National Institute for Health and Care Excellence

Version 1.0 Pre-consultation

Oesophago-gastric cancer

assessment and management in adults

Appendix G GRADE profiles 15 June 2017

Draft for Consultation

Developed by the National Guideline Alliance, hosted by the Royal College of Obstetricians and Gynaecologists

DRAFT FOR CONSULTATION

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Appendix G: GRADE Profiles

G.1 Radical treatment

What are the specific information and support needs before and after treatment for adults with oesophago-gastric cancer who are suitable for radical treatment and their carers?

Not applicable to this review.

G.2 Palliative management

What are the specific information and support needs of adults with oesophago-gastric cancer who are suitable for palliative treatments and care only?

Not applicable to this review.

G.3 MDT

What is the most effective organisation of local and specialist MDT services for adults with oesophago-gastric cancer? No evidence was identified for this review.

G.4 Surgical services

What is the optimal provision and organisation of surgical services for people with oesophago-gastric cancer? GRADE was not applicable for this review. See modified clinical evidence profile in the full guideline for evidence tables.

G.5 Staging investigations

What are the optimal staging investigations to determine suitability for curative treatment of oesophageal or gastro-oesophageal junctional cancer after diagnosis with endoscopy and whole-body CT scan?

GRADE was not used for this review. See modified clinical evidence profile in the full guideline for evidence tables.

G.6 Staging investigations

What are the optimal staging investigations to determine suitability for curative treatment of gastric cancer after diagnosis with endoscopy and whole-body CT scan?

GRADE was not used for this review. See modified clinical evidence profile in the full guideline for evidence tables.

G.7 HER2 testing in adenocarcinoma

Which people with adenocarcinoma of the stomach and oesophagus should have their tumours HER2 tested?

No evidence was identified for this review.

G.8 T1N0 oesophageal cancer

What is the optimal management of T1N0 oesophageal cancer?

Table 1: Clinical evidence profile: EMR versus oesphagectomy

Quality	assessment						Nº of patien	ts	Effect			
Nº of studie s	Study design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecisi on	Other consideratio ns	Endoscop ic mucosal resection	Surgica I resectio n	Relati ve (95% Cl)	Absolu te (95% Cl)	Qualit y	Importan ce
Overall	survival (fol	low up: ı	median 48 mon	iths)								

1

Quality	assessment						Nº of patien	ts	Effect			
Nº of studie s	Study design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecisi on	Other consideratio ns	Endoscop ic mucosal resection	Surgica I resectio n	Relati ve (95% Cl)	Absolu te (95% Cl)	Qualit y	Importan ce
1	observatio nal studies	not serio us	not serious	not serious	serious ¹	none	6/26 (23.1%)	6/44 (13.6%)	HR 1.60 (0.49 to 5.15)	5 year OS 85% with surgery vs 77% (43% to 92%) with EMR	VERY LOW	Important

CI: Confidence interval; HR: Hazard Ratio; OS: overall survival; EMR=Endoscopic mucosal resection 1. Downgraded one level for imprecision: event rate <300<u>HR</u> includes both default thresholds

Table 2: Clinical evidence profile: EMR versus ESD

Quality	assessment						Number patients	of	Effect			
Nº of studie s	Study design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecisi on	Other consideratio ns	EMR	ESD	Relative (95% Cl)	Absolu te (95% Cl)	Qualit y	Importan ce
Diseas	e free survival	(follow u	p: 12 months)									
1	observation al studies	seriou s ¹	not serious	not serious	serious ²	none	1/184 (0.5%)	0/116 (0.0%)	not estimabl e	-	VERY LOW	CRITICAL
Patholo	ogical margins	free (pos	st treatment)									
1	randomised trials	seriou s ¹	not serious	not serious	serious ²	none	144/18 4 (78.3%)	113/11 6 (97.4%)	RR 0.80 (0.74 to 0.87)	195 fewer per 1,000 (from 127 fewer to	VERY LOW	CRITICAL

Quality	assessment						Number patients	of	Effect			
Nº of studie s	Study design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecisi on	Other consideratio ns	EMR	ESD	Relative (95% Cl)	Absolu te (95% Cl)	Qualit y	Importan ce
										253 fewer)		
Stenosi	is (post treatm	ent)										
1	randomised trials	seriou s ¹	not serious	not serious	serious ²	none	17/184 (9.2%)	20/116 (17.2%)	RR 0.54 (0.29 to 0.98)	79 fewer per 1,000 (from 3 fewer to 122 fewer)	VERY LOW	CRITICAL
Overall	survival (follo	w up: 12	months)									
1	observation al studies	seriou s ¹	not serious	not serious	serious ²	none	NR/184	NR/116	not estimabl e	OS 85% at 1 year for both	VERY LOW	CRITICA
Perfora	tion (post trea	tment)										
1	observation al studies	seriou s ¹	not serious	not serious	serious ²	none	3/184 (1.6%)	3/116 (2.6%)	RR 0.63 (0.13 to 3.07)	10 fewer per 1,000 (from 23 fewer to 54 more)	VERY LOW	CRITICAL

CI: Confidence interval; RR: Risk ratio; OS: overall survival; EMR=Endoscopic mucosal resection; ESD=Endoscopic submucosal resection; NR=not reported 1. Tumours were on average 10mm larger in the ESD group 2. Downgraded one level for imprecision: <u>HR or RR includes both default thresholdsevent rate <300</u>

1

G.9 Surgical treatment of oesophageal cancer

What is the most effective operative approach for the surgical treatment of oesophageal cancer?

Quality	assessmen	it					No of patien	ts	Effect			
No of studie s	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecisio n	Other considerati ons	Transthora cic	Transhia tal	Relativ e (95% Cl)	Absolu te	Quality	Importanc e
Post-op	perative con	nplicatio	ns: Anastomo	tic leak - The	oracotomy+La	parotomy						
2	randomis ed trials	seriou s ¹	no serious inconsisten Cy	no serious indirectne ss	very serious ²	none	2/38 (5.3%)	4/35 (11.4%)	RR 0.52 (0.12 to 2.24)	55 fewer per 1000 (from 101 fewer to 142 more)	VERY LOW	CRITICAL
Post-op	perative con	nplicatio	ns: Anastomo	tic leak - The	oracotomy+La	aparotomy+Ce	rvical incision					
2	randomis ed trials	seriou s ¹ 1	serious ³	no serious indirectne ss	very serious ²	none	17/144 (11.8%)	28/151 (18.5%)	RR 0.48 (0.11 to 2.14)	96 fewer per 1000 (from 165 fewer to 211 more)	VERY LOW	CRITICAL
Overall	survival - T	horacoto	omy+Laparoto	my+Cervica	l incision							
1	randomis ed trials	seriou s ^{1<u>1</u>}	no serious inconsisten cy	no serious indirectne ss	serious ⁴	none	-	-	Not estimab le	-	LOW	CRITICAL

Quality	assessmer	nt					No of patien	ts	Effect			
No of studie s	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecisio n	Other considerati ons	Transthora cic	Transhia tal	Relativ e (95% Cl)	Absolu te	Quality	Importanc e
Intraop	erative bloc	od loss (n	nl) - Thoracot	omy+Laparo	tomy (Better i	indicated by lo	wer values)					
2	randomis ed trials	seriou s ^{1<u>1</u>}	very serious⁵	no serious indirectne ss	very serious ⁶	none	30	29	-	MD 8.98 higher (81.33 lower to 99.29 higher)	VERY LOW	CRITICAL
Intraop	erative bloc	od loss (n	nl) - Thoracot	omy+Laparo	tomy+Cervica	al incision (Bet	ter indicated k	oy lower val	ues)			
1	randomis ed trials	seriou s ^{1<u>1</u>}	no serious inconsisten cy	no serious indirectne ss	very serious ⁶	none	40	40	-	MD 16 higher (87.23 lower to 119.23 higher)	VERY LOW	CRITICAL
Length	of operatio	n (min) -	Thoracotomy	+Laparotom	y (Better indic	cated by lower	values)					
3	randomis ed trials	seriou s ¹	serious ⁷	no serious indirectne ss	serious ⁸	none	48	45	-	MD 30.68 lower (51.82 to 9.55 lower)	VERY LOW	IMPORTA NT
Length	of operatio	n (min) -	Thoracotomy	+Laparotom	y+Cervical ind	cision (Better i	ndicated by lo	wer values)				
1	randomis ed trials	seriou s ^{1<u>1</u>}	no serious inconsisten cy	no serious indirectne ss	very serious ⁹	none	40	47	-	MD 121.1 lower (152.37 to 89.83 lower)	VERY LOW	IMPORTA NT

Quality	assessmer	it					No of patien	ts	Effect			
No of studie s	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecisio n	Other considerati ons	Transthora cic	Transhia tal	Relativ e (95% Cl)	Absolu te	Quality	Importanc e
2	randomis ed trials	seriou s ¹	no serious inconsisten cy	no serious indirectne ss	very serious ²	none	8/38 (21.1%)	7/35 (20%)	RR 1.02 (0.24 to 2.29)	4 more per 1000 (from 152 fewer to 258 more)	VERY LOW	CRITICAL
Post-op	perative con	nplication	ns: Pneumoni	a - Thoracot	omy+Laparoto	omy+Cervical i	ncision					
2	randomis ed trials	seriou s ^{1<u>1</u>}	no serious inconsisten cy	no serious indirectne ss	very serious ²	none	7/52 (13.5%)	11/57 (19.3%)	RR 0.68 (0.29 to 1.62)	62 fewer per 1000 (from 137 fewer to 120 more)	VERY LOW	CRITICAL
Numbe	r of lymph r	odes res	ected - Thora	icotomy+Lap	arotomy+Cer	vical incision (Better indicat	ed by lower	values)			
1	randomis ed trials	seriou s ^{1<u>1</u>}	no serious inconsisten cy	no serious indirectne ss	no serious imprecision ¹ 0	none	94	111	-	MD 15 lower (18.18 to 11.82 lower)	MODER ATE	CRITICAL
Resecti	ion margin											
1	randomis ed trials	seriou s ¹	no serious inconsisten cy	no serious indirectne ss	no serious imprecision	none	92/282 (32.6%)	111/333 (33.3%)	RR 0.98 (0.82 to 1.17)	7 fewer per 1000 (from 60	MODER ATE	CRITICAL

Quality	assessmer	nt					No of patien	ts	Effect			
No of studie s	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecisio n	Other considerati ons	Transthora cic	Transhia tal	Relativ e (95% Cl)	Absolu te	Quality	Importanc e
										fewer to 57 more)		
Resecti	ion margin	- Thoraco	otomy+Laparo	tomy+Cervi	cal incision:R	0 resection						
1	randomis ed trials	seriou S ^{1<u>1</u>}	no serious inconsisten cy	no serious indirectne ss	no serious imprecision	none	68/94 (72.3%)	79/111 (71.2%)	RR 1.02 (0.86 to 1.21)	14 more per 1000 (from 100 fewer to 149 more)	MODER ATE	CRITICAL
Resecti	ion margin	- Thoraco	otomy+Laparo	tomy+Cervi	cal incision: F	1 resection						
1	randomis ed trials	seriou s ^{1<u>1</u>}	no serious inconsisten Cy	no serious indirectne ss	very serious ²	none	23/94 (24.5%)	28/111 (25.2%)	RR 0.97 (0.6 to 1.56)	8 fewer per 1000 (from 101 fewer to 141 more)	VERY LOW	CRITICAL
Resecti	ion margin	- Thoraco	otomy+Laparo	tomy+Cervi	cal incision: F	2 resection						
1	randomis ed trials	seriou s ^{1<u>1</u>}	no serious inconsisten cy	no serious indirectne ss	very serious ²	none	1/94 (1.1%)	4/111 (3.6%)	RR 0.3 (0.03 to 2.6)	25 fewer per 1000 (from 35 fewer to	VERY LOW	CRITICAL

	assessmen	Î.					No of patien	1	Effect		-	
No of studie s	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecisio n	Other considerati ons	Transthora cic	Transhia tal	Relativ e (95% Cl)	Absolu te	Quality	Importanc e
										58 more)		
Recurr	ence - Thora	acotomy	+Laparotomy									
1	randomis ed trials	seriou s ¹	no serious inconsisten cy	no serious indirectne ss	very serious ²	none	4/20 (20%)	6/19 (31.6%)	RR 0.63 (0.21 to 1.9)	117 fewer per 1000 (from 249 fewer to 284 more)	VERY LOW	IMPORTA NT
Recurr	ence - Thora	acotomy	+Laparotomy+	Cervical inc	ision							
1	randomis ed trials	seriou s ^{1<u>1</u>}	no serious inconsisten cy	no serious indirectne ss	serious ⁴	none	59/95 (62.1%)	59/110 (53.6%)	RR 1.16 (0.92 to 1.46)	86 more per 1000 (from 43 fewer to 247 more)	LOW	IMPORTA NT
Mortali	ty - Thoraco	tomy+La	aparotomy									
2	randomis ed trials	seriou s ¹	no serious inconsisten cy	no serious indirectne ss	very serious ²	none	2/52 (3.8%)	3/54 (5.6%)	not pooled	not pooled	VERY LOW	IMPORTA NT
30-day	mortality - T	Thoracot	omy+Laparot	omy+Cervica	al incision							
1	randomis ed trials	seriou s ^{1<u>1</u>}	no serious inconsisten cy	no serious indirectne ss	very serious ²	none	1/16 (6.3%)	1/16 (6.3%)	RR 1 (0.07 to 14.64)	0 fewer per 1000 (from	VERY LOW	IMPORTA NT

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Quality	assessmen	t					No of patien	ts	Effect			
No of studie s	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecisio n	Other considerati ons	Transthora cic	Transhia tal	Relativ e (95% Cl)	Absolu te	Quality	Importanc e
										58 fewer to 853 more)		
Progres	ssion-free s	urvival -	Thoracotomy [.]	+Laparotom	y+Cervical inc	ision						
1	randomis ed trials	seriou s ^{1<u>1</u>}	no serious inconsisten cy	no serious indirectne ss	very serious ²	none	-	-	Not estimab le	-	VERY LOW	CRITICAL

CI=Confidence interval; RR=relative risk; HR=Hazard ratio; MD=Mean difference; mI=millilitres; min=minutes

¹ <u>Chu 199, Goldminc 1993 -</u> Poor reporting of random sequence generation and allocation concealment.

² 95% CI crosses 2 default MID therefore downgraded by 2 levels

³ I2 73% therefore downgraded by 1 level

- ⁴ 95% CI crosses 1 default MID therefore downgraded by 1 level
- ⁵ I2 89% therefore downgraded by 2 levels
- ⁶ Default MID: +/-34.25: 95% CI crosses 2 default MIDs therefore downgraded by 2 levels
- ⁷ I2 71% therefore downgraded by 1 level

¹ I2 /1% therefore downgraded by 1 level
 ⁸ Default MID: +/-12.53: 95%Cl crosses 1 default MID therefore downgraded by 1 level
 ⁹ Default MID: +/-12.53: 95%Cl crosses 2 default MID therefore downgraded by 2 levels
 ¹⁰ Default MID: +/-7 therefore not downgraded for imprecision
 ¹¹Chou 2009, Jacobi 1997 - Poor reporting of random sequence generation and allocation concealment

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Table 4: Clinical evidence profile: Minimally invasive versus open oesophagectomy

Quality asses	sment					No of pat	tients	Effect			
No of Desig studi es	n Risk of bias	Inconsisten cy	Indirectne ss	Imprecision	Other consideratio ns	Minima Ily invasiv e	Open	Relati ve (95% CI)	Absolu te	Quality	Importance

Quality	y assessmen	t					No of pa	tients	Effect			
No of studi es	Design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecision	Other consideratio ns	Minima Ily invasiv e	Open	Relati ve (95% CI)	Absolu te	Quality	Importance
2	randomise d trials	Seriou s ²	no serious inconsistenc y	no serious indirectnes s	very serious ¹	none	8/170 (4.7%)	6/166 (3.6%)	RR 1.29 (0.44 to 3.54)	10 more per 1000 (from 20 fewer to 92 more)	VERY LOW	CRITICAL
Post-o	perative con	plication	s - Pulmonary	complication	าร							
2	randomise d trials	serious 2	serious ¹	no serious indirectnes s	serious ¹²	none	5/170 (2.9%)	11/16 6 (6.6%)	RR 0.45 (0.16 to 1.24)	36 fewer per 1000 (from 56 fewer to 16 more)	LOW	CRITICAL
Intraop	perative bloo	d loss (m	l) ³ (Better indi	cated by lowe	er values)							
2	randomise d trials	serious 2	very serious ⁴	no serious indirectnes s	very serious⁵	none	169	167	-	MD 109.43 lower (1061.1 2 lower to 842.26 higher)	VERY LOW	CRITICAL

Quality	/ assessmen	t					No of pa	tients	Effect			
No of studi es	Design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecision	Other consideratio ns	Minima Ily invasiv e	Open	Relati ve (95% Cl)	Absolu te	Quality	Importance
1	randomise d trials	serious 2	no serious inconsistenc y	no serious indirectnes s	serious ⁶	none	59	56	-	MD 10 higher (2.83 to 17.17 higher)	LOW	IMPORTAN T
Length	of operation	n (min) (B	letter indicated	l by lower val	ues)							
2	randomise d trials	serious 2	no serious inconsistenc y	no serious indirectnes s	serious ⁷	none	170	166	-	MD 48.06 higher (29.56 to 66.56 higher)	LOW	IMPORTAN T
Resect	tion margin -	R0										
1	randomise d trials	serious 2	no serious inconsistenc y	no serious indirectnes s	no serious imprecision	none	54/59 (91.5%)	47/56 (83.9 %)	RR 1.09 (0.92 to 1.16)	76 more per 1000 (from 67 fewer to 134 more)	MODERAT E	CRITICAL

Quality	y assessmen	t					No of pat	tients	Effect	-		
No of studi es	Design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecision	Other consideratio ns	Minima Ily invasiv e	Open	Relati ve (95% Cl)	Absolu te	Quality	Importance
1	randomise d trials	serious 2	no serious inconsistenc y	no serious indirectnes s	very serious ²	none	1/59 (1.7%)	5/56 (8.9%)	RR 0.19 (0.02 to 1.49)	72 fewer per 1000 (from 87 fewer to 44 more)	VERY LOW	CRITICAL
Numbe	er of lymph n	odes res	ected ⁸ (Better	indicated by	lower values)							
2	randomise d trials	serious 2	very serious ⁹	serious ¹⁰	no serious imprecision ¹¹	none	170	166	-	MD 19.32 lower (22.28 to 16.36 lower)	VERY LOW	CRITICAL
30 day	mortality											
1	randomise d trials	serious 2	no serious inconsistenc y	no serious indirectnes s	very serious ¹	none	1/59 (1.7%)	0/56 (0%)	RR 2.9 (0.12 to 72.62)	2 more per 1000 (from 1 fewer to 72 more)	VERY LOW	CRITICAL

CI=Confidence interval; RR=relative risk; MD=Mean difference; QoL=Quality of life; EORTC=European Organisation for Research and Treatment of Cancer; ml=millilitres; min=minutes

¹ 95% CI crosses both default MIDs therefore downgraded by 2

² <u>Biere 2012, Guo 2013 -</u> Poor reporting of random sequence generation and allocation concealment.

³ Mean (standard deviation) intraoperative blood loss in control arm (open oesophagectomy): 614.6 (490.3) ml

⁴ I2 98% therefore downgraded by 2
 ⁵ Default MID: +/- 245.15. 95% CI crosses both arms, therefore downgraded by 2

⁶ Default MID: +/- 10.5. 95% CI crosses 1 arm of default MID: therefore downgraded by 1 ⁷ Default MID: +/- 55.9. 95% CI crosses 1 arm, therefore downgraded by 1

⁸ Mean (standard deviation) number of lymph nodes resected in control arm (open oesophagectomy): 39.1 (11.5)

⁹ I2 99% therefore downgraded by 2

¹⁰ Inconsistency could be explained by variation in location of studies (China vs Netherlands), surgical practices and prevalence of oesophageal cancer.

¹¹ Default MID: +/- 5.75. 95% CI does not cross default MID therefore not downgraded

¹² 95%CI crossed one boundary of default MID and therefore downgraded by 1 level

Table 5: Clinical evidence profile: Hybrid versus open oesophagectomy

Quality	assessmen						No of p	atients	Effect			
No of studi es	Design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecisi on	Other considerati ons	Hybri d	Open	Relativ e (95% Cl)	Absolute	Quality	Importan ce
Major p	oost-operativ	e complie	cations - Pulm	onary compli	cation							
1	randomise d trials	no serious risk of bias ¹	no serious inconsistenc y	no serious indirectnes s	serious ²	none	18/103 (17.5 %)	31/104 (29.8 %)	RR 0.59 (0.33 to 0.97)	122 fewer per 1000 (from 9 fewer to 200 fewer)	MODERAT E	CRITICA L
Major p	oost-operativ	e complie	cations - Major	· post-operati	ve complica	tion						
1	randomise d trials	no serious risk of bias ¹	no serious inconsistenc y	no serious indirectnes s	no serious imprecisio n	none	37/103 (35.9 %)	67/104 (64.4 %)	RR 0.56 (0.38 to 0.77)	283 fewer per 1000 (from 148 fewer to 399 fewer)	HIGH	CRITICA L

Quality	assessment	t					No of p	atients	Effect			
No of studi es	Design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecisi on	Other considerati ons	Hybri d	Open	Relativ e (95% Cl)	Absolute	Quality	Importan ce
1	randomise d trials	no serious risk of bias ¹	no serious inconsistenc y	no serious indirectnes s	very serious ³	none	5/103 (4.9%)	5/104 (4.8%)	RR 1.01 (0.3 to 3.38)	0 more per 1000 (from 34 fewer to 114 more)	LOW	CRITICA L

CI=Confidence interval; RR=relative risk;

¹ Risk of bias assessment based on protocol and conference abstract. No full publication available.

² 95% CI crosses one default MIDs therefore downgraded by 1

³ 95% CI crosses both default MIDs therefore downgraded by 2

G.10 Lymph node dissection in oesophageal and gastric cancer

Does the extent of lymph node dissection influence outcomes in adults with oesophageal and gastric cancer?

 Table 4: Clinical evidence profile: D2 versus D1 lymphadenectomy for gastric cancer

Quality	assessment	t					No of pa	atients	Effect			
No of studie s	Design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecisio n	Other consideratio ns	D2	D1	Relativ e (95% Cl)	Absolut e	Qualit y	Importanc e
Overall	survival											
5	randomis ed trials	no serious	serious ¹	serious ²	serious ³	none	805	848	HR 0.91	lf 5yr OS is 49%	VERY LOW	CRITICAL

Quality	assessment	t					No of pa	atients	Effect			
No of studie s	Design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecisio n	Other consideratio ns	D2	D1	Relativ e (95% Cl)	Absolut e	Qualit y	Importanc e
		risk of bias							(0.71 to 1.17)	with D1 it is 52% with D2 (95%Cl 43% to 60%)		
Disease	e free surviv	al										
4	randomis ed trials	no serious risk of bias	serious ^{4,5}	No serious indirectnes s	No serious imprecision 6	none	642	690	HR 0.95 (0.84 to 1.07)	If 5yr DFS is 44% with D1 it is 46% with D2 (95%CI 42% to 50%)	LOW	IMPORTA NT
Postop	erative mort	ality										
7	randomis ed trials	serious 7	no serious inconsistenc y ⁸	serious ⁹	no serious imprecision ¹⁰	none	63/935 (6.7%)	33/978 (3.4%)	RR 2.02 (1.34 to 3.04)	34 more per 1000 (from 11 more to 69 more)	LOW	IMPORTA NT
Pancre	atic leak											
5	randomis ed trials	serious	no serious inconsistenc y ¹²	serious ¹³	no serious imprecision 14	none	23/855 (2.7%)	8/891 (0.9%)	RR 2.96 (1.32 to 6.65)	18 more per 1000 (from 3 more to	LOW	CRITICAL

Quality	assessmen	t					No of p	atients	Effect			
No of studie s	Design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecisio n	Other consideratio ns	D2	D1	Relativ e (95% Cl)	Absolut e	Qualit y	Importanc e
										51 more)		
Reoper	ation rate											
6	randomis ed trials	serious ¹⁵	no serious inconsistenc y ¹⁶	serious ¹⁷	very serious ¹⁸	none	79/734 (10.8%)	36/779 (4.6%)	RR 2.18 (1.32 to 3.6)	55 more per 1000 (from 15 more to 120 more)	VERY LOW	CRITICAL
Anasto	motic leak											
7	randomis ed trials	serious 7	no serious inconsistenc y ¹⁹	serious ²⁰	no serious imprecision ²¹	none	68/886 (7.7%)	32/922 (3.5%)	RR 2.12 (1.41 to 3.2)	39 more per 1000 (from 14 more to 76 more)	LOW	CRITICAL
Haemo	rrhage											
6	randomis ed trials	serious 7	no serious inconsistenc y ⁸	serious ²²	very serious ²³	none	18/963 (1.9%)	24/907 (2.6%)	RR 0.64 (0.34 to 1.2)	10 fewer per 1000 (from 17 fewer to 5 more)	VERY LOW	CRITICAL
Wound	infection											
5	randomis ed trials	serious 7	very serious ²⁴	very serious13	no serious imprecision 25	none	45/564 (8%)	25/820 (3%)	RR 3.51	77 more per 1000	VERY LOW	CRITICAL

Quality	assessmen	t i					No of pa	atients	Effect			
No of studie s	Design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecisio n	Other consideratio ns	D2	D1	Relativ e (95% Cl)	Absolut e	Qualit y	Importanc e
									(0.96 to 12.86)	(from 1 fewer to 362 more)		
Pulmor	nary complic	ation										
5	randomis ed trials	serious 7	no serious inconsistenc y ²⁶	serious ²⁷	no serious imprecision 28	none	73/795 (9.2%)	38/843 (4.5%)	RR 2.07 (1.41 to 3.03)	48 more per 1000 (from 18 more to 92 more)	LOW	CRITICAL
R0 rese	ection											
1	randomis ed trials	no serious risk of bias	no serious inconsistenc y	no serious indirectnes s	no serious imprecision 29	none	293/33 1 (88.5%)	339/38 0 (89.2%)	RR 0.99 (0.94 to 1.05)	9 fewer per 1000 (from 54 fewer to 45 more)	HIGH	CRITICAL

CI=Confidence interval; RR=relative risk; HR=Hazard ratio; OS=Overall survival; DFS=Disease free survival

¹ Heterogeneity: I2=64%

² Indirectness: increased mortality rates in those who underwent pancreatectomy and splenectomy might contribute to indirectness in interventions. Additionally, older trials might have been subject to relative inexperience in surgical techniques and post-operative care for D2 resection, thus confounding the results presented here.

³ Total 95% CI: 0.71, 1.17. Crosses one predetermined 0.80 MID, therefore downgraded by one point.

⁴ No clear reporting from systematic review of additional adjuvant or neoadjuvant treatments given therefore downgraded by 1 point.

⁵ Inconsistency: varying lengths of follow-up in included studies

⁶ Imprecision: 95% confidence interval does not cross the 0.80, 1.25 default MID thresholds

⁷ Risk of bias: Dent 1988 and Robertson 1994 have high risk of attrition bias, Li 2007 and Robertson have unclear risk of bias ratings.

⁸ Inconsistency: I-squared=0%

⁹ Indirectness: postoperative mortality could be affected by dissection of additional organs such as pancreatectomy and splenectomy, subgroup analyses have not been presented here. Older studies may not be comparable with newer studies where they may be better experience of surgical technique and post-operative care.

¹⁰ Imprecision: 95% confidence interval (1.34-3.04). No imprecision

¹¹ Risk of bias: Robertson 1994 has low sample size, Li 2007 and Robertson have unclear risk of bias ratings.

