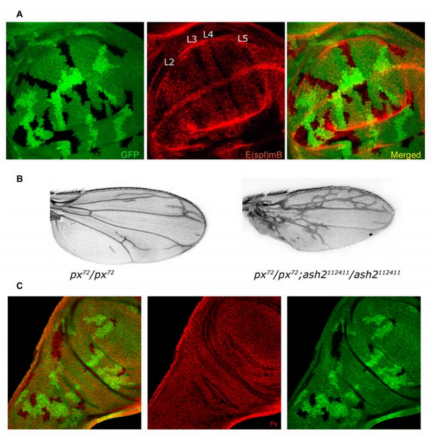
- wide prediction of Polycomb/Trithorax response elements in Drosophila melanogaster, Dev. Cell 5, 759-771.
- Roch, F., Baonza, A., Martin-Blanco, E. and Garcia-Bellido, A. (1998). Genetic interactions and cell behaviour in blistered mutants during proliferation and differentiation of the Drosophila wing. Development 125, 1823-1832
- Saget, O., Forquignon, F., Santamaria, P. and Randsholt, N. B. (1998). Needs and targets for the multi sex combs gene product in Drosophila melanogaster. Genetics 149, 1823-1838.
- Schnepp, B., Grumbling, G., Donaldson, T. and Simcox, A. (1996). Vein is a novel component in the Drosophila epidermal growth factor receptor pathway with similarity to the neuregulins. Genes Dev. 10, 2302-2313.
- Shearn, A. (1974). Complementation analysis of late lethal mutants of Drosophila melanogaster. Genetics 77, 115-125.
- Shearn, A. (1989). The ash-1, ash-2 and trithorax genes of Drosophila melanogaster are functionally related. Genetics 121, 517-525.
- Shearn, A., Rice, T., Garen, A. and Gehring, W. (1971). Imaginal disc abnormalities in lethal mutants of Drosophila. Proc. Natl. Acad. Sci. USA
- Shearn, A., Hersperger, E. and Hersperger, G. (1987). Genetic studies of mutations at low loci of Drosophila melanogaster which cause a wide variety of homeotic transformations, Roux's Arch. Dev. Biol. 196. 231-242
- Simon, J. (1995). Locking in stable states of gene expression: transcriptional control during Drosophila development. Curr. Opin. Cell Biol. 7, 376-385.
- Simon, J. A. and Tamkun, J. W. (2002). Programming off and on states in chromatin: mechanisms of Polycomb and trithorax group complexes. Curr. Opin. Genet. Dev. 12, 210-218.
- Strigini, M. and Cohen, S. M. (1999). Formation of morphogen gradients in the Drosophila wing. Semin. Cell Dev. Biol. 10, 335-344.
- Sturtevant, M. A. and Bier, E. (1995). Analysis of the genetic hierarchy guiding wing vein development in Drosophila. Development 121, 785-801.
- Sturtevant, M. A., Roark, M. and Bier, E. (1993). The Drosophila rhomboid gene mediates the localized formation of wing veins and interacts genetically with components of the EGF-R signaling pathway. Genes Dev. 7, 961-973
- Sturtevant, M. A., Biehs, B., Marin, E. and Bier, E. (1997). The spalt gene links the A/P compartment boundary to a linear adult structure in the Drosophila wing. Development 124, 21-32.
- Tabata, T. and Kornberg, T. B. (1994). Hedgehog is a signaling protein with a key role in patterning Drosophila imaginal discs. Cell 76, 89-102.
- Vervoort, M., Crozatier, M., Valle, D. and Vincent, A. (1999). The COE transcription factor Collier is a mediator of short-range Hedgehog-induced patterning of the Drosophila wing. Curr. Biol. 9, 632-639.
- Xu, T. and Rubin, G. M. (1993). Analysis of genetic mosaics in developing and adult Drosophila tissues. Development 117, 1223-1237.
- Zecca, M., Basler, K. and Struhl, G. (1995). Sequential organizing activities of engrailed, hedgehog and decapentaplegic in the Drosophila wing. Development 121, 2265-2278.

Supplemental Text and Figures

ASH2 specifically regulates some intervein promoting genes

ASH2 can play a role as a regulator of intervein promoting genes as shown by the downregulation of Blistered and *net* observed in *ash2* mutant clones and *ash2* mutant discs respectively. Both observations are consistent with the phenotypes arising from genetic interaccions. In order to asses if the downregulation of this kind of genes by ASH2 is a general effect, we carried out genetic interactions and analyzed the expression of other

intervein genes such as $E(spl)m\beta$ or px. Although the px^{72} al·lele, which displays by itself an extra vein phenotype, enhances the phenotype of $ash2^{112411}$ homozygous wings (Supp. Fig. 1B), we have not been able to detect a misregulation of neither Plexus nor $E(spl)m\beta$ in $ash2^{11}$ mutant clones (Supp. Fig. 1C and A). Taken together, this results indicate that ASH2 is not involved in the regulation of $E(spl)m\beta$ and Px and that the downregulation of other intervein promoting genes such as bs and net in ash2 mutant cells is a specific effect.



