Mef2 Interacts with the Notch Pathway during Adult Muscle Development in *Drosophila melanogaster*

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Abstract

Myogenesis of indirect flight muscles (IFMs) in *Drosophila melanogaster* follows a well-defined cellular developmental scheme. During embryogenesis, a set of cells, the Adult Muscle Precursors (AMPs), are specified. These cells will become proliferating myoblasts during the larval stages which will then give rise to the adult IFMs. Although the cellular aspect of this developmental process is well studied, the molecular biology behind the different stages is still under investigation. In particular, the interactions required during the transition from proliferating myoblasts to differentiated myoblasts ready to fuse to the muscle fiber. It has been previously shown that the Notch pathway is active in proliferating myoblasts, and that this pathway is inhibited in developing muscle fibers. Furthermore, the Myocyte Enhancing Factor 2 (Mef2), Vestigial (Vg) and Scalloped (Sd) transcription factors are necessary for IFM development and that Vg is required for Notch pathway repression in differentiating fibers. Here we examine the interactions between Notch and Mef2 and mechanisms by which the Notch pathway is inhibited during differentiation. We show that Mef2 is capable of inhibiting the Notch pathway in non-myogenic cells. A previous screen for Mef2 potential targets identified Delta a component of the Notch pathway. Dl is expressed in Mef2 and Sd-positive developing fibers. Our results show that Mef2 and possibly Sd regulate a Dl enhancer specifically expressed in the developing IFMs and that Mef2 is required for Dl expression in developing IFMs.

Introduction

Muscle development is a complex program that is evolutionarily well conserved. Muscle precursor cells are specified, then proliferate and fuse to form multinucleated myotubes which give rise to differentiated muscle fibers. The cellular changes that characterize this process have been well described whereas the molecular aspects have yet to be completely elucidated. *Drosophila melanogaster* has been shown to be a good model for understanding myogenesis, especially considering the conservation of the molecular aspects have yet to be completely elucidated. *Drosophila melanogaster* follows a well-defined cellular developmental scheme. During embryogenesis, a set of cells, the Adult Muscle Precursors (AMPs), are specified. These cells will become proliferating myoblasts during the larval stages which will then give rise to the adult IFMs. Although the cellular aspect of this developmental process is well studied, the molecular biology behind the different stages is still under investigation. In particular, the interactions required during the transition from proliferating myoblasts to differentiated myoblasts ready to fuse to the muscle fiber. It has been previously shown that the Notch pathway is active in proliferating myoblasts, and that this pathway is inhibited in developing muscle fibers. Furthermore, the Myocyte Enhancing Factor 2 (Mef2), Vestigial (Vg) and Scalloped (Sd) transcription factors are necessary for IFM development and that Vg is required for Notch pathway repression in differentiating fibers. Here we examine the interactions between Notch and Mef2 and mechanisms by which the Notch pathway is inhibited during differentiation. We show that Mef2 is capable of inhibiting the Notch pathway in non-myogenic cells. A previous screen for Mef2 potential targets identified Delta a component of the Notch pathway. Dl is expressed in Mef2 and Sd-positive developing fibers. Our results show that Mef2 and possibly Sd regulate a Dl enhancer specifically expressed in the developing IFMs and that Mef2 is required for Dl expression in developing IFMs.

Two waves of myogenesis have been described in *Drosophila*: the first occurs during embryogenesis to give rise to the larval muscles (for review, see [3,4]), the second takes place during pupariation and leads to the adult muscles (for review, see [4]). A particular set of adult muscle structures, the Indirect Flight Muscles (IFMs) provide a valuable tissue context to study muscle development. IFMs are composed of 7 dorsal ventral muscles (DVMs) and 6 dorsal longitudinal muscles (DLMs) per hemithorax [5]. They develop between 8 and 36 hours after puparium (APF) from adult muscle precursor cells (AMPs) named myoblasts that have been specified during embryogenesis and have proliferated on the wing imaginal disc during larval stages [6,7]. Interestingly, whereas DVMs are formed *de novo* during pupariation by myoblast fusion, DLMs are formed through fusion of myoblasts to larval scaffolds that escape histolysis [6].

The maintenance of the AMPs during the larval stages in a proliferative state requires the expression of the Twist bHLH transcription factor (Twi) and the activation of the Notch pathway. Twi has been primarily described for its role during mesoderm development in the embryo [8]. It is activated by the Notch pathway in AMPs and acts as an anti-differentiation signal (Figure S1, A) [9]. Interestingly, Twi activates the transcription of the Myocyte Enhancer Factor 2 gene (*Mef2*) in AMPs [10]. *Mef2* is a transcription factor essential for cardiac, visceral and somatic muscle development in the *Drosophila* embryo [11,12]. Furthermore, in the adult fly, *Mef2* mutants show severe defects in IFM differentiation [13,14] and *Mef2* overexpression in AMPs induces early differentiation, suggesting that *Mef2* is the major differentiation factor in IFM development [15]. Consistent with this, *Mef2* expression levels increase throughout IFM development, starting in the AMPs and reaching its maximal levels in the differentiating fibers [16]. Thus, the situation may appear paradoxical as Twi, the main anti-differentiation factor, activates the transcription of *Mef2*, the main pro-differentiation factor. In fact, while *Mef2* is expressed and *Mef2* protein is present in AMPs, its transcriptional activity is repressed. Indeed, Twi and Notch together activate the
**Holes in muscle** gene (Him) that encodes a repressor of Mef2 transcriptional activity [17–19]. Therefore, maintenance of AMPs in an undifferentiated state requires a subtle equilibrium directed by Notch between Mef2 quantity and its transcriptional activity. This predicts that differentiation could be triggered by a change in this equilibrium, either by over-activating Mef2 or by inhibiting the Notch pathway.