¹² Inconsistency: I-squared=0%.

¹³ Indirectness: Indirect intervention: patients undergoing pancreatectomy may be more likely to develop post-operative complications. Older studies may not be comparable to more recent studies due to improvements in training and experience with surgical technique and post-operative care.

¹⁴ Imprecision: 95% confidence interval: 1.36-7.41. No MIDs crossed

¹⁵ Risk of bias: Dent 1988 and Robertson 1994 have low sample sizes, Li 2007 and Robertson have unclear risk of bias ratings.

¹⁶ Heterogeneity: I2=7%

¹⁷ Indirectness: reoperation rate could be affected by dissection of additional organs such as pancreatectomy and splenectomy, subgroup analyses have not been presented here. Older studies may not be comparable with newer studies where there may be better experience of surgical technique and post-operative care.

¹⁸ 95% CI: 1.63-3.43. Very wide CI crossing both MIDs

¹⁹ Heterogeneity: I2=0%

²⁰ No explanation was provided

²¹ No imprecision. 95% CI: 1.47-3.29.

²² Indirectness: Haemorrhage poorly defined or not defined in most studies, therefore unclear of comparability across studies. Haemorrhage could be affected by dissection of additional organs such as pancreatectomy and splenectomy, subgroup analyses have not been presented here. Older studies may not be comparable with newer studies where there may be better experience of surgical technique and post-operative care.

²³ Imprecision: 95% CI: 0.39-1.26. Crosses two MIDs.

²⁴ Heterogeneity: I2=82%. Very serious imprecision

²⁵ 95% CI: 1.45-3.61. No imprecision as no MIDs crossed

²⁶ Heterogeneity: i2=0%

²⁷ Indirectness: Pulmonary complications poorly define in most studies. Unclear if exclusively refers to pneumonia or includes for instance pleural effusion and pulmonary embolus. Additionally, post-operative complications may have been higher in those who underwent pancreatectomy and splenectomy, older trials might have also been subject to relative inexperience in surgical techniques and post-operative care for D2 resection, thus confounding the results presented here.

²⁸ 95% CI: 1.44-3.06: No imprecision as no default MIDs crossed.

²⁹ 95% CI: 0.94-1.05. No imprecision as does not cross default MID.

Table 5: Clinical evidence profile: D3 versus D2 lymphadenectomy for gastric cancer

Quality assess	nent					No of p	atients	Effect			
No of Design studie s	Risk of bias	Inconsisten cy	Indirectne ss	Imprecisi on	Other consideratio ns	D3	D2	Relativ e (95% CI)	Absolu te	Quality	Importanc e

Overall survival

Quality	assessmen	t					No of pa	atients	Effect			
No of studie s	Design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecisi on	Other consideratio ns	D3	D2	Relativ e (95% Cl)	Absolu te	Quality	Importance
3	randomis ed trials	seriou s ¹	no serious inconsistenc y ²	serious ³	no serious imprecisio n ⁴	none	429	433	HR 0.99 (0.81 to 1.21)	If 5yr OS is 54% with D2 it would be 54% with D3 (95%CI 47% to 61%).	LOW	CRITICAL
Recurre	ence-free su	irvival										
1	randomis ed trials	no seriou s risk of bias	no serious inconsistenc y	no serious indirectnes s ⁵	no serious imprecisio n ⁶	none	99/260 (38.1 %)	100/26 3 (38%)	HR 1.08 (0.83 to 1.42)	5yr RFS 63% with D2 vs 60% with D3 (95%CI 51% to 68%).	MODERA TE	IMPORTA NT
Postop	erative mor	tality										
4	randomis ed trials	seriou S ¹	no serious inconsistenc y	serious ³	serious ⁷	none	14/563 (2.5%)	6/574 (1%)	RR 2.04 (0.78 to 5.35)	11 more per 1000 (from 2 fewer to 45 more)	VERY LOW	IMPORTA NT

Quality	assessmen	t					No of p	atients	Effect			
No of studie s	Design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecisi on	Other consideratio ns	D3	D2	Relativ e (95% Cl)	Absolu te	Quality	Importance
4	randomis ed trials	seriou s ¹	no serious inconsistenc y	serious ³	very serious ⁸	none	34/557 (6.1%)	30/567 (5.3%)	RR 1.15 (0.71 to 1.85)	8 more per 1000 (from 15 fewer to 45 more)	VERY LOW	CRITICAL
Anasto	motic leak											
4	randomis ed trials	seriou s ¹	no serious inconsistenc y	serious ³	very serious ⁹	none	27/557 (4.8%)	33/567 (5.8%)	RR 0.83 (0.51 to 1.36)	10 fewer per 1000 (from 29 fewer to 21 more)	VERY LOW	CRITICAL
Wound	infection											
2	randomis ed trials	no seriou s risk of bias	no serious inconsistenc y ¹⁰	serious ³	very serious ¹¹	none	8/262 (3.1%)	10/269 (3.7%)	RR 1.07 (0.18 to 6.45)	3 more per 1000 (from 30 fewer to 203 more)	VERY LOW	CRITICAL

Quality	assessmen	t					No of pa	atients	Effect			
No of studie s	Design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecisi on	Other consideratio ns	D3	D2	Relativ e (95% Cl)	Absolu te	Quality	Importane e
3	randomis ed trials	no seriou s risk of bias	no serious inconsistenc y	serious ³	serious ¹²	none	28/522 (5.4%)	38/532 (7.1%)	RR 0.75 (0.47 to 1.2)	18 fewer per 1000 (from 38 fewer to 14 more)	LOW	CRITICAL
Reoper	ation rate											
2	randomis ed trials	seriou s ¹	no serious inconsistenc y ¹³	serious ³	very serious ¹⁴	none	10/295 (3.4%)	5/298 (1.7%)	RR 1.77 (0.59 to 5.38)	13 more per 1000 (from 7 fewer to 73 more)	VERY LOW	IMPORTA NT
R0 rese	ection											
1	randomis ed trials	no seriou s risk of bias	no serious inconsistenc y	no serious indirectnes s	no serious imprecisio n ¹⁵	none	260/26 0 (100%)	261/26 3 (99.2 %)	RR 1.01 (0.99 to 1.02)	10 more per 1000 (from 10 fewer to 20 more)	HIGH	CRITICAL

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Quality	assessmen	nt					No of p	atients	Effect			
No of studie s	Design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecisi on	Other consideratio ns	D3	D2	Relativ e (95% Cl)	Absolu te	Quality	Importanc e
0	-	-	-	-	-	none	-	-	-	-		

CI=Confidence interval; RR=relative risk; HR=Hazard ratio; OS=overall survival; DFS=Disease free survival

¹ Risk of bias: Maeta 1999: inappropriate randomisation and attrition rate.

² Heterogeneity: i2=0%

³ Indirectness: postoperative complications could be affected by dissection of additional organs such as pancreatectomy and splenectomy (Yonemura 2008), subgroup analyses have not been presented here. Older studies may not be comparable with newer studies due to differences in surgical technique and experience and post-operative care. Differences in median follow-up time across included studies.

⁴ 95% CI: 0.81-1.21. No default MIDs crossed

- ⁵ Median follow-up 5.7 years
- ⁶ 95% CI: 0.83-1.42. One default MID crossed

⁷ 95% CI: 0.78-5.35. Wide CI crosses two default MIDs

⁸ 95% CI: 071-1.83. Two default MIDs crossed.

⁹ 95% CI: 0.51-1.36. Two default MIDs crossed

¹⁰ Heterogeneity: i2=40%

¹¹ 95% CI: 0.35-2.05. Two default MIDs crossed.

¹² 95% CI: 0.48-1.21. 1 default MID crossed

¹³ Heterogeneity: i2=3%
 ¹⁴ 95% CI: 0.69-5.35. Two default MIDs crossed.

¹⁵ 95% CI: 0.99-1.02.

Table 6: Clinical evidence profile: 3-field lymph node resection versus 2-field lymph node resection for oesophageal cancer

Quality	assessmei	nt				No of pat	ients	Effect				
No of studie s	Design	Risk of bias	Inconsisten cy	Indirectne SS	Imprecisi on	Other consideratio ns	Three field lymph node resectio n	Two field lymph node resectio n	Relativ e (95% CI)	Absolu te	Qualit y	Importanc e

	assessmen	1					No of pat		Effect			
No of studie s	Design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecisi on	Other consideratio ns	Three field lymph node resectio n	Two field lymph node resectio n	Relativ e (95% CI)	Absolu te	Qualit y	Importanc e
2	randomis ed trials	seriou s ¹	no serious inconsistenc y	very serious ²	no serious imprecisio n ³	none	5yr OS 61% (46% to 72%)	5yr OS 33% ¹³	HR 0.46 (0.3 to 0.71)	If 5yr OS is 33% with 2 field it would be 61% with 3 field (95%CI 46% to 72%).	VERY LOW	CRITICAL
	erative mort	ality										
2	randomis ed trials	seriou s ¹	no serious inconsistenc y	very serious ²	serious ⁴	none	3/109 (2.8%)	11/103 (10.7%)	RR 0.27 (0.08 to 0.94)	78 fewer per 1000 (from 6 fewer to 98 fewer)	VERY LOW	IMPORTA NT
Recurr	ent nerve pa	lsy										
2	randomis ed trials	seriou s ¹	very serious5	very serious ²	serious ⁶	none	29/109 (26.6%)	20/103 (19.4%)	RR 1.50 (0.32 to 7.08)	97 more per 1000 (from 132 fewer to	VERY LOW	CRITICAL

Quality	assessmen	t					No of pat	ients	Effect			
No of studie s	Design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecisi on	Other consideratio ns	Three field lymph node resectio n	Two field lymph node resectio n	Relativ e (95% CI)	Absolu te	Qualit y	Importanc e
										1000 more)		
Anasto	motic leak								-			
2	randomis ed trials	seriou S ¹	serious7	very serious ²	very serious ⁸	none	28/109 (25.7%)	23/103 (22.3%)	RR 0.80 (0.18 to 3.51)	45 fewer per 1000 (from 183 fewer to 560 more)	VERY LOW	CRITICAL
Pulmor	nary complic	ation										
1	randomis ed trials	seriou s ¹	no serious inconsistenc y	very serious ²	very serious ⁹	none	6/32 (18.8%)	5/30 (16.7%)	RR 1.13 (0.38 to 3.3)	22 more per 1000 (from 103 fewer to 383 more)	VERY LOW	CRITICAL
Chyloth	norax											
1	randomis ed trials	seriou s ¹	no serious inconsistenc y	serious ²	very serious ¹⁰	none	0/77 (0%)	3/73 (4.1%)	RR 0.14 (0.01 to 2.58)	35 fewer per 1000 (from 41 fewer to	VERY LOW	CRITICAL

Quality	assessmen	t					No of pat	ients	Effect			
No of studie s	Design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecisi on	Other consideratio ns	Three field lymph node resectio n	Two field lymph node resectio n	Relativ e (95% CI)	Absolu te	Qualit y	Importanc e
										65 more)		
Phrenic	c nerve pals	y										
1	randomis ed trials	seriou s ¹	no serious inconsistenc y	no serious indirectnes s	very serious ¹¹	none	4/32 (12.5%)	0/30 (0%)	RR 08.45 (0.47 to 150.66)	-	VERY LOW	CRITICAL
Trache	ostomy											
1	randomis ed trials	seriou s ¹	no serious inconsistenc y	serious ²	very serious ¹²	none	17/32 (53.1%)	3/30 (10%)	RR 5.31 (1.73 to 16.31)	431 more per 1000 (from 73 more to 1000 more)	VERY LOW	CRITICAL

CI=confidence interval; RR=relative risk; HR=Hazard ratio; OS=overall survival

¹ Risk of bias: Kato 1991 provides no details on randomisation method and allocation concealment. Nishihara 1998 also does not report randomisation method and may be subject to small sample size bias (n=62).

² Indirectness: Indirect populations. Kato 1991 includes patients with thoracic oesophageal carcinoma and Nishihara 1998 includes those with thoracic oesophageal carcinoma. Indirect interventions: lymphadenectomy described in Nishihara 1998 may not strictly follow definition in protocol and that defined in other included studies. Procedure and approach of lymphadenectomy would also presumably vary depending on site of primary tumour.

³ 95% CI: 0.30-0.71

⁴ 95% CI: 0.07-0.90. One default MID crossed.

⁵ Heterogeneity: i2=87% therefore very serious inconsistency.

⁶ 95% Cl: 0.82-2.27. Crosses 1 default MID.
 ⁷ Heterogeneity: i2=72%

⁸ 95% CI: 0.71-1.86. Crosses 2 default MIDs.

⁹ 95% CI: 0.38-3.30. Very wide CI, crosses both default MIDs.
 ¹⁰ 95% CI: 0.01-2.58. Very wide CI crosses both default MIDs.
 ¹¹ 95% CI: 0.47-150.66.
 ¹² 95% CI: 1.71-16.31
 ¹³ Assumed risk from Kato (1991)

Table 7: Clinical evidence profile: 3-field lymphadenectomy vs 2-field lymphadenectomy for oesophageal cancer: observational studies

Qualit	y assessmer	nt					No of patients		Effect			
No of stud ies	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecisi on	Other considerati ons	Three field lymphadenect omy	Two field lymphadenect omy	Relati ve (95% Cl)	Absol ute	Qual ity	Importa nce
5 year	· overall surv	ival										
2	observatio nal studies	seriou s ¹	no serious inconsisten cy	no serious indirectne ss	no serious imprecisio n	none	314/476 (66%)	43/86(50%)	-	5 yr. OS was from 13.6% to 38.2% better with 3- field	VER Y LO W	CRITICA L
Anast	omotic leak											
1	observatio nal studies	seriou s ¹	no serious inconsisten cy	no serious indirectne ss	serious ²	none	43/100 (43%)	164/410 (40%)	RR 1.07 (0.83 to 1.39)	28 more per 1000 (from 68 fewer to 156 more)	VER Y LO W	CRITICA L

Qualit	ty assessmer	nt	-	-	-		No of patients		Effect			
No of stud ies	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecisi on	Other considerati ons	Three field lymphadenect omy	Two field lymphadenect omy	Relati ve (95% Cl)	Absol ute	Qual ity	Importa nce
1	observatio nal studies	seriou s ¹	no serious inconsisten cy	no serious indirectne ss	no serious imprecisio n ³	none	15/100 (15%)	19/410 (4.6%)	RR 3.24 (1.71 to 6.14)	104 more per 1000 (from 33 more to 238 more)	VER Y LO W	CRITICA L
Woun	d infection											
1	observatio nal studies	seriou s ¹	no serious inconsisten cy	no serious indirectne ss	very serious ⁴	none	6/100 (6%)	19/410 (4.6%)	RR 1.29 (0.53 to 3.16)	13 more per 1000 (from 22 fewer to 100 more)	VER Y LO W	CRITICA L
Haem	orrhage											
1	observatio nal studies	seriou s ¹	no serious inconsisten cy	no serious indirectne ss	very serious⁵	none	0/100 (0%)	4/410 (0.98%)	RR 0.45 (0.02 to 8.33)	5 fewer per 1000 (from 10 fewer to 72 more)	VER Y LO W	CRITICA L

	ty assessmer	ıt					No of patients		Effect			
No of stud ies	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecisi on	Other considerati ons	Three field lymphadenect omy	Two field lymphadenect omy	Relati ve (95% Cl)	Absol ute	Qual ity	Importa nce
1	observatio nal studies	seriou s ¹	no serious inconsisten cy	no serious indirectne ss	very serious⁵	none	0/100 (0%)	4/410 (0.98%)	RR 0.45 (0.02 to 8.33)	5 fewer per 1000 (from 10 fewer to 72 more)	VER Y LO W	CRITICA L
Any p	ost-operative	complic	ation									
1	observatio nal studies	seriou s ¹	no serious inconsisten cy	no serious indirectne ss	serious ⁶	none	71/100 (71%)	248/410 (60.5%)	RR 1.17 (1.01 to 1.36)	103 more per 1000 (from 6 more to 218 more)	VER Y LO W	CRITICA L
Pneu	monia											
1	observatio nal studies	seriou S ¹	no serious inconsisten cy	no serious indirectne ss	very serious ⁷	none	10/100 (10%)	42/410 (10.2%)	RR 0.98 (0.51 to 1.88)	2 fewer per 1000 (from 50 fewer to 90 more)	VER Y LO W	CRITICA L

n=total number of participants; CI=confidence interval; RR=relative risk; OS=overall survival

Risk of bias: Tabira 1999: moderate overall risk of bias due to critical confounding bias. Kato 1991: serious risk of bias.
 95% CI: 0.83-1.39. Crosses 1 default MID
 95% CI: 1.71-6.14.
 95% CI: 0.53-3.16. Crosses two default MIDs
 95% CI: 0.02-8.33. Crosses two default MIDs
 95% CI: 1.01-1.36. Crosses 1 default MIDs
 95% CI: 1.01-1.36. Crosses 1 defaul MID
 7 Crosses two default MIDs

G.11 Localised oesophageal and gastro-oesophageal junctional adenocarcinoma

What is the optimal choice of chemotherapy or chemoradiotherapy in relation to surgical treatment for people with localised oesophageal and gastro-oesophageal junctional cancer?

Table 6:	Clinical evidence	profile: Compariso	on 1: Preoperative ch	emotherapy versus	postoperative chemotherapy

Quality assessmer	nt						No of patie	nts	Effect			
No of studies	Design	Risk of bias	Inconsist ency	Indirect ness	Impreci sion	Other considera tions	Preoperat ive CT	Postoper ative CT	Relati ve (95% Cl)	Abso lute	Quality	Importanc e
Overall survival												
1	randomi sed trials	serio us ¹	no serious inconsiste ncy	no serious indirectn ess	serious ²	none	54% (43% to 63%)	43%	HR 0.73 (0.54 to 0.99)	-	LOW	CRITICAL
R0 tumour resection	on rate											
1	randomi sed trials	serio us ¹	no serious inconsiste ncy	no serious indirectn ess	no serious impreci sion	none	157/164 (95.7%)	151/166 (91%)	RR 1.05 (0.99 to 1.12)	45 more per 1000 (from 9 fewer to 109	MODERA TE	IMPORTA NT

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Quality assessment						No of patients		Effect				
No of studies	Design	Risk of bias	Inconsist ency	Indirect ness	Impreci sion	Other considera tions	Preoperat ive CT	Postoper ative CT	Relati ve (95% CI)	Abso lute	Quality	Importanc e
										more)		
Progression free	survival											
1	randomi sed trials	serio us ¹	no serious inconsiste ncy	no serious indirectn ess	serious ²	none	45% (34% to 55%)	39%	HR 0.84 (0.63 to 1.12)	-	LOW	CRITICAL
Treatment relate	d mortality											
1	randomi sed trials	serio us ¹	no serious inconsiste ncy	no serious indirectn ess	very serious ³	none	1/153 (0.65%)	2/162 (1.2%)	RR 0.53 (0.05 to 5.78)	6 fewer per 1000 (from 12 fewer to 59 more)	VERY LOW	IMPORTA NT
Anastomotic lea	kage											
1	randomi sed trials	serio us ¹	no serious inconsiste ncy	no serious indirectn ess	very serious ³	none	19/153 (12.4%)	24/162 (14.8%)	RR 0.84 (0.48 to 1.47)	24 fewer per 1000 (from 77 fewer to 70 more)	VERY LOW	CRITICAL

Quality assessment							No of patients		Effect			
No of studies	Design	Risk of bias	Inconsist ency	Indirect ness	Impreci sion	Other considera tions	Preoperat ive CT	Postoper ative CT	Relati ve (95% Cl)	Abso lute	Quality	Importanc e
1	randomi sed trials	serio us ¹	no serious inconsiste ncy	no serious indirectn ess	very serious ³	none	16/153 (10.5%)	20/162 (12.3%)	RR 0.85 (0.46 to 1.57)	19 fewer per 1000 (from 67 fewer to 70 more)	VERY LOW	CRITICAL
Pulmonary comp	lication											
1	randomi sed trials	Serio us ¹²	no serious inconsiste ncy	no serious indirectn ess	very serious ³	none	24/153 (15.7%)	21/162 (13%)	RR 1.21 (0.7 to 2.08)	27 more per 1000 (from 39 fewer to 140 more)	VERY LOW	CRITICAL
Cardiovascular o	omplication	s										
1	randomi sed trials	serio us ¹	no serious inconsiste ncy	no serious indirectn ess	very serious ³	none	4/153 (2.6%)	3/162 (1.9%)	RR 1.41 (0.32 to 6.21)	8 more per 1000 (from 13 fewer to 96	VERY LOW	CRITICAL

Quality assessm	ent						No of patients		Effect			
No of studies	Design	Risk of bias	Inconsist ency	Indirect ness	Impreci sion	Other considera tions	Preoperat ive CT	Postoper ative CT	Relati ve (95% CI)	Abso lute	Quality	Importanc e
									CI)	more	Quality	е

CI=confidence interval; RR=relative risk; HR=Hazard ratio; CT=chemotherapy ¹ Unclear randomisation, allocation concealment and blinding ² 95%CI crossed 1 default MID.

³ 95%CI crossed 2 MIDs.

Table 7: Clinical evidence profile: Comparison 2: Preoperative chemotherapy versus surgery alone

Quality assess	uality assessment							nts	Effect			
No of studies	Design	Risk of bias	Inconsist ency	Indirect ness	Impreci sion	Other considera tions	Preoperat ive CT	Surg ery alon e	Relati ve (95% Cl)	Abso lute	Quality	Importanc e
Overall survival	(Histology su	ıbtype) - SC	C									
4	randomise d trials	serious ¹	no serious inconsiste ncy	no serious indirectn ess	serious ²	none	OS* 10% (7% to 16%)	OS* 16%	HR 0.83 (0.7 to 1)	-	LOW	CRITICAL
Overall survival	(Histology su	ıbtype) - Mi	xed									
1	randomise d trials	serious ¹	no serious inconsiste ncy	no serious indirectn ess	serious ²	none	5 year OS 19% (15% to 24%)	5 year OS 14%	HR 0.84 (0.72 to 0.98)	-	LOW	CRITICAL
Anastomotic lea	aks - SCC											
4	randomise d trials	serious ¹	no serious	no serious	very serious ³	none	13/199 (6.5%)	9/19 2	RR 1.38 (0.64	18 more per	VERY LOW	CRITICAL

Quality assessn	nent						No of patie	nts	Effect			
No of studies	Design	Risk of bias	Inconsist ency	Indirect ness	Impreci sion	Other considera tions	Preoperat ive CT	Surg ery alon e	Relati ve (95% CI)	Abso lute	Quality	Importanc e
			inconsiste ncy	indirectn ess				(4.7 %)	to 2.99)	1000 (from 17 fewer to 93 more)		
Anastomotic lea	aks - Mixed											
1	randomise d trials	serious ¹	no serious inconsiste ncy	no serious indirectn ess	very serious ³	none	23/400 (5.8%)	26/4 02 (6.5 %)	RR 0.89 (0.52 to 1.53)	7 fewer per 1000 (from 31 fewer to 34 more)	VERY LOW	CRITICAL
Anastomotic lea	iks - Cisplatin	+5-FU										
5	randomise d trials	serious ¹	no serious inconsiste ncy	no serious indirectn ess	very serious ³	none	36/599 (6%)	35/5 94 (5.9 %)	RR 1.02 (0.66 to 1.59)	1 more per 1000 (from 20 fewer to 35 more)	VERY LOW	CRITICAL
Cardiac complic	ations - SCC											
2	randomise d trials	serious ¹	no serious inconsiste ncy	no serious indirectn ess	very serious ³	none	21/122 (17.2%)	20/1 21 (16.5 %)	RR 1.04 (0.61	7 more per 1000	VERY LOW	CRITICAL

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Quality assessm	ient						No of patie	nts	Effect			
No of studies	Design	Risk of bias	Inconsist ency	Indirect ness	Impreci sion	Other considera tions	Preoperat ive CT	Surg ery alon e	Relati ve (95% Cl)	Abso lute	Quality	Importanc e
									to 1.77)	(from 64 fewer to 127 more)		
Cardiac complic	ations - Mixe	d										
1	randomise d trials	serious ¹	no serious inconsiste ncy	no serious indirectn ess	very serious ³	none	14/400 (3.5%)	15/4 02 (3.7 %)	RR 0.94 (0.46 to 1.92)	2 fewer per 1000 (from 20 fewer to 34 more)	VERY LOW	CRITICAL
Cardiac complic	ations - Cispl	latin+5FU										
3	randomise d trials	serious ¹	no serious inconsiste ncy	no serious indirectn ess	very serious ³	none	35/522 (6.7%)	35/5 23 (6.7 %)	RR 0.99 (0.65 to 1.53)	1 fewer per 1000 (from 23 fewer to 35 more)	VERY LOW	CRITICAL
Pulmonary com	plications - S	cc										
4	randomise d trials	serious ¹	no serious inconsiste ncy	serious	very serious ³	none	44/199 (22.1%)	50/1 92 (26%)	RR 0.86 (0.62	36 fewer per 1000	VERY LOW	CRITICAL

Quality assessme	ent						No of patie	nts	Effect			
No of studies	Design	Risk of bias	Inconsist ency	Indirect ness	Impreci sion	Other considera tions	Preoperat ive CT	Surg ery alon e	Relati ve (95% CI)	Abso lute	Quality	Importanc e
									to 1.21)	(from 99 fewer to 55 more)		
Pulmonary comp	lications - M	lxed										
1	randomise d trials	serious ¹	no serious inconsiste ncy	no serious indirectn ess	very serious ³	none	56/400 (14%)	58/4 02 (14.4 %)	RR 0.97 (0.69 to 1.36)	4 fewer per 1000 (from 45 fewer to 52 more)	VERY LOW	CRITICAL
Pulmonary comp	lications - Ci	splatine+5	FU									
5	randomise d trials		no serious inconsiste ncy	no serious indirectn ess	serious ²	none	100/599 (16.7%)	108/ 594 (18.2 %)	RR 0.92 (0.72 to 1.17)	15 fewer per 1000 (from 51 fewer to 31 more)	LOW	CRITICAL
Infectious compl												
2	randomise d trials	serious ¹	no serious inconsiste ncy	no serious indirectn ess	very serious ³	none	7/122 (5.7%)	10/1 21 (8.3 %)	RR 0.69 (0.27 to 1.76)	26 fewer per 1000 (from	VERY LOW	CRITICAL
			ncy	ess				%)				

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Quality assess	nent						No of patie	ents	Effect			
No of studies	Design	Risk of bias	Inconsist ency	Indirect ness	Impreci sion	Other considera tions	Preoperat ive CT	Surg ery alon e	Relati ve (95% Cl)	Abso lute	Quality	Importanc e
										60 fewer to 63 more)		
Infectious comp	lications - Mi	xed										
1	randomise d trials	serious ¹	no serious inconsiste ncy	no serious indirectn ess	serious ²	none	28/522 (5.4%)	42/5 23 (8%)	RR 0.67 (0.42 to 1.06)	27 fewer per 1000 (from 47 fewer to 5 more)	LOW	CRITICAL
Infectious comp	lications - Cis	splatin+5FL	I									
3	randomise d trials	serious ¹	no serious inconsiste ncy	no serious indirectn ess	serious ²	none	28/522 (5.4%)	42/5 23 (8%)	RR 0.67 (0.42 to 1.06)	27 fewer per 1000 (from 47 fewer to 5 more)	LOW	CRITICAL
Postoperative n	nortality - SCC	;										
3	randomise d trials	serious ¹	no serious inconsiste ncy	no serious indirectn ess	very serious ³	none	12/178 (6.7%)	13/1 71 (7.6 %)	RR 0.87 (0.41 to 1.85)	10 fewer per 1000 (from 45	VERY LOW	CRITICAL

