The glucose-sensing transcription factor MLX promotes myogenesis via myokine signaling

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Metabolic stress and changes in nutrient levels modulate many aspects of skeletal muscle function during aging and disease. Growth factors and cytokines secreted by skeletal muscle, known as myokines, are important signaling factors, but it is largely unknown whether they modulate muscle growth and differentiation in response to nutrients. Here, we found that changes in glucose levels increase the activity of the glucose-responsive transcription factor MLX (Max-like protein X), which promotes and is necessary for myoblast fusion. MLX promotes myogenesis not via an adjustment of glucose metabolism but rather by inducing the expression of several myokines, including insulin-like growth factor 2 (IGF2), whereas RNAi and dominant-negative MLX reduce IGF2 expression and block myogenesis. This phenotype is rescued by conditioned medium from control muscle cells and by recombinant IGF2, which activates the myogenic kinase Akt. Importantly, MLX-null mice display decreased IGF2 induction and diminished muscle regeneration in response to injury, indicating that the myogenic function of MLX is manifested in vivo. Thus, glucose is a signaling molecule that regulates myogenesis and muscle regeneration via MLX/IGF2/Akt signaling.

Keywords: MLX; IGF2; myokine; glucose; myogenesis; regeneration; skeletal muscle

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Unicellular and multicellular organisms have evolved several homeostatic adaptive strategies to regulate cell and organismal behavior in the face of fluctuating nutrient levels. Skeletal muscle is one of the tissues that are most profoundly remodeled in response to changes in the levels of glucose and other nutrients. Many aspects of skeletal muscle homeostasis are modulated by glucose, including myogenesis, the process by which mononucleated progenitors (myoblasts) fuse into syncytial muscle cells during development and regeneration. Several lines of evidence indeed point to glucose as an important mediator of myogenesis. First, high glucose levels promote myogenesis [Nedachi et al. 2008], and the expression of genes involved in glucose metabolism is up-regulated during myogenesis in vitro [Webster et al. 1990]. Second, glucose metabolism via glycolysis is necessary for myogenesis in the Drosophila embryo [Tixier et al. 2013]. Last, glucose restriction inhibits myogenesis by activating AMPK kinase and the sirtuin SIRT1, and partial loss of SIRT1 allows myogenesis to occur even at low glucose concentrations that are normally not permissive [Fulco et al. 2008]. Despite these studies, the mechanisms and signaling pathways sensing glucose and responsible for glucose-induced adaptive myogenesis are largely unknown.

A class of proteins that have recently gained attention for their many signaling roles is myokines; i.e., growth factors and cytokines secreted by skeletal muscle in response to a range of different stimuli [Pedersen and Febbraio 2012; Demontis et al. 2013a]. Myokines can act locally on the muscle itself as well as systemically on other tissues to regulate many physiological functions ranging from metabolic homeostasis to aging and disease-related processes [Boström et al. 2012; Demontis et al. 2014]. In addition to endocrine functions, there are many examples of autocrine and paracrine signaling of myokines on the muscle, including the modulation of myogenesis [Rai and Demontis 2015]. For example, leukemia inhibitory factor [LIF] is a

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myokine that regulates myoblast proliferation and differentiation and is produced in response to exercise (Hunt et al. 2010, 2011; Broholm et al. 2011). However, despite the important signaling roles of myokines, little is known about the transcription factors that modulate their expression in response to glucose and other nutrients.

Max-like protein X (MLX) is a member of the Myc-Max family of basic helix–loop–helix (bHLH) transcription factors [Billin et al. 1999] that, in response to high glucose levels, translocates from the cytoplasm to the nucleus (Peterson et al. 2010), where it regulates the expression of target genes with carbohydrate response element (CHORE) motifs in their promoter (Minn et al. 2005). MLX is ubiquitously expressed across tissues and regulates transcription by forming heterodimers with either MLX-interacting protein (MLXIP/MondoA) or MLXIP-like (MLXIP/CHREBP) that are primarily expressed in the muscle [Billin et al. 2000] and liver [Yamashita et al. 2001], respectively. Increased intracellular glucose leads to nuclear accumulation of MLX [along with MLXIP and MLXIPL] and increased DNA binding. Several lines of evidence support the hypothesis that glucose-6-phosphate is the primary metabolite responsible for MLX activation. First, MLX translocation to the nucleus and transcriptional activity can be stimulated by 2-deoxyglucose, an analog of glucose that accumulates in cells following phosphorylation but cannot be further metabolized [Stoltzman et al. 2008]. Second, phosphorylation of glucose by hexokinase is required for MLX activity [Stoltzman et al. 2008]. Last, phosphoglucomutase deficiency leads to glucose-6-phosphate buildup and enhances MLX activation [Stoltzman et al. 2011]. The precise mechanisms by which glucose-6-phosphate activates MLX are not completely understood but perhaps entail direct binding to MLXIP/MLXIPL [McFerrin and Atchley 2012]. In addition, glucose-dependent post-translational modifications can regulate MLXIP [Guinez et al. 2011], which can also be phosphorylated and inactivated by the energy-sensing kinase AMPK [Kawaguchi et al. 2002]. Thus, MLX and its coactivators can sense glucose via multiple mechanisms. Furthermore, because MLX is part of a large interactive network of transcriptions factors, including Myc, Max, Mxd, and MLXIP/MLXIPL, its transcriptional output may be determined by the context-specific organization of this network [Diotali et al. 2015].

In agreement with a fundamental role for MLX in glucose sensing, microarray studies indicate that ~75% of all transcriptional changes induced by glucose in hepatocytes are mediated by MLX [Ma et al. 2006]. For example, MLX induces the expression of enzymes involved in carbon metabolism [Ma et al. 2005, 2006; Iizuka et al. 2009] and proteins that regulate glucose uptake, including thioredoxin-interacting protein (TXNIP) [Parikh et al. 2007; Wu et al. 2013]. However, although MLX is important for glucose sensing and glucose metabolism in the liver [Postic et al. 2007], it is unknown whether it has similar roles in other tissues.

In this study, we examined the transcriptional and metabolic roles of MLX in skeletal muscle cells. We found that MLX transcriptional activity increases in response to glucose and that MLX is necessary for and promotes myogenic differentiation. However, rather than promoting glucose metabolism, MLX induces the expression of many myokines in response to glucose, including insulin-like growth factor 2 (IGF2), which recapitulates the effects of MLX on myogenic differentiation. These findings indicate that MLX modulates glucose-induced myoblast fusion and differentiation via myokine signaling.

Results

The transcription factor MLX induces target gene expression in muscle cells via glucose-induced DNA binding and histone H4 acetylation

To experimentally modulate MLX transcriptional activity in muscle cells, wild-type MLX (MLX-WT), dominant-negative MLX (MLX-DN), MLX RNAi (MLX shRNA), and N-terminally truncated MLX (MLX-ΔN121) constructs were generated and stably expressed in C2C12 mouse myoblasts via lentiviral-mediated transduction. Overexpression of MLX variants led to an ~60-fold increase in MLX mRNA levels compared with empty vector control, whereas MLX shRNA led to an 80% decrease in endogenous MLX mRNA levels compared with a nontargeting control shRNA (Fig. 1A; Supplemental Fig. S1A). MLX-WT, MLX-DN, and MLX-ΔN121 protein variants were detected by Western blot at the expected sizes (~20 kDa for MLX-ΔN121 and ~30 kDa for MLX-WT and MLX-DN) [Supplemental Fig. S1B] and by immunostaining of lentiviral-transduced cells that coexpress GFP [Supplemental Fig. S1C]. Both MLX-WT and MLX-DN were ubiquitously detected in the cells, whereas MLX-ΔN121 localized to nuclei [Supplemental Fig. S1C], consistent with previous evidence that the N-terminal domain of MLX harbors cytoplasmic retention sequences [Billin et al. 2000].

