UC Irvine UC Irvine Previously Published Works

Title

Subjective Global Assessment in chronic kidney disease: A review

Permalink

https://escholarship.org/uc/item/6m6472g0

Journal

Journal of Renal Nutrition, 14(4)

ISSN

1051-2276

Authors

Steiber, Alison L Kalantar-Zadeh, Kamyar Secker, Donna <u>et al.</u>

Publication Date

2004-10-01

DOI

10.1053/j.jrn.2004.08.004

Copyright Information

This work is made available under the terms of a Creative Commons Attribution License, available at https://creativecommons.org/licenses/by/4.0/

Peer reviewed

Subjective Global Assessment in Chronic Kidney Disease: A Review

Alison L. Steiber, PhD, RD, LD,* Kamyar Kalantar-Zadeh, MD, PhD, MPH,† Donna Secker, MS, RD,‡ Maureen McCarthy, MPH, RD, CSR, LD,§ Ashwini Sehgal, MD,¶ and Linda McCann, RD, LD

Nutritional assessment of patients with chronic kidney disease is a vital function of health care providers. Subjective Global Assessment (SGA) is a tool that uses 5 components of a medical history (weight change, dietary intake, gastrointestinal symptoms, functional capacity, disease and its relation to nutritional requirements) and 3 components of a brief physical examination (signs of fat and muscle wasting, nutrition-associated alternations in fluid balance) to assess nutritional status. SGA was originally used to predict outcomes in surgical patients; however, its use has gone beyond this function and population. In chronic kidney disease patients, SGA is incorporated into the complete nutritional assessment. Validation of SGA as a screening tool for surgical patients was done by Detsky et al in 1984. Since that time, SGA has been altered by different researchers and clinicians to better meet the needs of the patients they served. Validation of the altered SGA formats has not been thoroughly done. Further work in establishing validity and reliability of each version of SGA in different patient populations should be done to enable clinicians and researchers to properly use this nutritional assessment tool.

C UBJECTIVE GLOBAL ASSESSMENT

 \checkmark (SGA) is a tool used by health care providers to assess nutritional status and aid in the prediction of nutrition-associated clinical outcomes, such as postoperative infections¹ and/or mortality.² The tool has many strengths in the clinical and research setting: it is inexpensive; is rapid to conduct; can be used effectively by providers from different disciplines, such as nursing, dieti-

§McMinnville Kidney Center, McMinnville, OR.

Satellite Healthcare, Redwood City, CA.

© 2004 by the National Kidney Foundation, Inc.

doi:10.1053/j.jrn.2004.08.004

tians, and physicians; and in some studies has been found to be reproducible, valid, and reliable.^{3,4} Because of its strengths, SGA has been recommended by the National Kidney Foundation (NKF) Kidney Disease/Dialysis Outcomes and Quality Initiative (K/DOQI) for use in nutritional assessment in the adult dialysis population.⁵

However, for all its potential, SGA has yet to be thoroughly validated in the maintenance hemodialysis and peritoneal dialysis population. A study recently published disputed the validity and reliability of SGA in hemodialysis patients. Cooper et al⁶ examined SGA ratings between 2 observers and against total body nitrogen. These investigators concluded that SGA can detect the presence of malnutrition but not the degree of malnutrition.⁶ An additional complication in determining the usefulness of SGA in both the clinical and research arenas is the modification of the original tool. In the chronic kidney disease (CKD) literature, a minimum of 5 different SGA tools have been reported, 1,3,7-9 almost none of which have been tested in a large validation study.

^{*}Department of Nutrition, Case Western Reserve University, Cleveland, OH.

[†]Harbor-UCLA, UCLA David Geffen School of Medicine, Torrance, CA.

[‡]Hospital for Sick Children, Department of Clinical Dietetics, Division of Nephrology, Toronto, Ontario, Canada.

[¶]Division of Nephrology, MetroHealth Medical Center, Cleveland, OH.

Address reprint requests to Alison Steiber, PhD, RD, LD, Department of Nutrition, Case Western Reserve University, Dental Building, Room 201, 10900 Euclid Ave, Cleveland, OH 44106.

