The role of Dichaete in transcriptional regulation during Drosophila embryonic development

Jelena Aleksic1,2,3, Enrico Ferrero1,2, Bettina Fischer1,2, Shi Pei Shen1,4 and Steven Russell1,2*

Abstract

Background: Group B Sox domain transcription factors play conserved roles in the specification and development of the nervous system in higher metazoans. However, we know comparatively little about how these transcription factors regulate gene expression, and the analysis of Sox gene function in vertebrates is confounded by functional compensation between three closely related family members. In Drosophila, only two group B genes, Dichaete and SoxN, have been shown to function during embryonic CNS development, providing a simpler system for understanding the functions of this important class of regulators.

Results: Using a combination of transcriptional profiling and genome-wide binding analysis we conservatively identify over 1000 high confidence direct Dichaete target genes in the Drosophila genome. We show that Dichaete plays key roles in CNS development, regulating aspects of the temporal transcription factor sequence that confer neuroblast identity. Dichaete also shows a complex interaction with Prospero in the pathway controlling the switch from stem cell self-renewal to neural differentiation. Dichaete potentially regulates many more genes in the Drosophila genome and was found to be associated with over 2000 mapped regulatory elements.

Conclusions: Our analysis suggests that Dichaete acts as a transcriptional hub, controlling multiple regulatory pathways during CNS development. These include a set of core CNS expressed genes that are also bound by the related Sox2 gene during mammalian CNS development. Furthermore, we identify Dichaete as one of the transcription factors involved in the neural stem cell transcriptional network, with evidence supporting the view that Dichaete is involved in controlling the temporal series of divisions regulating neuroblast identity.

Keywords: Sox, Drosophila, Dichaete, Prospero, Nervous system, Neural stem cells

Background

Sox-box transcription factors play critical roles in the embryonic development of all metazoans in which they have been characterised [1]. The group B subgroup is of particular interest in vertebrate biology since it encodes a set of transcriptional regulators known to play critical roles in neural development and in the maintenance of embryonic stem cells [2-5]. In most vertebrates there are 5 group B genes: the group B1 genes Sox1, 2 and 3, which primarily act as transcriptional activators, and the group B2 genes Sox14 and 21, which act mainly as transcriptional repressors and appear to antagonise group B1 functions [6,7].

In Drosophila there are four group B genes; SoxNeuro (SoxN), Dichaete, Sox21a and Sox21b, the latter three clustered in the genome in a complex found in all insect genomes so far sequenced [8-10]. While there is a lack of clarity regarding the subdivision of group B proteins in insects [9,11], it is clear that, as with vertebrate group B1 proteins, both Dichaete and SoxN play critical roles in the early development of the embryonic nervous system. The functions of Sox21a and Sox21b are currently not known. However, neither gene is expressed in the embryonic CNS until very late in development, suggesting that in the fly, only two group B genes are involved in early CNS development. Functional studies using mouse Sox B1 genes to rescue fly mutant phenotypes indicate that the mammalian proteins are functionally related to both Dichaete and SoxN [12,13]. Sox2 in particular was able to rescue Dichaete functions, suggesting a high degree of functional conservation. The study of...
Dichaete may therefore provide useful insights into aspects of Sox2 function in the more complex vertebrate CNS.

*Dichaete* is expressed dynamically during embryogenesis where it is known to have functions in early segmentation, CNS development, hindgut morphogenesis and cuticle patterning [14-17]. In the CNS, *Dichaete* is expressed early in the ventral midline, the fly equivalent of the vertebrate floor plate, as well as the medial and intermediate columns of the neuroectoderm from the earliest stages of CNS specification. *SoxN* is also expressed in the neuroectoderm, in this case extending more laterally to include the lateral column, and is not expressed in the midline until the latter half of embryogenesis [18]. Thus, as is the case with group B genes in vertebrates, both Dichaete and SoxN are co-expressed in many of the cells of the early CNS and they exhibit extensive functional compensation with double mutants exhibiting severe neural hypoplasia in contrast to the comparatively mild phenotypes of single mutants [19,20]. Targeted expression of dominant negative forms of Dichaete or its mammalian orthologue Sox2 also show strong CNS phenotypes, further demonstrating that group B function is critical for normal CNS development [21].

At the molecular level, both Dichaete and SoxN are required for the correct expression of pro-neural genes in the Achaete-scute Complex, particularly *achaete (ac)* and *asense (ase)*, where they interact with the homeodomain proteins encoded by *intermediate neuroblasts defective (ind)* and *ventral neuroblasts defective (vnd)* [19,21-23]. Group B Sox proteins in the vertebrate CNS are also coexpressed with related homeodomain proteins along the ventro-medial axis of the developing neural tube, hinting at a deep conservation of Sox function in the CNS [24]. A role for Dichaete in the control of midline expression of the *slit* gene has been well characterised, with both genetic and molecular evidence demonstrating a critical interaction between Dichaete and the POU-domain protein Ventral veins lacking (Vvl) [13,25]. Mammalian Sox2 interacts with a related POU protein, Oct4, in the regulatory network controlling embryonic stem cell pluripotency and in primary neurogenesis [26-30]. Mouse Sox2 can also interact with Vvl in the fly, reinforcing the idea that Dichaete and Sox2 are functionally related [13].

More recently, a genomic analysis of embryos expressing dominant negative forms of Dichaete in combination with Dichaete binding data identified hundreds of potential target genes with annotated CNS functions, suggesting the *Drosophila* group B Sox proteins have widespread roles in CNS development [21]. Other studies have identified roles for Dichaete and SoxN in the regulation of *shavenbaby* in the epidermis, in part by repressing Wnt signalling through an interaction with the related HMG-domain protein Left1/Tcf [17,31]. Finally, more recent work in the embryonic and larval CNS has shown that Dichaete plays a role in the transcriptional hierarchy that controls the temporal specification of neuroblast fate, in particular controlling the decisions to terminate the self-renewal program and differentiate or die [32]. This bears a striking resemblance to the role group B1 Sox proteins play in the vertebrate neuroepithelium, where their down-regulation is required for cell-cycle exit and subsequent neural differentiation [33].