The Notch pathway is an evolutionarily conserved intercellular signaling pathway involved in numerous developmental processes such as cell-fate determination, neural development, and tissue homeostasis [20,21]. In *Drosophila*, a cell that expresses one of the pathway transmembrane ligands, Delta (Dl) or Serrate (Ser), signals an adjacent cell expressing the transmembrane receptor Notch. Notch affinity for Dl and Ser is modulated by the glycosyltransferase Fringe (Fng): glycosylation of Notch increases its affinity for Dl and decreases its affinity for Ser [22–24]. The interaction of Notch with its ligands induces two proteolytic cleavages of Notch leading to the release of its intracellular domain. This predicts that differentiation could be triggered by a change in an undifferentiated state requires a subtle equilibrium directed by Notch between Mef2 quantity and its transcriptional activity. The Notch pathway must be repressed by Vg. 

Regulation of the Notch pathway in differentiating IFMs is sufficient to activate Delta (Dl) or Serrate (Ser), with its cofactor Scalloped (Sd). Sd can bind DNA but does not possess a transactivation domain [33–35]. Sd-Vg dimer plays an important role during wing development [31,36–38]. Indeed, sd is expressed in AMPs during larval stages and vg is expressed in a subset of AMPs during embryogenesis and larval stages; they are both involved in muscular identity [38–40]. Moreover, in a *vg*null context, IFM differentiation is severely impaired, Notch is ectopically activated in developing fibers and the product is devoid of a DNA binding domain and is capable of activating transcription of its targets when associated with its cofactor Scalloped (Sd). Sd-Vg dimer plays an important role during wing, neural and muscle development [31,36–38]. Indeed, sd is expressed in AMPs during larval stages and vg is expressed in a subset of AMPs during embryogenesis and larval stages; they are both involved in muscular identity [38–40]. Moreover, in a *vg*null context, IFM differentiation is severely impaired, Notch is ectopically activated in developing fibers and vg expression is lost in IFMs [31,39]. Thus, in order for IFM differentiation to proceed, the anti-differentiation role of the Notch pathway must be repressed by Vg.

Altogether, data show that Sd, Vg and Mef2 are involved in IFM development. Moreover, Sd and Vg can both interact with Mef2 and Mef2 synergizes with Sd to activate a differentiation specific enhancer of vg [38,41]. Interestingly, mammalian orthologs of Sd and Mef2 interact to activate muscle-specific enhancers, suggesting that the roles of Sd/Mef2 during myogenesis could be conserved between different species [42,43]. In this study, we were interested in the mechanisms triggering Notch inhibition. If Vg is required, it is not sufficient and cannot be the triggering signal since it is expressed in AMPs where Notch is active [44]. We therefore focused on Mef2 for two reasons: i) Mef2 is capable of inducing differentiation [15], ii) Mef2 levels increase during development and its transcriptional targets depend on its activity level [16,19,45]. Here we show that Mef2 can repress the Notch pathway in various contexts. We propose that an aspect of this inhibition requires transcriptional regulation of Delta, a component of the Notch pathway. Indeed, we identified and characterized a 2.64 kb enhancer of Delta containing Mef2 and Sd binding sites and show that Mef2 is implicated in Delta regulation in the developing IFM muscle fibers. These results are consistent with the idea that Mef2 could repress the Notch pathway through Dl regulation.

**Materials and Methods**

**Fly stocks**

The following strains were used in this study: *UAS-Mef2* [11], *UAS-sd* [33], *1151-Gal4* [9], *UAS-H2B::YPF* [16], *UAS-Mef2[RNAi]* [47], *UAS-NDN* [48], *new*722-Gal4* [46], *sd111*, [49], *sf08* [49], patched-Gal4 [Bloomington #217]. The *UAS-H2B::YPF; tub-Gal80*, *neutralized*722-Gal4*/SM5CyO, TM6Tb* line was donated by Michel Ghigo.

**Over-expression in bristle cell lineage**

*UAS-H2B::YPF; tub-Gal80*, *neutralized*722-Gal4*/SM5CyO, TM6Tb* males were crossed to *UAS-Mef2, UAS-Mef2-UAS-sd, or UAS-NDN* females at 18°C. White pupae (0 hours after pupa formation, APF) were transferred to 29°C to allow GAL4 activity.

**Immunostaining and antibodies**

Anti-Twi and anti-Mef2 were generously donated by S. Roth, and B. Paterson respectively, and used at 1:200 and 1:5000 dilutions respectively. Chicken anti-GFP (1:1000 dilution) antibodies were purchased from AvesLab (Tigard, USA). Mouse anti-Cut, anti-Wingless, anti-Prospero and 22c10 were purchased from the Developmental Studies Hybridoma Bank, and were used at 1:200 (DSHB, University of Iowa, Department of Biology, Iowa City, IA 52242). Anti-Sens [50] was generously donated by H. Bellen and used at a 1:3000 dilution. Anti-Vg (1:200 dilution) was described in Goulev et al. [51]. Fluorescent-conjugated secondary antibodies were purchased either from Molecular Probes (Carlsbad, USA) or Jackson ImmunoResearch (West Grove, USA) and used at a 1:200 dilution. When needed, DAPI (Sigma-Aldrich, Saint-Louis, USA) was added with the secondary antibodies at 1 µg/ml concentration.

Pupae dissection was performed as previously described in Fernandes et al. [6]. Preparations were observed with a Zeiss 710 or Leica SP5 confocal microscope. Adult thoraces were observed with a Keyence VHX-2000 microscope.

**Plasmid constructions and transgenesis**

The *Dl2.6* sequence was amplified by PCR of genomic *Drosophila melanogaster* Canton S strain DNA (Oligonucleotides: D497_5_a2 5’CACTGGCGTATGCGACATCC3’ and D49 7_6_a2 5’ACAAGGCTTCAAGATTCCC’ and cloned into the pGEM-T easy vector (Promega, Madison, USA). *Dl2.6* was subcloned i) into the pGL3 plasmid with *EcoR1* and *Xho1* restriction enzymes (*Dl2.6-luc*) and ii) into the transgenic plasmid pGreen-H-Pelican [52] with *Kpn1-BglII* and *Kpn1-XhoI* restriction enzymes respectively (pGreen-H-Pelican-Dl2.6). pCasPer-hsp70-Mef2 and pCasPer-hsp70-sd plasmids are described in Bernard et al. [41]. Transgenesis was performed by BestGene (Chino Hills, USA).