^{1051-2276/04/1404-0002\$30.00/0}

To address these issues, a Subjective Global Assessment Consensus Conference was organized by the Department of Nutrition of the School of Medicine of Case Western Reserve University and held on November 7 and 8, 2003, in Cleveland, OH. The objectives of the conference were (1) to review the methods, techniques, and tools being used for SGA; (2) to examine the validity of SGA; and (3) to identify how and by whom SGA is being used in clinical practice and research. Attendance at this conference was by invitation only; announcements were placed in the Journal of Renal Nutrition, the American Journal of Kidney Disease, the Journal of the American Dietetic Association, and on an SGA website: http://www-.nephrology.rei.edu/sgahome.htm. The announcements requested applications from people interested in attending and/or presenting at the conference. Thirty individuals (physicians and dietitians) were invited to attend. During the day-and-a-half conference, presentations included original research results, experiences with SGA in clinical practice, and experiences with SGA in education programs for dietetics students. Throughout the conference, attendees participated in roundtable discussions to generate ideas

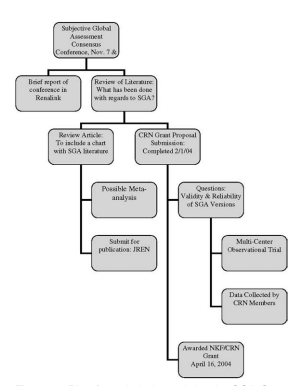


Figure 1. Plan for scholarly work by the SGA Consensus Conference Group.

for validating SGA within the renal population. The consensus of the group of professionals who attended this conference is that further study must be conducted to standardize and validate SGA for the CKD population. Figure 1 outlines the recommended plan for further scholarly work with SGA. This article is one component of that plan, and is intended to review current literature available on SGA and to make recommendations on work to be done.

History of SGA

Detsky et al^{1,10} published the first reports of a nutritional assessment tool, entitled SGA, that used clinical judgment to assess nutritional status in preoperative surgical patients and to predict postoperative infections; SGA had the best sensitivity and specificity for predicting infection after surgery. SGA was quickly used in other populations such as elderly patients, ^{11–13} patients with cancer¹⁴ or liver transplants, ¹⁵ and adult patients undergoing maintenance dialysis.^{2,3,6,16} The original SGA form (Fig 2) as reported by Detsky et al¹ had clinicians score 5 components of a medical history (ie, weight change, dietary intake, gastrointestinal symptoms, functional capacity, disease and its relation to nutritional requirements) and 3 components of a brief physical examination (ie, signs of fat and muscle wasting, nutrition-associated alternations in fluid balance). The patient is then assigned a rating of well nourished (A), moderately undernourished (B), or severely undernourished (C) by subjective consideration of the data collected in the 8 areas, without adhering to a rigid scoring system. From this original form, the tool has been modified by many others in an attempt to increase its predictive value and reproducibility.^{2,7,8} Hirsch et al validated SGA in 175 gastroenterology patients in 1990. That study found significant differences between well-nourished and moderately or severely undernourished patients in serum albumin, weight, midarm muscle circumference (MAMC), and triceps skinfold measurements.¹⁷

The first validation study in CKD patients occurred in 1993¹⁶ with continuous ambulatory peritoneal dialysis (CAPD) patients. SGA was performed on 23 CAPD and 36 hemodialysis patients, and significant correlations were seen between the subjects' SGA ratings and values for serum albumin, bioelectrical impedance,

Patient Name:			_Patier	Patient ID:			_Date: SGA Score			
Part 1: Medical History 1. Weight Change								1 22		
					Lene			Α	В	С
А.		all change in past 6			kgs.					
	mont	20-20-20-20-20-20-20-20-20-20-20-20-20-2	50/					_		
В.			< 5%	IOSS						
	gain									
	10%	5-								
	10%									
	10%	>								
C		nge in past 2 weeks:	increa	50				-		
0.	Unai	ige in past 2 weeks.	increa	30						
	20 V		no cha	ande						
			no one	inge						
			decrea	ase						
			000100							
2. Dietary In	take									
		all change:no								
	chan									
									6	
	-	change								
В.	Dura	tion:								
		weeks								
C.	Туре	of change:								
	-	suboptimal solid			full li	quid				
	diet				diet					
		hypocaloric liquid	5		starv	ation/				
3. Gastroint	estina	al (persisting	for >2	weeks)					
Symptoms										
nonenausea		diarrhe	ea		anorexia					
vomiting										
				n				-		
		airment (nutritional	ly relate	ed)				-		
А.	Over	all impairment:			none					
						erate				
P	0	in and Quarter			seve					
В.	Char	nge in past 2 weeks:				oved				
		3				hange			-	
					regre	essed		-		
Part 2. Phys	sical F	Examination				SGA	Score			
Fait 2. Filys				Norr	nal	Mild	Moder	ato	Sev	oro
5. Evidence	of	Loss of subcutaneous fat		NON	IIai	wind	Model	ale	Sev	ere
5. Evidence	01.	Muscle wasting			ò					
		Edema								
		Ascites (hemo only)								
Part 3. SGA	Ratin	ng (check one)			1.1					
A. Well-N			elv Mal	nourist	ned	C. S	Severely Ma	alnour	ished	
	5 anot		,ai				212.3.y .w.			

