

Published in final edited form as:

Nature. 2009 April 23; 458(7241): 987–992. doi:10.1038/nature07936.

Genome-wide analysis of Notch signalling in *Drosophila* by transgenic RNAi

Jennifer L. Mummery-Widmer^{1,*}, Masakazu Yamazaki^{1,*†}, Thomas Stoeger¹, Maria Novatchkova^{1,2}, Sheetal Bhalerao^{1,2}, Doris Chen³, Georg Dietzl², Barry J. Dickson², and Juergen A. Knoblich¹

¹Institute of Molecular Biotechnology of the Austrian Academy of Sciences (IMBA), Dr Bohr-Gasse 3, A-1030 Vienna, Austria

²Research Institute of Molecular Pathology (IMP), Dr Bohr-Gasse 7, A-1030 Vienna, Austria

³Max F. Perutz Laboratories (MFPL), Department of Biochemistry, Dr Bohr-Gasse 9, A-1030 Vienna, Austria

Abstract

Genome-wide RNA interference (RNAi) screens have identified near-complete sets of genes involved in cellular processes. However, this methodology has not yet been used to study complex developmental processes in a tissue-specific manner. Here we report the use of a library of *Drosophila* strains expressing inducible hairpin RNAi constructs to study the Notch signalling pathway during external sensory organ development. We assigned putative loss-of-function phenotypes to 21.2% of the protein-coding *Drosophila* genes. Using secondary assays, we identified 6 new genes involved in asymmetric cell division and 23 novel genes regulating the Notch signalling pathway. By integrating our phenotypic results with protein interaction data, we constructed a genome-wide, functionally validated interaction network governing Notch signalling and asymmetric cell division. We used clustering algorithms to identify nuclear import pathways and the COP9 signalosome as Notch regulators. Our results show that complex developmental processes can be analysed on a genome-wide level and provide a unique resource for functional annotation of the *Drosophila* genome.

Genome-wide RNAi screens have been performed in cultured cells¹⁻⁴ or by ubiquitous gene silencing in worms^{5,6} or planarians⁷. To study complicated developmental processes, however, genes need to be inactivated in a tissue-specific manner in intact animals. This has become possible through the creation of a transgenic RNAi library targeting 88% of the *Drosophila*⁸ protein-coding genes. To test the feasibility of this new approach, we focused on the Notch pathway, one of the most important regulators of development^{9,10}. Notch is activated by binding to its ligands Delta or Serrate. After ligand binding, Notch is cleaved by Presenilin and the intracellular domain acts in the nucleus as a transcriptional co-activator.

©2009 Macmillan Publishers Limited. All rights reserved

Correspondence and requests for materials should be addressed to J.A.K. (juergen.knoblich@imba.oeaw.ac.at).

[†]Present address: The Global COE program, Akita University School of Medicine, 1-1-1 Hondo, Akita 010-8543, Japan.

*These authors contributed equally to this work.

Author Contributions J.L.M.-W., M.Y. and J.A.K. designed the experiments. J.L.M.-W. and M.Y. carried out the genetic screen and secondary analysis. T.S. contributed to the secondary analysis. M.N. performed Bioinformatics data analysis. D.C. and S.B. contributed to generation of the second RNAi lines. G.D. and B.J.D. generated and provided the RNAi library. J.L.M.-W. and J.A.K. wrote the paper.

Author Information Reprints and permissions information is available at www.nature.com/reprints.

Supplementary Information is linked to the online version of the paper at www.nature.com/nature.

Full Methods and any associated references are available in the online version of the paper at www.nature.com/nature.

Several important Notch regulators were identified due to their characteristic phenotypes in *Drosophila* external sensory organs^{11, 12}. Adult external sensory organs are formed within the notum (dorsal thorax) during larval and pupal stages of *Drosophila* development (Fig. 1a). Within a group of epithelial cells called the proneural cluster, a single sensory organ precursor (SOP) is selected in a Notch-dependent process termed lateral inhibition. The SOP cell divides into pIIa and pIIb (Fig. 1a). While pIIa forms the two outer cells (the hair and socket), pIIb gives rise to an apoptotic glial cell and then to the two inner cells (the neuron and sheath) of the organ. During each division, the protein determinants Numb and Neuralized segregate into one of the two daughter cells where Numb inhibits the Notch receptor while Neuralized activates the Notch ligand Delta^{10, 13}. The resulting differences in Notch signalling establish unequal fates in the two daughter cells. Thus, Notch acts both in lateral inhibition and in asymmetric cell division (ACD) during external sensory organ development. Although the core components of the Notch pathway act in both processes, other components act specifically during lateral inhibition (E(spl)¹⁴) or ACD (Numb¹⁵, Sanpodo^{16, 17}).

Genome-wide RNAi screen

Hairpin constructs in the RNAi library are expressed under UAS/GAL4 control¹⁸. We tested *scabrous*-GAL4¹⁸, *pannier* (*pnr*)-GAL4¹⁹ and *fzIII*-GAL4 (also known as *MS248*-GAL4 or *P{GawB}MS248*)²⁰ using a set of 40 RNAi lines targeting 21 genes involved in external sensory organ development (Supplementary Table 1). Consistently, phenotypes were stronger and lethality lower with *pnr*-GAL4, and this line was selected for large-scale analysis.

A total of 20,262 transgenic RNAi lines were screened; these are predicted to target 11,619 of the 14,139 protein-coding genes (82.2%) in release 5.7 of the *Drosophila* genome²¹. Ten flies each were analysed and phenotypic abnormalities were recorded in a database (<http://bristlescreen.imba.oeaw.ac.at>). Because *pnr*-GAL4 is only expressed in a central region of the notum (Fig. 1b), lateral areas were unaffected and served as internal controls. Phenotypes were described using controlled vocabulary (Fig. 1b, Supplementary Table 2). Phenotypic strength (P_x) was expressed on a scale of 0 (not affected) to 10 (completely affected) as the fraction of the *pnr*-GAL4 expression area (or the fraction of external sensory organs present in that area) that displayed the respective phenotype. External sensory organ phenotypes were classified into 'gain' (P_{Gain}), 'loss' (P_{Loss}), 'empty or multiple sockets' (P_{Sockets}), 'hair cell duplication' ($P_{\text{Duplication}}$) and 'bristle morphology defects' (P_{BMorph}). In addition, orientation defects were recorded as 'planar polarity defects' (P_{Polarity}). Morphological notum defects were classified into 'notum malformation death' (P_{MDeath} ; loss of tissue), 'notum malformation migration' ($P_{\text{MMigration}}$; thorax closure defects), 'overproliferation' ($P_{\text{Overprolif}}$; notum enlargement) and 'colour' (P_{Colour}). GO term analysis showed that phenotype groups are significantly enriched for genes known to act in appropriate processes (Supplementary Table 3). The group of 211 genes for which $P_{\text{Overprolif}}$ is >0 contains tumour suppressors such as *Pten* and 5 of the 7 Hippo pathway members represented in the library. The group of 71 genes with P_{Polarity} of 4 contains four of the six core planar cell polarity (PCP) components, and four other PCP effectors. Visible phenotypes were scored for 19.6% of all RNAi lines (3,973 lines, 3,004 genes). A total of 1,803 lines (8.9% of all lines, 1,339 genes) were lethal whereas 496 lines (2.5% of all lines, 475 genes) were semi-lethal with no associated morphological defects. Sixty-nine per cent of lines (13,990 lines, 8,681 genes) gave no phenotype (Fig. 1c). Thus, our screen has assigned potential loss-of-function phenotypes to 21.2% (3,004 of 14,139) of the protein-coding genes in the *Drosophila* genome, more than half of which (1,597 genes, 2,067 lines) had not been characterized before.