**Cell culture experiments and transfection assays**

*Drosophila* S2 cells were maintained at 22°C in standard Schneider medium with fetal bovine serum (FBS 10% v/v) and antibiotics (streptomycin 100 µg/ml and penicillin 100 U/ml). *Drosophila* Dmd8 cells were also maintained at 22°C in standard Schneider medium with fetal bovine serum (FBS 5% v/v), antibiotics (streptomycin 100 µg/ml and penicillin 100 U/ml) and insulin (1%). Transfection assays were performed with Effectene reagent (QIAGEN, Valencia, USA) according to the manufacturer’s instructions. The Effectene: DNA ratio was 10:1.
and that Sd and Mef2 synergize to activate a differentiative pathway [53]. In each experiment, the mean and standard deviation were calculated on six independent transfection assays. Means were compared using the Student’s t-test.

Q-PCR experiments

Transfection assays were carried out in Dmd8 cells using 500 ng of each expression vector (pCasPer-hsp70-Mef2 or sd) adjusted to 1 μg with empty pCasPer-hsp70 vector. RNA extractions were done using the RNeasy kit (Qiagen, Valencia, USA). 100 ng of RNA from each extraction were retrotranscribed using the SuperScript III VILO (Life Technologies, Paisley, UK) according to the manufacturer’s protocol. Quantitative RT-PCR amplification mixture (10 μl) contained 1/20 of cDNA product, 10X SYBR Green 1 Master Mix buffer and 100 ng forward and reverse primers. The expression of Dl and C was quantified relative to rp49 housekeeping gene [54]. Each point was repeated three times. Primers for rp49 amplification were: forward primer 5’-CCGGCTTCAAGGACAGTGATCTG-, reverse primer 5’-CAGGTTGTGCAACCGAGACTT-. Dl amplification was carried out with: forward primer 5’-CCAGGCTCTTTGGTCACGAC-, reverse primer 5’-TGTTGCGCTTGATGTGCTTATTAG-. Reactions were run on a Light Cycler PCR machine (Roche). Cycle condition were 10 min at 95°C and 45 cycles at 95°C for 10 sec, 60°C for 10 sec. Quantification was done using the mathematical model described in Pfaffl et al. [55].

Results

The published data have demonstrated that vg and sd are expressed throughout IFM development, yet Vg-Sd is required to initiate Notch pathway repression at the DV boundary. However, data showed that Mef2 can interact with Sd during myogenesis to activate muscle genes or enhancers [38,41]. As Sd is expressed in the wing pouch [63], we cannot exclude that Notch repression at the DV boundary is due to interactions between Mef2 and endogenous Sd.

Mef2 can repress the Notch pathway at the DV boundary of the wing disc

Our first goal was to test whether Mef2 by itself can inhibit the Notch pathway. However, it has been shown that Mef2 overexpression in myogenic cells induces premature muscle differentiation [19], hence making it difficult to see whether Notch pathway repression could be the consequence of Mef2 expression or the consequence of the myogenic differentiation program. To circumvent this problem, we ectopically misexpressed Mef2 in non-myogenic cells in which the Notch pathway is activated. The Notch pathway is active in a number of developmental processes and has different transcriptional outputs depending on the cellular context. These outputs can be used to determine the activity of the pathway [25,26,29,30,61]. At the dorsal-ventral (DV) boundary of the wing imaginal disc, the Notch signaling induces cut (ct) expression [62]. We asked if Mef2 could repress the Notch pathway in the wing disc, by testing if its misexpression can prevent expression of ct. Normally (in ptc-gal4; UAS-GFP flies), Ct is expressed along the DV boundary (Fig. 1A-C). Mis-expression of Mef2 along the AP boundary using the ptc-gal4 driver (Fig. 1E), results in loss of Ct expression in the cells that ectopically express Mef2 (Fig. 1D, overlay in 1F). Two other Notch targets, Vg and Wingless (Wg) were tested and again a clear decrease in their expression was observed in Mef2 misexpressing cells (Fig. S2). Thus, Mef2 represses Notch pathway activity in non-myogenic cells at the DV boundary of the wing disc. However, data showed that Mef2 can interact with Sd during myogenesis to activate muscle genes or enhancers [38,41]. As Sd is expressed in the wing pouch [63], we cannot exclude that Notch repression at the DV boundary is due to interactions between Mef2 and endogenous Sd.
This cell undergoes a first asymmetric division to give rise to the pIIa cell in which the Notch pathway is active (N\textsuperscript{\Delta N}) and the pIIb cell where it is not (N\textsuperscript{\textnormal{null}}; Figure S3). The pIIb cell expresses Prospero (Pros) whereas the pIIa does not [63,66].

Asymmetric division of the pIIa cell then gives rise to the shaft and socket cells where the Notch pathway is only active in the socket cell. The pIIb cell undergoes an asymmetric division that gives a glial cell and the pIIb cell with an active Notch pathway. Finally, the pIIb cell divides asymmetrically to give rise to the sheath cell in which the Notch pathway is active and the neuron.

In a wild-type adult thorax, small bristles (microchaetes) are present (Fig. 2A, A’, arrows), showing a specific distribution pattern, aligned along the antero-posterior axis (Fig. 2A, dotted lines). Misexpression of a dominant-negative allele of Notch (N\textsuperscript{\Delta N}) between 0h and 30h APF in sensory organ precursors (SOPs) (Fig. 2B, arrows). When Sd and Mef2 were misexpressed alone using the same driver, no Notch phenotype was observed even if some bristles seemed shorter: external cells are present, bristles are still organized along the antero-posterior axis (Fig. 2A, dotted lines). Misexpression of a dominant-negative allele of Notch (NDN\textsuperscript{-}) or a dominant-negative allele of Notch (Dl\textsuperscript{-}) leads to a strongly disorganized microchaete arrangement (Fig. 3B, B’). Patches with no external cells (shaft or socket) are observed (Fig. 2B, B’, asterisks).

