Metabolomic profiling of amino acid alterations in anorexia nervosa: implications for appetite regulation and therapeutic strategies

K. DONATO^{1,2,3}, K. DHULI⁴, A. MACCHIA⁴, M.C. MEDORI⁴, C. MICHELL G. BONETTI^{4,5}, M.R. CECCARINI⁵, T. BECCARI⁵, P. CHIURAZZI⁶, S. STON V. BENFATTI⁸, L. DALLA RAGIONE^{8,9}, M. BERTELLI^{1,3,4}

Abstract. - OBJECTIVE: Anorexia nervos (AN), a severe psychiatric disorder primarily fecting adolescents and young adults, is d acterized by extreme dietary restriction and d torted body image. While the psychological a pects of AN are well-documented, it metabolic underpinnings remain ored. We think that metabolomic analy samples emerges as a promising to u complex physiological alterat This study aims to compre file ann ensi no acid concentrations air ` from AN patients and heal ontrols. A ally, it seeks to elucidate tial correlat is beand appetite dystween amino acid regulation in AN, ing light on the eby is of this a physiological 1 ing disorder.

PATIENT ND METHOD A total of 25 ge-matched healthy con-AN patier trols wg for this study. Hair samples we netabolites were extracted a sing high-resolution y-mass spectrometry. chro ochemical markers were data ` ered to aracterize participants' deso linical profiles.

tabolomic analysis revealed sigtalterations in amino acid concentrations in tients compared to healthy controls. Notably acciencies in essential amino acids (EAAs) and branched-chain amino acids (BCAAs) were observed, highlighting potential contributors to muscle wasting and appetite dysregulation. Further analysis identified specific amino acids as robust biomarkers capable of distinguishing AN patients with high sensitivity and specificity. lex metapolic disturbances associated with AN derscores the role of amino acid dysregthe the disorder's pathophysiology. The the disorder's pathophysiology is pathophysiology. The the disorder's pathophysiology is pathophysiology in the disorder's pathophysiology is pathophysiology. The disorder's pathophysiology is pathophysiology in the disorder's pathophysiology is pathophysiology. The disorder's pathophysiology is pathophysiology in the disorder's pathophysiology is pathophysiology. The disorder's pathophysiology is pathophysiology is pathophysiology. The disorder's pathophysiology is pathophysiology in the disorder's pathophysiology is pathophysiology. The disorder's pathophysiology is pathophysiology is pathophysiology is pathophysiology is pathophysiology. The disorder's pathophysiology is pathophysiology is pathophysiology is pathophysiology is pathophysiology is pathophysiology. The disorder's pathophysiology is pathophysiology is pathophysiology is pathophysiology is pathophysiology. The disorder's pathophysiology is pathophysiology is pathophysiology is pathophysiology is pathophysiology. The disorder's pathophysiology is pathophysiology

Kev Words:

Anorexia Nervosa, AN, Metabolomic, Chemoreceptors, EAA, BCAA, Valine, Leucine, APC.

Introduction

Anorexia nervosa (AN) is a debilitating psychiatric disorder characterized by extreme dietary restriction, distorted body image, and an intense fear of gaining weight¹. The disorder primarily affects adolescents and young adults, with a higher prevalence among females, and has a significant impact on physical health, psychological well-being, and overall quality of life². Despite its well-documented psychological aspects, AN is associated with complex physiological changes contributing to its etiology and maintenance³. The regulation of appetite and energy homeostasis is intricate and involves the interplay of various neuroendocrine, metabolic, and behavioral

¹MAGI EUREGIO, Bolzano, Italy

²Department of Health Sciences, University of Milan, Milan, Italy

³MAGISNAT, Atlanta Tech Park, Peachtree Corners, GA, USA

⁴MAGI'S LAB, Rovereto, Trento, Italy

⁵Department of Pharmaceutical Sciences, University of Perugia

⁶UOC Genetica Medica, Fondazione Policlinico Universitario "A. Cemelli & Istituto di Medicina Genomica, Università Cattolica del S. Cuore, Romanti IV

⁷I.S.B – Ion Source & Biotechnologies srl, Bresso, Milan, J

⁸Department of Eating Disorder, Palazzo Francisci Todi, 21 Umbriz Todi, Perugia, Italy

factors^{4,5}. Metabolomic analysis, a powerful tool that examines the comprehensive profile of small molecule metabolites in biological samples, has emerged as a promising approach to unraveling the underlying metabolic alterations in AN6,7. This approach has been successfully employed in various fields, including clinical diagnostics, drug development, and nutrition research⁸. Metabolomic analysis of biological samples, such as blood, urine, and hair, can offer insights into disease-specific metabolic perturbations and help uncover potential biomarkers for various disorders9. The utilization of hair metabolomics offers a unique advantage in the study of AN10; hair metabolomics captures long-term metabolic changes¹¹. Indeed, hair strands grow at an average rate of 1 cm per month, preserving metabolites over extended periods¹². This temporal dimension is particularly relevant for AN, a disorder characterized by chronic and persistent metabolic alterations¹³.

Recent advances in mass spectrometry and liquid chromatography have enabled high-throughput and high-resolution metabolomic analysis of hair samples¹¹. By profiling the metabolites present in hair, researchers can gain insights into the term biochemical processes associated v and potentially uncover novel metabolic pa linked to the disorder^{14,15}. These investigation have identified disturbances in lipid metabol amino acid metabolism, and ene duction AN patients¹⁶. Amino acids, a nponen n, play a of protein synthesis and eng metabo 17 Their pivotal role in maintaining rall h levels are tightly regu ve profound pathways, and any uptions 1 processes effects on physiol

proposed Recent rese indiviosa (AN), may also duals with A orexia experience alterations 1 air perception of erations change chemo sory stimuli. Thes sponse of taste receptors and regulators in th ential amino acids (EAAs) and of n amino ids (BCAAs)²⁰. brance