Quality control

To determine false-negative rates, we assembled a list of genes required for Notch signalling and/or ACD based on Flybase and more recent published information. Forty-two of the 50 selected genes are represented in the RNAi library (Supplementary Table 4) and 24 gave the expected phenotype. Eight genes could not be analysed owing to early lethality (six genes) or strong notum malformations (*cdc2* and *bre1*). The resulting false-negative rate of 29.4% (10 of 34 genes) is in good agreement with previously reported data for transgenic RNAi⁸.

Long double-stranded RNA hairpins are known to cause off-target effects²². Nonspecific phenotypes can also be generated by overexpression of genes near the transgene insertion site. We therefore generated a second hairpin construct or obtained a second RNAi line from the NIG-FLY stock centre for 73 genes (136 lines) that had resulted in visible phenotypes (Supplementary Table 5). Similar or identical phenotypes were observed for 63.0% of the genes (46 genes, 82 lines). Interestingly, however, we could confirm 88% (22 of 25 genes) of those phenotypes that were supported by more than one line in the library, although most of these lines are independent insertions of the same transgene. We also selected 43 non-essential genes (72 lines) with known adult phenotypes (or the absence thereof; Supplementary Table 6). For 93% of these (40 genes, 67 lines), our screen identified the correct phenotype. In three cases, the observed phenotypes were different from what has been described, but could be explained by the described cellular function (endocytosis for *orange*, Ras activity for *Neurofibromin 1*, and cAMP signalling for *dunce*) if RNAi causes phenotypes stronger than the described alleles. In any case, these data indicate that the false-positive rate of our screen is less than 7%.

Phenotypic categories

Notch acts twice during external sensory organ development²³. During lateral inhibition, loss and gain of Notch signalling increases and decreases the number of external sensory organs, respectively. During ACD, increased Notch signalling causes hair duplications or multiple sockets, whereas decreased Notch signalling causes all daughter cells to assume an internal fate and results in the apparent absence of external sensory organs. Thus, Notch regulators can be in the loss, gain, socket and duplication categories. We therefore used the phenotypes of genes known to be involved in Notch signalling and ACD (data not shown) to create compound categories that include all or most of these positive-control genes. The asymmetric cell division (ACD) category ($P_{\text{Loss}} > 7$ or $P_{\text{Duplication}} > 0$ or $P_{\text{Sockets}} > 0$) is made up of 226 genes (262 lines) and includes *bora*, *bazooka*, *Delta*, *musashi*, *Notch*, *numb*, *Presenilin*, *sanpodo*, *tramtrack* and *twins*. The lateral inhibition category includes all genes for which P_{Loss} is > 7 . In addition, we re-screened all images of genes where P_{Gain} is > 0 and retained only those where the extra external sensory organs are grouped together, indicating the formation of multiple SOP cells from one proneural cluster. The resulting lateral inhibition category contains 233 genes (264 lines). Analysis of all genes in these categories revealed (Supplementary Table 3) that the GO terms ‘asymmetric cell division’ and ‘nervous system development’ are significantly enriched in the ACD category. The lateral inhibition category over-represents the GO terms ‘neurogenesis’, ‘Notch signalling pathway’ and ‘cell fate commitment’. We therefore used these categories in various secondary assays to identify genes acting globally in the Notch pathway or more specifically during ACD.

Identification of Notch regulators

During wing development, Notch loss of function leads to wing notching or broader wing veins²⁴. To identify general Notch regulators, we expressed RNAi hairpins in the wing imaginal disc for 201 genes (226 lines) in the ‘lateral inhibition’ category. We analysed ten

flies each for wing vein and wing margin phenotypes (Fig. 2, Supplementary Fig. 1 and Supplementary Table 7). All other wing abnormalities were also recorded and separated into phenotypic classes (Supplementary Fig. 2). In addition to seven known Notch pathway members (*Notch*, *mind bomb 1*, *Delta*, *groucho*, *extra macrochaetae*, *Presenilin* and *O-fucosyltransferase 1 (O-fut1)*), this secondary analysis identified 23 genes not previously implicated in Notch signalling.

Notch signalling activates the transcription factor Suppressor of Hairless (Su(H)). The genes identified could participate in this signal transduction cascade, act in parallel to Su(H) or act as Su(H) targets. To distinguish these, we used a *lacZ* reporter for Su(H) activity²⁵ (Supplementary Table 7). In the developing wing disc, this reporter is expressed along the dorso-ventral compartment boundary (Fig. 2c). When we inactivated the potential Notch regulators along the anterior-posterior compartment boundary using *patched (ptc)*-GAL4, 8 genes (11 lines) caused a loss of β -galactosidase staining in the area where the boundaries overlap (Fig. 2d-k). Besides *Notch* itself and the known Notch regulators *mind bomb 1* and *O-fut1*, this identified five genes that function upstream of Su(H)-mediated transcriptional activation and have not previously been implicated in Notch signalling.

Notch signalling is tightly connected to endocytic trafficking¹³. Interestingly, one of the newly identified Notch regulators (CG5608) is homologous to *VAC14*, a yeast gene that controls trafficking to multivesicular bodies²⁶. Vac14 is an upstream activator of the PtdIns(3)P 5-kinase Fab1 (also known as PIKfyve)²⁶. Although the function of *Drosophila* CG5608 has not been analysed, Notch accumulates with Delta in intracellular vesicles in *fab1* mutant *Drosophila* cells²⁷. Further analysis of the CG5608 RNAi phenotype might help to understand the connection between phosphoinositide synthesis and Notch signalling. This is particularly interesting because *Vac14* loss results in neurodegeneration in mice²⁸, and multiple lines of evidence have connected the Notch pathway to neurodegeneration²⁹ before.

Genes regulating asymmetric cell division

Loss of external sensory organs can be caused either by increased Notch signalling during lateral inhibition or by reduced Notch signalling during ACD²³ (Fig. 3a). To resolve this complication, we assayed the presence of SOP cells by expressing green fluorescent protein (GFP) from the *phyllopod (phyl)* promoter (Fig. 3b). *phyl* is expressed in the proneural cluster and then restricted to the SOP³⁰. GFP was fused to the asymmetric localization domain of Partner of numb (Pon) so that the construct can also be used to analyse asymmetric protein segregation. We imaged at least 3 pupae each for 80 genes (91 lines) in the ACD category (Fig. 3b, Supplementary Fig. 3 and Supplementary Table 8). Thirty-three genes (33 lines) caused a complete loss of SOPs within the *pnr* expression domain whereas 6 genes (8 lines) caused a partial SOP loss. Eight genes (13 lines) increased SOP number, indicating that they are required for lateral inhibition. Importantly, 7 of these genes were tested with *MS1096*-GAL4 (also known as *P{GawB}Bx^{MS1096}*), and 5 caused Notch loss-of-function wing phenotypes (Supplementary Table 7), demonstrating that they are general regulators of Notch signalling. Finally, we identified 28 genes (31 lines) that may have a role in ACD because SOP cells were normal but no differentiated external sensory organs developed.

