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1 Title: Patient-Reported Outcome Measures in Carotid Artery Revascularisation: Systematic  
2 Review and Psychometric Analysis

3

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28 **Abstract**

29 **Objective:**

30 Patient-reported outcome measures (PROMs) provide a way to measure the impact of a  
31 disease and its associated treatments on the quality of life from the patients' perspective. The  
32 aim of this review was to identify PROMs that have been developed and/or validated in  
33 patients with carotid artery stenosis (CAS) undergoing revascularisation and to assess their  
34 psychometric properties and examine suitability for research and clinical use.

35 **Methods:**

36 Eight electronic databases including MEDLINE and CINAHL were searched using a two-  
37 stage search approach to identify studies reporting the development and/or validation of  
38 relevant PROMs in patients with CAS undergoing revascularisation. Supplementary citation  
39 searching and hand-searching reference lists of included studies were also undertaken. The  
40 Consensus-based Standards for the selection of health Measurement INstruments (COSMIN)  
41 and Oxford criteria were used to assess the methodological quality of the included studies and  
42 the psychometric properties of the PROMs were evaluated using established assessment  
43 criteria.

44 **Results:**

45 Five studies reporting on six PROMs were included: 36-Item Short Form Health Survey (SF-  
46 36), Euro-QoL-5-Dimension Scale (EQ-5D), Hospital Anxiety and Depression Scale  
47 (HADS), Dizziness Handicap Inventory (DHI), Quality of life for carotid artery disease scale  
48 and a disease-specific PROM for CAS. The rigour of the psychometric assessment of the  
49 PROMs was variable with most only attempting to assess a single psychometric criterion. No  
50 study reported evidence on construct validity and test-retest reliability. Evidence for  
51 acceptability for the use of SF-36, EQ-5D and the disease-specific PROM were rated good in  
52 most studies. Only one study reported a Cronbach alpha score >0.70 as evidence of internal

53 consistency. Overall, the psychometric evaluation of all included PROMs was rated as poor  
54 within the CAS population undergoing revascularisation.

55 **Conclusions:**

56 This review highlighted a lack of evidence in validated PROMs used for patients undergoing  
57 carotid artery revascularisation. As a result, the development and validation of a new PROM  
58 for this patient population is warranted in order to provide data which can supplement  
59 traditional clinical outcomes (stroke<30 days post-procedural, myocardial infarction and  
60 death) and capture changes in health status and quality of life to help inform treatment  
61 decisions.

62

63 **Keywords:** Carotid artery revascularisation; Patient-Reported Outcome Measures; PROMs;  
64 Quality of life; Vascular surgery; Psychometric.

65        **1. Introduction**

66        Thromboembolism from carotid artery stenosis (CAS) is a major cause of stroke, accounting  
67        for one in five cases of all strokes.<sup>1</sup> Patients with CAS can remain asymptomatic until the  
68        carotid arteries are severely narrowed or blocked and in some cases transient ischaemic attack  
69        or stroke is the first sign of the disease. Patients with severely narrowed or blocked arteries  
70        may undergo a surgical procedure to open the arteries and to prevent stroke and its  
71        complications from occurring, namely death or decrease in quality of life (QoL).

72  
73        Patient reported outcome measures (PROMs) are questionnaires completed by the patient in  
74        relation to their health and daily functioning. This provides a way of measuring the impact of  
75        a disease and its associated treatments on the health and QoL from the patients' perspective.<sup>2</sup>  
76        PROMs can be categorised as generic, disease-specific or dimensional specific (measure the  
77        effect of an intervention on a specific concept e.g. anxiety). Generic PROMs can be used in a  
78        variety of conditions and allows comparison across different patient groups. In contrast,  
79        disease-specific PROMs are specific to treatments and symptoms associated with a particular  
80        disease or condition. Both generic and disease-specific PROMs can be preference-based  
81        PROMs and can be used to estimate preference weights for calculating quality-adjusted life-  
82        years, from which an economic value of interventions can be assessed.<sup>3;4</sup>