In this study, we use genomics approaches, including DamID and transcriptional profiling, to gain insight into the role of Dichaete during embryonic CNS development. We also draw on previously published embryonic binding data generated by BDTNP [34] and the modENCODE project [35] to perform an integrative analysis of Dichaete function. We find that potential target genes are mainly downregulated in a Dichaete mutant, suggesting it may function largely as a transcriptional activator, and that Dichaete is one of the key factors in the *Drosophila* neural stem cell transcriptional network interacting with a temporal series of transcription factors regulating neuroblast fate. We also find evidence of an interaction with Prospero, which is apparently antagonistic in neuroblasts, but synergistic in the Ganglion Mother Cells. Together our findings support the view that Dichaete functionally acts as a group B1 Sox factor and further emphasises the striking conservation in Sox function in the CNS.

**Results**

**Generating a core set of Dichaete binding intervals**

Dichaete is important for a number of key developmental processes, including embryonic segmentation, hindgut morphogenesis and nervous system development. However, a thorough genomic analysis of the role of Dichaete during early *Drosophila* embryogenesis aimed at identifying target genes and regulatory networks has not been performed to date. Genome-wide binding data from blastoderm embryos is available from the Berkeley Drosophila Transcriptional Network Project [34] and data from later stages of embryogenesis was generated as part of the modENCODE project [35]. Our preliminary analysis indicated some discordance between these datasets. While some differences are to be expected due to different developmental stages used, the modENCODE data appeared to have weaker signal, and identified considerably fewer peaks than the DBTNP data. Since it was unclear whether this was due to false positives in the BDTNP data or false negatives in the modENCODE data, we elected to generate an independent genome-wide Dichaete binding dataset to identify high-confidence binding intervals supported by different analytical methods.

We used the DamID method [36] to generate Dichaete-Dam and control Dam profiles from three independent
biological replicates of stage 5–11 embryos. After normalisation and peak-calling we found that Dichaete, in common with other transcription factors, is associated with thousands of genomic regions in the embryo across development. We identified 6067 binding intervals at 1% FDR and over 16,000 at 25% FDR. We assessed enrichment for known transcription factor binding motifs within the 1% FDR binding intervals using i-cisTarget [37] and identified a Vielfältig motif (Vfl; E-score = 6.2) [38] followed by a series of Sox binding motifs (E-scores from 4.5 to 5.1), suggesting these DamID data identify bona fide Dichaete genomic occupancy. Assigning the 1% FDR binding intervals to genes, we found Dichaete associated with 3561 genes (Additional file 1: Table S1). This Dichaete-bound set of genes is significantly enriched for general developmental and regulation terms (p < 1E-50), and also more specifically for nervous system development (581 genes, p = 9.2E-31), neuron generation (374 genes, p = 2.9E-42) and neuron differentiation (336 genes, p = 5.9E-35). We also found enrichment for genes involved in brain development (69 genes, p = 1.6E-7), hindgut morphogenesis (39 genes, p = 2.6E-4) and segmentation (122 genes, p = 1.4E-10), all processes Dichaete is known to be involved in (the full set of enrichments is given in Additional file 2: Table S2).

The published ChIP datasets also identified large numbers of binding intervals and associated genes (BDTNP = 6452 binding intervals and modENCODE = 3520 binding intervals at FDR1%, mapping to approximately 6500 and 3500 genes, respectively). To focus on a set of high confidence Dichaete target genes and eliminate technique specific artefacts as far as possible, we combined our DamID dataset with the ChIP datasets to generate a set of core binding intervals. We identified intervals supported by at least one low stringency (25% FDR) and one high stringency (1% FDR) dataset from the independent experimental techniques; for example, all regions confirmed with 1% FDR DamID and 25% FDR ChIP were included, and vice versa (Figure 1A; Additional file 3: Table S3). Thus each of the intervals in the core dataset was supported by high stringency evidence in at least one dataset, and was also independently validated using an alternative experimental technique. These binding intervals were used for further analysis. While we recognise that this is a relatively stringent filter and will not include many bona fide Dichaete binding intervals, we contend that focusing on independently verified binding locations will allow us to make more confident inferences about Dichaete function in the embryo.

This amalgamation process generated a dataset of 6720 intervals associated with 4279 Drosophila genes (Additional file 4: Table S4), which were examined for general properties using the FlyMine data warehouse [39]. We found that almost half these genes (1925) are expressed in the larval CNS, an encouraging observation given the known role of Dichaete in larval neuroblasts [32]. We also found even stronger enrichment for genes associated with nervous system development (775 genes, p = 1.4E-67), neuron generation (461 genes, p = 2.1E-60), brain development (91 genes, p = 2.4E-17) and other nervous system terms (Additional file 5: Table S5). A large number of the Dichaete-bound genes have a role in the regulation of transcription (451 genes, 1.3E-47), with as many as two thirds (205 out of 294) of the FlyTF curated ‘trusted’ transcription factors featured in the gene list [40]. We also noticed a very strong enrichment for Dichaete binding at genes encoding specific transcription factor classes, including homeodomain proteins (99 homeodomain or homeodomain-like proteins, 3.0E-16); C2H2-type Zinc finger domain proteins (139 genes, 4.8E-5) and, to a lesser extent, fork head domain proteins (15 genes, 1.2E-2). Taken together, these data support the idea that Dichaete is a key developmental regulator controlling a battery of transcription factor genes involved in nervous system development and a variety of other developmental processes.

We examined the overall genomic distribution of the Dichaete core binding intervals, using the centre coordinate of each interval as an approximation of the Dichaete binding site. We found that two thirds of Dichaete binding (67%) maps to genic regions and of these, the majority (65%) are in introns. We also found an association between Dichaete binding and transcription start sites (TSS), observing a peak of binding intervals within 500 bp of a TSS, indicating a preference for this region (Figure 1B). However, this is a relatively weak predictor of overall Dichaete binding, with slightly less than 25% (1618 out of 6720) of the total Dichaete binding events found close to a TSS. We also found that Dichaete binding showed a significant preference for long introns (p-value < 2.2e-16, Wilcoxon rank sum test). The mean length of Dichaete-bound introns is 12.7 kb, while the mean length of all Drosophila introns excluding those bound by Dichaete is 1.5 kb (the median, which is less sensitive to outliers, still showed this difference: median Dichaete bound = 5.7 kb compared to 130 bp for the rest of the genome). While it is possible this observation is an artefact of larger regions of the genome being more likely to be bound in general, the high level of significance suggests this observation may well reflect an aspect of Dichaete function. Recent work does suggest that long and short introns are functionally distinct [41], with long introns being, for example, more likely to harbour regulatory sequences.

