Biochemical isolation of myonuclei employed to define changes to the myonuclear proteome that occur with aging

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Summary

Skeletal muscle aging is accompanied by loss of muscle mass and strength. Examining changes in myonuclear proteins with age would provide insight into molecular processes which regulate these profound changes in muscle physiology. However, muscle tissue is highly adapted for contraction and thus comprised largely of contractile proteins making the nuclear proteins difficult to identify from whole muscle samples. By developing a method to purify myonuclei from whole skeletal muscle, we were able to collect myonuclei for analysis by flow cytometry, biochemistry, and mass spectrometry. Nuclear purification dramatically increased the number and intensity of nuclear proteins detected by mass spectrometry compared to whole tissue. We exploited this increased proteomic depth to investigate age-related changes to the myonuclear proteome. Nuclear levels of 54 of 779 identified proteins (7%) changed significantly with age; these proteins were primarily involved in chromatin maintenance and RNA processing. To determine whether the changes we detected were specific to myonuclei or were common to nuclei of excitatory tissues, we compared aging in myonuclei to aging in brain nuclei. Although several of the same processes were affected by aging in both brain and muscle nuclei, the specific proteins involved in these alterations differed between the two tissues. Isolating myonuclei allowed a deeper view into the myonuclear proteome than previously possible facilitating identification of novel age-related changes in skeletal muscle. Our technique will enable future studies into a heretofore underrepresented compartment of skeletal muscle.

Key words: aging; brain; myonuclei; nuclear isolation; proteome; skeletal muscle.

Introduction

Skeletal muscle, which is essential for critical processes such as movement, swallowing, and breathing, dynamically adapts to the body’s needs by altering gene expression to accommodate physiological states such as muscle growth, regeneration, and aging. To date, investigations into molecular mechanisms driving shifts in gene expression have been largely candidate-based. Examining global changes in myonuclear proteins would provide insight into molecular processes regulating large-scale shifts in gene expression that accompany physiologic changes in muscle.

Aging leads to profound changes in skeletal muscle. Starting around age 30 and increasing after age 60, muscle mass and strength decline. The resulting frailty is associated with reduced mobility and greater risk of falls (Demontis et al., 2013). Global changes in RNA levels with age have been identified (Zahn et al., 2006; Kim et al., 2014; Su et al., 2015); however, large-scale proteomic analysis has been limited to the most abundant proteins. While many cell types support skeletal muscle function, the tissue is comprised primarily of myofibers, large, multinucleated cells densely packed with contractile proteins necessary for muscle function. Over half of the total protein in a muscle cell is either actin or myosin (Deshmukh et al., 2015). This lack of heterogeneity in muscle tissue makes proteomic studies that span a large dynamic range of protein abundances technically challenging.

Limited information exists on the dynamic alterations of myonuclear protein levels during aging or other physiologic states. Myonuclear proteins are difficult to study with an unbiased approach because (i) nuclear proteins make up a small minority of proteins in skeletal muscle tissue due to the high abundance of contractile proteins, (ii) myonuclei are challenging to isolate because skeletal muscle is difficult to disrupt without damaging myonuclei, and (iii) dense debris co-sediments with nuclei contaminating the nuclear fraction. In addition, muscle tissue is comprised of multiple cell types. Existing approaches do not provide sufficient enrichment to interrogate low abundance nuclear proteins, require large amounts of starting material, or necessitate genetic tagging of specific nuclei and have no mechanism of determining what proportion of isolated nuclei are myonuclei. An approach that would sufficiently enrich myonuclear proteins to allow unbiased interrogation would offer a new avenue to examine the molecular mechanisms regulating physiologic states in skeletal muscle.

By developing a method of purifying myonuclei from relatively small amounts of mouse muscle tissue, we were able to collect myonuclei for analysis by flow cytometry, biochemistry, and proteomics. Purification of myonuclei dramatically increased both the number and intensity of nuclear proteins detected compared to whole muscle tissue. Querying changes to the myonuclear proteome associated with aging revealed that nuclear levels of RNA processing proteins, nuclear transport proteins, and proteins regulating transcription increased with age. When compared to brain nuclei, many of the same processes were affected in both tissues with age, although the individual proteins that change differed between tissues. Our approach of isolating myonuclei will provide a useful tool for future studies of myonuclear changes in different physiologic states of skeletal muscle.

Results

Isolation of pure intact nuclei

To facilitate biochemical and molecular analyses of myonuclei, we optimized a technique to isolate intact nuclei with high purity from single

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mouse muscles (Fig. 1A). To isolate nuclei, the muscle was gently homogenized. After preparation of a crude nuclear fraction by filtration and low-speed centrifugation, a purified nuclear fraction was obtained by ultracentrifugation over a discontinuous two-step sucrose gradient. Nuclei collect at the interface between the two sucrose cushions with cytoplasmic material above the first cushion and the denser myofibrils under the second cushion. The nuclear fraction was collected and analyzed by downstream methods. The average time required for preparation of a nuclear sample was 5.5 h. The average yield from a single mouse gastrocnemius (GA) muscle was 1 \times 10^6 nuclei, whereas pooling two GA and two rectus femoris (RF) muscles yielded 6 \times 10^6 nuclei and 8 \mu g of total protein.