83  
84        The United States Food and Drug Administration (FDA) recommends the use of both generic  
85        and disease-specific measures in clinical trials<sup>5</sup> and in the United Kingdom the National  
86        Institute for Health and Care Excellence (NICE) use PROMs data to facilitate health  
87        technology assessments.<sup>6</sup> Since 2009 the NHS has made it a requirement to collect PROM  
88        data from patients before and after surgery in four surgical conditions: hip replacement, knee  
89        replacement, varicose vein treatment and groin hernia repair. Currently, PROMs are not

90 routinely used in carotid artery revascularisation. The addition of validated PROMs to the  
91 hard clinical outcomes (i.e. stroke < 30 days post-procedural, myocardial infarction and  
92 death) in patients undergoing carotid artery revascularisation, can provide information about  
93 the quality of care and the impact of treatment on a patient's QoL including wound  
94 complications, cranial nerve damage, drug side effects and anxiety associated with the  
95 condition and treatments.<sup>7</sup> It is important to use PROMs that have followed best practice in  
96 terms of their development and evaluation to ensure the PROMs are 'appropriate and  
97 comprehensive relative to its intended measurement concept, population, and use'.<sup>2</sup>

98

99 The aim of this review was to identify studies reporting on the development and/or validation  
100 of PROMs for use in patients with CAS undergoing revascularisation, critically appraise the  
101 psychometric properties of the PROMs, and examine its suitability for clinical and research  
102 use. This review forms part of a larger study funded by the NIHR examining the re-  
103 configuration of vascular services in the UK and identify targets for future research.

104

## 105 **2. Methods**

106 This systematic review was reported in accordance with the general principles recommended  
107 in the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA)  
108 statement.<sup>8</sup> A protocol was developed and registered on the PROSPERO international  
109 prospective register of systematic reviews  
110 ([http://www.crd.york.ac.uk/PROSPERO/display\\_record.php?ID=CRD42015023877](http://www.crd.york.ac.uk/PROSPERO/display_record.php?ID=CRD42015023877)).

111

### 112 **2.1 Data sources and searches**

113 Systematic searches were undertaken in eight electronic databases and research registers  
114 including MEDLINE, MEDLINE in Process, EMBASE, the Cochrane Library, CINAHL,

115 PROQOLID, PsychINFO and Web of Science. A two-stage search approach was used. The  
 116 first stage combined known generic and condition-specific terms for PROMs and CAS. The  
 117 studies were retrieved and examined for additional PROM terms used in CAS. Stage 2  
 118 incorporated PROM terms identified in stage 1 with a preliminary search strategy and a  
 119 methodological search filter for finding studies on measurement properties.<sup>9</sup> Databases were  
 120 searched from inception up to February 2015 (for stage 1) and up to May 2015 (for stage 2).  
 121 Both searches were updated in February 2017. No language or date restrictions were applied.  
 122 Searches were supplemented by hand-searching the reference lists of relevant reviews and  
 123 included studies, citation searching and contact with experts in the field. Details of the search  
 124 strategies are provided in Supplementary Appendix 1.

125

## 126 **2.2 Study selection**

127 All identified titles were examined for inclusion and any citations that clearly did not meet  
 128 the inclusion criteria were excluded (e.g. non-human, unrelated to CAS). All abstracts and  
 129 full text articles were then examined by at least two reviewers. Any disagreements in the  
 130 selection process were resolved by discussion, with involvement of a third reviewer when  
 131 necessary. A summary of the inclusion and exclusion criteria is presented in Table I.

132

133 **Table I: Study Selection Criteria**

	<b>Inclusion criteria</b>	<b>Exclusion criteria</b>
Population	A defined population of participants with a confirmed diagnosis of CAS (using ultrasonography, computed tomography, magnetic resonance imaging, or	Patients not diagnosed with CAS



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conventional angiography) who need, have had, or are undergoing revascularisation.

