Suboptimal nutrition during prenatal and early postnatal development is associated with increased risk for type 2 diabetes during adult life. A hallmark of such diabetes risk is altered body composition, including reduced lean mass and increased adiposity. Since stem cell number and activity are important determinants of muscle mass, modulation of perinatal nutrition could alter stem cell number/function, potentially mediating developmentally programmed reductions in muscle mass. Skeletal muscle precursors (SMP) were purified from muscle of mice subjected to prenatal undernutrition and/or early postnatal high-fat diet (HFD)—experimental models that are both associated with obesity and diabetes risk. SMP number was determined by flow cytometry, proliferative capacity measured in vitro, and regenerative capacity of these cells determined in vivo after muscle freeze injury. Prenatally undernourished (UN) mice showed significantly reduced SMP frequencies [Control (C) 4.8% ± 0.3% (% live cells) vs. UN 3.2% ± 0.4%, \( P = 0.015 \)] at 6 weeks; proliferative capacity was unaltered. Reduced SMP in UN was associated with 32% decrease in regeneration after injury ([C] 16% ± 3% of injured area vs. UN 11% ± 2%; \( P < 0.0001 \)). SMP frequency was also reduced in HFD-fed mice (chow 6.4% ± 0.6% vs. HFD 4.7% ± 0.4%, \( P = 0.03 \)), and associated with 44% decreased regeneration (chow 16% ± 2.7% vs. HFD 9% ± 2.2%; \( P < 0.0001 \)). Prenatal undernutrition was additive with postnatal HFD. Thus, both prenatal undernutrition and postnatal overnutrition reduce myogenic stem cell frequency and function, indicating that developmentally established differences in muscle-resident stem cell populations may provoke reductions in muscle mass and repair and contribute to diabetes risk.

**Introduction**

Nutritional or environmental stimuli acting during critical developmental windows can have a lasting impact on cellular structure/function and patterns of adult disease [1]. Indeed, a variety of prenatal nutritional stressors, including undernutrition, obesity, and placental insufficiency, may program metabolic adaptations that favor survival, but ultimately are detrimental to adult health. Moreover, prenatal nutritional stress is often followed by accelerated growth and fat accumulation during early childhood, which further increases risk for adult obesity, type 2 diabetes, and cardiovascular disease [2,3].

How suboptimal perinatal environments lead to adult disease has not been fully elucidated at a molecular level. However, a common physiological phenotype associated with developmentally mediated disease risk is altered body composition, with reduced lean body mass and increased fat mass observed in both humans and animal models [4–6].

Reductions in skeletal muscle mass may be particularly important as muscle is a key determinant of systemic metabolism and insulin sensitivity [7]. Moreover, humans with low birth weight, a marker of prenatal developmental history, demonstrate alterations in muscle fiber size and type and decreased oxidative capacity [8]. Similarly, experimental nutritional restriction during pregnancy alters offspring muscle mass and composition in sheep, in parallel with increased lipid accumulation and reduced oxidative capacity in muscle [9]. Such reductions in muscle mass and/or function may thus disrupt systemic metabolism and contribute to adult disease risk.

One determinant of skeletal muscle mass, as well as its appropriate maintenance with aging, is a quantitatively and qualitatively normal muscle stem cell population. Muscle stem cells, a subset of muscle satellite cells, are mononuclear cells residing between the plasma membrane and basal lamina of mature myofibers [10]; these cells are defined by their capacity to both self-renew and differentiate to generate...
mature, multinucleated muscle fibers, and are primary mediators of postnatal muscle regeneration [11]. Mouse models with impaired muscle stem cell function and number, such as Pax7-null mice, have significantly decreased muscle mass [12,13] and poor regenerative function after injury [14]. Moreover, satellite cells appear to be particularly sensitive to nutritional availability during periods of high growth. In poultry, early postnatal starvation yields reduced satellite cell proliferation, increased apoptosis, and sustained reductions in muscle mass despite normalization of food intake [15–17]. Reduced satellite cell number has also been observed in malnourished human children [18].

Given these interesting links between nutritional exposure during development, reduced muscle mass, and adult disease risk, an important question is whether muscle stem cell activity or regenerative capacity is altered as a function of perinatal nutrition and could contribute to adult disease in mammals.

Materials and Methods

Animal studies

Mice were housed in an Office of Laboratory Animal Welfare (OLAW)-approved facility, with controlled temperature, humidity, and light–dark cycle (07:00–19:00). Protocols were approved by the Joslin Diabetes Center Institutional Animal Use and Care Committee; “Principles of Laboratory Animal Care” (http://grants1.nih.gov/grants/olaw/references/phs000010.html) were followed.