To distinguish cell fate transformations from differentiation defects, we analysed 23 of these genes (26 lines) using cell-type-specific markers (Fig. 3c and Supplementary Table 9). Clear transformations of outer into inner cells were observed for six genes (six lines) whereas the others showed either normal lineages or abnormalities that can not be explained by cell fate transformations. Taken together with the 128 additional genes (146 lines) for which loss of

function caused P_{Sockets} of >0 or $P_{\text{Duplication}}$ of >0 , we have identified 134 regulators of ACD.

One of these is *capricious* (*caps*), a transmembrane protein implicated in cell–cell interactions during compartment boundary formation^{31,32} and neuronal targeting³³. On *caps* RNAi, SOP cells generate two pIIb cells, a phenotype typically observed for genes involved in Notch activation (Fig. 3c). Indeed, *caps*^{*c18fs*} mutant clones show phenotypes indicating a loss of Notch signalling (Supplementary Fig. 4), suggesting that Caps might be involved in generating cell adhesion between signal-sending and -receiving cells during Notch/Delta signalling.

A genome-wide network for Notch signalling

Genes or proteins in a signalling pathway often interact genetically or physically. Several attempts have been made to map such interactions on a genome-wide level^{34–36}, but these lack functional validation. To create a functionally validated map for Notch signalling, we first combined all genes in the ‘lateral inhibition’ or ‘asymmetric cell division’ categories and added genes previously implicated in Notch signalling that had not scored as positive in our screen. For each gene we searched for yeast two-hybrid, biochemical, genetic and other interaction data from various databases (see Methods). We only maintained interactions with partners also contained in our candidate set and removed all genes without interaction partners. The resulting gene network (<http://bristlescreen.imba.oeaw.ac.at>) includes 780 interactions between 177 genes, only 42 of which were previously known to act in Notch signalling during external sensory organ development (Fig. 4 and Supplementary Fig. 5).

To identify putative complexes, we used the Molecular Complex Detection (MCODE) and found nine clusters of highly interconnected nodes (Fig. 4). In protein–protein interaction networks, such clusters represent protein complexes and/or genetic pathways. Complex one is centred on Notch and Delta. It includes Notch-modifying enzymes, such as *O*-fucosyltransferase, and proteins in the Notch signalling pathway, such as the Presenilin complex and the nuclear protein Hairless. It also includes the guanine nucleotide exchange factor protein Trio, which has been shown to interact with Notch³⁷ but had not been functionally implicated in Notch signalling before. A second complex includes the known ACD regulators G- α 47A, Rapsynoid (also known as Pins) and Bazooka. Thus, network analysis of genetic data can successfully predict pathways that mediate even complex multicellular processes.

To validate the predictive capacity of our network, we tested complexes in which most members were not known to regulate Notch signalling before. Complex nine contains the importin α protein Karyopherin α 3 (Kap- α 3) as well as the importin α export receptor Cas (Fig. 5a). *Cas* mutants were previously shown to cause ACD defects³⁸. In *Kap- α 3* mutant clones (Fig. 5b, c), we observed strongly penetrant cell fate transformations similar to *Cas*. Neighbouring genes to complex nine include the nuclear pore components *Nup358* and *Nup50* (CG2158) (Fig. 5a). In all cases, RNAi causes cell fate transformations characteristic of increased Notch activity suggesting that nuclear import of an inhibitory component is rate limiting for the Notch pathway. Such a component has been postulated before³⁸ and the identification of the critical importin- α might facilitate its biochemical identification. Because the Kap- α 3–Cas complex also contains nuclear Lamin (Fig. 5a), our data might provide explanations for certain aspects of laminopathies. Interestingly, deregulated Notch signalling has previously been suggested to be crucially involved in certain laminopathies based on their stem cell phenotype³⁹.

The predicted complex five (Fig. 5d) contains subunits of the COP9 signalosome (CSN). RNAi of six of the seven CSN subunits screened resulted in occasional generation of two

pIIa cells (1 $P_{\text{Duplication}}$ 3), consistent with an inhibitory function in the Notch pathway (Fig. 5e, f and Supplementary Fig. 6). The phenotypes are also observed in mitotic clones mutant for *CSN4* or *CSN5* (Fig. 5g, h). CSN is an isopeptidase that removes Nedd8 conjugates from Cullin-RING class E3 ubiquitin ligases and prevents their degradation⁴⁰. Indeed, defects in lateral inhibition are also observed in *Cullin-3* (*gft*) mutants⁴¹ suggesting that *Cullin-3* might be the relevant CSN target during Notch signalling.

Conclusions

Our data show that genome-wide loss-of-function analysis by RNAi can be used to study complex developmental processes. From a comprehensive phenotypic database we extracted genes involved in Notch signalling by secondary screening or integration with other types of genome-wide analysis. Vac14 may regulate PtdIns(3,5)P₂ synthesis and trafficking of Notch or its ligand Delta into multivesicular bodies. Kap- α 3 is a cargo for Cas and might control the nuclear import of a rate limiting inhibitory signalling component. Finally, the COP9 signalosome regulates Cullin-1 (*lin19*) and Cullin-3 (ref. ⁴²), which in turn could regulate ubiquitylation of Notch or essential Notch pathway members. We describe an additional 23 uncharacterized genes and 5 putative protein complexes for which the mechanistic connection to Notch signalling is less clear. Most of those act downstream or in parallel to Su(H). So far, only a few functionally important nuclear targets of Notch have been identified. Our analysis could considerably expand the list of nuclear Notch targets and provide insight into how Notch regulates the various cell fate and morphology decisions in which it has been implicated.

In addition, we report a wide range of morphological phenotypes that can be used to derive groups of genes potentially involved in planar polarity, proliferation control, cell migration and apoptosis. The possibility to perform secondary screens on those groups illustrates the enormous power of transgenic RNAi now available to study development on a genome-wide level.

METHODS SUMMARY

For genome-wide and second RNAi line screening, crosses between UAS-hairpin RNAi males and *pnr*-GAL4 females were raised at 22 °C and shifted to 29 °C during larval stages. Ten flies were examined visually and all morphological abnormalities were recorded. Phenotypic classes are defined in Methods.

For secondary assays of wing morphology or Notch reporter activity, *MS1096*-GAL4 or *ptc*-GAL4; *Gbe*+*Su(H)*-*lacZ* were used, respectively (both at 25 °C). Lineage analysis was essentially as described¹⁵ using *pnr*-GAL4 at 29 °C. For live imaging⁴³, at least three UAS-RNAi; *pnr*-GAL4, *phyllopod*>>EGFP::Pon.LD pupae were imaged at 14 h APF (aged at 29 °C) on a Zeiss LSM510 confocal microscope. *phyllopod*>>EGFP::Pon.LD was generated by fusing enhanced GFP (EGFP) to the amino terminus of the Pon localization domain and placing it downstream of a 3.4 kb *phyllopod* promoter fragment³⁰. Mutant clones were generated using the *Ubx*-FLP/FRT system with (*Kap- α 3*^{D93}, *CSN4*^{null} and *CSN5*^{null} clones) or without (*caps*^{c18fs} clones) a cell lethal on the non-mutant chromosome arm¹⁷.

