Check for updates

Physical function endpoints in cancer cachexia clinical trials: Systematic Review 1 of the cachexia endpoints series

James McDonald^{1,2}, Judith Sayers^{1,2,3}, Stefan D. Anker^{4,5,6}, Jann Arends⁷, Trude Rakel Balstad^{8,9}, Vickie Baracos¹¹, Leo Brown³, Asta Bye¹², Olav Dajani¹², Ross Dolan¹³, Marie T. Fallon¹, Eilidh Fraser¹, Christine Griel⁷, Aleksandra Grzyb¹, Marianne Hjermstad¹², Mariam Jamal-Hanjani^{14,15,16}, Gunnhild Jakobsen¹⁷, Stein Kaasa¹², Donald McMillan¹³, Matthew Maddocks¹⁸, Iain Philips¹, Inger O. Ottestad¹⁰, Kieran F. Reid¹⁹, Mariana S. Sousa²⁰, Melanie R. Simpson¹⁷, Ola Magne Vagnildhaug^{10,21}, Richard J. E. Skipworth³, Tora S. Solheim^{10,21}, Barry J. A. Laird^{1,2*} 🝺 & On behalf of the Cancer Cachexia Endpoints Working Group

¹Edinburgh Cancer Research Centre, University of Edinburgh, Edinburgh, UK; ²St Columba's Hospice, Edinburgh, UK; ³Clinical Surgery, University of Edinburgh, Royal Infirmary of Edinburgh, Edinburgh, UK; ⁴Department of Cardiology (CVK), Berlin Institute of Health Center for Regenerative Therapies (BCRT), and German Centre for Cardiovascular Research (DZHK) partner site Berlin, Charité Universitätsmedizin, Berlin, Germany; ⁵Institute of Heart Diseases, Wroclaw Medical University, Wroclaw, Poland; ⁶German Centre for Cardiovascular Research (DZHK) partner site Berlin, Charité Universitätsmedizin Berlin, Berlin, Germany; ⁷Department of Medicine I, Medical Center – Universitätsmedizin Berlin, Berlin, Germany; ⁷ Freiburg, Faculty of Medicine, University of Freiburg, Freiburg im Breisgau, Germany; 8 Department of Clinical and Molecular Medicine, Faculty of Medicine and Health Sciences, NTNU–Norwegian University of Science and Technology, Trondheim, Norway; ⁹Department of Clinical Medicine, Clinical Nucleur, of Science and Technology, Trondheim, Norway; ¹⁰Cancer Clinic, St Olavs Hospital – Trondheim University Hospital, Trondheim, Norway; ¹¹Division of Palliative Care Medicine, Department of Oncology, University of Alberta, Edmonton, AB, Canada; ¹²Regional Advisory Unit for Palliative Care, Department of Oncology, Oslo University Hospital/ European Palliative Care Research Centre (PRC), and Institute of Clinical Medicine, University of Oslo, Oslo, Norway; 13 Academic Unit of Surgery, University of Glasgow, Glasgow Royal Infirmary, Glasgow, UK; ¹⁴Cancer Research UK Lung Cancer Centre of Excellence, University College London Cancer Institute, London, UK; ¹⁵Cheartment of Duclogy, University College London Hospitals, London, UK; ¹⁷Department of Public Laboratory, University College London Cancer Institute, London, UK; ¹⁶Department of Oncology, University College London Hospitals, London, UK; ¹⁷Department of Public Health and Nursing, Norwegian University of Science and Technology, Trondheim, Norway; ¹⁸Cicely Saunders Institute of Palliative Care, Policy and Rehabilitation, King's College London, London, UK; ¹⁹Laboratory of Exercise Physiology and Physical Performance, Boston Claude D. Pepper Older Americans Independence Center for Function Promoting Therapies, Brigham and Women's Hospital, Harvard Medical School, Boston, MA, USA; ²⁰Improving Palliative, Aged and Chronic Care through Clinical Research and Translation (IMPACCT), University of Technology Sydney, Sydney, NSW, Australia; ²¹Department of Clinical and Molecular Medicine, Norwegian University of Science and Technology, Trondheim, Norway

Abstract

In cancer cachexia trials, measures of physical function are commonly used as endpoints. For drug trials to obtain regulatory approval, efficacy in physical function endpoints may be needed alongside other measures. However, it is not clear which physical function endpoints should be used. The aim of this systematic review was to assess the frequency and diversity of physical function endpoints in cancer cachexia trials. Following a comprehensive electronic literature search of MEDLINE, Embase and Cochrane (1990–2021), records were retrieved. Eligible trials met the following criteria: adults (≥18 years), controlled design, more than 40 participants, use of a cachexia intervention for more than 14 days and use of a physical function endpoint. Physical function measures were classified as an objective measure (hand grip strength [HGS], stair climb power [SCP], timed up and go [TUG] test, 6-min walking test [6MWT] and short physical performance battery [SPPB]), clinician assessment of function (Karnofsky Performance Status [KPS] or Eastern Cooperative Oncology Group-Performance Status [ECOG-PS]) or patient-reported outcomes (physical function subscale of the European Organisation for the Research and Treatment of Cancer Quality of Life Questionnaires [EORTC QLQ-C30 or C15]). Data extraction was performed using Covidence and followed PRISMA guidance (PROSPERO registration: CRD42022276710). A total of 5975 potential studies were examined and 71 were eligible. Pharmacological interventions were assessed in 38 trials (54%). Of these, 11 (29%, n = 1184) examined megestrol and 5 (13%, n = 1928) examined anamorelin; nutritional interventions were assessed in 21 trials (30%); and exercise-based interventions were assessed in 6 trials (8%). The remaining six trials (8%) assessed multimodal interventions. Among the objective measures of physical function (assessed as primary or secondary endpoints), HGS was most commonly examined (33 trials, n = 5081) and demonstrated a statistically signif-

© 2023 The Authors. Journal of Cachexia, Sarcopenia and Muscle published by Wiley Periodicals LLC.

This is an open access article under the terms of the Creative Commons Attribution-NonCommercial-NoDerivs License, which permits use and distribution in any medium, provided the original work is properly cited, the use is non-commercial and no modifications or adaptations are made.

icant finding in 12 (36%) trials (n = 2091). The 6MWT was assessed in 12 trials (n = 1074) and was statistically significant in 4 (33%) trials (n = 403), whereas SCP, TUG and SPPB were each assessed in 3 trials. KPS was more commonly assessed than the newer ECOG-PS (16 vs. 9 trials), and patient-reported EORTC QLQ-C30 physical function was reported in 25 trials. HGS is the most commonly used physical function endpoint in cancer cachexia clinical trials. However, heterogeneity in study design, populations, intervention and endpoint selection make it difficult to comment on the optimal endpoint and how to measure this. We offer several recommendations/considerations to improve the design of future clinical trials in cancer cachexia.

Keywords cachexia; cancer; endpoints; physical function; trials

Received: 5 May 2023; Revised: 19 June 2023; Accepted: 2 August 2023

*Correspondence to: Barry J. A. Laird, Institute of Genetics and Cancer, University of Edinburgh, Edinburgh EH4 2XR, UK. Email: barry.laird@ed.ac.uk James McDonald and Judith Sayers are joint first authors.

Richard J. E. Skipworth, Tora S. Solheim and Barry J. A. Laird are joint senior authors.

Introduction

It is well established in oncology that a person's physical function is a critical component of physiological reserves and guides their assessment and management. This is best evidenced by the use of performance status (PS), which routinely informs decision making and is a robust indicator for survival and treatment stratification. PS also indicates the extent to which a patient is limited in terms of their ability to perform activities of daily living.

Since 1947, PS has been assessed using the Karnofsky Performance Status (KPS), a percentage score of physical function that is determined by the patient's clinician. KPS, however, has largely been superseded by the Eastern Cooperative Oncology Group-Performance Status criteria (ECOG-PS) since 1982, and is a reliable measure of physical functioning, prognosis and overall disease burden, and correlates highly with quality of life.^{1,2} ECOG-PS is easier to measure, has comparable sensitivity and specificity to KPS and is now used extensively in cancer care. Despite this, the role of measures of PS in cancer cachexia as a diagnostic criterion and/or outcome in clinical trials has not been reported.

Fearon and colleagues published a new consensus definition of cachexia in 2011. They proposed that cancer cachexia is a multifactorial syndrome defined by an ongoing loss of skeletal muscle mass (with or without loss of fat mass) that cannot be fully reversed by conventional nutritional support and leads to progressive functional impairment.³ In view of this complex definition, researchers are divided on how the syndrome is best assessed and have proposed various approaches that include physical, biochemical and patient-reported measures. Considering the role of PS in cachexia, Fearon et al. also noted that the measure was applicable in the refractory component, helping to identify those patients who are nearing the end of life.³ In addition, ECOG-PS has also been shown to be superior to measures of lean mass and nutritional intake in terms of survival prediction.4-6

The 2011 definition of cachexia raises an important question about how cancer cachexia is defined and therefore measured, particularly in the context of clinical trials. Regulatory bodies such as the Food and Drug Administration (FDA) now require that endpoints, be they patient-reported outcome measures (PROMs), clinician-determined or performance-based measures, should be related to how a patient feels, functions or lives/survives.⁷ Cachexia trials may focus on different aspects such as anorexia or quality of life and thus should use endpoints related to the potential mechanism of action of an intervention. For therapies that improve lean muscle mass, these should also demonstrate improvements in function.⁸

Cachexia trials have used a variety of different endpoints that aim to demonstrate changes in physical function. These may be categorized as assessments that are physician determined (e.g., ECOG-PS and KPS), reported by patients (e.g., functional subscales of quality of life assessments) or objective measures of physical function. The latter is the most diverse group and, among others, includes 'hand grip strength' (HGS), the 'timed up and go' (TUG) test, the '6-min walking test' (6MWT), the 'stair climb power' (SCP) test and the 'short physical performance battery' (SPPB).⁹

With such diversity in measures of physical function comes the challenge of interpreting clinical benefit of results from cachexia trials. Until there is an appraisal of the myriad of physical function endpoints and their use in cachexia, researchers will not be able to draw meaningful conclusions from their results. It is therefore imperative that cachexia researchers now take stock of what has been done and, more importantly, what may be meaningful.

The aim of this systematic review was to outline the frequency and diversity of physical function measures that are used in cancer cachexia trials. It is part of a series of reviews that examine a variety of endpoints used in cancer cachexia trials.

