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- 18 Title: From Mitochondria To Healthy Aging: The Role Of Branched-chain Amino Acids Treatment: MATeR
- 19 A Randomized Study
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37	Abstract

38 Rationale: Malnutrition often affects elderly patients and significantly contributes to the reduction in 39 healthy life expectancy, causing high morbidity and mortality. In particular, protein malnutrition is one of 40 the determinants of frailty and sarcopenia in elderly people. 41 Methods: To investigate the role of amino acid supplementation in senior patients we performed an open-42 label randomized trial and administered a peculiar branched-chain amino acid enriched mixture (BCAAem) 43 or provided diet advice in 155 elderly malnourished patients. They were followed for 2 months, assessing 44 cognitive performance by Mini Mental State Examination (MMSE), muscle mass measured by 45 anthropometry, strength measure by hand grip and performance measured by the Timed Up and Go (TUG) 46 test, the 30 seconds Chair Sit to Stand (30-s CST) test and the 4 meters gait speed test. Moreover we 47 measured oxidative stress in plasma and mitochondrial production of ATP and electron flux in peripheral 48 blood mononuclear cells. 49 Results: Both groups improved in nutritional status, general health and muscle mass, strength and 50 performance; treatment with BCAAem supplementation was more effective than simple diet advice in 51 increasing MMSE (1.2 increase versus 0.2, p=0.0171), ATP production (0.43 increase versus -0.1, p=0.0001), 52 electron flux (0.50 increase versus 0.01, p<0.0001) and in maintaining low oxidative stress. The 53 amelioration of clinical parameters as MMSE, balance, four meter walking test were associated to 54 increased mitochondrial function. 55 Conclusions: Overall, our findings show that sustaining nutritional support might be clinically relevant in 56 increasing physical performance in elderly malnourished patients and that the use of specific BCAAem 57 might ameliorate also cognitive performance thanks to an amelioration of mitochondria bioenergetics. 58 **Keywords** 59 Malnutrition, elderly patients, branched-chain amino acids, muscle mass and strength, mitochondrial 60 activity and biogenesis, oxidative stress.

61 Abbreviations

62 30-s CST, 30 seconds chair sit to stand test; ADL, activity of daily living; BCAAem, branched chain amino 63 acid enriched mixture; BCAAs, branched chain amino acids; BMI, body mass index; CIRS, cumulative index rating scale; COX-1 and 4, cytochrome C oxidase 1 and 4; FOXO, forkhead box O; GAPDH, glyceraldehyde 3-64 phosphate dehydrogenase; GDS, geriatric depression scale; GLM, linear regression models; MFN-1 and 2, 65 66 mitofusin-1 and 2; MMSE, mini-mental state examination; MNA, mini nutritional assessment test; mTOR, 67 mechanistic target of rapamycin; NRF-1, nuclear respiratory factor-1; NO, nitric oxide; OECD, organisation 68 for economic co-operation and development; PBMCs, peripheral blood mononuclear cells; ROS, reactive 69 oxygen species; RT-PCR, real time PCR; TBARs, thiobarbituric acid reactive substances; TFAM, mitochondrial 70 transcription factor A; TUG, timed up and go test.

71

### 72 Introduction

73 Thanks to an increased life expectancy, an improvement in health status and medical services, the older 74 population is constantly increasing. In 2015, 617 million (8.5%) people in the world were aged 65 and over 75 (older adults) and these numbers are estimated to rise to 1.6 billion by 2050 (1). Despite the increase in life 76 expectancy, there is no corresponding increase in healthy life expectancy: recent findings in 2015 show 77 that, despite a life expectancy at the age of 65 of 21.2 years for women and 17.9 years for men, only 9.4 78 years are healthy years (2). Concomitantly, health maintenance in older age will be one of the most 79 relevant societal challenges in the future years. Lifestyle changes appear to be fundamental in increasing 80 healthy life expectancy, and adequate nutrition is enormously important, given that malnutrition (i.e., 81 undernutrition), particularly as protein-energy deficit is very common amongst the elderly population. This 82 is due to the effects of aging per se that causes decreased salivation, difficulty swallowing, and delayed 83 emptying of the stomach and oesophagus, as well as slower gastrointestinal movement (3). Other 84 conditions associated with aging, such as drug use, loneliness, depression, lack of oral health, low quality of life, in addition to chronic non-communicable diseases, markedly increase the undernutrition risk (4). It has 85 86 been estimated that undernutrition affects between 20 and 50% of hospitalised patients (5) and 5 and 10% 87 of patients living at home in the community (6), the great variations reported in different studies depends

88 not only to differences in the population analysed, but also on the adopted definition. Recently a large 89 study (7), that applies harmonized criteria to define malnutrition in different clinical settings and 90 population, shows a great underestimation of the problem and confirms higher prevalence of malnutrition 91 in residents of nursing homes and hospitalized patients evaluated by Mini Nutritional Assessment test 92 (MNA). Malnutrition is more prevalent in patients affected by acute and chronic disorders with reduced 93 functional status (7) and is associated with poor clinical outcomes and prognosis. Malnutrition is associated 94 with reduced immune function, anaemia, impaired cognitive function, and higher hospitalisation rate and is 95 a strong independent predictor of mortality [for a review, see A. Granic et al, 2018 (8)]. Malnutrition causes 96 body weight loss and muscle atrophy, decreased muscle strength and function, impaired balance, and 97 increased fall and fall-related injuries. The malnutrition-linked muscle atrophy accelerates transition to 98 frailty (9) and has been considered as one of the determinants of sarcopenia (10,11). Importantly, 99 sarcopenia is defined as low muscle mass, with defective muscle strength (also named dynapenia) and 100 decline of physical performance (12) and is, per se, associated with increased morbidity and mortality (13). 101 Although several guidelines and consensus documents on nutritional care of malnourished elderly subjects 102 have been proposed (14), and protein needs established in the range from 0.8 g/kg/day (healthy adults) 103 and up to 1.5 g/kg/day (in some cases even higher) according to age, disease and degree of protein 104 depletion (15), the daily protein consumption in older subjects is often inadequate and undernutrition and 105 sarcopenia are underestimated and considered as one of the factors of aging (4, 7). 106 It has been suggested that the aging process significantly affects protein metabolism and enhances the 107 muscle wastage that accompanies undernutrition and sarcopenia. Some studies show lower plasma 108 concentrations of branched-chain amino acids (BCAAs) in elderly subjects (16,17), whereas others do not 109 (18,19). This may be due to the increased first-pass splanchnic extraction of amino acids in older people, 110 with a consequent decrease in delivery to the skeletal muscle tissue and availability for muscle tissue 111 anabolism (20). Most kinetics studies show no difference in the ability of older subjects to retain and 112 metabolise BCAAs (21-24).

These observations suggest that the dietary requirement of proteins and essential amino acids is higher in the elderly than in young adults (25) and that an increased intake of a mixture of amino acids or essential amino acids can increase amino acid availability and result in the stimulation of muscle protein anabolism (26).

A number of reports, including a recent well conducted meta-analysis concludes that dietary supplement of
 essential amino acids is more effective than non-essential amino acid or whole protein supplementations in
 malnourished patients (27).

120 Notably, amino acid mixtures enriched in BCAAs have been shown to promote mitochondrial biogenesis

and function, in addition to decrease oxidative stress via nitric oxide (NO) and mechanistic target of

rapamycin (mTOR) signalling pathways in middle-aged mice (28). A more recent study has shown the

stimulatory effect of leucine on mitochondrial respiration and ATP production in human macrophages (29).

124 These results are important because mitochondrial dysfunction is a hallmark of the aging processes and

age-related disorders, including sarcopenia and cognitive decline, are characterized by reduced

126 mitochondrial mass and function [for a comprehensive review, see N. Sun et al, 2016 (30)]. Dietary

127 supplementation of BCAA-enriched mixtures (BCAAem) may contribute to slow-down mitochondrial

decline and to ameliorate clinical status of malnourished elderly patients (31).

