Derangements of amino acids in cachetic skeletal muscle are caused by mitochondrial dysfunction

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Abstract

**Background** Cachexia is the direct cause of at least 20% of cancer-associated deaths. Muscle wasting in skeletal muscle results in weakness, immobility, and death secondary to impaired respiratory muscle function. Muscle proteins are massively degraded in cachexia; nevertheless, the molecular mechanisms related to this process are poorly understood. Previous studies have reported conflicting results regarding the amino acid abundances in cachectic skeletal muscle tissues. There is a clear need to identify the molecular processes of muscle metabolism in the context of cachexia, especially how different types of molecules are involved in the muscle wasting process.

**Methods** New in situ -omics techniques were used to produce a more comprehensive picture of amino acid metabolism in cachectic muscles by determining the quantities of amino acids, proteins, and cellular metabolites. Using matrix-assisted laser desorption/ionization (MALDI) mass spectrometry imaging, we determined the in situ concentrations of amino acids and proteins, as well as energy and other cellular metabolites, in skeletal muscle tissues from genetic mouse cancer models (n = 21) and from patients with cancer (n = 6). Combined results from three individual MALDI mass spectrometry imaging methods were obtained and interpreted. Immunohistochemistry staining for mitochondrial proteins and myosin heavy chain expression, digital image analysis, and transmission electron microscopy complemented the MALDI mass spectrometry imaging results.

**Results** Metabolic derangements in cachectic mouse muscle tissues were detected, with significantly increased quantities of lysine, arginine, proline, and tyrosine (P = 0.0037, P = 0.0048, P = 0.0430, and P = 0.0357, respectively) and significantly reduced quantities of glutamate and aspartate (P = 0.0008 and P = 0.0124). Human skeletal muscle tissues revealed similar tendencies. A majority of altered amino acids were released by the breakdown of proteins involved in oxidative phosphorylation. Decreased energy charge was observed in cachectic muscle tissues (P = 0.0101), which was related to the breakdown of specific proteins. Additionally, expression of the cationic amino acid transporter CAT1 was significantly decreased in the mitochondria of cachectic mouse muscles (P = 0.0133); this decrease may play an important role in the alterations of cationic amino acid metabolism and decreased quantity of glutamate observed in cachexia.

**Conclusions** Our results suggest that mitochondrial dysfunction has a substantial influence on amino acid metabolism in cachectic skeletal muscles, which appears to be triggered by diminished CAT1 expression, as well as the degradation of mitochondrial proteins. These findings provide new insights into the pathobiochemistry of muscle wasting.

**Keywords** Cancer cachexia; Mitochondrial dysfunctions; Amino acids; MALDI; Mass spectrometry imaging

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Introduction

Cachexia, which is often caused by cancer, leads to massive loss of total body mass, generalized inflammation, and pronounced muscle wasting, producing a significant decrease in the quality of life. Loss of skeletal muscle results in weakness, leading to immobility and eventual death secondary to impaired respiratory muscle function. Muscle wasting is thereby one of the most devastating characteristics of cachexia in late-stage cancer, for which no specific treatments are currently available. Several metabolic changes have been associated with loss of muscle mass in cachexia, including decreased protein synthesis, increased protein degradation, and deranged amino acid metabolism.

Until now, researchers have focused on understanding the molecular mechanisms behind protein and amino acid metabolism alterations in cachexia. On the one hand, the activation of the ubiquitin-dependent proteasome pathway seemed to play a major role in protein degradation in cancer cachexia. Specific ubiquitin ligase enzymes (E3) catalyzing the movement of the ubiquitin from the E2 enzyme to the substrate and are strongly up-regulated in an animal model of cachexia. Besides, several proteasome subunits are also up-regulated in a transcriptional manner in cachexia. On the other hand, protein synthesis is influenced, for example, by the activity of mechanistic (a.k.a., mammalian) target of rapamycin complex 1, which is progressively decreased in cachectic mice. All of these molecular processes are associated with a changed amino acid turnover. Meanwhile, only a few studies have reported altered quantities of amino acids in cachectic skeletal muscle, and their results have been conflicting. Furthermore, amino acid quantities in the context of protein degradation and synthesis and energy metabolism is a still unexploited topic. Of note, these issues are of utmost importance, as amino acids and proteins are linked and associated with loss of muscle mass in cachexia, including decreased protein synthesis, increased protein degradation, and deranged amino acid metabolism.

New in situ “-omics” approaches allow the identification of a broad and comprehensive range of possible alterations in amino acid and protein metabolism to help generate a more global picture of the molecular alterations associated with muscle wasting in cachexia. In the current study, we used matrix-assisted laser desorption/ionization (MALDI) mass spectrometry imaging as a sophisticated label-free, non-targeted, in situ method of examining skeletal muscles from cachetic and non-cachetic mice and patients. This is the first time amino acids, energy and other cellular metabolites, and proteins have been simultaneously examined in situ in the context of cancer cachexia.

Determination of the three classes of molecules enabled the investigation of functional relationships between and within amino acids, energy metabolites, and proteins. It allowed the identification of specific proteins that were potentially degraded and contributing to amino acid release in cachexia. Because cachexia is characterized by increased resting energy expenditure and low energy intake, the interplay between specific amino acids and energy metabolites was examined to detect molecular alterations. The further scope of the following study was the analysis of derangements of amino acids, to identify functional relationships with proteins, and simultaneously examine energy changes in skeletal muscles related to cancer cachexia.