Figure 2. A, B, and C original SGA.

MAMC, percent body fat, and normalized protein catabolic rate. This study's SGA methodology was used in the next major study in Canada and the United States (CANUSA) in the CKD population. CANUSA was a multicenter study conducted in Canada and the United States that investigated mortality and nutritional status in 680 patients on peritoneal dialysis.¹² This study changed Detsky's A, B, C method of rating SGA to a 7-point scale (Fig 3). The components assessed remained the same, but the rating scale was expanded. Using survival analysis, the relative risk of death was increased with worsening nutritional status as defined by SGA and loss of lean body mass.² A major outcome of the CANUSA study was that a 1-unit decrease in SGA equaled a 25% increase in mortality for CAPD patients. The 7-point rating scale has been pilot tested by Visser et al³ and Jones et al.⁴ The cross-sectional study by Visser et al³ on 13 hemodialysis and 9 peritoneal dialysis patients showed that SGA was positively correlated with body mass index (BMI), percent body fat, and MAMC. In a recently published article by Jones et al,⁴ both the A, B, C (3-point) scale and the 7-point scale SGA forms were conducted with 72 hemodialysis patients. Statistical differences were found between SGA scores (both A, B, C and 7-point scales) for MAMC and serum creatinine.⁴ The A, B, C scale was also statistically different between A and B groups with the serum C-reactive protein concentration.³

Kalantar-Zadeh et al,⁷ Stenvinkel et al,⁹ and Pifer et al⁸ have each studied different modified versions of SGA in samples ranging from 41 to 7,719 patients. Modifications in the rating scale (ie, from 7 points to 4^9 or 5^7 points) and the direction of data collection (ie, from prospective to retrospective⁸) have been made.

In 1999, Kalanter-Zadeh et al⁷ presented another version of the SGA that was originally referred to as modified quantitative SGA and in subsequent publications as the Dialysis Malnutrition Score (DMS).^{18,19} This fully quantitative version of SGA used the 7 original SGA components and created a quantitative scoring system. The scoring was a 5-point scale with 1 as normal and 5 as very severe malnutrition (Fig 4). The final score was the total sum of all 7 components. Each component was rated on a scale of 1 to 5 with a possible total range from 7 to 35. This method of SGA scoring produced high correlations with objective nutritional indicators such as total iron-binding capacity (TIBC) (r = 0 to 0.77) and MAMC (r = 0 to 0.66) and moderate correlations with serum albumin, BMI, bicep skinfold, age, and years on dialysis.⁷

The Malnutrition-Inflammation Score (MIS), developed by Kalantar-Zadeh, is a recently introduced, fully quantitative tool that is based on the 7 original SGA components and also includes 3 additional items (BMI and serum concentrations of albumin and serum TIBC).^{18,20} Each MIS component has 4 levels of severity from 0 (normal) to 3 (very severe). The sum of all 10 MIS components ranges from 0 to 30, denoting the increasing degree of severity (Fig 5). In a 2001 prospective study on 83 hemodialysis patients, MIS was compared with conventional SGA, its fully quantitative version (DMS), anthropometry, near-infrared measured body fat percentage, laboratory measures including serum C-reactive protein (CRP), and 12-month prospective hospitalization and mortality rates.¹⁸ MIS had significant correlations with prospective hospitalization and mortality as well as measures of nutrition, inflammation, and anemia in dialysis patients. The correlations were higher for MIS than either the conventional SGA or DMS with individual laboratory values as a predictor of outcome. In a 2004 recent multicenter study by the same group of investigators, the mortality and hospitalization predictability of the MIS was assessed in 378 hemodialysis patients; MIS was found to be comparable with serum CRP and serum interleukin-6.20 The MIS is currently being used in the multicenter Nutritional and Inflammatory Evaluation in Dialysis study (www.NIEDstudy.org).^{21,22}

In 1999, Stenvinkel et al⁹ published another version of the SGA. Although these researchers cited Detsky et al and Baker et al in their methods sections, Stenvinkel et al changed the scoring from the original A, B, C scale to a 4-point scale using 1 as normal nutritional status and 4 as severe malnutrition.⁹ Data on 109 adults with chronic kidney failure were analyzed by creating a bivariate variable with SGA scores 2 to 4 as one group and an SGA score of 1 as another group. In this manner they found those with scores between 2 and 4 were older, more frequently had a history