129 The MATeR study thus aimed to evaluate the efficacy of a specific BCAAem compared to diet advice to

130 promote mitochondrial function and improve clinical outcomes, particularly muscle and cognitive

131 performance, in malnourished elderly community-dwelling subjects.

132

#### 133 Materials and methods

134 *Study design*.

135 We conducted a parallel, randomized, controlled, open-label trial to determine the efficacy of dietary

136 BCAAem supplementation, as compared with diet advice, in the slow-down of both muscle and cognitive

137 deficit in malnourished community-dwelling men and women aged 80 years or older. Randomisation was

138 performed by computer generated tables to allocate treatments, with a simple randomization method; the

patients received a consecutive number after enrolment and were subsequently allocated to randomization
 list, according with Kim et al (32). The randomisation was carried out by the principal investigator, scientists
 performing lab measurement and statistical analyses were blind to treatment.

142 The inclusion criterion was malnutrition defined as MNA lower than 17. The MNA test is composed of 18 143 items that can be completed in less than 10 minutes. It provides a multidimensional assessment of senior 144 patients nutritional status taking into account four domains: anthropometry, general status, dietary habits, 145 and self-perceived health and nutrition states. Anthropometry includes the measurement of calf and arm 146 circumferences, Body Mass Index (BMI), calculated after the measurement of weight and height, and 147 questions about weight loss (4 items); general status comprehends 7 questions related to general health, 148 medication and mobility; the assessment of dietary habits comprehends 5 questions on the number of 149 meals, food and fluid intake and autonomy of feeding; 2 questions evaluate self-perceived health and 150 nutrition states (33,34).

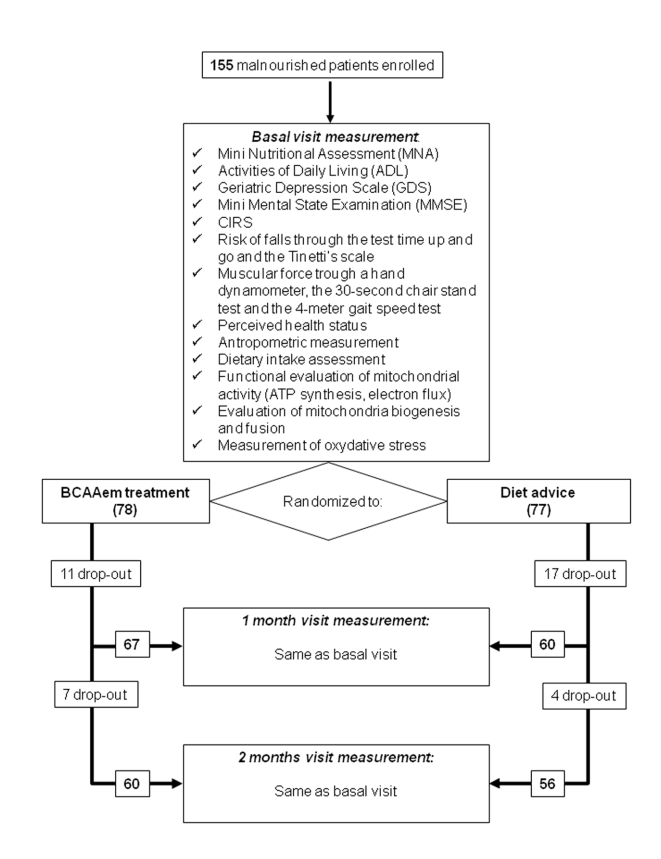
151 Exclusion criteria were known malignancy, life expectancy of less than two months, heart failure (NYHA IV), 152 end stage renal disease, liver cirrhosis (Child B-C), tube/percutaneous endoscopic gastrostomy feeding or 153 parenteral nutrition, Mini-Mental State Examination (MMSE) ≤ 18 and MNA>17. MMSE ≥ 18 identifies 154 patients with mild form of cognitive impairment, those patients generally do not have problems in 155 swallowing and are able to take drugs. We evaluated for inclusion 336 malnourished patients presenting to 156 our out-patients service for a geriatric evaluation, the evaluation was done by expert geriatricians, of the 157 evaluated patients 181 were excluded for presence of exclusion criteria; one hundred and fifty-five 158 malnourished elderly patients living at home in the community and who were admitted to the outpatients' 159 department of our Unit were enrolled. Patients were evaluated at baseline and randomised to receive diet 160 advice, summarised in an easy-to-use brochure for lay persons (77 patients) or to BCAAem supplements (78 161 patients, Aminotrofic®, kindly supplied by Errekappa Euroterapici S.p.A. and Professional Dietetics S.p.A, 2 sachets/day). Aminotrofic<sup>®</sup> is a BCAA enriched mixture, it contains Leucine (1,250 mg), Lysine (650 mg), 162 163 Isoleucine (625 mg), Valine (625 mg), Threonine (350 mg), Cystine (150 mg), Histidine (150 mg), 164 Phenylalanine (10 mg), Methionine (50 mg), Tyrosine (30 mg), Tryptophan (20 mg), Vitamin B 6 (0.1 mg),

Vitamin B1 (0.15 mg). We suggested the patients to take the BCAAem in the mid-morning and afternoon,
regardless of food ingestion. In order to check the compliance we asked the patients to bring back at center
the empty sachets at follow-up visit.

Diet advice comprised general advice on the meaning and consequences of malnutrition and dietary 168 169 recommendation based on the principle of "Food First" to maximize the patient nutritional intake from 170 regular food and drink according with the ESPEN Guidelines on Enteral Nutrition for geriatric patients (35). 171 According to the "Food First" approach we suggested the patients to increase the frequency of eating, 172 maximize the nutrient and energy density of food and drink and fortify food with the addition of fats and 173 sugars, suggested recommendations were given to the patients both orally by the physician and by the use 174 of a brochure; provided dietary recommendations are summarized in the Supplementary Table 1. 175 Period of patient enrolment: February 2013-September 2017. Patients were called back to the centre and 176 all the measurements were performed after 1 and 2 months. 116 patients completed the study (60 treated 177 with BCAAem and 56 with diet advice). In the BCAAem group, 2 patients died after the first month, 1 was 178 admitted to hospital within the first month, 10 patients did not return for the month-1 visit and 5 patients 179 did not return for the month-2 visit for personal reasons. In the diet advice group, 1 patient died during the 180 first month, 2 patients did not return for the month-1 visit since they were hospitalised, 14 patients did not 181 return for the month-1 visit and 4 patients for the month-2 visit for personal reasons, there were no 182 collateral effects. Only data from patients with complete follow up were included in the statistical analyses 183 (Fig. 1).

## 184 Fig.1. Diagram of the study design.

The diagram shows the study design and the number of patients at each visit in bold. The tests performedat each visit are specified.



189 The main outcome measures were muscle mass, strength and performance and mitochondrial ATP

190 production; secondary measurements were cognitive performance, nutritional status, health perceived

191 status, mitochondrial biogenesis and activity in peripheral blood mononuclear cells (PBMCs). The

192 measurements were carried out at baseline and after 1 and 2 months of treatment.

193 Clinical assessment.

194 Global clinical assessment: Self-sufficiency was measured by assessing the Katz Index of Activity of Daily 195 Living (ADL), which evaluates overall performance in six functions: bathing, dressing, going to toilet, 196 transferring, continence and feeding; cognitive performance was evaluated by MMSE that is a brief test 197 used to routinely track cognitive changes in an individual both cognitively intact or with severe cognitive 198 impairment over time. Patients' mood was evaluated by the short form of Geriatric Depression Scale (GDS), 199 the scale consists of 15 questions related to the patient's mood answered "yes" or "no". The cut-off point 200 adopted to define a patient as depressed is a GDS higher than 7 (36). Perceived health status was measured 201 by asking the patients to answer the question "How is your health in general?". The patients' answers: 202 "very good", "good", "fair", "bad" or "very bad" were rated from 5 (Very good) to 1 (Very bad), although 203 there is not yet full standardisation of the measurement of perceived health status across Organisation for 204 Economic Co-operation and Development (OECD) Countries, here we used a standard health interview 205 survey instrument used in the OECD Health Statistics 2007 (37), confirmed in OECD Health Statistics 2018 206 (available on line at http://www.oecd.org/els/health-systems/health-data.htm) and accepted in Italy. The 207 Cumulative Index Rating Scale (CIRS) was also recorded, this scale accounts for both the presence and the 208 severity of co-morbidities (38).

