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1	Effectiveness of nutritional and exercise interventions to improve body composition and
2	muscle strength or function in sarcopenic obese older adults: A systematic review
3	
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18	List of Abbreviations
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- 19
- 20 % BF; Percentage Body Fat
- 21 ACSM; American College of Sports Medicine
- 22 AHA; American Heart Association
- 23 BIA; Bioelectrical Impedance Analysis
- 24 BMI; Body Mass Index
- 25 CT; Computerized Tomography
- 26 DXA; Dual-energy X-ray Absorptiometry
- 27 EPIDOS; EPIDemiologie de l'OSteoporose
- 28 EWGSOP; European Working Group on Sarcopenia in Older People
- 29 FM; fat mass;
- 30 GFR; Glomerular Filtration Rate
- 31 HD; Habitual Diet
- 32 HG, Handgrip
- 33 HSC; High Speed Circuit
- 34 IGF-1; Insulin-like Growth Factor 1
- 35 IADL; Instrumental Activities of Daily Living
- 36 LM; Lean Mass
- 37 MRI; Magnetic Resonance Imaging
- 38 MPS; muscle protein synthesis
- 39 mTOR; mechanistic Target of Rapamycin
- 40 PRISMA; Preferred Reporting for Systematic Reviews and Meta-analyses
- 41 PROSPERO; International prospective register of systematic reviews

- 42 RCH+HD; Ricotta Cheese plus Habitual Diet
- 43 RCTs; Randomized controlled trials
- 44 RPE; Rates of Perceived Exertion
- 45 RM; Repetition Maximum
- 46 SD; Standard Deviation
- 47 SGOT; Serum Glutamic Oxaloacetic Transaminase
- 48 SGPT; Serum Glutamic-Pyruvic Transaminase
- 49 SPPB; Short Physical Performance Battery test
- 50 SMI; Skeletal muscle index; SO, Sarcopenic Obesity
- 51 SH; Strength Hypertrophy
- 52 TASM; Total appendicular skeletal muscle

53 Abstract

Although sarcopenic obesity (SO) poses a major public health concern, a robust approach for 54 the optimization of body composition and strength/function in SO has not yet been 55 established. The purpose of this systematic review was to assess the effectiveness of 56 57 nutritional (focusing on energy and protein modulation) and exercise interventions, either individually or combined, on body composition and strength/function in older adults with SO. 58 MEDLINE, the Cochrane Central Register of Controlled Trials, CINAHL and SPORTDiscus 59 were searched. Main inclusion criteria comprised sarcopenia as defined by the European 60 Working Group on Sarcopenia in Older People (EWGSOP) and obesity defined as % body 61 62 fat \geq 40 % (women) and \geq 28 % (men). Randomized controlled trials (RCTs), randomized controlled crossover trials and controlled clinical trials with older adults (mean age ≥ 65 63 years) following a nutritional regimen and/or an exercise training program were considered. 64 65 Out of 109 full text articles identified, only two RCTs (61 participants) met the inclusion criteria. One study was a nutritional intervention adding 15 g protein day⁻¹ (via cheese 66 consumption) to the participants' habitual diet. The second study was a high-speed circuit 67 resistance training intervention. Body composition did not change significantly in either of 68 the studies. However, the exercise intervention improved significantly muscle strength and 69 70 physical function. Although this review was limited by the small number of eligible studies, it provides evidence for the potential benefits of exercise and highlights the necessity for future 71 72 research to develop effective interventions including dietary and exercise regimens to combat 73 sarcopenic obesity.

74

75 Keywords: Aged; sarcopenia; obesity; dietary proteins; exercise; systematic review

76 1. Introduction

77

78 Sarcopenia is defined by the European Working Group on Sarcopenia in Older People 79 (EWGSOP) as the age-related decline of muscle mass and strength or function [1]. Low 80 strength and muscle mass are associated with poor functional status, physical impairments, 81 frailty, increased risk of falls, loss of independence and higher mortality risk [1][2]. It has 82 been suggested that in older people, strength is a stronger predictor of functional impairment 83 and mortality rates than absolute changes in muscle mass or lean mass alone [3-6]. Secondary 84 to functional impairments, muscle atrophy may also contribute to insulin resistance as muscle tissue plays the main role in glucose uptake and utilization [7]. According to a recent 85 systematic review, the prevalence of sarcopenia may vary from 1 % to 29 % in community-86 87 dwellers and 14-33 % in long-term care populations [8].

88

Another condition that can promote poor health is obesity, which is defined as 'abnormal or 89 90 excess body fat accumulation' [9], and is a growing concern due to its progressively rising 91 prevalence rates in older populations [10]. In 2010, 35 % and 28 % of the adults 65 years of age and older were reported to be obese in the US and the UK, respectively [11][12]. Similar 92 to sarcopenia, obesity can increase the risk of falls and mobility limitations in older age 93 94 [13][14], and when used in conjunction with indices of body composition and fat distribution (waist circumference or waist to hip ratio) it may be associated with adverse health effects, 95 96 such as cardiovascular disease, metabolic syndrome, diabetes mellitus and several cancers [15]. Furthermore, adipose tissue can infiltrate the muscle tissue [16] and mediate an 97 inflammatory response [17], which can result in muscle atrophy, mobility losses and lower 98 99 strength and muscle quality [16][18][19].