Mice exposed to maternal undernutrition were generated as previously described [6,19–21] (Supplementary Fig. S1; Supplementary Data are available online at www.liebertonline.com/scd). In brief, 6–8-week-old ICR females were housed with ICR males, and pregnancies dated by vaginal plug (day 0.5). Pregnant dams were allowed ad lib access to regular chow (Purina 9F, 21% of calories from protein, 21% from fat, and 58% from carbohydrate) until day 12.5, when dams were randomly assigned to ad lib feeding (C) or 50% food restriction (undernutrition, UN) calculated based on food intake of control dams for the specific day of pregnancy. At birth, litters were equalized to 8 pups. During suckling, dams were provided regular chow ad libitum. Both C and UN pups were weaned on day 21 to either Purina 9F chow or high-fat diet (HFD) containing 60% of calories from fat (Research Diets, Inc., Open Source Diets, D12492). For additional experiments, C57/BL6 mice with normal prenatal nutritional exposure were obtained from Jackson Laboratories at 3 weeks of age, and fed ad libitum with either chow or HFD for 3 weeks.

Blood glucose was measured between 9 and 11 AM (fed) or after a 16-h fast in tail vein samples from 8 week old mice (Ascensia Elite; Bayer). Insulin was measured using rat insulin enzyme-linked immunosorbent assay with mouse standards (Crystal Chem) (Supplementary Table S1).

Body composition was analyzed by dual energy X-ray absorptiometry (Hologic).

Stem cell isolation and measurement of myogenic colony formation

Muscle stem cells were isolated as previously described [22]. In brief, after anesthesia with pentobarbital (150 mg/kg), hindlimb muscles were dissected and digested with 0.2% (w/v) collagenase type II in Dulbecco’s modified Eagle’s medium (Gibco BRL) for 90 min at 37°C, and individual muscle fibers were dissociated by repeat pipetting. Isolated myofibers were then digested in 10 volumes of F10 medium containing 0.05 U/mL dispase (Gibco) and 0.012% collagenase type II for 30 min at 37°C with agitation to isolate myofiber-associated cells. Skeletal muscle precursor (SMP) cells were isolated by fluorescence activated cell sorting (FACS)-based purification using the markers Sca1-, CD45-, Mac1-, CXCR4+ and P1integrin+ as described [23]; a representative set of FACS plots is provided in Supplementary Fig. S2. To set detection thresholds for FACS sort and analysis, aliquots of interstitial muscle cells were stained with all antibodies minus one; these cells were analyzed by flow cytometry to set a gating threshold specific for each antibody. To assess myogenic colony formation, FACS-purified SMP cells were plated clonally in 96-well plates precoated with laminin (10 mg/mL; Gibco) and collagen (1 mg/mL; Sigma) and cultured in F10 at 37°C; the presence or absence of myogenic colonies was evaluated by light microscopy 5 days after plating, and expressed as percentage of wells containing colonies relative to total number of wells.

Muscle injury and regeneration

After anesthesia with Avertin (tribromoethanol; Sigma, 800 μL of 1:80 dilution intraperitoneally), dry ice was placed directly on the belly of the tibialis anterior muscle for 5 s. Three or 10 days after injury, animals were euthanized (pentobarbital), and injured muscles dissected. Muscles were fixed in 4% paraformaldehyde for 1 h and embedded in paraffin. Multiple sections were obtained at 60 μm increments, and slides stained with hematoxylin and eosin. Areas of muscle regeneration were assessed using ImageJ software. Specifically, nonoverlapping photos of each muscle were taken at 100× magnification, and a grid composed of small open dots was overlaid. Two blinded observers scored each dot in the injured area and assigned one of the following categories: regenerating fiber (centrally located nucleus), inflammatory cell (nucleus without underlying fiber), injured (fiber without central nucleus), or other (blood vessel/empty space).

Muscle fiber typing

After euthanasia, leg muscles were dissected, weighed, and frozen. Quadriceps sections were stained with succinyl dehydrogenase (Department of Pathology, Brigham and Women’s Hospital), and 7 photos were taken at 400×. A blinded observer scored each fiber as type I (dark staining), type 2a (medium staining), or type 2b (little to no staining). Incomplete fibers at the border of photos were excluded.