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Quality assessr	nent						No of patie	nts	Effect			
No of studies	Design	Risk of bias	Inconsist ency	Indirect ness	Impreci sion	Other considera tions	Preoperat ive CT	Surg ery alon e	Relati ve (95% Cl)	Abso lute	Quality	Importanc e
										fewer to 65 more)		
Postoperative n	nortality - Mix	ed										
1	randomise d trials	serious ¹	no serious inconsiste ncy	no serious indirectn ess	very serious ³	none	36/400 (9%)	40/4 02 (10%)	RR 0.9 (0.59 to 1.39)	10 fewer per 1000 (from 41 fewer to 39 more)	VERY LOW	CRITICAL
Postoperative n	nortality - Cisp	olatin+5-FU										
4	randomise d trials	serious ¹	no serious inconsiste ncy	no serious indirectn ess	very serious ³	none	48/578 (8.3%)	53/5 73 (9.2 %)	RR 0.90 (0.62 to 1.30)	9 fewer per 1000 (from 35 fewer to 28 more)	VERY LOW	CRITICAL
R0 tumour rese	ction rate - SC	CC 00										
4	randomise d trials	serious ¹	no serious inconsiste ncy	no serious indirectn ess	serious ²	none	70/200 (35%)	60/1 95 (30.8 %)	RR 1.14 (0.91 to 1.44)	43 more per 1000 (from 28 fewer	LOW	IMPORTA NT

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Quality assessm	nent						No of patie	nts	Effect			
No of studies	Design	Risk of bias	Inconsist ency	Indirect ness	Impreci sion	Other considera tions	Preoperat ive CT	Surg ery alon e	Relati ve (95% Cl)	Abso lute	Quality	Importanc e
										to 135 more)		
R0 tumour rese	ction rate - Mi	xed										
1	randomise d trials	serious ¹	no serious inconsiste ncy	no serious indirectn ess	no serious impreci sion	none	233/400 (58.3%)	215/ 402 (53.5 %)	RR 1.09 (0.96 to 1.23)	48 more per 1000 (from 21 fewer to 123 more)	MODERA TE	IMPORTA NT
R0 tumour rese	ction rate - Cis	splain+5FU										
4	randomise d trials	1	no serious inconsiste ncy	no serious indirectn ess	serious ²	none	303/600 (50.5%)	275/ 597 (46.1 %)	RR 1.10 (0.99 to 1.23)	46 more per 1000 (from 5 fewer to 106 more)	LOW	IMPORTA NT

CI=confidence interval; RR=relative risk; HR=Hazard ratio; OS=overall survival; 5FU=5-fluouracil; CT=chemotherapy; SCC=squamous cell carcinoma ¹ <u>Ancona 2001, Law 1997, Nygaard 1992, Schlag 1992a, MRC Allum 2009 -</u>Unclear randomisation<u>or/and</u>-allocation concealment and <u>no</u>blinding ² 95%CI crossed 1 default MID. ³ 95%CI crossed 2 default MIDs

Quality as:	sessment						No of patie	nts	Effect			
No of studies	Design	Risk of bias	Inconsist ency	Indirect ness	Impreci sion	Other considerati ons	Postoper ative CT	Surgery alone	Relati ve (95% CI)	Absol ute	Quali ty	Importan ce
Disease fr	ee survival											
1	randomise d trials	serious ¹	no serious inconsiste ncy	no serious indirectn ess	serious ²	none	5 year DFS 55% (43% to 66%)	5 year DFS 45%	HR 0.75 (0.53 to 1.07)	-	LOW	CRITICA L

Cl=confidence interval; HR=Hazard ratio; DFS=Disease free survival; CT=chemotherapy ¹ Unclear randomisation, allocation concealment and blinding ² 95%Cl crossed 1 default MID

Table 9: Clinical evidence profile. Comparison 4: Perioperative chemotherapy versus preoperative chemotherapy

Quality assessm	uality assessment								No of patients Effect Perioperat Preopera Relati Abs			
No of studies	Design	Risk of bias	Inconsist ency	Indirect ness	Impreci sion	Other considerat ions	Perioperat ive CT	Preopera tive CT	Relati ve (95% CI)	Abso lute	Quali ty	Importa nce
Overall survival												
1	randomised trials	serio us ¹	no serious inconsiste ncy	no serious indirectn ess	serious ²	none	5 year OS 30% (22% to 39%)	5 year OS 22%	HR 0.79 (0.62 to 1)	-	LOW	CRITICA L
Relapse free sur	vival											
1	randomised trials	serio us¹	no serious inconsiste ncy	no serious indirectn ess	serious ²	none	5 year RFS 36% (28% to 43%)	5 year RFS 19%	HR 0.62 (0.51 to 0.76)	-	LOW	CRITICA L

Cl=confidence interval; HR=hazard ratio; CT=confidence interval; OS=overall survival; RFS=relapse free survival ¹ Unclear randomisation, allocation concealment and blinding ² 95%Cl crossed 1 default MID.

Table 10: Clinical evidence profile. Comparison 5: Perioperative chemotherapy vs surgery alone

Quality assess	ment						No of pati	ents	Effect			
No of studies	Design	Risk of bias	Inconsi stency	Indirec tness	Impre cision	Other conside rations	Perioper ative CT	Sur ger y alo ne	Relat ive (95% CI)	Abs olut e	Quality	Importa nce
Overall surviva	al 👘 👘											
2	randomised trials	serious ¹	serious ²	no serious indirect ness	no seriou s imprec ision	none	5 year OS 25% (21% to 29%)	5 year OS 22%	HR 0.91 (0.81 to 1.03)	-	LOW	CRITICA L
Overall surviva	al - AC											
1	randomised trials	serious ¹	no serious inconsis tency	no serious indirect ness	seriou s ³	none	5 year OS 30% (25% to 35%)	5 year OS 24%	HR 0.85 (0.74 to 0.98)	-	LOW	CRITICA L
Overall surviva	al - Mixed											
1	randomised trials	serious ¹	no serious inconsis tency	no serious indirect ness	seriou s ³	none	5 year OS 18% (12% to 25%)	5 year OS 20%	HR 1.07 (0.87 to 1.32)	-	LOW	CRITICA L
Disease free s	urvival											
2	randomised trials	serious ¹	serious ²	no serious indirect ness	seriou s ³	none	5 year DFS 23% (18% to 29%)	5 year DFS 18%	HR 0.85 (0.72 to 1)	-	VERY LOW	CRITICA L

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Quality assess	ment						No of pati	ents	Effect			
No of studies	Design	Risk of bias	Inconsi stency	Indirec tness	Impre cision	Other conside rations	Perioper ative CT	Sur ger y alo ne	Relat ive (95% CI)	Abs olut e	Quality	Importa nce
Disease free su	ırvival - AC											
1	randomised trials	serious ¹	no serious inconsis tency	no serious indirect ness	seriou s ³	none	5 year DFS 34% (23% to 45%)	5 year DFS 24%	HR 0.65 (0.48 to 0.89)	-	LOW	CRITICA L
Disease free su	ırvival - Mixed											
1	randomised trials	serious ¹	no serious inconsis tency	no serious indirect ness	no seriou s imprec ision	none	5 year DFS 22% (16% to 29%)	5 year DFS 20%	HR 0.94 (0.77 to 1.13)	-	MODER ATE	CRITICA L
Any complicati	ons - AC											
1	randomised trials	serious ¹	no serious inconsis tency	no serious indirect ness	seriou s ³	none	28/113 (24.8%)	21/1 11 (18. 9%)	RR 1.31 (0.79 to 2.16)	59 mor e per 1000 (fro m 40 fewe r to 219 mor e)	LOW	CRITICA L
Postoperative I	mortality											
2	randomised trials	serious ¹	no serious	no serious	very seriou s ⁴	none	15/346 (4.3%)	18/3 45	RR 0.83 (0.43	9 fewe r per	VERY LOW	IMPORT ANT

Quality assess	ment						No of pati	ents	Effect			
No of studies	Design	Risk of bias	Inconsi stency	Indirec tness	Impre cision	Other conside rations	Perioper ative CT	Sur ger y alo ne	Relat ive (95% CI)	Abs olut e	Quality	Importa nce
			inconsis tency	indirect ness				(5.2 %)	to 1.62)	1000 (fro m 30 fewe r to 32 mor e)		
Postoperative	mortality - AC											
1	randomised trials	serious ¹	no serious inconsis tency	no serious indirect ness	very seriou s ⁴	none	5/113 (4.4%)	5/11 1 (4.5 %)	RR 0.98 (0.29 to 3.3)	1 fewe r per 1000 (fro m 32 fewe r to 104 mor e)	VERY LOW	IMPOR [®] ANT
Postoperative	mortality - Mixed											
1	randomised trials	serious ¹	no serious inconsis tency	no serious indirect ness	very seriou s ⁴	none	10/233 (4.3%)	13/2 34 (5.6 %)	RR 0.77 (0.35 to 1.73)	13 fewe r per 1000 (fro m 36 fewe r to 41	VERY LOW	IMPOR [®] ANT

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Quality assess	ment						No of pati	ents	Effect			
No of studies	Design	Risk of bias	Inconsi stency	Indirec tness	Impre cision	Other conside rations	Perioper ative CT	Sur ger y alo ne	Relat ive (95% CI)	Abs olut e	Quality	Importa nce
										mor e)		
R0 tumour res	ection rate											
2	randomised trials	serious ¹	serious ²	no serious indirect ness	seriou s ³	none	228/346 (65.9%)	216/ 345 (62. 6%)	RR 1.07 (0.92 to 1.25)	44 mor e per 1000 (fro m 50 fewe r to 157 mor e)	VERY LOW	IMPORT ANT
R0 tumour res	ection rate - AC											
1	randomised trials	serious ¹	no serious inconsis tency	no serious indirect ness	seriou S ³	none	95/113 (84.1%)	81/1 11 (73 %)	RR 1.15 (1 to 1.32)	109 mor e per 1000 (fro m 0 mor e to 234 mor e)	LOW	IMPORT ANT

Quality assess	ment						No of pati	ents	Effect			
No of studies	Design	Risk of bias	Inconsi stency	Indirec tness	Impre cision	Other conside rations	Perioper ative CT	Sur ger y alo ne	Relat ive (95% CI)	Abs olut e	Quality	Importa nce
1	randomised trials	serious ¹	no serious inconsis tency	no serious indirect ness	no seriou s imprec ision	none	133/233 (57.1%)	135/ 234 (57. 7%)	RR 0.99 (0.85 to 1.16)	6 fewe r per 1000 (fro m 87 fewe r to 92 mor e)	MODER ATE	IMPORT ANT

CI=confidence interval; RR=relative risk; HR=Hazard ratio; AC=adenocarcinoma; OS=overall survival; DFS=disease free survival; CT=chemotherapy ¹ <u>Ychou 2011, Kelsen 1998 -</u> Unclear randomisation; <u>or</u> allocation concealment and <u>unclear</u> blinding ² I2>50%

³ 95%CI crossed 1 default MID

⁴ 95%CI crossed 2 default MIDs

Table 11: Clinical evidence profile. Comparison 6: Preoperative chemoradiotherapy versus preoperative chemotherapy

Quality assessn	nent						No of pati	ents	Effect			
No of studies	Design	Risk of bias	Inconsi stency	Indirec tness	Impre cision	Other consider ations	Preoper ative CT	Pre ope rati ve CR T	Relat ive (95% Cl)	Abs olut e	Quality	Importar ce

Quality assessm	nent						No of pati	ents	Effect			
No of studies	Design	Risk of bias	Inconsi stency	Indirec tness	Impre cision	Other consider ations	Preoper ative CT	Pre ope rati ve CR T	Relat ive (95% CI)	Abs olut e	Quality	Importan ce
1	randomised trials	serious ¹	no serious inconsist ency	no serious indirect ness	very serious 2	none	45% (30% to 59%)	49%	HR 1.11 (0.74 to 1.67)	-	VERY LOW	CRITICA L
Post-operative of	complication: Anasto	omotic leak										
2	randomised trials	serious ¹	no serious inconsist ency	no serious indirect ness	very serious 2	none	12/129 (9.3%)	9/12 7 (7.1 %)	RR 1.32 (0.58 to 3.03)	23 more per 1000 (from 30 fewe r to 144 more)	VERY LOW	CRITICA L
Post-operative of	complication: Anasto	omotic leak - A	С									
1	randomised trials	serious ¹	no serious inconsist ency	no serious indirect ness	very serious 2	none	2/39 (5.1%)	2/36 (5.6 %)	RR 0.92 (0.14 to 6.21)	4 fewe r per 1000 (from 48 fewe r to 289 more)	VERY LOW	CRITICA L

Quality assessme	ent						No of pati	ents	Effect		
No of studies	Design	Risk of bias	Inconsi stency	Indirec tness	Impre cision	Other consider ations	Preoper ative CT	Pre ope rati ve CR T	Relat ive (95% CI)	Abs olut e	Quality
-	mplication: Anasto	motic leak - M	ixed								
1	randomised trials	serious ¹	no serious inconsist ency	no serious indirect ness	very serious 2	none	10/90 (11.1%)	7/91 (7.7 %)	RR 1.44 (0.58 to 3.63)	34 more per 1000 (from 32 fewe r to 202 more)	VERY LOW
Mortality											
2	randomised trials	serious ¹	no serious inconsist ency	no serious indirect ness	very serious 2	none	5/129 (3.9%)	2/12 7 (1.6 %)	RR 2.53 (0.5 to 12.69)	24 more per 1000 (from 8 fewe r to 184 more)	VERY LOW
Mortality - AC											
1	randomised trials	serious ¹	no serious inconsist ency	no serious indirect ness	no serious impreci sion	none	0/39 (0%)	0%	not poole d	not pool ed	MODER ATE

Importan ce

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Quality assessme	ent						No of pati	ients	Effect			
No of studies	Design	Risk of bias	Inconsi stency	Indirec tness	Impre cision	Other consider ations	Preoper ative CT	Pre ope rati ve CR T	Relat ive (95% CI)	Abs olut e	Quality	Importan ce
Mortality - Mixed												
1	randomised trials	serious ¹	no serious inconsist ency	no serious indirect ness	very serious 2	none	5/90 (5.6%)	2/91 (2.2 %)	RR 2.53 (0.5 to 12.69)	34 more per 1000 (from 11 fewe r to 257 more)	VERY LOW	
Wound infection	- AC											
1	randomised trials	serious ¹	no serious inconsist ency	no serious indirect ness	very serious 2	none	5/39 (12.8%)	1/36 (2.8 %)	RR 4.62 (0.57 to 37.64)	101 more per 1000 (from 12 fewe r to 1000 more)	VERY LOW	CRITICA L
R0 resection												
2	randomised trials	serious ¹	no serious inconsist ency	no serious indirect ness	serious 3	none	53/64 (82.8%)	45/6 1 (73. 8%)	RR 1.12 (0.93	89 more per 1000	LOW	IMPORT ANT

Quality assessn	nent						No of pati	ients	Effect			
No of studies	Design	Risk of bias	Inconsi stency	Indirec tness	Impre cision	Other consider ations	Preoper ative CT	Pre ope rati ve CR T	Relat ive (95% CI)	Abs olut e	Quality	Importan ce
									to 1.35)	(from 52 fewe r to 258 more)		
R0 resection - A	C											
1	randomised trials	serious ¹	no serious inconsist ency	no serious indirect ness	serious 3	none	33/39 (84.6%)	29/3 6 (80. 6%)	RR 1.05 (0.85 to 1.29)	40 more per 1000 (from 121 fewe r to 234 more)	LOW	IMPORT ANT
R0 resection - N	lixed											
1	randomised trials	serious ¹	no serious inconsist ency	no serious indirect ness	serious 3	none	20/25 (80%)	16/2 5 (64 %)	RR 1.25 (0.88 to 1.78)	160 more per 1000 (from 77 fewe r to 499	LOW	IMPORT ANT

Quality assessme	ent						No of pati	ents	Effect			
No of studies	Design	Risk of bias	Inconsi stency	Indirec tness	Impre cision	Other consider ations	Preoper ative CT	Pre ope rati ve CR T	Relat ive (95% CI)	Abs olut e	Quality	Importan ce
										more)		
Cardiac complica	itions											
2	randomised trials	serious ¹	no serious inconsist ency	no serious indirect ness	very serious 2	none	14/129 (10.9%)	10/1 27 (7.9 %)	RR 1.35 (0.63 to 2.88)	28 more per 1000 (from 29 fewe r to 148 more)	VERY LOW	CRITICA L
Cardiac complica	tions - AC											
1	randomised trials	serious ¹	no serious inconsist ency	no serious indirect ness	very serious 2	none	7/39 (17.9%)	6/36 (16. 7%)	RR 1.08 (0.4 to 2.9)	13 more per 1000 (from 100 fewe r to 317 more)	VERY LOW	CRITICA L
Cardiac complica	tions - Mixed											

Quality assessm	ent						No of pati	ients	Effect			
No of studies	Design	Risk of bias	Inconsi stency	Indirec tness	Impre cision	Other consider ations	Preoper ative CT	Pre ope rati ve CR T	Relat ive (95% CI)	Abs olut e	Quality	Importan ce
1	randomised trials	serious ¹	no serious inconsist ency	no serious indirect ness	very serious 2	none	7/90 (7.8%)	4/91 (4.4 %)	RR 1.77 (0.54 to 5.84)	34 more per 1000 (from 20 fewe r to 213 more)	VERY LOW	CRITICA L
Poor Tumour Re	gression Grade (TR	G >2 or Tumou	ur cells > 5	0%)								
2	randomised trials	serious ¹	no serious inconsist ency	no serious indirect ness	serious 3	none	64/129(4 9.6%)	99/1 27 (78 %)	RR 0.66 (0.49 to 0.90)	265 fewe r per 1000 (from 78 fewe r to 398 fewe r)	LOW	IMPORT ANT
Poor TRG - AC												
1	randomised trials	serious ¹	no serious inconsist ency	no serious indirect ness	serious 1	none	27/39 (69.2%)	33/3 6 (91. 7%)	RR 0.76 (0.60 to 0.95)	220 fewe r per 1000 (from	LOW	IMPORT ANT

Quality assessme	ent						No of pati	ents	Effect			
No of studies	Design	Risk of bias	Inconsi stency	Indirec tness	Impre cision	Other consider ations	Preoper ative CT	Pre ope rati ve CR T	Relat ive (95% CI)	Abs olut e	Quality	Importan ce
										46 fewe r to 367 fewe r)		
Poor TRG - Mixed	i i					-			-			
1	randomised trials	serious ¹	no serious inconsist ency	no serious indirect ness	serious 3	none	37/90 (41.1%)	66/9 1 (72. 5%)	RR 0.57 (0.43 to 0.75)	312 fewe r per 1000 (from 181 fewe r to 413 fewe r)	LOW	IMPORT ANT
Treatment-related	d morbidity: Any co	mplication (Mi	xed)									
1	randomised trials	serious ¹	no serious inconsist ency	no serious indirect ness	serious 3	none	42/90 (46.7%)	35/9 1 (38. 5%)	RR 1.21 (0.86 to 1.71)	81 more per 1000 (from 54 fewe r to 273 more)	LOW	IMPORT ANT

CI=confidence interval; RR=relative risk; HR=hazard ratio; TRG=tumour regression grade; AC=adenocarcinoma; CT=chemotherapy; CRT=chemoradiotherapy; ¹ <u>Burmeister 2011, Klevebro 2015 -</u> Unclear randomisation <u>and/or</u>, allocation concealment and <u>unclear</u> blinding ² 95%CI crossed 2 default MID ³ 95%CI crossed 1 default MID ⁴ I2>80%

Table 12: Clinical evidence profile. Comparison 7: Preoperative chemoradiotherapy versus surgery alone

Quality	assessment						No of patien	te	Effect				_	
No of studie s	Design	Risk of bias	Inconsisten cy	Indirectness	Impre cision	Othe r cons idera tions	Preoperati ve CRT	Surgery alone	Relat ive (95% CI)		Quality	Importa nce	F	ormatted Table
Post-op	erative com	plication: Anastor	notic leak									4	F	ormatted Table
6	randomis ed trials	serious ¹	no serious inconsistency	no serious indirectness	very seriou s ²	none	13/237 (5.5%)	10/255 (3.9%)	RR 1.44 (0.69 to 3.01)	17 mor e per 1000 (fro m 12 fewe r to 79 mor e)	VERY LOW	CRITICA L		
Post-op	erative com	plication: Anastor	notic leak - SCC	;								4	F	ormatted Table
5	randomis ed trials	serious ¹	no serious inconsistency	no serious indirectness	very seriou s ²	none	11/211 (5.2%)	10/229 (4.4%)	RR 1.26 (0.58 to 2.74)	11 mor e per 1000 (fro m 18 fewe r to 76	VERY LOW	CRITICA L		

Quality	assessment						No of patien	ts	Effect			4	Formatted Table
No of studie s	Design	Risk of bias	Inconsisten cy	Indirectness	Impre cision	Othe r cons idera tions	Preoperati ve CRT	Surgery alone	Relat ive (95% Cl)	Abs olut e	Quality	Importa nce	
										mor e)			
Post-op		plication: Anasto										4	Formatted Table
1	randomis ed trials	serious ¹	no serious inconsistency	no serious indirectness	very seriou s ²	none	2/26 (7.7%)	0/26 (0%)	RR 5 (0.25 to 99.3 4)	-	VERY LOW	CRITICA L	
Post-op	erative com	plication: Anasto	motic leak - =</td <td>40Gy RT</td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td>4</td> <td>Formatted Table</td>	40Gy RT								4	Formatted Table
5	randomis ed trials		no serious inconsistency		very seriou s ²	none	11/211 (5.2%)	10/229 (4.4%)	RR 1.26 (0.58 to 2.74)	11 mor e per 1000 (fro m 18 fewe r to 76 mor e)	VERY LOW	CRITICA L	
Post-op	erative com	plication: Anasto	motic leak - >40	Gy RT								4	Formatted Table
1	randomis ed trials	serious ¹	no serious inconsistency	no serious indirectness	very seriou s ²	none	2/26 (7.7%)	0/26 (0%)	RR 5 (0.25 to 99.3 4)	-	VERY LOW	CRITICA L	

Quality	assessment						No of patien	its	Effect			4
No of studie s	Design	Risk of bias	Inconsisten cy	Indirectness	Impre cision	Othe r cons idera tions	Preoperati ve CRT	Surgery alone	Relat ive (95% CI)	Abs olut e	Quality	Importa nce
4	randomis ed trials	serious ¹	no serious inconsistency	no serious indirectness	seriou S ³	none	90/289 (31.1%)	98/316 (31%)	RR 1.02 (0.8 to 1.29)	6 mor e per 1000 (fro m 62 fewe r to 90 mor e)	LOW	
Any pos	st-operative	complication - Si	ngle drug CT									•
1	randomis ed trials	serious ¹	no serious inconsistency	no serious indirectness	seriou s ³	none	45/138 (32.6%)	36/137 (26.3%)	RR 1.24 (0.86 to 1.79)	63 mor e per 1000 (fro m 37 fewe r to 208 mor e)	LOW	CRITICA L
Any pos	st-operative	complication - Do	ouble drug CT									٩
3	randomis ed trials	serious ¹	no serious inconsistency	no serious indirectness	very seriou s²	none	45/151 (29.8%)	62/179 (34.6%)	RR 0.88 (0.65	42 fewe r per 1000	VERY LOW	CRITICA L

Quality No of studie s	assessment Design	Risk of bias	Inconsisten cy	Indirectness	Impre cision	Othe r cons idera tions	No of patier Preoperati ve CRT	nts Surgery alone	Effect Relat ive (95% CI)	Abs olut e	Quality	Importa nce	Formatted Table
									to 1.2)	(fro m 121 fewe r to 69 mor e)			
Any pos	st-operative	complication - </td <td>=40Gy RT</td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td>•</td> <td>Formatted Table</td>	=40Gy RT									•	Formatted Table
2	randomis ed trials		no serious inconsistency	no serious indirectness	seriou s ²	none	59/173 (34.1%)	54/179 (30.2%)	RR 1.15 (0.84 to 1.55)	45 mor e per 1000 (fro m 48 fewe r to 166 mor e)	LOW	CRITICA L	
Any pos	st-operative	complication - >4	0Gy RT									4	Formatted Table
2	randomis ed trials	serious ¹	no serious inconsistency	no serious indirectness	very seriou s ²	none	31/116 (26.7%)	44/137 (32.1%)	RR 0.85 (0.58 to 1.25)	48 fewe r per 1000 (fro m 135 fewe r to	VERY LOW	CRITICA L	

Quality a	assessment						No of patien	ts	Effect			4	 Formatted Table
No of studie s	Design	Risk of bias	Inconsisten cy	Indirectness	Impre cision	Othe r cons idera tions	Preoperati ve CRT	Surgery alone	Relat ive (95% CI)	Abs olut e	Quality	Importa nce	
										80 mor e)			
30-day r	nortality											4	Formatted Table
3	randomis ed trials	serious ¹	no serious inconsistency	no serious indirectness	seriou s ³	none	11/151 (7.3%)	5/159 (3.1%)	RR 2.28 (0.82 to 6.34)	40 mor e per 1000 (fro m 6 fewe r to 168 mor e)	LOW	IMPORT ANT	
30-day n	nortality - S	cc										4	Formatted Table
2	randomis ed trials	serious ¹	no serious inconsistency	no serious indirectness	seriou s ³	none	10/131 (7.6%)	4/139 (2.9%)	RR 2.6 (0.85 to 8)	46 mor e per 1000 (fro m 4 fewe r to 201 mor e)	LOW	IMPORT ANT	

Quality	assessment						No of patien	ts	Effect			4	Formatted Table
No of studie s	Design	Risk of bias	Inconsisten cy	Indirectness	Impre cision	Othe r cons idera tions	Preoperati ve CRT	Surgery alone	Relat ive (95% CI)	Abs olut e	Quality	Importa nce	
1	randomis ed trials	serious ¹	no serious inconsistency	no serious indirectness	very seriou s ²	none	1/20 (5%)	1/20 (5%)	RR 1 (0.07 to 14.9)		VERY LOW	IMPORT ANT	
30-day r	mortality - </th <th>=40Gy RT</th> <th></th> <th></th> <th></th> <th></th> <th></th> <th></th> <th></th> <th></th> <th></th> <th>•</th> <th>Formatted Table</th>	=40Gy RT										•	Formatted Table
2	randomis ed trials		no serious inconsistency	no serious indirectness	very seriou s ²	none	5/70 (7.1%)	4/70 (5.7%)	RR 1.25 (0.35 to 4.46)	14 mor e per 1000 (fro m 37 fewe r to 198 mor e)	VERY LOW	IMPORT ANT	
30-day r	mortality - >4	40Gy RT										4	Formatted Table
1	randomis ed trials	serious ¹	no serious inconsistency	no serious indirectness	very seriou s ²	none	6/81 (7.4%)	1/89 (1.1%)	RR 6.59 (0.81 to	63 mor e per 1000	VERY LOW	IMPORT ANT	

Quality	assessment	t					No of patien	its	Effect			4	Formatted Table
No of studie s	Design	Risk of bias	Inconsisten Cy	Indirectness	Impre cision	Othe r cons idera tions	Preoperati ve CRT	Surgery alone	Relat ive (95% Cl)	Abs olut e	Quality	Importa nce	
									53.5 9)	(fro m 2 fewe r to 591 mor e)			
Blood lo	oss in surge	ry (ml) (SCC; doເ	uble; <=40Gy)) (E	Better indicated	by lower	values)					4	Formatted Table
1	randomis ed trials	serious ¹	no serious inconsistency	no serious indirectness	seriou s ⁴	none	50	50	-	MD 10 high er (1.9 2 to 18.0 8 high er)	LOW	CRITICA L	
R0/T0 r	esection rate)										-4	Formatted Table
8	randomis ed trials	serious ¹	very serious⁵	no serious indirectness	seriou s ³	none	508/672 (75.6%)	408/687 (59.4%)	RR 1.23 (1.08 to 1.40)	137 mor e per 1000 (fro m 48 mor e to 238 mor e)	VERY LOW	IMPORT ANT	

