Molecular Analysis of the Notch Repressor-Complex in *Drosophila*: Characterization of Potential Hairless Binding Sites on Suppressor of Hairless

Patricia Kurth¹, Anette Preiss¹, Rhett A. Kovall², Dieter Maier¹*

¹ Institut für Genetik (240), Universität Hohenheim, Stuttgart, Germany, ² Department of Molecular Genetics, Biochemistry and Microbiology, University of Cincinnati College of Medicine, Cincinnati, Ohio, United States of America

Abstract

The Notch signalling pathway mediates cell-cell communication in a wide variety of organisms. The major components, as well as the basic mechanisms of Notch signal transduction, are remarkably well conserved amongst vertebrates and invertebrates. Notch signalling results in transcriptional activation of Notch target genes, which is mediated by an activator complex composed of the DNA binding protein CSL, the intracellular domain of the Notch receptor, and the transcriptional coactivator Mastermind. In the absence of active signalling, CSL represses transcription from Notch target genes by the recruitment of corepressors. The Notch activator complex is extremely well conserved and has been studied in great detail. However, Notch repressor complexes are far less understood. In *Drosophila melanogaster*, the CSL protein is termed Suppressor of Hairless (Su(H)). Su(H) functions as a transcriptional repressor by binding Hairless, the major antagonist of Notch signalling in *Drosophila*, which in turn recruits two general corepressors – Groucho and C-terminal binding protein CtBP. Recently, we determined that the C-terminal domain (CTD) of Su(H) binds Hairless and identifies a single site in Hairless, which is essential for contacting Su(H). Here we present additional biochemical and in vivo studies aimed at mapping the residues in Su(H) that contact Hairless. Focusing on surface exposed residues in the CTD, we identified two sites that affect Hairless binding in biochemical assays. Mutation of these sites neither affects binding to DNA nor to Notch. Subsequently, these Su(H) mutants were found to function normally in cellular and in vivo assays using transgenic flies. However, these experiments rely on Su(H) overexpression, which does not allow for detection of quantitative or subtle differences in activity. We discuss the implications of our results.

Introduction

The Notch signalling pathway is highly conserved in metazoans, where it allows for intercellular communication during the specification of cell fates [1]. Notch encodes a single pass transmembrane receptor that is activated by transmembrane ligands presented by the signalling cell. As consequence of receptor activation, the intracellular Notch domain (ICN) is cleaved and migrates to the nucleus. There it binds to the CSL-type DNA-binding protein C-promoter binding factor 1 (CBF-1) in *H. sapiens*, [lag-1] in *C. elegans*, Suppressor of Hairless in *D. melanogaster* (Su(H)), and assembles, together with the coactivator Mastermind (Mam), a transcriptional activator complex (overview in [1–4]). Formation of the CSL-ICN-Mam ternary complex, in conjunction with other transcriptional components, results in the activation of Notch target genes, e.g. the *Hairy* and *Enhancer of split* (HES) family of genes. HES genes encode transcriptional repressors that function to shut down gene expression for genes that confer the primary cell fate, thereby enforcing a secondary fate within the signal-receiving cell [1–2].

The components of the activator complex (CSL-ICN-Mam) are highly conserved from worms and flies to humans in both primary sequence and the overall three-dimensional structure of this complex [5–6]. The central molecule of the activator complex is CSL, which contains three functional domains: the N-terminal domain (NTD), beta-trefoil domain (BTD), and C-terminal domain (CTD). Both the NTD and BTD contact DNA. The BTD and the CTD interact with ICN, whereby BTD forms a high-affinity interaction with the RAM domain of ICN and the CTD binds both the ankyrin repeats (ANK) of ICN and Mam [5–6], overview in [3].

In the absence of signal, CSL interacts with transcriptional corepressors to turn off transcription from Notch target genes. Similar to the activator complex, CSL is the central component of the repressor complex; however, in contrast to the activator complex, the structure of the repressor complex is still unknown. Human CBF-1 has been shown to interact with several different corepressors, e.g. SMRT/NCOR, MINT/SHARP, KyoT2, and CIR. Most of these corepressors contact a site within the BTD of CBF-1 that likely overlaps where the RAM domain of Notch binds. This has led to a model, in which the repression and activation of Notch target genes is mediated by the competition of ICN and corepressors for binding CBF-1 (overview in [7]).
melanogaster, the transcriptional corepressor Hairless is the major antagonist of Notch signalling (reviewed in [8]). Hairless binds to the CTD of Su(H) – the fly CSL protein - and recruits two additional corepressors, the C-terminal binding protein (CtBP) and Groucho (Gro). Together this repressor complex silences expression from Notch target genes ([9–12]). Hence, CSL plays a dual role in both activator and repressor complexes.

