A tumor suppressor, Rbf, promotes muscle growth and myofibrillogenesis. The loss of Rbf severely affects the myogenic transcriptional program. Rbf activates the transcription of myogenic genes in an E2F-dependent manner. Rbf genetically interacts with usp, ct, and Stat92E in late myogenesis. This work expands the understanding of the plethora of Rbf functions.
Rbf Activates the Myogenic Transcriptional Program to Promote Skeletal Muscle Differentiation

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SUMMARY

The importance of the retinoblastoma tumor suppressor protein pRB in cell cycle control is well established. However, less is known about its role in differentiation during animal development. Here, we investigated the role of Rbf, the Drosophila pRB homolog, in adult skeletal muscles. We found that the depletion of Rbf severely reduced muscle growth and altered myofibrillogenesis but only minimally affected myoblast proliferation. We identified an Rbf-dependent transcriptional program in late muscle development that is distinct from the canonical role of Rbf in cell cycle control. Unexpectedly, Rbf acts as a transcriptional activator of the myogenic and metabolic genes in the growing muscles. The genomic regions bound by Rbf contained the binding sites of several factors that genetically interacted with Rbf by modulating Rbf-dependent phenotype. Thus, our results reveal a distinctive role for Rbf as a direct activator of the myogenic transcriptional program that drives late muscle differentiation.

INTRODUCTION

The retinoblastoma (pRB) tumor suppressor protein is a multi-functional protein that is best known for its role in controlling cell cycle because it inhibits the E2F transcription factor. The functional inactivation of pRB is considered to be an obligatory event in human cancer and underscores the importance of pRB in tumorigenesis. However, whether the ability of pRB to promote cell cycle exit entirely accounts for its tumor suppressive property remains an unresolved question. This is mostly because the inactivation of pRB also reduces cell differentiation, including lipogenesis (Classon et al., 2000), myogenesis (de Bruin et al., 2003; Zacksenhaus et al., 1996), erythropoiesis (Spike et al., 2004), and osteogenesis (Thomas et al., 2001). However, deciphering the actual role of pRB in differentiation in vivo is complicated because the inactivation of RB often leads to ectopic cell proliferation and apoptosis, which may account for the differentiation defects. Thus, it is important to identify the in vivo settings where the bona fide role of pRB in differentiation can be investigated without the accompanied ectopic cell cycles.

The model organisms, such as Drosophila, are advantageous in providing insights into gene function during development. Mammals and flies share significant similarity in the arrangement of the E2F/pRB pathway, whereas the Drosophila E2F and pRB families are smaller. Therefore, redundancy and compensation among family members is attenuated. E2F is a heterodimer between an E2F subunit and a DP subunit. In Drosophila, there are two E2F genes, E2f1 and E2f2, a single gene encoding a DP subunit, and two pRB-related genes, Rbf and Rbf2. Either E2f1 or E2f2 requires Dp to bind to DNA. Therefore, the loss of Dp inactivates both E2Fs. In general, E2f1 is an activator of gene expression and is negatively regulated by Rbf, whereas the repressor E2f2 forms a repressive complex with either Rbf or Rbf2 (dREAM/Myb-MuvB). Unlike Rbf, the expression of Rbf2 is restricted to a few cell types, and it appears to be less significant than Rbf at late stages of development. The phenotype of Rbf mutants is highly reminiscent of the inactivation of RB in mammals and results in increased apoptosis and abnormal cell cycles (Du and Dyson, 1999). Genome-wide studies revealed that Rbf binds to chromatin in an E2F-dependent manner (Korenjak et al., 2012) and can either limit E2f1 activation or repress tissue-specific transcriptional programs in conjunction with E2f2 (Dimova et al., 2003; Lewis et al., 2004). Thus, Rbf is widely considered to be a transcriptional repressor.

Surprisingly, the loss of both E2Fs is permissive for most Drosophila development, and the patterns of cell proliferation and differentiation are largely normal. However, E2F is essential for animal viability because E2F-deficient animals die at the pupal stage because of severe defects in adult skeletal muscles (Zappia and Frolov, 2016) and in the fat body (Guarner et al., 2017). In skeletal muscles, E2F is needed for full activation of myogenic genes during muscle growth and myofibrillogenesis. E2f1 occupies the promoter region of several myogenic genes and directly regulates their expression. Interestingly, in addition to E2f1, Rbf occupancy is also enriched at the same genes. Because Rbf is a known inhibitor of E2f1, one could reason that Rbf may limit the activation of the myogenic E2F target genes and subsequently affect muscle development. However, the role of Rbf in myogenesis has not been studied.

Here, we investigated how the inactivation of Rbf affects the development of the adult skeletal muscles. Unexpectedly, we found that Rbf functions as an activator of the myogenic transcriptional program. The depletion of Rbf during myogenesis phenocopies the loss of E2F, i.e., severe reduction in muscle size, abnormal myofibrillogenesis, and incomplete activation of
the myogenic genes. Importantly, the differentiation defects of Rbf-depleted muscles occur without concomitant ectopic cell cycles. Computational analysis identified putative binding sites of ultraspireacle (usp), cut (ct), and signal-transducer and activator of transcription protein at 92E (Stat92E) in the promoters of the myogenic Rbf targets. The genetic-interaction experiments revealed that Rbf and these factors act synergistically. Thus, Rbf has a direct role in the differentiation during muscle development.

RESULTS

The Effect of Rbf Function in a Cell Culture-Based Model of Myogenic Differentiation Is Conserved between Drosophila and Mammals

Important insights into the role of the mammalian pRB were obtained using an in vitro myogenic differentiation model. Thus, we established a similar cell culture-based model using Drosophila. The Dmd8 cell line is derived from myoblasts of the wings discs of third instar larvae (Ui et al., 1987). We optimized the culturing conditions to robustly induce differentiation of Dmd8 myoblasts into multinucleated myotubes (see STAR Methods). The addition of the differentiation medium resulted in distinct morphological changes over time. This process was accompanied by a robust induction of the master myogenic transcription factor Myocyte enhancer factor 2 (Mef2; Figure 1A). At 6 days, the number of nuclei per cell was markedly increased, and some cells became highly elongated and resembled myotubes (Figure 1B).

To confirm that the observed morphological changes reflect the progression of myogenesis, we monitored the expression of several key myogenic genes that are routinely used as markers in vivo. Differentiation of myoblasts into adult skeletal muscles is accompanied by the upregulation of Mef2 with concomitant downregulation of twist (twi), heartless (htl), and stumps. We examined the expression of these genes at 0, 3, 6, and 9 days after the addition of differentiation medium to Dmd8 cells. Notably, these genes were downregulated through day 9 (Figure 1C, top). The expression of genes encoding for structural proteins, such as flightin, Myosin light chain2, and Tropomyosin1, were induced and increased over time (fln, Mlc2, and Tm1, respectively; Figure 1C, bottom). Overall, our data suggest that the myogenic transcriptional program is properly induced in the Dmd8 cell culture-based model and that the observed morphological changes indeed reflect the formation of myotubes and, in particular, fairly replicate the early phase of the indirect flight muscle development.

Having established the robust protocol of in vitro myogenic differentiation using Dmd8 cells, we set to probe the role of Rbf in this system. Dmd8 cells were incubated with Rbf-double-stranded RNA (dsRNA) to deplete the expression of Rbf by RNAi, and GFP-dsRNA was used as control. The efficiency of Rbf depletion, confirmed by western blot (Figure 1D), had almost no effect on cell proliferation at day 0 as revealed by anti-PhosphoHistone H3 (pH3) staining that marks mitotic cells (Figure 1E). In addition, the analyses at days 3 and 6, upon differentiation, indicated that Rbf-depleted cells exit the cell cycle on time (Figure 1E). However, Rbf-depleted cells were partially impaired in their ability to undergo myogenic differentiation (Figure 1F). The number of nuclei within syncytial myotube was slightly reduced compared with the control (Figure 1F, right). The depletion of Rbf was confirmed by measuring the expression of Rbf at the end of the experiment on day 6 (Figure 1G). Thus, the loss of Rbf in cultured Dmd8 myoblasts partially affects the ability of cells to form multinucleated myotubes but does not affect the cell cycle exit.

The Loss of Rbf Affects the Proliferation of the Adult Muscle Precursors In Vivo

The observation that the depletion of Rbf impairs myotube formation in vitro was intriguing. Thus, we investigated Rbf function in a more relevant in vivo system, such as the developing adult skeletal muscles in Drosophila. We focused on the indirect flight muscles (IFMs) because they most resemble mammalian skeletal muscles, and their development is described in great detail. There are two types of IFMs: the dorsal-longitudinal muscles (DLMs) and the dorsoventral muscles (DVMs). DLMs are formed by remodeling larval muscles that serve as templates to which myoblasts fuse, whereas DVMs are formed de novo by myoblast fusion (Dutta et al., 2004; Fernandes et al., 1991). The myoblasts that give rise to the IFMs are specified earlier during embryogenesis and are located in the adepithelial layer of the wing discs. Myoblasts can be marked with the twi-lacZ transgene that is expressed continuously in these cells until they fuse to form myotubes. Dividing myoblasts were visualized by immunofluorescence using the pH3 antibody. In the wild type, myoblasts actively proliferate during larval and early pupal stages as they
migrate to fuse with developing DLMs (Figure 2A). Around 16 h after pupa formation (APF), the three larval templates split, which results in six myotubes that initiate their compaction and attachment to tendons. Between 24 and 29 h APF, the number of unfused myoblasts (twi-lacZ positive), and consequently, pH3-positive cells, drop significantly. By 33 h APF, the fusion is complete, and no remaining myoblasts are detected (Figure 2A and 2B). Upon completion of fusion, myofibers are thickening and shortening. Then, from mid to late pupa muscles elongate and fill the entire adult thorax.

Such strict order of events during the development of IFMs allowed us to determine precisely the effect of Rbf depletion in muscle differentiation. The early larval lethality of the Rbf mutants precludes such analysis. Therefore, we used RNAi to deplete Rbf specifically in muscles by driving UAS-Rbf-RNAi with the muscle-specific Mef2-GAL4 driver. To confirm the efficiency of Rbf depletion, wing discs of third instar larvae were stained with Rbf antibody and counterstained with Mef2 antibody to label myoblasts. As shown in Figure 2C, Rbf was specifically depleted in myoblasts but not in the adjacent epithelial cells of the wing discs. Notably, the depletion of Rbf with Mef2-GAL4 resulted in fully penetrant pupal lethality, in which most animals died as pharate adults. We confirmed this phenotype with another muscle-specific GAL4 driver how248.

To assess the effect of Rbf depletion on myoblast proliferation, the number of mitotic myoblasts, labeled with pH3 antibody, was quantified at two time points: wing discs and 21 h APF. The number of pH3-positive myoblasts was slightly increased upon Rbf depletion compared with control wing discs (Figure 2D). However, the total number of myoblasts was not significantly affected, indicating that Rbf-depleted myoblasts likely have more-extended mitosis than wild-type cells have. Developing DLMs at 21 h APF were labeled with the antibody 22C10 (Figure 2E). Similarly, the loss of Rbf resulted in an elevated number of myoblasts undergoing mitoses (Figure 2H). However, the timing of the cell cycle exit was not significantly affected because the number of pH3-positive cells dropped dramatically at 29 h APF, which is similar to that of control animals, and was mostly undetectable at 33 h APF (Figures 2F–2H). Thus, the depletion of Rbf results in an increase in the number of proliferating myoblasts but does not delay cell cycle exit during the formation of the indirect flight muscles.

