Chronic erythropoietin treatment improves diet-induced glucose intolerance in rats

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Abstract

Erythropoietin (EPO) ameliorates glucose metabolism through mechanisms not fully understood. In this study, we investigated the effect of EPO on glucose metabolism and insulin signaling in skeletal muscle. A 2-week EPO treatment of rats fed with a high-fat diet (HFD) improved fasting glucose levels and glucose tolerance, without altering total body weight or retroperitoneal fat mass. Concomitantly, EPO partially rescued insulin-stimulated AKT activation, reduced markers of oxidative stress, and restored heat-shock protein 72 expression in soleus muscles from HFD-fed rats. Incubation of skeletal muscle cell cultures with EPO failed to induce AKT phosphorylation and had no effect on glucose uptake or glycogen synthesis. We found that the EPO receptor gene was expressed in myotubes, but was undetectable in soleus. Together, our results indicate that EPO treatment improves glucose tolerance but does not directly activate the phosphorylation of AKT in muscle cells. We propose that the reduced systemic inflammation or oxidative stress that we observed after treatment with EPO could contribute to the improvement of whole-body glucose metabolism.

Key Words
- glucose metabolism
- high-fat diet
- skeletal muscle
- myotubes

Introduction

Erythropoietin (EPO) is a glycoprotein synthesized by the kidney and plays a role in the regulation of red blood cell growth and survival. EPO, classically used for the treatment of anemia in chronic kidney disease, has recently emerged as a regulator involved in glucose metabolism. Results of early studies carried out on patients...
with end-stage renal disease (ESRD) indicated that EPO not only treated anemia but also improved insulin sensitivity (Borissova et al. 1993, Mak 1996). Results of other clinical studies of ESRD patients indicated that EPO treatment could improve fasting plasma insulin level (Tuozzu et al. 2004, Khedr et al. 2009) and control blood glucose (Spiaia et al. 2000, Khedr et al. 2009). The results of these studies have indicated that improvements in glucose homeostasis and insulin sensitivity occurred before, or independently of increased hematocrit, indicating a metabolic role for EPO that is dissociated from hematopoiesis (Spiaia et al. 2000). Animal models of long-term exposure to EPO showed clear positive effects on glucose metabolism. Transgenic mice constitutively overexpressing EPO displayed decreased body weight and improved glucose metabolism as compared with WT litter mates (Katz et al. 2010). Conversely, long-term exposure to EPO could prevent diet-induced obesity (Hojman et al. 2009). While a role of EPO in mediating muscle anabolism has been suggested, evidence that EPO directly targets skeletal muscle is still lacking.

Results of cell culture experiments indicated that EPO has the potential to directly target metabolic cells. Acute treatment with EPO protects pancreatic cells against the combined action of pro-apoptotic cytokines interleukin 1 (IL1β) tumor necrosis factor alpha (TNFα), and interferon gamma (IFNγ), suggesting indicative of its cytoprotective role (Maiese et al. 2005). In cultured adipocytes, EPO prevented dexamethasone-induced decrease in glucose uptake (Mikolás et al. 2012, Pan et al. 2013).

In this study, we investigated the effect of both acute and chronic EPO treatment on whole-body glucose metabolism and activation of the insulin signaling pathway in skeletal muscle. We found that EPO ameliorates diet-induced glucose intolerance and insulin signaling. We also show that EPO does not target the skeletal muscle cell indicating an indirect action through the improvement of several systemic factors known to be beneficial for whole-body glucose metabolism.

