Disrupted Anabolic and Catabolic Processes May Contribute to Alcohol-Accentuated SAIDS-Associated Wasting

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Background. Alcohol abuse is a comorbid factor in many human immunodeficiency virus (HIV)–infected patients. Previously, we demonstrated that chronic binge alcohol accentuates loss of body mass at terminal stage of simian immunodeficiency virus (SIV) infection. The purpose of this study was to investigate changes in pathways that may contribute to muscle wasting in chronic binge alcohol-fed SIV-infected macaques.

Methods. The impact of chronic binge alcohol during SIV infection on insulin signaling and the ubiquitin (Ub)–proteasome system—regulators of protein synthesis and degradation—was examined in SIV-infected macaques.

Results. SIV infection induced an inflammatory and pro-oxidative milieu in skeletal muscle, which was associated with decreased insulin-stimulated phosphatidylinositol 3-kinase (PI-3k) activity and upregulated gene expression of mTOR and atrogin-1, and protein expression of Ub-proteasome system 19S base. Chronic binge alcohol accentuated the skeletal muscle pro-oxidative milieu and 19S base expression. Additionally, chronic binge alcohol increased skeletal muscle protein expression of protein-tyrosine phosphatase 1B (a negative regulator of insulin signaling) and 19S proteasome regulator non-ATPase (Rpn) 6 subunit and Rpn12, and suppressed PI-3K activity. Animals that were alcohol-fed and SIV-infected for >15 months had increased Ub-proteasome system activity.

Conclusions. These data suggest negative modulation of insulin signaling coupled with enhanced Ub-proteasome system activity may be central mechanisms underlying chronic binge alcohol-induced accentuation of SIV-associated muscle wasting.

According to the Centers for Disease Control and Prevention, >400 000 people were living with AIDS in the United States at the end of 2003 [1]. The reduction in human immunodeficiency virus (HIV)–associated morbidity and mortality resulting from the use of highly active antiretroviral therapy (HAART) has made HIV infection a chronic disease, during which individuals are likely to engage in alcohol and drug abuse at rates comparable to those in the noninfected population [2]. Chronic alcohol consumption remains the most common and costly form of drug abuse with ~14 million Americans fulfilling the diagnostic criteria for alcohol abuse and/or alcoholism. Indeed, alcohol abuse and HIV infection frequently co-exist [3].

Muscle wasting remains an important determinant of increased morbidity and mortality observed in AIDS [4–7]. Chronic alcohol abuse is associated with skeletal muscle myopathy [8] resulting from decreased muscle protein synthesis [9–11] and possibly accelerated muscle...
proteolysis [12]. Additionally, indirect effects of alcohol such as alterations in nutritional state, micronutrient availability, and growth factor expression have been implicated in the etiology of alcohol-induced muscle wasting [13]. Because loss of lean body mass has a deleterious impact on the overall survival of HIV-infected individuals, factors that accelerate this process, such as chronic alcohol consumption, likely accelerate disease progression and development of AIDS-associated wasting [14–17].

We have previously shown that chronic binge alcohol-administered SIV-infected (ALC/SIV+) macaques had significantly lower body weight, body mass index (BMI), and limb muscle area compared with sucrose-treated SIV-infected (SUC/SIV+) animals [18]. In addition, our studies showed messenger RNA (mRNA) expression of IGF-1 is suppressed, atrogin-1 is increased, TNF-α is significantly increased, and myostatin is modestly elevated in skeletal muscle from ALC/SIV+ animals at terminal stage of SIV infection (SAIDS) [18]. Atrogin-1 is a muscle-specific E3 ubiquitin ligase and has been implicated as a causal factor in muscle wasting [19]. Tumor necrosis factor α (TNF-α) has been shown to exert anti-insulin effects in skeletal muscle [20, 21], and the negative regulator of skeletal muscle growth, myostatin [22], has been implicated in muscle wasting in HIV-infected men [23]. Together, our findings suggested upregulation of the ubiquitin (Ub)–proteasome system as a predominant mechanism underlying loss of skeletal muscle mass observed in those studies. However, it was not investigated whether parallel suppression in anabolic regulatory pathways was also involved. Moreover, whether localized skeletal muscle inflammation was associated with an oxidative milieu further contributing to dysregulation in muscle mass was not investigated. The present study extends our observations to integrate the role of disrupted anabolic signaling as an additional, potentially synergistic mechanism that is responsible for disruption in the balance between the synthetic and catabolic mechanisms leading to erosion of muscle mass in chronic binge alcohol-administered SIV-infected macaques.

