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The muscle metabolome differs between healthy and frail older adults

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ABSTRACT:

Populations around the world are aging rapidly. Age-related loss of physiological functions negatively affects quality of life. A major contributor to the frailty syndrome of aging is loss of skeletal muscle. In this study we assessed the skeletal muscle biopsy metabolome of healthy young, healthy older and frail older subjects to determine the effect of age and frailty on the metabolic signature of skeletal muscle tissue. In addition, the effects of prolonged whole-body resistance-type exercise training on the muscle metabolome of older subjects were examined. The baseline metabolome was measured in muscle biopsies collected from 30 young, 66 healthy older subjects and 43 frail older subjects. Follow-up samples from frail older (24 samples) and healthy older subjects (38 samples) were collected after 6 months of prolonged resistance-type exercise training. Young subjects were included as a reference group. Primary differences in skeletal muscle metabolite levels between young and healthy older subjects were related to mitochondrial function, muscle fiber type, and tissue turnover. Similar differences were observed when comparing frail older subjects with healthy older subjects at baseline. Prolonged resistance-type exercise training resulted in an adaptive response of amino acid metabolism, especially reflected in branched chain amino acids and genes related to tissue remodeling. The effect of exercise training on branched-chain amino acid-derived acylcarnitines in older subjects points to a downward shift in branched-chain amino acid catabolism upon training. We observed only modest correlations between muscle and plasma metabolite levels, which pleads against the use of plasma metabolites as a direct read-out of muscle metabolism and stresses the need for direct assessment of metabolites in muscle tissue biopsies.

KEYWORDS: muscle biopsy, frailty, aging, tissue remodeling

INTRODUCTION

Populations around the world are aging rapidly. It has been estimated that people older than 60 years of age will make up 22% of the world population, whereas people older than 80 years of age will account for 4.4% of the world population in 2050¹. Age-related loss of physiological functions compromises independence at older age. A major contributor to the frailty syndrome of aging is skeletal muscle loss, which can lead to increased disability in the older population. In most people muscle mass and strength start to decline around the age of 35 with more progressive muscle loss observed after the age of 65²⁻⁶.

Exercise training has been applied to counter the effects muscle mass loss, such as loss of strength ^{7, 8}. We recently demonstrated that prolonged resistance-type exercise training partially shifts the skeletal muscle transcriptome of older subjects toward an expression pattern observed in muscle tissue of young subjects, with changes in gene expression related to vascularisation, tissue remodelling and glucose metabolism ⁹. The transcriptome analysis also revealed substantial differences between healthy young men, healthy older subjects and frail older subjects before any intervention was undertaken. In particular, genes related to young subjects.

The age-related differences in gene expression level in muscle are expected to be reflected in local levels of metabolites. However, limited information is available on the effect of aging on muscle metabolite levels. A recent study in mice showed that aging affects muscle glucose and fatty acid metabolism¹⁰, whereas a study in rats indicated that aging leads to muscle group-specific perturbations in lipid and glucose metabolism consistent with mitochondrial dysfunction ¹¹. In humans, it was found that changes in lipid content and oxidative activity in skeletal muscle during aging are related to a shift in muscle fiber type¹². Interestingly, exercise training led to reprogramming of mitochondrial function and intermediary

metabolism in insulin-insensitive obese subjects ¹³. In both human studies, the metabolic profiling platforms that were used had limited coverage and focused on specific submetabolomes. A major bottleneck in achieving extended coverage has been the limited amount of muscle tissue material that can be sampled from human subjects.

On the basis of the observed shift in transcriptome profiles after training we hypothesized that a similar shift towards the young phenotype occurs in the muscle metabolome after resistance-type exercise training in older subjects. The recent development of a comprehensive targeted metabolic profiling platform optimized and validated for small muscle biopsies makes it possible to elaborately verify this hypothesis ¹⁴. Here we aim to investigate the effect of aging and frailty on the skeletal muscle metabolome. In addition, we examined the impact of prolonged resistance-type exercise training on the metabolome of frail and healthy older subjects.

MATERIALS AND METHODS

Experimental design

Muscle biopsies (*m. vastus lateralis*), plasma and serum samples were collected from frail older ^{15, 16}, healthy older ^{17, 18} and young subjects ¹⁹. The frail group included both frail and pre-frail subjects and are hereafter referred to as frail. Medical history of all subjects was evaluated by medical questionnaires, which were analyzed by a physician. Subjects who were unable to participate in the exercise program due to pain were excluded prior to starting the intervention. Included subjects who showed severe discomfort during the training sessions were excluded from further participation in the study. Baseline metabolite levels were measured in skeletal muscle tissue of 30 young, 66 healthy and 43 frail older subjects (Table 1). We also measured metabolites in plasma and serum for 50 young, 76 healthy and 62 frail

 older subjects (Supporting Information Table S1). The transcriptome of the muscle samples was measured in an earlier study ⁹.