Amino as as a firal players in this paserve and glocks for proteins and entires, particulate in neurotransmitter synthesis and contribute to energy production througenesis and the citric acid cycle²¹. Vsregulation of amino acid metabolism has implicated in a range of metabolic disorders, in any diabetes, obesity, and cardiovascular diseases²². Our study has two main objectives: firstly, to comprehensively profile the amino acid concentrations in hair samples from AN patients

and healthy controls; and secondly, to elucidate potential correlations between amino acid alterations and appetite dysregulation in AN. By examining the metabolic profile of AN we aim to identify specific amino ac significantly altered and understar now these perturbations contribute to the ca ex metabolic milieu associated with the di Understanding the link between ap o acid olism and appetite regulation, influence ould provide va chemosensory pathway gical insights into the phy s of AN. vs involand, we elucidating the metabol rgets f may uncover no herap ne treatment of AM ntributes oreover, th to the broa of metabol s, shedding of hair metavolomics as a pote light on t tool to investigate co and chronic disorders If the chemosensory with er apprecia rpinnings of appetite gulation.

emosensol unction

gnals provide us with mulmosensory duri social interactions²³. Chemotipl ons can have effects on response starvation, indeed individuals with anorexia (AN) exhibit degrees of chemosensory on¹⁹. Studies²⁴ on animals in which the availability of suitable food is limited show EAAs regulate food intake. This response is triggered by the detection of nutrients within a brain region, the anterior piriform cortex (APC)²⁵⁻²⁷. When meals with low essential amino-acid content are consumed, their concentrations decrease in plasma and brain²⁸. As the concentration of intracellular EAAs decreases, the corresponding transfer RNA (tRNA) becomes deacylated²⁹. Subsequently, the enzyme general control nonderepressible kinase 2 (GCN2) phosphorylates eukaryotic initiation factor 2 (eIF2), resulting in a slowdown of protein synthesis. This favors the translation of mRNA, regulating gene expression^{30,31}. This regulatory process takes longer than the rapid decrease in protein synthesis. Following studies²⁴ on mice, both APC and GCN2 have been identified as EAA chemosensors. APC exhibits sensory function when EAA levels are depleted³², with GCN2 playing a pivotal role by detecting meals lacking in EAA, binding to deacylated tRNA, and phosphorylating eIF2, consequently impairing protein synthesis³³. Numerous studies^{24,34} highlight that the reduced food intake of EAAs involves mechanisms in both the hypothalamus and the APC, and research on mice showed a direct connection between these two regions. The hypothalamus contains appetite regulators, like neuropeptide Y (NPY)³⁵. Starvation triggers an increase in ghrelin, the hunger-stimulating hormone, which activates NPY³⁶. Recent research³⁷ involved subjecting mice to a valine-deficient diet, resulting in significant reductions in food intake and body weight. When a diet rich in valine is reintroduced, normal food intake levels return. This suggests that the taste of a valine-deficient diet might deter mice from consuming an EAA-deficient diet³⁷. Similar results were observed in pigs³⁸, a valine-deficient diet reduced food intake, and this effect was enhanced when an excessive dose of leucine was administered. This could be attributed to the adverse effects of an imbalanced diet in branched-chain amino acids (BCAA), including valine, leucine, and isoleucine, leading to an anorexic response as a protective measure. This response occurs rapidly, likely to safeguard against neuronal pathologies resulting from the imbalance of these three BCAAs³⁷. Recent studies³⁸ involving piglets and pigs have focused on optimizing body weight gain through the careful consideration of dietary valine-to-lysine ratios³⁸. Additionally, research has explored centered around valine, isoleucine, and try to enhance growth performance, while all mining the potential negative impacts on g associated with high concentrations of leucine lysine³⁹. Tryptophan, a precurse rotonin neurotransmitter known for gulatin ldies^{37,38} n these food intake, is a key play in bran-It is well-established that mbala ched-chain amino aci (Bu This is due brain's ability to a o trypto to the shared tra rters between e neutral amino acids, yptophan, BCAA. Consequently, his imb e can lead to reduced serotonin thesis and, c wently, a decrease ake40. This finding in food hold significant ons for individuals with eating disorders, impl a nervosa (AN)⁴¹. Therefore, theinc re is for fur investigation into the way no ac olved in these processes.

Me bolism ways of AA

tne-deficient, threonine-deficient, and lysicts cause mice to cease eating before aching sarlety⁴²⁻⁴⁴ rats can detect EAA-deficient within just 20-30 minutes, resulting in melination⁴⁵. However, when exposed to an EAA-balanced diet after such an experience, their eating rate increases rapidly⁴⁶. Hence, the biochemical and neurological mechanisms responsible

for detecting the presence of EAAs are remarkably swift²⁴. The anterior piriform cortex (APC) has been identified as the brain region responsible for sensing EAAs. An EAA-imbalanced rapidly reduce the levels of limiting a in the APC by up to 56% in just 21 nutes²⁸. A deficiency in EAAs leads to an mulation of deacylated tRNA, resulting in suc levels of aminoacylated tRNA that n thesis cannot be initiated²⁹. This d mechan lets involves an cognizing EAA-deficier new p tial enzyme in initia in synthe. orylates IF2⁴⁷. the GCN2 enzyme wh charge To further suppg ne role NA in detecting EA eficient die njections of tRNA sy inhibitors (L no alcohols) rat APC, resulting in rewere peri ned duced food intake a 20 minutes, simulating ient diet⁴⁸. L-amino the of an EAA nois inhibit the synthesis of charged tRNA, bring the synthesis of uncharged tRNA. The midal cells layer II in the APC serve as itput neur . These cells are excitatory tl ergic r ons regulated by GABAergic glu he absence of new protein synprotein esis, inhibitory proteins critical for controlling 's output circuit are quickly lost from al membrane⁵⁰. Consequently, the APC cannot maintain its normal balance between stimulatory and inhibitory neurons within the circuit. This leads to the liberation of excitatory glutamatergic neurons, which transmit signals to various brain areas involved in heightened motor activity and the rejection of EAA-deficient diets, resulting in reduced food intake and impaired growth³⁴. Leucine activates the mammalian target of rapamycin complex 1 (mTORC1) signaling pathway. This EAA stimulates ribosomal protein S6 kinase (S6K1) and inhibits eukaryotic translation initiation factor 4E binding protein-1 (4EBP1)51. mTOR pathway regulation depends on leucine transport⁵², and it triggers protein synthesis and cell growth in the presence of EAA-balanced diets. As mentioned earlier, valine-deficient diets decrease food intake, and this effect is exacerbated when leucine intake is elevated²⁶. High levels of leucine cause excessive mTOR signaling, adversely affecting normal growth by reducing food intake⁵³. However, since BCAAs (valine, leucine, isoleucine) share the same transporters, valine has been shown to hinder the transport of leucine across the blood-brain barrier, mitigating the excessive stimulation of mTOR³⁹. Excessive leucine concentrations stimulate the catabolism of all BCAAs (valine, leucine, isoleucine) and not just leucine³⁹. A deficiency in these amino acids can alter the expression of growth hormone-insulin-like growth factor-1 (*GH-IGF-1*). GH promotes the secretion of IGF-1, but this process appears to be contingent on the availability of valine, which can, in turn, inhibit the expression of IGF-1. Consequently, as dietary leucine levels increase, food intake decreases^{54,55}. However, adding valine and isoleucine to the diet can counteract the negative effects of high leucine concentrations³⁹. It is crucial, though, to maintain an appropriate balance and not overconsume leucine. Problems related to food intake and growth are associated with an unbalanced intake of valine, leucine, and isoleucine. An EAA-balanced diet not only increases appetite but also enhances growth. Therefore, a direct signaling mechanism operates to maintain a balanced EAA profile in the diet⁵². The metabolomic pathways of BCAA, leucine, valine, and isoleucine are represented in Figure 1.

