Adaptive Modifications of Muscle Phenotype in High-Altitude Deer Mice Are Associated with Evolved Changes in Gene Regulation

Graham R. Scott,*1 Todd S. Elogio,1 Mikaela A. Lui,1 Jay F. Storz,2 and Zachary A. Cheviron*,3

1Department of Biology, McMaster University, Hamilton, ON, Canada
2School of Biological Sciences, University of Nebraska, Lincoln
3Department of Animal Biology, University of Illinois, Urbana-Champaign

*Corresponding author: E-mail: scottg2@mcmaster.ca; cheviron@illinois.edu.

Abstract

At high-altitude, small mammals are faced with the energetic challenge of sustaining thermogenesis and aerobic exercise in spite of the reduced O2 availability. Under conditions of hypoxic cold stress, metabolic demands of shivering thermogenesis and locomotion may require enhancements in the oxidative capacity and O2 diffusion capacity of skeletal muscle to compensate for the diminished tissue O2 supply. We used common-garden experiments involving highland and lowland deer mice (Peromyscus maniculatus) to investigate the transcriptional underpinnings of genetically based population differences and plasticity in muscle phenotype. We tested highland and lowland mice that were sampled in their native environments as well as lab-raised F1 progeny of wild-caught mice. Experiments revealed that highland natives had consistently greater oxidative fiber density and capillarity in the gastrocnemius muscle. RNA sequencing analyses revealed population differences in transcript abundance for 68 genes that clustered into two discrete transcriptional modules, and a large suite of transcripts (589 genes) with plastic expression patterns that clustered into five modules. The expression of two transcriptional modules was correlated with the oxidative phenotype and capillarity of the muscle, and these phenotype-associated modules were enriched for genes involved in energy metabolism, muscle plasticity, vascular development, and cell stress response. Although most of the individual transcripts that were differentially expressed between populations were negatively correlated with muscle phenotype, several genes involved in energy metabolism (e.g., Ckmt1, Ehhadh, Acaa1a) and angiogenesis (Notch4) were more highly expressed in highlanders, and the regulators of mitochondrial biogenesis, PGC-1α (Ppargc1a) and mitochondrial transcription factor A (Tfam), were positively correlated with muscle oxidative phenotype. These results suggest that evolved population differences in the oxidative capacity and capillarity of skeletal muscle involved expression changes in a small suite of coregulated genes.

Key words: capillarity, hypoxia adaptation, muscle fiber type, oxygen transport, physiological genomics, RNA-seq.

Introduction

High-altitude environments provide fertile ground for examining the integrative physiological and genomic mechanisms of adaptation. Patterns of animal diversity change substantially across elevational gradients (Sánchez-Cordero 2001; McGuire et al. 2014), and a high degree of species turnover occurs in part from adaptation to local environments (Jankowski et al. 2012). The concurrent declines in temperature and oxygen tension with elevation are a particular challenge to small highland mammals, which must sustain high rates of aerobic metabolism to support thermogenesis and locomotion in spite of a diminished oxygen supply (Hayes 1989). Both genotypic specialization and phenotypic plasticity in the physiological systems that mediate oxygen transport and utilization could be important for meeting this challenge in highland natives, but we are just beginning to understand the mechanisms involved (Storz, Scott, et al. 2010; Scott 2011; Cheviron and Brumfield 2012).

Evolved changes in skeletal-muscle phenotype have occurred in several highland vertebrate taxa. The locomotory muscle of high-altitude birds that are sampled in their native environment (León-Velarde et al. 1993; Hepple et al. 1998; Mathieu-Costello et al. 1998) or raised in captivity at sea level (Scott, Egginton, et al. 2009; Scott, Richards, et al. 2009) is typically highly capillarized and more oxidative than that of lowland birds. Highland mammals also exhibit increased oxidative capacity in the locomotory, respiratory, and/or cardiac muscles (Sheafor 2003; Cheviron et al. 2012, 2014; Schippers et al. 2012). These derived trait changes should increase the capacities for oxygen diffusion and aerobic metabolism in the muscle, and may therefore enhance fitness-related physiological performance under hypoxia.

Changes in gene regulation have been shown to contribute to evolved differences in locomotory muscle phenotypes that are related to organismal performance. For example, mice selected for high levels of voluntary wheel running exhibit...
increased aerobic exercise performance and possess more oxidative and capillarized gastrocnemius muscle (Audet et al. 2011; Templeman et al. 2012), which is associated with differential expression of many genes involved in muscle plasticity and calcium signaling (Burniston et al. 2013). Similarly, rats selected for high running endurance exhibit upregulation of several genes involved in lipid metabolism in comparison to low capacity runners (Bye et al. 2008). In high-altitude populations of deer mice, an enhanced thermogenic capacity in hypoxia is associated with higher activities of oxidative enzymes as well as both constitutive and plastic changes in gene expression in locomotory muscle compared with lowland deer mice (Cheviron et al. 2012, 2014). Although changes in capillarity and fiber composition of skeletal muscle have evolved in several highland taxa (León-Velarde et al. 1993; Scott, Egginton, et al. 2009), the transcriptomic underpinnings of these convergent changes in phenotype are not known.

The deer mouse (*Peromyscus maniculatus*) has the broadest altitudinal distribution of any North American mammal, stretching from below sea level in Death Valley, California to over 4,300 m above sea level in numerous mountain ranges (Hock 1964). Population genetic studies of deer mice in western North America have demonstrated that highland mice are often genetically distinct from lowland conspecifics, and estimates of gene flow between highland and lowland populations are very low across the interface between the western Great Plains and the Front Range of the Southern Rocky Mountains (<0.01 migrants per generation) (Storz et al. 2012; Natarajan et al. 2015). Survivorship studies of high-altitude deer mice have documented strong directional selection for increased thermogenic capacity under hypoxia (Hayes and O’Connor 1999). Such selection is likely to have influenced the metabolic capacities of skeletal muscle because shivering thermogenesis requires high levels of muscle activity (Oufara et al. 1987; Van Sant and Hammond 2008).