Intervention	Any surgical treatment indicated for CAS e.g. carotid endarterectomy, carotid artery stenting and angioplasty	Non-surgical interventions for CAS
Outcomes	PROMs (including generic, disease-specific, preference-based, functional and symptoms) used to assess quality of life in patients with CAS undergoing revascularisation	Outcome measures of patient satisfaction or experience in the relevant population  PROMs from Proxy
Study design	Any	
Publication type	Published or unpublished full-text peer reviewed journal articles including structured abstracts with all relevant information	Reviews, Editorial and Opinion pieces
Language	English	Non-English

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CAS, carotid artery stenosis; PROMs, Patient reported outcome measures;

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### 135 **2.3 Data abstraction**

136 Data relating to study design, patient characteristics, type of surgical treatment, type of

137 PROM used, methods and outcomes were extracted by one reviewer into a standardised data

138 extraction form, and independently checked for accuracy by a second. Any discrepancies  
139 were resolved by discussion, with involvement of a third reviewer, if required.

140

#### 141 **2.4 Psychometric evaluation**

142 The methodological quality and the psychometric properties of the included PROMs were  
143 assessed by two independent reviewers. Any disagreements were resolved by discussion and  
144 when needed with the involvement of a third reviewer. Criteria used to appraise the PROMs  
145 (see Table II) were adapted from published recommendations.<sup>10-16</sup> These criteria have been  
146 successfully applied previously<sup>17;18</sup> and are consistent with the FDA guidance.<sup>2</sup> The  
147 instruments were examined for their reliability (the degree to which measures are  
148 reproducible and consistent over time in patients with a stable condition); validity (the degree  
149 to which the instrument measures what it is supposed to measure); responsiveness (the degree  
150 to which the instrument detects meaningful change over time if a change truly exists) and  
151 acceptability (the degree to which the instrument is acceptable to the patients). As no gold  
152 standard exists for QoL, criterion validity was not assessed.

153

154 **Table II: Appraisal criteria for assessing the psychometric properties of patient-**  
155 **reported outcome measures**

<b>Domain</b>	<b>Sub-domain</b>	<b>Criteria</b>
Reliability	Test re-test	The intra-class correlation/ weighted kappa score should be $\geq 0.70$ for group comparisons and $\geq 0.90$ if scores are going to be used for decisions about an individual based on their score. <sup>10</sup>  The mean difference (paired t test or Wilcoxon signed-

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rank test) between time point 1 (T<sub>1</sub>) and time point 2 (T<sub>2</sub>) and the 95% CI should also be reported.

Internal consistency		A Cronbach's alpha score of $\geq 0.70$ is considered good and it should not exceed $\geq 0.92$ for group comparisons as this is taken to indicate that items in the scale could be redundant. Item total correlations should be $\geq 0.20$ . <sup>13</sup>
Validity	Content validity	This is assessed qualitatively during the development of an instrument. To achieve good content validity, there must be evidence that the instrument has been developed by consulting patients, experts as well as undertaking a literature review.  Patients should be involved in the development stage and item generation. The opinion of patient representatives should be sought on the constructed scale. <sup>10;12;13</sup>
	Construct validity	A correlation co-efficient of $\geq 0.60$ is taken as strong evidence of construct validity. Authors should make specific directional hypotheses and estimate the strength of correlation before testing. <sup>10;13;14</sup>
Responsiveness	Responsiveness	There are a number of methods to measure this including t-tests, effect size, standardised response means or responsiveness statistics Guyatts' responsiveness index. <sup>16</sup> There should be statistically significant changes in score of an expected

magnitude.<sup>15</sup>

Acceptability      Floor-ceiling effects      A floor or ceiling effect is considered if 15% of respondents are achieving the lowest or the highest score on the instrument.<sup>14</sup>

Acceptability      Acceptability was measured by the completeness of the data supplied. 80% or more of the data should be complete.<sup>12</sup>

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156

### 157      **3. Results**

158      A total of 1,670 records were identified, of which 126 full-text articles were considered  
159      eligible for inclusion. Following detailed examination, five studies (reporting on the  
160      development and/or validation of six PROMs) were included in this review. All the included  
161      studies reported the validation or development of PROMs in patients with symptomatic  
162      and/or asymptomatic CAS undergoing surgical treatment. The majority of the excluded  
163      studies did not present data evaluating the measurement properties of PROMs and only  
164      reported the use of PROMs in patients with CAS undergoing revascularisation. A summary  
165      of the process for identifying and selecting the relevant literature is presented in Figure 1.