We have initiated a detailed analysis of the Notch repressor complex in *Drosophila*. Recently, we have shown that Hairless and Su(H) form a high affinity complex, and that mutations within Su(H) that affect binding of ICN have no effect on Hairless binding. Nonetheless, Hairless and Notch compete for Su(H) in vitro, despite the disparities in affinities of ICN and Hairless for the CTD of Su(H). Moreover, we have mapped a single residue in Hairless that is crucial for binding Su(H) ([10]). To further our understanding of Notch signalling and the repressor complex in *Drosophila*, we have analysed 17 single, double, and triple mutations in the CTD of Su(H) for their involvement in the binding of Hairless using a yeast two-hybrid assay. A double mutation was identified that strongly reduces interactions with

![Figure 1. Fine mapping of the Hairless contact sites on Su(H) CTD. A) Surface representation of CSL-DNA structure with the NTD, BTD, and CTD coloured in dark and light grey, respectively. The DNA is in a stick representation with carbon, oxygen, nitrogen, and phosphorous atoms coloured yellow, red, blue, and orange, respectively. Notch ANK is coloured in blue and MamN in red, and represented as transparent ribbons. The residues on the CTD that interacted with ICN and Mam in the yeast two-hybrid assay are coloured green. Residues that were mutated in the course of this work are coloured magenta. B) Primary sequence of the Su(H)-CTD construct; the CTD is shown in bold. Amino acids shown to contact ANK/Mam are depicted in green; red are those tested for Hairless binding, and brown depicts the AR2 mutation that disrupts the CTD fold. C) Mutant CTD constructs were tested in a yeast two-hybrid assay for binding to full length Hairless (HFL) and to intracellular Notch (ICN I). Moreover, the mutant CTD constructs were tested in a yeast three-hybrid assay for their potential to assemble the ternary activator complex with ANK and MamN. Empty vectors served as negative controls. Relative position of mutations within the CTD is indicated. Note that binding of Hairless but not of ICN I to LEWA or WA is reduced and is nearly abolished in WARE and WVR. However, all these mutants display normal binding to Notch and are capable of forming a ternary activator complex. The constructs GD, GH, VEAD and CR bear mutations at the CTD-ANK and CTD-Mam interfaces, consistent with a strongly reduced binding to ICN I. Hence, Notch and Hairless contact different sites on Su(H). Mutant AR2, in which residues within the hydrophobic core of CTD are mutated, is likely compromised for folding, and fails to bind either HFL or ICN I.

doi:10.1371/journal.pone.0027986.g001
Hairless, but neither affected DNA nor Notch binding by Su(H).
In spite of this reduction in binding, overexpression of the Su(H)
double mutant in a transciplational cell culture assay, as well as in
the fly, revealed little to no changes in function compared to wild
type Su(H). These results were unexpected and we consider two
possibilities: (1) potentially other residues in Su(H) contribute to
the binding of Hairless, which allows for a sufficiently strong
interaction in vivo; or (2) alternatively, the presence of endogenous
Su(H) in our cellular and in vitro assays distorts our results.

Results

Identification of potential Hairless binding sites in the
CTD of Su(H)

Recently, we have identified the C-terminal domain of Su(H) as
the binding domain for Hairless (CTD, amino acids 417–528).
Binding to Hairless was enhanced by the presence of the N-
terminal z-helix (amino acids 1–119), which helps to stabilize the
folding of the CTD. Mutations that affect binding to ICN did not
interfere with the binding to Hairless, suggesting that ICN and
Hairless do not compete for the same contact sites in Su(H) CTD
[10].

To identify the amino acids in CTD responsible for interaction
with Hairless, a total of 17 single, double or triple amino acid
substitutions were introduced by in vitro mutagenesis. The main
criterion for the changes was (1) whether the amino acids were
surface exposed, which was based on the orthologous mammalian
and C. elegans CSL structures; and/or (2) within a putative protein-
protein interaction domain that was determined computationally
(http://spider.cchmc.org) (Fig. 1A,B). The sites of mutation were
changed to residues that would likely interfere with Hairless
binding (Fig. 1B,C). The mutant constructs were tested in a yeast
two-hybrid assay using Hairless or ICN I as bait (Fig. 1 C). In
addition, we assayed for the formation of the ternary activator
complex consisting of Su(H), Notch Ank and MamN (Fig. 1C;[10]).
For the majority of mutants examined no changes in binding
were detected. However, four mutations showed reductions in
Hairless binding: CTDLEWA (L490E/W491A), CTDWA (W491A),
CTDWARE (W491A/R493E) and CTD WVR (W491A/V492R/
R493E). Together, this assay revealed that the residues Trypto-
phan 491 and Arginine 493 are likely important for the binding of
Hairless, because the combined mutation CTD WARE nearly
abolished Hairless binding in the yeast assay. In addition, Leucine
490 appeared to contribute since CTDLEWA bound less well than
CTDWA, whereas mutation of Valine 492 did not further reduce
binding in CTD WVR. The single mutations CTD RE (R493E) and
CTD VR (V492R) were without effect (Fig. 1C).