The relationship between sarcopenia and obesity is complex, with the development/ 101 progression of one condition being closely connected to the other (Figure 1). The condition 102 103 where sarcopenia and obesity occur together has been termed sarcopenic obesity (SO) [20]. It has been suggested that SO can predispose older individuals to more physical disabilities, gait 104 and balance abnormalities, and an increased risk of falls compared with either of the two 105 conditions alone [21]. Individuals with SO are exposed to ~2.5 times higher risk of reporting 106 107 Instrumental Activities of Daily Living (IADL) disabilities compared with adults without obesity but with sarcopenia, or adults with obesity but without sarcopenia [22]. This negative 108 synergistic effect of sarcopenia and obesity is in accordance with the findings from the 109 EPIDOS (EPIDemiologie de l'OSteoporose) study, which reported that among a cohort of 110 1,308 women divided in four groups: 1) without sarcopenia or obesity 2) with obesity but not 111 112 sarcopenia 3) with sarcopenia but not obesity and 4) with SO, the latter was the poorest in terms of performing physical activities that required strength [23]. According to a meta-113 114 analysis of 12 prospective cohort studies with a total number of 35,287 participants, the adults with SO had a 24 % higher risk of all-cause mortality compared with their healthy 115 counterparts [24]. 116

117

Figure 1 Relationship between sarcopenia and obesity and associated outcomes

119

Although SO has gained significant attention from the scientific community in recent years,
and a plethora of existing definitions and cut-offs for sarcopenia and obesity exist, there is no
universally accepted definition for SO [1][25][26]. Depending on the definition criteria and

cut-offs used the prevalence rates of SO can vary up to 26-fold, which makes detection and
management of the condition challenging for healthcare practitioners [27]. Moreover, there
are operational challenges around the management of SO. While exercise training can be
beneficial for both obesity and sarcopenia, the dietary management of obesity may require
energy restriction, whilst management of sarcopenia requires an increased intake of
macronutrients, especially protein [28].

129

This has resulted in a growing body of evidence highlighting potentially beneficial nutritional 130 and exercise strategies, aiming to reverse or attenuate the negative effects of ageing on body 131 composition and physical function [29-31]. Particular focus has been placed on protein 132 133 intake, energy modulation and resistance exercise [32][33]. With regard to protein intake, there seems to be a consensus for the benefits of increased protein intake, ranging from 1.0 g. 134 kg bw⁻¹ · day⁻¹ to 1.5 g · kg bw⁻¹ · day⁻¹, with the higher values appropriate for those older 135 136 adults with chronic conditions, sarcopenia and malnutrition, or when combined with resistance exercise [28][34][35]. 137

138

However, there are relatively few intervention studies utilizing exercise training and/or 139 nutritional regimens for older adults with SO [26][36]. It appears that most intervention trials 140 have aimed to attenuate muscle loss at an early stage rather than try to 'reverse' an 141 established condition related to advanced ageing such as sarcopenia or SO, which would be 142 143 far more challenging [37]. Furthermore, to the best of our knowledge, there has been no systematic review to date assessing the effectiveness of nutritional and exercise strategies, 144 145 alone or combined, to improve body composition and strength/function indices in older 146 individuals with SO. Therefore, the purpose of this systematic review was to assess the

evidence for the use of diets modulating energy and protein (or amino acids) content, exercisetraining regimens, or diet and exercise training combined, in older adults with SO.

149

The focus of this systematic review was to determine the effectiveness of protein or energy-150 modulating regimens, with or without exercise training on body composition and function in 151 152 adults, 65 years of age and older with SO. In particular, our aims were to 1) determine changes in absolute muscle mass, total appendicular skeletal muscle (TASM), skeletal muscle 153 index (SMI), fat mass, % body fat, body weight and body mass index (BMI), 2) assess 154 changes in muscle strength and/or physical function (including muscle strength, power, gait 155 speed and balance and 3) evaluate the effect of these interventions on quality of life, 156 metabolic profile, activities of daily living, adverse effects of supplementation or food 157 choices, compliance rates and changes in habitual dietary intake during or after the 158 interventions. 159

160

161 2. Approach

162

This systematic review was performed according to the Preferred Reporting for Systematic
Reviews and Meta-analyses (PRISMA) guidelines [38]. The protocol was registered with the
International prospective register of systematic reviews (PROSPERO registration number:
CRD42015017311).

167

168 2.1. Search Strategy

170 The Cochrane Central Register of Controlled Trials, MEDLINE (via EBSCOhost Research Databases), CINAHL and SPORTDiscus were searched up to and including May 2016. The 171 last search was conducted on 22 May 2016. No limits were applied for date of publication. 172 Combinations of key terms with Medical Subject Headings (MeSH) and Boolean operators 173 were used. The main keywords and terms used were: Age*/ Adult*/ Old*/ Elderly/ Senior, 174 Sarcopeni*/ Lean/ Frail/ Atrophy/ Weakness, Obes*/ Overweight/ Body Mass Index, 175 Exercise/ Training/ Strength/ Muscle/ Mass/ Hypertrophy/ Size/ Body Composition, Diet/ 176 Supplements/ Protein/ Amino Acids/ Energy, Life Quality/ Intervention. The search limiters 177 178 were English language and studies with human participants [the complete search strategy is presented in supplementary file 2]. 179

180

181 2.2. Inclusion Criteria

182

We included randomized control trials (RCTs), randomized control crossover trials and 183 controlled clinical trials using prospective nutritional and/or exercise interventions to 184 attenuate/ reverse the loss of muscle mass, reduce adipose tissue and optimize muscle 185 strength or function. Given that there are no universally adopted definition criteria for SO, 186 some authors may have used different terms to define the participants, e.g. 'weak and 187 overweight' or 'obese frail' etc. Such studies were included only if the participants had a 188 sarcopenic phenotype based on the definition criteria and cut-off scores recommended by the 189 190 EWGSOP [1]. Therefore, studies were included only if they presented data for a) body composition (data on absolute muscle mass, appendicular muscle mass, Total Appendicular 191 Skeletal Muscle (TASM) or Skeletal Muscle Index (SMI) assessed by Dual-energy X-ray 192 Absorptiometry (DXA), Bioelectrical Impedance Analysis (BIA), Computerised Tomography 193 194 (CT) or Magnetic Resonance Imaging (MRI) and b) muscular strength and/ or physical