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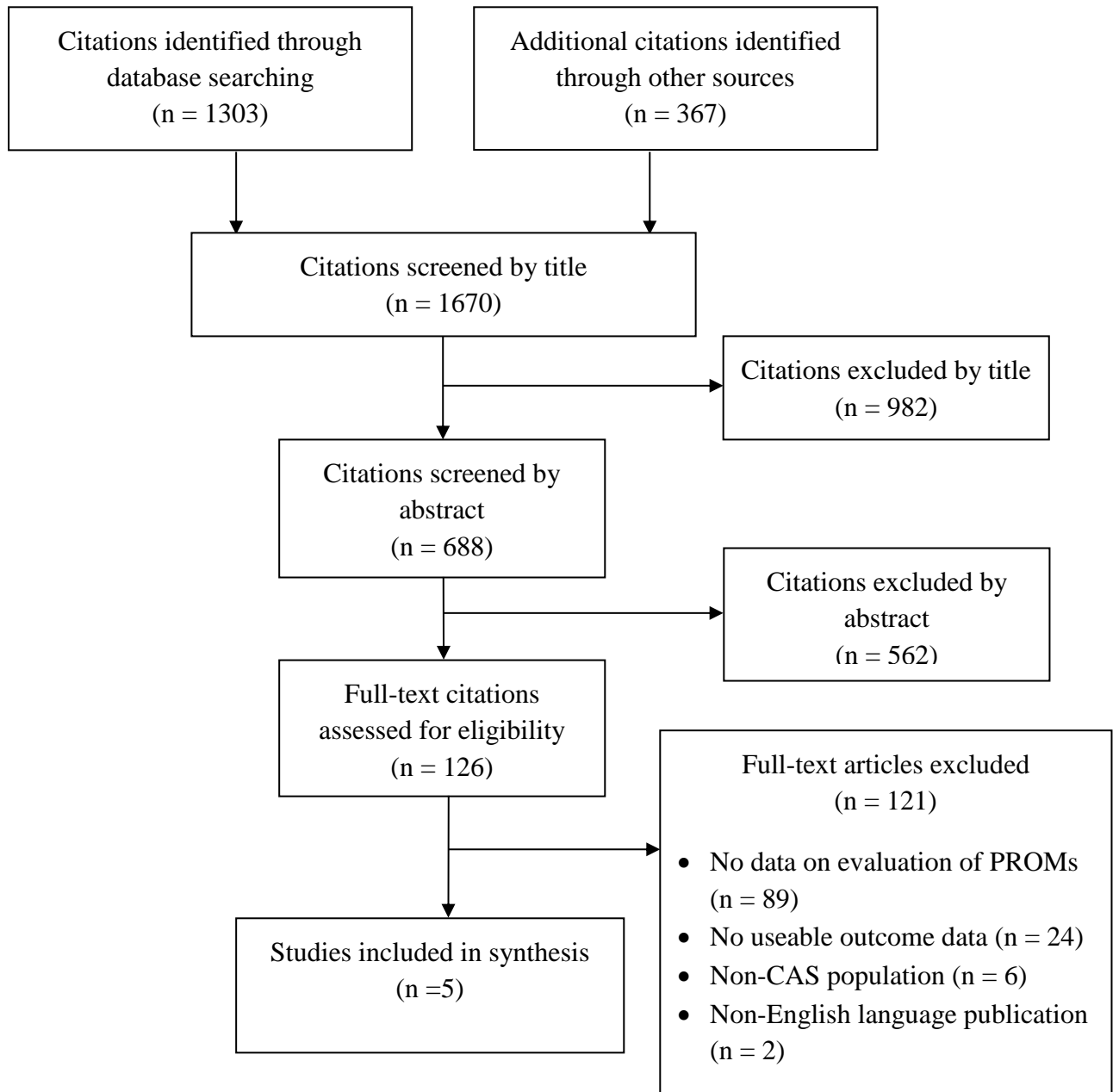
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175 **Figure 1: Study flow chart (adapted) of study selection**

