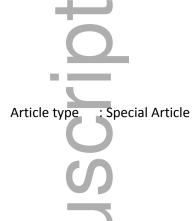
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A North American Expert Opinion Statement on Sarcopenia in Liver Transplantation

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Abstract:

Loss of muscle mass and function, or sarcopenia, is a common feature of cirrhosis and contributes significantly to morbidity and mortality in this population. Sarcopenia is a main indicator of adverse outcomes in this population, including poor quality of life, hepatic decompensation, mortality in patients with cirrhosis evaluated for LT, longer hospital and intensive care unit stay, higher incidence of infection following LT, and higher overall health care cost. While it is clear that muscle mass is an important predictor of LT outcomes, many questions remain, including the best modality for assessing muscle mass, the optimal cut-off values for sarcopenia, the ideal timing and frequency of muscle mass assessment, and how to best incorporate the concept of sarcopenia into clinical decision making. For that reason, we assembled a group of experts to form the North American Working Group on Sarcopenia in Liver Transplantation to use evidence from the medical literature to address these outstanding questions regarding sarcopenia in LT. We believe sarcopenia assessment should be considered in all patients with cirrhosis evaluated for liver transplantation. Skeletal muscle index (SMI) assessed by computed tomography constitutes the best studied technique for assessing sarcopenia in patients with cirrhosis. Cut-off values for sarcopenia, defined as SMI <50 cm²/m² in male and <39 cm²/m² in female patients constitute the validated definition for sarcopenia in patients with cirrhosis. The management of sarcopenia requires a multi-pronged approach including nutrition, exercise and additional pharmacological therapy as deemed necessary. Future studies should evaluate whether recovery of sarcopenia with nutritional management in combination with an exercise program is sustainable, and how improvement in muscle mass might be associated with improvement in clinical outcomes.

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Introduction

Sarcopenia, the disproportionate loss of muscle mass, is common in patients with cirrhosis awaiting liver transplantation (LT). Sarcopenia has been shown to be a significant risk factor for waitlist mortality, postoperative complications, and post-LT death (1-5). The new international statistical classification of diseases and related health problems, 10th revision (ICD-10-CM) (M62.84) code for sarcopenia represents a significant recognition of sarcopenia as a disease (6). While it is clear that muscle mass is an important predictor of LT outcomes, many questions remain, including the best modality for measuring muscle mass, the optimal cut-off values for sarcopenia, the ideal timing and frequency of muscle mass assessment, and how to best incorporate the concept of sarcopenia into clinical decision making.

Therefore, in 2018, we convened a group of experts to form the North American Working Group on Sarcopenia in Liver Transplantation to use evidence from the medical literature to address these outstanding questions regarding sarcopenia in LT. This document represents the product of our efforts to develop a statement by experts in the field on the current state of knowledge and key gaps for future high impact research on sarcopenia in the LT setting.

The importance of sarcopenia as a construct

Sarcopenia is a term derived from the Greek *sarco* (flesh) and *penia* (deficiency). Sarcopenia was initially defined as age-related loss of skeletal muscle (7), but has since been expanded to reflect low muscle mass leading to negative effects on physical performance and clinical outcomes across a broad range of disease states outside of geriatric populations.

Conceptually, sarcopenia is only one – but likely the dominant – component of the larger construct of global physical dysfunction that is prevalent in patients with cirrhosis that has most recently been described with the term "frailty". While frailty is the manifesting <u>symptom</u> of impaired global physical functioning, loss of muscle mass is an obvious <u>sign</u> that frailty may be present. Sarcopenia is the metric of clinically-relevant skeletal muscle depletion that can be objectively measured in clinical practice and is least likely to be affected by acute illness or alterations in cognitive function. For this reason, our working group chose to define sarcopenia using only muscle mass, although we acknowledge that muscle *function* (e.g., grip strength) is incorporated into definitions of sarcopenia by other groups including the European Working Group on Sarcopenia in Older People and Asian Working Group for Sarcopenia (8, 9).

Although multiple definitions of sarcopenia for patients with cirrhosis exist in the literature, low muscle mass, regardless of how it is measured, is a powerful predictor of clinically relevant adverse outcomes, including poor quality of life (10), hepatic decompensation (11), mortality in patients with cirrhosis on the LT waitlist (3, 11-14), longer hospital and intensive care unit stay (2,

15), higher incidence of infection following LT (2, 16), higher overall health care cost (17), and post-LT mortality (3, 18).