To assess the transcriptional activity of transgenic MLX variants, we examined changes in the expression of Txnip and Arrdc4, two genes known to be induced by glucose and MLX [Stoltzman et al. 2008; Yu et al. 2009]. Both MLX-DN and MLX shRNA prevented glucose-induced expression of Txnip and Arrdc4, whereas MLX-WT significantly enhanced their expression [Fig. 1B]. Moreover MLX-ΔN121 blunted Txnip and Arrdc4 induction by glucose, indicating that N-terminal truncation impairs MLX transcriptional activity, although it leads to constitutively nuclear localization [Supplemental Fig. S1A–C].

To further corroborate these findings, we assessed the DNA-binding capacities of MLX variants to the CHORE-containing Txnip promoter by chromatin immunoprecipitation (ChIP) combined with quantitative PCR (qPCR). Expression of MLX-WT allowed for glucose-dependent recruitment of MLX and MLXIP [MondoA] to the Txnip promoter, whereas this was blocked by MLX-DN but retained with MLX-ΔN121 [Supplemental Fig. S1D]. Furthermore, histone H4 acetylation, an epigenetic mark induced by glucose and indicative of open chromatin and active transcription, was increased at the Txnip promoter in response to glucose and MLX-WT but inhibited by
Therefore, nuclear localization of MLX is insufficient for transcriptional activity and requires recruitment of additional transcriptional coactivators, such as histone acetyltransferases, as previously described (Davies et al. 2008; Peterson et al. 2010). Taken together, these findings indicate that the transcription factor MLX modulates glucose-induced transcriptional responses in muscle cells and that both its N-terminal domain and DNA binding are required for histone H4 acetylation to activate target gene expression in muscle cells.

Regulation of MLX target genes in muscle cells may also stem from the interaction of MLX with other bHLH transcription factors at target gene promoters. In particular, MLX has recently been shown to coordinately regulate a subset of target genes with c-Myc (Shen et al. 2015) and to be necessary for survival of Myc-driven tumors (Carroll et al. 2015). We found that c-Myc occupancy at the Txnip promoter is reciprocal to MLX and likely mediated by E-boxes contained within the CHORE motifs (Supplemental Fig. S1D). Therefore, antagonistic promoter occupancy by MLX and c-Myc may also contribute to determining the transcriptional output of MLX target genes in muscle cells.

**MLX regulates myogenesis in response to glucose**

C2C12 myoblasts form syncytial muscle cells (myotubes) when grown to confluence in cell culture medium containing 10% fetal bovine serum (FBS) and subsequently cultured for 8 d in differentiation-inducing medium containing 2% horse serum. As measured by immunostaining (Fig. 1C), control myoblasts differentiated efficiently and reached maximal fusion by 4 d of differentiation due to MLX-WT overexpression and impairment of myogenesis due to loss of MLX activity (MLX-DN overexpression and MLX shRNA) compared with controls (empty vector [EV] and a nontargeting [NT] shRNA). Conversely, MLX-DN and shRNA decrease Txnip expression and prevent their glucose-induced up-regulation. The histograms show the mean of three biological replicates ±SEM. ∗ P < 0.05. See also Supplemental Figures S1. C) Phase contrast and α-actin immunostaining demonstrate increased myotube formation at day 4 of differentiation due to MLX-WT overexpression and impairment of myogenesis due to loss of MLX activity (MLX-DN overexpression and MLX shRNA) compared with controls (empty vector [EV] and a nontargeting [NT] shRNA). See also Supplemental Figures S2 and S3. (D) The fusion index (i.e., the number of nuclei within syncytial myotubes) indicates that MLX-WT increases myoblast fusion, whereas MLX-DN and shRNA decrease myoblast fusion. Myoblasts stably expressing MLX-WT have increased glucose sensitivity and increased fusion index at lower glucose concentrations that are normally not permissive. Conversely, myoblasts stably expressing MLX-DN and MLX shRNA cannot fuse regardless of glucose concentrations. ∗ P < 0.05, SEM. n = 3.

**Figure 1.** The transcription factor MLX regulates glucose-induced myogenesis. (A) Increased Mlx expression is detected by quantitative PCR (qPCR) in muscle cells stably expressing MLX-WT and MLX-DN compared with control cells expressing an empty vector [EV] control. MLX shRNA decreases Mlx levels compared with a nontargeting (NT) control shRNA. (B) MLX protein variants and MLX shRNA regulate the expression of MLX target genes known to be glucose-responsive. MLX-WT increases the expression of Txnip and Arrdc4, which are also induced by glucose. Conversely, MLX-DN and MLX shRNA decrease Txnip and Arrdc4 expression and prevent their glucose-induced up-regulation. The histograms show the mean of three biological replicates ±SEM. ∗ P < 0.05. See also Supplemental Figure S1.
myotube size, which is defined as the area of the myotubes divided by the number of myotube nuclei (Supplemental Fig. S2A). Interestingly, the proportional myotube size did not significantly change in response to MLX-WT overexpression, indicating that myotube hypertrophy is primarily due to the higher number of myoblasts fusing to form myotubes.

Because MLX transcriptional activity is regulated by glucose, we next probed their interaction by examining the extent of myoblast fusion at day 4 of differentiation in response to different levels of MLX activity and glucose. Myoblast fusion was maximal at 10–25 mM glucose concentrations in control cells (Fig. 1D), whereas myoblasts overexpressing MLX-WT appeared more glucose-sensitive and had maximal fusion at relatively low glucose concentrations (5 mM). Conversely, myoblasts overexpressing MLX-DN or MLX shRNA were glucose-insensitive and unable to fuse even at high glucose concentrations (40 mM). These findings indicate that MLX and glucose are indeed required for myoblast fusion and that MLX activity dictates cellular sensitivity to glucose.

To dissect the mechanisms involved, we tested whether MLX modulates cell cycle exit, myogenic commitment (nuclear detection of myogenin), and expression of genes encoding proteins that mediate myoblast fusion. However, none of these myogenic processes were modulated by MLX (Supplemental Fig. S2A,B). Moreover, myoblasts overexpressing MLX-DN or MLX shRNA were glucose-insensitive and unable to fuse even at high glucose concentrations (40 mM). These findings indicate that MLX and glucose are indeed required for myoblast fusion and that MLX activity dictates cellular sensitivity to glucose.

To identify MLX target genes relevant for myogenesis, we focused on a set of 80 genes that were significantly and concordantly regulated by MLX-WT, MLX-DN, and MLX shRNA (see Materials and Methods for selection criteria). Of these, 62 genes were transcriptionally activated by MLX, whereas 18 genes were repressed. Enriched gene clusters were identified by DAVID (Database for Annotation Visualization and Integrated Discovery) and PIR (Protein Information Resource) annotation keywords and indicated that there was a significant enrichment (31 out of 67 annotated genes) for muscle-secreted proteins (i.e., myokines) (Fig. 3B), indicating an unanticipated role of MLX in regulating their expression.

MLX regulates myokine expression

In order to discover novel genes regulated by MLX that could be responsible for its effects on myogenesis, we performed RNA sequencing (RNA-seq) of cells with MLX gain and loss of function (Supplemental Table S1). RNA-seq analyses found many genes significantly regulated in muscle cells overexpressing MLX-WT, MLX-DN, and MLX shRNA compared with their respective controls (Fig. 3A). To identify MLX target genes relevant for myogenesis, we focused on a set of 80 genes that were significantly and concordantly regulated by MLX-WT, MLX-DN, and MLX shRNA (see Materials and Methods for selection criteria). Of these, 62 genes were transcriptionally activated by MLX, whereas 18 genes were repressed. Enriched gene clusters were identified by DAVID (Database for Annotation Visualization and Integrated Discovery) and PIR (Protein Information Resource) annotation keywords and indicated that there was a significant enrichment (31 out of 67 annotated genes) for muscle-secreted proteins [i.e., myokines] (Fig. 3B), indicating an unanticipated role of MLX in regulating their expression.