195 function identified by one of the following tests: handgrip strength, knee flexion/extension, peak expiratory flow, gait speed, the Short Physical Performance Battery test (SPPB), the 196 timed up-and-go test or the stair climb power test . The mean age cut-off for inclusion was \geq 197 198 65 years based on how 'old age' is defined in the joint recommendations from the American College of Sports Medicine (ACSM) and the American Heart Association (AHA) [39]. The 199 inclusion criterion for obesity was defined as mean percentage body fat (% BF) ≥ 28 % in 200 men and ≥ 40 % in women [22] or in the absence of % BF data, a BMI ≥ 27 kg·m⁻². For any 201 given BMI, a person with sarcopenia will have by definition more body fat compared with 202 203 their counterparts without sarcopenia, therefore, adults with sarcopenia can present highadiposity at BMIs substantially lower than 30 kg·m⁻² [40]. Moreover, it is not uncommon for 204 205 studies to recruit participants who would be classified as overweight or obese based on a BMI cut-off ranging from 25 to 28 kg \cdot m⁻² when the focus is on sarcopenic obesity and/or when 206 participants come from a non-Caucasian ethnic group [25][26]. Studies that presented neither 207 the % BF nor BMI were included only if these indices could be derived from the weight, 208 209 height and body fat mass values, or if the authors of the study when contacted provided the essential information. 210

211

Nutritional interventions aiming to promote muscle hypertrophy by macronutrient profile modification or weight loss via energy restriction were of primary interest. Studies providing extra macronutrients (especially proteins or amino acids and their metabolites) either in the form of whole foods or dietary supplements administered through the oral route only were considered. Exercise regimens including resistance, balance, aerobic and mixed exercise protocols influencing lean mass, fat mass, muscle hypertrophy, strength, power, speed and/ or physical functional were also of primary interest.

220 **2.3. Exclusion Criteria**

221

222	Studies were excluded if the protocol involved administration of any kind of prescription
223	only/pharmaceutical agents, or any type of supplementation administered via a route other
224	than oral. Studies including participants with cachexia or with serious mental and cognitive
225	conditions prohibiting adherence to a structured exercise/ nutrition regimen, such as
226	Alzheimer's or dementia, were excluded.
227	
228	2.4. Study Selection
229	
230	The titles and abstracts were screened for eligibility (CT), and the full text copies of
231	potentially eligible articles were obtained for further inspection. The full-text articles were
232	independently assessed for eligibility by two reviewers (CT and JJ). The reference lists of
233	eligible articles and review papers as well as journals specializing in older age were hand
234	searched for potential articles. Any disagreement between the two reviewers was resolved by
235	a third reviewer (CAG).
236	
237	2.5. Data Extraction

238 Data were extracted from each eligible article by two reviewers (CT and CAG). Any

disagreements between the two reviewers were resolved by discussion until consensus was

240 reached. Demographic (age, sex, ethnicity/host country and habitation), methodological

241 (study design, sample sizes, duration, nutritional/dietary and/or exercise intervention plan,

supplement type, dosing/frequency of administration, exercise training

type/frequency/volume, assessment method, blinding) and outcome data (changes within and
between groups, significance, dropout rates, compliance, adverse effects) were compiled in a
standardized Excel spreadsheet.

2.6. Quality Assessment

249	The quality of the studies was assessed by two independent reviewers using a modified
250	version of the Downs and Black rating scale [41][42]. The Downs and Black scale is one of
251	the most credible instruments for the quality assessment of randomized [43] and non-
252	randomized intervention trials [44]. Modified scoring for Question 27 was performed as
253	detailed by Eng et al. [42]: the original scale had a maximum score of 32 but in this review
254	Question 27 was modified to score either 0 or 1 point instead of the original 0-5 points.
255	Therefore, the maximum total score for the five sections of the scale (reporting, external
256	validity, internal validity/bias, internal validity/confounding, power) was 28.
257	
257	
257	2.7. Principal Summary Measures
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258	2.7. Principal Summary Measures The primary outcome measures were 1) differences in mean of skeletal muscle mass (either
258 259	
258 259 260	The primary outcome measures were 1) differences in mean of skeletal muscle mass (either

3. Results

3.1. Description of studies

268	Our search strategy resulted in 1,440 potential articles. After the exclusion of 1,331 articles
269	based on titles and abstracts, 109 full-text articles reporting 109 studies were retrieved and
270	assessed for eligibility. The detailed flow chart of the selection process is presented in Figure
271	2. The authors of two potentially eligible studies [45][46] were contacted for further
272	information, but retrieval of all the essential body composition data was not possible for
273	reasons unrelated to this review, therefore, the articles were excluded. A total of $n=2$ studies
274	[47][48] including n= 61 participants met the inclusion criteria and were included in the
275	review. Study A[47] was a nutritional intervention and study B[48] an exercise training
276	intervention; neither of the studies combined exercise with diet.
277	
278	Figure 2. Information flow through the phases of the systematic review according to
279	PRISMA guidelines.
280	
281	3.2. Quality Assessment
282	
283	The two studies were randomized control trials of moderate methodological quality based on
284	the modified Downs and Black rating scale [41][42]. The total score for each study was 18
285	out of 28. The summary key information of the methodological strengths and limitations is
286	presented in table 1 (supplementary file 1 presents the complete breakdown of the scoring in
287	the different subsections of the scale). Both studies performed power calculations to
288	determine the population sample size prior to recruitment, however, study B[48] was
289	underpowered; target was $n=21$ per group, but the final analysis was conducted with $n=9$ and
	underpowered, target was $n-21$ per group, but the initial analysis was conducted with $n-2$ and

n=8 for the control and intervention group, respectively. In study A[47] only the testers were