The Dichaete core dataset was also found to overlap with a sizeable proportion of mapped enhancer regions. Of a total of 8975 reported enhancers (1862 from RedFly and 7113 from FlyLight), Dichaete binding was associated with 2400 (27%), including almost half of the
experimentally validated enhancers reported in RedFly (887 out of 1862) [42]. The 887 Dichaete-bound RedFly enhancers are associated with a total of 229 genes, including vvl, encoding a known Dichaete midline cofactor, the neuroectoderm column specifying gene vnd, as well as genes for transcription factors with roles in neuroblast pluripotency and differentiation such as hb, Kr, nub, grh and mira [43,44]. Substantial overlap was found with the enhancers mapped by the FlyLight project [45,46], particularly with those showing expression in neuroblasts (28/30), the midline (114/201 early and 476/1072 late) and in large subsets of the CNS (478/811) (Additional file 6: Table S6). Together these data indicate that Dichaete associates with both core promoter regions and with classical enhancers.

As noted above, our DamID dataset was enriched for Vfl and Sox-related binding motifs. A similar analysis with the BDTNP and modENCODE binding intervals again identified strong enrichment for Vfl motifs (E-scores of 9.4 and 8.1 respectively) and weaker enrichment for Sox-like motifs (E-scores 2.6 and 2.8). In our core dataset the top enriched motifs were Vfl (E-score 9.4), Trithorax-like (Trl, E-score 4.9), Stat92E (E-score 4.8) and Sox motifs (E-score 4.1) (Figure 1C). Using a previously published Dichaete motif from a bacterial 1-hybrid screen for a motif scan [47], we found that 82% of all intervals contain matches to the motif, supporting the view that the core binding intervals represent specific Dichaete binding sites.

Taken together, these observations suggest a role for Dichaete in the transcriptional regulatory network active in the developing CNS, including its previously reported function in the midline, as well as a role in neuroblasts. Binding associated with hundreds of mapped enhancer regions and an association with transcription start sites indicates that Dichaete may display two distinct modes of operation at enhancers and promoters. Alternatively,
this may reflect aspects of higher order chromatin organisation where Dichaete bound enhancers are brought into close proximity to core promoter regions.

**Dichaete may act as a regulatory hub during embryogenesis**

It is known that essential proteins in densely interconnected biological processes such as development are more likely to be genetic network hubs - regulators connected to a large number of network nodes [48,49]. A number of developmental transcription factors fit this role, and between themselves these also tend to be highly interconnected. In the case of Dichaete, the large number of binding regions in the genome suggested a potential role as a hub, supported by the finding that the binding is frequently associated with other transcriptional regulators. To investigate whether Dichaete does indeed appear to have a regulatory role for a large number of targets, we performed a gene expression analysis in *Dichaete* mutant null embryos.

We have previously performed targeted gene expression analysis of group B Sox function in the CNS using dominant negative constructs [21]. To gain a more comprehensive view of Dichaete function during development we performed microarray expression profiling with RNA extracted from *Dichaete* mutants. We isolated RNA from stage 10–11 *Dichaete* null mutant embryos and identified genes with significantly changed expression compared to heterozygous siblings. Remarkably, we found that the majority of genes in the genome appeared to change expression and consequently our usual data normalisation methods were inadequate. We therefore utilised a normalization approach that relied on identifying the minority of invariant genes and using these to normalise the remainder [50]. With moderate stringency statistical thresholds (M-value > 0.5 or < -0.5, p-value < 0.05), 9120 genes were differentially expressed: approximately 65% of the genes in the *Drosophila* genome. In order to focus on a reliable set of differentially expressed genes, we applied a more stringent threshold (M-value > 1 or < -1, p-value < 0.01), identifying 4518 differentially expressed genes, and these were used for further analysis (Additional file 7: Table S7). The vast majority of these (>90%, 4182 genes) were found to be downregulated, with only 336 genes upregulated in the mutant. While some of these gene expression changes are likely to be pleiotropic effects reflecting the early role of Dichaete in segmentation, encouragingly we still found significant enrichment for genes involved in segmentation (p = 7.2E-9) and neurogenesis (252 genes, p = 5.3E-6) and submitting the gene list to the FlyExpress database generated a Genome-wide Expression Map that highlights the developing CNS, supporting the view that *Dichaete* regulates genes in the CNS (Figure 1D) [51]. While there is some overlap between genes involved in segmentation and those involved in neurogenesis, segmentation genes alone do not account for all the neurogenesis genes affected, and pleiotropy is unlikely to explain all the changes we observe. Previous studies have found that regions of the embryo such as the thorax, which are largely unaffected by the segmentation phenotype, still display strong neural phenotypes [13]. Furthermore, at the level of the individual embryo, segmentation defects are highly variable with many individuals showing comparatively weak phenotypes [14,15]. In contrast, we focus here on very strong changes in gene expression that are likely to reflect more widespread consequences of *Dichaete* loss. Although we identify a strong CNS signature in the GO enrichments, the most enriched GO term in these data was for Proteolysis (293 genes, p = 1.2E-10), which may possibly reflect a generic stress response, suggesting that many of the identified genes may change expression as an indirect response to the loss of Dichaete function.

To focus on likely direct target genes we intersected the core binding intervals with the mutant gene expression data. This generated a set of 1373 genes that show significant expression changes in *Dichaete* mutant embryos and are bound by Dichaete in independent binding studies: we refer to these as Dichaete target genes (Additional file 8: Table S8, which for reference also includes previously reported expression in dominant negatives [21]). We emphasise that both the binding and the expression data were analysed with relatively stringent cut-offs and that the true number of Dichaete targets is likely to be higher. The Dichaete target genes are highly enriched for developmental processes (3E-54), and nervous system specific annotations (generation of neurons, p = 2.4E-39; nervous system development, p = 1.4E-32; neuron differentiation, p = 1.8E-33) (Additional file 9: Table S9). Again we found the list of targets was strongly enriched for transcription factors, with approximately half (152) of the 296 genes present in the FlyTF ‘trusted’ list of transcription factors [40] differentially expressed in the Dichaete null mutant. This was reflected in enrichment for gene transcription/expression GO terms (p = 1.3E-34), and for genes encoding Homeodomain (50 genes, p = 5.8E-18), fork head (12 genes p = 2.1E-4) and C2H2 Zinc finger (65 genes, p = 2.1E-4) genes. These high-confidence Dichaete target genes further support the view that Dichaete plays a key role in embryonic development and suggests it may act as a regulatory hub.