Considering that genes encoding muscle-secreted signaling proteins [myokines] constitute a major class of...
MLX-regulated genes, we tested whether conditioned media from differentiating control cells can rescue the fusion defects of myoblasts overexpressing MLX-DN. Interestingly, the addition of this milieu of secreted factors stimulated fusion of MLX-DN myoblasts, resulting in the formation of myotubes after 4 days in culture (Fig. 3C), suggesting that myokines may be required for glucose-induced myogenesis.

From the RNA-seq data of MLX-regulated myokines, we selected the chemokines Ccl2 and Ccl7, the insulin-like growth factor Igf2, the interleukin-1 receptor antagonist Il1rn, the lectin-binding protein Lgals3bp, the matricellular CCN family gene Nov, and the thrombospondin Thbs2 and examined them further because of their potential association with myogenesis (Florini et al. 1991a; Iruela-Arispe et al. 1993; Sakamoto et al. 2002; Yahiaoui et al. 2008).

First, by qPCR, we validated their expression in response to MLX activity and found that MLX-WT increases their mRNA levels, whereas MLX-DN and MLX shRNA decrease their expression relative to controls (Fig. 3D). Second, we tested whether MLX-induced myokines are modulated during differentiation and by changes in glucose levels. The expression of Ccl2, Ccl7, Igf2, Il1rn, and Nov increased in response to higher glucose levels (Fig. 3D), whereas the expression of most myokines did not increase following differentiation [Supplemental Fig. S4A], with the exception of Igf2 expression, which increased [Fig. 3E]. These findings indicate that glucose and MLX regulate the expression of myokines produced by myoblasts.

**MLX promotes myokine gene transcription by increasing histone H4 acetylation**

Glucose influences gene expression by inducing chromatin remodeling via epigenetic modifications such as acetylation of histones H3 and H4 [Lee et al. 2014]. Considering that MLX increases H4 acetylation in proximity to Txnip [Supplemental Fig. S1D; Cha-Molstad et al. 2009], we performed ChiP of acetylated histone H4 (H4ac) from MLX-WT-expressing and MLX-DN-expressing cells followed by sequencing (Fig. 4) to determine whether MLX generally regulates H4 acetylation in proximity to its target genes. Myoblasts with MLX-DN showed overall increased H4ac near all gene promoters (Fig. 4A), which could be explained by increased glucose uptake by these cells [Fig. 2A]. However, MLX-WT myoblasts displayed increased H4ac specifically in the promoter regions of genes up-regulated by MLX-WT (as determined from RNA-seq data) [Fig. 3] and decreased H4ac in the promoter regions of down-regulated genes [Fig. 4A; Supplemental Table S2]. Interestingly, there...
was an overall correlation between the relative levels of H4ac and the presence of predicted MLX-binding sites; i.e., CHORE-like motifs (Fig. 4B; Supplemental Table S3). For example, the glucose- and MLX-regulated gene *Arrdc4* showed a strong H4ac signal in MLX-WT but not in MLX-DN cells surrounding the transcriptional start site (TSS), which also contains a CHORE motif. Similarly, the myokine *Lgals3bp* displayed strong H4ac levels at the +1 nucleosome position in MLX-WT compared with MLX-DN. RNA-seq tracks show transcript levels, and computationally identified MLX-binding CHOREs (denoted with an asterisk) that lie in close proximity to the H4ac peaks are identified below. (C) ChIP-qPCR validation of H4ac enrichment at the promoter regions of MLX target genes in response to overexpression of MLX-WT (red) compared with MLX-DN (green). Arrows in B indicate regions displaying H4ac peaks that have been further validated by ChIP-qPCR. See also Supplemental Figure S5 and Supplemental Tables S2 and S3.

Figure 4. Histone H4 acetylation occurs specifically at MLX target genes and is dependent on MLX DNA binding. (A) ChIP-seq (ChIP combined with deep sequencing) data for H4ac demonstrate overall H4ac enrichment around the TSSs of genes in MLX-WT and MLX-DN myoblasts and the presence of nucleosome-free regions in close proximity to the TSSs. Overall, a global increase in H4ac levels is observed in MLX-DN cells compared with MLX-WT. However, genes specifically up-regulated by MLX-WT (and having decreased expression with MLX-DN and MLX shRNA) show increased H4 acetylation, whereas genes down-regulated by MLX show decreased H4 acetylation. (B) Examples of MLX target genes that show increased H4ac levels in MLX-WT compared with MLX-DN. RNA-seq tracks show transcript levels, and computationally identified MLX-binding CHOREs (denoted with an asterisk) that lie in close proximity to the H4ac peaks are identified below. (C) ChIP-qPCR validation of H4ac enrichment at the promoter regions of MLX target genes in response to overexpression of MLX-WT (red) compared with MLX-DN (green). Arrows in B indicate regions displaying H4ac peaks that have been further validated by ChIP-qPCR. See also Supplemental Figure S5 and Supplemental Tables S2 and S3.
IGF2 is a myokine regulated by MLX and promotes myogenesis

We tested whether exogenous addition of any of these MLX-induced myokines regulates myoblast fusion. Recombinant mouse myokines were added to differentiation media of control and MLX-DN-overexpressing myoblasts at concentrations of 1, 10, and 100 ng/mL, and myotube formation was monitored. No recombinant myokine was able by itself to induce fusion of MLX-DN myoblasts (Supplemental Fig. S4B), except for 10 and 100 ng/mL IGF2 (Fig. 5A). Concentrations of 100 ng/mL IGF2 also enhanced fusion of control myoblasts (Fig. 5A).

Given the capacity of IGF2 to rescue the myoblast fusion defects due to MLX-DN, we tested whether IGF2 transcriptional changes induced by MLX indeed result in corresponding changes in IGF2 protein levels. To this purpose, we measured IGF2 protein levels by ELISA from undifferentiated myoblasts and differentiating cultures at days 2 and 4. Measurement of IGF2 levels in conditioned medium and cell lysates indicated that IGF2 protein levels increase during myogenesis and in response to MLX-WT overexpression, particularly in the conditioned medium but also in the cell lysates. Conversely, IGF2 protein levels were low in myoblasts overexpressing MLX-DN and MLX shRNA (Fig. 5B). This confirms that changes in IGF2 expression lead to corresponding changes in intracellular and, most notably, secreted IGF2 protein levels.

To test whether changes in IGF2 protein levels induce corresponding changes in IGF2 signaling, we monitored by Western blot phospho-Akt [Ser473] levels, which increase in response to exogenous IGF2. As predicted by the progressive increase in IGF2 expression and IGF2 protein levels during myogenesis, phosphorylation of Akt increases during the course of muscle differentiation. Moreover, increased Akt phosphorylation was found in response to MLX-WT overexpression, whereas myoblasts with MLX-DN and MLX shRNA had lower phospho-Akt levels [Fig. 5C], consistent with IGF2 mRNA and protein changes elicited by MLX gain and loss of function. IGF2 has been previously shown to be produced by myoblasts (Florini et al. 1991a; Erbay et al. 2003) and stimulate myogenic differentiation via Akt (Florini et al. 1991b; Yoon and Chen 2008; Markljung et al. 2009). To test whether Akt is indeed a key mediator of myoblast fusion downstream from MLX/IGF2 signaling, we used pharmacological inhibitors of Akt and found that they prevented MLX and IGF2-induced myoblast fusion following induction of differentiation (Supplemental Fig. S6A–C).

We then tested the responsiveness of MLX-WT-overexpressing and MLX-DN-overexpressing cells to IGF2. In control cells, concentrations of IGF2 between 50 and 100 ng/mL increased the fusion index from control levels of 25% up to 40%. When exposed to exogenous IGF2, myoblasts overexpressing MLX-DN gradually increased their fusion index from 0% (with 1 ng/mL IGF2) up to curve of the myoblast fusion index in response to IGF2 levels indicates that while exogenous addition of recombinant mouse IGF2 to culture medium stimulates fusion in control cells, myoblasts overexpressing MLX-WT are insensitive and appear to have already reached a maximum level of myoblast fusion. In addition, IGF2 rescues myoblast fusion defects due to MLX-DN overexpression. Myoblasts overexpressing IGF2 have increased fusion at low levels of glucose that are normally not permissive for fusion, similar to MLX-WT overexpression (shown in D). The mean of three biological replicates ±SEM is shown, with P < 0.05 compared with the respective control at the same concentration of IGF2/glucose.