The isolated nuclei were stained by 4',6-diamidino-2-phenylindole (DAPI) and were free of visible debris when examined by light microscopy (Fig. 1B). The nuclear envelope must remain intact to retain nuclear proteins for examination after purification. To determine whether the nuclear envelope was breached in the purification process, we challenged the nuclei with FITC-conjugated 500 kDa dextran, which is excluded from nuclei with intact nuclear envelopes (D’Angelo et al., 2009). Nuclei were incubated with the dextran and analyzed by fluorescence microscopy. We found that 93% of isolated nuclei excluded the fluorescent dextran and thus had intact nuclear envelopes (Fig. 1C). To further examine whether soluble nuclear proteins were retained during isolation, and to determine the enrichment of nuclear proteins and depletion of proteins from other cellular organelles in the nuclear fractions, we immunoblotted total muscle tissue, as well as cytoplasmic and nuclear fractions (Fig. 1D, for full blots, see Fig. S1, Supporting information). Proteins from various nuclear compartments were enriched in the nuclear fraction including the nuclear envelope (Nup 214, a nuclear pore protein), the soluble fraction (RNA-binding protein HuR), and chromatin (Histone H3). Purified nuclei were also enriched in endoplasmic reticulum markers (ER) but depleted of cytoplasmic (tubulin) markers and depleted for the ER marker calnexin, although calnexin depletion was variable. Together, these results indicate that nuclei isolated using this method retain nuclear proteins, are depleted of other organelles, and have intact nuclear envelopes.

Isolated nuclei are predominantly myonuclei

As muscle tissue contains multiple cell types, we assessed what percentage of nuclei purified from muscle tissue were myonuclei by performing flow cytometry using myonuclear-specific markers. TMEM38A is an outer nuclear envelope transmembrane protein expressed in excitation cells with highest levels in myofibers (Bleunven et al., 2008). We took advantage of this property to use TMEM38A as a marker of myonuclei. Immunofluorescence analysis of isolated nuclei revealed that while all intact nuclei were labeled with a marker of the nuclear pore complex (mAb414), not all nuclei examined were positive for TMEM38A (Fig. 2A). Subsequently, we analyzed immunostained nuclei by flow cytometry. Singlet nuclei were identified by DAPI labeling and side scatter (SSC) and were gated for further analysis (Fig. 2B). To determine what percentage of intact nuclei was TMEM38A positive (TMEM38A+), we gated on mAb414-positive (mAb414+ nuclei. Of this mAb414+ population of nuclei, 96.4% \pm 3.2 SE were TMEM38A+ (Fig. 2C). While a bright and a dim population of TMEM38A+ nuclei were observed, both populations were distinct from nuclei labeled with control IgG (inset Fig. 2C). As TMEM38A localizes both to the nucleus and the sarcoplasmic reticulum (SR), the bright and dim populations of TMEM38A+ nuclei are likely the result of variable amounts of SR associated with isolated nuclei.

To confirm that the TMEM38A antibody specifically labels myonuclei, we took advantage of genetic labeling using the nTnG mouse model which contains a transgene with nuclear-targeted tdTomato and eGFP (Prigge et al., 2013). As illustrated in Fig. 2D, the transgene is composed of a floxed nuclear localization signal (NLS)-containing tdTomato gene followed by a stop codon; outside the floxed region is an NLS-containing eGFP gene. Prior to Cre-mediated recombination, the ubiquitously

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**Fig. 1** Workflow for isolating nuclei from skeletal muscle. (A) Workflow: 1) Skeletal muscle was dissected from mice and then 2) minced and homogenized. 3) The homogenate was filtered and nuclei pelleted at low speed. 4) The crude nuclear fraction was resuspended, layered over sucrose cushions, and 5) ultracentrifuged. 6) The nuclear fraction was collected, diluted, and pelleted at low speed. 7) The nuclear pellet was collected for downstream applications. (B) Purified nuclei were stained with DAPI and examined by microscopy. Singlet nuclei were identified by DAPI labeling and side scatter (SSC) and were gated for further analysis (Fig. 2B). To determine what percentage of intact nuclei was TMEM38A positive (TMEM38A+), we gated on mAb414-positive (mAb414+) nuclei. Of this mAb414+ population of nuclei, 96.4% \pm 3.2 SE were TMEM38A+ (Fig. 2C). While a bright and a dim population of TMEM38A+ nuclei were observed, both populations were distinct from nuclei labeled with control IgG (inset Fig. 2C). As TMEM38A localizes both to the nucleus and the sarcoplasmic reticulum (SR), the bright and dim populations of TMEM38A+ nuclei are likely the result of variable amounts of SR associated with isolated nuclei.
experienced tdTomato labels all nuclei. Upon recombination, the tdTomato gene and stop codons are excised and eGFP labels the nuclei of recombined cells. We bred these nTnG mice to mice expressing Cre recombinase under control of the muscle creatine kinase promoter (Ckmm) (Bruning et al., 1998). In the offspring of this cross, myonuclei are eGFP positive (eGFP*), while all other nuclei are tdTomato positive (tdTomato*). To confirm the specificity of myonuclear labeling in these offspring, we isolated single myofibers from GA muscles and visualized by fluorescence microscopy (Fig. 2E). All myonuclei within single myofibers were eGFP* with only a few tdTomato* cells on each myofiber. The fluorescent labels were retained during nuclear purification (Fig. 2F), making them reliable markers for subsequent flow cytometry analyses.

To compare the specificity of the two myonuclear markers, nuclei were isolated from the muscles of Cre+ nTnG mice and wild-type mice, immunostained for TMEM38A or control IgG, and analyzed by flow cytometry (Fig. 2G). On average, 85.5% ± 5.2 SE of total nuclei were positive for both TMEM38A and eGFP, indicating that the vast majority of purified nuclei were myonuclei. As previously observed (Fig. 2C), both bright and dim populations of TMEM38A were distinct from the control IgG-labeled wild-type nuclei (inset Fig. 2G). To rule out the possibility that a subset of nonmyonuclei were labeled with TMEM38A, we further analyzed the TMEM38A+ nuclei and found that 99.7% ± 0.03 SE were eGFP*, indicating that TMEM38A labels myonuclei (Fig. 2H). Taken together, these results indicate that nuclei isolated by this method from skeletal muscle tissue are at least 85.5% ± 5.2 SE myonuclei.