Patients and Methods

Study Design and Participant Recru

The study was designed to investigate the metabolic alterations in anorexia nervosa through hair metabolomic analysis. A total offemale AN patients and 25 age and bed health female controls were recruited from the cital centers in Italy. The AN patients metabolomic analysis and DSM-V criteria for the disorder theorem of the cital part of the disorder to the cital part of the cital part of the disorder to the cital part of the ci

in eating disorders⁵⁶. Healthy controls were selected based on normal or underweight BMI and matched for age and gender. All participants provided written informed consent, and the was conducted following ethical guide.

Clinical Data Collection

Clinical data were collected oth AN patients and healthy controls their chara demographic and clinical racteristics eight, BMI, pro ta included age, height beha of comorbidities, di s, bioche cal markers (glycemia, creatini uric um, m acid, sodium, p ssium, esium, roid-stivitamin D3, c sterol, trigly mulating b glutamic-02 tic transasuvate transaminase), and minase, *c*ama. other relevant paran

Har Sample Collection and Preparation

flair samples were collected from participants leaviting a 1 diameter hair strand, starting from the school base and extending 4-5 cm town the tire ach hair sample was cleaned, dried, a matter of 1 cm long pieces. These hair seces were placed in glass vials and subjected bolite extraction¹¹.

Metabolite Extraction

Metabolites were extracted from the hair samples using a multi-step procedure⁵⁷. First, 30 mg of hair pieces were immersed in 2 mL of methanol in glass vials. The vials were vortexed and incubated at 50°C for 1 hour. After cooling

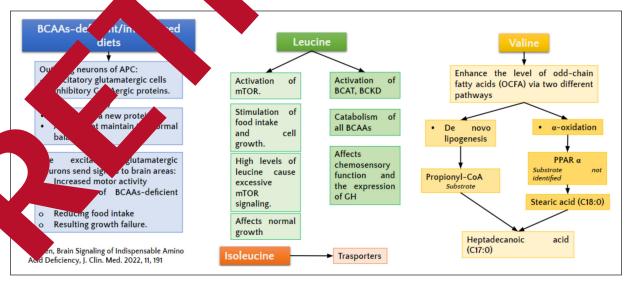


Figure 1. Representation of metabolic pathways of BCAA, leucine, valine, and isoleucine⁵².

to room temperature, 2 mL of acetonitrile was added, and the vials were vortexed, followed by centrifugation at 13,000 g for 10 minutes. The organic layers were collected, combined, and dried under N2 gas. The remaining hair residue was further processed by adding water and adjusting the pH to extract both acidic and basic metabolites. The obtained extracts were then reconstituted in H₂O:CH₂OH (70:30).

Liquid Chromatography-Mass Spectrometry (LC-MS) Analysis

Metabolomic analysis was performed using liquid chromatography (LC) coupled with mass spectrometry (MS)⁵⁸. LC separation was carried out using a Phenomenex Jupiter C18 column (50 x 2.1 mm, 5 µm particle size). Binary gradient elution with mobile phases consisting of water with 0.2% formic acid (mobile phase A) and acetonitrile (mobile phase B) was employed. The LC system was coupled to an HCT Ultra high-capacity ion trap mass spectrometer equipped with an electrospray ionization (ESI) source. The mass spectrometer was operated in both positive and negative ion modes.

Data Analysis

The acquired mass spectrometry data we cessed and analyzed using the SANIST sof suite, a tool specifically designed for metabolo data analysis⁵⁹. Statistical com betwe AN patients and healthy cont onducte values using appropriate tests, and e calculated to assess the signif of d nces in amino acid concentra < 0.05). Ratios of Afic ami were calculated to provid ghts into the play and potential impli appetite reg on.

Statistic Inalysis

In the context of our result, we employed multiplicate analysis techniques, specifically Price Corponent Analysis (PCA) and Receiver and Charteristic (ROC) analysis, in a sign ance the shold set at p-value < 0.05

to evaluate the robustness of our findings. The ratio of patients with Anorexia Nervosa (AN) to healthy controls (CTR) was calculated based on amino acid concentrations in hair, expre nanograms per milligram. Additiona Least Squares Regression (PLSR) s utilizea dy correlato identify biomarkers most st ted with anorexia nervosa by cong patient and control data. Furthern lucted ROC analysis on selected netabolites. ting maximum sensitivi and specificity and th associated cut-off po rea under curve (AUC). These in cal appaches ght int provided a com nensiv ภgnifibetween cant differen in metabol and healthy individuals orexia nerv our understanding of the controls, tribu pathophysiology of L ndition.

Results

analysis of hair samples from another and healthy controls revealed significations in amino acid concentrations that exhibited a key role in the metabolic disturssociated with anorexia nervosa (AN).