Methods

This systematic review was reported as described in the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) statement.¹⁰

Search strategy

The search for studies published from January 1990 until 2 June 2021 was conducted by a research librarian (University of Oslo, Norway) using the databases MEDLINE (Ovid), Embase (Ovid) and Cochrane Central Register of Controlled Trials (*Appendix S1*). It was registered on the International Prospective Register of Systematic Reviews (PROSPERO registration: CRD42022276710) where further detail is available.¹¹

Eligibility criteria

Articles were considered eligible if they were controlled trials investigating interventions that aim to treat or attenuate cachexia and associated conditions (as defined in the PROSPERO register) in adult patients with cancer. There were no restrictions in the type of intervention (pharmacological, nutritional, exercise, multimodal etc.) nor the type of comparator. To reduce bias and focus on outcomes with most clinical impact, articles were excluded if the study included fewer than 40 patients and/or the intervention lasted <14 days. Studies were included if they were published in full text from 1990 and were written in English.

Data selection and extraction

This systematic review is part of a comprehensive collaboration including reviews examining different endpoints in cachexia (body composition, oncology, physical function, PROMs, systemic inflammation and nutritional). As most controlled trials in cachexia explore several different endpoints (as primary or secondary), articles were divided evenly among the review team for data extraction.

All articles identified were transferred to Covidence software.¹² Article selection based on titles was carried out by two independent reviewers (O. D. and B. L.). Articles selected by their titles had their abstracts read and selected by two independent reviewers (T. S. S. and B. L.). Any uncertainties in assessing the eligibility of the studies were discussed among the authors until a consensus was reached.

A data extraction table was developed, pilot-tested and refined within the review group before data were extracted from each article by two independent authors from the review group. Articles relevant to each systematic review were then identified from the data. For this review, relevant articles were those that assessed the specified physical function endpoints noted in this review.

Assessing risk of bias

The methodological quality of each study was systematically assessed by four independent reviewers (J. M., J. S., O. D.

and B. L.) using the modified Downs and Black checklist.¹³ Among other criteria, the tool assesses study design, external and internal validity, estimate of variance reporting and whether the outcome was defined and robust.

Endpoints

Endpoints investigated in this review were measures of physical function in cancer cachexia. These included objective assessments of physical function: HGS,¹⁴ 6MWT,¹⁵ SCP,¹⁶ SPPB¹⁷ and TUG.¹⁸ The review also included measures of physical function that are assessed by clinicians (ECOG-PS and KPS) and patient-reported assessments of physical function (physical function subscales of the European Organisation for the Research and Treatment of Cancer Quality of Life Questionnaires [EORTC QLQ-C30]).¹⁹

Data analysis

As expected, the retrieved studies were heterogeneous in terms of interventions and patient characteristics, and the variety of outcome measures studied large. As such, meta-analysis of the effect of the interventions was not relevant, and these data were summarized narratively. In studies where the sample size was more than 100, raw data on objective measures and corresponding variability of measures were extracted and presented in keeping with PRISMA guidelines.¹⁰

Results

After removal of duplicates, 5975 records were reviewed by title or abstract (the abstract was assessed where the title was insufficient), resulting in 369 records being appraised in full. Following appraisal, 250 records were further excluded, leaving 116 that were eligible for the systematic review database. Of these, 71 studies examined physical function endpoints and thus were eligible to be included in the review. This is detailed in *Figure 1*.

The key characteristics of eligible trials are presented in *Table 1*. As predicted, the trials were heterogeneous in terms of intervention and tumour site studied. Trials also varied in sample size from n = 40 to n = 929 patients. Pharmacological interventions were assessed in 38 (54%) trials, and of these, 11 (29%) examined megestrol (n = 1184) and 5 (13%) examined anamorelin (n = 1928). Twenty-one (30%) trials examined nutritional interventions (n = 2340), six (8%) trials examined exercise-based interventions (n = 430) and six (8%) trials examined combination/multimodal interventions (n = 1422).

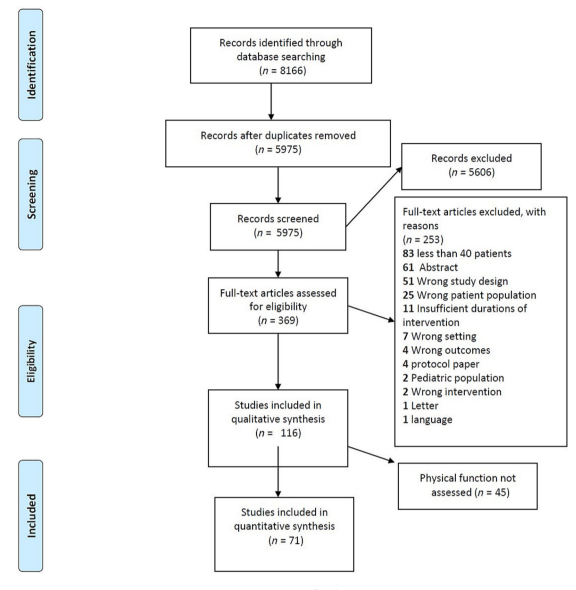


FIGURE 1 PRISMA flowchart.

In total, measures of physical function were used as a primary endpoint in 21 (30%) trials. Such measures include objective measures of physical function (HGS, 6MWT, TUG, SCP and SPPB), clinician assessment of function (ECOG-PS and KPS), which were each used in 7 (10%) trials, and PROMs of physical function (EORTC QLQ-C30 physical function [PF] subscale) in 10 (14%) trials. The remainder of the studies used physical function measures as secondary endpoints. Where primary and secondary endpoints were not clear, all outcomes were assumed to be secondary.

Table 2 summarizes the number and combined size of eligible trials that report each functional endpoint. Of note, HGS was examined in 33 trials (n = 5081) and demonstrated a statistically significant finding in 12 (36%) trials (n = 2099). The 6MWT was assessed in 12 trials (n = 1074) and demonstrated a statistically significant finding in 4 (36%) trials (n = 403). SCP, TUG and SPPB were each assessed in three trials (n = 371, 279 and 320, respectively). For clinician assessment of function, statistically significant changes in ECOG-PS were noted in 7 out of 9 (78%, n = 890) studies, while statistically significant changes in KPS were noted in 5 out of 16 studies (32%, n = 711). For patient-reported assessments of function, the physical function subscale of the EORTC QLQ-C30 was statistically significant in 11 out of 25 trials (44%, n = 1794).

Figure 2 shows the relationship between study interventions and functional measures across eligible trials. Most

	2	((с
Quality	-	Design	<i>u</i> 1	Cancer	Intervention	Comparator	Primary outcome	Function measure ^a
7 F	Ľ.	RCT	58	Head and neck	Individual nutritional counselling	Usual standard of care	EORTC QLQ-C30	EORTC QLQ-C30-PF
6 R	2	RCT	190	Mixed	Megestrol acetate + dexamethasone	Placebo	Appetite	KPS
7 9 R	~~~~	RCT	120 120	Mixed Lung	Mirtazapine Dietary counselling/fish oil	Placebo Usual standard of care	Appetite Feasibility, compliance,	HGS TUG
							רט-טטא סד, weight, skeletal muscle, measures of physical function	
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	Ξ. Ψ	ג <u>ר</u>	47 40	Mixed Mixed	Herbal combination Multimodal	Placebo + megestrol Usual standard of care	Weight gain Feasibility	HGS + ECOG-PS HGS
	Ř	RCT	100	Gastrointestinal	Individual nutritional	General dietary advice	PG-SGA	HGS
6 RCT	RC	Б	96	Lung	therapy Artificial nutrition	Usual standard of care	Nutritional status, EORTC QLQ-C30, incidence of complications	EORTC QLQ-C30-PF
9 R(	Ř	RCT	190	Mixed	Machine-based resistance exercise	Progressive muscle relaxation	Multiple	EORTC QLQ-C30-PF
7 RCT	Å	h	40	Pancreatic	Resistance exercise	Usual standard of care	Mobility, muscle	6MWT, 400-m walk, chair rise
10 RCT 7 RCT	8 8 0 0	55	52 111	Mixed Mixed	Multimodal Parenteral nutrition	Usual standard of care Oral feeding	SPPB EORTC QLQ-C15-PAL,	SPPB, HGS, TUG ECOG-PS
					with provitamins, trace elements and electrolytes		deterioration-free survival	
7 RCT 8 RCT	$\mathcal{O}$	<b>⊢</b> ⊢	40 166	Gastrointestinal Mixed	Captopril Nutritional counselling + whey	Placebo Nutritional counselling	EORTC QLQ-C30 Phase angle	EORTC QLQ-C30-PF HGS
7 RCT	^R C	⊢	47	Mixed	Home parenteral nutrition	Usual standard of care	Fat-free mass	HGS, 6MWT
8 RCT	RC	L	55	Lung	Supplement with protein + carbohydrate	Isocaloric supplement	Adverse events, clinical signs, changes in medication, routine blood tests	HGS
8 RCT 5 RCT	RC RC		44 43	Mixed Pancreatic	Walking Supervised resistance training	Usual standard of care Usual standard of care	SPPB Feasibility, effectiveness, muscle strength CPET, 6MWT	SPPB 6MWT
8 8 8CT	S S	<u> </u>	174 90	Lung Mixed	Anamorelin Megestrol acetate + celecoxib	Placebo Megestrol acetate + blacebo	Lean body mass Weight	HGS, 6MWT HGS, ECOG-PS
8 RCT 7 RCT 8 RCT		666	307 125 54	Head and neck Pancreatic Lung	Multimodal Usual standard of care Trial drug Placebo Thalidomide + cinobufagin Usual standard of care	Usual standard of care Placebo I Usual standard of care	PG-SGA Survival Arm circumference, weight, albumin, EORTC QLQ-C30, side	EORTC QLQ-C30-PF HGS, SCP, 6MWT, TUG HGS
							ettects, Hus	(Continues)