Increased depth of proteomic detection in myonuclear proteome

We next compared the proteomes of whole muscle tissue samples and isolated nuclear samples using mass spectrometry. Nuclei were purified from GA and RF muscles pooled from three mice. Of 1771 proteins detected in purified nuclear samples, 906 were not detected in whole muscle samples; similarly, 921 of 1786 proteins detected in the whole muscle samples were not detected in isolated myonuclei. We compared the average peptide LC peak intensity area for biological duplicates of nuclear isolation procedure with downstream analysis by mass spectrometry. Muscles were pooled from two mice and five replicates were analyzed for each age. After imputation and correction for batch effect, samples were compared based on p-value and mean fold change. As in the previous experiment, 60% of the proteins detected were classified as nuclear by GO.

The levels of 54 of 779 identified proteins (7%) changed significantly with age (Fig. 4A, full list in Table S2, Supporting information). Because the samples are fractionated, we cannot distinguish overall changes in steady state protein levels from a change in nuclear distribution. Of the significantly changed proteins (P < 0.05, fold change > 1.5), 43 were more abundant in nuclei from old muscle, 1.5- to 7-fold higher than young muscle levels, and 11 were less abundant, 1.5- to 4.5-fold lower than young muscle levels (Table 1). The two categories with the most changed proteins were chromatin maintenance and RNA processing (Fig. 4B). The changes, both increases and decreases, in levels of proteins related to chromatin maintenance suggest general epigenetic changes with aging. In addition to age-related increases in levels of RNA processing proteins, levels of proteins associated with transcriptional regulation increased, suggesting overall changes in transcript production and/or processing. Nuclear transport protein levels also increased, potentially indicating alterations in nuclear transport and potentially

Fig. 2 Isolated nuclei are predominately myonuclei. (A) Purified nuclei were stained with DAPI and immunostained with anticoregernuclearexpression complex antibody (Mab414) to label all nuclear envelopes and TMEM38A antibody to label myonuclei. Pictured are representative images of TMEM38A+ and TMEM38A- nuclei. Bar = 10 μm. (B) Isolated nuclei were analyzed by flow cytometry by side scatter (SSC) and DAPI; intact DAPI+ singlets were selected for further analysis (red gate). (C) IgG control immunostained nuclei (gray) were compared to experimental nuclei immunostained with Mab414 and TMEM38A antibodies (magenta). Myonuclei were defined to be positive for both TMEM38A and eGFP, indicating that the vast majority of purified nuclei were myonuclei. As previously observed (Fig. 2C), both bright and dim populations of TMEM38A were distinct from the control IgG-labeled wild-type nuclei (inset). (D) Transgenic mice with a cassette containing nuclear-targeted tdTomato and eGFP reporter proteins (nTnG) were crossed with mice expressing Cre recombinase from a skeletal muscle-specific promoter (Ckmm) to genetically label myonuclei and nonmyonuclei with distinct fluorescent markers in the offspring. All nuclei from wild-type mice (WT) are nonfluorescent. All nuclei from nTnG- Ckmm Cre- mice (Cre-) are fluorescent red. Myonuclei from nTnG- Ckmm Cre+ mice (Cre+) are fluorescent green, while nonmyonuclei are fluorescent red. (E) Single myofibers were isolated from Cre- and Cre+ mice retained their fluorescent label after isolation. Bar = 10 μm. (F) Nuclei isolated from a Cre- mouse (purple) and a WT mouse (gray) were analyzed by TMEM38A and eGFP fluorescence; 85.5% ± 5.2 SE of nuclei from Cre+ mice were positive for both TMEM38A and eGFP (n = 4). Bright and dim TMEM38A populations were distinct from control IgG immunostained nuclei (inset). TMEM38A+ nuclei were selected for further analysis (red gate). (H) TMEM38A+ nuclei were analyzed for eGFP fluorescence; 99.7% ± 0.03 SE of TMEM38A+ nuclei were eGFP*, confirming that TMEM38A labels myonuclei.
consequential mislocalization of proteins. Ribosomal protein levels also increased with age, which could be related to changes in ribosome assembly and altered nuclear transport or increased rough ER contribution to the aged nuclei fraction. Of the changed proteins from non-nuclear cellular compartments, most proteins were classified as plasma membrane proteins and mitochondrial proteins. That these proteins change could reflect age-related changes in the overall protein abundance or a differential association with young and old myonuclei. These data show that nuclear levels of markers of diverse nuclear processes including chromatin organization, RNA processing, and nuclear transport are altered during aging of mouse skeletal muscles.

**Myonuclei share common aging pathways with brain nuclei**

To investigate whether the changes to the myonuclear proteome reflect general age-related changes or muscle-specific changes, we compared the age-related changes in myonuclei to those observed in brain nuclei, another excitatory, long-lived, postmitotic tissue. We isolated and processed brain nuclei from young (3-month) and old (24-month) mice as described above for myonuclei. By immunoblotting, isolated brain nuclei were free of contamination from mitochondrial and cytoplasmic compartments (Fig. S2, Supporting Information). Similar to the myonuclear samples, 61% of the proteins identified from brain nuclei were classified as nuclear by GO. Analysis revealed that levels of 32 of 845 identified proteins (4%) changed significantly (FDR corrected, p < 0.05, >1.5-fold change) between young and old brain nuclei (Fig. 5A). Of the changed proteins, 21 were increased in abundance in aged brain nuclei, 1.5- to 3-fold compared to young brain levels, while 11 decreased, 1.5- to 10-fold compared to young brain levels (Table 2). Similar to the changes detected in myonuclei with age, the majority of these proteins were related to chromatin maintenance and RNA processing (Fig. 5B).

