

# SETDB1 modulates the TGF $\beta$ response in Duchenne muscular dystrophy myotubes

## Authors

Alice GRANADOS<sup>1</sup>, Maeva ZAMPERONI<sup>1</sup>, Roberta RAPONE<sup>1,2</sup>, Maryline MOULIN<sup>1</sup>, Ekaterina BOYARCHUK<sup>1</sup>, Costas BOUYIOUKOS<sup>1</sup>, Laurence DEL MAESTRO<sup>1</sup>, Véronique JOLIOT<sup>1</sup>, Elisa NEGRONI<sup>3</sup>, Myriame MOHAMED<sup>1</sup>, Sandra PIQUET<sup>1</sup>, Anne BIGOT<sup>3</sup>, Fabien LE GRAND<sup>4</sup>, Sonia ALBINI<sup>1,5,\*</sup>, Slimane AIT-SI-ALI<sup>1\*</sup>

## Affiliations

<sup>1</sup>Université Paris Cité, CNRS, Epigenetics and Cell Fate, UMR7216, F-75013 Paris, France

<sup>2</sup>Current affiliation: Institut de Biologie de l'Ecole Normale Supérieure (IBENS), Ecole Normale Supérieure, CNRS, INSERM, PSL Research University, 75005 Paris, France

<sup>3</sup>Sorbonne Université, Inserm, Institut de Myologie, Centre de Recherche en Myologie, Paris, France

<sup>4</sup>Université Claude Bernard Lyon 1, CNRS UMR 5261, INSERM U1315, Institut NeuroMyoGène, Pathophysiology and Genetics of Neuron and Muscle (PGNM) Unit 69008 Lyon, France

<sup>5</sup>Current affiliation: Genethon, 91000, Evry, France; Université Paris-Saclay, Univ Evry, Inserm, Génomique, Integrare research unit UMR\_S951, 91000, Evry-Courcouronnes, France

*In memory of our collaborator and friend Armando Felsani*

## Correspondence

\*[slimane.aitsiali@u-paris.fr](mailto:slimane.aitsiali@u-paris.fr); [salbini@genethon.fr](mailto:salbini@genethon.fr)

## Key results

- TGF $\beta$  induces nuclear accumulation of SETDB1 in healthy myotubes
- SETDB1 is enriched in DMD myotube nuclei with intrinsic TGF $\beta$  pathway overactivation
- SETDB1 LOF in DMD myotubes attenuates TGF $\beta$ -induced pro-fibrotic response
- Secretome of TGF $\beta$ -treated DMD myotubes with SETDB1 LOF is less deleterious on myoblast differentiation

## SUMMARY

Overactivation of the TGF $\beta$  signaling in Duchenne muscular dystrophy (DMD) is a major hallmark of disease progression, leading to fibrosis and muscle dysfunction. Here, we investigated the role of SETDB1, a histone lysine methyltransferase involved in muscle differentiation. Our data show that, following TGF $\beta$  induction, SETDB1 accumulates in the nuclei of healthy myotubes, while being already present in the nuclei of DMD myotubes where TGF $\beta$  signaling is constitutively activated. Interestingly, transcriptomics revealed that depletion of SETDB1 in DMD myotubes leads to downregulation of TGF $\beta$ -target genes coding for secreted factors involved in extracellular matrix remodeling and inflammation. Consequently, SETDB1 silencing in DMD myotubes abrogates the deleterious effect of their secretome on myoblast differentiation by impairing myoblast pro-fibrotic response. Our findings indicate that SETDB1 potentiates the TGF $\beta$ -driven fibrotic response in DMD muscles, providing a new axis for therapeutic intervention.

## INTRODUCTION

Histone lysine methylation/demethylation is a major epigenetic mechanism that controls chromatin states (1), with lysine 9 of histone 3 trimethylation (H3K9me3) being a hallmark of repressed chromatin (2). SETDB1 (SET Domain, Bifurcated 1, also called KMT1E, or ESET in the mouse) (3) is one of the H3K9 SUV39 KMT family. SETDB1 regulates many cellular states, such as stemness and terminal differentiation, including skeletal muscle terminal differentiation (1, 3). Although SETDB1 is mainly nuclear in proliferating myoblasts, where it represses terminal differentiation genes through H3K9me3 deposition, we have uncovered an original mechanism whereby murine Setdb1 is exported to the cytoplasm of differentiating myoblasts in a Wnt-mediated manner, allowing the de-repression of late differentiation genes and the formation of mature multinucleated myotubes (4). SETDB1 is associated with many human diseases including inflammatory bowel disease (IBD) (5), many cancer types (6), neuropsychiatric, genetic and cardiovascular diseases (3).

Dystrophinopathies are a spectrum of muscle genetic diseases caused by alterations in the *Dystrophin* gene, and Duchenne muscular dystrophy (DMD) is the most severe form, caused by total absence of the coded protein (7, 8). Dystrophin is essential for the maintenance of muscle membrane integrity by providing a strong mechanical link between the actin cytoskeleton and the extracellular matrix (ECM) (9, 10). Thus, the lack of functional Dystrophin is deleterious to membrane integrity, at the origin of the continuous degeneration of skeletal muscles (11), leading to exhaustion of muscle stem cells (MuSCs), progressive muscle wasting, chronic inflammation and severe fibrosis (i.e., accumulation of ECM components such as Collagens), which is the most prominent feature of DMD. Fibrosis is defined as pathological wound healing with an abnormal deposition of ECM resulting in the replacement of functional tissue by fibrotic connective tissue, which leads to an alteration of tissue and organ homeostasis (12, 13). Fibrosis involves intracellular and extracellular soluble effectors also called secretome, including pro- and anti-inflammatory cytokines, growth factors and ECM remodelers such as TGF $\beta$ , Matrix metalloproteinases (MMP)/Tissue Inhibitors of MMPs (TIMPs) and Collagens (14). The main pathway that sustains this excessive pro-fibrotic response is the TGF $\beta$  pathway, which is overactivated in DMD (15, 16).

In DMD patients, TGF $\beta$  levels, monitored by the nuclear accumulation of phosphorylated SMAD2/3 (pSMAD2/3) transcriptional modulators, are elevated in both plasma and muscle, and were shown to contribute to pathological fibrosis (17). However, the signaling pathways contributing to human DMD fibrosis, and the signals coordinating diverse cell types to maintain or restore muscle tissue homeostasis, remain largely unknown. Yet, TGF $\beta$  pathway overactivation is the common feature observed in Dystrophin-deficient myotubes *in vitro* independently of the type of mutation (15, 16) and exacerbated TGF $\beta$ /SMAD pathway has been shown to lead to fusion defects in *in vitro* DMD myotubes. Moreover, TGF $\beta$  pathway has been described as a molecular brake of myoblast fusion

and, thus, muscle regeneration, while its inhibition leads to the formation of giant myofibers *in vivo* (18).

SETDB1 has been shown to regulate the TGF $\beta$  response in cancer and in T-cells (19-22) and pulmonary fibrosis (23) contexts. Here we investigated the SETDB1/TGF $\beta$  interplay in a model of DMD myotubes derived from hiPSCs or immortalized myoblasts (15, 24).

We show that SETDB1 contributes to the deregulated TGF $\beta$  signaling in DMD, indicating that SETDB1 may participate in muscle dysfunction and increased fibrosis. Indeed, TGF $\beta$  induces nuclear accumulation of SETDB1 in healthy myotubes, while SETDB1 is constantly accumulated in DMD myotube nuclei with intrinsic overactivated TGF $\beta$  pathway. Moreover, transcriptomics showed that SETDB1 silencing attenuates the TGF $\beta$ -dependent pro-fibrotic and anti-differentiation response in DMD myotubes. Interestingly, many TGF $\beta$ /SETDB1-dependent genes code for secreted proteins involved in ECM remodeling and inflammation. Conditioned medium assays show that TGF $\beta$ -treated myotubes with SETDB1 LOF produce a secretome which reduces the negative impact of TGF $\beta$  response on myoblast differentiation and impairs myoblast pro-fibrotic response. Our findings point to a role of SETDB1 in the fine-tuning of the TGF $\beta$  response in muscles, which is deregulated in DMD, participating in fibrosis.

## RESULTS

### SETDB1 relocalizes in the nuclei of healthy differentiated human myotubes in response to TGF $\beta$ pathway activation

All our experiments were conducted in human models generated from either human induced pluripotent stem cells (hiPSCs, which we have already described (15), or from immortalized myoblasts derived from healthy or DMD patients (24). hiPSCs can be directly induced to differentiate into myotubes using a transgene based epigenetic reprogramming strategy (15).

We have already showed that in murine myoblasts Setdb1 is both cytoplasmic and nuclear, and during terminal muscular differentiation Setdb1 is exported to cytoplasm in a Wnt-dependent manner, allowing de-repression of muscle genes (4). Thus, we have first investigated, and quantified (as depicted in **Figure S1A**), the subcellular localization of SETDB1 in proliferating *versus* differentiating healthy human myoblasts, and in myotubes in response to TGF $\beta$ /SMAD pathway activation (**Figure 1A**). TGF $\beta$ /SMAD pathway is constitutively activated in human proliferating myoblasts while nuclear phospho-SMAD3 (p-SMAD3) levels decrease upon differentiation (**Figures 1B, S1B**), as already shown in murine muscle cells (18). Differentiated myotubes are still responsive to TGF $\beta$ /SMAD pathway activation since p-SMAD3 nuclear levels are restored upon TGF $\beta$ 1

treatment (**Figure 1F, S1B**). TGF $\beta$  treatment efficiency was evidenced by the activation of many TGF $\beta$  target genes, including *TGF $\beta$*  gene itself, concomitant to a decrease in many differentiation genes (**Figure S1D**).

We have confirmed in human muscle cells that SETDB1 is also highly enriched in the nuclei of proliferating myoblasts while it is depleted in nuclei of differentiated myotubes (**Figures 1B, 1C**), as in murine cells (4). Interestingly, TGF $\beta$ /SMADs pathway activation in muscle cells is correlated with SETDB1 presence in nuclei (**Figures 1B, 1C**) and we were able to trigger SETDB1 nuclear enrichment in differentiated myotubes after TGF $\beta$ 1 treatment (**Figures 1B, 1C**), in absence of significant changes in SETDB1 mRNA (**Figure 1D**) and protein levels (**Figures 1E, S1C**), neither during terminal differentiation nor in response to 24h-TGF $\beta$ 1 treatment. This result was confirmed in our healthy human hiPSCs-derived model (15) of muscle cells (**Figure S2C** upper panels). We next used the ALK4/5/7 inhibitor SB-431542 in myotubes to block TGF $\beta$  pathway activation prior to TGF $\beta$ 1 addition (**Figure 1G**). The SB-431542 efficiency was evidenced by its effect on phospho-SMAD3 nuclear levels and TGF $\beta$  target and *MYOD1* gene expression (**Figures 1H, 1J, S1E**). Autocrine TGF $\beta$  signal blocking does not change SETDB1 localization in myotubes in basal condition but blocks SETDB1 nuclear relocalization in healthy myotubes in response to TGF $\beta$ 1 (**Figures 1H, 1I**).

Altogether, these data showed that SETDB1 is excluded to the cytoplasm upon terminal differentiation of human myoblasts, and TGF $\beta$  pathway activation is directly responsible for SETDB1 nuclear relocalization in myotube nuclei.

### **SETDB1 is accumulated in DMD myotube nuclei in TGF $\beta$ -dependent manner**

We next investigated SETDB1 level and subcellular localization in the context of DMD in which the TGF $\beta$  pathway is overactivated both *in vivo* (17, 25) and in *in vitro* cellular models (15, 16).

First, immunohistochemistry assays show that SETDB1 co-localizes with phospho-SMAD3 in the centrally localized nuclei in DMD patient muscle histological sections (**Figure 2A**). To further confirm this observation in a more quantitative manner, we efficiently differentiated three healthy myoblasts or hiPSCs lines and three DMD lines carrying different *Dystrophin* mutations (myoblasts with exons 45-52 deletion (**Figure 2B**), myoblasts with point mutation in exon 59 (**Figure S2B**), and hiPSCs with exon 45 deletion (**Figure S2C**). Compared to healthy differentiated myotubes, DMD myotubes display overactivated TGF $\beta$ /SMAD pathway, as depicted by *TGF $\beta$ 1* mRNA expression (**Figure S2A**) and nuclear phospho-SMAD3 protein level (**Figures 2D, 2F**, and (15, 16)), as expected.

The quantification of SETDB1 nuclear/cytoplasmic ratio in healthy myotubes shows that SETDB1 translocated in the nuclei after 6h of TGF $\beta$ 1 treatment and this relocalization tends to decrease after 24h (**Figure 2B-C, S2A-B**). Interestingly, in DMD myotubes, SETDB1 is accumulated in the nuclei at steady state and is persistent in response to TGF $\beta$  pathway activation (DMD exon 45 deletion in myoblasts or hiPSCs, **Figures 2B, 2C**; DMD point mutation, **Figure S2B**) and is associated with a higher activation of *TGF $\beta$ 1* expression (**Figure S2A**). The effect of TGF $\beta$  pathway activation on the subcellular localization of SETDB1 is further confirmed by the western blotting of nuclear and cytoplasmic fractions of healthy and DMD myotubes (**Figures 2D, 2E, 2F**) and also in our hiPSC-derived DMD myotubes (exon 45 deletion, **Figure S2C**). We also blocked the autocrine signal of TGF $\beta$  in DMD myotubes with SB-431542 and found that it leads to decreased SETDB1 levels in nuclei with and without TGF $\beta$ 1 treatment (**Figure 2G**). Overall, these data showed an effect of TGF $\beta$  on SETDB1 nuclear relocalization in at least 3 different DMD patient myotubes with different *Dystrophin* gene mutations.

To elucidate the molecular mechanisms underlying SETDB1 relocalization in DMD myotubes, we first tested if SETDB1 nuclear translocation in response to TGF $\beta$  was dependent on SMAD3, which is the effector translocating to the nucleus in response to TGF $\beta$ . Our results show that SETDB1 nuclear enrichment in response to TGF $\beta$  is not dependent on the presence of SMAD3 protein (**Figure S2D-F**).

SETDB1 is known to bear numerous post-translational modifications including many phosphorylations (<https://www.phosphosite.org/proteinAction.action?id=9498&showAllSites=true>). We thus hypothesized that the phosphorylation status of SETDB1 might correlate with its subcellular localization. To check this, we tested the migration profile of SETDB1 in several subcellular compartments. Our data showed that SETDB1 from the cytosolic and peri-nuclear fractions migrates higher compared to nuclear SETDB1 (**Figure S2G**), indicating possible hyperphosphorylation in these compartments. However, these observations require further investigation.

Taken together, these data showed that TGF $\beta$  induces SETDB1 nuclear enrichment in normal and DMD myotubes, regardless of the genetic background. This suggested that SETDB1 could modulate the cellular response to TGF $\beta$ .

### **SETDB1 loss-of-function attenuates the TGF $\beta$ response in human DMD myotubes**

To investigate the role of SETDB1 in the TGF $\beta$  response, we performed SETDB1 loss-of-function (LOF) in both healthy and DMD myotubes treated or not with TGF $\beta$ 1 (as depicted in **Figure 3A**). While the siRNA mediated SETDB1 knockdown (KD, more than 80%) was efficient (**Figures 3B, S3A**), it did not show any effect on the levels of SMAD2/3 proteins nor their phosphorylation levels (**Figure 3B**). However, SETDB1 LOF led to a decrease in the TGF $\beta$  response, as depicted by



the expression of *ad hoc* TGF $\beta$  target genes involved in pro-fibrotic response and inflammation, such as *TGF $\beta$ 1*, *TIMP1* (*Tissue Inhibitor of Metalloproteinases 1*), *IL6* (*Interleukin 6*), *FN1* (*Fibronectin 1*) and *CTGF* (*Connective Tissue Growth Factor*) (**Figure 3C, S3C**). Note that SETDB1 KD alone, in absence of TGF $\beta$ , does not affect or only slightly the expression of these genes (**Figure S3D-E**). Since these TGF $\beta$  target genes are downregulated in SETDB1 KD condition, we hypothesized that in normal condition SETDB1 could target and repress inhibitors of the TGF $\beta$  pathway. A gene candidate approach showed that, indeed, SETDB1 LOF led to an increase in *SKIL* and *SMAD7* genes only in DMD (**Figures 3D, S3B**), two known inhibitors and fine tuners of the TGF $\beta$  pathway, while in absence of TGF $\beta$ , SETDB1 KD alone does not affect or only slightly the expression of these genes (**Figures S3D, S3F**). Of note, concomitant to the attenuation of the TGF $\beta$  response, SETDB1 LOF in 3-days differentiated myotubes induced a significant increase in muscle early differentiation marker *Myogenin* and regeneration marker *MYH3* (*embryonic Myosin Heavy Chain*), but not in the muscle late differentiation marker *MYH1* (**Figures 3E, S3B**). Of note, SETDB1 KD alone also showed an effect on the expression of muscle differentiation genes (**Figure 3E, S3G**).

Thus, SETDB1 LOF attenuates the TGF $\beta$ 1-induced fibrotic response while promoting regeneration and could be beneficial in DMD patients.

### SETDB1 alters the TGF $\beta$ regulated secretome in human DMD myotubes

To deepen the role of SETDB1 in the TGF $\beta$  response at the global level, we performed RNA-seq comparing DMD myotubes with healthy myotubes in response to TGF $\beta$ , with or without SETDB1 LOF. Principal component analysis (PCA) of gene expression levels showed that the triplicates and duplicates were distinctively separated and that the main source of variability between samples are the cell line source, healthy *versus* DMD. The second source seems to be the effect of the TGF $\beta$  treatment and SETDB1 silencing (**Figure S4A**).

To track changes in gene expression, first in response to TGF $\beta$ , we performed differential gene expression analysis (**Figure 4A**). We identified 1043 TGF $\beta$ -dependent differentially expressed genes (DEGs) in healthy myotubes (489 up- and 554 down-regulated genes) and only 480 in DMD myotubes (363 up- and 117 down-regulated genes) (Top genes in **Table S1, S2**). These data showed that while the TGF $\beta$  pathway is already intrinsically activated in DMD myotubes it is not saturated and can still be further activated *in vitro* (see also **Figure 2 & 3**, and (15)). Interestingly, only TGF $\beta$ -dependent 184 TGF $\beta$ -dependent genes (corresponding to 18% of healthy and 39% of DMD DEGs) are commonly deregulated in healthy and DMD conditions (**Figure 4A**).

Gene ontology (GO) analysis of the TGF $\beta$ -responsive genes in healthy myotubes showed a significant enrichment in terms related to the regulation of MAPK cascade, negative regulation of locomotion, cellular component movement, cell adhesion, SMAD protein phosphorylation, migration,

motility, chemotaxis, vessel and nervous development (**Figure S4B**). While in DMD myotubes, GO showed that the main categories are related to cytokine-mediated signaling, extra cellular matrix (ECM) remodeling and organization, inflammation, SMAD protein phosphorylation and positive regulation of cell motility (**Figures 4B, S4C**), reminiscent of disease traits. Collectively, these data validated the responsiveness of both healthy and DMD myotubes to the TGF $\beta$  treatment.

Then, we have checked the DEGs in SETDB1 LOF condition, both in healthy and DMD myotubes treated with TGF $\beta$ . We found that SETDB1 LOF affected 320 genes in response to TGF $\beta$  in healthy myotubes (**Figure 4A**, top genes on **Table S3**). To determine the signature of the transcriptional changes, we applied gene set enrichment analysis (GSEA) and found that the SETDB1-dependent genes belong to ECM and cell surface categories (**Figure S4D**). Together, these data already suggest that SETDB1 LOF affects genes coding for secreted factors.

