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Yazan Duwayri, Emory University
CW Hicks, Johns Hopkins University
AK Vavra, Northwestern Feinberg
E Goldsborough, Johns Hopkins University
M Rebuffatti, University of California Los Angeles
J Almeida, University of Miami
M Haurani, Ohio State University
CB Ross, Piedmont Healthcare
SK Shah, University of Florida
PK Shireman, Univ Texas Hlth

Only first 10 authors above; see publication for full author list.

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Caitlin W. Hicks, MD, MS, Ashley K. Vavra, MD, MS, Earl Goldsborough III, BS, Michelle Rebuffatti, BS, Jose Almeida, MD, Yazan M. Duwayri, MD, Mounir Haurani, MD, MPH, Charles B. Ross, MD, Samir K. Shah, MD, MPH, Paula K. Shireman, MD, MS, MBA, Christopher J. Smolock, MD, Jeniann Yi, MD, MCS, Karen Woo, MD, PhD

aDivision of Vascular Surgery and Endovascular Therapy, Department of Surgery, Johns Hopkins University School of Medicine, Baltimore
bDivision of Vascular Surgery, Department of Surgery, Northwestern Feinberg School of Medicine, Chicago
cJohns Hopkins University School of Medicine, Baltimore
dDivision of Vascular Surgery, Department of Surgery, David Geffen School of Medicine at University of California Los Angeles, Los Angeles
eMiami Vein and Division of Vascular and Endovascular Surgery, University of Miami Miller School of Medicine, Miami
fDivision of Vascular Surgery and Endovascular Therapy, Emory University, Atlanta
gDivision of Vascular Diseases and Surgery, The Ohio State University Wexner Medical Center, Columbus
hVascular Center of Excellence, Piedmont Heart and Vascular Institute, Piedmont Healthcare, Atlanta
iDivision of Vascular Surgery, University of Florida, Gainesville
jDivision of Vascular and Endovascular Surgery, Long School of Medicine, University of Texas Health San Antonio
kDepartment of Surgery, South Texas Veterans Health Care System, San Antonio
lDepartment of Vascular Surgery, Sydell and Arnold Miller Family Heart, Vascular, and Thoracic Institute, The Cleveland Clinic, Cleveland
mDivision of Vascular Surgery and Endovascular Therapy, Department of Surgery, University of Colorado, Aurora.

Correspondence: Karen Woo, MD, PhD. Division of Vascular Surgery, Department of Surgery, David Geffen School of Medicine at University of California Los Angeles, 200 UCLA Medical Plaza Ste 526, Los Angeles, CA 90095 (kwoo@mednet.ucla.edu).

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Abstract

A previously published review focused on generic and disease-specific patient-reported outcome measures (PROMs) relevant to vascular surgery but limited to arterial conditions. The objective of this project was to identify all available PROMs relevant to diseases treated by vascular surgeons and to evaluate vascular surgeon perceptions, barriers to widespread implementation, and concerns regarding PROMs. We provide an overview of what a PROM is and how they are developed, and summarize currently available PROMs specific to vascular surgeons. We also report results from a survey of 78 Society for Vascular Surgery members serving on committees within the Policy and Advocacy Council addressing the barriers and facilitators to using PROMs in clinical practice. Finally, we report the qualitative results of two focus groups conducted to assess granular perceptions of PROMs and preparedness of vascular surgeons for widespread implementation of PROMs. These focus groups identified a lack of awareness of existing PROMs, knowledge of how PROMs are developed and validated, and clarity around how PROMs should be used by the clinician as main subthemes for barriers to PROM implementation in clinical practice.

Keywords

Patient-reported outcomes; Patient-reported outcome measures; Vascular surgery

The success of vascular surgery interventions is most commonly judged on objective measures defined by physicians. For example, the Society for Vascular Surgery (SVS) reporting standards for endovascular interventions to treat lower extremity peripheral artery disease (PAD) define key procedural outcomes including technical success, periprocedural complications, sustained hemodynamic improvement, patency, and freedom from repeat interventions. Although objective measures of success are an important component of healthcare, patient perceptions of intervention outcomes are equally important. As a result, there has been increasing interest on the part of the healthcare community in patient-reported outcomes (PROs) for measuring treatment effectiveness and evaluating quality of care.

PROs are defined by the US Food and Drug Administration as “any report of the status of a patient’s (or person’s) health condition, health behavior, or experience with healthcare that comes directly from the patient, without interpretation of the patient’s response by a clinician or anyone else.” PRO measures (PROMs) are the tools that are used to collect PROs and can measure outcomes in a variety of domains including quality of life, mood and physical function among others (Tables I through V provide examples of specific domains). The Centers for Medicare & Medicaid Services (CMS) recently defined PROMs as a high priority, suggesting that, “although patient reports of their health and experience with care are not the only outcomes that should be measured, they certainly are an important component.”

ANATOMY OF A PROM

PROMs often take the form of a questionnaire that can either be completed by the patient on their own, or are administered to the patient by someone else. Each item in the questionnaire is grouped into a “domain,” which represents a general category of assessment...
included in the PROM, such as pain, psychological impact, effect on social activities, and effect on physical activities (Tables I-V). Each PROM has a unique scoring mechanism that is relevant to the topic of the PROM.

**Satisfaction versus health-related quality of life.**

Two broad categories of PROMs are health-related quality of life (HRQOL) measures, which can be general or disease specific; and satisfaction measures, which focus on the patient experience of receiving health care. HRQOL measures assess how a disease and its treatment affect the physical, psychologic, and/or social aspects of life. HRQOL can be measured with objective assessments of functioning or health status (eg, frequency of pain) or more subjective evaluation of health (eg, extent to which pain hinders ability to engage in social activities).

Satisfaction differs from HRQOL in that it is entirely subjective. With respect to health care, patient satisfaction generally refers to the extent that the patient believes that high-quality health care was delivered. Thus, satisfaction could potentially be defined differently by different people and satisfaction could be defined differently by the same person at different times. One of the most well-known satisfaction instruments is the Hospital Consumer Assessment of Healthcare Providers and Systems survey, which was developed by the CMS and the Agency for Healthcare Research and Quality. The Hospital Consumer Assessment of Healthcare Providers and Systems is administered to a random sample of patients discharged from a hospital on a monthly basis and addresses topics including nurse and doctor communication, and the cleanliness and quietness of the hospital environment.