Because Rbf-depleted myoblasts showed an increased mitotic index at 21 h APF, we asked whether some of these cells were eliminated by apoptosis. Developing DLMs were stained with antibodies against the death caspase-1 (Dcp-1), a critical executioner of apoptosis, throughout muscle development (21, 24, 29, 33, 43, 72, and 96 h APF; Figure S1). No Dcp-1 staining was detected in Rbf-depleted muscles or in the control, suggesting that the loss of Rbf does not appear to induce a detectable level of apoptosis in myoblasts. However, other apoptotic or nonapoptotic mechanisms, independent of Dcp-1, may be activated by the loss of Rbf.

**Rbf Regulates Myofibril Formation and Muscle Growth**

During myofibrillogenesis, the developing myotubes are gradually converted into myofiber by forming myofibrils and assembling presarcromeric structures, which dynamically mature into highly ordered repetitive contractile units, known as sarcomeres, as muscles grow and elongate (Fernandes et al., 1991; Reddy and Beall, 1993). A mild defect was already evident upon Rbf depletion at the onset of myofibrillogenesis (Figures 2F and 2G). DLMs at 29 and 33 h APF were dissected and stained with phalloidin to visualize myofibril formation. Premyofibrils structure assembles first at the myofiber surface and then spread throughout the interior of the fiber (Sparrow and Schöck, 2009). Although the surface of developing muscles showed proper formation of myofibrils, the interior plane of DLMs, taken 4 μm below the fiber surface, lacked the repetitive myofibril organization, and some myofibrils were missing (Figures 3A and 3B). In addition, the presarcromeric structures, which had an orderly display in control myofibrils, were abnormally organized in Rbf-depleted muscles. The precise process of assembling a highly regular array of myofibrils and sarcomere depends on the mechanical tension mediated by muscle attachment (Weitkunat et al., 2014). Because assembly was abnormal in Rbf-depleted muscle fiber, we examined the compaction of the fiber by measuring the length of muscle fibers at 29 h APF. Consistent with the observed defects, Rbf-depleted fibers were significantly longer than those of the control, implying a defect in tension (Figure 3C).

Given that Mef2>Rbf-RNAi animals die as pharate adults, we examined the formation of adult muscle at late pupa development (96 h APF) before the lethal stage. Persistence of Rbf knockdown at this developmental stage was confirmed by...
Figure 3. The Loss of Rbf Impairs Myofibrillogenesis and Muscle Growth

(A and B) Two confocal section images of developing DLMs stained using phalloidin at 29 h APF (A) and 33 h APF (B). Interior plane was taken 4 μm below the surface plane.

(C) Comparison of fiber length (μm) between Mef2>mCherry-RNAi and Mef2>Rbf-RNAi at 33 h APF. * indicates a statistically significant difference.

(D) Immunoblot analysis showing the expression levels of Rbf and β-Tubulin in Mef2>mCherry-RNAi and Mef2>Rbf-RNAi.

(E) Mef2>mCherry-RNAi and Mef2>Rbf-RNAi stained with phalloidin and β-PS integrin.

(F) Dual staining with DAPI and phalloidin to visualize myofibrils and nuclei in Mef2>mCherry-RNAi and Mef2>Rbf-RNAi.

(G) mRNA levels of various myogenic genes in Mef2>mCherry-RNAi and Mef2>Rbf-RNAi.

(H) Quantification of number of nuclei, area of DLM4, and average size of nuclei per section of DLM4.

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western blot (Figure 3D). The overall structure of IFM was analyzed by staining both transverse and sagittal cross-sections of thoraces. Notably, we found a severe reduction in muscle size in Rbf-depleted muscles, indicating defective muscle growth (Figure 3E). Additionally, myofibrils were not properly aligned and showed signs of actin clumps and defrayed phenotype (Figure 3F). This indicates that Rbf knockdown affects the proper regulation of muscle growth, as well as myofibrillogenesis, which is likely independent of its role in cell cycle control.

To further characterize the effect of Rbf depletion, we extracted RNA from IFMs and measured the expression of several known muscle-related genes (Figure 3G). In agreement with the phenotype described, the expression of genes encoding for components of sarcomeric structure and myofibrils, such as fln, sarcomere length short (sals), Tropomyosin2 (Tm2), Myosin heavy chain (Mhc), Actin88F (Act88F), and Mlc2, were significantly reduced, along with the myogenic regulators held out wings (how), Mef2, and spalt major (salm).

Because Rbf-depleted myoblasts showed an increased mitotic index at 21 h APF, we asked whether these extra myoblasts were fusing to the developing myotubes. We stained dissected muscles with DAPI and quantified the total number of nuclei per cross section of mature DLM. Notably, the number of nuclei was not significantly changed in Rbf-depleted muscles. Thus, despite an increase in pH3-positive myoblasts at the early stages of myogenesis (Figures 2D, 2E, and 2H), there is no significant effect on the total number of fused nuclei. Thus, our data indicate that either Rbf-depleted myoblasts take longer to complete mitosis or the extra myoblasts are eliminated by an alternative mechanism that is not mediated by Dcp1.

Overall, our data suggest that Rbf has an important role in skeletal-muscle development and that the loss of Rbf affects muscle growth and myofibrillogenesis, which is likely caused by reduced expression of the muscle-specific genes.

**Rbf Activates the Myogenic Transcriptional Program during Muscle Development**

As described above, the loss of Rbf in muscle results in two distinct phenotypes. First, there is increased proliferation in Rbf-depleted myoblasts before their fusion to myotubes. We stained dissected muscles with DAPI and quantified the total number of nuclei per cross section of mature DLM. Notably, the number of nuclei was not significantly changed in Rbf-depleted muscles. Thus, despite an increase in pH3-positive myoblasts at the early stages of myogenesis (Figures 2D, 2E, and 2H), there is no significant effect on the total number of fused nuclei. Thus, our data indicate that either Rbf-depleted myoblasts take longer to complete mitosis or the extra myoblasts are eliminated by an alternative mechanism that is not mediated by Dcp1.

Overall, our data suggest that Rbf has an important role in skeletal-muscle development and that the loss of Rbf affects muscle growth and myofibrillogenesis, which is likely caused by reduced expression of the muscle-specific genes.

We compared the wild-type transcriptomes between two developmental stages: proliferating myoblasts in wing discs and fully developed muscles in pharate adults (Figure 4A; Table S1). To enrich for the myoblast cell population, third-instar larval wing discs were dissected, and the notum, which contains myoblasts, was cut and used to isolate RNA. We note that the sample also contained wing disc epithelial cells that may mask the expression of myoblast-specific genes. In total, we have identified, by RNA sequencing (RNA-seq), 27,238 individual transcripts that were expressed in either sample. As previously reported, myogenesis is accompanied by profound transcriptional changes (Spletter et al., 2015, 2018). Consistently, we found that the expression of 10,446 transcripts was activated over time, whereas 11,376 transcripts were repressed between the two developmental points (Figure 4A; Table S2; false-discovery rate [FDR] < 0.05). To determine which of these transcripts are direct targets of Rbf and, hence, can be regulated by Rbf, we isolated chromatin from dissected IFMs of pharate adults and performed chromatin immunoprecipitation sequencing (ChIP-seq) using an Rbf antibody (Table S3). As expected, Rbf occupancy is highly enriched in the promoter and intronic regions of the gene (Figure 4B). In the promoter region, Rbf is preferentially present immediately upstream of the transcription start site (TSS; Figure 4C), which is in agreement with previous findings (Korenjak et al., 2012). We defined Rbf direct targets as transcripts in which Rbf is present nearby the TSS (region 1 kb upstream and 1 kb downstream of the TSS). Using this criterion, we found that, in mature muscles, 4,146 transcripts are direct Rbf targets (FDR < 0.05; fold enrichment ≥ 4; Table S3). Strikingly, during the normal myogenic program almost half of these mRNAs are repressed, and one-third of these mRNAs are activated (2,033 transcripts and 1,432 transcripts, respectively; Figure 4D; Table S2).

Because, in *Drosophila*, the recruitment of Rbf to chromatin is exclusively dependent on E2F (Korenjak et al., 2012), we determined the genomic occupancy of E2F in the mature muscles by ChIP-seq using the Dp antibody (Figures 4B and 4C; Table S4). Remarkably, more than 94% of transcripts identified as Rbf targets were also found to be Dp targets (Figure 4E). Accordingly, the profile of the genomic occupancy of Dp was strikingly similar to the Rbf profile (Figures 4B and 4C). This suggests that both Rbf and Dp are present near the promoters of transcripts that change their expression during normal myogenesis.

To determine the functional importance of Rbf occupancy on these promoters, we isolated RNA from the IFMs at the pharate stage after Rbf depletion with the *Mef2-GAL4* driver. We performed RNA-seq and compared the transcriptome with its
Figure 4. Rbf Activates the Myogenic Transcriptional Program

(A) Heat map representing log2-fold change expression levels of transcripts in the flight muscles at 96 h APF relative to the notum of the third-instar larvae in wild-type animals using Genesis. Total of 27,238 transcripts (muscles versus myoblasts). n = 3 samples of wing disc notums; n = 2 samples of mature muscles.

(B) Genomic occupancy profile of Rbf and Dp in flight muscles of wild-type animals at 96 h APF annotated using Cis-Regulatory Element Annotation System (CEAS); 4,146 transcripts defined as Rbf targets, n = 2 independent samples per antibody and two input samples.

(C) Average profile of Rbf (red, 3,470 peaks) and Dp (blue, 5,148 peaks) mapped reads in 50 bp plotted near the TSS in kilobase using CEAS.

(D) Overlap between Rbf-direct targets (4,146 transcripts) and transcripts that are either repressed (11,376 transcripts) or activated (10,446 transcripts) during myogenic program in wild type (WT; muscles versus myoblasts).

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To identify biological processes that were affected by the loss of metabolic process (Figures 5A and 5B). These results are consis-
tent with the well-known role of Rbf as a negative regulator of cell cycle, DNA replication, and purine and pyrimidine metabolism (Dimova et al., 2003; Lewis et al., 2004; Nicolay et al., 2013).

Matching control (Table S1). We specifically focused on the transcripts that were both direct targets of Rbf and differentially expressed upon Rbf depletion (FDR < 0.05). Among the 2,033 mRNAs that were direct targets of Rbf and were repressed during normal myogenesis, 1,146 transcripts were differentially expressed in Rbf-depleted muscles when compared with the control. Conversely, out of the 1,432 activated-Rbf direct targets, 932 transcripts were differentially expressed upon Rbf depletion (Figures 4F and 4G; Table S2). The latter was unexpected because Rbf is thought to function primarily as a repressor of gene expression, yet the depletion of Rbf prevented the activation of a large number of Rbf targets during muscle development. To analyze how the expression of these transcripts was affected in Rbf-depleted muscles, we grouped the profiles based on their change in expression by hierarchical clustering (Figures 4F and 4G). We found that many Rbf direct targets failed to be repressed (986 versus 1,146 transcripts; Figure 4F; Table S2) and failed to be activated (851 versus 932 transcripts, Figure 4G; Table S2) in Rbf-depleted muscles. The occupancy of both Rbf and Dp near the TSS of the 1,146 transcripts (Figure 4F, right) and 932 transcripts (Figure 4G, right) was visualized with ChIP-tag density plots and compared.