In DMD myotubes, SETDB1 LOF affected 212 genes in response to TGF $\beta$  (**Figure 4A, 4C**, top genes on **Table S4**), 120 were less abundant and 92 more abundant. In particular, SETDB1 LOF induced less abundant mRNAs associated with ECM remodeling, inflammation and fibrosis, including *THBS1* (*Thrombospondin-1*), *Myostatin*, *BMPR2* (*Bone Morphogenetic Protein Receptor type 2*), *ADAMTS8* (*ADAM Metalloproteinase with Thrombospondin type 1 motif 8*), *MMP14* (*Matrix Metalloproteinase 14*), *LIF* (*Leukemia Inhibitory Factor*), *SERPINE1* (*Plasminogen Activator Inhibitor Type 1 or PAI1*), *WNT5A*, and *MAMDC2* (*MAM domain containing 2*) (**Figures 4C, 4F, S4E**). All these genes, gene categories and pathways are notably involved in the DMD disease traits. Furthermore, enrichment analysis for biological processes and gene network of biological pathway analyses highlighted that the SETDB1-dependent genetic programs in DMD myotubes in response to TGF $\beta$  are mainly involved in inflammation, signaling and cell surface (**Figure 4D-E**). SETDB1 KD alone, in absence of TGF $\beta$ , affects at a lesser extent the expression of the tested genes (**Figure S3D and S3H**).

Altogether, here we validated the efficacy of TGF $\beta$  to induce fibrotic and inflammation genetic programs and highlighted that SETDB1 silencing alters the transcription of many secreted factors (secretome) and ECM components, attenuating the deleterious effect of TGF $\beta$  overactivation in DMD myotubes.

### **SETDB1 LOF in myotubes has a beneficial impact on the TGF $\beta$ -induced secretome: it improves muscle differentiation and reduces fibrosis**

So far, our data showed that SETDB1 LOF in myotubes could potentially modulate the impact of TGF $\beta$  on surrounding cells and influence the process of muscle regeneration. To test this, we produced conditioned medium (CM) in healthy or DMD myotubes with or without SETDB1 LOF, treated or not with TGF $\beta$  (**Figure 5A**). Then, CM from healthy myotubes was applied on confluent healthy myoblasts, and CM from DMD myotubes on DMD myoblasts (**Figure 5A**).



As a control, we first checked whether CM could impact myoblast differentiation as compared to fresh differentiation medium and found no changes (**Figure S5A-C**). Thus, receiving myoblasts treated with fresh medium or CM supplemented with TGF $\beta$ 1 showed a less efficient differentiation (**Figure S5A-C**). In absence of medium change, CM from TGF $\beta$ -treated myotubes completed impaired fusion, as expected, meaning that TGF $\beta$ 1 is not consumed by the cells and remains high in the CM and though hides the myotube response (18).

We then included medium change after acute TGF $\beta$ 1 treatment to study the intrinsic myotube pro-fibrotic response (**Figure 5A**). Our results showed that DMD myoblasts, but not healthy ones, have a decreased fusion index when differentiated with CM from TGF $\beta$ 1-treated myotubes, but myoblasts incubated with CM from TGF $\beta$ 1-treated and SETDB1 LOF myotubes show a more normal fusion index (**Figures 5B-C**). SETDB1 KD alone in myotubes, in absence of TGF $\beta$ , does not affect the fusion index of the myoblasts receiving the conditioned medium (**Figure S5E**).

The discrepancy between the effects of the CM in healthy versus DMD myoblasts could be due to the higher differentiation potential of the healthy myoblasts (**Figure S5D**).

Interestingly, the receiving myoblasts incubated with CM from TGF $\beta$ -treated and SETDB1 LOF myotubes showed a significant decrease in fibrotic markers *SERPINE1* and *MSTN* (*Myostatin*) (**Figure 5D**). Concomitantly, they displayed an increase in differentiation markers expression such as *MYOD1*, *Myogenin*, *MCK* (*Muscle Creatine Kinase*) and *MYH1* (**Figures 5E-F**). SETDB1 KD alone in myotubes, in absence of TGF $\beta$ , does not affect the expression of muscle differentiation genes in the myoblasts receiving the conditioned medium (**Figure S5F**).

Together, these data show that the TGF $\beta$  pathway activation could be attenuated by targeting SETDB1 in DMD patients. Thus, SETDB1 LOF in myofibers could be beneficial for the myofibers themselves but also to the differentiation of the surrounding myoblasts and potentially limit fibrosis.

## DISCUSSION

Tissue repair, such as muscle regeneration after injury, involves many cell types which communicate together through secreted molecules, the so-called secretome, to orchestrate replacement of damaged myofibers with new functional ones and re-establish tissue homeostasis. Thus, in addition to their contractile properties, myofibers have also a central role in cell-cell communication since they are able to secrete the so-called myokines especially during regeneration and differentiation, including dedicated vesicles acting on muscle adaptation to damage or exercise (26). This myofibers secretome plays important roles in intercellular communication of the muscle

resident cells, i.e., MuSCs, myofibers themselves, the fibro-adipogenic progenitors (FAPs) and macrophages (27).

Although FAPs and macrophages have been described as major TGF $\beta$  sources, and as fibrosis and inflammation effectors, myofibers are also responsive to TGF $\beta$ 1 and have been shown to participate in TGF $\beta$ /SMAD response during muscle regeneration *in vivo* (28, 29). TGF $\beta$ -induced fibrosis is a key pathological feature in muscle disease, such as DMD, and TGF $\beta$  overactivation worsens muscle degeneration by preventing proper repair. Moreover, even though myoblasts display a TGF $\beta$ /SMAD autocrine signal required for proliferation, TGF $\beta$  pathway prevents myoblast fusion by controlling actin remodeling and its activation decreases upon muscle differentiation. Interestingly, DMD muscle cellular models have been shown to recapitulate TGF $\beta$ /SMAD overactivation pathological feature *in vitro* associated with muscle defects and fibrosis (15, 16). Hence, targeting TGF $\beta$ -induced pro-fibrotic response in DMD to slow down disease progression appears as a promising therapeutic approach.

TGF $\beta$  signaling nuclear endpoint mediated by the two major TGF $\beta$  downstream transcription factors SMAD2 and SMAD3, also involves many chromatin-modifying enzymes including SETDB1. SETDB1 regulates different cell fates including stemness and terminal differentiation (2). SETDB1 is also involved in fibrosis (23), inflammatory response and diseases (5, 30-32) and has been linked to TGF $\beta$  response in many non-muscle contexts (19-23). SETDB1 has been previously described by us and others as a major negative regulator of muscle terminal differentiation through repression of muscle gene expression in myoblast nuclei (4, 33). SETDB1 is responsive to Wnt signaling that is known to have a pro-differentiation effect on myoblasts and causes SETDB1 export to the cytoplasm allowing muscle gene de-repression. Interestingly, SETDB1 has also been shown to be a regulator of TGF $\beta$ /SMAD pathway in cancer (19-22) and pulmonary fibrosis (23) contexts. In this study, we have shown that SETDB1 nuclear localization follows TGF $\beta$ /SMAD activation in human muscle cells. SETDB1 is exported to the cytoplasm during muscle terminal differentiation, while TGF $\beta$ /SMAD pathway activation decreases, but it can be transiently relocalized in myotube nuclei in response to TGF $\beta$  pathway activation.

In the effort to identify new key players of TGF $\beta$  response whose dysregulation is a key driver of DMD progression, we hypothesized that SETDB1 could participate in TGF $\beta$  response in muscle and, though, could be involved in deregulation of TGF $\beta$  pathway in DMD. We found that SETDB1 is constitutively accumulated in DMD myotube nuclei correlating with phospho-SMAD3 high nuclear levels *in vivo* and *in vitro*. As expected, DMD myotubes are more responsive to TGF $\beta$  activation as compared to healthy ones. Interestingly, we were able to block SETDB1 nuclear localization by inhibiting intrinsic autocrine TGF $\beta$  signal in DMD myotubes. We also found that SETDB1 nuclear translocation is not dependent on SMAD3 protein and might involve phosphorylation events. We also performed a loss-of-function of SETDB1 and studied TGF $\beta$ -induced pro-fibrotic response. We

found that healthy but especially DMD myotubes lacking SETDB1 display an attenuated response to TGF $\beta$ /SMAD pathway activation without changing phospho-SMAD3 levels. These results point to unprecedented link between SETDB1 and TGF $\beta$  response in muscle, and more especially in DMD context. Nonetheless, the exact mechanisms through which SETDB1 localization is controlled by either TGF $\beta$  or Wnt pathways are not yet understood.

To better understand how SETDB1 impacts TGF $\beta$  response in myotubes, we performed global transcriptomic analysis on healthy and DMD myotubes. We found that SETDB1 loss-of-function leads to a global decrease of RNA levels of gene coding for factors involved in ECM remodeling, inflammation and TGF $\beta$  and Wnt pathway signaling that respond to TGF $\beta$ . Since SETDB1 is known for its repressive action on gene expression, we made the hypothesis that SETDB1 could repress TGF $\beta$  inhibitors as already described in cancer context (20). Interestingly, we found that SETDB1 loss-of-function leads to an increased expression of the TGF $\beta$  inhibitor genes *SKIL* and *SMAD7*. The precise mechanism through which SETDB1 controls the expression of these genes remain unclear. Chromatin immunoprecipitation assays on SETDB1 and H3K9me3 marks would be required to confirm that SETDB1 represses TGF $\beta$  inhibitor expression but remain challenging in myotubes. In general, the mechanisms of action of SETDB1 at the chromatin level are well-documented (34, 35). Therefore, we can speculate that the enrichment of SETDB1 in the nucleus in DMD myotubes might affect chromatin accessibility and 3D genome organization, the points which would be interesting to address in future.

Finally, since most of SETDB1 direct or indirect targets are coding for secreted factors involved in fibrotic response and that TGF $\beta$  is known for its anti-differentiation role, we tested the effect of myotube secretome on myoblasts. Moreover, DMD muscle cells display fusion defects in some DMD models of hiPSC-derived myotubes and our model of immortalized myoblast-derived myotubes. Here, TGF $\beta$ -treated myotube secretome negatively impacts myoblast differentiation and more specifically fusion, but SETDB1 LOF in DMD myotubes has a beneficial cell non-autonomous effect on myoblast fusion. We show here the effect of SETDB1 on myotube secretome that affects envoning cells. Targeting SETDB1 in post-mitotic myotubes does not seem to have deleterious effect on cell viability but its targeting in muscle tissue would be challenging since we do not know the impact on other cell types such as FAPs and macrophages. Nevertheless, we highlight interesting, secreted targets displaying aberrant expression in DMD myotubes that could play a role in deleterious environment of DMD. Moreover, some of deregulated targets have an unknown function such as *ANKRD33B* but could be involved in muscle regeneration process.

In summary, this study highlights the role of SETDB1 in post-mitotic muscle cells in adaptation to the environmental cues (**Figure S6**). Even though the mechanism(s) involved in SETDB1 regulation and the requirement of its enzymatic activity remains elusive, we unraveled a functional effect of its LOF on muscle cell differentiation and shed light on gene networks under SETDB1 control that would be worthy to study individually in the context of DMD. At last, we did not

investigate Wnt and TGF $\beta$  interaction in this study but it seems that SETDB1 acts as a mediator of these pathway communication in muscle. Even though Wnt and TGF $\beta$  are described as having opposite effects on muscle terminal differentiation, some studies also described their collaboration in muscle (36, 37). Here, we showed *in vitro* the cell non-autonomous effect of SETDB1 inhibition in myotubes on the surrounding myoblast differentiation through the control of secreted factor expression induced by TGF $\beta$ /SMAD pathway. Nevertheless, it is unclear if this effect could be beneficial *in vivo* during regeneration since some of the targets have been described in literature for their double-edged effect on muscle regeneration and this could depend on the cell type involved. A fine-tuned balance has to be established in muscle tissue to allow a proper regeneration and SETDB1 could be involved in this mechanism by controlling muscle differentiation, but also by regulating myofiber response to environmental cues.

## MATERIALS & METHODS

### Establishment of stable cell lines and cell culture

The human induced pluripotent stem cell lines, control and DMD, generated from skin fibroblasts from Coriell (**Table 1**), were obtained from the Marseille Stem Cells platform in Marseille Medical Genetics laboratory and described in (38). iPSCs were maintained in culture in mTesR1 medium and dissociated using ReLeSRTM (StemcellTM). They were transfected with epB-Puro-TT-mMyoD and epB-Bsd-TT-FlagBaf60c2 by electroporation using the Neon Transfection System as described in (15, 16). Selection was performed using 5  $\mu$ g/ml of Puromycin and 10  $\mu$ g/ml of Blasticidin at the same time.

Human control and DMD immortalized myoblasts were obtained from AFM-MyoLine. Immortalization was performed as described in: (24), using human telomerase-expressing and cyclin-dependent kinase 4-expressing vectors. They were cultured on Gelatin-coated plates and propagated in DMEM high glucose GlutaMAX™ (Invitrogen, 61965-026)/Medium 199 (Invitrogen, 41150020) 4:1 mixture, supplemented with 20% FBS (Sigma, F7524), 50  $\mu$ g/mL Fetuin, 5  $\mu$ g/mL insulin, 0.5 ng/mL bFGF, 5 ng/mL hEGF and 0.2  $\mu$ g/mL Dexamethasone (Sigma D4902).

### Myogenic differentiation of iPSCs and human immortalized myoblasts

iPSCs stably expressing MyoD and Baf60C Tet-ON inducible transgenes were propagated in mTeSR1 on Matrigel-coated wells. To induce myogenic differentiation starting from iPSC colonies, doxycycline (200 ng/mL) was added in cells maintained in mTeSR1 (day 0). After 24 h of treatment with doxycycline (day1), cells were dissociated as single cells using TrypLE and plated 25.10<sup>3</sup>/cm<sup>2</sup> in growth media (GM) (knockout DMEM (Invitrogen) supplemented with 1 mM L-glutamine, 20% knockout serum replacement medium (KOSR, Invitrogen), Glutamax 1X, 0.1 mM nonessential aminoacids (NEAA, Invitrogen), 50 U/mL streptomycin (Invitrogen) plus hES cell recovery

supplement (10  $\mu$ M) (Stemgent) and doxycycline. On day 3, GM medium was switched to differentiation media (DM) (knockout serum-free DMEM containing 1X Insulin-Transferrin-Selenium (ITS) (SIGMA)) supplemented with doxycycline until day 7.

Human control and DMD immortalized myoblasts were grown at > 80% confluence and the medium was switched for DM media.

### **Myotube transfection with siRNAs**

Three days-differentiated myotubes cultured in differentiation medium (DM) were transfected with siRNAs at a final concentration of 70 nM, employing the Lipofectamin™ RNAiMAX transfection agent (Invitrogen™, #13778100). Cells were kept 2 days in transfection medium before switching with fresh DM. We used the ON-TARGETplus Human SETDB1 siRNA (Dharmacon™; L-020070-00-0010); *SMAD3* siRNA (Sens: GUGUGAGUUCGCCUUCUAUUAU; Antisens: AUAUUGAAGGCGAACUCACAC), and the ON-TARGETplus Non-targeting Control Pool (Dharmacon™; D-001810-10) as a scrambled control siRNA.

### **Immunofluorescence on cells and tissue**

Muscle histological sections were obtained from muscle biopsies of 15-years old DMD patient paravertebral muscle and a healthy 17-years old individual. Frozen samples were fixed in ice-cold acetone for 1 min and then air-dried. They were blocked in 4% BSA-PBS solution for 45 min at room temperature (RT) and then incubated with the primary antibodies at 4°C overnight. Secondary antibodies were applied for 45 min at RT in the dark before mounting.

Cells were grown on Matrigel- and Gelatin-coated coverslips for iPSCs and human myoblasts respectively. They were fixed with 4% PFA for 20 min and they were saturated with 50 mM NH<sub>4</sub>Cl for 10 min at RT. Then, they were permeabilized with 0,5% Triton X-100 for 5 min. Cells were washed 3 times with PBS between each steps. Cells were next incubated with primary antibodies at the concentration indicated in the antibody list below. Alexa-488 or 555 or 647 were used as secondary antibodies (Invitrogen). Nuclei were counterstained with DAPI.

Images were acquired with a Leica DMI-6000B microscope and analysis was performed using Fiji software. In-house macro was used to performed nuclear and cytoplasmic signal quantification. Acquisition of iPSCs-derived myotubes was performed using ImageXpress Micro High Content Screening System. All the conditions were acquired with identical settings and were analyzed with an in-house macro on ImageJ/Fiji to quantify nuclear and cytoplasmic signal. The used antibodies are listed on **Table 2**.

### **RNA and Quantitative Reverse Transcription-PCR (RT-qPCR)**

Total RNA was extracted using RNeasy micro-kit (Qiagen) following manufacturer's procedures. DNase (Qiagen) treatment was performed to remove residual DNA. 1  $\mu$ g of total RNA was reverse transcribed with High-Capacity cDNA Reverse Transcription Kit (Applied Biosystems). Real-time quantitative PCR was performed to analyze relative gene expression levels using SYBR Green



Master mix (Applied Biosystems) following manufacturer indications. Relative expression values were normalized to the housekeeping genes mRNA *PPIA* or *UBC*. Primers are listed in **Table 3**.

### **Nuclear and cytoplasmic fractionation**

Cells were scraped directly in 3 volumes of buffer A (20 mM HEPES pH7.0, 0.15 mM EDTA, 0.15 mM EGTA, 10 mM KCl) supplemented with 0.15 mM Spermine, 0.5 mM Spermidine, protease inhibitor cocktail 1X (SIGMAFAST™) and phosphatase inhibitor cocktail 1X (Sigma). Cells were lysed with 0.5% NP-40 and mixed gently by inversion. 0.88 volume of buffer SR (50 mM HEPES pH7, 0.25 mM EDTA, 10 mM KCl, 7% sucrose) supplemented with Spermine, Spermidine, protease and phosphatase inhibitor cocktails was added before centrifugating the cells for 5 min at 2000 x g at 4°C and the supernatant was collected as the cytoplasmic fraction. The core nuclei pelleted at 2000 x g was resuspended in 1 volume of Low Salt buffer (20 mM Tris pH7.65, 25% glycerol, 1.5 mM MgCl<sub>2</sub>, 0.2 mM EDTA, 20 mM NaCl) with 3X of protease and phosphatase inhibitor cocktails. One volume of High Salt buffer (same as Low Salt but with 900 mM NaCl) supplemented with 5 mM ATP while vortexing. The nuclei samples were kept on ice for 30 min and mix by inversion every 5 min. 1 volume of sucrose buffer (20 mM Tris pH7.65, 60 mM NaCl, 15 mM KCl and 0.34 M sucrose) was added before treating the cells with 0.0025 U/μL of MNase for 10 min at 37°C and 1 mM CaCl<sub>2</sub>. Reaction was stopped with 4 mM EDTA. Samples were next sonicated for 10 min and ultracentrifuged at 40000 rpm at 4°C for 30 min. Supernatant was collected as nuclear fraction.

### **Western blot**

Cells were lysed in RIPA buffer (20 mM Tris pH 7.65, 150 mM NaCl, 0.1% SDS, 0.25% NaDoc and 1% NP-40) supplemented with protease inhibitor 1X (SIGMAFAST™) and phosphatase inhibitor 1X (Sigma) and kept on ice for 30 min. Cell lysates were sonicated at 4°C for 10 min (30 sec ON, 30 sec OFF) at medium frequency (Bioruptor Diagenode). Then, the lysates were centrifugated for 10 min at 4°C at maximum speed and the supernatants were kept as the samples. Extracts were resolved on pre-cast polyacrylamide gel cassettes (NuPAGE® 4-12% Bis-Tris) (Invitrogen) and 1X NuPAGE MOPS SDS Running Buffer and transferred into nitrocellulose membrane (Amersham) in 20 mM phosphate transfer buffer (pH 6.7). Membrane was blocked in 5% skim milk in PBST Buffer (1X PBS, 0.2% Tween 20) and incubated overnight at 4°C with the primary antibody (see Table of antibodies). Membranes were washed twice 5 minutes in PBST, incubated with appropriate secondary antibody IRDye® (Li-Cor) in PBST, washed twice 10 minutes in PBST, once 10 minutes in PBS and then imaged on Odyssey® Imaging System. The used antibodies are listed on **Table 2**.