Table 1	Table 1 (continued)								
Year	Author	Quality	Design	u	Cancer	Intervention	Comparator	Primary outcome	Function measure ^a
2018	Schink ⁴³	თ	Ь	131	Mixed	Supervised physical exercise + nutritional advice	Nutritional advice only	Skeletal muscle mass	HGS, EORTC QLQ-C30, 6MWT, KPS
2018	Uster ⁴⁴	6	RCT	58	Mixed	Multimodal	Usual standard of care	Global health status, FORTC OLO-C30	EORTC QLQ-C30-PF, HGS 6MMT
2018	Turcott ⁴⁵	7	RCT	33	Lung	Nabilone	Placebo	FAACT	EORTC QLQ-C30-PF
2017	Jatoi ⁴⁶	00	RCT	263	Mixed	Creatine	Placebo	Weight gain	HGS
2017	Currow ^{4/}	00	RCT	513	Lung	Anamorelin	Placebo	Safety, tolerability	HGS
2017	Leedo ⁴⁰	ø	RCT	40	Lung	Delivery of protein-rich	Usual standard of care	EORTC QLQ-C30	ECOG, HGS
C 1 U C	collocim49	o	Ľ	11	Mixed	meals	and to be be at a low		
2017	Solneim Warnar ⁵⁰	0 1	ב ב מ	- 4 	IVIIXEU Pancroatic	Nuturnoaat Fish oil	Usual standard OI care Marina nhocnholinide	reasibility Waiabt appatita	BINIVI, HGS FORTC OI O-CRO-DF
2017	Zietarska ⁵¹	. 9	RCT 2	95	Colorectal	Oral nutritional	Usual standard of care	Treatment toxicity	KPS
	<del>+</del> -	c	FU C	007		supplement	-		
91.02	Lakayama T	×	לב צ	180	Lung	Anamorelin	Placebo	Lean body mass, Hos	
2016	lemei Gavazzi ⁵⁴	2 X	<u>ר</u> ב	70	Mived	Anamorelin Home enteral nutrition	Nutritional councelling	נפאר Lean body mass, אט אפומא <del>ו</del>	עסא אקא נפא
2016	Woo ⁵⁵	- 6	ג <u>ל</u>	67	Pancreatic	Pancreatic exocrine	Usual standard of care	Body weight	EORTC QLQ-C30-PF
	ł					replacement therapy		)	
2016 2016	Coats ⁵⁶ Canozzi ⁵⁷	10 %	בא גע	87 60	Mixed	Espindolol Immediata lifectula	Placebo Delaved lifectule	Rate of weight change Body composition	HGS, SCP, SPPB, 6MWT 6MMT
2		)		2		intervention	intervention		
2015	Focan ⁵⁸	7	RCT	53	Mixed	Mindfulness/dietician	Usual standard of care	Weight loss, BMI,	EORTC QLQ-C30-PF
								QLQ-C30, total daily	
								calorie intake, FFMQ, satisfaction with	
		г	Ľ	ç				intervention	
2012 2014	arcia Hond ⁶⁰	~ 00	RCI Phase I	22	Mixed	Anamorelin Monoclonal antibodv	Placebo Dose comparison	Lean body mass Safety, tolerability	HGS EORTC
-	ñ	)	5						QLQ-C30-PF
2014 2014	Poulsen ^{ol} Pottel ⁶²	ωœ	RCT	61 85	Mixed Head and neck	Multimodal Echium oil	Usual standard of care Usual standard of care	Weight Weight loss	EORTC QLQ-C30-PF EORTC QLQ-C30-PF,
2013	Dahrila-Dintiniana ⁶³	199 7		679	[astrointoctinal	lebomiti.M	Iterial etabard of caro	INN suitets leaditistud	
				070				Nottingham Screening Tool, ECOG-PS	
2013	Dobs ⁶⁴	8	RCT	159	Mixed	Enobosarm	Placebo	Lean body mass	6MWT, HGS + SCP
2012	Maccio ⁶⁵	∞ ^c	۲ ۲	124 77	Gynaecological	Multimodal	Megestrol acetate	Lean body mass, REE	HGS + ECOG-PS
7107	Nait	2		77	רמוורו במוור	r-Calillule	LIACEDO	BIA, EORTC QLQ-C30, BFI	
2012	Wen ⁶⁷	5	RCT	63	Mixed	Megestrol acetate + thalidomide	Megestrol acetate	questionnaire, survival Weight, fatigue (MFSI-SF scale) muality of life	ECOG-PS, HGS
								(EORTC QLQ-C30),	
								sarety, adverse events	(Continues)

135321900009, 0, Downloaded from https://onlinelibury.wiley.com/doi/10.102/j.Sml.13321 by University College London UCL Library Services, Wiley Online Library on [1109/2023], See the Terms and Conditions (https://onlinelibury.wiley.com/doi/10.102/j.Sml.13321 by University College London UCL Library Services, Wiley Online Library for uses of use; OA articles are governed by the applicable Cranity Common License

Table 1	<b>Fable 1</b> (continued)								
Year	Author	Quality	Design	<i>u</i> 1	Cancer	Intervention	Comparator	Primary outcome	Function measure ^a
2012	Madeddu ⁶⁸	7	RCT	60	Mixed	L-Carnitine + celecoxib + megestrol	L-Carnitine + celecoxib	Lean body mass, daily physical activity	HGS, 6MWT
2011	Baldwin ⁶⁹	8	RCT	358	Mixed	Multimodal	Nutritional	Survival	HGS
2011	Silander ⁷⁰	9	RCT	134	Mixed	PEG feeding before	encouragement Usual standard of care	Weight loss	EORTC QLQ-C30-PF
2010	Mantovani ⁷¹	7	RCT	332	Mixed	treatment + individual nutritional support Multiple (five arms)	Multiple	Lean body mass,	HGS, <b>ECOG-PS</b> , EORTC
2008 2006	Wiedenmann ⁷² Fearon ⁷³	8	RCT	89 518	Pancreatic Mixed	Infliximab Eicosapentaenoic aci	Placebo acidPlacebo	REE, fatigue Lean body mass Weight	QLQ-C30-PF 6MWT, <b>KPS</b> KPS, <b>EORTC QLQ-C30-PF</b>
2005	Ravasco ⁷⁴	7	RCT	75	Head and neck	supplement Nutritional supplements + nutritional	Nutritional supplements	PG-SGA, energy intake	EORTC QLQ-C30-PF
2005 2004	Gordon ⁷⁵ Isenring ⁷⁶	10	גר גר	50	Pancreatic Mixed	counselling Thalidomide Intensive nutritional		Weight Weight, free-fat mass,	HGS, EORTC QLQ-C30-PF EORTC QLQ-C30-PF
2003	Fearon ⁷⁷	00	RCT	200	Pancreatic	counselling Oral supplement + N-3 fatty acid	Oral supplement	PG-SGA, EUKIC QLQ-C30 KPS, EORTC QLQ-C30, energy intake,	KPS, EORTC QLQ-C30-PF
2003 2000	Bruera ⁷⁸ Erkurt ⁷⁹	7 8	RCT	60 100	Mixed Mixed	Multimodal Megestrol acetate	Placebo Usual standard of care	protein intake Appetite Weight, ECOG-PS, appetite, malnutrition,	KPS ECOG-PS
1999	McMillan ⁸⁰	7	RCT	73	Mixed	Megestrol	Placebo	loss of taste and smell Weight gain, QoL	EORTC QLQ-C30-PF
1999	Westman ⁸¹	7	RCT	255	Mixed	acetate + Ibuproten Megestrol acetate	Placebo	Quality of life: EORTC	EORTC QLQ-C30-PF
1998	Catalina ⁸²	ß	RCT	107	Mixed	Megestrol acetate	Megestrol acetate	Weight, nutritional	KPS
1998 1997	De Conno ⁸³ Neri ⁸⁴	6 4	RCT	42 225	Mixed Mixed	now dose) Megestrol acetate Medroxyprogesterone	(mgn aose) Placebo Usual standard of care	status, quanty of me, Nro Appetite Multiple patient-reported	KPS KPS
1996	Lissoni ⁸⁵	7	RCT	86	Mixed	acetate Supportive case 4 molatonin	Supportive care	outcomes Weight loss, TNF	KPS
1996	Gebbia ⁸⁶	9	RCT	122	Mixed	Megestrol acetate	Dose comparison	Appetite, weight, food intake, KPS, extension PS, pain, energy, depression, survival,	KPS
1996	Simons ⁸⁷	7	RCT	134	Mixed	Medroxyprogesterone acetate	Placebo	toxicity Appetite, weight	EORTC QLQ-C30-PF
									(Continues)

Year	Year Author	Quality	Quality Design <i>n</i>	ч	Cancer	Intervention	Comparator	Primary outcome	Function measure ^a
1994 Lai ⁸⁸	Lai ⁸⁸	ы	RCT 52	52	Mixed	Megestrol acetate + prednisolone	Placebo	Appetite, weight, KPS, tolerance to radiation,	KPS
1993	1993 Downer ⁸⁹	<del></del>	RCT	60	Mixed	Medroxyprogesterone acetate	Placebo	Appetite, weight, mid-arm KPS circumference, triceps	KPS
								SKIN TOID	
Abbrev Cooper FAACT, Multidi QoL, qu	iations: 6MWT, 6-mi ative Oncology Grou Functional Assessm mensional Fatigue S uality of life; RCT, rar	in walking p-Performi lent of And symptom II	test; BFI, ance Stat orexia/Ca nventory controlled	Big Five us; EOR1 chexia T -Short F d trial; R	e Inventory; BIA, bioel TC QLQ-C30-PF, Europ Therapy; FFMQ, Five-F orm; PEG, percutane 'EE, resting energy exi	ectrical impedance analysis ean Organisation for the Re acet Mindfulness Question ous endoscopic gastroston penditure; SCP, stair climb	; BMI, body mass index; t search and Treatment of naire; HGS, hand grip s my; PG-SGA SF, Patient- power; SPPB, short phys	Abbreviations: 6MWT, 6-min walking test: BFI, Big Five Inventory: BIA, bioelectrical impedance analysis: BMI, body mass index; CPET, cardiopulmonary exercise test: ECOG-PS, Eastern Cooperative Oncology Group-Performance Status; EORTC QLQ-C30-PF, European Organisation for the Research and Treatment of Cancer Quality of Life Questionnaires physical function; FAACT, Functional Assessment of Anorexia/Cachexia Therapy; FFMQ, Five-Facet Mindfulness Questionnaire; HGS, hand grip strength; KPS, Karnofsky Performance Status; MFSI-SF, Multidimensional Fatigue Symptom Inventory-Short Form; PEG, percutaneous endoscopic gastrostomy; PG-SGA SF, Patient-Generated Subjective Global Assessment Short Form; QoL, quality of life; RCT, randomized controlled trial; REE, resting energy expenditure; SCP, stair climb power; SPPB, short physical performance battery; TNF, tumour necrosis factor;	ise test; ECOG-PS, Eastern onnaires physical function; ormance Status; MFSI-SF, I Assessment Short Form; F, tumour necrosis factor;

**Fable 1** (continued)

QoL, quality of life; RCT, randomized controlled trial; REE, resting energ TUG, timed up and go test. ^aWhere functional measure is in bold, this denotes a significant finding studies assessed pharmacological interventions, followed by nutritional intervention and then exercise/lifestyle and multimodal interventions. HGS was assessed most commonly, particularly in the larger trials and in those where a pharmaceutical intervention was assessed. These aggregated data do not allow us to assess which measurements are more useful in terms of being an accurate re-

surements are more useful in terms of being an accurate reflection of physical functioning or their sensitivity to changes in functioning. However, we explored the relationships between specific endpoints looking at the 29 studies that report more than one measure of physical function. HGS reported was one of the measures in 25 of these studies and most commonly assessed together with 6MWT (k = 9), ECOG-PS (k = 5), KPS (k = 5) or EORTC-PF (k = 5) (*Table 3* and *Figure 3*).