Next, we performed gene-ontology (GO) enrichment analysis to identify biological processes that were affected by the loss of Rbf during myogenesis (Table S5). Genes that are normally repressed by Rbf, hence, those that were increasing in Rbf-depleted muscles (Figure 4F, vertical cyan bar), were significantly enriched for the cell cycle-related categories: DNA repair, response to stress, protein-DNA complex assembly, and DNA metabolic process (Figures 5A and 5B). These results are consistent with the well-known role of Rbf as a negative regulator of cell cycle, DNA replication, and purine and pyrimidine metabolism (Dimova et al., 2003; Lewis et al., 2004; Nicolay et al., 2013).

Interestingly, Rbf target genes that are activated by Rbf and were downregulated in Rbf-depleted muscles (Figure 4G, vertical magenta bar) were significantly enriched for metabolic-related categories, including generation of energy and oxidation-reduction processes, which are known to be important for muscle development and function (Figures 5A and 5C, magenta; P5CDh1, Cyt-c-p, and Zasp66 (Figure 5F, top and bottom)). Significantly, the enrichment was largely lost in Dp-depleted animals, thus further confirming the specificity of the signal. The reduced recruitment was not due to altered levels of Rbf expression in Mef2>Dp-RNAi animals (Figure 5G). Note, that because Dp expression was knocked down only in muscles, there was a substantial contribution of the remaining wild-type tissue in the chromatin samples, which may explain why the loss of Rbf and Dp enrichment in the promoters was only partial after Dp depletion.

Overall, our data support the interpretation that, unlike in mammals, the role of Rbf in activating the expression of the muscle-dependent transcripts is mediated by E2F.

The Expression of Rbf Dynamically Changes throughout Muscle Development

To get insights into the function of Rbf during muscle differentiation, we systematically investigated the expression of Rbf at
multiple time points during flight-muscle development with an anti-Rbf antibody (Figure 6A). Rbf was highly expressed in proliferating myoblasts at early stages of pupal development as myoblasts were migrating to fuse with forming myotubes. Strikingly, the levels of Rbf expression between the myoblasts and the developing myotubes were dramatically different. At 14 h APF, the levels of Rbf were considerably lower in myotubes compared with the surrounding pool of migrating myoblasts. The expression of Rbf remained low in myotubes until fusion was complete, and no myoblasts were detectable at 33 h APF.

To validate that the observed staining accurately reflected Rbf expression, we depleted Rbf throughout muscle development and stained dissected DLMs at 29 h APF with an anti-Rbf antibody. Rbf staining was observed in neither myoblasts nor myotubes, and yet, a strong signal was consistently detected in wild-type sample (Figure 6B). Furthermore, unlike Rbf, both E2f2 and Dp were readily detected in both proliferating myoblasts and in the nuclei of developing myotubes with no difference in their relative expression (Figures 6C and 6D). We note that the Dp antibody also revealed a nonspecific cytoplasmic staining in myotubes because it persisted in Dp-depleted muscles (Figure 6C, right). Hence, the low level of Rbf staining in myotubes is not due to the inability to detect the expression of proteins by immunofluorescence. Next, we examined Rbf expression at later stages of muscle development when the expression of sarcomeric components increases over time, as indicated with the Mhcc-tauGFP reporter (Figure 6E). The Rbf signal was detected in dissected DLMs at 52 h APF and remained present at 80 and 96 h APF (Figure 6E).

We concluded that actively proliferating myoblasts express high levels of Rbf, whereas the expression of Rbf is dramatically reduced as myoblasts undergo fusion to form myotubes.

**Rbf Is Required to Regulate Muscle Maturation in Late Myogenesis**

The results described above show that Rbf expression persists throughout myogenesis. To compare the importance of Rbf between early stages and late stages of myogenesis, the expression of Rbf was temporally and spatially restricted using the TARGET system (McGuire et al., 2004). Rbf was depleted in muscles using the *Mef2-GAL4* driver at different time points with the temperature-sensitive GAL4 repressor (GAL80<sup>ts</sup>; Figure 7A, left). Rbf-RNAi was induced at 31°C by inactivating GAL80<sup>ts</sup>, and thus, allowing GAL4 expression. Conversely, at 18°C, GAL80<sup>ts</sup> is functional, and therefore, Rbf-RNAi is no longer active. The animals with *Mef2>Rbf-RNAi, Dicer2*, and GAL80<sup>ts</sup> raised at 31°C throughout development showed about 50% lethality during pupa development (Figure 7A, right). When the system was switched OFF (Rbf-RNAi) was no longer induced at 18°C at either mid or early pupa development, animals displayed the phenotype, suggesting that Rbf is required during pupal development. However, when the system was switched OFF earlier, immediately preceding pupa formation (larvae were moved to low temperature at late third-instar development), no lethality was detected, thus implying that the loss of Rbf in proliferating myoblasts during larval stages is not detrimental to development. Hence, Rbf function appears to be more relevant at later stages of myogenesis than at earlier stages.

To confirm the importance of Rbf during late myogenesis, we depleted Rbf using a late muscle driver, *Act88F-GAL4*. This driver is expressed in the postfusion IFMs, at the onset of myofibril formation (Bryantsev et al., 2012). The overall structure of myofibrils was examined in sagittal sections of *Act88F-Rbf-RNAi;Dicer2* adults and compared to controls. Interestingly, myofibrils failed to display a repetitive organization in Rbf-depleted myofibers (Figure 7B). Most thoraces showed disorganized and wavy myofibrils, and in some cases, a frayed phenotype was detected, in which sarcomeres were barely visible (Figure 7B, bottom). To determine the significance of these defects, we assessed overall muscle function in Rbf-depleted myofibers with a flight test assay. In this assay, animals that fly land on the top of the cylinder walls, whereas flightless flies land at the bottom. Notably, Rbf-depleted animals were flightless compared with controls (Figure 7C). Thus, Rbf is required during the stage of muscle maturation to promote proper myofibril and sarcomere assembly, which is needed for proper muscle function.

**Rbf Interacts with Other Transcription Factors during Myogenesis**

In mammals, pRB interacts with tissue-specific transcription factors other than E2F to promote differentiation (Novitch et al., 1999; Schneider et al., 1994; Thomas et al., 2001). In addition...
to the E2F binding sites, several additional DNA motifs were identified in the sequences around Rbf peaks. The motifs were matched against Drosophila and vertebrate databases of known motifs using the motif comparison tool TOMTOM. Among the transcripts repressed by Rbf (Figure 6D, left), we identified the binding site for the DNA replication-related element factor (Dref), which functions synergistically with an E2F-binding site and is commonly found in the promoters of proliferation-related genes (Hirose et al., 1993). We also found a motif match for the insulin promoter BEAF-32. BEAF-32 cooperates with an E2f2/Rbf repressor complex dREAM to block the activity of the adjacent enhancers (Korenjak et al., 2014).

When the parameters were adjusted to allow for any number of repeats per sequence, and the motif length was between 6 and 14 bp, the binding sites of ESR1/2 and CUX1 were identified in the promoters of Rbf-activated transcripts (Figure 7D, right). The Drosophila orthologs are estrogen-related receptor (ERR) and hormone receptor 83 (Hr83) for the former motif, and cut (ct) for the latter. To further investigate what other transcription factor binding sites are differentially enriched between activated and repressed transcripts, we used the bioinformatics tool STORM to determine the occurrence of the regulatory motif near the Rbf peak summit. The occurrence of the motifs in the promoter of usp, signal-transducer and activator of transcription at 92E (Stat92E), and delta (Di) was about two times greater in the Rbf-activated transcripts compared with the Rbf-repressed transcripts (Figure 7E).

The importance of these sequence motifs in the promoters of Rbf targets was functionally evaluated with genetic-interaction tests. We asked whether knocking down any of the transcription factors that we identified above enhanced the lethal phenotype of Mef2->Rbf-RNAi animals. To increase the range of the Rbf-dependent phenotype, we removed UAS-Dicer2 to allow Mef2->Rbf-RNAi animals to survive through adulthood (Figure 7F). The knockdown of usp, ct, and Stat92E resulted in the lethality of otherwise viable Mef2->Rbf-RNAi animals (Figure 7F). Significantly, the lethal phenotype was enhanced compared with the effects of the knockdown of each transcription factor alone without Rbf (Figure 7F). In contrast, other transcription factors, including Di and ERR, did not enhance Rbf-dependent phenotype in this assay (Figure S2). To control for RNAi specificity, these tests were repeated with several independent RNAi lines.

These results suggest that usp, ct, and Stat92E are important in adult skeletal muscles. To further explore this idea, we depleted these transcription factors with Act88F-GAL4 and performed a flight test as a readout of flight-muscle function. Notably, the loss of usp led to lethality, and the loss of ct significantly impaired flight ability (Figure 7G), whereas Stat92E and Di affected the adults to a lesser extent, and ERR did not affect them. Because the animals Act88F->Rbf-RNAi are already flightless (Figure 7C), it precluded us from testing whether Rbf genetically interacts with those transcription factors in this assay.

We concluded that usp, ct, and Stat92E are important for Rbf function during late myogenesis. Given that the binding sites for these factors occur more frequently at the promoters of transcripts activated by Rbf, we suggest that the interaction between Rbf and these DNA-binding factors may have a role in Rbf-dependent gene activation.

**DISCUSSION**

In this study, we addressed a long-standing question about the role of pRB in terminal differentiation during animal muscle development. We show that the inactivation of Rbf, the Drosophila pRB homolog, results in severe abnormalities in skeletal muscle formation without concomitant ectopic cell cycles. Notably, these structural defects are accompanied by the failure to activate expression of late-myogenic genes during muscle maturation. Our data suggest that Rbf is required for the activation of the myogenic transcriptional program and genetically interacts with other transcription factors, such as usp, ct, and Stat92E. Thus, in addition to the canonical function of Rbf as a transcriptional repressor, we identified the specific developmental context in which Rbf operates as an activator of gene expression (Figure 7H).

There is ample evidence that the ablation of RB affects muscle development. For example, Rb1<sup>-/-<sup> mouse embryos that are partially rescued to birth by low levels of Rb1 expression or by a wild-type placenta exhibit reduced skeletal muscle mass and disorganized myofibril structure (de Bruin et al., 2003; Zacksenhaus et al., 1996). However, this defect is accompanied by high levels of both apoptosis and ectopic cell cycles, which complicate the interpretation of the phenotype. In flies, the inactivation of Rbf similarly affects cell proliferation, but these defects are restricted to early stages of myogenesis when myoblasts are actively dividing. Rbf-depleted myoblasts exited the cell cycle on time, and no ectopic cell divisions were detected thereafter. Thus, the number of nuclei per muscle was not significantly impaired. Therefore, the loss of Rbf during myoblast proliferation has no significant effect in myogenesis. This observation is consistent with previous reports showing that cell cycle exit upon terminal differentiation remains largely unperturbed in Rbf-mutant cells or in cells that overexpress E2f1 (Butitta et al., 2007; Firth and Baker, 2005). Thus, Drosophila provides
a powerful system to examine the *bona fide* role of Rbf in myogenesis without unwanted effects associated with ectopic cell proliferation.