Two mutations CTDVEAD (V442E/A443D) and CTD CR (G471R)
nearly abolished binding to ICN I, and two other mutations CTD GD
(G438D) and CTD GH (G439H CTD CR) reduced binding to ICN I;
however, none of these mutations affected binding to Hairless
(Fig. 1C), CTD CR, CTD GD, and CTD GH lie within the region
known to contact the Notch Ankyrin repeats, whereas CTD VEAD is
in the vicinity of ANK and MAM, but does not make direct contact
with these proteins. Two controls were included, the empty vector and
the double mutant CTD MR2 (W459A/F460A). The CTD MR2 mutant
affects amino acids buried within the hydrophobic core of CTD and
is hence predicted to disrupt CTD folding. As expected, both controls
did not bind to either Hairless or ICN I, and failed to assemble the
ternary complex with N-Ank and MamN (Fig. 1C).

Based on the structures of mammalian and worm CSL proteins,
Su(H) is expected to bind DNA with its N-terminal and beta-trefoil
domains (NTD, BTD; Fig. 1A). Accordingly, mutations in CTD
should not interfere with DNA binding, which was confirmed by
an electrophoretic mobility shift assay (EMSA) with the relevant
Su(H) mutants (Fig. 2). To this end, R496E, W491A, L490E/
W491A and W491A/R493E mutations were introduced into full
length Su(H) cDNA that was in vitro transcribed and translated.
The E(spl) m8 oligo nucleotide containing a Su(H) binding site
(m8-S1, [44]) was used as the target DNA. No difference in DNA-
binding was observed between wild type and mutant Su(H) protein
(Fig. 2).

The CTD WARE double mutation fails to bind truncated forms
of Hairless

Based on the near complete loss of binding to full length
Hairless, the double mutation CTD WARE (W491A/R493E) was
chosen for further analysis. Previously, we defined a subdomain
of Hairless, termed NTCT (amino acids 171–375; [10]), which
recapitulated all of the binding of Hairless to Su(H) in vitro.
Unexpectedly, we found that CTD WARE bound to NTCT
similarly to full length Su(H) and only slightly weaker than wild
type CTD (Fig. 3). In addition, we tested the ability of two Hairless
NTCT mutants, the NT-deletion NTCT ANT and the single site
mutant NTCT LD (L235D) for binding to the Su(H) constructs.
Both NTCT mutants fail to bind to full length Su(H) and showed a
markedly reduced binding activity towards CTD (Fig. 3; [10]).
However, NTCT ANT and notably NTCT LD completely failed to
bind to CTD WARE, strongly indicating that the affected amino
acids are involved in the binding of Su(H) and Hairless (Fig. 3).

Figure 2. DNA binding is not altered in Su(H) mutants.
Electromobility shift assay for binding of Su(H) protein variants to the
radiolabelled E(spl)m8-S1 oligo [44]. Control, no protein added (lane 1).
The binding of the mutant Su(H) WARE, Su(H) RE, Su(H) LEWA, or Su(H) WARE
proteins (lane 3–6) to DNA was similar as the wild-type Su(H) protein
(lane 2).

doi:10.1371/journal.pone.0027986.g002
Su(H)\(^{\text{WARE}}\) gives a normal response in a transcriptional assay

Thus far, our data indicated that the W491A/R493E amino acid substitutions had an effect on the Su(H)-Hairless interaction. In order to test the effect of this mutation in a more physiological setting, it was introduced into the Su(H) full length cDNA [Su(H)\(^{\text{WARE}}\)] and transiently expressed in S2 cells. Subsequently, we analysed the transcriptional activation and repression of a luciferase reporter construct bearing Su(H) binding sites (NRE-reporter; [13]). We reasoned that Su(H)\(^{\text{WARE}}\) should behave as a transcriptional activator together with ICN, similar to wild type Su(H), because binding of Notch was unaffected by the mutation. This was indeed observed – transfection of ICN alone strongly activates the NRE-reporter, via endogenous Su(H), which was taken as 100% to normalize the other results [13]. Addition of the wild type Su(H) construct resulted in about three- to four-fold increase of luciferase activity (Fig. 4), which is in agreement with earlier observations [10,14]. A likewise increase in reporter activity was obtained by adding Su(H)\(^{\text{WARE}}\), indicating that the mutant protein can efficiently assemble an activator complex in S2 cells (Fig. 4).

Assembly of the repressor complex was tested by cotransfecting the S2 cells with Hairless and ICN. This caused a strong downregulation of ICN mediated transcriptional activation of the NRE-reporter to about 40% [10,12], because Hairless can assemble a repressor complex with endogenous Su(H) on the NRE promoter. Interestingly, Hairless is able to abrogate the strong activation mediated by the addition of exogenous Su(H) to near completion (Fig. 4; [10]), indicating that Hairless and Su(H) interact with each other. To our surprise, the same degree of repression was observed with Su(H)\(^{\text{WARE}}\) (Fig. 4). Apparently, Hairless binds the Su(H) mutant with sufficiently enough affinity to repress transcription as efficiently as the wild type Su(H) protein. This result was unexpected since the yeast two-hybrid data suggested a near complete lack of binding of Su(H)\(^{\text{WARE}}\) to Hairless. However, at this stage of our analysis, it was unclear whether this effect was specific to S2 cell culture.