In Table 3, we summarize the number of studies that identified statistically significant effects of one, both or neither measure of physical function for each pair of measures. While it was common that both HGS and the other measures of physical function were found to be either statistically significant or not in these trials, 6MWT, ECOG-PS and KPS were more often statistically significant in trials where HGS was reported not to be statistically significant (Table 3). It was of interest that HGS generally decreased in the intervention and also in the control groups of the reported trials, although in two cases, the intervention group had stable or improved HGS, respectively. Where the 6MWT was assessed, there was no discernible difference between the control and intervention groups. Limited inference can be drawn from these observations; however, the effect of any cachexia intervention could be to attenuate decline and clearly may be dependent on the population being examined.

Table S1 details the raw values of HGS, 6MWT and TUG in selected two arm trials ( $n \ge 100$ ) where these were reported. All trials were pharmacological interventions including studies of anamorelin, enobosarm and antimyostatin therapies. There were no trials where either SPPB or SCP was assessed in which the sample size was more than 100. These data demonstrate that raw values were broadly comparable although there was a limit to data reported.

Table S2 details the raw values of HGS, 6MWT and TUG in selected multi-arm trials ( $n \ge 100$ ). Little inference can be made from these data as only a small number of trials used a multi-arm design.

Of the 71 studies examining physical function included in this review, 27 were designed as randomized trials that included a placebo control group. The details of these studies are presented in *Table S3*. As the risk of bias is lower in this group, a subset analysis of the randomized, placebo-controlled studies was performed, with results presented in *Table S4*. These studies included a total of 4594 participants. Broadly, functional endpoints demonstrated results with statistical significance at a similar proportion as in the overall cohort, but analysis is limited by the small sample sizes for each functional endpoint.

					כווחכם אוווורמוולויב לוושטוזכוושוב ווזוגה כבוחחיב	gniricant results	
Endpoint	Number of studies	Year published	Total sample size	Type of intervention	Number of studies (percentage of studies)	Sample size (percentage of total sample size)	Type of intervention
Objective measures of physical function Hand grip strength 33	ysical function 33	1998–2021	5081	Pharmacological: 20 Nutritional: 8 Exercise/lifestyle ^a : 0	12 (36%)	2099 (41%)	Pharmacological: 9 Nutritional: 2 Exercise/lifestyle ^a : 0
Six-minute walking test	12	2008–2019	1074	Pharmodan. 5 Pharmacological: 6 Nutritional: 1 Exercise/lifestyle ^a : 2 Mutricoscal: 2	4 (33%)	403 (38%)	Pharmacological: 1 Nutritional: 0 Exercise/lifestyle ^a : 0
Stair climb power	m	2013–2018	371	Pharmacological: 3 Nutritional: 0 Exercise/lifestyle ^a : 0 Multimodal: 0	1 (33%)	159 (43%)	Pharmacological: 1 Nutritional: 0 Exercise/lifestyle ^a : 0
Timed 'up and go' test	m	2018–2020	279	Pharmacological: 1 Nutritional: 1 Exercise/lifestyle ^a : 0 Multimodal: 1	0 (0.00%)	0 (0.00%)	Pharmacological: 0 Nutritional: 0 Exercise/lifestyle ^a : 0 Multimodal: 0
SPPB	m	2016–2020	320	Pharmacuan. 1 Nutritional: 0 Exercise/lifestyle ^a : 1 Multimodal: 1	0 (0.00%)	0 (0.00%)	Pharmacological: 0 Nutritional: 0 Exercise/lifestyle ^a : 0 Multimodal: 0
Clinician-assessed measures of physical function ECOG 9 2000-2	es of physical fi 9	<b>unction</b> 2000–2020	1570	Pharmacological: 7 Nutritional: 2 Exercise/lifestyle ^a : 0 Mutritional-10	7 (78%)	890 (57%)	Pharmacological: 4 Nutritional: 2 Exercise/lifestyle ^a : 0
KPS	16	1993–2021	2236	Pharmacological: 11 Nutritional: 6 Exercise/lifestyle ^a : 0 Multimodal: 1	5 (31%)	711 (32%)	Pharmacological: 4 Nutritional: 0 Exercise/lifestyle ^a : 0 Multimodal: 1
ECOG + KPS	25	1993–2021	2806	Pharmacological: 18 Nutritional: 6 Exercise/lifestyle ^a : 0 Multimodal: 1	12 (48%)	1601 (57%)	Pharmacological: 8 Nutritional: 2 Exercise/lifestyle ^a : 0 Multimodal: 2
PROMS OT PNYSICAI TUNCTION EORTC QLQ-C30 physical function	25 25	1996–2022	3167	Pharmacological: 11 Nutritional: 10 Exercise/lifestyle ^a : 2 Multimodal: 2	11 (44%)	1794 (57%)	Pharmacological: 3 Nutritional: 7 Exercise/lifestyle ^a : 1 Multimodal: 0

Table 2 Frequency of use of functional endpoints in eligible trials

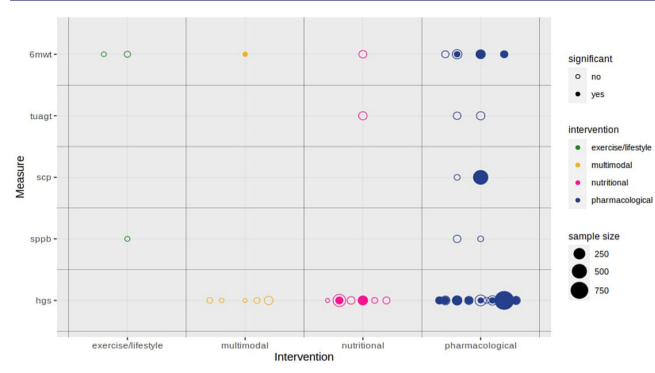


FIGURE 2 The relationship between study interventions and functional measures across eligible trials. 6MWT, 6-min walking test; HGS, hand grip strength; SCP, stair climb power; SPPB, short physical performance battery; TUG, timed up and go test.

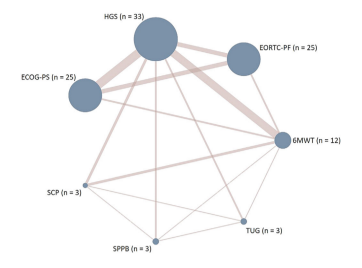
Table 3	Pairs of measures	of physical function	reported in included studies
---------	-------------------	----------------------	------------------------------

			1	lumber of studies		
Measure 1	Measure 2	Assessing both measures	Both measures statistically significant	Measure 1 statistically significant	Measure 2 statistically significant	Neither measure statistically significant
HGS	6MWT	9	1	1	3	4
HGS	ECOG-PS	5	2	0	3	0
HGS	KPS	5	0	1	2	2
HGS	EORTC-PF	5	0	1	0	4
HGS	SCP	3	0	1	1	1
6MWT	SCP	3	0	2	1	0
EORTC-PF	KPS	3	0	1	1	1
EORTC-PF	ECOG-PS	2	0	0	1	1
EORTC-PF	6MWT	2	0	0	1	1
HGS	SPPB	2	0	2	0	0
HGS	TUG	2	0	1	0	1
KPS	6MWT	2	1	1	0	0
SCP	SPPB	1	0	0	0	1
SCP	TUG	1	0	0	0	1
6MWT	SPPB	1	0	1	0	0
6MWT	TUG	1	0	1	0	0
TUG	SPPB	1	0	0	0	1

Abbreviations: 6MWT, 6-min walking test; ECOG-PS, Eastern Cooperative Oncology Group-Performance Status; EORTC-PF, European Organisation for the Research and Treatment of Cancer physical function; HGS, hand grip strength; KPS, Karnofsky Performance Status; SCP, stair climb power; SPPB, short physical performance battery; TUG, timed up and go test.

# Discussion

This is the first systematic review of physical function endpoints in clinical trials examining interventions for cancer cachexia. It was noted that a broad range of interventions showed varying levels of efficacy, assessed with different outcome measures. Objective measures such as HGS and, to a lesser extent, the 6MWT have been studied in the largest trials, usually in the context of a pharmacological intervention.



**FIGURE 3** Network diagram of the reporting of physical function measures in included trials. Size of nodes reflects the number of studies reporting each measure and the thickness of the connecting edges reflects the number of studies reporting each pair of measures (numerical details found in *Table 3*). 6MWT, 6-min walking test; ECOG-PS, Eastern Cooperative Oncology Group-Performance Status; EORTC-PF, European Organisation for the Research and Treatment of Cancer physical function; HGS, hand grip strength; SCP, stair climb power; SPPB, short physical performance battery; TUG, timed up and go test.