Figure 5. MLX promotes myogenesis via IGF2 signaling. (A) Of several MLX-induced myokines, only IGF2 induces myogenesis and rescues myoblast fusion defects caused by MLX-DN overexpression. Immunostaining for α-actin (red) and DAPI (4′,6-diamidino-2-phenylindole) (blue) is shown. See also Supplemental Figure S4. (B) ELISA measurement of IGF2 levels in conditioned cell culture medium and cell lysates indicates that IGF2 protein levels increase as differentiation proceeds and in response to MLX activity (MLX-WT), whereas loss of MLX function (MLX-DN and MLX shRNA) decreases IGF2 protein levels. The histograms show the mean of three different experiments ±SEM. [*] P < 0.05. (C) Treatment of serum-deprived muscle cells [–] with recombinant mouse IGF2 [+I] increases phospho-Akt levels [Ser473]. Akt phosphorylation also increases during myogenesis [Mb] myoblast, [D2] day 2 of differentiation, [D4] day 4 of differentiation coincident with increased IGF2 mRNA and protein levels. Loss of MLX function [MLX-DN and MLX shRNA] decreases Akt phosphorylation, whereas MLX-WT increases it, indicating that MLX regulates IGF2/Akt signaling. (D) The dose response
normal control levels [25%) with 25 ng/mL IGF2 and further increased to 40% at the maximum IGF2 concentration of 100 ng/mL. The fusion index of myoblasts overexpressing MLX-WT remained steady at ~40% with all concentrations of IGF2 ranging from 1 to 100 ng/mL, suggesting that myoblast fusion is already maximal in these cells due to MLX-induced Igf2 expression (Fig. 5D). Similarly, myoblasts overexpressing IGF2 displayed a higher fusion index than control cells at all glucose concentrations, including low glucose levels that are typically not permissive for myoblast fusion [Fig. 5E]. Therefore, there is strong interdependence between MLX, IGF2, and glucose levels during myogenesis. Altogether, our data are consistent with an important role for autocrine and paracrine IGF2/Akt signaling in stimulating myoblast fusion and differentiation downstream from MLX.

Muscle regeneration is impaired in muscles of MLX-null mice

To further investigate the role of MLX in myogenesis, we analyzed the muscles of MLX-null mice (Mlx−/−). The tibialis anterior muscles from 12-wk-old mice lacking MLX appeared essentially normal when compared with wild-type (Mlx+/+) controls, with no signs of pathology or impaired muscle development [Fig. 6A], suggesting that fetal myogenesis is unaffected by loss of MLX. Measurement of muscle and myofiber size revealed that, indeed, there were no changes in the proportional area of tibialis anterior muscles from control and Mlx−/− mice, although there was a slight increase in the proportion of larger myofibers in Mlx−/− muscles [Fig. 6B]. In response to injury, myogenesis can also occur in adults in order to regenerate damaged muscles. Importantly, Igf2 is a key myokine known to regulate adult muscle regeneration [Ge et al. 2009; Ikemoto-Uezumi et al. 2015]. We therefore tested whether MLX regulates muscle regeneration via IGF2 signaling. When injected with cardiotoxin, tibialis anterior muscles of wild-type mice efficiently removed necrotic myofibers and displayed regenerating centrally nucleated myotubes 7 d after injury, whereas MLX-null mice had persistent necrotic myofibers and reduced fusion of myotubes, indicating impaired regeneration [Fig. 6A]. Remarkably, cardiotoxin-injured tibialis anterior muscles from Mlx−/− mice displayed impaired regeneration exemplified by an overall decrease in muscle area accompanied by an increased proportion of smaller-sized myotubes compared with isogenic controls [Fig. 6B]. Consistent with reduced myoblast fusion due to the absence of MLX, the expression of its transcriptional targets was also perturbed in regenerating muscle. Decreased Mlx mRNA levels coincided with a severe loss of Txnip expression, which was not influenced by cardiotoxin [Fig. 6C]. Igf2 mRNA, on the other hand, was dramatically up-regulated following cardiotoxin injury, but the increase in Igf2 expression was significantly blunted in Mlx−/− muscles, similar to what is seen in myoblasts lacking MLX activity [Figs. 3,4]. Other myokines regulated by MLX that were identified from RNA-seq [Fig. 3], including Thbs2, Col6a1, Col6a2, and Nov, showed transcriptional regulation similar to that of Igf2 (Supplemental Fig. S7), suggesting that they may also contribute to muscle regeneration. Altogether, these findings indicate that MLX is necessary for efficient muscle regeneration in vivo and that this is due to its capacity to regulate the expression of Igf2 and other myokines.

Discussion

Insulin resistance and diminished intracellular uptake of glucose are components of both aging and diabetes, but it is largely unknown whether they are causally associated with the decline in the regeneration of muscle and other tissues that is observed in these conditions. One possibility is that limited glucose uptake leads to lower ATP production and thus indirectly compromises regeneration by affecting cellular resources available for this process. Alternatively, glucose may act as a signaling molecule that directly modulates pathways responsible for regeneration. This study demonstrates a previously unanticipated role for the glucose-sensing transcription factor MLX in regulating the differentiation and growth of muscle cells via myokine signaling [Fig. 7], suggesting that altered MLX signaling may contribute to decreased regeneration in many disease conditions.

Much of the research on MLX to date has focused on its role in sensing changes in glucose levels in hepatocytes and concordantly adjusting glucose metabolism by increasing the expression of target genes involved in glycolysis and negatively regulating glucose uptake [Ma et al. 2005; Postic et al. 2007; Stoltzman et al. 2008; Peterson et al. 2010]. However, we found that MLX does not regulate glycolysis in muscle precursor cells. Moreover, although MLX increases the expression of Txnip [a negative regulator of glucose uptake] in muscle cells, we observed no changes in the expression of other reported MLX target genes associated with glucose metabolism. Therefore, glucose metabolism does not account for the modulation of myogenesis by MLX gain and loss of function. However, muscle progenitor cells have low glucose uptake and metabolism compared with contracting muscle cells, suggesting that MLX may regulate glucose metabolism in other circumstances. In line with this model, muscle-specific loss of MLXIP does not affect glucose metabolism in the resting state but leads to higher lactate levels in response to short-term fasting and exercise [Minako et al. 2014]. This paradoxical effect, opposite to what is expected, is due to the interaction of MLXIP with PGC-1α and does not require CHORE motifs and MLX [Minako et al. 2014]. Future studies will determine whether MLX can influence glucose metabolism in muscles under specific circumstances.

Surprisingly, we found that MLX regulates the expression of several muscle-secreted signaling proteins [myokines] that are also induced by glucose. Among these, IGF2 recapitulated the effects of MLX in modulating glucose-induced myogenesis by activating the kinase Akt, which is known to be required for myoblast fusion downstream from IGF2 [Jiang et al. 1999]. In glucose-restricted
conditions, myoblasts with increased MLX activity and IGF2 levels indeed fused to a much greater degree than control myoblasts, indicating that glucose-induced signaling mediated by MLX/IGF2/Akt can promote myogenesis in unfavorable conditions. These findings are in line with a previous study that has shown that glucose can promote...
myogenesis by decreasing the activity of the energy-sensing AMPK kinase [Fulco et al. 2008]. In this respect, it is interesting to note that AMPK can also inhibit nuclear translocation and transcriptional activity of MLX heterodimers via phosphorylation [Kawaguchi et al. 2002], suggesting that low glucose can block myogenesis by inhibiting MLX/IGF2 signaling via AMPK. Moreover, AMPK inhibits mTOR, with which both MLX (Carroll et al. 2015) and MLXIP [Kaadige et al. 2015] functionally interact, suggesting multiple levels of cross-talk between MLX and nutrient-sensing pathways. We therefore propose that the glucose-sensing transcription factor MLX and myokine signaling may encompass a novel part of the molecular pathway necessary for glucose-induced adaptive myogenesis. Given its widespread expression, MLX may modulate the transcription of secreted signaling proteins in response to glucose in many cell types, which was also implicated by a recent study in colorectal cancer cells [Nam et al. 2015]. Therefore, our study indicates that MLX has more diverse roles than previously anticipated, presumably in concert with the large network of positive and negative transcriptional regulatory factors to which it belongs [Dioiai et al. 2015].