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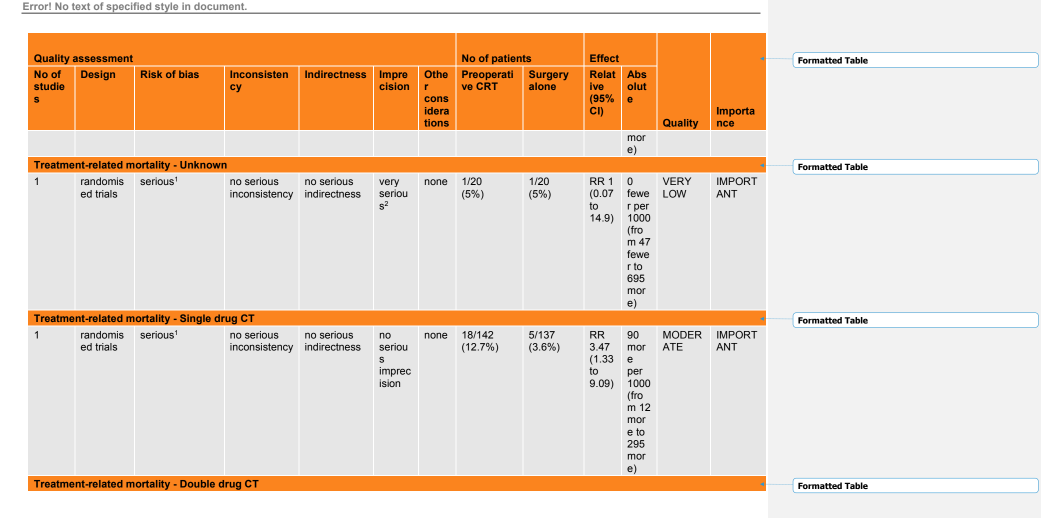
Quality	assessment	t in the second s					No of patien	nts	Effect			4	Formatted Table	
No of studie s	Design	Risk of bias	Inconsisten cy	Indirectness	Impre cision	Othe r cons idera tions	Preoperati ve CRT	Surgery alone	Relat ive (95% CI)	1	Quality	Importa nce		
R0/T0 r/	esection rate	e - SCC										4	Formatted Table	
5	randomis ed trials	serious ¹	no serious inconsistency	no serious indirectness	seriou S ³	none	221/347 (63.7%)	189/358 (52.8%)		95 mor e per 1000 (fro m 32 fewe r to 253 mor e)	LOW	IMPORT ANT		
R0/T0 r/	esection rate	e - AC										-	Formatted Table	
1	randomis ed trials	serious ¹		no serious indirectness	seriou s ³	none	36/36 (100%)	32/40 (80%)	1.24 (1.09 to 1.42)	192 mor e per 1000 (fro m 72 mor e to 336 mor e)	LOW	IMPORT ANT	٤T	
R0/T0 r	esection rate	e - Mixed										•	Formatted Table	
2	randomis ed trials	serious ¹	no serious inconsistency	no serious indirectness	seriou s ³	none	251/289 (86.9%)	187/289 (64.7%)	1.34 (1.24	220 mor e	LOW	IMPORT ANT	۲۲	

Juality	assessment						No of patient	its	Effect			4	Formatted Table
No of studie s	Design	Risk of bias	Inconsisten cy	Indirectness	cision	Othe r cons idera tions	ve CRT	Surgery alone	Relat ive (95% Cl)	Abs olut e	Quality	Importa nce	
									to 1.45)	per 1000 (fro m 155 mor e to 291 mor e)			
R0/T0 r	esection rate	e - Single drug CT										4	Formatted Table
1	randomis ed trials	serious ¹	no serious inconsistency	no serious indirectness	no seriou s imprec ision	none	29/112 (25.9%)	0/94 (0%)	49.6 (4.8 to 512. 16)	-	MODER ATE	IMPORT ANT	
RO/TO r	esection rate	e - Double drug C	r										Formatted Table
7	randomis ed trials	serious ¹	serious ⁶	no serious indirectness	seriou S ³	none	479/560 (85.5%)	408/593 (68.8%)	1.21 (1.09 to 1.33)	144 mor e per 1000 (fro m 62 mor e to 227 mor e)	VERY LOW	IMPORT ANT	

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Quality	assessment						No of patien		Effect			4	Formatted Table
lo of tudie	Design	Risk of bias	Inconsisten cy	Indirectness	Impre cision	Othe r cons idera tions	Preoperati ve CRT	Surgery alone	Relat ive (95% CI)	Abs olut e	Quality	Importa nce	
ŀ	randomis ed trials	serious ¹	very serious⁵	no serious indirectness	seriou s ³	none	213/359 (59.3%)	141/349 (40.4%)	1.49 (1.01 to 2.17)	198 mor e per 1000 (fro m 4 mor e to 473 mor e)	VERY LOW	IMPORT ANT	
0/T0 re	esection rate	e - >40Gy RT										-	Formatted Table
k	randomis ed trials	serious ¹	very serious⁵	no serious indirectness	seriou s ³	none	295/313 (94.2%)	267/338 (79%)	1.17 (1.04 to 1.32)	134 mor e per 1000 (fro m 32 mor e to 253 mor e)	VERY LOW	IMPORT ANT	
reatme	ent-related m	nortality										4	Formatted Table
3	randomis ed trials	serious ¹	no serious inconsistency	no serious indirectness	seriou s ³	none	34/417 (8.2%)	16/410 (3.9%)	RR 2.03 (1.16	40 mor e per	LOW	IMPORT ANT	

Quality No of studie s	assessment Design	Risk of bias	Inconsisten cy	Indirectness	Impre cision	Othe r cons idera tions	No of patien Preoperati ve CRT	nts Surgery alone	Effect Relat ive (95% CI)	Abs olut e	Quality	Importa nce	Formatted Table
									to 3.55)	1000 (fro m 6 mor e to 100 mor e)			
Treatme	ent-related m	ortality - SCC										Formatted Table	
6	randomis ed trials	serious ¹	no serious inconsistency	no serious indirectness	seriou s ³	none	32/369 (8.7%)	14/364 (3.8%)	RR 2.17 (1.2 to 3.91)	45 mor e per 1000 (fro m 8 mor e to 112 mor e)	LOW	IMPORT ANT	
Treatme	1	ortality - Mixed										4	Formatted Table
1	randomis ed trials	serious ¹	no serious inconsistency	no serious indirectness	very seriou s ²	none	1/28 (3.6%)	1/26 (3.8%)	RR 0.93 (0.06 to 14.0 9)	3 fewe r per 1000 (fro m 36 fewe r to 503	VERY LOW	IMPORT ANT	



Quality	assessment			-	-		No of patien	Effect			4	Formatted Table	
No of studie s	Design	Risk of bias	Inconsisten cy	Indirectness	Impre cision	Othe r cons idera tions	Preoperati ve CRT	Surgery alone	Relat ive (95% CI)	Abs olut e	Quality	Importa nce	
	randomis ed trials	no serious risk of bias	no serious inconsistency	no serious indirectness	very seriou s ²	none	16/275 (5.8%)	11/273 (4%)	RR 1.28 (0.61 to 2.66)	11 mor e per 1000 (fro m 16 fewe r to 67 mor e)	LOW	IMPORT ANT	
reatme	ent-related m	nortality - =40Gy</td <td>/ RT</td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td>4</td> <td>Formatted Table</td>	/ RT									4	Formatted Table
5	randomis ed trials	serious ¹	no serious inconsistency	no serious indirectness	seriou s ³	none	31/338 (9.2%)	14/336 (4.2%)	RR 2.11 (1.17 to 3.82)	46 mor e per 1000 (fro m 7 mor e to 118 mor e)	LOW	IMPORT ANT	
Freatme	ent-related m	ortality - >40Gy F	RT									4	Formatted Table
2	randomis ed trials	serious ¹	no serious inconsistency	no serious indirectness	seriou s ³	none	3/79 (3.8%)	2/74 (2.7%)	RR 1.4 (0.24	11 mor e	LOW	IMPORT ANT	

Quality	assessment						No of patien		Effect			1	Formatted Table
No of studie s	Design	Risk of bias	Inconsisten cy	Indirectness	Impre cision	Othe r cons idera tions	ve CRT	Surgery alone	Relat ive (95% Cl)	olut e	Quality	Importa nce	
										1000 (fro m 21 fewe r to 194 mor e)			
ntraope	arative treat	ment-related mor	rbidity: Haemorrh	1age (>300 mL)								4	Formatted Table
1	ed trials	serious ¹	no serious inconsistency	no serious indirectness	seriou S ³	none	8/80 (10%)	2/80 (2.5%)	(0.88 to 18.2 6)	75 mor e per 1000 (fro m 3 fewe r to 432 mor e)	LOW	CRITICA L	
Overall	survival (OS	ه)										•	Formatted Table
9	randomis ed trials	serious ¹	serious ⁶	no serious indirectness	seriou s ³	none	OS* 38% (33% to 42%)	OS* 27%	HR 0.75 (0.67 to 0.84)	-	VERY LOW	CRITICA L	

Quality assessment								No of patients				•	Formatted Table
No of studie s	Design	Risk of bias	Inconsisten cy	Indirectness	Impre cision	Othe r cons idera tions	Preoperati ve CRT	Surgery alone	Relat ive (95% Cl)	Abs olut e	Quality	Importa nce	
7	randomis ed trials	serious ¹	no serious inconsistency	no serious indirectness	seriou s ³	none	OS* 35%(29% to 40%)	OS* 26%	HR 0.79 (0.68 to 0.92)	-	LOW	CRITICA L	
OS - AC	;											4	Formatted Table
2	randomis ed trials	serious ¹	no serious inconsistency	no serious indirectness	seriou s ³	none	5 year OS 44% (35% to 53%)	5 year OS 28%	HR 0.64 (0.5 to 0.82)	-	LOW	CRITICA L	
OS - Miz	xed											4	Formatted Table
2	randomis ed trials	serious ¹	no serious inconsistency	no serious indirectness	seriou s ³	none	5 year OS 31% (21% to 40%)	5 year OS (21%)	HR 0.76 (0.59 to 0.99)	-	LOW	CRITICA L	
OS - Sir	ngle drug CT											4	Formatted Table
1	randomis ed trials	serious ¹	no serious inconsistency	no serious indirectness	very seriou s ²	none	5 year OS 23% (14% to 34%)	5 year OS 22%	HR 0.96 (0.72 to 1.28)	-	VERY LOW	CRITICA L	
OS - Do	uble drug C	Т										4	Formatted Table
8	randomis ed trials	serious ¹	no serious inconsistency	no serious indirectness	no seriou s imprec ision ³	none	OS* 38% (34% to 43%)	OS* 25%	HR 0.69 (0.61 to 0.78)	-	MODER ATE	CRITICA L	

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Disease free survival - Double drug CT

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Quality	assessment			No of patien	ts	Effect							
No of studie s	Design	Risk of bias	Inconsisten cy	Indirectness	Impre cision	Othe r cons idera tions	Preoperati ve CRT	Surgery alone	Relat ive (95% CI)	Abs olut e	Quality	Importa nce	
OS - =40Gy RT</th													
5	randomis ed trials	serious ¹	no serious inconsistency	no serious indirectness	seriou s ³	none	5 year OS 29% (24% to 34%)	5 year OS 20%	HR 0.77 (0.67 to 0.89)	-	LOW	CRITICA L	
OS - >40)Gy RT											-	
4	randomis ed trials	serious ¹	serious ⁶	no serious indirectness	seriou s ³	none	OS* 52% (45% to 58%)	OS* 36%	HR 0.65 (0.54 to 0.79)	-	VERY LOW	CRITICA L	
Disease	Disease free survival - SCC												
3	randomis ed trials	serious ⁵	no serious inconsistency	no serious indirectness	seriou s ³	none	DFS 46% (40% to 52%)	DFS* 34%	HR 0.77 (0.63 to 0.95)	-	LOW	CRITICA L	
Disease	free surviva	al - Single drug CT										<	

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46%(40%

DFS* 33%

(23% to

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to 52%)

DFS* 34% HR

DFS* 31%

-

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(0.47 to 0.86)

HR

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(0.70

Quality a	assessment	t					No of patien	its	Effect			4	Formatted Table
No of studie s	Design	Risk of bias	Inconsisten cy	Indirectness	Impre cision	Othe r cons idera tions	Preoperati ve CRT	Surgery alone	Relat ive (95% CI)	Abs olut e	Quality	Importa nce	
									to 1.25)				
Disease		al - =40Gy RT</td <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td>•</td> <td>Formatted Table</td>										•	Formatted Table
1	ed trials	serious ¹	no serious inconsistency	no serious indirectness	seriou s ³	none	5 year DFS 40% (29% to 51%)	5 year DFS 24%	HR 0.64 (0.47 to 0.86)	-	LOW	CRITICA L	
Disease	free surviva	al - >40Gy RT										4	Formatted Table
2	randomis ed trials	serious ¹	no serious inconsistency	no serious indirectness	very seriou s ²	none	DFS* 33% (23% to 44%)	DFS* 31%	HR 0.94 (0.70 to 1.25)	-	VERY LOW	CRITICA L	
Post-op	erative com	plication: stenos	is									4	Formatted Table
1	randomis ed trials	serious ¹	no serious inconsistency	no serious indirectness	very seriou s ²	none	2/80 (2.5%)	1/80 (1.3%)	RR 2 (0.19 to 21.6 2)	13 mor e per 1000 (fro m 10 fewe r to 258 mor e)	VERY LOW	CRITICA L	

CI=confidence interval; RR=relative risk; HR=hazard ratio; OS=overall survival; DFS=disease free survival; AC=adenocarcinoma; SCC=squamous cell carcinoma; CRT=chemoradiotherapy; CT=chemotherapy; RT=radiotherapy

*OS/DFS was calculated from survival rate at 5 years or, if it was less than 5 years, the survival rate from the last year available. ¹ <u>Apinop 1994, Bass 2014, Bosset 1997, Lee 2004, Lv 2010, Marietter 2014, van Hagen 2012, Burmeister 2005, Tepper 2008 - Unclear randomisation and/or-allocation</u> concealment and <u>unclear</u> blinding

² 95%CI crossed 2 default MIDs

³ 95%CI crossed 1 default MID ⁴ Default MID: +/-7.5ml; 95% CI crossed 1 MID

⁵ 12>80%

6 12>50%

Table 13: Clinical evidence profile. Comparison 8: Postoperative chemoradiotherapy versus postoperative chemotherapy

Quality ass	essment						No of patier	its	Effect			
No of studies	Design	Risk of bias	Inconsisten cy	Indirect ness	Impreci sion	Other considera tions	Postopera tive CRT	Postoper ative CT	Relati ve (95% Cl)	Abso lute	Qual ity	Importa nce
Overall sur	vival											
1	randomised trials	serious ¹	no serious inconsistenc y	no serious indirectn ess	very serious ²	none	5-years OS 37% (9% to 67%)	5-years OS 38%	HR 1.02 (0.42 to 2.44)	-	VER Y LOW	CRITICA L

Cl=confidence interval; HR=hazard ratio; OS=overall survival; CT=chemotherapy; CRT=chemoradiotherapy;

¹ Unclear randomisation, allocation concealment and blinding

² 95%CI crossed 2 default MIDs

Table 14: Clinical evidence profile. Comparison 9: Postoperative chemoradiotherapy versus sugery alone

Quality assessm	nent						No of patier	nts	Effect			
No of studies	Design	Risk of bias	Inconsis tency	Indirect ness	Imprec ision	Other consider ations	Postopera tive CRT	Sur gery alon e	Relati ve (95% CI)	Abso lute	Quality	Importan ce
Number going for	or radical resection											
1	randomised trials	serio us¹	no serious	no serious	no serious	none	61/78 (78.2%)	64/8 0	RR 0.98	16 fewer	MODERA TE	CRITICAL

Quality assess	nent						No of paties	nts	Effect			
No of studies	Design	Risk of bias	Inconsis tency	Indirect ness	Imprec ision	Other consider ations	Postopera tive CRT	Sur gery alon e	Relati ve (95% CI)	Abso lute	Quality	Importan ce
			inconsist ency	indirectn ess	impreci sion			(80 %)	(0.83 to 1.15)	per 1000 (from 136 fewer to 120 more)		
Treatment relate	ed mortality											
1	randomised trials	serio us¹	no serious inconsist ency	no serious indirectn ess	no serious impreci sion	none	0/78 (0%)	0/80 (0%)	No event in either arm	-	MODERA TE	IMPORTA NT
Overall survival												
1	randomised trials	serio us¹	no serious inconsist ency	no serious indirectn ess	serious 2	none	16% (7% to 27%)	10- year OS 6%	HR 0.66 (0.47 to 0.94)	-	LOW	CRITICAL

CI=confidence interval; RR=relative risk; HR=Hazard ratio; CRT=chemoradiotherapy; OS=overall survival ¹ Unclear randomisation, allocation concealment and blinding ² 95%CI crossed 1 default MID.

G.12 Gastric Cancer

What is the optimal choice of chemotherapy of chemoradiotherapy in relation to surgical treatment for gastric cancer?

Quality	/ assessmen	t					No of patien	ts	Effect			
No of studi es	Design	Risk of bias	Inconsist ency	Indirect ness	Impreci sion	Other considera tions	Post-op chemother apy	Post-op chemora diothera py	Relative (95% CI)	Abso lute	Quali ty	Importan ce
Overal	l survival											
6	Randomis ed trials	Serious 1,2,3,4,5,6	No serious inconsiste ncy	No serious indirectn ess	Serious ⁷	None	5-year OS 55% (49% to 61%)	5-year OS 52%	HR 0.91 (0.76 to 1.09)	-	LOW	CRITICA L
Diseas	e-free Surviv	al										
6	Randomis ed trials	Serious 1,2,3,4,5,6	No serious inconsiste ncy	No serious indirectn ess	Serious ⁷	None	5 year DFS 61% (56% to 66%)	5-year DFS 52%	HR 0.75 (0.63 to 0.88)	-	LOW	CRITICA L
Neutro	penia: Grade	3-4										
5	Randomis ed trials	Serious 1,2,3,5,6	No serious inconsiste ncy	No serious indirectn ess	Serious ⁷	None	165/552 (29.9%)	129/527 (24.5%)	RR 1.25 (1.04 to 1.51)	61 more per 1000 (from 10 more to 125 more)	LOW	CRITICA L

Table 15: Clinical evidence profile: Post-operative chemoradiotherapy versus post-operative chemotherapy

Cl=confidence interval; RR=relative risk; HR=hazard ratio; OS=overall survival; DFS=disease free survival;

¹ Bamias 2010: unclear random sequence generation

² Yu 2012: unclear random sequence generation and allocation concealment

³ Kwon 2010: unclear random sequence generation and allocation concealment
 ⁴ Kim 2010: unclear random sequence generation and allocation concealment
 ⁵ Zhu 2012: unclear random sequence generation and allocation concealment
 ⁶ Lee 2012: unclear random sequence generation and allocation concealment
 ⁷ Effect estimate crosses 1 default MID
 ⁸ Effect estimate crosses 2 default MIDs

Table 16: Clinical evidence profile. Post-operative chemotherapy versus surgery alone

Quality as	sessment						No of patients	5	Effect			-	Formatted Table
No of studies	Design	Risk of bias	Inconsiste ncy	Indirectne ss	Impreci sion	Other consider ations	Post-op chemothera py	Surgery alone	Relative (95% CI)	Absol ute	Qualit y	Importan ce	
Overall Su	ırvival											•	Formatted Table
5	Randomis ed trials	Serious 1,2,3,4	Serious5	No serious indirectnes s	No serious imprecis ion ⁶	None	5-year OS 50% (43% to 56%)	5-year OS 39%	HR 0.74 (0.61 to 0.9)	-	LOW	CRITICAL	
Disease-fr	ee survival*											•	Formatted Table
3	Randomis ed trials	Serious ^{1,3}	No serious inconsisten cy	No serious indirectnes s	Serious ⁸	None	5-year DFS 57% (51% to 62%)	5-year DFS 46%	HR 0.73 (0.62 to 0.87)	-	LOW	CRITICAL	
Any toxici	ty: Grade 3-4	L .										•	Formatted Table
1	Randomis ed trials	serious risk of bias	No serious inconsisten cy	No serious indirectnes S	No serious imprecis ion	None	279/496 (56.3%)	30/478 (6.3%)	RR 8.96 (6.28 to 12.78)	500 more per 1000 (from 331 more to 739 more)	HIGH	CRITICAL	
Neutroper	nia: Grade 3-4	4										-	Formatted Table
1	Randomis ed trials	No serious risk of bias	No serious inconsisten cy	No serious indirectnes s	No serious imprecis ion	None	107/496 (21.6%)	1/478 (0.21%)	RR 103.12 (14.45 to 735.8)	214 more per 1000	HIGH	CRITICAL	

Quality as	sessment						No of patients		Effect			
No of studies	Design	Risk of bias	Inconsiste ncy	Indirectne ss	Impreci sion	Other consider ations	Post-op chemothera py	Surgery alone	Relative (95% CI)	Absol ute	Qualit y	Importan ce
										(from 28 more to 1000 more)		
Treatmen	t-related mor	tality										
3	Randomis ed trials	Serious 1,2,3	No serious inconsisten cy	No serious indirectnes s	Serious ⁸	None	7/350 (2%)	1/364 (0.27%)	RR 4.22 (0.91 to 19.59)	9 more per 1000 (from 0 fewer to 51 more)	LOW	IMPORTA NT

95%CI=95% Confidence interval; OS=Overall survival; DFS=Disease free survival; RR=relative risk; HR=Hazard ratio;

³ Bouche 2005: unclear random sequence generation and allocation concealment
 ² Chipponi 2004: unclear allocation concealment
 ³ Di Costanzo 2008: high risk of attrition bias, unclear random sequence generation and allocation concealment,
 ⁴ Neri 2001: unclear random sequence generation and allocation concealment

⁵ *I-squared statistic* > 50%

⁶ Statistical significance used as MID
 ⁷ No explanation was provided
 ⁸ HR crosses one default MID

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Table 17: Clinical evidence profile. Pre-operative chemotherapy versus surgery alone

Quality	assessmer	nt					No of patients	i	Effect			
No of studie s	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecis ion	Other considerati ons	Pre-op chemothera py	Surg ery alone	Relativ e (95% Cl)	Absol ute	Qualit y	Importance
Overall	survival											
1	Randomi sed trials	Serious ¹	No serious inconsisten cy	No serious indirectne ss	Very serious ²	None	5-year OS 54% (37% to 68%)	5- year OS 48%	HR 0.84 (0.53 to 1.35)	-	VERY LOW	CRITICAL
Progres	ssion-free s	urvival										
1	Randomi sed trials	Serious ¹	No serious inconsisten cy	No serious indirectne ss	Serious ³	None	5-year PFS 48% (32% to 62%)	5- year PFS 38%	HR 0.76 (0.5 to 1.17)	-	LOW	CRITICAL
Death a	at end of fol	low-up										
3	Randomi sed trials	Serious 1,4,5	No serious inconsisten cy	No serious indirectne ss	Serious ⁶	None	84/193 (43.5%)	48.6 %	RR 0.92 (0.74 to 1.14)	39 fewer per 1000 (from 126 fewer to 68 more)	LOW	CRITICAL
R0 rese	ection											
2	Randomi sed trials	Serious ^{1,4}	Serious ⁷	No serious indirectne ss	Serious ⁶	None	133/163 (81.6%)	114/1 52 (75%)	RR 1.09 (0.87 to 1.36)	68 more per 1000 (from 97 fewer	VERY LOW	IMPORTAN T

Quality	assessmer	nt					No of patients	;	Effect			
No of studie s	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecis ion	Other considerati ons	Pre-op chemothera py	Surg ery alone	Relativ e (95% Cl)	Absol ute	Qualit y	Importance
										to 270 more)		
Toxicity	y: Grade 3-4	ļ.										
1	Randomi sed trials	Serious ⁴	No serious inconsisten cy	No serious indirectne ss	Very serious ⁸	None	5/27 (18.5%)	0/1 (0%)	RR 0.79 (0.06 to 9.71)	-	VERY LOW	CRITICAL
Post-op	o complicati	ion (any)										
1	Randomi sed trials	Serious ¹	No serious inconsisten cy	No serious indirectne ss	Serious ⁶	None	19/70 (27.1%)	11/68 (16.2 %)	RR 1.68 (0.86 to 3.26)	110 more per 1000 (from 23 fewer to 366 more)	LOW	CRITICAL
Anasto	motic Leak											
2	Randomi sed trials	Serious ¹	No serious inconsisten cy	No serious indirectne ss	Very serious ⁸	None	3/117 (2.6%)	2/84 (2.4%)	RR 1.46 (0.25 to 8.45)	11 more per 1000 (from 18 fewer to 177 more)	VERY LOW	CRITICAL

Quality	assessmen	it					No of patients		Effect			
No of studie s	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecis ion	Other considerati ons	Pre-op chemothera py	Surg ery alone	Relativ e (95% Cl)	Absol ute	Qualit y	Importance
2	Randomi sed trials	Serious ^{1,9}	No serious inconsisten cy	No serious indirectne ss	Very serious ⁸	None	3/117 (2.6%)	1/84 (1.2%)	RR 1.57 (0.24 to 10.29)	7 more per 1000 (from 9 fewer to 111 more)	VERY LOW	CRITICAL
Post-op	o pneumonia	a										
1	Randomi sed trials	Serious ⁹	No serious inconsisten cy	No serious indirectne ss	Very serious ⁸	None	0/47 (0%)	1/16 (6.3%)	RR 0.12 (0.01 to 2.76)	55 fewer per 1000 (from 62 fewer to 110 more)	VERY LOW	CRITICAL
Transfu	usion											
1	Randomi sed trials	Serious ¹	No serious inconsisten cy	No serious indirectne ss	Serious ⁶	None	10/70 (14.3%)	4/68 (5.9%)	RR 2.43 (0.8 to 7.37)	84 more per 1000 (from 12 fewer to 375 more)	LOW	CRITICAL

Quality	assessmen	it					No of patients		Effect			
No of studie s	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecis ion	Other considerati ons	Pre-op chemothera py	Surg ery alone	Relativ e (95% Cl)	Absol ute	Qualit y	Importance
1	Randomi sed trials	Serious ¹	No serious inconsisten cy	No serious indirectne ss	Very serious ⁸	None	3/70 (4.3%)	1/68 (1.5%)	RR 2.91 (0.31 to 27.33)	28 more per 1000 (from 10 fewer to 387 more)	VERY LOW	IMPORTAN T

95%CI=95% Confidence interval; OS=Overall survivalP DFS=Progressionse free survival; RR=relative risk; HR=Hazard ratio; ¹ Schuhmacher 2009: unclear random sequence generation and allocation concealment

² HR crosses 2 MIDs

⁵ HR crosses 2 MIDs
 ³ HR crosses 1 default MID
 ⁴ Kobayahsi 2000: unlcear random allocation
 ⁵ Wang 2000: inadequate allocation concealment, unclear random allocation
 ⁶ Effect estimate crosses 1 MID

⁷ I-squared statistic> 50%

⁸ Effect estimate crosses 2 default MIDs
 ⁹ Imano 2010: unclear random sequence generation

Table 18: Clinical evidence profile. Post-operative chemoradiotherapy versus surgery alone

Quality a	assessmei	nt					No of patients		Effect			
No of studie s	Design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other consideratio ns	Post-op chemoradiotherap y	Surg ery alon e	Relative (95% CI)	Absol ute	Quality	Importan ce