**PROM development.**

The US Food and Drug Administration dictates five steps in the development of a PROM. The first step is to develop an appropriate conceptual model. The specific concept of interest that the PROM aims to assess should be carefully defined along with boundaries for what is going to be assessed. For example, if the objective is to assess pain, what are the components that will be measured (eg, intensity, quality, variability) and what are the components that are outside the scope of the proposed PROM (eg, treatment of the pain, effects of the pain on physical function and/or mental health)? Second, the conceptual framework is adjusted by gathering patient and stakeholder input, often in the form of focus groups and/or individual interviews. In the third step, a draft instrument is developed and assessed by administering the instrument to a diverse group of patients who would be in the target population for the PROM. These patients are interviewed individually after they complete the draft instrument to ensure readability and a uniform understanding of the items, as well a to examine whether additional domains should be included in the assessment.

The last two steps of PROM development are more relevant to HRQOL instruments than satisfaction instruments. The fourth step involves having large numbers (hundreds or thousands) of patients from diverse backgrounds and health circumstances complete the instrument and confirm that it measures what it intends to measure by comparing the responses with objective measures of health. The instrument’s sensitivity to change is also
assessed by determining whether the instrument’s score changes appropriately with changes in the patient’s health status. Finally, the instrument undergoes translation and cultural adaptation and repeat of step four after those changes. The development of a PROM is meant to ensure its application to a broad range of patients regardless of race, sex, and ethnicity, although the extent to which this is true may vary by individual PROMs.

Objectives.

A previously published review focused on generic and disease-specific PROMs relevant to vascular surgery but limited to arterial conditions. The objective of this project was to identify all available PROMs relevant to diseases treated by vascular surgeons and to evaluate vascular surgeon perceptions, barriers to widespread implementation, and concerns regarding PROs.

OVERVIEW OF PROMS IN VASCULAR SURGERY

We performed a comprehensive literature search of PubMed using a series of search terms for PROMs (“patient reported outcomes” AND vascular AND surgery NOT heart NOT breast). Ninety articles were identified, all of which were reviewed in a semi-structured manner by members of the SVS Performance Measures Committee. We identified 30 PROMs specific to vascular surgery disease processes.

PAD-specific PROMs.

We identified 14 PROMs that address PAD (Table I). Seven PROMs are general PAD instruments, and seven are specific to claudication. Of the general PAD instruments, the VascuQoL is widely used in research. The VascuQoL is composed of 25 questions assessing 5 QOL domains (pain, symptoms, activities, social impact, and emotional impact of PAD). It is brief and detects postintervention change in PAD severity better than generic HRQOL instruments. However, the VascuQoL is better used as an assessment of global QOL, as opposed to an instrument used for assessing functional status. The VascuQoL-6 is an abridged version of the VascuQoL and can be completed very quickly; however, its brevity calls into question the comprehensiveness of the instrument.

The Peripheral Artery Disease Quality of Life Questionnaire was validated using other established instruments that measured functional elements of PAD, including community-based walking ability in the Walking Impairment Questionnaire and health status in the 36-Item Short Form Health Survey (SF-36). The tool consists of 38 questions assessing 5 QOL domains (social relationships and interactions, self-concept and feelings, symptoms and limitations in physical functioning, fear and uncertainty, and positive adaptation) and can be completed in less than 10 minutes. However, the Peripheral Artery Disease Quality of Life Questionnaire has limited testing in some populations (Table I). The Peripheral Artery Questionnaire is capable of assessing PAD-specific QOL, overall QOL, and treatment impacts on patient QOL. The questionnaire is comprised of 20 questions assessing 7 PAD-specific QOL domains (assessment of the most symptomatic leg, change in symptoms, physical limitation, social function, treatment satisfaction, and overall QOL); however, it is not the most comprehensive PAD-specific QOL instrument.
Occlusive Disease 86-Item Questionnaire has been used comprehensively to evaluate the QOL effects of combinations of pharmacologic and exercise on patients with PAD; however, the questionnaire takes approximately 20 minutes to complete, limiting its widespread use.\textsuperscript{12,13,16-20,94}

The Patient Benefit Index for Peripheral Arterial Disease (PBI-PAD) evaluates the severity of impairment of PAD-specific symptoms and sequelae\textsuperscript{24,25} The PBI-PAD consists of two questionnaires—the Patient Needs Questionnaire and the Patient Benefit Questionnaire—with the former administered before treatment and the latter administered 3 months after treatment.\textsuperscript{24} Although the PBI-PAD enables clinicians to calculate preprocedure and postprocedure differences, mitigating the likelihood of response shift and recall bias, the questionnaire is administered over a time span of a few months, creating challenges with feasibility.\textsuperscript{24}

The FLeQKI was originally used to measure QOL in patients with critical limb ischemia.\textsuperscript{10,11} Its validity and reliability are comparable to that of the SF-36; however, it is only validated in German.\textsuperscript{10,94}

Two claudication-specific instruments are used to identify patients with claudication: the World Health Organization (WHO)/Rose Questionnaire and the Edinburgh Claudication Questionnaire (ECQ). The WHO/Rose tool was developed in 1962 to assess patients with intermittent claudication and readapted in 1977 to satisfy requirements to become an official PRO.\textsuperscript{8,36} The questionnaire uses binomial scoring to identify individuals as either “claudicant” or “nonclaudicant.” Further adaptation allowed for differentiation of possible claudicants into grade 1 versus grade 2 claudication exhibiting increased sensitivity compared with its preadapted version.\textsuperscript{37} Although the WHO/Rose questionnaire is supported by the WHO, it is not a true QOL tool.\textsuperscript{94} The ECQ is used to identify patients with claudication in general populations\textsuperscript{8,25} and provides binary outcome analysis (claudicant vs nonclaudicant), of which claudicant can be further sub-divided into definite claudicant and atypical claudicant. The ECQ is not intended to be used as a QOL measure.\textsuperscript{9,94}

Two surveys are useful for PAD severity but do not comprehensively measure QOL. The Claudication Scale (CLAU-S) exhibited strong associations with objective evaluators of PAD severity. However, while CLAU-S is useful for a quick functional assessment in claudicants, it is not a global assessment of QOL.\textsuperscript{5-7,94} Similarly, the Walking Impairment Questionnaire provides results that are strongly correlated with previously established, objective measures of PAD severity.\textsuperscript{38-40}