Aside from being an important marker of pre and post-LT mortality, sarcopenia is associated with additional important clinical parameters independent of scoring systems for severity of liver dysfunction. Skeletal muscle plays an integral role in ammonia detoxification, and sarcopenia has been identified as an independent risk factor for hepatic encephalopathy (HE) in patients with cirrhosis (19). Other complications such as ascites and infections are more common in sarcopenic patients compared to non-sarcopenic patients with cirrhosis (2). Modalities to Evaluate Muscle Mass in the Liver Transplant Candidate

There currently exists significant heterogeneity in the metrics used to define sarcopenia in

the published domain. The great challenge in identifying a single standard is that many modalities exist for muscle mass quantification, including anthropometry, bioelectrical impedance analysis (BIA), dual-energy X-ray absorptiometry (DEXA), ultrasound (US), magnetic resonance imaging (MRI) and computed tomography (CT). The advantages and limitations of each modality as they relate to measurement of muscle mass in patients with end-stage liver disease (ESLD) are summarized in Table 1.

We advocate for the use of CT-based skeletal muscle area, measured at the third lumbar vertebra on an abdominal CT scan (either non-contrast or contrast-enhanced scans are acceptable), in patients with cirrhosis for the following reasons (Please see example in Figure 1). First, cross-sectional imaging is commonly used in transplant centers to monitor for hepatocellular carcinoma (HCC) and to evaluate the vascular and biliary anatomy for surgical planning. While preliminary reports suggest that MRI-based imaging yields equivalent results to CT scans (20), CT scans tend to be more widely available, of lower cost, and more rapidly performed in clinical practice. Second, the majority of published reports on sarcopenia in LT have used standard-of-care CT imaging, summarized in Table 2. Several key questions remain regarding the use of CT-based estimation of skeletal muscle mass, including the use of values below specific percentiles (*i.e.* 5th percentile) of age and sex-matched population or optimal cut-points for mortality discrimination, sensitivity of changes over time as well as the validity of measurement of psoas alone *versus* the total muscle area (21).

Defining sarcopenia

A large, multi-center study of 396 patients with cirrhosis from five North American liver transplant centers established standardized cut-off values of skeletal muscle index (SMI) at <50 cm²/m² and <39 cm²/m² to define sarcopenia in men and women with cirrhosis, respectively (22).

These sex-specific SMI cut-points were strongly associated with pre-LT mortality independent of age and MELD score (15).

Although muscle mass has been shown in multiple studies to be associated with post-LT mortality (3, 18, 23), data reporting pre-LT muscle indexes associated with post-LT adverse outcome is limited. Recently, SMI<48 cm²/m² in acutely ill men undergoing urgent evaluation and LT, was associated with higher post-LT mortality (24).

When assessing sarcopenia by SMI on an abdominal CT scan, there does not appear to be a large difference between measurements at L3 *versus* L4 vertebra (25). There is also excellent agreement between the various software programs (*i.e.* SliceOmatic, ImageJ, etc.) with respect to measurements of abdominal skeletal muscle area (26).

SMI seems to be a more complete and robust measurement than individual measurement of the psoas muscle or the psoas muscle index (PMI), especially in men with cirrhosis. In addition, low PMI identifies an incomplete subset of patients at increased risk of mortality indicated by low SMI (21).

While sarcopenia has classically been associated with increased mortality in both men and women with cirrhosis, emerging evidence suggests that sarcopenia is associated with disproportionately higher rates of mortality in men as compared to women (27). This emphasizes the importance of survival analysis stratification by sex rather than simply adjusting multivariable models for sex. Furthermore, the role of ethnicity in baseline muscle mass and muscle loss has not yet been determined in patients with cirrhosis (28). While more data are needed, the prevalence of sarcopenia within each BMI category is another consideration when building a definition to incorporate into clinical practice. Lastly, divergent study outcomes such as overall mortality in evaluated patients, waitlist mortality in listed patients, post-LT mortality in patients undergoing LT, and short term *vs.* long-term outcomes, confound the comparison between published studies and development of generalized definitions.

Key points

- 1. Standardized tools and validation between techniques are important considerations for the evaluation of sarcopenia in patients with end-stage liver disease awaiting LT.
- 2. CT constitutes the best-studied technique for measuring sarcopenia in patients with cirrhosis.
- FLEXIT (Fitness, Life Enhancement, and Exercise in Liver Transplantation Consortium) cut-off values (SMI <50 cm²/m² in men and <39 cm²/m² in women)

constitute the most robust definition for sarcopenia in patients with end-stage liver disease awaiting LT.

Recommendations

1. We recommend the use of SMI as a marker of sarcopenia for outcome prediction.

- 2. We recommend the use of FLEXIT cut-off values to define sarcopenia in cirrhosis in order to characterize cohorts of patients for prospective clinical trials.
- 3. Future studies to define sarcopenia should be established for use in clinical practice with consideration of sex, age and ethnicity.