We discovered a role for MLX in the production of IGF2 and other myokines by myoblasts in response to glucose. Although IGF2 expression was known to increase during muscle differentiation [Kou and Rotwein 1993] and regeneration [Levinovitz et al. 1992] and to modulate these processes, there have not been any transcriptional activators of IGF2 described to date, and only negative regulators of IGF2 expression were known [Markljung et al. 2009; Duputié et al. 2012]. We found that glucose and MLX induce IGF2 transcription, whereas MLX-DN and MLX shRNA reduce its expression. Our study therefore addresses this important area of research by demonstrating that MLX transcriptionally activates IGF2 expression in a glucose-dependent manner during myogenesis. Although the presence of MLX-binding sites [CHORE motifs] and modulation of histone H4 acetylation suggest a direct role for MLX in regulating IGF2 expression, it remains possible that indirect mechanisms are also involved.

Consistent with our findings that MLX regulates myoblast fusion and differentiation in vitro, we found that muscle regeneration is impaired in Mlx-null mice. Compared with controls, loss of MLX resulted in persistence of necrotic myofibers, reduced myotube fusion, lower proportional muscle area, and decreased expression of Igf2 and other injury-induced myokines. Both the reduced myotube fusion and the persistence of necrotic myofibers observed in Mlx-null muscles undergoing regeneration could be explained by reduced Igf2 expression because regenerating aged muscles [which have reduced Igf2 expression] of wild-type mice display similar alterations, and these can be rescued by IGF2 administration [Ikemoto-Uezumi et al. 2015]. Therefore, IGF2 is a key mediator of the myogenic functions of MLX. However, MLX induces the expression of several other myokines in addition to IGF2, and therefore it is possible that regeneration deficits observed in muscles of Mlx-null mice derive also from reduced transcription of these myokines.

The remarkable similarity between the impaired regeneration of aged wild-type mice and young Mlx-null mice suggests a possible role for MLX in the maintenance of muscle homeostasis throughout aging via regulation of myokine expression. On this basis, MLX could delay the progression of sarcopenia [the age-related loss of skeletal muscle mass and function] and promote muscle growth and regeneration in the context of many debilitating diseases characterized by muscle atrophy [Demontis et al. 2013b; Piccirillo et al. 2014].

In summary, we found that MLX is a transcriptional sensor of glucose levels in muscle cells and that it orchestrates adaptive glucose-induced myogenesis by regulating the expression of several myokines. IGF2 is a key MLX-induced myokine that stimulates myoblast fusion and differentiation via Akt in response to glucose and muscle injury. Considering that the activity of glucose-responsive pathways changes during development, aging, and diseases characterized by metabolic dyshomeostasis, our findings provide a possible mechanistic basis for understanding how glucose acts as a signaling molecule to control muscle cell fate in different physiologic and pathologic contexts.

Materials and methods

Cell culture methods

The mouse myoblast C2C12 cell line [American Type Culture Collection, CRL-1772] was maintained in standard conditions, i.e., in high-glucose [4.5 g/L, equivalent to 25 mM] DMEM containing 10% FBS and supplemented with GlutaMax (Life Technologies) in a 5% CO2 incubator at 37°C. To induce differentiation, C2C12 cells were grown to near confluence in 10% FBS-containing growth medium and subsequently in...
C-shLentiviral vector.

Lentiviral transduction

Lentiviral-mediated transduction of myoblasts was achieved by cotransfecting 293FT cells with the lentiviral long-terminal repeat-containing expression vectors alongside the packaging vectors pCMV8.9 and pVSV-G to produce VSV-G pseudo-typed lentiviral vectors. Viral titers were measured by serial dilution and transduction of 293FT cells. C2C12 cells were then transduced with lentiviral vectors at a multiplicity of infection of 100 with 8 µg/mL polybrene for 24 h and expanded. pCDH-EF1-MCS-T2A-GFP lentiviral transduced cells were 100% GFP-positive and maintained transgenic GFP and MLX expression persistently. Following transduction, shRNA transduced cells were further selected with 2 µg/mL puromycin for 7 d, and knockout was tested by qPCR. TG513600A/GIS50306 MLX shRNA resulted in a >70% decrease in MLX mRNA levels relative to the nontargeting shRNA (TR30021) control [Fig. 1], while the other three shRNA were not efficacious in knocking down MLX levels [data not shown].

ELISA

For measuring IGF2 levels in conditioned cell culture medium and cell lysates, the mouse IGF2 Duoset ELISA kit (RND Biosystems, DY792) was used following the manufacturer’s instructions. Conditioned medium was collected after 8 h of incubation with cells, whereas cell lysates were obtained by washing cells with PBS with protease inhibitors and lysing them in PBS by freezing–thawing three times. The conditioned medium and lysate samples were centrifuged, and the supernatants were used for measuring IGF2 levels.

Western blot

Four days after differentiation, muscle cells cultured in 12-well plates were washed with cold PBS and lysed on ice with 200 µL of RIPA buffer with protease and phosphatase inhibitors. Protein concentrations were quantitated with the Bio-Rad protein assay, and all samples were standardized in order to load 20 µg of protein per polyacrylamide gel lane. PAGE was performed, and proteins were transferred to PVDF membranes for probing. Membranes were blocked with 5% milk powder and incubated with primary antibodies overnight at 4°C before incubation with secondary antibodies for 2 h and chemiluminescence imaging. Primary antibodies used were rabbit anti-phospho-Akt [Ser473] D9E [Cell Signaling, 4060], rabbit anti-phospho-Erk antibody [Cell Signaling, 4060], rabbit anti-MLX (Abcam, ab167286), and mouse polyclonal anti-MLX (Abcam, ab167286; this weak-affinity antibody detects MLX upon overexpression but not its endogenous levels).

Immunostaining

Cells were fixed by adding 1 vol of 4% paraformaldehyde to 1 vol of culture medium and incubating for 15 min at room temperature. Cells were then permeabilized with 0.1% Triton X-100, blocked with 2% BSA and 2% horse serum for 2 h, and then incubated with primary antibody overnight at 4°C followed by washes and incubation with Alexa fluor 488- and 555-conjugated secondary antibodies and DAPI. The primary antibodies used were mouse polyclonal anti-MLX (Abcam, ab167286), mouse IgM anti-a-actin 5C5 (Santa Cruz Biotechnology, sc-58670), and mouse anti-myogenin F5D [Dako, IR067], all at a dilution of 1:100. Myogenin levels were assessed by immunostaining after differentiation because myoblasts are myogenin-negative prior to addition of differentiation medium. EDU staining was
performed following EDU labeling for 2 h in culture and following the recommended instructions using the Click-it EdU Alexa fluor 555 imaging kit (Life Technologies, C10338).

Image analysis
Image analysis was done with ImageJ [Schneider et al. 2012]. ImageJ was used to derive the fusion index by counting the nuclei present in syncytial myotubes and within mononucleated myoblasts, the percentage of total nuclei within myotubes, and the percentage of EDU- and myogenin-positive nuclei. ImageJ was also used to quantitate the area of myotubes to derive the myotube size to nuclei ratio.