The analysis of the Rbf-depleted muscles led to two important findings. First, the depletion of Rbf results in the reduction of the muscle size along with defects in the formation of the sarcomeric structures. Although this phenotype may arise from defective tension during muscle attachment at the onset of myofibrillogenesis, the results of Rbf depletion with the late muscle driver argue that Rbf is directly involved in late-muscle maturation. Second, we show that Rbf activates the myogenic transcriptional program. This conclusion is supported by integration of RNA-seq and ChIP-seq, which identified a large group of direct Rbf targets that failed to be induced upon Rbf depletion. Among them are genes encoding components of the sarcomeric complexes and myogenic regulators, which may help to explain the observed defects in muscle maturation. Rbf was also required for the proper activation of the metabolic genes. In concordance with our results, pRB has been linked to the regulation of the metabolic function in other systems (Dyson, 2016). pRB is required for the expression of genes involved in mitochondrial oxidative metabolism, glucose, and fatty acid metabolism in differentiated C2C12 myotubes (Petrov et al., 2016) and was shown to regulate oxidative phosphorylation genes in muscle and brown adipose tissue in mice (Blanchet et al., 2011). In *Drosophila*, the loss of Rbf was shown to have a profound effect on glutamine metabolism (Niclay et al., 2013). Altogether, our data provide a simple explanation that the reduced size of Rbf-depleted muscles is likely due to the failure to properly upregulate the myogenic transcriptional program.

In *Drosophila*, Rbf is thought to operate primarily as a transcriptional repressor (Dimova et al., 2003; Georlette et al., 2007; Korenjak et al., 2012). These genome-wide studies profiled either the entire animals or cultured cells. Thus, a particular differentiation program executed in a few cells could be missed. Here, we used dissected adult skeletal muscles for both RNA-seq and ChIP-seq. This allowed us to identify an Rbf-specific transcriptional program that operates *in vivo* during muscle differentiation.

One of the outstanding questions is how Rbf activates transcription during myogenesis. Mammalian pRB was implicated in potentiating the activity of tissue-specific factors, such as myogenin and MEF2C in cultured myotubes cells (Novitch et al., 1999; Schneider et al., 1994) and CBFA-1 in Saos-2 cells (Thomas et al., 2001), and to interact with the histone demethylase KDM5A (Benevolenskaya et al., 2005). These interactions were shown to be important for the expression of late-differentiation markers. The analysis of Rbf-bound regions from our ChIP-seq experiment revealed an E2F binding site as the prevailing motif for both the activated and the repressed transcripts. Consistently, there was a remarkable overlap between the occupancy profiles of Dp and Rbf in muscles. In addition, the occupancy of Rbf at several myogenic genes was largely abolished in Dp-depleted muscle (this work and Zappia and Frolov, 2016). Thus, in *Drosophila*, the activation of the myogenic genes by Rbf is likely to be E2F dependent. This is consistent with previous work, in which genomic targeting of Rbf occurred entirely via E2F/Dp (Korenjak et al., 2012). In agreement with this idea, the inactivation of E2F by depleting Dp phenocopies the inactivation of Rbf in muscles (Zappia and Frolov, 2016). However, we cannot completely exclude the possibility that the effects of Dp and Rbf depletion may be different on some promoters. One surprising implication of these results is that E2f1 alone, in the absence of Rbf, is unable to activate the myogenic genes. We note that E2f1 can still activate transcriptions at this time point because the cell cycle E2F targets are upregulated in Rbf-depleted muscles. Therefore, the role of Rbf as a positive regulator of gene expression appears to be limited to genes involved in differentiation. Intriguingly, mammalian pRB functions by reversing the repression of the histone demethylase KDM5A at the promoters of the nuclear encoded-mitochondrial genes during differentiation (Várailaj et al., 2015). Because we found that Rbf is similarly needed for the activation of the metabolic genes including OXPHOS genes during muscle differentiation, it may be interesting to explore whether this mechanism is conserved.

Remarkably, there is an abrupt change in the levels of Rbf protein between proliferating myoblasts and growing myotubes. This raises the possibility that Rbf is degraded upon myoblasts

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**Figure 7. Rbf Affects the Maturation of Flight Muscles in Late Development and Interacts with Other Transcription Factors**

(A) Rbf depletion in muscles was temporally induced with tub-GAL80stop by switching ON the system at 31°C to inactivate GAL80stop and OFF (18°C) at midpupa development, early pupa development, and late third-instar larva. Percentage of animals scored at each developmental stage, n = 52–130 flies/genotype. Genotypes are..., Mef2-GAL4, and w-, tub-GAL80stop UAS-Dicer2; Mef2-GAL4 UAS-Rbf1-RNAi.

(B) Rbf depleted using Act88F-GAL4 in flight muscles at late-pupa development. Confocal sections of sagittal sections of adult thoraces stained with phalloidin. Scale bar, 20 μm. Myofibril morphology quantified by scoring percentage of thoraces displaying phenotype. n = 16 and 22 hemithoraces/genotype.

(C) Flight ability scored as percentage of flies landing on each section of the column, n = 80 and 110 flies/genotype. Genotypes are..., Act88F-GAL4, UAS-Actr2c and w-,... UAS-Rbf1-RNAi.

(D) De novo motif discovery MEME. Sequences repressed by Rbf (986 transcripts and 575 peaks) and activated by Rbf (851 transcripts and 404 peaks), TOMTOM matched the motif against Drosophila (left) and vertebrate (right) databases.

(E) STORM-scanned sequences near Rbp peak in the two groups of transcripts (repressed and activated by Rbf).

(F) Genetic interactions between Rbf and the transcription factors, ultraspiracle (usp), cut (ct), and Signal-transducer and activator of transcription protein at 92E (Stat92E). Knockdown of each gene was driven by Mef2-GAL4. Lethality during development was scored as in (A), n = 140–474 flies/genotype.

(G) Flight test as in (C), n = 180–505 flies/genotype. Genotypes are (F) Mef2-GAL4 UAS-mCherry-RNAi, UAS-Rbf-RNAi/Mef2-GAL4, (G) Act88F-GAL4 UAS-EFGP-RNAi, (H) Model representing how Rbf regulates flight muscle development in vivo. Canonical function of Rbf during myoblasts proliferation (left), Rbf represses the activity of E2F in promoter of cell cycle genes. Role of Rbf as an activator of flight muscle development (right). Rbf directly activates and represses the expression of genes in growing muscles, which is mediated by E2F and other transcription factors, including usp, ct, and Stat92E.

See also Table S6 and Figure S2.
fusión y, posiblemente, una nueva población emergente de Rbf que activa la transcripción durante el crecimiento muscular y la maduración. En efecto, un degradación proteasómica mediada de Rbf ha sido previamente sugerido para ser vinculado con señales de desarrollo (Acharya et al., 2010). Notably, otro regulador muscular, el Rbf proteínas que se desplazan al núcleo de DLMs en el inicio de la maduración muscular (Spletter et al., 2015). Así, cambios en los niveles o localización de reguladores musculares podrían ser un特征。

Interesantemente, sobre tres cuartos de los transcriptos que son directamente regulados por Rbf en DLMs también son blanco de Rbf en larvas (Korenjak et al., 2012). Conciliador, hemos detectado Rbf en la TSS de varios genes específicos de músculo, incluyendo Tm1, how, fln, Mef2, Lmpt, and sald en larva (Zappia and Frolov, 2016). Por lo tanto, ocupación Rbf en genes musculares no aparece correlacionar con el tiempo de su activación, sugiriendo que la presencia de Rbf sola no es suficiente para activar su transcripción. Una posibilidad es que Rbf puede asistir a la transcripción de otros factores en el proceso regresivo de los registros musculares. Esta idea es impulsada por la identificación de los sitios de enlace de cinco factores de transcripción en los espacios de Rbf. En la transcripción programada. Este modelo es respaldado por la asociación de los sitios de enlace de tres factores transcriptuales con respecto a los niveles de Rbf. Intrigante, tres de estos genes, usp, ct, y DI, fueron identificados en el gran escenario de los genes requeridos para la regresión muscular y la función (Schonrrer et al., 2010), aunque su precisa función no ha sido determinada. Otros estudios informaron que el rol de Stat92E en el desarrollo muscular embriónico (Liu et al., 2009) y la implicación de ct en la diversificación de mioblastos que dan lugar a diferentes tipos de músculos (Sudarsan et al., 2001). Por lo tanto, la supresión de usp, ct, y DI con el gen muscular Act88F-GAL4, impide la capacidad de animales para el vuelo, lo que sugiere su implicación en la diversificación de mioblastos y la capacidad de volar, lo que indica su importancia en la capacidad de volar.

Para animales, en los cuales pRB directamente interactúa con estos factores para potenciar su actividad, en Drosophila, la recaptura de Rbf a genes musculares es más mediada por E2F. No obstante, el orden de aparición es similar, tanto en pRB y Rbf función para ayudar a la activación de registros transcriptuales para promover diferenciación. Dado tal conservación notable, Drosophila puede ser un sistema poderoso para investigar el papel del retinoblastoma en el contexto de la regulación de diferenciación.

**STAR METHODS**

Detalladas las métodos se proporcionan en el formato en línea de este artículo y incluyen los siguientes:

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  - Quantification of number of nuclei in muscles
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**SUPPLEMENTAL INFORMATION**

Suplemento de información incluye dos figuras y siete tablas y puede encontrarse en línea en https://doi.org/10.1016/j.celrep.2018.12.080.

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**AUTHOR CONTRIBUTIONS**

M.P.Z. y M.V.F. concebieron el proyecto, diseñaron los experimentos, analizaron los datos, y redactaron el manuscrito. M.P.Z. realizó la mayoría de los experimentos con apoyo importante de A.R., que generó todos los datos relacionados con las Dmd8 células y la knockdown con Act88F patron. A.B.M.K.I. realizó la bioinformática data analysis of high-throughput sequencing data sets. Todos los autores revisaron y editaron el manuscrito.

**DECLARATION OF INTERESTS**

Los autores declaran no tener interés alguno.

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REFERENCES

Acharya, P., Raj, N., Buckley, M.S., Zhang, L., Duperon, S., Williams, G., Henry, R.W., and Arnos, D.N. (2010). Paradoxical instability-activity relationship defines a novel regulatory pathway for retinoblastoma proteins. Mol. Biol. Cell 21, 3989–3901.

Althammer, S., Gonzalez-Vallinas, J., Ballaré, C., Beato, M., and Eyras, E. (2011). Pyicos: a versatile toolkit for the analysis of high-throughput sequencing data. Bioinformatics 27, 3333–3340.

Anders, S., and Huber, W. (2010). Differential expression analysis for sequence count data. Genome Biol. 11, R106.

