Quality of life in restorative *versus* non-restorative resections for rectal cancer: systematic review

Samuel Lawday (b) 1.2.3.*, Nicholas Flamey¹, George E. Fowler (b) 1.2, Matthew Leaning¹, Nadine Dyar¹, Ian R. Daniels (b) 1, Neil J. Smart^{1,3} and Christopher Hyde³

*Correspondence to: Samuel Lawday, Bristol Centre for Surgical Research, Canynge Hall, 58 Whiteladies Road, Clifton, Bristol BS8 2PL, UK (e-mail: samuel.lawday@bristol.ac.uk)

Abstract

Background: Low rectal cancers could be treated using restorative (anterior resection, AR) or non-restorative procedures with an end/permanent stoma (Hartmann's, HE; or abdominoperineal excision, APE). Although the surgical choice is determined by tumour and patient factors, quality of life (QoL) will also influence the patient's future beyond cancer. This systematic review of the literature compared postoperative QoL between the restorative and non-restorative techniques using validated measurement tools

Methods: The review was registered on PROSPERO (CRD42020131492). Embase and MEDLINE, along with grey literature and trials websites, were searched comprehensively for papers published since 2012. Inclusion criteria were original research in an adult population with rectal cancer that reported QoL using a validated tool, including the European Organization for Research and Treatment of Cancer QLQ-CR30, QLQ-CR29, and QLQ-CR38. Studies were included if they compared AR with APE (or HE), independent of study design. Risk of bias was assessed using the Risk Of Bias In Non-Randomized Studies of Interventions (ROBINS-I) tool. Outcomes of interest were: QoL, pain, gastrointestinal (GI) symptoms (stool frequency, flatulence, diarrhoea and constipation), and body image.

Results: Nineteen studies met the inclusion criteria with a total of 6453 patients; all papers were observational and just four included preoperative evaluations. There was no identifiable difference in global QoL and pain between the two surgical techniques. Reported results regarding GI symptoms and body image documented similar findings. The ROBINS-I tool highlighted a significant risk of bias across the studies.

Conclusion: Currently, it is not possible to draw a firm conclusion on postoperative QoL, pain, GI symptoms, and body image following restorative or non-restorative surgery. The included studies were generally of poor quality, lacked preoperative evaluations, and showed considerable bias in the data.

Introduction

The treatment for rectal cancer has changed significantly over the past 20 years with the introduction of MRI-based, multidisciplinary team-directed, individualized patient care and the selective use of neoadjuvant therapies¹. For most patients with rectal cancer, surgery continues to be mainstay of curative treatment. Surgical techniques are based on total mesorectal excision (TME), and comprise either a restorative rectal resection (anterior resection, AR) with anastomosis or an excisional rectal technique with an end/permanent stoma (Hartmann's, HE; or abdominoperineal excision, APE)². Over the past 100 years, debate has existed regarding which surgical technique (restorative or excisional) provides the best outcome for the patient, with trends towards restorative surgery³. It is recognized that each operative approach is different and both techniques are not suitable for all patients⁴. Tumour stage, morphology, and

clinical presentation all influence patient outcomes, and identifying the true impact of individual operations can be difficult. Clinical research, systematic reviews, and meta-analyses of rectal cancer outcomes have concentrated on the technical elements and technology used to perform the procedure. These include reviews on robotic versus laparoscopic^{5,6}, open versus laparoscopic^{7,8}, and transanal TME ^{9,10} surgery, most of which focused on demonstrating surgical and oncological equivalence or cost–benefit of the procedural approach. There has been little focus on comparing quality of life (QoL) or patient-reported outcome measures (PROMs).

A number of tools, such as EQ-5DTM (EuroQol Group, Rotterdam, the Netherlands), Short Form 36, Functional Assessment of Cancer Therapy—Colorectal (FACT-C), and European Organization for Research and Treatment of Cancer (EORTC) QLQ-CR29, QLQ-CR30, and QLQ-CR38, have all been validated to accurately reflect changes in patients' QoL^{11–15}.

¹HeSRU, Royal Devon and Exeter Hospital, Exeter, UK

²Bristol Centre for Surgical Research, University of Bristol, Bristol, UK

³College of Medicine and Health, University of Exeter, Exeter, UK

Some of these tools are generic and global; EQ-5DTM measures overall health status and is applicable in any condition. EORTC QLQ-CR29 and QLQ-CR30 questionnaires are global QoL tools specific for colorectal cancer. These tools have been shown offer validity and reliability in population groups to ensure that the results reflect true patient experience. A paucity of high-quality comparative PROM and QoL data following rectal cancer surgery to aid in patients' decision-making between surgical options was reported in 2012¹⁶. Comparison of these two surgical techniques is challenging, and there is currently no one PROM that can aid this. Indeed, social interaction, body image, and overall QoL influence the patient's future in living with and beyond cancer¹⁷ and, when obtaining informed consent, patients should be advised objectively about the treatment options available 18. This systematic review of the literature on low rectal cancer compared restorative with non-restorative resection, focusing on validated QoL measures.

Methods

This systematic review was registered on PROSPERO (CRD42020131492), and complies with PRISMA and AMSTAR guidelines 19,20. Using a PICO search, the population of interest comprised patients with low rectal cancer undergoing an intervention of AR or restorative bowel resection compared with those who had APE or non-restorative bowel resection, with an outcome measured using a validated QoL tool. The review question was established a priori, with inclusion and exclusion criteria, and the risk-of-bias tool chosen before completion of the search. A comprehensive search of Embase and MEDLINE was completed. The search strategy is available in supplementary material (Appendix S1). This built on the published Cochrane systematic review¹⁶, using only papers published from this date. References of included articles were screened for suitable papers. Grey literature was searched in the British Library Thesis repository and Grey Literature search engine. ISRCTN and ClinicalTrials.gov were screened for suitable trials. The search was completed on 20 June 2020. Published data comparing validated QoL outcomes in adults undergoing radical surgery for rectal cancer were included. Transabdominal surgical techniques with curative intent were included. There was no limit based on follow-up time. Randomized and observational studies were included, although randomization between the two groups was thought to be unlikely.

Inclusion criteria were: surgical intervention for rectal cancer in adults aged over 18 years; surgery performed with curative intent; comparison of different surgical techniques (without restriction by study design)—AR (restorative resection with or without temporary ileostomy) versus APE/TME or Hartmann's (non-restorative resection with permanent/end stoma); QoL data provided by means of a validated tool; and published since last Cochrane review in 2012¹⁶.