In mice, we found higher quantities of cationic amino acids and lower quantities of glutamate and aspartate in skeletal muscle tissues during cachexia, whereas patient tissues provide first insights in similar molecular changes. Several protein expression changes in the mitochondria of cachectic tissues were associated with alterations in energy and amino acid metabolism. To the best of our knowledge, we herein present the first in situ study describing molecular alterations in cachexia caused by functional changes in the mitochondria of skeletal muscle tissues. Furthermore, we determined that cationic amino acid transporter 1 (CAT1), a mitochondria-associated protein, is involved in this process.

Methods

Collection of tissue samples

This study was approved by the Ethics Committee of the Medical Faculty of the Technical University of Munich (4916/S), and written consent was obtained from all patients before surgery. The compound mutant mouse line (pVillin-KRASV12G x ApcM1638N), with the inbred C57BL/6N background, was bred at the Zentrum für Präklinische Forschung at Klinikum rechts der Isar, TUM, Munich. The animals were assessed at specific timepoints for the presence of intestinal tumours, as previously described, and their body weights were determined at the time of sacrifice (see Table 1 for baseline characteristics). The non-cachetic group consisted of tumour-free wild-type and single transgenic littermates from the same breedings. Cachetic mice were defined as

| Characteristic | Non-cachetic | Cachetic |
|---------------|-------------|----------|
| Number | 10 | 11 |
| Gender (n) | 9 | 5 |
| Male | 1 | 6 |
| Female | 8.7 | 6.6 |
| Range | 4–15 | 4–10 |
| Weight (g) | Mean | 31.7 |
| Range | 27.4–36.0 | 21.7–32.0 |
animals with at least a 10% reduction in body weight, compared with the median weight of the non-cachectic group. All mice were maintained with a 12 hr light-dark cycle and fed a standard diet and water ad lib. Overall, 21 mice were analysed, and 11 tumour-bearing mice were classified as cachectic. Ten mice represented the non-cachectic group. Mice were euthanized via cervical dislocation. The quadriceps muscles were subsequently resected and then immediately shock frozen and stored until further use in liquid nitrogen.

Patient tissues (samples from the musculus rectus abdominis) were obtained from patients diagnosed with pancreatic ductal adenocarcinoma (see Table 2 for baseline characteristics) during routine surgery. The patients underwent surgery at the Department of Surgery, TUM, from 2008 until 2015. Samples were obtained from 36 patients, 13 of whom exhibited a weight loss of at least 7% within the 6 months before surgery and were thereby defined as cachectic patients. The remaining 23 patients, with a weight loss of less than 5%, were considered non-cachetic. Definition of cachexia was according to an international consensus. Tissues from six patients (three non-cachetic and three cachectic patients; Supporting Information, Appendix S1 for baseline characteristics) were stored in liquid nitrogen as fresh-frozen samples until MALDI mass spectrometry imaging analysis. Tissues from the remaining 30 patients were fixed in formalin and embedded in paraffin and then used to construct two-tissue microarrays, which were stored at room temperature until immunohistochemistry (IHC) analysis.

Matrix-assisted laser desorption/ionization mass spectrometry imaging

Frozen muscle tissue samples were cryosectioned into 12 μm sections using Microm560 (Microm International, Walldorf, Germany) and thaw mounted onto indium tin oxide-coated conductive slides (Bruker Daltonics, Bremen, Germany). The slides were pre-treated with 1:1 poly-L-lysine (Sigma-Aldrich, Munich, Germany) and 0.1% Nonidet P-40 (Sigma) before mounting. Briefly, the samples were covered with 10 mg/mL 9-aminoacridine (9AA) matrix (Sigma-Aldrich) in 70% methanol for metabolite analysis or 10 mg/mL 1,5-diaminonaphthalene (DAN) matrix (Sigma-Aldrich) in 70% acetonitrile for amino acid analysis, using a SunCollect sprayer (Sunchrom, Friedrichsdorf, Germany). The following preferences were used for the automatic sprayer (for both 9AA and DAN): vial distance of 0.50 mm for the X direction and 2.00 mm for the Y direction, 20 mm Z position and offset of the spray head, and medium X/Y speed. The matrix was deposited in eight layers using variable increasing spray rates, as previously reported by Ly et al. For metabolite analysis, data were acquired in negative ion mode using a Bruker Solarix 7.0 T Fourier-transform ion cyclotron resonance (FTICR) mass spectrometer (Bruker Daltonics) over a mass range of 50–1000 m/z and at a lateral resolution of 100 μm. Amino acids were analysed over a mass range of 50–250 m/z. Because DAN matrix is not stable in a vacuum, a faster measurement protocol was established by changing the lateral resolution for amino acids to 150 μm. For protein analysis, 10 mg/mL sinapinic acid matrix (Sigma-Aldrich) in 60% acetonitrile and 0.2% trifluoroacetic acid was deposited onto the sections using an ImagePrep automated sprayer (Bruker Daltonics). Linear positive ion mode over a mass range of 2000–25 000 m/z was used for protein measurements with the Bruker Ultraflex III MALDI-TOF/TOF MS (Bruker Daltonics). The lateral resolution for examining intact proteins was predefined as 60 μm. The Smartbeam-II Nd:YAG laser (355 nm) fired 300 times for protein ablation with a frequency of 200 Hz and a sample rate of 0.50 GS/s. External calibration was performed with Protein Calibration Standard I (Bruker Daltonics), mixed 1:1 (v/v) with the matrix solution, and spotted onto the slide.