In some cases, duplications of the shaft are seen (Fig. 2B, B’; arrowheads). However, some microchaetes look normal, showing that the N\textsuperscript{\Delta N} allele is not fully penetrant (Fig. 2B, arrows). When sd or Mef2 were misexpressed alone using the same driver, no Notch phenotype was observed even if some bristles seemed shorter: external cells are present, bristles are still organized along the antero-posterior axis, and no shaft duplications were observed (Fig. 2C, C’, D, D’, dotted lines). In contrast, when sd and Mef2 were simultaneously misexpressed, no external cells were observed (Fig. 2E, E’), consistent with the idea that either the pII cell is not specified or that the Notch pathway is inhibited after the pII cell specification, giving rise to two pIIb cells after cell division (see Figure S3). To test these two hypotheses, we looked at the developing nota at 21h APF. At this time point, in wild type flies (neur-Gal4; UAS-H2B::YFP genetic context) SOPs are already specified (Fig. 3B) and express the transcription factor Senseless, a marker of the SOP lineage (Sens, Fig. 3C, [50]). Moreover, as expected, in two cell clusters (Fig. 3A-D, arrowheads), one cell corresponding to the pIIb expresses Pros (Fig. 3D, arrowhead b) and the other cell corresponding to pIIa does not (3D, arrowhead a). When Sd and Mef2 are misexpressed (Fig. 3E-H) SOPs are specified and express Sens (Fig. 3F-G). In two cell clusters, the two cells express Pros, signifying that they all adopt a pIIb fate (Fig. 3H, arrowhead b). This phenotype is observed in Notch loss of function genetic background [66]. Therefore, when Sd and Mef2 are together misexpressed in SOPs, the Notch pathway is inhibited, at least during the first asymmetric division, and this repression is not due to a change in cell fate. This inhibition during first asymmetric division is consistent with phenotypes observed in adults. However, Mef2 or Sd alone overexpression do not show Notch-like adult phenotypes, suggesting that Sd and Mef2 are required to inhibit the Notch pathway.

Identification of a putative Sd/Mef2 response element in the Delta

Our data show that Mef2 and probably Sd interact with the Notch pathway. As Sd/Mef2 acts as a transcription factor, we wondered whether Sd/Mef2 could directly activate genes involved in the Notch pathway. A previous study identified 670 genomic regions bound by Mef2 [67]. We decided to resequence these 670 regions for clusters of Sd and Mef2 binding sites using the Cluster-Buster software (Cluster Buster, http://zlab.bu.edu/cluster-buster/). This led us to identify two overlapping genomic regions associated to the Notch pathway ligand Dl gene (clones D497_5_a2 and D497_6_a2; mef2 ChIP data http://furlonglab.

embl.de/). As Dl is expressed during IFM development [31] in the differentiating fibers, this makes it a potential Sd/Mef2 target.

**DI2.6 activation profile in vivo**

In order to ask whether DI could be regulated by Sd and/or Mef2, we generated transgenic fly lines in which 2640 bp DI genomic fragment (DI2.6) was used to drive GFP expression. DI2.6 sequence spans the two clones identified to be bound by Mef2 (see above) and containing 2 Sd and 9 Mef2 predicted binding sites (Figure S4).

**DI2.6-GFP expression starts in developing fibers at 16 h APF (Fig. 4A-D). Activation persists in fibers at 24 h APF (Fig. 4E-H) and 30 h APF (Fig. 4I-L). No activation is detected in the swimming myoblasts around the fibers at 24 h APF (Fig. 4E-H, arrowheads) or during late differentiation stages from 36 h APF (Fig. 4M-P). In conclusion, the DI2.6 enhancer drives the GFP expression in developing DLMs during the early stages of their development (16 h–30 h APF). We next decided to test whether the DI2.6 enhancer is activated by Mef2 and Sd. When Mef2 levels are decreased by overexpressing an RNAi-Mef2 transgene in 21 h APF pupae (1151-Gal4, UAS-RNAi-Mef2), DI2.6 activation is lower than in wild-type genetic context (Figure 3F). This result shows that Mef2 is required in vivo for DI2.6 activation. As null sd mutants are early lethal, we decided to test DI2.6 activation in sd\textsuperscript{111} and sd\textsuperscript{81L} pupal lethal alleles [49]. We observed a significant activation of DI2.6 in these genetic contexts (Figure S5) implying that either that Sd is not required or that these mutant alleles are not strong enough to interfere with DI2.6 activation.

**Regulation of DI2.6 in vitro**

In order to examine its regulation, we cloned DI2.6 in a luciferase reporter vector to measure its activity in cell culture experiments. We used two Drosophila cell lines to analyze enhancer activation by Sd and or Mef2, as both have putative binding sites located in the enhancer. The first is the classical Schneider 2 (S2) cell line, derived from a primary culture of embryonic Drosophila cells [68]. The second is the Dm86 cell line, derived from myoblasts located on the wing disc in the third instar larval stage of Drosophila development [17,69,70]. We co-transfected the DI2.6-luc reporter plasmid with various combinations of Mef2 and Sd expression vectors (see Material and Methods) and quantified the luciferase activity. In S2 cells, the DI2.6 enhancer is not activated when Sd is transfected alone (Fig. 6A; Table S1). When Mef2 is overexpressed, we observe a significant activation (Fig. 6A, p = 0.003; Table S1). This activation is further substantially increased when Sd and Mef2 are overexpressed together (Fig. 6A, p = 8×10\textsuperscript{-4} compared to control, p = 3×10\textsuperscript{-3} compared to Mef2 overexpression; Table S1). In the Dm86 cell line, the DI2.6 enhancer is also not activated by Sd alone. In contrast, it is significantly activated by Mef2 alone (Fig. 6B, p = 7.6×10\textsuperscript{-6}; Table S2). When both Sd and Mef2 are present, this activation is significantly decreased relative to Mef2 alone (Fig. 6B, p = 4.3×10\textsuperscript{-6}; Table S2), but still significantly higher than the baseline expression (Fig. 6B; compared to empty plasmids p = 2.3×10\textsuperscript{-4}; Table S2).