Exclusion criteria were: lack of specific rectal cancer data (mixed data with colonic cancers but no subgroup data provided); unresectable disease or palliative surgery; local excision techniques; inclusion of surgery for inflammatory or benign growth; and not available in the English language 21,22 .

Data analysis

Titles and abstracts of each article were screened before the whole paper being requested. Included papers underwent review.

Patients who had undergone resection with bowel continuity restored were included in the AR group. The inclusion of patients with a temporary ileostomy vaired between papers, but this group included patients with a temporary ileostomy and those who had the ileostomy reversed. The APE group included all patients who had undergone non-restorative resection, including APE and HE.

Authors of articles with data presented in graphical form were contacted in an attempt to obtain numerical data; if there was no response from the corresponding author, the data were included in a summative analysis but not in the tables.

One reviewer extracted data into an electronic data collection sheet. A second independent reviewer checked this, with discrepancies resolved by consensus. Data extracted included: study-related data (first author, year of publication, journal, study design, duration of follow-up, outcomes measured, funding), patient characteristics (surgical technique, tumour site, neoadjuvant therapy) and outcome data (validated QoL measure results). Risk of bias was assessed using the Risk Of Bias In Non-Randomized Studies of Interventions (ROBINS-I) tool²³; differences were resolved by consensus.

Narrative summary and qualitative analysis were planned, with comparison of the results between studies both during short- and long-term follow-up. Quantitative analysis through meta-analysis was considered inappropriate owing to clinical heterogeneity in interventions, the non-normal distribution of results, study design, and the variety of validated QoL outcome measures used.

Outcome measures

The primary measure was difference in average global QoL. Other health-related items investigated were: gastrointestinal (GI) symptoms (stool frequency, flatulence, diarrhoea, and constipation), pain, and body image.

Results

Of 21 074 abstracts screened, 76 full papers were scrutinized and 19 included in the final review (Fig. 1) $^{24-42}$. Nineteen studies met the inclusion criteria (Table 1) with a total of 6453 patients (range 43-1608). All articles described observational studies, although one study 37 included patients from the National Surgical Adjuvant Breast and Bowel Project randomized trial (NSABP-R-04) from the USA. No patients were randomized between surgical techniques. Patient follow-up varied from 6 months to 5 years. Only four^{30,34,35,37} of the included studies provided preoperative QoL data and then followed patients up; two other papers^{26,41} provided serial QoL measurements, but not preoperative data. Thirteen studies provided only one measure of postoperative QoL, with no preoperative data. Fourteen studies used QLQ-CR30, eight used QLQ-CR29, and seven used QLQ-CR38 (Table 1). All studies compared outcomes for patients with rectal cancer; seven studies considered only rectal cancer within 4-6 cm of the anal verge, although it was not always stated how this was measured (Table S1). Surgical approaches and main findings are summarized in Table 2. Quantitative analysis was not completed because of the skewed data distribution, variety of QoL tools used, use of median (range), and the lack of standard deviation reporting⁴³.

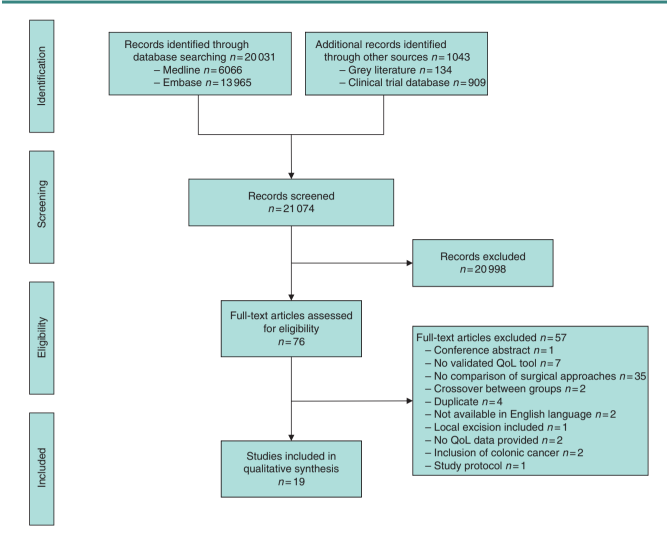


Fig. 1 PRISMA diagram showing selection of articles for review QoL, quality of life.

Global quality of life

Only two studies included in the review identified a statistically significant difference in global QoL between restorative/AR and non-restorative/APE surgery. A single-centre study³⁵ identified better QoL in patients who had undergone AR with colonic pouch formation at 13 months (P = 0.009), although no preoperative data were available. A population-based, cross-sectional study²⁷ identified better global OoL in patients who had undergone APE (P = 0.026) at a median of 4.4 years after surgery, but again there were no preoperative QoL data. Extracted global QoL data are reported in Table 3. A difference of 10 on the EORTC 1-100 scale was used to compare the two surgical approaches; with the exception of one study³⁵, no difference in global QoL between the two groups was noted^{28,29,32,37}. Direction-of-effect analysis (based on whether a score at any time point after baseline favours AR or APR) found that four studies demonstrated better global QoL in patients who had undergone APE, four favoured AR, and three identified no difference. No link was identified when patients were separated by length of follow-up.

Pain

Extracted data for the pain domain of the validated QoL scores are presented in Table 4. Two studies identified a statistically significant reduction in pain after APE among patients who had follow-up longer than 2 years. Other studies with a clinically relevant difference on long-term follow-up also demonstrated reduced pain in the APE group. In long-term direction-of-effect analysis, five of the seven studies found reduced pain in the APE group. One study⁴⁰ demonstrated increased pain in the APE group during follow-up of less than 12 months, but presented no preoperative data. No other studies identified a statistically significant or clinically relevant difference between the two groups. Direction-of-effect analysis showed that three studies identified no difference, although two with follow-up of less than 12 months favoured AR. Others^{29,31,32} with no numerical data available also showed no difference in postoperative pain. In one study²⁸, multiple logistic regression showed that both AR (odds ratio (OR) 1.39, 95 per cent c.i. 1.01 to 1.90) and APE (OR 1.71, 1.19 to 2.44) were associated with chronic pelvic pain at a median of 4.2 years of compared with partial mesorectal excision.