### **Transcriptome analysis and bioinformatics**

RNA was isolated as described above. Two or three independent biological replicates were sequenced depending on the cell conditions. Libraries were generated using the Ion AmpliSeq™ Transcriptome Human Gene Expression Kit (A24325; Ion Torrent™). Sequencing was performed on an Ion S5 sequencer (Ion Torrent™). The reads were analyzed using the plugin AmpliseqRNA of Torrent Suite (release 5.10) and mapped on the panel of human AmpliseqRNA which reveals the



expression of 20813 genes. Out of these 18185 genes were uniquely mapped on human reference gene names (retrieved from ENSEMBL\_102 release).

Using the raw count Table generated, genes having low counts were filtered out based on the log2 of Counts Per Million (logCPM). We kept genes that have at least a 1 logCPM mean expression level in at least one of the experimental conditions, leaving 12 060 expressed genes for all the downstream analyses. The differential expression analysis of the filtered data was performed using the edgeR package of R and the limma-trend method described in (39). For each experimental setting Differentially Expressed Genes (DEGs) were further analyzed for all the available enrichments. Enrichment analysis and visualization of the DEGs were performed using the clusterProfiler R package (40), using Gene Ontologies, DOSE, the MSigDB database and GSEA (41).

## Statistical analyses

Statistical analyses were carried out using Excel and R. Data are represented as mean +/- SEM as described in the figure legend. Graphs are prepared using Excel and R. Double tail *t* test was used for statistical analysis, \**p*<0.05, \*\**p*<0.01, \*\*\**p*<0.001.

## REFERENCES

1. C. Mozzetta, E. Boyarchuk, J. Pontis, S. Ait-Si-Ali, Sound of silence: the properties and functions of repressive Lys methyltransferases. *Nat Rev Mol Cell Biol* **16**, 499-513 (2015).
2. J. Padeken, S. P. Methot, S. M. Gasser, Establishment of H3K9-methylated heterochromatin and its functions in tissue differentiation and maintenance. *Nat Rev Mol Cell Biol* **23**, 623-640 (2022).
3. M. Markouli, D. Strepkos, C. Piperi, Structure, Activity and Function of the SETDB1 Protein Methyltransferase. *Life (Basel)* **11**, (2021).
4. S. Beyer, J. Pontis, E. Schirwis, V. Battisti, A. Rudolf, F. Le Grand, S. Ait-Si-Ali, Canonical Wnt signalling regulates nuclear export of Setdb1 during skeletal muscle terminal differentiation. *Cell Discov* **2**, 16037 (2016).
5. L. Juznic, K. Peuker, A. Strigli, M. Brosch, A. Herrmann, R. Hasler, M. Koch, L. Matthiesen, Y. Zeissig, B. S. Loscher, A. Nuber, G. Schotta, V. Neumeister, T. Chavakis, T. Kurth, M. Lesche, A. Dahl, A. von Massenhausen, A. Linkermann, S. Schreiber, K. Aden, P. C. Rosenstiel, A. Franke, J. Hampe, S. Zeissig, SETDB1 is required for intestinal epithelial differentiation and the prevention of intestinal inflammation. *Gut* **70**, 485-498 (2021).
6. D. Strepkos, M. Markouli, A. Klonou, A. G. Papavassiliou, C. Piperi, Histone Methyltransferase SETDB1: A Common Denominator of Tumorigenesis with Therapeutic Potential. *Cancer Res* **81**, 525-534 (2021).
7. E. P. Hoffman, R. H. Brown, Jr., L. M. Kunkel, Dystrophin: the protein product of the Duchenne muscular dystrophy locus. *Cell* **51**, 919-928 (1987).
8. J. R. Mendell, L. Rodino-Klapac, Z. Sahenk, V. Malik, B. K. Kaspar, C. M. Walker, K. R. Clark, Gene therapy for muscular dystrophy: lessons learned and path forward. *Neurosci Lett* **527**, 90-99 (2012).
9. E. P. Hoffman, The discovery of dystrophin, the protein product of the Duchenne muscular dystrophy gene. *The FEBS journal* **287**, 3879-3887 (2020).
10. B. J. Petrof, J. B. Shrager, H. H. Stedman, A. M. Kelly, H. L. Sweeney, Dystrophin protects the sarcolemma from stresses developed during muscle contraction. *Proc Natl Acad Sci U S A* **90**, 3710-3714 (1993).
11. K. A. Lavidas, R. Kakkar, E. M. McNally, The dystrophin glycoprotein complex: signaling strength and integrity for the sarcolemma. *Circ Res* **94**, 1023-1031 (2004).

12. K. L. Walton, K. E. Johnson, C. A. Harrison, Targeting TGF-beta Mediated SMAD Signaling for the Prevention of Fibrosis. *Front Pharmacol* **8**, 461 (2017).
13. T. H. Kao, H. F. Liao, D. Wolf, K. Y. Tai, C. Y. Chuang, H. S. Lee, H. C. Kuo, K. Hata, X. Zhang, X. Cheng, S. P. Goff, S. K. Ooi, T. H. Bestor, S. P. Lin, Ectopic DNMT3L triggers assembly of a repressive complex for retroviral silencing in somatic cells. *J Virol* **88**, 10680-10695 (2014).
14. A. L. Serrano, C. J. Mann, B. Vidal, E. Ardite, E. Perdiguero, P. Munoz-Canoves, Cellular and molecular mechanisms regulating fibrosis in skeletal muscle repair and disease. *Current topics in developmental biology* **96**, 167-201 (2011).
15. L. Caputo, A. Granados, J. Lenzi, A. Rosa, S. Ait-Si-Ali, P. L. Puri, S. Albini, Acute conversion of patient-derived Duchenne muscular dystrophy iPSC into myotubes reveals constitutive and inducible over-activation of TGFbeta-dependent pro-fibrotic signaling. *Skelet Muscle* **10**, 13 (2020).
16. I. Y. Choi, H. Lim, K. Estrellas, J. Mula, T. V. Cohen, Y. Zhang, C. J. Donnelly, J. P. Richard, Y. J. Kim, H. Kim, Y. Kazuki, M. Oshimura, H. L. Li, A. Hotta, J. Rothstein, N. Maragakis, K. R. Wagner, G. Lee, Concordant but Varied Phenotypes among Duchenne Muscular Dystrophy Patient-Specific Myoblasts Derived using a Human iPSC-Based Model. *Cell Rep* **15**, 2301-2312 (2016).
17. P. Bernasconi, E. Torchiana, P. Confalonieri, R. Brugnoli, R. Barresi, M. Mora, F. Cornelio, L. Morandi, R. Mantegazza, Expression of transforming growth factor-beta 1 in dystrophic patient muscles correlates with fibrosis. Pathogenetic role of a fibrogenic cytokine. *J Clin Invest* **96**, 1137-1144 (1995).
18. F. Girardi, A. Taleb, M. Ebrahimi, A. Datye, D. G. Gamage, C. Peccate, L. Giordani, D. P. Millay, P. M. Gilbert, B. Cadot, F. Le Grand, TGFbeta signaling curbs cell fusion and muscle regeneration. *Nat Commun* **12**, 750 (2021).
19. D. Du, Y. Katsuno, D. Meyer, E. H. Budi, S. H. Chen, H. Koeppen, H. Wang, R. J. Akhurst, R. Derynck, Smad3-mediated recruitment of the methyltransferase SETDB1/ESET controls Snail1 expression and epithelial-mesenchymal transition. *EMBO Rep* **19**, 135-155 (2018).
20. T. Y. Ryu, K. Kim, S. K. Kim, J. H. Oh, J. K. Min, C. R. Jung, M. Y. Son, D. S. Kim, H. S. Cho, SETDB1 regulates SMAD7 expression for breast cancer metastasis. *BMB Rep* **52**, 139-144 (2019).
21. Y. Wakabayashi, T. Tamiya, I. Takada, T. Fukaya, Y. Sugiyama, N. Inoue, A. Kimura, R. Morita, I. Kashiwagi, T. Takimoto, M. Nomura, A. Yoshimura, Histone 3 lysine 9 (H3K9) methyltransferase recruitment to the interleukin-2 (IL-2) promoter is a mechanism of suppression of IL-2 transcription by the transforming growth factor-beta-Smad pathway. *The Journal of biological chemistry* **286**, 35456-35465 (2011).
22. P. C. Wu, J. W. Lu, J. Y. Yang, I. H. Lin, D. L. Ou, Y. H. Lin, K. H. Chou, W. F. Huang, W. P. Wang, Y. L. Huang, C. Hsu, L. I. Lin, Y. M. Lin, C. K. Shen, T. Y. Tzeng, H3K9 histone methyltransferase, KMT1E/SETDB1, cooperates with the SMAD2/3 pathway to suppress lung cancer metastasis. *Cancer Res* **74**, 7333-7343 (2014).
23. T. Liu, P. Xu, S. Ke, H. Dong, M. Zhan, Q. Hu, J. Li, Histone methyltransferase SETDB1 inhibits TGF-beta-induced epithelial-mesenchymal transition in pulmonary fibrosis by regulating SNAI1 expression and the ferroptosis signaling pathway. *Arch Biochem Biophys* **715**, 109087 (2022).
24. K. Mamchaoui, C. Trollet, A. Bigot, E. Negroni, S. Chaouch, A. Wolff, P. K. Kandalla, S. Marie, J. Di Santo, J. L. St Guily, F. Muntoni, J. Kim, S. Philippi, S. Spuler, N. Levy, S. C. Blumen, T. Voit, W. E. Wright, A. Aamiri, G. Butler-Browne, V. Mouly, Immortalized pathological human myoblasts: towards a universal tool for the study of neuromuscular disorders. *Skelet Muscle* **1**, 34 (2011).
25. M. Yamazaki, S. Minota, H. Sakurai, K. Miyazono, A. Yamada, I. Kanazawa, M. Kawai, Expression of transforming growth factor-beta 1 and its relation to endomysial fibrosis in progressive muscular dystrophy. *Am J Pathol* **144**, 221-226 (1994).
26. D. C. Bittel, J. K. Jaiswal, Contribution of Extracellular Vesicles in Rebuilding Injured Muscles. *Front Physiol* **10**, 828 (2019).

27. M. C. Le Bihan, I. Barrio-Hernandez, T. P. Mortensen, J. Henningsen, S. S. Jensen, A. Bigot, B. Blagoev, G. Butler-Browne, I. Kratchmarova, Cellular Proteome Dynamics during Differentiation of Human Primary Myoblasts. *J Proteome Res* **14**, 3348-3361 (2015).
28. F. Accornero, O. Kanisicak, A. Tjondrokoesoemo, A. C. Attia, E. M. McNally, J. D. Molkentin, Myofiber-specific inhibition of TGFbeta signaling protects skeletal muscle from injury and dystrophic disease in mice. *Hum Mol Genet* **23**, 6903-6915 (2014).
29. M. M. G. Hillege, A. Shi, R. A. Galli, G. Wu, P. Bertolino, W. M. H. Hoogaars, R. T. Jaspers, Lack of Tgfb1 and Acvr1b synergistically stimulates myofibre hypertrophy and accelerates muscle regeneration. *Elife* **11**, (2022).
30. R. Hachiya, T. Shiihashi, I. Shirakawa, Y. Iwasaki, Y. Matsumura, Y. Oishi, Y. Nakayama, Y. Miyamoto, I. Manabe, K. Ochi, M. Tanaka, N. Goda, J. Sakai, T. Suganami, Y. Ogawa, The H3K9 methyltransferase Setdb1 regulates TLR4-mediated inflammatory responses in macrophages. *Sci Rep* **6**, 28845 (2016).
31. E. Johnson, K. Salari, S. Yang, SETDB1: A perspective into immune cell function and cancer immunotherapy. *Immunology* **169**, 3-12 (2023).
32. R. Wang, H. Li, J. Wu, Z. Y. Cai, B. Li, H. Ni, X. Qiu, H. Chen, W. Liu, Z. H. Yang, M. Liu, J. Hu, Y. Liang, P. Lan, J. Han, W. Mo, Gut stem cell necroptosis by genome instability triggers bowel inflammation. *Nature* **580**, 386-390 (2020).
33. Y. J. Song, J. H. Choi, H. Lee, Setdb1 Is Required for Myogenic Differentiation of C2C12 Myoblast Cells via Maintenance of MyoD Expression. *Mol Cells* **38**, 362-372 (2015).
34. V. V. Zakharova, M. D. Magnitov, L. Del Maestro, S. V. Ulianov, A. Glentis, B. Uyanik, A. Williard, A. Karpukhina, O. Demidov, V. Joliot, Y. S. Vassetzky, R. M. Mege, M. Piel, S. V. Razin, S. Ait-Si-Ali, SETDB1 fuels the lung cancer phenotype by modulating epigenome, 3D genome organization and chromatin mechanical properties. *Nucleic Acids Res*, (2022).
35. Y. Jiang, Y. E. Loh, P. Rajarajan, T. Hirayama, W. Liao, B. S. Kassim, B. Javidfar, B. J. Hartley, L. Kleofas, R. B. Park, B. Labonte, S. M. Ho, S. Chandrasekaran, C. Do, B. R. Ramirez, C. J. Peter, W. J. C, B. M. Safaie, H. Morishita, P. Roussos, E. J. Nestler, A. Schaefer, B. Tycko, K. J. Brennand, T. Yagi, L. Shen, S. Akbarian, The methyltransferase SETDB1 regulates a large neuron-specific topological chromatin domain. *Nat Genet*, (2017).
36. S. Biressi, E. H. Miyabara, S. D. Gopinath, P. M. Carlig, T. A. Rando, A Wnt-TGFbeta2 axis induces a fibrogenic program in muscle stem cells from dystrophic mice. *Sci Transl Med* **6**, 267ra176 (2014).
37. A. Rudolf, E. Schirwis, L. Giordani, A. Parisi, C. Lepper, M. M. Taketo, F. Le Grand, beta-Catenin Activation in Muscle Progenitor Cells Regulates Tissue Repair. *Cell Rep* **15**, 1277-1290 (2016).
38. C. Badja, G. Maleeva, C. El-Yazidi, E. Barruet, M. Lasserre, P. Tropel, B. Binetruy, P. Bregestovski, F. Magdinier, Efficient and cost-effective generation of mature neurons from human induced pluripotent stem cells. *Stem Cells Transl Med* **3**, 1467-1472 (2014).
39. C. W. Law, M. Alhamdoosh, S. Su, X. Dong, L. Tian, G. K. Smyth, M. E. Ritchie, RNA-seq analysis is easy as 1-2-3 with limma, Glimma and edgeR. *F1000Res* **5**, (2016).
40. T. Wu, E. Hu, S. Xu, M. Chen, P. Guo, Z. Dai, T. Feng, L. Zhou, W. Tang, L. Zhan, X. Fu, S. Liu, X. Bo, G. Yu, clusterProfiler 4.0: A universal enrichment tool for interpreting omics data. *Innovation (Camb)* **2**, 100141 (2021).
41. A. Liberzon, C. Birger, H. Thorvaldsdottir, M. Ghandi, J. P. Mesirov, P. Tamayo, The Molecular Signatures Database (MSigDB) hallmark gene set collection. *Cell Syst* **1**, 417-425 (2015).

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## **Supplementary Materials:** 6 Supplementary figures and 4 supplementary tables

Figure S1: SETDB1 localization depends on TGF $\beta$ /SMAD pathway activation during muscle terminal differentiation

Figure S2: SETDB1 translocate into muscle cell nuclei in response to TGF $\beta$ /SMAD pathway activation and show more persistent nuclear signal in DMD myotubes irrespectively of the type of DMD mutation

Figure S3: Efficient siRNA-mediated SETDB1 knockdown in TGF $\beta$ -treated DMD myotubes leads to a decrease in TGF $\beta$  target gene expression and an increase in pro-myogenic factors *SKIL* and *MYH3*

Figure S4: Healthy and DMD myotubes respond differently to TGF $\beta$ /SMAD pathway activation but display some SETDB1 target gene signatures in common

Figure S5: TGF $\beta$ /SMAD pathway activation leads to fusion defects in muscle cells

Figure S6: Graphical abstract

Table S1: Top DEGs + *versus* – TGF $\beta$ 1 in healthy myotubes

Table S2: Top DEGs + *versus* – TGF $\beta$ 1 in DMD myotubes

Table S3: Top DEGs siSETDB1 *versus* siCTL during TGF $\beta$  response in WT myotubes

Table S4: Top DEGs siSETDB1 *versus* siCTL during TGF $\beta$  response in DMD myotubes



## FIGURE LEGENDS

### Figure 1: SETDB1 is excluded from myotube nuclei upon differentiation and can be relocated in response to TGF $\beta$ pathway activation

- A.** Scheme of experimental design. Cells were collected in proliferating phase (myoblasts) or after 3 days of differentiation (myotubes) with or without TGF $\beta$ 1 treatment at 20 ng/mL.
- B.** Immunostaining of SETDB1 (red) and phospho-SMAD3 (green). Nuclei were stained with DAPI (blue). Scale bar, 10  $\mu$ M.
- C.** Quantification of SETDB1 nuclear/cytoplasmic signal ratio.
- D.** RT-qPCR of *SETDB1* in healthy proliferating myoblasts and differentiated myotubes +/- TGF $\beta$ 1.
- E-F.** Quantification of SETDB1 and phospho-SMAD3 protein levels in proliferating myoblast and myotube treated or not with TGF $\beta$ 1 in total extracts. Alpha tubulin was used as a loading control to normalize samples.
- G.** Scheme of experimental design with TGF $\beta$  inhibitor SB-413542.
- H.** Immunostaining of SETDB1 (red) and phospho-SMAD3 (green). Nuclei were stained with DAPI (blue). SB-413542 blocks SETDB1 relocalization in myotube nuclei upon TGF $\beta$ 1 treatment. Scale bar, 10  $\mu$ M.
- I-J.** Quantification of SETDB1 nuclear/cytoplasmic signal ratio (I), and phospho-SMAD3 nuclear signal (J).
- For all panels:** Statistics were performed on  $\geq 3$  biological replicates ( $>100$  nuclei for immunostaining quantification) and data are represented as average  $\pm$  SEM \* $p < 0.05$ ; \*\* $p < 0.01$ ; \*\*\* $p < 0.001$  (unpaired Student's t test).

### Figure 2: DMD differentiated muscle cells display a constitutive relocalization of SETDB1 in the nuclei and a higher activation of TGF $\beta$ /SMAD pathway

- A.** Immunostaining of SETDB1 (green), pSMAD3 (magenta) and laminin (red) on histological slides from healthy individual or DMD patient paravertebral muscles. Nuclei were stained with DAPI (blue). White arrows identify centrally localized nuclei of damaged myofibers in DMD muscle section. Scale bar, 10  $\mu$ M.
- B.** Immunostaining of SETDB1 (red) in healthy#1 and DMD del ex 45 myotubes. Nuclei were stained with DAPI (blue). Scale bar, 10  $\mu$ M.
- C.** Quantification of SETDB1 nuclear/cytoplasmic signal ratio.
- D.** Western blot of nuclear and cytoplasmic fractions of healthy and DMD myotubes in response to TGF $\beta$ 1 showing protein levels of SETDB1 and phospho-SMAD3. RNA polymerase II and alpha tubulin were used as loading controls of nuclear and cytoplasmic fractions respectively.



**E-F.** Quantification of SETDB1 nuclear/cytoplasmic ratio (E), and phospho-SMAD3 nuclear signal (F).

**G.** Immunostaining of SETDB1 (red) and phospho-SMAD3 (green) in DMD del ex45 myotubes and quantification of SETDB1 nuclear/cytoplasmic signal ratio. Nuclei were stained with DAPI (blue). Scale bar, 10  $\mu$ M.