Quality No of studie	assessme r Design	nt Risk of	Inconsiste ncy	Indirectne ss	Imprecisi	Other	No of patients Post-op chemoradiotherap	Surg	Effect Relative (95%	Absol		
S		bias				ns	у	alon e	CI)		Quality	Importan ce
1	Randomi sed trials	Serio us¹	No serious inconsisten cy	No serious indirectne ss	Serious ²	None	6-year OS 15%(9% to 21%)	6- year OS 24%	HR 1.35 (1.09 to 1.67)	-	LOW	CRITICAL
Relaps	e-free survi	ival										
1	Randomi sed trials	Serio us ¹	No serious inconsisten cy	No serious indirectne ss	No serious imprecisi on	None	6-year RFS 11%(7% to 17%)	6- year RFS 24%	HR 1.52 (1.23 to 1.89)	-	MODERAT E	CRITICAL

95%CI=95% Confidence interval; OS=Overall survival; RFS=Relapse free survival; RR=relative risk; HR=Hazard ratio ¹ MacDonald 2001: unclear allocation concealment and random sequence generation ² HR crosses 1 MID

Table 19: Clinical evidence profile. Perioperative chemotherapy versus surgery alone

Quality	v assessmer	nt					No of patients	;	Effect			
No of studi es	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecis ion	Other considerati ons	Peri-op chemothera py	Surg ery alone	Relativ e (95% CI)	Absol ute	Quality	Importance
Overal	l survival											
1	Randomi sed trials	Seriou s ¹	No serious inconsisten cy	No serious indirectne ss	Serious ²	None	5-year OS 35% (28% to 44%)	5- year OS 25%	HR 0.75 (0.6 to 0.93)	-	LOW	CRITICAL
Diseas	e-free surviv	val										

Quality	v assessmer	nt					No of patients	5	Effect			
No of studi es	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecis ion	Other considerati ons	Peri-op chemothera py	Surg ery alone	Relativ e (95% CI)	Absol ute	Quality	Importance
1	Randomi sed trials	Seriou s ¹	No serious inconsisten cy	No serious indirectne ss	Serious ²	None	5-year PFS 31%(23% to 39%)	5- year PFS 17%	HR 0.66 (0.53 to 0.82)	-	LOW	CRITICAL
Curativ	ve resection											
1	Randomi sed trials	Seriou s ¹	No serious inconsisten cy	No serious indirectne ss	No serious imprecisi on	None	169/244 (69.3%)	166/2 50 (66.4 %)	RR 1.04 (0.92 to 1.18)	27 more per 1000 (from 53 fewer to 120 more)	MODERAT E	IMPORTAN T

95%CI=95% Confidence interval; OS=Overall survivalP DFS=Progressionse free survival; RR=relative risk; HR=Hazard ratio ¹ Cunningham 2006: random sequence generation not described ² HR crosses 1 default MID

Table 20 Clinical evidence profile. Perioperative chemotherapy versus Perioperative chemoradiotherapy (postoperative radiation only)

Quality	assessment						No of pa	tients	Effect			
No of studie s	Design	Risk of bias	Inconsisten Cy	Indirectne ss	Imprecisi on	Other consideratio ns	Peri-op CT	Post- op CRT	Relativ e (95% CI)	Absolu te	Qualit v	Importanc e

Quality	assessment						No of par	tients	Effect			
No of studie s	Design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecisi on	Other consideratio ns	Peri-op CT	Post- op CRT	Relativ e (95% Cl)	Absolu te	Qualit y	Importanc e
1	randomise d trials	very serious 1	no serious inconsistenc y	no serious indirectnes s	no serious imprecisio n	none	162/393 (41.2%)	162/39 5 (41%)	RR 1.01 (0.85 to 1.19)	4 more per 1000 (from 62 fewer to 78 more)	LOW	CRITICAL
Haemat	tological toxic	city (grade	e 3 or higher)									
1	randomise d trials	very serious 1	no serious inconsistenc y	no serious indirectnes s	serious ²	none	173/393 (44%)	134/39 5 (33.9%)	RR 1.3 (1.09 to 1.55)	102 more per 1000 (from 31 more to 187 more)	VERY LOW	CRITICAL
GI toxic	ty (grade 3 c	or higher)										
1	randomise d trials	very serious 1	no serious inconsistenc y	no serious indirectnes s	serious ²	none	145/393 (36.9%)	166/39 5 (42%)	RR 0.88 (0.74 to 1.04)	50 fewer per 1000 (from 109 fewer to 17 more)	VERY LOW	CRITICAL

95%CI=95% confidence interval; CT=chemotherapy; CRT=chemoradiotherapy; RR=relative risk; GI=gastrointestinal; post-op=postoperative; peri-op=perioperative ¹ Randomisation method was not described in details and all the outcomes considered were not reported. ² 95%CI crossed one boundary of default MID

Table 21: Clinical evidence profile. Peri-operative chemotherapy versus Perioperative chemoradiotherapy alone (preoperative radiation only)

		- 11										
Quality	assessme	nt					No of patients		Effect			
No of studi es	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecis ion	Other considerati ons	Peri-op chemoradiother apy	Che mot hera py alon	Relative (95% CI)	Absol ute	Quality	Importan
Surgio	ol complico	tional or	astamotic lea	ok				е			Quality	се
1	Randomi sed trial	No seriou s	No serious inconsisten cy	No serious indirectne ss	Very serious ¹	None	4/51	3/54	RR 1.41 (0.33 to 6.00)	23 more per 1000 (from 37 fewer to 278 more)	LOW	CRITICAL
Surgica	al complica	tions: ch	est infection									
1	Randomi sed trial	No seriou s	No serious inconsisten Cy	No serious indirectne ss	Very serious ¹	None	5/51	5/54	RR 1.06 (0.33 to 3.44)	6 more per 1000 (from 62 fewer to 226 more)	LOW	CRITICAL
Surgica	al complica	tions: ov	verall									
1	Randomi sed trial	No seriou s	No serious inconsisten cy	No serious indirectne ss	Very serious ¹	None	11/51	12/5 4	RR 0.97 (0.47 to 2.00)	7 fewer per 1000 (from 118	LOW	CRITICAL

Quality	assessme	nt					No of patients		Effect			
No of studi es	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecis ion	Other considerati ons	Peri-op chemoradiother apy	Che mot hera py alon e	Relative (95% Cl)	Absol ute	Quality	Importan ce
										fewer to 222 more)		
Haema	tological co	omplicati	ons: neutrop	enia								
1	Randomi sed trial	No seriou s	No serious inconsisten cy	No serious indirectne ss	Very serious ¹	None	27/60	24/6 0	RR 1.13 (0.74 to 1.71)	52 more per 1000 (from 104 fewer to 284 more)	LOW	CRITICAL
Haema	tological co	omplicati	ions: overall									
1	Randomi sed trial	No seriou s	No serious inconsisten cy	No serious indirectne ss	Very serious ¹	None	31/60	30/6 0	RR 1.03 (0.73 to 1.47)	15 more per 1000 (from 135 fewer to 235 more)	LOW	CRITICAL
Gastro	intestinal co	omplicat	ions: overall									
1	Randomi sed trial	No seriou s	No serious inconsisten cy	No serious indirectne ss	Very serious ¹	None	18/60	19/6 0	RR 0.95 (0.55 to 1.62)	16 fewer per 1000 (from	LOW	CRITICAL

Quality	assessme	nt					No of patients		Effect			
No of studi es	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecis ion	Other considerati ons	Peri-op chemoradiother apy	Che mot hera py alon e	Relative (95% CI)	Absol ute	Quality	Importan ce
										143 fewer to 196 more)		

95%CI=95% confidence interval; CT=chemotherapy; RR=relative risk;

¹ Leong 2017: RR crosses both MIDs

Table 22: Clinical evidence profile. Intraperitoneal chemotherapy (IPC) versus surgery alone

Quality	assessment						No of patients	5	Effect			
No of studie s	Design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecisi on	Other consideratio ns	IPC	Surg ery alon e	Relativ e (95% Cl)	Absolu te	Quality	Importance
Periope	rative mortal	lity										
3 <u>1</u>	Randomis ed trials	Seriou s ⁶	No serious inconsistenc y	No serious indirectnes s	Very serious ⁴	None	4 <u>3/269</u> <u>135</u> (<u>1.52.</u> <u>2</u> %)	2 <u>1</u> /2 22 <u>133(</u> 0. <u>97</u> <u>5</u> %)	RR <u>1.82.96</u> (0.39 <u>31</u> to <u>8.4328.</u> 05)	7- <u>15</u> more per 1000 (from 5 fewer to 67 <u>203</u> more)	VERY LOW	IMPORTAN T

2	Randomis ed trials	Seriou s ⁷	No serious inconsistenc y	No serious indirectnes s	Serious ² N	lone	12/134 (9%)	(1.1 (RR 6.53 D.87 to 8.94)	62 more per 1000 (from 1 fewer to 539 more)	LOW	CRITICAL
Overall	survival rate											
5	randomised trials	Seriou 6 ⁷	Scrious⁸	no scrious indirectness	Serious ⁴	none	146/ 30 (63.(%)	2	1.8	277 more per 1000 (from 80 more to 570 more)	VERY LOW	CRITICAL
Overall	survival rate		nermic intraper	ative IPC								
4 <u>3</u>	randomised trials	Seriou s ⁷	no serious inconsistenc y	no serious indirectness	no serious imprecision	none	75/1 8 (63.6 %)	(25.6		330 more per 1000 (from 74 more to 785 more)	MODE RATE	CRITICAL
Overall	survival rate	- Hyperth	ermic intraopei	rative IPC								
3	randomised trials	Seriou s ⁹	no serious inconsistenc y	no serious indirectness	Serious⁴	none	71/1 2 (63.4 %)	(45.8		160 more per 1000 (from 5 fewer to 376 more)	LOW	CRITICAL

Diseas	e free survival	rate - No	rmothermic int	raoperative CT	•							
1	randomised trials	Seriou s ³	no serious inconsistenc y	no serious indirectness	Serious ⁴	none	78/13 5 (57.8 %)	74/13 3 (55.6 %)	RR 1.04 (0.84 to 1.28)	22 more per 1000 (from 89 fewer to 156 more)	LOW	CRITICAL

RR=relative risk; 95%CI=95%confidence interval;IPC=intraperitoneal chemotherapy; CT=chemotherapy

⁴ Unclear on attrition rate

² 95%CI crossed two boundries of MID

³ Not intention to treat analysis

⁴ 95%CI crossed one boundary of MID

⁵ one study was not intention to treat analysis and two studies were unclear on attrition rates

⁶ unclear attrition rateone study unclear on attrition rate and one other study was not intention to treat analysis

⁹-<u>All three studies</u>Eujimura 1994, Hamazoe 1994, Yonemura 2001 - unclear randomisation and intention to treat analysis

Table 23: Clinical evidence profile. Intraperitoneal chemotherapy (IPC) versus intravenous chemotherapy (IVC)

Quality	assessm	ent					No of patier		Effect			
No of studie s	Desig n	Risk of bias	Inconsisten cy	Indirectne ss	Impreci sion	Other considerati ons	IPC	IVC	Relative (95% CI)	Absolute	Quali ty	Importa nce
Periope	rative mo	ortality										
1	Rando mised trial	Serious ¹	No serious inconsistenc y	No serious indirectnes s	Very serious ²	None	0/39 (0%)	1/44 (2.3 %)	RR 0.38 (0.02 to 8.95)	-	VER Y LOW	IMPORT ANT
Treatme	ent-relate	d morbidity:	Neutropenia									
1	Rando mised trials	Serious ¹	No serious inconsistenc y	No serious indirectnes s	Very Serious ²	None	8/39 (20. 5%)	11/4 4(25 %)	RR 0.82 (0.37 to 1.83)	-	VER Y LOW	CRITICA L
Overall	survival I	rate										

Quality	assessme	ent					No of patier		Effect			
No of studie s	Desig n	Risk of bias	Inconsisten cy	Indirectne ss	Impreci sion	Other considerati ons	IPC	IVC	Relative (95% CI)	Absolute	Quali ty	Importa nce
<u>54</u>	rando mised trials	Serious ⁴	no serious inconsistenc y	no serious indirectness	Serious 3	none	345 <u>2</u> 61/5 77 442 (59.8 1%)	<u>18</u> / 5 90 457	RR <u>1.21.27</u> (1. <u>02-05</u> to 1.44 <u>54</u>)	101 more per 1000 (from 10 more to 208 more)	LOW	CRITIC AL
Overall	survival r	ate - Normot	hermic intraop	erative IPC								
<u> </u>	rando mised trials	Serious ⁴	serious	no serious indirectness	Serious ³	none	261 <u>1</u> 77/2 9342 8 (61%)	<u>291</u>	RR 1.24 53 (0.95 83 to 2.791.62)	125 more per 1000 (from 26 fewer to 323 more)	VERY LOW	CRITIC AL
Overall	survival r	ate - Hyperth	ermic intraope	erative IPC								
2	rando mised trials	Serious ⁴	no serious inconsistenc y	no serious indirectness	Serious 3	none	84/1 49 (56.4 %)	78/1 66 (47 %)	RR 1.2 (0.96 to 1.48)	94 more per 1000 (from 19 fewer to 226 more)	LOW	CRITIC AL

RR=relative risk; 95%CI=95%confidence interval;IPC=intraperitoneal chemotherapy; CT=chemotherapy ¹ unclear on blinding and selective outcome reporting ² 95%CI crossed two boundries of MID

³ 95%Cl crossed one boundary of MID ⁴ All five-four studies (Kang 2014, Shimoyama 1999, Fujimoto 1999, Ikeguchi 1995) were unclear/inappropriate randomisation method and no/unclear blinding

⁵ I2 > 50%

1

G.13 Squamous cell carcinoma of the oesophagus

What is the most effective curative treatment of squamous cell carcinoma of the oesophagus?

studi es Postoperat	ative mor	Risk of bias rtality Serious ^{1,2,3,4,5,6,7,8}	Inconsis tency	Indirect ness	Imprec ision	Other consider ations	Chemoradiothe rapy followed	Surg ery	Relati	Abso		
8 ran sed	indomi s						by surgery	alon e	ve (95% CI)	lute	Quality	Importanc e
	als		no serious inconsist ency	no serious indirectn ess	serious 9	none	44/524 (8.4%)	23/5 45 (4.2 %)	RR 1.9 (1.18 to 3.07)	38 more per 1000 (from 8 more to 87 more)	LOW	CRITICAL
Postoperat	ative moi	rtality - Concomitan	nt									
sed trial	ed als	Serious ^{1,2,3,4,6,7,8} rtality - Sequential	no serious inconsist ency	no serious indirectn ess	no serious impreci sion	none	33/442 (7.5%)	15/4 65 (3.2 %)	RR 2.25 (1.26 to 4.02)	40 more per 1000 (from 8 more to 97 more)	MODERA TE	CRITICAL

Quality	/ assessm	ent					No of patients		Effect			
No of studi es	Design	Risk of bias	Inconsis tency	Indirect ness	Imprec ision	Other consider ations	Chemoradiothe rapy followed by surgery	Surg ery alon e	Relati ve (95% Cl)	Abso lute	Quality	Importanc e
2	randomi sed trials	serious⁵	no serious inconsist ency	no serious indirectn ess	very serious ¹⁰	none	11/82 (13.4%)	8/80 (10 %)	RR 1.26 (0.54 to 2.97)	26 more per 1000 (from 46 fewer to 197 more)	VERY LOW	CRITICAL
Postop	perative m	ortality - Transhiata	l									
1	randomi sed trials		no serious inconsist ency	no serious indirectn ess	very serious ¹⁰	none	5/50 (10%)	6/50 (12 %)	RR 0.83 (0.27 to 2.55)	20 fewer per 1000 (from 88 fewer to 186 more)	VERY LOW	CRITICAL
Postop	perative m	ortality - 2-stage ap	oroach									
1	randomi sed trials	serious⁵	no serious inconsist ency	no serious indirectn ess	very serious ¹⁰	none	8/47 (17%)	5/38 (13.2 %)	RR 1.29 (0.46 to 3.63)	38 more per 1000 (from 71 fewer	VERY LOW	CRITICAL

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Quality	y assessm	ient					No of patients		Effect			
No of studi es	Design	Risk of bias	Inconsis tency	Indirect ness	Imprec ision	Other consider ations	Chemoradiothe rapy followed by surgery	Surg ery alon e	Relati ve (95% Cl)	Abso lute	Quality	Importanc e
										to 346 more)		
Postop	perative m	ortality - 2 or 3 stag	ge approach									
3	randomi sed trials	Serious ^{6,7,8}	no serious inconsist ency	no serious indirectn ess	no serious impreci sion	none	27/254 (10.6%)	9/27 4 (3.3 %)	RR 3.16 (1.51 to 6.6)	71 more per 1000 (from 17 more to 184 more)	MODERA TE	CRITICAL
Postop	perative m	ortality - Left thora	cotomy									
1	randomi sed trials	serious ¹	no serious inconsist ency	no serious indirectn ess	no serious impreci sion	none	0/118 (0%)	0/11 8 (0%)	not poole d	not poole d	MODERA TE	CRITICAL
Postop	perative m	ortality - Not repor	ted surgical a	pproach								
2	randomi sed trials	serious ^{2,4}	no serious inconsist ency	no serious indirectn ess	very serious ¹⁰	none	4/55 (7.3%)	3/65 (4.6 %)	RR 1.53 (0.39 to 5.9)	24 more per 1000 (from 28 fewer to 226	VERY LOW	CRITICAL

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Quality	/ assessm	ent					No of patients		Effect			
No of studi es	Design	Risk of bias	Inconsis tency	Indirect ness	Imprec ision	Other consider ations	Chemoradiothe rapy followed by surgery	Surg ery alon e	Relati ve (95% CI)	Abso lute	Quality	Importanc e
										more)		
30-day	mortality									,		
3	sed trials	serious ^{1,5,8}	no serious inconsist ency	no serious indirectn ess	serious 9	none	14/246 (5.7%)	6/24 5 (2.4 %)	RR 2.07 (0.85 to 5.03)	26 more per 1000 (from 4 fewer to 99 more)	LOW	CRITICAL
30-day		- Concomitant										
2	sed trials	serious ^{1,8}	no serious inconsist ency	no serious indirectn ess	very serious ¹⁰	none	6/199 (3%)	1/20 7 (0.48 %)	RR 6.59 (0.81 to 53.59)	27 more per 1000 (from 1 fewer to 254 more)	VERY LOW	CRITICAL
30-day		- Sequential										
1	randomi sed trials	serious ⁵	no serious inconsist ency	no serious indirectn ess	very serious	none	8/47 (17%)	5/38 (13.2 %)	RR 1.29 (0.46 to 3.63)	38 more per 1000 (from	VERY LOW	CRITICAL

Quality	y assessm	ent					No of patients		Effect			
No of studi es	Design	Risk of bias	Inconsis tency	Indirect ness	Imprec ision	Other consider ations	Chemoradiothe rapy followed by surgery	Surg ery alon e	Relati ve (95% Cl)	Abso lute	Quality	Importanc e
										71 fewer to 346 more)		
30-day	mortality	- 2-stage approac	h									
1	randomi sed trials	serious ⁵	no serious inconsist ency	no serious indirectn ess	very serious ¹⁰	none	8/47 (17%)	5/38 (13.2 %)	RR 1.29 (0.46 to 3.63)	38 more per 1000 (from 71 fewer to 346 more)	VERY LOW	CRITICAL
30-day	mortality	- 2 or 3 stage app	roach									
1	randomi sed trials	serious ⁸	no serious inconsist ency	no serious indirectn ess	very serious ¹⁰	none	6/81 (7.4%)	1/89 (1.1 %)	RR 6.59 (0.81 to 53.59)	63 more per 1000 (from 2 fewer to 591 more)	VERY LOW	CRITICAL

Quality	/ assessm	ent					No of patients		Effect			
No of studi es	Design	Risk of bias	Inconsis tency	Indirect ness	Imprec ision	Other consider ations	Chemoradiothe rapy followed by surgery	Surg ery alon e	Relati ve (95% Cl)	Abso lute	Quality	Importanc e
1	randomi sed trials	serious ¹	no serious inconsist ency	no serious indirectn ess	no serious impreci sion	none	0/118 (0%)	0/11 8 (0%)	not poole d	not poole d	MODERA TE	CRITICAL
Treatm	nent-relate	d mortality - 2-stage	approach									
1	randomi sed trials	serious ¹¹	no serious inconsist ency	no serious indirectn ess	very serious ¹⁰	none	5/35 (14.3%)	5/34 (14.7 %)	RR 0.97 (0.31 to 3.06)	4 fewer per 1000 (from 101 fewer to 303 more)	VERY LOW	CRITICAL
Treatm	ent-relate	d mortality - 2 or 3-s	tage approa	ch						,		
2		serious ^{6,7}	no serious inconsist ency	no serious indirectn ess	no serious impreci sion	none	20/193 (10.4%)	6/18 5 (3.2 %)	RR 3.21 (1.32 to 7.79)	72 more per 1000 (from 10 more to 220 more)	MODERA TE	CRITICAL

Quality	/ assessm	ent					No of patients		Effect			
No of studi es	Design	Risk of bias	Inconsis tency	Indirect ness	Imprec ision	Other consider ations	Chemoradiothe rapy followed by surgery	Surg ery alon e	Relati ve (95% CI)	Abso lute	Quality	Importanc e
1	randomi sed trials	serious ¹	no serious inconsist ency	no serious indirectn ess	no serious impreci sion	none	0/118 (0%)	0/11 8 (0%)	not poole d	not poole d	MODERA TE	CRITICAL
Treatm	ent-relate	d mortality - Left or	right thorac	otomy								
1	randomi sed trials	serious ¹²	no serious inconsist ency	no serious indirectn ess	very serious	none	3/80 (3.8%)	0/80 (0%)	RR 7 (0.37 to 133.3 6)	-	VERY LOW	CRITICAL
Treatm	ent-relate	d mortality - Not rep	orted surgio	cal approa	ch							
2	randomi sed trials	serious ^{2,4}	no serious inconsist ency	no serious indirectn ess	very serious ¹⁰	none	4/61 (6.6%)	3/65 (4.6 %)	RR 1.37 (0.35 to 5.32)	17 more per 1000 (from 30 fewer to 199 more)	VERY LOW	CRITICAL
Treatm	ent-relate	d mortality										
7	randomi sed trials	serious ^{1,2,4,6,7,11,12}	no serious inconsist ency	no serious indirectn ess	serious 9	none	32/487 (6.6%)	14/4 82 (2.9 %)	RR 2.17 (1.2 to 3.91)	34 more per 1000 (from 6 more to 85	LOW	CRITICAL

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Quality	/ assessm	ent					No of patients		Effect			
No of studi es	Design	Risk of bias	Inconsis tency	Indirect ness	Imprec ision	Other consider ations	Chemoradiothe rapy followed by surgery	Surg ery alon e	Relati ve (95% Cl)	Abso lute	Quality	Importanc e
										more		
Treatm	ent-relate	d mortality (Concon	nitant))		
6	randomi sed trials	Serious ^{1,2,4,6,7,11,12}	no serious inconsist ency	no serious indirectn ess	no serious impreci sion	none	29/448 (6.5%)	11/4 40 (2.5 %)	RR 2.43 (1.27 to 4.63)	36 more per 1000 (from 7 more to 91 more)	MODERA TE	CRITICAL
Treatm	ent-relate	d mortality - Sequer	ntial									
1	sed trials	serious ²	no serious inconsist ency	no serious indirectn ess	very serious ¹⁰	none	3/39 (7.7%)	3/42 (7.1 %)	RR 1.08 (0.23 to 5.02)	6 more per 1000 (from 55 fewer to 287 more)	VERY LOW	CRITICAL
Overal	l survival	rate										
7	randomi sed trials	SeriouS 2,7,8,11,12,13,14	no serious inconsist ency	no serious indirectn ess	serious 9	none	95/389 (24.4%)	68/4 00 (17 %)	RR 1.42 (1.09 to 1.84)	71 more per 1000 (from	LOW	CRITICAL

Quality	y assessm	ient					No of patients		Effect			
No of studi es	Design	Risk of bias	Inconsis tency	Indirect ness	Imprec ision	Other consider ations	Chemoradiothe rapy followed by surgery	Surg ery alon e	Relati ve (95% Cl)	Abso lute	Quality	Importanc e
										15 more to 143 more)		
Overal	I survival	rate (Concomitant)										
6	sed trials	serious ^{7,8,11,12,13,14}	no serious inconsist ency	no serious indirectn ess	serious 9	none	87/350 (24.9%)	61/3 53 (17.3 %)	RR 1.42 (1.08 to 1.87)	73 more per 1000 (from 14 more to 150 more)	LOW	CRITICAL
Overal	I survival	rate (Sequential)										
1	randomi sed trials	serious ²	no serious inconsist ency	no serious indirectn ess	very serious ¹⁰	none	8/39 (20.5%)	7/47 (14.9 %)	RR 1.38 (0.55 to 3.46)	57 more per 1000 (from 67 fewer to 366 more)	VERY LOW	CRITICAL

Quality	y assessm	ent					No of patients		Effect			
No of studi es	Design	Risk of bias	Inconsis tency	Indirect ness	Imprec ision	Other consider ations	Chemoradiothe rapy followed by surgery	Surg ery alon e	Relati ve (95% CI)	Abso lute	Quality	Importanc e
1	randomi sed trials	serious ¹¹	no serious inconsist ency	no serious indirectn ess	very serious ¹⁰	none	8/35 (22.9%)	3/34 (8.8 %)	RR 2.59 (0.75 to 8.95)	140 more per 1000 (from 22 fewer to 701 more)	VERY LOW	CRITICAL
Overall survival rate- 2-stage or transhiatal approach												
1	randomi sed trials	serious ¹⁴	no serious inconsist ency	no serious indirectn ess	very serious ¹⁰	none	8/41 (19.5%)	4/43 (9.3 %)	RR 2.1 (0.68 to 6.44)	102 more per 1000 (from 30 fewer to 506 more)	VERY LOW	CRITICAL
Overal	l survival	rate - 2 or 3 stage ap	proach									
2	randomi sed trials	serious ^{7,8}	no serious inconsist ency	no serious indirectn ess	serious 9	none	43/149 (28.9%)	40/1 46 (27.4 %)	RR 1.05 (0.76 to 1.46)	14 more per 1000 (from 66 fewer	LOW	CRITICAL

Quality	/ assessm	ent					No of patients		Effect			
No of studi es	Design	Risk of bias	Inconsis tency	Indirect ness	Imprec ision	Other consider ations	Chemoradiothe rapy followed by surgery	Surg ery alon e	Relati ve (95% CI)	Abso lute	Quality	Importanc e
										to 126 more)		
Overal 1		rate - Left or right ti serious ¹²	no serious inconsist ency	no serious indirectn ess	serious 9	none	20/80 (25%)	10/8 0 (12.5 %)	RR 2 (1 to 4)	125 more per 1000 (from 0 more to 375 more)	LOW	CRITICAL
Overal	l survival	rate - Not reported	surgical app	roach								
2	randomi sed trials	serious ^{2,13}	no serious inconsist ency	no serious indirectn ess	serious 9	none	16/84 (19%)	11/9 7 (11.3 %)	RR 1.69 (0.83 to 3.45)	78 more per 1000 (from 19 fewer to 278 more)	LOW	CRITICAL
Diseas	e free sur	vival rate (Concomi	itant)									