291	blinded but not the participants. In study B[48] the two groups were exercising at different
292	times, therefore, participants were partially blinded. Study A[47] reported and tested for a
293	range of potential confounders, but failed to report essential information regarding the
294	participants' dietary intake at baseline and follow-up. The results in Study A were based on
295	an intention-to-treat analysis, whereas in Study B the analysis conducted was per-protocol.
296	
297	Table 1. Summary key points of the included study designs.
298	
299 300	3.3. Participant Characteristics
301	Participants in study A[47] were physically-independent individuals living in Mexico. Their
	Participants in study A[47] were physically-independent individuals living in Mexico. Their mean \pm SD age and TASM were 76 \pm 5.4 years and 15.5 \pm 2.9 kg, respectively. The mean %
301 302 303	
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302 303	mean \pm SD age and TASM were 76 \pm 5.4 years and 15.5 \pm 2.9 kg, respectively. The mean % BF of men and women was 33.3 \pm 6.2 % and 47.8 \pm 6.6 %, respectively. At baseline two men
302 303 304	mean \pm SD age and TASM were 76 \pm 5.4 years and 15.5 \pm 2.9 kg, respectively. The mean % BF of men and women was 33.3 \pm 6.2 % and 47.8 \pm 6.6 %, respectively. At baseline two men had a % BF < 28 % and three women < 40 %. All participants in study B[48] were
302 303 304 305	mean \pm SD age and TASM were 76 \pm 5.4 years and 15.5 \pm 2.9 kg, respectively. The mean % BF of men and women was 33.3 \pm 6.2 % and 47.8 \pm 6.6 %, respectively. At baseline two men had a % BF < 28 % and three women < 40 %. All participants in study B[48] were independent-living community dwellers from South Miami (USA). The mean \pm SD age and

309 3.4. Study Design

310

311 The aim of study A[47] was to assess whether the addition of a protein rich food to the

312 habitual diet could increase TASM and strength in older individuals with sarcopenia. The

- study was a 3-month RCT with a control (habitual diet; HD) and an intervention group
- 314 (habitual diet + 210 g ricotta cheese per day; RCH+HD). The cheese provided 15.7 g extra
- protein (including 8.6 g of essential amino acids), 10.4 g carbohydrate, 18.4 g fat and a total

of 267 kcal per day. Cheese was divided into three 70 g portions and participants were
instructed to consume each portion along with their usual breakfast, lunch and dinner. Dualenergy x-ray absorptiometry was used to measure TASM and body composition changes.

Study B[48] was a 15-week single blind RCT, which aimed to assess the effectiveness of a 320 321 novel exercise regime based on a high speed circuit (HSC) resistance training program (intervention) on body composition, muscular performance and IADL compared with a 322 conventional strength hypertrophy (SH) regime (control group) in community-dwellers with 323 324 SO. Body composition was assessed by single frequency BIA. Both groups performed exercises at 11 pneumatic gym machines twice per week. The SH protocol involved three sets 325 326 of 10-12 repetitions at 70 % of 1RM with a 1-2 min recovery break between sets. Participants 327 were instructed to keep a similar speed of contraction for both the concentric and eccentric phase (2 seconds per phase). The HSC group performed 10-12 repetitions at the same 11 328 exercises, but in a circuit pattern (i.e. moving from one exercise to the other) with no break in 329 330 between exercises, unless one full circuit was complete. Three full circuits were performed in total. The resistance load was selected based on maximum power output for each machine. 331 The concentric phase was performed as fast as possible while the eccentric in 2 seconds. No 332 dietary or nutritional element was introduced in the study, and neither dietary patterns nor 333 334 intakes were reported.

335

336 **3.5. Outcomes**

337

338 **3.5.1. Body composition**

340 No significant changes were seen in body composition, in either experimental or control groups. In study A[47] the addition of ricotta cheese resulted in no significant changes in lean 341 mass, TASM or body fat in the intervention or control group (Table 2). Secondary analysis 342 343 by sex showed that although men (n=8) in the intervention group experienced an increase in TASM by 490 g, this was not significantly different either from baseline or when compared 344 against the control group (p=0.42), which gained a non-significant 220 g of TASM. 345 Similarly, in study B[42] no statistically significant differences were detected in any of the 346 body composition indices, regardless of the exercise regimen (Table 2). Skeletal muscle 347 index (SMI) increased non-significantly in both groups (from 6.5 ± 0.66 kg·m⁻² to 6.6 ± 0.59 348 kg·m⁻² in HSC and from 6.7 \pm 0.45 kg·m⁻² to 6.8 \pm 0.42 kg·m⁻² in SH). 349 350 351
Table 2. Summary of the included studies
 352 3.5.2 Strength and/ or function 353 354 In study A[47], the group receiving the extra protein noted a non-significant trend towards an 355 increase in strength (+0.9 % relative increase). Although the control group experienced a drop in strength (-3.5 %), the difference between the two groups did not achieve statistical 356 significance (p=0.06). 357 358 Study B[48] reported significant improvements in several aspects of strength and function in 359 both exercise groups (Table 2). In particular, the strength-hypertrophy (SH) control group 360 experienced significant improvements in leg press 1RM by 22 % (p<0.01), chest press 1RM 361 by 16 % (p=0.03), leg press peak power by 19 % (p=0.03) and chest press peak power by 362 15% (p<0.01) whereas a non-significant increase of 12% was detected in hand grip strength 363

364 (from 17.3 ± 2.7 kg to 19.4 ± 4.6 kg; p>0.05). The HSC group had a significant improvement

in chest press 1RM by 21 % (p<0.01), leg press peak power by 41% (p<0.01) chest press peak power by 24 % (p<0.01) but hand grip strength did not change significantly (increased by 10 %, from 17.7 \pm 7.8 kg to 19.4 \pm 6.6 kg; p>0.5). Between group differences were detected only for leg press peak power, with the HSC group performing better than the control by 158 W [95 % CI (2, 315), p=0.005].