209 Nutritional status assessment: We assessed patients' dietary intake using the PROGEO software (Progeo 210 S.r.l., Italy), it provides an extensive food database and allows to record and to accurately estimate 211 patients' average nutritional intake. The Photo Intake tool helps patients to recognize the amount of food 212 by the visual weight method, ingested showing pictures of food in 3 portions, food quantities can also be recorded as conventional standard units (spoon, glass, cup, etc.). The software provides a large food 213 214 database and automatically displays patients' average daily calories and nutrients intake. The interview was 215 based on the recall method on 7 days making reference to the "standard week" as suggested by the 216 manufacturer. The interview was done by geriatricians trained by a nutritionist. 217 BMI was assessed by weighing patients by a precision scale and measuring their height using an altimeter

218 wall, BMI was calculated as weight in kg/height in meters squared. Percentage of fat mass was measured

219 using a plicometer (Mahr GMBH Esslingen), the Pollock, Schmidt and Jackson's formula on three sites 220 (triceps, subscapular and abdomen) was applied (39,40). Skinfold thickness measurements were performed 221 by trained staff according to standard technique: the skinfold thickness was measured by lifting a fold of 222 skin and subcutaneous fat away from the underlying muscle and bone, the skinfold thickness was measured 223 in duplicate with the plicometer. When a difference between the first and the second measurement 224 exceeded 6 mm, a third measurement was taken. The plicometer is applied 1cm from the ridge of skin; take 225 reading 3 seconds after application, to standardise any effects produced by deformation of tissues. The 226 triceps skinfold was measured at the back of the left arm, midway between the acromial process of the 227 scapula and the olecranon process of the ulna. The subscapular skinfold is picked up just under the lower 228 angle of the scapular lifted horizontally below the tip of right scapula. The abdominal skinfold was lifted 229 diagonal midway between umbilicus and right anterior superior iliac spine.

230 Muscle mass, strength and performance: Appendicular muscle mass was measured using arm and calf 231 circumference. Arm circumference was measured in duplicate to the nearest 0.001 m at a point midway 232 between the lateral projection of the acromion process of the scapula and the inferior margin of the 233 olecranon process of the ulna. The mean of the two measurements was used in the analyses. The calf 234 circumference was measured to the nearest 0.001 m on the left leg with the participant standing straight, 235 feet 20 cm apart, body weight equally distributed on both feet and at the level of the widest circumference 236 of the calf; measurements were taken according to the Longitudinal Aging Study Amsterdam (LASA -237 http://www.lasa-vu.nl/themes/physical/anthropometry.htm).

Muscle strength was measured via the hand grip test, using a hydraulic hand dynamometer (MSD, Europe)
(41) to assess muscle performance and mobility the Timed Up and Go (TUG) test, 30 seconds Chair Sit to

240 Stand (30-s CST) test and the 4 meters gait speed test were performed.

TUG is a simple test used to assess a person's mobility and requires both static and dynamic balance. TUG is performed by measuring the time that the patient takes to rise from a chair, walk three meters, turn around, walk back to the chair and sit down. TUG performed in ten seconds or less indicate normal mobility; 11–20 seconds are within normal limits for frail, elderly and disabled patients and greater than 20

seconds means that the person needs assistance outside and indicates further examination and intervention. A score of 30 seconds or more suggests that the person may be prone to falls (42). The 30-s CST allows the evaluation of lower body strength and to assess the fatigue effect due to the number of sitto-stand repetitions. It is performed with a chair without arms, the patient seated in the middle of the chair with the arms crossed over his/her chest, then is instructed to stand up as quickly as possible safely without using his/her arms. The number of stands the patients completed in 30 seconds is manually recorded (43).

The 4-meter gait speed test was performed using a stopwatch and measures the time, in seconds, the patients take to complete a 4-meter walk.

The risk of falls further was evaluated using the Tinetti Gait and Balance Instrument as follows: to test the patient's balance, the patient has to sit in a hard, armless chair and is asked to rise and stay standing, then turn 360° and sit back down. Next, the patient walks a few meters at a normal speed, turns, walks back and sits down. The evaluator observes several features and scores the patient's performance: the higher the score, the better the performance. The maximum score for Gait is 12 points, while the maximum for Balance is 16 points, with a total maximum for the overall Tinetti Instrument of 28 points. Score Interpretation: <19 high risk of falls, 19-28 low risk of falls.

260 Laboratory tests.

261 Functional evaluation of mitochondrial activity: In order to isolate mitochondrial fractions, blood cells were washed twice in ice-cold PBS, then lysed in 0.5 mL buffer A (50 mMTris, 100 mMKCl, 5 mM MgCl2, 1.8 mM 262 263 ATP, 1 mM EDTA, pH 7.2), supplemented with protease inhibitor cocktail III (Calbiochem), 1 mM PMSF and 264 250 mMNaF. Samples were clarified by centrifuging at 650×g for 3 min at 4°C, and the supernatant was 265 collected and centrifuged at 13000×g for 5 min at 4°C. This supernatant was discarded and the pellet 266 containing mitochondria was washed in 0.5 mL buffer A and suspended in 0.25 mL buffer B (250 mM 267 sucrose, 15 mM K2HPO4, 2 mM MgCl2, 0.5 mM EDTA, 5% w/v BSA). A 50 μL aliquot was sonicated and used 268 for the measurement of protein content, as reported in Campia et al, 2009 (44); the remaining part was 269 diluted to a protein concentration of 10  $\mu$ g/ $\mu$ L and stored at -80°C until the use. The activity of Complex I– 270 III was measured on 10  $\mu$ L of non-sonicated mitochondrial samples (44), suspended in 0.59 mL buffer C (5

mM KH2PO4, 5 mM MgCl2, 5% w/v BSA). Then 0.38 mL buffer D (25% w/v saponin, 50 mM KH2PO4, 5 mM
MgCl2, 5% w/v BSA, 0.12 mM cytochrome c-oxidized form, 0.2 mM NaN3) was added for 5 min at room
temperature. The reaction was started with 0.15 mM NADH and was followed for 5 min. The absorbance
was read using a Synergy HT Multi-Mode Microplate Reader (Bio-Tek Instruments, Winooski, VT). Results
were expressed as nmol reduced cytochrome C/min/mg mitochondrial proteins. In each experimental set,
the complex I inhibitor rotenone (100 µM) was added as an internal negative control. In the presence of
rotenone, the electron flux was reduced to below 5%.

278 The amount of ATP was measured on 20 µg of mitochondrial extracts using the ATP Bioluminescent Assay

279 Kit (FL-AA, Sigma Aldrich Co., St. Louis, MO). Data were converted into nmol/mg mitochondrial proteins,

using a previously set calibration curve.

Under these experimental conditions, the rate of cytochrome C reduction, expressed as nmol cytochrome C
 reduced/min/mg cell protein, was dependent on the activity of both Complex I and Complex III.

283 Real time PCR and assessment of mitochondria biogenesis and fusion: Real time PCR (RT-PCR) was used to

evaluate the mRNA levels of Cytochrome C Oxidase 1 and 4 (COX-1 and COX-4), Mitofusin-1 and 2 (MFN-1

and MFN-2), Nuclear Respiratory Factor-1 (NRF-1) and Mitochondrial Transcription Factor A (TFAM) from

whole blood nucleated cells.