Our data show that Mef2 activates the DI2.6 enhancer and that, in S2 cells, Mef2 synergizes with Sd for this activation. However, the level of activation by Sd/Mef2 in S2 cells is close to the level of activation by Mef2 alone in Dm86 cells. We know that Dm86 cells derive from 3\textsuperscript{rd} instar wing disc AMPs and are therefore likely to express sd [69]. Thus, this high activation is possibly due to interactions between endogenous Sd and transfected Mef2. A possible explanation for the reduced activation when Sd and Mef2 are together transfected in Dm86 cells could be a “squelching”
phenomenon [71]. Accordingly, in S2 cells where there is no Sd or Mef2, co-transfection of Mef2 and Sd shows a significantly increased activation compared to Mef2 alone.

We next determined whether this activation of the Dl2.6 enhancer by Sd and Mef2 is associated with an increase of endogenous Dl mRNA levels. We quantified Dl mRNAs by Q-RT-PCR in Dmd8 cell culture experiments. When Dmd8 cells are transfected by Mef2 and sd expressing plasmids, we observed a significant increase in Dl mRNA levels (2.32 fold, 5% confidence interval: [1.66–3.23]; 1.84-fold, 5% confident interval: [1.10–3.06]; Table S3). Together with ex vivo enhancer regulation, this suggests that Dl is regulated by Mef2 and possibly together with Sd via the Dl2.6 enhancer.

Sd and Mef2 activate Dl in SOPs when they are both misexpressed

Our ex vivo data show that Sd and Mef2 synergize to activate the Dl2.6 enhancer in S2 cells and that Sd and Mef2 overexpression in Dmd8 cells induces a increase in Dl mRNA levels. Therefore, to test whether Sd and Mef2 could induce Dl overexpression in vivo, we overexpressed Sd and Mef2 in SOPs using the neur-Gal4 driver (Fig. 7). In controls (neur-Gal4; UAS-H2B::YFP), SOP clusters express H2B::YFP (Fig. 7B, arrowheads) and Dl immunoreactivity is homogeneously present at low levels in SOPs (Fig. 7C, arrowheads) and non-SOP cells (Fig. 7C, asterisk). When Sd and Mef2 are ectopically misexpressed in SOPs (neur-}

Figure 2. Notch repression by Sd and Mef2 in SOPs. A’–E’ are magnifications of A–E. In wild type thoraxes, bristles (A, A’ arrows) are aligned along the antero-posterior axis (A’, dashed lines). When a dominant negative allele of Notch is expressed using the neur-Gal4 driver, different Notch phenotypes are observed on thoraxes, spanning from wild-type bristles (B, B’, arrows), to complete absence of external cells, shaft and socket (B, B’, asterisks). Duplicated bristles resulting from Notch inhibition during the pI/pII or pII/pIII asymmetric divisions are also present (B’, arrowheads). When sd (C, C’) or Mef2 (D, D’) are expressed alone, no Notch phenotypes are observed: bristles are present and align along the antero-posterior axis (C’, D’, dotted line). When sd and Mef2 are expressed together, no external cells are observed, corresponding to a strong Notch phenotype (E, E’).

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Figure 3. The Notch pathway is repressed during the first asymmetric division of SOPs upon Sd/Mef2 ectopic overexpression. YFP (A–D) or YFP, sd, and Mef2 (E–H) were overexpressed using the neur-Gal4 driver. In wild type genetic context, cells expressing the YFP (B) express Sens, a marker of the SOP lineage, detected with a specific antibody (C). In two cell clusters (A–D, arrowheads), only one cell corresponding to the pilb cell expresses Pros (D, arrowhead b). The second cell corresponding to the pilla cell does not express Pros (D, arrowhead a). When Sd, Mef2 and YFP are co-overexpressed, cells expressing YFP (F), as in wild type, express Sens (G) showing that these cells maintain their SOP identity. The cells of the two cell clusters (E–H, arrowheads) both express Pros (H, arrowhead b), signifying that they correspond to two pilb cells. A and E correspond to DAPI staining.

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Gal4, UAS-H2B::YFP, UAS-sd, UAS-Mef2 genetic background; Fig. 7E-H), Dl expression is increased in SOP clusters (Fig. 7G, arrows; SOP clusters are indicated by arrowheads). Therefore, we conclude that Sd and Mef2 can activate Dl when they are together misexpressed.

Mef2 is required for Dl expression in developing DLMs

We showed that Mef2 is required for Dl activation in developing fibers (Fig. 5) and that Mef2 can activate Dl in cultured cells (Fig. 6) and in SOPs when misexpressed with Sd (Fig. 7). To test whether Mef2 is required for Dl expression in 24 h APF developing fibers, we decided to knock down Mef2 expression using RNA interference. We drove UAS-RNAi-Mef2 transgene with 1151-Gal4 which is expressed in myoblasts and developing fibers [9]. In 24 h APF pupae, developing fibers can be detected using a DAPI staining: they exhibit a specific alignment of the nuclei (Fig. 8A, D, m). In wild-type pupae, Dl is expressed in fibers (Fig. 8B, B', m). In a previous publication, we detected Dl only in developing fibers [31]. However, we think that this difference is due to technical issues. Indeed, Dl expression has been recently observed in myoblasts [72] and our Dl labeling is very similar to Dl labeling in epithelial cells [73]. In 1151-Gal4, UAS-RNAi-Mef2 genetic context (Fig. 8D-F, magnification in 8D', F', m). Dl expression in myoblasts seems close to Dl expression in controls (Fig. 8E', m compared to 8B', m). In contrast, it is clearly reduced in developing fibers (Fig. 8E, asterisks compared to control 8B, asterisks; Fig. 8E', asterisks, compared to control 8B', asterisks). Therefore, we concluded that Mef2 is required for Dl expression in developing fibers, but not in the swarming myoblasts.