In vivo transcriptional response of Notch target genes during wing development

To analyse the in vivo activity of the mutant Su(H) protein, transgenic flies were established using the PhiC31 method [15]. This system avoids position effects and hence allows the direct comparison of different transgenes at the same location. Su(H)\(^{\text{WARE}}\) was cloned into an appropriate UAS-vector and integrated at the 96E landing site for comparison with the accordant Su(H) construct [10]. Moreover, the transgenic Su(H)\(^{\text{WARE}}\) line was crossed with full length Hairless HFL and with mutant Hairless H\(^{\text{L}}\), each integrated at 68E, to allow for a combined overexpression. The latter completely failed to bind wild type Su(H) [10]. The wild type and mutant Su(H) and Hairless transgenes were locally overexpressed using the Ga4/UAS-system [16].

First we analysed the consequences on the expression of the Notch target gene wingless (wg). Wg is expressed in the developing wing imaginal disc in a ring outlining the presumptive wing pouch and along the dorso-ventral boundary, which eventually forms the margin of the wing (Fig. 5) [17,18]. The constructs were induced singly or in combination in a central area of the wing disc. Overexpression of either Su(H) or Su(H)\(^{\text{WARE}}\) effects an overproliferation of the affected tissue, which is typical for Notch gain of function, suggesting that both caused the activation of Notch target genes. Accordingly, a subtle expansion of Wg expression was observed compared to the control (Fig. 5). The ectopic Wg expression in the inner and outer rings was not anticipated since wg, according to several publications, is not a Notch target in this part of the tissue [19–21]. As expected, overexpression of Hairless HFL antagonized the expression of Wg at the intersection of the HFL expression domain and the presumptive margin and led to less tissue due to cell death.

Figure 3. Su(H)\(^{\text{WARE}}\) binding capacity to Hairless mutants. Yeast two-hybrid assay to test for binding activity of wild type Su(H), CTD and mutant CTD\(^{\text{WARE}}\) constructs with Hairless NTCT, NTCT\(^{\text{L}}\) and NTCT\(^{\text{LD}}\); empty vector served as a negative control. Note the lack of binding of CTD\(^{\text{WARE}}\) with the \(\Delta N\) deletion or the NTCT\(^{\text{L}}\) mutation. doi:10.1371/journal.pone.0027986.g003

Figure 4. Activation and repression of a Notch reporter gene by Su(H) variants. Effects of the mutant Su(H)\(^{\text{WARE}}\) (WARE) on Notch ICN mediated expression from the NRE-reporter (luciferase reporter; [13]). We reasoned that Su(H)\(^{\text{WARE}}\) should behave as a transcriptional activator together with ICN, similar to wild type Su(H), because binding of Notch was unaffected by the mutation. This was indeed observed – transfection of ICN alone strongly activates the NRE-reporter, via endogenous Su(H), which was taken as 100% to normalize the other results [13]. Addition of the wild type Su(H) construct resulted in about three- to four-fold increase of luciferase activity (Fig. 4), which is in agreement with earlier observations [10,14]. A likewise increase in reporter activity was obtained by adding Su(H)\(^{\text{WARE}}\), indicating that the mutant protein can efficiently assemble an activator complex in S2 cells (Fig. 4).

Figure 4. Activation and repression of a Notch reporter gene by Su(H) variants. Effects of the mutant Su(H)\(^{\text{WARE}}\) (WARE) on Notch ICN mediated expression from the NRE-reporter (luciferase reporter; [13]). We reasoned that Su(H)\(^{\text{WARE}}\) should behave as a transcriptional activator together with ICN, similar to wild type Su(H), because binding of Notch was unaffected by the mutation. This was indeed observed – transfection of ICN alone strongly activates the NRE-reporter, via endogenous Su(H), which was taken as 100% to normalize the other results [13]. Addition of the wild type Su(H) construct resulted in about three- to four-fold increase of luciferase activity (Fig. 4), which is in agreement with earlier observations [10,14]. A likewise increase in reporter activity was obtained by adding Su(H)\(^{\text{WARE}}\), indicating that the mutant protein can efficiently assemble an activator complex in S2 cells (Fig. 4).

Assembly of the repressor complex was tested by cotransfecting the S2 cells with Hairless and ICN. This caused a strong downregulation of ICN mediated transcriptional activation of the NRE-reporter to about 40% [10,12], because Hairless can assemble a repressor complex with endogenous Su(H) on the NRE promoter. Interestingly, Hairless is able to abrogate the strong activation mediated by the addition of exogenous Su(H) to near completion (Fig. 4; [10]), indicating that Hairless and Su(H) interact with each other. To our surprise, the same degree of repression was observed with Su(H)\(^{\text{WARE}}\) (Fig. 4). Apparently, Hairless binds the Su(H) mutant with sufficiently enough affinity to repress transcription as efficiently as the wild type Su(H) protein. This result was unexpected since the yeast two-hybrid data suggested a near complete lack of binding of Su(H)\(^{\text{WARE}}\) to Hairless. However, at this stage of our analysis, it was unclear whether this effect was specific to S2 cell culture.
However, induction of the mutant HLD was indistinguishable from wild type confirming complete loss of Su(H) binding in this mutant.