Materials and methods

Chemicals, ELISA kits, and antibodies

Palmitic acid endotoxin-free cell culture grade, anti-protease cocktail (P8340), and BSA (cell culture grade) were purchased from Sigma–Aldrich. Fetal bovine serum (FBS), minimum essential media alpha (MEMα), and penicillin/streptomycin (P/S) antibiotic cocktail (15070-063) were from purchased Gibco. The experiments were conducted with human recombinant EPO (Janssen Cilag, Sydney, NSW, Australia). Primary antibodies were from Stressgen (heat-shock protein 72 (HSP72), SPA-810, 1/2000, Victoria, BC, Canada) or Cell Signaling (AKT (9272), P-AKT\(^{\text{Ser473}}\) (9271), Danvers, MA, USA), and secondary antibodies were purchased from Sigma–Aldrich (sheep anti-mouse, A5906) or Cell Signaling (goat anti-rabbit, 7074). Serum insulin and TNFz were measured using rat ultrasensitive ELISA Kits (Alpco, Laboratory Diagnostics (Caringbah, NSW, Australia) and R&D Systems, Inc. (Minneapolis, MN, USA) respectively). Triglycerides (TGs) and cholesterol were measured using Thermo Scientific (Waltham, MA, USA) and Stanbio Kits respectively (Boerne, TX, USA). Blood glucose levels were measured using an AcuCheck II glucometer (Roche, Castle Hill, NSW).

Cell culture

L6 and 3T3L1 cells were obtained from Lonza (Stockholm, Sweden). Human muscle primary cell cultures were established from muscle biopsies collected from male participants. The protocol was approved by the Local Ethics Committee (03/10/GESE, Montpellier, France) and informed and written consent was obtained from all participants. Myoblasts isolation, purification, and culture were performed as described previously (Barto et al. 2010). Muscle cells were grown in medium containing MEMα supplemented with 10% FBS and 50 μg/ml P/S at 5% CO2 and 37 °C. Once at confluence, cells were grown in MEMα, 2% FBS, and 50 μg/ml P/S. All experiments were conducted at the myotube stage of differentiation. The medium was changed to MEMz, 1% BSA, P/S, with serum 12 h before insulin stimulation. As a model of insulin-resistance, muscle cells were treated for 48 h with 500 μM palmitate. The cells incubated with EPO (10 U/ml final concentration) or palmitate were stimulated with 0, 10, and 100 nM insulin for 5 min. In the different experiments, cells were incubated with EPO for 5 min, 15 min, or 48 h. The cells were then washed twice in ice-cold phosphoprotein buffer (20 mM Tris, pH 7.4, 150 mM NaCl, 5 mM EDTA, 150 mM NaF, 2 mM Na\(_2\)VO\(_4\), and 10 mM pyrophosphate) and lysed (lysis buffer: 100 μl 0.5 mM phenylmethylsulfonyl fluoride (PMSF), protease inhibitor, and 1% Triton X-100 in protein buffer). The cell lysates were centrifuged for 10 min at 4 °C. The soluble fraction was collected and suspended in 100 μl of a Laemmlli buffer/β-mercaptoethanol mixture and stored at −20 °C.

Glycogen synthesis

Glycogen synthesis was measured using a modified version of a previously described method (Al-Khalili et al. 2003).
Briefly, L6 myotubes were incubated with EPO (10 mU/ml) for 24 h, and then serum starved for 4 h. After stimulation with insulin at 0 or 120 nM for 30 min, cells were supplemented with [3H]-deoxyglucose (0.1 mM and 0.5 mM) BSA. Glucose transport was determined by the addition of insulin (100 nM) for 20 min in the presence of Krebs–Ringer phosphate buffer supplemented with 0.1% BSA. Glucose transport was determined by the addition of glucose–insulin index was calculated as previously reported (Marchionne et al. 2012). The animals were returned to their cage and received the final EPO injection. Two days later, after an overnight fast, animals were assigned to a nonstimulated (INS−) or insulin-stimulated (INS+) condition (1 IU insulin i.p./kg body weight). Soleus and extensor digitorum longus (EDL) muscles were collected 10 min after injection. In both acute and chronic EPO treatment studies, animals were killed by a lethal injection of Lethabarb (325 mg/ml pentobarbitone sodium) followed by surgical removal of the heart.