Incorporating Sarcopenia into Clinical Practice

Although *severe* sarcopenia may be easy to identify visually (i.e., the "eyeball test"), objective and validated measures are needed to detect early stages of muscle loss, when interventions to slow the progression may be more effective. Ascites, obesity, and body fat distribution may render sarcopenia less apparent, and early stages of muscle loss are often not visually obvious. Furthermore, quantification of muscle mass provides objective data, which is especially critical in the setting of LT.

One of the advantages of determination of muscle mass is that it can be done on crosssectional imaging (e.g., abdominal CT or MRI), which is often performed in LT patients as standard of care. Radiologists can provide total abdominal skeletal muscle area at the L3 vertebral level from these routine scans, from which SMI can easily be calculated. We anticipate that this will enable more widespread incorporation of the information provided from muscle mass measurement into clinical practice.

Objective assessment of sarcopenia in the LT candidate is important to two major reasons: 1) clinical decision-making and 2) intervention. We expand upon both of these areas here. *Incorporating sarcopenia into clinical decision-making*

It is important to note that the North American Working Group on Sarcopenia in Liver Transplantation recommends that sarcopenia should **not** be the **sole** criterion for declining or delisting candidates for LT. A significant limitation of utilizing sarcopenia for risk stratification is the lack of a threshold value of muscle loss that predicts health outcomes prohibitive for a particular intervention such as LT. Furthermore, the threshold of "futility" is not universally defined but, instead, varies by center, clinician, and patient values. Rather, we advocate that an objective metric of sarcopenia be taken into consideration within the full context of the other medical, physical, functional and psychosocial factors of each individual patient with respect to their transplant candidacy as well as their own values of what a successful transplant looks like.

That being said, sarcopenia has important implications for a patient awaiting LT. At the minimum, in the outpatient setting, a patient with sarcopenia should be counseled that his or her higher risk of waitlist mortality exceeds that predicted by the MELDNa score and that they have a higher risk of complication after LT. This information may help to motivate the patient to seek a faster path to transplant, including living donor LT or to accept higher-risk donor livers. In addition, this information may motivate patients and providers to engage in interventions that might mitigate muscle loss (see next section).

With respect to clinical decision-making, sarcopenia may hold a unique place within the <u>inpatient</u> setting where performance-based assessments of frailty may be misleading due to acute changes in functional performance that may not accurately reflect their underlying "steady state" physiologic reserve. An objective assessment of muscle mass may indicate that the patient has good underlying physiologic reserve that will support a full peri-operative recovery. For a patient <u>without</u> multiple risk factors for poor post-LT outcomes, sarcopenia *alone* is not sufficient to deny LT but may guide the decision about the quality of liver to accept in attempt to minimize liver-related complications and optimize overall patient recovery (29). For a patient *with* multiple co-morbidities that may also negatively impact post-LT outcomes, identification of sarcopenia - in combination with these other medical risk factors - may be added objective evidence to *not* proceed with LT.

Incorporating sarcopenia into management and treatment of the LT candidate

We recommend that measurement of muscle mass best be integrated into clinical practice to identify patients for "prehabilitation" programs focused on optimizing nutrition and physical activity (30). The management of sarcopenia requires a multi-pronged approach including nutrition, exercise and additional pharmacological therapy as deemed necessary. The incorporation of behavioral change strategies while delivering nutrition (31) and exercise prescriptions (32) encourages exploration of the patient's individual personal and social factors to motivate and increase the likelihood of patient engagement (33). We recommend the following strategies based on the current available evidence:

a. A nutrition prescription. At minimum, this consists of three major components:

 A target caloric intake recommendation. For non-obese individuals (BMI <30 kg/m²), the latest European Association for the Study of the Liver (EASL) Clinical Practice Guidelines on nutrition in chronic liver disease recommend an optimal daily energy intake of at least 35 kcal/kg of actual body weight corrected for fluid retention. In obese patients, a

tailored moderately hypocaloric diet (with a reduction of 500 to 800 kcal/day) has been suggested (34). The International Society for Hepatic Encephalopathy and Nitrogen Metabolism (ISHEN) Consensus (35) provides additional BMI-stratified target caloric recommendations based on an ideal body weight (also corrected for fluid retention), including BMI ranges of 20-30, 30-40 and >40 kg/m². Supplemental enteral nutrition should

be considered in hospitalized patients who are unable to meet calorie intake targets with oral intake alone (34, 36).

2. A target protein intake recommendation. Protein restriction is not necessary in patients with hepatic encephalopathy (37). Guidelines recommend a daily protein intake of 1.2-1.5 grams/kg protein (34, 35) with further study required as to whether dairy/vegetable protein may have a benefit over meat protein in the setting of hepatic encephalopathy (34) and the variable impact of protein across the spectrum of liver disease severity.