Several cellular processes changed with age in nuclei from both brain and muscle (Fig. 5C). Brain and muscle nuclei shared increases in markers of transcription, and RNA processing with age as well as overall changes in chromatin maintenance markers, supporting the concept that these are common pathways affected in aging. Two pathways were uniquely changed in nuclei from aged muscle versus brain. Nuclear lamin levels increased in aged brain nuclei but not myonuclei, while ribosomal proteins increased with age in myonuclei, but not brain nuclei. Despite the similarity in changed processes, the only protein significantly changed with age in both tissues was RCC1 (Fig. 5D), which increased more than 2-fold in aged nuclei in both tissues. RCC1 is the major guanine nucleotide exchange factor for Ran, helping to establish the Ran GTP gradient, which is the driving force in nucleocytoplasmic transport (Izaurralde et al., 1997). This increase in RCC1 in both aged brain and muscle nuclei may suggest age-related changes in nuclear transport (Fig. 5C) in both tissues.

While some of the same processes were affected by aging in both brain and muscle nuclei, the specific proteins that changed differed between the two tissues. Most of the proteins that changed in the nuclei of either tissue were detected in both tissues but only changed in one (Fig. 5D): 49 of 54 proteins (91%) that changed in muscle were detected...
Thus, our biochemical isolation has the advantage of isolating a high percentage of myonuclei without genetic labeling.

Using biochemical fractionation, we dramatically enriched the nuclear proteome from skeletal muscle. In many tissues, nuclear proteins are readily detectable by proteomics in unfraccionated tissue. Nuclear proteins in skeletal muscle are underrepresented because they comprise a small percentage of the overall muscle proteome (Table S3, Supporting information). Consistent with a previous report (Deshmukh et al., 2015), we find that >60% of the peptide reads identified by mass spectrometry of whole muscle tissue correspond to contractile proteins. Others have employed various approaches to increase proteomic detection of noncontractile proteins in skeletal muscle including depletion of contractile proteins (Carberry et al., 2014; Gueugneau et al., 2014) and enriching less detected proteins (Gannon & Ohlendieck, 2012). While these approaches have enriched some noncontractile proteins, they identified few nuclear proteins. Our study offers the first systematic examination of the myonuclear proteome, identifying 535 annotated nuclear proteins by GO, and provides tools for others to interrogate changes in this important cellular compartment.

As muscles age, muscle mass and function decrease and myofibers undergo fiber-type switching. Proteomic studies of aging muscle have described alterations in the most abundant proteins in muscle: contractile and metabolic proteins (Baraibar et al., 2014; Su et al., 2015). To investigate how aging affects myonuclei at the protein level, we compared the myonuclear proteomes of young and old mice. Some of the changes we detected in protein levels mirror changes in transcript levels: transcript levels of RNA-binding proteins Hnnpr (Welle et al., 2003), Srsf5, Paf1, and Ddx41 (Su et al., 2015), a chromatin maintenance protein Ssrp1 (Su et al., 2015), and a nuclear transport protein Kpnb1 (Swindell, 2009) increase with age consistent with changes we observed in protein levels. Protein levels of the RNA-binding protein Khdrbs1 also increase with age (Laohavinij, 2015). In addition to these specific proteins, transcript levels for RNA processing machinery genes in general increase with age (Gheorghe et al., 2014). Similarly, we identified an increase in RNA-binding protein levels within the nucleus with age. Our data showed an increase in ribosomal proteins in myonuclei with age. Others have characterized increases in transcript levels of ribosomal proteins (Calura et al., 2008; Fedorov et al., 2014) and decreased ribosome biogenesis (Kirby et al., 2015) with age. While our results could reflect increased ER contribution to the nuclear fraction

Discussion

In this study, we present a method optimized to yield high-purity myonuclei from relatively small samples for downstream analysis by flow cytometry, biochemistry, and mass spectrometry. We exploited this approach to interrogate the myonuclear proteome and investigate changes to the myonuclear proteome that occur with age.

Isolating nuclei from skeletal muscle has historically been difficult. Several researchers have developed approaches to isolate nuclei from skeletal muscle, each technique optimized for a downstream application like ChIP (Ohkawa et al., 2012), analysis of nuclear envelopes (Wilkie & Schirmer, 2008), or comparison of multiple cellular compartments (Dimarou et al., 2012). Our goal was to develop an isolation technique compatible with downstream analysis by flow cytometry and mass spectrometry using relatively small sample sizes to preserve biological variability. Our approach results in high-purity myonuclei isolated from a single GA or RF muscle sufficient for analysis by flow cytometry and pooled GA and RF muscles from two mice sufficient for analysis by mass spectrometry. The isolated nuclei are impermeable to large dextrans, indicating that the nuclear envelopes remain intact, and soluble nuclear proteins such as transcription factors and RNA processing enzymes are readily detectable. Both results indicate that the nuclei do not undergo major loss of protein during purification.

Genetic labeling approaches could complement our biochemical fractionation strategy. Fluorescent labeling using nTnG transgenic mice as used here combined with fluorescent activated nuclear sorting or a recently developed approach termed isolation of nuclei tagged in specific cell types (INTACT) (Deal & Henikoff, 2011; Jankowska et al., 2016) could increase purity of isolated myonuclei. However, both methods require generation of new mouse strains for each cell type of interest.
Table 1: Myonuclear proteins that change significantly with age