Western blotting analysis

The cell lysates and tissue samples were analyzed by western blotting as described previously (Barrés et al. 2009). Briefly, the tissues were homogenized in a 10:1 (volume-to-weight) ratio of ice-cold extraction buffer (10 mM Tris–HCl (pH 7.4); 100 mM NaCl; 1 mM each of EDTA, EGTA, NaF, and PMSF; 2 mM Na3VO4; 20 mM Na2P2O7; 1% Triton X-100; 10% glycerol; 0.1% SDS; 0.5% deoxycholate; and 10 μg/ml protease inhibitor cocktail). Solubilized proteins were suspended in Laemmli buffer, heated for 5 min at 95 °C or 20 min at 65 °C (for phosphorylated proteins). Proteins (20–90 μg) were separated by SDS–PAGE and transferred onto nitrocellulose membranes. The membranes were blocked, washed, and incubated overnight at 4 °C with primary antibody (P-AKT and total AKT 1/2000 in TBST 5% BSA; HSP72, 1/2000 1% milk). Following incubation with HRP-conjugated secondary antibodies, proteins were detected by ECL (Millipore, Christensen et al. 2012). The tissues were collected 60, 90, and 180 min following injection, snap–frozen in liquid nitrogen and stored at −80 °C until analysis was performed. Blood glucose levels were measured in a separate cohort of adult rats.
Bayswater, VIC, Australia) and visualized using the ChemiDoc XRS+ and Image Lab Software (Bio-Rad).

RNA extraction and RT-PCR

RNA extraction and RT-PCR was performed using a modified version of a previously described method (Barres et al. 2012). In brief, RNA from rat bone marrow, rat skeletal muscle, and L6 cells was purified using TRIzol reagent according to the recommendations of the manufacturer. cDNA synthesis was performed using the iScript cDNA Synthesis Kit (Bio-Rad). PCR was performed in a final volume of 10 μl, consisting of diluted cDNA, SYBR Green PCR Master Mix, EPO receptor (EPOR) primer, and nuclease-free water. The samples were analyzed in triplicates. The primers were designed using Primer3 as follows: sense, AGGTGGACGTGTCAGCAGGC and antisense, CCCGCAGGTTGCTCAGGAC.

Oxidative stress

F2-isoprostanes (F2-IsoPs) and isofurans (IsoFs), formed in vivo by free radical nonenzymatic oxidation of arachidonic acid (C20:4n-6), were quantified (Morrow et al. 1990, Henriksen et al. 2011). F2-IsoPs and IsoFs in soleus muscle were measured by gas chromatography–mass spectrometry using electron-capture negative ionization as previously described (Mas et al. 2011).

Statistical analyses

The results are presented as mean ± S.D. All data were tested for normality and equality of variance. Data derived from western blots in the acute EPO studies were analyzed using a two-way ANOVA when only three groups were analyzed (CTL, HFD, and HFD/EPO). Insulin stimulation in vivo was analyzed with a two-way ANOVA (ins×groups). Following ANOVA, post hoc analysis was performed using the Bonferroni’s test for multiple comparisons. For cell culture experiments, a two-way ANOVA (insulin×EPO) was used for analysis of AKT activation following incubation with insulin and EPO once data were normalized via appropriate transformation. Nonparametric (Kruskal–Wallis or Wilcoxon–Mann–Whitney U) tests were used for palmitate, glycogen synthesis, and glucose uptake experiments. The correlations were investigated using the Pearson’s product-moment coefficient. The significance level for all comparisons was set at P < 0.05. The statistical analysis was conducted with SPSS (IBM).