# Objective measures of physical function

Among the endpoints assessed in the present review, HGS was assessed in the largest number of studies and had the largest total sample size. The use of HGS is congruent with definitions of cachexia⁹⁰ and supports work by Song and co-workers.⁹¹ In over 1400 patients with cancer cachexia, they demonstrated that low HGS at baseline was an independent risk factor for cachexia and associated with reduced 1-year survival. These findings corroborated earlier work by Zhuang and co-workers.⁹²

Our results have also identified the 6MWT as a candidate for functional assessment in cachexia. The advantage of it is that it does not require any specialized equipment and may be performed in any healthcare setting. Despite being commonly used as a measure of physical function, there are limited data to support its use in cachexia. LeBlanc and co-workers examined the 2011 consensus definition of cachexia and compared it with key measures including the 6MWT. They failed to show a relationship between cachexia stage and 6MWT.93 Nonetheless, when assessed alongside other measures of physical function, the 6MWT was more frequently found to be statistically significant. Interpreting this observation cautiously, 6MWT may be more sensitive to changes in physical function, may be impacted by other symptoms such as dyspnoea or have less variability between participants, meaning that smaller changes are associated with greater statistical certainty. Considering that few eligible studies have assessed SCP, TUG and SPPB (three studies each), no firm conclusions can be drawn from our results.

# Performance status

Our findings highlighted the potential importance of ECOG-PS in the assessment of cachexia. As mentioned, ECOG-PS has largely succeeded KPS as the measure of PS in cachexia trials, which may be due to fewer categories within the former. We also see in our results that ECOG-PS may be more sensitive to changes in physical function than HGS given that ECOG-PS alone was found to be statistically significant in three of five studies assessing both measures.

#### Patient-reported measure of physical function

The EORTC QLQ-C30-PF subscale was one of the most widely assessed PROMs in the present review, used in 33 studies. We have, however, focused on those that specifically reported physical function data (EORTC QLQ-C30-PF) (25 studies). With only eight of these studies also assessing another measure of physical function, it is unclear whether changes in objective measures of physical function consistently are reflected in changes in EORTC QLQ-C30-PF. Further studies are needed to determine the relationship between objective and PROMs of physical function, along with an assessment of which approach reflects the most clinically meaningful assessment and how other factors (such as frailty) may impact these. In addition, the relationship between lean mass and function may not be linear and may have a ceiling. Bye and co-workers highlighted this by demonstrating that above a certain cut-off of lean mass, physical function plateaued. Therefore, that relationship between lean mass and physical

J. McDonald et al.

function may be dependent on absolute value of the former.⁹⁴ Another critical consideration is that different patient-reported measures of physical function assessment may in fact reflect different contexts of day-to-day living and, as such, one size may not fit all.⁹⁵

#### Implications

It is important to note here that statistical significance, or lack thereof, reflects the degree of effectiveness of the intervention on a particular measure, the variance of that measure and the sample size of the study. Furthermore, statistical significance does not necessarily equate to clinical significance. Ultimately, the choice of outcome measures must be guided by a consensus as to the most clinically important outcome and, where possible, quantitative evidence identifying which measure most accurately reflects this outcome. The most appropriate outcome chosen should reflect the aim and mechanism of the intervention.

While there is an increasing use of functional endpoints in cachexia trials, there is, however, no consensus as to which is the optimal measure to use.⁹⁶ Indeed, endpoints are inconsistent across cachexia trials and therefore difficult to translate into clinical practice. This is perhaps best illustrated by the ROMANA trials, which examined anamorelin and used a co-primary endpoint of lean muscle mass and HGS.⁵³ While the trials showed statistically significant results for lean mass, these did not translate to changes in HGS. It is unsurprising as anamorelin is directed at a receptor on Agouti-related protein (AgRP)/neuropeptide Y (NPY) neurons in the hypothalamus, where its primary activity is to enhance appetite and feeding behaviour, and may not have been a relevant outcome for an appetite stimulant.

In the POWER trials (enobosarm), which used SCP as an endpoint, researchers were then criticized that it is not a pure measure of physical function and may be influenced by other parameters such as cardiovascular disease or osteoarthritis.⁹⁷ Similarly, in the other trials, function was measured using step count and results are awaited from trials that are underway regarding the impact on this choice of endpoint.^{98,99}

Understanding which endpoints are best used in cachexia trials will help develop appropriate treatments. Presently, there remains a cascade of endpoints that can be used, ranging between mortality, quality of life, exercise capacity, clinician assessment of function and PROMs, which can include measures of physical function. However, some alignment is needed. This matters not only for regulatory approval but also to patients, particularly when endpoints such as HGS have a clear relationship with prognosis.

Regarding the former, regulatory agencies consider cancer cachexia as a seriously debilitating and/or life-threatening condition. For a drug to reach regulatory approval, it must be efficacious and deliver a meaningful therapeutic effect,¹⁰⁰

which could be changes in lean mass and/or functional capacity vis-à-vis how a patient feels, forms and functions. Any meaningful change should correlate with morbidity or mortality, be related to the disease, have a plausible biological mechanism of action and be related to baseline function. These criteria lend themselves well to HGS being regarded as core functional assessment in cachexia trials—as either a primary or co-primary endpoint. HGS is widely established as being an indicator of muscle strength and general health status, particularly in older people, so the observations of this review are in keeping with this.

Despite the widespread use of HGS, it is notable that other measures including 6MWT and ECOG-PS and KPS were more often found to be statistically significant in trials where HGS was not included. Once again, we cautiously interpret these findings to suggest that these other measures may be more sensitive to changes in physical function, have a lower variability and/or be a reflection of the multiple morbidities that may be present in populations with cancer cachexia. In sarcopenia (as in cachexia), HGS is well recognized for its excellent screening/diagnostic use,¹⁰¹ but sarcopenia trials have shifted away from using it as an endpoint as it is similarly inconsistent or nonresponsive to interventions.¹⁰² Variability in HGS may in part be due to the lack of standardized protocols.¹⁰³ Key to this would be powering trials to this endpoint, and understanding the minimal clinically important difference (MCID) needs to be taken into account. To date, establishing the MCID for grip strength remains a priority. Bohannon conducted a systematic review to answer this. However, as only four studies were eligible, it was difficult to reach firm conclusions.¹⁰⁴ Despite this, he proposed that a change of 5.0-6.5 kg would represent a meaningful change in HGS.

Often, in clinical trials in cachexia, measures of lean mass are assessed alongside measures of function. This is based on the hypothesis that improvements in lean mass would translate to changes in physical function. Both the enobosarm (POWER) and anamorelin (ROMANA) trials adopted this approach, though neither showed improvements in function alongside the improvements in lean mass. There may be several reasons why this is the case including the assumption, perhaps wrong, that muscle mass and function have a linear relationship. Ramage and Skipworth propose a sigmoid relationship between muscle mass and physical function where in the early stages of disease, function is preserved in the presence of loss of muscle (akin to precachexia).¹⁰⁵ Yet when more substantial loss of muscle is present, functional decline is accelerated to the points where recovery is challenging (refractory cachexia). It is key therefore that patient populations for trials are chosen appropriately so that interventions targeting muscle mass and function are given the optimal chance to be efficacious.

In keeping with the theme of improving trial endpoints, there is also the opportunity for functional measures to be combined with biomarkers. Using the systemic inflammatory response as a biomarker would be a good starting point as it is now regarded as being central to the genesis of cancer cachexia and is associated with quality of life and survival.¹⁰⁶ The observations by Song et al. that combining function (HGS) with markers of the systemic inflammatory response has improved utility are of interest and worthy of further exploration.⁹¹ It is also clear that HGS has consistency with ECOG and EORTC-PF both within the clinical trials reviewed and in the literature more generally.¹⁰⁷ HGS may give objective credibility to subjective ECOG-PS and EORTC-PF. This consistency of association and their relationship with survival make HGS, ECOG-PS and EORTC-PF the starting point for the assessment of accelerometer measurements in future studies. It must also be considered that HGS, ECOG-PS and EORTC-PF measure distinctly different elements of physical function and are assessed in different ways. Therefore, it may not be appropriate to assign superiority and may depend upon the intervention and the goal/aim of the intervention.

It is challenging to compare the prognostic utility of different physical functional tests as rarely have these been compared in the same population. ECOG-PS remains the most widely assessed and validated prognostic factor across cancer trials^{5,108} although HGS has been assessed in a large cohort of patients with cancer cachexia⁹¹; however, the lack of work examining the other measures in a prognostic capacity does not mean that they are less useful. As well as the prognostic utility of these different measures, another consideration is patient acceptability. HGS is relatively easy to perform with minimal patient burden compared with a 6MWT, for example, which may cause considerable fatigue and may also cause other symptoms (e.g., breathlessness). These aspects will also influence the choice of endpoint in cachexia trials.

For future research, there are also some important design considerations. Detailed baseline characterization of patients, including co-morbidities, is needed, particularly when they may impact on desired study outcomes. Observed trial design issues could be overcome with standardization of protocols of physical function endpoints, appropriate training, certification, monitoring and recertification of staff conducting endpoint assessments. When performance-based, clinicianassessed and patient-assessed measures of physical function are being assessed, it is important that they are aligned with the mechanism of action of the intervention. Emerging digital technologies could also be used to develop better endpoints¹⁰⁹ and are being assessed by the European Medicines Agency.¹¹⁰

We posit that the appropriate endpoint(s) should be purposely selected to accurately and meaningfully capture patient-important functional improvements in cancer cachexia. The choice of PRO, PS or PF endpoint (or combination of these endpoints) should also be aligned with the underlying mechanism of action of a specific treatment or intervention. For PF endpoints, particular consideration should be given to what aspect of human performance the test actually measures and how an intervention may need to be specifically tailored to ultimately elicit benefit on the selected PF endpoint. For example, if a study seeks to test the efficacy of an anabolic agent on lower extremity muscle strength or power, then the selection of a specific PF test that accurately assesses lower extremity muscle performance should be considered. Similarly, if investigators seek to utilize an exercise intervention to improve deficits in lower extremity physical performance as determined by the SPPB test, the design of the exercise intervention should have training components that specifically target balance, gait and lower extremity muscle performance impairments.

# Strengths and limitations

The review has several strengths. First, a prospective design was adopted (PROSPERO) and a thorough literature search was undertaken including the last three decades of cachexia trials. Second, a robust strategy for appraisal and data extraction was adopted using multiple independent reviewers. Third, we used a validated quality appraisal tool (modified Downs and Black) that enables an appreciation of the quality of included studies to be assessed. Finally, data were presented to allow an appreciation of the broad range of studies and highlight which measures have been used most frequently and with the potential for the greatest sensitivity. A key limitation is that aggregated data do not allow a detailed assessment of the relationship between different measures of physical function and it is therefore not possible to make definitive statements about whether particular measures are preferable because of their accuracy or sensitivity to changes in physical function. Within studies, there were also likely to be significant individual patient differences and there may be an assumption that the cancer is causing weight loss, reduced function and loss of lean mass when it is possible that it may be multifactorial.