| log2 fold change | P-value | Uniprot ID | Gene Symbol | Protein name | Role |
|------------------|---------|------------|-------------|--------------|------|
| −2.19            | 0.009   | P43276     | Hist1h1b    | Histone H1.5 | Chromatin maintenance |
| −2.12            | 0.014   | Q70V5-2    | Synm        | Synemin      | Cytoskeletal |
| −1.73            | 0.019   | Q6PIC6     | Atplalpha3  | Sodium/potassium-transporting ATPase subunit alpha-3 | Membrane ion pump |
| −1.67            | 0.040   | OS4944     | Smarc1      | SWNSNP-related matrix-associated actin-dependent regulator of chromatin subfamily E member 1 | Chromatin maintenance |
| −1.46            | 0.018   | Q91XV3     | Basp1       | Brain acid soluble protein 1 | Transcription |
| −1.40            | 0.047   | P49813     | Tmd1        | Tropomodulin-1 | Sarcomeric |
| −1.38            | 0.036   | Q9D6R2-2   | Idh3a       | Isocitrate dehydrogenase [NAD] subunit alpha, mitochondrial | Metabolism |
| −1.24            | 0.039   | P62242     | Rps8        | 40S ribosomal protein S8 | Ribosomal |
| −1.22            | 0.032   | AZAUC9     | Kif4d1      | Kelch-like protein 41 | Myofibril assembly |
| −1.08            | 0.025   | Q9ESU6     | Brd4        | Bromodomain-containing protein 4 | Chromatin maintenance |
| −0.73            | 0.016   | P15637     | Cav3        | Caveolin-3 | Membrane protein scaffold |
| 0.69             | 0.024   | Q810A7-2   | Ddx42       | ATP-dependent RNA helicase DDX42 | RNA processing |
| 0.70             | 0.041   | G3UX35     | Smarc4      | Isoform of Q3TKT4, Transcription activator BRG1 | Chromatin maintenance |
| 0.82             | 0.031   | P62908     | Rps3        | 40S ribosomal protein S3 | Ribosomal |
| 0.84             | 0.038   | Q8K4Q8     | Colec12     | Collectin-12 | Cell signaling |
| 0.84             | 0.021   | Q6O8S3     | Caprin1     | Caprin-1 | RNA processing |
| 0.86             | 0.013   | Q64S11     | Top2b       | DNA topoisomerase 2-beta | Transcription |
| 0.87             | 0.046   | E9Q7Q3     | Tpm3        | Isoform of P21107, Tropomyosin alpha-3 chain | Sarcomeric |
| 0.93             | 0.040   | Q8K278     | Rfa1        | RNA polymerase II-associated factor 1 homolog | Transcription |
| 0.97             | 0.027   | Q64S22     | Hist2h2ab   | Histone H2A type 2-B | Chromatin maintenance |
| 1.00             | 0.002   | S4R1C4     | Atp2b2      | Isoform of Q9RK7, Calcium-transporting ATPase | Membrane ion pump |
| 1.03             | 0.025   | P62309     | Snpg        | Small nuclear ribonucleoprotein G | RNA processing |
| 1.07             | 0.032   | Q8BH74     | Nup107      | Nuclear pore complex protein Nup107 | Nuclear Transport |
| 1.13             | 0.029   | Q3UN8B     | Mcpt4       | Isoform of P21812, Mast cell protease 4 | Cytoskeletal protease |
| 1.26             | 0.026   | Q9Z29P     | Glyr1       | Putative oxidoreductase GLYR1 | Chromatin maintenance |
| 1.35             | 0.033   | G3U22      | Syncrip     | Isoform of Q7TMK5, Heterogeneous nuclear ribonucleoprotein Q | RNA processing |
| 1.35             | 0.049   | Q99Y0      | Hadhb       | Trifunctional enzyme subunit beta, mitochondrial | Metabolism |
| 1.36             | 0.002   | P08113     | Hsp90b1     | Endoplasm | ER chaperone |
| 1.36             | 0.040   | Q8CJF7     | Ahtc1f1     | Protein ELYS | Nuclear Transport |
| 1.38             | 0.006   | P32067     | Sdb         | Lupus La protein homolog | RNA processing |
| 1.40             | 0.040   | P68040     | Gnb2l1      | Guanine nucleotide-binding protein subunit beta-2-like 1 | Cell signaling |
| 1.43             | 0.025   | P47857     | Pfkm        | ATP-dependent 6-phosphofructokinase, muscle type | Metabolism |
| 1.43             | 0.002   | Q9Y1Q5     | Rpn1        | Dolichyl-diphosphooligosaccharide-protein glycosyltransferase subunit 1 | ER glycosylation |
| 1.44             | 0.048   | Q60749     | Khdbs1      | KH domain-containing, RNA-binding, signal transduction-associated protein 1 | RNA processing |
| 1.44             | 0.023   | Q08539-2   | Bin1        | Myc box-dependent-interacting protein 1 | Apoptotic process |
| 1.46             | 0.030   | Q8QZT1     | Acat1       | Acetyl-CoA acetyltransferase, mitochondrial | Metabolism |
| 1.50             | 0.016   | Q35326     | Srsl5       | Serine/arginine-rich splicing factor 5 | RNA processing |
| 1.51             | 0.002   | P62858     | Rps28       | 40S ribosomal protein S28 | Ribosomal |
| 1.52             | 0.001   | P10852     | Slic3a2     | 4F2 cell-surface antigen heavy chain | Membrane transporter |
| 1.57             | 0.025   | Q9D8E6     | Rp4         | 60S ribosomal protein L4 | Ribosomal |
| 1.59             | 0.028   | O35343     | Kpn4a       | Importin subunit alpha-3 | Nuclear Transport |
| 1.61             | 0.016   | P42669     | Pura        | Transcriptional activator protein Puf-alpha | RNA processing |
| 1.61             | 0.015   | Q99MR6-3   | Srrt        | Serrate RNA effector molecule homolog | RNA processing |
| 1.78             | 0.004   | A2AW05     | Srp1        | Isoform of Q08943, FACT complex subunit SSRP1 | Transcription |
| 1.78             | 0.019   | E9PYL9     | Gm10036     | Protein Gm10036 | RNA processing |
| 1.81             | 0.008   | Q8VHM5     | Hnmpr       | Heterogeneous nuclear ribonucleoprotein R | RNA processing |
| 1.84             | 0.002   | P30275     | Ckmt1       | Creatine kinase U-type, mitochondrial | Metabolism |
| 1.85             | 0.046   | Q62WY9     | Hist1h2bc   | Histone H2B type 1-C/E/G | Chromatin maintenance |
| 1.90             | 0.047   | P70168     | Kpn1        | Importin subunit beta-1 | Nuclear Transport |
| 1.90             | 0.035   | P62315     | Snrgd1      | Small nuclear ribonucleoprotein Sm D1 | RNA processing |
| 1.94             | 0.026   | Q9CQF3     | Nusht1      | Cleavage and polyadenylation specificity factor subunit 5 | RNA processing |
| 2.05             | 0.023   | D3YW2X     | Ylp1m       | Isoform of Q9RK7, YLP motif-containing protein 1 | RNA processing |
| 2.33             | 0.013   | Q8VE37     | Rct1        | Regulator of chromosome condensation | Nuclear Transport |
| 2.89             | 0.005   | Q99KK2     | Cmas        | N-aclylneuraminate cytidylytransferase | Metabolism |