287 Red cells were lysed in all peripheral blood samples, total nucleated cells were collected and dissolved in

288 TRIzol reagent (TRISure, Bioline Reagents Ltd, UK) and frozen at -80 °C until RNA extraction. RNA was

isolated using chloroform extraction and subsequent isopropanol precipitation according to the

290 manufacturer's protocol. 1 μg of RNA was reverse-transcribed to single-stranded cDNA using the SensiFAST

291 cDNA Synthesis Kit (Bioline Reagents Ltd, UK). RT-PCR was performed using the SensiFAST SYBR Hi-ROX Kit

292 (Bioline Reagents Ltd, UK). The housekeeping control gene was β-actin, and gene expression was quantified

using the  $2^{-\Delta\Delta Ct}$  method. The primers used (Invitrogen, California, USA) are shown in **Supplementary Table** 

294 **2**.

All the lab experiments were performed in duplicate, data presented are averages of the duplicates. Thecoefficient of variation intra-operator ranges between 0.03 and 1.00 for all the measurements.

Oxidative stress: To determine the oxidative stress level, we measured plasma thiobarbituric acid reactive
 substances (TBARs), as indicators of lipid peroxidation, using ELISA (TBARS Assay Kit, Cayman Chemical, MI,
 USA), according to the manufacturer's protocol.

300 Statistical analyses: The sample size was calculated on both clinical and lab outcomes; amongst clinical 301 outcome muscle mass was used; in particular sample size provide an 85% power (p<0.05), 50 patients per 302 group have to be enrolled to detect a difference (alpha error = 0.05) of at least 2% variation in muscle mass, 303 based on a study on the effect of BCAAs administration on muscle mass and performance in humans (45). 304 As the patients were old and frail we assumed a possible 35% of drop out at the follow-up. In order to 305 calculate sample size for lab tests we considered as significant an increase of 1.5 fold in ATP production as 306 shown by D'Antona et al with BCAAem in aged mice (28), data on ATP production in humans by PBMCs 307 derives from Avis et al (46) based on this analyses sample size was calculated to provide 95% power 308 (p<0.05) to detect a 1.5-fold difference in ATP production was 13 patients per group.

All the analysed variables were tested for normality by the kurtosis test, TUG, 30-s CST, 4 meters walking
test, TBARs, electron flux were non-Gaussian.

To evaluate possible differences between patients treated with BCAAem or diet advice at baseline the
 patients were compare by one-way ANOVA for Gaussian variables and by the Mann-Whitney U test for
 non-Gaussian ones. Gender was compared amongst patients treated with BCAAem or diet adviceby χ2 test.

The effect of treatment was evaluated per protocol using the two-way ANOVA for repeated measurements
for Gaussian variables, non-Gaussian variables were evaluated after logarithmic transformation. To
evaluate possible influences of mitochondrial function on muscle and cognitive performance six linear
regression models (GLM) were fitted, between ATP and electron flux and TUG, 30-s CST, Tinetti and 4
meters walking test, hand grip and MMSE, non-Gaussian variables were logarithmically transformed.
SPSS 24.0 were used for the analyses and p<0.05 was considered statistically significant. Graphs were</li>

drawn using GraphPad 7.0 for Windows.

321 Ethics Committee approval and consent to participate.

322 The study was approved by the Ethics Committee of our Hospital ("Comitato Etico Interaziendale A.O.U.

323 Città della Salute e della Scienza di Torino - A.O. Ordine Mauriziano - A.S.L. TO1", protocol number

0002637), in accordance with the ethical standards of the Declaration of Helsinki and its subsequent

amendments. Informed consent was obtained from all individual participants included in the study. The full

- 326 protocol is available upon request to the corresponding author.
- 327

# 328 Results

329 Patients treated with BCAAem or diet advice were comparable for all the clinical variables analysed, this

excludes possible selection bias that could influence our results (**Supplementary Table 3**). Compliance to

- BCAAem was good, none of the patients have a compliance lower than 75%, the compliance ranges
- 332 between 75% and 90%.
- 333 Treatment significantly improves general health and cognitive performance.
- Patients' general health measured by perceived health status equally improved in both treatment groups
- and significantly correlated with nutritional status (MNA: R=0.50, p<0.0001; fat percentage: R=0.26,

p=0.005) at the end of the follow-up period. Also, the mood measured by GDS significantly improved. The

337 level of independence was not significantly influenced by treatment.

338 Patients' overall cognitive performance measured by MMSE significantly improved in patients treated with

BCAAem, not in patients treated with diet advice; MMSE significantly correlated with MNA (R=0.28,

p=0.002) as well as GDS (R=-0.32, p<0.0001) at the end of the follow-up period (**Table 1**).

## 341 Treatment significantly improves nutritional status.

342 Patients adhered to the dietary recommendation as shown by the increased caloric intake. Caloric intake

343 increased in both groups and was particularly consistent in the group treated with diet advice, where

dietary recommendations were reinforced by the use of an easy-to-use illustrated brochure, in this group

- also protein intake was significantly higher. During treatment, we observed a significant improvement in
   nutritional status measured by MNA, fat mass and BMI in both groups (Table 2).
- 347 BCAAem treatment increases muscle mass, strength and performance.
- 348 Muscle mass measured by calf and arm circumferences significantly increased in both groups as well as
- 349 muscular strength measured by hand grip strength (**Table 3**).
- 350 To evaluate whether muscular performance was influenced by treatment, we performed the TUG, the 30-s
- 351 CST test to evaluate both performance and resistance to fatigue. Muscular performance improved with
- treatment as did muscle mass. Risk of falls was measured using the Tinetti scale and the TUG, mobility was
- 353 measured using the 4-meter gait speed test, these tests improved equally in the two groups (**Table 3**).
- 354 BCAAem improve bioenergetic capacity of PBMCs.
- 355 Here we show that ATP production and electron flux significantly increased over time only in mitochondria
- 356 from patients treated with BCAAem, and that BCAAem maintain oxidative stress at baseline values,
- 357 whereas, in patients treated with diet advice, oxidative stress increased over time (**Table 4**).
- 358 To model the relationship between mitochondria stimulation on PBMCs and effects of BCAAem or diet
- advice on muscular and cognitive performance we applied a linear regression approach.
- 360 Our data showed that mitochondrial ATP production significantly predicts balance measured by Tinetti
- after 2 months of treatment and 4 meters walking test after 1 month of treatment (**Table 5**). MMSE is
- 362 significantly predicted by ATP after 1 month and by electron flux after 2 months of treatment (**Table 6**).
- 363 We also evaluated the effect of BCAAem treatment on some of the main mitochondrial biogenesis and
- fusion markers. Our data showed that treatment increases the expression of COX-1 and COX-4 and TFAM,
- 365 whereas NRF-1 shows only a non-significant trend towards the increase, significant differences versus
- 366 baseline levels were measured only in patients treated with BCAAem. The expression of MFN-1 and MFN-2

was increased although not significantly by treatment; in the BCAAem treated patients we observed an
 increased expression of these two molecules after one month of therapy (Table 7).

369

370 Discussion

As life expectancy increases, adequate nutrition is fundamental for successful aging, here, we confirm that amelioration of nutritional status is associated with improvement in general health status, muscle and cognitive performance in old malnourished patients, and show that this may be due to an improvement of their mitochondrial bioenergetics profile and decreasing oxidative stress.

375 Here we show that the diagnosis of malnutrition and its treatment, albeit using different approaches, is 376 fundamental in improving patients' general health and nutritional status. Indeed, in both our treatment 377 groups, there was an improvement in MNA, and weight gain, however the increase in caloric and protein 378 intake was higher in patients treated only with diet advice; the use of this approach instead of the use of an 379 isonitrogenous mix of non-essential aminoacids with a double blind design may be considered as a 380 limitation of the study, however the use of the "food first" approach underlines the role of the physician's 381 counselling in patients' adherence to diet. Another possible limitation of the study is the use of a "non-382 standard" recall method for diet evaluation, here we use a recall on the 7 days before the interview as our 383 old patients usually follow a standard diet with little variation day by day, the evaluation of 7 days allow us 384 to make a more comprehensive evaluation asking the patients "what is your usual meal (breakfast, lunch, 385 dinner and snacks)? Do you change your meal during one week? If yes when and how during the previous 386 week?". The improvement in cognitive performance, measured by MMSE, was significant in patients taking 387 BCAAem supplements: indeed, patients treated with BCAAem gain, on average, 1.2 points of MMSE. This 388 result is in accordance with a previous study in a cohort of patients with severe chronic obstructive 389 pulmonary disease. However, in this study, the increase in cognitive performance was associated with 390 improved pulmonary gas exchange and there was no mechanistic explanation (47). The effect of the 391 administration of BCAAs in cognitive function was also previously assessed in severely brain damaged

392 patients with hepatic encephalopathy or traumatic brain injuries: in these patients, intravenous BCAAs 393 improve consciousness, assessed using the Glasgow Coma Scale and performance, assessed using the 394 Disability Rating Scale. However, no cognitive tests were performed in these studies and the underlying 395 mechanism was not investigated (48–52). A mechanistic explanation has been provided by the study of 396 animal models of brain injury, demonstrating the efficacy of dietary supplementation with BCAAs in 397 promoting cognitive performance, by restoring hippocampal function, given that BCAAs act as glutamate 398 and GABA precursors (53). In humans, a recent, very large retrospective study showed an association 399 between serum level of isoleucine, leucine and valine with a lower risk of dementia. This study does not 400 report any causal mechanism (54).