# Conclusions

This systematic review has examined key measures of physical function in cachexia clinical trials. Multiple measures have been studied and are in use. HGS stands out as being studied most often and provides an important means of comparing results between trials. However, other measures may be more sensitive to changes in physical function.

# **Conflict of interest statement**

S. D. A. has received grants and personal fees from Vifor and Abbott Vascular, and personal fees for consultancies, trial

committee work and/or lectures from Actimed, Amgen, AstraZeneca, Bayer, Boehringer Ingelheim, BioVentrix, Brahms, Cardiac Dimensions, Cardior, Cordio, CVRx, Cytokinetics, Edwards, Farraday Pharmaceuticals, GSK. HeartKinetics, Impulse Dynamics, Novartis, Occlutech, Pfizer, Repairon, Sensible Medical, Servier, Vectorious and V-Wave. He is named co-inventor of two patent applications regarding MR-proANP (DE 102007010834 and DE 102007022367), but he does not benefit personally from the related issued patents. M. F. has received personal fees from Pfizer. M. J.-H. has received funding from CRUK, NIH National Cancer Institute, IASLC International Lung Cancer Foundation, Lung Cancer Research Foundation, Rosetrees Trust, UKI NETs and NIHR. M. J.-H. has consulted for, and is a member of, the Achilles Therapeutics Scientific Advisory Board and Steering Committee, has received speaker honoraria from Pfizer, Astex Pharmaceuticals, Oslo Cancer Cluster and Bristol Myers Squibb and is co-inventor on a European patent application relating to methods to detect lung cancer (PCT/US2017/028013). R. J. E. S. has received personal fees for consultancy from Artelo, Actimed, Faraday and Helsinn; B. L. has received personal fees for consultancy from Artelo, Actimed, Faraday, Kyona Kirin and Toray.

# **Online supplementary material**

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

# References

- Karnofsky DA, Abelmann WH, Craver LF, Burchendal JH. The use of the nitrogen mustards in the palliative treatment of carcinoma. With particular reference to bronchogenic carcinoma. *Cancer* 1948;1: 634–656.
- Dewys WD, Begg C, Lavin PT, Band PR, Bennett JM, Bertino JR, et al. Prognostic effect of weight loss prior to chemotherapy in cancer patients. Eastern Cooperative Oncology Group. Am J Med 1980;69: 491–497.
- Fearon K, Strasser F, Anker SD, Bosaeus I, Bruera E, Fainsinger RL, et al. Definition and classification of cancer cachexia: an international consensus. *Lancet Oncol* 2011;**12**:489–495.
- Laird BJ, Kaasa S, McMillan DC, Fallon MT, Hjermstad MJ, Fayers P, et al. Prognostic factors in patients with advanced cancer: a comparison of clinicopathological factors and the development of an inflammation-based prognostic system. *Clin Cancer Res* 2013;19:5456–5464.
- Simmons CP, Koinis F, Fallon MT, Fearon KC, Bowden J, Solheim TS, et al. Prognosis in advanced lung cancer—a prospective study examining key clinicopathological factors. *Lung Cancer* 2015;88:304–309.
- Abbass T, Dolan RD, MacLeod N, Horgan PG, Laird BJ, McMillan DCJCNE. Comparison of the prognostic value of MUST, ECOG-PS, mGPS and CT derived body composition analysis in patients with advanced lung cancer. *Clin Nutr ESPEN* 2020;40:349–356.
- 7. Administration FD. 2022.
- Fearon K, Argiles JM, Baracos VE, Bernabei R, Coats A, Crawford J, et al. Request for regulatory guidance for cancer cachexia intervention trials. J Cachexia Sarcopenia Muscle 2015;6:272–274.
- Maddocks M, Granger CL. Measurement of physical activity in clinical practice and research: advances in cancer and

chronic respiratory disease. *Curr Opin Support Palliat Care* 2018;**12**:219–226.

- Salameh JP, Bossuyt PM, McGrath TA, Thombs BD, Hyde CJ, Macaskill P, et al. Preferred Reporting Items for Systematic Review and Meta-Analysis of diagnostic test accuracy studies (PRISMA-DTA): explanation, elaboration, and checklist. BMJ 2020;370:m2632.
- Solheim T, Laird B, Skipworth RJ, Fallon M, Kaasa S, Dajani O. A systematic literature review examining endpoints for cancer cachexia trials CRD42022276710. PROSPERO; 2022.
- 12. Veritas Health Innovation. *Covidence evidence review software*. Melbourne, Australia.
- Downs SH, Black N. The feasibility of creating a checklist for the assessment of the methodological quality both of randomised and non-randomised studies of health care interventions. J Epidemiol Community Health 1998;52:377–384.
- Mendes J, Afonso C, Moreira P, Padrao P, Santos A, Borges N, et al. Association of anthropometric and nutrition status indicators with hand grip strength and gait speed in older adults. JPEN J Parenter Enteral Nutr 2019;43:347–356.
- Laboratories ATSCoPSfCPF. ATS statement: guidelines for the six-minute walk test. Am J Respir Crit Care Med 2002; 166:111–117.
- Bean JF, Kiely DK, LaRose S, Alian J, Frontera WR. Is stair climb power a clinically relevant measure of leg power impairments in at-risk older adults? Arch Phys Med Rehabil 2007;88:604–609.
- Guralnik JM, Simonsick EM, Ferrucci L, Glynn RJ, Berkman LF, Blazer DG, et al. A short physical performance battery assessing lower extremity function: association with self-reported disability and prediction of mortality and nursing home admission. J Gerontol 1994;49:M85–M94.

- Bohannon RW. Reference values for the five-repetition sit-to-stand test: a descriptive meta-analysis of data from elders. *Percept Mot Skills* 2006;103:215–222.
- Fayers P, Aaronson N, Bjordal K, Groenvold M, Curran D, Bottomley A. *The EORTC QLQ-C30 scoring manual*. Brussels: European Organisation for the Research and Treatment of Cancer; 2001. p 5–15.
- Kutz LM, Abel J, Schweizer D, Tribius S, Krull A, Petersen C, et al. Quality of life, HPV-status and phase angle predict survival in head and neck cancer patients under (chemo)radiotherapy undergoing nutritional intervention: results from the prospective randomized HEADNUT-trial. Radiother Oncol 2022;166:145–153.
- Currow DC, Glare P, Louw S, Martin P, Clark K, Fazekas B, et al. A randomised, double blind, placebo-controlled trial of megestrol acetate or dexamethasone in treating symptomatic anorexia in people with advanced cancer. *Sci Rep* 2021;11: 2421.
- Hunter CN, Abdel-Aal HH, Elsherief WA, Farag DE, Riad NM, Alsirafy SA. Mirtazapine in cancer-associated anorexia and cachexia: a double-blind placebo-controlled randomized trial. J Pain Symptom Manage 2021;62:1207–1215.
- Tobberup R, Carus A, Rasmussen HH, Falkmer UG, Jorgensen MG, Schmidt EB, et al. Feasibility of a multimodal intervention on malnutrition in patients with lung cancer during primary anti-neoplastic treatment. *Clin Nutr* 2021;40:525–533.
- Famil-Dardashti A, Haijigholami A, Badri S, Yekdaneh A, Moghaddas A. The role of Trigonella, Cichorium, and Foeniculum herbal combination in the treatment of cancer-induced anorexia/cachexia: a quasi-experimental study. *INt J Cancer Manag* 2020;13.
- 25. Balstad TR, Brunelli C, Pettersen CH, Schonberg SA, Skorpen F, Fallon M, et al.

Power comparisons and clinical meaning of outcome measures in assessing treatment effect in cancer cachexia: secondary analysis from a randomized pilot multimodal intervention trial. *Front Nutr* 2020;7:602775.

- Movahed S, Toussi M, Pahlavani N, Motlagh A, Eslami S, Nematy M, et al. Effects of medical nutrition therapy compared with general nutritional advice on nutritional status and nutrition-related complications in esophageal cancer patients receiving concurrent chemoradiation: a randomized controlled trial. *Mediterrean J Nutr Metab* 2020;13: 265–276.
- Qiu Y, You J, Wang K, Cao Y, Hu Y, Zhang H, et al. Effect of whole-course nutrition management on patients with esophageal cancer undergoing concurrent chemoradiotherapy: a randomized control trial. *Nutrition* 2020;69:110558.
- Hong Y, Wu C, Wu B. Effects of resistance exercise on symptoms, physical function, and quality of life in gastrointestinal cancer patients undergoing chemotherapy. *Integr Cancer Ther* 2020; 19:1534735420954912.
- Kamel FH, Basha MA, Alsharidah AS, Salama AB. Resistance training impact on mobility, muscle strength and lean mass in pancreatic cancer cachexia: a randomized controlled trial. *Clin Rehabil* 2020; 34:1391–1399.
- Storck LJ, Ruehlin M, Gaeumann S, Gisi D, Schmocker M, Meffert PJ, et al. Effect of a leucine-rich supplement in combination with nutrition and physical exercise in advanced cancer patients: a randomized controlled intervention trial. *Clin Nutr* 2020;**39**:3637–3644.
- Bouleuc C, Anota A, Cornet C, Grodard G, Thiery-Vuillemin A, Dubroeucq O, et al. Impact on health-related quality of life of parenteral nutrition for patients with advanced cancer cachexia: results from a randomized controlled trial. Oncologist 2020;25:e843–e851.
- Dehghani M, Mirzaie M, Farhadi P, Rezvani A. The effect of ACE inhibitor on the quality of life amongst patients with cancer cachexia. *Asian Pac J Cancer Prev* 2020;21:325–330.
- Cereda E, Turri A, Klersy C, Cappello S, Ferrari A, Filippi AR, et al. Whey protein isolate supplementation improves body composition, muscle strength, and treatment tolerance in malnourished advanced cancer patients undergoing chemotherapy. *Cancer Med* 2019;8: 6923–6932.
- Obling SR, Wilson BV, Pfeiffer P, Kjeldsen J. Home parenteral nutrition increases fat free mass in patients with incurable gastrointestinal cancer. Results of a randomized controlled trial. *Clin Nutr* 2019; 38:182–190.
- Laviano A, Calder PC, Schols A, Lonnqvist F, Bech M, Muscaritoli M. Safety and tolerability of targeted medical nutrition for cachexia in non-small-cell lung cancer: a

randomized, double-blind, controlled pilot trial. *Nutr Cancer* 2020;**72**:439–450.