Assessment of systemic and local inflammation

Systemic inflammation was assessed in plasma samples obtained via intracardiac puncture at 3 and 10 days after muscle injury. Tumor necrosis factor alpha (TNF-α), interleukin-6 (IL-6), monocyte chemotactic protein 1 (MCP-1), and plasminogen activator inhibitor-1 (PAI-1) were measured using multiplex enzyme-linked immunosorbent assay (Millipore). Local muscle inflammation (quadriceps) was assessed by both immunohistochemistry for the macrophage marker Mac2 (Cedarlane Labs) and quantitative real-time (RT)–polymerase
chain reaction. In brief, RNA was extracted (Trizol; Invitrogen), cDNA generated from 1 μg RNA (Applied Biosystems), and RT–polymerase chain reaction analysis performed using SYBRGreen detection, with GAPDH as endogenous control (Applied Biosystems). Primer sequences are available in Supplementary Table S2.

**Statistical analyses**

Data are presented as mean ± standard error of the mean. Multigroup comparisons of body or muscle weight, hormonal parameters, muscle area, and SMP frequencies were assessed by analysis of variance (Statview), with post hoc pairwise comparisons assessed by Fisher’s protected least significant difference. Multigroup comparisons of regeneration and fiber type were assessed by χ² analysis. P < 0.05 was considered significant.

**Results**

**Prenatal undernutrition alters body composition and reduces muscle mass**

To determine the effect of antenatal or early postnatal nutrition on myogenic stem cell number and function, mice exposed to maternal undernutrition in utero were studied [6] (Supplementary Fig. S1). In this model, maternal food restriction during the final week of gestation results in a 14%–20% reduction in birth weight in UN pups as compared with controls (C).

Despite reduced weight at birth, UN mice have similar weights to C mice by 6–7 weeks of age, reflecting catch-up growth in UN mice (C 38.0 ± 0.7 g vs. UN 35.8 ± 1 g; P = 0.07). However, body composition analysis (dual energy X-ray absorptiometry) revealed that UN mice had persistent reductions in lean body mass as compared with C (C 30.4 ± 0.7 g vs. UN 28.4 ± 0.2 g; P = 0.03) (Fig. 1A). Similarly, the combined weight of the quadriceps, gastrocnemius, soleus, tibialis anterior, and extensor digitalis longus muscles was significantly decreased in UN mice (C 0.88 ± 0.02 g vs. UN 0.74 ± 0.03 g; P = 0.01) (Fig. 1B).

**UN mice have altered muscle fiber type**

Alterations in fiber number or type have been previously described in low birth weight (LBW) humans and in humans at risk for diabetes based on obesity or diabetes family history [8,24,25]. Mice with UN exposure had no differences in fiber number per cross section of quadriceps muscle at either 6–7 weeks or 9 months of age. However, at 6–7 weeks, UN mice exhibited a 16% decrease in type I (slow oxidative) fibers (P < 0.001), a 20% increase in type IIA (fast oxidative) fibers (P < 0.001), and 6% increase in type IIB (fast glycolytic) fibers as compared with C (P < 0.05) (Supplementary Table S3). At 9 months of age, type I fiber number did not differ, but UN mice had a 34% increase in type IIA and a 23% decrease in type IIB fibers (P < 0.05 for both). Fiber number and type were also assessed in 9-month-old control and UN mice fed a HFD from weaning (C-HFD and UN-HFD, respectively). Neither fiber number nor fiber type distribution differed between C-HFD and UN-HFD muscle (Supplementary Table S3).

**Mice exposed to maternal UN have decreased muscle stem cell numbers with normal ex vivo myogenic capacity**

Muscle stem cells are crucial for muscle development, growth, and repair after injury [26]. To determine whether muscle stem cell number and/or function are altered in this experimental model, SMP cells were isolated from the myofiber-associated cell compartment of the hindlimbs of 6-week-old mice using FACS-based purification [22,23] based on the following markers: CD45−, Sca1+, Mac1+, CXCR4+, and β1 integrin+. Cells isolated using this protocol can self-renew, produce mature myotubes [23], and functionally engraft to enhance muscle function in mdx mice, a mouse model of Duchenne muscular dystrophy [22]. In UN mice (which show decreased muscle mass), SMP cell frequency was reduced by ~33% (C 4.8% ± 0.3% vs. UN 3.2% ± 0.4%, P = 0.02, percentage of live cells isolated) (Fig. 2A, B).