Baseline samples from healthy young male subjects served as reference and were derived from several studies performed within our group, in which exactly the same techniques and processing were used for sample collection. Samples from the frail and healthy older subjects were obtained from two study centers. More details of the studies can be found in the papers published previously ^{15, 18}. In addition, muscle biopsies were obtained after 6 months of resistance-type exercise training for 38 healthy and 24 frail older subjects. All muscle biopsies and blood samples were obtained in the morning after an overnight fast. Subjects ate a standardized meal the evening before sample collection. Subjects refrained from doing any strenuous physical activity for 3 days prior to muscle biopsy collection.

Fried criteria ²⁰ were used to assess the frailty in older subjects, which takes into account unintentional weight loss, weakness, self-reported exhaustion, slow walking speed, and low physical activity. Based on the above mentioned criteria, the healthy older subjects were not considered frail or pre-frail at the start of the intervention study ²¹.

In our study population, all older subjects regardless of their health status (frail or healthy) improved in muscle performance following 6 months of resistance-type exercise training (Supporting Information Figure S1 Table S2), as illustrated by significantly increased leg extension and leg press strength after training (*P*-value < 0.01)^{9, 15, 21}. Both healthy older and frail older subjects followed similar progressive full-body resistance type exercise training. In brief, the training consisted of a 5 minute warm-up on a cycle ergometer, followed by 4 sets on the leg-press and leg-extension machines. In addition, 3 sets of chest press, lat pulldown, pec-dec and vertical row machines were performed (Technogym, Rotterdam, The Netherlands). The healthy older subjects trained 3 times per week and frail subjects trained 2 times per week. Moreover, subjects received a protein or control supplement during the study.

Healthy older subjects took 15 gram milk protein or control drink at breakfast and frail older subjects took a similar 15 gram drink (milk protein or control) at breakfast and lunch every day throughout the entire 6 months intervention. Full details can be found in the earlier papers on the phenotypical impact of training on our study population ^{15, 21}. All studies were approved by The Medical Ethics Committee of either Wageningen University or Maastricht University and comply with the Declaration of Helsinki.

Metabolomics analysis of circulating metabolites

Amino acids and biogenic amines were derivatized (Acc-Tag) in 5 µL aliquots of plasma. Samples were analyzed using an ACQUITY UPLC system with autosampler (Waters, Etten-Leur, The Netherlands) coupled with a Xevo Tandem quadrupole mass spectrometer (Waters) operated using QuanLynx data acquisition software (version 4.1; Waters). An Accq-Tag Ultra column (Waters) was used. The Xevo TQ was used in the positive-ion electrospray mode and all analytes were monitored in Multiple Reaction Monitoring (MRM) using nominal mass resolution. Acquired data were evaluated using TargetLynx software (Waters), by integration of assigned MRM peaks and normalization using proper internal standards²²,

Acylcarnitines, trimethylamine-N-oxide, choline, betaine, deoxycarnitine and carnitine were analyzed in 5 μ L plasma, spiked with an internal standard, using a UPLC-MS/MS. Also here an Accq-Tag Ultra column was used. The Xevo TQ was used in the positive-ion electrospray mode and all analytes were monitored in MRM using nominal mass resolution. In-house developed algorithms ²³ were applied using the pooled QC samples to compensate for shifts in the sensitivity of the mass spectrometer over the batch.

Organic acids were measured by GC-MS using 50 µL of plasma sample prepared using a twostep derivatization procedure with subsequent oximation using methoxyamine hydrochloride (MeOX) and silylation using N-Methyl-N-(trimethylsilyl)trifluoroacetamide (MSTFA. Samples were measured on an Agilent GC (7890A) coupled to Agilent Quadrupole-MS with

EI source (Agilent MSD 5975C). Separation was performed using a HP-5MS column (30 m x 0.25 m x 0.25 μ m; Agilent). The raw data were pre-processed using Agilent MassHunter Quantitative Analysis software for GC-MS (Agilent, Version B.04.00), and quantitation of metabolite response was calculated as the peak area ratios of the target analyte to the respective internal standard. In-house developed algorithms were applied using the pooled QC samples to compensate for shifts in the sensitivity of the mass spectrometer over the batch. Serum metabolite concentrations determined by NMR were measured as described by Mihaleva et al ²⁴. In short, serum samples were ultrafiltrated and automated quantum mechanical line shape fitting of ¹H NMR spectra was performed using PERCH.