From older mice, they could also support abnormalities in ribosome biogenesis or export. With our method of enriching nuclear proteins, we were able to investigate the myonuclear proteome at greater depth than previously possible and identify novel myonuclear age-related changes. To determine whether the age-related changes we detected in the myonuclear proteome were common to nuclei of other excitatory postmitotic tissues, we compared aging in brain and muscle nuclei. Previous comparisons of transcript levels from whole skeletal muscle and...
neuronal tissues with aging predominantly identified common changes in metabolism (Capitanio et al., 2009; Gheorghe et al., 2014), chromatin maintenance, and RNA processing (Gheorghe et al., 2014; Su et al., 2015). We likewise detected age-related changes in proteins involved in chromatin maintenance and RNA processing in nuclei from both brain and muscle. In addition, we identified changes in transcription. While the same processes were affected in nuclei from both tissues, Rcc1 was the only protein that increased with age in nuclei from both tissues. RCC1 helps establish the Ran GTP gradient necessary for nucleocytoplasmic transport (Izaurralde et al., 1997). Increased Rcc1 activity increases DNA damage repair and reduces senescence, suggesting a role in aging (Cekan et al., 2016). While pathways were shared between nuclei from both tissues, the individual proteins that changed differed, which is consistent with a previous study in which aging affected the same pathways in two hind-limb muscles but via different proteins (Chaves et al., 2013).

In summary, our approach to isolating myonuclei allowed a deeper view into the myonuclear proteome than previously possible. Muscle-specific changes that occur with aging in nuclear proteins could offer new insight into processes contributing to age-related muscle loss. Isolating and interrogating myonuclei will also allow investigation into nuclear processes involved in muscle growth, regeneration, and response to disease. In addition to examining changes in the proteome, one could isolate myonuclei to interrogate...
the nascent transcriptome and other nuclear processes difficult to detect over cytoplasmic background. Our technique will enable future studies into a heretofore underrepresented compartment of skeletal muscle.

Experimental procedures

Mice

Wild-type C57BL6 mice were obtained from Charles River laboratories (Wilmington, MA) and the National Institute of Aging. nTnG mice containing an allele for nuclear-targeted tdTomato and eGFP reporter proteins (B6N.129S6-Gt(ROSA)26Sortm1(CAG-tdTomato*,EGFP*)Eesc/J) (Prigge et al., 2013) were purchased from Jackson Laboratory (Bar Harbour, ME). These mice were bred to mice expressing Cre recombinase under control of the muscle creatine kinase promoter (B6.FVB(129S4)-Tg(ME). These mice were bred to mice expressing Cre recombinase under control of the muscle creatine kinase promoter (B6.FVB(129S4)-Tg(ME). These mice were bred to mice expressing Cre recombinase under control of the muscle creatine kinase promoter (B6.FVB(129S4)-Tg(ME). These mice were bred to mice expressing Cre recombinase under control of the muscle creatine kinase promoter (B6.FVB(129S4)-Tg(ME), 1998), also purchased from Jackson Laboratory. The genotype of the offspring was determined using PCR protocols available on the Jackson Laboratory website.

All experiments were performed using tissues from 3- to 24-month-old male C57BL6 mice or 3- to 6-month-old male mice homozygous for the nTnG transgene and heterozygous for Cre recombinase. Experiments were performed in accordance with approved guidelines and ethical approval from Emory University’s Institutional Animal Care and Use Committee and in compliance with the National Institutes of Health.

Nuclear isolation

The procedure described below was extensively modified from (Wilkie & Schirmer, 2008) to achieve sufficient sample quantity and quality for analysis by flow cytometry (a single mouse GA or RF) and mass spectrometry (pooled GA and RF from two to three mice). All steps were carried out at 4 °C.

Whole brain or GA and RF muscles were dissected, minced, and suspended in 10 mL homogenization buffer 1 (10 mM HEPES, 60 mM KCl, 0.5 mM spermidine, 0.15 mM spermine, 2 mM EDTA, 0.5 mM EGTA, 300 mM sucrose, 5 mM MgCl2, 2 mM dithiothreitol (DTT), and 5% complete mini protease inhibitors (CMP) (Roche Diagnostics, Risch-Rotkreuz, Switzerland)). Subsequently, muscles were homogenized with 20–25 strokes using a 15-mL PTFE tissue grinder with clearance 0.15–0.25 mm (VWR, Radnor, PA, USA). The homogenate was filtered and centrifuged at 1000 × g for 10 min, yielding a crude nuclear pellet. The pellet was resuspended in 3 mL 1.7 M sucrose with two strokes of the dounce homogenizer and loaded over a two-step sucrose cushion: 2.8 M sucrose and 2.0 M sucrose in 50 mM HEPES, 15 mM KCl, and 5 mM MgCl2. The sample was centrifuged for 3 h at 18673 × g in a SW41Ti rotor in a Beckman Optima LE-80k ultracentrifuge. For larger