Ashburner, M., Ball, C.A., Blake, J.A., Botstein, D., Butler, H., Cherry, J.M., Davis, A.P., Dolinski, K., Dwight, S.S., Eppig, J.T., et al.: The Gene Ontology Consortium (2000). Gene ontology: tool for the unification of biology. Nat. Genet. 25, 25–29.

Bailey, T.L., and Elkan, C. (1994). Fitting a Mixture Model by Expectation Maximization to Discover Motifs in Biopolymers (AAAI Press).

Benjamini, Y., and Hochberg, Y. (1995). Controlling the false discovery rate: a practical and powerful approach to multiple testing. J. R. Stat. Soc. B Stat. Methodol. 57, 289–300.

Blanchet, E., Annicotte, J.-S., Lagarrigue, S., Aguilar, V., Clapé, C., Chavey, C., Fritz, V., Casas, F., Apparailly, F., Auwerx, J., and Fajas, L. (2011). E2F transcription factor-1 regulates oxidative metabolism. Nat. Cell Biol. 13, 1146–1152.

Bryanstev, A.L., Baker, P.W., Lovato, T.L., Jaramillo, M.S., and Cripps, R.M. (2012). Differential requirements for Myocyte Enhancer Factor-2 during adult myogenesis in Drosophila. Dev. Biol. 361, 191–207.

Buttitta, L.A., Katzaroff, A.J., Perez, C.L., de la Cruz, A., and Edgar, B.A. (2007). A double-assurance mechanism controls cell cycle exit upon terminal differentiation in Drosophila. Dev. Cell 12, 631–643.

Classon, M., Kennedy, B.K., Mulloy, R., and Harlow, E. (2000). Opposing roles of pRB and p107 in adipocyte differentiation. Proc. Natl. Acad. Sci. USA 97, 10826–10831.

de Bruin, A., Wu, L., Saavedra, H.J., Wilson, P., Yang, Y., Rosol, T.J., Weinstein, M., Robinson, M.L., and Leone, G. (2003). Rb function in extraembryonic lineages suppresses apoptosis in the CNS of Rb-deficient mice. Proc. Natl. Acad. Sci. USA 100, 6546–6551.

Dimova, D.K., Stevaux, O., Frolow, M.V., and Dyson, N.J. (2003). Cell cycle-dependent and cell cycle-independent control of transcription by the Drosophila E2F/RB pathway. Genes Dev. 17, 2308–2320.

Du, W., and Dyson, N. (1999). The role of RBF in the introduction of G1 regulation during Drosophila embryogenesis. EMBO J. 18, 916–925.

Du, W., Vidal, M., Xie, J.-E., and Dyson, N. (1996). RBF, a novel RB-related gene that regulates E2F activity and interacts with cyclin E in Drosophila. Genes Dev. 10, 1206–1218.

Dutta, D., Anant, S., Ruiz-Gomez, M., Bate, M., and VijayRaghavan, K. (2004). Founder myoblasts and fibre number during adult myogenesis in Drosophila. Development 131, 3761–3772.

Dyson, N.J. (2016). RB1: a prototype tumor suppressor and an enigma. Genes Dev. 30, 1492–1502.

Fernandes, J., Bate, M., and VijayRaghavan, K. (1991). Development of the indirect flight muscles of Drosophila. Development 113, 67–77.

Firth, L.C., and Baker, N.E. (2005). Extracellular signals responsible for spatially regulated proliferation in the differentiating Drosophila eye. Dev. Cell 8, 541–551.

Georlette, D., Ahn, S., MacAlpine, D.M., Cheung, E., Lewis, P.W., Beall, E.L., Bell, S.P., Speed, T., Manak, J.R., and Botchan, M.R. (2007). Genomic profiling and expression studies reveal both positive and negative activities for the Drosophila Myb MuVbdRE-complex in proliferating cells. Genes Dev. 21, 2880–2896.

Guarner, A., Morris, R., Korenjak, M., Boukhal, M., Zappia, M.P., Van Rechem, C., Whetstone, J.R., Ramaswamy, S., Zou, L., Frolov, M.V., et al. (2017). E2F/ DP Prevents Cell-Cycle Progression in Endocycling Fat Body Cells by Suppressing dATM Expression. Dev. Cell 43, 689–703.e6.

Heinz, S., Benner, C., Spann, N., Bertolino, E., Lin, Y.C., Latio, P., Cheng, J.X., Murre, C., Singh, H., and Glass, C.K. (2010). Simple combinations of lineage-determining transcription factors prime cis-regulatory elements required for macrophage and B cell identities. Mol. Cell. 38, 576–589.

Hirose, F., Yamaguchi, M., Handa, H., Inomata, Y., and Matsukage, A. (1993). Novel 8-base pair sequence (Drosophila DNA replication-related element) and specific binding factor involved in the expression of Drosophila genes for DNA polymerase α and proliferating cell nuclear antigen. J. Biol. Chem. 268, 2092–2099.

Hubbard, T.J.P., Aken, B.L., Beal, K., Ballester, B., Caccamo, M., Chen, Y., Clarke, L., Coates, G., Cunningham, F., Cutts, T., et al. (2007). Ensembl 2007. Nucleic Acids Res. 35, D610–D617.

Korenjak, M., Anderssen, E., Ramaswamy, S., Whetstone, J.R., and Dyson, N.J. (2012). RBF binding to both canonical E2F targets and noncanonical targets depends on functional dE2F/dDP complexes. Mol. Cell. Biol. 32, 4375–4387.

Korenjak, M., Kwon, E., Morris, R.T., Anderssen, E., Amzallag, A., Ramaswamy, S., and Dyson, N.J. (2014). dREAM co-operates with insulator-binding proteins and regulates expression at divergently paired genes. Nucleic Acids Res. 42, 8939–8953.

Langmead, B., Trapnell, C., Pop, M., and Salzberg, S.L. (2009). Ultrafast and memory-efficient alignment of short DNA sequences to the human genome. Genome Biol. 10, R25.

Lewis, P.W., Beall, E.L., Fleischer, T.C., Georlette, D., Link, A.J., and Botchan, M.R. (2004). Identification of a Drosophila Myb-E2F/RBF transcriptional repressor complex. Genes Dev. 18, 2929–2940.

Liao, Y., Smyth, G.K., and Shi, W. (2013). The Subread aligner: fast, accurate and scalable read mapping by seed-and-vote. Nucleic Acids Res. 41, e108.

Lilly, B., Zhao, B., Ranganayakulu, G., Paterson, B.M., Schulz, R.A., and Olson, E.N. (1995). Requirement of MADs domain transcription factor D-MEF2 for muscle formation in Drosophila. Science 267, 688–693.

Liu, Y.H., Jakobsen, J.S., Valentin, G., Amarantos, I., Gilmour, D.T., and Furlong, E.E.M. (2009). A systematic analysis of Timm function reveals Eya and JAK-STAT signaling as essential regulators of muscle development. Dev. Cell 16, 280–291.

McGuire, S.E., Mao, Z., and Davis, R.L. (2004). Spatiotemporal gene expression targeting with the TARGET and gene-switch systems in Drosophila. Sci. STKE 2004, pl6.

Nicolai, B.N., Gameiro, P.A., Tschöp, K., Korenjak, M., Heilmann, A.M., Asara, J.M., Stephanopoulos, G., Iliopoulos, O., and Dyson, N.J. (2013). Loss of RBF1 changes glutamine catabolism. Genes Dev. 27, 182–196.

Novitch, B.G., Spicer, D.B., Kim, P.S., Cheung, W.L., and Lassar, A.B. (1999). pRb is required for MEF2-dependent gene expression as well as cell-cycle arrest during skeletal muscle differentiation. Curr. Biol. 9, 449–459.

Perez-Llamas, C., and Lopez-Bigas, N. (2011). Gitools: analysis and visualisation of genomic data using interactive heat-maps. PLoS ONE 6, e19541.

Petrov, P.D., Ribot, J., Lopez-Mejia, I.C., Fajas, L., Palou, A., and Bonet, M.L. (2016). Retinoblastoma Protein Knockdown Favors Oxidative Metabolism and Glucose and Fatty Acid Disposal in Muscle Cells. J. Cell. Physiol. 231, 708–718.

Quinlan, A.R., and Hall, I.M. (2010). BEDTools: a flexible suite of utilities for comparing genomic features. Bioinformatics 26, 841–842.

Reddy, M.C., and Beall, C. (1993). Ultrastructure of developing flight muscle in Drosophila, I: Assembly of Myofibrils. Dev. Biol. 160, 443–465.
Robinson, J.T., Thorvaldsdóttir, H., Winckler, W., Guttman, M., Lander, E.S., Getz, G., and Mesirov, J.P. (2011). Integrative genomics viewer. Nat. Biotechnol. 29, 24–26.

Schneider, J.W., Gu, W., Zhu, L., Mahdavi, V., and Nadal-Ginard, B. (1994). Reversal of terminal differentiation mediated by p107 in Rb/C0/C0 muscle cells. Science 264, 287–291.

Schones, D.E., Smith, A.D., and Zhang, M.Q. (2007). Statistical significance of cis-regulatory modules. BMC Bioinformatics 8, 19.

Shin, H., Liu, T., Manrai, A.K., and Liu, X.S. (2009). CEAS: cis-regulatory element annotation system. Bioinformatics 25, 2605–2606.

Sparrow, J.C., and Schock, F. (2009). The initial steps of myofibril assembly: integrins pave the way. Nat. Rev. Mol. Cell Biol. 10, 293–298.

Zappia, M.P., and Frolov, M.V. (2016). E2F function in muscle growth is necessary and sufficient for viability in Drosophila. Nat. Commun. 7, 10509.