In tissue metabolome analysis

Metabolites were extracted from 10 mg of wet muscle tissue. This tissue was further lyophilized and weighted to determine the dry tissue mass. After pulverizing the tissue, metabolites were extracted using methanol/chloroform/water (MCW). The extraction method used in this study has extensively been described and characterized elsewhere ¹⁴.

Amines, acylcarnitines and oxylipins were measured using the platforms also used for measurement of these metabolites in plasma. The validation of these methods for human tissue biopsies is described in detail elsewhere ²³. In-house developed algorithms were applied to compensate for shifts in the sensitivity of the mass spectrometer over multiple batches of measurements ²³. The metabolite response was determined by the peak area ratio of the target analyte to the appropriate internal standard. These response ratios were used in the subsequent data analysis. ATP, ADP, creatine and phosphocreatine were determined using established enzymatic assays. For the first 3 metabolites, commercially available fluorimetric assay kits were used (BioVision cat.# K354-100, K355-100 and K635-100) following the manufacturer's instructions. Phosphocreatine was measured according to a colorimetric assay

kit protocol described by Szas et al.²⁵. In total 96 metabolites including amine, acylcarnitines, organic acids, oxylipins and a number of nucleotides were measured.

Statistical analysis

Statistical analysis was performed on log-transformed data. We used analysis of variance (ANOVA) for between group comparisons at baseline. *P*-value <0.05 was considered statistically significant. We used linear mixed models for assessment of the training effect. Our model included exercise training, subject, sex, protein supplementation and within subject correlation. Analyses of the training effect was performed separately for frail and healthy older subjects due to differences in training frequency. To summarize acylcarnitines into one single metabolite (eigenmetabolite), the Non-linear Iterative partial least squares (NIPALS) algorithm ²⁶ of the mixOmics R library was used to calculate the Singular Value Decomposition (SVD) of acylcarnitines. To integrate metabolomics and transcriptomics data the multilevel sparse partial least squares (sPLS) module from the mixOmics R library was used ²⁷. We applied canonical correlation (cut-off => 0.8), which highlights the strongest correlations between the two data sets. FactoMineR was used to perform principal component analysis (PCA) ²⁸. All analyses were done using R (version 3.02).

ASCA (ANOVA Simultaneous Component Analysis) was performed to determine global differences on metabolites. ASCA is a multivariate method that partitions variation in the data and enables to interpret these partitions by simultaneous component analysis ²⁹. Analysis was performed separately for the frail and healthy older subjects, using delta values of each metabolite for each individual (value after training - value before training) with supplement, sex and their interaction in the model as factors. Analysis was done under Matlab (version R2012a).

RESULTS

Baseline comparisons between healthy older, frail older and young subjects

Comparison of the plasma metabolite profiles of young and healthy older subjects showed marked differences in levels of many metabolites between the groups. Analysis of variance (ANOVA) showed that differences in plasma metabolite levels between older and young subjects were mainly accounted for by the amino acids and acylcarnitines. These differences were in line with previous observations of age-related effects on the plasma metabolome ^{10, 30, 31}.

Figure 1 shows a principal component analysis (PCA) plot of the biogenic amine profiles of the muscle biopsies obtained from young, healthy older and frail older subjects, before and after exercise. We observed clear age-related differences in the biogenic amine profiles of skeletal muscle, as well as a difference between healthy and frail older subjects. The PCA plot also revealed an effect of prolonged resistance-type exercise training on amine concentrations in muscle tissue. PCA revealed a similar effect for organic acids in muscle (Supporting Information Figure S2).

ANOVA yielded a series of skeletal muscle metabolites that differ significantly between healthy older subjects and young subjects. Many of these metabolites are amino acids and organic acids (Table 2). The outcome of ANOVA modeling of muscle metabolites in healthy vs. frail older subjects is presented in Table 3.

Effect of prolonged resistance-type exercise training in frail and healthy older subjects

In the PCA plot in Figure 1 we can observe that prolonged resistance-type exercise training has an effect on the biogenic amine profile of skeletal muscle tissue of frail and healthy older subjects. The trends in the PCA plot suggest that upon training both healthy and frail older subjects shift towards a younger phenotype (see also the plot for the average value of isoleucine in Supporting information Figure S3,a). The variation in this PCA plot may not

only be due to phenotype (young, healthy and frail old) and exercise, but also to sex and protein supplementation. Accordingly, we performed ANOVA Simultaneous Component Analysis (ASCA) to account for these different sources of biological variation. We observed that prolonged resistance-type exercise training had a weak effect on muscle tissue metabolite levels. No significant interaction of protein supplementation with training in both frail and healthy older subjects could be observed. Interaction of sex with training was not significant in healthy older subjects, but was significant in frail older subjects (*P*-value=0.03).