3. A late-evening snack recommendation. In an attempt to shorten the overnight fasting period, patients are advised to take a snack shortly before bedtime or during nighttime hours and to eat breakfast (34, 38). We would support the recommendation of a late-night snack across Child Pugh classes. The optimal composition of the late-evening snack is not clearly defined, with recommendations varying from branched chain amino acids (BCAA) containing supplements (39) to snacks containing ~50 grams of carbohydrates and 10-20 grams of protein (31).

b) Exercise. Although an evidence-based exercise prescription is still not available for patients with cirrhosis, extrapolating current knowledge from the 11 published studies on exercise in patients with cirrhosis (please refer to Table 4) into practice, it is recommended for patients to perform moderate intensity exercise for no less than 30 minutes per day, starting with a brief warm-up (5-10 minutes) and ending with a stretching / cool-down phase (5-10 minutes), 3-5 times per week, aiming for a total of 150-200 minutes of exercise per week (32). Exercise bouts should have a duration of no less than 5-10 minutes, depending on disease status and tolerance, with patients commonly having to undergo multiple exercise bouts per day in order to accomplish these goals. In general, it is recommended that aerobic and resistance training be combined in a 3:2 ratio, although for the purpose of improving sarcopenia this ratio should favor resistance training (40).

c) Pharmacotherapy (Table 3)

1. Vitamin D3. Vitamin D deficiency is highly prevalent in cirrhosis and is a well-defined contributor to sarcopenia. Detection and repletion with cholecalciferol is standard practice in transplant hepatology (41).

2. Ammonia lowering treatments. Excess ammonia delivered to muscle is the pre-eminent metabolic driver of sarcopenia (42). It is reasonable to expect that current and potentially future ammonia lowering treatments (43, 44) to prevent encephalopathy will also be helpful in minimizing sarcopenia.

3. Hormonal therapy. Standard hormonal therapy to sustain a euthyroid and euglycemic state defends muscle mass and function. In addition, the majority of men with advanced cirrhosis have hypogonadism as measured by decreased total and free testosterone levels. A randomized trial showed that intramuscular androgen therapy of hypogonadal cirrhotic men improved muscle mass, bone mass and hemoglobin A1C (45); transdermal therapy also appears effective with fewer potential adverse events.

4. L-Carnitine. L-carnitine is a quaternary amine (3-hydroxy-4-N-trimethylaminobutyrate) needed for fatty acid oxidation. Binding of L-carnitine to acetyl groups enables movement of acetylated fatty acids into mitochondria and their oxidation to generate energy in the form of ATP. Carnitine supplementation was recently reported to suppress muscle loss in an initial study of cirrhotic patients (46).

Key Points

1. We believe sarcopenia should not be a sole criterion on which to determine candidacy for LT.

2. Attaining nutritional goals appropriate for patients with cirrhosis is currently the predominant treatment for sarcopenia.

Recommendations

1. We recommend that the presence or absence of sarcopenia be considered as part of the multidisciplinary assessment in cirrhosis.

- 2. Until more evidence is available, we recommend for patients with cirrhosis and sarcopenia to exercise 150-200 minutes per week, including both aerobic and resistance
- training (ratio favoring the latter), along with a nutritional intervention with an adiposity-tailored caloric intake that favors protein with key amino acids and aims to prevent starvation.
- 3. Male patients with cirrhosis and sarcopenia are potentially eligible for pharmacologic treatment with testosterone replacement.

Sarcopenia in Children

Children with irreversible liver disease offer a complex challenge to clinical care teams that is distinct from adults since they have a time-limited opportunity for growth and development and may suffer lifelong consequences if not expeditiously transplanted.

The PELD (Pediatric End-Stage Liver Disease) score does not adequately capture waitlist and peri-operative risk nor the extent to which ESLD impairs functional development. Clinical (weight, height, anthropometrics) and biochemical (serum total protein and albumin) data fail to fully characterize malnutrition in chronically ill children/those with end-stage organ/liver failure and are often confounded. Objective nutritional biomarkers are lacking for children with ESLD.

Children with ESLD suffer from malnutrition, muscle wasting, and deconditioning that result from the insidious effects of underlying hepatic synthetic dysfunction and systemic inflammation. Broader non-laboratory assessment metrics are needed to more fully capture the extent of ill health status associated with chronic or end-stage liver disease in children. Pilot studies have demonstrated that children with ESLD have smaller psoas muscle areas than healthy controls, and that the psoas muscle area does not correlate with weight z-scores nor PELD score (47, 48). Significant gaps exist, including the (Table 5) absence of unifying definition of sarcopenia in growing children, the paucity of evidence to date of HOW to measure sarcopenia in children, with different modalities utilized, the evolving validated reference data for total psoas muscle area (tPMA) (via CT abdominal images) in children, the near term impact of interventions such as nutritional support, exercise programs etc. limited by challenges of longitudinal/serial evaluation of sarcopenia due to cumbersome (DEXA) or radiation heavy (CT) assessment methods, the lack of full understanding of how the presence of sarcopenia pre-LT affect outcomes post LT, and whether sarcopenia be considered in clinical decision-making.