The Intermittent Claudication Questionnaire assesses QOL in intermittent claudicants and highlights the effects of claudication on tasks such as performing errands.\textsuperscript{14} The Intermittent Claudication Questionnaire is easy to administer and has been validated in English and Turkish. However, it has only been studied in the context of exercise programs and requires validation through other PAD studies.\textsuperscript{12-15,94}

The Claudication Symptom Instrument was developed by the Comparative Effectiveness Research Translation Network Collaborative in 2010 to compare the response of symptoms to medical versus surgical treatment of claudication.\textsuperscript{3} A mean of the intensity score for
the five evaluated symptoms is used to track symptoms over time or compare response to intervention.\textsuperscript{3} The Sickness Impact Profile—Intermittent Claudication is an abridged version of the Sickness Impact Profile, which has 11-fold more questions.\textsuperscript{26-28} Although the Sickness Impact Profile—Intermittent Claudication is a brief questionnaire and is easily scored, use of the disease-specific measures outside the context of the longer questionnaire requires further validation.\textsuperscript{29,30,94}

**Aortic aneurysm-specific PROMs.**

We identified four instruments specific to aortic aneurysms (Table II). The Aneurysm-Dependent Quality of Life (AneurysmDQoL) questionnaire assesses condition-specific QOL for patients with abdominal aortic aneurysm (AAA).\textsuperscript{42-44} The AneurysmDQoL assesses effect of AAA-specific symptoms on QoL and saliency of the symptoms.\textsuperscript{44,45} The Aneurysm Symptom Rating Questionnaire (AneurysmSRQ) assesses patient perception of the severity of AAA-specific symptoms.\textsuperscript{42-44} The AneurysmSRQ, in conjunction with the AneurysmDQoL, provides the most comprehensive assessments of fear of rupture, ability to forget about condition, and size of the aneurysm.\textsuperscript{43} However, for both AneurysmSRQ and AneurysmDQoL, the responsiveness to change (ability to detect a change in clinical symptoms or condition over time) has yet to be assessed.\textsuperscript{44}

The Aneurysm Treatment Satisfaction Questionnaire addresses the lack of condition-specific PROM for patients with AAA.\textsuperscript{42-44} The instrument evaluates patient satisfaction, assessing multiple aspects of patient QOL and attitudes relating to medical treatment (eg, information, postoperative follow-up, convenience, results feedback, and side effects). The questionnaire assesses two domains. The first domain focuses on monitoring/preintervention aspects of AAA and is applicable to all patients. The second domain focuses on postintervention treatment and, therefore, is only relevant for patients who have undergone aneurysm repair. The Aneurysm Treatment Satisfaction Questionnaire assesses patient satisfaction presurgical and postsurgical intervention; however, its responsiveness to change has yet to be assessed.\textsuperscript{44}

The Consequences of Screening questionnaire is an AAA-specific tool for assessing the psychosocial effects of screening in asymptomatic patients.\textsuperscript{46} The tool consists of two parts that assess 28 QOL dimensions ranging from anxiety, to uncertainty about the result of an ultrasound, to sexuality. Although the Consequences of Screening questionnaire has high content validity, responsiveness, and reliability among participants, the study was conducted using men and the content validity and reliability of this measure have not been tested among women.\textsuperscript{46}

**Thoracic outlet syndrome-specific PROMs.**

There were four thoracic outlet syndrome (TOS) specific instruments identified (Table III). The Cervical-Brachial Symptom Questionnaire (CBSQ) has been used in a battery of PROMs for patients presenting for evaluation of TOS and measures functional upper extremity disturbances related to the performance of certain common physical activities.\textsuperscript{38,48,49} Although the CBSQ is useful in the initial evaluation for TOS, its ability to predict the response to successful thoracic outlet decompression is unclear.\textsuperscript{49}
The Disabilities of the Arm, Shoulder and Hand (DASH) questionnaire was jointly developed by the American Academy of Orthopedic Surgeons, the Council of Musculoskeletal Specialty Societies, and Toronto’s Institute for Work and Health in 1996. The DASH assesses problems associated with daily tasks. There are four optional items to assess QOL dimensions specific to workers, athletes, and musicians. However, its length brings into question the likelihood of patient adherence. The Quick DASH was developed as an abridged version of the DASH with same functionality as the original DASH. The Quick DASH exhibits greater precision in differentiating various intensities of disability; however, it is not as comprehensive as the original DASH.

The Neurogenic Thoracic Outlet Symptom (NTOS) Index is a composite score that combines the validated DASH and CBSQ with a 10-point visual analog scale for pain. Constituent tools are scored, and the final scores are transformed onto a scoring range from 0 to 100, with a higher score suggesting greater degrees of disability. Although the NTOS Index is highly comprehensive, no clear advantage has been identified for using the NTOS Index as opposed to the composite tools independently.

Venous-specific PROMs.

We identified five PROs specific to venous disease (Table IV). The Aberdeen Varicose Vein Questionnaire (AVVQ) encompasses predominantly physical domains and social functioning aspects of QOL in patients with varicose veins. The AVVQ was validated in a prospective study using the self-administered SF-36 in patients undergoing varicose vein surgery. The AVVQ has since been used in multiple randomized studies as a measure to compare different venous treatment options. Its specificity often fails to assess the side effects of venous interventions; however, when used in conjunction with the SF-36, the AVVQ provides a more detailed evaluation.

The Chronic Venous Insufficiency (CIVIQ20) questionnaire identifies aspects of quality of life affected by venous insufficiency beyond physical discomfort, including psychological, physical functioning, social functioning, and pain. The CIVIQ20 has high content validity, internal consistency, and reliability in clinical research projects. In addition, it can be self-administered and had high sensitivity to change over time (responsiveness). However, the CIVIQ20 questionnaire offers a less thorough assessment of mental impacts of having varicose veins.