**For all panels:** Statistics were performed on  $\geq 3$  biological replicates ( $> 100$  nuclei for immunostaining quantification) and data are represented as average  $\pm$  SEM \* $p < 0.05$ ; \*\* $p < 0.01$ ; \*\*\* $p < 0.001$  (unpaired Student's t test).

### Figure 3: SETDB1 silencing leads to a decreased response to TGF $\beta$ 1 in DMD myotubes while key myogenic markers increase

**A.** Scheme of experimental design. Cells were treated with siRNAs scrambled (CTL) or against SETDB1 for 2 days after 3 days of differentiation (myotubes) and then treated or not with TGF $\beta$ 1 at 20 ng/mL.

**B.** Western blot showing SETDB1, phospho-SMAD3 and total SMAD3 protein levels. H3 was used as a loading control.

**C.** RT-qPCR of TGF $\beta$ /SMADs pathway known targets *TGF $\beta$ 1*, *IL6* and *TIMP1* in healthy and DMD myotubes +/- siSETDB1 +/- TGF $\beta$ 1.

**D.** RT-qPCR of TGF $\beta$ /SMADs pathway inhibitors *SKIL* and *SMAD7* in healthy and DMD myotubes +/- siSETDB1 +/- TGF $\beta$ 1.

**E.** RT-qPCR of myogenic markers *Myogenin*, *MYH1* and *MYH3* (embryonic MHC).

**For all panels:** Statistics were performed on  $\geq 3$  biological replicates and data are represented as average  $\pm$  SEM \* $p < 0.05$ ; \*\* $p < 0.01$ ; \*\*\* $p < 0.001$  (unpaired Student's t test).

### Figure 4: SETDB1 silencing in DMD myotubes leads to a decreased expression of TGF $\beta$ -dependent genes involved in ECM remodeling, receptor signaling transduction and inflammation

**A.** Venn diagram shows that a different set of TGF $\beta$ 1-responsive genes were differentially expressed upon SETDB1 silencing in healthy and DMD myotubes.

**B.** Volcano plot of differentially expressed genes in DMD myotubes +/- TGF $\beta$ 1.

**C.** Heatmap of expression level z-scores for DEGs in DMD myotubes for TGF $\beta$  versus TGF $\beta$ +siSETDB1 comparison. Some genes found to be increased upon TGF $\beta$ 1 treatment decrease upon SETDB1 silencing (*IGFBP3*, *THBS1*, *IL6ST*).

**D-E.** Enriched categories tree plot (D) and enriched gene-concept network for biological pathways deregulated for TGF $\beta$  *versus* TGF $\beta$ +siSETDB1 in DMD myotubes.

**F.** mRNA levels of gene validated by RT-qPCR involved in ECM-remodeling (*THBS1*, *SERPINE1*) and inflammation (*LIF*).

**For all panels:** Statistics were performed on  $\geq 3$  biological replicates and data are represented as average  $\pm$  SEM \* $p < 0.05$ ; \*\* $p < 0.01$ ; \*\*\* $p < 0.001$  (unpaired Student's t test).

### **Figure 5: Secretome of SETDB1 deficient myotubes in response to TGF $\beta$ 1 reduces the negative impact of TGF $\beta$ treatment on myoblast differentiation**

**A.** Diagram of the conditioned medium experiments.

**B.** Immunofluorescence of myosin heavy chain (red) and nuclei staining with DAPI (blue). Scale bar, 50  $\mu$ M.

**C.** Quantification of fusion index after 6 days of differentiation in the conditioned medium.

**D.** RT-qPCR of fibrotic markers *SERPINE1* and *MSTN* after 3 days in the conditioned medium. **E.**

RT-qPCR of early myogenic markers *MYOD1* and *MYOGENIN*.

**F.** RT-qPCR of late myogenic markers *MYH1* and *MCK*.

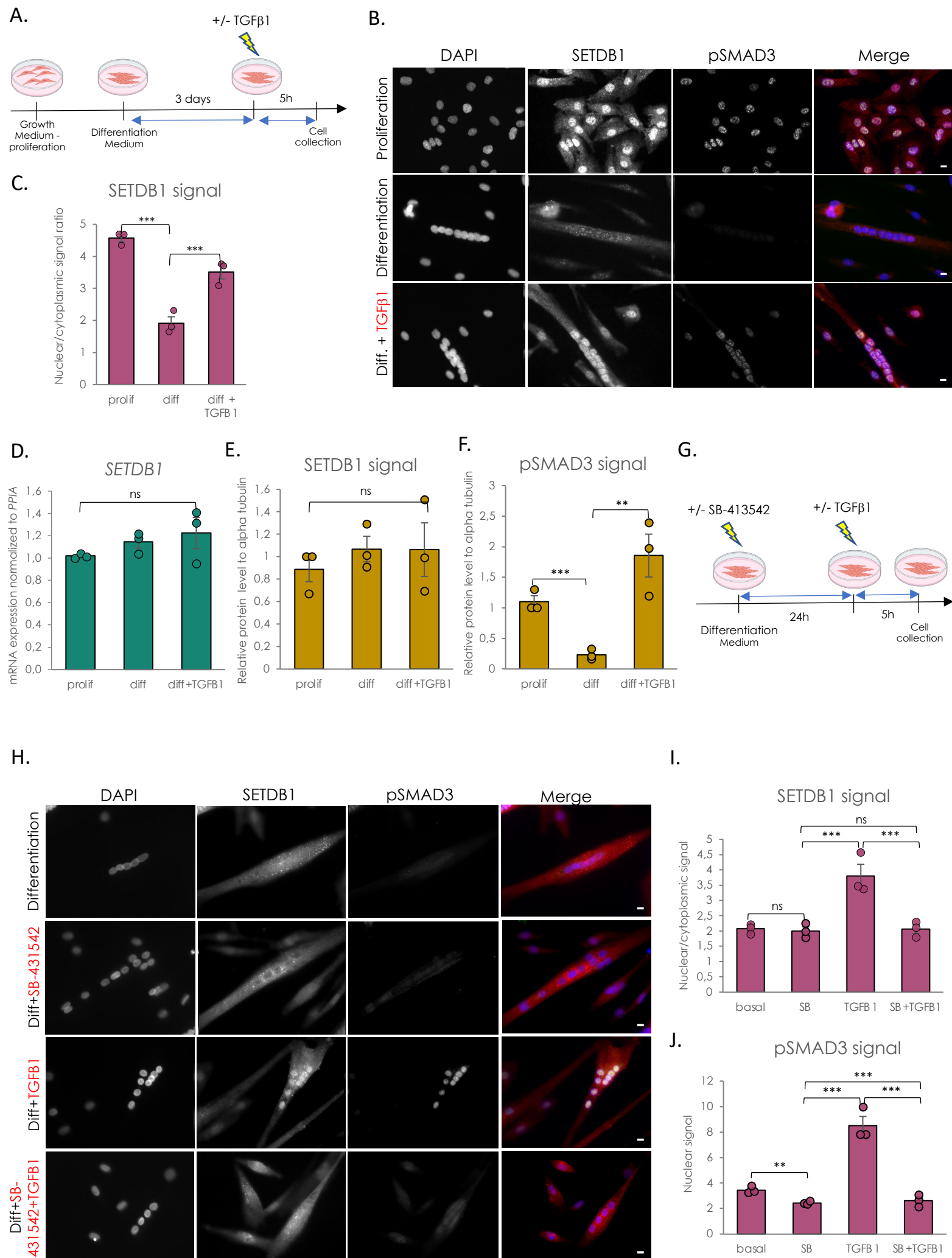
**For all panels:** Statistics were performed on  $\geq 3$  biological replicates ( $> 100$  nuclei for immunostaining quantification) and data are represented as average  $\pm$  SEM \* $p < 0.05$ ; \*\* $p < 0.01$ ; \*\*\* $p < 0.001$  (unpaired Student's t test).

**Table 1:** Cell line list

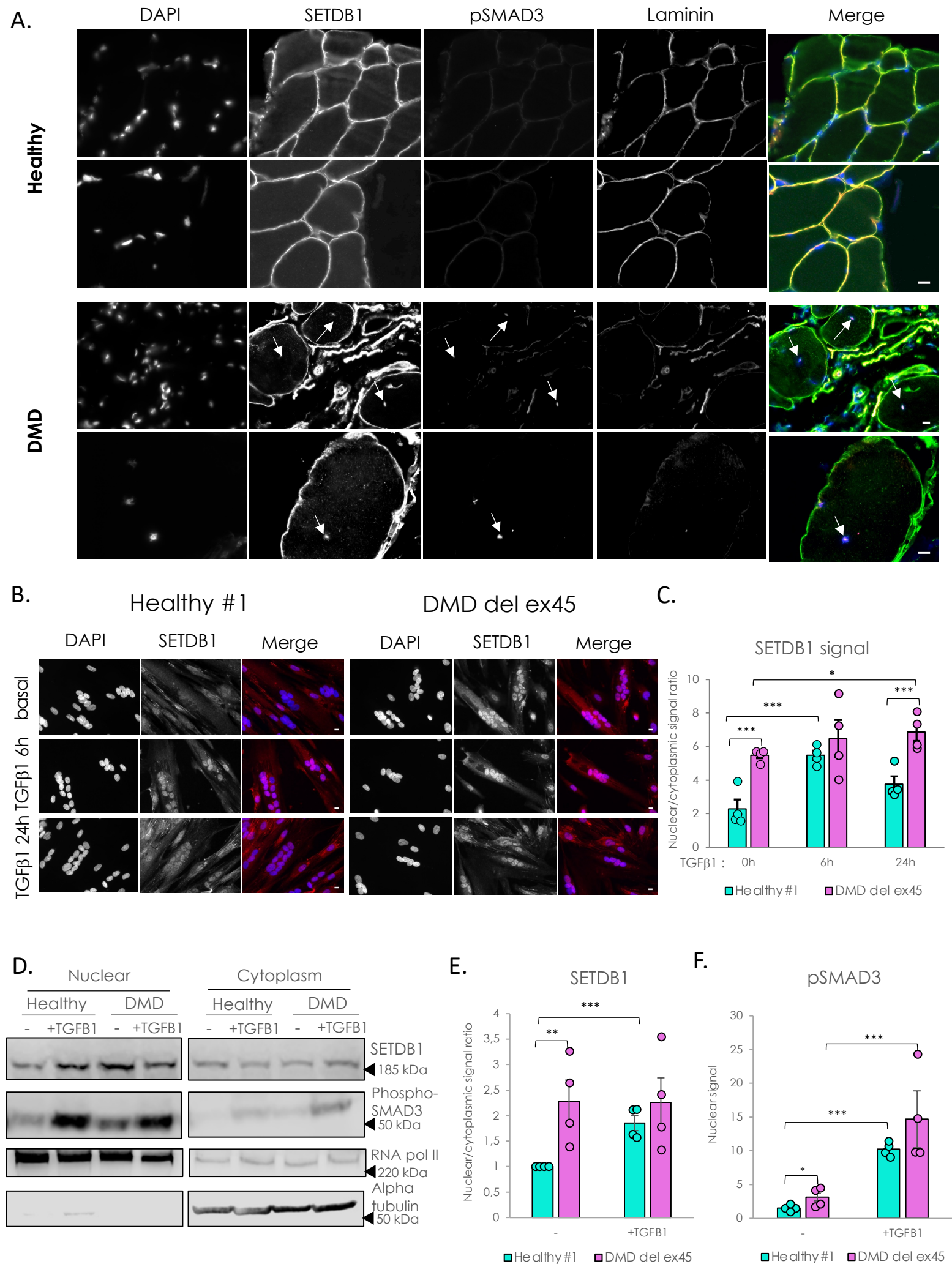
**Table 2:** Antibodies list

**Table 3:** Primer list

## Figure 1, Granados et al.



**Figure 2, Granados et al.**



**Figure 2, Granados et al.**

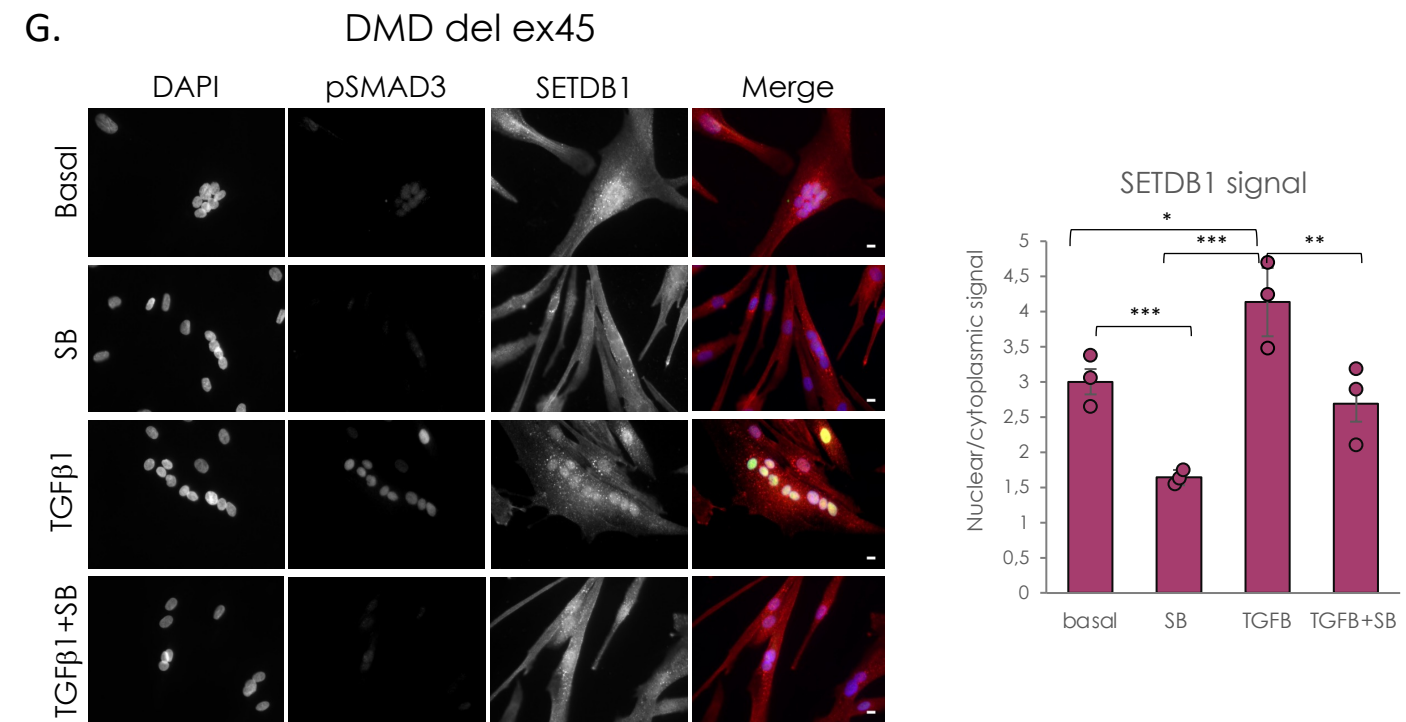




Figure 3, Granados et al.