Zhang, Y., Liu, T., Meyer, C.A., Eckhoude, J., Johnson, D.S., Bernstein, B.E., Nusbaum, C., Myers, R.M., Brown, M., Li, W., and Liu, X.S. (2008). Model-based analysis of ChIP-Seq (MACS). Genome Biol. 9, R137.
**STAR★METHODS**

**KEY RESOURCES TABLE**

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Antibodies**      |        |            |
| mouse anti-RBF (IF) | Du et al., 1996 | DX2 |
| mouse anti-RBF (WB) | Du et al., 1996 | DX3 |
| mouse anti-RBF (ChIP) | Du et al., 1996 | DX3:DX5 |
| rabbit anti-DP      | Dimova et al., 2003 | 212 |
| rabbit anti-pH3     | Millipore | 06-570 |
| mouse anti-β-PS-integrin | DSHB | CF.6G11; RRID:AB_528310 |
| mouse anti-futsch   | DSHB | 22c10; RRID:AB_528403 |
| mouse anti-β-tubulin| DSHB | E7; RRID:AB_528499 |
| mouse anti-β-Gal    | DSHB | 40-1a; RRID:AB_528100 |
| rabbit anti-Mef2    | Lilly et al., 1995 | N/A |
| **Critical Commercial Assays** | | |
| SensiFast SYBR No-ROX Mix | Bioline | BIO-98020 |
| Sensifast cDNA Synthesis Kit | Bioline | BIO-65053 |
| **Deposited Data**  |        |            |
| Rbf and Dp ChIP-seq in muscles | This paper | GEO: GSE102043 |
| RNA-seq of muscles and notum in wild type and Me2 > Rbf-RNAi | This paper | GEO: GSE102104 |
| Drosophila Ensembl Biomart version 77, BDGP5.77 | Ensembl (Hubbard et al., 2007) | N/A |
| Database of known motifs Eurakyote DNA (Vertebrates in vivo and in silico) and FLY DNA | Tomtom | http://meme-suite.org/tools/tomtom |
| Transfac database, insect metrix (professional version release 2009.4) | Transfac | N/A |
| **Experimental Models: Cell Lines** | | |
| Drosophila melanogaster cell line: ML-Dmd8 | DGRC | Flybase: FBtc0000092; RRID:CVCL_Z754 |
| **Experimental Models: Organisms/Strains** | | |
| Act88F-GAL4 | Richard. M. Cripps | Bryantsev et al., 2012 |
| P[twi:: bgal] | Richard. M. Cripps | N/A |
| vgAMe-lacZ | Alexis Lalouette | N/A |
| P[GAL4-Mef2.R]3 | Bloomington Drosophila Stock Center | Flybase: FBst0027390 |
| UAS-Rbf-RNAi | Vienna Drosophila Resource Center | Flybase: FBst0450139 |
| UAS-Dp-RNAi | Vienna Drosophila Resource Center | Flybase: FBst0450633 |
| P[VALIUM20-mCherry]attP2 | Bloomington Drosophila Stock Center | Flybase: FBti0143385 |
| P[tubP-GAL80][ts]20 | Bloomington Drosophila Stock Center | Flybase: FBti0027796 |
| UAS-usp-RNAi<sup>GD1554</sup> | Vienna Drosophila Resource Center | Flybase: FBst0452461 |
| UAS-usp-RNAi<sup>HP00546</sup> | Bloomington Drosophila Stock Center | Flybase: FBst0027258 |
| UAS-ctl-RNAi<sup>RD1237</sup> | Vienna Drosophila Resource Center | Flybase: FBst0470085 |
| UAS-Stat92E-RNAi<sup>HM600095</sup> | Bloomington Drosophila Stock Center | Flybase: FBst0033637 |
| UAS-Stat92E-RNAi<sup>HP01265</sup> | Bloomington Drosophila Stock Center | Flybase: FBst0031317 |
| P[UAS-Dcr-2.D]1, w1118; P[GAL4-Mef2.R]R1 | Bloomington Drosophila Stock Center | Flybase: FBst0025756 |
| **Oligonucleotides** | | |
| Primer sequence | This paper | Table S7 |

(Continued on next page)
CONTACT FOR REAGENT AND RESOURCE SHARING

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Maxim V. Frolov (mfrolov@uic.edu).

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Fly stocks

The line Act88F-GAL4 (Bryantsev et al., 2012) and P{twi- bgal} were provided by Richard. M. Cripps. The stock vgAME-lacZ was used to mark the developing indirect flight muscles. The GAL4 drivers P{GawB}how24B and P{GAL4-Mef2.R}3, and the GFP reporter for Mhc expression P{Mhc-tauGFP}2 were obtained from Bloomington Drosophila Stock Center (Bloomington, IN, USA). The lines UAS-Rbf-RNAi and UAS-Dp-RNAi were obtained from the library RNAi-GD (ID 10696 and 2722, respectively) at the Vienna Drosophila Resource Center (Vienna, Austria). The stocks P{VALIUM20-mCherry}attP2, and P{VALIUM20-EGFP.shRNA.3}attP40 from TRiP collection was used as control, and P{UAS-Dcr-2.D} was used to help processing long, partially double-stranded endogenous transcripts (hairpin RNAs) into endo-siRNAs. In all experiments the transgene UAS-Dicer-2 was added to enhance the Rbf-dependent phenotype, except Figure 7F. These stocks were obtained from Bloomington Drosophila Stock Center. All fly crosses were made at 25°C in vials containing standard cornmeal-agar medium.

Rbf was depleted in muscles using Mef2-GAL4 and regulated over time with the TARGET system (McGuire et al., 2004). P{tubP-GAL80[ts]} was inactivated in water bath at 31°C whereas it was fully active in incubator set at 18°C. Females laid eggs for a period of 24 h at 18°C. Animals were kept at 18°C during embryogenesis. Then, vials were placed at 31°C during larval development to switch ON the TARGET system to inactivate GAL80[ts] and release the inhibition on GAL4 activity. The system was switch OFF (back at 18°C) at mid pupa development (approx. 36 h APF), early pupa development (around 0 h APF), and late third instar larva. Additionally, the system was kept either ON (31°C) or OFF (18°C) during development to control GAL80 activity.

Two RNAi lines were used for usp (1: UAS-usp-RNAiGD1554 and 2: UAS-usp-RNAiJF02546), one for UAS-ct-RNAiGD1237, and two for Stat92E (1: UAS-Stat92E-RNAiHM20035 and 2: UAS-Stat92E-RNAiJF01265), three for ERR (1: UAS-ERR-RNAiHM23520, 2: UAS-ERR-RNAiHF02431, and 3: UAS-ERR-RNAiHM20035), and two for Di (1: UAS-DI-RNAiGD1238 and 2: UAS-DI-RNAiJF02825). All these RNAi lines were obtained from Bloomington Drosophila Stock Center except UAS-usp-RNAiGD1554, UAS-ct-RNAiGD1237 and UAS-DI-RNAiGD1238, obtained at the Vienna Drosophila Resource Center (Vienna, Austria).

Cell culture, dsRNA and myotube differentiation

Drosophila ML-Dmd8 cells (RRID:CVCL_Z754) were maintained at 25°C in Schneider’s Drosophila Medium (GIBCO, 2018-03) supplemented with 10% v/v fetal bovine serum (FBS, HyClone) and 10 µg/ml insulin (Sigma, I0516).

dsRNA knockdown was done by soaking cells with 50 µg of dsRNA targeting GFP and Rbf in serum-free media for 4 h in six-well plates, after which complete media was added. Two days after dsRNA treatment, cells were re-plated into 24-well plates and re-soaked with dsRNA. Then growth medium was switched to differentiation medium to induce differentiation. Sequences of primers used to synthesize dsRNA are listed in Table S7.

Differentiation into myotubes was induced by seeding cells at 5,000 cells/mm² with the growth medium, which contains 10% FBS supplemented with 10 µg/ml insulin in Schneider’s. Media was changed 2 hours after seeding when cells reached 100% confluence. Differentiation medium composition was Schneider medium with 2% v/v horse serum (GIBCO, 16050-122), 10 µg/ml insulin, and...
et al., 1996), rabbit polyclonal anti-Dp (#212, 1:300)(Dimova et al., 2003), rabbit anti-E2f2 antibodies (#79), rabbit anti-Cleaved e3, and 0.1% Triton X-100 in PBS. After washing three times with PBS, coverslips were mounted with FluorSave Reagent (Calbiochem, 345789) on glass slides.

Tissues were dissected and immediately fixed in 4% formaldehyde in PBS for 30 min, permeabilized in 0.3% Triton X-100 in PBS twice for 10 min each and blocked in 2% bovine serum albumin (BSA) with 0.3% Triton X-100 in PBS for 60 min. The DLMs of pupa staged at 15, 20, 21, 24, 29 and 33 h APF were dissected as in (Weitkunat and Schnorrer, 2014), fixed for 20 min, and washed in PBS with 0.1% Triton X-100. Thoraces of 2- to 5-day-old adult or pharate pupa staged at 43, 52, 72, 80 and 96 h APF were fixed for 1 h in relaxing buffer. Thoraces were bisected in the sagittal plane, and were then fixed for an additional 15 min as described in (Schnorrer et al., 2010). In the case of transverse plane sectioning, flies were snap-frozen in liquid nitrogen and cut twice with a razor. The transverse sections were fixed for 1 h. A minimum of five to eight animals per genotype was dissected per experiment, except for the experiment that required quantitative analysis, in which case the staining was carried out two or three times.

Samples were incubated with antibodies overnight at 4°C in 2% BSA and 0.1% Triton X-100 in PBS. After washing four times for 10 min each in 0.1% Triton X-100 (in PBS), samples were incubated with appropriate Cy3- or Cy5-conjugated anti-rabbit secondary antibodies (Jackson Immunoresearch Laboratories, used at 1:500) for 30 min in 1% BSA. After washing three times with PBS, coverslips were mounted with FluorSave Reagent (Calbiochem, 345789) on glass slides.

Immunofluorescence and confocal microscopy
ML-Dmd8 cells seeded in 12mm poly-L-lysine coated round coverslip (Fisher Scientific, 08-774-384) in a 24-well plate were harvested at day 0, 3 and 6 post-differentiation. Cells were washed with PBS 1×, fixed in 4% formaldehyde for 15 min, washed with PBS, and blocked in 3% bovine serum albumin (BSA) with 0.5% Triton X-100 for 30 min. Cells were incubated with antibodies for 2 h in a humid chamber at 4°C. After washing three times for 5 min each in PBS, cells were incubated with appropriate Cy3- or Cy5-conjugated anti-rabbit secondary antibodies (Jackson Immunoresearch Laboratories, used at 1:500) for 30 min in 1% BSA. After washing three times with PBS, coverslips were mounted with FluorSave Reagent (Calbiochem, 345789) on glass slides.

Tissues were dissected and immediately fixed in 4% formaldehyde in PBS for 30 min, permeabilized in 0.3% Triton X-100 in PBS twice for 10 min each and blocked in 2% bovine serum albumin (BSA) with 0.3% Triton X-100 in PBS for 60 min. The DLMs of pupa staged at 15, 20, 21, 24, 29 and 33 h APF were dissected as in (Weitkunat and Schnorrer, 2014), fixed for 20 min, and washed in PBS with 0.1% Triton X-100. Thoraces of 2- to 5-day-old adult or pharate pupa staged at 43, 52, 72, 80 and 96 h APF were fixed for 1 h in relaxing buffer. Thoraces were bisected in the sagittal plane, and were then fixed for an additional 15 min as described in (Schnorrer et al., 2010). In the case of transverse plane sectioning, flies were snap-frozen in liquid nitrogen and cut twice with a razor. The transverse sections were fixed for 1 h. A minimum of five to eight animals per genotype was dissected per experiment, except for the experiment that required quantitative analysis, in which case the staining was carried out two or three times.

Samples were incubated with antibodies overnight at 4°C in 2% BSA and 0.1% Triton X-100 in PBS. After washing four times for 10 min each in 0.1% Triton X-100 (in PBS), samples were incubated with appropriate Cy3- or Cy5-conjugated anti-mouse and anti-rabbit secondary antibodies (Jackson Immunoresearch Laboratories, used at 1:300) for 90 min in 10% normal goat serum and 0.1% Triton X-100 in PBS. After washing with 0.1% Triton X-100, tissues were stored in glycerol with antifade, and mounted on glass slides. All steps were performed at room temperature, unless otherwise stated.