370

371 The Short Physical Performance Battery (SPPB) test improved significantly over time only

within the HSC group from 8.0 ± 1.5 to 9.6 ± 1.2 (p=0.02). Between group differences

favored the HSC group [mean difference 1.1 (95 % CI (-0.1, 2.4), p=0.08], although this was
not statistically significant.

375

376 **3.5.3 Secondary Outcomes**

377

Consumption of ricotta cheese in study A[47], resulted in significantly lower fasting insulin 378 379 levels in men (p=0.05) but not in women. There were no other significant changes in hepatic 380 markers (Serum Glutamic Oxaloacetic Transaminase (SGOT), Serum Glutamic-Pyruvic Transaminase (SGPT) and Alkaline Phosphatase), kidney function (blood urea, uric acid, 381 creatinine and Glomerular Filtration Rate (GFR)), anabolism (Insulin-like Growth Factor-1 382 (IGF-1)) or insulin resistance. No cases of microalbuminuria were present in the RCH+HD 383 group after the intervention period. In the intervention group, 25 % of women reported early 384 satiety after the consumption of ricotta cheese, however, dietary intakes were not reported. 385 Eight participants from the intervention group dropped out; five were due to personal health 386 issues, two could not eat the entire portion of ricotta cheese, and one had to relocate. In the 387 control group three people dropped out (two for personal reasons and one for modifying the 388

habitual diet). However, all participants were measured pre- and post-intervention accordingto an intention-to-treat analysis.

391

392 The exercise intervention in study B[48] resulted in acute joint pain only in the SH group. In addition, the HSC group reported significantly lower rates of perceived exertion (RPE) with a 393 mean difference of -1.5 (95 % CI -2.0,-0.12, p=0.04). Adherence rates were similar in the two 394 groups; 81 % in HSC and 85 % in SH. Regarding the Instrumental Activities of Daily Living 395 (IADLs) there were significant improvements within both groups (pre vs post); namely, time 396 397 needed for jacket on and off (from 11.5 ± 3.5 s to 10.2 ± 2.0 s; p=0.04), scarf pick-up (5.2 ± 1.1 s to 4.7 \pm 0.91 s; p<0.01 and pan carry (4.9 \pm 0.61 s to 3.9 \pm 0.77 s; p<0.01) improved 398 399 significantly within the control group, while the HSC group experienced significant 400 improvements in time for sit-to-stand (from 16.1 ± 5.7 s to 13.4 ± 3.9 s; p=0.02) and pan carry (5.4 \pm 1.3 s to 4.5 \pm 1.2 s; p<0.01). No differences were observed between the two 401 groups in the aforementioned IADLs. 402

403

In summary, neither of the studies had a significant effect on body composition. The 404 introduction of ricotta cheese in the habitual diet of participants in study A[47] aimed to 405 increase their protein intake but it was not reported whether or not this was achieved, and if 406 so to what extent. In the same study, there was a trend for an increase in strength in the 407 408 intervention group, but this was not significant. The only significant improvement reported was the fasting insulin levels, but that was reported only in men in the intervention group. 409 Despite the lack of body composition changes, in study B[48], the high speed circuit 410 411 resistance training and strength hypertrophy resistance training protocols significantly improved strength, power and function indices. Finally, due to the limited data extracted and 412

diversity of methodologies, statistical pooling was not feasible and therefore, a narrativeanalysis was conducted.

415 4. Discussion

416

The aim of this systematic review was to assess the effectiveness of nutritional and/or 417 exercise interventions on body composition and strength or function in older adults with 418 419 obesity and sarcopenia. Although only two studies were identified, the lack of intervention 420 trials clearly highlights the need for more research in this area, especially trials combining exercise with nutritional approaches targeting this population group. With regard to the main 421 outcomes, neither an increase in protein intake by $15g \cdot day^{-1}$ nor a 15-week resistance 422 423 exercise protocol produced significant improvements in body composition indices. However, the exercise intervention (both the control group following a strength-hypertrophy resistance 424 exercise protocol and the intervention group utilizing a high-speed power-orientated circuit 425 resistance training) reported significant improvements in both strength and function. 426

427

428 4.1 Effects of protein intake on body composition and function in sarcopenic obesity
 429

Study A[47] attempted to utilize the effects of protein on increased skeletal muscle mass 430 accretion rates. Although the authors acknowledged that the suggested recommendations for 431 protein intake in older individuals with sarcopenia are 1.2- 1.5 g \cdot kg bw⁻¹ \cdot day⁻¹ [47], they did 432 not report the participants' daily protein intake, therefore, it was not corroborated whether 433 434 such intakes were achieved. It has been suggested that maximal muscle protein synthesis (MPS) rates in older adults can be achieved using \sim 35- 40 g protein \cdot meal⁻¹ [49-51] or 0.4 g 435 protein \cdot kg bw⁻¹ \cdot meal⁻¹ [51]. A valid question would be whether a daily addition of 210 g 436 437 ricotta cheese (delivering 15.7 g protein [47]) to the habitual diet could practically augment

438	muscle mass in older adults with sarcopenia. It is important to note that the cheese servings
439	were not consumed in one meal, instead they were spread over the three main meals, that is,
440	70 g cheese (~5 g of extra protein per meal) consumed with breakfast, lunch and dinner.
441	Protein intakes in Study A[47] were not reported, but if we extrapolate data from studies in
442	similar population cohorts [52], it has been suggested that older individuals are not likely to
443	consume an adequate amount of protein during all main meals. Tieland et al. [52] reported
444	mean protein intakes of ~8 g, ~18 g and ~29 g for breakfast, lunch and dinner, respectively.
445	Therefore, it is uncertain whether the addition of 5 g protein in the main meals in study A[47]
446	was enough to significantly augment MPS.
447	
448	Another confounder may have been the potential impact of the addition of cheese on the
449	habitual diet given the fact that 25 % of women in Study A[47] experienced early satiety. It
450	could be consequently speculated that women's habitual diet was modified with the addition
451	of ricotta, potentially displacing the intake of other foods. However this cannot be confirmed
452	as the habitual diet was not reported.
453	
454	In study A [47], even though there was a trend towards increased strength, it could be argued
455	that higher -and perhaps different distributions of- protein intake [31] were needed to enhance
456	muscle strength and accretion of skeletal muscle mass. It should be also noted that the power
457	calculation for sample size was based on lean mass as the primary outcome, rather than
458	muscle strength. Therefore, it is unknown whether a larger sample size was needed to reveal
459	a significant change in handgrip strength.