MATERIALS AND METHODS

All experiments were approved by the Institutional Animal Care and Use Committee at both Tulane National Primate Research Center (TNPRC) in Covington, Louisiana, and Louisiana State University Health Sciences Center in New Orleans, Louisiana, and adhered to National Institutes of Health guidelines for the care and use of experimental animals. Four- to six-year-old male macaques (Macaca mulatta) obtained from TNPRC breeding colonies were studied. Excellent health of animals was determined by (1) complete physical examinations by a veterinarian; (2) complete blood counts and serum chemistries; and (3) negative serological status for simian retrovirus (confirmed by DNA amplification by nested polymerase chain reaction [PCR], enzyme immunoassay [EIA], and Western immunoblot analysis) and simian T-lymphotropic virus type 1 (EIA and Western immunoblot analysis) based on assays performed by the Pathogen Detection Laboratory (California National Primate Research Center, Davis, CA). Age- and body-weight-matched animals were randomized to either isocaloric SUC/SIV+ or ALC/SIV+ groups. Skeletal muscle samples were obtained at necropsy from a group of SIV-negative, healthy control macaques (age, 7.4 ± 1.10 years; weight [mean +/- SEM], 11.1 ± 1.51 kg) and used as reference values for comparison of analyzed variables. A total of 28 animals were studied: 9 in the SUC/SIV+ group, 11 in the ALC/SIV+ group, and 8 in the control group. Animals were individually housed in a Biosafety Level 2 containment building.

Experimental Protocol

The gastric catheter placement for alcohol delivery, the alcohol delivery protocol, and the route of SIV infection have been described elsewhere [18, 24]. Animals were exposed to chronic alcohol by intragastric administration of a mean of 13–14 grams of ethanol (30% w/v water) per kilogram of body weight per week beginning 3 months prior to SIV infection and continued throughout the duration of study. This approach of intragastric delivery was selected to reduce experimental variability and ensure chronic binge-like intoxicating blood alcohol concentrations of 50–60 mmol/L. Total calories provided by alcohol and sucrose averaged 15%. Animals were provided monkey chow ad libitum (Lab Fiber Plus Primate diet DT; PMI Nutrition International, St. Louis, MO) and supplemented with fruits, vitamins, and Noyes treats (Research Diets, New Brunswick, NJ). Three months after initiating chronic binge alcohol administration, animals were inoculated intravenously with 10,000 times the 50% infective dose (ID50) of SIVmac251, provided by Preston Marx. Intravenous inoculation was performed at the conclusion of an ethanol or sucrose session when blood alcohol levels were elevated to simulate infection during an alcohol binge. The progression of SIV disease was monitored throughout the study period through clinical, biochemical, and immunological parameters (CD4/CD8 lymphocyte ratios) in addition to plasma viral kinetics (SIV gag RNA levels) as described elsewhere [18, 24, 25]. Skeletal muscle (gastrocnemius) samples were obtained at necropsy when animals met criteria for euthanasia based on the following: (1) loss of 25% of body weight from maximum body weight since assignment to protocol; (2) major organ failure or medical conditions unresponsive to treatment; (3) surgical complications unresponsive to immediate intervention; or (4) complete anorexia for 4 days. Because of the large range in time of SIV infection within the groups, initial data analyses grouped results based on time after SIV infection, as either early infection or late infection. The late infection group is defined as the group with time to euthanasia that was ≥2 SD from the mean time to euthanasia in the early infection group. In the SUC/SIV+ group, the mean time (± SD) of SIV infection for the early...
(n = 6) and late (n = 3) infection groups was 5.2 ± 2.3 months and 24.3 ± 8.2 months, respectively. In the ALC/SIV + group, the mean time (± SD) of SIV infection for the early (n = 8) and late (n = 3) infection groups was 5.3 ± 3.0 months and 25.4 ± 5.9 months, respectively. No significant differences were found in outcome measures obtained in these animals on analysis according to early or late infection, with the exception of proteasome activity. Thus, with the exception of proteasome activity, for which both combined and late infection values are shown, results represent the combined data of both early and late subgroups. In addition, no significant differences were noted in viral load (~6.3 log copies/mL) or CD4/CD8 cell ratio (~0.7) at time of necropsy, as reported elsewhere [18].