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209 **3.1 Study characteristics**

210 Table III presents the study characteristics of the five included studies. All the included  
 211 studies used PROMs to assess the health related quality of life (HRQoL) or functional status  
 212 of patients undergoing revascularisation and reported aspects of the methodological details of  
 213 the PROMs development and/or validation. The studies were prospective in design and were  
 214 undertaken in the USA,<sup>19</sup> Germany,<sup>20</sup> USA/ Canada,<sup>21</sup> Taiwan<sup>22</sup> and Latvia.<sup>23</sup> The studies  
 215 were published between 2010<sup>19</sup> and 2015,<sup>23</sup> and the majority of the studies were of a small to  
 216 moderate size with the number of participants ranging from 61<sup>22</sup> to 2502.<sup>21</sup> Adults of either  
 217 sex were recruited with the proportion of men ranging between 55%<sup>23</sup> to 84%<sup>22</sup> and the mean  
 218 age range between 69 years<sup>21</sup> and 73 years.<sup>22</sup>

219

220 The patients' clinical diagnosis varied across studies: four studies<sup>19-22</sup> included patients with  
 221 both symptomatic and asymptomatic carotid artery stenosis, whilst one study, Ivanova et al<sup>23</sup>  
 222 only included asymptomatic patients. The types of surgical treatment reported for carotid  
 223 revascularisation included carotid endarterectomy (CEA),<sup>20;23</sup> carotid artery stenting (CS)<sup>22</sup>  
 224 and in two studies<sup>19;21</sup> both CS and CEA were used.

225

226 **Table III: Study and patient characteristics of included studies reporting validation of**  
 227 **PROMs in patients**

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Author, year	Country	Diagnosis (Sample size)	Age, years (mean ±SD)	Gender n/N (%) males)	Reported PROM(s)	Timing of PROM(s) assessment	Treatment
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Stolker 2010 <sup>19</sup>	USA (SAPPHIRE Trial)	High risk patients symptomatic and asymptomatic <b>(N=310)</b>	72 ( $\pm$ 8)	211/310 (68.1)	EQ-5D SF-36 Disease- specific PROM	Baseline, 2 weeks, 1,6 and 12 months post-surgery	CEA versus CS in high risk patients
Attigah 2011 <sup>20</sup>	Germany	Symptomatic and asymptomatic carotid stenosis <b>(N=102)</b>	Median age (range): 70 (42- 86)	70/102 (68.6)	HADS EQ-5D	1 day before and 2 days post-surgery	Local anaesthetic in CEA
Cohen 2011 <sup>21</sup>	USA &Canada (CREST Trial)	Symptomatic and asymptomatic carotid stenosis <b>(N=2,502)</b>	69 (NR)	1626/250 2 (65)	SF-36 Disease- specific PROM	Baseline, 2 weeks, 1 month and 1 year post- surgery	CEA versus CS
Hsu 2014 <sup>22</sup>	Taiwan	Symptomatic and asymptomatic carotid stenosis <b>(N=61)</b>	73.3 ( $\pm$ 10.5)	51/61 (83.6)	SF-36 DHI	1 week before, 1 and 6 months post-surgery	CS