Results

Acute and chronic EPO injection in lean animals is associated with an increased phosphorylation of AKT in muscle

We tested the effect of EPO on the activation of glucose metabolism in skeletal muscle by measuring the phosphorylation of AKT, a signaling node in the metabolic action of insulin. After an acute i.p. injection of EPO, AKT phosphorylation at serine 473 was elevated in soleus muscle with peak activation at 90 min (T0 vs 90 min: t = −2.806, P < 0.05; Fig. 1A and B). Despite enhanced phosphorylation of AKT in skeletal muscle, acute EPO did not affect glycemia within a 90 min period following injection (Fig. 1C), indicating that EPO-induced AKT signaling activation in vivo in skeletal muscle is not sufficient to alter whole-body glucose homeostasis. Treatment of lean animals with EPO for 2 weeks also resulted in increased basal AKT phosphorylation in soleus (t = −2.21, P = 0.045; Fig. 1D and E).

Figure 1

EPO activates AKT phosphorylation in rat soleus muscle. (A) Quantification of P-AKT/total AKT in soleus muscle following acute i.p. injection of 1300 IU EPO/kg (n = 4) and (B) western blot images. (C) Blood glucose time course following i.p. injection of 1300 IU EPO/kg (n = 4). (D) Quantification of P-AKT/total AKT following chronic EPO treatment (500 U/kg, three times a week for 2 weeks) in chow-fed animals (n = 8) and (E) western blot images. *P < 0.05 indicates a statistically significant difference between groups.

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EPO improves glucose tolerance and partially restores insulin-stimulated AKT phosphorylation in high-fat-fed rats

The activation of AKT signaling following injection of EPO prompted us to determine the effect of repeated EPO treatment on glucose metabolism. Lean and diet-induced-obese rats were chronically treated with EPO for 2 weeks. We controlled for the fact that, at initiation of the EPO treatment, animals fed HFD were bigger than chow-fed animals and exhibited signs of impaired glucose metabolism, as shown by higher blood glucose (P<0.05; Table 1). At the completion of the study, the 6-week HFD resulted in increased fasting insulin and glucose levels (P<0.05; Fig. 2A and B). EPO lowered fasting blood glucose in HFD/EPO animals (diet×EPO interaction: F(1,32)=6.1, P<0.05; Fig. 2A) compared with HFD, while serum insulin levels were not significantly affected (diet×EPO interaction: F(1,28)=2.1, P=0.15; Fig. 2B). Animals fed with a HFD and treated with EPO also showed improved glycemic control as indicated by reduced blood glucose levels during an IPGTT (F(1,28)=3.09, P=0.040; Fig. 2C) with a significant difference between HFD and HFD/EPO at 15 min. Glucose area under the curve (AUC; Fig. 2D) was increased in the HFD group and normalized in the HFD/EPO group, while insulin AUC increased with HFD and was unchanged by EPO treatment (Fig. 2E).

Accordingly, the glucose–insulin index was increased in the HFD group and reduced in the HFD/EPO group (P<0.05; Fig. 2F).

Energy intake was similar during the course of the study (Fig. 3A). Chronic EPO injection did not modify weight gain or fat accumulation in EPO or HFD/EPO rats compared with CTL and HFD respectively, as measured by body or retroperitoneal fat pad weights (Fig. 3B and C). These results indicate that EPO ameliorates glucose metabolism independently from changes in body weight or retroperitoneal fat mass. While circulating TGs were unchanged, total-cholesterol levels were decreased after EPO treatment (Table 1).

To determine if EPO treatment ameliorated insulin-sensitivity in skeletal muscle, we injected insulin into fasting CTL, HFD, and HFD/EPO animals and assessed AKT phosphorylation in oxidative and glycolytic muscles. In soleus, there was an interaction between insulin and treatment groups (F(1,46)=3.60, P=0.045; Fig. 4A and B). Insulin markedly stimulated AKT phosphorylation in CTL (t=−4.76, P<0.01), while HFD inhibited its action (Fig. 4A and B). Although EPO treatment restored insulin-stimulated AKT phosphorylation in HFD/EPO compared with CTL, we did not find a significant difference compared with baseline level (Ins− versus Ins+, P>0.05). Increased baseline levels of P-AKT in this group could explain this result (Fig. 4A). In EDL, insulin-induced AKT phosphorylation was inhibited in HFD, yet EPO did not restore insulin signaling (P>0.05; Fig. 4C and D).