461 It has been previously reported that protein supplementation can enhance function in older462 adults. Namely, Tieland et al. [53] provided an oral supplement delivering 15 g of protein

twice daily (with breakfast and lunch) to older frail adults. This addition resulted in
significantly enhanced physical performance. The potential for high protein meals to maintain
or increase muscle mass and strength in older adults has been recently reported by Loenneke
et al. [54] who showed that one or two meals containing 30-45 g protein · day⁻¹ were
associated with higher lean mass and strength compared with those who did not consume any
meals over the threshold of 30g protein.

469

470 4.2 Effects of exercise training on body composition and function in sarcopenic obesity 471

The mechanisms underpinning the effects of exercise on body composition and function in older age are mainly accounted for by regulation of genes, circulating hormone levels (e.g. testosterone, IGF-1) and metabolic pathways (especially by activating the mechanistic target of rapamycin (mTOR), which is a pathway also activated by leucine-rich protein meals [31]) and have been reviewed in detail by Garatachea et al. [55] and McGlory and Phillips [56]. However, it is still unclear whether the modulation of these pathways can translate into realworld benefits for adults with SO.

479

In study B[48], the aim was to assess the effect of high-speed resistance exercise training on 480 indices of SO. In spite of possible methodological limitations, the improvements in strength, 481 power and IADL reported in study B[42], provide some evidence that exercise can improve 482 several domains of physical performance such as strength and power in older adults with SO. 483 This is in agreement with previous reports supporting the benefits of resistance exercise 484 485 training on clinically important outcomes, even in the absence of increased muscle mass [8][31][57]. This may be partly accounted for by the adaptive plasticity in the neuromuscular 486 487 system and skeletal muscle tissue in response to resistance exercise even in advanced older

488 age [58]. A significant improvement particularly in power can be very important for individuals with SO since muscle power can be a predictor of mobility skills and a more 489 influential indicator of physical capacity compared with absolute changes in strength [59]. 490 491 Another interesting finding from study B[48] was the large effect size observed in peak leg power achieved by exercising at 50 % 1RM. To a certain extent, this finding may be 492 explained by the novel aspect of the study design, that is, the resistance exercise progression 493 494 protocol. Resistance load would increase only when a power plateau was reached [48]. 495 Therefore, the protocol was designed in such a way as to favor maximum power output.

The lack of significant changes in lean mass or muscle mass after exercise training in adults 497 498 with sarcopenia, (which has also been reported elsewhere [8]), may be accounted for by 499 protocol-specific differences such as: duration, type, intensity, volume and frequency of exercise, as well as the availability of adequate nutrients (protein/amino acids), which are 500 needed to elicit an anabolic response and consequently muscle hypertrophy [31]. One 501 502 limitation of study B[48] was the lack of control for dietary intake, which could have partly explained the lack of effect on body composition. It has been shown that a bout of resistance 503 exercise can stimulate muscle protein synthesis (MPS) to a higher degree than protein 504 breakdown, however, in the absence of post-workout provision of nutrients (especially 505 protein) it can result in negative net muscle protein balance [60][61], and is a limitation of 506 507 study designs to date.

508

496

509 These data support the potential benefit of a resistance exercise program within lifestyle
510 intervention protocols due to its positive effect on muscle strength, power and function in
511 older adults with SO. Although no statistically significant body-composition changes were
512 reported in the included studies, the significant improvements in strength, power and function

513 may be more important for the quality of life of adults with SO than absolute changes in body514 fat or lean mass per se.

515

516 **4.3 Recommendations for future research**

517

518 More intervention trials should be undertaken to identify effective lifestyle strategies in adults 519 with SO, that will inform more robust approaches to combat this condition. Future research 520 should also bridge the gap in knowledge with respect to multimodal approaches combining 521 resistance exercise training with dietary strategies modulating protein intakes, in order to 522 augment muscle mass and strength [32] or fat free mass [62], and potentially alongside an 523 energy-deficit diet to promote fat loss [33].

524

It is important to note that although the need to augment muscle mass is paramount, a 525 reduction in fat mass and especially fat infiltrating the muscle tissue is equally important, 526 527 since intermuscular fat can result in mobility limitations [16]. Exercise training can preferentially reduce intermuscular adipose tissue more effectively than caloric-restriction 528 alone [63]. However, a combination of exercise with caloric-restriction can lead to greater 529 losses of total fat mass, which in turn may result in greater improvements in physical 530 function, sometimes even at the expense of lean tissue [64][65]. Nevertheless, it is currently 531 unknown whether this loss of lean mass may be detrimental in the long term for the life 532 quality of an older individual with sarcopenia who has already experienced a large decline in 533 muscle mass and strength. 534

535

To our knowledge only three studies to date, have reported significantly increased muscle orlean mass while concomitantly reducing fat mass [66-68]. The pilot study conducted by