The Venous Clinical Severity Score (VCSS) was developed by the American Venous Forum in 2000 as part of the three-part Venous Severity Score (VSS). The VSS also includes the Venous Segmental Disease Score and the Venous Disability Score. The VCSS was developed to expand the CEAP classification by using a 0 to 3 scaling system for symptoms and findings that are progressive, measuring changes over short periods of time. VCSS scores correlate with the extent of the diseases anatomically. Additionally, the validation study illustrated the sensitivity of VCSS and VSS to changes after superficial venous surgery. Although the VCSS and VSS are not technically PROs, because both instruments include a physical examination component, both are a reliable way of tracking changes over time and used as a measure for comparison in several randomized trials.
The patient-reported Villalta (PRV) scale was developed as an adaptation of the original Villalta scale, which is used to diagnose post-thrombotic syndrome.\textsuperscript{71} Although the Villalta scale requires a clinical visit to perform a physical examination of an affected limb, the PRV scale was developed as a self-reported tool to assess symptoms and signs of post-thrombotic symptoms. The PRV has been shown to have very good agreement with the original Villalta scale,\textsuperscript{72} and has been used to enable remote assessment of PTS in a recent large clinical study.\textsuperscript{73}

The VVSymQ Instrument assesses unpleasant symptoms of varicose veins.\textsuperscript{74} The instrument is useful in assessing patient experience of varicose vein symptoms before and after the intervention; however, its brevity brings into question the comprehensiveness of the instrument.\textsuperscript{75}

**Hemodialysis access-specific PROMs.**

The Vascular Access Questionnaire (VAQ) assesses patient perception and attitude surrounding vascular access-related issues\textsuperscript{76} (Table V). The VAQ can be used to assess the saliency of patient vascular access-related concerns.\textsuperscript{76,77} However, the initial study outlining its development and clinical usefulness was limited, sampling only from a pool of Canadian dialysis patients within a single-payer health care system.\textsuperscript{76}

**Wound-specific PROMs.**

The Wound-QoL instrument measures wound-related QOL in patients with chronic wounds of varying etiology (Table V). The Wound-QoL incorporates components of three different preexisting PRO surveys—the Freiburg Life quality Assessment for wounds, the Cardiff Wound Impact Schedule, and the Wurzburg Wound Score—and has been validated for use in all patients with chronic wounds.\textsuperscript{78,79} Compared with these preexisting instruments, the Wound-QoL also evaluates multiple dimensions of quality of life; it is validated in English and is shorter in length, decreasing the patient burden of responding and improving the chances of obtaining high-quality data.\textsuperscript{80}

**Lymphedema-specific PROMs.**

The Lymphedema of the Limbs Quality of Life (LYMQOL) instrument is a lymphedema-specific instrument that has been adapted to both arm- and leg-specific lymphedema\textsuperscript{81} (Table V). The face and content validity for both Lymphedema of the Limbs Quality of Life have been demonstrated as well as for the four domains.\textsuperscript{82} The instrument has been evaluated in several non-English languages\textsuperscript{83-86}; however, construct validity and responsiveness have yet to be demonstrated.\textsuperscript{82}

**Selecting PROMs.**

Although recommendations for specific PROMs for vascular surgery patients are outside the scope of this review, there are some general considerations and resources to assist when deciding whether to start collecting PROs and how to select specific measures. The first consideration when selecting a specific measure is the intended use. For example, if the goal is to improve the diagnosis of patients with claudication, an instrument such as the ECQ would be most appropriate (Table I). Alternatively, if the goal is to assess
change in symptoms over time in response to treatment, then a measure designed to assess symptom severity would be most appropriate (eg, the Claudication Symptom Instrument) (see Tables I-V for the advantages and disadvantages of specific measures). Additional criteria to consider are whether the measure is validated (see anatomy of a PROM) and whether it is appropriate for the patient population selected. For example, there are measures that are validated general measures for quality of life for assessing change across a diverse group of patients (SF-36) and those that are specific to a particular disease process (see Tables I-V). Measure item length and availability in multiple languages can also have significant implications for response rate and limit adequate sampling of a patient population. Finally, factors that influence feasibility of implementation should be considered. Mode (self-administration vs interviewer administration) and method for collection (eg, electronic medical record and paper) and tools for analysis and reporting (eg, ePRO) will vary based on the measure and can significantly impact the cost and support available at a particular institution.

At present, there is limited consensus for use of particular measures for patients with vascular diseases. However, there are some resources that can help to guide selection for subsets of vascular patients. For example, the SVS reporting guidelines for TOS recommend the use of the QuickDash and CBSQ scores (see Table III) in the assessment of patient response to treatment for neurogenic TOS.50 However, a review of additional vascular-related guidelines and policy statements from large professional societies patients with PAD,1,95 venous disease,96 and cardiovascular disease97 did not yield any recommendations for specific endorsed PROMs. Additional resources for general guidelines to selection and best practices for PROM implementation include Health-Measures98 and the National Quality Forum.99

**VASCULAR SURGEON PERCEPTIONS OF PROMS: SURVEY DATA**

A survey was designed by the members of the Patient Reported Outcomes subcommittee of the SVS Performance Measures Committee to address the barriers and facilitators to using PROMs in clinical practice (Supplementary Table, online only). The survey was distributed to 106 SVS members serving on committees within the Policy and Advocacy Council. Of the 78 respondents completing the survey (response rate of 73.6%), 80.8% had heard of PROMs. All respondents (100%) who had heard of PROMs felt PROMs could be useful in assessing vascular surgery patients, particularly for patients with venous disease, PAD, and TOS (Fig). Only 23.1% of respondents indicated that their practice or institution used PROMs, although 80.0% indicated that their institution supported the use of PROMs. Of those respondents that actively used PROMs (n = 10 [12.8%]), the most common reason for collecting PROM data was for research and/or quality improvement initiatives (70.0%), followed by fulling an institutional requirement (50.0%), and quality reporting (40.0%). Nearly all respondents (90.0%) indicated they would consider using PROMs if they had the ability to incorporate the results into clinical practice, and 70.0% of respondents indicated they would consider using PROMs if they were incorporated into the electronic medical record. Reasons for not collecting PROMs were varied, and included concerns about available PROMs not being specific to patient problems and an inability to obtain results or analyze the collected data.
BARRIERS TO THE IMPLEMENTATION OF PROMs IN VASCULAR SURGERY: FOCUS GROUP DATA

Among those SVS members who completed the survey, a subset volunteered to participate in focus groups. Two 1-hour-long focus groups were conducted to assess granular perceptions of PROMS and preparedness of vascular surgeons for widespread implementation of PROMs. Focus group topics targeted physician awareness and knowledge of PROMs, the potential advantages and disadvantages of using PROMs in a vascular surgery practice, and any barriers that would impede PROM collection and interpretation.