Muscle mRNA and Protein Expression
Muscle samples were frozen in liquid nitrogen and stored at −80°C until analyses. Samples were homogenized and total RNA was isolated using an RNeasy Mini kit (Qiagen, Valencia, CA) according to the manufacturer’s instructions. Muscle tissue lysates were dissected and homogenized using a PRO 200 homogenizer (PRO scientific, Oxford, CT) in buffer A (25 mmol/L HEPES [pH, 7.4], 1% Nonidet P-40 [NP-40], 137 mmol/L NaCl, 1 mmol/L PMSF, 10 µg/mL aprotinin, 1 µg/mL pepstatin, and 5 µg/mL leupeptin). Samples were centrifuged at 14 000 g for 20 minutes at 4°C, and protein concentrations were determined by a BCA protein assay kit (Bio-Rad Laboratories, Hercules, CA).

Real-time Reverse-Transcription PCR Primers and Probes
The amplification primers and probes used for determination of TNF-α, IFN-γ, IL-6, IL-1β, and atrogin-1 were purchased from Applied Biosystems (Carlsbad, CA) and used according to the manufacturer’s instructions. The amplification primers and FAM-labeled probes used for determination of RPS13 and mTOR are as follows: RPS13 forward primer, 5’-TCTGAC-GAGCTGAAGGAGGAGATT; RPS13 reverse primer, 5’-TCTC TCAGGATCACACCGATTT; RPS13 probe, 5’-AAACTGGCA CAAGAAGGGGCTGACTCCCT; mTOR forward primer, 5’-A GTGTCCTTGCTCTGTTGTCAGGCT; mTOR reverse primer, 5’-AAATTAGCTGGGACATGTTGTTG; mTOR probe, 5’-AA GCAAATCTCTTGCCCTAGGCTCCCAA.

Real-time Quantitative Reverse-Transcription PCR
Total RNA was reverse transcribed and real-time PCR was performed with a Superscript III Platinum 2-step quantitative reverse-transcription PCR (RT-qPCR) kit according to the manufacturer’s instructions (Invitrogen, Carlsbad, CA). All reactions were performed on a CFX96 system (Bio-Rad Laboratories, Hercules, CA). The primer and probe concentrations were 500 and 150 nmol/L, respectively, for RPS13 and mTOR, respectively, and RT-qPCR data were analyzed using the ΔΔCt method. Target genes were compared with the endogenous control, RPS13, and normalized to control values or to SUC/SIV + values if the target was not detectable in control samples. RPS13 was chosen as the endogenous control based on a meta-analysis of 13 629 human gene array samples to determine the most stably expressed genes [26].

Western Blot Analysis
Muscle proteins (50 µg) were resolved by sodium dodecyl sulfate polyacrylamide gel electrophoresis and subjected to immunoblot analysis. Protein abundance was detected with antibodies against insulin receptor substrate 1 (IRS-1), IRS-2, phosphatidylinositol 3 kinase (PI3-K), Akt1, Akt2, protein-tyrosine phosphatase 1B (PTP1B), and 19S base S5A subunit (Upstate, Lake Placid, NY); insulin receptor β subunit (IRβ; Santa Cruz Biotechnology, Santa Cruz, CA); glucose transporter type 4 (GLUT4; R&D Systems, Minneapolis, MN); 19S proteasome regulator non-ATPase 6 subunit (Rpn6) and Rpn12 (Biomol International, Plymouth Meeting, PA); and β-actin (Sigma-Aldrich, St. Louis, MO), using Chemiluminescence Reagent Plus (PerkinElmer Life Science, Boston, MA), and quantified with a densitometer. Proteins were normalized to β-actin.