Quality	y assessm	ent					No of patients		Effect			
No of studi es	Design	Risk of bias	Inconsis tency	Indirect ness	Imprec ision	Other consider ations	Chemoradiothe rapy followed by surgery	Surg ery alon e	Relati ve (95% CI)	Abso lute	Quality	Importanc e
5	randomi sed trials	serious ^{6,7,8,12,13}	serious ¹⁵	no serious indirectn ess	ອ ອ	none	190/386 (49.2%)	103/ 370 (27.8 %)	RR 1.69 (1.18 to 2.4)	192 more per 1000 (from 50 more to 390 more)	VERY LOW	CRITICAL
Disease free survival rate - 2 or 3 stage approach												
3	randomi sed trials	serious ^{6,7,8}	no serious inconsist ency	no serious indirectn ess	serious 9	none	145/261 (55.6%)	82/2 40 (34.2 %)	RR 1.45 (0.87 to 2.41)	154 more per 1000 (from 44 fewer to 482 more)	LOW	CRITICAL
Diseas	e free sur	vival rate - Left or r	ight thoracot	omy								
1	randomi sed trials	serious ¹²	no serious inconsist ency	no serious indirectn ess	serious 9	none	15/80 (18.8%)	5/80 (6.3 %)	RR 3 (1.14 to 7.86)	125 more per 1000 (from 9 more	LOW	CRITICAL

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Quality							No of notion to		Effect			
No of studi es	y assessm Design	Risk of bias	Inconsis tency	Indirect ness	Imprec ision	Other consider ations	No of patients Chemoradiothe rapy followed by surgery	Surg ery alon e	Relati ve (95% CI)	Abso lute	Quality	Importanc e
										to 429 more)		
Diseas	e free sur	vival rate - Not repo	rted surgica	I approach	1							
1	randomi sed trials	no serious risk of bias ¹³	no serious inconsist ency	no serious indirectn ess	no serious impreci sion	none	30/45 (66.7%)	16/5 0 (32 %)	RR 2.08 (1.32 to 3.28)	346 more per 1000 (from 102 more to 730 more)	HIGH	CRITICAL
Any po	ost-operat	ive complication										
5	randomi sed trials	serious ^{2,5,6,7,8}	no serious inconsist ency	no serious indirectn ess	serious 9	none	106/336 (31.5%)	111/ 354 (31.4 %)	RR 1.01 (0.81 to 1.27)	3 more per 1000 (from 60 fewer to 85 more)	LOW	IMPORTA NT
Any po		ive complication - C	oncomitant									
3	randomi sed trials	serious ^{2,6,7,8}	no serious	no serious	serious 9	none	76/254 (29.9%)	80/2 74	RR 1.04	12 more per	LOW	IMPORTA NT

Quality	/ assessm						No of patients		Effect			
No of studi es	Design	Risk of bias	Inconsis tency	Indirect ness	Imprec ision	Other consider ations	Chemoradiothe rapy followed by surgery	Surg ery alon e	Relati ve (95% Cl)	Abso lute	Quality	Importanc e
			inconsist ency	indirectn ess				(29.2 %)	(0.8 to 1.35)	1000 (from 58 fewer to 102 more)		
Any po	ost-operati	ive complication - S	equential									
2	randomi sed trials	serious ⁵	no serious inconsist ency	no serious indirectn ess	very serious ¹⁰	none	30/82 (36.6%)	31/8 0 (38.8 %)	RR 0.96 (0.65 to 1.43)	16 fewer per 1000 (from 136 fewer to 167 more)	VERY LOW	IMPORTA NT
Any po	ost-operati	ive complication - 2-	stage appro	ach								
2	randomi sed trials	serious⁵	no serious inconsist ency	no serious indirectn ess	very serious ¹⁰	none	16/47 (34%)	13/3 8 (34.2 %)	RR 1 (0.55 to 1.8)	0 fewer per 1000 (from 154 fewer to 274 more	VERY LOW	IMPORTA NT

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Quality	/ assessm	ent					No of patients		Effect			
No of studi es	Design	Risk of bias	Inconsis tency	Indirect ness	Imprec ision	Other consider ations	Chemoradiothe rapy followed by surgery	Surg ery alon e	Relati ve (95% Cl)	Abso lute	Quality	Importanc e
Any po	st-operati	ive complication - 2	or 3-stage a	pproach								
3	sed trials	serious ^{6,7,8}	no serious inconsist ency	no serious indirectn ess	serious 9	none	76/254 (29.9%)	80/2 74 (29.2 %)	RR 1.04 (0.8 to 1.35)	12 more per 1000 (from 58 fewer to 102 more)	LOW	IMPORTA NT
Any po	ost-operati	ive complication - No	ot reported	surgical ap	proach							
1	randomi sed trials	serious ²	no serious inconsist ency	no serious indirectn ess	very serious ¹⁰	none	14/35 (40%)	18/4 2 (42.9 %)	RR 0.93 (0.55 to 1.59)	30 fewer per 1000 (from 193 fewer to 253 more)	VERY LOW	IMPORTA NT
Post-o	perative c	omplication: Anasto	motic leak									
7	randomi sed trials	serious ^{1,2,3,4,5,11,12}	no serious inconsist ency	no serious indirectn ess	very serious ¹⁰	none	16/376 (4.3%)	13/3 85 (3.4 %)	RR 1.32 (0.67 to 2.59)	11 more per 1000 (from 11	VERY LOW	IMPORTA NT

Quality	/ assessm	lent					No of patients		Effect			
No of studi es	Design	Risk of bias	Inconsis tency	Indirect ness	Imprec ision	Other consider ations	Chemoradiothe rapy followed by surgery	Surg ery alon e	Relati ve (95% Cl)	Abso lute	Quality	Importanc e
										fewer to 54 more)		
		omplication: Anasto			ant							
5	randomi sed trials	serious ^{1,2,3,4,11,12}	no serious inconsist ency	no serious indirectn ess	very serious ¹⁰	none	9/294 (3.1%)	8/30 5 (2.6 %)	RR 1.23 (0.52 to 2.93)	6 more per 1000 (from 13 fewer to 51 more)	VERY LOW	IMPORTA NT
Post-o	perative c	omplication: Anasto	motic leak -	Sequentia	ıl							
2	sed trials	serious⁵	no serious inconsist ency	no serious indirectn ess	very serious ¹⁰	none	7/82 (8.5%)	5/80 (6.3 %)	RR 1.47 (0.5 to 4.33)	29 more per 1000 (from 31 fewer to 208 more)	VERY LOW	IMPORTA NT
Post-o		omplication: Anasto	motic leak -	Transhiat	al approa	ch						
1	randomi sed trials	serious ³	no serious	no serious	very serious	none	0/50 (0%)	1/50 (2%)	RR 0.33 (0.01	13 fewer per	VERY LOW	IMPORTA NT

Quality	/ assessm	ent					No of patients		Effect			
No of studi es	Design	Risk of bias	Inconsis tency	Indirect ness	Imprec ision	Other consider ations	Chemoradiothe rapy followed by surgery	Surg ery alon e	Relati ve (95% Cl)	Abso lute	Quality	Importanc e
			inconsist ency	indirectn ess					to 7.99)	1000 (from 20 fewer to 140 more)		
Post-o	perative c	omplication: Anasto	motic leak -	2-stage a	oproach							
2	randomi sed trials	serious ^{5,11}	no serious inconsist ency	no serious indirectn ess	very serious ¹⁰	none	3/73 (4.1%)	4/72 (5.6 %)	RR 0.74 (0.17 to 3.26)	14 fewer per 1000 (from 46 fewer to 126 more)	VERY LOW	IMPORTA NT
Post-o	perative c	omplication: Anasto	motic leak -	Left thora	cotomy							
1	randomi sed trials	serious ¹	no serious inconsist ency	no serious indirectn ess	very serious ¹⁰	none	3/118 (2.5%)	1/11 8 (0.85 %)	RR 3 (0.32 to 28.43)	17 more per 1000 (from 6 fewer to 232 more	VERY LOW	IMPORTA NT

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Quality	y assessm	ent					No of patients		Effect			
No of studi es	Design	Risk of bias	Inconsis tency	Indirect ness	Imprec ision	Other consider ations	Chemoradiothe rapy followed by surgery	Surg ery alon e	Relati ve (95% Cl)	Abso lute	Quality	Importance
Post-o	perative c	omplication: Anas	stomotic leak -	Left or rig	ht thorac	otomy						
1	randomi sed trials	serious ¹²	no serious inconsist ency	no serious indirectn ess	very serious	none	1/80 (1.3%)	0/80 (0%)	RR 3 (0.12 to 72.56)	-	VERY LOW	IMPORTA NT
Post-o	perative c	omplication: Anas	stomotic leak -	Not repor	ted surgio	al approach						
2	randomi sed trials	serious ^{2,4}	no serious inconsist ency	no serious indirectn ess	very serious ¹⁰	none	9/55 (16.4%)	7/65 (10.8 %)	RR 1.51 (0.61 to 3.76)	55 more per 1000 (from 42 fewer to 297 more)	VERY LOW	IMPORTA NT
Post-o	perative c	omplication: Infec	tion							,		
2		serious ^{5,8}	no serious inconsist ency	no serious indirectn ess	serious 9	none	34/128 (26.6%)	20/1 30 (15.4 %)	RR 1.57 (1 to 2.45)	88 more per 1000 (from 0 more to 223 more)	LOW	IMPORTA NT

Quality	y assessm	ent					No of patients		Effect			
No of studi es	Design	Risk of bias	Inconsis tency	Indirect ness	Imprec ision	Other consider ations	Chemoradiothe rapy followed by surgery	Surg ery alon e	Relati ve (95% CI)	Abso lute	Quality	Importanc e
1	randomi sed trials	serious ⁸	no serious inconsist ency	no serious indirectn ess	very serious ¹⁰	none	8/81 (9.9%)	5/89 (5.6 %)	RR 1.76 (0.6 to 5.16)	43 more per 1000 (from 22 fewer to 234 more)	VERY LOW	IMPORTA NT
Post-o	perative c	omplication: Infectio	on - Sequent	tial								
1	randomi sed trials		no serious inconsist ency	no serious indirectn ess	serious 9	none	26/47 (55.3%)	15/4 1 (36.6 %)	RR 1.51 (0.94 to 2.44)	187 more per 1000 (from 22 fewer to 527 more)	LOW	IMPORTA NT
		omplication: Infection	on - 2-stage	approach								
1	randomi sed trials	serious ⁵	no serious inconsist ency	no serious indirectn ess	serious 9	none	26/47 (55.3%)	15/4 1 (36.6 %)	RR 1.51 (0.94 to 2.44)	187 more per 1000 (from 22 fewer	LOW	IMPORTA NT

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Quality	/ assessm	ent					No of patients		Effect		
No of studi es	Design	Risk of bias	Inconsis tency	Indirect ness	Imprec ision	Other consider ations	Chemoradiothe rapy followed by surgery	Surg ery alon e	Relati ve (95% Cl)	Abso lute	Quality
										to 527 more)	
Post-o	perative c	omplication: Infectio	n - 2 or 3 st	age appro	ach						
1	randomi sed trials	serious ⁸	no serious inconsist ency	no serious indirectn ess	very serious ¹⁰	none	8/81 (9.9%)	5/89 (5.6 %)	RR 1.76 (0.6 to 5.16)	43 more per 1000 (from 22 fewer to 234 more)	VERY LOW
Post-o	perative c	omplication: stenosi	s (Concomi	tant; Left o	or right the	oracotomy)					
1	randomi sed trials	serious ¹²	no serious inconsist ency	no serious indirectn ess	very serious ¹⁰	none	2/80 (2.5%)	1/80 (1.3 %)	RR 2 (0.19 to 21.62)	13 more per 1000 (from 10 fewer to 258 more)	VERY LOW

Importanc e

IMPORTA NT

IMPORTA NT

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Blood loss in surgery (ml) (Concomitant; Transhiatal) (Better indicated by lower values)

Quality	/ assessm	ent					No of patients		Effect			
No of studi es	Design	Risk of bias	Inconsis tency	Indirect ness	Imprec ision	Other consider ations	Chemoradiothe rapy followed by surgery	Surg ery alon e	Relati ve (95% Cl)	Abso lute	Quality	Importanc e
1	randomi sed trials		no serious inconsist ency	no serious indirectn ess	serious ¹⁶	none	50	50	-	MD 10 highe r (1.92 to 18.08 highe r)	LOW	IMPORTA NT
Intraop	perative tre	eatment-related mor	bidity: Haen	norrhage (>	>300 mL)	(Concomitar	it; Left or right tho	racoto	my)			
1	randomi sed trials	serious ¹²	no serious inconsist ency	no serious indirectn ess	serious 9	none	8/80 (10%)	2/80 (2.5 %)	RR 4 (0.88 to 18.26)	75 more per 1000 (from 3 fewer to 432 more)	LOW	IMPORTA NT
Diseas	e free sur	vival – Concomitant	CRT and 2	or 3 stage	open oese	ophagectom	у					
3	randomi sed trials	serious ^{6,7,8}	no serious inconsist ency	no serious indirectn ess	serious 9	none	DFS* 41% (33% to 48%)	31%	HR 0.77 (0.63 to 0.95)	-	LOW	CRITICAL
Overal	l survival	(2-stage approach)										
1	randomi sed trials	serious ¹¹	no serious	no serious	very serious	none	5-years OS 16% (5% to 33%)	10%	HR 0.8 (0.48	-	VERY LOW	CRITICAL

Quality	, assessm	ent							Effect			
No of studi es	Design	Risk of bias	Inconsis tency	Indirect ness	Imprec ision	Other consider ations	Chemoradiothe rapy followed by surgery	Surg ery alon e	Relati ve (95% CI)	Abso lute	Quality	Importanc e
			inconsist ency	indirectn ess					to 1.34)			
Overal	l survival	(2 or 3-stage approa	ch)									
2	randomi sed trials	serious ^{6,7,8}	no serious inconsist ency	no serious indirectn ess	no serious impreci sion	none	OS* 41%(33% to 48%)	39%	HR 0.96 (0.79 to 1.18)	-	MODERA TE	CRITICAL
Overal	l survival	(2-stage or transhiat	al approach	I)								
1	randomi sed trials	serious ¹⁴	no serious inconsist ency	no serious indirectn ess	serious 9	none	5-years OS 62%(40% to 77%)	34%	HR 0.45 (0.24 to 0.84)	-	LOW	CRITICAL
Overal	l survival	(surgical approach –	unspecifie	d)								
1	randomi sed trials	serious ¹³	no serious inconsist ency	no serious indirectn ess	serious 9	none	5-years OS 29%(19% to 40%)	25%	HR 0.89 (0.67 to 1.19)	-	LOW	CRITICAL

95% CI = 95% Confidence interval; CRT= chemoradiotherapy; DFS = Disease free survival; OS = overall survival; RR=relative risk; HR=Hazard ratio;

¹ Cao 2009 - Unclear randomisation, allocation concealment and blinding

² Le Prise 1994 - Unclear randomisation, allocation concealment and blinding

³ Mashhadi 2015 - Unclear allocation concealment and blinding

⁴ Natsugo 2006 - Unclear randomisation, allocation concealment and blinding

⁵ Nygaard 1992 - Unclear randomisation, allocation concealment and blinding
 ⁶ Bosset 1997 - Unclear randomisation, allocation concealment and blinding

⁷ Lee 2004 - Unclear randomisation, allocation concealment and blinding

⁸ Mariette 2014 - Unclear allocation concealment and blinding

9 95% CI crossed 1 default MID

¹⁰ 95%CI crossed 2 default MIDs

¹¹ Apinop 1994 - Unclear randomisation, allocation concealment and blinding
 ¹² Lv 2010 - Unclear allocation concealment and blinding
 ¹³ Burmeister 2015 - appropriate randomisation and adequate allocation concealment and blinding of research staff and investigators
 ¹⁴ van Hagen 2012 - unclear randomisation, allocation concealment and blinding

¹⁵ I2>50%

¹⁶ Default MID: +/-7.5 ml; 95% CI crossed 1 MID

¹⁷ I2>75%

*OS/DFS was calculated from survival rate at 5 years or, if it was less than 5 years, the survival rate from the last year available.

Table 25: Clinical evidence profile. Chemoradiotherapy followed by surgery versus chemoradiotherapy alone

Quality	y assessmen	t					No of patients Effect					
No of studi es	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecis ion	Other considerati ons	Chemoradiothe rapy followed by surgery	CRT alon e	Relati ve (95% CI)	Abso lute	Quality	Importanc e
Overal	I mortality es	stimates (2-stage appro	bach)								
1	randomise d trials	seriou s ¹	no serious inconsisten cy	no serious indirectne ss	no serious imprecisi on	none	69/86 (80.2%)	75/8 6 (87.2 %)	RR 0.92 (0.81 to 1.05)	70 fewer per 1000 (from 166 fewer to 44 more)	MODERA TE	CRITICAL
Treatm	nent related n	nortality	(2-stage appro	oach)								
1	randomise d trials	seriou s ¹	no serious inconsisten cy	no serious indirectne ss	serious ²	none	11/86 (12.8%)	3/86 (3.5 %)	RR 3.67 (1.06 to 12.68)	93 more per 1000 (from 2 more to 407 more)	LOW	CRITICAL

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Quality	/ assessmen	t					No of patients		Effect			
No of studi es	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecis ion	Other considerati ons	Chemoradiothe rapy followed by surgery	CRT alon e	Relati ve (95% CI)	Abso lute	Quality	Importanc e
3-years	s overall surv	vival rate	(surgical app	roach – uns	pecified)							
1	randomise d trials	seriou S ³	no serious inconsisten cy	no serious indirectne ss	very serious ⁴	none	23/129 (17.8%)	25/1 30 (19.2 %)	RR 0.93 (0.56 to 1.55)	13 fewer per 1000 (from 85 fewer to 106 more)	VERY LOW	CRITICAL
Overal	l survival (O	S) – Conc	omitant CRT	and any type	e of surgica	l approach						
2	randomise d trials	Seriou s ^{1,3}	no serious inconsisten cy	no serious indirectne ss	serious ²	none	OS* 18% (12% to 26%)	18%	HR 0.99 (0.79 to 1.24)	-	LOW	CRITICAL
Overal	l survival – 2	stage oe	sophagecton	ıy								
1	randomise d trials	seriou s ¹	no serious inconsisten cy	no serious indirectne ss	serious ²	none	5-years OS 10% (4% to 19%)	13%	HR 1.15 (0.82 to 1.61)	-	LOW	CRITICAL
Overal	l survival – s	urgical a	pproach unsp	ecified								
1	randomise d trials	seriou s ³	no serious inconsisten cy	no serious indirectne ss	serious ²	none	4-years OS 26% (16% to 37%)	22%	HR 0.89 (0.66 to 1.2)	-	LOW	CRITICAL

Quality	/ assessmen	t					No of patients		Effect			
No of studi es	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecis ion	Other considerati ons	Chemoradiothe rapy followed by surgery	CRT alon e	Relati ve (95% CI)	Abso lute	Quality	Importanc e
1	randomise d trials	seriou s ³	no serious inconsisten cy	no serious indirectne ss	serious ⁵	none	25	37	-	MD 0.95 highe r (0.2 lower to 2.1 highe r)	LOW	IMPORTA NT

95% CI = 95% Confidence interval; CRT= chemoradiotherapy; DFS = Disease free survival; OS = overall surviva; RR=relative risk; HR=Hazard ratio ¹ Stahl 2005/2008 - Unclear randomisation and allocation concealment; unblinded

² 95%CI crossed 1 default MID

³ Bonnetain 2006/Bedenne 2007 - Unclear randomisation and blinding

⁴ 95%CI crossed 2 MIDs

⁵ Default MID: +/- 1.29; 95%CI crossed 1 MID

*OS was calculated from survival rate at 5 years or, if it was less than 5 years, the survival rate from the last year available.

Table 26: Clinical evidence profile. Chemoradiotherapy followed by surgery versus chemotherapy followed by surgery

Quality	/ assessme	nt					No of patients		Effect			
No of studi es	Design	Risk of bias	Inconsist ency	Indirect ness	Impreci sion	Other considerat ions	CRT followed by surgery	CT follo wed by Surg ery	Relati ve (95% CI)	Absol ute	Quality	Importanc e
Mortali	ity											
3	randomi sed trials	serious 1,2,3	no serious inconsiste ncy	no serious indirectn ess	very serious ⁴	none	13/255 (5.1%)	8/25 1 (3.2 %)	RR 1.49 (0.65	16 more per 1000	VERY LOW	CRITICAL

Quality	/ assessme	ent					No of patients		Effect			
No of studi es	Design	Risk of bias	Inconsist ency	Indirect ness	Impreci sion	Other considerat ions	CRT followed by surgery	CT follo wed by Surg ery	Relati ve (95% CI)	Absol ute	Quality	Importanc e
									to 3.39)	(from 11 fewer to 76 more)		
Mortal	ity - Conco	mitant										
2	randomi sed trials	serious _{2,3}	no serious inconsiste ncy	no serious indirectn ess	very serious ⁴	none	5/208 (2.4%)	2/21 0 (0.95 %)	RR 2.53 (0.5 to 12.69)	15 more per 1000 (from 5 fewer to 111 more)	VERY LOW	CRITICAL
Mortal	ity - Sequer	ntial										
1	randomi sed trials	serious ¹	no serious inconsiste ncy	no serious indirectn ess	very serious ⁴	none	8/47 (17%)	6/41 (14.6 %)	RR 1.16 (0.44 to 3.07)	23 more per 1000 (from 82 fewer to 303 more)	VERY LOW	CRITICAL
	ity - 2-stage											
2	randomi sed trials	serious 1,2	no serious inconsiste ncy	no serious	very serious ⁴	none	8/165 (4.8%)	6/16 0	RR 1.16 (0.44	6 more per	VERY LOW	CRITICAL

Quality	/ assessme	nt					No of patients		Effect			
No of studi es	Design	Risk of bias	Inconsist ency	Indirect ness	Impreci sion	Other considerat ions	CRT followed by surgery	CT follo wed by Surg ery	Relati ve (95% CI)	Absol ute	Quality	Importanc e
				indirectn ess				(3.8 %)	to 3.07)	1000 (from 21 fewer to 78 more)		
Mortalit	y - 2 or 3- st	tage approa	ich									
1	randomi sed trials	serious ³	no serious inconsiste ncy	no serious indirectn ess	very serious ⁴	none	5/90 (5.6%)	2/91 (2.2 %)	RR 2.53 (0.5 to 12.69)	34 more per 1000 (from 11 fewer to 257 more)	VERY LOW	CRITICAL
Any po	stoperativ	e mortality										
2	randomi sed trials	serious 1,2	no serious inconsiste ncy	no serious indirectn ess	very serious ⁴	none	8/165 (4.8%)	6/16 0 (3.8 %)	RR 1.16 (0.44 to 3.07)	6 more per 1000 (from 21 fewer to 78 more)	VERY LOW	CRITICAL

Quality	assessme	nt					No of patients		Effect			
No of studi es	Design	Risk of bias	Inconsist ency	Indirect ness	Impreci sion	Other considerat ions	CRT followed by surgery	CT follo wed by Surg ery	Relati ve (95% CI)	Absol ute	Quality	Importanc e
1	randomi sed trials	serious ²	no serious inconsiste ncy	no serious indirectn ess	no serious imprecis ion	none	0/118 (0%)	0/11 9 (0%)	No event in either arm	-	MODERAT E	CRITICAL
Any po	stoperative	e mortality	- Sequential									
1	randomi sed trials	serious ¹	no serious inconsiste ncy	no serious indirectn ess	very serious ⁴	none	8/47 (17%)	6/41 (14.6 %)	RR 1.16 (0.44 to 3.07)	23 more per 1000 (from 82 fewer to 303 more)	VERY LOW	CRITICAL
Any po	stoperative	e mortality	(2-stage app	roach)								
2	randomi sed trials	serious 1,2	no serious inconsiste ncy	no serious indirectn ess	very serious ⁴	none	8/165 (4.8%)	6/16 0 (3.8 %)	RR 1.16 (0.44 to 3.07)	6 more per 1000 (from 21 fewer to 78 more)	VERY LOW	CRITICAL
3-years	s overall su	rvival rate	(Concomitan	t)								
2	randomi sed trials	serious _{2,3}	no serious inconsiste ncy	no serious	serious⁵	none	101/143 (70.6%)	81/1 44	RR 1.26	146 more per	LOW	CRITICAL

Quality	/ assessme	nt					No of patients		Effect			
No of studi es	Design	Risk of bias	Inconsist ency	Indirect ness	Impreci sion	Other considerat ions	CRT followed by surgery	CT follo wed by Surg ery	Relati ve (95% Cl)	Absol ute	Quality	Importanc e
				indirectn ess				(56.3 %)	(1.05 to 1.5)	1000 (from 28 more to 281 more)		
3-years	s overall su	rvival rate	- 2-stage app	roach								
1	randomi sed trials	serious ²	no serious inconsiste ncy	no serious indirectn ess	serious ⁵	none	87/118 (73.7%)	68/1 19 (57.1 %)	RR 1.29 (1.07 to 1.56)	166 more per 1000 (from 40 more to 320 more)	LOW	CRITICAL
3-years	s overall su	rvival rate	- 2 or 3-stage	approach								
1	randomi sed trials	serious ³	no serious inconsiste ncy	no serious indirectn ess	very serious ⁴	none	14/25 (56%)	13/2 5 (52%)	RR 1.08 (0.65 to 1.8)	42 more per 1000 (from 182 fewer to 416 more)	VERY LOW	CRITICAL

Quality	, assessme	ent					No of patients		Effect			
No of studi es	Design	Risk of bias	Inconsist ency	Indirect ness	Impreci sion	Other considerat ions	CRT followed by surgery	CT follo wed by Surg ery	Relati ve (95% CI)	Absol ute	Quality	Importanc e
1	randomi sed trials	serious ³	no serious inconsiste ncy	no serious indirectn ess	very serious ⁴	none	5-years OS 69% (38% to 87%)	49%	HR 0.52 (0.2 to 1.36)		VERY LOW	CRITICAL
Progre	ssion-free	survival rat	e (Concomit	ant; 2 or 3 s	tage appro	oach)						
1	randomi sed trials	serious ³	no serious inconsiste ncy	no serious indirectn ess	very serious ⁴	none	14/25 (56%)	13/2 5 (52%)	RR 1.08 (0.65 to 1.8)	42 more per 1000 (from 182 fewer to 416 more)	VERY LOW	CRITICAL
Treatm	ent-related	l morbidity:	Any complie	cation (Seq	uential; 2-s	tage approac	h)					
1	randomi sed trials	serious ¹	no serious inconsiste ncy	no serious indirectn ess	very serious ⁴	none	16/47 (34%)	14/4 1 (34.1 %)	RR 1 (0.56 to 1.78)	0 fewer per 1000 (from 150 fewer to 266 more)	VERY LOW	IMPORTAN T
Post-o	perative co	mplication:	Anastomoti	c leak								
2	randomi sed trials	serious	serious ⁶	no serious indirectn ess	very serious ⁴	none	5/165 (3%)	3/16 0 (1.9 %)	RR 1.53 (0.13	10 more per 1000	VERY LOW	IMPORTAN T

Quality	assessme	ent					No of patients		Effect			
No of studi es	Design	Risk of bias	Inconsist ency	Indirect ness	Impreci sion	Other considerat ions	CRT followed by surgery	CT follo wed by Surg ery	Relati ve (95% CI)	Absol ute	Quality	Importanc e
									to 17.89)	(from 16 fewer to 317 more)		
Post-o	perative co	mplication:	Anastomoti	c leak - Cor	ncomitant							
1	randomi sed trials	serious ²	no serious inconsiste ncy	no serious indirectn ess	very serious ⁴	none	3/118 (2.5%)	0/11 9 (0%)	RR 7.06 (0.37 to 135.18)	-	VERY LOW	IMPORTAN T
Post-o	perative co	mplication:	Anastomoti	c leak - Sec	quential							
1	randomi sed trials	serious ¹	no serious inconsiste ncy	no serious indirectn ess	very serious ⁴	none	2/47 (4.3%)	3/41 (7.3 %)	RR 0.58 (0.1 to 3.31)	31 fewer per 1000 (from 66 fewer to 169 more)	VERY LOW	IMPORTAN T
		1	Anastomoti	c leak (2-st	age approa	ich)						
2	randomi sed trials	serious _{1,2}	serious ⁶	no serious indirectn ess	very serious ⁴	none	5/165 (3%)	3/16 0 (1.9 %)	RR 1.53 (0.13 to 17.89)	10 more per 1000 (from 16	VERY LOW	IMPORTAN T