A combined overexpression of Hairless and Su(H) led to a remarkable loss of tissue and repression of \(wg\) expression (Fig. 5), which is in accordance with earlier observations and can be explained by the formation of a large surplus of repressor complexes formed [10–12,22]. The mutant HLD fails to bind to Su(H), therefore, the combined overexpression resembled the phenotype of the sole Su(H) overexpression (Fig. 5). Again, in combination with HFL, Su(H)WARE behaved largely identical as wild type Su(H), indicating the normal formation of repressor complexes (Fig. 5), which also confirms our S2 cell culture results.

Activity Su(H) and Hairless protein variants during eye development of the fly

To substantiate these results we extended our analysis to the Drosophila eye, where Notch signalling is required at multiple, subsequent steps (reviewed in [23]). We used the gmr-Gal4 line that drives expression in the differentiating retina [24]. As reported earlier [25], overexpression of Su(H) is characterized by an overproliferation of eye tissue, as expected for a gain of Notch activity (Fig. 6). A likewise phenotype was induced by the overexpression of Su(H)WARE in accordance with its ability to assemble an activator complex together with Notch (Fig. 6a). In contrast, expression of the antagonist Hairless resulted in small, irregular eyes by interference with several Notch dependent processes and subsequent induction of apoptosis (Fig. 6b) [26–28]. In combination, Su(H) and Hairless overexpression led to almost eyeless flies: only small slits remained lacking ommatidial structures or eye color (Fig. 6b). Again, a similar result was observed with Su(H)WARE indicating little differences compared to wild type Su(H). In contrast, overexpression of the mutant HLD had little biological effect and did not influence the activity of the Su(H) constructs (Fig. 6b). Similar results were observed in the process of lateral inhibition during bristle formation on the thorax ([10] and not shown).

The small eyes resultant from Hairless overexpression are partly due to apoptosis induced by the repression of several Notch target genes and the concomitant downregulation of EGFR signalling activity [27–29]. We wondered whether the primary cause of the extreme adult phenotypes seen with the combined overexpression of HFL and Su(H) was also due to apoptosis. This was confirmed by staining for the cleaved, active form of Caspase-3 – the final effector Caspase in the apoptotic cascade [30] – which was dramatically increased in the eye discs of the relevant combinations (Fig. 7). In contrast no apoptosis was seen upon ectopic expression of the mutant HLD (Fig. 7). This result is in agreement with the adult eye phenotype. Because HLD fails to bind to Su(H), HLD cannot be recruited to the respective promoters to assemble the respective repression complex, explaining the absence of apoptosis.

Discussion

While we have detailed knowledge of the structure of the ternary activator complex (CSL-ICN-MAM), repression of Notch two proteins. Overexpression of mutant Su(H)\(^{WARE}\) results also in a slight overproliferation of the wing disc and subtle \(Wg\) expansion. Also in combination with Hairless, Su(H)\(^{WARE}\) strongly impedes proliferation and \(Wg\) expression. However, no such effect of HLD on Su(H)\(^{WARE}\) can be observed.

doi:10.1371/journal.pone.0027986.g005
respectively, using gmr-Gal4 as driver line. As a control, lacZ was overexpressed which gives a wild type looking eye. Su(H) and Su(H)WARE induce a slight overgrowth of tissue resulting in enlarged eyes. Overexpression of HLD causes a nearly wild type eye. A combination of Su(H) with HFL results in a complete loss of the ommatidia: only a small eye slit remains that is totally smooth and devoid of any red eye pigment. In contrast, eye specific overexpression of HFL causes smaller eyes with irregular arrangement of the ommatidia, giving a rough appearance. Overexpression of HLD causes nearly wild type eye. A combination of Su(H) with HFL results in a complete loss of the ommatidia; only a small eye slit remains that is totally smooth and devoid of any red eye pigment. In contrast, a combined overexpression of Su(H) and HLD causes enlarged eyes similar to the sole Su(H) overexpression, demonstrating lack of protein binding. Overexpression of Su(H)WARE alone or in combination with Hairless variants gives similar results suggesting again that the WARE mutation does not influence repressor complex formation.