Ivanova 2015 <sup>23</sup>	Latvia	Asymptomatic carotid artery stenosis (N=120)	Median age (range): 69.3 (42-84)	66/120 (55)	Quality of life for carotid artery disease	1,3,6,9,12 months before entry and 4 months until total of 24 months	CEA
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CS, carotid artery stenting; CEA, carotid endarterectomy; CREST, Carotid Revascularisation Endarterectomy Versus Stenting Trial; DHI, Dizziness Handicap Impact; SAPPHERE, Stenting and Angioplasty with Protection in Patients at High Risk for Endarterectomy; PROMs, patient reported outcome measure; SD, standard deviation; NR, not reported

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231  
232

### 233 3.2 PROMs data and psychometric evaluation

234 Five studies reported data relating to the psychometric evaluation of PROMs in patients  
235 undergoing carotid revascularisation. Of these, two were generic PROMs: 36-item Short  
236 Form Health Survey (SF-36)<sup>19;21;22</sup> and Euro-QoL 5 Dimension Scale (EQ-5D).<sup>19;20</sup> Two were  
237 dimension-specific PROMs: Hospital Anxiety & Depression scale (HADS)<sup>20</sup> - a mental  
238 health specific PROM and Dizziness Handicap Inventory (DHI)<sup>22</sup>. Two were condition-  
239 specific PROMs: Quality of life for carotid artery disease scale designed by Ivanova et al<sup>23</sup>  
240 and a disease-specific PROM for CAS<sup>19</sup> which was designed for use in the SAPPHERE trial  
241 (Stenting and Angioplasty with Protection in Patients at High Risk for Endarterectomy)<sup>19</sup> and  
242 was further adapted and used in the CREST study (Carotid Revascularization Endarterectomy  
243 versus Stenting Trial).<sup>21</sup>

244



245 The timings of administering the PROMs were different between the five studies. The  
 246 shortest post intervention follow-up was two days and the longest was 24 months. The rigour  
 247 of the psychometric assessment of the PROMs was variable, with most only attempting to  
 248 assess a single psychometric criterion. The evaluation was generally poor across all the  
 249 included studies in this review. The results of the psychometric evaluation are presented in  
 250 Table IV. In brief, the quality of each psychometric criterion was based on: 1) using the  
 251 appropriate statistical test for a specific criterion and 2) the results of the test fulfilled the  
 252 criteria mentioned in the methods section and Table II. Each criterion was evaluated  
 253 independently and objectively by two independent reviewers.

254

255 **Table IV: Summary of the psychometric properties of patient-reported outcome**  
 256 **measures**

PROM	Internal consistency	Test re-test Reliability	Content validity	Construct validity	Responsiveness	Floor/ceiling	Acceptability
<b>SF-36</b>							
Cohen 2011 <sup>21</sup>	0	0	0	0	+/-	0	+
Stolker 2010 <sup>19</sup>	0	0	-	0	+/-	0	+
Hsu 2014 <sup>22</sup>	?	0	0	0	0	0	0
<b>EQ-5D</b>							
Stolker 2010 <sup>19</sup>	0	0	-	0	+/-	0	+

Attigah	0	0	0	0	?	0	0
2011 <sup>20</sup>							

#### Disease-Specific PROM

Cohen	0	0	-	0	?	0	+
2011 <sup>21</sup>							
Stolker	0	0	-	0	?	0	+
2011 <sup>19</sup>							

#### Quality of Life for Carotid Artery Disease

Ivanova	0	0	+/-	0	-	-	0
2015 <sup>23</sup>							

#### DHI

Hsu	0	0	0	0	?	0	0
2014 <sup>22</sup>							

#### HADS

Attigah	0	0	0	0	?	0	0
2011 <sup>20</sup>							

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DHI, Dizziness Handicap Impact; EQ-5D, EuroQol 5 dimensions; HADS, hospital anxiety and depression scale; PROMs, patient reported outcome measures; SF-36, 36-item Short Form Medical Outcomes Study

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#### Psychometric and operational criteria:

0	Not reported (no evaluation completed)
-	Evidence not in favour
-/+	Weak evidence
+	Evidence in favour
?	Methodology questionable