401 Poor nutrition and, in particular, protein-energy deficit further accelerate the loss of muscle mass and 402 function associated with age: sarcopenia (55,56). Sarcopenia increases the risk of falls, disability, frailty, loss 403 of independence and death increasing healthcare costs (57,58). Here, we show that intervention on 404 malnutrition can improve muscle mass and performance. In particular, we show an improvement in 405 balance by the Tinetti balance test and the TUG test, with a consequent reduction of the risk of falls and 406 fall-related injuries in a population with increased risk of falls (average TUG higher than 14 seconds). 407 Treatment increases muscle performance, increasing mobility, and resistance to fatigue; the improved 408 performance in the 4-meter gait speed test observed during treatment demonstrates an overall increase in 409 health and the performance status of the patients according to previous study (62). Also, muscle strength 410 measured using hand dynamometer increased in both treatment groups.

Taken together, these data demonstrate that dietary intervention can counteract the decrease in physical and cognitive performance in old malnourished patients and that loss of muscle mass and function can be countered using an appropriate treatment strategy in a rapidly aging population.

Several studies suggest a major role of mitochondrial dysfunction as a major contributor to aging and age related diseases [for a review, see Cedikova et al, 2016 (63)]. Mitochondrial biogenesis, ATP production and
 oxidative phosphorylation capacity decrease with aging and production of reactive oxygen species,

417 damaged mitochondrial DNA and protein increase [for a review, see Cedikova et al, 2016 (63)]. In this 418 study, we further investigate the mechanisms underlying the better clinical effects obtained with the 419 administration of BCAAem, besides diet advice, by evaluating the treatment effects on mitochondrial 420 function, biogenesis and fusion, as well as oxidative stress. In animal models, the administration of BCAAs 421 has been shown to be effective in increasing the number and the function of mitochondria (28). In a small 422 cohort of young healthy obese and non-obese subjects, the infusion of a mixture of aminoacids increased ATP production rates of muscular mitochondria in lean, but not in obese subjects (64). The increase in size, 423 424 number and energy production of mitochondria that follows the administration of BCAAem was associated 425 with better muscle performance, on both at skeletal and cardiac muscle level in mice (28). Here we 426 evaluated mitochondria from PBMCs, it has been previously shown that mitochondria isolated from 427 skeletal muscle and from PBMCs have a similar bioenergetics profile and are associated with gait speed in 428 older adults (65), as regards neurological tissues other authors measured mitochondrial function in PBMCs 429 and correlates its reduction with neurodegeneration (66). According with these observations we show that 430 mitochondrial activity measured using ATP production and electron flux increase is associated with both 431 cognitive and muscular performance amelioration in elderly patients; these data are particularly interesting 432 as they suggest a possible non-invasive and reliable measure to follow up treatment outcomes. We show 433 that the increase of TFAM, mitochondrial respiratory chain (COX-1 and COX-4) and mitofusin gene 434 expressions correlate with an increase in ATP production and electron flux, and suggest that BCAAem 435 treatment induces biogenesis, activity and fusion of mitochondria, stimulating at the end the bioenergetic 436 capacity of PBMCs.

437 Mitochondrial biogenesis and fusion are particularly increased after the first month of treatment.

Afterwards, there is a plateau with no further increase, whereas mitochondrial activity increases during the entire follow-up period. This suggests that, after an increase in the number and fusion of mitochondria, the organelles maintain an increased function without any further increase in their number.

Other than the effect on muscle mass and function, treatment of malnutrition is associated with improved
general health. This may be due to better energy production associated with an increase in respiratory

chain activity and a decrease in oxidative stress, in an experimental model of progeroid aging characterised
by increased DNA damage, boosting mitochondrial preserves mammalian health and increases longevity
(67).

It is well known that oxidative damage is one of the components of aging: the increase in free radical production and the decrease in the defence against oxidative stress cause molecular alteration and functional decay; the free radical theory of aging suggests that oxidative stress is an important factor in age-associated diseases (68). Here, we show that BCAAem supplementation lowers the levels of oxidative stress in elderly patients.

In conclusion, this study, for the first time, suggests that BCAAem treatment in old malnourished patients
 could be a good strategy able to ameliorate the bioenergetic capacity of PBMCs, this effect may partially
 explain the positive trend on muscle and cognitive performance in these patients.

454

## 455 **Conflict of interest statement and funding**

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460

## 461 Statement of authorship

FS and IB performed the lab experiments, acquired and analysed the lab data, and participated in drafting and critically revising the manuscript. CRavetta, GC, FD, CF, FGP and PP performed the clinical evaluation of patients and managed the dataset. MM, EN, CRiganti, CRuocco and GCI participated in the study design and were major contributors in writing the manuscript. PD designed the study, performed the statistical analyses and wrote the paper. All authors read and approved the final manuscript.

# 469 Availability of data and materials

- 470 The datasets generated and/or analysed during the current study are not publicly available but are
- 471 available from the corresponding author upon reasonable request.

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# **Table 1. Global clinical assessment at baseline and follow-up according to treatment.**

- 658 The table shows multiple T-test and Two-way ANOVA for multiple measure results. Values are shown as
- 659 mean ± SE and 95% CI of the difference between basal, 1 and 2 months of treatment.
- 660 \* Denotes differences between baseline and 1 month; \*\* Denotes differences between baseline and 2

661 months; \$ Denotes differences between 1 and 2 months; Significant values are in bold.

	Perceived health status						
	BCAAem <sup>1</sup>		Diet advice		Two-way ANOVA		
	Mean±SE	95% CI of difference	<i>Mean±SE</i>	95% CI of difference	Effect of	p	
baseline	3±0.1	-0.56 to -0.02*	2.9±0.1	-0.56 to -0.02*	Time	< 0.0001	
1 month	3.3±0.1	-0.5 to 0.02\$	3.2±0.1	-0.6 to -0.04\$	Treatment	0.8437	
2 months	3.5±0.1	-0.8 to -0.3**	3.5±0.2	-0.8 to -0.3**	Interaction	0.9300	
		MMSE <sup>2</sup>					
	B	CAAem <sup>1</sup>	Die	t advice	Two-way	ANOVA	
	Mean±SE	95% CI of difference	Mean±SE	95% CI of difference	Effect of	р	
baseline	25.1±0.4	-1.1 to -0.04*	25.6±0.4	-0.2 to 0.8*	Time	0.0139	
1 month	25.7±0.4	-0.8 to 0.2\$	25.3±0.5	-1.0 to 0.06\$	Treatment	0.9422	
2 months	26±0.4	-1.4 to -0.3**	25.8±0.4	-0.7 to 0.4**	Interaction	0.0171	

	GDS <sup>3</sup>					
	BCAAem <sup>1</sup>		Diet advice		Two-way ANOVA	
	Mean±SE	95% CI of difference	<i>Mean±SE</i>	95% CI of difference	Effect of	p
baseline	6.1±0.3	-0.4 to 0.8*	6.0±0.4	-0.2 to 0.9*	Time	0.0084
1 month	5.9±0.4	-0.3 to 0.9\$	5.7±0.4	-0.3 to 0.8\$	Treatment	0.7448
2 months	5.6±0.4	-0.1 to 1.1**	5.4±0.3	0.01 to 1.2**	Interaction	0.9184
			AD	L <sup>4</sup>		
	B	CAAem <sup>1</sup>	Die	t advice	Two-way	ANOVA
	Mean±SE	95% CI of difference	Mean±SE	95% CI of difference	Effect of	p
baseline	10.2±0.3	-0.2 to 0.4*	9.7±0.3	-0.2 to 0.5*	Time	0.3569
1 month	10.1±0.3	-0.2 to 0.5\$	9.5±0.3	-0.5 to 0.1\$	Treatment	01130
2 months	10.0±0.3	-0.1 to 0.6**	9.7±0.3	-0.3 to 0.30**	Interaction	0.2294

<sup>1</sup>Branched Chain Amino Acid Enriched Mixture; <sup>2</sup>Mini-Mental State Examination; <sup>3</sup>Geriatric Depression Scale;

664 <sup>4</sup>Activity of Daily Living.