538 Maltais et al. [68] was the only one that recruited older overweight adults with a low appendicular lean mass index. The authors concluded that 16-weeks of a whole body 539 resistance exercise regimen (at 80% 1RM) followed by the consumption of ~13 g dairy 540 supplement (chocolate milk with added milk powder) increased lean mass and reduced fat 541 mass even at the absence of caloric restriction (n=8). In the same study, the group that 542 received a soy-based beverage (matched for energy, protein, essential amino acids, 543 carbohydrate and calcium content) (n=8), reported significant increases only in lean mass but 544 no changes in fat mass. Similar results, but in pre-menopausal women, have been reported by 545 546 Josse et al. [66]. After 16 weeks of mixed exercise training (aerobic and resistance) combined with a caloric restriction (500 kcal daily deficit), only the high protein group (30% of total 547 energy intake came from protein, half of which was derived from dairy products) experienced 548 549 lean gains concomitant with fat losses. Albeit in younger adults, a recent four-week intervention combining 2.4 g protein \cdot kg bw⁻¹ \cdot day⁻¹ (achieved by consuming 3-4 whey-550 based protein beverages daily) with a mixed resistance, plyometric and high-intensity interval 551 552 training alongside an energy deficit regimen resulted in significantly higher lean mass and lower body fat [67]. The control group which differed only in protein intake (1.2 g protein · 553 kg bw⁻¹ · day⁻¹) did not experience a significant change in lean mass [67]. 554

555

Although the effectiveness of the aforementioned paradigms needs to be evaluated in older
adults with SO, they provide the framework for an initial approach to combat this condition.
What is primarily lacking from the literature is trials recruiting older adults with SO.
Additionally, protocols combining a high protein diet (potentially using whey or dairy
proteins) with a modest caloric deficit along with a well structured exercise training regimen
could be adopted. Moreover, long-term and follow-up studies with adults with SO who have

intentionally lost weight should be undertaken, in order to assess the impact of weight loss(especially if it comes from lean tissue) on life quality and other comorbidities.

564

565 **4.4 Limitations**

566

The main limitation of this review is the scarcity of data and studies undertaken in older 567 groups with SO. We reviewed only studies with participants presenting the sarcopenic 568 phenotype, as defined by the EWGSOP [1], using an appropriate methodology to assess body 569 570 composition. Although the EWGSOP reached a consensus in 2010 on the definition and assessment of sarcopenia, adopting criteria for low muscle and low strength or function, there 571 are recent studies [68][69] that use solely the criterion of low muscle mass to define 572 573 sarcopenia as it was initially proposed [70], without taking into consideration low strength or 574 function. In addition, one study conducted before 2010 was excluded because muscle mass was assessed using urinary creatinine [71], a method not included in the EWGSOP definition. 575 Regarding the two studies included in this review, study B[48] aimed to recruit specifically 576 participants with sarcopenic obesity. Study A[47] used sarcopenia as an inclusion criterion, 577 578 and although the mean % BF values met our cut-off criteria for obesity, after personal communication with the authors it was reported that although the majority of participants had 579 the sarcopenic obesity phenotype, a small proportion (5 out of 40) had a % BF below our cut-580 off. 581

Although of vital importance, there is an apparent lack of interventions with older adults with SO. This is in accordance with Finger et al. [62] who commented that interventions may refer to or discuss sarcopenia, however, the number of studies recruiting adults specifically with sarcopenia is very limited. This is even more complex with respect to studies on sarcopenic obesity, with a number of reviews [26][36][33][72] presenting interventions with participants

who either had none or only one of the conditions (sarcopenia or overweight/obesity but not both) and extrapolate these results to propose ways to improve the sarcopenic obesity phenotype. An example of an intervention study is that by Gadelha et al. [73] investigating the effect of exercise training on changes in the sarcopenic obesity index (assessed by the residuals method initially presented by Newman et al. [40] and adopted by the EWGSOP) of older Brazilian women. However, older age was the main inclusion criterion and no criteria specific for sarcopenia or obesity were adopted.

594

595 **4.5 Conclusion**

596

597 This review assessed studies investigating the effectiveness of exercise or nutritional interventions to improve the body composition and strength/function of older adults with 598 sarcopenia and obesity. None of the included studies significantly reduced body fat or 599 600 increased either skeletal muscle mass or lean mass. Although the number of included studies was low, it is evident that exercise training can elicit significant improvements in aspects of 601 physical fitness such as muscle strength and power, and consequently improve performance 602 603 in activities of daily living in adults with SO. The addition of 15 g protein day^{-1} to the habitual diet via cheese consumption revealed a non-statistically significant trend towards 604 605 increased handgrip strength, and a significantly better insulin response in men, but not in women. The lack of published data highlights the necessity for new research adopting 606 universally accepted cut-offs for sarcopenic obesity, with the inclusion of appropriately 607 designed exercise programs and dietary regimens, and with detailed assessments of dietary 608 patterns and protein intakes for the targeted population group. 609

610

611	Supplementary material
612	Supplementary file 1 (.pdf) S1 Table. Assessment of the methodological quality of the
613	included studies with the modified Downs and Black Scale.
614	Supplementary file 2 (.pdf) S1 Appendix. Search strategy
615	Supplementary file 3 (.doc) PRISMA Checklist
616	
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623	
624	CT, CAG, EB and JJ designed the study protocol and contributed to the writing of the
625	manuscript. CT conducted the search and screening of titles, abstracts, full-text articles, the
626	study selection, data extraction and quality assessment and prepared the manuscript. JJ
627	screened the full-text articles and assessed the eligibility of the studies. CAG conducted the
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629	authors read the final version of the manuscript.
630	
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632	
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641

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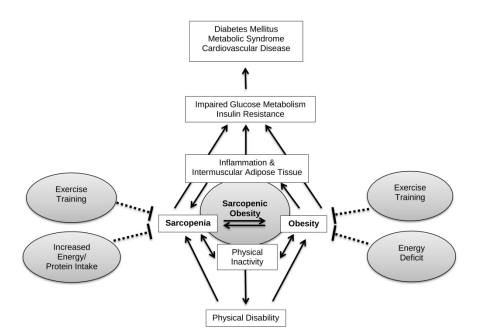


Figure 1. Relationship between sarcopenia and obesity and associated risks as well as
management strategies. Notes: Solid arrow: direct and positive association; Dashed line
management strategy attenuating/reversing the condition:

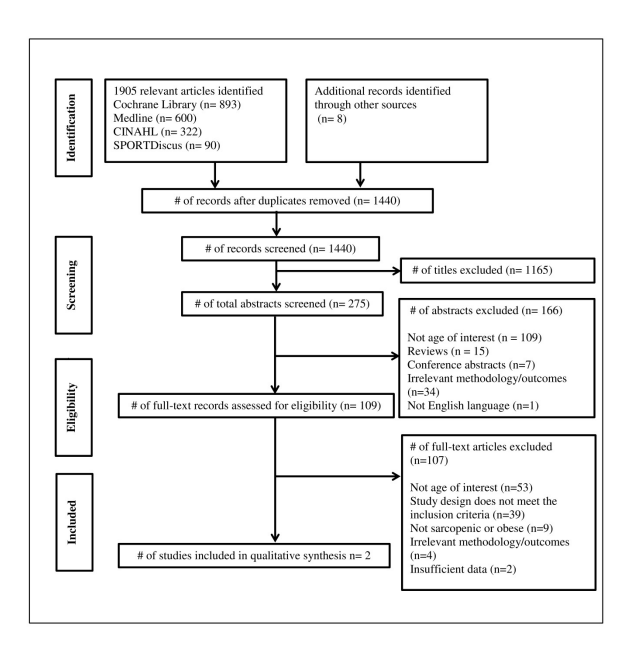


Figure 2. Information flow through the phases of the systematic review according to
PRISMA guidelines.

Study	Summary	Strengths	Limitations	
Study A	o Nutritional	• Intention to treat	• Lack of baseline	
Aleman-	Intervention	analysis	and follow up	
Mateo et al.	o 40 participants	 Body composition 	dietary intake and	
[47]	o 3 months	by dual-energy xray	physical activity	
	• Habitual diet plus	absorptiometry	data	
	210g Ricotta cheese	o Physically-		
	$\cdot day^{-1}$	independent		
	(intervention) vs	participants.		
	habitual diet	• Baseline and follow		
	(control)	up clinical tests for		
		kidney and liver		
		function		
		 Blinded personnel 		
		delivering the		
		assessment tests.		
Study B	0 Exercise	 Independent living 	 No allocation 	
Balachandran	Intervention	community-	concealment	
et al. [48]	o 21 participants	dwellers.	 Per-protocol 	
	o 15 weeks	 Participants were 	analysis	
	 High speed circuit 	partially blinded to	 Underpowered 	
	resistance (HSC)	the intervention.	 Characteristics of 	
	training	 Testing personnel 	participants lost to	
	(intervention) vs	blinded	follow-up not	
	strength hypertrophy	 All sessions 	described	
	(SH) resistance	supervised by 2	 No description of 	
	training (control)	physiology majors	the exercise setting	

889 Table 2. Summary key points of the included study designs.

Study	Setting/ Study Design/ Duration	Group	Participants Mean Age (SD)/ characteristics	Exercise Training	Nutritional Intervention	Sample Size (n) Drop-out (DO n) Female (F n) Adherence (%)	Assessment of a)body composition b) strength or function	Outcome Measure
Study A (Aleman-Mateo et al. [47])	Mexico/ RCT: two arms, one control, one intervention / 3 months	Control	76.7 (5.8) / physically- independent, sarcopenic based on low TASM and strength, obese based on %BF	No	Habitual diet (HD)	Baseline n=20, Final n= 12 DO n= 3 F n=12	a) DXA	$\begin{array}{l} \text{TASM} \rightarrow, \\ \text{FM} \rightarrow, \text{LM} \rightarrow, \\ \text{HG} \rightarrow \end{array}$
	montins	.115	/0 D1			F n=12 N/A	b) HG strength	
		Intervention	75.4 (5.0)/ independent living sarcopenic based on low TASM and strength, obese based on %BF	No	HD plus 210 g of ricotta cheese/day, (providing 15.7 gr extra protein/day)	Baseline n=20, Final n=17 DO n=8		TASM \rightarrow , FM \rightarrow , LM \rightarrow , HG \rightarrow
						F n=11		
						N/A		
Study B (Balachandran et al. [48])	USA/ RCT: Two arms, one control, one intervention/ 15 weeks	Control	71 (8.2)/ independent living community dwellers from South Miami, sarcopenic based on SMI and strength, obese based on %BF and BMI	Strength-hypertrophy (SH) training, 11 exercises, 3 sets of 10- 12 reps per set at 70% 1RM	No	Baseline n=10, Final n=9 DO n= 1 F n=8 85%	 a) BIA b) HG strength, SPPB, Leg press 1RM, Chest press 1RM, Leg press power, Chest press 	SMI→, %BF→, SPPB→, Leg 1RM↑**, Leg Power↑*, Chest 1RM↑*, Chest Power↑**, HG→
		Intervention	71.6 (7.8)/ independent living community dwellers from South Miami	High speed circuit (HSC) training, 11 exercises: 3 circuits of 10-12 reps per exercise at loads that maximised peak power output	No	Baseline n=11, Final n= 8 DO n= 3 F n =8 81%	power,	SMI→, %BF→, SPPB↓*, Leg 1RM→, Leg Power↑** [‡] , Chest 1RM↑**, Chest power↑**, HG→

890 Table 2. Summary of the included studies

- 891 Notes: \rightarrow no significant change, \uparrow significant increase, \downarrow significant decrease, *<0.05, **<0.01, \ddagger significantly better than the control group;
- 892 %BF, percent body fat; BMI, body mass index; DXA, dual-energy xray absorptiometry; FM, fat mass; HG, handgrip; LM, lean mass; RCT, randomised control Trial; RM,
- 893 repetition maximum; SPPB, short physical performance battery test; TASM, total appendicular skeletal muscle.