The objective of this study was to determine whether the evolved increases in aerobic capacity and oxidative enzyme activities in the skeletal muscle of highland deer mice (Cheviron et al. 2012, 2014) are associated with a more capillarized and oxidative muscle phenotype. We then aimed to determine the relationship between various phenotypic traits and gene expression profiles in the skeletal muscle using whole transcriptome shotgun sequencing (RNA sequencing [RNA-seq]) in order to elucidate the regulatory changes that underlie adaptive modifications of muscle physiology in high-altitude mice. Because induced changes in muscle phenotype may occur at different ontogenetic stages, we used an experimental design that allowed us to account for the combined effects of developmental plasticity and physiological plasticity during adulthood. We made comparisons between highland (Mt. Evans CO, 4,350 m) and lowland (Lincoln NE, 430 m) mice that were sampled at their native elevations, and we also compared F1 progeny of wild-caught mice that were born and reared under common garden conditions at low elevation.

## Results

### Muscle Phenotype Differs between Highlanders and Lowlanders

High-altitude mice exhibited an enhanced oxidative phenotype of the locomotory (gastrocnemius) muscle (figs. 1 and 2). This result was reflected in significant main effects of population altitude on several oxidative phenotypes, including the areal and numerical densities of oxidative fibers. In pairwise comparisons between populations, highlanders exhibited greater areal and numerical densities of oxidative fibers (both slow oxidative fiber density as well as total oxidative fiber density) when mice were sampled in their native environment, and these differences were mirrored by nonsignificant trends in the F1 progeny of wild-caught mice that were reared in the common-garden lab environment. The total proportion of oxidative fibers in the muscle was greater in highland deer mice by approximately 14–25% as a proportion of the total transverse area of the muscle and it was greater by approximately 5–14% as a proportion of total fiber number (figs. 1 and 2). The main cause of these differences was a 23–83% higher abundance by area of slow oxidative (type I) muscle fibers (figs. 1 and 2) with no change in the abundance of fast oxidative (type Ila) muscle fibers (table 1). The population differences in areal density were generally greater than the differences in numerical density because slow oxidative fibers were larger in highland deer mice. There were no differences in the size of fast oxidative and fast glycolytic fibers between highland and lowland mice (table 1).

Highland mice also had an increased capillarity in the gastrocnemius muscle in comparisons involving both native and captive-reared individuals. Many indices of capillarity were higher in highland mice compared with lowland mice, including capillary surface density (~23–37% higher), capillary density (~10–21% higher), the ratio of capillary surface to muscle fiber surface (~19–36% higher), and the capillary to fiber ratio (~8–17% higher) (figs. 1 and 3). There was also a noticeably higher vessel tortuosity in the highland mice, as reflected by the greater discrepancy in capillary surface density than in capillary density and the qualitative difference in the pattern of capillary staining (fig. 3). The overall difference in capillarity between highland and lowland mice was greater than the difference between native and F1 mice with the same population-of-origin, indicating that the observed population difference is not attributable to phenotypic plasticity.

There was a positive correlation between capillary surface density and the areal density of oxidative fibers (fig. 4A). Thus, the increase in capillarity in highland deer mice could have resulted solely from the greater oxygen demands imposed by the increase in oxidative fibers in highland mice, and not to provide an additional improvement of oxygen diffusion capacity under hypoxia. To explore this possibility, we used the residuals from the regression of capillary surface density against the areal density of oxidative fibers to test for differences in capillarity while controlling for the difference in fiber composition between highland and lowland deer mice. Residuals from this regression were significantly greater in
highlanders than in lowlanders (fig. 4B), suggesting that the enhanced capillarity of highland mice should increase oxygen diffusion capacity in hypoxia.

**Population Differences and Plasticity in the Muscle Transcriptome**

We used a general linear model to identify genes that exhibited persistent expression differences between highland and lowland populations across rearing environments and those that exhibited significant plasticity (as revealed by within-population comparisons between mice sampled in their native environment vs. F1 progeny of wild-caught mice). In total, 657 genes (5.4% of measured transcripts) exhibited significant expression differences between populations and/or between rearing environments (native vs. lab F1). Of these variable transcripts, the vast majority exhibited significant plasticity in expression as indicated by differences between rearing environments, without a significant population effect (89.7%—589 genes). A smaller number of genes were differentially expressed between populations, without significant effects of rearing environment (10.3%—68 genes). Twenty-nine genes (4.4% of variable transcripts) exhibited significant expression differences between populations and between rearing environments. We focused on this subset of 657 variable transcripts to investigate the transcriptomic basis of population differences in muscle phenotypes (supplementary tables S1 and S2, Supplementary Material online).

We calculated correlation coefficients for all pairwise comparisons of transcript abundance and we then used modulated modularity clustering (MMC) (Stone and Ayroles 2009) to identify transcriptional modules of coregulated genes (Rockman 2008; Ayroles et al. 2009). This analysis revealed a high degree of correlational structure among differentially expressed transcripts (fig. 5). The 68 genes with significant population effects clustered into two modules, the largest of which (module P2) contained 50 genes (fig. 5A). We recovered a similar pattern for the genes with significant differences in expression between rearing environments (i.e., native vs. lab F1) and also consistent with the MMC analysis. However, we did find a significant number of genes (10 genes) that exhibited consistent expression differences between populations in both native and lab F1 environments.
The 589 genes with significant effects of rearing environment clustered into a total of five modules, the largest of which (module T5) contained nearly 38% of the environmentally sensitive transcripts (224 genes).