Key Points

- 1. The majority of children are too young to undergo functional or performance-based (frailty) testing before time of liver transplantation, underscoring the special role and importance of muscle mass assessment in this vulnerable patient population.
- 2. Pilot studies demonstrate that total psoas muscle area (tPMA) increases over time until late adolescence and is smaller in pediatric ESLD than healthy controls.

Recommendations

Sarcopenia assessment should be considered in children with ESLD. Amongst those
pediatric patients requiring clinically indicated CT imaging, the availability of recent
pediatric specific novel tools providing age- and sex-specific reference growth curves
will facilitate targeted interventions and their near-term impacts.

2. Targeting non-invasive assessment strategies for children is a high needs research area.

Sarcopenia in Special Patient Populations

- a. Sarcopenic obesity can be present in 20-40% of cirrhotic LT candidates (49, 50).
 Patients with sarcopenia are more likely to be obese and those with sarcopenic obesity are more likely to have non-alcoholic fatty liver disease as the etiology of liver disease (50). Of particular concern is the fact that sarcopenia can be difficult to recognize in the presence of obesity. Future research should include measures of obesity beyond BMI such as CT measures of adipose tissue.
- b. Chronic kidney disease (CKD) alone can affect muscle mass. Sarcopenia occurs in roughly 10% of patients with CKD alone and is associated with increased mortality (51). In renal transplant recipients, sarcopenia may persist for years after transplantation, with age and duration of dialysis being important predictors (52). Little is known about the relationship between sarcopenia and CKD in LT candidates. Normal renal function in patients with cirrhosis has been correlated with higher measures of muscle mass (53). Data on measures such as duration of renal insufficiency and renal replacement therapy in LT candidates with sarcopenia are not available.

Future directions for research

Sarcopenia has become a topic of prolific exploration in patients with ESLD over the last few years. Currently, the evaluation of sarcopenia in patients with cirrhosis appears in over 1,500 publications. Important research questions that merit exploration for sarcopenia assessment in patients with ESLD include 1) the availability of reliable, accessible and practical tools to be used in clinical practice, 2) the optimal frequency of measurement over time, 3) how to assess the clinical meaning of changes over time, whether they have prognostic value independent of other measurements, and 4) how they can best be used to engage patients in self-motivation.

One of the most important knowledge gaps in cirrhosis care is our limited understanding of the time frame associated with the efficacy of interventions. We lack information on whether recovery of sarcopenia with nutritional management in combination with an exercise program is sustainable, and how an improvement in muscle mass might be associated with improvement in clinical outcomes.

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Table 1. Quantitative Tools Evaluating Sarcopenia in Cirrhosis

Modalities	Experience	Advantages/Limitations
Dual-energy X-	• 36-month mortality in 144	Advantages:
ray	men with cirrhosis (54)	• Safe, inexpensive and readily available
absorptiometry		Reproducible
(DEXA)	Concordance between various	Low radiation exposure
	muscle mass assessment	Limitations:
	techniques in 59 patients with	• Failure to differentiate water from muscle, therefore
	cirrhosis listed for LT (55)	affected by lower limb edema
		• Weak concordance between DEXA and CT in
		identification of sarcopenia in cirrhosis
Bioelectrical	Prognosis of sarcopenia in	Advantages:
impedance	161 patients with cirrhosis	• Safe, rapid, easy, inexpensive
analysis (BIA)	(56)	Limitations:
		• Affected by fluid retention, diuretic use, liquid and
		food intake before the test, physical activity, and
		BMI
Ultrasound	• Development of sarcopenia	Advantage:
(thigh muscle	model in 159 patients	• Safe, easy, inexpensive
thickness)	evaluated for LT (57)	No radiation exposure
		• High intraobserver and interobserver reliability
		Limitations:
		Reproducibility is unknown
Mid-arm	Concordance between various	Advantage:
muscle	muscle mass assessment	• Safe, inexpensive and readily available
	i	1

circumference	techniques in 59 patients with	Limitations:
	 cirrhosis listed for LT (55) Nutritional assessment by Royal Free Hospital Global Assessment (RFH-GA) in 232 patients with cirrhosis listed for LT (23) 	 Low intraobserver and interobserver reliability Affected by subcutaneous adipose tissue loss Weak correlation with CT determined muscle mass
CT/MRIOSOUS	 MRI-based muscle assessment in 166 patients with decompensated cirrhosis treated with transjugular intrahepatic portosystemic shunt (TIPS) (58) Development of sarcopenia model in 159 patients evaluated for LT (57) 	 Advantage: No radiation exposure (MRI) Fast, accurate Ability to differentiate 3 main body compartments, i.e., muscle, visceral and subcutaneous adipose tissue Ability to identify muscle radiodensity to determine ectopic fat accumulation in muscle Appears not to be affected by the presence of ascites or edema Reduction in price and radiation exposure to only 2.6 millisieverts by single slice CT (45) Limitations: Cost High ionizing radiation exposure makes whole body CT scan unsuitable for longitudinal assessments