The focus groups were conducted over video conference with a facilitator using a semistructured interview guide and were recorded. The recordings were professionally transcribed and transcripts were independently analyzed by three researchers who also participated in the focus group (C.H., A.V., and K.W.) and an additional analyst (M.R.). Transcripts were analyzed independently by at least two researchers for each transcript, who used open coding, resolved discrepancies with triangulation, and applied thematic analysis. The analysis identified four themes from the focus group data: (1) knowledge gaps, (2) the usefulness of PROMs in vascular surgery, (3) barriers to use of PROMs, and (4) concerns regarding unintended consequences of using PROMs in measuring quality of care.

Knowledge gaps.

Three subthemes were identified in knowledge gaps: a lack of (1) awareness of existing PROMs, (2) knowledge of how PROMs are developed and validated, and (3) clarity around how PROMs should be used by the clinician. Regarding knowledge of existing PROMs, most participants were aware of types of measurements (e.g., quality of life), but few were able to name specific measures and most were unaware that there were numerous validated PROMs relevant to vascular surgery patients. In addition, participants frequently conflated patient experience and satisfaction measures with HRQOL. Participants also described PROMs as a less rigorous form of data and felt that data reported by patients reflected opinion rather than outcome.

Focus group participants lacked an understanding of the rigorous development process that a validated PROM must undergo. In particular, there was a sentiment that PROM development and validation occurs in the absence of clinician participation. The participants believed that, if clinicians are not consulted or fail to participate in development or validation of PROMs, then it will be more challenging to ensure PROMs are applied and interpreted appropriately. However, participants did think that if recommendations for use of specific PROMs or guidelines on implementation and interpretation were released from known and respected specialty societies, they would be more likely to adopt and accept PROMs in their practice.

Finally, even among participants who had some experience with collection of PROMs, very few knew how to use PRO results to guide the care of individual patients in the way that traditional outcomes are used. The groups pointed out that the data depend not only on which PROM is used, but also when and how the information is collected. Although some
participants had experience with PROMs in a research setting, there was little experience or knowledge of how PROMs might be reported to clinicians and how the reports could impact patient care. However, having acknowledged this limitation, surgeons agreed that the prevalence of PROMs in clinical practice is evolving and will likely take time before best practices can be established.

**Usefulness of PROMs in vascular surgery.**

Two subthemes were identified in the usefulness of PROMs in vascular surgery: (1) the importance of incorporating the patient voice to define value, and (2) whether existing PROMs are capable of accounting for some of the confounding effect of patient morbidity on the outcome of care for vascular surgery patients. Participants agreed that the historical approach to patient care in medicine has failed to incorporate patient centeredness. Although the groups acknowledged that it may take time to determine what PROMs are most appropriate, the consensus was that failure to study the patient’s perspective on their health outcomes will impede our ability to provide truly valuable care.

However, although participants felt strongly that PROMs could help to overcome physician bias for or against how interventions or treatments impact a patient’s health, the groups were also concerned about whether PROMs would be applicable universally, given the complexity and variability of vascular surgery patients. For example, the group expressed concern that PROMs may be less applicable in patients who have limited treatment options such as those with end-stage PAD with no potential for revascularization. In addition, participants felt strongly that risk adjustment must be applied to PROs given the prevalence of significant comorbidities in vascular surgery patients and variation in outcomes by region and institution.

**Barriers to the use of PROMs.**

The focus groups identified two significant barriers to widespread use of PROMs: (1) logistical challenges of collecting and using PRO data, and (2) mistrust of outside oversight (eg, payors, employers) of outcome metrics. In the theme of logistical challenges, specific process barriers were identified that can be categorized under workflow and infrastructure. An example of infrastructure barriers is a lack of resources required to use an electronic device to capture responses. The group agreed that requiring patients to fill out paper forms that require manual entry into a database would likely require additional personnel or strain existing clinical staff. Even in centers where there was a potential for use of tablets or other electronic capture methods, surgeons acknowledged that the vascular surgery patient population is largely composed of elderly individuals who may have limited knowledge of or access to technology, therefore, potentially limiting the response rate in a typical practice. Furthermore, there may be a limited ability to capture these data within the electronic medical record itself, decreasing the physician’s ability to use the data for individual patients.

The groups expressed recurring concerns about how PROs would be interpreted not just by physicians but also by payers and policy makers. There was general concern about how PRO results would be used to measure the quality of care provided by physicians. Participants
acknowledged that although most postoperative outcomes in vascular surgery are a direct consequence of the procedure, many short- and long-term outcomes are also directly impacted by comorbidities and the nature of the disease, and are independent of sound decision-making and/or a well-executed operation. There was hope, but also skepticism, regarding whether payers and policymakers would acknowledge the need for further study before determining how best to hold providers accountable to PRO results.

Unintended consequences.

There were two subthemes were identified in the unintended consequences that centered around the ethics of patient and procedure selection: (1) risk avoidance and (2) the appropriateness of procedures. The concept of risk avoidance, specifically physician avoidance of caring for high-risk patients, is not unique to PROs, but is a concern that physicians have expressed in response to public reporting of outcomes. The surgeons in the focus groups indicated that use of PROMs to measure quality of care may be even more likely to influence this practice than traditional outcomes.

Participants also suggested that PROMs may not correlate with the appropriateness of a procedure. For example, there are procedures in vascular surgery such as interventions for claudication or varicose veins that may be associated with positive short-term impacts on a patient’s health, but can be associated with high rates of recurrence or complications that negatively impact a patient’s health in the long term. If PROs are considered in isolation without considering the appropriateness of the procedure, a positive result reported by a patient may inadvertently drive increases in certain procedures without consideration for appropriateness.

OVERCOMING THE BARRIERS

Based on our assessment of available vascular surgery-related PROMs, and feedback from the vascular surgery community, a number of actions are required to facilitate the widespread implementation of vascular surgery-related PROMs.