Gastrointestinal symptoms

GI symptoms were measured using a variety of tools (EORTC QLQ-CR30, QLQ-CR30, QLQ-CR30) alongside the Faecal Incontinence Quality of Life Scale and Wexner scale after restorative surgery^{44,45} (Table 5). The use of specific tools to

Table 1 Details of included studies

Reference	Year	Setting	Country	Research design	Duration of follow-up	No. of patients	QoL measure used
1. Mrak et al. ³⁵	2011	Single centre	Austria	Observational, prospectively maintained database	Minimum 3 years	59	EORTC QLQ-C30, EORTC QLQ CR29
2. How et al. ³⁰	2012	Single centre	UK	Observational, prospective	2 years	62	EORTC QLQ-C30, EORTC QLQ-CR38
3. Konanz et al. ³³	2013	Single centre, university- affiliated hospital	Germany	Observational, prospective database	Minimum 12 months	124	EORTC QLQ-C30, EORTC QLQ-CR38
4. Digennaro et al. ⁴²	2013	Multicentre	Italy	Observational, retrospective	Median 26.5 months (APE), 52.5 months (AR)	60	EORTC QLQ-C30, EORTC QLQ- CR38, Short Form 36
5. Arraras et al. ²⁴	2013	Single centre	Spain	Observational, prospective	Minimum 12 months	84	EORTC QLQ-C30, EORTC QLQ-CR29
6. Penchev et al. ³⁶	2014	Single centre, complex cancer centre	Bulgaria		Minimum 6 months	71	EORTC QLQ-CR38: sexual function
7. Russell et al. ³⁷	2015	Multicentre	USA	Observational, patients recruited to chemotherapy RCT	12 months	1608	FACT-C, EORTC QLQ-CR38
8. Feddern et al. ²⁸	2015	Population database	Denmark	Observational, cross-sec- tional survey	Minimum 4.2 years	1369	Brief Descriptive, Danish Pain Questionnaire (McKill)
9. Honda et al. ²⁹	2016	Single centre, cancer institute hospital	Japan	Observational, cross-sec- tional survey	Minimum 2 years	291	EORTC QLQ-C30, EORTC QLQ- CR29, modified FIQL
10. Monastyrska et al. ³⁴	2016	Single centre, oncology centre	Poland	Observational, prospective	6 months	100	EORTC QLQ-C30, EORTC QLQ-CR29
11. Klose <i>et al</i> . ³¹	2017	Single centre, university- affiliated hospital	Germany	Observational, prospectively maintained database	58 months	143	EORTC QLQ-C30, EORTC QLQ-CR29
12. Wani et al. ⁴⁰	2017	Single centre, Kashmir	India	Observational, prospective	12 months	130	EORTC QLQ-C30, EORTC QLQ-CR29
13. Costa et al. ²⁵	2018	Single centre	Portugal	Observational, retrospective	21 months	43	International Index of Erectile Function
14. Koeter et al. ³²	2018	Population database	the Netherlands	Observational, longitudinal, prospective population- based survey	5.1 years	905	EORTC QLQ-C30, EORTC QLQ-CR38
15. Trenti et al. ³⁹	2018	Two centres	Spain	Observational, prospective	4.5 years	224	EORTC QLQ-C30, EORTC QLQ-CR29
16. Silva et al. ³⁸	2018	Single centre	Brazil	Observational, retrospective	3.84 years	125	EORTC QLQ-CR29 EORTC QLQ-CR29
17. Du et al. ²⁶	2019	Single centre	China	Observational, retrospective	12 months after surgery	43	EORTC QLQ-C30, EORTC QLQ-CR38
18. Feddern et al. ²⁷	2019	Single centre	China	Observational, cross-sec-	Median 4.4 years	898	EORTC QLQ-C30
19. Ding et al. ⁴¹	2020	Single centre	China	tional survey Observational, prospective	12 months	114	FIQL

Funding: 1. Not funded and no relationships to declare; 2. Pelican Cancer Foundation, Basingstoke; Colorectal Research Unit, North Hampshire Hospital, Basingstoke; 3. Not stated; 4. Not stated; 5. Grant 2443/2009 from Departamento de Salud del Gobierno de Navarra (Navarre Government Health Department), Spain; 6. Not stated; 7. NCI-U10-CA-12027, U10-CA-37377, U10-CA-69974, U10-CA-69651, and U10-CA-21661, by Roche; 8. Funding from private foundation of Daehnfeldt; 9. Japanese Foundation for Research and Promotion of Endoscopy; 10. Not stated; 11. Not stated; 12. Not stated; 14. Data collection funded by a VENI Grant (no. 45110-041) from Netherlands Organization for Scientific Research awarded to F. Mols, together with a Medium Investment Grant from the Netherlands Organization for Scientific Research (NWO no. 480-08-009). D. Schoormans supported by a Social Psychology Fellowship from the Dutch Cancer Society (no. UVT2013-5893); 15. Not stated; 16. Not stated; 17. Grants from scientific research project of Gansu health industry (GSWST2013-03); 18. Danish Cancer Society; 19. Hunan Provincial Nature Science Foundation (2016JJ2106), Hunan Provincial Nature Science Foundation (2019J340510).