EPO treatment is associated with reduced inflammation, and oxidative stress, and increases HSP72 expression

To determine the potential contributors of EPO-induced amelioration of glucose metabolism or AKT phosphorylation in muscle, we investigated the effect of chronic EPO on inflammation levels, oxidative stress, and expression of HSP72, a chaperone protein known to protect insulin signaling (Chung et al. 2008). Systemic levels of the

### Table 1 Parameters assessed in this study

<table>
<thead>
<tr>
<th>Parameter</th>
<th>CTL</th>
<th>EPO</th>
<th>HFD</th>
<th>HFD/EPO</th>
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<td>n=16</td>
<td>n=16</td>
<td>n=16</td>
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<td>Body weight week 4 (g)</td>
<td>336±24</td>
<td>392±35*</td>
<td>464±34†</td>
<td>466±49†</td>
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<td>FBG (mM)</td>
<td>5.59±0.65</td>
<td>6.48±0.47*</td>
<td>7.0±0.53†</td>
<td>7.2±0.67†</td>
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<tr>
<td>Animal characteristics at the end of the study</td>
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<td>n=8</td>
<td>n=8</td>
<td>n=8</td>
</tr>
<tr>
<td>Body weight (g)</td>
<td>367±14</td>
<td>377±44</td>
<td>464±34†</td>
<td>466±49†</td>
</tr>
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<td>TNFα (pg/ml)</td>
<td>0.70±0.67†</td>
<td>0.77±0.67†</td>
<td>2.03±0.53†</td>
<td>1.06±0.67†</td>
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<tr>
<td>Hct (%)</td>
<td>49.2±1.2</td>
<td>67.9±2.5†</td>
<td>47.3±1.2</td>
<td>61.9±1.9†</td>
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<tr>
<td>TGs (mM)</td>
<td>0.29±0.11†</td>
<td>0.29±0.08†</td>
<td>0.43±0.20†</td>
<td>0.43±0.16†</td>
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<tr>
<td>Total cholesterol (mg/dl)</td>
<td>69.0±13.7</td>
<td>48.4±17.2</td>
<td>75.6±11.2</td>
<td>59.6±15.1†</td>
</tr>
</tbody>
</table>

FBG, fasting blood glucose; Hct, hematocrit; TGs, triglycerides; TNFα, tumor necrosis factor alpha; CTL, control, chow-fed treated with saline; HFD, high-fat diet treated with saline; EPO, chow-fed treated with erythropoietin; HFD/EPO, HFD treated with EPO. Statistical analysis was conducted using a two-way ANOVA (diet×EPO). A Bonferroni correction was used for multiple comparisons and specific effects between groups. Animal characteristics at the start of EPO treatment: *P<0.05, CTL versus HFD. Animal characteristics at the end of the study: †P<0.05, ‡versus CTL and †versus HFD.
inflammatory cytokine TNFα were increased with HFD and reduced with EPO treatment (diet×EPO interaction: F(1,26)=5.54, P=0.026, with a significantly lower levels in HFD/EPO compared with HFD; Table 1), indicating that EPO may participate in the decrease in systemic inflammation. To determine the role of EPO in oxidative stress in our model of HFD-induced glucose intolerance, we quantified both F2-IsoPs and IsoFs levels in muscle. Measurement of F2-IsoPs or IsoFs is a state-of-the-art approach for assessing oxidative stress in vivo in diverse disorders including obesity (Montuschi et al. 2004). EPO treatment decreased oxidative stress in soleus muscle as indicated by decreased IsoFs levels in the HFD/EPO group compared with HFD (P<0.05; Fig. 5A). The same trend was observed for F2-IsoP without reaching significance (Fig. 5B). We also showed that EPO treatment rescued the HFD-induced reduction in HSP72 expression in soleus (F(1,26)=4.98, P=0.035, HFD lower than all other groups, P<0.05; Fig. 5C and D).