Amino Acid Concentrations

Comparative analysis of amino acid concentrations in hair samples showed distinct differences between AN patients and healthy controls (Table I). Notably, propionyl-carnitine and carnitine concentrations were significantly lower in anorexic patients compared to controls (p < 0.001), indicating potential disruptions in lipid metabolism. Propionyl-carnitine and carnitine are involved in fatty acid metabolism, and their reduced levels may suggest altered lipid utilization in anorexic individuals. With the results obtained, it was possible to define the true metabolomic profiles of anorexic patients and of controls. Their comparison enables the discrimination of an affected person from a healthy one, as shown in Figure 2.

Tat ... Amino Acid Concentrations in AN Patients and Healthy Controls. The ratio is between healthy controls (CTR), where we have a concentration in AN Patients and Healthy Controls. The ratio is between healthy controls (CTR), where we have a concentration in AN Patients and Healthy Controls. The ratio is between healthy controls (CTR), where

Amino Acid	Healthy Controls [Ng/Mg]	AN Patients [Ng/Mg]	Ratio CTR/AN	<i>p</i> -value
nyl-Carnitine	1.45784	0.00788	185.01	1.76E-21
Carnitine	0.08468	0.00396	21.38	6.12E-14
Leucine/Isoleucine	3.47448	1.33596	2.60	6.08E-08
Valine	9.04788	3.92512	2.30	4.38E-07

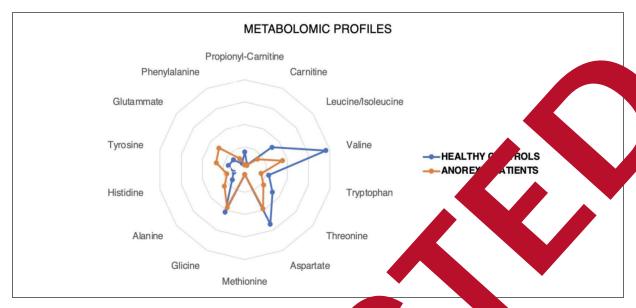


Figure 2. Metabolomic profiles of anorexic patients and controls.

Leucine, isoleucine, and valine are branched-chain amino acids (BCAAs) crucial for muscle maintenance and protein synthesis 60 . AN patients exhibited significantly lower concentration of leucine (p < 0.001), isoleucine (p < 0.001) compared to controls, suggesting compromised protein metabolism and muscle servation in AN. This observation aligns with known catabolic state associated approach as p = 1.000

Alanine, tyrosine, and ph concei trations were found to be ner in patients than in controls. Alanine arbohyplaye drate metabolism⁶², m degradation or alt metabo thways linlism in Al ked to protein q sine and phenylalanine rsors of ne ransmitters⁶³, and then elevate Is might contribute to often observed in the neuro mical imbala AN. Fi 3 and Figure 4 ill. e the alterations acid concentrations in anorexic patients in a to 1 thy controls. Figure 3 shows the con differe oolomic ofiles between patients hile cipal component analysis cont e 4) si patients and controls clustegroups. Significant differences two distr bserved for propionyl-carnitine, carnitine, ine, valine, and other amino acids. ese alterations point to potential metabolic diions associated with anorexia nervosa.

Essential Amino Acid Deficiency

An important finding is the significant deficiency of essential amino acids (EAAs) in AN patients.

total concert ion of EAAs was notably lower in Lipatients (Lipatients (Lipatie

Tryptophan Ratio

The ratio of tryptophan to valine, leucine, and isoleucine was elevated in AN patients (0.27) compared to controls (0.19), resulting in a ratio of 0.70 (CTR/AN). This finding suggests that anorexic individuals preferentially transport tryptophan over these BCAAs due to their shared transporters⁴⁰.

The observed alterations in amino acid concentrations provide insights into the metabolic disruptions in AN. The deficiencies in BCAAs and EAAs point to the importance of these amino acids in maintaining muscle mass and regulating appetite.

These findings have potential clinical implications for AN treatment. Restoring BCAA and EAA levels through targeted supplementation may support muscle preservation and address appetite dysregulation⁶⁵. Further investigations are warranted to explore the mechanisms underlying these metabolic alterations and to develop personalized interventions for AN patients⁶⁶⁻⁶⁸ (Table II).

Machine Learning Algorithms

We employed a machine learning model, specifically Partial Least Squares Regression⁶⁹, to explore the potential utility of classification models

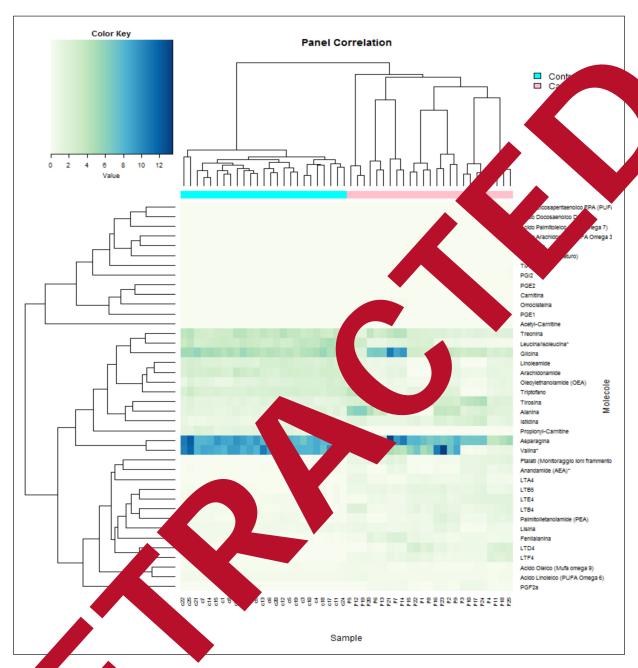


Fig. Heat to compare metabolomic profiles of patients and controls. Patients are shown in red, and controls are shown.

conte. . . . , as shown in Table III. In ntext, the features represent molecules, all and us to identify those of utmost importune are the robustness of our findings, e conducted 100 iterations of analysis⁷⁰. In each tion, we randomly divided the samples into a transport of the data and a testing set (comprising 25% of the data). The outcomes of this analysis were promising, with the model consistently achieving a high correlation

of 98%. The top 10 molecules that consistently emerged as the most important throughout these iterations hold a particular interest in the context of AN. These molecules serve as potential markers, demonstrating their exceptional ability to distinguish between different groups within our dataset. However, it is important to note that while these molecules are effective classifiers within our dataset, further investigations are required to elucidate the nature of this variation.