Also, occasional large nuclei corresponding to the larval template nuclei can be observed (Fig. 8A', n). Cells between the developing fibers correspond to myoblasts (Fig. 8A', F', m). In wild-type pupae, Dl is expressed in fibers (Fig. 8B, B', asterisks) and in surrounding myoblasts (Fig. 8B, B', m). In a previous publication, we detected Dl only in developing fibers [31]. However, we think that this difference is due to technical issues. Indeed, Dl expression has been recently observed in myoblasts [72] and our Dl labeling is very similar to Dl labeling in epithelial cells [73]. In 1151-Gal4, UAS-RNAi-Mef2 genetic context (Fig. 8D-F, magnification in 8D', F', m). Dl expression in myoblasts seems close to Dl expression in controls (Fig. 8E', m compared to 8B', m). In contrast, it is clearly reduced in developing fibers (Fig. 8E, asterisks compared to control 8B, asterisks; Fig. 8E', asterisks, compared to control 8B', asterisks). Therefore, we concluded that Mef2 is required for Dl expression in developing fibers, but not in the swarming myoblasts.

Figure 4. Activation of the Dl2.6 enhancer in vivo monitored by the expression of the Dl2.6-GFP transgene. E–H' are magnifications of the dotted squares in E–H. In 16 h, 24 h, 30 h and 36 h APF pupae, the 22c10 antibody labels muscles and nerves (B, F, F', J and N respectively), Dl enhancer is activated in developing fibers of 16 h, 24 h and 30 h APF pupae (C, G, G', K respectively) but not in myoblasts at 16 h APF (A–D) and 24 h APF (E–F, magnification in E'–H', arrowheads). Dl enhancer is not activated in developing fibers of 36 h APF pupa (M–P). DAPI-GFP overlay is shown in D, H, and H', L and P. Scale bar: 100 μm.

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Discussion

In this paper, we explored the relationship between the Mef2 myogenic factor and the Notch pathway during muscle fiber development. Previous studies have shown that the Notch pathway activation is required in swarming myoblasts i) to express the anti-differentiation factor Twi [9] and ii) together with Twi, to activate specific targets, such as *Holes in muscle* (Him, [17]). Moreover, it has been shown that persistent activation of the Notch pathway in developing muscles induces muscle degeneration [9]. Together, these data show that the Notch pathway is needed in myoblasts but must be repressed in developing fibers. In myoblasts, Notch interacts with Mef2: it cooperates with Twi to activate Him [17] which is a repressor of Mef2 activity [18,19]. Moreover, muscle degenerations observed when Notch is ectopically activated in fibers is due to persistent activation of Him and inhibition of Mef2 activity [19]. Thus, the Notch pathway is activated and Mef2 activity is repressed in undifferentiated myoblasts whereas the Notch pathway is repressed and Mef2 is active in differentiating fibers. To understand how the Notch pathway is repressed in

![Figure 5. Reduced activation of the *Dl2.6* enhancer *in vivo* monitored by the expression of the *Dl2.6-GFP* transgene.](image)

In 21 h APF pupae, muscle fibers are visualized with DAPI staining by the specific arrangement of their nuclei (A, D, asterisks). In a wild-type pupae, *Dl2.6* enhancer is activated in developing fibers (B, overlay in C). In pupae overexpressing an RNAi-Mef2 transgene, *Dl2.6* enhancer activation is significantly lower than in wild-type (E, overlay in F).

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![Figure 6. Activation of the *Dl2.6* enhancer in S2 and Dmd8 cells by Sd and Mef2.](image)

A: In S2 cells, the *Dl2.6* enhancer was not activated by Sd alone. In contrast, it was significantly activated by Mef2 (*p* = 0.0003) and Sd/Mef2 (*p* = 0.0008). Activation with Sd/Mef2 was significantly higher than activation by Mef2 alone (*p* = 0.003). B: In Dmd8 cells, as in S2 cells, *Dl2.6* is not activated by Sd alone. It is significantly activated by Mef2 (*p* = 7.6 × 10⁻⁶) and Sd/Mef2 (*p* = 4.3 × 10⁻⁶). However, activation by Sd/Mef2 was lower than activation by Mef2 alone (*p* = 4.3 × 10⁻⁶).

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Figure 7. Activation of Dl by Sd/Mef2 in 28 h APF pupa SOPs. In SOPs, visualized with YFP (green) expressed under the control of the neur-Gal4 driver (arrowheads in A–D), Dl (red) is detectable at the same levels as in non-SOP cells (asterisks in A–D) of the notum. When YFP, sd and Mef2 are overexpressed in SOPs using the same driver (E–H), Dl is up-regulated in SOPs (arrows in G) relative to non-SOP cells (asterisks in G) of the notum. DAPI labeling is shown in gray (A,E). Overlays are shown in D and H.

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Figure 8. Expression of Dl in IFMs of 24 h APF pupae. Expression of Dl was detected with an anti-Dl antibody in wild-type B, B’–C,C’) and 1151-Gal4, UAS-RNAiMef2 (E, E’–F,F’) 21 h APF pupae. A’–F’ show magnified areas indicated by the dotted squares in A–F respectively. Muscles are visualized with DAPI staining by the specific arrangement of their nuclei (asterisks in A, D, A’, D’; muscles are located between the two dotted lines in A’ and D’). Large nucleus (n in A’) is a larval template nucleus. Myoblasts are located between muscles (m in A’, D’). In wild type pupae, Dl is detected in myoblasts and developing fibers (B, B’; overlay in C, C’). Expression levels in myoblasts and fibers are close (B’). Overexpression of a RNAi-Mef2 construct in myoblasts and fibers using the 1151-Gal4 driver (D–F, D’–F’) lowers Dl expression in developing fibers compared to myoblasts (E’, compare m to asterisks) and in 1151-Gal4, UAS-RNAiMef2 muscles compared to wild type muscles (compare E’, asterisks to B’, asterisk).