The Notch interaction network was generated by combining interaction data from the DroID, STRING⁴⁴ and BioGRID⁴⁵ databases. Interactions were only maintained if both interactors were contained in the ‘lateral inhibition’ or ‘asymmetric cell division’ categories or a list of previously characterized Notch regulators. The resulting network was drawn using Cytoscape⁴⁶. The MCODE⁴⁷ algorithm was used to identify sub-network complexes.

A list of fly stocks and antibodies used as well as further methodological details are in the Supplementary Information.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

Antibodies, plasmids and fly stocks were provided by the Vienna *Drosophila* RNAi Center, the Bloomington *Drosophila* Stock Center, the National Institute of Genetics Fly Stock Center (NIG-FLY), the Developmental Studies Hybridoma Bank, C.-T. Chien, R. J. Fleming, T. Klein, R. Lehmann, A. Nose, G. S. Suh and F. Wirtz-Peitz. We thank C. Cowan for comments on the manuscript, V. Rolland for help with secondary analysis, E. Kleiner, G. Haas, S. Reiter, T. Pritz, Z. Topalovic, S. Farina Lopez, S. Wculek and Ö. Copur for technical assistance in constructing the custom second RNAi line collection, C. Gallagher for generating the Su(H) antibody, P. Pasierbek for bio-optics support, F. Stocker for graphics assistance and P. Serrano Drozdowskyj for creating the online database. M.Y. was supported by a European Union Marie Curie Mobility Fellowship. Work in J.A.K.'s laboratory is supported by the Austrian Academy of Sciences, the Wiener Wissenschafts-, Forschungs- und Technologiefonds (WWTF), the Austrian Science Fund (FWF) and the EU network ONCASYM.

METHODS

Fly strains and constructs

The following fly strains were used: *pnr*-GAL4 (MD237; Bloomington *Drosophila* Stock Center (BDSC)); *sca*-GAL4 (G537.4)¹⁸; *fzIII*-GAL4 (MS248)^{20, 48}; *MS1096*-GAL4 (BDSC); *ptc*-GAL4 (G559.1)¹⁸; *Gbe+Su(H)-lacZ*²⁵ (a gift from T. Klein); FRT82B, *Kap-α3^{D93}* (a gift from R. J. Fleming); *CSN4^{null}* and FRT82B, *CSN5^{null}* (gifts from G. S. Suh); FRT2A, *caps^{c18fs}* (a gift from A. Nose); and *pnr*-GAL4, *phyllopod*>>EGFP::Pon.LD. All RNAi lines came from the Vienna *Drosophila* RNAi Center (VDRC) library and their details are published⁸, except second RNAi lines, which were custom made (see below) or supplied by the National Institute of Genetics Fly Stock Center (NIG-FLY). VDRC UAS-driven inverted repeat (UAS-IR) construct target prediction mapping to release 5.7 of the *Drosophila* genome (Flybase release FB2008_04) was carried out essentially as described⁸, except that all genes targeted by 80% of 19-base polymers were considered as on-target genes. For this reason, one construct may have more than one target. When no target was identified by automated mapping efforts, constructs were annotated manually and identified as 'manual mapping' (Supplementary Table 2 and <http://bristlescreen.imba.oeaw.ac.at>). Only protein-coding gene targets were considered when calculating phenotypic rates. Previously uncharacterized genes are those identified only with CG numbers in genome release 5.7. Flies were raised on standard media at 25 °C unless otherwise stated. Note that the numbers of lines re-tested in each secondary assay do not always match those identified in the primary screen because certain lines have been replaced after the completion of the RNAi library. To generate *phyllopod*>> EGFP::Pon.LD flies, using standard cloning techniques, the nuclear GFP and polyA cassette from pStinger-phy1^{3,4}-GFP (a gift from C.-T. Chien³⁰) was removed, an SV40 polyA cassette (PCR amplified from the pUAST vector) was added, and the vector was converted to an *attR*-site-containing Gateway Destination vector (pStinger-phy1^{3,4}-Gateway) where the first 14 amino acids plus gateway linkers are fused in frame to the coding sequence of interest after an LR clonase (Invitrogen) reaction to generate an expression clone. EGFP::Pon.LD⁴⁹ was then PCR amplified with primers that add Gateway 5' and 3' *attB* recombination sites and three 3' in-frame STOP codons from the template pBS-TATA-EGFP::Pon.LD-SV40 (gift from F. Wirtz-Peitz), and inserted into pDONR221 (Invitrogen) using a BP Clonase (Invitrogen) recombination reaction. An LR Clonase (Invitrogen) reaction was used to insert EGFP::Pon.LD from the entry clone into

pStinger-phy1^{3,4}-Gateway. The final vector was confirmed by sequencing before injection into *w¹¹¹⁸* embryos for the production of transgenic flies.

Second RNAi lines

We generated or obtained from the NIG-FLY stock centre second UAS-IR fly lines for 73 genes that showed a phenotype in the primary bristle screen. The second line collection of transgenic fly stocks were independently created and designed in such a way that there was a maximum of 70% overlap between the primary construct in the original library and the new IR sequence. Most IR sequences overlap the original construct by less than 50%. Primer pairs for second custom hairpin constructs were designed as previously published⁸, with the following additional constraints/exceptions. Two hundred primer pairs were designed for each target using Primer3 (ref. ⁵⁰). Only those that were predicted to have a single primary target, an s19 score ≥ 0.9 and a maximum CAN count of five were chosen. Restriction sites of the preferred enzymes EcoRI and XbaI were avoided; in rare cases BglII was used as an alternative. Genomic DNA as template was preferred over cDNA. In case of no suitable primers, PCR conditions submitted to Primer3 were loosened to a product size range of 120 to 600 bp, melting temperature of 62 to 72 °C and higher maximum allowed self complementarity. Detailed primer information is provided in Supplementary Table 10. The UAS-IR constructs were cloned using published methods⁸ into a custom designed vector, pKC26, and then inserted into a second chromosome site (43.4) using a customized ϕ 31 system⁵¹.

RNAi bristle screen

The pre-screen was carried out using 40 UAS-IR lines⁸ targeting 21 genes known to be involved in asymmetric cell division and the Notch pathway using *pnr*-GAL4, *sca*-GAL4 and *fzIII*-GAL4 at 18, 25 and 29 °C. The GAL4/UAS system is temperature-dependent, and higher temperatures result in stronger expression but also increase the chances of lethality. Phenotypic assessment was basically the same as that in the genome-wide RNAi bristle screen (see below).