Development of vascular surgery-specific PROMs.

Future vascular surgery-related PROMs should assess issues surrounding areas where there are no existing PROMs, including carotid disease, aortic dissection, chronic limb-threatening ischemia, and mesenteric disease. Currently, these issues are not addressed in existing vascular surgery PROMs and their assessment is critical to a comprehensive understanding of the effect that the spectrum of commonly performed vascular surgery operations and interventions have on a patient’s HRQOL and satisfaction.

Recommendations for vascular surgery PROM best practices.

Recommendations for best practices in areas including PROM selection, administration, recording of results, and the use of the results at both the individual and population levels should be developed. Recommendations should consider varying practice settings and available resources.
PROM education for the vascular surgery community.

Education for the vascular surgery community is required about how PROMs are developed, available vascular surgery-related PROMs, the distinction between HRQOL and satisfaction, and how PROMs can be integrated into clinical practice to optimize patient outcomes and experience. Potential platforms for education include webinars, live events at professional meetings, web-based tutorials and printed materials such as newsletter articles and reviews in journals. The educational opportunities must be widely accessible and accommodate various preferences for learning.

Partnering with stakeholders.

The most important stakeholder in addressing the issues surrounding vascular surgery PROMs is the patient. Future development and implementation efforts must include the patient voice and partner with patients to ensure success. Professional societies with shared interests in disease processes and treatments should work together in developing PROMs, best practices, and educational programming.

To incorporate PROs into reimbursement models, collaboration with payers, including the CMS, is required to optimize the method of implementation for both the clinician and the patient. Including PROs into reimbursement models should occur in a stepwise fashion with initial introduction as a process measure, to allow clinicians to acclimate to new practice elements.

CONCLUSIONS

PROs are gaining increased attention in all fields of medicine. A number of validated disease or procedure specific PROMs exist that are relevant to vascular surgery, including those for PAD, AAA, TOS, venous disease, wounds, and lymphedema. Based on survey and focus group data, there is strong support for the use of PROs in vascular surgery practice as a means to provide truly valuable care for our patients. However, several barriers exist to widespread implementation of PROs in vascular surgery. PRO collection is resource intensive and the widespread lack of education about the development, use, and potential harms related to PROM collection and reporting will significantly impede successful adoption. Further research is required to develop PROMs for all common vascular diseases and to ensure best practices around collection and interpretation. Societal leadership will play a pivotal role in defining how PROs may be best used in vascular surgery and collaboration with physicians, patients and payors will be vital to optimize patient care and improve patient-centered outcomes without encouraging risk avoidance or inappropriate care.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.
REFERENCES


J Vasc Surg. Author manuscript; available in PMC 2023 January 12.
88. Weldring T, Smith S. Patient-reported outcomes (PROs) and patient-reported outcome measures (PROMs). Health Services Insights 2013;6:61–8. [PubMed: 25114561]


Fig.
Summary of survey response answers to question about what patient groups patient-reported outcome measures (PROMs) may be helpful. CLTI, Chronic limb-threatening ischemia; TOS, thoracic outlet syndrome.
Table I.
Peripheral arterial disease-related patient-reported outcomes measures (PROM)