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Mef2, likely together with Sd, inhibits the Notch pathway

To investigate Mef2 capacity to inhibit the Notch pathway, we ectopically expressed Mef2 in two non-myogenic contexts. Indeed, it has been shown that Mef2 overexpression induces early muscle differentiation [15] and thus Notch repression could be due to the differentiation context rather than to Mef2 expression. Our experiments in the wing disc showed that Mef2 overexpression inhibits activation of Notch known targets at the wing margin (et, vg, vgo; Fig. 1, Figure S1). However, when Mef2 was ectopically expressed in SOPs, we did not observe classical Notch loss of function phenotypes (duplication of the shaft, absence of external cells, [75]). A strong Notch-like phenotype, with no external cells, was only observed when Mef2 and Sd were co-overexpressed (Fig. 2). Moreover, we show that Sd and Mef2 overexpression inhibits the Notch pathway during the first asymmetric division of SOPs (Fig. 3). It seems therefore that inhibition of the Notch pathway by Mef2 in SOPs requires Sd. Differences between these results is likely due to differences in sd endogenous expression: whereas sd is expressed in the wing pouch [63], we did not detect any sd expression in SOPs (data not shown). Thus, it is likely that in the wing pouch, ectopic Mef2 interacts with endogenous Sd and therefore Sd is always required together with Mef2 to inhibit the Notch pathway. This result is consistent with previous studies: Mef2 and Sd can interact physically in Drosophila [38], Mef2 and Sd are required for IFM differentiation [41, 76] and Mef2 and Sd cooperate to activate a differentiation specific enhancer in IFMs [41]. Moreover, it has been shown that mammalian orthologs of sd and Mef2 can activate muscle-specific promoters [43].

Mef2 activates Dl through a specific enhancer

We next wanted to determine how the Notch pathway is inhibited by Mef2 and possibly Sd. Since Mef2 and Sd are transcription factors, we hypothesized that they inhibit the pathway by regulating target genes. A previous study isolated 670 cis-regulatory modules (CRMs) bound by Mef2 during embryogenesis [67]. We therefore adopted a bioinformatics approach to rescreen these 670 CRMs for clusters of Sd/Mef2 binding sites, postulating that Sd/Mef2 targets during embryonic myogenesis are also Sd/Mef2 targets during IFM development. We isolated a 2.6 kb sequence, Dl2.6, which was predicted to be a Dl enhancer (see supplementary materials in [67]). Using a transgenic approach, we showed that Dl2.6 is activated in IFMs between 16 h and 36 h APF (Fig. 4) where Sd and Mef2 activate the vestigial adult muscle enhancer [41].

We show that Mef2 can activate Dl2.6 in myogenic and non-myogenic cultured cells and that Mef2 in vivo knock-down induces a decrease in Dl2.6 activation as well as Dl expression in muscle fibers (Fig. 5, 8). Interestingly, when Mef2 is down-regulated in the entire IFM myogenic lineage, a significant Dl decrease is observed in developing IFMs whereas Dl expression in myoblasts is not significantly changed (myoblasts and fibers; Fig. 8). This shows that Mef2 is required for Dl expression only in developing IFMs, likely through the activation of the Dl2.6 enhancer. We tried to test whether Sd is involved in Dl2.6 activation and Dl expression. In vivo experiments did not allow us to conclude on this question. Indeed in sd11L and sd1152 mutants, Dl2.6 is activated. However, these alleles are not as early lethal as sd null alleles and are likely to retain a part of Sd activity [49]. Therefore we cannot exclude that this remnant activity is sufficient to activate the Dl2.6 enhancer. Ex vivo experiments seemed to be more conclusive: Sd and Mef2 synergize to activate Dl2.6 in S2 cells (Fig. 6A). On the other hand an inhibitory effect of Sd is observed in Dmef2 cells (Fig. 6B). This inhibition may be due to a squelching effect [71]. Such phenomenon has been previously described with the Sd/Vg transcription factor. Indeed, the SRF-A enhancer is activated by Sd and Vg, but activation by Sd/Vg diminished with increasing Sd concentrations [34]. This is interpreted as a titration of the Vg factor by Sd: when Sd is in excess, Sd-Vg dimers are replaced on promoters by Sd monomers. In the same way, an excess of Sd could titrate the Mef2 factor and therefore diminish the activation of the Dl2.6 enhancer. Thus, considering i) cultured cell results, ii) that Sd-Mef2 ectopic expression in SOPs induces Dl and iii) that Sd-Mef2 overexpression in Dmef2 cells induces an increase in Dl mRNA levels, we favored the hypothesis that Sd is also involved in Dl2.6 activation and Dl expression. However, further studies are needed to conclude on this point.

Role of Mef2 and Dl in Notch inhibition

Together these data show that Mef2 interacts with the Notch pathway during IFM differentiation, at least by activating Dl. However, if Dl can participate in mechanisms involved in Notch inhibition, it is not the only factor responsible for it. Indeed, Dl is expressed in myoblasts (Fig. 8) were the Notch pathway is activated [9], suggesting that Dl activation by Mef2 in developing fibers is a mechanism for maintaining Dl expression in the myogenic lineage. Moreover, Dl expression is not sufficient to inactivate the pathway: Dl over-activation in SOPs or myoblasts using the neu-Gal4 or 1151-Gal4 drivers respectively failed to reproduce Mef2 overexpression or Notch phenotypes (data not shown). This last observation is similar to previously published results of Dl overexpression in SOPs [77].