# **Table 2. Patients' nutritional status at baseline and follow-up according to treatment.**

- 668 The table shows multiple T-test and Two-way ANOVA for multiple measure results. Values are shown as
- 669 mean ± SE and 95% CI of the difference between basal, 1 and 2 months of treatment.
- 670 \* Denotes differences between baseline and 1 month; \*\* Denotes differences between baseline and 2
- 671 months; \$ Denotes differences between 1 and 2 months; Significant values are in bold.

			Caloric Inta	ake (Kcal/day)		
	B	CAAem <sup>1</sup>	Die	Diet advice		ANOVA
	Mean±SE	95% CI of	Mean±SE	95% CI of	Effect of	р
		difference		difference		
baseline	1095±36	-137 to -15.8*	1042±29	-169 to -48*	Time	<0.0001
1 month	1172±36	-80 to 42.4\$	1151±31	-168 to -47\$	Treatment	0.9740
2 months	1189±36	-156 to -34.1**	1259±38	-277 to -156**	Interaction	0.0031
		C	aily protein In	take (g/Kg weight)		
	B	CAAem <sup>1</sup>	Die	et advice	Two-way	ANOVA
	Mean±SE	95% CI of	Mean±SE	95% CI of	Effect of	p
		difference		difference		
baseline	0.85±0.02	-0.03 to 0.09*	0.87±0.02	-0.20 to -0.04*	Time	0.0874
1 month	0.83±0.02	-0.06 to 0.06\$	0.97±0.02	-0.04 to 0.08\$	Treatment	0.0001

2 months	0.83±0.02	-0.03 to 0.09**	0.96±0.02	-0.14 to -0.02**	Interaction	0.0007	
			N	1NA <sup>2</sup>			
	B	CAAem <sup>1</sup>	Di	et advice	Two-way	ANOVA	
	Mean±SE	95% CI of	Mean±SE	95% CI of	Effect of	p	
		difference		difference			
baseline	14.8±0.26	-0.03 to 0.09*	14.9±0.28	-0.16 to -0.04*	Time	<0.0001	
1 month	18.0±0.43	-0.06 to 0.06\$	17.8±0.38	-0.04 to 0.08\$	Treatment	0.9057	
2 months	18.9±0.5	-0.03 to 0.09**	18.8±0.47	-0.14 to -0.02**	Interaction	0.8366	
			Fat n	nass (%)			
	B	CAAem <sup>1</sup>	Diet advice Two		Two-way	way ANOVA	
	Mean±SE	95% CI of	Mean±SE	95% CI of	Effect of	p	
		difference		difference			
baseline	18.8±0.94	-1.68 to 0.61*	20.2±0.88	-1.7 to 0.6*	Time	0.0009	
1 month	19.3±0.92	-2.16 to 0.13\$	20.8±0.88	-1.6 to 0.7\$	Treatment	0.1708	
2 months	20.3±0.94	-2.70 to -0.41**	21.2±0.88	-2.2 to 0.1**	Interaction	0.6438	
			E	3MI <sup>3</sup>			
	B	CAAem <sup>1</sup>	Di	et advice	Two-way	ANOVA	
	Mean±SE	95% CI of	Mean±SE	95% CI of	Effect of	p	
		difference		difference			

baseline	20.5±0.42	-1.42 to 0.10*	20.8±0.41	-1.38 to 0.14*	Time	0.0010
1 month	21.2±0.68	-0.92 to 0.60\$	21.5±0.42	-0.81 to 0.71\$	Treatment	0.7239
2 months	21.3±0.66	-1.57 to -0.05**	21.5±0.41	-1.43 to 0.09**	Interaction	0.9482

673 <sup>1</sup>Branched Chain Amino Acid Enriched Mixture; <sup>2</sup>Mini Nutritional Assessment test; <sup>3</sup>Body Mass Index.

# Table 3. Muscle mass, strength and performance at baseline and follow-up according to treatment.

- 677 The table shows multiple T-test and Two-way ANOVA for multiple measure results. Values are shown as
- 678 mean ± SE and 95% CI of the difference between basal, 1 and 2 months of treatment.
- <sup>679</sup> \* Denotes differences between baseline and 1 month; \*\* Denotes differences between baseline and 2
- 680 months; \$ Denotes differences between 1 and 2 months; Significant values are in bold.

		Muscle mass							
		Calf circumference (cm)							
	BCAAem <sup>1</sup>		Die	Diet advice		ANOVA			
	Mean±SE	95% CI of	Mean±SE	95% CI of	Effect of	p			
		difference		difference					
baseline	30.4±0.35	-1.14 to -0.04*	30.7±0.43	-1.08 to 0.02*	Time	0.0004			
1 month	31.0±0.38	-0.87 to 0.23\$	31.2±0.40	-0.54 to 0.56\$	Treatment	0.8560			
2 months	31.3±0.39	-1.45 to -0.36**	31.19±0.39	-1.07 to 0.03**	Interaction	0.4521			
			Arm circum	ference (cm)					
	В	CAAem <sup>1</sup>	Die	et advice	Two-way	ANOVA			
	Mean±SE	95% CI of	Mean±SE	95% CI of	Effect of	p			
		difference		difference					
baseline	22.7±0.36	-0.75 to 0.23*	23.0±0.40	-0.83 to 0.15*	Time	0.0045			
1 month	23.0±0.37	-0.77 to 0.21\$	23.3±0.40	-0.54 to 0.44\$	Treatment	0.6754			

2 months	23.3±0.38	-1.03 to -0.04**	23.4±0.44	-0.88 to 0.10**	Interaction	0.7351
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			Muscle	e strength		
			Hand	grip (Kg)		
	B	CAAem <sup>1</sup>	Die	et advice	Two-way ANOVA	
	Mean±SE	95% CI of	Mean±SE	95% CI of	Effect of	p
		difference		difference		
baseline	17.9±1.0	-2.01 to 0.47*	17.9±1.0	-1.84 to 0.65*	Time	0.0474
1 month	18.5±1.0	-1.6 to 0.87\$	18.7±0.9	-1.0 to 1.50\$	Treatment	0.7796
2 months	18.3±1.0	-2.40 to 0.10**	19.1±1.0	-1.59 to 0.90**	Interaction	0.5231
			TUG	TUG (sec) <sup>2</sup> Diet advice Two-way AN		
	B	CAAem <sup>1</sup>	Die			ANOVA
	Mean±SE	95% CI of	Mean±SE	95% CI of	Effect of	р
		difference		difference		
baseline	19.8±2.14	1.5 to 7.6*	20.5±1.5	-1.2 to 4.9*	Time	0.0001
1 month	15.2±1.0	-2.9 to 3.2\$	18.7±1.6	-2.1 to 4.0\$	Treatment	0.2780
2 months	15.1±1.1	1.6 to 7.8**	17.7±1.7	-0.3 to 5.9**	Interaction	0.3215
			30-	30-s CST <sup>3</sup>		
	BO	CAAem <sup>1</sup>	Diet	t advice	Two-way /	ANOVA
	Mean±SE	95% CI of	Mean±SE	95% Cl of	Effect of	p

		difference		difference		
baseline	6.8±0.5	-2.6 to -0.7*	6.0±0.5	-2.4 to -0.5*	Time	<0.0001
1 month	8.4±0.6	-1.0 to 0.88\$	7.4±0.5	-1.6 to 0.3\$	Treatment	0.3328
2 months	8.5±0.7	-2.7 to -0.7**	8.1±0.6	-3.0 to -1.1**	Interaction	0.5810