<table>
<thead>
<tr>
<th>PROM</th>
<th>Year</th>
<th>No. of items</th>
<th>Domains</th>
<th>Time to complete, min</th>
<th>Validated</th>
<th>Advantages</th>
<th>Disadvantages</th>
</tr>
</thead>
<tbody>
<tr>
<td>CSI — Vascular&lt;sup&gt;1,4&lt;/sup&gt;</td>
<td>2016</td>
<td>5</td>
<td>Symptoms</td>
<td>5</td>
<td>Yes</td>
<td>Assess specific symptoms, as opposed to global symptoms; designed and validated for assessment of outcomes to treatment</td>
<td>Limited testing in diverse populations; study was not designed for optimal evaluation of CSI sensitivity to change (responsiveness to change in patient’s symptoms)</td>
</tr>
<tr>
<td>CLAU-S&lt;sup&gt;5-7&lt;/sup&gt;</td>
<td>1995</td>
<td>47</td>
<td>Everyday life, pain, effect on social activities, illness-specific fears, psychological impact</td>
<td>5</td>
<td>Yes</td>
<td>Strongly associated with objective measures of PAD severity</td>
<td>Functional assessment with limited use as global QOL tool</td>
</tr>
<tr>
<td>ECQ&lt;sup&gt;8&lt;/sup&gt;</td>
<td>1992</td>
<td>6</td>
<td>Identification of claudication</td>
<td>Not reported</td>
<td>Yes</td>
<td>Tantamount to WHO/Rose but more specific</td>
<td>Not a true QOL assessment</td>
</tr>
<tr>
<td>FLeQKI&lt;sup&gt;10,11&lt;/sup&gt;</td>
<td>2007</td>
<td>35</td>
<td>Comorbidity, effect on physical activities, effect on social activities, pain, psychological impact</td>
<td>Not reported</td>
<td>Yes</td>
<td>Correlate with SF-36</td>
<td>Only validated in German</td>
</tr>
<tr>
<td>Intermittent Claudication Questionnaire&lt;sup&gt;12-15&lt;/sup&gt;</td>
<td>2002</td>
<td>16</td>
<td>Pain, effect on physical activities, effect on social activities, psychological impact</td>
<td>3.7</td>
<td>No</td>
<td>Brief</td>
<td>Requires further validation</td>
</tr>
<tr>
<td>Peripheral Artery Disease Quality of Life Questionnaire&lt;sup&gt;16,17&lt;/sup&gt;</td>
<td>2012</td>
<td>38</td>
<td>Social relationships and interactions, self-concept and feelings, symptoms, effect on physical activities, psychological impact, positive adaptation</td>
<td>&lt;10</td>
<td>Yes</td>
<td>Demonstrates physical and emotional consequences of PAD on patient QOL</td>
<td>Limited testing in diverse populations, distinct age and race subgroups, and impact of intervention on disease progression</td>
</tr>
<tr>
<td>Peripheral Artery Questionnaire&lt;sup&gt;18,19&lt;/sup&gt;</td>
<td>2004</td>
<td>20</td>
<td>Symptoms, change in symptoms, effect on physical activities, effect on social activities, treatment satisfaction, and overall QOL</td>
<td>Not reported</td>
<td>Yes</td>
<td>Holistic assessment of QOL and treatment effect</td>
<td>Less comprehensive than other PAD-specific QOL tools</td>
</tr>
<tr>
<td>Peripheral Artery Occlusive Disease 86-Item Questionnaire&lt;sup&gt;12,13,16-23&lt;/sup&gt;</td>
<td>1995</td>
<td>86</td>
<td>Functional status, pain, general complaints, mood, anxiety, social life, evaluation of treatment for PAD</td>
<td>20</td>
<td>Yes</td>
<td>Extensive</td>
<td>Length limits adherence</td>
</tr>
<tr>
<td>PBI-PAD&lt;sup&gt;24-25&lt;/sup&gt;</td>
<td>2018</td>
<td>12</td>
<td>Everyday life, working life, therapy, leisure time, body, psychological impact</td>
<td>Not reported</td>
<td>Yes</td>
<td>Calculates pre—post differences</td>
<td>Feasibility; 2 questionnaires over 3 months</td>
</tr>
<tr>
<td>Sickness Impact Profile—Intermittent Claudication&lt;sup&gt;26-30&lt;/sup&gt;</td>
<td>1975</td>
<td>12</td>
<td>Sleep and rest, home management, ambulation, mobility, social interaction and alertness, behavior</td>
<td>Not reported</td>
<td>Yes</td>
<td>Brief; uses simple scoring scheme</td>
<td>Largely bereft in clinical spaces</td>
</tr>
<tr>
<td>VascuQoL&lt;sup&gt;31-34&lt;/sup&gt;</td>
<td>2001</td>
<td>25</td>
<td>Pain, symptoms, effect on physical activities, effect on social activities, psychological impact</td>
<td>9</td>
<td>Yes</td>
<td>Highlights PAD treatments effects on QOL</td>
<td>Relationship with functional status is lacking</td>
</tr>
<tr>
<td>VascuQoL-6&lt;sup&gt;35&lt;/sup&gt;</td>
<td>2014</td>
<td>6</td>
<td>Pain, symptoms, effect on physical activities, effect on social activities, psychological impact</td>
<td>1.4</td>
<td>Yes</td>
<td>Derived from VascuQoL but shorter</td>
<td>Limited comprehensiveness</td>
</tr>
<tr>
<td>PROM</td>
<td>Year</td>
<td>No. of items</td>
<td>Domains</td>
<td>Time to complete, min</td>
<td>Validated</td>
<td>Advantages</td>
<td>Disadvantages</td>
</tr>
<tr>
<td>-------------------------------------------</td>
<td>------</td>
<td>--------------</td>
<td>----------------------------------------</td>
<td>-----------------------</td>
<td>-----------</td>
<td>-----------------------------------------------------------------------------</td>
<td>---------------------------------------------</td>
</tr>
<tr>
<td>WHO/Rose Questionnaire,8,36,37</td>
<td>1962</td>
<td>8</td>
<td>Identification of claudication</td>
<td>Not reported</td>
<td>Yes</td>
<td>Global utilization; endorsed by WHO</td>
<td>Not a true QOL assessment</td>
</tr>
<tr>
<td>Walking Impairment Questionnaire,34,38,41</td>
<td>1990</td>
<td>22</td>
<td>Pain, distance, walking speed, and stair climbing</td>
<td>5</td>
<td>Yes</td>
<td>Strong correlation with objective measures of PAD severity</td>
<td>Functional tool only, not a global assessment of QOL</td>
</tr>
</tbody>
</table>

CLA-U-S, Claudication Scale; CSI, Claudication Symptom Instrument; ECQ, Edinburgh Claudication Questionnaire; PAD, peripheral artery disease; PBI-PAD, Patient Benefit Index for Peripheral Arterial Disease; QOL, quality of life; SF-36, Short Form-36; WHO, World Health Organization.
### Table II.

Aneurysm-related patient-reported outcomes measures (*PROM*)

<table>
<thead>
<tr>
<th>PROM</th>
<th>Year</th>
<th>No. of items</th>
<th>Domains</th>
<th>Time to complete, minutes</th>
<th>Validated</th>
<th>Advantages</th>
<th>Disadvantages</th>
</tr>
</thead>
<tbody>
<tr>
<td>AneurysmDQoL</td>
<td>2016</td>
<td>24</td>
<td>Physical symptoms, psychological symptoms, treatment satisfaction</td>
<td>Not reported</td>
<td>Yes</td>
<td>Provides most comprehensive assessments of fear of rupture, control, ability to forget about condition and size of aneurysm</td>
<td>Responsiveness to change not assessed</td>
</tr>
<tr>
<td>AneurysmSRQ</td>
<td>2016</td>
<td>44</td>
<td>Symptoms</td>
<td>Not reported</td>
<td>Yes</td>
<td>Provides most comprehensive assessments of fear of rupture, control, ability to forget about condition and size of aneurysm</td>
<td>Responsiveness to change not assessed</td>
</tr>
<tr>
<td>Aneurysm Treatment Satisfaction Questionnaire</td>
<td>2016</td>
<td>11</td>
<td>Monitoring/preintervention satisfaction, postoperative treatment satisfaction</td>
<td>Not reported</td>
<td>Yes</td>
<td>Assesses patient satisfaction presurgical and postsurgical intervention</td>
<td>Responsiveness to change not assessed</td>
</tr>
<tr>
<td>Consequences of Screening</td>
<td>2018</td>
<td>62</td>
<td>Psychological, effects on social activities</td>
<td>Not reported</td>
<td>Yes</td>
<td>High content validity, responsiveness, and reliability</td>
<td>Limited testing in gender diverse populations — untested content validity and reliability</td>
</tr>
</tbody>
</table>

*AneurysmDQoL*, Aneurysm-Dependent Quality of Life; *AneurysmSRQ*, Aneurysm Symptom Rating Questionnaire.
### Table III.