Table 2. Summary of studies investigating the CT-determined low muscle mass in patients with ESLD

Author/	Study	Sarcopenia	Muscle Included	Frequency
Year	Population /	Definition		of Sarcopenia
	Center			
	(Single,			
	Multi)			
Englesbe	163 LT	Lowest TPA quartile:	Total psoas areas	25%
et al.,	recipients/			

2010(1)	Cinala	TPA < 1420 mm ² at the		
2010 (1)	Single			
		level of the fourth lumbar		
		vertebra (L4)		
Montano-	112 patients	Mortality associated SMI	Total cross-sectional	50% Males
Loza et	with	cut-offs in cancer* (60)	areas of psoas, erector	18% Females
al.,	cirrhosis		spinae, quadratus	
2012 (59)	evaluated		lumborum, transversus	
	for LT/		abdominis, external and	
	Single		internal obliques, and	
			rectus abdominis	
			(Normalized to height)	
Tandon et	142 patients	Mortality associated SMI	Total cross-sectional	54% Males
al.,	listed for	cut-offs in cancer [*] (60)	areas of psoas, erector	21% Females
2012 (61)	LT/		spinae, quadratus	
	Single		lumborum, transversus	
	$\mathbf{\nabla}$		abdominis, external and	
			internal obliques, and	
			rectus abdominis	
			(Normalized to height)	
Meza-	116 patients	Mortality associated SMI	Total cross-sectional	31% Males
Junco et	with HCC	cut-offs in 1475 patients	areas of psoas, erector	28% Females
al.,	evaluate for	with solid tumors ^{\pm} (63)	spinae, quadratus	
2013(62)	LT/		lumborum, transversus	
	Single		abdominis, external and	
			internal obliques, and	
			rectus abdominis	
-	5		(Normalized to height)	
Krell et	207 LT	Lowest TPA	Total psoas areas	33%
al.,	recipients/	Tertile:		
2013 (16)	Single			
		TPA <1499 mm ² for men		
		and <954 mm ² for women		
DiMartini	338 LT	Mortality associated SMI	Total cross-sectional	76% Males

et al.,	recipients/	cut-offs in cancer* (60)	areas of psoas, erector	51% Females
2013 (14)	Single		spinae, quadratus	51701 emaies
2013 (14)	Single			
			lumborum, transversus	
_			abdominis, external and	
1 7			internal obliques, and	
			rectus abdominis	
			(Normalized to height)	
Masuda	204 patients	PMA	Sum of the areas of the 2	58% Males
et al.,	undergoing	below the 5th percentile	psoas	36% Females
2014 (15)	LT/	for each sex:		
	Single			
		PMA <800 cm ² for men		
-		and <380 cm ² for women		
Durand et	562 patients	Optimal cut-offs of	Right psoas muscle	NA
al.,	listed for	TPMT/height to		
2014 (12)	LT/	discriminate waiting list		
	Single	mortality		
		Transversal psoas muscle		
		thickness [TPMT/height		
		(mm/m)] at the level of		
		umbilicus≤ 16.8 mm/m		
Yadav et	213 patients	Mortality associated SMI	Total cross-sectional	28% Males
al., 2015	listed for	cut-offs in cancer* (60)	areas of psoas, erector	13% Females
(4)	LT/		spinae, quadratus	
	Single		lumborum, transversus	
			abdominis, external and	
			internal obliques, and	
-			rectus abdominis	
	1		(Normalized to height)	
Carey et	396 patients	Optimal cut-offs of SMI	Total cross-sectional	50% Males
al.,	listed for	to discriminate waiting	areas of psoas, erector	33% Females
2017 (22)	LT/Multi	list mortality:	spinae, quadratus	
			lumborum, transversus	