EPO does not directly target AKT phosphorylation in muscle cells

We aimed to determine if EPOR was expressed in muscle cells and whether it could mediate metabolic action in vivo and in vitro. While we could not detect EPOR in soleus or EDL muscles, we found that EPOR was expressed at substantial levels in L6 myoblasts and myotubes (Fig. 6A). We then demonstrated that EPO had no significant effect on glucose uptake or glycogen synthesis in L6 myotubes incubated with EPO alone or co-incubated with EPO and insulin (P>0.05; Fig. 6B and C). Given the previous results indicating that EPO activates the PI3K/AKT pathway in metabolic cells (Mikolás et al. 2012), we investigated the effect of EPO on AKT phosphorylation in differentiated rat muscle cell lines and human primary cells. We then first confirmed the existence of a synergy between EPO and insulin towards AKT phosphorylation using 3T3-L1 adipocytes treated with insulin and EPO (P>0.05; Fig. 7A and B). We then showed that EPO alone did not activate AKT in L6 myotubes acutely or chronically incubated with 10 U/ml EPO. When cells were stimulated with submaximal or maximal doses of insulin, EPO did not affect insulin-stimulated AKT phosphorylation (Fig. 7C and D). However, EPO did improve AKT phosphorylation in rat myotubes exposed to palmitate (Fig. 7E and F). We also showed that, in human primary myotubes, EPO did not activate AKT and did not affect the action of insulin on
P-AKT levels (Fig. 7G and H). Taken together, these results provide evidence that while EPOR is expressed in rat myotubes, EPO does not directly potentiate insulin action in muscle cells in vitro. However, EPO can rescue palmitate-induced inhibition of AKT activation, indicating that specific pathway activated by palmitate might have played a role. As EPOR was not detectable in soleus, our results also indicate that changes observed in vivo in soleus from EPO-treated animals resulted from an indirect action of EPO on this tissue.

Figure 3
Short-term EPO treatment does not alter body weight gain or fat tissue mass. (A) Energy intake and (B) body weight during the course of the study. (C) Retroperitoneal fat pad mass at the end of the study. Statistical significance: *P<0.05 between chow- or HFD-fed animals and **P<0.01 between groups. CTL, chow and saline; EPO, chow and EPO; HFD, high fat diet and saline; HFD/EPO, HFD and EPO; n=8 in each group.

Figure 4
Chronic EPO treatment partially restores insulin-stimulated AKT phosphorylation in soleus muscle. Quantification and western blotting images showing P-AKT/total AKT in nonstimulated (Ins–) or in insulin-stimulated (Ins+) conditions in (A and B) soleus and (C and D) EDL muscles. P-AKT/total AKT is expressed as fold of the value for the control nonstimulated condition. *P<0.05 indicates a statistically significant difference between conditions. CTL, control and saline; EPO, control and EPO; HFD, high-fat diet and saline; HFD/EPO, HFD and EPO; n=8 in each group.
We report that EPO treatment improves glucose tolerance in diet-induced-glucose-intolerant rats. We further show that EPO modulates AKT phosphorylation in soleus, and provide evidence that EPO does not directly target the muscle cell to trigger this effect. We observed that EPO treatment is associated with a reduced systemic inflammation, lowered muscle oxidative stress, and increased muscle HSP72 expression, indicating that the improvement in glucose metabolism induced by EPO results from the amelioration of systemic factors.