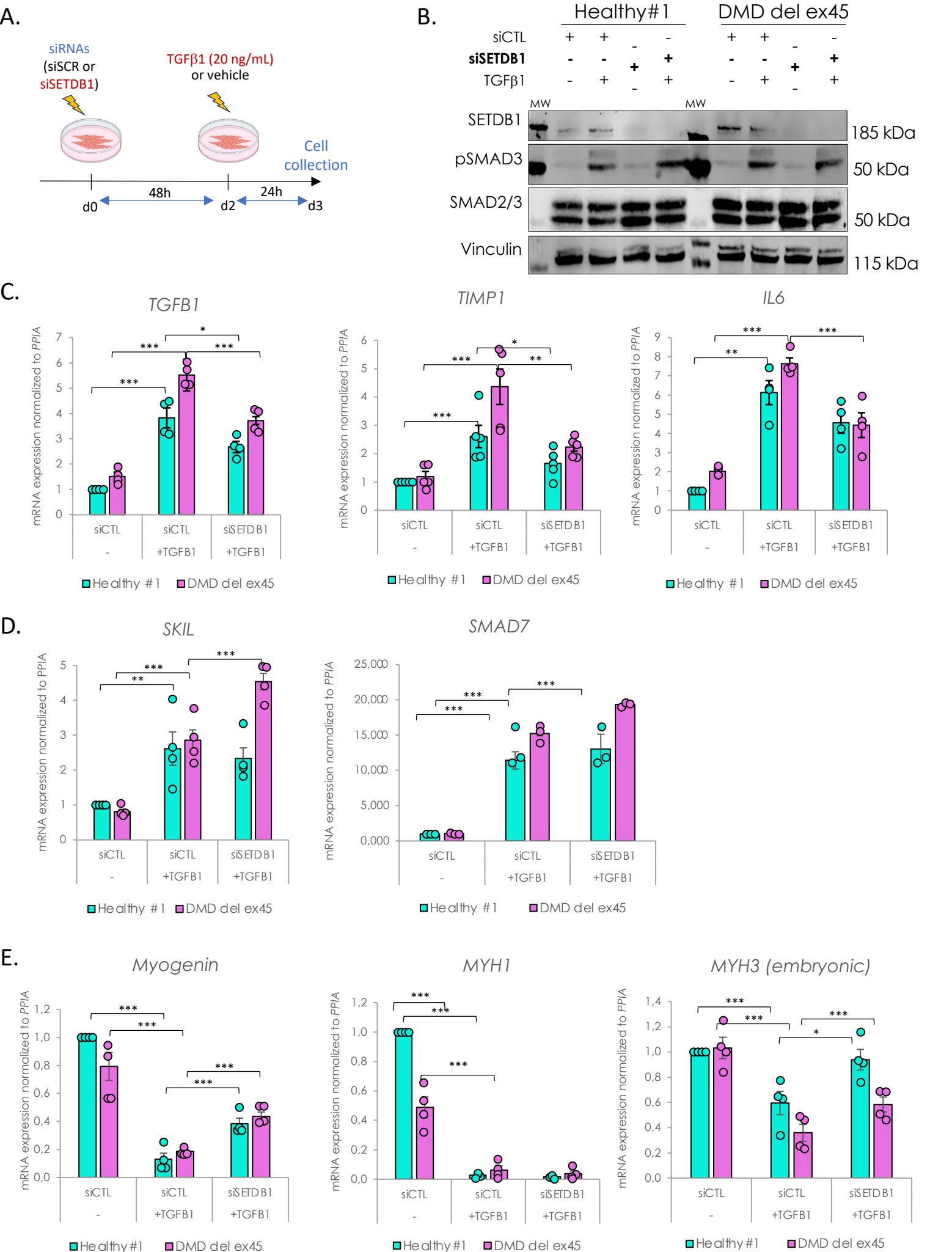
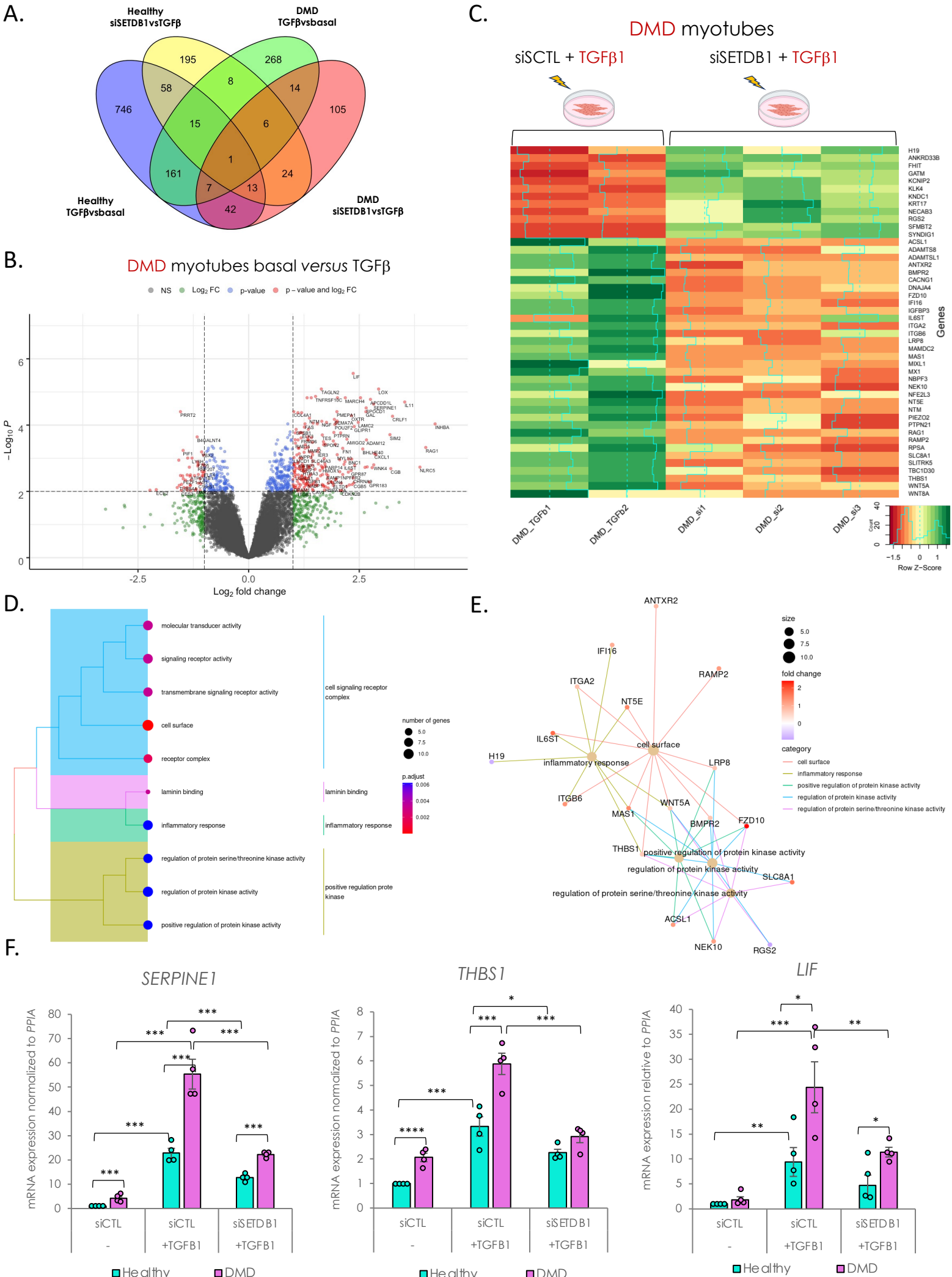
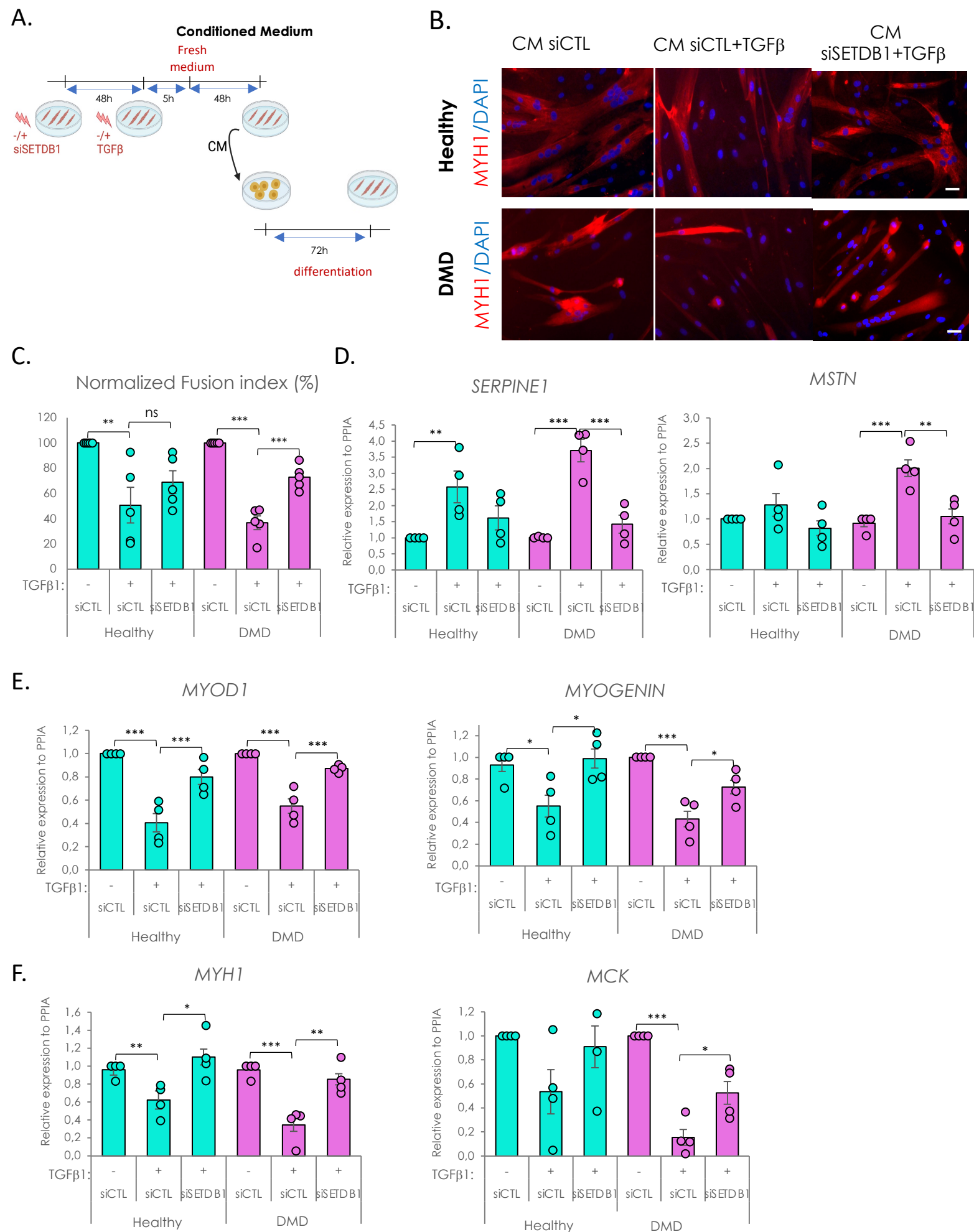




Figure 4, Granados et al.



## Figure 5, Granados et al.



**Table 1: Cell line list**

<i>Cells</i>	<i>Reference</i>	<i>Patient</i>
Induced pluripotent stem cells (iPSCs)	AG08C5 Coriell	Skin fibroblast, 1 year old healthy male
	GM25313	Skin fibroblast, 13-year-old DMD male patient, deletion ex45
Immortalized myoblasts (hTERT/CDK4) (hMB)	AB1190C16PV	Paravertebral, 16-year-old healthy male
	AB1167C20TFL	Fascia lata, 20-year-old healthy male
	AB1023DMD11Q	Quadriceps, 11-year-old DMD male
	AB1071DMD13PV	Paravertebral, 13-year-old DMD male

**Table 2: Antibodies list**

<i>Antibody</i>	<i>reference</i>	<i>Use &amp; dilution</i>
SETDB1 (rabbit)	Santa Cruz sc-66884	Immunofluorescence (1:200)
SETDB1 (mouse)	Abcam ab107225	Western blot (1:1000)
SETDB1 (mouse)	Thermo Fisher Scientific MA5-15722	Immunofluorescence (1:200)
Myosin heavy chain, MYH1E	DSHB	Immunofluorescence (1:10), Western blot (1:200)
Phospho-SMAD3 (rabbit)	Abcam ab52903	Immunofluorescence (1:200), Western blot (1:1000)
SMAD3 (mouse)	Proteintech 66516-1-Ig	Western blot (1:1000)
PAI1 (rabbit)	Proteintech 13801-1-AP	Western blot (1:1000)
RNA polymerase II (rabbit)	Abcam ab5095	Western blot (1:1000)
Lamin A/C (rabbit)	Proteintech 10298-1-AP	Western blot (1:1000)
H3 (C-16) (goat)	Santa Cruz sc-8654	Western blot (1:1000)

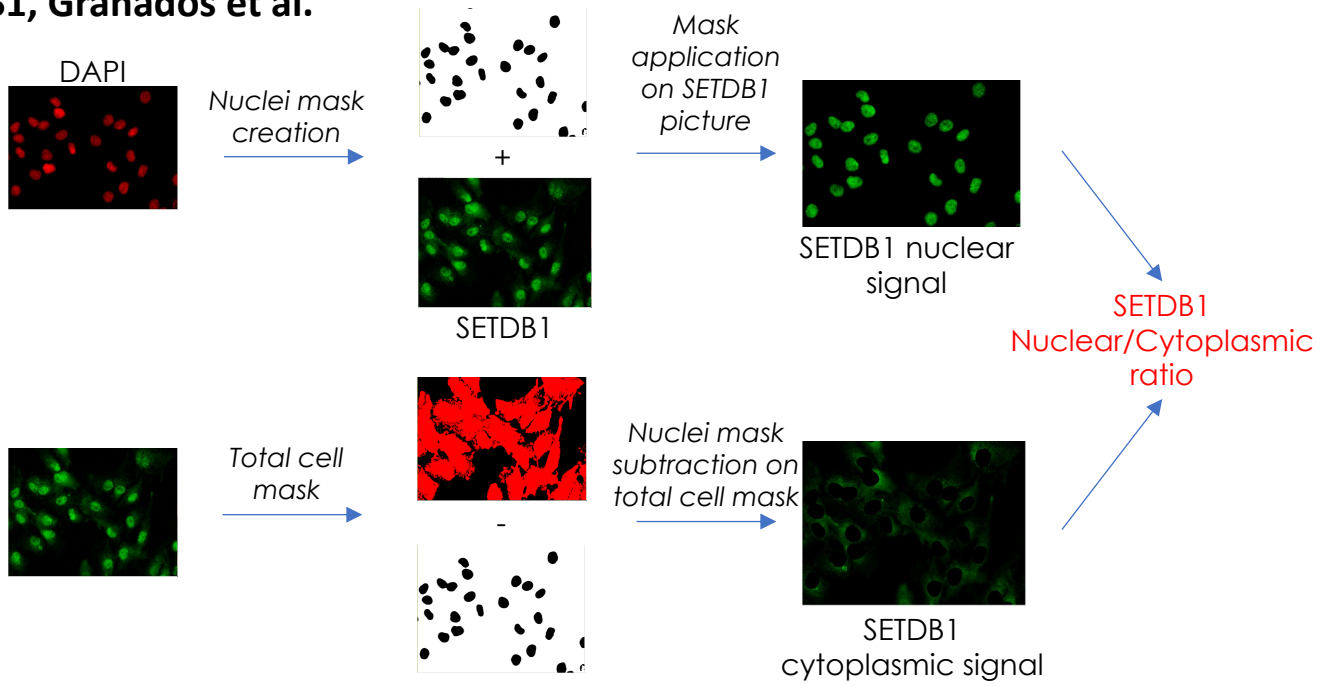
**Table 3: Primer list**

<i>Targets</i>	<i>Forward</i>	<i>Reverse</i>
<i>SETDB1</i>	CAATACCGGGACAGTAGCTC	TCTGGTCTTTTGGAGTTCTGC
<i>TGFB1</i>	GCCTGAGGCCGACTACTA	CTGTGTGTACTCTGCTTGAAC
<i>TIMP1</i>	TTCTGCAATTCCGACCTCG	TCATAACGCTGGTATAAGGTGG
<i>IL6</i>	TAGTGAGGAACAAGCCAGAGC	TGGGTCAGGGGTGGTTATTG
<i>SKIL</i>	GCCCCAAATGTGTCACTTAC	TCCCACTTTGTTTATGTCTCTGA
<i>SMAD7</i>	GTGTTGCTGTGAATCTTACGG	TCGGGTATCTGGAGTAAGGAG
<i>MYOGENIN</i>	AATGCAGCTCTCACAGCGCCTC	TCAGCCGTGAGCAGATGATCC
<i>MYH1</i>	CTGTTGCAGTTTCTCATTGGTG	CCAGGCAGTACTTCATTGGG
<i>MYH3 (embryonic)</i>	CAAGAGTTCTCAGGATGGGAAG	GCCATGTCTTCGATCCTGTC
<i>FN1</i>	AGCCGAGGTTTTAACTGCGA	CCCACTCGGTAAGTGTTCCC
<i>CTGF</i>	TGTGCACCGCCAAAGAT	GCACGTGCACTGGTACTT
<i>SERPINE1</i>	TCCACAAATCAGACGGCAGC	TCGTAGTAATGGCCATCGGG
<i>THBS1</i>	CTGGCCCAATGAGAACCTGG	GCCCTGAGTTGGGAAGGTTG
<i>LIF</i>	CCAAGTGGCACAGCTCAATG	CTTGTCCAGGTTGTTGGGGA

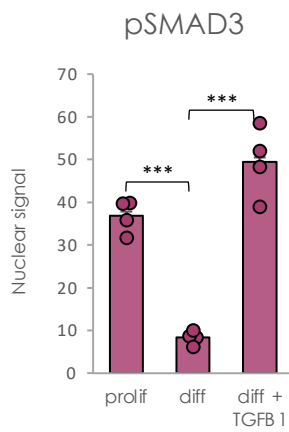
<i>MSTN</i>	AGCGATGGCTCTTTGGAAGAT	TTTGGGTTTTCCATCCACTTGC
<i>BMPR2</i>	GCAGGTTCTCGTGTCTAGGG	CCTGGTCCCAACAGTCTTCG
<i>WNT5A</i>	GCAGCACTGTGGATAACACC	GCTCACCGCGTATGTGAAGG
<i>ADAMTS8</i>	GCAGCGCCATGTATCTCACA	TGTGGGGAGGGGCAGG
<i>ANKRD33B</i>	GGGGAACACAGCCCTAATCA	GCGTTCCTCCTTTCAAGGTCA
<i>MMP14</i>	GGCTGCCTACCGACAAGATT	GCCCTGAGCTCTTCGTTGAA
<i>MYOD1</i>	TGCTCCGACGGCATGATGGACTA	TTGTAGTAGGCGCCTTCGTAGCAGTT
<i>MCK</i>	TGGAGAAGCTCTCTGTGGAAGCTC	TCCGTCATGCTCTTCAGAGGGTAGTA
<i>SMAD3</i>	TGAGGCTGTCTACCAAGTTGACC	GTGAGGACCTTGTCAAGCCACT

## Figure S1, Granados et al.

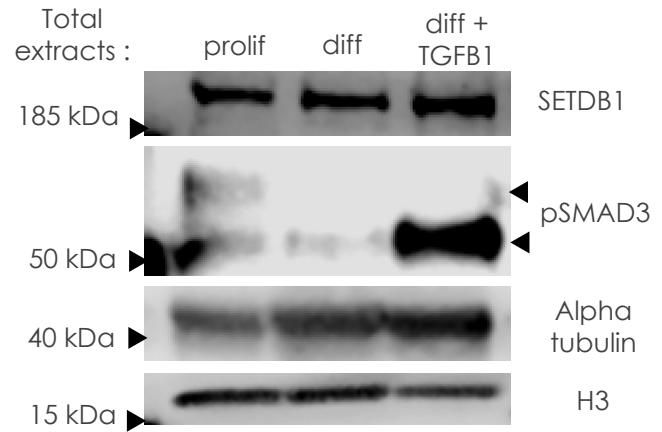
A.



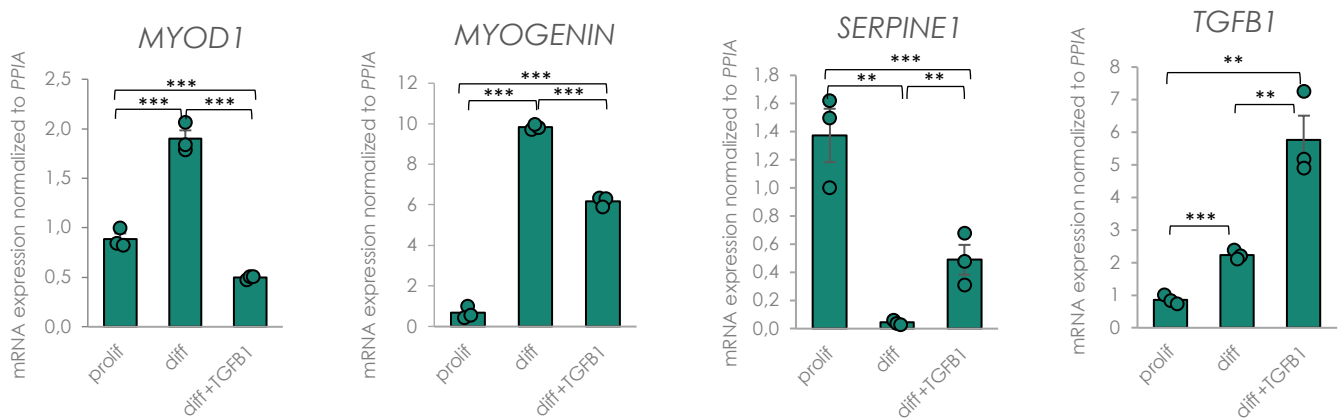
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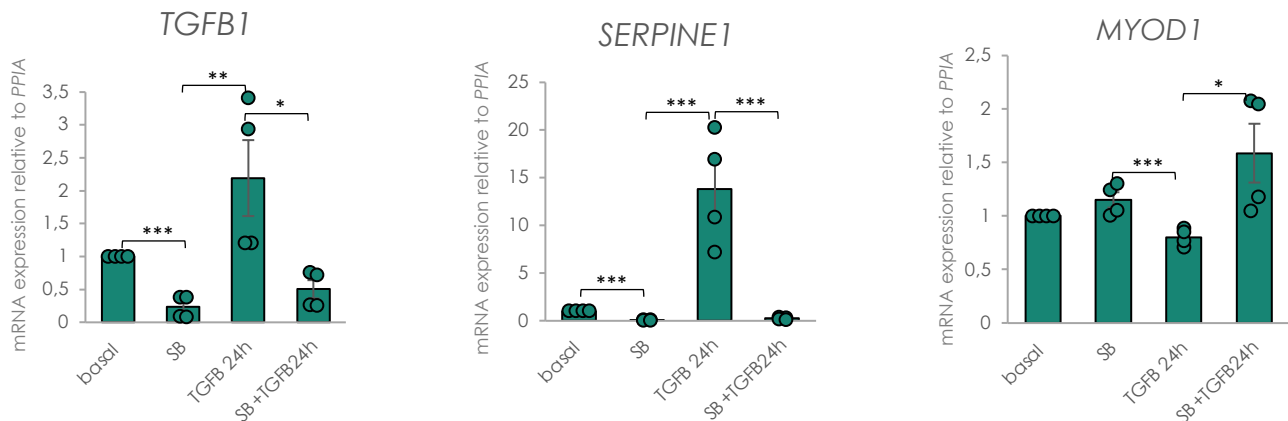
C.



D.



E.