Figure 6. Overexpression of Su(H) and Hairless protein variants affect Drosophila eye development. A) Ectopic expression of Su(H) (lane 2) and Su(H)WARE (lane 3) cause an increase of eye size compared with a control lac-Z (con, lane 1) ectopic expression using the gmr-Gal4 driver line. Eye size of male flies was measured from 19 to 20 individuals of each genotype. Average area is given in kilo pixel (kpx). Error bar represents standard deviation. B) The UAS transgenes Su(H) and Su(H)WARE were expressed singly or in combination together with full length Hairless HFL or HLD, respectively, using gmr-Gal4 as driver line. As a control, lacZ was overexpressed which gives a wild type looking eye. Su(H) and Su(H)WARE induce a slight overgrowth of tissue resulting in enlarged eyes. In contrast, eye specific overexpression of HFL causes smaller eyes with irregular arrangement of the ommatidia, giving a rough appearance. Overexpression of HLD causes a nearly wild type eye. A combination of Su(H) with HFL results in a complete loss of the ommatidia; only a small eye slit remains that is totally smooth and devoid of any red eye pigment. In contrast, a combined overexpression of Su(H) and HLD causes enlarged eyes similar to the sole Su(H) overexpression, demonstrating lack of protein binding. Overexpression of Su(H)WARE alone or in combination with Hairless variants gives similar results suggesting again that the WARE mutation does not influence repressor complex formation.

doii:10.1371/journal.pone.0027986.g006

signal transduction is less well understood (overview in [4,7]). We have started a more detailed analysis on the Notch repression complex in Drosophila which contains the CSL-type DNA binding protein Su(H), the bridging platform protein Hairless and the two general corepressors, Groucho and C-terminal binding protein [9,11,12,31]. We have shown recently that Su(H) and Hairless form a high affinity complex that involves the CTD of Su(H) and the NT-domain of Hairless [10]. Moreover, our work demonstrated that Notch can outcompete Hairless for the binding of Su(H). This observation is startling for two reasons: firstly, both Notch and Hairless show comparable affinity for Su(H) which is in the nanomolar range, and secondly, the two molecules contact different sites in Su(H), excluding a simple competition scenario [10]. Presumably, the switch between activator and repressor status is more complicated and may involve structural changes in Su(H) [32,33].

In order to provide the molecular basis for a deeper understanding of these processes we have started to map the Su(H)-Hairless sites of interaction. In Hairless, we have been able to determine a single amino acid that is crucial for the binding of Su(H) without overtly disturbing Hairless structure [10]. Here we identify two residues W491 and R493 in the CTD of Su(H) that likely contribute to the binding of Hairless based on our yeast two-hybrid data. Strikingly, the WARE mutant behaved similar to wild type in our cellular and in vivo assays. How could this discrepancy in our results be resolved? The simplest explanation is the involvement of one or more additional contact sites in Su(H) located elsewhere that sufficiently stabilize the binding of the full length Hairless and Su(H) proteins in vivo, but not the interaction between CTD and Hairless in the yeast assay. As we have already mutated most of the surface exposed residues in Su(H) CTD without affecting Hairless binding, we must conclude that single mutations are not disruptive and that we have not fortuitously hit upon the right combination of multiple amino acids in Su(H) to completely disrupt binding. Certainly the determination of the Su(H)-Hairless complex crystal structure will clarify the role of these residues in Hairless binding, as well as define other important interaction regions.

However, we also need to consider the quantitative differences in the approaches. In the yeast the molecules are tested in a near 1:1 molar ratio (assuming equal expression and stability of the proteins), whereas both in vivo approaches were based on overexpression and hence assayed with an excess of Su(H). S2 cells lack Notch but express both Su(H) and Hairless [13,34,35]; and the endogenous levels of Su(H) are sufficient for a strong response to experimental ICN doses [13,14,36]. Addition of Su(H) enhances the ICN response nearly fourfold [10,14,36], similar to the in vivo situation, where overexpression of Su(H) elicits Notch gain of function phenotypes [12,22,37]. We have no information on the amount of Su(H) in a cell that is freely available for binding to either Notch or Hairless, and the above observations suggest that Su(H) is limiting. Clearly, Su(H) occupancy on Notch target gene promoters is highly dynamic and enhanced by the presence of ICN [35]. However, Drosophila cells express high levels of Su(H) in the cytoplasm which is rather unconventional for a transcription factor [22,38,39]. The mechanisms underlying Su(H) nuclear import/export are little understood and may involve Notch signalling and repression, respectively [38–41]. Most likely
cytoplasmic Su(H) is unavailable for transcriptional complex formation, be it repressor or activator complex. Moreover, Su(H) from the cytoplasm may resupply the nucleus once Su(H) is bound in complexes such that a steady level of free nuclear Su(H) is achieved. Overexpression of Su(H) may raise this level considerably, explaining the increase of Notch output in the cell culture as well as in fly tissue. Hairless would access the same free pool of Su(H), however, only bind to wild type protein to build up repressor complexes and silence Notch target genes, whereas Notch could access both mutant and wild type Su(H). Assuming a large enough pool of wild type Su(H), then repressor complex formation might in fact be as efficient in the presence of mutant as of wild type overexpressed Su(H) (Fig. 8). This rather speculative model would be in agreement with an in vivo reduced or lack of binding between Hairless and Su(H)WARE and concomitant repression by a complex consisting primarily of Hairless and endogenous Su(H). Experiments addressing the exact composition of the repressor complexes, i.e. the presence of mutant Su(H), may help support this model.