ChIP
ChIP was performed from C2C12 cells cultured in 15-cm dishes following fixation in 1% formaldehyde for 10 min, quenching with 125 mM glycine for 5 min, washing with PBS, and collection of the cells by centrifugation. Subsequently, Active Motif’s ChIP-IT Express kit (catalog no. 53008) was used following the manufacturer’s instructions. In brief, collected cells were lysed with kit buffers and dounce-homogenized. The nuclear pellet was then sonicated, and chromatin was sheared using a Diagenode Bioruptor with 10 pulses each of 17 sec on and 43 sec off. Chromatin shearing efficiency was assessed by sample purification and gel electrophoresis and consistently yielded sheared chromatin within the range of 200–1000 base pairs (bp), with intense foci around 300–400 bp. Once chromatin shearing was confirmed, immunoprecipitation was performed by incubating 3 µg of antibody with 40 µL of chromatin sample and 25 µL of protein G magnetic beads in a total volume of 100 µL overnight at 4°C on a tube rotator. The beads were then washed and eluted, and the eluted sample was reverse-cross-linked and digested with proteinase K. The eluted DNA was purified using the ReliaPrep FFPE gDNA mini-preparation system [Promega] for real-time qPCR. The fold enrichment was calculated as the proportional change from the naïve IgG control without glucose treatment. The antibodies used were naïve rabbit IgG [EMD Millipore, 12-370] and sc-14705 [Santa Cruz Biotechnology], sc-133397 [Santa Cruz Biotechnology], 13-2500 [clone 9E10, Thermo Fisher Scientific], and 06-598 [EMD Millipore] for MLX, MLXIP, MYC, and H4ac, respectively. Note that the anti-MLX and anti-MLXIP antibodies used were naïve rabbit IgG (EMD Millipore, 12-370) and sc-133397 (Santa Cruz Biotechnology) (Hunt et al. 2011). For ChIP-qPCR experiments appear to have low affinity because, in ChIP-seq (ChIP combined with deep sequencing) experiments, peaks could be called only at the Txnip promoter but nowhere else in the genome. Oligonucleotides used for ChIP-qPCR experiments are listed in Supplemental Table S4, and we confirmed that enrichment was not present at a control locus distal from the promoter (Txnip flanking).

ChIP-seq and data analysis
For ChIP-seq, libraries were prepared from 10 ng of DNA using the NEB Next ChIP-seq library preparation reagent set for Illumina with NEB Next high-fidelity 2× PCR master mix according to the manufacturer’s instructions [New England Biolabs] with the following modifications: A second 1:1 Ampure cleanup was added after adaptor ligation, the Ampure size selection step prior to PCR was eliminated, and the 72°C extension step of the PCR was lengthened to 45 sec. Completed libraries were analyzed for insert size distribution on a 2100 BioAnalyzer high-sensitivity kit [Agilent Technologies] or Caliper LabChip GX DNA high-sensitivity reagent kit [PerkinElmer]. Libraries were quantified using the Quant-it PicoGreen dsDNA assay [Life Technologies] and Kapa library quantification kit [Kapa Biosystems] or low-pass sequencing on a MiSeq nano kit [Illumina]. Fifty-cycle single-end sequencing was performed on an Illumina HiSeq 2500 for DNA obtained from MLX-WT and MLX-DN samples immunoprecipitated for H4ac with three replicates for each sample, including input chromatin. For all of these ChIP-seq data, we used BWA [version 0.5.9-r26-dev, default parameter] to align the reads to mouse genome mm9 [MGSCv37 from Sanger] and then marked duplicated reads with Picard [version 1.65 [1160]], with only nonduplicated reads kept by Samtools [parameter “-q 1 -F 1024” version 0.1.18 [i982:295]]. To control the quality of the data and estimate the fragment size, the nonduplicated version of SPP [version 1.11] was used to draw cross-correlation with support of R [version 2.14.0] using the packages caTools [version 1.17] and bitops [version 1.0–6]. All of our data passed quality control following ENCODE criterion. Upon manually inspecting the cross-correlation plot generated by SPP, the best fragment size estimates (the smallest fragment size estimated by SPP in all of our cases) were used to extend each read and combine replicates to generate bigwig files to view on IGV [version 2.3.40]. After we observed the high consistency between replicates, we scaled the bigwig files and normalized them to 60 M reads so that the peaks of samples are roughly comparable. We generated the normalized H4ac signal on the bigwig files by bigWigSummary from University of California at Santa Cruz and averaged it across gene sets [Kent et al. 2010]. In order to identify regions of variable H4 acetylation with MLX genotypes, we considered the read count ratio versus input normalized by total reads; i.e., (Rep1 + Rep2 + Rep3 + 1)/(Input1 + Input2 + Input3 + 1)/([TotalRep1 + TotalRep2 + TotalRep3]/[TotalInput1 + TotalInput2 + TotalInput3]) for 4-kb regions centered around the TSS from the RefSeq database. This allowed for identification of differentially acetylated promoter regions that was further confirmed by ChIP-qPCR. ChIP-seq results are available at the Gene Expression Omnibus (GEO) with GEO accession number GSE74678.

Computational prediction of CHORE motifs
To predict potential CHORE-like sequences in proximity to H4ac peaks, we used putative CHORE sequences previously identified from ChIP-seq of MLXIPL (CHREBP) in human adipose and liver tissue (Pourngvarin et al. 2015) to generate a position weight matrix. This matrix was used to scan the motif matches on the mouse mm9 genome by STORM [Schones et al. 2007], with a P-value cutoff of 1 × 10⁻⁶.

Real-time qPCR
RNA was extracted from cultured cells with Trizol reagent (Life Technologies) followed by RNeasy minikit (Qiagen) cleanup and DNase treatment. RNA was quantitated, and 500 ng was reverse-transcribed into cDNA using iScript cDNA synthesis kit (Bio-Rad). The resulting cDNA was used for qPCR with SYBR Green (Bio-Rad) with validated oligonucleotides for specific transcripts. Normalized expression of qPCR data was obtained by following the ΔΔCt method and using the housekeeper genes Ppia (for cell culture samples) and Hprt [for regeneration studies], which were previously found to be reliable normalization control genes in myoblast culture conditions and muscle regeneration experiments [Hunt et al. 2013]. Oligonucleotide sequences used for qPCR are listed in Supplemental Table S4.

RNA-seq and data analysis
RNA was extracted as described above. Three biological replicates were prepared for RNA-seq with the TruSeq stranded
mRNA library preparation kit (Illumina) and sequenced on the Illumina HiSeq 2000 platform. FastQ sequences were mapped to the mouse mm9 genome by StrongARM, developed for the Pediatric Cancer Genome Project (Downing et al. 2012). Mapped reads were counted with HTSeq (Anders et al. 2015), and gene-level FPKM (fragments per kilobase per million mapped fragments) values were computed. All sample data were collated into a matrix using R (3.0.1) and log start-transformed $\log_2$(FPKM + 1) in STATA/MP 11.2. Genes were statistically tested by class ($t$-test, unequal variance $t$-test per gene). Bonferroni correction and false discovery rates (FDRs) were determined to allow both strict and modest multiple comparison filtering (Partek Genomics suite 6.6). Six-hundred-fifty-five genes were selected that had an absolute value log ratio of at least 1 in at least one comparison and passed FDR at 5% (vs. respective controls). We then filtered for genes that showed concordant regulation with MLX-WT, MLX-DN, and MLX shRNA; i.e., genes that change in the same directions in MLX-DN and MLX shRNA samples and do not have the same directional change in the MLX-WT samples. The 80 genes concordantly regulated by MLX were identified because they satisfied a minimum of two out of three of the following requirements: a significant change of at least 1.5-fold compared with control for (1) MLX-WT, (2) MLX-DN, and (3) MLX shRNA; that is, the 80 selected genes were either regulated consistently by both MLX-DN and MLX shRNA or in an opposite fashion by MLX-WT and MLX-DN and/or MLX shRNA. These were analyzed for enriched gene sets by using the SwissProt PIR (SP_PIR) and DAVID. The results of RNA-seq mapping and quantitation, the 80 selected genes, and the enriched gene clusters are presented in Supplemental Table S1. RNA-seq results are available at the Gene Expression Omnibus with accession number GSE70028.