| log2 fold change | P-value | Uniprot ID | Gene Symbol | Protein name | Role |
|-----------------|---------|------------|-------------|--------------|------|
| −3.40           | 0.001   | P10854     | Hist1h2bm   | Histone H2B type 1-M | Chromatin maintenance |
| −3.19           | 0.003   | P84228     | Hist1h3b    | Histone H3.2 | Chromatin maintenance |
| −2.03           | 0.005   | D37784     | Syt1        | Isoform of P46096, Synaptotagmin-1 | Vesicular trafficking |
| −1.77           | 0.024   | Q9CFW2     | Tubb2b      | Tubulin beta-2B chain | Cytoskeletal |
| −1.54           | 0.008   | Q8QZY9      | Sf3b4       | Splicing factor 3B subunit 4 | RNA processing |
| −1.32           | 0.035   | Q9QYG0      | Ndrg2       | Protein NDRG2 | Signal transduction |
| −1.32           | 0.032   | Q9ES97-3    | Rtn3        | Reticulon-3 | Beta amyloid regulation |
| −0.86           | 0.005   | A2AR02      | Ppg         | Peptidyl-prolyl cis-trans isomerase G | Lamin |
| −0.79           | 0.024   | Q9QYX7-2    | Pcd          | Protein piccolo | Cytoskeletal |
| −0.70           | 0.014   | P28659-2    | Cell1       | CUGBP Elav-like family member | RNA processing |
| −0.53           | 0.019   | Q0P678      | Zc3 h18     | Zinc finger CCCH domain-containing protein 18 | RNA processing |
| 0.53            | 0.023   | P0C065      | H2afz       | Histone H2A.Z | Chromatin maintenance |
| 0.57            | 0.026   | Q6PDMD2     | Srsf1       | Serine/arginine-rich splicing factor 1 | RNA processing |
| 0.64            | 0.011   | P21619      | Lmnb2       | Lamin-B2 | Lamin |
| 0.69            | 0.045   | Q9CX86      | Hnmpa0      | Heterogeneous nuclear ribonucleoprotein A0 | RNA processing |
| 0.71            | 0.046   | P70372      | Elav1       | ELAV-like protein 1 | RNA processing |
| 0.75            | 0.013   | Q64525      | Hist2h2bb   | Histone H2B type 2-B | Chromatin maintenance |
| 0.75            | 0.013   | A0A087WRG2  | U2surp      | Isoform of O6N8V3, U2 snRNP-associated SURF motif-containing protein | RNA processing |
| 0.80            | 0.021   | P70288      | Hdc2        | Histone deacetylase 2 | Chromatin maintenance |
| 0.84            | 0.048   | Q91V95      | Ddx1        | ATP-dependent RNA helicase DDX1 | RNA processing |
| 0.88            | 0.026   | Q9CU62      | Smc1a       | Structural maintenance of chromosomes protein 1A | Chromatin maintenance |
| 0.98            | 0.039   | G3U234      | Eftud2      | Isoform of O08810, 116 kDa US small nuclear ribonucleoprotein component | RNA processing |
| 1.09            | 0.036   | Q62318      | Trim28      | Transcription intermediary factor 1-beta | Translation |
| 1.11            | 0.005   | Q9CQ7      | Snrb2       | U2 small nuclear ribonucleoprotein B\* | RNA processing |
| 1.18            | 0.042   | P63158      | Hmgb1       | High mobility group protein B1 | Chromatin maintenance |
| 1.27            | 0.018   | P10126      | Eef1a1      | Elongation factor 1-alpha-1 | Lamin |
| 1.30            | 0.049   | Q61191      | Hcf1        | Host cell factor 1 | Lamin |
| 1.33            | 0.003   | P48678-2    | Lmna        | Prelamin-A/C | Lamin |
| 1.37            | 0.019   | Q8VE37      | Rcc1        | Regulator of chromosomal condensation | Nuclear Transport |
| 1.37            | 0.006   | Q6Z3P8-2    | Zc3b4       | Zinc finger CCCH domain-containing protein 4 | RNA processing |
| 1.39            | 0.037   | Q7NT2-2     | Far2        | Fatty acyl-CoA reductase 2 | Metabolism |
| 1.49            | 0.005   | A0A0G2J095  | Rbn1        | Isoform of Q80T69, Round spermatid basic protein 1 | Translation |

Table 2 Brain nuclear proteins that change significantly with age

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preparations for proteomics, nuclei were isolated with the same gradient but centrifuged with a SW32Ti rotor in a Beckman Optima LE-80K centrifuge at 175587 × g for 195 min and buffers were supplemented with 1× HALT protease and phosphatase inhibitor (Thermo Scientific, Waltham, MA, USA).

After ultracentrifugation, the nuclei concentrated at the interface between the 2.0 m and 2.8 m sucrose layers were collected, diluted 1:15 with resuspension buffer (20 mM HEPES, 10 mM KCl, 1.5 mM MgCl₂, 0.5 mM spermidine, 0.15 mM spermine, 0.2 mM EDTA, and 5% CMP (Roche Diagnostics)), and mixed thoroughly by inverting. For optimal yield, the nuclear fraction was collected into conical tubes pretreated with 1% bovine serum albumin (BSA) to prevent nuclei sticking to tube walls. The nuclei were pelleted at 3000 × g for 15 min. The small clear pellet was washed in resuspension buffer and pelleted at 2000 × g for 5 min. For biochemistry and mass spectrometry, the pellet was washed two more times to remove residual BSA.

To assess the integrity of isolated nuclei, a 10-μl aliquot of nuclei was stained with 1 μg mL⁻¹ DAPI and 60 μg mL⁻¹ 500kD fluorescein isothiocyanate (FITC)-conjugated dextran (Sigma-Aldrich, St. Louis, MO, USA) and examined by fluorescence microscopy. Nuclei were scored for exclusion of the dextran by fluorescence microscopy (D’Angelo et al., 2009).