Table 2  Clinical characteristics of patients with cancer

| Characteristic          | Non-cachetic | Cachectic |
|-------------------------|--------------|-----------|
| Number                  | 23           | 13        |
| Gender (n)              |              |           |
| Male                    | 15           | 4         |
| Female                  | 8            | 9         |
| Age (years)             | 67           | 65        |
| Range                   | 45–88        | 44–83     |
| Weight (kg)             | 77           | 64        |
| Range                   | 40–106       | 41–98     |
| Body mass index (kg/m²) | 25.6         | 22.5      |
| Mean                    | 16.2–34.2    | 15.8–28.4 |
| Weight loss (%)         | 1.0          | 13.3      |
| Range                   | 0–4.8        | 7.7–25.5  |

Tissue staining and co-registration

After acquisition of the mass spectrometry data, the matrix was removed with 70% ethanol, and the tissue sections were stained with haematoxylin and eosin. The sections were transferred to dH₂O for 1 min and then haematoxylin (Carl Roth, Karlsruhe, Germany) for 1 min; after washing in tap water for approximately 5 min, they were transferred to eosin Y (Sigma-Aldrich) for 1 min. The sections were subsequently dehydrated using an increasing alcohol solution series [70%, 90%, and 100% ethanol (Merck) and isopropanol (Merck); 30 s each], transferred to xylene (Carl Roth) for at least 2 min, coverslipped, scanned with a Mirax desk slide scanner (Zeiss, Göttingen, Germany) using a 20× magnification.
objective, and co-registered with the respective mass spectrometry imaging data using fleximaging v. 4.0 (Bruker Daltonics).

Matrix-assisted laser desorption/ionization mass spectrometry imaging data analysis

fleximaging software was used for normalization against the root mean square (FTICR) or total ion count [time of flight (TOF)] of all data points. “Virtual microdissection” was performed, defining regions of interest (ROIs) of cross-sectioned muscle fibres and excluding non-muscle tissue components. Specification of ROIs and exportation of spectral data for each ROI were similarly performed using fleximaging.

MATLAB script, including the bioinformatics and image processing toolboxes (v.7.10.0, MathWorks, Natick, MA, USA), was used for subsequent FTICR data processing. In this script, fleximaging-exported spectra were processed using the LIMPIC algorithm. This algorithm includes a baseline subtraction (100 data points window size), resampling (0.001 m/z bin width), and smoothing (Kaiser filter with factor 3) to exclude chemical and electronic noise before peak picking. Peak picking itself was performed with a minimum peak width of 5.0E – 4 m/z, a signal-to-noise threshold of 2, and a minimum intensity threshold of 0.01% for the base peak of each spectrum. To enable direct comparisons between peak lists from different patients, peaks were clustered with a specific mass tolerance (5.0E – 8 * (m/z)^2, TOF peak lists were generated using SCiLS Lab (v2019a, Bremen, Germany). Metabolites, amino acids, and proteins were identified by matching accurate mass data with Metlin (http://metlin.scripps.edu/index.php), the Human Metabolome Database (http://www.hmdb.ca), and the MaTisse database, allowing a window of 4 ppm for metabolites or amino acids.

Ion images were exported by fleximaging as single colour visualizations with a black background. The colour was detected entirely by image processing software and was visualized over the corresponding scanned haematoxylin and eosin-stained tissue. Adenosine monophosphate (AMP), adenosine diphosphate (ADP), adenosine triphosphate (ATP), and energy charge visualization was performed by an in-house developed python script based on the previously published MSIdV tool for mass spectrometry imaging. The remaining heatmap visualizations were received from SCiLS Lab.

Transmission electron microscopy

For transmission electron microscopy (TEM) analysis, fresh-frozen tissues were fixed with 2.5% glutaraldehyde in 0.1 M sodium cacodylate buffer, pH 7.4, and TEM fixation buffer (Electron Microscopy Sciences, Hatfield, USA) at 4 °C. The samples were subsequently post-fixed in 2% aqueous osmium tetraoxide 44, dehydrated in increasing concentrations of ethanol (30–100%) and propylene oxide, embedded in Epon (Merck, Darmstadt, Germany), and dried for 24 h at 60 °C. Semithin sections were cut and stained with toluidine blue. Ultrathin sections of 50 nm were collected on 200-mesh copper grids and stained with uranyl acetate and lead citrate before examination by TEM (Zeiss Libra 120 Plus, Carl Zeiss NTS GmbH, Oberkochen, Germany). Images were acquired using a Slow Scan CCD camera and iTEM software (Olympus Soft Imaging Solutions, Münster, Germany). The number of mitochondria was determined by manually counting the mitochondria per field of view in longitudinally sectioned fibres, viewed at 1600x magnification.