Thoracic outlet syndrome (TOS)-related patient-reported outcomes measures (PROM)

<table>
<thead>
<tr>
<th>PROM</th>
<th>Year</th>
<th>No. of Items</th>
<th>Domains</th>
<th>Time to complete, minutes</th>
<th>Validated</th>
<th>Advantages</th>
<th>Disadvantages</th>
</tr>
</thead>
<tbody>
<tr>
<td>CBSQ[38,48-50]</td>
<td>2007</td>
<td>14</td>
<td>Symptoms</td>
<td>Not reported</td>
<td>Yes</td>
<td>Useful in initial evaluation for TOS</td>
<td>Unclear ability to predict response to successful thoracic outlet decompression</td>
</tr>
<tr>
<td>DASH[51,52]</td>
<td>1996</td>
<td>30</td>
<td>Symptoms, effect on physical activities, effect on social activities</td>
<td>Not reported</td>
<td>Yes</td>
<td>Assesses problems associated with daily tasks</td>
<td>Length of questionnaire</td>
</tr>
<tr>
<td>Quick DASH[50,52-54]</td>
<td>2008</td>
<td>11</td>
<td>Symptoms, effect on physical activities, effect on social activities</td>
<td>Not reported</td>
<td>Yes</td>
<td>Same principles as DASH and less time</td>
<td>Not as comprehensive as DASH</td>
</tr>
<tr>
<td>NTOS[49,50,52,55,56]</td>
<td>2013</td>
<td>45</td>
<td>Symptoms, effect on physical activities, effect on social activities, pain</td>
<td>&lt;20</td>
<td>No</td>
<td>Comprehensive, comprised of 3 tools (CBSQ, DASH, and 10-point scale for pain)</td>
<td>No clear advantage to this instrument over the CBSQ and DASH as individual measures</td>
</tr>
</tbody>
</table>

CBSQ, Cervical-Brachial Symptom Questionnaire; DASH, Disabilities of the Arm, Shoulder and Hand; NTOS, Neurogenic Thoracic Outlet Symptom Index.
Table IV.

Venous disease-related patient-reported outcomes measures (PROM)

<table>
<thead>
<tr>
<th>PROM</th>
<th>Year</th>
<th>No. of items</th>
<th>Domains</th>
<th>Time to complete, min</th>
<th>Validated</th>
<th>Advantages</th>
<th>Disadvantages</th>
</tr>
</thead>
<tbody>
<tr>
<td>AVVQ&lt;sup&gt;57-65&lt;/sup&gt;</td>
<td>1993</td>
<td>13</td>
<td>Symptoms, effect on physical activities, use of compression, body image/appearance</td>
<td>&lt;5</td>
<td>Yes</td>
<td>Varicose vein specific; can be used to compare different venous treatment options</td>
<td>Specificity often fails to assess side effects of venous interventions; more detailed evaluation of QOL when used in conjunction with the SF-36</td>
</tr>
<tr>
<td>CIVIQ20&lt;sup&gt;66-68&lt;/sup&gt;</td>
<td>1996</td>
<td>20</td>
<td>Psychological, physical functioning, social functioning, and pain</td>
<td>Not reported</td>
<td>Yes</td>
<td>High content validity, reliability, internal consistency; high responsiveness by patients with self-administration</td>
<td>Limited assessment of mental impacts of having varicose veins</td>
</tr>
<tr>
<td>VCSS&lt;sup&gt;09-30&lt;/sup&gt;</td>
<td>2000</td>
<td>10</td>
<td>Symptoms, physical examination</td>
<td>Not reported</td>
<td>Yes</td>
<td>Tracks change over time; especially after superficial venous surgery</td>
<td>Not designed to directly measure healthcare-related QOL</td>
</tr>
<tr>
<td>PRV&lt;sup&gt;71-73&lt;/sup&gt;</td>
<td>2016</td>
<td>13</td>
<td>Symptoms, pain, body image/appearance</td>
<td>Not reported</td>
<td>Yes</td>
<td>Sensitive tool for diagnosing post-thrombotic syndrome</td>
<td>Text-based tool is not as accurate as the visually assisted tool</td>
</tr>
<tr>
<td>VVSymQ&lt;sup&gt;74-75&lt;/sup&gt;</td>
<td>2014</td>
<td>5</td>
<td>Symptoms</td>
<td>Not reported</td>
<td>Yes</td>
<td>Assesses patient experience of symptoms before and after the intervention</td>
<td>Limited comprehensiveness</td>
</tr>
</tbody>
</table>

AVVQ, Aberdeen Varicose Vein Questionnaire; CIVIQ20, Chronic Venous Insufficiency; PRV, patient-reported Villalta; QOL, quality of life; SF-36, Short Form 36; VCSS, Venous Clinical Severity Score.
Table V.
Hemodialysis access, wounds and lymphedema-related patient-reported outcomes measures (PROM)

<table>
<thead>
<tr>
<th>PROM</th>
<th>Year</th>
<th>No. of Items</th>
<th>Domains</th>
<th>Time to complete, min</th>
<th>Validated</th>
<th>Advantages</th>
<th>Disadvantages</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hemodialysis access</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>V AQ</td>
<td>2008</td>
<td>17</td>
<td>Symptoms, body image/appearance, function, effect on physical activities, effect on social activities, and mood</td>
<td>Not reported</td>
<td>Yes</td>
<td>Assesses saliency of patient vascular access-related concerns</td>
<td>Initial study used limited testing in diverse populations</td>
</tr>
<tr>
<td>Wounds</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Wound QoL</td>
<td>2014</td>
<td>17</td>
<td>Everyday life, body, psyche</td>
<td>2.4</td>
<td>Yes</td>
<td>Validated for use in all patients with chronic wounds</td>
<td>Validated in English-speaking patients</td>
</tr>
<tr>
<td>Lymphedema</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lymphedema of the Limbs Quality of Life—arm</td>
<td>2010</td>
<td>22</td>
<td>Symptoms, body image/appearance, function, and mood</td>
<td>Not reported</td>
<td>Yes</td>
<td>Evaluated in several non-English languages</td>
<td>Construct validity and responsiveness not yet demonstrated</td>
</tr>
<tr>
<td>Lymphedema of the Limbs Quality of Life—leg</td>
<td>2010</td>
<td>23</td>
<td>Symptoms, body image/appearance, function, and psychological</td>
<td>Not reported</td>
<td>Yes</td>
<td>Evaluated in several non-English languages</td>
<td>Construct validity and responsiveness not yet demonstrated</td>
</tr>
</tbody>
</table>

V AQ, Vascular Access Questionnaire.