Table II. ROC analysis of selected metabolites. The maximum sensitivity and specificity values and their correlated cut-off are shown. Then, we were able to calculate the area under the curve (AUC).

Molecules	Cut-off	Sensibility	Specificity	AUC
Propionyl-Carnitine	0.5825	1	1	
Carnitine	0.0255	1	1	1
PGE2	0.022	1	1	1
PGE1	0.007	1	1	1
Arachidonamide	1.7975	1	1	
Linoleamide	1.5595	1	1	
Leucine/Isoleucine	2.3875	0.88	1	0.
OEA	1.308	1	0.84	0.92
Valine	7.3105	0.88	1	0.94
LTB4	0.475	0.88	1	0.0
LTB5	0.525	1		
LTA4	0.149	1	1	
LTF4	0.293	1	1	1
PGI2	0.0055	1	96	0.98



Figure 4. Principal component analysis (PCA). Metabolic Alterations in Amino Acid Concentrations. Patients are shown in red, and controls are shown in light blue.

Discussions

The present study was carried out to study metabolic alterations in anorexia nervosa (AN) through the analysis of amino acid concentrations in hair samples. The results provide valuable insights into the intricate biochemical changes associated with this complex and debilitating disorder. However, it is essential to consider the role of the chemosensory pathway in these metabolic changes. This pathway plays a pivotal role in shaping eating behaviors and is influenced by various amino acids. The observed alterations in amino acid concentrations underscore the extent of metabolic disturbances in AN and other metabolic disorders⁷¹. The reduced levels of propionyl-carnitine and carnitine are of particular interest, as they reflect potential disruptions in lipid metabolism⁷². These two molecules play a crucial role in fatty acid transportation into mitochondria for energy production⁷³. The scarcity of propionyl-carnitine and carnitine in anorexic patients may suggest altered lipid utilization pathways, aligning with the energy conservation strategies often observed in individuals with AN⁷⁴. The creased concentrations of leucine, isoleuc valine in AN patients underscore the pr nd alterations in protein metabolism and muscl servation in AN⁷⁵. Branched-chain amino a (BCAAs) are vital for maintain scle ma and their deficiency in AN ibute 1 muscle wasting and catab eduction of BCAAs is a noteworth rovides a biochemical basis patient commonly seen in

The elevated of alanine ine, and otential phenylalanine tients indica metabolism⁷⁸. Alanidisturbances in amino ne's role carbohydrate bolism makes its AN a plausible increas ction of altered c pathways associated with protein catameta ry, the elevation of tyrosine and bol aligns wheir role as precursors pheny mitter nthesis⁶³, suggesting that neur ances might contribute to chemi c features of AN. The marked iropsych ency of essential amino acids (EAAs) in AN s significant implications⁷⁹. EAAs e not only building blocks for proteins but also regulators of appetite and food intake⁸⁰. The decrease in total EAAs in AN patients points to a potential mechanism contributing to appetite dysregulation⁸¹, and the overall malnourished state observed in AN. The increased ratio

Table III. Partial Least Squares Regression to identify the biomarkers more correlated to anorexia nervosa, comparing patients and controls data.

Molecules	Importance_mea
Propionyl-Carnitine	100
Carnitine	93.04
LTB5	92.93
Arachidonamide	92.11
PGE2	91,7
Linoleamide	8
LTA4	.41
LTF4	\$2.30
Leucine/Isoleucine	01
OEA	
PGI2	
LTB4	77.2
PGE1	77.06
VALINE	71.64

yptophan to valine, Licine, and isoleucine cates preferential transport of tryptophan in hese amino acids share the patients. A this alteration might lead to transporte lance the central nervous system's production⁸². The preferential neurou ensport of tryptophan over the BCAAs could to decreased appetite and the perpe-If the catabolic state⁸³. Finally, the metabolic profile of anandamide and oleoylethanolamide showed interesting results. According to the literature84, anandamide is positively correlated with excessive physical activity. These data are supported by our findings. Indeed, anandamide was overexpressed in AN patients. On the contrary, oleoylethanolamide was more expressed in controls, while in literature seems to be positively correlated with weight loss⁸⁵.

While the findings of this study offer valuable insights into the metabolic alterations associated with AN, it is crucial to acknowledge the role of the chemosensory pathway in shaping the appetite and eating behaviors of individuals with this disorder. The deficiencies in BCAAs and EAAs underscore the potential benefits of targeted nutritional interventions that aim to restore these amino acid levels⁸⁶. Such interventions could have a dual effect of preserving muscle mass and modulating appetite⁶⁵.

Limitations

However, it is essential to acknowledge certain limitations of the study. The cross-sectional design limits the ability to establish causality and does not capture the dynamic changes that might occur over time. Additionally, the sample size is modest, warranting larger studies to validate these findings and explore potential factors influencing these metabolic alterations, such as disease duration, severity, and comorbidities.

Conclusions

The metabolomic analysis of hair samples from AN patients and healthy controls provides valuable insights into the metabolic disturbances associated with this complex disorder. The alterations in amino acid concentrations, especially EAAs and BCAAs, shed light on potential mechanisms underlying muscle wasting, appetite dysregulation, and neuropsychiatric features in AN. Further research is warranted to unravel the intricate interplay of these metabolic changes and to develop early diagnosis and targeted therapeutic strategies that address the multifaceted nature of anorexia nervosa.

Conflict of Interest

The authors declare no conflict of interest.

Informed Consent

All subjects gave their informed consent for inclusion fore they participated in the study.

Ethics Approval

The study was conduct of accords to the Deciaration of Helsinki, and roved by the Committee of Azienda Sanita (Approval No. 132-201).

Fund

This purch we middle by the Provincia Autonoma di Bolza. We me framework of LP 14/2006.

prs' Če ons

Construalization, 3.; Methodology, S.C.; Investigation V.B. and L.D.R.; Writing- original draft preparation, K. D. G.B., Writing, review and editing, K. Dhuli, J.B., M.R.C., T.B. and P.C.; Project admination, M.B.; Funding acquisition, M.B. All authors have ad agreed to the published version of the manuscript.