		SMI $<39 \text{ cm}^2/\text{m}^2$ for	abdominis, external and	
		women and $<50 \text{ cm}^2/\text{m}^2$	internal obliques, and	
		for men	rectus abdominis	
_	_		(Normalized to height)	
Tandon et	159 patients	Mortality associated SMI	Total cross-sectional	57% Males
al.,	evaluated	cut-offs in cancer* (60)	areas of psoas, erector	25% Females
2016 (57)	for LT/		spinae, quadratus	
1 7	Single		lumborum, transversus	
			abdominis, external and	
			internal obliques, and	
			rectus abdominis	
			(Normalized to height)	
Van Vugt	585 patients	Mortality associated SMI	Total cross-sectional	42% Males
et al.,	listed for	cut-offs in 1475 patients	areas of psoas, erector	47% Females
2017 (64)	LT/Multi	with solid tumors ^{\pm} (63)	spinae, quadratus	
			lumborum, transversus	
	U		abdominis, external and	
			internal obliques, and	
			rectus abdominis	
			(Normalized to height)	
Van Vugt	224 patients	Lowest sex-specific	Total cross-sectional	25% Males
et al.,	listed for	quartile of SMI:	areas of psoas, erector	24% Females
2018 (17)	LT/		spinae, quadratus	
	Single	L3 SMI <44.1 for men	lumborum, transversus	
(and <37.9 for women	abdominis, external and	
	_		internal obliques, and	
			rectus abdominis	
	5		(Normalized to height)	
			- /	

Abbreviations: PMA, Psoas Muscle Area, PMI, Psoas Muscle Index, SMI, Skeletal Muscle Index; TPA, Total Psoas Area; TPMT, Transversal Psoas Muscle Thickness

*L3 SMI \leq 38.5 cm²/m² for women and \leq 52.4 cm²/m² for men

^{*}Defined as L3 SMI <=41 cm²/m² for women and <=53 cm²/m² for men with body mass index (BMI) >=25 and <=43 cm²/m² in patients with BMI<25.

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Table 3. Pharmacotherapy for Cirrhotic Sarcopenia

Author/	Intervention	Typical Dosing	Comment
Year			

Corey al., 2014	Cholecalciferol	2000 IU/d	Deficiency common in
(41)			cirrhosis
Davuluri et al.,	Leucine	7.5 g/d typically in divided	Included in many
2016 (43)		doses with additional amino	nutritional supplements
Tsien et al., 2015		acids	
(44)			
Holecek et al.,	2-hydroxymethyl	1 g TID	Metabolite of leucine
2017 (65)	butyrate (HMB)		
			Nutritional supplement
			with anti-catabolic
			action
Sinclair et al.,	Testosterone in	Testosterone undecanoate	Gel preferred for
2016 (45)	androgen deficient	1000 mg IM, schedule per	sustained physiologic
	men	RCT; or transdermal gel 50	levels, concerns for
		mg/d (66)	thrombosis and prostate
			cancer
Ohara et al., 2018	L-Carnitine	1000 mg/d or BID	Essential nutrient for
(46)			fatty acid metabolism
			One fourth is
			synthesized in the
			kidney and liver
	I	1	

Author/	-	Main	Dietary	Skeletal	N of s	ubjects	Primary	Skeletal Muscle
Year	Design	Exercise Intervention	Intervention	Muscle Assessment	Active	Control	aim(s)	Mass Outcomes
Konishi <i>et al</i> , 2011(67)	ONCT	Walking (6 months)	Optimal kCal	BIA lean mass (muscle weight)	3	NA	HOMA-IR change	22.5 kg (before) vs. 22.8 kg (after)
Patullo <i>et al</i> , 2013 (68)	ONCT	Walking (24 weeks)	Optimal kCal/protein	Handgrip strength	7	NA	HOMA-IR change	45 lbs (before) vs. 44 lbs (after)
Roman <i>et al</i> , 2014 (69)	Open RCT	Cycle ergometry (12 weeks)	Leucine	Arm and thigh circumference s	10	10	6MWT, 2MST, muscle mass, and HRQoL	MAMC $\downarrow 0.5 \text{ cm}$ in active & $\uparrow 0.6$ cm in control; TC $\uparrow 5 \text{ cm in active}^{\parallel}$ & $\downarrow 1 \text{ cm in}$ control
Zenith <i>et al</i> , 2014 (70)	Open RCT	Cycle ergometry (8 weeks)	Optimal kCal/protein	Thigh US & circumference	9	10	Peak VO ₂ , muscle mass and HRQoL	Thigh US [*] ↑ 0.05 cm/m ² in active [¶] & no change in control; TC ↑ 1.2 cm in active [¶] & 0.2 cm in control
Debette- Gratien <i>et</i> <i>al</i> , 2015 (71)		Cycle ergometry (12 weeks)	NA	Quadriceps Strength	13	NA	Acceptabilit y of exercise program	30 kg (before) to 37 kg (after)¶
Macias- Rodriguez <i>et al</i> , 2016 (72)	Open RCT	Cycle ergometry (14 weeks)	Optimal kCal/protein	BIA phase angle	14	15	HRQoL and HVPG	Phase angle ↑ 0.3 in active¶ & no change in control

Table 4. Exercise clinical trials investigating the effect on muscle mass or function