Next we used linear mixed models on univariate metabolite levels to account for phenotype, exercise, protein supplementation and sex as sources of biological variation. We observed that prolonged resistance-type exercise training affected muscle levels of acylcarnitines in both the healthy older and frail older subjects (Table 4 and 5, respectively). These include acylcarnitines that are produced during oxidation of branched chain amino acids (propionyl (C3), methylmalonyl (C4-DC), and isovaleryl (C5) acylcarnitines) and acylcarnitines that are produced during fatty acid oxidation (C6-C20 acylcarnitines) ³². Because the exercise effect on several acylcarnitines was nearly significant (*P*-value >0.05), we used the singular value decomposition to summarize the levels of fatty acid derived and amino acid derived acylcarnitines (denoted as FAAC and AAAC, respectively in Figure 4 and 5). Subsequently, we also performed linear mixed model analysis to determine the effect of training on the fatty acid derived and amino acid derived acylcarnitines levels were significantly decreased after training in both healthy and frail older subjects (see example of the effects for propionylcarnitine in Supporting information Figure S3,b).

We used multilevel sPLS to integrate changes in the muscle transcriptome and muscle metabolome after training. Here the goal was to investigate the interaction of two matched datasets and the selection of subsets of either positively or negatively correlated variables across all subjects. Only those genes were selected that were significantly changed by training

(FDR<0.05) in both frail and healthy older subjects. The resulting correlation network in Figure 2 shows that especially branched chain amino acids correlate with genes related to connective tissue/extracellular matrix such as collagen, laminin and Secreted Protein, Acidic, Cysteine-Rich (SPARC).

Correlation between muscle and plasma metabolites levels

Next we investigated to what extent baseline plasma metabolite levels reflect muscle metabolite levels. To that end we first constructed separate correlation heatmaps for muscle and plasma metabolites within each compartment. Figure 3a-b shows that metabolites that belong to the same group of metabolites (e.g. amino acids, acylcarnitines) are correlated to each other, which is observed for both plasma and muscle. However, the heatmap of the correlation between muscle and plasma metabolites (Figure 3c) showed only minor to moderate correlations (Pearson correlation between 0.3 and 0.5). The correlation networks (Figure 3d) revealed the strongest correlations for 3-hydroxybutyric acid, 4-hydroxyproline, proline, branched chain amino acids and several acylcarnitines. Correlations between serum and muscle metabolites are presented in Supporting Information Figure S4. The results are generally similar to the correlations observed between plasma and muscle metabolites.

DISCUSSION

Comprehensive metabolic profiling of muscle biopsies: differences with age and effect of resistance-type exercise training in a heterogeneous study population

Comprehensive metabolic profiling of muscle biopsies revealed pronounced differences between the muscle biopsy metabolomes of healthy young, healthy older and frail older subjects. Furthermore, distinct effects of prolonged resistance-type exercise training could be observed in both groups of older subjects..

A key asset of our analysis was the deployment of five robust profiling platforms using a single and thoroughly validated muscle-biopsy extraction procedure ¹⁴. The analytical variation in these profiles was around 20-30%, which was smaller than the biological variation between the muscle biopsies. In order to separate the sources of biological variation in our heterogeneous study population, we used univariate linear mixed models. Interestingly, the metabolic effects of supplementation and differences between males and females were relatively minor compared to the differences between old and young and between healthy and frail. The effects were also minor compared to the response to prolonged resistance-type exercise training. A multivariate approach (ASCA) did not show any significant effects of training, probably because not all sources of biological variation could be adequately accounted for ³³. In the following discussion we focus on the univariate linear mixed model approach since it allows us to more precisely account for protein supplementation and sex in our study population.

Baseline comparisons

Interestingly, in muscle biopsies baseline levels of metabolites from the tricarboxylic acid cycle (succinic acid, fumaric acid, 2-ketoglutaric acid) were lower in the healthy older subjects compared to young subjects. These differences were accompanied by lower levels of ATP, ADP, branched chain amino acids and acylcarnitines in the healthy older subjects, suggesting impaired mitochondrial function or a lower number of mitochondria in the muscle of the older subjects, which in turn may be a consequence of the lower habitual physical activity of the older subjects. These observations are in line with data on the muscle transcriptomic profiles of these subjects, which show decreased expression of genes related to mitochondrial function and oxidative phosphorylation in the older subjects compared to young subjects, with the lowest expression observed in the frail older subjects ⁹. These

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findings are in agreement with other studies ^{10, 34-38}, which also showed a down-regulation of the mitochondrial electron-transport chain pathway in older as compared to young subjects.