**Metabolic analyses**

Extracellular lactate from cell culture medium was quantitated with the glycolysis cell-based assay kit (Cayman Chemical). Glucose uptake was measured with the glucose uptake cell-based assay kit (Cayman Chemical) following the plate-reading method indicated in the manufacturer's instructions. Seahorse analysis was done using a Seahorse XF24 analyzer, the XF glycolysis stress test kit, and the XF cell mito stress test kit after optimization of drug concentrations for C2C12 cells (the concentrations used are indicated in Fig. 2).

**Statistical analyses and calculations**

Data shown represents the mean ± SEM with a minimum of three biological replicates. Statistical comparisons between groups were done using GraphPad Prism software and one-way or two-way ANOVA with Dunnet’s multiple comparison tests for assays with one and two variables, respectively. Z-score calculations, RNA-seq graphs, and heat maps were generated with Spotfire DecisionSite software.

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**References**

Alzhanov DT, McInerney SF, Rotwein P. 2010. Long range interactions regulate Igf2 gene transcription during skeletal muscle differentiation. *J Biol Chem* 285: 38969–38977.

Anders S, Pyl PT, Huber W. 2015. HTSeq—a Python framework to work with high-throughput sequencing data. *Bioinformatics* 31: 166–169.

Billin AN, Eilers AL, Queva C, Ayer DE. 1999. Mlx, a novel Max-like BHLLHZip protein that interacts with the Max network of transcription factors. *J Biol Chem* 274: 36344–36350.

Billin AN, Eilers AL, Coulter KL, Logan JS, Ayer DE. 2000. MondoA, a novel basic helix–loop–helix-leucine zipper transcriptional activator that constitutes a positive branch of a max-like network. *Mol Cell Biol* 20: 8845–8854.

Boström P, Wu J, Jedyshowsky MF, Korde A, Ye L, Lo JC, Rasbach KA, Boström EA, Choi JH, Long JZ. 2012. A PGC1-a dependent myokine that drives brown-fat-like development of white fat and thermogenesis. *Nature* 481: 463–468.

Broholm C, Laye MJ, Brandt C, Vadalasetty R, Pilegaard H, Pedersen BK, Scheele C. 2011. LIF is a contraction-induced myokine stimulating human myocyte proliferation. *J Appl Physiol* 111: 251–259.

Carroll PA, Diolaiti D, McFerrin L, Gu H, Djkovic D, Du J, Cheng PF, Anderson S, Ulrich M, Hurley JB. 2015. Deregulated Myc requires MondoA/Mlx for metabolic reprogramming and tumorigenesis. *Cancer Cell* 27: 271–285.

Cha-Molstad H, Saxena G, Chen J, Shalev A. 2009. Glucose-stimulated expression of Tmxp is mediated by carbohydrate response element-binding protein, p300, and histone H4 acetylation in pancreatic β cells. *J Biol Chem* 284: 16986–16995.

Davies MN, O’Callaghan BL, Towle HC. 2008. Glucose activates ChREBP by increasing its rate of nuclear entry and relieving repression of its transcriptional activity. *J Biol Chem* 283: 24029–24038.

Demontis F, Piccirillo R, Goldberg AL, Perrimon N. 2013a. The influence of skeletal muscle on systemic aging and lifespan. *Aging Cell* 12: 943–949.

Demontis F, Piccirillo R, Goldberg AL, Perrimon N. 2013b. Mechanisms of skeletal muscle aging: insights from *Drosophila* and mammalian models. *Dis Model Mech* 6: 1339–1352.

Demontis F, Patel VK, Swindell WR, Perrimon N. 2014. Intertissue control of the nucleolus via a myokine-dependent longevity pathway. *Cell Rep* 7: 1481–1494.

Diolaiti D, McFerrin L, Carroll PA, Eisenman RN. 2015. Functional interactions among members of the MAX and MLX transcriptional network during oncogenesis. *Biochim Biophys Acta* 1849: 484–500.

Downing JR, Wilson RK, Zhang J, Mardis ER, Pui C-H, Ding L, Ley TJ, Evans WE. 2012. The pediatric cancer genome project. *Nat Genet* 44: 619–622.

Duputé A, Antoine E, Apelt N, Milligan L, Carbonell F, Lelay-Taha M-N, Piette J, Weber M, Montarras D, Pinset C. 2012. H19 antisense RNA can up-regulate Igf2 transcription by...
activation of a novel promoter in mouse myoblasts. *PLoS One* 7: e37923.

Erbay E, Park I-H, Nuzzi PD, Schoenherr CJ, Chen J. 2003. IGF-II transcription in skeletal myogenesis is controlled by mTOR and nutrients. *J Cell Biol* 163: 931–936.

Florini J, Magri K, Ewton D, James P, Grindstaff K, Rotwein P. 1991a. ‘Spontaneous’ differentiation of skeletal myoblasts is dependent upon autocrine secretion of insulin-like growth factor-II. *J Biol Chem* 266: 15197–15193.

Florini JR, Ewton DZ, Magri KA. 1991b. Hormones, growth factors, and myogenic differentiation. *Annu Rev Physiol* 53: 201–216.

Fulco M, Cen Y, Zhao P, Hoffman EP, McBurney MW, Sauve AA, Sartorelli V. 2008. Glucose restriction inhibits skeletal myoblast differentiation by activating SIRT1 through AMPK-mediated regulation of Nampt. *Dev Cell* 14: 661–673.

Ge Y, Wu A-L, Warness C, Liu J, Zhang C, Kawasome H, Terada N, Boppard MD, Schoenherr CJ, Chen J. 2009. mTOR regulates skeletal muscle regeneration in vivo through kinase-dependent and kinase-independent mechanisms. *Am J Physiol Cell Physiol* 297: C1434–C1444.

Guinez C, Filhoulaud G, Rayah-Benhamed F, Marmier S, Dubuquoy C, Dentin R, Moldes M, Burnol A-F, Yang X, Lefebvre T, et al. 2011. O-GlcNAcylation increases ChREBP protein content and transcriptional activity in the liver. *Diabetes* 60: 1399–1413.

Hunt L, Tudor E, White J. 2010. Leukemia inhibitory factor-dependent increase in myoblast cell number is associated with phosphotyidylinositol 3-kinase-mediated inhibition of apoptosis and not mitosis. *Exp Cell Res* 316: 1002–1009.

Hunt LC, Upadhaya A, Jazayeri JA, Tudor EM, White JD. 2011. Caspase-3, myogenic transcription factors and cell cycle inhibitors are regulated by leukemia inhibitory factor to mediate inhibition of myogenic differentiation. *Skeletal Muscle* 1: 17.

Hunt L, Gorman C, Kintakas C, McCulloch D, Mackie E, White JD. 2011. Dominant negative Mlx improves metabolic profile in diabetic and kinase-independent mechanisms. *Am J Physiol Endocrinol Metab* 291: C3168–C3175.

Iizuka K, Takeda J, Horikawa Y. 2009. Hepatic overexpression of Ikemoto-Uezumi M, Uezumi A, Tsuchida K, Fukada S-i, Yamada H, Kuroki T, et al. 2008. Warburg effect regulated by amphiregulin in the development of colorectal cancer. *Cancer Med* 4: 575–587.

Iruela-Arispe ML, Liska DJ, Sage EH, Bornstein P. 1993. Differential expression of thrombospondin 1, 2, and 3 during murine development. *Dev Dyn* 197: 40–56.

Jiang B-H, Aoki M, Zheng JZ, Li J, Vogt PK. 1999. Myogenic signaling of phosphatidylinositol 3-kinase requires the serine-threonine kinase Akt/protein kinase B. *Proc Natl Acad Sci U S A* 96: 2077–2081.