- <b>-</b>		
	ine	etti

	BCAAem <sup>1</sup>		Diet	Diet advice		ANOVA
	Mean±SE	95% CI of	Mean±SE	95% CI of	Effect of	p
		difference		difference		,
baseline	20.4±0.8	-2.1 to -0.1*	18.3±0.8	-3.2 to -1.1*	Time	< 0.0001
1 month	21.5±0.7	-1.7 to 0.3\$	20.4±0.8	-1.3 to 0.7\$	Treatment	0.1503
2 months	22.2±0.7	-2.8 to -0.8**	20.7±0.9	-3.4 to -1.4**	Interaction	0.2076

4 meters walking test (sec)

	BCAAem <sup>1</sup>		Diet advice		Two-way ANOVA	
	Mean±SE	95% CI of	Mean±SE	95% CI of	Effect of	p
		difference		difference		
baseline	8.2±0.6	-0.3 to 1.7*	9.8±0.7	0.4 to 2.3*	Time	<0.0001
1 month	7.5±0.6	-0.7 to 1.3\$	8.4±0.7	-0.6 to 1.4\$	Treatment	0.1684
2 months	7.2±0.6	0.04 to 2.0**	8.0±0.7	0.8 to 2.8**	Interaction	0.3955

## **Table 4. Mitochondrial activity and oxidative stress at baseline and follow-up according to treatment.**

- 684 The table shows multiple T-test and Two-way ANOVA for multiple measure results. Values are shown as
- 685 mean ± SE and 95% CI of the difference between basal, 1 and 2 months of treatment.
- <sup>686</sup> \* Denotes differences between baseline and 1 month; \*\* Denotes differences between baseline and 2
- 687 months; \$ Denotes differences between 1 and 2 months; Significant values are in bold.

			ATP (chang	es vs baseline)		
	BCAAem <sup>1</sup>		Diet advice		Two-way ANOVA	
	Mean±SE	95% CI of	Mean±SE	95% CI of	Effect of	p
		difference		difference		
baseline	1.00±0.00	-0.45 to -0.15*	1.00±0.00	-0.13 to 0.17*	Time	0.0001
1 month	1.30±0.09	-0.28 to 0.016\$	0.98±0.01	-0.16 to 0.14\$	Treatment	0.0005
2 months	1.43±0.10	-0.58 to -0.28**	0.99±0.02	-0.14 to 0.16**	Interaction	0.0001
			Electron flux (ch	anges vs baseline)		
	B	BCAAem <sup>1</sup>	Die	t advice	Two-way	ANOVA
	Mean±SE	95% CI of	Mean±SE	95% CI of	Effect of	р
		difference		difference		
baseline	1.00±0.00	-0.38 to -0.13*	1.00±0.00	-0.13 to 0.13*	Time	< 0.0001
1 month	1.26±0.05	-0.37 to -0.12\$	1.00±0.01	-0.14 to 0.12\$	Treatment	< 0.0001

2 months	1.50±0.09	-0.62 to -0.38**	1.01±0.04	-0.14 to 0.12**	Interaction	< 0.0001
			TBAR	s² (μg/M)		
	B	CAAem <sup>1</sup>	Die	et advice	Two-way	ANOVA
	Mean±SE	95% CI of	Mean±SE	95% CI of	Effect of	p
		difference		difference		
baseline	2.3±0.4	-2.8 to 1.2*	4.1±0.7	-3.02 to 0.97*	Time	0.0007
1 month	3.0±0.57	-2.4 to 1.6\$	4.5±0.9	-4.61 to -0.61\$	Treatment	0.0289
2 months	3.2±0.70	-3.1 to 0.85**	6.7±1.3	-5.64 to -1.64**	Interaction	0.0332

<sup>1</sup>Branched Chain Amino Acid Enriched Mixture; <sup>2</sup>Thiobarbituric Acid Reactive Substances.

- **Table 5. Muscle performance and balance.**
- 693 The table shows the results for linear regression models (GLM), non-Gaussian variables indicated by \* were
- 694 logarithmically transformed. Significant values are in bold.

Dependent variable	Co-variate	β±SE	t	р	95% CI
Tinetti baseline	Slope	18.9±2.7	7.0	0.000	13.5 to 24.4
	Electronflux baseline	-0.02±0.01	-1.4	0.179	-0.05 to 0.009
	ATP baseline	0.05±0.03	1.5	0.134	-0.016 to 0.119
Tinetti 1 month	Slope	20.0±2.9	6.8	0.000	14.1 to 25.8
	Electronflux 1 month*	0.000±0.013	0.02	0.984	-0.03 to 0.26
	ATP 1 month	0.011±0.04	0.32	0.752	-0.06 to 0.08
Tinetti 2 months	Slope	48.9±13.5	3.6	0.001	21.8 to 75.9
	Electronflux 2 months*	0.05±0.04	1.5	0.131	-0.02 to 0.122
	ATP 2 months	-13.5±6.2	-2.2	0.033	-25.9 to -1.1
Dependent variable	Co-variate	β±SE	t	р	95% CI
4 meters walking test	Slope	0.89±0.08	11.2	0.000	0.7 to 1.0
baseline*	Electronflux baseline	0.000±0.000	0.8	0.394	0.000 to 0.001
	ATP baseline	-0.001±0.001	-1.4	0.175	-0.003 to 0.001

4 meters walking test 1	4 meters walking test 1 Slope		9.6	0.000	0.76 to 1.16	
month*	Electronflux 1 month*	0.000±0.000	0.40	0.691	-0.001 to 0.001	
	ATP 1 month	-0.002±0.001	-2.0	0.045	-0.005 to -6.1E <sup>-5</sup>	
4 meters walking test 2	Slope	0.42±0.46	0.92	0.361	-0.49 to 1.33	
months*	Electronflux 2 months*	-0.002±0.001	-1.73	0.089	-0.004 to 0.000	
	ATP 2 months	0.23±0.21	1.08	0.283	-0.19 to 0.644	

# 697 **Table 6. Cognitive performance.**

698 The table shows the results for linear regression models (GLM), non-Gaussian variables indicated by \* were

699 logarithmically transformed. Significant values are in bold.

Dependent variable	Co-variate	β±SE	t	р	95% CI
MMSE <sup>1</sup> baseline	Slope	24.3±1.4	17.2	0.000	21.5 to 27.2
	Electronflux baseline	0.002±0.007	0.22	0.829	-0.01 to 0.02
	ATP baseline	0.02±0.018	0.99	0.327	-0.02 to 0.05
MMSE <sup>1</sup> 1 month	Slope	22.7±0.17	12.9	0.000	19.2 to 26.20
	Electronflux 1 month*	-0.002±0.08	-0.27	0.790	-0.02 to 0.013
	ATP 1 month	0.048±0.02	2.29	0.026	0.006 to 0.09
MMSE <sup>1</sup> 2 months	Slope	29.9±7.9	3.8	0.000	14.01 to 45.7
	Electronflux 2 months*	0.05±0.02	2.5	0.014	0.01 to 0.09

		ATP 2 months	-3.31±3.6	-0.9	0.366	-10.6 to 3.9
700	<sup>1</sup> Mini-Mental State Exam	ination.				

Table 7. Mitochondria biogenesis and fusion at baseline and follow-up according to treatment. The table
 shows multiple T-test and Two-way ANOVA for multiple measure results. Values are shown as mean ± SE
 and 95% CI of the difference between basal, 1 and 2 months of treatment.