To assess whether reduced number of myogenic precursor cells in UN mice was accompanied by alterations in their ability to enter a myogenic program, SMPs were plated at 1 cell per well, and myogenic colonies were evaluated 5 days later by microscopic inspection. The number of myogenic colonies arising from the clonally plated SMPs did not differ between groups, suggesting that, on a per cell basis, muscle stem cells from UN mice retain their normal capacity to survive and initiate myogenesis, as assessed in a surrogate ex vivo culture environment.

**In vivo regeneration after muscle injury is decreased in low-birth-weight mice**

Because muscle stem cells play an important role in muscle repair, assessment of regeneration after injury is used as a measure of stem cell function [27]. To determine the functional consequence of reduced SMP numbers in UN mice, muscle histology and regeneration were assessed in 9-month-old UN or C mice at 3 or 10 days after in vivo freeze injury (representative sections, Fig. 3A). Consistent with differences in muscle weight observed as early as 6 weeks of life, overall muscle size was decreased in injured UN mice at...
9 months of age, as determined by cross-sectional area of the tibialis anterior (day 3: C 10.1 ± 0.7 × 10⁶ vs. UN 6.1 ± 0.7 × 10⁶ pixels, P = 0.004). The area of muscle injury was similar between experimental groups on both days 3 and 10 (Supplementary Fig. S3). However, at 3 days after injury, there was a 32% reduction in the number of regenerating fibers (Fig. 3B, left) in UN mice (C 16% ± 2.6% vs. UN 11% ± 2.2% of injured area; P < 0.0001), and an 18% reduction at 10 days after injury (C: 74% ± 3% vs. UN 61% ± 2.2%; P < 0.001) (Fig. 3B, right).

Since both inadequate and excessive inflammatory responses can hinder muscle regeneration [28,29], inflammation was assessed at both the local and systemic level in 9-month-old mice 3 days after cold injury. There were no significant differences in (1) serum levels of inflammatory markers (MCP1, TNF-α, IL-6, or PAI-1), (2) quadriceps muscle inflammatory gene expression (IL-1β, F4/80, MCP1, TNF-α, IL-6, and PAI-1), or (3) inflammatory cell infiltration in injured muscle, as indicated by immunohistochemical analysis of Mac2 (Supplementary Fig. S4A–C).

Since increased adiposity is a dominant phenotype in UN mice, observed as early as 3 weeks of age [21], it is possible that early life obesity per se could contribute to reduced muscle stem cell frequency. SMP frequency was measured in hindlimb muscle from mice with normal antenatal nutrition fed a HFD from ages 3 to 6 weeks postnatally. HFD-fed mice were heavier (chow 23.3 ± 0.5 g vs. HFD 25.4 ± 0.8 g, P = 0.001), as expected, but had reduced muscle mass (hindlimb muscle weight: chow 1.29 ± 0.04 g, HFD 1.21 ± 0.03 g, P = 0.008). Interestingly, SMP frequencies were reduced by 27% in HFD-fed mice (chow 6.4% ± 0.6% vs. HFD 4.7% ± 0.4%, expressed as percentage of live cells, P = 0.03, Fig. 4).

The impact of high-fat feeding in both C and UN mice on stem cell-initiated myogenesis was also assessed using in vivo regeneration assays. Muscle regeneration 3 days after cold injury was reduced by 42% in control mice fed a HFD
Strikingly, regeneration was reduced to an even greater extent in HFD-fed UN mice (64% reduction, UN 11% vs. UN-HFD 4%, $P<0.0001$) (Fig. 5).

**Discussion**

Both prenatal undernutrition and early postnatal overnutrition are associated with reduced muscle stem cell number and reduced muscle regenerative capacity. These defects, while sustained during early life, persist into adulthood and may contribute to developmentally mediated reductions in muscle mass and altered body composition. Given that muscle mass is an important mediator of insulin-stimulated glucose uptake and systemic metabolism, reductions in stem cell number may contribute to associations between early life nutrition and developmental risk for adult disease.

Muscle growth and maintenance depend on adequate stem cell availability and repair. Muscle stem cells are normally quiescent, but upon muscle injury, can be activated to proliferate, self-renew, and differentiate into myoblasts. These myoblasts subsequently fuse with other myoblasts, as well as damaged muscle fibers, to form new functional muscle. This process may be particularly vulnerable to postnatal nutrition, as starvation reduces satellite cell proliferation and increases apoptosis in poultry [15–17] and satellite cell number is reduced in malnourished human children [18]. Reduced function of genes critical for postnatal satellite cell survival, such as Pax7, can cause accelerated muscle wasting soon after birth [14]. Thus, the reductions in stem cell number observed in UN mice may contribute directly to the decreased muscle mass observed during postnatal growth and in adulthood—a key phenotype in both UN mice and LBW humans [4,5]. Indeed, these experimental data confirm the hypothesis suggested by Cianfarani, who proposed that stem cells are critical for normal tissue maintenance and function and that prenatal malnutrition would decrease these populations leading to early tissue malfunction and contribute to LBW-associated disease phenotypes [30].