Ex Vivo Antioxidant Capacity Determinations
Antioxidant capacity was determined in skeletal muscle homogenates according to the method of Klemm et al [27]. Briefly, snap-frozen samples of tissue were ptempically homogenized and mixed with Dulbecco phosphate-buffered saline containing 3.5 mmol/L glucose (pH, 7.4; concentration, 1:10) on ice. The samples were incubated (shaking water bath at 37°C) in the presence or absence of 10 mmol/L N-ethylmaleimide (to prevent glutathione recycling) for 30 minutes. At time 0, CPH (1-hydroxy-3-carboxy-2,2,5,5-tetramethylpyrrolidine HCl) and tertiary butyl hydroperoxide were added to the samples at final concentrations of 2.0 and 0.4 mmol/L, respectively, and the mixture was incubated in the shaking water bath for 30 minutes. The samples were then snap-frozen and stored in liquid nitrogen until X-band electron paramagnetic resonance (EPR) spectroscopy was performed. On analysis, samples were quickly thawed in a water bath at 37°C, aspirated into glass capillaries (ID, 0.8 mm), and read at room temperature using a Bruker EMX (Bruker BioSpin, Billerica, MA) with the following settings: microwave power, 40 mW; frequency, 9.79 GHz; center field, 3485 G; sweep width, 100 G; time constant, 10.3 milliseconds; sweep time, 10 seconds (5 sweeps); and receiver gain, 7.9 × 103. Scans from samples incubated in the absence of N-ethylmaleimide were subtracted from the corresponding ones in its presence with the derivative data subjected to double integration using WINEPR software (Bruker BioSpin, Billerica, MA); values are expressed as a percentage of antioxidant capacity.

PI3-K Activity Assay
IRS-1–associated PI3-K activity levels were assessed as described elsewhere [28, 29]. Briefly, 500 µg of muscle lysate was immunoprecipitated with 3 µg of IRS-1 antibody and protein

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Agarose. IRS immune complexes were incubated (10 minutes at 22°C) in 50 µL of 20 mmol/L Tris/HCl buffer (pH 7.0) containing 50 µmol/L [γ-32P] ATP (5 µCi; Perkin Elmer, Boston, MA), 10 mmol/L MgCl₂, 2 mmol/L MnCl₂, 100 mmol/L NaCl, 2 mmol/L EDTA, and 10 µg of phosphatidylinositol. After thin layer chromatography, radiolabeled phosphatidylinositol 3-phosphate was visualized by autoradiography and quantitated by densitometry (BioRad, Hercules, CA).

20S Proteasome Activity Assay

20S proteasome activity levels were measured using a 20S Proteasome Activity assay kit (Chemicon International, Temecula, CA) according to the manufacturer’s instructions. Briefly, 20 µg of muscle lysate was incubated (30 minutes at 37°C) with fluorophore 7-amino-4-methylcoumarin (AMC)–labeled specific 20S proteasome substrates LLVY-AMC. The free AMC fluorescence was quantified using a 380/460 nm filter set in a fluorometer (BioTex, Winooski, VT). The AMC standard curve was generated with a dilution series of purified 20S proteasome. Each sample/substrate combination was assayed in duplicate, both in the presence and in the absence of the proteasome inhibitor MG132 (20 µmol/L; Calbiochem, La Jolla, CA).

Data Analysis

All data are presented as the mean (± SEM) in each experimental group. The sample size is included in the legend of each figure. Treatment effects were established using 1-way analysis of variance with a Kruskal-Wallis test followed by a Dunn multiple comparisons test. Results for which P ≤ .05 were considered to be statistically significant. Statistical analyses were performed using Prism 5 software (GraphPad Software, San Diego, CA).

RESULTS

SIV Infection and Chronic Binge Alcohol Consumption Favor Skeletal Muscle Inflammation and Oxidative Stress

Skeletal muscle mRNA expression of inflammatory cytokine genes TNF-α, IFN-γ, IL-6, and IL-1β were markedly increased in SIV-infected animals (Figure 1). IL-6 mRNA expression was significantly greater in the SIV-infected animals (P < .05) (Figure 1). A trend of higher expression of all inflammatory cytokines was observed in skeletal muscle of ALC/SIV+ animals. However, this trend did not reach statistical significance. Only IL-6 and IL-1β mRNA expression was detected in skeletal muscle obtained from control animals. Skeletal muscle antioxidant capacity was significantly reduced in SUC/SIV+ animals by 2.5-fold (P < .05) and in ALC/SIV+ animals by 7-fold (P < .01) (Figure 1).