In a previous study, we have shown that vg is also required for Notch inhibition in developing DLMs and thus for a proper muscle differentiation. We showed that in absence of vg, fng is not expressed in DLMs and that forcing fng expression in the vg mutant is sufficient to rescue the muscle phenotype [31]. fng encodes a glycosyltransferase that modifies the Notch receptor, increasing its affinity for its ligand Dl [22–24]. In wild-type flies, it is expressed in developing fibers but not in myoblasts [31]. It seemed therefore that the modifications of Notch by Fng, and thus the modification of Notch affinity for Dl, is a critical step in Notch repression in developing DLMs and therefore for DLM differentiation. Here we show that Mef2 can inhibit the Notch pathway and that Mef2 is responsible for Dl expression in developing fibers. Therefore, Notch inhibition is associated with fng and Dl expression in developing fibers. These results may appear contradictory as Dl activates the Notch pathway. However, Dl is expressed in the signal sending cell. Moreover, in the wing disc, the Notch pathway is inactive in cells expressing Notch and Dl. Indeed, a mechanism, called cis-inhibition, involving the Notch receptor and its ligands in the repression of the pathway has been previously described [for review, see [78]]. In the wing pouch of the wing imaginal disc, Notch is ubiquitously expressed, Dl is only expressed in the ventral part and fng is only expressed in the dorsal part. When Dl overexpressing clones are induced in the dorsal part, the Notch pathway is activated at the boundary of the clones, but not in the clonal cells where Dl and fng are expressed.
[79]. A similar result is observed when fng overexpressing clones are induced in the ventral part [80]. Thus, in cells expressing N, Dl and fng, the Notch pathway is inactivated. Therefore, activation of Dl by Mef2 in developing fibers, where fng is expressed, could be necessary for Notch inactivation through a cis-inhibition mechanism. Interestingly, fng is also expressed in the wing disc [81] and SOPs (data not shown) where Mef2 ectopic expression inhibits the Notch pathway.

Since fng is required for Notch inhibition and Mef2 inhibits the Notch pathway, we also asked whether Mef2 can activate fng as well. However, using the fng<sup>SM12-LacZ</sup> transgene as a reporter, we never observed any activation of fng by Mef2. Moreover, we did not find Sd/Mef2 binding site clusters in and around the fng gene. However, we showed that Sd and Mef2 can activate a muscle specific enhancer of vg [41] and that Vg is required for fng expression [31]. Therefore, even if fng is not directly activated by Sd/Mef2, it is possible that Sd and Mef2 may be involved in fng activation through vg activation.

### Triggering differentiation: a switch in Mef2-Notch interactions?

The Notch pathway is active in myoblasts [9] where it activates Him, a repressor of Mef2 transcriptional activity [18,19]. In contrast, in developing IFMs, Notch is not active [9]. Him is not expressed [19] and therefore Mef2 is active [76]. It has been shown that Mef2 activates different target genes according to its activity level [16,45] and that Mef2 overexpression in myoblasts induces early differentiation [15]. Soler et al. proposed that a balance of Him and Mef2 activities could regulate muscle differentiation: high levels of Him repress Mef2 activity and consequently differentiation, while high levels of Mef2 overwhelm repression by Him and induce muscle differentiation [19]. Here we propose that one effect of Mef2 high activity levels could be to trigger the repression of the Notch pathway, switching from a stable undifferentiated state in which the Notch pathway represses Mef2 activity through Him (Figure S1, A) to a stable differentiating state in which Mef2 activity remains high through a positive feedback loop and represses the Notch pathway (Figure S1, B). This inactivation could occur through a cis-inhibition mechanism, since Mef2 activates Dl in developing fibers.

### Supporting Information

#### Figure S1

A: in AMPs, the Notch pathway activates twi (1) that inhibits muscle differentiation (2). The pro-differentiation gene Mef2 is activated by Twi (4) but the transcriptional activity of Mef2 is repressed by Him (5), a target of Twi and Notch (3). B: in differentiating fibers, Notch is not active and therefore Him and Twi are absent. Mef2, for which levels increase due to a positive feedback, is transcriptionally active and triggers muscle differentiation. (TIF)

#### Figure S2

Notch repression by Mef2 in the wing disc. Panels B and H show Vg and Wg expression patterns in wild type third instar larva wing disc (DAPI in A, G). When Mef2 is overexpressed along the AP boundary of the third instar larva wing disc using the ptc-Gal4 driver (D, J), neither Vg (E, arrowhead) or Wg (K, arrowheads), two known targets of the Notch pathway, are detected where Mef2 is ectopically expressed (overlay in F, L, DAPI in C, I). (TIF)

#### Figure S3

Schematic representation of SOP development. Notch pathway activity is represented by red outlining. Prospero expression is represented in blue. A: In normal developmental circumstances, SOP microchaete development starts by specification of the pI cell by lateral inhibition. At 17 h APF, this cell will undergo an asymmetrical division to give rise to the pIIa and pIIb cells. The pIIa will give rise to the external shaft and socket cells. The pIIb cell will divide asymmetrically to give rise to the pIIIa cell and a glial cell that will degenerate. The pIIb cell gives rise to the internal cells of the sensory organ, the shaft cell and the neuron. Prospero staining at 21 h APF (purple rectangle) reveals one prospero positive cell in which the Notch pathway is not active. B: Theoretically, Notch pathway repression after pl specification and throughout SOP development should induce two pIIb cells after pl division and only glial cells after plIIb division. (TIF)

#### Figure S4

Schematic representation of the Dl2.6 enhancer (3R: 15,142,950...15,145,589). Predicted Mef2 binding sites are shown as yellow arrowheads. Predicted Sd binding sites are shown as gray arrowheads. (TIF)

#### Table S1

Raw data for Dl2.6 activity in S2 cells. (XLSX)

#### Table S2

Raw data for Dl2.6 activity in Dmd8 cells. (XLSX)

#### Table S3

Q-RT-PCR were performed on cells transfected by an empty pCasPer vector or on cells transfected by pCasPer-hsp70-Mef2 and pCasPer-hsp70-sd vectors. Each point was repeated 3 times (see material and methods; Casper1-3, SDMEF1-3). dCp (Cp RP49 - Cp Dl) was calculated to normalize data according to the RP49 housekeeping gene expression. The mean<sup>1</sup> and the confidence interval<sup>2</sup> were calculated in each condition. dddCp<sup>3</sup> was obtained subtracting dCp (SDMEF condition) from dCp (Casper condition). Confidence interval<sup>4</sup> is calculated adding or removing the confidence interval in<sup>2</sup>. Increase<sup>5</sup> and the confidence interval<sup>6</sup> are obtained by the formula: Increase = 2<sup>ddCp</sup>. (Pfaffl, 2001). (XLSX)

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Author Contributions
Conceived and designed the experiments: CC AL JS. Performed the experiments: CC PK AL. Analyzed the data: CC AL. Contributed reagents/materials/analysis tools: AL JS. Wrote the paper: CC AL.

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