**Dichaete in the hindgut**

We have previously shown that Dichaete is required for correct morphogenesis of the embryonic hindgut, where it positively regulates *dpp* and negatively regulates *hh* [16]. We examined the Dichaete binding and expression
data to determine the extent of Dichaete involvement in gut development. We found many of the genes known to be involved in aspects of hindgut development were bound and apparently controlled by Dichaete. Hindgut development is modulated by a cascade of transcription factors and signalling pathways, starting with the activity of the terminal genes \( \text{tll} \) and \( \text{hhb} \), through a highly conserved "gastrulation cassette" (\( \text{cad}, \text{byn}, \text{fkh} \) and \( \text{wg} \)) to a set of patterning and morphogenesis genes (\( \text{bowl}, \text{dpp}, \text{drm}, \text{en}, \text{hh}, \text{lin}, \text{upd} \) and Dichaete itself) [52]. Of 25 genes known to be involved in hindgut development, we found that 17 were strongly downregulated in Dichaete mutant embryos and three showed weak upregulation (Figure 2). While \( \text{hh} \) is repressed by Dichaete in the hindgut, our expression profiling shows downregulation of \( \text{hh} \) in Dichaete mutants, this is likely a reflection of loss of \( \text{hh} \) expression in the CNS. In addition, we observed Dichaete core binding intervals associated with 21 of these genes, including at sites overlapping known hindgut enhancer elements in the \( \text{dpp} \) and \( \text{fkh} \) genes as well as enhancers showing embryonic gut expression in the FlyLight collection (\( \text{dpp}, \text{hh}, \text{wgl}, \text{retn}, \text{crb} \)) [45,46] (Figure 2).

**Known Dichaete targets in the CNS**

Previous work has shown that Dichaete directly regulates genes of the AS-C in the neuroectoderm and \( \text{sli} \) in the midline [21,23,25] and we found all four genes in the complex downregulated at least 2-fold in Dichaete mutants. We identified substantial Dichaete binding across the AS-C and almost all of the core binding intervals in this region overlap with known RedFly enhancers (\( \text{ac}_\text{pg7} \) and \( \text{ase}_\text{F2.0} \)) or regulatory elements identified by the FlyLight project [45,46] (Figure 3A). We also noticed a prominent binding interval 3' to \( l(1)sc \), where no embryonic regulatory regions are reported in the FlyLight set, however there is a VT enhancer region reported here [53].

While Dichaete has been shown to affect the expression of \( \text{ac}, \text{sc} \) and \( \text{ase} \), there have been no reports relating to \( l \)
Dichaete in the neuroectoderm

Dichaete is dynamically expressed in the neuroectoderm, in neuroblasts, GMCs and in differentiating neural cells, and our Dichaete target gene set strongly suggests that it regulates several hundred genes in these lineages. In contrast, Dichaete mutant phenotypes are relatively weak in the ventral nerve cord due to functional compensation by the related group B gene SoxN. To gain insights into Dichaete function in the ventral nerve cord we focused our attentions on some of the core regulatory pathways controlling neural differentiation. In the embryonic and larval CNS, Dichaete has been reported to play a role in regulating the switch from neuroblast self-renewal to cell cycle exit and differentiation or apoptosis [32]. We therefore examined the Dichaete target set to seek evidence for similar activity in the embryonic nervous system. Strikingly, we found extensive binding at all of the genes implicated in the temporal transcription factor cascade (svp, hb, Kr, nub, pdm2 and cas) with most of these downregulated more than 2-fold in Dichaete mutants (hb and nub were downregulated approximately 1.5-fold). There are cross-regulatory interactions between the transcription factors governing neuroblast fate and we therefore compared Dichaete core binding with available data for Hb and Kr. We found extensive overlap between Dichaete core binding

\((1)sc\) in \textit{Dichaete} mutants: while we find that ventral nerve cord \(l(1)sc\) expression is apparently unaffected in \textit{Dichaete}\null mutant embryos, we noticed a loss of expression in specific populations of cells in the deuteocerebral and protocerebral regions of the embryonic brain (Figure 3B-C), [54]). Together these observations strongly suggest that Dichaete interacts with many of the known regulatory elements in the \textit{AS-C} and that at least some of this binding is functionally relevant to embryonic expression of the pro-neural genes in the Complex.

In the case of \textit{sli}, which Dichaete regulates in midline glia, we find some evidence of Dichaete binding, however, most of the intervals are not identified in the core set, presumably because midline glia comprise a very small fraction of the whole embryo. Previous work showed that Dichaete binding can be detected at the midline enhancer with specific PCR assays despite an apparent lack of binding in BDTNP and modENCODE datasets [21,25] and we note that a low stringency DamID peak overlaps the \textit{GMR32A06}\null regulatory region encompassing the \textit{sli} midline enhancer. There is a core Dichaete binding interval within the \(^{1}\text{st}\) \textit{sli} intron, where a regulatory element controlling expression in the posterior gut is reported [55], and \textit{sli} expression is downregulated in Dichaete mutants (2.6 fold). Thus our data support the view that Dichaete more broadly regulates \textit{sli} expression (Figure 3D).

**Figure 3** Dichaete at known target genes. A) Achaete-scute Complex: Dichaete Dam ID binding profiles with 1% and 25% FDR intervals. Dark blue tracks represent FlyLight Enhancers with expression categorised as: small subset of the CNS at stage 16 (st16,CNS_SS), any CNS in germ band extended embryos (GBE) and large subset of the CNS at stage 16 (st16,CNS_LS). Grey bars represent other mapped enhancer elements. Core (red) represents the regions defined as core binding intervals overlapping DamID and ChIP data. The black regions represent gene models and the genome coordinates. B and C) antibody staining revealing \(l(1)sc\) expression in wild type (B) and \(D^{7}/D^{3}\) \(Lgs1-a\) (C) stage 16 embryos. P, D and T refer to the protocerebrum, deutocerebrum and tritocerebrum. Red arrowheads indicate loss of \(l(1)sc\) expression in the duetocerebrum and tritocerebrum. D) \textit{sli} tracks as above with FlyLight enhancers identified as expressed in the midline at stage 16.
intervals and the high stringency (1% FDR) Kr and Hb binding intervals determined by BDTNP at all of the core temporal transcription factors (Figure 4). Although there are no available binding data for Svp, Nub/Pdm2 or Cas, a genetic interaction between Nub/Pdm2 and Dichaete has been reported [56].