The genome-wide screen was carried out using a collection of transgenic fly lines carrying UAS-IR constructs⁸. To drive expression of IRs for each gene on the notum, males from each UAS-IR fly line were crossed with *pnr*-GAL4 virgin females. Virgin females were obtained in large numbers using a 'virginizer' line where the *pnr*-GAL4 stock also contained an *hs-hid* transgene (originally from R. Lehmann) on the Y chromosome (*w¹¹¹⁸/Y, hs-hid, pnr-GAL4/TM3, Sb*). Stocks were maintained at 22 °C. Once third instar larvae were observed in the stock bottles, the parents were discarded and the bottles heat-shocked at 37 °C in a water bath for one hour on two consecutive days to kill progeny carrying the *hs-hid* chromosome (males). A similar strategy was used to obtain appropriate GAL4 virgin females for other assays in this study. Mating to UAS-IR males and egg laying were carried out at 22 °C for two days, then the crosses were shifted to 29 °C to enhance the expression and activity of GAL4. About 14 days after mating, 10 males (or females if not enough males of the correct genotype were available, and in the case of X chromosome UAS-IR insertion lines) were screened for phenotypes on the notum. The number of flies screened for each line was sometimes less than 10 when there was associated lethality. We judged each phenotype under a normal stereo microscope (Stemi 2000, Zeiss) equipped with a digital camera (PowerShot G6, Canon), and if there was any phenotype took several representative pictures for each transgenic line. Each phenotype was scored in a semiquantitative manner based on the affected area within the *pnr*-GAL4 expression zone, ranging from 0 to 10 (0 means no phenotype; 10 means 90 to 100% of the area was affected by the given phenotype). Areas of the notum where *pnr*-GAL4 was not expressed served as internal

controls for each fly. Phenotypic classes are defined as: ‘gain’ for any number of extra external sensory organs, ‘loss’ for missing external sensory organs, ‘empty or multiple sockets’ for external sensory organs consisting of one or more sockets but no hair, ‘hair cell duplication’ for duplication of hairs, with or without concomitant duplication of the socket, ‘bristle morphology defects’ for other morphological abnormalities of the external sensory-organ hair, ‘planar polarity defects’ for deviations from the posterior orientation of external sensory organs, ‘notum malformation death’ for loss or reduction of the *pnr*-GAL4-expressing region, ‘notum malformation migration’ for thorax closure defects where the two heminota remain separated, ‘overproliferation’ for enlargement of the *pnr*-GAL4-expression area and ‘colour’ for any colour difference between the central, *pnr*-GAL4-expression area and lateral regions. Phenotypes that do not fall into any of these categories were recorded as ‘other’. Adult fly cuticles were prepared using standard methods and mounted in Hoyer’s media. Cuticle preparation images were processed from multiple image stacks using Helicon Focus software (Helicon Soft). All phenotypic information, including stereo microscope and cuticle preparation images, was stored in a custom database. An online database with all results from the primary screen, including images for each phenotype, is available at <http://bristlescreen.imba.oeaw.ac.at>.

Wing RNAi screen and Su(H) reporter assay

Males from 226 UAS-IR lines with a ‘lateral inhibition’ external sensory organ phenotype were crossed with *MS1096*-GAL4 virgin females. At least ten wings from ten different adult flies (preferably male, but also female when necessary) were checked under a normal stereo microscope (Stemi 2000, Zeiss). If any phenotype or abnormality was found, the wings were dissected and mounted on a slide, and were observed under a histology microscope to investigate detailed phenotypes. Definitions for phenotypes recorded in the *MS1096*-GAL4 secondary screen are as follows: notched margin (invagination of the wing margin, excluding invaginations near ectopic vein material), wider vein (wider delta-like-vein phenotypes, considered weak if only wider near the wing margin), bent up (wings bent dorsally), bent up or lethal (wings bent dorsally or lethal (could not distinguish between CyO and non-CyO flies in the same vial)), bent down (wings bent ventrally), wrinkled (general wrinkled structure/appearance), furrow (wrinkled proximal-distal ‘furrows’ but not generally wrinkled), wrinkled margin (wrinkled wing margins, but not generally wrinkled), blistered wings (bubbles between the epithelial layers), necrotic (brown patches of necrotic-looking tissue), tufts of trichomes (multiple wing hairs arranged in clusters), supernumary trichomes (general increase in density of trichomes), ectopic bristles (ectopic structures on the wing blade resembling wing margin bristles), and size (grossly smaller or larger than control wings (subjective, not quantified)). For the lacZ reporter assay, males from UAS-IR lines that showed wing notching or delta-like vein phenotypes in the wing RNAi screen were crossed with *ptc*-GAL4; *Gbe*+*Su(H)*-lacZ/TM6b virgin females. Wing discs from at least five third instar (non balancer) larvae were dissected and stained with 5-bromo-4-chloro-3-indolyl- β -D-galactoside (X-gal) solution using standard protocols and imaged by light microscopy.

Antibodies

The antibodies used in this study are as follows: mouse anti-cut (2B10, 1:500, DSHB), rat anti-Su(H) (1:100, made after ref. ⁵²), mouse anti-Prospero (MR1A, 1:10, DSHB), rabbit anti-Prospero (1:1,000; ref. ⁵³), mouse anti-Elav (9F8A9, 1:30, DSHB) and rat anti-Elav (7E8A10, 1:100, DSHB).

GO term enrichment

Genes with different single or combinations of phenotypes in the primary screen were grouped together and checked for significant enrichment of associated GO terms. Data were analysed using Gostat (T. Beissbarth, <http://gostat.wehi.edu.au/cgi-bin/goStat.pl>), searching against the Flybase database of GO and corrected for multiple testing using the False Discovery Rate (Benjamini correct method).

Orthology prediction

All orthology predictions for candidate genes were made using InParanoid (<http://inparanoid.sbc.su.se/cgi-bin/index.cgi>)⁵⁴.

Interaction map

Publicly available interaction data were integrated to obtain a unified and more complete *Drosophila* interactome network with which to work. The data resources used were DroID (www.droidb.org) (version 2 and version 3), STRING⁴⁴ (version 7.0 and version 7.1) and BioGRID⁴⁵ (version 2.0.40). The inclusion of redundant information from different data sets was avoided by using the PubMed identifying number, where available, as a unique tag to describe the interaction between two entities. Genetic and interolog links were only imported from one data source per entity pair. STRING text mining data were only included if there was at least one other evidence type to support the suggested interaction. Finally, interactions were included into further consideration only if both interactors belonged to the set of 'lateral inhibition' or 'asymmetric cell fate' or were contained in a list of previously characterized Notch regulators. The resulting network was drawn using Cytoscape⁴⁶ (<http://www.cytoscape.org>). Socket and duplication phenotypes are dominant over loss and gain phenotypes for determining node shape. The MCODE⁴⁷ algorithm was used to identify subnetwork complexes using default parameters (loops excluded, haircut selected, fluff not selected, K-Core value 2 and maximum depth 100), except for 'node score cutoff', which was changed to 0.5. Detailed files for import into Cytoscape are available (<http://bristlescreen.imba.oeaw.ac.at>).