Materials and Methods

Yeast two-hybrid experiments

Single, double or triple missense mutations in CTD were introduced using the QuickChange II XL site-directed mutagenesis kit (Stratagene). All mutants were sequence verified (StarSeq, Mainz). CTD mutants were cloned into pJG vector [42] and tested for protein interactions with pEG constructs as described previously [10]. Primer sequences are available upon request. The yeast three-hybrid experiments were performed with the N-ANK domain cloned in the pESC-Leu vector (Stratagene) and pEG-MamN as outlined in [10].

The CTDWARE mutant DNA was excised with MscI and EcoRI from the CTDWARE pJG-construct and reintroduced in likewise digested Su(H) cDNA to generate the mutant full length construct Su(H)WARE. It was shuttled into pRmHa-3 [43] and pUAST-attB- vectors [15] for subsequent in vivo analyses.

Electro-mobility shift assays - EMSA

DNA binding assays of Su(H) and Su(H) mutants were performed according to standard protocols using a double stranded DNA-oligomer (made by hybridization of primers 5'GGT TCT TTC AGC TCG GTT CCC ACG CCA CGA GCC AC 3' and 5'TTG GGT GGC TCG TGG CGT GGG AAC CGA GCT GAA AG 3' and labelled with Klenow polymerase) containing the E(spl)m8-S1 Su(H) binding site [44] and Su(H) proteins produced from cDNA by in vitro transcription/translation using the TNT coupled Reticulocyte Lysate System (Promega).

Cell culture assays

For cell culture experiments Drosophila Schneider S2 cells, obtained from the Drosophila Genomics Resource Centre DGRC (Indiana University, Bloomington USA), were transfected with the respective constructs and the activity measured with a Notch responsive luciferase reporter (NRE-reporter) as described previously [13]. Renilla expression plasmid (tk-Renilla; Promega) was cotransfected as internal control. Reporter activation elicited by transfection with pMT-ICN was taken as 100% [45]. Cotransfection with Su(H) and Hairless constructs were analysed as described before [10]. CuSO₄ was used to induce protein expression 6 h after transfection. Luciferase activity was measured 18 h later in duplicate (Lumat LB 9507, EG & Salem, MA) using the dual-luciferase reporter assay system (Promega).

Analysis of mutant Hairless and Su(H) transgenes in vivo

Transgenic Su(H)WARE flies were generated with the PhiC31 integrase-based integration system [15] to avoid position effects and allow for a direct comparison with likewise integrated wild type Su(H) [10]. For co-overexpression experiments, the Su(H)-
Figure 8. Model of a Su(H) cytoplasmic pool. A) The large cytoplasmic pool of Su(H) is not available for complex formation in the nucleus, however, is used to replenish nuclear Su(H). This explains the apparent limitation of Su(H) when ICN is overexpressed. The export mechanism is currently unknown and may represent a further level of Su(H) regulation. B) Overexpression of Su(H) increases the cytoplasmic protein pool, thereby raising the level of available nuclear Su(H). This may cause Notch gain of function phenotypes in tissues, where Notch signalling takes place, or in the presence of exogenous ICN. The darker colour represents mutant Su(H) protein to exemplify the model; wild type Su(H) would behave identical. C) Combined overexpression of Su(H) and Hairless allows formation of repressor complex, as long as endogenous wild type Su(H) is available from the cytoplasmic pool. Lack of binding of Hairless to mutant Su(H) predicts an enrichment of activator complexes containing the mutant Su(H) and ICN. doi:10.1371/journal.pone.0027986.g008

Acknowledgments

We acknowledge S. Artavanis-Tsakonas, S. Bray and C. Delidakis for sharing DNA-constructs and F. Karch for sending the Phil3 fly lines and vectors. We thank A. Schulz and I. Wech for invaluable technical help.

Author Contributions

Conceived and designed the experiments: AP RAK DM. Performed the experiments: PK DM. Analyzed the data: PK AP RAK DM. Contributed reagents/materials/analysis tools: PK AP RAK DM. Drafted and revised the manuscript: AP RAK DM. Critically read the article: PK.
19. Del Álamo Rodríguez DA, Terriente J, Galindo ML, Cosuo JP, Díaz-Benjumea FJ (2008) Different mechanisms initiate and maintain wingless expression in the Drosophila wing hinge. Development 129: 3995–4004.

20. Perea D, Terriente J, Díaz-Benjumea FJ (2009) Temporal and spatial windows delimit activation of the outer ring of wingless in the Drosophila wing. Dev Biol 328: 445–453.

21. Terriente J, Perea D, Suzanne M, Díaz-Benjumea FJ (2008) The Drosophila gene zfh2 is required to establish proximo-distal domains in the wing. Dev Biol 320: 102–112.