Table 2 Surgical intervention and study conclusions

Reference	Surgical in	tervention	Conclusions/recommendations			
	AR (restorative)	APE (non-restorative)	Global QoL	Symptoms		
Mrak et al. ³⁵	Ultralow TME anterior resection and colonic J-pouch anastomosis	APE and end colostomy	QoL better after AR than APE in several respects	After AR, patients had better physical, role, cognitive, and social functioning with better body image. After APE, patients had significantly higher urinary frequency and were significantly more embarrassed by their condition		
How et al. ³⁰	AR	APE and end colostomy	No difference in global QoL	There was more diarrhoea after AR and more pain at 2 years after operation. Better sexual and social func- tioning after AR		
Konanz et al. ³³	ISR or LAR	APE and end colostomy	No difference in global QoL	Physical functioning significantly better after AR. Symptom scores for diarrhoea and constipation worse after AR		
Digennaro et al. ⁴²	CAA (sewn and stapled)	APE and end colostomy	No difference in global QoL	APE group had worse sexual function, whereas most patients in AR group had faecal incontinence and sometimes obstructed defaecation, with an important impact on		
Arraras et al. ²⁴	LAR (with colorectal anastomosis or CAA)	APE and end colostomy	No difference in global QoL	QoL Higher stool frequency and incontinence in AR group, but better emotional functioning		
Penchev et al. ³⁶	AR	APE and end colostomy	Not compared	Sexual dysfunction worse in men after APE than AR		
Russell et al. ³⁷	Sphincter-sparing surgery	Non-sphincter-sparing surgery	No difference in global QoL	AR group had better body image, male sexual enjoyment, and micturition symptoms. APR group had better GI tract symptoms and less weight loss. No difference in FACT-C		
Feddern et al. ²⁸	AR (with TME or PME)	APE; Hartmann's in- cluded but separate	Not compared	No association between pain intensity and type of surgery		
Honda et al. ²⁹	Very low AR or ISR	APE and end colostomy	No difference in global QoL	Worse constituetion, defaecation problems and anxiety in ISR group		
Monastyrska et al. ³⁴	LAR (without stoma)	APE and end colostomy	No difference in global QoL	Physical, cognitive, and emotional functioning better in AR group		
Klose et al. ³¹	ISR	APE and end colostomy	No difference in global QoL	ISR group had better cognitive functioning and weight gain, and less nausea and		

(continued)

Table 2. (continued)

Reference	Surgical in	tervention	Conclusions/recommendations			
	AR (restorative)	APE (non-restorative)	Global QoL	Symptoms		
				vomiting, pain, dyspnoea, appetite loss, and embarassment. APE group had less diarrhoea, stool frequency, buttock		
				pain, bloating, sore skin, and faecal incontinence		
Wani et al. ⁴⁰	LAR	APE and end colostomy	No difference in global QoL	Nausea and vomiting worse in AR group, but urinary frequency, abdominal pain and embarassment worse in APE group		
Costa et al. ²⁵	AR	APE and end colostomy	Not compared	APE and not AR is a risk factor for de novo ED		
Koeter et al. ³²	LAR	APE and end colostomy	Not compared	No differences in physical activity between the two groups. Physical and role functioning seemed worse in APE group		
Trenti et al. ³⁹	AR	APE and Hartmann's	No difference in global QoL	Faecal incontinence worse in AR group and body image worse in		
Silva et al. ³⁸	Sphincter-saving surgery with closure of temporary ileostomy	APE and end colostomy	No difference in global QoL	APR group APE group had significantly better functional and		
Du et al. ²⁶	AR with anal reconstruction	APE and end colostomy	No difference in global QoL	symptom scale scores Emotional and social functioning better in		
Feddern et al. ²⁷	LAR	APE and end colostomy	Global health status worse in AR group	AR group AR group had worse problems with diarrhoea and constipation		
Ding et al. ⁴¹	Ultralow AR (Dixon) and modified CAA (modified Parks)	Miles APE and end colostomy	Not compared	At 12 months, AR group had better scores in all four criteria of FIQL score		

AR, anterior resection; APE, abdominoperineal excision; QoL, quality of life; TME, total mesorectal excision; ISR, intersphincteric resection; LAR, low anterior resection; CAA, coloanal anastomosis; GI, gastrointestinal; FACT-C, Functional Assessment of Cancer Therapy—Colorectal; PME, partial mesorectal excision; ED, Erectlie Dysfunction; FIQL, Faecal Incontinence Quality of Life Scale.

compare the two groups is challenging because of the difference in symptoms experienced by patients in the AR and APE groups. Long-term follow-up of greater than 2 years demonstrated favourable outcomes for APE over AR. In the domains of stool frequency, flatulence, GI symptoms, diarrhoea, and constipation, patients in the AR group had worse symptoms than those in the APE group. Only one study³⁴, at 6 months but with no preoperative comparator, demonstrated better outcomes in the AR group; all the other studies either showed no difference or reported better outcomes in patients who had undergone APE.

Body image and sexual function

Five studies 26,32,35,37,39 identified higher rates of negative body image in the APE group than the AR group, although preoperative data were not available. The other studies reported no difference; none reported better body image in the APE group (Table 6). Sexual function was worse in the APE group, but most studies

that measured this did not have preoperative data (Table 7). Nine papers^{24–26,30,33,35,37,38,40} reported worse sexual functioning and/ or interest in the APE group; five^{29,31,32,34,39} identified no difference between the two groups. One study³⁴ identified worse functioning in the APE group, but this difference was present before operation and may reflect a difference in patient and tumour characteristics.

Risk of bias

The ROBINS-I risk-of-bias assessment was completed for all studies. All bar one had at least a low/moderate risk of bias. Twelve had a serious risk of bias in at least one domain and five had a critical risk of bias in at least one domain (Fig. 2). Two had a low risk of bias in more than six domains, both of which found no difference in global QoL. The reason for high risk of bias varied between studies. Recurring themes included the nonreporting of patient characteristics including co-morbidities, different disease profiles, different preoperative chemoradiotherapy

Table 3 Global quality-of-life measure using EORTC QLQ-CR30

	n	Surgical	Global QoL score*					Direction of effect
		procedure	Baseline	0–5 months	6–11 months	1–2 years	> 2 years	
How et al. ³⁰ †	62	APE AR	83 (33–100) 79 (17–100)			79 (33–100) 71 (33–100)	83 (17–100) 75 (33–100)	Trend favours APE but n.s.
Mrak et al. ³⁵	59	APE AR	,3 (1, 100)			, 1 (33 100)	60.4 (20.1) 75.7 (20.1)	Favours AR (P < 0.050)‡
Konanz et al. ³³	124	APE AR AR (ISR)					59.2 65.9 58.1	Trend favours AR but n.s.
Monastyrska et al. ³⁴	100	APE AR	51.7 61.3			60.5 69	30.1	Trend favours AR but n.s.
Wani et al. ⁴⁰	130	APE AR	01.5			67.9(21.2) 59.3(23.7)		Trend favours APE but n.s.
Arraras et al. ²⁴	84	APE AR				71.8(25.7) 70.9(28.0)		Trend favours APE but n.s.
Du et al. ²⁶	43	APE AR		69.0(6.3) 74.3(7.9)	69.4(6.4) 75.4(8.9)	70.8(10.9) 75.9(8.9)		Trend favours AR but n.s.
Trenti et al. ³⁹	224	APE AR AR (CAA)		(:)	()	()	67.3 (21.4) 69.8 (24.6) 65.6 (23.4)	No difference
Silva et al. ³⁸ †	125	APE AR					75 (0–100) 75 (0–100)	No difference
Digennaro et al. 42†	60	APE AR					66.6 (50–100) 66.6 (16.7–100)	No difference

*Values are mean(s.d.) unless indicated otherwise; †values are median (range). The European Organization for Research and Treatment of Cancer (EORTC) QLQ-CR30 score has a range of 1–100, where 0 represents the best quality of life attainable. A score difference or change of 10 is claimed to be clinically important. Values are rounded to one decimal place. Articles with data represented graphically are not included in this table. *P < 0.050 was considered statistically significant. QoL, quality of life; APE, abdominoperineal excision; AR, anterior resection; n.s., not statistically significant; ISR, intersphincteric resection; CAA, coloanal anastomosis.

regimens between the groups, and variation in questionnaire completion.