The termination of the neuroblast programme is marked by the segregation of Prospero and Miranda to the GMC, the nuclear localisation of Prospero and the decision to differentiate or die. We found extensive Dichaete binding across the pros gene (Figure 5A), and Dichaete core binding associated with mira (Figure 5B) and insc (not shown) among other genes encoding components of the asymmetric cell division machinery. To determine whether Dichaete regulates pros, we examined Pros expression in Dichaete mutants and observed an increase in Pros levels at stage 10, particularly in the medial column of the neuroectoderm (Figure 5D and E), and by stage 11 Pros was seen at high levels in the neuroectoderm of mutant embryos (Figure 5F and G). Whether this reflects an increase in the number of Pros-expressing cells in the embryo or precocious activation of Pros remains to be determined. By stage 16 Pros levels were considerably reduced compared to wild type (not shown). While it is possible that loss of Pros is due to an indirect consequence of Dichaete loss, our data are consistent with a model where Dichaete is involved in repressing pros expression early during neurogenesis but in activating or maintaining pros at later stages. Since the expression results could be confounded by indirect effects or by SoxN functional compensation, we examined Pros expression in embryos expressing dominant negative Sox constructs in the neuroectoderm. Whether using Dichaete constructs deleted for the DNA binding domain (ΔHMG, Figure 5I), Engrailed-repressor fusions (Figure 5J) or mouse Sox2 Engrailed-repressor fusions (Figure 5K and L) we found downregulation of Pros in the neuroectoderm from late stage 10 onwards, supporting the view that Dichaete can directly regulate pros.

Particularly relevant to the decision to exit the cell cycle to differentiate or undergo apoptosis, we found Dichaete binding at W, grim and rpr (Figure 5C), indicative of regulatory input into the apoptotic pathway, as well as key cell cycle regulators such as Cyclin E. Combined with our findings relating to the AS-C and temporal NB program described above, the binding and expression data indicate that Dichaete plays a key role in the entire pathway from neural specification, through determination of NB identity to the generation of differentiated neural cells.

**The role of Dichaete in the neuroblast regulatory network**

Our analysis of various CNS pathways suggested Dichaete involvement in many different processes and to elaborate

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**Figure 4 Temporal neuroblast cascade.** A-F) Dichaete DamID profiles (blue) and core binding intervals (red) along with the 1% FDR binding intervals for Kr (dark blue) and Hb (light blue) from the BDTNP at the 5 genes of the temporal neuroblast cascade and the Dichaete region.
Figure 5 (See legend on next page.)
this further we compared our core Dichaete binding intervals with a range of binding data generated by the BDTNP and modENCODE projects. We selected binding datasets available for 41 different transcription factors encompassing the first 12 hrs of embryogenesis and used a resampling-based method [57] to identify highly significantly overlapping binding profiles. We performed an exhaustive pairwise comparison and identified 14 significant overlaps (Figure 6A, Additional file 10: Table S10). Of these, 8 included overlaps with Twist, a mesoderm-specific TF that also shows significant binding at HOT regions in the Drosophila genome [58]. We found that Dichaete significantly overlapped with Prospero and Twist. The overlap with Pros is in line with our findings presented above. The overlap with Twist, on the other hand, is unlikely to be of specific functional significance, since there is little overlap between Dichaete and Twist expression during embryogenesis. Rather, the significant overlap may be a reflection of the tendency of both transcription factors to bind at HOT regions.

The relationship between Dichaete and Pros was further explored by comparing the Dichaete core binding intervals with Pros-DamID and expression data in more detail [59]. We again found extensive binding overlap, with 902 of 1478 Prospero intervals also Dichaete core intervals. The Prospero/Dichaete overlaps were associated with 704 genes (Additional file 11: Table S11), including 17 genes involved in axon guidance pathways (p = 3.6E-5), as well as a collection of genes involved in the regulation of developmental processes, including cell fate commitment (106 genes, p = 3.1E-39). We also compared gene expression data from Dichaete and pros mutants and found examples of genes with opposite expression changes (down in Dichaete, up in pros) as well as genes with similar expression changes (Figure 6B), suggesting Dichaete and Pros may both collaborate and antagonise. Pros promotes neural differentiation and represses self-renewal via a regulatory network involving Ase, Dpn and Sna [60]. We found that Dichaete binding intervals were associated with almost all of the genes in this network (34/35), with over half (20/35) identified as high confidence Dichaete targets (Figure 7A). We also found that 65% (23/35) showed overlapping Dichaete and Prospero binding, including Prospero itself (Figure 7B).

As we note above, Dichaete and mammalian Sox2 show considerable functional similarity and recently, a genomic study identified over ~1400 Sox2 target genes in neural stem cells [61]. We converted this list of mouse genes to their Drosophila orthologues (1235 genes) and compared these with genes in the core Dichaete binding set and our list of high confidence direct target genes, identifying 523 and 217 genes respectively (Additional file 12: Table S12 and Additional file 13: Table S13). Over half of the 271 conserved targets are highly enriched for regulation-associated GO terms (P < E-30), 35% annotated as being involved in transcriptional regulation.
35% in the generation of neurons (p = 5.7E-30). Remarkably, 13 out of the 34 Dichaete-bound neuroblast regulatory network genes are also Sox2 targets in the mouse neural system, found to be mainly genes connected with Notch signalling (4/6), neuroblast cell fate (3/8) and neuronal differentiation (3/7). This remarkable overlap hints at a deep conservation in the regulatory networks driving the development of neural cells between distantly related species, and points to a key role for group B transcription factors in this process.

Discussion

The *Drosophila* Dichaete gene encodes a group B Sox domain protein previously implicated in diverse developmental pathways, including key aspects of early segmentation and CNS development [14,15]. We have combined genome-wide binding analysis and gene expression profiling to identify a set of over a thousand high confidence Dichaete targets in the fly genome. Our conservative target list reflects genes bound by Dichaete in independent binding assays and that show very significant expression changes in Dichaete mutant embryos. It is likely this represents an underestimate of the true extent of Dichaete activity in the fly genome for several reasons. First, we employed stringent cut-off criteria for both the binding and the gene expression analysis. Second, loss of Dichaete function is frequently compensated by the related Group B protein SoxN, thus many genes showing high confidence binding may not exhibit significant expression changes in Dichaete mutants [19,20]. This is supported by recent work expressing dominant negative forms of Dichaete and its vertebrate orthologue Sox2 in the CNS where we identified over 300 additional target genes [21]. In addition, we have recently analysed the genomics of SoxN in the *Drosophila* embryo and find very substantial overlap between Dichaete and SoxN binding as well as a repertoire of genes uniquely bound by each gene (EF and SR, in preparation). Taken together our analysis indicates that Dichaete directly regulates a substantial fraction of genes in the fly genome, particularly in the CNS, suggesting it may act as a regulatory hub. Consistent with this view, we found that over half of the transcription factors encoded in the fly genome are bound by Dichaete and misregulated in the mutant, indicating that Dichaete is likely to be involved in many of the regulatory networks driving CNS development.