Relationships between Muscle Phenotype and Transcriptomic Variation

Population-specific differences in gene expression were significantly associated with the observed differences in muscle phenotype. We used a combination of principal components analysis (PCA) and regression analysis to determine which transcriptional modules were most strongly associated with muscle phenotypes. Because genes within transcriptional modules are, by definition, highly correlated, PCA can be used to summarize expression patterns of particular transcriptional modules (Cheviron et al. 2014; Stager et al. 2015). For each of the transcriptional modules, the first principal component (PC1) accounted for 32.4–99.7% of the variance in gene expression among individuals.

Table 1. Fiber Types in the Gastrocnemius Muscle of Deer Mice.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Tr.</th>
<th>Lowlanders</th>
<th>Highlanders</th>
<th>F</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>N_{ox,m}</td>
<td>Native</td>
<td>0.299 ± 0.024</td>
<td>0.288 ± 0.041</td>
<td>Pop 0.042</td>
<td>0.838</td>
</tr>
<tr>
<td></td>
<td>Lab F1</td>
<td>0.316 ± 0.027</td>
<td>0.340 ± 0.029</td>
<td>RE 1.308</td>
<td>0.260</td>
</tr>
<tr>
<td>N_{ox,m}</td>
<td>Native</td>
<td>0.353 ± 0.030</td>
<td>0.328 ± 0.044</td>
<td>Pop 0.556</td>
<td>0.461</td>
</tr>
<tr>
<td></td>
<td>Lab F1</td>
<td>0.383 ± 0.023</td>
<td>0.359 ± 0.032</td>
<td>RE 0.885</td>
<td>0.353</td>
</tr>
<tr>
<td>Type I area</td>
<td>Native</td>
<td>1,082 ± 114</td>
<td>1,503 ± 127*</td>
<td>Pop 5.092</td>
<td>0.030</td>
</tr>
<tr>
<td></td>
<td>Lab F1</td>
<td>1,329 ± 65</td>
<td>1,436 ± 136</td>
<td>RE 0.593</td>
<td>0.446</td>
</tr>
<tr>
<td>Type IIa area</td>
<td>Native</td>
<td>1,448 ± 114</td>
<td>1,574 ± 134</td>
<td>Pop 0.948</td>
<td>0.337</td>
</tr>
<tr>
<td></td>
<td>Lab F1</td>
<td>1,491 ± 115</td>
<td>1,598 ± 110</td>
<td>RE 0.077</td>
<td>0.783</td>
</tr>
<tr>
<td>Type IIb area</td>
<td>Native</td>
<td>2,277 ± 178</td>
<td>2,404 ± 130</td>
<td>Pop 0.152</td>
<td>0.699</td>
</tr>
<tr>
<td></td>
<td>Lab F1</td>
<td>2,561 ± 109</td>
<td>2,313 ± 161</td>
<td>RE 0.380</td>
<td>0.541</td>
</tr>
</tbody>
</table>

Note.—Transverse area of each fiber type (slow oxidative, type I; fast oxidative, type IIa; fast glycolytic, type IIb) is reported in μm². A_{ox,m}, areal density of fast oxidative fibers; N_{ox,m}, numerical density of fast oxidative fibers; one degree of freedom for each main effect variable (Pop, population altitude; RE, rearing environment) and 36 for the residual.

*Significant pairwise difference between highlanders and lowlanders within an environment.
Therefore, we used PC1 scores as an index of overall module expression to identify transcriptional modules that were significantly associated with muscle phenotypes. Of the two modules composed of genes with significant population effects, only one (P2) was significantly associated with any of the measured phenotypic traits. Module P2 expression scores were negatively associated with the total areal ($R^2 = 0.51$, $P = 0.00026$) and numerical ($R^2 = 0.29$, $P = 0.00011$) densities of oxidative fibers, capillary density ($R^2 = 0.23$, $P = 0.00223$), capillary surface density ($R^2 = 0.46$, $P = 0.00009$), and the ratio of capillary surface area to fiber surface area ($R^2 = 0.51$, $P = 0.00004$) (table 2). The relationship between module P2 expression scores and capillary to fiber ratio were marginally significant ($R^2 = 0.16$, $P = 0.0513$). Correspondingly, highland mice exhibited significantly lower module P2 expression scores than lowland mice, regardless of rearing environment (fig. 6). PC2 scores for module P1 were associated with several phenotypic traits, but again, this axis accounted for little of the explained variance in gene expression within modules (3.3% and 5.6%, respectively) (supplementary table S3, Supplementary Material online).

Enrichment analyses revealed that the transcriptional modules associated with muscle phenotypes contain a diverse assemblage of genes with a variety of molecular functions, many of which are relevant for metabolism, muscle fiber-type differentiation, and angiogenesis (supplementary table S4, Supplementary Material online). Module P2—the larger module comprising genes that were differentially expressed between populations regardless of rearing environment—was enriched (at a $P$ value corrected for the false discovery rate [FDR], $q < 0.10$) for genes annotated with four gene ontology (GO) terms associated with metabolic processes, “secondary metabolic process” ($q = 0.034$), “oxidation—reduction process” ($q = 0.064$), “toxin metabolic process” ($q = 0.016$), and “organic acid metabolic process” ($q = 0.094$). Module T5 was significantly enriched for genes annotated with several GO terms related to muscle...
corresponding with the observed differences in muscle phenotype (fig. 7 and supplementary table S1, Supplementary Material online). Highland deer mice exhibited higher expression of several genes that are associated with oxidative energy metabolism, including enoyl-CoA hydratase (Ehhadh) and acetyl-coenzyme A acyltransferase (Acaa1a) (enzymes involved in \( \beta \)-oxidation), mitochondrial creatine kinase (Ckm1), a mitochondrial ribosomal protein (Mrpl22), and mannose receptor C type 1 (Mrc1). Expression of the latter is positively associated with mitochondrial gene expression in the muscle of humans (Moreno-Navarrete et al. 2013). Highlanders also exhibited higher expression of two aldehyde dehydrogenase paralogs (Aldha1 and Aldha1a7), which encode enzymes that can protect against ischemia-reperfusion injury by reducing oxidative stress, autophagy, and apoptosis during hypoxia (Ma et al. 2011; Contractor et al. 2013; Zhang et al. 2013). The expression of two genes that have been associated with angiogenesis, cadherin-7 (Cdh7) (Hayward et al. 2011) and Notch-4 (Notch4) (Lv et al. 2013), was also more highly expressed in highlanders. Most of the remaining genes with high expression in highlanders are uncharacterized (supplementary table S1, Supplementary Material online).