Immunohistochemistry staining and digital image analysis

For IHC analyses, parts of the mouse muscle tissues were fixed in 4% (v/v) neutral-buffered formalin and embedded in paraffin by an automatic processor. CAT1 analysis of patient tissues was performed on the formalin-fixed paraffin-embedded tissues. Expression levels of CAT1 and voltage-dependent anion channel (VDAC) were analysed on 3 μm consecutive sections by IHC staining with the CAT1 antibody [14195-1-AP (Proteintech, Manchester, UK), 1:20] and VDAC antibody [#4866 (Cell Signaling Technology, Frankfurt am Main, Germany), 1:100]. For validation purposes of mitochondrial proteins, cytochrome c oxidase subunit (COX) 7C [PA5-51284 (Invitrogen, Carlsbad, CA, USA), 1:10] and cytochrome c [GTX108585 (GeneTex, Inc., Alton Pkwy Irvine, CA, USA), 1:25] stainings were performed on 12 μm cryosectioned mouse skeletal muscle tissues. Used tissues are representing consecutive tissue sections as analysed in MALDI imaging. The automated slide processing system Ventana DISCOVERY XT System (Ventana Medical Systems, Inc., Tucson, AZ, USA) was used in accordance with the manufacturer’s instructions. All stained slides were digitalized at 20x objective magnification using the Mirax desk slide scanner. IHC staining results were quantified using the image analysis software Definiens Developer XD2 (Definiens AG, Munich, Germany). This software allows quantification of IHC staining intensities within a user-specified ROI. ROIs were defined in longitudinally sectioned muscle fibres. Algorithms were developed, modified specifically for CAT1, VDAC, COX7C, and cytochrome c, and established in semantic and context-based segmentation processes, which included staining intensity, shape, area, colour features, and neighbourhood. In addition, the algorithms were optimized to consider only mitochondrial staining. The quantified parameters for CAT1, VDAC, COX7C, and cytochrome c were values representing a point on a continuous spectrum of protein expression in relative units.
**Immunofluorescence analysis of myosin heavy chain expression**

Immunofluorescence analysis of myosin heavy chain (MHC) expression was achieved using primary antibodies against MHC-I (BA-F8, 1:50), MHC-Ila (SC-71, 1:600), and MHC-Ilb (BF-F3, 1:100) according to Bergmeister et al. Primary antibodies were purchased from the Developmental Hybridoma Bank (University of Iowa), whereas secondary antibodies (AF633 IgG2b, AF488 IgG1, and AF555 IgM, all 1:250) were purchased from Invitrogen. Slides were digitalized at 20× objective magnification using an Axio Scan.Z1 (Zeiss). For fibre-type analysis, all fibres within an ROI (as described for MALDI mass spectrometry imaging) were characterized. Quantification was performed by digital image analysis in Definiens Developer XD2.

**Statistical analysis**

To determine significant differences between cachectic and non-cachectic tissues, the rank-based Mann–Whitney U test was used. Further statistical testing for functional relationships between and within metabolites, amino acids, and proteins was investigated using Spearman’s rank correlation analysis. Correlation plots and calculations were generated and performed using R (The R Foundation, Vienna, Austria; “corrplot” package). P values equal to or less than 0.05 were considered statistically significant.

**Results**

**Amino acid derangements were detected by in situ metabolomics in cachexia**

Sixteen amino acids were detected in situ in mouse skeletal muscle tissues. Lysine ($P = 0.0037$), arginine ($P = 0.0048$), tyrosine ($P = 0.0357$), and proline ($P = 0.0430$) were significantly more abundant in cachetic mouse muscle tissues than in non-cachetic mice. Alanine, asparagine, glutamine, leucine/isoleucine, methionine, phenylalanine, threonine, tryptophan, and valine analysis revealed a higher relative mean intensity in cachetic mice than in non-cachetic ones, but the differences were not statistically significant (Figure 1A). In contrast, quantities of glutamate ($P = 0.0008$) and aspartate ($P = 0.0124$) were significantly less in cachetic mouse skeletal muscle tissues. All intensities and P values of determined amino acids in non-cachetic and cachetic mice can be obtained from Supporting Information, Appendix S2. Figure 1B shows the results of false colour visualization of amino acids, demonstrating the differences in skeletal muscle tissues between the cachetic and non-cachetic mice.

Results in skeletal muscle tissues of cachectic patients revealed similar amino acid profiles to those observed in mice (Supporting Information, Appendix S3); however, the amino acid differences between cachectic and non-cachectic patients did not reach statistical significance. As the amino acid alterations could be due to protein breakdown processes, our subsequent investigations examined this issue.

**Amino acids are linked to protein breakdown processes**

To detect possible protein breakdown targets, protein expression was determined by MALDI-TOF and correlated with the quantities of amino acids. A negative correlation between amino acids and proteins suggests the presence of protein breakdown.

We determined the correlations between all detected proteins and the quantities of amino acids. Spearman’s rank correlation analysis revealed 15 annotated proteins, which have significant negative correlations with amino acid quantities (Supporting Information, Appendix S4). The proteins for which significant correlations were detected included thymosin beta-4, individual proteins for oxidative phosphorylation (OXPHOS), histone H2B, and glutathione S-transferase P. Specifically, thymosin beta-4 (Figure 2A) correlated negatively with the quantity of lysine ($P = 0.0090$), arginine ($P = 0.0026$), proline ($P = 0.0049$), and threonine ($P = 0.0431$). Cytochrome c oxidase subunit (COX) 6B1 correlated negatively with lysine ($P = 0.0405$), arginine ($P = 0.0027$), glutamine ($P = 0.0209$), and tryptophan ($P = 0.0368$). Figure 2B shows an example of the detected proteins: false colour visualization demonstrated lower expression of COX6B1 in cachetic muscle tissues than in non-cachetic mice. Statistical analysis of the visualization results showed significantly decreased COX6B1 in cachexia (Figure 2C; $P = 0.0008$). Supporting Information, Appendix S5 shows the analyses for three OXPHOS proteins: COX7C, ATPase F6, and cytochrome c. At this, the abundances of COX7C and ATPase F6 were significantly decreased ($P = 0.0127$ and $P = 0.0048$, respectively), determined by MALDI mass spectrometry imaging. Interestingly, a majority of detected proteins that correlated negatively with amino acids were associated with the mitochondria. In addition, validation experiments were performed by IHC stainings and confirmed MALDI mass spectrometry imaging analysis results for COX7C as well as for cytochrome c (Supporting Information, Appendix S5). COX7C abundance detected by IHC was also significantly decreased ($P = 0.0159$) in cachetic mouse skeletal muscle tissues in comparison with non-cachetic mice.