The core Dichaete binding intervals we identified are enriched for Sox binding motifs but we also found significant overrepresentation of binding motifs for Vf, the GAGA-binding factor Trl and the JAK-STAT pathway transcription factor Stat92E. All three of these factors have been identified as key elements in the regulatory
programme that drives the onset of zygotic gene expression in the blastoderm embryo [38,62-64]. Dichaete also plays a key role in early zygotic gene expression, regulating the correct expression of pair rule genes [14,15], and we found overlapping Vf/Dichaete binding at eve, h, and run stripe enhancers. While most of the work on Vf has focused on understanding its function during the maternal to zygotic transition, the gene is expressed more widely after cellularisation, particularly in the CNS [64]. Indeed recent work has shown a specific role for Vf in the CNS midline [65], a tissue where Dichaete is known to be active [13,25] and we found overlapping Vf/Dichaete binding associated with sli and comm, known Dichaete midline targets. Post cellularisation functions for Trl and Stat92E are well established.

More recently, these three factors, particularly Vf and Trl, have been strongly associated with enhancer activity driven by Highly Occupied Target (HOT) regions [58]. HOT regions have been identified in large scale studies of the Drosophila, C. elegans and human genomes, and represent genomic sites where many functionally unrelated transcription factors bind, frequently in the absence of specific binding motifs [66]. The finding that Dichaete binding locations are marked by overrepresentation of binding motifs for factors defining HOT regions, coupled with the widespread gene expression effects of Dichaete mutations, suggests that Dichaete may also play a role in regulatory interactions at HOT enhancers. It is notable that Dichaete, in common with all other characterised Sox proteins, is known to bind DNA upon binding [56]. It is possible that Dichaete activity at HOT regions is mediated by this binding activity, helping to bring together complexes of other regulators. In this view, Dichaete would assist binding of factors at non-canonical target sites by favouring protein-protein interactions. In one of the bona fide Dichaete regulatory elements that have been studied in detail, the sli midline enhancer, Dichaete helps coordinate interactions between the POU factor Vf and a Sim/Tango heterodimer [25].

Aside from a proposed role at HOT regions, our analysis indicated Dichaete binds to and is active at many characterised regulatory elements. Almost half the enhancers catalogued by RedFly and a substantial fraction of neural enhancers identified by the FlyLight project [45,46] show evidence of Dichaete regulation. Along with this, we observed an association between Dichaete binding and transcriptional start sites, suggesting to us one of two possibilities. Either Dichaete directly engages with core promoter elements or looping interactions between Dichaete bound enhancers and the transcriptional machinery results in ChIP or DamID assays capturing these interactions. In this respect we note that Dichaete binds in the minor groove of DNA, perhaps making it more likely to capture indirect interactions.

Whether Dichaete acts at defined tissue-specific enhancers, HOT regions, core promoters, or all three, our analysis uncovered widespread involvement in specific developmental processes in the embryo. For example, our previous studies highlighted a role for Dichaete in hindgut morphogenesis and identified dpp as a likely target gene, since targeted dpp expression in the hindgut of Dichaete mutants was able to partially rescue the phenotype [16]. Our new analysis implicates Dichaete in the regulation of many of the key factors responsible for hindgut specification and morphogenesis, with most of the characterised transcription factors or signalling pathway components known to be important for hindgut development [52] bound and regulated by Dichaete. This further emphasises the view that Dichaete plays a hub-like role in controlling regulatory networks. We note that hindgut phenotypes and gene expression are unlikely to be functionally compensated by other Sox factors. While the group E gene Sox100B is also expressed in the embryonic hindgut [67], we have not seen evidence for synergistic interactions between Dichaete and Sox100B mutants (SR unpublished observations) and thus functional compensation by Sox100B is less likely. On the other hand, the related group B gene Sox21b is expressed in the hindgut and partially overlaps with Dichaete [9]. Although deletions encompassing Sox21b show no obvious phenotype, assessing possible functional compensation of Dichaete functions is difficult due to the close proximity of the two genes (~40 kb). It has recently been reported that human SOX2 is involved in gut development where it interacts antagonistically with CDX2 [68]. Caudal is a Drosophila orthologue of CDX2 and we found Dichaete binding and associated repression of cad, hinting at further levels of regulatory network conservation across metazoa.

In common with vertebrate group B genes, Dichaete plays a prominent role in the CNS. Many previous studies focused on single genes have shown that Dichaete is involved in neural specification via the regulation of proneural genes in the Achaete-scute complex [21-23] and our current analysis provides a genomic perspective on this, identifying extensive Dichaete binding across the complex. Importantly, much of this binding coincides with mapped regulatory elements and we found changes in the expression of complex genes in Dichaete mutants. Dichaete is involved in the temporal cascade that confers specific identities to neuroblasts and their progeny [32] and our analysis provides considerable insights into this role. We found Dichaete binding associated with all of the characterised genes in the temporal cascade, as well as considerable overlapping binding between Dichaete, Hb and Kr, strongly supporting the idea that cross-regulatory interactions between these genes is important for correct neural specification. For example, maintenance of Hb or loss of Cas, the first and last genes in the
cascade, lead to prolonged expression of Dichaete and cells remain in a neuroblast state [32]. Our analysis suggests that Dichaete may help maintain the temporal cascade expression in the neuroblast.