SIV Infection and Chronic Binge Alcohol Consumption Disrupt Anabolic Signaling Pathways

Insulin-receptor signaling cascade proteins and downstream targets were assessed by Western blot analysis. There were no statistical differences in the protein content of IRS-1, IRS-2, IRβ, PI3-K, Akt1, Akt2, and GLUT4 between groups (Figure 2). The PTP1B protein content was not altered by SIV infection but was significantly increased in the ALC/SIV+ animals (P < .01).
However, overall skeletal muscle PTP1B activity did not differ between groups (Figure 3). IRS-1–associated PI3-K activity was not changed by SIV infection but was significantly reduced in skeletal muscle of ALC/SIV animals ($P < .05$) (Figure 4). Expression of mTOR mRNA was significantly reduced in skeletal muscle of both SUC/SIV$^+$ and ALC/SIV$^+$ animals ($P < .05$) (Figure 4). In contrast, atrogin-1 mRNA expression was markedly increased in skeletal muscle from both SIV-infected groups ($P < .01$) (Figure 4).

**SIV Infection and Chronic Binge Alcohol Consumption Alter Protein Expression of Ub-Proteasome System Components and Increase Proteasome Activity**

SIV infection did not affect skeletal muscle expression of the 20S catalytic core ($\beta 21$) of the 26S proteasome (Figure 5). However, SIV infection increased protein expression of the 19S base (S5A subunit) in both sucrose-administered macaques ($P < .05$) and alcohol-administered macaques ($P < .01$), but it did not produce significant alterations in protein expression of 19S lid subunits Rpn6 and Rpn12 (Figure 5). In skeletal muscle of ALC/SIV$^+$ macaques an additional marked increase in protein expression of both Rpn6 and Rpn12 was observed ($P < .01$) (Figure 5). Proteasome activity in skeletal muscle isolated from SIV-infected animals did not reflect any significant changes (Figure 5). However, when stratified by time after SIV infection, skeletal muscle obtained from the late infection ALC/SIV$^+$ group showed significant increases in 20S proteasome activity compared with the late infection SUC/SIV$^+$ group ($P < .05$) (Figure 5).

![Figure 2.](https://academic.oup.com/jid/article-abstract/204/8/1246/815932)
The present study examined the impact of chronic binge alcohol intake on anabolic and catabolic mechanisms of muscle wasting in SIV-infected rhesus macaques. SIV infection created an inflammatory and oxidative milieu in skeletal muscle. These changes were associated with increased atrogin-1 and decreased mTOR mRNA expression. Chronic binge alcohol augmented skeletal muscle oxidative stress, dysregulated Ub-proteasome system subunit expression, and increased proteasome activity in late infection animals. Moreover, chronic binge alcohol increased PTP1B protein expression, and decreased IRS-1–associated PI3-K activity in SIV-infected macaques. Interestingly, in previous studies chronic binge alcohol for up to 13 months in uninfected macaques failed to alter skeletal muscle mRNA expression of IL-6, TNF-α, and atrogin-1 [25]. Taken together, these findings suggest that alcohol amplifies catabolic and blunts anabolic skeletal muscle mechanisms in SIV-infected macaques, contributing to muscle wasting.