SUBJECTIVE GLOBAL ASSESSMENT RATING FORM						
Patient Name: ID #: Date:						
HISTORY						
WEIGHT/WEIGHT CHANGE: (Included in K/DOQI SGA) 1. Baseline Wt:	Rate 1-7					
DIETARY INTAKE No Change(Adequate) No Change(Inadequate) 1. Change: Sub optimal Intake: Protein Kcal Duration Full Liquid: Hypocaloric Liquid Starvation						
GASTROINTESTINAL SYMPTOMS (Included in K/DOQI SGA-anorexia or causes of anorexia) Symptom: Frequency:* Duration:+ None						
FUNCTIONAL CAPACITY Description Duration: No Dysfunction	b					
DISEASE STATE/COMORBIDITIES AS RELATED TO NUTRITIONAL NEEDS Primary Diagnosis Comorbidities Normal requirements Increased requirements Decreased requirements Acute Metabolic Stress: None Low Moderate PHYSICAL EXAM						
Loss of subcutaneous fat (Below eye, triceps,Some areasAll areasbiceps, chest) (Included in K/DOOI SGA)Muscle wasting (Temple, clavicle, scapula, ribs,Some areasAll areasquadriceps, calf, knee, interosseous (Included in K/DOOI SGA)Edema (Related to undernutrition/use to evaluate weight change)OVERALL SGA RATING Very mild risk to well-nourished=6 or 7 most categories or significant, continued improvement. Mild-moderate = 3, 4, or 5 ratings. No clear sign of normal status or severe malnutrition. Severely Malnourished = 1 or 2 ratings in most categories/significant physical signs of malnutrition.						

Figure 3. The 7-point scale SGA form.

of tobacco use, and had significantly lower BMI, serum creatinine, serum albumin, urine urea, and lean body mass (measured by dual-energy x-ray absorptiometry).⁹

The Dialysis Outcomes and Practice Patterns Study (DOPPS) study created m-SGA that was graded retrospectively using a patient interview. The score was based on the caregiver's ratings

1- Weight ch	ang	e (overall change in	۱p	oast 6 months)					
1		2	Π	3		4		5	
no weight chang gain	e or	minor Wt loss (<5%)		Wt loss 5 to 10 %		Wt loss 10 to 15%		Wt loss > 15% in	
2- Dietary int	ake		Ĭ						
1		2		3		4		5	
no change		sub-optimal solid diet		full liquid diet or moderate overall decrease		hypo-caloric liquid		starvation	
3- Gastrointe	estin	al symptoms							
1	2			3		4		5	
no symptoms		nausea		vomiting or moderate GI symptoms		diarrhea		severe anorexia	
4- Functiona	l ca	oacity (nutritional	ly	related functional i	impai	rment)			
1		2		3		4		5	
none (improved)		difficulty with ambulation	Ī	difficulty with norr activity	mal	light activity		bed/chair-ridden with no or little activity	
5- Co-morbic	lity								
1		2		3		4		5	
dialysis<12 months and healthy otherwise		dialysis 1-2 yrs or mild comorbidity						very severe multiple comorbidit	
(B) Physic	al E	xam:	_				_		
1- Decreased	l fat	stores or loss	0	f subcutaneou	is fa	t (below eyes, triceps,	bic	eps, chest)	
1	Π	2		3		4		5	
none (no change)				moderate				severe	
2- Signs of m	nusc	le wasting (tem	pl	e, clavicle, scapula	, ribs,	quadriceps, knee, inte	ros	seous)	
1		2		3		4		5	
			Ń	moderate				severe	

Figure 4. The fully quantitative version of the SGA, also known as modified SGA or DMS. Five scale parameters are used, and the values are summed. A value of 7 is normal, and 35 is the most severe malnutrition.

relative to weight loss, visual somatic store loss, appetite, nausea and vomiting, energy level, and disease burden. The rating for m-SGA is normal,

moderate (any 3 areas rated as a moderate or severe level), or severe (at least 3 areas at severe level). Those patients who rated a severe m-SGA