22. Furriols M, Bray S (2006) Dissecting the mechanisms of Suppressor of Hairless function. Dev Biol 292: 520–532.

23. Voas MG, Rebay I (2004) Signal integration during development: insights from the Drosophila eye. Dev Dyn 229: 162–175.

24. Hay BA, Maile R, Rubin GM (1997) P element insertion-dependent gene activation in the Drosophila eye. Proc Natl Acad Sci U S A 94: 5195–5200.

25. Müller D, Nagel AC, Preiss A (2006) A molecular link between Hairless and Pros26.4, a member of AAA-ATPase subunits of the proteasome 19S regulatory particle in Drosophila. J Cell Sci 119: 250–258.

26. Müller D, Kugler SJ, Preiss A, Maier D, Nagel AC (2005) Genetic modifier screens on Drosophila. Genetics 168: 441–449.

27. Müller D, Kugler SJ, Preiss A, Maier D, Nagel AC (2005) Genetic modifier screens on Hairless gain-of-function phenotypes reveal genes involved in cell differentiation, cell growth and apoptosis in Drosophila. Genetics 171: 1157–1152.

28. Nagel AC, Preiss A (2001) Fine tuning of Notch signaling by differential co-repressor recruitment during eye development of Drosophila. Hereditas 140: 77–84.

29. Prozter CE, Wech I, Nagel AC (2008) Hairless induces cell death by downregulation of EGFR signaling activity. J Cell Sci 121: 3167–3176.

30. Wech I, Nagel AC (2005) Mutations in zfh2 promote cell-type specific apoptosis in the Drosophila eye. Cell Death Differ 12: 145–152.

31. Yu SY, Yoo SJ, Yang L, Zapata C, Srinivasan A, et al. (2002) A pathway of signals regulating effector and initiation caspases in the developing Drosophila eye. Proc Natl Acad Sci U S A 99: 15480–15485.

32. Friedmann DR, Wilson JJ, Kovall RA (2008) RAM-induced allostery facilitates assembly of a Notch pathway active transcription complex. J Biol Chem 283: 14781–14791.

33. Koval RA (2008) More complicated than it looks: assembly of Notch pathway transcription complexes. Oncogene 28: 5099–5109.

34. Felton RG, Kooh PJ, Rebay I, Regan CL, Xin T, et al. (1990) Molecular interactions between the protein products of the neurogenic loci, Notch and Delta, two EGF-homologous genes in Drosophila. Cell 61: 523–534.

35. Krejci A, Bray SJ (2007) Notch activation stimulates transient and selective binding of SuH/CsL to target enhancers. Genes Dev 21: 1322–1327.

36. Eastman DS, Sée R, Skufoš E, Bangalore L, Bray S, et al. (1997) Synergy between Suppressor of Hairless and Notch in regulation of Enhancer of split new split and midsplit expression. Mol Cell Biol 17: 5629–5638.

37. Schweiguth F, Posakony JW (1994) Antagonistic activities of Suppressor of Hairless and Hairless control alternative cell fates in the Drosophila adult epidermis. Development 120: 1433–1441.

38. Fortini ME, Artavanis-Tsakonas S (1994) The Suppressor of Hairless protein participates in Notch receptor signaling. Cell 79: 273–282.

39. Gho M, Lecourtois M, Gerault G, Posakony JW, Schweiguth F (1996) Subcellular localization of Suppressor of Hairless in Drosophila sense organ cells during Notch signalling. Development 122: 1673–1682.

40. Frise E, Knoblich JA, Younger-Shepherd S, Jan LY, Jan YN (1996) The Drosophila Numb protein inhibits signaling of the Notch receptor during cell-cell interaction in the sensory organ lineage. Proc Natl Acad Sci U S A 93: 11925–11932.

41. Maier D, Nagel AC, Johannes R, Preiss A (1999) Subcellular localization of Hairless protein shows a major focus of activity within the nucleus. Mech Dev 89: 195–199.

42. Gyuris J, Golemis E, Chertkov E, Brent R (1993) Gdi1, a human G1 and S phase protein phosphatase that associates with cdk2. Cell 75: 791–803.

43. Bunch TA, Grinblat Y, Goldstein LS (1998) Characterization and use of the Drosophila metallothionein promoter in cultured Drosophila melanogaster cells. Nucleic Acids Res 16: 1043–1061.

44. Bailey AM, Posakony JW (1995) Suppressor of Hairless directly activates transcription of Enhancer of split Complex genes in response to Notch receptor activity. Genes Dev 9: 2609–2622.

45. Matsuno K, Diederich RJ, Go MJ, Blaumueller CM, Artavanis-Tsakonas S (1995) Deltaex acts as a positive regulator of Notch signaling through interactions with the Notch ankyrin repeats. Development 121: 2633–2644.

46. Maier D, Nagel AC, Preiss A (2002) Two isoforms of the Notch antagonist Hairless are produced by differential translation initiation. Proc Natl Acad Sci U S A 99: 15480–15485.