Quality	/ assessme	nt	Quality assessment									
No of studi es	Design	Risk of bias	Inconsist ency	Indirect ness	Impreci sion	Other considerat ions	CRT followed by surgery	CT follo wed by Surg ery	Relati ve (95% CI)	Absol ute	Quality	Importanc e
										fewer to 317 more)		
Post-o	perative co	mplication:	stenosis (Co	oncomitant	; 2-stage a	pproach)						
1	randomi sed trials	serious ²	no serious inconsiste ncy	no serious indirectn ess	very serious ⁴	none	2/118 (1.7%)	0/11 9 (0%)	RR 5.04 (0.24 to 103.91	-	VERY LOW	IMPORTAN T

95% CI = 95% Confidence interval; CRT= chemoradiotherapy; OS = overall survival; RR=relative risk;HR=Hazard ratio

¹ Nygaard 1992 - Unclear randomisation, allocation concealment and blinding
 ² Cao 2009 - Unclear randomisation, allocation concealment and blinding
 ³ Klevebro 2015 - Unclear randomisation and allocation concealment and blinding

⁴ 95% CI crossed 2 default MID

⁵ 95% CI crossed 1 default MID

⁶ 12>50%

Table 27: Clinical evidence profile. Surgery followed by chemoradiotherapy versus surgery alone

Quality assessment		No of patients Effect								
No of Design Ris studi of es bia	ncy	Indirectn ess	Impreci sion	Other considerati ons	Surgery followed by Chemoradiother apy	Surg ery	Relati ve (95% Cl)	Absol ute	Quality	Importance

Quality	/ assessme	nt					No of patients		Effect			
No of studi es	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Impreci sion	Other considerati ons	Surgery followed by Chemoradiother apy	Surg ery	Relati ve (95% Cl)	Absol ute	Quality	Importance
1	randomis ed trials	seriou s ¹	no serious inconsiste ncy	no serious indirectn ess	serious ²	none	19/78 (24.4%)	10/80 (12.5 %)	RR 1.95 (0.97 to 3.92)	119 more per 1000 (from 4 fewer to 365 more)	LOW	CRITICAL
10-yea	r progressio	on free s	urvival rate									
1	randomis ed trials	seriou s ¹	no serious inconsiste ncy	no serious indirectn ess	serious ²	none	14/78 (17.9%)	5/80 (6.3 %)	RR 2.87 (1.09 to 7.59)	117 more per 1000 (from 6 more to 412 more)	LOW	CRITICAL

95%CI = 95% confidence interval; CRT = chemoradiotheray; RR=relative risk; ¹ Lv 2010 - Unclear allocation concealment and blinding ² 95% CI crossed 1 default MID

³ 95% CI crossed 2 default MIDs

Table 28: Clinical evidence profile. Chemoradiotherapy alone versus surgery alone

			-									
Quality	assessmen	t					No of patients		Effect			
No of studie s	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecisi on	Other considerati ons	CRT alone	Surg ery alone	Relativ e (95% CI)	Absol ute	Qualit y	Importanc e
Overall	mortality es	stimates										
1	randomis ed trials	seriou s ¹	no serious inconsisten cy	no serious indirectne ss	very serious ²	none	15/36 (41.7%)	20/44 (45.5 %)	RR 0.92 (0.55 to 1.52)	36 fewer per 1000 (from 205 fewer to 236 more)	VERY LOW	CRITICAL
30-day	mortality											
1	randomis ed trials	seriou s ¹	no serious inconsisten cy	no serious indirectne ss	very serious ²	none	0/36 (0%)	3/44 (6.8%)	RR 0.17 (0.01 to 3.26)	57 fewer per 1000 (from 68 fewer to 154 more)	VERY LOW	CRITICAL
Overall	survival rat	e at 2-yea	ars									
1	randomis ed trials	seriou s ¹	no serious inconsisten cy	no serious indirectne ss	very serious ²	none	21/36 (58.3%)	24/44 (54.5 %)	RR 1.07 (0.73 to 1.57)	38 more per 1000 (from 147 fewer to 311 more)	VERY LOW	CRITICAL

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Quality	assessmen	t					No of patients		Effect			
No of studie s	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecisi on	Other considerati ons	CRT alone	Surg ery alone	Relativ e (95% Cl)	Absol ute	Qualit y	Importanc e
Overall	survival rat	e at 5-ye	ars									
1	randomis ed trials	seriou s ¹	no serious inconsisten cy	no serious indirectne ss	serious ³	none	17/36 (47.2%)	10/44 (22.7 %)	RR 2.08 (1.09 to 3.96)	245 more per 1000 (from 20 more to 673 more)	LOW	CRITICAL
Overall	survival (O	S) – Con	comitant CRT	and 2 or 3 st	tage sugery				-	-		
1	randomis ed trials	seriou s ¹	no serious inconsisten cy	no serious indirectne ss	very serious ²	none	5-years OS 50% (26% to 70%)	47%	HR 0.92 (0.47 to 1.79)	-	VERY LOW	CRITICAL
Disease	e-free surviv	val rate a	t 2-years									
1	randomis ed trials	seriou s ¹	no serious inconsisten cy	no serious indirectne ss	very serious ²	none	20/36 (55.6%)	24/44 (54.5 %)	RR 1.02 (0.68 to 1.52)	11 more per 1000 (from 175 fewer to 284 more)	VERY LOW	CRITICAL
5-years	disease-fre	e surviva	al rate									
1	randomis ed trials	seriou s ¹	no serious inconsisten cy	no serious indirectne ss	serious ³	none	17/36 (47.2%)	12/44 (27.3 %)	RR 1.73 (0.96 to 3.13)	199 more per 1000	LOW	CRITICAL

Quality	assessmen	ıt			No of patients		Effect					
No of studie s	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecisi on	Other considerati ons	CRT alone	Surg ery alone	Relativ e (95% Cl)	Absol ute	Qualit y	Importanc e
										(from 11 fewer to 581 more)		

95%CI = 95% confidence interval; CRT = chemoradiotherapy; OS = Overall survival; RR=relative risk; HR=Hazard ratio ¹ Chiu 2005/Teoh 2012 - Unclear randomisation, allocation concealment and blinding ² 95% CI crossed 2 default MIDs ³ 95% CI crossed 1 default MID

Table 29: Clinical evidence profile. Surgery alone versus radiotherapy alone

Quality	assessmen	t					No of patient	s	Effect			
No of studie s	Design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other considerati ons	Surgery alone	RT alone	Relativ e (95% Cl)	Absol ute	Qualit y	Importanc e
Treatm	ent-related r	nortality										
2	randomis ed trials	Serious 1,2	serious ³	no serious indirectne ss	very serious ⁴	none	6/83 (7.2%)	7/80 (8.8%)	RR 1.23 (0.08 to 20.09)	20 more per 1000 (from 80 fewer to 1000 more)	VERY LOW	CRITICAL

Quality No of studie s	assessmen Design	t Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other considerati ons	No of patient Surgery alone	s RT alone	Effect Relativ e (95% Cl)	Absol ute	Qualit y	Importanc e
1	randomis ed trials	serious ¹	no serious inconsistenc y	no serious indirectne ss	very serious ³	none	3/44 (6.8%)	0/43 (0%)	RR 6.84 (0.36 to 128.68)	-	VERY LOW	CRITICAL
Treatm	ent-related n	nortality - 3	-stage approa	ch								
1	randomis ed trials	serious ²	no serious inconsistenc y	no serious indirectne ss	very serious ⁴	none	3/39 (7.7%)	7/37 (18.9 %)	RR 0.41 (0.11 to 1.46)	112 fewer per 1000 (from 168 fewer to 87 more)	VERY LOW	CRITICAL
Overall	survival rate	e - 2-stage	approach									
1	randomis ed trials	serious ¹	no serious inconsistenc y	no serious indirectne ss	serious ⁵	none	24/44 (54.5%)	14/43 (32.6 %)	RR 1.68 (1.01 to 2.78)	221 more per 1000 (from 3 more to 580 more)	LOW	CRITICAL
Overall	survival rate	e										
2	randomis ed trials	serious 1,2	no serious inconsistenc y	no serious indirectne ss	serious⁵	none	30/83 (36.1%)	17/78 (21.8 %)	RR 1.7 (1.05 to 2.74)	153 more per 1000 (from 11 more	LOW	CRITICAL

Quality	assessmen	t					No of patients	5	Effect			
No of studie s	Design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other considerati ons	Surgery alone	RT alone	Relativ e (95% Cl)	Absol ute	Qualit y	Importanc e
										to 379 more)		
Overall	survival rat	e - 3-stage	approach									
1	randomis ed trials	serious ²	no serious inconsistenc y	no serious indirectne ss	very serious ⁴	none	6/39 (15.4%)	3/35 (8.6%)	RR 1.79 (0.48 to 6.64)	68 more per 1000 (from 45 fewer to 483 more)	VERY LOW	CRITICAL
Overall	survival (OS	S) – 3 stage	approach									
1	randomis ed trials	serious ²	no serious inconsistenc y	no serious indirectne ss	no serious imprecisio n	none	5-years OS 31% (15% to 49%)	7%	HR 0.44 (0.27 to 0.72)	-	MOD ERTA TE	CRITICAL

95%CI = 95% confidence interval; CRT = chemoradiotherapy; OS = Overall survival;RR=relative risk; HR=Hazard ratio

¹ Badwe 1998 - Unclear randomisation and blinding
 ² Fok 1994 - Unclear randomisation, allocation concealment and blinding

³ 12>50%

⁴ 95% CI crossed 2 default MIDs
 ⁵ 95% CI crossed 1 default MID

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Quality assessment No of patients Effect Surg No of Design **Risk of bias** Inconsist Indirect Impreci Other CT followed Relati Absol considerat studi ency ness sion by surgery ery ve ute (95% es ions alon Importanc ĊI) Quality е е **30-day mortality** serious 1,2,3,4 4 randomi 10/303 12/3 RR 6 VERY CRITICAL no serious no very none serious⁵ (3.3%) 11 0.84 fewer LOW sed trials inconsiste serious (3.9 (0.38 indirectn ncy per ess %) to 1000 1.86) (from 24 fewer to 33 more) 30-day mortality - 2-stage approach 1 randomi serious¹ 6/41 5/38 RR 14 VERY CRITICAL no serious no very none sed trials inconsiste serious serious⁵ (14.6%) (13.2 1.11 more LOW indirectn %) (0.37 ncy per 1000 ess to 3.35) (from 83 fewer to 309 more) 30-day mortality - 2 stage or transhiatal approach 2 randomi serious 2,4 no serious no 4/143 7/15 RR 19 VERY CRITICAL very none (2.8%) 5 0.57 fewer LOW sed trials inconsiste serious serious⁵ indirectn (4.5 (0.05 ncy per %) 1000 ess to 6.57) (from 43 fewer to 252 more)

Table 30: Clinical evidence profile. Chemotherapy followed by surgery versus surgery alone

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Quality	/ assessme	nt					No of patients	5	Effect			
No of studi es	Design	Risk of bias	Inconsist ency	Indirect ness	Impreci sion	Other considerat ions	CT followed by surgery	Surg ery alon e	Relati ve (95% Cl)	Absol ute	Quality	Importanc e
30-day	mortality -	Left thoracoton	ny									
1	randomi sed trials	serious ³	no serious inconsiste ncy	no serious indirectn ess	no serious imprecis ion	none	0/119 (0%)	0/11 8 (0%)	not pooled	not poole d	MODERAT E	CRITICAL
Treatm	ent-related	mortality										
6	randomi sed trials	Serious 2,3,4,6,7,8	no serious inconsiste ncy	no serious indirectn ess	very serious ⁵	none	17/365 (4.7%)	11/3 63 (3%)	RR 1.48 (0.73 to 3.03)	15 more per 1000 (from 8 fewer to 62 more)	VERY LOW	CRITICAL
Treatm	ent-related	mortality - 3 sta	age approach	1 I								
2	randomi sed trials	serious ^{6,7}	no serious inconsiste ncy	no serious indirectn ess	very serious⁵	none	3/68 (4.4%)	2/68 (2.9 %)	RR 1.4 (0.29 to 6.87)	12 more per 1000 (from 21 fewer to 173 more)	VERY LOW	CRITICAL
Treatm	ent-related	l mortality - 2 or	3 stage appr	oach								
1	randomi sed trials	serious ⁸	no serious inconsiste ncy	no serious indirectn ess	very serious ⁵	none	4/24 (16.7%)	0/22 (0%)	RR 8.28 (0.47	-	VERY LOW	CRITICAL

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Quality	/ assessme	nt					No of patients	5	Effect			
No of studi es	Design	Risk of bias	Inconsist ency	Indirect ness	Impreci sion	Other considerat ions	CT followed by surgery	Surg ery alon e	Relati ve (95% Cl)	Absol ute	Quality	Importanc e
									to 145.5)			
Treatm	ent-related	mortality - 2-st	age or transh	iatal appro	ach							
2	randomi sed trials	serious ^{2,4}	no serious inconsiste ncy	no serious indirectn ess	very serious⁵	none	10/154 (6.5%)	9/15 5 (5.8 %)	RR 1.11 (0.47 to 2.66)	6 more per 1000 (from 31 fewer to 96 more)	VERY LOW	CRITICAL
Treatm	ent-related	mortality - Left	thoracotomy	1								
1	randomi sed trials	serious ³	no serious inconsiste ncy	no serious indirectn ess	no serious imprecis ion	none	0/119 (0%)	0/11 8 (0%)	not pooled	not poole d	MODERAT E	CRITICAL
Postop	erative mo	rtality										
6	randomi sed trials	serious 1,2,3,4,6,7	no serious inconsiste ncy	no serious indirectn ess	very serious⁵	none	17/364 (4.7%)	16/3 79 (4.2 %)	RR 1.1 (0.57 to 2.09)	4 more per 1000 (from 18 fewer to 46 more)	VERY LOW	CRITICAL

Quality	assessme	nt					No of patients	5	Effect			
No of studi es	Design	Risk of bias	Inconsist ency	Indirect ness	Impreci sion	Other considerat ions	CT followed by surgery	Surg ery alon e	Relati ve (95% Cl)	Absol ute	Quality	Importance
1	randomi sed trials	serious ¹	no serious inconsiste ncy	no serious indirectn ess	very serious⁵	none	6/41 (14.6%)	5/38 (13.2 %)	RR 1.11 (0.37 to 3.35)	14 more per 1000 (from 83 fewer to 309 more)	VERY LOW	CRITICAL
Postop	erative mo	rtality - 3-stage	approach									
2	randomi sed trials	serious ^{6,7}	no serious inconsiste ncy	no serious indirectn ess	very serious⁵	none	2/61 (3.3%)	2/68 (2.9 %)	RR 1.1 (0.19 to 6.36)	3 more per 1000 (from 24 fewer to 158 more)	VERY LOW	CRITICAL
Postop	erative mo	rtality - 2 stage	or transhiata	l approach								
2	randomi sed trials	serious ^{2,4}	no serious inconsiste ncy	no serious indirectn ess	very serious⁵	none	9/143 (6.3%)	9/15 5 (5.8 %)	RR 1.09 (0.44 to 2.65)	5 more per 1000 (from 33 fewer to 96 more)	VERY LOW	CRITICAL

Quality	assessme	ent					No of patients	5	Effect			
No of studi es	Design	Risk of bias	Inconsist ency	Indirect ness	Impreci sion	Other considerat ions	CT followed by surgery	Surg ery alon e	Relati ve (95% Cl)	Absol ute	Quality	Importanc e
1	randomi sed trials	serious ³	no serious inconsiste ncy	no serious indirectn ess	no serious imprecis ion	none	0/119 (0%)	0/11 8 (0%)	not pooled	not poole d	MODERAT E	CRITICAL
Overal	l survival ra	ate										
3	randomi sed trials	serious ^{6,8,9}	no serious inconsiste ncy	no serious indirectn ess	very serious⁵	none	23/194 (11.9%)	16/1 93 (8.3 %)	RR 1.39 (0.78 to 2.49)	32 more per 1000 (from 18 fewer to 124 more)	VERY LOW	CRITICAL
Overal	l survival ra	ate - 3 stage app	oroach									
1	randomi sed trials	serious ⁶	no serious inconsiste ncy	no serious indirectn ess	very serious⁵	none	7/47 (14.9%)	3/47 (6.4 %)	RR 2.33 (0.64 to 8.48)	85 more per 1000 (from 23 fewer to 477 more)	VERY LOW	CRITICAL
Overal	l survival ra	ate - 2 or 3 stage	e approach									
1	randomi sed trials	serious ⁸	no serious inconsiste ncy	no serious indirectn ess	very serious⁵	none	7/24 (29.2%)	8/22 (36.4 %)	RR 0.8 (0.35 to 1.85)	73 fewer per 1000 (from 236	VERY LOW	CRITICAL

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Quality	, assessme	ent					No of patients	S	Effect			
No of studi es	Design	Risk of bias	Inconsist ency	Indirect ness	Impreci sion	Other considerat ions	CT followed by surgery	Surg ery alon e	Relati ve (95% Cl)	Absol ute	Quality	Importanc e
										fewer to 309 more)		
Overal	l survival ra	ate - Unspecified	i									
1	randomi sed trials	serious ⁹	no serious inconsiste ncy	no serious indirectn ess	very serious⁵	none	9/123 (7.3%)	5/12 4 (4%)	RR 1.81 (0.63 to 5.26)	33 more per 1000 (from 15 fewer to 172 more)	VERY LOW	CRITICAL
Overal	l survival (OS) – Any type o	of surgical ap	proach								
2	randomi sed trials	Serious ^{2,9}	no serious inconsiste ncy	no serious indirectn ess	serious ¹	none	5-years OS 22% (15% to 29%)	13%	HR 0.75 (0.6 to 0.93)	-	LOW	CRITICAL
Overal	l survival –	2 stage or trans	hiatal oesop	hagectomy	,							
1	randomi sed trials	serious ²	no serious inconsiste ncy	no serious indirectn ess	serious ¹ º	none	5-years OS 26% (16% to 38%)	15%	HR 0.71 (0.51 to 0.98)	-	LOW	CRITICAL
Overal	l survival –	unreported surg	gical approad	ch								
1	randomi sed trials	serious ⁹	no serious inconsiste ncy	no serious indirectn ess	serious ¹	none	5-years OS 19% (11% to 29%)	12%	HR 0.78 (0.58 to 1.04)	-	LOW	CRITICAL

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Quality	assessme	ent					No of patients	S	Effect			
No of studi es	Design	Risk of bias	Inconsist ency	Indirect ness	Impreci sion	Other considerat ions	CT followed by surgery	Surg ery alon e	Relati ve (95% CI)	Absol ute	Quality	Importanc e
Diseas	e free surv	ival rate (2 stage	e or transhiat	tal)					-			
1	randomi sed trials	serious ²	no serious inconsiste ncy	no serious indirectn ess	serious ¹	none	19/85 (22.4%)	9/84 (10.7 %)	RR 2.09 (1 to 4.34)	117 more per 1000 (from 0 more to 358 more)	LOW	CRITICAL
Diseas	e free survi	ival (DFS) – 2 st	age or transl	niatal								
1	randomi sed trials	serious ²	no serious inconsiste ncy	no serious indirectn ess	serious ¹ 0	none	5-years DFS 23% (13% to 35%)	13%	HR 0.72 (0.52 to 1)	-	LOW	CRITICAL
Anasto	motic leak	age										
6	randomi sed trials	Serious 1.2,3,4,6,7	no serious inconsiste ncy	no serious indirectn ess	very serious ⁵	none	21/364 (5.8%)	19/3 79 (5%)	RR 1.15 (0.65 to 2.02)	8 more per 1000 (from 18 fewer to 51 more)	VERY LOW	IMPORTA NT
Anasto	motic leak	age - 2-stage ap	proach									
1	randomi sed trials	serious ¹	no serious inconsiste ncy	no serious indirectn ess	very serious⁵	none	3/41 (7.3%)	2/38 (5.3 %)	RR 1.39 (0.25	21 more per 1000	VERY LOW	IMPORTA NT

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Quality	assessme	nt					No of patients	5	Effect			
No of studi es	Design	Risk of bias	Inconsist ency	Indirect ness	Impreci sion	Other considerat ions	CT followed by surgery	Surg ery alon e	Relati ve (95% Cl)	Absol ute	Quality	Importanc e
									to 7.87)	(from 39 fewer to 362 more)		
Anasto	motic leak	age - 3-stage ap	proach									
2	randomi sed trials	serious ^{6,7}	no serious inconsiste ncy	no serious indirectn ess	very serious ⁵	none	7/61 (11.5%)	7/68 (10.3 %)	RR 1.03 (0.41 to 2.61)	3 more per 1000 (from 61 fewer to 166 more)	VERY LOW	IMPORTA NT
Anasto	motic leak	age - 2-stage or	transhiatal a	ppraoch								
2	randomi sed trials	serious ^{2,4}	no serious inconsiste ncy	no serious indirectn ess	very serious⁵	none	11/143 (7.7%)	9/15 5 (5.8 %)	RR 1.31 (0.58 to 2.97)	18 more per 1000 (from 24 fewer to 114 more)	VERY LOW	IMPORTA NT
Anasto	motic leak	age - Left thorac	cic									
1	randomi sed trials	serious ³	no serious inconsiste ncy	no serious indirectn ess	very serious⁵	none	0/119 (0%)	1/11 8 (0.85 %)	RR 0.33 (0.01 to 8.03)	6 fewer per 1000 (from	VERY LOW	IMPORTA NT

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Quality	assessme	ent					No of patients	s	Effect			
No of studi es	Design	Risk of bias	Inconsist ency	Indirect ness	Impreci sion	Other considerat ions	CT followed by surgery	Surg ery alon e	Relati ve (95% Cl)	Absol ute	Quality	Importanc e
										8 fewer to 60 more)		
Treatm	ent-related	I morbidity: bloo	od loss (2-sta	ge or trans	hiatal appr	oach) (Better	indicated by lo	ower va	lues)			
1	randomi sed trials	serious⁴	no serious inconsiste ncy	no serious indirectn ess	no serious imprecis ion	none	60	69	-	MD 62 higher (45.7 1 to 78.29 higher)	MODERAT E	IMPORTA NT
Treatm	ent-related	l morbidity: wou	and infection	(2-stage or	transhiata	l approach)						
1	randomi sed trials	serious ⁴	no serious inconsiste ncy	no serious indirectn ess	very serious⁵	none	4/60 (6.7%)	7/69 (10.1 %)	RR 0.66 (0.2 to 2.14)	34 fewer per 1000 (from 81 fewer to 116 more)	VERY LOW	IMPORTA NT
Post-o	perative tre	eatment related	morbidity: Ar	nastomotic	leakage (2	stage or tran	shiatal)					
1	randomi sed trials	serious ²	no serious inconsiste ncy	no serious indirectn ess	very serious⁵	none	8/85 (9.4%)	9/84 (10.7 %)	RR 0.88 (0.36 to 2.17)	13 fewer per 1000 (from 69 fewer	VERY LOW	IMPORTA NT

Quality	Quality assessment								Effect			
No of studi es	Design	Risk of bias	Inconsist ency	Indirect ness	Impreci sion	Other considerat ions	CT followed by surgery	Surg ery alon e	Relati ve (95% Cl)	Absol ute	Quality	Importanc e
										to 125 more)		

95%CI = 95% confidence interval; CRT = chemoradiotherapy; DFS = Disease free survival; OS = Overall surviva; RR=relative risk; HR=Hazard ratio

¹ Nygaard 1992 - Unclear randomisation, allocation concealment and blinding

² Boonstra 2011 - Unclear allocation concealment and blinding

³ Cao 2009 - Unclear randomisation, allocation concealment and blinding

⁴ Law 1997 - Unclear randomisation, allocation concealment and blinding

⁵ 95%CI crossed 2 default MIDs

⁶ Ancona 2001 - Unclear allocation concealment and blinding

⁷ Baba 2000 - Unclear randomisation, allocation concealment and blinding

⁸ Maipang 1994 - Unclear randomisation, allocation concealment and blinding

⁹ MRC 2002 - Unclear randomisation and blinding
 ¹⁰ 95% CI crossed 1 default MID

¹¹ Schlag 1992 - Unclear randomisation, allocation concealment and blinding

Table 31: Clinical evidence profile. Chemoradiotherpy versus radiotherapy alone

Quality	/ assessm	ent					No of patients		Effect			
No of studi es	Design	Risk of bias	Inconsist ency	Indirect ness	Impreci sion	Other considera tions	CRT alone	RT alon e	Relati ve (95% Cl)	Abso lute	Quality	Importanc e
Treatm	nent related	d mortality (concomitant))									
8	randomi sed trials	serious ^{1,2,3,4,5,6,7,8}	no serious inconsiste ncy	no serious indirectn ess	very serious ⁹	none	8/322 (2.5%)	7/33 0 (2.1 %)	RR 1.17 (0.47 to 2.9)	4 more per 1000 (from 11 fewer	VERY LOW	CRITICAL

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Quality	/ assessm	ent					No of pa	atients	Effect			
No of studi es	Design	Risk of bias	Inconsist ency	Indirect ness	Impreci sion	Other considera tions	CRT alone	RT alon e	Relati ve (95% Cl)	Abso lute	Quality	Importanc e
										to 40 more)		
Overal	l survival r	ate (sequential)								more)		
2	randomi sed trials	serious ^{11,12}	serious ¹⁰	no serious indirectn ess	very serious ⁹	none	20/70 (28.6%)	26/7 6 (34.2 %)	RR 0.4 (0.02 to 8.14)	205 fewer per 1000 (from 335 fewer to 1000 more)	VERY LOW	CRITICAL
Overal	l survival r	ate at 1 year (Concomita	ant)									
8	randomi sed trials	Serious 1.2.3.7.8,13,14,15	serious ¹⁰	no serious indirectn ess	serious ¹	none	256/43 3 (59.1%)	215/ 436 (49.3 %)	RR 1.21 (0.99 to 1.48)	104 more per 1000 (from 5 fewer to 237 more)	VERY LOW	CRITICAL
		ate at 3 years (Concomi	1									
8	randomi sed trials	Serious ^{1,2,3,7,8,13,14,15}	no serious inconsiste ncy	no serious indirectn ess	no serious imprecis ion	none	117/43 3 (27%)	65/4 36 (14.9 %)	RR 1.82 (1.4 to 2.37)	122 more per 1000 (from 60	MODERA TE	CRITICAL

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Quality	assessmo	ent					No of pa	atients	Effect			
No of studi es	Design	Risk of bias	Inconsist ency	Indirect ness	Impreci sion	Other considera tions	CRT alone	RT alon e	Relati ve (95% Cl)	Abso lute	Quality	Importanc e
										more to 204 more)		
Overal	l survival r	ate at 5 years (Conc	omitant)									
6	randomi sed trials	serious ^{1,2,3,7,8,14}	no serious inconsiste ncy	no serious indirectn ess	no serious imprecis ion	none	58/332 (17.5%)	25/3 30 (7.6 %)	RR 2.33 (1.51 to 3.58)	101 more per 1000 (from 39 more to 195 more)	MODERA TE	CRITICAL
Overal	l survival (OS) - Concomitant										
4	randomi sed trials	Serious ^{1,2,3,6}	no serious inconsiste ncy	no serious indirectn ess	no serious imprecis ion ¹⁷	none	OS* 13% (0% to 19%)	4%	HR 0.63 (0.51 to 0.77)	-	MODERA TE	CRITICAL
Overal	l survival (OS) - Sequential										
1	randomi sed trials	serious ¹¹	no serious inconsiste ncy	no serious indirectn ess	serious ¹ 7	none	5- years OS 3%(1% to 11%)	6%	HR 1.21 (0.77 to 1.9)	-	LOW	CRITICAL

Quality	/ assessmi	ent					No of pa	atients	Effect			
No of studi es	Design	Risk of bias	Inconsist ency	Indirect ness	Impreci sion	Other considera tions	CRT alone	RT alon e	Relati ve (95% CI)	Abso lute	Quality	Importanc e
2	randomi sed trials	serious ^{2,3}	very serious ¹⁸	no serious indirectn ess	very serious ⁹	none	51/97 (52.6%)	67/1 02 (65.7 %)	RR 0.88 (0.48 to 1.63)	79 fewer per 1000 (from 342 fewer to 414 more)	VERY LOW	CRITICAL
Diseas	e free surv	vival (DFS) - concomitant										
2	randomi sed trials	Serious ^{2,3}	serious10	no serious indirectn ess	serious ¹ 7	none	1-year DFS 72%(6 3% to 79%)	55%	HR 0.56 (0.4 to 0.78)	-	VERY LOW	CRITICAL
Treatm	ent related	I morbidity - concomitan	t				,					
6	randomi sed trials	Serious ^{1,2,6,7,13,14}	no serious inconsiste ncy	no serious indirectn ess	serious ¹ 7	none	95/306 (31%)	88/3 06 (28.8 %)	RR 1.09 (0.88 to 1.36)	26 more per 1000 (from 35 fewer to 104 more)	LOW	IMPORTA NT

95%CI = 95% confidence interval; CRT = chemoradiotherapy; DFS = Disease free survival; OS = Overall survival;RR=relative risk; HR=Hazard ratio ¹ Araujo 1991 - Unclear randomisation, allocation concealment, blinding and unclear outcome report ² Cooper 1999- Unclear randomisation, allocation concealment and blinding

³ Gao 2002 - Unclear randomisation, allocation concealment and blinding

⁴ Kaneta 1997 - Unclear randomisation, allocation concealment and blinding ⁵ Slabber 1998 - Unclear randomisation, allocation concealment and blinding

- ⁶ Zhu 2000 Unclear randomisation, allocation concealment and blinding
- ⁷ Zhao 2005 Unclear allocation concealment and blinding
- ⁸ Smith 1998 Unclear blinding
- ⁹ 95%CI crossed 2 default MIDs
- ¹⁰ I2>50%

¹¹ Hatlevoll 1992 - Unclear randomisation, allocation concealment and blinding

- ¹² Hishikawa 1991 Unclear randomisation, allocation concealment and blinding
- ¹³ Han 2012 Unclear randomisation, allocation concealment and blinding
- ¹⁴ Kumar 2007 Unclear randomisation, allocation concealment and blinding
- ¹⁵ Herskovic 1992/AI-Sarraf 1997 Unclear randomisation, allocation concealment and blinding
- ¹⁶ 95%CI crossed 1 default MID

¹⁷ I2=75%

*OS was calculated from survival rate at 5 years or, if it was less than 5 years, the survival rate from the last year available.