		17.		
(A) Patients' related medic	al history:			
1- Change in end dialysis	dry weight (overall change i	in past 3-6 months):		
0	1	2	3	
No decrease in dry weight or weight loss <0.5 kg	Minor weight loss (≥0.5 kg but <1 kg)	Weight loss more than one kg but <5%	Weight loss >5%	
2- Dietary intake:				
0	1	2	3	
Good appetite and no deterioration of the dietary intake pattern	Somewhat sub-optimal solid diet intake	Moderate overall decrease to full liquid diet	Hypo-caloric liquid to starvation	
3- Gastrointestinal (GI) sy	nptoms:		•	
0	1	2	3	
No symptoms with good appetite	Mild symptoms, poor appetite or nauseated occasionally	Occasional vomiting or moderate GI symptoms	Frequent diarrhea or vomiting or severe anorexia	
4- Functional capacity (nu	tritionally related functional			
0	1	2	3	
Normal to improved functional capacity, feeling fine	Occasional difficulty with baseline ambulation, or feeling tired frequently	Difficulty with otherwise independent activities (e.g. going to bathroom)	Bed/chair-ridden, or little to no physical activity	
	number of years on Dialysi			
0	1	2	3	
On dialysis less than one year and healthy otherwise	Dialyzed for 1-4 years, or mild co-morbidity (excluding MCC*)	Dialyzed >4 years, or moderate co-morbidity (including one MCC*)	Any severe, multiple co- morbidity (2 or more MCC*)	
(B) Physical Exam (accord	ling to SGA criteria):			
6- Decreased fat stores or	loss of subcutaneous fat (b	elow eyes, triceps, biceps, ches	t):	
0	1	2	3	
Normal (no change)	mild	moderate	Severe	
	g (temple, clavicle, scapula, rit	os, quadriceps, knee, interosseo	us):	
0	1	2	3	
Normal (no change)	mild	moderate	Severe	
(C) Body mass index:		·		
8- Body mass index: BMI =	= Wt(kg) / Ht ² (m)			
0	1	2	3	
BMI≥20 kg/m ²	BMI: 18-19.99 kg/m ²	BMI: 16-17.99 kg/m ²	BMI<16 kg/m ²	
(D) Laboratory Parameters	5:			
9- Serum albumin:			- 7-	
0	1	2	3	
Albumin≥ 4.0 g/dL	Albumin: 3.5-3.9 g/dL	Albumin: 3.0-3.4 g/dL	Albumin: <3.0 g/dL	
10- Serum TIBC (total Iron				
0	1	2	3	
TIBC> 250 mg/dL	TIBC: 200-249 mg/dL	TIBC: 150-199 mg/dL	TIBC: <150 mg/dL	
	above 10 components			

Figure 5. MIS. *Major comorbid conditions include congestive heart failure class III or IV, full-blown AIDS, severe coronary artery disease, moderate to severe chronic obstructive pulmonary disease, major neurologic sequelae, and metastatic malignancies or s/p recent chemotherapy. **&** Suggested equivalent increments for serum transferrin are >200 (0), 170 to 200 (1), 140 to 170 (2), and <140 mg/dL.

level had a relative risk of 1.33 for mortality compared with those with a moderate or normal rating,⁸ which was statistically significant.

Although each of the versions has strengths, their lack of uniformity makes it difficult both to compare research results on nutritional status from one study to the next and to provide consistent methodology guidance for clinicians wishing to use this tool. Currently the NKF regularly offers training sessions at its Clinical Nephrology meetings to train renal dietitians in the use of the 7-point SGA. No other formal training forum currently exists. Therefore, it is assumed that the majority of renal dietitians currently conducting SGA are using the version recommended by K/DOQI and studied by Visser et al³ and Jones et al.⁴

Current Literature With SGA as a Nutritional Assessment Tool

Table 1 includes studies that used SGA as a method of nutritional status determination for further comparisons against a dependent variable (eg, mortality). From this table it is clear that SGA, using either the A, B, C or the 7-point scale, detects the presence of malnutrition; however, the controversy appears when SGA is correlated with serum albumin. In some studies serum albumin was significantly lower in the SGA malnourished group,^{9,23-25} whereas in others, serum albumin was not significantly different between the normal and the malnourished groups.^{4,26} Serum albumin is one of the most commonly used indicators for malnutrition in the CKD population, and although it is affected by several other factors including inflammation, this inconsistency has raised questions about the validity of SGA. To that end, incorporating serum laboratory markers for malnutrition may be a solution, as done in the MIS.

Studies have shown significant differences between SGA categories for many other nutritionrelated variables, ie, BMI, MAMC, serum prealbumin, TIBC or transferrin, ferritin, insulin-like growth factor 1, phase angle (bioelectrical impedance analysis), percent body fat, lean body mass, comorbidity state (diabetes, cardiovascular disease, etc), c-reactive protein and cytokines, and creatinine clearance.^{4,7,9,24-31}

The risk of mortality has been assessed by CANUSA,² Lawson et al,²⁷ Davies et al,²⁸ Kalantar-Zadeh,^{18,20} and Pifer et al,⁸ with all showing a statistically significant increase in risk or rate of mortality with the presence of malnutrition as determined by SGA.

Interventional trials using kilocalorie and protein supplements, such as in the studies by Caglar et al³² and Steiber et al,³³ have shown varying effects on changes in an individual's pre-SGA and post-SGA rating, depending on the intervention duration. The trial by Caglar et al³² was 6 months, included 85 patients, and used the 7-point scale. They were able to show an improvement in the 7-point SGA over time; however, Steiber et al³³ did not see a significant change in pre-SGA and post-SGA scores over a 3-month period when the A, B, C rating system was used in 22 patients.