Both acute and chronic EPO treatment in vivo led to AKT activation in soleus muscle from lean animals. A single injection of a high dose of EPO induced transient phosphorylation of AKT without altering systemic levels of glucose, indicating that EPO-induced phosphorylation of AKT in skeletal muscle has a limited role in acute glucose control in animals with normal blood glucose levels. In contrast, results described in a previous report indicated that acute EPO injection in streptozotocin-induced diabetic rats with elevated blood glucose levels (approximately 22 mM) led to a small (6%) but significant decrease in blood glucose levels; however, the tissues involved have not been identified (Mikolás et al. 2012). Initial blood glucose levels may thus be critical in the acute effect of EPO, consistent with the study reporting that hyperglycemia is necessary for stimulation of EPO-induced glucose uptake in adipocytes (Mikolás et al. 2012). As we have also shown that rat soleus muscle did not express EPOR, AKT activation in our model is likely to be indirect via a still unknown mechanism (Wang et al. 2014). In human, acute i.v. injection of EPO was without effect on phosphorylation of AKT, AMP-activated protein kinase, or acetyl Co-A carboxylase, while insulin sensitivity and blood glucose levels were not affected (Christensen et al. 2012). The absence of AKT activation following acute EPO injection in that study may be due to the use of a much lower dose of EPO (200 and 400 IU/kg versus 1300 IU/kg in the current study), to the timing of the collection of muscle biopsies, to the fiber type composition of the muscle analyzed, or species-specific differences.

In our study, high-fat-fed animals presented increased body weight and retroperitoneal fat mass, elevated fasting

**Figure 5**

Chronic EPO reduces oxidative stress and restores HSP72 levels in soleus muscle. (A) Oxidative stress assessed by measurement of IsoFs and (B) F2-IsoP levels in soleus muscle. (C) Quantification of HSP72 protein level. (D) Representative western blot image. *P < 0.05 indicates a statistically significant difference between groups. CTL, control and saline; EPO, control and EPO; HFD, high-fat diet and saline; HFD/EPO, HFD and EPO; n = 8 in each group.

**Discussion**

We report that EPO treatment improves glucose tolerance in diet-induced-glucose-intolerant rats. We further show that EPO modulates AKT phosphorylation in soleus, and provide evidence that EPO does not directly target the muscle cell to trigger this effect. We observed that EPO treatment is associated with a reduced systemic inflammation, lowered muscle oxidative stress, and increased muscle HSP72 expression, indicating that the improvement in glucose metabolism induced by EPO results from the amelioration of systemic factors.

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**Figure 6**

While EPOR is expressed in muscle cells, EPO does not stimulate glucose uptake or glycogen synthesis in cultured myotubes. (A) EPOR expression in bone marrow, muscle (soleus and EDL), myoblasts, and myotubes. (B) Insulin-stimulated glycogen synthesis and (C) glucose uptake in L6 myotubes incubated with or without EPO. *P < 0.05 indicates a statistically significant difference between groups. NS, nonsignificant.
Figure 7
EPO does not activate AKT phosphorylation in cultured myotubes. (A) Quantifications and (B) representative western blot of P-AKT in response to insulin in 3T3L1 adipocytes under control or EPO-treated conditions for 48 h (n = 3). (C) Quantifications and (D) representative western blot of P-AKT/total AKT in response to insulin in L6 myotubes incubated under control or EPO-treated conditions for 5 min, 15 min, or 48 h (n = 4). (E) Quantification and (F) representative western blot image of P-AKT/total AKT in response to insulin in L6 myotubes incubated with EPO, palmitate, or EPO/palmitate. (G) Quantifications and (H) representative western blot image of P-AKT/total AKT in response to 0 or 1 nM insulin in human primary myotubes incubated with or without EPO (n = 3). *P < 0.05 between conditions as shown on figure. EPO5, EPO15, and EPO48: incubation with EPO for 5 min, 15 min, or 48 h respectively. Palm, incubation with palmitate; Palm/EPO, incubation with palmitate and EPO.