**Immunoblotting**

GA and RF muscles were dissected and either homogenized in homogenization buffer 2 (150 mM NaCl, 1% NP-40, 0.5% sodium deoxycholate, 0.1% sodium dodecyl sulfate, 50 mM Tris-pH 8, 5% CMP (Roche Diagnostics)) or processed to isolate nuclei as described above. The cytoplasmic fraction was retained after the first centrifugation step. Equal protein content (Bradford, 1976) was resolved by SDS-PAGE electrophoresis. Proteins were transferred to nitrocellulose membranes and detected with antibodies (Table S4, Supporting information) and enhanced chemiluminescence.

**Single myofiber isolation**

GA muscles were dissected and processed as described previously (Pichavant & Pavlath, 2014). Briefly, muscles were enzymatically digested, single myofibers transferred to multiwell plates and fixed with paraformaldehyde.

**Flow cytometry**

Isolated nuclei were incubated with antibodies (see Table S4, Supporting information for antibodies and dilutions) or appropriate isotype controls for 30 min on ice, then washed, and incubated with secondary antibodies for 30 min on ice. Nuclei were stained with 1 μg mL⁻¹ DAPI immediately before analysis. After gating on DAPI⁺ nuclei, 20 000 nuclear events from each sample were analyzed for Texas Red, AF647, and eGFP fluorescence using a BD LSRII flow cytometer. Analyses of flow nuclear events from each sample were analyzed for Texas Red, AF647, and eGFP fluorescence using a BD LSRII flow cytometer. Analyses of flow cytometry data were performed using FlowJo LLC, Ashland, Oregon, USA (version X 10.0.7i2).

**LC-MS/MS analysis**

Samples were processed following established protocols (Wang et al., 2016). For a detailed description, see supplemental methods. Briefly, 50 μg from each sample was subjected to in-solution trypsin digest (Herskovitz et al., 2010). Peptide mixtures were separated by a NanoAcquity UHPLC (Waters, Milford, FA, USA) and monitored on a Orbitrap Fusion mass spectrometer (ThermoFisher Scientific, San Jose, CA, USA). The mass spectrometer cycle was programmed for “top speed acquisition” with a cycle time of 3 s.

**Label-free proteomic quantification**

For a detailed description of quantification, see supplemental methods (Data S1). Briefly, for analysis of purified myonuclei compared to whole muscle samples, raw files were searched against the mouse UniPort reference database using SEQUEST algorithm through Proteome Discoverer 2.0 platform (Thermo Scientific, Bremen, Germany). The embedded Percolator algorithm (Kall et al., 2007) was used to filter the peptide spectral matches to achieve a false discovery rate (FDR) of <1%. For analysis of age-related proteomic changes in myonuclei and brain nuclei, data were analyzed using MAXQUANT, freely distributed by J. Cox of Max Planck Institute of Biochemistry, Munich, Germany (v1.5.2.8) with Thermo Foundation 2.0 for RAW file reading capability (Cox and Mann 2008). The label-free quantification (LFQ) algorithm in MaxQuant (Cox et al., 2014, Luber et al., 2010) was used for protein quantitation. Imputation of missing values using Perseus (Tyanova et al., 2016) was followed by batch effect correction using ComBat (Johnson et al., 2007). After calculating fold change and p-value for each protein, data were analyzed by GO term using DAVID (da Huang et al., 2008a,b) or KEGG pathway by GO-ELITE v1.2.5 (Zambon et al., 2012).

**Image acquisition**

For analysis of isolated nuclei, images were obtained using an Axioplan microscope (Carl Zeiss MicroImaging, Oberkochen, Germany) with either a 0.3 NA 10× Plan-Neofluar objective (Carl Zeiss MicroImaging) or a 0.8 NA 25× Plan-Neofluar objective (Carl Zeiss MicroImaging) and were recorded with a camera (Carl Zeiss MicroImaging) and Scion Image 1.63 (Scion Corporation, Torrance, CA, USA) software. For analysis of single myofibers, images were obtained using a Nikon Eclipse TE2000-U confocal microscope (Nikon, Tokyo, Japan) with a 0.5 NA Plan Fluor 20× objective (Nikon) and were recorded with a SensiCam QE ( Cooke, Campbell, CA, USA) with IPLab 4.0 (Scanalytics, Fairfax, VA, USA) software. All images were assembled and equally processed using Adobe Photoshop CS4 version 11.0.2 (Adobe, San Jose, CA, USA).

**Statistical analysis**

Data were analyzed for statistical significance based on fold change, P-value, or z-score. z-score enrichment was determined using GOElite (Zambon et al., 2012). A P-value of < 0.05 or z-score of >1.96 was considered statistically significant. Relevant changes were considered >1.5-fold.

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Author contributions

AAC participated in conception and design, collection and assembly of data, data analysis and interpretation, financial support, manuscript writing, and final approval of manuscript. ED and DD were involved in sample preparation, data analysis and interpretation of proteomic experiments, and final approval of manuscript. NTS carried out data analysis and interpretation of proteomic experiments, manuscript writing, and final approval of manuscript. AHC took part in conception and design, data analysis and interpretation, manuscript writing, and final approval of manuscript. GP participated in conception and design, data analysis and interpretation, financial support, manuscript writing, and final approval of manuscript.

Conflict of interest

None declared.

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For sources cited in supplemental materials see supplementary references (Data S2).

Supporting Information

Additional Supporting Information may be found online in the supporting information tab for this article. Also, the raw proteomics files are available at http://www.proteomexchange.org/.

Fig. S1 Full blots of antibodies used in Figure 1.

Fig. S2 Biochemical purity of nuclei isolated from the brain.

Table S1 Comparison of proteins detected in whole muscle and purified myonuclei samples.

Table S2 Comparison of protein levels detected in nuclei isolated from brain and muscle from young and old mice.

Table S3 Summary of nuclear proteins detected in proteomic studies of aging skeletal muscle.

Table S4 Antibodies used.

Data S1 Methods: Detailed descriptions of LC-MS/MS analysis and label-free quantification.

Data S2 Supplementary references.