At this stage, the combination of two different measurement techniques—high-resolution mass spectrometry by FTICR and analysis of proteins by TOF—enabled the detection of biological relationships and thus, for example, the
Figure 1  *In situ* analysis of amino acids in cachectic mouse skeletal muscle tissues. (A) Quantities of amino acids in skeletal muscles from cachectic and non-cachectic mice. All intensity values were determined by mass spectrometry imaging. Quantities of lysine, arginine, proline, and tyrosine were significantly higher in cachectic mice than in non-cachectic ones. The intensities for alanine, asparagine, glutamine, leucine/isoleucine, methionine, phenylalanine, threonine, tryptophan, and valine also revealed a higher relative mean intensity in cachectic mice than in non-cachectic ones. Glutamate and aspartate intensities were significantly decreased in cachectic mouse muscles. Whiskers of the boxplots illustrate the minimal and maximum intensity values. (B) False colour visualization of amino acids in mouse skeletal muscle tissues. Lysine, arginine, and proline were increased, and glutamate and aspartate were decreased in cachectic mouse muscles, compared with non-cachectic ones. *P* < 0.05, **P** < 0.01, and ***P** < 0.005. H&E, haematoxylin and eosin.
potential degradation of OXPHOS proteins. Four of these proteins were highlighted by correlation analysis and seemed to have enhanced breakdown in cachexia. The involvement of OXPHOS proteins suggested the possibility of alterations in energy in cachexia and thereby focused our further investigations on the mitochondria.

**Energy charge is reduced in cachectic muscle tissues**

Because the correlation between amino acid and protein data suggested degradation of specific OXPHOS enzymes, we conducted metabolite measurements to examine energy alterations in cachectic mouse skeletal muscle tissues. Because the quadriceps muscle is composed of mixed fibre types, we examined non-cachectic and cachectic mouse skeletal muscle tissues for fibre-type composition. This issue is important for regarding energy aspects, due to the adaption of specific fibre types to oxidative or glycolytic metabolism. No differences could be detected regarding the individual fibre type composition between non-cachectic and cachectic mouse skeletal muscle tissues (Supporting Information, *Appendix S6*).

Adenosine monophosphate, ADP, and ATP concentrations were determined as representatives of the energy state in muscle tissues. To determine the available energy in tissues, the energy charge was calculated from the AMP, ADP, and ATP concentrations, using the formula shown in *Figure 3A*. The energy charge was lower in cachectic mice than in non-cachectic mice (*P* = 0.0101), indicating that energy stores were reduced in cachectic mouse skeletal muscle tissues. AMP was slightly increased in the cachectic mouse skeletal muscle tissues, when compared with non-cachectic muscle tissues (*Figure 3B*). ADP concentrations were similar in cachectic and non-cachectic mice, and ATP concentration was marginally decreased in cachexia.

Reduced energy in cachectic mouse skeletal muscle tissues could result in changed function of the tricarboxylic acid (TCA) cycle as a compensatory alteration or as a co-influencing factor. In cell metabolomics analysis, the majority of detected TCA cycle molecules did not differ significantly between cachectic and non-cachectic mouse muscle tissues (Supporting Information, *Appendix S7*). However, the quantity of oxaloacetate was significantly decreased (*P* = 0.0448) (*Figure 3C*) and the quantity of malate was significantly increased (*P* = 0.0295) in cachexia. These results indicate that cachexia was not associated with a generalized change in regulation of the TCA cycle, but it did seem to be associated with alterations of the dicarboxylic acid part.

We next used the energy charge results to examine underlying molecular mechanisms. A total of 58 proteins correlated significantly with the calculated energy charge (*Figure 4*). Specifically, COX7C (*P* = 0.0257) and COX6B1 (*P* = 0.0052) were all significantly and positively correlated with the energy charge. Thymosin beta-4 (*P* = 0.0054), histone H2B (*P* ≤ 0.0001), and redox Cu/Zn superoxide dismutase (*P* = 0.0059) also correlated positively with the energy charge. Notably, expression of ubiquitin, which is a marker for protein degradation,
correlated negatively \((P = 0.0009)\) with the energy charge, as well as with COX7C \((P = 0.0216)\) and other proteins. These proteins may be tagged for degradation.

Five amino acids were also significantly correlated with the energy charge, suggesting a functional relationship. For example, lysine \((P = 0.0122)\) and arginine \((P = 0.0086)\), the quantities of which differed significantly between cachectic and non-cachectic mouse tissues, were correlated negatively with the energy charge (Figure 4). Glutamine \((P = 0.0108)\) and leucine/isoleucine \((P = 0.0324)\) were likewise negatively correlated with the energy charge. Glutamate, which was significantly decreased in the cachectic state, correlated positively with the energy charge \((P = 0.0437)\).

**Cationic amino acid transporter 1 is changed in cachexia**

We next focused on examining changes in CAT1, which was previously identified as an essential factor for L-arginine transport into mitochondria.\(^\text{25}\) The potential role of this transport protein in cachexia was supported by our findings of prominent alterations in arginine and lysine, both of which are cationic amino acids.