Previously we demonstrated that SUC/SIV* animals did not show significant alterations in BMI, although the observed BMI in ALC/SIV* animals was 20% lower in animals with SAIDS [18]. In addition, our studies demonstrated that ALC/SIV* macaques had significantly lower limb muscle area compared with SUC/SIV* macaques with SAIDS [18]. Moreover, limb, abdominal, and hip circumferences were all 5%–10% smaller in ALC/SIV* animals, but skinfolds were not significantly smaller. These findings suggested that these changes were not due to loss of subcutaneous fat but due to loss in muscle mass [18]. Weight loss, particularly loss of metabolically active lean tissue, has been associated with accelerated disease progression, loss of muscle protein mass, and impairment of strength and functional status in HIV-infected individuals [30]. The findings from the present study provide insight into the mechanisms responsible for the greater loss in skeletal muscle. Contrary to what we expected, with the exception of proteasome activity, the derangements in skeletal muscle components of both anabolic and catabolic pathways were already evident during the first 10 months of infection. These findings suggest that changes in the skeletal muscle milieu leading to muscle wasting in SIV infection occurs throughout the course of infection and is possibly enhanced as the disease progresses.

Previous studies in patients with AIDS wasting have provided evidence for the involvement of the Ub-proteasome system in loss of skeletal muscle mass, and it has been suggested that this pathway is activated by pro-inflammatory cytokines such as TNF-α and glucocorticoids, leading to AIDS-associated wasting [31]. Mild oxidative stress causes increased protein catabolism and increased expression of Ub-proteasome system components in skeletal myotubes [32]. Oxidative stress also increases muscle-specific E2 and E3 proteins (enzymes involved in activating and attaching ubiquitin to substrates for degradation by the 26S
proteasome) that regulate muscle catabolism [33]. The 26S proteasome is responsible for the degradation of intracellular proteins and is composed of a 20S catalytic core and 19S polar caps. The 19S caps are made up of a lid and a base composed of several regulatory protein subunits. The results from the present study support those findings. A lower antioxidant capacity was observed in skeletal muscle of ALC/SIV animals, which was associated with increased protein expression of the 19S base (S5A). Although SUC/SIV animals showed no change in measured Rpn subunit expression, ALC/SIV animals had significantly increased protein expression of Rpn6 and Rpn12. The function of the Rpn subunits is not completely understood. However, studies report that Rpn6 is essential for the proper assembly of the 19S lid component as well as the proper functioning of the 26S proteasome [34].

We have previously shown that skeletal muscle IGF-1 mRNA expression is decreased in both SUC/SIV+ and ALC/SIV+ animals with SAIDS, suggesting suppression of anabolic regulation of skeletal muscle mass [18]. The results of the present study provide further support for that hypothesis. Our results show that the protein content of a specific phosphatase, PTP1B, is significantly increased in ALC/SIV+ macaques. PTP1B is a negative regulator of cellular insulin signaling and is responsible for insulin receptor dephosphorylation, resulting in termination of insulin signaling. PTP1B also dephosphorylates IRS-1, resulting in deactivation of this insulin signaling mediator [35]. Additionally, PTP1B has been shown to regulate IGF-1 receptor kinase activity, and upregulation of PTP1B decreased both IGF-1 receptor and Akt activation [36]. Thus, we predict that increased expression PTP1B results in impaired insulin- and IGF-1–mediated anabolic effects, and that this is further enhanced in ALC/SIV+ animals.

In parallel, our results show that PI3-K activity is significantly decreased in skeletal muscle of ALC/SIV+ animals. This may be due to increased PTP1B expression decreasing insulin and IGF-1 signaling. Decreased PI3-K activity would be expected to result in decreased Akt activation and consequently decreased mTOR and forkhead box (FoxO) phosphorylation, as illustrated in Figure 6. Both mTOR and FoxO are important transcription factors involved in muscle anabolic and catabolic responses through their modulation of protein synthesis (mTOR) and muscle-specific E3 ligase (atrogen-1 and MuRF-1) gene expression. When FoxO

Figure 4. Skeletal muscle phosphatidylinositol 3 kinase (PI3-K) activity, mTOR expression, and atrogin 1 expression. A, PI3-K activity normalized to levels in control animals (n = 8) in sucrose-treated simian immunodeficiency virus (SIV)–infected (SUC/SIV+) animals (n = 9) and chronic binge alcohol-administered SIV-infected (ALC/SIV+) animals (n = 11). B, C, Levels of mTOR gene and atrogin-1 gene messenger RNA (mRNA) expression, respectively, expressed as fold change of control values (n = 5) in SUC/SIV+ (n = 9) animals and ALC/SIV+ animals (n = 10). P values were obtained using 1-way analysis of variance with a Kruskal-Wallis test followed by a Dunn multiple comparisons test. *P < .05 versus control; **P < .01 versus control.
proteins are phosphorylated they are unable to translocate to the nucleus and promote transcription of the E3 ligases. These muscle-specific ubiquitin ligases target muscle proteins for degradation by the Ub-proteasome system. We speculate that decreased PI3-K activity leads to decreased Akt-mediated phosphorylation of mTOR and FoxO family members (Figure 6). Our results show significantly decreased mTOR and increased atrogin-1 mRNA expression in skeletal muscle of SIV-infected