Most of the genes that were differentially expressed between populations were expressed at a lower level in the highland mice (supplementary table S1, Supplementary Material online). Individual genes that encode proteins involved in the sarcomere (Actn1), peptide transport (Slc15a2), and muscle growth (Tgf) were expressed at lower levels in highland mice, possibly because expression of these genes is more prevalent in fast glycolytic fibers (which are less abundant in the gastrocnemius of highlanders; figs. 1 and 2). Several transcripts for genes involved in cellular protection from reactive \( \mathrm{O}_2 \) species or xenobiotics were also lower in highlanders, such as transferrin (Tfr), ceruloplasmin (CP), glutathione S-transferase (Gstt3), cytochrome P450 enzymes (9030605E09Rik), and flavin-containing monoxygenase (Fmo1). Curiously, some genes involved in oxidative energy metabolism (Atp6, Nd3, Nd4, Ndr1, and Pank1) as well as a potent vasodilator of arteries (urocortin, Ucn) (Diaz and Smani 2013) were also expressed at lower levels in highlanders.

Relationships between Muscle Phenotype and the Expression of Candidate Genes

In addition to the discovery-driven analyses of transcriptional variation, we tested for associations between muscle phenotype and expression level for a number of candidate genes involved in regulating energy metabolism, mitochondrial biogenesis, and/or angiogenesis (Lin et al. 2005; Lanza and Sreekumaran Nair 2010; Gustafsson 2011) (see Materials and Methods). There were functional similarities between our set of candidate genes and the set of genes that were overrepresented in the enrichment analyses, which is not surprising given that both sets of genes are associated with metabolism, muscle fiber-type differentiation, and angiogenesis. Six of the candidate genes exhibited positive associations
with at least one phenotypic trait. Expression of the peroxisome proliferator-activated receptors (PPAR) γ coactivator 1α (PGC-1α) gene Ppargc1a, a purported master regulator of mitochondrial biogenesis, was associated with the numerical density of oxidative fibers in the muscle ($R^2 = 0.203$, $P = 0.046$) and the ratio of capillary surface to fiber surface ($R^2 = 0.239$, $P = 0.034$) (table 3). The association between Tfam, a key activator of mitochondrial gene expression, and the areal density of oxidative fibers was marginally significant ($R^2 = 0.197$, $P = 0.050$). Several of the growth factors (Angpt1, Pdgfa, Pdgfd, and Vegfc) were positively associated with the size of a particular fiber type (table 3) (although only slow oxidative fiber area differed between populations; table 1).

The majority of significant associations between candidate gene expression and muscle phenotype was negative (table 3). Fifteen of the 43 candidate genes showed significant negative associations with at least one phenotypic trait.
Deer mice at high altitudes sustain high metabolic rates to support locomotion and thermogenesis (Hayes 1989) and enhanced performance capacities appear to be adaptive (Hayes and O’Connor 1999). Highland deer mice have therefore evolved an elevated aerobic capacity in hypoxia relative to their lowland counterparts (Cheviron et al. 2012, 2014). Here, we show that this is partly due to a substantial increase in capillarity and oxidative fiber abundance in the skeletal muscle. We also show that the derived muscle phenotype of highland deer mice is associated with significant variation in gene expression across the skeletal muscle transcriptome. For several candidate genes, population differences in expression levels were associated with specific muscle phenotypes, and many of the differentially expressed genes are known to play a role in energy metabolism, muscle fiber composition, and vascular development.

**Discussion**

The highly oxidative phenotype of skeletal muscle in highland deer mice (figs. 1 and 2), which occurs in conjunction with an increase in the activity of several oxidative enzymes (Cheviron et al. 2012, 2014), could have multiple potential benefits at high altitudes. Highland animals must cope with cold temperatures, so the evolution of a more oxidative muscle phenotype could enhance the capacity for shivering and possibly nonshivering thermogenesis (Mineo et al. 2012). A highly oxidative muscle phenotype should also increase the total mitochondrial respiration of an entire muscle when intracellular O2 tensions fall (Hochachka 1985; Scott, Richards, et al. 2009). This is because mitochondrial respiration can be limited by O2 during intracellular hypoxia (Gnaiger 2001), which would reduce the maximum attainable respiration of individual muscle fibers. Therefore, having more oxidative fibers should counterbalance the inhibitory effects of hypoxia on individual muscle fibers. The adaptive significance of having a more oxidative phenotype could involve one or both of these mechanisms in highland taxa that have evolved this derived trait (León-Velarde et al. 1993; Mathieu-Costello et al. 1998; Scott, Egginton, et al. 2009).