The quantity of mitochondrial CAT1 in skeletal muscle tissues was lower in cachectic mice than in non-cachectic ones (Figure 5A). Using digital image analysis of stained mouse skeletal muscle tissues, significantly less CAT1 was observed in the cachectic state (Figure 5B; \(P = 0.0133)\). There was a similar tendency towards decreased CAT1 in muscle tissues of cachectic patients, compared with non-cachectic patients, but the difference did not reach statistical significance \((P = 0.5498; \text{Figure 5C})\). For comparing an equal group size of patients without gender influence, we also calculated CAT1 expression for female patients alone, which is marginally improving the statistics \((P = 0.2261)\). CAT1 expression was significantly correlated with the quantity of lysine \((P = 0.0122; \text{negative correlation})\), proline \((P = 0.0266; \text{negative correlation})\), and glutamate \((P = 0.0106; \text{positive correlation})\), suggesting an important influence of CAT1 on amino acid quantities in cachectic tissues.

The CAT1 expression results could have been influenced by a change in the number of mitochondria, as fewer mitochondria would lead to reduced quantities of mitochondrial proteins. Therefore, TEM was performed to determine the number and possible morphological alterations of mitochondria.
mitochondria in mouse skeletal muscle tissues. No differences in mitochondrial morphology were observed between mitochondria in cachectic and non-cachectic mice; we did not observe electron-lucent areas, swelling, and fragmentation of cristae, which were previously reported in the muscles of mice with cancer cachexia. Similarly, the number of mitochondria was not significantly decreased in the cachectic state (Figure 5D and 5E, \(P = 0.1051\)). Additionally, there were no systematic ultrastructural changes between cachectic and non-cachectic mice.

To provide further evidence of a comparable number of mitochondria in cachectic and non-cachectic mice, VDAC expression was determined by IHC. The quantity of VDAC correlated significantly with the number of mitochondria determined by electron microscopy (Figure 5F; \(P = 0.0033\)) and revealed no significant differences between cachectic and non-cachectic mouse skeletal muscle tissues (Figure 5G; \(P = 0.5648\)). Thus, TEM and VDAC expression confirmed the presence of a similar number of mitochondria in cachectic and non-cachectic mouse skeletal muscle tissues, thereby suggesting a functional—not quantitative—change in mitochondria in cachectic tissues. CAT1 also seemed to be essential for specific aspects of amino acid metabolism in skeletal muscle cells, with significantly lower expression in cachectic mouse skeletal muscle tissues.

**Discussion**

In this study, intact skeletal muscle tissues from a murine genetic cancer model were examined for molecular alterations associated with cancer cachexia. For the first time, a combination of three different MALDI mass spectrometry imaging methods was used to explore in situ associations between amino acids, proteins, and energy and other cell metabolites in cachectic skeletal muscle tissues. Significant amino acid alterations were detected in cachectic mice, with higher quantities of lysine, arginine, proline, and tyrosine and lower quantities of glutamate and aspartate (Figure 1A).

Changes in the quantities of glutamine, isoleucine, leucine, and valine in cachectic skeletal muscle tissues were reported in previous studies. However, the specific results were conflicting, as individual studies found lower and/or higher quantities of the same amino acids in cachectic skeletal muscle tissues. Changes of detected abundances could occur because...
Figure 5 Changes in CAT1 expression in cachexia. (A) Digital image analysis of the CAT1 immunohistochemistry (IHC) results. Definiens Software Developer XD2 was used to detect the CAT1 expression in all tissue sections. (B) The detected CAT1 abundance was significantly lower in skeletal muscles of cachectic mice than in non-cachectic ones ($P = 0.0133$). Whiskers of the boxplots represent the lowest and highest CAT1 expression in each group. (C) CAT1 expression in skeletal muscle tissues from humans with cancer for all and only female patients. CAT1 was slightly lower in patients with cachexia, compared with non-cachectic patients, but the difference did not reach statistical significance. Horizontal lines represent the mean intensity of each group. (D) Number of mitochondria in mouse skeletal muscle tissues determined by transmission electron microscopy (TEM) at 1600× magnification. The red colour represents the mitochondria. Intact mitochondria were observed in tissues from both cachectic and non-cachectic mice. (E) Boxplots illustrating the number of mitochondria in mouse skeletal muscle tissues. No significant difference was detected between cachectic and non-cachectic mice. (F) Correlation analysis between the number of mitochondria determined by electron microscopy and the intensity of voltage-dependent anion channel (VDAC) staining detected by IHC. The number of mitochondria was significantly correlated with VDAC expression ($P = 0.0033$). (G) Statistical analysis of VDAC staining in muscle tissues from cachectic and non-cachectic mice. No significant difference was detected. $P < 0.05$. 

Derangements of amino acids in cachectic skeletal muscle

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of different analytic platforms, resulting in specific sample preparations, and the individual selected skeletal muscles. Our results in quadriceps muscle tissue and MALDI analysis showed no significant changes towards a higher quantity of all of these amino acids in cachectic skeletal muscle tissues (Figure 1A). Furthermore, among all amino acids detected in our study, significant changes were especially noted with cationic amino acids and glutamate. Because lysine, arginine, and glutamate are connected by a biochemical pathway related to mitochondria,27,28 we focused on mitochondria to understand the detected amino acid alterations. Of note, mitochondrial enzymes catalyze the synthesis of glutamate by transferring the amino group from amino acids such as lysine and arginine to α-ketoglutarate.27,28 Nonetheless, lysine and arginine must first enter the mitochondria.