Discussion

Overall, there was no improvement in global QoL across the studies; restorative surgery was not found to improve QoL compared with a permanent stoma. However, some caution is required in interpretation of the research because published data were at significant risk of bias and no high-quality papers existed to allow accurate analysis of the difference in QoL. Different symptom profiles were identified; when studies reported differences of statistical value, patients who underwent APE had worse body image and those who underwent AR had worse GI symptoms.

The ROBINS-I assessment of all the studies reflected this high risk of bias and demonstrated the paucity of good-quality studies aimed at assessing this clinically relevant question. This should be considered when assessing the conclusions of the review. The risk of bias was contributed to by the paucity of preoperative QoL data, the use of single-point QoL scores, and failure to control for the location of the rectal cancer. The distance of the cancer from the anal verge is paramount in deciding surgical and neoadjuvant treatments, and therefore has a significant impact on postoperative outcomes. Two studies included in the review had a low risk of bias in 6 or more domains. One report found no difference in global QoL, but reported better cognitive and social functioning with fewer symptoms of pain, diarrhoea, sleep disturbance, and constipation in patients who had undergone APE. Another documented no difference in global QoL between the two surgical techniques; however, no raw data were included in the publication

and it was not therefore included in any Tables. The same authors identified worse sexual function and micturition symptoms in patients who had undergone APE, but they had better GI symptom profiles.

Postoperative differences in QoL measures may have been present before surgery and therefore cannot be explained simply by differences in surgical technique. The importance of ensuring that the disease profile is matched should be highlighted; controlling for tumour height and neoadjuvant therapy is important as these have an impact on patient QoL. This may explain the results of many of the studies included in this review. The lack of preoperative QoL measures to identify any possible differences being caused by variation in surgical indication rather than surgical technique introduces significant potential bias into most studies included in the review. It is therefore challenging to identify whether disease location, preoperative differences in QoL or operative approach is the reason for a difference in postoperative QoL. Variation in duration of follow-up may reflect different aspects of the patient journey. Long-term follow-up may miss significant shortterm variation in QoL and will miss patients with short postoperative survival; however, longer follow-up allows good assessment of function. The variability of follow-up and grouping of patients across these time brackets may mean a mixed picture is provided across the included studies. Many studies included patients with a large range of follow-up times and it is therefore difficult to draw specific conclusions regarding changes in QoL over time.

The high search volume reflects the large amount of work being done regarding QoL outcomes after rectal cancer surgery. Many articles were excluded, as they did not offer a comparison between restorative and non-restorative rectal procedures.

Table 4 Validated measures of pain

	Surgical procedure			Pain score*			Direction of effect
	procedure	Baseline	0-5 months	6–11 months	1–2 years	> 2 years	enect
QLQ-CR30: pain							
How et al. ³⁰ †	APE	0 (0-67)			0 (0-67)	0 (0-33)	Favours APE
·	AR	0 (Ò-10Ó)			17 (0–89)	33 (0–67)	(P < 0.050)‡
Mrak et al. ³⁵	APE	,			,	24.4	Trend favours
	AR					17.5	AR but n.s.
Konanz et al. ³³	APE					25.3	Trend favours
	AR					17.5	AR but n.s.
	AR (ISR)					22.7	
Monastyrska et al. ³⁴	APE	27			2		Trend favours
Wioriabeyrbika ee al.	AR	23			9		APE but n.s.
Wani et al. 40	APE	23			18.5(21.9)		Trend favours
wani ci ai.	AR				26.3(29.9)		APE but n.s.
Arraras et al. ²⁴	APE				23.8(26.1)		Trend favours
Miaias et ui.	AR				17.9(25.5)		AR but n.s.
Du et al. ²⁶	APE		10.3(4.3)	10.0(4.0)	9.1 (4.7)		No difference
Du et al.	AR		10.1(4.7)	10.6(4.5)	9.8(5.8)		No difference
Trenti et al. ³⁹	APE		10.1(4.7)	10.0(4.3)	3.0(3.0)	10 1 (01 6)	Trend favours
Henti et al.	AR					12.1 (21.6)	APE but n.s.
						13.5 (20.9)	APE DUI II.S.
C:l at al 38±	AR (CAA)					14.9 (21.1)	Tuon d formanno
Silva et al. ³⁸ †	APE					0 (0–100)	Trend favours
010 0000 11 11	. AR					16.7 (0–100)	APE but n.s.
QLQ-CR29: abdominal pa				00.4(05.4)			T 16
Wani et al. ⁴⁰	APE			32.1(26.4)			Trend favours
- 24	AR			9.3(22.2)			AR but n.s.
Arraras et al. ²⁴	APE			12.7(24.7)			No difference
00	AR			12.5(21.6)			
Silva et al. ³⁸ †	APE					0 (0–66.7)	No difference
	AR					0 (0–100)	
QLQ-CR29: buttock pain							
Wani et al. ⁴⁰	APE			25.0(28.1)			Trend favours
	AR			12.1(18.0)			AR but n.s.
Arraras et al. ²⁴	APE			15.9(27.2)			No difference
	AR			16.7(24.6)			
Trenti et al. ³⁹	APE			. ,		11.7 (21.9)	Trend favours
	AR					17.2 (27.5)	APE but n.s.
	AR (CAA)					18.9 (27.2)	
Silva et al. ³⁸ †	APE					0 (0–100)	Favours APE
	AR					0 (0–100)	(P < 0.050)‡

*Values are mean(s.d.) unless indicated otherwise; †values are median (range). The European Organization for Research and Treatment of Cancer (EORTC) QLQ-CR30 and QLQ-CR29 scores have a range of 1–100, where 0 represents the lowest symptom burden. A score difference or change of 10 is claimed to be clinically important. Values are rounded to one decimal place. Articles with data represented graphically are not included in this table. ‡P < 0.050 was considered statistically significant. APE, abdominoperineal excision; AR, anterior resection; n.s., not statistically significant; ISR, intersphincteric resection; CAA, coloanal anastomosis.