Availability of Data and Materials

All the data are within the test.

ORCID ID

Kevin Donato: 0000-0001-6527-3753 Gabriele Bonetti: 0000-0002-0666-9616

Maria Rachele Ceccarini: 0000-0002-5344-4890

Tommaso Beccari: 0000-0001-9637-6579 Pietro Chiurazzi: 0000-0001-5104-1521 Simone Cristoni: 0000-0001-9579-7953 Matteo Bertelli: 0000-0002-7646-2872

Refer ices

- Viapiana O, G ve R, To D, Rossini M, a V, Ido Frace ⊆, Adaal density mi S. Mar increases in and bio markers of b nover in paweight. Bone tients an nervosa gain. 2007; 40: 1073
- 2) Archys J, Mitchell Males J, Nielsen S. Morates in patients anorexia nervosa and other eating disorders. A meta-analysis of 36 studies. Arch Oen Psychiatry 2011; 68: 724-731.
 - Berner LA, B. TA, Lavender JM, Lopez E, Wienga CE, K. WH. Neuroendocrinology of rein and a nervosa and bulimia nervosa: and ghrelin. Mol Cell Endocrinol 2019, 1911. 110320.
 - Sevlin MJ, Walsh BT. "The neuroendocrinology of ita nervosa." Neuroendocrinology of mood. In, Heidelberg: Springer Berlin Heidelberg 1988; 291-307.
- Selby EA, Panza E, Plasencia M. Positive emotion dysregulation in eating disorders and obesity. In J Gruber (Ed.) The Oxford handbook of positive emotion and psychopathology. Oxford University Press, 2019; pp. 424-443.
- Shih PB. Metabolomics Biomarkers for Precision Psychiatry. Adv Exp Med Biol 2019; 1161: 101-113
- Qiu S, Cai Y, Yao H, Lin C, Xie Y, Tang S, Zhang A. Small molecule metabolites: discovery of biomarkers and therapeutic targets. Signal Transduct Target Ther 2023; 8: 132.
- 8) Villas-Bôas SG, Rasmussen S, Lane GA. Metabolomics or metabolite profiles? Trends Biotechnol 2005; 23: 385-386.
- Mussap M, Loddo C, Fanni C, Fanos V. Metabolomics in pharmacology - a delve into the novel field of pharmacometabolomics. Expert Rev Clin Pharmacol 2020; 13: 115-134.
- Jang WJ, Choi JY, Park B, Seo JH, Seo YH, Lee S, Jeong CH, Lee S. Hair Metabolomics in Animal Studies and Clinical Settings. Molecules 2019; 24: 2195.
- Chen Y, Guo J, Xing S, Yu H, Huan T. Global-Scale Metabolomic Profiling of Human Hair for Simultaneous Monitoring of Endogenous Metabolome, Short- and Long-Term Exposome. Front Chem 2021; 9: 674265.

- 12) Eisenbeiss L, Steuer AE, Binz TM, Baumgartner MR, Kraemer T. (Un)targeted hair metabolomics: first considerations and systematic evaluation on the impact of sample preparation. Anal Bioanal Chem 2019; 411: 3963-3977.
- Cobo-Golpe M, Baumgartner MR, Binz TM, Kraemer T, Steuer AE. Detection of hair metabolome changes in cocaine users using untargeted metabolomics. Toxicol Anal et Clin 2022; 34: S34.
- 14) Rosenberg AM, Rausser S, Ren J, Mosharov EV, Sturm G, Ogden RT, Patel P, Kumar Soni R, Lacefield C, Tobin DJ, Paus R, Picard M. Quantitative mapping of human hair greying and reversal in relation to life stress. Elife 2021; 10: e67437.
- 15) Föcker M, Cecil A, Prehn C, Adamski J, Albrecht M, Adams F, Hinney A, Libuda L, Bühlmeier J, Hebebrand J, Peters T, Antel J. Evaluation of Metabolic Profiles of Patients with Anorexia Nervosa at Inpatient Admission, Short- and Long-Term Weight Regain-Descriptive and Pattern Analysis. Metabolites 2020; 11: 7.
- 16) Mayo-Martínez L, Rupérez FJ, Martos-Moreno GÁ, Graell M, Barbas C, Argente J, García A. Unveiling Metabolic Phenotype Alterations in Anorexia Nervosa through Metabolomics. Nutrients 2021; 13: 4249.
- Lopez MJ, Mohiuddin SS. Biochemistry, Essential Amino Acids. In: StatPearls Treasure (FL): StatPearls Publishing; 2023.
- 18) Church DD, Hirsch KR, Park S, Kim IY, Carlo, Pasiakos SM, Wolfe RR, Ferrando AA. Es Amino Acids and Protein Synthesis: Insight to Maximizing the Muscle and Whole-Body sponse to Feeding. Nutrients 2 2 3717.
- 19) Kinnaird E, Stewart C, Tchanna and te sensitivity in anorexia nervosa systema eview. Int J Eat Disord 2018; 51: 184.
- 20) Anthony TG, Gietze DW.
 id deprivation in the central system. Jury
 Opin Clin Nutral ab Care 20: 26-101.
- 21) Gutiérrez-Para Romero H, Tarak M. An evolutiona perspandon amino acus. Nat Edu 2010; 3-29.
- 22) Zhu a Bai M, Xie X, Wang C, Dai H, Cha J, Han F, Lin W. Imparad Amino Acid Melism and Its Correlation With Diabetic Kidney ase Procession in Type 2 Diabetes Mellitus.
- Steven RJ. An all evaluation of the functions f huma. The Chem Senses 2010; 35: 3-20.
- 24) etzen Dv. o S, Anthony TG. Mechanisms of od intake repression in indispensable amino acceptancy. Annu Rev Nutr 2007; 27: 63-78.
- A, Cherasse Y, Zeng H, Zhang Y, Harding HP, Con D, Fafournoux P. The GCN2 kinase biases eding behavior to maintain amino acid homeostasis in omnivores. Cell Metab 2005; 1: 273-277.
- Gloaguen M, Le Floc'h N, Corrent E, Primot Y, van Milgen J. Providing a diet deficient in va-