**Figure S1: SETDB1 localization depends on TGF $\beta$ /SMAD pathway activation during muscle terminal differentiation**

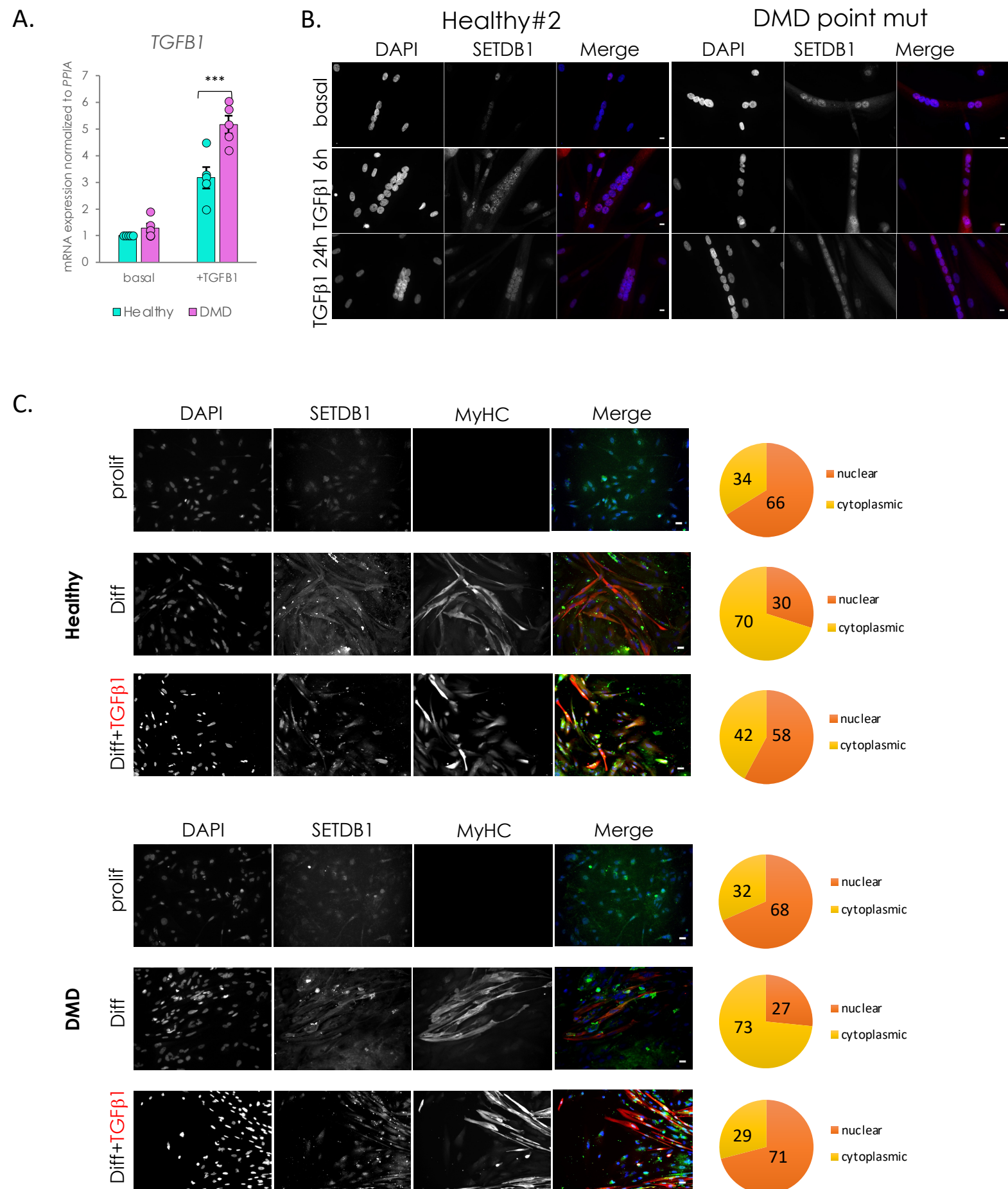
**A.** Scheme of microscopy analysis and nuclear/cytoplasmic signal quantification. Nuclear masks were setup from DAPI pictures and applied on SETDB1 pictures and total cell masks were selected on SETDB1 pictures. Cytoplasmic quantification was performed by subtracting nuclei mask from total cell masks. **B.** Quantification of phospho-SMAD3 nuclear fluorescence in proliferating myoblasts, myotubes treated or not with TGF $\beta$ 1. **C.** Protein levels of SETDB1 and phospho-SMAD3 in total extracts of myoblasts and myotubes treated or not with TGF $\beta$ 1.

**D.** RT-qPCR of early myogenic (*MYOD1*, *Myogenin*) and TGF $\beta$ -related genes (*TGFB1*, *SERPINE1*) in myoblasts and myotubes treated or not with TGF $\beta$ 1. **E.** RT-qPCR of early myogenic marker *MYOD1* and TGF $\beta$ -related genes (*SERPINE1*, *TGFB1*) in myotubes treated or not with TGF $\beta$ 1 and/or its inhibitor SB-431542.

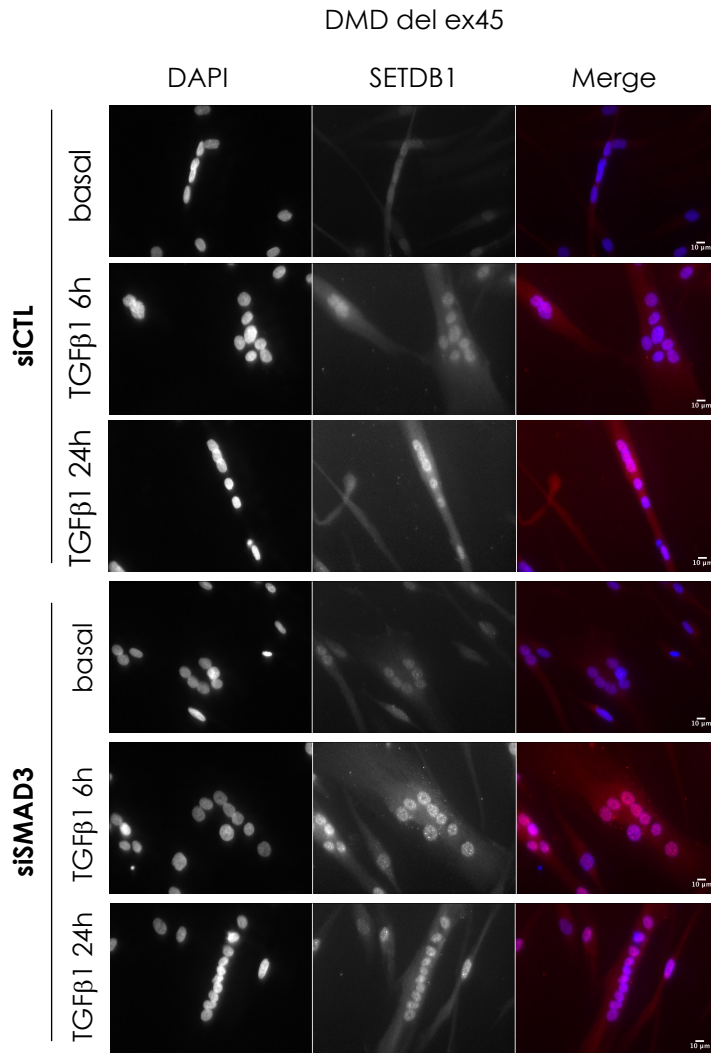
**For all panels:** Statistics were performed on  $\geq 3$  biological replicates ( $>100$  nuclei for immunostaining quantification) and data are represented as average  $\pm$  SEM \* $p < 0.05$ ; \*\* $p < 0.01$ ; \*\*\* $p < 0.001$  (unpaired Student's t test).



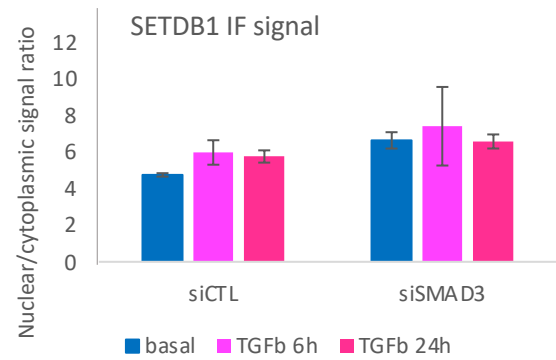
**Figure S2, Granados et al.**



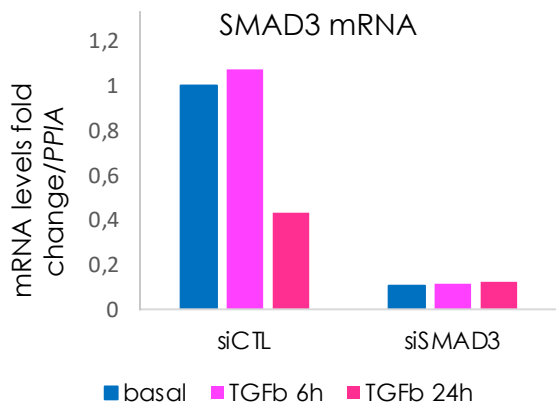
D



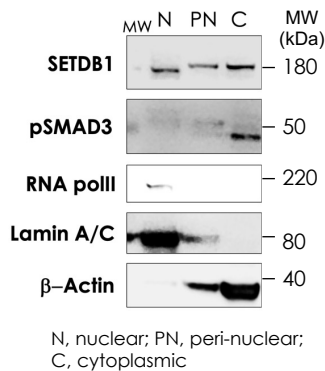
E



F



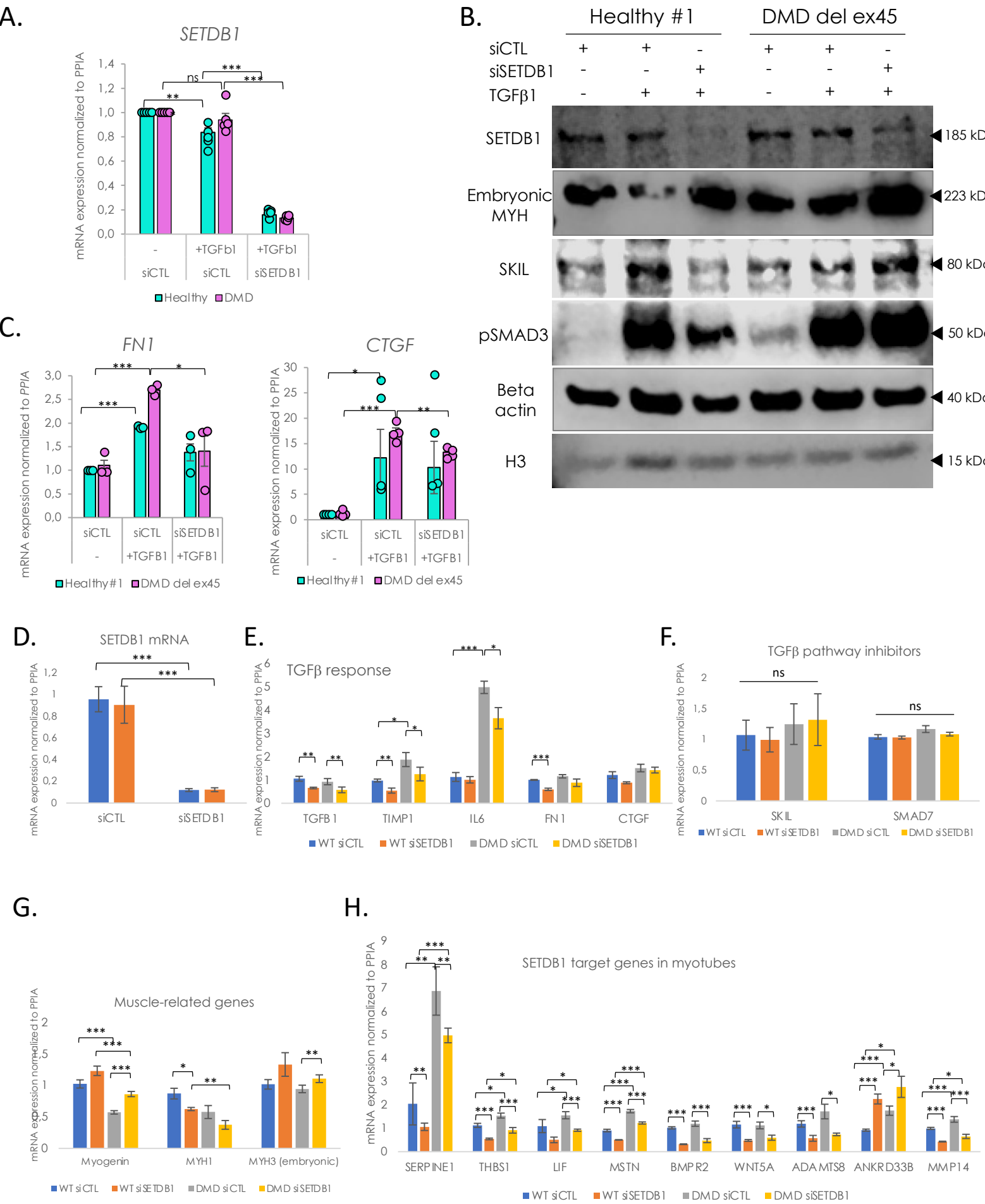
G



**Figure S2: SETDB1 translocate into muscle cell nuclei in response to TGFβ/SMAD pathway activation and show more persistent nuclear signal in DMD myotubes irrespectively of the type of DMD mutation**

**A.** RT-qPCR of *TGFB1* shows a higher response of DMD myotubes to TGFβ1 treatment as compared to WT cells. **B.** Immunostaining of SETDB1 (red) in healthy#2 and DMD point mutation myotubes. Nuclei were stained with DAPI (blue). Scale bar, 10 μM. **C.** SETDB1 and MyHC immunostaining in proliferating muscle cells and in myotubes treated or not with TGFβ1 derived from iPSCs of healthy or DMD individuals. Nuclei were stained with DAPI (blue). Scale bar, 10 μM. Diagrams represent the percentage of SETDB1 signal intensity measure in nuclei and cytoplasm of the cells. **D-F.** DMD del ex45 differentiating myotubes (48 h differentiation) were transfected with control scrambled (siCTL) or SMAD3 siRNA (siSMAD3). 2 days later, myotubes were treated with TGFβ and after 1 day, myotubes were subjected to different assays, as follows: **F.** Immunostaining of SETDB1 (red), and nuclei staining with DAPI (blue). Scale bar, 10 μm. **G.** Quantification of SETDB1 (n=2) nuclear/cytoplasmic IF signal ratio. **H.** RT-qPCR of *SMAD3* mRNA in control siRNA (siCTL) and siRNA against SMAD3 (siSMAD3) conditions to check the siRNA efficiency (representative of n=2). **G.** Western blot showing SETDB1 and pSMAD3 protein migration profiles in nuclear (N), peri-nuclear (PN) and cytoplasmic (C) fractions of healthy myotubes. RNA polymerase II and Lamin A/C were used as controls for nuclear fraction and beta-Actin as a control of cytoplasmic fraction. **For all panels:** Statistics were performed on ≥3 biological replicates (>100 nuclei for immunostaining quantification) and data are represented as average ± SEM \*p<0.05; \*\*p<0.01; \*\*\*p<0.001 (unpaired Student's t test).

Figure S3, Granados et al.



**Figure S3: Efficient siRNA-mediated SETDB1 knockdown in TGF $\beta$ -treated DMD myotubes leads to a decrease in TGF $\beta$  target gene expression and an increase in pro-myogenic factors *SKIL* and *MYH3***

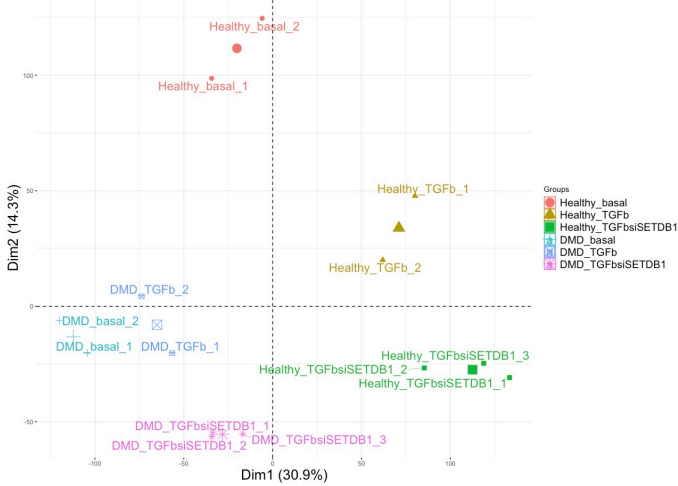
**A.** RT-qPCR of *SETDB1* shows an acute decrease (>80%) of *SETDB1* expression upon siRNAs-mediated silencing. *SETDB1* expression decreases upon TGF $\beta$ 1 treatment in healthy myotubes but not in DMD myotubes. **B.** Western blot showing protein levels of SETDB1, SKIL, embryonic MYH and phospho-SMAD3 in healthy *versus* DMD myotubes upon SETDB1 silencing and TGF $\beta$ 1 treatment. **C.** RT-qPCR of TGF $\beta$ /SMADs pathway known targets *FN1* and *CTGF* in healthy and DMD myotubes +/- siSETDB1 +/- TGF $\beta$ 1. TGF $\beta$ -related gene expression is decreased in SETDB1-silenced *versus* siCTL DMD myotubes in response to TGF $\beta$ 1. **D-H.** RT-qPCR of *SETDB1* (**D**), TGF $\beta$  known target genes *TGFB1*, *TIMP1*, *IL6*, *FN1* and *CTGF* (**E**), of TGF $\beta$  pathway inhibitors *SKIL* and *SMAD7* (**F**), muscle differentiation genes *Myogenin*, *MYH1* and *MYH3* (**G**) and of SETDB1 target genes such as *SERPINE1*, *THBS1*, *LIF*, *MSTN*, *BMP2*, *WNT5A*, *ADAMTS8*, *ANKRD33B* and *MMP14* (**H**) in healthy and DMD myotubes +/- siSETDB1 in basal condition (without TGF $\beta$  treatment).

**For all panels:** Statistics were performed on  $\geq 3$  biological replicates and data are represented as average  $\pm$  SEM \* $p$ <0.05; \*\* $p$ <0.01; \*\*\* $p$ <0.001 (unpaired Student's  $t$  test).

Figure S4, Granados et al.

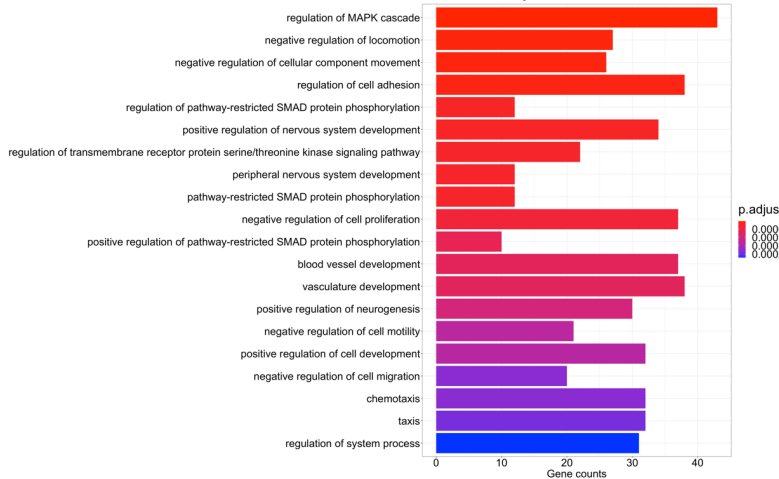
A.

Principal component analysis



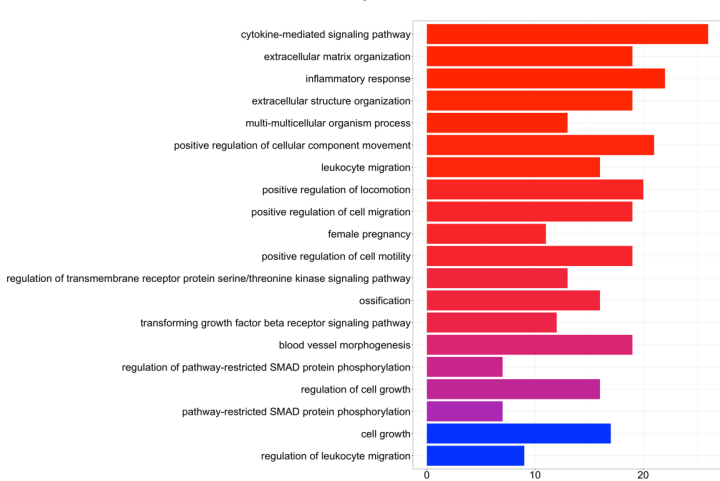
B.