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257 The SAPPHERE trial<sup>19</sup> included high-risk patients with symptomatic carotid stenosis of >50%  
258 and patients with asymptomatic CAS with >80% stenosis. Patients were randomised to either  
259 the CS arm (159 patients) or the CEA arm (151 patients). HRQoL was assessed at baseline,  
260 two weeks and one, six and 12 months using SF-36, EQ-5D and a disease-specific PROM  
261 with six questions asking about difficulty with walking, eating/swallowing, driving,  
262 headache, neck pain and leg pain. The study did not report any qualitative evidence to  
263 support the content validity of the disease-specific PROM. Only four- subscales of the SF-36  
264 were used (physical function, role limitations, pain, vitality), the authors justified this  
265 decision that only these four dimensions were sensitive to differences between CS versus  
266 CEA and provided no further evidence. However, data on three of these subscales were not  
267 sensitive at all and did not show any statistically significant change from baseline, only the  
268 physical scale of SF-36 showed some responsiveness at two weeks. The disease-specific  
269 PROM in this study did not undergo further psychometric analysis to assess its  
270 responsiveness. The strongest feature of PROMs used in this study was acceptability with  
271 data completeness being above 80%.

272

273 The CREST trial<sup>21</sup> included data from 2,502 patients with symptomatic and asymptomatic  
274 CAS. 1,262 patients were assigned to CS and 1,240 to the CEA arm of the trial. HRQoL was  
275 assessed at baseline, two weeks, one month and one year post intervention using SF-36 and  
276 an adapted version of the disease-specific PROM from the SAPPHERE trial.<sup>19</sup> The disease-  
277 specific PROM included eight questions (including difficulty in walking, difficulty in  
278 swallowing/eating, driving, neck pain, headaches, leg pain, level of overall pain and the  
279 number of times pain medications were needed). No qualitative evidence for content validity,  
280 internal consistency and reliability of either instrument was provided. However, both  
281 instruments showed good acceptability with data completion rates of 85 to 90%. The SF-36

282 scores improved across five out of eight dimensions of health (P value < 0.01) at two weeks  
283 for patients undergoing CS versus CEA.

284

285 Attigh et al<sup>20</sup> assessed HRQoL in 102 patients undergoing CEA under local anaesthetic. The  
286 SF-36 and HADS were used to assess HRQoL. Evidence on validity, reliability, acceptability  
287 and consistency were not reported for either PROM. The psychometric evaluation only  
288 concentrated on responsiveness using univariate comparisons and multivariate analysis,  
289 neither of which was suitable for assessing the responsiveness of the PROMs.

290 The CAS specific PROM was developed by Ivanova et al.<sup>23</sup> The initial version was based on  
291 generic and neurovascular specific HRQoL questionnaires. This was reviewed by patients  
292 with CAS and clinicians. The final draft included 17 domains each with four choices. The  
293 PROM was assessed in 120 patients with asymptomatic CAS, one to three days before CEA  
294 and six to seven months after that. The authors reported improved physical, functional,  
295 psychological and social function but these were not statistically significant. Furthermore,  
296 many domains had floor/ceiling effects of more than 28.5% raising questions regarding the  
297 relevance of some of the questions included in this PROM.

298

299 Hsu et al<sup>22</sup> assessed the effect of CS on HRQoL in patients with CAS suffering with  
300 dizziness. Of the 178 patients who underwent CS, only 61 complained of dizziness. HRQoL  
301 was assessed using SF-36 and DHI. The SF-36 showed evidence of internal consistency  
302 (Cronbach's alpha score >0.70) but the statistical assessment of responsiveness was based on  
303 non-parametric measures and no evidence was presented regarding the completeness of the  
304 data for each of the domains.