Finally, our analysis uncovered a striking relationship between Dichaete and Pros, with Dichaete negatively regulating pros expression early in neural development. In addition, both proteins show an extensive and highly significant overlap in their binding profiles. The gene expression data indicate that Dichaete and Pros may have antagonistic interactions since we find genes encoding neuroblast functions (e.g. ase, insc, mira and dpm) are downregulated in Dichaete mutants but upregulated in pros mutants [59]. However, we also find that genes involved in aspects of neuronal differentiation (e.g. esc, zfh1 and Lim1) are positively regulated by both factors. Taken together it is tempting to speculate that in neuroblasts, when Pros is cytoplasmic, Dichaete positively regulates genes required to maintain the self-renewal state and keeps pros levels down. In the GMC, Dichaete function must be downregulated to allow cells to exit the cell cycle and differentiate [32], consequently pros expression would be upregulated and the protein translocated to the nucleus by the well-established asymmetric division mechanism, repressing neuroblast genes and promoting differentiation. While Dichaete appears to be uniformly expressed in the neuroectoderm, its expression in neuroblasts is dynamic with many neuroblasts expressing Dichaete transiently [23,32]. In addition, and related to the subcellular partitioning of Pros, Dichaete is reported to shuttle between cytoplasm and nucleus, at least early in CNS development [56]. Furthermore, Dichaete is dynamically expressed in GMCs and their progeny, consistent with the proposed interaction with Pros [16]. These observations are consistent with the view that control of Dichaete is important for first determining self-renewal versus differentiation, followed by a role in aspects of neuronal differentiation.

The emerging view from our studies and previous work with Dichaete is of a transcriptional regulator with multifaceted roles in development. We have previously shown that mammalian Sox2 can provide Dichaete function, rescuing Dichaete mutant phenotypes. However, the designation of Dichaete as a group B1 protein based on functional arguments is considered by some to be inconsistent with phylogenetic arguments that firmly place Dichaete in the B2 group [11]. In vertebrates, group B2 proteins act as transcriptional repressors, antagonising group B1 functions [6,7]. Since we see very few genes upregulated in Dichaete mutants, our analysis suggests that Dichaete may be acting primarily as a transcriptional activator. However, this type of mutant expression study is prone to pleiotropic effects, so further investigation of specific targets and tissues is needed. In vertebrates the group B1 proteins play critical roles in the specification and maintenance of neural stem cells, exactly the functions described for Dichaete. The observed correspondence between Dichaete and Sox2 target genes show that these proteins are not only conserved at the functional level when assayed in mutant rescue experiments but also, remarkably, at the level of the gene regulatory networks they control in the fly and mouse nervous system.

One possible explanation for these disparate findings regarding the classification of Dichaete as a group B1 or B2 protein may be provided by the role of Dichaete in the regulation of proneural genes and its early activity on pros. In these specific cases, Dichaete acts to repress these genes in the neuroectoderm while SoxN acts as an activator [19-22]. It is therefore possible that, in the last common ancestor of the vertebrates and invertebrates when the B1- B2 split occurred, the ancestral Dichaete gene had an limited B2-like repressor role as well as more prominent B1-like activator role in the CNS. As the lineages diverged the vertebrate B2 genes evolved specialised repressor functions while, in the invertebrates, they maintained more basal activator function. Support for the idea that insect Sox genes represent conserved basal functions of more diverged vertebrate family members comes from experiments replacing the mouse group E gene, Sox10, with the fly Sox100B coding sequence. In these studies the fly gene is able to provide substantial Sox10 function in the developing embryo, more so than the Sox8 gene, which is far closer to Sox10 at the sequence level [69].

Conclusions
In summary, we present a rigorous analysis of the genomics of the Drosophila group B transcription factor Dichaete, highlighting regulatory input into several key developmental pathways. Our studies provide a baseline for more detailed analysis of highly conserved aspects of group B Sox function in neural stem cells and in neuronal differentiation.

Methods

DamID

Ecoli Dam was introduced into a pUAST plasmid carrying a full-length Dichaete cDNA to produce a Dam_Dichaete fusion [15] and this vector used to generate transgenic fly lines via standard P-element transformation [70]. A Dam only vector was prepared in parallel. Embryos from homozygous Dichaete-Dam and Dam only lines were collected from 2-7 hrs after egg laying (stages 5-11) and DNA processed for DamID according to the method of Vogel et al. [71]. Biological triplicates of enriched DNA from experimental and control lines were amplified by PCR and Cy3/ Cy5 dyes incorporated by random priming with Klenow polymerase. Combined DNA samples were loaded onto Nimblegen D.mel ChIP 2.1 M tiling arrays (GEO platform GPL15057), hybridized overnight at 42°C, then washed
and scanned in an Axon GenePix scanner the following day. Scanned arrays were quantile normalised separately for samples and controls, and peak calling was performed using RINGO [72] to identify binding intervals at different False Discovery Rates. Full DamID data is available from the NCBI Gene Expression Omnibus as part of Super Series GSE49095.

Gene expression
Stage 10–11 embryos (5 and 7.5 hours after egg laying) from a cross between $D^{72}$/TM3, twi-GAL4 UAS-Gfp and $D^{513}$/TM3, twi-GAL4 UAS-Gfp, were hand picked under a fluorescence dissecting microscope. GFP negative homozygous Dichaete mutant embryos and their heterozygous single GFP positive siblings were collected and approximately 150 embryos per sample were stored frozen in Trizol. Following RNA extraction, reverse transcription, Klenow amplification and labelling, samples were hybridised to INDAC FL003 (GEO:GPL14121) Drosophila gene expression arrays using our standard protocols (http://www.flychip.org.uk/). Four biological replicates were performed for each experiment, with 2 dye swaps incorporated into the experimental design to control for bias. Arrays were quality checked manually, removing spots affected by high levels of background or artefacts. Our standard data analysis pipeline was employed (http://www.flychip.org.uk/) using Dapple for spotfinding and quantifying signal intensities [73]. The normalisation step was performed using invariant normalisation [50] to address the fact that the majority of genes change expression, with the later analysis stages performed with limma. The thresholds used to find differentially expressed genes were average M-value $< -1$ or $> 1$, and p-value $< 0.01$. All gene expression data is available from the NCBI Gene Expression Omnibus as part of Super Series GSE49095.

Dichaete core binding
Existing ChIP data was obtained from the BDTNP website [34], where we used the published binding intervals, and from modENCODE ([35]; DCCid modENCODE_2571). In the latter case we reprocessed the raw data using the quantile normalisation and peak-calling approaches described above. To create the core dataset, we used intervals supported by at least one low stringency (25% FDR) and one high stringency (1% FDR) dataset from independent experimental techniques. This meant that all intervals confirmed by at least one 1% FDR ChIP and the 25% FDR DamID, and vice versa, were included in the final Dichaete core set (Figure 1A). A .bed file of Dichaete core binding intervals is provided as Additional file 14: Table S14.