- line but with excess leucine results in a rapid decrease in feed intake and modifies the postprandial plasma amino acid and α -keto acid concentrations in pigs. J Anim Sci 2012; 90: 3135-3142.
- 27) Hawkins RA, O'Kane RL, Simpson IA Structure of the blood-brain barrier and its in the transport of amino acids. J 2006; 136: 218S-226S.
- 28) Koehnle TJ, Russell MC, Morin to liets deficient in indispensable are acids and y decrease the concentration of the limit and sino acid in the anterior pinn on cortex of rats 2004; 134: 2365-23
- 29) Huynh LN, Thanga Charl. Linking tRNA localization with actival attritional ass responses. Compact yole 2016. 12-31
- 30) Baird TD RC. Eukary on factor 2 phosph of translation ontrol in metabolic Adv 212; 3: 307-321.
- 31) Kilberg MS, Balas Manian M, Fu L, Shan J. scription factors twork associated with me assino acid response mammalian cells. Adv Nutr 2012; 3: 295-306.
- Hansen HS ana V. Non-endocannabinoid ines and 2-monoacylglycerols in J Pharmacol 2019; 176: 1443-
- 33) Hao e, marp JW, Ross-Inta CM. Uncharged tR-NA and sensing of amino acid deficiency in mamber piriform cortex. Science 2005; 307: 1776-
- 34) Gietzen DW, Aja SM. The brain's response to an essential amino acid-deficient diet and the circuitous route to a better meal. Mol Neurobiol 2012; 46: 332-348.
- 35) Aponte Y, Atasoy D, Sternson SM. AGRP neurons are sufficient to orchestrate feeding behavior rapidly and without training. Nature Neurosci 2011; 14: 351-355.
- 36) Cummings DE, Purnell JQ, frayo RS, Schmidova K, Wisse BE, Weigle DS. A preprandial rise in plasma ghrelin levels suggests a role in meal initiation in humans. Diabetes 2001; 50: 1714-1719.
- Goto S, Nagao K, Bannai M. Anorexia in rats caused by a valine-deficient diet is not ameliorated by systemic ghrelin treatment. Neurosci 2010; 166: 333-340.
- Siebert D, Khan DR, Torrallardona D. The Optimal Valine to Lysine Ratio for Performance Parameters in Weaned Piglets. Animals (Basel) 2021; 11: 1255.
- 39) Kerkaert HR, Cemin HS, Woodworth JC, DeRouchey JM, Dritz SS, Tokach MD, Goodband RD, Haydon KD, Hastad CW, Post ZB. Improving performance of finishing pigs with added valine, isoleucine, and tryptophan: validating a meta-analysis model. J Anim Sci 2021; 99: skab006.
- Fernstrom JD. Large neutral amino acids: dietary effects on brain neurochemistry and function. Amino Acids 2013; 45: 419-430.

- 41) Hu J, Le Q, Zhang M, Kuang S, Gu W, Sun Y, Jean Jacques K, Zhang Y, Li Y, Sun J, Yang Y, Wang Y, Xu S. and Yan, X. Effects of amino acids on olfactory-related receptors regulating appetite in silver pomfret. Aquac Res 2021; 52: 2528-2539.
- 42) Gietzen DW. Time course of food intake and plasma and brain amino acid concentrations in rats fed amino acid-imbalanced or-deficient diets. Interaction of the Chemical Senses with Nutrition 1986; 415-456.
- 43) Mori M, Kawada T, Ono T, Torii K. Taste preference and protein nutrition and L-amino acid homeostasis in male Sprague-Dawley rats. Physiol Behav 1991; 49: 987-995.
- Koehnle TJ, Russell MC, Gietzen DW. Rats rapidly reject diets deficient in essential amino acids. J Nutr 2003: 133: 2331-2335.
- 45) Feurté S, Nicolaidis S, Berridge KC. Conditioned taste aversion in rats for a threonine-deficient diet: demonstration by the taste reactivity test. Physiol Behav 2000; 68: 423-429.
- 46) Rogers QR, Leung PM. The influence of amino acids on the neuroregulation of food intake. Fed Proc 1973; 32: 1709-1719.
- 47) Pezeshki A, Chelikani PK. Low Protein Diets and Energy Balance: Mechanisms of Action on Energy Intake and Expenditure. Front Nutr 2021; 8: 67
- 48) Gietzen DW, Hao S, Anthony TG. "A deid-sensing mechanisms: biochemistry a pehavior". Handbook of Neurochemistry and ular Neurobiology. Springer US, 2007; 249-2
- 49) Ekstrand JJ, Domroese ME, Johnson DM, P SL, Knodel SM, Behan M, B. A ne subdivision of anterior pirit corte d associ ated deep nucleus with el feature of interest for olfaction and epile. Comp 434: 289-307.
- 50) Sharp JW, Rose La CM, Barria L, Payne JA, Rudell JB, Gira DW. Effects antial amino acid definition wn-regulation and action the GABA eceptaristic piriform cortex. J Net. 2013; 127: 520-530.
- 51) Sax RA, Sabatini D, TOR Signaling in Gran, Metabolism, and sease. Cell 2017; 960-97 Erratum in: Cell 2017; 169: 361-371.
- 52) An Brain Signaling of Indispensable Deficier J Clin Med 2021; 11: 191.
 - Cemils okar anD, Dritz SS, Woodworth JC, eRouch coodband RD. Meta-regression alysis to certificate the influence of branched-chain and large neutral amino acids on growth performal series. J Anim Sci 2019; 97: 2505-2514.
- Winter MK, Pfaffl MW, Roth FX. The effects of branched-chain amino acid interactions on rowth performance, blood metabolites, enzyme netics and transcriptomics in weaned pigs. Br J Nutr 2010; 103: 964-976.
- 55) Kwon WB, Touchette KJ, Simongiovanni A, Syriopoulos K, Wessels A, Stein HH. Excess di-