\* Denotes differences between baseline and 1 month; \*\* Denotes differences between baseline and 2

707 months; \$ Denotes differences between 1 and 2 months; Significant values are in bold.

	COX-1 <sup>1</sup> (changes vs baseline)								
	BCAAem <sup>2</sup>		Die	Diet advice		Two-way ANOVA			
	Mean±SE	95% Cl of	Mean±SE	95% CI of	Effect of	р			
		difference		difference					
baseline	1.0±0.0	-23.8 to -1.9*	1.0±0.0	-11.4 to 10.6*	Time	0.1155			
1 month	13.9±8.5	-4.4 to 17.5\$	1.4±0.3	-13.2 to 8.7\$	Treatment	0.1967			
2 months	7.3±3.6	-17.3 to 4.7**	3.7±1.2	-13.6 to 8.3**	Interaction	0.1409			
			COX-4 <sup>3</sup> (chan	ges vs baseline)					
	BCAAem <sup>2</sup> Diet advice		Two-way A	ANOVA					
	Mean±SE	95% CI of	Mean±SE	95% CI of	Effect of	p			
		difference		difference					
baseline	1.0±0.0	-2.3 to -0.10*	1.0±0.0	-1.39 to 0.76*	Time	0.0459			
1 month	2.2±0.7	-0.7 to 1.47\$	1.3±0.09	-1.1 to 1.04\$	Treatment	0.2373			

2 months	1.8±0.5	-1.9 to 0.3**	1.3±0.19	-1.4 to 0.73**	Interaction	0.3786		
			TFAM <sup>4</sup> (chan	ges vs baseline)				
	BCAAem <sup>2</sup> Diet advice Two-way ANOVA							
	Mean±SE	95% CI of	Mean±SE	95% CI of	Effect of	p		
		difference		difference				
baseline	1.0±0.0	-6.9 to -0.6*	1.0±0.0	-3.8 to 2.5*	Time	0.0178		
1 month	4.8±1.0	-2.4 to 3.9\$	1.7±0.5	-4.5 to 1.8\$	Treatment	0.0932		
2 months	4.2±1.15	-6.2 to 0.1**	3.0±1.5	-5.1 to 1.2**	Interaction	0.2235		
			NRF-1 <sup>5</sup> (chan	ges vs baseline)				
	B	CAAem <sup>2</sup>		ges vs baseline) t advice	Two-way A	ANOVA		
			Die	t advice				
	BC Mean±SE	C <b>AAem<sup>2</sup></b> 95% CI of difference			<b>Two-way A</b> Effect of	ANOVA p		
baseline		95% CI of	Die	t advice 95% Cl of				
baseline 1 month	Mean±SE	95% CI of difference	Die Mean±SE	<b>t advice</b> 95% Cl of difference	Effect of	p		
	Mean±SE 1.0±0.0	95% CI of difference -20.2 to 3.5*	Die Mean±SE 1.0±0.0	t advice 95% Cl of difference -13.5 to 10.2*	Effect of Time	р 0.6599		

MFN-1 <sup>6</sup> (changes vs baseline)								
BCAAem <sup>2</sup>		Diet	Diet advice		Two-way ANOVA			
Mean±SE	95% Cl of	Mean±SE	95% CI of	Effect of	p			
	difference		difference					
1.0±0.0	-22.4 to -2.1*	1.0±0.0	-10.8 to 9.4*	Time	0.0746			
13.2±7.7	-7.0 to 13.3\$	1.7±0.3	-10.3 to 10.0\$	Treatment	0.1648			
10.1±6.1	-19.3 to 1.0**	1.8±0.3	-11.0 to 9.3**	Interaction	0.1320			
		MFN-2 <sup>7</sup> (chan	ges vs baseline)					
BCAAem <sup>2</sup> Diet advice				Two-way A	NOVA			
Mean±SE	95% Cl of	Mean±SE	95% CI of	Effect of	p			
	difference		difference					
1.0±0.0	-11.6 to -1.1*	1.0±0.0	-6.0 to 4.5*	Time	0.0772			
7.3±4.1	-1.9 to 8.6\$	1.7±0.3	-5.9 to 4.6\$	Treatment	0.2046			
3.9±1.6	-8.2 to 2.3**	2.4±0.5	-6.6 to 3.9**	Interaction	0.1810			
	Mean±SE 1.0±0.0 13.2±7.7 10.1±6.1 BC Mean±SE 1.0±0.0 7.3±4.1	Mean±SE       95% Cl of         difference         1.0±0.0       -22.4 to -2.1*         13.2±7.7       -7.0 to 13.3\$         10.1±6.1       -19.3 to 1.0**         BCAAem <sup>2</sup> Mean±SE       95% Cl of         difference       difference         1.0±0.0       -11.6 to -1.1*         7.3±4.1       -1.9 to 8.6\$	BCAAem <sup>2</sup> Diet         Mean±SE       95% Cl of       Mean±SE         difference       difference       1.0±0.0         1.0±0.0       -22.4 to -2.1*       1.0±0.0         13.2±7.7       -7.0 to 13.3\$       1.7±0.3         10.1±6.1       -19.3 to 1.0**       1.8±0.3         10.1±6.1       -19.3 to 1.0**       1.8±0.3         MEN-2 <sup>7</sup> (change)       MEN-2 <sup>7</sup> (change)         Mean±SE       95% Cl of       Mean±SE         0.10±0.0       -11.6 to -1.1*       1.0±0.0         1.0±0.0       -11.6 to -1.1*       1.0±0.0         7.3±4.1       -1.9 to 8.6\$       1.7±0.3	BCAAem <sup>2</sup> Diet advice           Mean±SE         95% Cl of         Mean±SE         95% Cl of           difference         difference         difference           1.0±0.0         -22.4 to -2.1*         1.0±0.0         -10.8 to 9.4*           13.2±7.7         -7.0 to 13.3\$         1.7±0.3         -10.3 to 10.0\$           10.1±6.1         -19.3 to 1.0**         1.8±0.3         -11.0 to 9.3**           MEN-2 <sup>7</sup> (changes vs baseline)         MEN-2 <sup>7</sup> (changes vs baseline)           Mean±SE         95% Cl of         Mean±SE         95% Cl of           Mean±SE         95% Cl of         Mean±SE         95% Cl of           1.0±0.0         -11.6 to -1.1*         1.0±0.0         -6.0 to 4.5*           1.0±0.0         -11.6 to -1.1*         1.0±0.0         -6.0 to 4.5*	BCAAem <sup>2</sup> Diet advice       Two-way A         Mean±SE       95% Cl of       Mean±SE       95% Cl of       Effect of         difference       difference       difference       difference       Interaction         1.0±0.0       -22.4 to -2.1*       1.0±0.0       -10.8 to 9.4*       Time         13.2±7.7       -7.0 to 13.3\$       1.7±0.3       -10.3 to 10.0\$       Treatment         10.1±6.1       -19.3 to 1.0**       1.8±0.3       -11.0 to 9.3**       Interaction         MEN-2 <sup>7</sup> (changes vs baseline)       MEN-2 <sup>7</sup> (changes vs baseline)       Interaction         Mean±SE       95% Cl of       Mean±SE       95% Cl of         Mean±SE       95% Cl of       Mean±SE       95% Cl of         Mean±SE       95% Cl of       Mean±SE       95% Cl of         1.0±0.0       -11.6 to -1.1*       1.0±0.0       -6.0 to 4.5*       Time         1.0±0.0       -11.6 to -1.1*       1.0±0.0       -6.0 to 4.5*       Time         7.3±4.1       -1.9 to 8.6\$       1.7±0.3       -5.9 to 4.6\$       Treatment			

<sup>1</sup>Cytochrome C Oxidase-1; <sup>2</sup>Branched Chain Amino Acid Enriched Mixture; <sup>3</sup>Cytochrome C Oxidase- 4;

<sup>4</sup>Mitochondrial Transcription Factor A; <sup>5</sup>Nuclear Respiratory Factor-1; <sup>6</sup>Mitofusin-1; <sup>7</sup>Mitofusin-2.