Alterations in stem cell number or function can also affect the ability of an organism to repair [14]. For example, mice with mutations in the dystrophin gene (mdx mice) have poor integrity of muscle fibers, increased vulnerability to mechanical stress, and thus need for perpetual repair, but also have reduced frequency of SMPs [22]. Together with the early observations of Schultz and colleagues that repeated stresses reduce the proliferative capacity of satellite cells [31], it is also possible that reductions in regeneration in UN mice could reflect not only early life reductions in SMP number, but also subclinical increases in muscle damage accumulated during life, which further magnify age-related reductions in repair capacity.

Tissue stem cell number and function are highly dependent on features of the systemic and local microenvironment (niche), as demonstrated for aging-related dysfunction. For example, in muscle, repair responses may convert to favor fibrogenic, rather than myogenic, processes with age [32]. Regeneration capacity can be restored in aging mice by exposure to the circulation of young mice, in part via reactivation of Notch signaling [27,33]. In accord with this concept, the present data suggest that alterations in the intrauterine or early postnatal nutritional/metabolic environ-
ment also affect muscle regenerative function. Since functional impairment was detected in vivo, reflected by reduced regeneration after muscle injury, but not during ex vivo myogenic colony formation assays, it is likely that reduced availability of myogenic stem cells in UN mice contributes to this process.

Since undernutrition in mice is associated with early onset adiposity and progressive glucose intolerance with aging, it is possible that obesity per se or other features of a diabetogenic microenvironment might contribute to reductions in stem cell frequency or function. Interestingly, early life obesity (produced by high-fat feeding) was associated with a 27% reduction in SMP frequency and reduced regeneration after muscle injury. Moreover, the effects of HFD to reduce regeneration after muscle injury were additive with prenatal undernutrition. Thus, an adverse prenatal metabolic environment, early life onset of nutritional obesity (or both), and chronic obesity may all be detrimental for stem cell activity and repair.

While the specific mechanisms mediating the effects of both the prenatal and postnatal nutrient environment on stem cell number and/or function remain unclear at this time, stem cell-independent mechanisms may also contribute to our findings of decreased regeneration in UN and/or HFD-fed mice, including the size of the injury, extent of inflammation, and other aspects of the systemic or local tissue milieu. While larger injuries, whether in absolute size or as a percent of the muscle, could slow the regenerative process (fewer satellite cells to repair the injury), there were no differences in the area of injury (percentage of cross-sectional area) in this model. Similarly, either reduced or excessive inflammation could also impair muscle growth, regeneration, and injury responses [28,34–36]. However, systemic or local inflammation was not altered in UN mice. While elevated glucose levels may alter differentiation of muscle stem cells [37], circulating glucose levels are consistently normal in our models at the age when stem cell number was assessed [6]. Whether nutritional or obesity-related alterations in amino acids, other metabolites, or nutritionally responsive growth factors critical for satellite cell development (e.g., insulin like growth factor 1 [IGF1]) could also contribute is an important question for future study [16,38]. Additional developmental signals or components of the systemic or tissue microenvironment, for example, stem cell niche [39,40], or growth factors produced locally by muscle fibers [16,41,42] could also contribute to reductions in stem cell frequency, function, and/or differences in myogenic versus adipogenic lineage development during regeneration [43], in the setting of UN exposure and obesity.

In summary, our studies demonstrate that low birth weight induced by prenatal undernutrition is associated with decreased frequency and in vivo functionality of muscle stem cells, as assessed by regeneration after injury, reduced muscle mass, and altered fiber type. Further, a HFD during early life is also associated with reduced stem cell number and decreased regenerative capacity, in both normal- and low-birth-weight mice. Thus, decreased number of stem cells and associated changes in regenerative capacity can result from adverse metabolic environments during both prenatal and early postnatal life. Such reductions in muscle stem cell number and function may contribute to alterations in muscle mass and body composition associated with developmentally mediated risk for adult disease.

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Author Disclosure Statement

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References


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