The augmented capillarity in the locomotory muscle of highland mice (figs. 1 and 3) should increase the diffusion capacity for oxygen from the blood and should therefore confer an advantage for sustaining aerobic performance in hypoxia (Wagner 1996; Scott and Milsom 2006; Cano et al. 2013). There is generally a strong relationship between the mitochondrial oxygen demands of a tissue and the capacity for oxygen supply from the microcirculation (Hepple 2000). Interestingly, the magnitude of the increase in capillarity in highlanders appears to be greater than the increase in oxidative capacity (fig. 4). This suggests that oxygen diffusion capacity is further enhanced in highlanders to improve oxygen transport in hypoxia. This should also be advantageous in highland deer mice given that they have evolved an increased hemoglobin-O2 affinity (Chappell and Snyder 1984; Storz et al. 2009; Storz, Runck, et al. 2010; Natarajan et al. 2013, 2015), as the increased muscle diffusion capacity mitigates the

| Table 2. Transcriptic Modules Associated with Muscle Phenotypes. |
|---------------------------------|------------------|
| Trait                             | Associated modules |
| Total areal density of oxidative fibers | P2*, T5         |
| Total numerical density of oxidative fibers | P2, T5         |
| Areal density of fast oxidative fibers | T5             |
| Numerical density of fast oxidative fibers | T5             |
| Capillary density                 | P2              |
| Capillary surface density         | P2              |
| Capillary surface per fiber surface | P2*             |

*Note.—The genes that comprise each module are presented in supplementary tables S1 and S2, Supplementary Material online. *Module phenotype associations that remain significant after Bonferroni correction for multiple tests (P < 0.00045).
The trade-off between pulmonary O₂ loading (which is important for safeguarding arterial O₂ saturation under hypoxia) and O₂ unloading in the peripheral circulation.

Despite considerable differences in a range of environmental factors between the two rearing environments (e.g., temperature, ambient partial pressure of O₂, humidity, food type, food availability, and biotic interactions), population differences in capillarity persisted in comparisons of both native and lab-reared F₁ individuals. Similarly, the difference in the proportional abundance of oxidative fibers was generally more pronounced in comparisons between highland and lowland deer mice than in comparisons between native versus lab-reared F₁ individuals with the same population-of-origin, although this trait may have also been affected by rearing environment (as revealed by a lack of significant pairwise differences between populations in the F₁ mice).

Although we cannot entirely rule out transgenerational epigenetic effects, these results suggest that genetic differences between highland and lowland deer mice may underlie adaptive variation in muscle phenotype. It remains possible that these traits can be further modified by phenotypic plasticity; however, the effects of cold acclimation on oxidative capacity are not consistent across studies of domestic mice. Increases in the activity of oxidative enzymes in the muscle occur in response to cold acclimation in some studies (Mineo et al. 2012) but not others (Beaudry and McClelland 2010), and the respiratory capacity of isolated mitochondria does not seem to be altered by cold acclimation (Meyer et al. 2010; Mineo et al. 2012). Likewise, combining cold and hypoxia has no effect on the activity of oxidative enzymes in the gastrocnemius muscle in domestic mice (Beaudry and McClelland 2010). Furthermore, muscle capillarity and oxidative capacity often do not change or even decrease in response to high-altitude hypoxia exposure in lowland humans and other mammals (Mathieu-Costello 2001; Levett et al. 2012; Jacobs et al. 2013). Although muscle disuse (e.g., limb immobilization) is associated with substantial changes in muscle phenotype (Clark 2009), some wild rodents experience minimal changes in muscle form and function in response to large seasonal variations in activity during hibernation or torpor.
Therefore, evolved genotypic specializations may play a more important role than phenotypic plasticity in creating the muscle phenotype of native highlanders.

Transcriptomic Basis of the High-Altitude Phenotype

The evolved differences in gene expression between highland and lowland populations suggest that regulatory changes in genes involved in aerobic metabolism have contributed to high-altitude adaptation (fig. 7). Expression of the transcriptional coactivator PGC-1α (Ppargc1a), which activates the expression of genes involved in mitochondrial metabolism (Lin et al. 2005; Lanza and Sreekumaran Nair 2010), was positively associated with the abundance of oxidative fibers in the muscle (table 3), and was expressed at higher levels in highlanders than lowlanders. Mitochondrial creatine kinase (Ckmt1), which is prevalent in slow oxidative fibers and is important for controlling oxidative phosphorylation to promote energy supply-demand coupling and metabolite stability (Hochachka 1993; Ventura-Clapier et al. 1998), was expressed at higher levels in highlanders than lowlanders. The potential importance of this enzyme in regulating mitochondrial respiration under hypoxia is supported by experimental studies in humans (Ponsot et al. 2006), rats (Walsh et al. 2006), and birds (Scott, Richards, et al. 2009). The mitochondrial isoform of creatine kinase is associated with mitochondria and is highly expressed in tissues with high ATP demands (Wallimann et al. 1992), so its upregulation in highlanders could be associated with the observed variation in expression of Ppargc1a and Tfam. Genes that encode the metabolic enzymes acetyl-coenzyme A acyltransferase (Acaa1a) and aldehyde dehydrogenase 1A1 (Aldh1a1), which are upregulated in mice selected for high levels of voluntary wheel running (Barniston et al. 2013), were also more highly expressed in highlanders than in lowlanders. Expression of Acaat1 is known to be induced by PPAR transcription factors (Guo et al. 2007), for which PGC-1α is a coactivator, suggesting a potential relationship between the patterns of variation in expression of Ppargc1a and Acaat1. However, not all genes involved in oxidative energy metabolism were upregulated. Some mitochondrially encoded transcripts, including two of complex I of the electron transport system (ND3 and ND4, transcripts of two of the 44 subunits of NADH dehydrogenase) and one of ATP synthase (Atp6, one of nine subunits of the FO region of ATP synthase), were downregulated in highlanders. The meaning of these changes in expression is unclear, because NADH dehydrogenase and ATP synthase are multisubunit enzymes whose functions require coordinated expression of all polypeptide subunits (Duggan et al. 2011; Suarez and Moyes 2012). Nevertheless, our findings suggest that high-altitude adaptation involves substantial transcriptomic restructuring in association with a more oxidative muscle phenotype.