Recommendations

A review of the literature indicates that use of SGA as a nutrition assessment tool for CKD patients is growing, in both the clinical and research settings. However, given the variability of published results, SGA cannot be considered a gold standard in nutrition assessment for CKD patients. The validity and reliability of SGA must be proven in a large, multicenter trial with sufficient power to be able to prevent type I and II errors. Additionally, the study's sample must represent the current CKD population. One of the difficulties associated with conducting a study such as this is choosing which version of SGA to test. It may be that different SGA versions are appropriate for different patient disease states, different age stages, or different clinical purposes (eg, screening preoperatively versus full assessment of maintenance hemodialysis patients). Another difficulty is data collection. To get a representative sample, data would need to be collected from all areas of the country in a random manner. This could be done in a way similar to that of Beto et al³⁴ in a nationally collaborative research project through the National Kidney Foundation's Council on Renal Nutrition (CRN). Using this model, registered dietitians from local CRN groups throughout the United States could randomly collect data on patients in their dialysis centers.

Many of the studies reviewed collapsed the SGA scores into 2 groups (normal and malnourished) for analyses. For instance, Julien et al, 30 Lawson et al, 27 Abdullah et al, 35 and Jones et al, 26 used the A, B, C rating system and all dichotomized the final results by merging the B and C groups together for comparison against the A-rated group. Davies et al²⁸ used the 7-point scale and collapsed it into 6 to 7, 3 to 5, and 1 to 2 for analysis, and then grouped those with a 5 or less into a "malnourished group" and compared those patients with the 6 to 7 group. This method of analysis substantiates the conclusion of Cooper et al,6 who found that SGA detects the presence of malnutrition but not the degree. It is possible that the need for the collapsed groups in such studies has more to do with inadequately powered studies or analytical tools (eg, logistic regression) than the lack of detectable precision of SGA. When presenting results of SGA in aggregate, it may be useful to show them in both a full and a collapsed or aggregated format. This would highlight any linear relationships as well as show differences between those with and without malnutrition.

First Author	Journal*	Year; Volume (No.): Page Range	Rating Method	Main Comparison Variable(s)	n	Results
Maiorca	Nephrol Dial	1995;10	ABC	Survival	578	No difference in survival
Cianciaruso	Transplant AJKD	1995;26(3)	ABC	Age	487	between SGA groups Older patient ↓ SGA
Maggiore A	Kidney Int	1996;50(6)	ABC	Bioelectrical impedance	131	score SGA ↑ as phase angle ↑, not predictive in patients
Jones CH	Nephrol Dial Transplant	1997	ABC	analysis Nutrition parameters	76	with worst SGA rating LBM, CrCl, BMI, MAMC, handgrip, weight ↓ in B
Abdullah	Miner Electrolyte Metabolism	1997;23(3-6)	ABC	IGF-1, TNF α	20	and C groups B and C groups \downarrow IGF-1 and \uparrow TNF α
Noh H Kalantar-Zadeh K	Perit Dial Int AJKD	1998;18(4) 1998;31(2)	ABC	Mortality Laboratory parameters	106 59	C group has ↓ TIBC
Kalantar-Zadeh K	Nephrol Dial Transplant	1999;14(7): 1732-1738	5-point	Alb, TIBC, anthropometry	41 hemodialysis	Fully quantitative SGA had good correlation with laboratory and anthropometric
Biesenbach G	Nephrol Dial	1999;14(3)	ABC	Diabetic versus Nondiabetic	30	nutritional markers No difference between
Passadakis P	Transplant Adv Perit Dial	1999;15		Bioelectrical impedance analysis	47	SGA groups Correlation between phase angle and SGA
Visser R	Adv Perit Dial	1999;15: 222-225	7-point	BMI, anthropometry, albumin	13 hemodialysis 9 peritoneal dialysis	7-point SGA scale is a valid and reliable tool for assessing nutritional status among end-stage
Davies SJ	Kidney Int	2000;57(4)	7-point		141	renal disease patients
Kalantar-Zadeh K	AJKD	2001;38(6): 1251-1263	4-point, plus 3 new items	CRP, mortality, hospitalization	83 hemodialysis	MIS predicted clinical outcome
Lawson J	JREN	2001;11(1)	ABC	Mortality	87	↑ mortality in B and C groups
Sezer S	Adv Perit Dial	2001;17		Alb	100	Alb ↓ in malnourished patients
Julien J	EDTNA	2001;27(4)	ABC	Alb, prealb	32	Prealb ↑ in A versus B and C groups
Cooper BA	AJKD	2002;40(1): 126-132	ABC	Total body nitrogen	76	and c groups SGA differentiated severely malnourished patients from those with normal nutrition, but was not a reliable predictor of degree of malnutrition
Caglar K	Kidney Int	2002;62	7-point	Time-dependent change	85	SGA ↑ over 6 mo
Bakewell A	Q J Med	2002;95(12)	7-point	Incidence of malnutrition	70	SGA \downarrow over time (NS)
Steiber A	JREN	2003;13(3)	ABC	HD-PNI	22	HD-PNI ↓ in B and C
Kalantar-Zadeh K	Nephrol Dial Transplant	2004;19(6): 1507-1519	4-point, plus 3 new items	CRP, cytokines, mortality, hospitalization	378 hemodialysis	groups (NS) MIS was superior to albumin and was similar to CRP and IL-6 in predicting clinical outcome