Gene ontology – Healthy myotubes basal versus TGFB



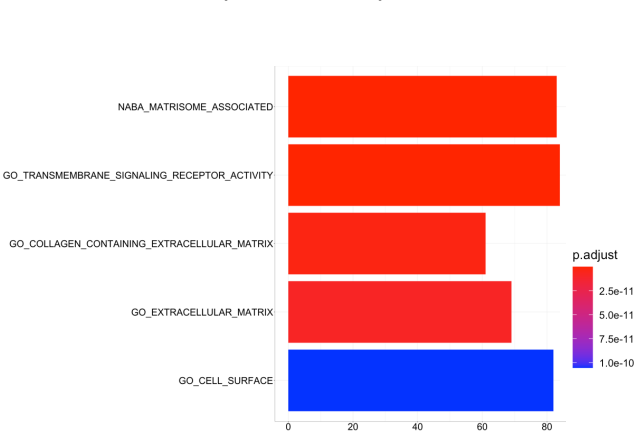
C.

Gene ontology – DMD myotubes basal versus TGFB



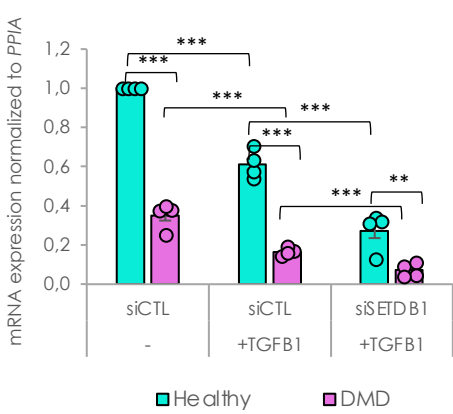
D.

GSEA - Healthy myotubes TGFB versus TGFB+siSETDB1

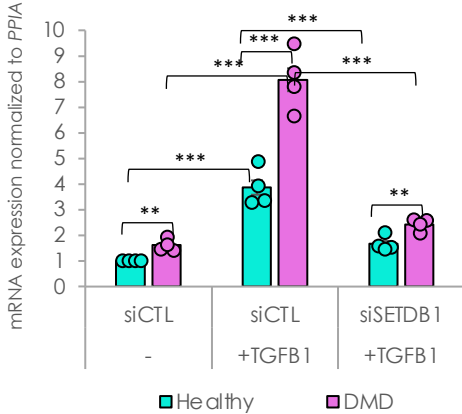


E.

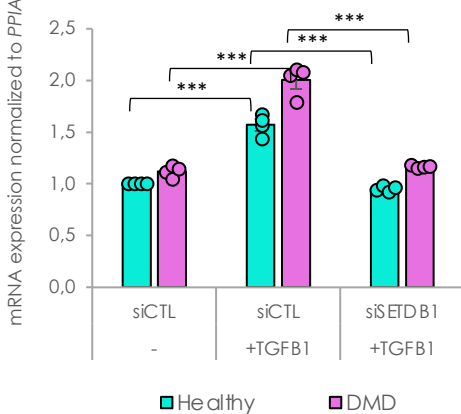
MSTN



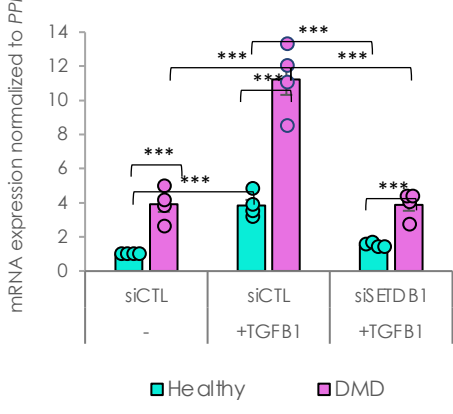
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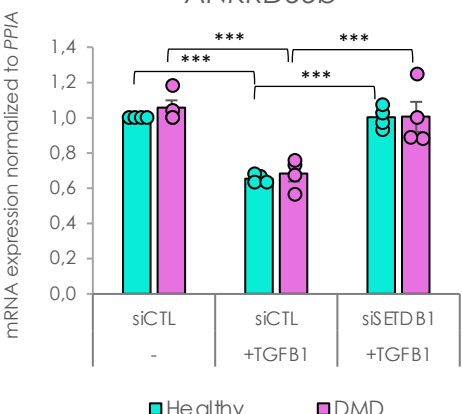
WNT5A



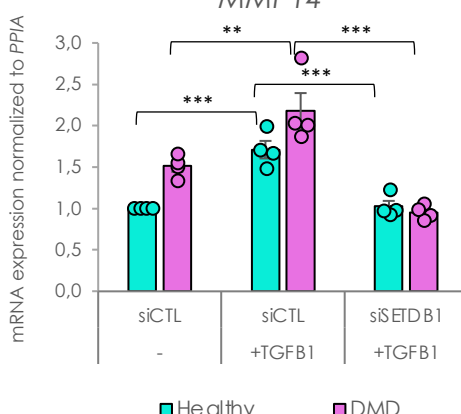
ADAMTS8



ANKRD33b



MMP14





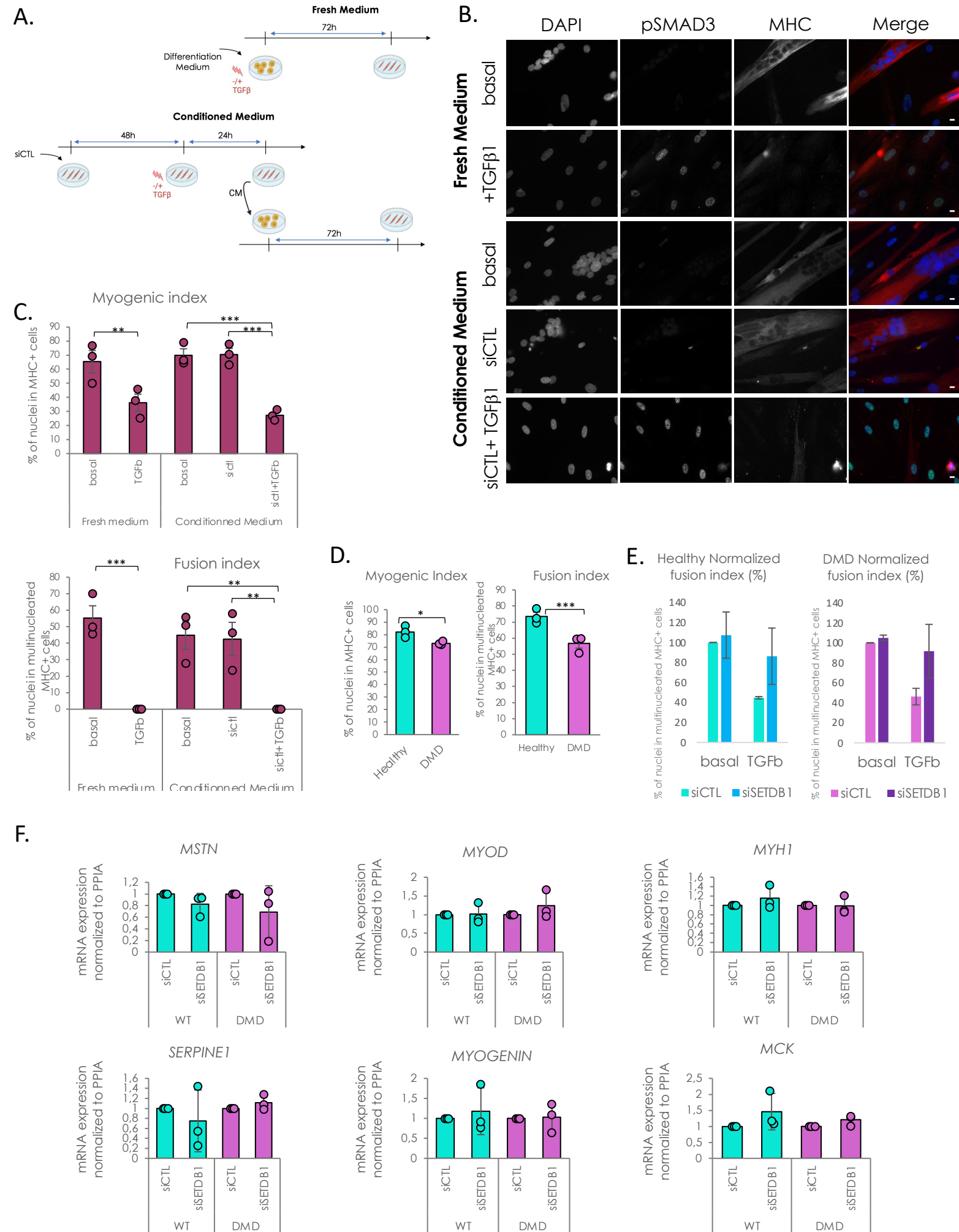
**Figure S4: Healthy and DMD myotubes respond differently to TGF $\beta$ /SMAD pathway activation but display some SETDB1 target gene signatures in common**

**A.** Principal component analysis of the filtered (see Materials & Methods) RNA-seq data, depicting a well grouping of all the samples for each of the 6 experimental conditions of the study. The variability captured by the first PC corresponds to the healthy vs. DMD conditions and the one of the second PC to the TGF $\beta$  treatment. **B.** Gene Ontology enrichment of DEGs from healthy myotubes basal *versus* TGF $\beta$  comparison. **C.** Gene Ontology enrichment of DEGs from DMD myotubes basal *versus* TGF $\beta$  comparison. **D.** Gene Set Enrichment Analysis (GSEA) of DEGs from healthy myotubes TGF $\beta$  +/- siSETDB1. **E.** Validation by RT-qPCR of genes coding for proteins involved in ECM-remodeling (*ADAMTS8*, *MMP14*), TGF $\beta$ /BMP and Wnt pathway (*MSTN*, *BMP2*, *WNT5A*) and unknown function but predicted as regulator of muscle differentiation (*ANKRD33B*).

**For all panels:** Statistics were performed on  $\geq 3$  biological replicates and data are represented as average  $\pm$  SEM \* $p < 0.05$ ; \*\* $p < 0.01$ ; \*\*\* $p < 0.001$  (unpaired Student's t test).



## Figure S5, Granados et al.

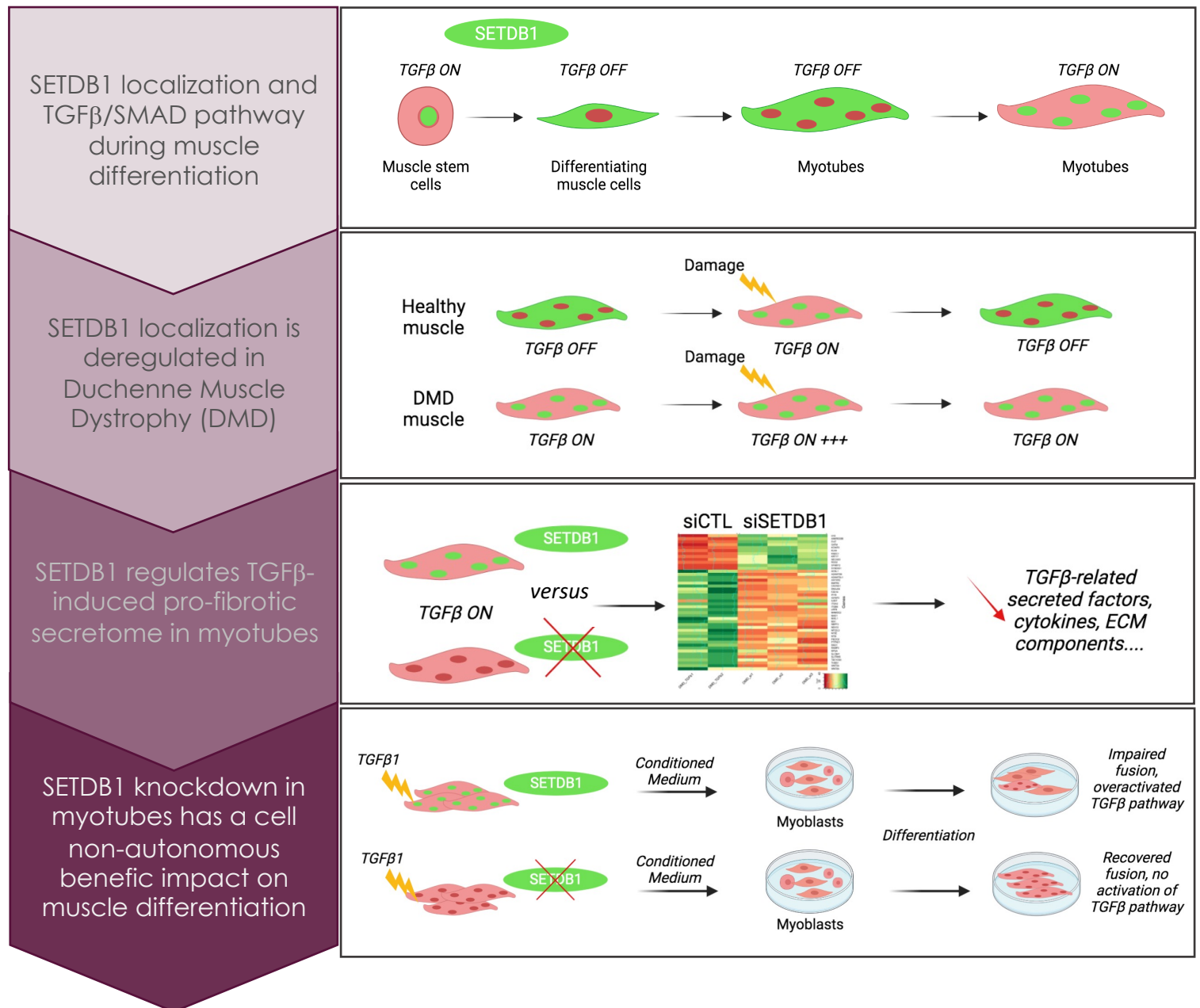


## Figure S5: TGF $\beta$ /SMAD pathway activation leads to fusion defects in muscle cells

**A.** Diagram of the experimental design. **B.** Immunofluorescence of pSMAD3 (green) and MHC (red) in myoblasts differentiated in fresh or conditioned medium +/- TGF $\beta$ 1. Nuclei were stained with DAPI (blue). Scale bar, 10  $\mu$ M. **C.** Myogenic and fusion index of the myoblasts differentiated in fresh or conditioned medium +/- TGF $\beta$ 1. **D.** Raw myogenic and fusion index in basal conditioned medium show difference in differentiation rate between healthy and DMD myotubes. **E.** Quantification of fusion index after 6 days of differentiation in conditioned medium produced by healthy or DMD myotubes +/- siSETDB1 +/- TGF $\beta$ 1. **F.** RT-qPCR of pro-fibrotic genes, *MSTN* and *SERPINE1*, and muscle-related genes *MYOD1*, *Myogenin*, *MYH1* and *MCK* after 3 days of differentiation in conditioned medium produced by healthy or DMD myotubes +/- siSETDB1.

**For all panels:** Statistics were performed on  $\geq 3$  biological replicates ( $>100$  nuclei for immunostaining quantification) and data are represented as average +/- SEM \* $p < 0.05$ ; \*\* $p < 0.01$ ; \*\*\* $p < 0.001$  (unpaired Student's t test).

**Figure S6, Granados et al.**



**Figure S6: Graphical Abstract**

- TGFβ induces nuclear accumulation of SETDB1 in healthy myotubes
- SETDB1 is enriched in DMD myotube nuclei with intrinsic TGFβ pathway overactivation
- SETDB1 LOF in DMD myotubes attenuates TGFβ-induced pro-fibrotic response
- Secretome of TGFβ-treated DMD myotubes with SETDB1 LOF is less deleterious on myoblast differentiation

**Table 1:** Top DEGs + versus – TGFB1 in healthy myotubes

Genes	LogFC	Adj p-value	Genes	LogFC	Adj p-value	Genes	LogFC	Adj p-value
TAGLN2	2,117414122	0,0000029	FZD1	-1,709895607	0,000820576	CYGB	1,91232727	0,004636356
PMAIP1	1,822825158	0,0000295	CPT1A	1,892601801	0,000847677	CHRNA9	2,131251441	0,004655553
SPSB1	1,954444533	0,0000431	FLRT3	-1,666029718	0,000907678	HES6	1,511256537	0,004686457
LEFTY2	2,378015571	0,0000633	SPOCD1	2,072190972	0,000955143	BHLHE40	1,577516443	0,004688974
MYL10	2,775733677	0,000102766	KY	1,741592873	0,000987349	DUSP15	1,504729237	0,004688974
NGF	1,657626522	0,000102766	MMP24	1,785794877	0,001060517	PRTG	-1,939105695	0,004688974
HSD11B2	2,056381007	0,00014734	PTPRN	1,572910987	0,001060517	TAS2R50	-1,617874683	0,004707551
CPEB1	2,008260893	0,00014734	SPHK1	1,553357986	0,001060517	CAPG	1,617433777	0,004761555
GRIK3	1,897772958	0,00014734	SLC6A15	-1,604281321	0,001060517	IL15	1,632420492	0,004800381
CAMK2B	1,66141848	0,00014734	RGS10	1,775216817	0,001091293	RCSD1	1,50536248	0,005066626
LIF	1,609459933	0,00014734	ACAP1	-1,518147984	0,001091293	BCHE	-1,514123565	0,00508091
FAS	1,65594723	0,000176389	CDH22	1,952879439	0,001160065	HAPLN1	-1,939131798	0,005275934
COMP	3,079879046	0,000193172	VDR	2,38037933	0,001185628	TRAPPC11	-1,683727235	0,005288437
KRT17	2,249228221	0,000193172	MADCAM1	2,012788312	0,001218815	E2F2	-1,769263425	0,005304981
CYSLTR1	-1,686154285	0,000193172	IL11	1,720981756	0,00125013	ABHD1	-1,604574292	0,005424992
EFNA5	-1,539212907	0,000193172	ATF3	1,505914322	0,001406572	EDN3	-2,42948784	0,005452562
CBFA2T3	1,585761128	0,00021232	BNC1	1,727769999	0,001463008	KCNG3	-2,006427044	0,005469823
TINAGL1	2,040560934	0,000214261	UGT8	-2,091670121	0,001463008	MT2A	2,078060952	0,005630917
ST6GALNAC2	2,621131534	0,000278225	MDGA1	-1,57495199	0,001500196	MMRN1	-5,596523239	0,005630917
MYL7	1,928998074	0,000278225	LRRN3	-2,023789769	0,001564602	NPPB	2,22275891	0,005782479
RAB11FIP4	1,68799875	0,000278225	APCDD1L	1,573817321	0,001603986	SFRP1	-1,766514568	0,005851596
CADM2	-1,756947008	0,000278225	LEFTY1	2,948081363	0,001646694	NUPR1	2,251896144	0,006190195
KCNH3	1,526606981	0,000294724	INSC	1,984068026	0,001654329	MAPK10	-1,765559007	0,006190195
SH2D4A	1,542140036	0,000304292	FGFBP3	-1,960631592	0,001750743	TENM1	-1,823947329	0,006245981
BMP8A	1,66627264	0,000311971	LRP2	-2,613936968	0,001789815	ABCA7	-1,6412153	0,006245981
UBASH3B	1,556460437	0,000311971	KCNK3	2,416533797	0,001890051	ADAMTS3	-1,554488887	0,006363145
GAL	2,095209135	0,000339293	DSCAML1	2,603034024	0,001909552	SPATA22	1,86416685	0,006490852
FZD5	-2,217424853	0,000339293	LPAR4	-2,840036833	0,002027283	MAB21L2	-2,344727736	0,006490852
ERBB4	-1,608195924	0,000339293	BMP6	1,797677275	0,002067634	PREX2	-1,895214593	0,00654097
JUNB	1,651100338	0,000363745	DRD2	1,911310104	0,002119699	LRRIQ1	-2,230304268	0,006747866
SYTL1	2,38078558	0,000381256	INHBA	1,943531045	0,002198855	PLXDC1	1,720148106	0,006814196
APOBEC3C	1,532547029	0,000381256	HHIPL2	-3,371136543	0,002252125	CBR1	2,746093702	0,006991214
SNAI2	1,80924925	0,000391447	CRLF1	1,941126358	0,002294054	PTPRZ1	-2,289835098	0,006997138
RASL10A	1,877442599	0,000415202	ADAMTSL2	1,594546261	0,002388538	GATA3	1,58890351	0,007566377
PDK4	2,83974063	0,00043132	FZD3	-1,5642083	0,00246262	SLC6A3	-1,725541298	0,007724626
FBLN1	-1,588350862	0,000444297	MKI67	-1,620940441	0,002506658	EPHA4	-1,949119966	0,007737852
BMP3	-1,918899291	0,000449801	CXCR2	1,854163088	0,00256305	LTBP4	-1,510220625	0,008205986
LAMP5	1,686374304	0,000451403	FOSL1	1,871421395	0,002601514	RASD1	1,776910188	0,008344487
NDRG1	1,551820455	0,000451403	ZNF385B	-1,81912287	0,002601514	FAT4	-1,540323008	0,008352329
KCNT2	-1,6565026	0,00046383	SLC15A3	-1,776559618	0,002623801	TAS2R3	-2,803397109	0,008662833
PCDH18	-1,908015374	0,00048586	GPR26	2,554885167	0,002627384	FAM228B	1,920625724	0,008786462
C1QL3	2,750006723	0,000496238	PHLDA2	1,912747543	0,002627384	CGB5	1,997764466	0,008933737
SUMO4	-1,986991477	0,000506112	RRAD	2,053971838	0,002728733	DUSP2	1,65487497	0,008983737
ARHGDIB	1,850929182	0,000509686	FREM2	-1,565177808	0,002728733	ROBO1	-1,537260889	0,008983737
PCSK6	1,711984469	0,000509686	CAV1	1,914603656	0,002899024	GPR183	2,015068932	0,009086429
IRF1	1,590773748	0,000509686	WNK4	1,997056036	0,002931427	TTF1	-1,94369392	0,009313694
KIAA1211L	2,76892938	0,000517694	UGT3A1	-1,686489198	0,003345925	TAS2R31	-2,897454085	0,009353729
BBC3	1,535319531	0,000517694	MAPRE2	1,610130633	0,003380818	CD34	-3,228526136	0,009577181
ROBO2	-2,353637478	0,000534263	KDR	-1,538782171	0,003406179	ZNF493	-1,597943058	0,009582422
MSTN	-2,419217607	0,000572993	TFAP2C	1,719741735	0,003623613	OSGIN1	2,246915673	0,009631694
SFRP2	-2,547991562	0,00060178	PODN	1,685594321	0,003623613	ANK1	1,501084248	0,009652588
ZNF488	1,613702593	0,000668475	ASPH	-1,680643217	0,003696591	SCN3A	-1,978797967	0,009826333
CYP26C1	-2,60465348	0,000697753	NPR1	-1,93870055	0,00402619	SAC3D1	1,586634217	0,009949286
CDH6	-1,630541842	0,000707553	HES5	-5,038583814	0,004080276	TMEM168	-2,739006335	0,009993603
ADAMTS5	-1,875618352	0,000718764	SRGAP2	-1,614350913	0,004106754			
MX1	3,4674981	0,000725331	ISG15	1,862365951	0,004133911			
HMOX1	1,659405071	0,000725331	BIK	1,73069843	0,00451334			
HIST1H4D	-1,571617314	0,000814482	CACNG7	-1,885120997	0,004522138			
CDK5R2	1,65690867	0,000820304	IRS4	-2,193265101	0,004527926			
BMP5	-1,950914501	0,000820304	SLC7A8	-1,852506781	0,004527926			