The comparison of observational data between single-arm studies further increases the risk of bias, which was therefore avoided by exclusion of papers with no direct comparison between groups. Selection bias in observational studies in the review will have been increased further owing to patient selection for different techniques. The use of non-validated tools for postoperative QoL was also commonplace and did not allow accurate and reliable conclusions to be drawn from the data. An example is the use of QoL questions that had not been assessed to demonstrate validity, reliability or to ensure that they provided a true reflection of the patient experience. There was variation between studies, with heterogeneity in results identified. The variety of inclusion criteria, differing levels of neoadjuvant therapy, differences in follow-up time, and range of surgical techniques is likely to be reflected in the differences in results.

Most studies did not specify location of the rectal cancer as an inclusion criterion; higher rectal tumours suitable for AR and not for APE will create selection bias because of a lower risk of developing low anterior resection syndrome, and produce more favourable outcomes in the AR group. The cohort of patients who underwent AR in the included studies often excluded those who had not undergone ileostomy reversal. The non-closure rate of defunctioning ileostomies 18 months after AR was 25.1–30 per cent^{46,47} and these patients are considered to have permanent loop ileostomies. The exclusion of these patients, therefore, is not reflective of clinical practice. These patients may have had their QoL improved by having an end colostomy at initial operation rather than living with a loop ileostomy and its attendant challenges of dietary restrictions, skin irritation, and renal impairment, although this was not addressed in the present analysis. Some studies excluded patients who had a postoperative anastomotic leak. Such leaks have a significant impact on long-term QoL and therefore introduce significant bias into the relevant studies. The exclusion of patients with a permanent ileostomy, patients at higher risk of low anterior resection syndrome, and those who had an anastomotic leak may reflect favourably on patients who have

Table 5 Validated scores for gastrointestinal symptoms

	Surgical			Score*			Direction of
	procedure	Baseline	0–5 months	6–11 months	1–2 years	>2 years	effect
QLQ-CR29: flatulence							
Trenti et al. ³⁹	APE					31.5(26.8)	Favours APE
	AR					42.1(30.0)	(P < 0.050)‡
Wani et al. ⁴⁰	AR (CAA)			22 1/27 0\		56.7(30.5)	No difference
wani et ai.	APE AR			32.1(27.9) 33.7(28.3)			No difference identified
Arraras et al. ²⁴	APE			33.3(26.5)			No difference
THI THI HOUSE CO. S.	AR			34.0(26.5)			identified
QLQ-CR29: stool frequency				,			
Trenti et al. ³⁹	APE					21.8(22.3)	Trend favours
	AR					31.8(25.1)	APE but n.s.
Wani et al. ⁴⁰	AR (CAA) APE			29.8(26.6)		40.0(26.5)	No difference
vv aiii et ui.	AR			28.4(24.3)			identified
Arraras et al. ²⁴	APE			14.3(18.5)			Favours APE
	AR			33.3(23.6)			(P < 0.050)‡
QLQ-CR28: GI tract sympton	ms						, , , , ,
Du et al. ²⁶	APE		20.1(8.4)	18.3(7.4)	15.1(5.5)		No difference
December 21 37	AR	04.4	15.9(4.0)	15.5(3.6)	14.2(3.5)		identified
Russell et al. ³⁷	APE AR	21.4 16.8			15.2 18.9		No difference identified
Konanz et al. ³³	APE	10.0			10.9	23.6	Trend favours
Konanz et al.	AR					32.5	APE but n.s.
	AR (ISR)					37.8	
	APE			6.7			No difference
	AR			0			identified
EORTC QLQ-CR30: nausea/		0 (0, 00)			0 (0, 22)	0 (0, 00)	NI - 3:66
How et al. ³⁰ †	APE AR	0 (0–33) 0 (0–33)			0 (0–33) 0 (0–33)	0 (0–33) 0 (0–33)	No difference identified
Mrak et al. ³⁵	APE	0 (0–33)			0 (0–33)	6.7	No difference
	AR					3.8	identified
Konanz et al. ³³	APE					2.3	No difference
	AR					4.9	identified
134	AR (ISR)	44.01			40.71	4.6	
Monastyrska et al. ³⁴	APE	11.3‡			13.7‡		Favours AR
Wani et al. ⁴⁰	AR APE	4.7‡			7.4‡ 8.1(18.1)		(P < 0.050)‡ No difference
wan et ui.	AR				7.3(17.2)		identified
Arraras et al. ²⁴	APE				3.9(18.2)		No difference
	AR				5.2(17.5)		identified
Du et al. ²⁶	APE		7.7(5.6)	7.8(5.5)	5.8(5.9)		No difference
	AR		6.2(5.4)	6.1(4.4)	5.0(5.6)	4 = (4 = 0)	identified
Trenti et al. ³⁹	APE AR					4.5 (15.3) 2.8 (8.4)	No difference identified
	AR (CAA)					2.3 (7.4)	idelitilled
Silva et al. ^{38†}	APE					(0–100)	No difference
	AR					(0–100)	identified
EORTC QLQ-CR30: diarrhoe							
How et al. ³⁰ †	APE	33 (0–67)			0 (0–67)‡	0 (0–67)‡	Favours APE
Mrak et al. ³⁰	AR	0 (0–100)			33(0–100)‡	33 (0–67)‡	(P < 0.050)‡
Mrak et at.	APE AR					16.7 26.1	Trend favours APE but n.s.
Konanz et al. ³³	APE					16.7‡	Favours APE
ixoriariz et ar.	AR					34.1‡	(P < 0.050)‡
	AR (ISR)					45.5‡	71
Monastyrska et al. ³⁴	APE	30.7			38.7‡		Favours AR
. 140	AR	32			0.7‡		(P < 0.050)‡
Wani et al. ⁴⁰	APE				15.0(25.1)		No difference
Arraras et al. ²⁴	AR APE				16.7(32.1) 11.1(19.2)		identified Trend favours
Allalas et ul.	AR				21.4(27.3)		APE but n.s.
Du et al. ²⁶	APE		9.8(8.0)	8.9(7.8)	8.7(7.8)		No difference
	AR		12.3(9.4)	11.8(7.3)	9.3(6.5)		identified
Trenti et al. ³⁹	APE		` /	` '	` '	17.1(24.6)	Trend favours
	AR					22.9(25.8)	APE but n.s.
0'1 1 00 ⁺	AR (CAA)					27.8(27.8)	27 1166
Silva et al.38 [†]	APE					0 (0-66.7)	No difference
	AR					0 (0-100)	identified