References

1. Kiger AA, et al. A functional genomic analysis of cell morphology using RNA interference. *J. Biol.* 2003; 2:27. [PubMed: 14527345]
2. Boutros M, et al. Genome-wide RNAi analysis of growth and viability in *Drosophila* cells. *Science.* 2004; 303:832–835. [PubMed: 14764878]
3. Bettencourt-Dias M, et al. Genome-wide survey of protein kinases required for cell cycle progression. *Nature.* 2004; 432:980–987. [PubMed: 15616552]
4. Goshima G, et al. Genes required for mitotic spindle assembly in *Drosophila* S2 cells. *Science.* 2007; 316:417–421. [PubMed: 17412918]
5. Kamath RS, et al. Systematic functional analysis of the *Caenorhabditis elegans* genome using RNAi. *Nature.* 2003; 421:231–237. [PubMed: 12529635]
6. Gonczy P, et al. Functional genomic analysis of cell division in *C. elegans* using RNAi of genes on chromosome III. *Nature.* 2000; 408:331–336. [PubMed: 11099034]
7. Reddien PW, Bermange AL, Murfitt KJ, Jennings JR, Sanchez Alvarado A. Identification of genes needed for regeneration, stem cell function, and tissue homeostasis by systematic gene perturbation in planaria. *Dev. Cell.* 2005; 8:635–649. [PubMed: 15866156]
8. Dietzl G, et al. A genome-wide transgenic RNAi library for conditional gene inactivation in *Drosophila*. *Nature.* 2007; 448:151–156. [PubMed: 17625558]

9. Artavanis-Tsakonas S, Rand MD, Lake RJ. Notch signaling: cell fate control and signal integration in development. *Science*. 1999; 284:770–776. [PubMed: 10221902]
10. Schweisguth F. Notch signaling activity. *Curr. Biol*. 2004; 14:R129–R138. [PubMed: 14986688]
11. Bang AG, Hartenstein V, Posakony JW. Hairless is required for the development of adult sensory organ precursor cells in *Drosophila*. *Development*. 1991; 111:89–104. [PubMed: 2015800]
12. Schweisguth F, Posakony JW. Antagonistic activities of Suppressor of Hairless and Hairless control alternative cell fates in the *Drosophila* adult epidermis. *Development*. 1994; 120:1433–1441. [PubMed: 8050354]
13. Le Borgne R, Bardin A, Schweisguth F. The roles of receptor and ligand endocytosis in regulating Notch signaling. *Development*. 2005; 132:1751–1762. [PubMed: 15790962]
14. Nagel AC, Maier D, Preiss A. Su(H)-independent activity of hairless during mechano-sensory organ formation in *Drosophila*. *Mech. Dev*. 2000; 94:3–12. [PubMed: 10842054]
15. Rhyu MS, Jan LY, Jan YN. Asymmetric distribution of numb protein during division of the sensory organ precursor cell confers distinct fates to daughter cells. *Cell*. 1994; 76:477–491. [PubMed: 8313469]
16. O'Connor-Giles KM, Skeath JB. Numb inhibits membrane localization of sanpodo, a four-pass transmembrane protein, to promote asymmetric divisions in *Drosophila*. *Dev. Cell*. 2003; 5:231–243. [PubMed: 12919675]
17. Hutterer A, Knoblich JA. Numb and α -Adaptin regulate Sanpodo endocytosis to specify cell fate in *Drosophila* external sensory organs. *EMBO Rep*. 2005; 6:836–842. [PubMed: 16113648]
18. Brand AH, Perrimon N. Targeted gene expression as a means of altering cell fates and generating dominant phenotypes. *Development*. 1993; 118:401–415. [PubMed: 8223268]
19. Calleja M, et al. Generation of medial and lateral dorsal body domains by the pannier gene of *Drosophila*. *Development*. 2000; 127:3971–3980. [PubMed: 10952895]
20. Cavodeassi F, Rodríguez I, Modolell J. Dpp signalling is a key effector of the wing-body wall subdivision of the *Drosophila* mesothorax. *Development*. 2002; 129:3815–3823. [PubMed: 12135920]
21. Wilson RJ, Goodman JL, Strelets VB. FlyBase: integration and improvements to query tools. *Nucleic Acids Res*. 2008; 36:D588–D593. [PubMed: 18160408]
22. Kulkarni MM, et al. Evidence of off-target effects associated with long dsRNAs in *Drosophila melanogaster* cell-based assays. *Nature Methods*. 2006; 3:833–838. [PubMed: 16964256]
23. Hartenstein V, Posakony JW. A dual function of the Notch gene in *Drosophila* sensillum development. *Dev. Biol*. 1990; 142:13–30. [PubMed: 2227090]
24. Blair SS. Wing vein patterning in *Drosophila* and the analysis of intercellular signaling. *Annu. Rev. Cell Dev. Biol*. 2007; 23:293–319. [PubMed: 17506700]
25. Furriols M, Bray S. A model Notch response element detects Suppressor of Hairless-dependent molecular switch. *Curr. Biol*. 2001; 11:60–64. [PubMed: 11166182]
26. Dove SK, et al. Vac14 controls PtdIns(3,5)P(2) synthesis and Fab1-dependent protein trafficking to the multivesicular body. *Curr. Biol*. 2002; 12:885–893. [PubMed: 12062051]
27. Rusten TE, et al. Fab1 phosphatidylinositol 3-phosphate 5-kinase controls trafficking but not silencing of endocytosed receptors. *Mol. Biol. Cell*. 2006; 17:3989–4001. [PubMed: 16837550]
28. Zhang Y, et al. Loss of Vac14, a regulator of the signaling lipid phosphatidylinositol 3,5-bisphosphate, results in neurodegeneration in mice. *Proc. Natl Acad. Sci. USA*. 2007; 104:17518–17523. [PubMed: 17956977]
29. Koo EH, Kopan R. Potential role of presenilin-regulated signaling pathways in sporadic neurodegeneration. *Nature Med*. 2004; 10(Suppl):S26–S33. [PubMed: 15272268]
30. Pi H, Huang SK, Tang CY, Sun YH, Chien CT. phyllopod is a target gene of proneural proteins in *Drosophila* external sensory organ development. *Proc. Natl Acad. Sci. USA*. 2004; 101:8378–8383. [PubMed: 15148389]
31. Milan M, Perez L, Cohen SM. Boundary formation in the *Drosophila* wing: functional dissection of Capricious and Tartan. *Dev. Dyn*. 2005; 233:804–810. [PubMed: 15830355]