Figure 5. Skeletal muscle Ub-proteasome system subunit expression and proteasome activity. A, Representative Western blot and mean (± SEM) densitometric values of proteasome subunit expression in skeletal muscle of control animals (n = 8), sucrose-treated simian immunodeficiency virus (SIV)-infected (SUC/SIV⁺) animals (n = 11). Expression was normalized to β-actin and expressed as fold change of control values. B, Proteasome activity in skeletal muscle of control animals (n = 8), SUC/SIV⁺ animals (n = 9), and ALC/SIV⁺ animals (n = 11). C, Proteasome activity in control (n = 8) and late infection animals, SUC/SIV⁺ animals (n = 3), and ALC/SIV⁺ animals (n = 3). Pvalues were obtained by 1-way analysis of variance with a Kruskal-Wallis test followed by a Dunn multiple comparisons test. *P < .05 versus control; **P < .01 versus control. AMC, 7-amino-4-methylcoumarin.

Figure 6. Pathways involved in protein turnover. Protein turnover involves a delicate balance of synthetic and catabolic mechanisms. Insulin and insulin-like growth factor 1 (IGF-1) signaling lead to protein synthesis through activation of the mTOR pathway, Akt signaling, and phosphorylation of forkhead box (FoxO). Protein degradation occurs via major cellular proteolytic pathways, which can be enhanced by cytokines and reactive oxygen species. Dysregulated insulin signaling, increased cytokine expression, decreased antioxidant capacity, and upregulation of the Ub-proteasome system can all lead to an imbalance in protein turnover, favoring degradation over synthesis. Circled pathway members, with arrows indicating direction of change, are the results presented in this article. Dashed arrows and crosses show what we predict occurs to other pathway members based on the data we have presented. Together, these changes may lead to decreased protein synthesis and enhanced protein breakdown.
animals. In another study, we found that MuRF-1 mRNA levels were increased in both SUC/SIV+ and ALC/SIV+ animals with SAIDS [18].

Taken together, the results of our studies demonstrate that chronic alcohol consumption during SIV infection may lead to decreased insulin anabolic regulation of skeletal muscle mass, as reflected by the increased PTP1B expression, decreased PI3-K activity, reduced mTOR expression, and increased muscle-specific E3 ubiquitin ligase expression (Figure 6). We propose that this dysfunctional insulin signaling is a mechanism that likely contributes to SIV- and HIV-associated muscle wasting. Thus, these findings suggest that the decrease in muscle mass in animals with SAIDS is likely the result of a combination of a pro-inflammatory, pro-oxidative, and anabolic imbalance that favors catabolic processes leading to loss of muscle mass and impaired muscle growth. These findings are in agreement with previous reports in the literature from clinical studies indicating that inflammation impacts the responsiveness of muscle tissue to an anabolic stimulus [37] and is associated with marked alterations of protein metabolism [38, 39].

In conclusion, our results show that SIV infection results in marked alterations in the skeletal muscle milieu that are characterized by inflammation and oxidative stress. These are associated with significant derangements in components of catabolic and anabolic pathways responsible for maintenance of skeletal muscle mass. In addition, we show that chronic alcohol accentuates the pro-oxidative and pro-inflammatory skeletal muscle milieu, further disrupting both anabolic and catabolic regulatory mechanisms and promoting loss of skeletal muscle mass. In combination, the altered signaling mechanisms and inflammatory and oxidative milieu are likely central mechanisms for the accentuated SAIDS wasting of chronic binge alcohol-treated SIV-infected animals previously reported by our laboratory.

Notes

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