(continued)

Table 5 (continued)

	Surgical			Score*			Direction of
	procedure	Baseline	0–5 months	6–11 months	1–2 years	>2 years	effect
EORTC QLQ-CR30: consti	ipation						
How et al. ³⁰ †	APE	0 (0-100)			0 (0-67)‡	0 (0-33)	No difference
	AR	0 (0–67)			0 (0-100)‡	0 (0–67)	identified
Mrak et al. ³⁵	APE	, ,			, , , ,	14	No difference
	AR					21.6	identified
Konanz et al. ³³	APE					12‡	Favours APE
	AR					25.2‡	(P < 0.050)‡
	AR (ISR)					20.2‡	, , , , , ,
Monastyrska et al. ³⁴	APE ´	36.67			16‡		Favours AR
,	AR	23.3			0‡		(P < 0.050)‡
Wani et al. ⁴⁰	APE				15.5 (27.9)		No difference
	AR				15.4 (27.2)		identified
Arraras et al. ²⁴	APE				20.6 (24.7)		No difference
	AR				26.8 (33.9)		identified
Du et al. ²⁶	APE		13.9 (9.0)	13.7 (7.1)	12.1 (4.8)		No difference
	AR		15.8 (8.9)	14.5 (6.5)	13.0 (5.2)		identified
Trenti et al. ³⁹	APE		, ,	, ,	, ,	8.1 (19.2)‡	Favours APE
	AR					28.4 (32.1)‡	(P < 0.050)‡
	AR (CAA)					20.0 (24.1)‡	, , , , , ,
Silva et al. ³⁸ †	ÀPE ′					0 (0–100)	No difference
•	AR					0 (0–100)	identified

^{*}Values are mean(s.d.) unless indicated otherwise; † values are median (range). The European Organization for Research and Treatment of Cancer (EORTC) QLQ-CR30, QLQ-CR38, and QLQ-CR29 scores have a range of 1–100, where 0 represents the lowest symptom burden. A score difference or change of 10 is claimed to be clinically important. Values are rounded to one decimal place. †P < 0.050 was considered statistically significant. APE, abdominoperineal excision; AR, anterior $resection; \'{C}AA, \'{c}oloanal\ an astomosis; n.s., not\ statistically \^significant; ISR, intersphincteric\ resection.$

Table 6 Validated measures of body image

	Surgical	, ,						Direction of
	procedure	Baseline	0–5 months	6–11 months	1–2 years	> 2 years	effect	
Du et al. ²⁶	APE		75.1(11.4)	77.4(11.6)	79.9(9.4)		Favours AR	
QLQ-CR38	AR		81.1(11.5)	84.4(8.9)	86.5(10.6)		(P < 0.050)‡	
Mrak et al. ³⁵	APE		,	, ,	, ,	63.7(30.1)	Trend favours	
QLQ-CR29	AR					79.2(23.9)	AR but n.s.	
How et al. ³⁰ †	APE	100 (50-100)			75 (25-100)	75 (2Š–10Ó)	No difference	
QLQ-CR38	AR	92 (33–100)			83 (0-100)	75 (33–100)	identified	
Konanz et al. ³³	APE	, ,			,	62.4	Trend favours	
QLQ-CR38	AR					75.3	AR but n.s.	
	AR (ISR)					72.7		
Arraras et al. ²⁴	APE				92.1(11.7)		No difference	
QLQ-CR29	AR				85.4(21.8)		identified	
Wani et al. ⁴⁰	APE				84.1(15.0)		No difference	
QLQ-CR29	AR				83.6(13.9)		identified	
Trenti et al. ³⁹	APE					68.0(27.8)	Favours AR	
QLQ-CR29	AR					81.9(26.2)	(P < 0.050)‡	
	AR (CAA)					81.5(21.7)		
Silva et al. ³⁸ †	APE					86.1 (0–100)	No difference	
QLQ-CR29	AR					88.9 (0–100)	identified	

*Values are mean(s.d.) unless indicated otherwise; †values are median (range). The European Organization for Research and Treatment of Cancer (EORTC) QLQ-CR38 and QLQ-CR29 scores have a range of 1–100, where 0 represents the lowest symptom burden. A score difference or change of 10 is claimed to be clinically important. Values are rounded to one decimal place. Articles with data represented graphically are not included in this table. †P < 0.050 was considered statistically significant. APE, abdominoperineal excision; AR, anterior resection; n.s., not statistically significant; ISR, intersphincteric resection; CAA, coloanal anastomosis

undergone AR and not reflect clinical practice. This, therefore, does not allow surgeons to provide patients with accurate information.

The results of this systematic review are in keeping with previously published work. The Cochrane review¹⁶ published in 2012 found equipoise in QoL outcomes and was also unable to recommend AR over APE. A previous meta-analysis⁴⁸ from 2007 also identified no difference in QoL outcomes after AR versus APE for rectal cancer. The present systematic review supports these findings in studies that have been published since the Cochrane

review in 2012. Data published since this date should also allow for the introduction of enhanced recovery after surgery protocols, the use of preoperative MRI, and should not include the laparoscopic learning curve. These subsequent studies may therefore be more relevant to current practice. Another study⁴⁹ published since completion of the search also identified no difference in global QoL; however, although patients who had undergone APE had lower preoperative QoL, this was not accounted for in the conclusion that patients should undergo restorative surgery for low rectal cancer.