Changes in the expression of genes involved in regulating the capillary network also appear to have contributed to evolved phenotypic differences between highland and lowland populations. Cadherin-7 and Notch-4 in particular are expressed at higher levels in highlanders than in lowlanders. Although cadherin-7 is a N-type cadherin that is found primarily in neural tissue (Faulkner-Jones et al. 1999), its expression increases in association with angiogenesis in response to ischemia in the brain (Hayward et al. 2011) and E-type cadherins are known to be extremely important cell adhesion molecules in angiogenesis. The Notch signaling pathway is important for coordinating cell behaviors that lead to the creation of stable and patent blood vessels during angiogenesis, and the notch-4 receptor is restricted to the vascular system (Sainson and Harris 2008; Lv et al. 2013). When notch-4 is bound by its ligand (delta-like 4), signaling by the

(Cotton and Harlow 2010). Therefore, evolved genotypic specializations may play a more important role than phenotypic plasticity in creating the muscle phenotype of native highlanders.

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**Fig. 8.** There was a strong negative association between the expression of fibroblast growth factor receptor 2 (Fgfr2) and the abundance of slow oxidative fibers. There was a significant linear correlation between Fgfr2 transcript abundance and both the numerical ($R^2 = 0.574, P = 0.0001$) and areal ($R^2 = 0.446, P = 0.0013$) densities of slow oxidative fibers in the gastrocnemius muscle. Dashed lines represent the 95% confidence intervals of the regression. Symbols are as follows: F1 lab-raised lowlanders, black upwards triangles; native lowlanders, white upwards triangles; F1 lab-raised highlanders, dark gray downwards triangles; native highlanders, light gray downwards triangles.
receptor inhibits vascular endothelial growth factors (VEGF) expression as well as endothelial cell proliferation and migration (Williams et al. 2005). This is notable in light of the negative association we observed between capillarity and the expression of Vegfa and other growth factors involved in promoting the proliferative and migratory stages of angiogenesis (table 3). It is possible that the highly capillarized phenotype of skeletal muscle in highlanders is initiated by growth factor signaling at an earlier stage of development, and that it is maintained during adulthood by an upregulation of notch-4 expression to stabilize the mature capillary network.

There was a negative association between muscle phenotypes and the expression of many genes. PC1 scores of module P2 expression were lower in highlanders than in lowlanders (fig. 6) (corresponding to lower abundance in highlanders for roughly half of the transcripts in module P2) and were negatively associated with several of the oxidative and capillarity phenotypes. A similar observation was made for PC1 of module T5 expression. Collectively, these modules are enriched for genes assigned to several GO processes involved in metabolism, muscle plasticity, and vascular development. The majority of significant associations between muscle phenotypes and the expression of candidate genes involved in energy metabolism, mitochondrial biogenesis, or angiogenesis was also negative (table 3). These results were surprising and unexpected because the protein products of many of the genes that were expressed at lower levels in highlanders have been shown to contribute positively to capillarity or oxidative capacity in rodents and other mammals (Lin et al. 2005; Lanza and Sreekumaran Nair 2010; Gustafsson 2011). These seemingly paradoxical results may result from the effects that differences in muscle phenotype should have on intracellular signals of oxygen and energy homeostasis. For example, differential expression of a small number of other genes (such as Notch-4) could maintain the highly capillarized phenotype of highlanders, which would increase cellular O2 supply relative to lowlanders and would presumably dampen signaling by the factors that drive the responses to oxygen limitation (such as hypoxia-inducible factors and VEGF). A negative association between muscle phenotypes and expression could thus result for genes that do not cause the differences in muscle phenotype but are sensitive to its effects.

As the increased oxidative capacity and capillarity of skeletal muscle in highland mice were not primarily attributable to phenotypic plasticity, we reasoned that the underlying genes would not exhibit plasticity in expression. Nevertheless, a large number of genes were differentially expressed between mice in the native and lab environments (figs. 5 and 6). Most of the expression differences were not associated with any muscle phenotypes, and may instead reflect transcriptomic changes associated with different aspects of natural and lab environments (table 2). Indeed, environmentally sensitive modules that were not associated with muscle phenotype were enriched for genes involved in signaling, cell migration, cell turnover, immune function, and cell stress, which may reflect the substantial differences between native and lab environments in features such as food access, diet quality, exposure to pathogens, and a range of other factors in addition to differences in O2 tension. Determining which of these environmentally sensitive genes are responding to changes in elevation will require controlled reciprocal acclimation experiments.

Conclusions

One of the goals of evolutionary physiology is to elucidate the mechanistic basis of adaptive variation in organismal performance (Garland and Carter 1994; Dalziel et al. 2009). Populations and species that have diverged across altitudinal gradients are well suited to this endeavor. They provide an opportunity to elucidate how genotypic specialization, phenotypic plasticity, and their interaction can contribute to adaptive enhancements of physiological performance in challenging environments, particularly as cold and hypoxia have well-defined impacts on the physiological systems important for exercise and thermogenesis. In this study, we show that highland deer mice have a highly oxidative and capillarized muscle phenotype that is associated with population differences in expression for a small suite of genes involved in metabolism, muscle plasticity, and vascular development. The capacity for oxygen transport and utilization in the muscle is extremely important for thermogenesis (Oufara et al. 1987; Van Sant and Hammond 2008)—an organismal performance trait with demonstrated fitness benefits in high-altitude deer mice (Hayes and O’Connor 1999)—suggesting that the unique muscle phenotype and transcriptional profile of highland deer mice are adaptive. Consistent with previous work in deer mice (Cheviron et al. 2012, 2014), these results suggest that differences in both evolved and plastic gene expression patterns contribute to fitness-related variation in physiological performance at high altitudes by altering phenotypic traits at several levels of biological organization.