The primary antibodies were mouse anti-β-Gal (40-1a, 1:200, Developmental Studies Hybridoma Bank (DSHB)), mouse anti-β-PS Integrin (CF.6G11, 1:50, DSHB), mouse 22c10 (1:30, DSHB), rabbit anti-Mef2 (1:1,000) (Lilly et al., 1995), mouse RBF (DX2, 1:20, (Du et al., 1996), rabbit polyclonal anti-Dp (#212, 1:300)(Dimova et al., 2003), rabbit anti-E2f2 antibodies (#79), rabbit anti-Cleaved Drosophila Dcp-1 (A6216, 1:500, Cell Signaling), rat anti-Kettin (MAC155, 1:1000, Babraham Institute) and rabbit anti-pH3 (Millipore, 1:200), Rhodamine-Phalloidin or fluorescein isothiocyanate–Phalloidin were used to counterstain and stain thin filaments, and 4,6-diamidino-2-phenylindole (DAPI) for nucleus staining. Mef2>2p-RNAi was used as negative control because of the non-specific background signal detected in the cytoplasm of myotubes.

Fluorescent images were acquired with the laser-scanning confocal microscope (Zeiss LSM Observer.Z1) using x20/0.8, x40/1.2 and x100/1.45 objectives. Images were processed using Photoshop (Adobe Systems). All images are confocal single-plane images, unless otherwise stated. Only representative image is shown per experiment.

Western blot
Six or eight pharate pupae (96 h APF) were collected per sample by removing the pupa case. Flight muscles were dissected from 15 - 20 thoraces of pharate pupa (96 h APF) in relaxing buffer, snap-frozen in dry ice, and homogenized in 40 μL or 100 μL of lysis buffer (50mM HEPES[pH8.0], 100mMKCl, 2mMEDTA, 10%glycerol, 10 mM NaF, 1 mM diithiothreitol [DTT], 1 mM phenylmethylsulfonfyl fluoride [PMSF], 0.1% NP-40, protease inhibitor cocktail (Sigma)).
ML-Dmd8 cells were plated in a 12-well plate and soaked with 20 μg dsRNA as previously described here. Cells were harvested 4 days post-treatment, span down for 2 min at 4000 rpm in a refrigerated centrifuge. Samples were homogenized in 80 μL of lysis buffer. Bradford method was used to estimate protein content in each sample. Between 15 and 20 μg of sample was loaded by well using laemmli loading buffer. Proteins were resolved by electrophoresis on a sodium dodecyl sulfate 8% and 10% polyacrylamide gel electrophoresis gel. Membranes were probed with mouse anti-E7 (β-tubulin, 1:7000, DSHB), or mouse anti-RBF (DX3, 1:5000). Goat anti-mouse secondary antibody conjugated with horseradish peroxidase was used (1:5000). The blots were developed with Pierce ECL kit (Thermofisher).

**RNA-seq and RT–qPCR**

**Sample preparation**

Wandering third instar larvae were harvested, wing discs were dissected, and notum was cut and collected. Flight muscles were dissected from female pharate animals at 96 h APF. At least three to five pharates, and between 11 and 15 third instar larvae were dissected per sample. ML-Dmd8 cells were seeded in a 24-well plate, and harvested after 0, 3, 6 and 9 days of differentiation using TRIZol™ Reagent (Invitrogen). Two or three independent biological samples were collected for each genotype and developmental stage, or treatment condition. Total RNA was isolated with TRIZol.

**RNA-seq libraries and sequencing**

Samples were submitted to the Genomics Facility at the University of Chicago, Chicago, IL, for DNase treatment, generation of total RNA libraries (Ribo-Zero depletion) using Illumina TruSEQ Stranded kits, and sequencing using Illumina HiSEQ4000. Illumina bcl2fastq2-v2.17.1.14 software was used for basecalling.

**Read Quality Assessment and Filtering**

Basic assessment of Illumina output reads (FastQ) quality including GC bias were checked by FastQC program (https://www.bioinformatics.babraham.ac.uk/projects/fastqc/). Poor quality reads were eliminated before mapping based on default quality flag by Illumina pipeline in FastQ file.

**Mapping of reads**

Mapping of reads was done with TopHat (tophat version 2.1.1 with Bowtie version 2.3.2) (Trapnell et al., 2009). Identified 50bp short reads were uniquely aligned allowing at best two mismatches to the reference genome from Ensembl database (BDGP5.77) (Hubbard et al., 2007). Sequence matched exactly more than one place with equally quality were discarded to avoid bias. The reads that were not mapped to the genome were utilized to map against the transcriptome (junctions mapping). EnsEMBL version 77 (Hubbard et al., 2007) gene model was used for this process.

**Read count, normalization and differential expression analysis**

After mapping, we used SubRead package featureCount (version 2.21) (Liao et al., 2013) to calculate absolute read abundance (read count, rc) for each transcript/gene associated to EnsEMBL (v77) genes of D. melanogaster. For differential expression (DE) analysis by popular bioconductor package DESeq2 (version 1.14.1) with R version 3.3.2 (2016-10-31) (Anders and Huber, 2010) that uses a model based on the negative binomial distribution. To avoid false positive, we considered only those transcripts where at least 10 reads are annotated in at least one of the samples used in this study. Heatmap were generated using Genesis platform (Sturn et al., 2002). A hierarchical unsupervised clustering was applied using Pearson correlation distance. List of genes and transcripts were compared using Venn diagram from Venny 2.1.

**RT–qPCR**

Reverse transcription to measure standard mRNAs was performed using the SensiFAST cDNA Synthesis Kit (Bioline) according to the manufacturer’s specifications. qPCR was performed with the SensiFast SYBR No-ROX Mix (Bioline) on a LightCycler 480 (Roche). The reference genes RpL32, RpL30 were validated as stable control genes. Normalization was calculated using the geometric mean of these reference genes. Primer sequences are listed in Table S7. Each sample was measured twice. In all experiments normalized data is relative to control (Day 0, GFP–dsRNA, Mef2>mCherry–RNAi).

**ChIP-seq**

**Sample preparation**

Flight muscle tissues were dissected out of at least 35 pupae staged at pharate (96 h APF) and homogenized. Control genotype was used Mef2>mCherry–RNAi. Chromatin was extracted using homogenizer with 60 mM KCl, 15 mM NaCl, 4 mM MgCl2, 15 mM HEPES (pH 7.6), 0.5% Triton X-100, 0.5 mM dithiothreitol (DTT), 10 mM sodium butyrate and protease inhibitor cocktail (Complete, Roche). Samples were crosslinked for 15 min at room temperature in 1.8% formaldehyde, and 225 mM Glycine was added. Then, cells were lysed with 15 mM HEPES at pH 7.6, 140 mM NaCl, 1 mM EDTA, 0.5 mM EGTA, 0.1% sodium deoxycholate, 1% Triton X-100, 0.5 mM DTT, 0.1% SDS, 0.5% lauroylsarcosine and 10 mM sodium butyrate with protease inhibitor cocktail (Complete, Roche), and were sheared using a Branson 450 Sonifier. Chromatin was immunoprecipitated with mouse anti-RBf (DX3/DX5, ratio 1:1) (Du et al., 1996) and anti-DP (#212, (Dimova et al., 2003)). Complexes were pulled down with Protein G Dynabeads (Invitrogen), washed with lysis buffer four times, washed twice with TE (pH 8), eluted and decrosslinked overnight at 65°C; RNA was degraded with RNase A (Sigma) for 1 h at 37°C and protein was digested with proteinase K for 2 h at 50°C. DNA was purified by phenol-chloroform extraction, followed by overnight ethanol precipitation. Two independent biological samples were used.
**ChIP-seq libraries and sequencing**

Samples were submitted to the Genomics Facility at the University of Chicago, Chicago, IL. ChIP-seq libraries were generated using an Illumina ChiP-SEQ kit, and sequenced using Illumina HiSEQ2500 (v4 chemistry). Illumina bcl2fastq-1.8.3 software was used for basecalling.

**ChIP-seq data analysis**

Illumina pipeline analyzed short reads were uniquely aligned allowing at best two mismatches to the reference *Drosophila melanogaster* genome version BDGP 5.77, using the program BOWTIE (version 2.3.2) (Langmead et al., 2009). Peak caller algorithm MACS (version 1.4.2) (Zhang et al., 2008) was used to determine enriched peak region with parameters:--nomodel,--tsize = 50,--shift-size = 160. Shift size was determined using Pyicos (Althammer et al., 2011) strand correlation method.

Enriched peaks were annotated to all EnsEMBL (Hubbard et al., 2007) transcripts (for *Drosophila* EnsEMBL Biomart version 77) using bedTools (Quinlan and Hall, 2010) within a 5Kb window.

Distribution of enriched reads along the genome and transcription start site (TSS) of RefSeq transcripts were determined using CEAS (Shin et al., 2009). Read density graph or average profile of reads around TSS or any particular genomic location is based on “sitepro” script of the CEAS tool.

Rbf transcript targets were defined as direct targets if Rbf occupancy is enriched within −1 Kb to + 1 Kb from TSS (Transcription start site). Dp direct targets were defined the same way.

Tag density heatmaps around the transcript start site (TSS) were prepared using program HOMER (version 4.9) (Heinz et al., 2010) with parameter (-size /C0 5000,5000 -hist 25 -ghist) and image visualize using gitools (Perez-Llamas and Lopez-Bigas, 2011).

**ChIP-qPCR**

Animals staged at pharate (96 h APF) were dissected out of their pupa case and homogenized. Genotype used were *Mef2>*mCherry-RNAi and *Mef2>*Dp-RNAi. Chromatin was extracted and immunoprecipitated as mentioned earlier. In addition to mouse anti-Rbf (DX3/DX5, ratio 1:1) (Du et al., 1996) and anti-DP (#212, (Dimova et al., 2003), both mouse anti-Myc (9e10) and rabbit IgG (Sigma) were used as non-specific antibodies. The input genomic DNA (before precipitation), as well as immunoprecipitated DNA, was quantified using qPCR as described earlier. Primer sequences are in Table S7. A negative sequence site that does not contain any predicted E2F-binding sites was used as control. The protein enrichment was calculated as the percentage of immunoprecipitated DNA relative to input DNA (prior DNA precipitation) for each antibody. Data presented are relative to the negative binding site for each ChIP. Each sample was measured twice.

**Next-generation sequencing data visualization**

Data from ChIP-seq and RNA-seq were integrated and visualized using Integrative Genomics Viewer (IGV_2.3.79, (Robinson et al., 2011)). The reads assembled were uploaded as BedGraph for RNA-seq and as bigWig format for ChIP-seq. DmBDGP5.77 used as genome annotation.

**De novo motif discovery**

To determine an over-represented short sequence motif in enriched peak, we used total 100 nucleotide sequence (from peak summit, 50bp down and 50bp up). We used MEME (Bailey and Elkan, 1994) program for finding enriched motif. For E2F motifs zero or one occurrence per sequence was set and motif length was 6-10 bp (Figure 5E). To discover new motifs we used zero or one occurrence per sequence and motif length 6-10 bp in transcripts repressed by Rbf; any number of occurrences per sequence and motif length 6-10 bp and 6-14 bp for transcripts activated by Rbf (Figure 7D). Tomtom was used to compare the motifs against a database of known motifs from both Eurakyote DNA (Vertebrates in vivo and in silico) and FLY DNA. E values lower than 0.01 were considered as a match.