Table 7 Validated measures of sexual function

	Surgical procedure	Baseline	0–5 months	6–11 months	1–2 years	> 2 years	Direction of effect
Sexual interest Mrak et al. ³⁵ QLQ-CR29	APE					23.8(25.2) F 33.6(33.2) M	Trend in women
	AR					40.7(27.8) F 51.2(35.7) M	favours AR but n.s. Favours AR in men (P < 0.050)‡
Wani et al. ⁴⁰ QLQ-CR29	APE				41.0(20.0) F 45.2(24.8) M		Trend favours AR but n.s
	AR				56.6(34.8) F 64.4(36.2) M		
Trenti et al. ³⁹ QLQ-CR29	APE				11.1(28.0) F 40.5(31.5) M		Trend favours AR but n.s.
	AR				20.0(25.8) F 42.3(33.2) M		
	AR (CAA)				14.3(17.8) F 46.0(30.7) M		
Silva et al. ³⁸ † QLQ-CR29	APE				66.7 (0–100) F 33.3 (0–100) M		No difference identified
	AR				66.7 (33.3–100) F 33.3 (0–100) M		
Sexual enjoyment How et al. ³⁰ † QLQ-CR38	APE AR	16.5 (0–100) 50 (0–100)			17 (0–100) 67 (0–100)	33 (0–67) 16 (0–67)	Trend favours APE at 2 years but
Konanz et al. ³³ QLQ-CR38	APE AR AR (ISR)					56.4 53.7 75.9	n.s. Favours AR (P < 0.050, ISR versus APE)‡
Penchev et al. ³⁶ QLQ-CR38 Du et al. ²⁶ QLQ-CR38 Sexual functioning	APE AR APE AR		41.7(16.0) 52.1(12.3)	27.7(31.2) 18.9(15.1) 46.7(16.0) 56.7(12.1)	51.7(14.2) 60.1(11.6)		Trend favours APE but n.s. Favours AR (P < 0.050)‡
How et al. ³⁰ † QLQ-CR38 Konanz et al. ³³ QLQ-CR38	APE AR APE AR AR (ISR)	0(0–83) 33(0–83)			0(0–67) 33(0–67)	0(0-33) 33(0-100) 21.7 31.7 44.4	Favours AR (P < 0.050)‡ Favours AR (P < 0.050, ISR versus APE)‡
Arraras et al. ²⁴ QLQ-CR29	APE AR				3.5(10.5) F 31.2(35.4) M	4.8(12.6) F 36.1(38.8) M	No difference identified
Penchev et al. ³⁶ QLQ-CR38	APE AR			9.6(15.9) 10.36(16.3)	J 1.2 (JJ. 1) 1VI	50.1(50.0) 141	No difference identified
Du et al. ²⁶ QLQ-CR38	APE AR		38.8(17.2) 46.6(13.3)	42.1(17.9) 52.0(13.1)	46.6(14.5) 56.1(14.2)		Favours AR (P < 0.050, ISR versus APE)‡

*Values are mean(s.d.) unless indicated otherwise; †values are median (range). The European Organization for Research and Treatment of Cancer (EORTC) QLQ-CR38 and QLQ-CR29 scores have a range of 1–100, where 0 represents the lowest symptom burden. A score difference or change of 10 is claimed to be clinically important. Values are rounded to one decimal place. Articles with data represented graphically are not included in this table. $^{\dagger}P < 0.050$ was considered statistically significant. APE, abdominoperineal excision; AR, anterior resection; n.s., not statistically significant; ISR, intersphincteric resection.

These results should be discussed with patients as part of shared decision-making and consenting for operative management of rectal cancer, although this review cannot recommend one surgical approach over another for improved QoL. Future studies should record detailed clinical factors alongside properly validated preoperative QoL measures used for patients undergoing both surgical approaches. These studies should include only patients with low rectal cancer, as previously defined in the literature^{50,51}, to allow direct comparison between techniques and reduce selection bias. Patients should be followed up adequately with the same QoL measures used after surgery, and both short- and long-term data collected. The use of the collaborative research model may provide a framework for this work. The Colostomy Impact Score (CIS) and the Low Anterior Resection Score (LARS) both now have validated convergence on the EORTC QLQ-C30, and may therefore be useful in allowing a direct comparison between the two surgical techniques^{52–54}. The impact of ileostomy on patients' QoL should

a Risk of bias of individual studies

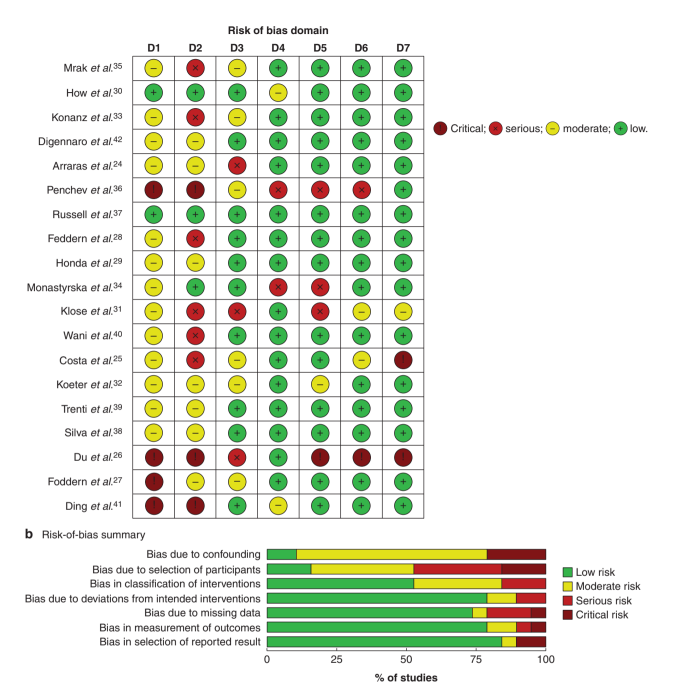


Fig. 2 Risk-of-bias assessment

a Risk of bias in individual trials. D1, bias due to confounding; D2, bias due to selection of participants; D3, bias in classification of interventions; D4, bias due to deviations from intended interventions; D5, bias due to missing data; D6, bias in measurement of outcomes; D7, bias in selection of reported result. b Risk-of-bias summary.

be considered, and may not be assessed accurately by the CIS and LARS. Additional work is required to understand the process by which surgeons decide which operations to offer the individual patient.

Disclosure. The authors declare no conflict of interest.

Supplementary material

Supplementary material is available at BJS Open online.

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