Blood glucose, and insulin levels as well as impaired glucose tolerance. It is noteworthy that while EPO treatment improved glucose control, it did not change body weight and retroperitoneal fat mass. This indicates that EPO ameliorates glucose metabolism independently from changes in body composition. Such effect can be explained via direct action on metabolic tissues, or indirectly through EPO anti-inflammatory effects (Li et al. 2006, Alnaeeli et al. 2014). Partial recovery of glucose tolerance was accompanied by recovery of baseline glucose levels in the HFD/EPO group while insulin levels were not yet affected. It is tempting to speculate that we observed early adaptations, and that longer EPO treatment or larger doses may be necessary to normalize insulin levels and to restore normal glucose metabolism (Katz et al. 2010). Our results also indicated that EPO treatment restored insulin-stimulated AKT activation in skeletal muscles in response to insulin in the HFD/EPO group in oxidative muscle (soleus); however, this did not lead to a statistically significant increase in P-AKT levels when compared with HFD due to increased baseline levels in this group. In contrast, EDL, a muscle characterized by glycolytic metabolism, did not show any response to EPO treatment. As EPOR mRNA was undetectable in soleus or EDL, we conclude that any effect of EPO treatment on muscle AKT phosphorylation is indirect and results from lower systemic inflammation, reduced muscle oxidative stress, or increased HSP72 expression, all known to positively influence insulin signaling. On the other hand, other metabolic tissues, such as liver (Meng et al. 2013) or adipose tissue (Mikolás et al. 2012, Wang et al. 2013), which have been shown to express EPOR and respond to EPO, can also participate in improved glucose control in our model.

Our results from cultured cells confirm that EPO can stimulate AKT activation in synergy with insulin in 3T3L1 adipocytes and demonstrate the lack of direct action of EPO on muscle cells. Indeed, we show that EPO did not affect AKT phosphorylation when rat L6 or human primary myotubes were incubated with EPO or co-incubated with both EPO and insulin, indicating that EPO did not directly target AKT in differentiated skeletal muscle cells. In agreement with the lack of effect of EPO on AKT activation in myotubes, we also demonstrate that EPO did not increase glucose uptake or glycogen synthesis. However, we showed that EPO had the potential to rescue AKT phosphorylation when cells were subjected to palmitate treatment, a model used to mimic in vitro the deleterious effect of HFD on insulin signaling. While the underlying mechanism cannot be determined from our results, these indicate that EPO signaling in myotubes may provide some protection to the insulin signaling cascade. Expression of the EPOR gene has been long thought to be restricted to the erythroid lineage but has since been identified in C2C12 myoblasts (Ogilvie et al. 2000, Launay et al. 2010, Jia et al. 2012) and myotubes...
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Conversely, overexpression or restoration of this protein improves insulin sensitivity (Chung et al. 2008, Gupta et al. 2009). The mechanisms involved in the reduced expression of HSP72 in skeletal muscle in T2D are still unclear, but seem to be driven by a systemic inflammatory state (Chung et al. 2008).

In conclusion, we show that a 2-week EPO treatment in rats fed a HFD can improve glucose tolerance independently of body weight and retroperitoneal fat content. We observed incomplete restoration of insulin-stimulated AKT activation in muscle in vivo; however, given the absence of EPOR expression in rat soleus and EDL muscles and the absence of AKT activation in myotubes exposed to EPO, we conclude that EPO does not target AKT directly in skeletal muscle. The observed reduced systemic TNFα level, and oxidative stress, as well as the restoration of HSP72 expression in soleus may act in synergy to provide protection against the diet-induced insulin resistance in metabolic tissues and contribute to the positive action of EPO on glucose homeostasis.

Declaration of interest
The authors declare that there is no conflict of interest that could be perceived as prejudicing the impartiality of the research reported.

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Author contribution statement
C C, R B, and D S designed the studies. C C, H A, E M, M M, A N M, and P R conducted the experiments. C C, R B, H A, J M, C B, T A M, and S T analyzed the results. C C, R B, and D S wrote the manuscript and all authors approved the final version of the manuscript.

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