32. Milan M, Weihe U, Perez L, Cohen SM. The LRR proteins capricious and Tartan mediate cell interactions during DV boundary formation in the *Drosophila* wing. *Cell*. 2001; 106:785–794. [PubMed: 11572783]
33. Shinza-Kameda M, Takasu E, Sakurai K, Hayashi S, Nose A. Regulation of layer-specific targeting by reciprocal expression of a cell adhesion molecule, capricious. *Neuron*. 2006; 49:205–213. [PubMed: 16423695]
34. Giot L, et al. A protein interaction map of *Drosophila melanogaster*. *Science*. 2003; 302:1727–1736. [PubMed: 14605208]
35. Gavin AC, et al. Functional organization of the yeast proteome by systematic analysis of protein complexes. *Nature*. 2002; 415:141–147. [PubMed: 11805826]
36. Gavin AC, et al. Proteome survey reveals modularity of the yeast cell machinery. *Nature*. 2006; 440:631–636. [PubMed: 16429126]
37. Crouner D, Le Gall M, Gates MA, Giniger E. Notch steers *Drosophila* ISNb motor axons by regulating the Abl signaling pathway. *Curr. Biol*. 2003; 13:967–972. [PubMed: 12781136]
38. Tekotte H, et al. Dcas is required for importin- α 3 nuclear export and mechano-sensory organ cell fate specification in *Drosophila*. *Dev. Biol*. 2002; 244:396–406. [PubMed: 11944946]
39. Meshorer E, Gruenbaum Y. Gone with the Wnt/Notch: stem cells in laminopathies, progeria, and aging. *J. Cell Biol*. 2008; 181:9–13. [PubMed: 18378774]
40. Bosu DR, Kipreos ET. Cullin-RING ubiquitin ligases: global regulation and activation cycles. *Cell Div*. 2008; 3:7. [PubMed: 18282298]
41. Mistry H, Wilson BA, Roberts IJ, O’Kane CJ, Skeath JB. Cullin-3 regulates pattern formation, external sensory organ development and cell survival during *Drosophila* development. *Mech. Dev*. 2004; 121:1495–1507. [PubMed: 15511641]
42. Wu JT, Lin HC, Hu YC, Chien CT. Neddylation and deneddylation regulate Cul1 and Cul3 protein accumulation. *Nature Cell Biol*. 2005; 7:1014–1020. [PubMed: 16127432]
43. Bellaiche Y, Gho M, Kaltschmidt JA, Brand AH, Schweisguth F. Frizzled regulates localization of cell-fate determinants and mitotic spindle rotation during asymmetric cell division. *Nature Cell Biol*. 2001; 3:50–57. [PubMed: 11146626]
44. von Mering C, et al. STRING 7—recent developments in the integration and prediction of protein interactions. *Nucleic Acids Res*. 2007; 35:D358–D362. [PubMed: 17098935]
45. Breitkreutz BJ, et al. The BioGRID Interaction Database: 2008 update. *Nucleic Acids Res*. 2008; 36:D637–D640. [PubMed: 18000002]
46. Shannon P, et al. Cytoscape: a software environment for integrated models of biomolecular interaction networks. *Genome Res*. 2003; 13:2498–2504. [PubMed: 14597658]
47. Bader GD, Hogue CW. An automated method for finding molecular complexes in large protein interaction networks. *BMC Bioinformatics*. 2003; 4:2. [PubMed: 12525261]

References

48. Gallagher CM, Knoblich JA. The conserved c2 domain protein lethal (2) giant discs regulates protein trafficking in *Drosophila*. *Dev. Cell*. 2006; 11:641–653. [PubMed: 17084357]
49. Lu B, Ackerman L, Jan LY, Jan YN. Modes of protein movement that lead to the asymmetric localization of partner of Numb during *Drosophila* neuroblast division. *Mol. Cell*. 1999; 4:883–891. [PubMed: 10635314]
50. Rozen S, Skaletsky H. Primer3 on the WWW for general users and for biologist programmers. *Methods Mol. Biol*. 2000; 132:365–386. [PubMed: 10547847]
51. Groth AC, Fish M, Nusse R, Calos MP. Construction of transgenic *Drosophila* by using the site-specific integrase from phage phiC31. *Genetics*. 2004; 166:1775–1782. [PubMed: 15126397]
52. Gho M, Lecourtois M, Geraud G, Posakony JW, Schweisguth F. Subcellular localization of Suppressor of Hairless in *Drosophila* sense organ cells during Notch signalling. *Development*. 1996; 122:1673–1682. [PubMed: 8674407]
53. Vaessin H, et al. *prospero* is expressed in neuronal precursors and encodes a nuclear protein that is involved in the control of axonal outgrowth in *Drosophila*. *Cell*. 1991; 67:941–953. [PubMed: 1720353]

54. Remm M, Storm CE, Sonnhammer EL. Automatic clustering of orthologs and in-paralogs from pairwise species comparisons. *J. Mol. Biol.* 2001; 314:1041–1052. [PubMed: 11743721]

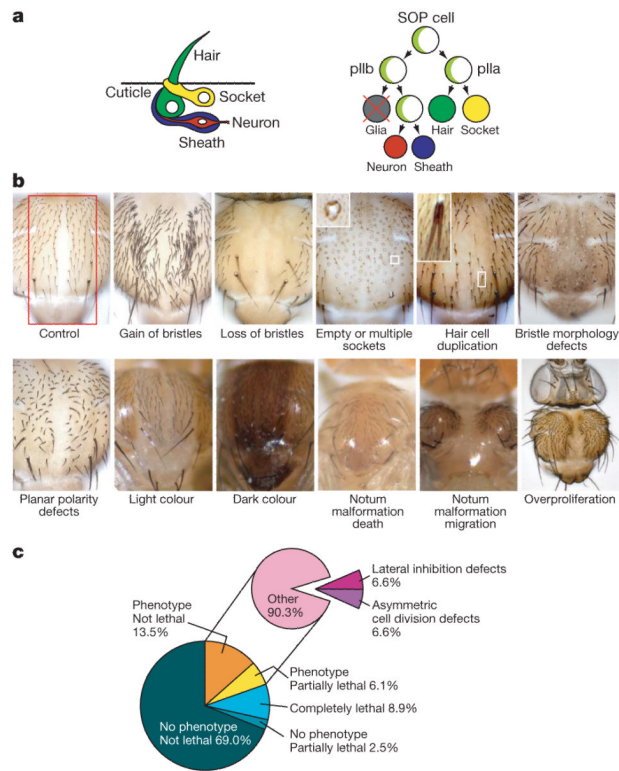


Figure 1. Phenotypic categories and screen results

a, Schematic view of external sensory organ morphology and lineage (Numb and Neuralized, light green). **b**, Typical examples of phenotypic categories. The *pannier*-GAL4 expression area is indicated (red) in the control panel. Examples are *Delta* (gain of bristles), *O-fut1* (loss of bristles), *numb* (empty or multiple sockets), *bazooka* (hair cell duplication), CG31961 (bristle morphology defects), *starry night* (planar polarity defects), CG18095 (light colour), CG34331 (dark colour), CG5462 (notum malformation death), CG7904 (notum malformation migration) and *mob as tumor suppressor* (overproliferation). Original magnification, $\times 10$ (except overproliferation, $\times 5$). **c**, Relative frequency of phenotypes in the entire primary genome-wide screen (left) and among visible phenotypes (right).