Materials and Methods

Populations of Highland and Lowland Deer Mice

Adult deer mice were live trapped on the summit of Mount Evans (Clear Creek County, CO at 39°35’18”N, 105°38’38”W, 4,350 m above sea level) (P. m. rufinus) and at low altitude in the Great Plains (Nine Mile Prairie, Lancaster County, NE at 40°52’12”N, 96°48’20.3”W, 430 m above sea level) (P. m. nebrascensis) as previously described (Cheviron et al. 2013, 2014). One set of mice, the in situ group, from each locality was sampled on the day of capture at their native elevation for histology and transcriptomics. Another set of mice was transported to a common-garden lab environment at the University of Nebraska (elevation 360m) and used as a parental stock to establish captive bred highland and lowland lines. From each of these lines, we sampled full-sibling F1 progeny from one highland and one lowland breeding pair. All F1 mice were born and raised to adulthood in a common normoxic environment, and were then sampled for histology and transcriptomics. Lab-raised mice were held in standard holding conditions at 23°C with unlimited access to mouse chow and water (12 h light:12 h dark photoperiod). All experimental protocols were approved by the University of Nebraska...
Muscle Histology

Muscle capillarity and oxidative phenotype were assessed as previously described (Scott, Egginton, et al. 2009; Scott and Johnston 2012). An entire gastrocnemius muscle was dissected, coated in embedding medium, frozen in liquid N₂-cooled isopentane, and stored at −80 °C. The muscle was sectioned (10 μm) transverse to muscle fiber length in a cryostat at −20 °C. Alkaline phosphatase activity was used to identify capillaries by staining in assay buffer (concentrations in mM: 1.0 nitroblue tetrazolium, 0.5 5-bromo-4-chloro-3-indoxyl phosphate, 28 NaBO₂, 7 MgSO₄; pH 9.3) for 1 h at room temperature. Succinate dehydrogenase activity was used to identify oxidative muscle fibers (both slow and fast), also by staining for 1 h at room temperature (assay buffer concentrations in mM: 0.6 nitroblue tetrazolium, 2.0 KH₂PO₄, 15.4 Na₃HPO₄, 16.7 sodium succinate). Slow myosin immunoreactivity using the S58 antibody was used to identify oxidative muscle fibers (Developmental Studies Hybridoma Bank, Iowa City, IA) as follows. After an initial fixation in acetone for 10 min, sections were blocked in 10% normal goat serum (made up in phosphate-buffered saline [PBS] [0.15 mol L⁻¹, pH 7.4] containing 1% Triton X-100 and 1.5% BSA [PBS/TX/BSA]) for 1 h. They were then incubated overnight at 2 °C in S58 antibody solution (1:10 dilution in PBS/TX/BSA). The next morning, sections were treated for 10 min with Peroxidase Blocking Reagent (Dako, Burlington, ON, Canada), incubated in secondary antibody (antimouse biotin IgA; Southern Biotech, Birmingham, AL) solution (1:20 dilution in PBS/TX/BSA) for 1 h, incubated in ExtrAvidin-Peroxidase solution (1:50 dilution in PBS/TX/BSA; Sigma-Aldrich, Oakville, ON, Canada) for 30 min, and finally developed in 0.4 mg mL⁻¹ 3-amino-9-ethyl-carbazole solution (containing 0.02% H₂O₂ in 0.05 M sodium acetate buffer, pH 5.0) for approximately 5 min. Sections were well rinsed in PBS between each of the above steps. Images were collected using light microscopy, and stereological quantification methods were used to make unbiased measurements (Weibel 1979; Egginton 1990). We analyzed a sufficient number of images for each sample to account for heterogeneity, determined by the number required to yield a stable mean value.

There is generally a strong relationship between the capacity for oxygen diffusion from capillaries and the mitochondrial oxygen demands they are meant to support (Hepple 2000). Therefore, our expectation was that differences in muscle oxidative capacity would be associated with corresponding differences in muscle capillarity. If increases in muscle capillarity arose in highland mice to enhance O₂ transport in hypoxia, then capillarity would be greater in highland mice than predicted by the normal relationship between capillarity and oxidative capacity. To assess whether there was variation in capillarity that was independent of the variation in muscle oxidative phenotype we first regressed capillary surface density to the areal density of oxidative fibers, and we then used residuals from this regression for all subsequent statistical comparisons (see below).

Histology data are generally reported as means ± standard error (except when data points from individual samples are shown). Two-factor analysis of variance and Bonferroni multiple comparisons tests were used as appropriate to assess the main effects and interactions of population and acclimation environment, and significance level of P < 0.05 was used.

Muscle Transcriptomics

We utilized previously published gastrocnemius RNA-seq data (Cheviron et al. 2014) to quantify genome-wide patterns of gene expression for 24 individuals that were also used for muscle phenotyping to determine the transcriptomic basis of variation in muscle phenotypes (native elevation group: Highland mice, n = 6; lowland mice, n = 6; Lab F1 group: Highland mice, n = 6; lowland mice, n = 6) (NCBI SRA accession number SRA091630). These data were originally used to identify the genomic basis of thermogenic performance differences between high and low elevation populations. In this study, we present a reanalysis of a subset of these data to test for associations between transcriptomic profiles and muscle phenotypes (see supplementary table S5, Supplementary Material online, for the individuals that were used from this larger data set). We isolated mRNA from gastrocnemius muscle using a micro PolyA purist kit (Ambion), and generated Illumina sequencing libraries following standard protocols (available upon request). Libraries were sequenced as 76 nt single-end reads on the Illumina Genome Analyzer Ix, five individuals were multiplexed using Illumina index primers, and were sequenced in a single lane of a flow cell. Image analysis and base calling were performed using Illumina pipeline software. Our sequencing strategy produced an average of 5.9 million reads per individual (range = 1.05–15.7 million reads/individual).