- Stuecher K, Bolling C, Vogt L, Niederer D, Schmidt K, Dignass A, et al. Exercise improves functional capacity and lean body mass in patients with gastrointestinal cancer during chemotherapy: a single-blind RCT. Support Care Cancer 2019;27: 2159–2169.
- Wiskemann J, Clauss D, Tjaden C, Hackert T, Schneider L, Ulrich CM, et al. Progressive resistance training to impact physical fitness and body weight in pancreatic cancer patients: a randomized controlled trial. *Pancreas* 2019;**48**:257–266.
- Katakami N, Uchino J, Yokoyama T, Naito T, Kondo M, Yamada K, et al. Anamorelin (ONO-7643) for the treatment of patients with non-small cell lung cancer and cachexia: results from a randomized, double-blind, placebo-controlled, multicenter study of Japanese patients (ONO-7643-04). *Cancer* 2018;**124**:606–616.
- Kouchaki B, Janbabai G, Alipour A, Ala S, Borhani S, Salehifar E. Randomized double-blind clinical trial of combined treatment with megestrol acetate plus celecoxib versus megestrol acetate alone in cachexia-anorexia syndrome induced by GI cancers. *Support Care Cancer* 2018; 26:2479–2489.
- 40. Britton B, Baker AL, Wolfenden L, Wratten C, Bauer J, Beck AK, et al. Eating as treatment (EAT): a stepped-wedge, randomized controlled trial of a health behavior change intervention provided by dietitians to improve nutrition in patients with head and neck cancer undergoing radiation therapy (TROG 12.03). Int J Radiat Oncol Biol Phys 2019;103:353–362.
- Golan T, Geva R, Richards D, Madhusudan S, Lin BK, Wang HT, et al. LY2495655, an antimyostatin antibody, in pancreatic cancer: a randomized, phase 2 trial. J Cachexia Sarcopenia Muscle 2018;9: 871–879.
- Xie M, Chen X, Qin S, Bao Y, Bu K, Lu Y. Clinical study on thalidomide combined with cinobufagin to treat lung cancer cachexia. J Cancer Res Ther 2018;14: 226–232.
- 43. Schink K, Herrmann HJ, Schwappacher R, Meyer J, Orlemann T, Waldmann E, et al. Effects of whole-body electromyostimulation combined with individualized nutritional support on body composition in patients with advanced cancer: a controlled pilot trial. BMC Cancer 2018;18:886.
- Uster A, Ruehlin M, Mey S, Gisi D, Knols R, Imoberdorf R, et al. Effects of nutrition and physical exercise intervention in palliative cancer patients: a randomized controlled trial. *Clin Nutr* 2018;37: 1202–1209.
- 45. Turcott JG, Del Rocio Guillen Nunez M, Flores-Estrada D, Onate-Ocana LF, Zatarain-Barron ZL, Barron F, et al. The effect of nabilone on appetite, nutritional status, and quality of life in lung cancer patients: a randomized, double-blind clin-

ical trial. *Support Care Cancer* 2018;**26**: 3029–3038.

- Jatoi A, Steen PD, Atherton PJ, Moore DF, Rowland KM, Le-Lindqwister NA, et al. A double-blind, placebo-controlled randomized trial of creatine for the cancer anorexia/weight loss syndrome (N02C4): an Alliance trial. Ann Oncol 2017;28: 1957–1963.
- Currow D, Temel JS, Abernethy A, Milanowski J, Friend J, Fearon KC. ROMANA 3: a phase 3 safety extension study of anamorelin in advanced nonsmall-cell lung cancer (NSCLC) patients with cachexia. Ann Oncol 2017;28: 1949–1956.
- 48. Leedo E, Gade J, Granov S, Mellemgaard A, Klausen TW, Rask K, et al. The effect of a home delivery meal service of energy- and protein-rich meals on quality of life in malnourished outpatients suffering from lung cancer: a randomized controlled trial. Nutr Cancer 2017;69: 444–453.
- 49. Solheim TS, Laird BJA, Balstad TR, Stene GB, Bye A, Johns N, et al. A randomized phase II feasibility trial of a multimodal intervention for the management of cachexia in lung and pancreatic cancer. J Cachexia Sarcopenia Muscle 2017;8: 778–788.
- 50. Werner K, Kullenberg de Gaudry D, Taylor LA, Keck T, Unger C, Hopt UT, et al. Dietary supplementation with n-3-fatty acids in patients with pancreatic cancer and cachexia: marine phospholipids versus fish oil—a randomized controlled double-blind trial. *Lipids Health Dis* 2017; 16:104.
- Zietarska M, Krawczyk-Lipiec J, Kraj L, Zaucha R, Malgorzewicz S. Nutritional status assessment in colorectal cancer patients qualified to systemic treatment. *Contemp Oncol (Pozn)* 2017;21:157–161.
- 52. Takayama K, Katakami N, Yokoyama T, Atagi S, Yoshimori K, Kagamu H, et al. Anamorelin (ONO-7643) in Japanese patients with non-small cell lung cancer and cachexia: results of a randomized phase 2 trial. *Support Care Cancer* 2016; 24:3495–3505.
- Temel JS, Abernethy AP, Currow DC, Friend J, Duus EM, Yan Y, et al. Anamorelin in patients with non-smallcell lung cancer and cachexia (ROMANA 1 and ROMANA 2): results from two randomised, double-blind, phase 3 trials. *Lancet Oncol* 2016;**17**:519–531.
- 54. Gavazzi C, Colatruglio S, Valoriani F, Mazzaferro V, Sabbatini A, Biffi R, et al. Impact of home enteral nutrition in malnourished patients with upper gastrointestinal cancer: a multicentre randomised clinical trial. *Eur J Cancer* 2016;64: 107–112.
- 55. Woo SM, Joo J, Kim SY, Park SJ, Han SS, Kim TH, et al. Efficacy of pancreatic exocrine replacement therapy for patients with unresectable pancreatic cancer in a randomized trial. *Pancreatology* 2016;**16**: 1099–1105.

1353921906009, 0, Downloaded from https://onlinelibrary.wiley.com/doi/10.1002/jssm.13321 by University College London UCL Library Services, Wiley Online Library on [11.09/2023]. See the Terms and Conditions

(https://onlinelibrary

.wiley

conditions) on Wiley Online Library for rules of use; OA articles are governed by the applicable Creative Commons

- 56. Stewart Coats AJ, Ho GF, Prabhash K, von Haehling S, Tilson J, Brown R, et al. Espindolol for the treatment and prevention of cachexia in patients with stage III/IV non-small cell lung cancer or colorectal cancer: a randomized, doubleblind, placebo-controlled, international multicentre phase II study (the ACT-ONE trial). J Cachexia Sarcopenia Muscle 2016;7:355–365.
- Capozzi LC, McNeely ML, Lau HY, Reimer RA, Giese-Davis J, Fung TS, et al. Patientreported outcomes, body composition, and nutrition status in patients with head and neck cancer: results from an exploratory randomized controlled exercise trial. *Cancer* 2016;**122**:1185–1200.
- Focan C, Houbiers G, Gilles L, Van Steeland T, Georges N, Maniglia A, et al. Dietetic and psychological mindfulness workshops for the management of cachectic cancer patients. A randomized study. Anticancer Res 2015;35: 6311–6315.
- Garcia JM, Boccia RV, Graham CD, Yan Y, Duus EM, Allen S, et al. Anamorelin for patients with cancer cachexia: an integrated analysis of two phase 2, randomised, placebo-controlled, doubleblind trials. *Lancet Oncol* 2015;16: 108–116.
- 60. Hong DS, Hui D, Bruera E, Janku F, Naing A, Falchook GS, et al. MABp1, a first-inclass true human antibody targeting interleukin- $1\alpha$  in refractory cancers: an openlabel, phase 1 dose-escalation and expansion study. *Lancet Oncol* 2014;**15**: 656–666.
- Poulsen GM, Pedersen LL, Osterlind K, Baeksgaard L, Andersen JR. Randomized trial of the effects of individual nutritional counseling in cancer patients. *Clin Nutr* 2014;**33**:749–753.
- 62. Pottel L, Lycke M, Boterberg T, Pottel H, Goethals L, Duprez F, et al. Echium oil is not protective against weight loss in head and neck cancer patients undergoing curative radio (chemo)therapy: a randomised-controlled trial. BMC Complement Altern Med 2014;14:382.
- Dobrila-Dintinjana R, Trivanovic D, Zelic M, Radic M, Dintinjana M, Petranovic D, et al. Nutritional support in patients with colorectal cancer during chemotherapy: does it work? *Hepatogastroenterology* 2013;**60**:475–480.
- Dobs AS, Boccia RV, Croot CC, Gabrail NY, Dalton JT, Hancock ML, et al. Effects of enobosarm on muscle wasting and physical function in patients with cancer: a double-blind, randomised controlled phase 2 trial. *Lancet Oncol* 2013;14: 335–345.
- 65. Maccio A, Madeddu C, Gramignano G, Mulas C, Floris C, Sanna E, et al. A randomized phase III clinical trial of a combined treatment for cachexia in patients with gynecological cancers: evaluating the impact on metabolic and inflammatory profiles and quality of life. *Gynecol Oncol* 2012;**124**:417–425.