Roman <i>et al</i> , 2016 (73) Berzigotti	Open RCT	Cycle ergometry (12 weeks) Aerobic &	NA kCal	DXA, Thigh circumference	15	10	Peak VO ₂ , fat/lean body mass, and risk of falls HVPG and	Lean mass \uparrow 1.05 kg in active & \downarrow 0.05 kg in control 56.7 kg (before)
<i>et al</i> , 2017 (74)	ONCT	resistance (16 weeks)	reduction	BIA lean mass	60	NA	body weight	vs. 55.6 kg (after)
Nishida <i>et al</i> , 2017(75)	ONCT	Bench step- ups (12 months)	BCAA	Intramuscular adipose content	9	NA	Anaerobic threshold, fat in liver and muscle, and glycemic control	-0.47 (before) vs 0.42 (after)
Hiraoka <i>et al</i> , 2017(76)	ONCT	Walking (3 months)	BCAA (as late night snack)	BIA-SMI, handgrip and leg strength	33	NA	Muscle volume and function	 ↑ 13% in muscle volume[¶], ↑ 5% in leg strength[¶], and ↑ 10% in handgrip[¶]
Kruger <i>et al</i> , 2018 (77)	Open RCT	Cycle ergometry (8 weeks)	Optimal kcal/protein	Thigh US & circumference	20	20	Peak VO ₂	Thigh US ^{**} ↑ 0.06 cm/m ² in active¶ & 0.06 in control; TC ↑ 1.8 cm in active¶ & 0.6 cm in control

[¶]Denotes comparison was statistically significant

*Average feather index showed was significant in active group, whereas average compression index was not.

**Average feather index showed a p=0.05 in active group, whereas average compression index was not significant.

6MWT, six-minute walk test; **2MST**, 2-min step tests; **BCAA**, branched-chain amino acids; **BIA**, bioelectrical impedance analysis; **DXA**, dual energy X-ray absorptiometry; **HOMA-IR**, Homeostatic Model Assessment of Insulin Resistance, **HRQoL**, health-related quality of life; **HVPG**, hepatic vein

pressure gradient; **kCal**, kilocalories (nutritional requirement); **MAMC**, mid-arm muscle circumference; **NA**, not available; **ONCT**, open non-controlled clinical trial; **RCT**, randomized clinical trial; **SMI**, skeletal muscle index; **TC**, lower thigh circumference; **US**, ultrasound

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Table 5. Summary of Published Literature on Sarcopenia in Pediatric Liver Disease

Article	Design	Study Population	Definition of	Key Findings/Clinical Outcomes
			Sarcopenia	

Mager et al, 2018 (78)	Retrospective	N=41 children post-LT (age 0.5 – 17 years)	DEXA – to measure appendicular skeletal muscle mass (SMM) z- score <-2	 SMM z-score <-2 in 41% of cohort, persisting up to 8 yrs post LT Sarcopenia associated with increased hospital duration and ventilator dependency (total PICU stay), higher readmission rates/LOS readmission, and younger age, female sex, impaired catch- up growth at 1-year post LT
Lurz et al, 2018 (79)	Retrospective	N=23 children with ESLD with clinically indicated CT Control: 2:1 age and sex matched healthy controls (trauma victims with CT)	CT - PMSA at L3/4 and L4/5	 Children with ESLD have a smaller PMSA than healthy controls. The PMSA is independent of anthropometric markers in children with ESLD.
Mangus et al. 2017 (48)	Retrospective	N=81 subjects with organ failure – ESLD (35), ESRD (20), IF (26) Control (39)	CT - PMSA at L2/3	Reduction in muscle mass (ESLD 23%, ESRD 19%, IF 24%) vs. healthy controls Serum total protein and albumin, and BMI, fail to fully characterize malnutrition in chronically ill children

CT, computed tomography; **DEXA**, dual energy X-ray absorptiometry; **ESLD**, end stage liver disease; **ESRD**, end stage renal disease; **IF**, intestine failure; **LOS**, length of stay; **LT**, liver transplantation; **PICU**, Pediatric Intensive Care Unit; **PMSA**, psoas muscle surface area

Figure Legends

Figure 1. Total Muscle Area Quantification at the Level of 3^{rd.} Lumbar Vertebra using Abdominal CT Images from Two Male Patients with Cirrhosis.

Figure 1 A and B, respectively, present a patient who had low SMI (46 cm^2/m^2) and high SMI (60 cm^2/m^2) as indicated by the red shading.

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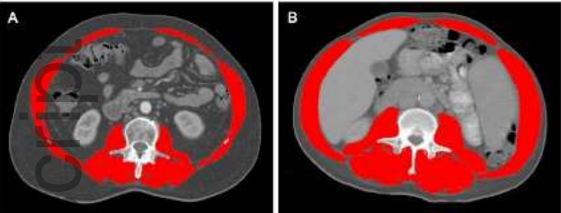
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Figure 1.



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