**STORM**

Possible occurrence of transcription factor motif in 350bp or 550 bp of peak summit centered location (175 bp up and 175 bp down, and 275 bp up and 275 bp down) were predicted with STORM algorithm (Schones et al., 2007) with a p value cutoff determined based on the size of the input sequence as (1/100xsequence-size), and using position frequency matrices (PFM) from Transfac database (insect metrix) (professional version release 2009.4). The percentage of motif matching with each transcription factor was calculated for each set of sequence (activated and repressed by Rbf). The ratio of % transcription factor in activated over repressed was determined.

**Functional Enrichment Analysis**

The list of transcripts repressed and activated by Rbf, as defined in Figure 5, were annotated to genes. Functional annotation is based on Gene Ontology (GO) (Ashburner et al., 2000; http://www.geneontology.org) as extracted from EnsEMBL (Hubbard et al., 2007). Accordingly, all genes are classified into the ontology: genes involved in Biological Process (BP). We have taken only the GO/pathway categories that have at least 10 genes annotated. We used Gitools for enrichment analysis (Perez-Llamas and Lopez-Bigas, 2011) (http://www.gitools.org). Resulting p values were adjusted for multiple testing using the Benjamin and Hochberg’s method of False Discovery Rate (FDR) (Benjamini and Hochberg, 1995). Enrichment map were visualized using Cytoscape v3.3.0 platform (Shannon et al., 2013).
et al., 2003). The parameters were set as $p < 0.005$, FDR $< 0.05$, Similarity threshold between nodes:0.3. Flybase was used for online resources on Database of Drosophila Genes & Genomes.

**QUANTIFICATION AND STATISTICAL ANALYSIS**

Scripts were used in ImageJ for automatic and unbiased quantification, and are available upon request

Graphs were generated with the GraphPad Prism version 7.0 (Graphpad Software). The group means were analyzed for overall statistical significance using two-way analysis of variance and non-parametric analysis (Mann–Whitney).

**Quantification of mitotic myoblasts and total number**
The total number of mitotic myoblasts was determined using phosphorylated histone H3 antibody. The myoblast-positive area and developing DLMs area were marked with twi-lacZ reporter in the wing disc in late third instar larvae (120 h after egg-laying), and in pupa thoraces staged at 15, 20, 24, 29 and 33 h APF. Developing DLMs area was also labeled using 22C10 marker in pupa thoraces staged at 21 h APF, and Phalloidin in 29 and 33 h APF. Images were taken with confocal microscopy in a z stack to cover all nuclei in the tissue, and then were projected into a single plane. The area covered by the adult myoblasts and developing DLMs was quantified using the ImageJ software (1.48v, NIH, USA). The plugin Image-based Tool for Counting Nuclei was used to determine the number of pH3-positive nuclei in the myoblast-positive area and developing DLM. To count the total number of myoblasts in the wing discs a mask was created with twi-lacZ marker around the area of the adult muscle precursors to limit DAPI count to myoblasts. Minimum of 5 animals per genotype and developmental stage were scored. Two independent experiments, only one shown, except for Rbf-depleted DLMs staged at 29 and 33 h APF. More details regarding the statistical analysis and sample size for each experiment can be found in legend of Figure 2.

**Quantification of number of nuclei in muscles**
The number of nuclei, the average size of nuclei and the cross-section area of DLM in transverse sections of pharate was quantified using the Analyze particle function of ImageJ Student’s. Briefly, the full area of DLM number 4 was selected as the region of interest using β-PS integrin. then, an automatic threshold was applied to DAPI channel. Watershed segmentation was then performed to separate overlapping spots, and the total number of particles was counted along with the average size of particles and the area selected. Nine animals per genotype were scored. More details regarding the statistical analysis and sample size for each experiment can be found in legend of Figure 3.

**Quantification of fiber length**
ImageJ was also used to quantify muscle fiber length of DLMs in pupa thoraces staged at 29 h APF. Minimum of 5 animals per condition were scored. Two independent experiments were done, only one representative experiment shown. More details regarding the statistical analysis and sample size for each experiment can be found in legend of Figure 3.

**Quantification of myofibril morphology**
Images of the DLM sagittal section were blindly scored as either frayed, trapezoid or wild-type regarding myofibril morphology as in (Schnorrer et al., 2010). At least 6 hemithoraces were scored per genotype/gender/experiment. Two independent experiments were conducted. More details regarding the statistical analysis and sample size for each experiment can be found in legend of Figure 7.

**Quantification of Dmd8 cells**
The number of Mef2-positive nuclei and pH3-positive nuclei in differentiating ML-Dmd8 cells were counted using the Analyze particle function of ImageJ Student’s. Briefly, automatic thresholds were applied to red channels (either Mef2 or pH3), and the green channel (Phalloidin) was inverted and subtracted from the red channel to select only Mef2 and pH3 spots within cells stained with Phalloidin. Watershed segmentation was then performed to separate overlapping spots, and total number of particles was counted. Same procedure was applied to blue channel (DAPI) to count total number of nuclei to normalize data. Total number of nuclei per myotube, defined as the fusion index to characterize the efficacy of differentiation, was also quantified using ImageJ. A mask was created using Phalloidin around each myotube to limit particle count to each myotube. At least ten images per treatment/time point/experiment were processed. Three independent experiments were done. More details regarding the statistical analysis and sample size for each experiment can be found in legend of Figure 1.

**DATA AND SOFTWARE AVAILABILITY**

Both ChIP-seq and RNA-seq data from this publication have been deposited to the GEO database (https://www.ncbi.nlm.nih.gov/geo/) and assigned the identifier Series GSE102105 (https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE102105), which is composed of GSE102043 and GSE102104.
Supplemental Information

Rbf Activates the Myogenic Transcriptional Program
to Promote Skeletal Muscle Differentiation

Maria Paula Zappia, Alice Rogers, Abul B.M.M.K. Islam, and Maxim V. Frolov
Figure S1: The loss of Rbf does not induce apoptosis mediated by depl in developing indirect flight muscles. Related to Figure 2.

Developing DLMs stained over time from 21 to 96 h APF. Apoptotic cells were marked using anti-dep1 antibody (right panel). Overall structure was analyzed using DAPI and Phalloidin or 22c10 to mark developing IFM or Kettin to stain sarcomere depending on the developmental stages (left panel). Single confocal sections are shown. Scale 40 µm. Full genotypes are w-, UAS-Dicer2; +; Mef2-GAL4/UAS-mCherry-RNAi (left panel) and w-, UAS-Dicer2; +; Mef2-GAL4/UAS-Rbf-RNAi (right panel)
Figure S2: Rbf interacts with other transcription factors. Related to Figure 7.

Genetic interactions between Rbf and candidates were analyzed by knocking down the expression of each gene with or without Rbf-RNAi in muscles using Mef2-GAL4 driver. The candidates tested are estrogen-related receptor (ERR) and Delta (Dl). Lethality during pupa development was analyzed as % animals scored at each developmental stage. Adults: light grey, Pharate: grey, Early pupa: dark grey. N= 94 - 319 flies per genotype. Two independent experiments. Genotypes are Mef2-GAL4/UAS-mCherry-RNAi, UAS-Rbf-RNAi/Mef2-GAL4, three RNAi lines were used for ERR (1: UAS-ERR-RNAiHMJ23520, 2: UAS-ERR-RNAiJF02441, and 3: UAS-ERR-RNAiHMC01087), and two for Dl (1: UAS-Dl-RNAiGD1238 and 2: UAS-Dl-RNAiP02825).
Table S7: Primer sequence. Related to STAR Methods

| Gene       | Forward primer sequence | Reverse primer sequence | Application       |
|------------|-------------------------|-------------------------|-------------------|
| *twi*      | AGACTCAGCACAGATCCCAGAC | GAGCTGGCCGATCCTACACG    | RT-qPCR           |
| *Mef2*     | CCGATATCATGAGCCCTACAC  | CTGTTACCACCTGATCCTAC   | RT-qPCR           |
| *htl*      | AAGCGTCCGAAACCTCCAC    | TGGTCGCTTCTGTTGACCT    | RT-qPCR           |
| *stumps*   | AACAGGTCGACTGCTCACC    | ACTGTCGACTGCTGATC      | RT-qPCR           |
| *Act88F*   | ATGGTGGTTGATCGTGAGCG   | CTTCTCCATGTCCTCCAGT    | RT-qPCR           |
| *Mhc*      | AGAACGACCTGAGGAGGAC    | TCAGCCTCTGCTGCTGCT     | RT-qPCR           |
| *fln*      | GGCAAGAGGACGACGATC    | TCTGCTCCAGATCGATC      | RT-qPCR           |
| *Rpl32*    | TACAGGCGCAATGCTGAGC   | GACGCCTCTGCTGATC       | RT-qPCR           |
| *Rpl30*    | GCAAAATCTGCTGCTGC    | ACTCTGCTGCTGCTGCT      | RT-qPCR           |
| *Lmpt-transcript K* | AGTGGCTGCCCTAAACAA    | ATGGTCCCTCTGCTGATC     | RT-qPCR           |
| *sals*     | GCAAGGCATGAGAAGAAAC   | TCAGCTCTAGCTGACTG      | RT-qPCR           |
| *Tm1-transcript H* | ATAAAGCAGCCACGCAA    | TTCTGCTGCTCGCTCTG      | RT-qPCR           |
| *Tm2*      | TCTCCAGGCGCTTCTGAGC   | TTCTCAGGCGCTTCTGAGC    | RT-qPCR           |
| *Mlc2*     | TTCTCTGCTTCTCCAGAGC   | TTGTCGAGCTCCATGCAGG    | RT-qPCR           |
| *how*      | ATCTGTCGATGACCTGCATG  | TTAGCTCACTTCTGCTGCG    | RT-qPCR           |
| *saln*     | ACTACAGGACCGACCAACAG  | ATGGTCTGCTGCTGCTCG     | RT-qPCR           |
| *Rbf*      | AACAGGACAGGACGAGCTG   | ACTCTGCTGCTGCTGCTG     | RT-qPCR           |
| *P5CDh1 157-40* | TTTCTGCTCCAGCTTCTG    | TTAGCTCTGCTGCTGCTG     | RT-qPCR           |
| *Cyt-c-p -496/-372* | AAAAGATGTCTTGCTGCAGT  | TTAGCTCTGCTGCTGCTG     | RT-qPCR           |
| *Zasp66 -847/-712* | TGGGACAGAAGTTTATTGACC | TTAGCTCTGCTGCTGCTG     | RT-qPCR           |
| *Negative site* | TTAGCTCTGCTGCTGCTG    | TTAGCTCTGCTGCTGCTG     | RT-qPCR           |
| *Rbf*      | CGAACAGGACGAGCTG      | ACTCTGCTGCTGCTGCTG     | RT-qPCR           |
| *GFP*      | CTAATACGACTACTAAGGAGG | CTAATACGACTACTAAGGAGG  | dsRNA             |
|           | AGCACGGAGACGCTTCC    | GCAGGACAGCAGATGATCCTTG | dsRNA             |
|           | TAAACGGCCGCAAGGAT    | GCAGGACAGCAGATGATCCTTG | dsRNA             |

* T7 sequence is underlined

**Table S7: Primer sequences.** Sequence of primers used to measure gene expression (RT-qPCR), genomic occupancy (ChIP-qPCR) and primers used to make dsRNA.