305

#### 4. Discussion

This review identified six PROMs in five studies<sup>19-23</sup> that reported details on the development and/or validation of PROMs for use in patients with CAS undergoing revascularisation. The quality of the instruments was variable with respect to their development and psychometric properties. None of the identified PROMs had undergone rigorous psychometric validation in patients undergoing carotid artery revascularisation. Validation of basic psychometric criteria such as construct validity and test-retest reliability had not been undertaken. Only one study, Hsu et al<sup>22</sup> attempted to assess the internal consistency of SF-36 although the methodology they used was questionable. Based on the findings of our review it is not possible to recommend a PROM for use in patients with CAS undergoing revascularisation.

The strength of the review lies on our comprehensive and extensive search strategy which was used to identify relevant studies. In addition, to minimise bias two reviewers undertook the screening, data coding, data extraction and psychometric analysis of all the studies, and the review covered all types of study designs. The methodological quality assessment criteria were developed from published studies as per FDA PROMs development guidance.<sup>2</sup> However, there are a number of limitations to our review which warrant caution to its application. The patient population included in this systematic review were heterogeneous in terms of the type of CAS, the stage of disease, and treatment pathway. For example, the Quality of life for carotid artery disease scale, reported by Ivanova et al<sup>23</sup> was developed in a Latvian population and the PROMs reported in Hsu et al (DHI and SF-36)<sup>22</sup> underwent validation in a Chinese population. As a result, the application of the findings from these studies to English speaking people is uncertain due to language validation and cross-cultural adaptation of PROMs.<sup>11</sup> It is important to note that these limitations are principally sourced in the evidence base, rather than the methods used to interrogate and evaluate it.

331

332 It is recommended that PROMs data is collected and evaluated as part of randomised  
333 controlled trials (RCTs) and service analysis.<sup>3;24-26</sup> Evidence from this review shows that  
334 most PROMs used in previous carotid trials lacked validation. Another tool occasionally  
335 used to assess functional HRQOL outcomes following CS or CEA in clinical trials is the  
336 modified Rankin scale<sup>27</sup> (a functional assessment scale for assessing handicap in stroke  
337 patients).<sup>28</sup> However, the Rankin score was not included in this review as it does not capture a  
338 patient's subjective perception of their QoL, and thus cannot be considered to be a true  
339 PROM.<sup>29</sup> The benefits of supplementing clinical outcome data with a well-developed, valid,  
340 consistent, reliable and responsive instrument could help provide more targeted data on  
341 aspects such as how patients feel after specific interventions, treatment efficacy, and  
342 identification of patients most likely to benefit from the procedure. Particularly since the  
343 intervention procedure is frequently done in patients who might be asymptomatic. Hence,  
344 having a universal accepted PROM measure for assessing QoL in patients undergoing carotid  
345 revascularisation will be valuable to the patients, clinicians and decision makers to guide  
346 them in providing an efficient and cost-effective treatment plan.

347

348 Some of the issues noted in this review maybe addressed by either developing a disease-  
349 specific PROM or developing a set of questions specific to CAS which can be added to  
350 complement a generic PROM (e.g. SF-36 or EQ-5D) as recommended by regulating bodies.<sup>6</sup>  
351 However, when developing a PROM questionnaire it is important to use qualitative methods  
352 involving patients and clinicians and insure the questionnaire captures both the breadth of the  
353 patient experience and the instrument to be reliable, valid, responsive and acceptable to  
354 patients. The questionnaire should be easy to administer and attention should be given to its  
355 format, setting and time required for completion. In addition, research exploring how to

356 integrate PROMs into the patient pathway needs to be undertaken, including when and at  
357 what time-points should the PROM be administered.

358

## 359 **5. Conclusion**

360 This review highlights a lack of evidence for valid, reliable, responsive and acceptable  
361 PROMs for use in patients undergoing carotid artery revascularisation. As a result, the  
362 development and validation of a new PROM for this patient population is warranted in order  
363 to provide data which can supplement traditional clinical outcomes (stroke<30 days post-  
364 procedural, myocardial infarction and death) and capture changes in health status and quality  
365 of life to help inform treatment decisions.

366

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## 373 **Conflicts of interest**

374 The authors declare no conflicts of interest.

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