CAT1, a transporter molecule in mammalian mitochondria,25 carries arginine and lysine through membranes.29 The presence of CAT1 in mitochondria was confirmed by western blot analysis of isolated cardiac mitochondria and by confocal microscopy, and it seemed to be involved in cationic amino acid transport, mitochondrial stress, survival, and diminished reactive oxygen species production.25 In our study, we detected higher quantities of the cationic amino acids lysine and arginine and a lower quantity of glutamate in the muscles of cachetic mice, compared with non-cachetic ones. We also observed lower expression of CAT1 in cachetic mice. Correlation analysis confirmed significant correlations between CAT1 and lysine, as well as glutamate: low CAT1 expression was associated with more lysine and less glutamate in muscle tissues. These findings may be due to enhanced protein breakdown in cachexia,30 leading to increased levels of certain amino acids, such as arginine and lysine (Figure 6A). Reduced metabolic conversion because of diminished CAT1 expression could also account for the significantly higher quantities of arginine and lysine [Figure 6B, (1)]. Of note, our findings are consistent with the results of previous studies, which showed that arginine and lysine were increased in the gastrocnemius of cachetic mice32 and glutamate was reduced in quadriceps muscles of cachetic C26 hosts.31 Decreased production of glutamate from arginine and lysine because of reduced transamination reactions could account for the significantly lower quantity of glutamate in cachetic mouse tissues [Figure 6B, (2)]. Glutamate production could also be hampered by a low NADH/NAD⁺ ratio in mitochondria, which would favour oxidative decarboxylation of α-ketoglutarate to succinyl-CoA.34 An alteration in an electron shuttle system through the inner mitochondrial membrane, for example, could be an explanation for an altered ratio. This may lead to a lower amount of NADH inside mitochondria.

Reduced expression of CAT1 in the muscles of cachetic mice and patients could arise from several mechanisms. For instance, a lower number of mitochondria would reduce mitochondrial CAT1 expression. However, examination of the tissues with TEM (the gold standard for measuring mitochondrial content33) revealed no evidence of a reduced number or altered structural integrity of mitochondria in cachexia (Figure 5E). Likewise, examination of mitochondrial content by staining for VDAC (the most abundant protein in the outer membrane of mitochondria33) did not show significant differences between cachetic and non-cachetic mice (Figure 5G). These results indicate that decreased CAT1 expression is not the result of an altered number of mitochondria; instead, it is due to a specific decrease in expression of the CAT1 protein itself (Figure 5B). Thus, decreased mitochondrial CAT1 expression has been described for the first time in cancer cachexia, which may be highly relevant to the amino acid metabolism changes observed in this setting, as discussed earlier.

Manifestations of cachexia can differ according to the type of tumour.34 In the current study, colorectal and pancreatic cancer revealed similar molecular changes during cachexia. Lysine, arginine, and proline quantities were increased, and glutamate and aspartate were decreased in skeletal muscle tissues of both types of tumours during cachexia. Notably, human material offered only preliminary results in the current study due to the small patient number and further studies, including more patients, are necessary for confirming these molecular alterations. A recent study comparing lung carcinoma and colon adenocarcinoma models under cachectic conditions also found molecular similarities between the two types of malignancies.35 Despite different food intake in the two models, hypothalamic expression of the orexigenic neuropeptide Y was increased in both tumour types, compared with controls. At this point, our data suggest tumour type-independent molecular alterations, at least for colorectal and pancreatic cancers. Additional studies are necessary to further explore the influence of tumour type on the molecular manifestations of cachexia.

Protein breakdown is highly discussed in the context of cachexia.3 In the present study, simultaneous examination of proteins and amino acids enabled the identification of proteins that may be selectively degraded in cachetic skeletal muscle tissues. A majority of detected and degraded proteins in this study were OXPHOS proteins, which were negatively correlated with an overabundance of certain amino acids (Figure 2A), thereby suggesting the presence of a degradation process. Of note, only small undigested proteins (<25 kDa) are determined by MALDI imaging protein measurement and released amino acids from muscle cannot be set into relation with proteins and metabolites in the current study. A recent study of rat gastrocnemius muscles revealed an accumulation of oxidatively modified mitochondrial proteins in cachexia.36 Oxidative modifications can be repaired enzymatically or lead to protein degradation. Both processes are essential for maintaining homeostasis and survival of cells.37 Oxidative modifications might also initiate degradation of the OXPHOS proteins.
detected in the current study. A previous study found reduced quantity of COX6B1 protein in cachectic mouse tissues, compared with controls, which is consistent with our results (Figure 2C). COX6B1 could be altered by oxidative modification and therefore degraded in cachectic skeletal muscle tissues. This degradation could account for the lower quantity of COX6B1 in cachectic skeletal muscle tissues (Figure 2C) and the negative correlation between COX6B1 and individual amino acids (Figure 2A). Of interest, CAT1 is also involved in reducing oxidative stress. Normal CAT1 expression could help maintain normal protein signatures in mitochondria by preventing oxidative damage and thereby protecting proteins from degradation.