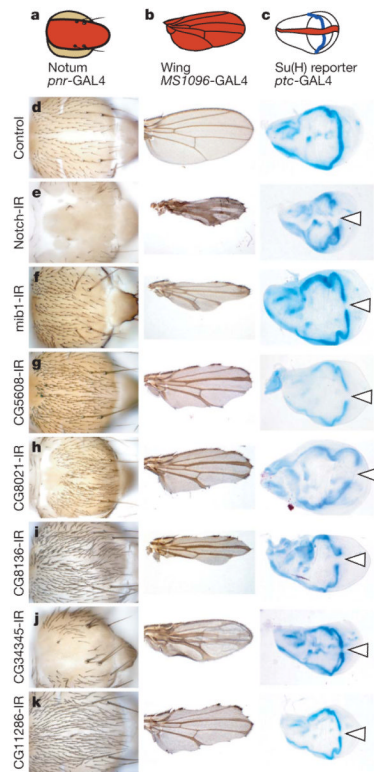


Figure 2. Secondary assays for general components of Notch signalling

Genes with a Notch-like phenotype on the notum were re-screened for wing morphology or expression of a transcriptional Notch reporter. **a–c**, Schematics with GAL4 expression domains shown in red. **d–k**, Appropriate controls and new genes scoring positive in all assays are shown. Transformant IDs (TIDs) for each UAS-RNAi line are as follows: 1112 (*Notch*), 27526 (*mib1*), 45569 (CG5608), 23675 (CG8021), 23748 (CG8136), 20105 (CG34345) and 17541 (CG11286). Arrowheads mark loss of β -galactosidase staining. IR, inverted repeat. Original magnification, $\times 10$ (nota, left column), $\times 2.5$ (wings, middle column) and $\times 5$ (wing discs, right column).

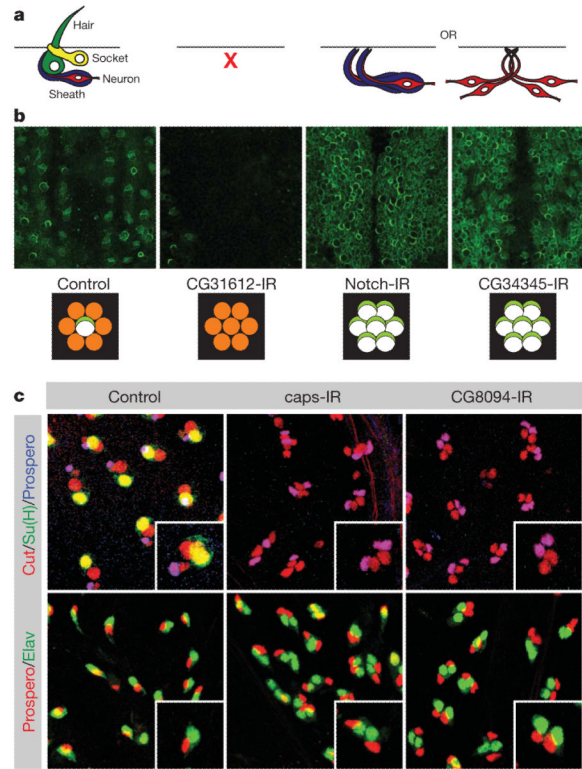


Figure 3. Secondary assays for asymmetric cell division genes

a, Loss of bristle phenotypes can be caused by defects in SOP specification or survival, or by cell fate transformations. **b**, Decreases (CG31612) and increases (*Notch*, CG34345) in SOP cell number were observed. Schematics below each image show interpretations for the number of SOP cells (white with light green crescent) formed from proneural clusters (orange). **c**, Antibody lineage staining identified cell fate transformation phenotypes. All SOP daughter cells are marked by Cut, socket cells by Su(H), sheath cells by Prospero and neurons by Elav. TIDs are 27228 (*Notch*), 20105 (CG34345), 30545 (CG31612), 3046 (*caps*) and 35338 (CG8094). Original magnification, $\times 40$.

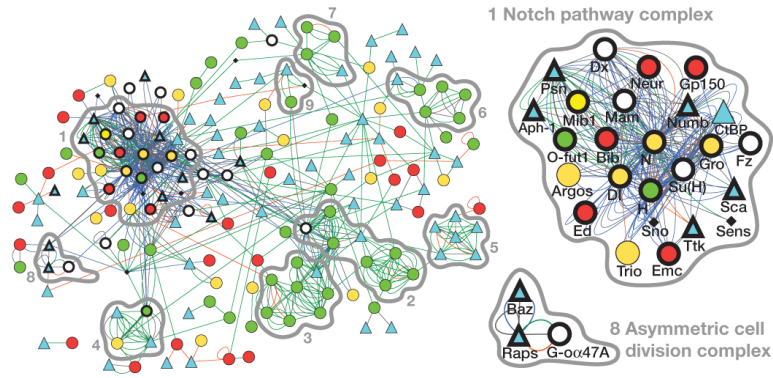


Figure 4. An interaction network for Notch signalling

'Lateral inhibition' and 'asymmetric cell division' category genes with interactions are displayed as a network map. Genes are shown as nodes (node shape/colour denotes phenotype: gain, red circle; loss, green circle; both gain and loss, yellow circle; socket or hair duplication, blue triangle; completely lethal, black diamond; control genes with no phenotype in gain, loss, socket or duplication categories, white circle; positive control genes, bold outline), and relationships between nodes as edges (edge colour denotes interaction type: genetic, blue; interolog, green; yeast two-hybrid, orange; other, grey). MCODE-identified complexes are outlined in grey and rank is indicated. Enlargements of positive control complexes for the Notch pathway and asymmetric cell division are shown.

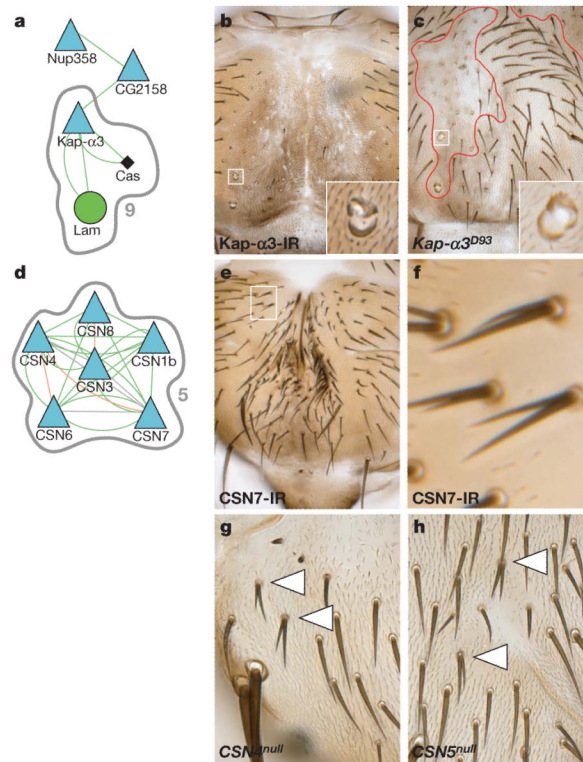


Figure 5. Nuclear import and the COP9 signalosome regulate Notch signalling and asymmetric cell division

a, MCODE complex nine contains Cas, Kap- α 3 and Lam (grey outline). Selected neighbouring nodes from the main interaction network have been included. **b, c**, Both *Kap- α 3* RNAi and *Kap- α 3* mutant clones show multiple socket phenotypes. **d**, MCODE complex five shows nodes for the COP9 signalosome. **e–h**, Knockdown of *CSN7* (**e**, close up in **f**) and both *CSN4* (**g**) and *CSN5* (**h**) mutant clones cause hair cell duplication phenotypes. Original magnification, $\times 10$ (**b, c, e**) and $\times 20$ (**g, h**).