- etary leucine in diets for growing pigs reduces growth performance, biological value of protein, protein retention, and serotonin synthesis1. J Anim Sci 2019; 97: 4282-4292.
- 56) Vo M, Accurso EC, Goldschmidt AB, D. The Impact of DSM-5 on Eating Disease Diagnoses. Int J Eat Disord 2017; 50: 581.
- 57) Ramírez Fernández MDM, William IR, Jankowski D, Hill V, Samyn N. Develope and f an UP-LC-MS/MS method for the palysis and extraction of the polydrug history in intanyl analogus. Forensic Sci Int 2021, 37: 110177.
- 58) Marcos A, León C, lández M, Cás-, Nozal I tro-Rubio F, G mbrorido-Untarg c study sio E, Creg etabo ometry in by liquid matographyned cocaine brain tis the effects of 10l ministration and o male and female young rate hromatogr A 2023; 1700:
- 59 I. J. J. S. Dusi G, Brs. J. Ja P. SANIST: optimization of a technology for compound identification based on the European Union directive with applications prensic, pharmaceutical and food nalyses. Jo. al of Mass Spectrometry 2017; 16-21. At able at: https://boa.unimib.it/han-355
- 60) Wolfe nn. Branched-chain amino acids and mussle protein synthesis in humans: myth or reality? Coc Sports Nutr 2017; 14: 30.
- Disclop CA, Machate T, Henning T, Henkel J, Püschel G, Weber D, Grune T, Klaus S, Weitkunat K. Detrimental effects of branched-chain amino acids in glucose tolerance can be attributed to valine induced glucotoxicity in skeletal muscle. Nutr Diabetes 2022; 12: 20
- 62) Sookoian S, Pirola CJ. Alanine and aspartate aminotransferase and glutamine-cycling pathway: their roles in pathogenesis of metabolic syndrome. World J Gastroenterol 2012; 18: 3775-3781.
- 63) Fernstrom JD, Fernstrom MH. Tyrosine, phenylalanine, and catecholamine synthesis and function in the brain. J Nutr 2007; 137: 1539S-1547S.
- 64) Rigamonti AE, Tamini S, Cicolini S, De Col A, Caroli D, Mai S, Rondinelli E, Saezza A, Cella SG, Sartorio A. Evaluation of an Amino Acid Mix on the Secretion of Gastrointestinal Peptides, Glucometabolic Homeostasis, and Appetite in Obese Adolescents Administered with a Fixed-Dose or ad Libitum Meal. J Clin Med 2020; 9: 3054.
- 65) VanDusseldorp TA, Escobar KA, Johnson KE, Stratton MT, Moriarty T, Cole N, McCormick JJ, Kerksick CM, Vaughan RA, Dokladny K, Kravitz L, Mermier CM. Effect of Branched-Chain Amino Acid Supplementation on Recovery Following Acute Eccentric Exercise. Nutrients 2018; 10: 1389.
- 66) Bifari F, Nisoli E. Branched-chain amino acids differently modulate catabolic and anabolic states in

- mammals: a pharmacological point of view. Br J Pharmacol 2017; 174: 1366-1377.
- Hajian-Tilaki K. Receiver Operating Characteristic (ROC) Curve Analysis for Medical Diagnostic Test Evaluation. Caspian J Intern Med 2013; 4: 627-635.
- 68) Habibzadeh F, Habibzadeh P, Yadollahie M. On determining the most appropriate test cut-off value: the case of tests with continuous results. Biochem Med 2016; 26: 297-307.
- 69) Boulesteix AL, Strimmer K. Partial least squares: a versatile tool for the analysis of high-dimensional genomic data. Brief Bioinform 2007; 8: 32-44.
- Cassel C, Hackl P, Westlund AH. Robustness of partial least-squares method for estimating latent variable quality structures. J Appl Stat 1999; 26: 435-446.
- 71) Chen C, Hou G, Zeng C, Ren Y, Chen X, Peng C. Metabolomic profiling reveals amino acid and carnitine alterations as metabolic signatures in psoriasis. Theranostics 2021; 11: 754.
- 72) Bene J, Hadzsiev K, Melegh B. Role of carnitine and its derivatives in the development and management of type 2 diabetes. Nutrition 2018; 8: 8.
- 73) Virmani MA, Cirulli M. The Role of I-Carnitine in Mitochondria, Prevention of Metabolic Inflexibility and Disease Initiation. Int J Mol Sci 2022 23: 2717.
- 74) Wu T, Guo A, Shu Q, Qi Y, Kong Y, Sun Z, T S, Fu Z. L-Carnitine intake prevents irregula ing-induced obesity and lipid metabolism der. Gene 2015; 554: 148-154.
- 75) Adibi SA. Metabolism of branch hain amacids in altered nutrition. 1976; 2: 1287-1302.
- 76) Mann G, Mora S, Madu goke O Aranchedchain Amino Acids: Satas

- and Implications for Muscle and Whole-body Metabolism. Front Physiol 2021; 12: 702826.
- 77) Shimomura Y, Murakami T, Nakai N, Nagasaki M, Harris RA. Exercise promotes BCAA cetabolism: effects of BCAA supplementation etal muscle during exercise. J Nutrico 04; 15 1583S-1587S.
- 78) Miflin B, Lea P. Amino acid lism. Annu Rev Plant Physiol 1977; 28: 299-
- 79) Xiao F, Guo F. Impacts ssentia o acids on energy balance followed Metab 57 101393.
- 80) Wang W, Xu Y, Chang P, K, Song F. Acter y lysine reculate with performance via the nutrice sensing aling part dys in largemouth as (Microp. alm s). Front Mar Sci 2 7: 595682.
- 81) Moris C., Blades C., Aoh Y. Nutrient-Based App Regulation. J Obes Metab Syndr 2022; 31: 10
- 82) d E, Øverli Ø, verg S. Tryptophan Metabolic Pathways and Bran Serotonergic Activity: A Comparative Review. Front Endocrinol 2019; 10: 158.
- 8 larper AE, N r RH, Block KP. Branched-chain abolism. Annu Rev Nutr 1984; 4:
- 34) Dietrich A, McDaniel WF. Endocannabinoids and exercise. Br J Sports Med 2004; 38: 536-541.
 - S, Arnold M, Krieger JP, Wolfstädter B, McJer U, Langhans W, Mansouri A. Oleoylethanolamide-induced anorexia in rats is associated with locomotor impairment. Physiol Rep 2018; 6: e13517.
- 86) Nie C, He T, Zhang W, Zhang G, Ma X. Branched Chain Amino Acids: Beyond Nutrition Metabolism. Int J Mol Sci 2018; 19: 954.