One of our findings is that levels of 4-hydroxyproline and proline are lower in the healthy older subjects compared to the young subjects. Both of these amino acids have been associated with collagen turnover ^{39, 40}. Muscle levels of two precursors of proline, ornithine and arginine, are higher in the older subjects, which together with lower levels of 4-hydroxyproline and proline may be due to dysfunction of the mitochondrial ornithine aminotransferase ⁴¹, leading to accumulation of arginine and ornithine. This notion is in line with slightly lower expression of genes related to tissue remodeling, such as collagen in the older subjects ⁹.

We observed higher levels of β -isoamino butyric acid (BAIBA) in muscle tissue of healthy older subjects compared to young subjects. This metabolite has been suggested to be produced upon exercise by expression of PGC-1 α and has been proposed as a myokine stimulating browning of white adipose tissue and hepatic β -oxidation ⁴². In our study population we observed lower expression of PGC-1 α target genes in healthy older subjects compared to young subjects, and in frail older subjects compared to healthy older subjects ⁹. The activity of PGC-1 α is modulated post-translationally by NAD⁺ dependent deacetyltation by SIRT1 (Figure 4-b) ⁴³. Since NAD⁺ levels decrease with age ⁴⁴, we expect a concurrent downregulation of PGC-1 α ⁴⁵. However, lower levels of BAIBA were only observed when comparing frail to healthy older subjects, whereas BAIBA levels were higher in healthy older subjects aujects compared to young subjects. Assuming that aging lead to down-regulation of PGC-1 α activity. Thus, we cannot confirm the notion that BAIBA acts as a PGC-1 α induced myokine. A possible explanation for the discrepancy is that in our study

differences in PGC-1 α expression are caused by age and frailty-dependent processes, whereas previously described PGC-1 α -mediated effects on BAIBA were caused by acute exercise.

Two polyamines, spermine and spermidine, were found to be significantly different between the frail in comparison to the healthy older subjects. Polyamines are involved in tissue regeneration and cell proliferation and differences are associated with both exercise and muscle pathology ⁴⁶⁻⁴⁸. The genes directly involved in the polyamine pathway are however not differentially expressed between frail and healthy older subjects. Hence the observed differences in polyamine levels between the frail and healthy older subjects are more likely to be attributed to effects at the level of enzyme activity or metabolite transport. Previous studies have shown that perturbations in polyamine metabolism are associated with neuromuscular disorders ^{49, 50}. On the transcriptome level we indeed found indications of neuromuscular perturbations in the frail older subjects ⁹. In addition, increased spermine levels were recently linked to skeletal muscle atrophy ⁵¹. However, we observed lower levels of spermine in frail subjects in comparison to healthy older subjects, even though frail older subjects generally have less skeletal muscle and are likely to exhibit more extensive muscle atrophy. Ost et al. recently reported that spermidine is increased in the skeletal muscle of mice overexpressing uncoupling protein 1. The authors proposed that this might be an adaptive response to cope with the additional oxidative stress ⁵². This would suggest that also in the frail elderly the increase in spermidine is a response to reduced oxidative capacity and increase in oxidative stress.

The level of carnosine is decreased in healthy older subjects in comparison to young subjects, and in frail in comparison to healthy older subjects. Carnosine is an abundant metabolite in muscle where it plays an important role in intracellular pH buffering ⁵³. Carnosine has also been associated with chelation of metal ions and antioxidant activity ⁵⁴. Furthermore, carnosine levels are higher in type II muscle fibers compared to type I. A likely explanation

for the significantly lower levels of carnosine in healthy and frail older subjects is therefore a decrease of the muscle fiber II/I ratio with respectively age and lack of exercise⁵⁵.

Several oxylipins derived from linoleic acid (LA) and α -linoleic acid (ALA) occur at higher levels in the muscle of the healthy older subjects compared to young subjects. On the other hand, levels of metabolites derived from the Δ -6 desaturase product dihomo- γ -linoleic acid (DGLA) are lower in healthy older subjects. We postulate that due to reduced Δ -6 desaturase activity linoleic acid and α -linoleic acid accumulate in the muscle of the healthy older subjects, whereas downstream Δ -6 desaturase product dihomo- γ -linoleic acid products are depleted ⁵⁶.