Genome version and annotations
All genomic coordinates are in Genome Release 5 (dm3), and the genome annotations used were FlyBase R5.48, obtained from FlyMine v36.0 [39]. Where relevant, binding coordinates were converted to Genome Release 5 (dm3) using the UCSC liftover tool. Unless indicated otherwise, analysis was performed custom-written Perl scripts (v5.12.4), with the graphs and statistical tests done in R 2.15.2. The gene assignments to intervals were performed as follows: all genes directly overlapping a particular binding interval were assigned as hits to that interval. If an interval was found to overlap no genes, the closest gene would instead be assigned as a hit, up to the maximum of a 10 kb range around both sides of the interval. Drosophila orthologues of mouse Sox2 target genes were identified with FlyMine using data from TreeFam release 8.0.

Gene list analysis
All gene lists were analysed using FlyMine [39] to obtain basic summary statistics and enrichment analysis, including GO term and domain enrichment. For enrichment, FlyMine uses a hypergeometric distribution, and the Holm-Bonferroni correction was applied to get the adjusted p-values. In the case of the binding data, a gene length correction was also applied. The expression heatmap of genes differentially expressed in Dichaete null mutants was generated using the Fly Express 6.0 GEMs tool [51]. Data on which genes are likely to be transcription factors was obtained from FlyTF v2 [40].

Motif analysis
The de novo motif finding was performed using i-CisTarget [37]. The RSAT matrix-scan tool was used for motif scanning: 1st order Markov Model was used to generate the background model from the input sequences, and the sequence matches with the weight score $> = 4$ were retrieved [74].

Binding interval location
For the purposes of this analysis, a 1 nucleotide coordinate in the centre of each binding interval was considered to be an approximation of the location of the Dichaete binding event. An overlap with known genome features was then determined using gene, exon and intron coordinates from FlyBase R5.48, obtained via FlyMine. For the transcription start site (TSS) analysis, the start coordinate for each gene was taken as an approximation of the TSS location. Enhancer locations were obtained from the REDFly database version 3.2 [42], and the intervals from the FlyLight project were used as published [45,46]. The overlaps were found using Bedtools v2.17 [75].

Transcription factor binding overlaps
For the broader analysis of the transcription factor network in embryogenesis, the binding data was not reanalysed. Instead, the peak interval files provided were used
for the analysis (a list of the datasets used is provided on Additional file 15: Table S15). In the cases where more than one dataset falling within the 0-12 h period was available for a particular transcription factor, a union of the datasets was created for that TF, and this was used in the co-occurrence analysis [57].

Additional files

Additional file 1: Table S1. List of genes with Dichaete binding intervals from 1% FDR Dichaete DamID.

Additional file 2: Table S2. Gene Ontology Biological Process enrichment analysis of Dichaete-bound genes from 1% FDR DamID.

Additional file 3: Table S3. Cooccur Analysis of TF overlaps. Headers: tf1, Dichaete dataset 1; tf2, 1 Dichaete dataset 1; overlapped_in_tf1, number of TF1 binding interval overlapping TF2; n_tf1, number of binding intervals in TF1 dataset; percent_in_tf1, fraction of TF1 dataset overlapping with TF2; overlapped_in_tf2, number of TF2 binding intervals overlapping with TF1; n_tf2, number of TF2 binding intervals in dataset; percent_in_tf2, fraction of TF2 intervals overlapping TF1; n_samples, number of resampling runs; pvalue, raw p value; padj, adjusted p value.

Additional file 4: Table S4. Core Dichaete bound genes from the amalgamation of DamID and ChIP datasets.

Additional file 5: Table S5. Gene Ontology Biological Process enrichment analysis of Core Dichaete-bound genes.

Additional file 6: Table S6. RedFly and FlyLight enhancers associated with Dichaete Binding.

Additional file 7: Table S7. Genes differentially expressed comparing Dichaete mutant embryos to wild type. Column headers: Transcript, FlyBAse transcript ID; Fbgn, FlyBase gene ID; D_mutant_invar_limma p-value, p-value for expression change; D_mutant_invar_log2 fold expression change, Log2 fold expression change; D_mutant_invar_limma p-value, p-value for expression change.

Additional file 8: Table S8. Expression changes of Dichaete target genes. Column headers: FlyBase transcript ID; Fbgn, FlyBase gene ID; Gene, FlyBase gene symbol; D_intergenic, number of TF1 binding intervals in dataset; percent_in_tf1, fraction of TF1 dataset overlapping with TF1; n_samples, number of resampling runs; p_value, raw p value; padj, adjusted p value; Gene 1, TF 1 gene name; Gene 2, TF2 gene name.

Additional file 9: Table S9. Gene Ontology Biological Process enrichment analysis of Dichaete target genes.

Additional file 10: Table S10. Cooccur Analysis of TF overlaps. Headers: tf1, transcription factor 1 dataset; tf2, transcription factor 2 dataset; overlapped_in_tf1, number of TF1 binding interval overlapping TF2; n_tf1, number of binding intervals in TF1 dataset; percent_in_tf1, fraction of TF1 dataset overlapping with TF2; overlapped_in_tf2, number of TF2 binding intervals overlapping with TF1; n_tf2, number of TF2 binding intervals in dataset; percent_in_tf2, fraction of TF2 intervals overlapping TF1; n_samples, number of resampling runs; pvalue, raw p value; padj, adjusted p value; Gene 1, TF 1 gene name; Gene 2, TF2 gene name.

Additional file 11: Table S11. Genes bound by Dichaete and Prospero.

Additional file 12: Table S12. Drosophila orthologues of mouse Sox2 bound genes in the Dichaete core binding set.

Additional file 13: Table S13. Drosophila orthologues of mouse Sox2 bound genes in the Dichaete target gene set.

Additional file 14: Table S14. Dichaete core binding intervals in bed format.

Additional file 15: Table S15. Table of transcription factor binding data used in overlap analysis.

Competing interests
The authors declare that they have no competing interests.

Authors’ contributions
JA performed the gene expression experiments and the majority of the data analysis. EF performed the DamID experiments. BF assisted with microarray data analysis. SPS performed the pros mutant analysis. SR conceived the study, provided the funding and analysed the data. JA and SR wrote the paper. All authors read and approved the manuscript.

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Author details
1Department of Genetics, University of Cambridge, Cambridge, UK.
2Cambridge Systems Biology Centre, University of Cambridge, Cambridge, UK. 3Present address: Wellcome Trust-Medical Research Council Cambridge Stem Cell Institute, Tennis Court Road, Cambridge CB2 1QR, UK. 4Present address: Meridigen Biotech, Taipei 114, Taiwan.

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