Mitochondria, as well as OXPHOS proteins, are associated with ATP production and, therefore, energy generation in muscle tissues. Given our findings of OXPHOS protein degradation and altered mitochondrial protein signatures during cachexia, we examined the energy status of cachectic skeletal muscle tissues. The energy charge, defined as the mole fraction of ATP plus half the mole fraction of ADP, is linearly related to the amount of temporarily stored metabolically available energy. We found that energy charge was significantly decreased in cachexia. Furthermore, correlation analysis indicated that higher energy charge was associated with a lower quantity of individual amino acids, higher expression of OXPHOS proteins, and lower expression of ubiquitin. Nonetheless, glutamate correlated positively with the energy charge (Figure 4). A reduced rate of ATP synthesis (determined by 31P nuclear magnetic resonance spectroscopy) has been previously described in cancer cachexia. Degradation of mitochondrial OXPHOS proteins could at least partially explain the reduced energy stores in cachectic muscles. As ubiquitin, a marker for protein degradation, can initiate proteasomal degradation, lower ubiquitin tagging could reduce degradation of OXPHOS proteins and the quantity of amino acids but increase the energy charge. The positive correlation between glutamate and energy charge could underlie the anaplerotic processes of glutamate in cancer cachexia.

Reduced energy availability in muscle tissues could lead to higher glycolytic activity, because two NADH and two ATP molecules are produced from one glucose molecule, and the NADH can be utilized for ATP synthesis in mitochondria. However, the inner mitochondrial membrane is impermeable to NADH, so cytosolic NADH must be transported into the mitochondrial matrix by transport systems, such as the malate-aspartate shuttle. High glycolytic activity has been described in cachexia, which could lead to high quantities of cytosolic NADH (Figure 6B, (3)). Pin et al. formally described an increased reliance on glycolytic metabolism and a
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Conflict of interest

None declared.

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Online supplementary material

Additional supporting information may be found online in the Supporting Information section at the end of the article.

Appendix S1. Advanced baseline characteristics for patients, which is separating patient material in individual fixation methods. FFPE tissues were used for immunohistochemical staining, and fresh frozen tissues were used for the determination of amino acids.

Appendix S2. Intensities of amino acids in skeletal muscle tissues of cachectic and non-cachectic mice. P values were calculated by Mann–Whitney U test.

Appendix S3. In situ amino acid quantities in skeletal muscle tissues from cachectic patients with cancer. Changes in lysine, arginine, proline, glutamate, and aspartate with cachexia in humans revealed similarities to the changes observed in cachectic mouse skeletal muscle tissues. The horizontal lines represent the mean intensity of each group.

Appendix S4. P values (Spearman’s rank correlation analysis) of significant negative correlations between proteins and amino acids. Included proteins were may degraded in skeletal muscle tissues of cachectic mice.

Appendix S5. Analysis of mitochondrial proteins in mouse skeletal muscle tissues (A) Statistical analysis for the OXPHOS-related proteins COX7C, cytochrome c, and ATPase F6 determined by MALDI mass spectrometry imaging. COX7C (P = 0.0127) and ATPase F6 (P = 0.0048) expression was significantly decreased in cachexia, compared with non-cachectic skeletal muscle tissues. Cytochrome c was also decreased in cachectic mouse skeletal muscle tissues, but the differences did not reach statistical significance (P = 0.1145). (B) Immunohistochemistry (IHC) results confirmed changes of mitochondrial proteins detected by MALDI mass spectrometry imaging. Quantification of the IHC, performed by digital image analysis, revealed a lower expression of COX7C in cachectic mouse skeletal muscle tissues (P = 0.0159) and a similar change of cytochrome c without reaching significance level (P = 0.2512). *P < 0.05, **P < 0.01.

Appendix S6. Immunofluorescence analysis of myosin heavy chain (MHC) expressions. (A) Statistical analysis of the expression of myosin heavy chains in cachetic and non-cachetic mouse skeletal muscle tissues. MHC1 expressions were in both comparison groups 0.02%. The Mann–Whitney U test was performed for all fibre types and revealed no significant change between non-cachetic and cachectic mouse comparison groups. (B) Exemplary pictures for immunofluorescence stained cross-sectioned mouse skeletal muscle tissues. The left tissue section is representing a non-cachetic mouse,

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Reduced flux of glycolysis-derived pyruvate into the TCA cycle, which might also be followed by reducing energy in cachectic skeletal muscle tissues. Malate dehydrogenase 1 enzyme can utilize NADH to reduce oxaloacetate to malate for transporting electrons from NADH across the inner membrane via the malate-α-ketoglutarate transporter [Figure 6B, (4)]. This could account for the low oxaloacetate and high malate quantities found in the current study (Figure 3C). High levels of cytosolic NADH may reduce a high amount of oxaloacetate to malate, available for carrying electrons across the inner mitochondrial membrane for ATP synthesis.

To conclude, our data revealed mitochondrial dysfunction in cachetic skeletal muscle tissue, which appeared to have a substantial influence on amino acid metabolism and protein breakdown, based on several lines of evidence. Firstly, we observed significant accumulation of lysine and arginine in cachexia, which can be metabolized in mitochondria to glutamate. Secondly, indices of degradation suggested the presence of degradation of mitochondrial OXPHOS proteins. Thirdly, the energy charge was reduced in cachexia, which may be at least partly attributed to mitochondrial dysfunction. Finally, cachetic muscle tissues exhibited significantly reduced expression of the mitochondrial protein CAT1, which may play an important role in the observed molecular changes associated with cachexia.
whereby the right tissue is belonging to a cachectic mouse. Shown are type I (blue), type IIA (green), type IIB (red), and type IIX (unstained) fibres.

Appendix S7. Statistical analysis regarding changes of molecules in the tricarboxylic acid (TCA) cycle. Peak intensity was significantly higher for malate ($P = 0.0295$) and lower for oxaloacetate ($P = 0.0448$) in tissues of cachectic mice, compared with non-cachectic ones. Boxplot whiskers represent the lowest and highest peak intensities in each group. * $P < 0.05$.

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