- Kraft M, Kraft K, Gartner S, Mayerle J, Simon P, Weber E, et al. L-Carnitine-supplementation in advanced pancreatic cancer (CARPAN)—a randomized multicentre trial. *Nutr J* 2012;**11**:52.
- Wen HS, Li X, Cao YZ, Zhang CC, Yang F, Shi YM, et al. Clinical studies on the treatment of cancer cachexia with megestrol acetate plus thalidomide. *Chemotherapy* 2012;58:461–467.
- Madeddu C, Dessi M, Panzone F, Serpe R, Antoni G, Cau MC, et al. Randomized phase III clinical trial of a combined treatment with carnitine + celecoxib ± megestrol acetate for patients with cancer-related anorexia/cachexia syndrome. *Clin Nutr* 2012;**31**:176–182.
- 69. Baldwin C, Spiro A, McGough C, Norman AR, Gillbanks A, Thomas K, et al. Simple nutritional intervention in patients with advanced cancers of the gastrointestinal tract, non-small cell lung cancers or meso-thelioma and weight loss receiving chemotherapy: a randomised controlled trial. J Hum Nutr Diet 2011;24:431–440.
- Silander E, Nyman J, Bove M, Johansson L, Larsson S, Hammerlid E. Impact of prophylactic percutaneous endoscopic gastrostomy on malnutrition and quality of life in patients with head and neck cancer: a randomized study. *Head Neck* 2012; 34:1–9.
- Mantovani G, Maccio A, Madeddu C, Gramignano G, Serpe R, Massa E, et al. Randomized phase III clinical trial of five different arms of treatment for patients with cancer cachexia: interim results. *Nutrition* 2008;**24**:305–313.
- Wiedenmann B, Malfertheiner P, Friess H, Ritch P, Arseneau J, Mantovani G, et al. A multicenter, phase II study of infliximab plus gemcitabine in pancreatic cancer cachexia. J Support Oncol 2008;6:18–25.
- Fearon KC, Barber MD, Moses AG, Ahmedzai SH, Taylor GS, Tisdale MJ, et al. Double-blind, placebo-controlled, randomized study of eicosapentaenoic acid diester in patients with cancer cachexia. J Clin Oncol 2006;24:3401–3407.
- Ravasco P, Monteiro-Grillo I, Marques Vidal P, Camilo ME. Impact of nutrition on outcome: a prospective randomized controlled trial in patients with head and neck cancer undergoing radiotherapy. *Head Neck* 2005;**27**:659–668.
- Gordon JN, Trebble TM, Ellis RD, Duncan HD, Johns T, Goggin PM. Thalidomide in the treatment of cancer cachexia: a randomised placebo controlled trial. *Gut* 2005;54:540–545.
- Isenring EA, Capra S, Bauer JD. Nutrition intervention is beneficial in oncology outpatients receiving radiotherapy to the gastrointestinal or head and neck area. *Br J Cancer* 2004;91:447–452.
- 77. Fearon KC, Von Meyenfeldt MF, Moses AG, Van Geenen R, Roy A, Gouma DJ, et al. Effect of a protein and energy dense N-3 fatty acid enriched oral supplement on loss of weight and lean tissue in cancer

cachexia: a randomised double blind trial. *Gut* 2003;**52**:1479–1486.

- Bruera E, Strasser F, Palmer JL, Willey J, Calder K, Amyotte G, et al. Effect of fish oil on appetite and other symptoms in patients with advanced cancer and anorexia/cachexia: a double-blind, placebocontrolled study. J Clin Oncol 2003;21: 129–134.
- Erkurt E, Erkisi M, Tunali C. Supportive treatment in weight-losing cancer patients due to the additive adverse effects of radiation treatment and/or chemotherapy. J Exp Clin Cancer Res 2000;19: 431–439.
- McMillan DC, Wigmore SJ, Fearon KC, O'Gorman P, Wright CE, McArdle CS. A prospective randomized study of megestrol acetate and ibuprofen in gastrointestinal cancer patients with weight loss. Br J Cancer 1999;**79**:495–500.
- Westman G, Bergman B, Albertsson M, Kadar L, Gustavsson G, Thaning L, et al. Megestrol acetate in advanced, progressive, hormone-insensitive cancer. Effects on the quality of life: a placebo-controlled, randomised, multicentre trial. *Eur J Cancer* 1999;**35**:586–595.
- Vadell C, Segui MA, Gimenez-Arnau JM, Morales S, Cirera L, Bestit I, et al. Anticachectic efficacy of megestrol acetate at different doses and versus placebo in patients with neoplastic cachexia. *Am J Clin Oncol* 1998;**21**:347–351.
- De Conno F, Martini C, Zecca E, Balzarini A, Venturino P, Groff L, et al. Megestrol acetate for anorexia in patients with far-advanced cancer: a double-blind controlled clinical trial. *Eur J Cancer* 1998; 34:1705–1709.
- Neri B, Garosi VL, Intini C. Effect of medroxyprogesterone acetate on the quality of life of the oncologic patient: a multicentric cooperative study. *Anticancer Drugs* 1997;8:459–465.
- Lissoni P, Paolorossi F, Tancini G, Barni S, Ardizzoia A, Brivio F, et al. Is there a role for melatonin in the treatment of neoplastic cachexia? *Eur J Cancer* 1996;**32A**: 1340–1343.
- Gebbia V, Testa A, Gebbia N. Prospective randomised trial of two dose levels of megestrol acetate in the management of anorexia-cachexia syndrome in patients with metastatic cancer. *Br J Cancer* 1996; 73:1576–1580.
- Simons JP, Aaronson NK, Vansteenkiste JF, ten Velde GP, Muller MJ, Drenth BM, et al. Effects of medroxyprogesterone acetate on appetite, weight, and quality of life in advanced-stage non-hormonesensitive cancer: a placebo-controlled multicenter study. J Clin Oncol 1996;14: 1077–1084.
- Lai YL, Fang FM, Yeh CY. Management of anorexic patients in radiotherapy: a prospective randomized comparison of megestrol and prednisolone. J Pain Symptom Manage 1994;9:265–268.
- 89. Downer S, Joel S, Allbright A, Plant H, Stubbs L, Talbot D, et al. A double blind

placebo controlled trial of medroxyprogesterone acetate (MPA) in cancer cachexia. *Br J Cancer* 1993;**67**: 1102–1105.

- Vanhoutte G, van de Wiel M, Wouters K, Sels M, Bartolomeeussen L, De Keersmaecker S, et al. Cachexia in cancer: what is in the definition? *BMJ Open Gastroenterol* 2016;**3**:e000097.
- Song M, Zhang Q, Tang M, Zhang X, Ruan G, Zhang X, et al. Associations of low hand grip strength with 1 year mortality of cancer cachexia: a multicentre observational study. J Cachexia Sarcopenia Muscle 2021;12:1489–1500.
- Zhuang CL, Zhang FM, Li W, Wang KH, Xu HX, Song CH, et al. Associations of low handgrip strength with cancer mortality: a multicentre observational study. J Cachexia Sarcopenia Muscle 2020;11: 1476–1486.
- LeBlanc TW, Nipp RD, Rushing CN, Samsa GP, Locke SC, Kamal AH, et al. Correlation between the international consensus definition of the cancer anorexia-cachexia syndrome (CACS) and patient-centered outcomes in advanced non-small cell lung cancer. J Pain Symptom Manage 2015;49: 680–689.
- 94. Bye A, Sjoblom B, Wentzel-Larsen T, Gronberg BH, Baracos VE, Hjermstad MJ, et al. Muscle mass and association to quality of life in non-small cell lung cancer patients. J Cachexia Sarcopenia Muscle 2017;8:759–767.
- 95. Schurr T, Loth F, Lidington E, Piccinin C, Arraras JI, Groenvold M, et al. Patient-reported outcome measures for physical function in cancer patients: content comparison of the EORTC CAT Core, EORTC QLQ-C30, SF-36, FACT-G, and PROMIS measures using the International Classification of Functioning, Disability and

Health. BMC Med Res Methodol 2023; 23:21.

- 96. Balstad TR, Solheim TS, Laird BJ. Endpoints in clinical trials in cancer cachexia: where to start? 2018.
- Jatoi A, Ritter HL, Dueck A, Nguyen PL, Nikcevich DA, Luyun RF, et al. A placebocontrolled, double-blind trial of infliximab for cancer-associated weight loss in elderly and/or poor performance non-small cell lung cancer patients (N01C9). Lung Cancer 2010;68:234–239.
- Solheim TS, Laird BJA, Balstad TR, Bye A, Stene G, Baracos V, et al. Cancer cachexia: rationale for the MENAC (Multimodal-Exercise, Nutrition and Anti-inflammatory medication for Cachexia) trial. *BMJ Support Palliat Care* 2018;8:258–265.
- 99. Hall CC, Skipworth RJE, Blackwood H, Brown D, Cook J, Diernberger K, et al. A randomized, feasibility trial of an exercise and nutrition-based rehabilitation programme (ENeRgy) in people with cancer. J Cachexia Sarcopenia Muscle 2021; In press.
- 100. FDA. Development and approval process. 2022.
- Bhasin S, Travison TG, Manini TM, Patel S, Pencina KM, Fielding RA, et al. Sarcopenia definition: the position statements of the sarcopenia definition and outcomes consortium. J Am Geriatr Soc 2020;68: 1410–1418.
- Tieland M, Verdijk LB, de Groot LC, van Loon LJ. Handgrip strength does not represent an appropriate measure to evaluate changes in muscle strength during an exercise intervention program in frail older people. Int J Sport Nutr Exerc Metab 2015;25:27–36.
- McGrath R, Cawthon PM, Clark BC, Fielding RA, Lang JJ, Tomkinson GR. Recommendations for reducing heterogeneity

in handgrip strength protocols. *J Frailty Aging* 2022;**11**:143–150.

- Bohannon RW. Minimal clinically important difference for grip strength: a systematic review. J Phys Ther Sci 2019;31: 75–78.
- 105. Ramage MI, Skipworth RJE. The relationship between muscle mass and function in cancer cachexia: smoke and mirrors? *Curr Opin Support Palliat Care* 2018;12: 439–444.
- Laird BJ, Fallon M, Hjermstad MJ, Tuck S, Kaasa S, Klepstad P, et al. Quality of life in patients with advanced cancer: differential association with performance status and systemic inflammatory response. *J Clin Oncol* 2016;**34**:2769–2775.
- 107. Lopez-Bueno R, Andersen LL, Koyanagi A, Nunez-Cortes R, Calatayud J, Casana J, et al. Thresholds of handgrip strength for all-cause, cancer, and cardiovascular mortality: a systematic review with doseresponse meta-analysis. Ageing Res Rev 2022;82:101778.
- 108. Simmons CPL, McMillan DC, McWilliams K, Sande TA, Fearon KC, Tuck S, et al. Prognostic tools in patients with advanced cancer: a systematic review. J Pain Symptom Manage 2017;**53**:962–70 e10, 962, 970.e10.
- 109. Manta C, Patrick-Lake B, Goldsack JC. Digital measures that matter to patients: a framework to guide the selection and development of digital measures of health. *Digit Biomark* 2020;4:69–77.
- Smail EJ, Alpert JM, Mardini MT, Kaufmann CN, Bai C, Gill TM, et al. Feasibility of a smartwatch platform to assess ecological mobility: Real-time Online Assessment and Mobility Monitor (ROAMM). J Gerontol A Biol Sci Med Sci 2023;**78**:821–830.