Table 1. Studies Using SGA as a Tool in Their Methodology

Abbreviations: SGA, subjective global assessment; LBM, lean body mass; CrCl, creatinine clearance; BMI, body mass index; MAMC, midarm muscle circumference; IGF-1, insulin growth factor-1; TNF α , tumor necrosis factor alpha; TIBC, total iron-binding capacity; alb, serum albumin; prealb, serum prealbumin; HD-PNI, Hemodialysis Prognostic Nutritin Index; MIS, Malnutrition-Inflammation Score; CRP, C-reactive protein NS, not significant. *Medline abbreviations used.

In a large study with sufficient power, SGA may be able to detect differences between all 7/5 points or A, B, and C. Similarly, a continuous score may resolve the issue independent of sample size. Theoretically, with careful methodology and statistical analysis, a large, nationally representative study could be designed to determine the validity and reliability of SGA within the diverse United States CKD population. Until the issue of which form of SGA is best suited to the hemodialysis population is determined, clinicians who are currently using one of the forms of SGA should continue to perform SGA. SGA is without a doubt a useful tool for nutritional assessment. However, as with all of the available tools, it should be used in conjunction with anthropometric, laboratory, and dietary intake measures to form a comprehensive nutritional assessment.

References

1. Detsky AS, McLaughlin JR, Baker JP, et al: What is subjective global assessment of nutritional status? JPEN J Parenter Enteral Nutr 11:8-13, 1987

2. Canada-USA (CANUSA) Peritoneal Dialysis Study Group. Adequacy of dialysis and nutrition in continuous peritoneal dialysis: Association with clinical outcomes. J Am Soc Nephrol 7:198-207, 1996

3. Visser R, Dekker FW, Boeschoten EW, et al: Reliability of the 7-point subjective global assessment scale in assessing nutritional status of dialysis patients. Adv Perit Dial 15:222-225, 1999

4. Jones CH, Wolfenden RC, Wells LM: Is subjective global assessment a reliable measure of nutritional status in hemodialysis? J Ren Nutr 14:26-30, 2004

5. K/DOQI, National Kidney Foundation: Clinical practice guidelines for nutrition in chronic renal failure. Am J Kidney Dis 35:S1-140, 2000 (suppl 2)

6. Cooper BA, Bartlett LH, Aslani A, et al: Validity of subjective global assessment as a nutritional marker in end-stage renal disease. Am J Kidney Dis 40:126-132, 2002

7. Kalantar-Zadeh K, Kleiner M, Dunne E, et al: A modified quantitative subjective global assessment of nutrition for dialysis patients. Nephrol Dial Transplant 14:1732-1738, 1999

8. Pifer TB, McCullough KP, Port FK, et al: Mortality risk in hemodialysis patients and changes in nutritional indicators: DOPPS. Kidney Int 62:2238-2245, 2002

9. Stenvinkel P, Heimburger O, Paultre F, et al: Strong association between malnutrition, inflammation, and atherosclerosis in chronic renal failure. Kidney Int 55:1899-1911, 1999

10. Detsky AS, Baker JP, Mendelson RA, et al: Evaluating the accuracy of nutritional assessment techniques applied to hospitalized patients: Methodology and comparisons. JPEN J Parenter Enteral Nutr 8:153-159, 1984

11. Christensson L, Unossons M, Ek AC: Measurement of perceived health problems as a means of detecting elderly people at risk of malnutrition. J Nutr Health Aging 7:257-262, 2003

12. Christensson L, Unosson M, Ek AC: Evaluation of nutritional assessment techniques in elderly people newly admitted to municipal care. Eur J Clin Nutr 56:810-818, 2002

13. Covinsky KE, Covinsky MH, Palmer RM, et al: Serum albumin concentration and clinical assessments of nutritional status in hospitalized older people: Different sides of different coins? J Am Geriatr Soc 50:631–637, 2002