Effect of prolonged resistance-type exercise training

We compared the effect of prolonged resistance-type exercise training in both healthy and frail older subjects on all measured metabolites using multilevel sPLS. There was a profound correlation between the adaptive response to training between the transcriptome and amino acids in the muscle metabolome (canonical correlations between 0.7 and 0.8). Particularly high correlations were observed between expression changes of extracellular matrix genes and amino acids. Although it is unlikely that there is a direct link between expression of these genes and levels of these metabolites, it does imply that these changes in amino acid levels are part of the adaptive response to resistance-type exercise training.

At the metabolite level, the most striking effects of resistance-type exercise training in frail and healthy older subjects were observed for the C3 (propionyl) and C5 (isovaleryl) muscle acylcarnitines which are derived from branched chain amino acids. After training, the amino acid derived acylcarnitines showed a significant decrease both in the healthy and frail older subjects, accompanied by an increase of branched chain amino acids. A likely explanation is that the flux-determining mitochondrial branched chain α -keto acid hydrogenase (BCKDH)

complex ⁵⁷ has a compromised response to prolonged resistance-type exercise training. As is schematically depicted in Figure 4-a, the BCKDH complex can respond to exercise via different mechanisms. PGC-1 α is a known activator of BCKDH, but training did not have an effect on its gene expression in our study. Exercise is known to increase NAD⁺ levels ⁴⁵ and could thus activate PGC-1 α in a post-translational manner via SIRT1. Apparently this mechanism is also not activated by training in the older subjects. These effects are specific for branched chain amino acids oxidation and no significant effects on fatty acid derived acylcarnitines were found. A decrease in branched chain amino acids oxidation may stimulate mTOR related pathways activation and protein synthesis or attenuate muscle protein breakdown^{58, 59}, which could be beneficial for older subjects. As this mechanism occurs at enzymatic level, the available metabolomics and transcriptomics data can however not confirm this hypothesis and in future studies proteomics analysis would be called for.

Correlation of plasma and muscle metabolome

The weak correlations between plasma and muscle metabolite levels indicate that plasma levels only partially reflect muscle metabolism, even though skeletal muscle is one of the largest metabolically active tissues in the human body. This seems to suggest that these metabolites are also produced by other metabolic compartments. In a recent study, lack of correlation between acylcarnitine levels in plasma and tissues in mice was attributed to differences in turnover in plasma and muscle compartments, and contribution of other compartments than muscle to plasma acylcarnitine levels ⁶⁰. The same rationale very likely also applies to amino acids involved in collagen metabolism (proline, 4-hydroxyproline), which can also be formed in bone. Adipose tissue is also a metabolically active compartment with regard to branched chain amino acids besides muscle ⁶¹, and this will likely weaken plasma-muscle level correlations. As a consequence, correlations between plasma metabolites

and muscle metabolites are too modest to support their use as direct read-outs of muscle metabolism ^{62, 63}.

Conclusion

The major differences in muscle metabolome of healthy older and young subjects relate to mitochondrial function, fiber-type composition, and tissue turnover. Similar differences were observed when comparing frail older subjects to healthy older subjects. Prolonged resistance-type exercise training resulted in a correlative adaptive response of amino acids and genes responsible for tissue remodeling. The effect of exercise on amino acid derived acylcarnitines in healthy and frail older subjects points towards decreased branched chain amino acids catabolism, likely due to attenuated activation of the flux-determining mitochondrial branched chain α -keto acid hydrogenase complex in older subjects. Only modest correlations between muscle metabolite and plasma levels were found, which prohibits the use of the latter as read-outs of muscle metabolism.

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ASSOCIATED CONTENT

Supporting Information

Figure S1. Change of leg extension 1RM after resistance type exercise training. Figure S2. PCA plot of metabolites detected by the organic acid platform in muscle tissue. Figure S3. Group means with 95 percent confidence intervals for isoleucine and propionylcarnitine. Figure S4. Pearson correlation heatmap between muscle and serum metabolites. Table S1. Characteristics of subjects of which plasma samples were studied. Table S2. Characteristics of the older subjects of which samples were obtained before and after training.