Read Mapping, Normalization, and Statistical Analysis of RNA-seq Data

We performed a series of sequential filtering steps to remove low-quality reads and base calls as well as technical artifacts stemming from library preparation. First, reads with mean Phred quality scores less than 30 were removed from the data set. Second, low-quality bases (Phred score < 30) were trimmed from these remaining high-quality sequences using the “Trim Sequences” tool in CLC Genomics Workbench 6.0.4 (Trimming settings: Trim using quality scores limit: 0.001). Finally, reads were scanned for exact matches to the known Illumina adaptor sequences using the “Trim Adaptors” tool in CLC Genomics Workbench, and if detected, they were trimmed from the sequence read. We estimated transcript abundance by mapping sequence reads to the Mus musculus genome, build 36.1, using CLC Genomics workbench (mapping settings: minimum length fraction = 0.9, minimum similarity fraction = 0.8, and maximum number of hits for a read = 10). We chose to use the well-annotated and established Mus genome as reference instead of the draft P. maniculatus genome to avoid issues associated with poor
assembly and annotations that are common in draft genome assemblies (Denton et al. 2014), but this may have prevented us from estimating expression of rapidly evolving genes that are highly divergent between Mus and Peromyscus. Finally, we excluded genes with less than an average of five reads per individual because genes with low count values are typically subject to increased measurement error and reduced power to detect differential expression (Robinson and Smyth 2007). Filtering low coverage transcripts has the benefit of increasing statistical power to detect differential expression for well-sampled (i.e., high coverage) genes by reducing the number of independent tests that are performed. This filtering strategy could potentially result in bias against short, lowly transcribed genes, but we are unlikely have sufficient power to detect differential expression of such genes anyway because of their low sequence coverage. These trimming and filtering steps resulted in a final data set of 12,175 detected genes and an average of 2.6 million sequence reads per individual (range of 0.5–6.7 million reads).

We used the function calcNormFactors in the program edgeR (Robinson et al. 2010; Robinson and Oshlack 2010) to normalize read counts among individuals, and to control for differences in the total library size (number of total reads) among individuals. Following this normalization procedure, we tested for differences in transcript abundance between populations (highland vs. lowland), and rearing environments (in situ and F1) using a generalized linear model approach in edgeR. Population and rearing environment were included as the main effects, and we included a term for their interaction. We estimated model dispersion for each gene separately using the function estimateDisp in edgeR (McCarthy et al. 2012), and tested for genes that exhibited significant expression differences between populations and/or rearing environments using a general linear model (GLM) likelihood ratio test implemented in edgeR. We controlled for multiple tests by enforcing a genome-wide FDR of 0.05 (Benjamini and Hochberg 1995).

We assessed the degree of correlation in transcript abundance among the genes with significant population or environment effects (FDR < 0.05; n = 657 genes; see Results) to define putative transcriptional modules of coexpressed genes (Ayroles et al. 2009). To do this, we calculated Pearson correlation coefficients for all pairwise gene expression values, and then used MMC to group genes according to their pairwise correlation coefficients (Ayroles et al. 2009; Stone and Ayroles 2009). To determine which of these transcriptional modules were most strongly associated with muscle phenotypes, we used PCA to summarize overall module expression. Because the first two principal component axes (PC1 and PC2) accounted for 47–100% of the explained gene expression variance within modules, PC1 and PC2 scores were used as a proxy measure to summarize overall module expression. Module-specific PC1 and PC2 scores were then used in linear regression analyses to test for associations between module expression and each of the measured phenotypes. PCA and linear regressions were performed in R (ver. 3.1.1; R Development Core Team 2014), and we corrected for multiple tests using Bonferroni adjusted P values.

Once trait-associated modules were identified, we used a suite of functional annotation tools to identify specific “biological process” GO terms associated with each module. First, we used MGI GO Term Mapper (http://www.informatics.jax.org/tools.html, last accessed March 25, 2015) to identify GO Slim terms associated with each gene in a given module, then we used the MGI GO Term Mapper and GOrilla (Eden et al. 2009) to test for functional enrichment of specific terms within modules, and we visualized the GOrilla output using the program REVIGO (Supek et al. 2011).

In addition to these discovery-driven enrichment analyses, we tested for associations between muscle phenotype and expression level for a number of candidate genes (using linear regression analyses, as described above). We considered gene families involved in regulating energy metabolism and/or mitochondrial biogenesis, including the PPAR (Ppara, Pparg, and Pparg2), the PPARγ coactivators (PGC; Ppargc1a, Ppargc1b, and Ppargc1c), nuclear respiratory factors 1 (Nrf1) and 2 (Nfe2l2), and mitochondrial transcription factor A (Tfam) (Lin et al. 2005; Lanza and Sreekumaran Nair 2010). We also considered a number of gene families in which one or more members are involved in angiogenesis, including VEGF (Vegfa, Vegfb, and Vegfc) and their receptors (VEGFR1, Flt1, and VEGFR2, Kdr), angiopoietins (Angpt1, Angpt2, and Angpt4) and their receptors (Tie1 and Tek), fibroblast growth factors (FGF; Fgfl, Fgfl6, Fgfl7, Fgfl10, Fgfl12, Fgfl13, and Fgfl14) and their receptors (FGFR; Fgfr1, Fgfr2, Fgfr3, and Fgfr11), platelet-derived growth factors (PDGF; Pdgfa, Pdgfb, Pdgfd, and Pdgfd1) and their receptors (PDGFR; Pdgfra, Pdgfrb, and Pdgfrd), and matrix metalloproteinases (MMP; Mmp2, Mmp9, Mmp13, Mmp14, Mmp15, and Mmp24) (Gustafsson 2011). We targeted all members of each particular family that were expressed in the muscle, including genes whose influence on muscle phenotype has not been previously documented.

**Supplementary Material**

Supplementary tables S1–S5 are available at Molecular Biology and Evolution online (http://www.mbe.oxfordjournals.org/).

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