Kaadige MR, Yang J, Wilde BR, Ayer DE. 2015. MondoA–Mlx transcriptional activity is limited by mTOR–MondoA interaction. *Mol Cell Biol* 35: 101–110.

Kawaguchi T, Osatomi K, Yamashita H, Kabashima T, Uyeda K. 2002. Mechanism for fatty acid ‘sparing’ effect on glucose-induced transcription regulation of carbohydrate-responsive element binding–protein by AMP-activated protein kinase. *J Biol Chem* 277: 3829–3835.

Kent WJ, Zweig AS, Barber G, Hinrichs AS, Karolchik D. 2010. BigWig and BigBed: enabling browsing of large distributed datasets. *Bioinformatics* 26: 2204–2207.

Kou K, Rotwein P. 1993. Transcriptional activation of the insulin-like growth factor-II gene during myoblast differentiation. *Mol Endocrinol* 7: 291–302.

Lee JD, Carrier A, Shah S, Snyder NW, Wei S, Venneti S, Worth AJ, Yuan Z-F, Lim H-W, Liu S. 2014. Akt-dependent metabolic reprogramming regulates tumor cell histone acetylation. *Cell Metab* 20: 306–319.

Levinovitz A, Jennische E, Oldfors A, Edwall D, Norstedt G. 1992. Activation of insulin-like growth factor II expression during skeletal muscle regeneration in the rat: correlation with myotube formation. *Mol Endocrinol* 6: 1227–1234.

Ma L, Tsatsos NG, Towlle HC. 2005. Direct role of ChREBP-Mlx in regulating hepatic glucose-responsive genes. *J Biol Chem* 280: 12019–12027.

Ma L, Robinson LN, Towlle HC. 2006. ChREBP-Mlx is the principal mediator of glucose-induced gene expression in the liver. *J Biol Chem* 281: 28721–28730.

Markljung E, Jiang L, Jaffe JD, Mikkelson TS, Wallerman O, Latham M, Zhang X, Wang L, Saen- Vash Y, Gnrkire A. 2009. ZBED6, a novel transcription factor derived from a domesticated DNA transposon regulates IGF2 expression and muscle growth. *PLoS Biol* 7: e1000256.

McFerrin LG, Atchley WR. 2012. A novel N-terminal domain may dictate the glucose response of Mondo proteins. *PLoS One* 7: e34803.

Minako I, Benny H-JC, Motoyuki K, Ming L, Byoungsoon H, Heinrich T, Robert AH, Lawrence C. 2014. MondoA deficiency enhances sprint performance in mice. *Biochem J* 464: 35–48.

Minn AH, Hafele C, Shaley A. 2005. Thioredoxin-interacting protein is stimulated by glucose through a carbohydrate response element and induces β-cell apoptosis. *Endocrinology* 146: 2397–2405.

Nam SO, Yotsumoto F, Miyata K, Fukagawa S, Yamada H, Kuroki M, Miyamoto S. 2015. Warburg effect regulated by amphiregulin in the development of colorectal cancer. *Cancer Med* 4: 575–587.

Nedachi T, Kadotani A, Ariga M, Katagiri H, Kanazaki M. 2008. Ambient glucose levels qualify the potency of insulin myogenic actions by regulating SIRT1 and FoxO3a in C2C12 myocytes. *Am J Physiol Endocrinol Metab* 294: E668–E678.

Parikh H, Carlsson E, Chutkwak WA, Johansson LE, Storgaard H, Poulsen P, Saxena R, Ladd C, Schulze PC, Mazzini MJ. 2007. TXNIP regulates peripheral glucose metabolism in humans. *PLoS Med* 4: e158.

Pedersen BK, Febbraio MA. 2012. Muscles, exercise and obesity: skeletal muscle as a secretory organ. *Nat Rev Endocrinol* 8: 457–465.

Peterson CW, Stoltzman CA, Sighinolfi MP, Han K-S, Ayer DE. 2010. Glucose controls nuclear accumulation, promoter binding, and transcriptional activity of the MondoA–Mlx heterodimer. *Mol Cell Biol* 30: 2887–2895.

Piccirillo R, Demonis F, Perrimon N, Goldberg AL. 2014. Mechanisms of muscle growth and atrophy in mammals and *Drosophila*. *Dev Dyn* 243: 201–215.

Postic C, Dentin R, Denechau P-D, Girard J. 2007. ChREBP, a transcriptional regulator of glucose and lipid metabolism. *Annu Rev Nutr* 27: 179–192.

Pourgharib N, Chang B, Imamura M, Chen J, Mooluwan K, Sae-Lee C, Li W, Chan L. 2015. Genome-wide analysis of ChREBP binding sites on male mouse liver and white adipose chromatin. *Endocrinology* 156: 1982–1994.
Glucose induces myogenesis via myokines

Rai M, Demontis F. 2015. Systemic nutrient and stress signaling via myokines and myometabolites. *Annu Rev Physiol* doi: 10.1146/annurev-physiol-021115-105305.

Sakamoto K, Yamaguchi S, Ando R, Miyawaki A, Kabasawa Y, Takagi M, Li CL, Perbal B, Katsube K-i. 2002. The nephroblastoma overexpressed gene (NOV/ccn3) protein associates with Notch1 extracellular domain and inhibits myoblast differentiation via Notch signaling pathway. *J Biol Chem* 277: 29399–29405.

Schneider CA, Rasband WS, Eliceiri KW. 2012. NIH Image to ImageJ: 25 years of image analysis. *Nat Methods* 9: 671–675.

Schones DE, Smith AD, Zhang MQ. 2007. Statistical significance of cis-regulatory modules. *BMC Bioinformatics* 8: 19.

Shen L, O’Shea JM, Kaadige MR, Cunha S, Wilde BR, Cohen AL, Welm AL, Ayer DE. 2015. Metabolic reprogramming in triple-negative breast cancer through Myc suppression of TXNIP. *Proc Natl Acad Sci* 112: 5425–5430.

Stoltzman CA, Kaadige MR, Peterson CW, Ayer DE. 2011. MondoA senses non-glucose sugars: regulation of thioredoxin-interacting protein (TXNIP) and the hexose transport curb. *J Biol Chem* 286: 38027–38034.

Stixier V, Bataillè L, Etard C, Jańska T, Weger M, DaPonte JP, Strähle U, Dickmeis T, Jańska K. 2013. Glycolysis supports embryonic muscle growth by promoting myoblast fusion. *Proc Natl Acad Sci* 110: 18982–18987.

Webster KA, Gunning P, Hardeman E, Wallace DC, Kedes L. 1990. Coordinate reciprocal trends in glycolytic and mitochondrial transcript accumulations during the in vitro differentiation of human myoblasts. *J Cell Physiol* 142: 566–573.

Wu N, Zheng B, Shaywitz A, Dagon Y, Tower C, Bellinger G, Shen C-H, Wen J, Asara J, McGraw TE. 2013. AMPK-dependent degradation of TXNIP upon energy stress leads to enhanced glucose uptake via GLUT1. *Mol Cell* 49: 1167–1175.

Yahiaoui L, Gvozdic D, Danialou G, Mack M, Petrof BJ. 2008. CC family chemokines directly regulate myoblast responses to skeletal muscle injury. *J Physiol* 586: 3991–4004.

Yamashita H, Takenoshita M, Sakurai M, Bruick RK, Henzel WJ, Shillinglaw W, Arnot D, Uyeda K. 2001. A glucose-responsive transcription factor that regulates carbohydrate metabolism in the liver. *Proc Natl Acad Sci* 98: 9116–9121.

Yoon M-S, Chen J. 2008. PLD regulates myoblast differentiation through the mTOR–IGF2 pathway. *J Cell Sci* 121: 282–289.

Yu F-X, Goh S-R, Dai R-P, Luo Y. 2009. Adenosine-containing molecules amplify glucose signaling and enhance txnip expression. *Mol Endocrinol* 23: 932–942.