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TABLES

Table 1. Characteristics of subjects of which skeletal muscle tissue biopsies were studied

	Young	Healthy older	Frail older	
N (male / female)	30/0	47/19	25/18	
Age (years)	21.7 ± 2.5	71.7 ± 5.2	77.5 ± 8.0	
Height (m)	1.83 ± 0.06	1.72 ± 0.08	1.67 ± 0.09	
Weight (kg)	76.7 ± 11.8	75.9 ± 13.3	77.5 ± 11.1	
$BMI^{a} (kg / m^{2})$	22.6 ± 2.7	25.5 ± 3.0	27.5 ± 3.7	
Body Fat (%)	14.9 ± 4.9	24.5 ± 5.6	32.1 ± 8.8	

Data was presented as mean \pm SD. a: body mass index,

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Table 2. Muscle metabolites that are significantly different between healthy older and young males ^a

Metabolite	<i>P</i> -value	FC ^b (Older/Young)
TCA Cycle		
Succinic acid	0.02	0.76
2-ketoglutaric acid	0.03	0.76
Fumaric acid	0.04	0.82
Lactic acid	0.05	0.69
Energy		
ATP	< 0.01	0.75
ADP	0.01	0.88
Branched chain amino acids		
Valine	< 0.01	0.81
Leucine	< 0.01	0.81
Isoleucine	0.03	0.84
Acylcarnitines		
Acetylcarnitine (C2)	< 0.01	0.49
Malonylcarnitine (C3-DC)	< 0.01	0.46
Intracellular buffering		
Carnosine	< 0.01	0.7
Arginine, Proline Pathway		
Ornithine	< 0.01	1.55
Arginine	< 0.01	1.34
4-hydroxy-proline	0.01	0.69
Proline	0.02	0.84
Glycylglycine	0.05	0.87
Methionine	< 0.01	0.8
Other amino acids		
Lysine	< 0.01	1.44
Aspartic acid	< 0.01	1.45
Oxylipins		
LA (LOX)		
9-HODE	< 0.01	1.4
13-HODE	0.01	1.37
13-KODE	0.03	1.3
LA(CYP450)		
9,10-EpOME	0.04	1.39
ALA (LOX)		
9-HOTrE	0.01	1.71
DGLA (LOX)		
15S-HETrE	0.02	0.83
8-HETrE	0.03	0.85

Aminobutyric acids

β amino isobutyric acid	< 0.01	1.61
α-aminobutyric acid	0.01	0.82

a: Metabolites are presented that significantly (P-value =<0.05) differ between young and old subjects according to univariate ANOVA models. b: Fold Change

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Table 3. Muscle metabolites that are significantly different between frail and healthy older subjects.^a

Metabolite	Group	Sex	Interaction	FC ^b (Frail/Healthy)
TCA Cycle				
Citric acid	<0.01	NS ^c	NS	0.54
Acylcarnitines				
Isovalerylcarnitine (C5)	< 0.01	NS	NS	0.42
Octenoylcarnitine (C8)	0.03	0.03	NS	0.77
Malonylcarnitine(C3-DC)	0.02	NS	NS	0.77
Carnitine (C0)	0.01	NS	NS	0.75
Intracellular buffering				
Carnosine	0.01	NS	NS	0.8
Oxylipins				
LA (CYP450)				
12,13DiHOME	0.04	NS	0.03	1.18
DGLA (LOX)				
8HETrE	0.03	NS	NS	0.81
15SHETrE	< 0.01	NS	NS	0.77
Polyamine metabolism				
Spermidine	0.01	0.02	NS	1.24
Spermine	0.04	NS	NS	0.9
Other amino acids				
Histidine	<0.01	NS	NS	0.79
Asparagine	0.01	NS	NS	0.81
Taurine	0.01	NS	NS	0.79
Serine	0.01	NS	NS	0.86
Glycine	0.02	NS	NS	0.81
oacetylserine	0.02	NS	NS	0.9
Homoserine	0.02	NS	NS	0.85
Tyrosine	0.02	NS	NS	0.83
Tryptophan	0.02	0.04	NS	0.83
Methionine	0.02	NS	NS	0.83
Glutamine	0.03	NS	NS	0.82
Pyroglutamic acid	0.03	NS	NS	0.83

Glutamic acid	0.04	NS	NS	0.82
Glycylglycine	<0.01	NS	NS	0.77
Aminobutyric acids				
β amino isobutyric acid	0.05	NS	NS	0.79

a: Metabolites are presented that significantly (P-value =<0.05) differ between frail and healthy older subjects according to a univariate ANOVA models that accounted for Group effect (Frail vs. Healthy), Sex effect and Group-Sex interaction; Significance of Group (Frail vs. Healthy older subjects) effects and Group-Sex interactions has been indicated; b :fold change; c: not significant

Table 4. Training effect on muscle metabolites in healthy older subjects^a

Metabolite	Training	FC(post/pre training) ^b
Pipecolic acid	0.002	1.64
Isovalerylcarnitine (C5)	0.005	0.56
Linoleylcarnitine (C18:2)	0.01	0.61
Oleylcarnitine (C18:1)	0.01	0.7
Propionylcarnitine (C3)	0.01	0.73
Palmitoylcarnitine (C16)	0.02	0.75
11.12.EpETrE	0.03	1.26
Tetradecenoylcarnitine (C14:1	0.03	0.47
AAAC ^c	0.02	0.77
FAAC ^d	NS	0.96

a: Metabolites are presented that significantly (*P*-value =<0.05) differ pre- and post-training in healthy older subjects according to univariate linear mixed models. We note that we constructed also linear mixed models that account for other factors and their interactions, but these were not found to be significant b: fold change, c,d: Single Value Decomposition (SVD) were calculated for amino acid and fatty acid acylcarnitines, denoted as AAAC and FAAC, respectively.