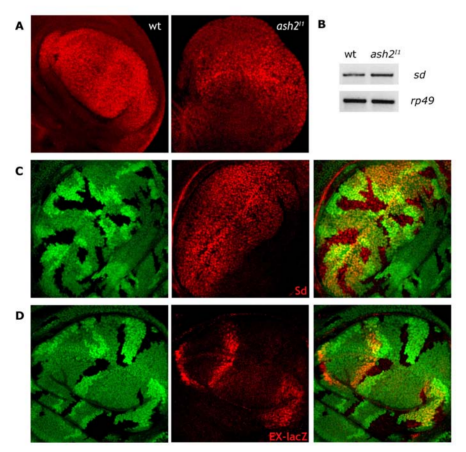
Supp. Fig. 1 - (A) E(spl)mβ-lacZ is not misexpressed in $ash2^{l1}$ mutant clones. (B) Interaction between px^{l2} and $ash2^{l124l1}$ mutants results in an enhancement of the phenotypes. (C) Px protein levels are not altered in $ash2^{l1}$ mutant clones. In A and C, homozygous $ash2^{l1}$ cells are black, heterozygous cells are light green and wt cells are bright green.

ASH2 inhibits knirps expression beyond L2

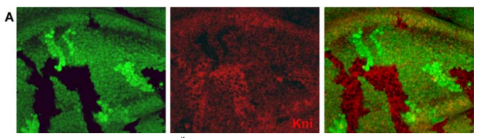
Since the de-repression of the vein organizing gene *kni* in *ash2* mutant clones (Fig. 7) can be direct or mediated by the misregulation of other genes controling *kni* expression, we have further analyzed this subject. The characterization of the L2 enhancer of *kni* revealed that it harbours binding sites for the Scalloped/Vestigial complex (Sd/Vg), En, Brk and Salr (Lunde et al., 2003). The ASH2-induced repression of *kni* is independent of Brk (Fig. 6) and Sd (Supp. Fig. 2A-C) since we could not detect an alteration of their expression in

ash2 mutant clones. We could neither detect an alteration of the expression of the minimal L2 enhancer (EX-lacZ) (Supp. Fig. 2D).

Salm is weakly downregulated in *ash2*¹¹ mutant clones (Fig. 6) but we have demonstrated that this is not the cause of Kni derepression, since the latter is upregulated even when Salm is overexpressed (Fig.8). Furthermore, we confirmed that ASH2 mediated *kni* derepression is also Salr-independent (Supp. Fig. 3A). Taken together, these results suggest that the ASH2 regulatory function is independent of the Sal-C and of the L2 enhancer.



Supp. Fig. 2 - (A) Sd pattern of expression is not altered in the wing pouch of wild-type and $ash2^{II}$ mutant wing imaginal discs. (B) Semiquantitative RT-PCR shows that levels of the sd transcript are not altered in $ash2^{II}$ mutant larvae. (C) Sd protein levels are not remarkably affected in $ash2^{II}$ mutant clones. (D) EX-lacZ enhancer expression is not significantly affected in $ash2^{II}$ mutant clones. In C and D, homozygous $ash2^{II}$ cells are black, heterozygous cells are light green and wt cells are bright green.



Supp. Fig. 3 - (A) Kni is upregulated in $ash^{2^{lt}}$ clones even when salr is overexpressed with nub-gal4. Homozygous $ash^{2^{lt}}$ cells are black, heterozygous cells are light green and wt cells are brigth green. The overexpression of Salr can be infered because the L2 vein is missing, as it happens with adult flies.

ash2 is more crucial for vein pattern formation than other trxG genes

To gain further insight into *ash2* function we compared its mutant phenotypes with those of other trxG mutant aleles through clonal analysis in

the adult wing. We produced Minute and Twin clones by irradiating the progeny larvae of the following crosses with X rays at 80 ± 12h and 60± 12h.

TWIN CLONES

 f^{36a} ; mwh bld cn $f^{+98D}/TM1$,mwh x tr $x^{E2}/TM6b$ f^{36a} ; mwh jv $f^{+77A}/TM2$ x ash $1^{B1}/TM6c$ f^{36a} ; mwh jv $f^{+77A}/TM2$ x brm $^2/TM6b$ f^{36a} ; mwh bld cn $f^{+98D}/TM1$, mwh x osa $^2/TM6b$ f^{36a} ; mwh bld cn $f^{+98D}/TM1$, mwh x mor $^1/TM6b$

MINUTE CLONES

f^{36a}; f^{+87D}M(3) w¹²⁴ /TM3 x trx^{E2}/TM6b f^{36a}; M(3) f^{+77A}/TM3 x ash1^{B1}/TM6c f^{36a}; M(3) f^{+77A}/TM3 x brm²/TM6b f^{36a}; f^{+87D}M(3) w¹²⁴ /TM3 x osa²/TM6b f^{36a}; f^{+87D}M(3) w¹²⁴ /TM3 x mor¹/TM6b