14. Bauer J, Capra S, Ferguson M: Use of the scored Patient-Generated Subjective Global Assessment (PG-SGA) as a nutrition assessment tool in patients with cancer. Eur J Clin Nutr 56:779-785, 2002

15. Stephenson GR, Moretti EW, El Moalem H, et al: Malnutrition in liver transplant patients: Preoperative subjective global assessment is predictive of outcome after liver transplantation. Transplantation 72:666-670, 2001

16. Enia G, Sicuso C, Alati G, et al: Subjective global assessment of nutrition in dialysis patients. Nephrol Dial Transplant 8:1094-1098, 1993

17. Hirsch S, de Obaldia N, Petermann M, et al: Subjective global assessment of nutritional status: Further validation. Nutrition 7:35-37, 1991

18. Kalantar-Zadeh K, Kopple JD, Block G, et al: A malnutrition-inflammation score is correlated with morbidity and mortality in maintenance hemodialysis patients. Am J Kidney Dis 38:1251-1263, 2001

19. Kalantar-Zadeh K, Rodriguez RA, Humphreys MH: Association between serum ferritin and measures of inflammation, nutrition and iron in haemodialysis patients. Nephrol Dial Transplant 19:141-149, 2004

20. Kalantar-Zadeh K, Kopple JD, Humphreys MH, et al: Comparing outcome predictability of markers of malnutritioninflammation complex syndrome in hemodialysis patients. Nephrol Dial Transplant 2004 (in press)

21. Kalantar-Zadeh K, Block G, Humphreys MH, et al: A low, rather than a high, total plasma homocysteine is an indicator of poor outcome in hemodialysis patients. J Am Soc Nephrol 15:442-453, 2004

22. Kalantar-Zadeh K, Block G, McAllister CJ, et al: Appetite and inflammation, nutrition, anemia and clinical outcome in hemodialysis patients. Am J Clin Nutr 2004 (in press)

23. Sezer S, Ozdemir FN, Akman B, et al: Predictors of serum albumin level in patients receiving continuous ambulatory peritoneal dialysis. Adv Perit Dial 17:210-214, 2001

24. Passadakis P, Sud K, Dutta A, et al: Bioelectrical impedance analysis in the evaluation of the nutritional status of continuous ambulatory peritoneal dialysis patients. Adv Perit Dial 15:147-152, 1999

25. Kalantar-Zadeh K, Kleiner M, Dunne E, et al: Total iron-binding capacity—estimated transferrin correlates with the nutritional subjective global assessment in hemodialysis patients. Am J Kidney Dis 31:263-272, 1998

26. Jones CH, Newstead CG, Will EJ, et al: Assessment of nutritional status in CAPD patients: Serum albumin is not a useful measure. Nephrol Dial Transplant 12:1406-1413, 1997

27. Lawson JA, Lazarus R, Kelly JJ: Prevalence and prognostic significance of malnutrition in chronic renal insufficiency. J Ren Nutr 11:16-22, 2001

28. Davies SJ, Phillips L, Griffiths AM, et al: Analysis of the effects of increasing delivered dialysis treatment to malnourished peritoneal dialysis patients. Kidney Int 57:1743-1754, 2000

29. Biesenbach G, Debska-Slizien A, Zazgornik J: Nutritional status in type 2 diabetic patients requiring haemodialysis. Nephrol Dial Transplant 14:655-658, 1999

30. Julien JP, Combe C, Lasseur C: Subjective global assessment of nutrition a useful diagnostic tool for nurses? EDTNA ERCA J 27:193-196, 2001

31. Qureshi AR, Alvestrand A, Danielsson A, et al: Factors predicting malnutrition in hemodialysis patients: A cross-sectional study. Kidney Int 53:773-782, 1998

32. Caglar K, Fedje L, Dimmitt R, et al: Therapeutic effects of oral nutritional supplementation during hemodialysis. Kidney Int 62:1054-1059, 2002

33. Steiber AL, Handu DJ, Cataline DR, et al: The impact of nutrition intervention on a reliable morbidity and mortality indicator: The hemodialysis-prognostic nutrition index. J Ren Nutr 13:186-190, 2003

34. Beto JA, Bansal VK, Hart J, et al: Hemodialysis prognostic nutrition index as a predictor for morbidity and mortality in hemodialysis patients and its correlation to adequacy of dialysis. Council on Renal Nutrition National Research Question Collaborative Study Group. J Ren Nutr 9:2-8, 1999

35. Abdullah MS, Wild G, Jacob V, et al: Cytokines and the malnutrition of chronic renal failure. Miner Electrolyte Metab 23:237-242, 1997