Metabolite	Training	Supplement	Sex	Interaction	FC(post/pre training) ^b
Propionylcarnitine (C3)	< 0.01	NS ^c	NS	NS	0.75
Glucose	< 0.01	NS	< 0.01	NS	1.35
Lactic acid	0.01	NS	< 0.01	NS	1.55
Tetradecenoylcarnitine (C14:1)	0.03	NS	NS	NS	2.00
Methionine	0.04	< 0.01	NS	NS	1.22
Tryptophan	0.04	NS	NS	NS	1.21
β Alanine	0.05	NS	NS	0.01	0.92
Isoleucine	0.05	< 0.01	0.02	NS	1.21
Myristoylcarnitine (C14)	0.05	NS	NS	NS	1.73
AAAC ^d	0.01	NS	NS	NS	0.76
FAAC ^e	NS	NS	NS	NS	1.95

Table 5. Training effect on muscle metabolites in frail older subjects ^a

a: Metabolites are presented that significantly (*P*-value =<0.05) differ pre- and post-training in frail older subjects according to univariate linear mixed models that account for Supplement, Sex and (Training and Supplement) Interaction. We note that also linear mixed models were constructed that account for other interactions, but these were not found to be significant; b: fold change, c: Not Significant, d & e: Single Value Decomposition (SVD) were calculated for amino acid and fatty acid acylcarnitines (AAAC and FAAC, respectively).

FIGURE LEGENDS

Figure 1. Principal component analysis (PCA) plot of biogenic amines detected in muscle biopsies. To visualize whether groups are significantly different from each other, confidence ellipses (95% Confidence Interval) were drawn around them.

Figure 2. Correlation network of muscle metabolites and genes. Only significantly changed genes were selected (FDR<0.05). Metabolite canonical correlation cutoff <= 0.80. Circle: gene, rectangle: metabolite, Green : positive correlation.

Figure 3. Pearson correlation heatmap of muscle to muscle metabolites (a), plasma to plasma metabolites (b), muscle to plasma metabolites (c). (d) Pearson correlation network of the most strongly correlated muscle and plasma metabolites. Red and blue indicate positive and negative correlations, respectively. Thick lines: correlation ~0.5, thin lines: 0.3 <correlation <0.5. Pink nodes: muscle tissue (T) metabolites and yellow nodes: plasma (P) metabolites,

Figure 4. (a) Schematic representation of mitochondrial oxidation of free fatty acids (FFA) and branched chain amino acids (BCAA). Arrows indicate effect of prolonged resistance-type exercise training on older subjects (healthy and frail) on BCAA (increase) and acylcarnitines (decrease) as well as the proposed (dashed-arrows) downregulation of the branched chain α-keto acid hydrogenase (BCKDH) complex. (b) Schematic representation of age-related NAD⁺ dependent acetylation of PGC1α. Dashed arrows indicate NAD⁺ and SIRT1 dependent downregulation of PGC-1α and β-isoamino butyric acid (BAIBA).

Figure 1.





Figure 3.



.

Muscle metabolites

Glycolic acid (P)

L-4-hydroxy-pro

L-4-hydroxypro

L-Proline (T)

(P)

L-proline (P)

3-Hydroxybutyric acid (P)

3-Hydroxybutyric acid_1 (T)

FA (18:1) (P)

m

Amines

Acylcarnitines Organic acids

L-lysine (P)

L-homoserine (P)

Acetylcarnitine (P)

(P)

Fatty acids

Oxylipins

(c)

Plasma metabolites

L-carnosine (T)

L-valine (P)

2-hydroxybutyric acid (P)

Butyrylcamitine (T)

L-Methionine (T)

L-leucine (P)

L-alpha-

Lactic acid (T)

L-isoleucine (P)

nobutyric

(P)

Malonylcamitine (T)

Glyceric acid (P)

Acetylcarnitine (T)



- 49 50 51 52 53 54 55 56
- 57 58
- 59 60