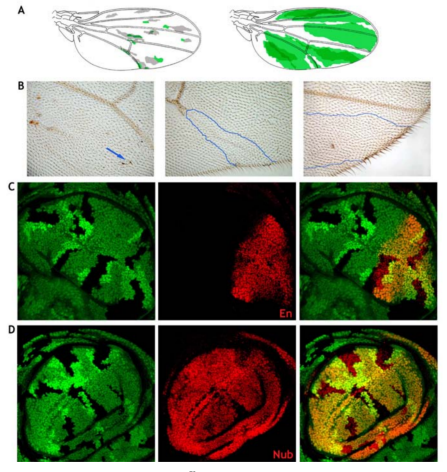
As previously described, *brm, mor* and *osa* are essential for cell viability (Brizuela and Kennison, 1997; Elfring et al., 1998; Treisman et al., 1997) since we were not able to obtain mutant clones for the alleles studied. Thus, we focused on the effects of *trx* and *ash1* mutant alleles, for which we could indeed obtain mutant clones. Supplemental Tables 1 and 2 indicate the

frequency of the clones obtained and a summary of the phenotypes observed (Supp. Figs. 4 and 5) in trx and ash1 mutant clones respectively. These results show that the phenotypes observed in ash2 are different or more evident, in terms of vein patterning, than the ones found in trx and ash1 mutant clones.

trxE2 mutant clones

	Frequence(clones/wing)	Effects
Twin clones	13/76 (0,17%)	- Alteration of cell viability
Minute clones	9/149 (0,06%)	- Vein disruption
		- Anteriorization of the posterior
		compartment, with differenciated socketed
		bristles that ressemble the ones from the
		anterior wing margin.

Supp. Table 1. Frequency of tnx^{E2} mutant clones obtained with the Twin clone and the *Minute* techniques. The most common phenotypes observed in both cases are summarized. Supp. Fig 4 shows some examples of these phenotypes.



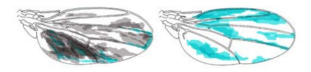
Supp. Fig. 4 - (A) Plot of the localization of tnx^{E2} mutant clones (green) and twin clones (grey) in the wing. (B) The loss of tnx activity results in anterioritzation of the posterior compartment as seen by the appearance of socketed bristles in intervein E (blue arrow, right pannel) and in the posterior margin (left pannel). Loss of L5 vein tissue is also observed (middle pannel). The levels of En (C) and Nub (D) proteins (in red) are decreased in tnx^{E2} mutant clones. In (C) and (D), homozygous tnx^{E2} cells are black, heterozygous cells are light green and wt cells are brigth green.

ash1^{B1} mutant clones

	Frequence(clones/wing)	Effects
Twin clones	34/221 (0,15%)	- Survival and cell proliferation defects
Minute clones	24/362 (0,06%)	- Triple Row disruption
		- Extra pigmentation that reflects a
		failure in cell differentiation
		- Size reduction of the compartment
		where the clone is localized

Supp. Table 2. Frequency of *ash1*^{B1} mutant clones obtained with the Twin clone and the *Minute* techniques. The most common phenotypes observed in both cases are summarized. Supp. Fig 5 shows some examples of these phenotypes.

A)Plot of ash181 mutant clones



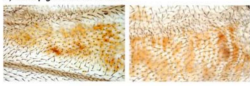
B) Reduction of the compartment size where the clone is localized



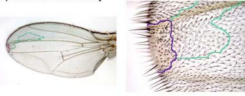
C) Triple Row disruption



D) Extra pigmentation defects



E) Effects on cell viability



Supp. Fig. 5 - (A) Plot of $ash1^{B1}$ mutant clones (blue) and twin clones (grey).(B),(C),(D) and (E) show different effects of loss of ash1 activity.

Supplemental References

Brizuela, B. J. and Kennison, J. A. (1997). The Drosophila homeotic gene moira regulates expression of engrailed and HOM genes in imaginal tissues. *Mech Dev* 65, 209-20. Elfring, L. K., Daniel, C., Papoulas, O., Deuring, R., Sarte, M., Moseley, S., Beek, S. J., Waldrip, W. R., Daubresse, G., DePace, A. et al. (1998). Genetic analysis of brahma: the Drosophila homolog of the yeast chromatin remodeling factor SWI2/SNF2. *Genetics* 148, 251-65.

Lunde, K., Trimble, J. L., Guichard, A., Guss, K. A., Nauber, U. and Bier, E. (2003). Activation of the knirps locus links patterning to morphogenesis of the second wing vein in Drosophila. *Development* **130**, 235-48.

Treisman, J. E., Luk, A., Rubin, G. M. and Heberlein, U. (1997). eyelid antagonizes wingless signaling during Drosophila development and has homology to the Bright family of DNA-binding proteins. *Genes Dev* **11**, 1949-62.