**Table 2:** Top DEGs + versus – TGFB1 in DMD myotubes

Genes	LogFC	Adj p-value	Genes	LogFC	Adj p-value
TGFB1	2,75040849	5,16E-08	BNC1	2,22770933	0,001244942
IL32	2,163620694	2,67E-07	QPCT	1,556242386	0,001244942
LIF	2,335338653	4,04E-06	MYL10	1,972655907	0,00146367
TAGLN2	1,614876963	5,10E-06	C6orf15	1,678899961	0,001517905
LOX	2,910516076	1,49E-05	ACTN3	-1,617310124	0,00159821
IL11	3,500409887	1,56E-05	S100A11	1,542630473	0,001754586
ANGPTL4	2,505569257	1,56E-05	HMOX1	1,644698523	0,002038795
MARCH4	2,157892488	1,56E-05	PARP14	1,696207068	0,002076847
ITGA2	1,980632839	1,56E-05	FSTL3	2,029350269	0,002148251
APCDD1L	2,725494981	1,81E-05	NLRC5	3,843801654	0,00225234
CCL4	5,438924629	2,45E-05	WNK4	2,802240205	0,00225234
PRRT2	-1,567243856	3,16E-05	SLCO2B1	2,732760296	0,00225234
SPOCD1	2,629199572	4,54E-05	ARHGDIB	2,226199038	0,002446559
GAL	2,598934135	6,15E-05	PTHLH	4,711087434	0,002450317
PMEP1	1,967218429	6,15E-05	IL6ST	2,123553616	0,002450317
SEMA7A	1,911459216	6,15E-05	F2RL1	1,701706227	0,002546264
SIM2	3,167334171	6,29E-05	CPA4	1,642588269	0,002547328
SERPINE1	2,798165062	6,57E-05	TRIML2	2,65612546	0,002803042
INHBA	4,186346592	7,54E-05	RRAD	2,356494017	0,002830075
GDF15	1,973998395	8,06E-05	CAV2	1,944528389	0,002907175
JUNB	1,952571749	8,06E-05	DUSP5	1,614895079	0,002978339
OXTR	2,296104211	8,32E-05	LTBP2	2,603423885	0,003245881
LAMC2	2,448525099	0,000121314	RAMP1	1,731342839	0,003662479
CRLF1	3,222009526	0,00013696	TAC1	1,590107658	0,003700528
AMIGO2	2,19537543	0,000188579	GPR87	2,290047196	0,003705592
POU2F2	1,920477198	0,000199134	EBI3	1,834331335	0,003890978
ANXA4	1,524080736	0,000203676	CLCA2	1,568271593	0,004203243
NGF	1,575477534	0,000214729	MX1	2,151667583	0,004673777
ADAM12	2,627168827	0,000241171	NPFFR2	2,071848324	0,004673777
PTPRN	1,898344697	0,000278524	FOSL1	2,033533552	0,004848003
TNFRSF11B	2,052407425	0,000306379	C1QL1	-1,518618864	0,00567038
FN1	2,097273737	0,000316043	CHRNA9	2,341930935	0,006317128
LDLRAD4	1,885583009	0,000366423	PROKR2	2,187361985	0,006317128
SPON2	1,664023037	0,000367141	CDKN2B	2,078064312	0,006317128
TES	1,638095057	0,000367141	C1QL3	2,251176908	0,006486544
LEFTY2	1,665034244	0,000405062	CLDN4	1,75335273	0,006486544
RAG1	3,975923848	0,000483132	MSTN	-1,615087924	0,006486544
MMP9	1,928522542	0,000499646	HSPA12A	1,709492127	0,006879139
PIF1	-1,50786107	0,000529838	CGB5	2,351065776	0,007003126
CXCL1	2,816696053	0,000539842	ANKRD1	1,667202737	0,00716927
NT5E	2,187868134	0,000665576	SRPX2	1,946225219	0,007347603
TAP1	1,706154724	0,000665576	GPR26	1,770766319	0,007671595
BHLHE40	2,552101744	0,000728008	COMP	1,563971118	0,007671595
IER3	1,535483982	0,00095399	GPR183	2,697987123	0,008448068
GLIPR1	2,359671334	0,001126099	CYGB	1,810473194	0,008551238



**Table 3:** Top DEGs siSETDB1 versus siCTL during TGFB response in WT myotubes

Genes	LogFC	Adj p-value	Genes	LogFC	Adj p-value	Genes	LogFC	Adj p-value
MAMDC2	-1,520498038	5,65E-05	ARSJ	-1,061549661	0,008451464	RPS6KL1	-1,267489634	0,023290191
SETDB1	-2,312260334	6,09E-05	FLT1	-1,332461615	0,008613958	CLCA2	-1,650785368	0,024689191
ACSL1	-1,455424424	0,000224437	HAS2	-1,247511382	0,008751755	KCNG3	-2,004431023	0,024951235
THBS1	-1,301394066	0,000224437	HMCN1	-1,044961141	0,008751755	APLNR	-2,306740676	0,027475948
BMPR2	-1,176640246	0,000224437	GPC6	-1,294102823	0,008794726	LAMA1	-1,105272513	0,028027906
ATP8B2	-1,05885524	0,000224437	ITGB6	-1,119913491	0,008794726	GSDMA	-1,111110953	0,028371421
PTGFRN	-1,53323687	0,000474681	LRP8	-1,042762389	0,008794726	GRK4	1,686838586	0,028644611
LIMA1	-1,175745808	0,000501383	ST8SIA2	-1,054476505	0,009159413	FREM1	-1,676226071	0,029134271
IGFBP3	-1,471141886	0,00057845	SCN11A	-1,758429933	0,009303076	MAN1A1	-1,378856426	0,029134271
TMED7	-1,443559104	0,00057845	DOK3	-1,010817943	0,009434878	NAALADL2	-1,525850019	0,029709578
TMEM109	-1,281138773	0,00057845	SUSD5	-1,206994107	0,009994918	LAMA2	-1,445964152	0,029886349
IGSF22	-1,179795818	0,00057845	MPZL2	-2,408914913	0,010148671	RPL13	1,177052671	0,030194496
NTM	-1,097552605	0,000596109	LRRN3	-2,423741639	0,010347444	LRRC2	-1,122755043	0,030247404
GRIK3	-1,382794973	0,000633272	DSCAML1	-1,583808697	0,010432176	PCDH18	-1,142379056	0,030682091
PRSS23	-1,21234796	0,000633272	HMOX1	1,01627009	0,010534753	CLDN22	-1,451975925	0,030829524
PARM1	-1,196752186	0,000723086	SLC39A11	-1,348927723	0,011182178	SCN2A	-1,301431103	0,031312572
RAMP2	-1,709990009	0,00072624	ENPP6	-1,329673444	0,011182178	DNAJC22	-1,866109729	0,031497413
PCDHA4	-1,179801452	0,000866001	SLC27A1	-1,039664553	0,011182178	PRKCQ	-1,748274946	0,031643005
SLC8A1	-2,168674612	0,001035925	KY	-1,136090745	0,011569641	TLL2	-1,382705171	0,031643005
IGSF3	-1,452586062	0,001260586	HOXA1	-1,869923761	0,011589371	HPDL	-1,0274529	0,031691539
UGGT2	-1,373806653	0,001260586	WNT8A	-1,723404066	0,011605214	CLGN	-1,00964704	0,031709166
CD109	-1,206613739	0,001260586	DCX	-1,172252082	0,011605214	SMARCD1	-2,674991532	0,032096292
ZFYVE27	-1,146034744	0,001260586	CHRM3	-1,328414995	0,01184128	C1QL3	-1,110617004	0,032165845
AKR1C3	1,196726021	0,001288236	ARSB	-1,05339903	0,01184128	KIAA1549L	-1,301619635	0,032682881
ARHGAP26	-1,44235698	0,001370776	RAG1	-1,520711756	0,0120045	TMED8	-1,039140935	0,033560494
GPR137C	-1,350324674	0,001387303	LRFN2	-1,269349114	0,012103738	OAF	-1,149040279	0,033574106
FGF13	-1,003563077	0,001387303	NFIA	-1,56438338	0,01222235	ISM1	-1,131304626	0,033574106
NRP2	-1,303642909	0,001435207	SLC4A8	-1,423660225	0,012780684	LRRC37A	-1,179829418	0,033755698
OR2H2	-3,608774047	0,001483697	ITGA4	-1,093763889	0,012860251	PER3	-1,000955169	0,034677493
TMEM87B	-1,158116859	0,001779461	MMP25	-1,194320365	0,013314621	LRFN5	-1,607087431	0,035473366
SMTNL1	-1,069024474	0,001779461	OLFML1	-1,985558064	0,013852692	DPY19L2	-1,043176127	0,035841864
TAL2	-1,110677244	0,001884083	CNTNAP3	-1,246093716	0,014099073	MAST1	-1,282552641	0,036121551
CHDH	-1,49571412	0,002016351	CLDN1	-1,078499463	0,014200411	CCL4	1,885798209	0,037313963
PAK3	-1,059555572	0,002178155	CACNG4	-1,004577351	0,01422263	FAT3	-1,074630898	0,037313963
RIMS1	1,002106455	0,002207039	MKI67	-1,476461877	0,014255468	SPIB	-1,503325119	0,037761996
PLCD4	-1,709870369	0,002291852	MXRA8	-1,068346915	0,014255468	CDH4	-1,671716499	0,037870918
POLR3D	-1,05688535	0,002378671	GPLD1	-1,024911621	0,014268962	CBR1	-2,167825525	0,038060638
FRMPD1	-1,092657246	0,002688295	KCNK3	-1,637967552	0,014865901	ADAM33	-1,124290334	0,039108986
CACNA1E	-1,119064387	0,003649151	FAM71F1	1,003725779	0,015144667	UBE2Q2	-1,050446746	0,03956611
GRPR	-1,333005464	0,003905121	PAPPA	-1,235866455	0,015144667	DESI2	-1,177585114	0,040182971
SLC37A3	-1,035129517	0,003905121	GARNL3	-1,06505093	0,017145732	PODN	-1,010497684	0,040182971
PRIMA1	-2,433823631	0,004160811	RNF220	-1,073998499	0,017340312	GJC1	-1,132522857	0,041033645
DKK2	-1,476262445	0,004326207	ABCC9	-1,238756215	0,017696163	COL22A1	1,691834247	0,041291631
GAL3ST4	-1,296668125	0,004399025	CXCL1	-1,582775383	0,018516113	SCN4B	-1,181753524	0,042784027
ZNF107	-1,129045081	0,004399025	BRCA2	-1,44530328	0,018516113	HIST2H3D	-1,10116306	0,042784027
DPY19L3	-1,009527364	0,004399025	DOC2B	-1,133045331	0,018516113	YTHDC2	-1,074434231	0,044167886
KRT80	-1,272269317	0,005432878	THBD	-1,04413245	0,018516113	SLC6A3	1,32166898	0,044288292
TMEM182	-1,076767897	0,005750453	GAS2L1	-1,029502553	0,018516113	CNTN5	-1,150597818	0,044381061
CAMK4	-1,51980501	0,005834327	KDR	-1,412431822	0,019245021	SCN7A	-1,467305844	0,044542055
SPTLC3	-2,155845794	0,006288784	ROR1	-1,044621628	0,019603878	GRIN2A	-1,412058254	0,047051083
FSD1L	-1,190564652	0,006555192	PRDM8	-1,214084634	0,019779272	PCDH17	-1,143147116	0,049034992
KSR2	-1,802189364	0,00723519	VCAN	-1,192075825	0,019779272	TSPAN11	-1,52753589	0,049416366
SEZ6	-1,161203502	0,00723519	CDH6	-1,046088047	0,020401625	DNAJA4	-2,230979822	0,049955982
KIAA1211	-1,005076581	0,00723519	CNTN2	-1,109787194	0,020995772	FAXC	-1,046733564	0,049974705
APCDD1L	1,149403017	0,007527773	NEK10	-1,131980357	0,021155723			
ARNT2	-1,077264685	0,007527773	RRAD	1,152195301	0,021663764			
TENM3	-1,003246377	0,00807358	THEMIS2	-1,141961836	0,021663764			
TENM2	-1,01075277	0,008088306	SPTBN5	-1,200082097	0,022154881			
NT5E	-1,337136771	0,008215045	CARNS1	-1,189226427	0,022569992			



**Table 4:** Top DEGs siSETDB1 versus siCTL during TGFB response in DMD myotubes

Genes	LogFC	Adj p-value	Genes	LogFC	Adj p-value
MAMDC2	-1,637289048	1,59E-05	TBC1D30	-1,656334075	0,017074581
NTM	-1,190266731	0,000624532	TMEM109	-0,852260252	0,019233362
CACNG1	-1,183738436	0,001856085	MIXL1	-1,554596572	0,019233362
MAS1	-1,521127791	0,001856085	WNT8A	-2,267660056	0,020236735
IGFBP3	-1,376670225	0,00219827	RGS2	0,830392825	0,020236735
SLC8A1	-1,793845414	0,002530987	PRSS23	-0,872430004	0,021751043
ACSL1	-1,287676377	0,003389855	HEXB	-0,59923785	0,022743141
KLK4	1,787192625	0,003389855	OLIG3	-1,219844213	0,022795667
NT5E	-1,535868776	0,003691319	ANO2	-0,809761055	0,023330054
RAG1	-3,601343309	0,003691319	CXCL12	-1,052378458	0,023330054
KCNIP2	1,004157391	0,003691319	IL6ST	-1,816664708	0,023330054
ADAMTSL1	-1,233489406	0,003799356	DNAJA4	-1,462582406	0,0237548
ANTXR2	-0,898772223	0,004091009	C6orf118	-1,270419296	0,025251564
LRP8	-0,874823587	0,004452942	UCP2	0,591231944	0,025251564
FZD10	-2,477744095	0,010340651	FHIT	0,974006361	0,025251564
IFI16	-1,095725152	0,010354658	PCDHB5	-0,847006813	0,026603158
ADAMTS8	-0,749833348	0,010652676	MOXD1	-1,300307077	0,026603158
ITGA2	-0,833861933	0,010652676	NECAB3	0,57770924	0,026603158
ANKRD33B	1,309182508	0,010772045	MX1	-1,671442415	0,026673574
VEZT	-0,59658059	0,010948764	IGSF22	-0,898697771	0,027562876
LIMA1	-0,805583317	0,010948764	SETDB1	-1,0498379	0,028675644
ITGB6	-1,363673345	0,011201951	FBLN1	-0,829507714	0,02996258
POLN	-0,950024758	0,011211256	RPSA	-1,790736619	0,033731882
RAMP2	-1,299151192	0,011334712	MATN2	-0,849749358	0,035265666
NBPF3	-1,164405324	0,011446775	MLIP	-1,211291173	0,035922112
CANX	-0,679162696	0,012181954	PGPEP1	0,632953024	0,035922112
BMPR2	-0,753326146	0,012181954	SFMBT2	0,683128284	0,035922112
ZFYVE27	-0,809883445	0,012181954	RAP1GAP	0,71600913	0,035922112
NFE2L3	-0,939940084	0,012181954	NKAIN2	-1,137475403	0,039898636
KNDC1	0,708511646	0,012238511	MARCH2	-0,595011539	0,039948549
WNT5A	-0,703188943	0,013030024	KRT17	0,948663616	0,042060417
TMEM30A	-0,705417259	0,013030024	FRY	-0,58182675	0,042984885
THBS1	-0,73807264	0,013030024	PTGFRN	-0,743069331	0,042984885
NEK10	-1,269427146	0,013030024	RTL1	-0,890963897	0,042984885
PTPN21	-0,775583543	0,013160275	ARRDC2	-1,191091084	0,042984885
FAM20A	-0,869507338	0,013160275	AMHR2	-2,339438416	0,042984885
KIT	-0,917148702	0,013160275	GATM	0,521594077	0,042984885
PIEZO2	-1,18208656	0,013160275	FXYP4	3,180179625	0,042984885
SLITRK5	-1,053308863	0,015640425	ZFP28	-1,866547884	0,045979659
SYNDIG1	0,893507731	0,015866611	DDOST	-0,696025419	0,047060967
LIF	-0,657896325	0,017074581	PRDM8	-1,346055025	0,047060967
LRTM1	-0,907971475	0,017074581	SLA	1,287085305	0,048993455