G.14 Non-metastatic oesophageal cancer not suitable for surgery

What is the optimal treatment for adults with non-metastatic disease in the oesophagus who are not suitable for surgery?

Qualit	y assessme	nt					No of patients					
No of studi es	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecisi on	Other considerati ons	Radiotherapy	Chemo- radiother apy	Relati ve (95% CI)	Absol ute	Quality	Importanc e
Overa	I Survival a	t 3 years	(assessed with	th: Kaplan-M	leier Overall	Survival)						
<u>33</u>	randomis ed trials	seriou s ¹	no serious inconsisten cy	no serious indirectne ss	no serious imprecisi on	none	14% at three years ¹²	21% at three years (from 15% to 28%)	HR 0.8 (0.65 to 0.97)	-	MODERA TE	CRITICAL

Table 32: Clinical evidence profile. Comparison 1: Radiotherapy versus chemoradiotherapy

	y assessme	nt					No of patients		Effect	1		
No of studi es	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecisi on	Other considerati ons	Radiotherapy	Chemo- radiother apy	Relati ve (95% CI)	Absol ute	Quality	Importanc e
1	randomis ed trials	seriou s ¹	no serious inconsisten cy	no serious indirectne ss	very serious ⁷	none	2/57 (3.5%)	5/54 (9.3%)	RR 0.38 (0.08 to 1.87)	57 fewer per 1000 (from 85 fewer to 81 more)	VERY LOW	IMPORTA NT
One-Y	ear Progres	sion Fre	e Survival rate	(follow-up '	l years)							
2	randomis ed trials	seriou s ⁸	very serious ⁹	no serious indirectne ss	very serious ¹⁰	none	42/146 (28.8%)	48/143 (33.6%)	RR 0.93 (0.3 to 2.89)	23 fewer per 1000 (from 235 fewer to 634 more)	VERY LOW	CRITICAL
Three	Year Progre	ession Fr	ee Survival ra	te (follow-up	o 3 years)							
1	randomis ed trials	seriou s ¹	no serious inconsisten cy	no serious indirectne ss	very serious ⁷	none	8/111 (7.2%)	9/110 (8.2%)	RR 0.87 (0.32 to 2.35)	10 fewer per 1000 (from 54 fewer to 91 more)	VERY LOW	CRITICAL

Qualit	y assessme	nt					No of patients		Effect			
No of studi es	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecisi on	Other considerati ons	Radiotherapy	Chemo- radiother apy	Relati ve (95% Cl)	Absol ute	Quality	Importanc e
2	randomis ed trials	seriou s ²	serious ¹¹	no serious indirectne ss	no serious imprecisi on	none	1/144 (0.69%)	14/145 (9.7%)	RR 0.11 (0.02 to 0.55)	86 fewer per 1000 (from 43 fewer to 95 fewer)	LOW	IMPORTA NT
Treatn	nent-Related	d Toxicity	y - Esophagitis	s (assessed	with: Grade	2-4)						
2	Randomis ed trials	Seriou s ¹	No serious inconsisten cy	No serious indirectne ss	Serious ⁶	none	37/93 (39.8%)	49/100 (49%)	RR 0.81 (0.6 to 1.09)	93 fewer per 1000 (from 196 fewer to 44 more)	LOW	IMPORTA NT

CI=confidence interval; RR=relative risk; HR=hazard ratio; ¹ <u>Wobbes 2001, Kumar 2007, Lui 2012 -</u> Unclear reporting of allocation concealment and randomisation process.

¹ Woodes 2001, Kumar 2007, Lur 2012 - Unclear reporting of allocation concearment and randomisation process.
 ² Due to inadequate reporting of randomisation process and blinding. Gao 2009: very limited details on methodology.
 ³ I-squared statistic >75
 ⁴ Effect estimate cross one MID
 ⁵ Unclear reporting of allocation concealment and randomisation process.
 ⁶ i-squared statistic between 50-75%
 ⁷ Very continue information of Concealment and randomisation process.

1

⁷ Very serious imprecision as 95% CI cross two default MIDs.

⁸ No explanation was provided

⁹ Very serious heterogeneity. I-squared> 75%. Also presented by subgroup (chemotherapy class) due to heterogeneity.
 ¹⁰ Serious impresion. 95% CI crosses one default MID.

¹¹ Downgraded for serious inconsistency. I-squared statistic 50-74.99. ¹²3 year overall survival taken from RT arm of Kumar 2007

			-	-								
Qualit	v assessme	nt					No of patients		Effect			
No of studi es	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecisi on	Other considerati ons	5-FU-based chemo- radiotherapy (CRT)	Non- 5-FU- base d CRT	Relati ve (95% Cl)	Absol ute	Quality	Importanc e
1-Year 1	Overall Sur randomis ed trials	no serio us risk of bias	te no serious inconsisten cy	no serious indirectne ss	very serious ¹	none	9/37 (24.3%)	11/35 (31.4 %)	RR 0.77 (0.37 to 1.64)	72 fewer per 1000 (from 198 fewer to 201 more)	LOW	CRITICAL
2-Year	Overall Sur	vival ra	te									
1	randomis ed trials	no serio us risk of bias	no serious inconsisten cy	no serious indirectne ss	serious ²	none	29/37 (78.4%)	23/35 (65.7 %)	RR 1.19 (0.89 to 1.6)	125 more per 1000 (from 72 fewer to 394 more)	MODERAT E	CRITICAL
Treatn	nent-Related	Mortal	ity (assessed	with: Mortal	ity due to tre	eatment-related	d toxic effects)					
1	randomis ed trials	no serio us risk of bias	no serious inconsisten cy	no serious indirectne ss	very serious ³	none	1/37 (2.7%)	2/35 (5.7%)	RR 0.47 (0.04 to 4.99)	30 fewer per 1000 (from 55 fewer	LOW	IMPORTAN T

Table 33: Clinical evidence profile. Comparison 2: 5-FU-based chemoradiotherapy versus non-5-FU-based chemoradiotherapy

Qualit	y assessme	nt					No of patients		Effect			
No of studi es	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecisi on	Other considerati ons	5-FU-based chemo- radiotherapy (CRT)	Non- 5-FU- base d CRT	Relati ve (95% Cl)	Absol ute	Quality	Importanc e
										to 228 more)		
Treatn	nent-Related	d Morbio	dity: Grade 4/5	5 Toxicity (as	sessed with	n: WHO Toxicit	y Grading)					
1	randomis ed trials	no serio us risk of bias	no serious inconsisten cy	no serious indirectne ss	very serious ³	none	11/37 (29.7%)	15/35 (42.9 %)	RR 0.69 (0.37 to 1.3)	133 fewer per 1000 (from 270 fewer to 129 more)	LOW	IMPORTAN T

CI=confidence interval; RR=relative risk; 5-FU=5-Fluouracil; CRT=chemoradiotherapy

¹ Effect estimate crosses two MIDs

² Effect estimate crosses one MID

³ Very serious imprecision. 95% CI crosses two default MIDs.

G.15 First-line palliative chemotherapy

What is the optimal palliative first-line systemic chemotherapy for locally advanced and/or metastatic oesophago-gastric cancer?

Qualit	y assessme	nt					No of patie	nts	Effect			
No of studi es	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecisi on	Other considerati ons	Combinat ion CT	Single- agent CT	Relati ve (95% CI)	Absol ute	Quality	Importanc e
Overa	ll survival (a	ssessed w	ith: Kaplan M	eier Mortality	/ estimates)							
4	randomis ed trials	serious _{1,2}	no serious inconsisten cy	no serious indirectne ss	no serious imprecisi on	none	-	-	HR 0.77 (0.65 to 0.91)	-	MODERAT E	CRITICAL
Treatn	nent-related	death										
4	randomis ed trials	serious 1,2	no serious inconsisten cy	no serious indirectne ss	very serious ³	none	6/337 (1.8%)	3/223 (1.3%)	RR 1.31 (0.39 to 4.34)	4 more per 1000 (from 8 fewer to 45 more)	VERY LOW	IMPORTA NT
Treatn	nent-related	toxicity: N	ausea and Vo	miting (asse	ssed with: \	WHO Grade 3/4)					
2	randomis ed trials	no serious risk of bias	no serious inconsisten cy	no serious indirectne ss	very serious ³	none	16/175 (9.1%)	11/174 (6.3%)	RR 1.44 (0.69 to 3.02)	28 more per 1000 (from 20 fewer to 128 more)	LOW	CRITICAL

 Table 34: Clinical evidence profile. Single agent chemotherapy versus combination chemotherapy

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Qualit	y assessme	nt					No of patie	nts	Effect			
No of studi es	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecisi on	Other considerati ons	Combinat ion CT	Single- agent CT	Relati ve (95% Cl)	Absol ute	Quality	Importanc e
Treatn	nent-related	toxicity: D	iarrhoea (asse	essed with: V	VHO Grade	3/4)						
2	randomis ed trials	no serious risk of bias	serious inconsisten cy ⁴	no serious indirectne ss	very serious ³	none	5/175 (2.9%)	5/174 (2.9%)	RR 1.28 (0.07 to 21.75)	8 more per 1000 (from 27 fewer to 596 more)	LOW	CRITICAL

CI=confidence interval; RR=relative risk; HR=hazard ratio; CT=chemotherapy;

¹ Colucci- unclear allocation concealment, no intention to treat analysis
 ² Lutz- single-therapy arm was closed earlier (Simon 2-stage minimax design)
 ³ 95% CI crosses 2 default MIDs
 ⁴ 12 > 50%

Table 35: Clinical evidence summary. 5-FU/cisplatin/anthracycline combinations versus 5-FU/cisplatin combinations without anthracyclines

Qualit	ty assessm	ent					No of patien	its	Effect			
No of stud ies	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecis ion	Other considerati ons	5- FU/cisplati n/anthracy cline combinati ons	5-FU/cisplatin combinations (without anthracylines)	Relat ive (95% Cl)	Absol ute	Quality	Importa nce
Overa	Il survival											
3	randomis ed trials	no seriou s risk of bias	no serious inconsisten cy	no serious indirectne ss	serious ¹	none	-	-	HR 0.70 (0.43 to 1.15)	-	MODERA TE	CRITICA L

Quali	ty assessme	ent					No of patien	ts	Effect			
No of stud ies	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecis ion	Other considerati ons	5- FU/cisplati n/anthracy cline combinati ons	5-FU/cisplatin combinations (without anthracylines)	Relat ive (95% Cl)	Absol ute	Quality	Importa nce
Progr	ession-Free	e Surviva	ıl			·	·	·				
1	randomis ed trials	seriou s ²	no serious inconsisten cy	no serious indirectne ss	very serious ³	none	-	-	HR 0.95 (0.58 to 1.57)	-	VERY LOW	CRITICA L

CI=confidence interval; RR=relative risk; HR=hazard ratio; 5-FU=5-fluouracil

² 95% CI crosses 2 default MID boundaries

Table 36: Clinical evidence summary. 5-FU/cisplatin/anthracycline combinations versus 5-FU/anthracycline combinations (without cisplatin

Qualit	ty assessme	ent					No of patients		Effect			
No of stud ies	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecis ion	Other considerati ons	5- FU/cisplatin/a nthracycline combinations	5- FU/anthracy cline combination s (without cisplatin)	Relat ive (95% Cl)	Absol ute	Quality	Importa nce
Overa	II survival											
2	randomis ed trials	seriou s ¹	no serious inconsisten cy	no serious indirectne ss	no serious imprecisi on	none	-	-	HR 0.7 (0.54 to 0.89)	-	MODERA TE	CRITICA L

CI=confidence interval; RR=relative risk; HR=hazard ratio; 5-FU=5-fluouracil ¹ Roth- no ITT analysis, no information on follow-up of participants

Table 37: Clinical evidence summary. Irinotecan containing regimes versus non-irinotecan containing regimes

			• • • • • • • • • • • • • • • • • • •		. J.,	,			J	- J		
Qualit	y assessmer	nt					No of pati	ents	Effect			
No of studi es	Design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecisi on	Other consideratio ns	Irinoteca n containi ng regimes	non- irinot ecan conta ing regim es	Relati ve (95% CI)	Absolu te	Quality	Importance
Overa	ll survival											
4	randomise d trials	serious 1	no serious inconsistenc y	no serious indirectnes s	serious ²	none	-	-	HR 0.87 (0.73 to 1.05)	-	LOW	CRITICAL
Progre	ession-free s	urvival										
3	randomise d trials	serious	no serious inconsistenc y	no serious indirectnes s	serious ²	none	-	-	HR 0.83 (0.68 to 1.01)	-	LOW	CRITICAL
Treatn	nent-related	death										
3	randomise d trials	no serious risk of bias	no serious inconsistenc y	no serious indirectnes s	serious ²	none	1/268 (0.37%)	8/258 (3.1%)	RR 0.21 (0.05 to 0.98)	24 fewer per 1000 (from 1 fewer to 29 fewer)	MODERAT E	IMPORTAN T
Treatm	nent discont	inuation o	due to toxicity									

Qualit	y assessmer	nt					No of patie	ents	Effect			
No of studi es	Design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecisi on	Other consideratio ns	Irinoteca n containi ng regimes	non- irinot ecan conta ing regim es	Relati ve (95% CI)	Absolu te	Quality	Importance
3	randomise d trials	no serious risk of bias	no serious inconsistenc y	no serious indirectnes s	serious ²	none	32/272 (11.8%)	53/26 3 (20.2 %)	RR 0.65 (0.34 to 1.24)	71 fewer per 1000 (from 133 fewer to 48 more)	MODERAT E	CRITICAL

CI=confidence interval; RR=relative risk; HR=hazard ratio; ¹ Park- unclear randomization, allocation concealment and blinding of assessors ² 95% CI crosses one default MID boundary

Table 38: Clinical evidence summary. Docetaxel containing regimes versus non-docetaxel containing regimes

Qualit	y assessme	nt					No of pat	ients	Effect			
No of studi es	Design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other considerati ons	Docetax el containi ng regimes	Non- docet axel- conta ining regim es	Relativ e (95% CI)	Absol ute	Quality	Importanc e
Overa	II survival											

Qualit	v assessme	nt					No of pat	ients	Effect			
No of studi es	Design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other considerati ons	Docetax el containi ng regimes	Non- docet axel- conta ining regim es	Relativ e (95% CI)	Absol ute	Quality	Importanc e
4	randomis ed trials	no serious risk of bias	no serious inconsistenc y	no serious indirectne ss	serious ¹	none	-	-	HR 0.87 (0.76 to 1.01)	-	MODERAT E	CRITICAL
Treatm	nent-related	death										
5	randomis ed trials	serious 2,3	no serious inconsistenc y	no serious indirectne ss	very serious ⁴	none	9/550 (1.6%)	12/51 7 (2.3%)	RR 0.75 (0.34 to 1.65)	6 fewer per 1000 (from 15 fewer to 15 more)	VERY LOW	IMPORTAN T
Time t	o progressi	on										
3	randomis ed trials	serious5	very serious ⁶	no serious indirectne ss	very serious ⁴	none	-	-	HR 0.85 (0.56 to 1.29)	-	VERY LOW	CRITICAL
Treatm	nent discont	tinuation d	ue to toxicity									
5	randomis ed trials	serious 3,5	no serious inconsistenc y	no serious indirectne ss	serious ¹	none	84/478 (17.6%)	95/44 6 (21.3 %)	RR 0.85 (0.65 to 1.1)	32 fewer per 1000 (from 75 fewer	LOW	CRITICAL

Qualit	y assessme	nt					No of pat	ients	Effect			
No of studi es	Design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other considerati ons	Docetax el containi ng regimes	Non- docet axel- conta ining regim es	Relativ e (95% CI)	Absol ute	Quality	Importanc e
										to 21 more)		
Treatr	nent-related	toxicity: di	arrhoea									
1	randomis ed trials	serious⁵	no serious inconsistenc y	no serious indirectne ss	serious ^{1,7}	none	15/121 (12.4%)	0/122 (0%)	RR 31.25 (1.89 to 516.54)	-	LOW	CRITICAL
Treatr	nent-related	toxicity: N	ausea and von	niting								
1	randomis ed trials	serious ⁵	no serious inconsistenc y	no serious indirectne ss	very serious ⁴	none	9/121 (7.4%)	14/12 2 (11.5 %)	RR 0.65 (0.29 to 1.44)	40 fewer per 1000 (from 81 fewer to 50 more)	VERY LOW	CRITICAL
Qualit	y of Life: Ph	ysical Fund	ctioning (Bette	r indicated b	y lower valu	ues)						
1	randomis ed trials	serious ⁸	no serious inconsistenc y	no serious indirectne ss	serious ¹	none	44	41	-	MD 1.8 lower (7.84 lower to 4.24 higher)	LOW	IMPORTAN T

Qualit	y assessme	nt					No of pat	ients	Effect			
No of studi es	Design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other considerati ons	Docetax el containi ng regimes	Non- docet axel- conta ining regim es	Relativ e (95% CI)	Absol ute	Quality	Importanc e
1	randomis ed trials	serious ⁸	no serious inconsistenc y	no serious indirectne ss	serious ¹	none	44	41	-	MD 2.13 higher (4.97 lower to 9.23 higher)	LOW	IMPORTAN T
Qualit	y of Life: En	notional Fu	nctioning (Bet	ter indicated	by lower va	lues)						
1	randomis ed trials	serious ⁸	no serious inconsistenc y	no serious indirectne ss	serious ¹	none	44	41	-	MD 8.06 higher (2.85 to 13.27 higher)	LOW	IMPORTAN T
Qualit	y of Life: Co	gnitive Fur	nctioning (Bett	er indicated	by lower va	lues)						
1	randomis ed trials	serious ⁸	no serious inconsistenc y	no serious indirectne ss	seriou ^{s1}	none	44	41	-	MD 3.6 lower (10.08 lower to 2.88 higher)	LOW	IMPORTAN T
Qualit	y of Life: So	cial Functi	oning (Better i	ndicated by	lower values	5)						
1	randomis ed trials	serious ⁸	no serious inconsistenc y	no serious indirectne ss	serious ¹	none	44	41	-	MD 7.5 higher (1.39 to	LOW	IMPORTAN T

Qualit	y assessme	nt					No of pati	ients	Effect			
No of studi es	Design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other considerati ons	Docetax el containi ng regimes	Non- docet axel- conta ining regim es	Relativ e (95% CI)	Absol ute	Quality	Importanc e
										13.61 higher)		
Qualit	y of Life: Glo	obal Quality	y of Life (Bette	r indicated b	y lower valu	Jes)						
1	randomis ed trials	serious ⁸	no serious inconsistenc y	no serious indirectne ss	serious ¹	none	44	41	-	MD 7.3 higher (0.64 to 13.96 higher)	LOW	IMPORTAN T

CI=confidence interval; RR=relative risk; HR=hazard ratio; MD=mean difference;

¹ 95% CI cross one deafult MID

² Al-Batran: allocation concealment unclear

³ Roth- Docetaxel dose reduced due to toxicity ⁴ 95% CI cross two default MIDs

⁵ Wang- unclear blinding of outcome assessors
 ⁶ I-squared statistic for heterogeneity > 75%

⁷ 0 events in one arm

⁸ Sadighi- only 71 participants included in QOL analysis (15 did not complete baseline questionnaire)

v assessme	nt					No of patients		Effect			
Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecisi on	Other considerati ons	Oral 5-FU prodrug (capecitabine) containing regime	IV 5- FU conta ining regim es	Relati ve (95% Cl)	Absol ute	Quality	Importanc e
I Survival											
randomis ed trials	no serio us risk of bias	no serious inconsisten cy	no serious indirectne ss	serious ¹	none	-	-	HR 0.87 (0.77 to 0.99)	-	ODERATE	CRITICAL
ssion-free	survival										
randomis ed trials	no serio us risk of bias	no serious inconsisten cy	no serious indirectne ss	serious ¹	none	-	-	HR 0.89 (0.79 to 1.01)	-	MODERAT E	CRITICAL
nent-related	death										
randomis ed trials	no serio us risk of bias	no serious inconsisten cy	no serious indirectne ss	very serious ²	none	1/156 (0.64%)	2/155 (1.3%)	RR 0.5 (0.05 to 5.42)	6 fewer per 1000 (from 12 fewer to 57 more)	LOW	IMPORTA NT
	Design I Survival randomis ed trials ession-free randomis ed trials	I Survival randomis ed trials ression-free survival randomis ed trials no ed trials randomis ed trials randomis ed trials no serio us risk of bias	DesignRisk of biasInconsiste ncyDesignRisk of Inconsiste ncyI Survivalno 	DesignRisk of biasInconsiste ncyIndirectn essI Survivalno serio us risk of biasno serious inconsisten cyno serious indirectne ssI Survivalno serio us risk of biasno serious inconsisten cyno serious indirectne ssrandomis ed trialsno serio us risk of biasno serious inconsisten cyno serious indirectne ssrandomis ed trialsno serio us risk of biasno serious inconsisten cyno serious indirectne ssnent-related death us risk of biasno serious inconsisten cyno serious indirectne ss	Design DesignRisk of biasInconsiste ncyIndirectn essImprecisi onI Survivalrandomis ed trialsno serio us risk of biasno serious inconsisten cyno serious indirectne 	Design DesignRisk of biasInconsiste ncyIndirectn essImprecisi onOther considerati onsI Survivalrandomis ed trialsno serio us risk of biasno serious inconsisten cyno serious indirectne ssserious1 nonenonerandomis ed trialsno serio us risk of biasno serious inconsisten cyno serious indirectne ssserious1 nonenonerandomis ed trialsno serio us risk of biasno serious inconsisten cyno serious serious indirectne ssserious1 nonenonerandomis ed trialsno serio us risk of biasno serious no serious inconsisten cyno serious1 serious2nonenent-related death randomis ed trialsno serio us risk ofno serious sindirectne ssno serious2 serious2none	Design DesignRisk of biasInconsiste ncyIndirectn essImprecisi onOther considerati onsOral 5-FU prodrug (capecitabine) containing regimeI Survivalrandomis ed trialsno serio us risk of biasno serious inconsisten cyno serious indirectne ssserious1none-randomis ed trialsno serio us risk of biasno serious inconsisten cyno serious indirectne ssserious1none-randomis ed trialsno seriou serio us risk of biasno serious indirectne ssserious1none-randomis ed trialsno seriou serio us risk of biasno serious indirectne ssserious1none-randomis ed trialsno serious inconsisten cyno serious ssserious1none-randomis ed trialsno serious inconsisten cyno serious ssserious2none1/156 (0.64%)	Design biasRisk of biasInconsiste ncyIndirectn essImprecisi onOther considerati onOral 5-FU prodrug (capecitabine) containing regimeIV 5- FU containing regimeI Survivalno serious serio us of biasno serious inconsisten cyno serious serious indirectne ssno serious1 no serious1nonerandomis ed trialsno serio us of biasno serious inconsisten cyno serious2 serious1nonerandomis ed trialsno serious inconsisten cyno serious2 indirectne ssserious1 serious1nonerandomis ed trialsno serious inconsisten cyno serious2 indirectne ssnonerandomis ed trialsno serio us risk of biasno serious2 indirectne ssno serious2 serious1nonerent-related death randomis ed trialsno serious2 inconsisten cyno serious2 serious2none1/156 (0.64%)2/155 (1.3%)	Design biasRisk of biasInconsiste ncyIndirectn essImprecisi onOther considerati onsOral 5-FU prodrug (capecitabine) containing regimeV 5- FU containing regimeRelati ve (95% CI)I SurvivalII </td <td>Design biasRisk of biasInconsiste ncyIndirectn essImprecisi onOther considerati onsOral 5-FU prodrug (capecitabine) containing regimeIV 5- FU containing regimeRelati ve (95%) (1)Absol uteI Survivalrandomis of serio us risk of biasno serious indirectne ssno serious seriousi no serious indirectne ssnoneHR 0.87 (0.77 to 0.99)-randomis ed trialsno serio us risk of biasno serious indirectne ssserious1 serious indirectne ssnoneHR 0.87 (0.77 to 0.99)-randomis ed trialsno serio us risk of biasno serious indirectne ssserious1 serious1noneHR 0.87 (0.77 to 0.99)-rent-related death randomis ed trialsno serious inconsisten cyno serious serious2serious2 serious2none1/156 (0.64%)2/155 (1.3% (1.3%)RR fewer per fuon 12 fewer to 57</td> <td>Design biasRisk ncyInconsiste ncyIndirectn essImprecisi onOther considerati onsOral 5-FU 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serious indirectne ssserious1 serious indirectne ssnoneHR 0.87 (0.77 to 0.99)-randomis ed trialsno serio us risk of biasno serious indirectne ssserious1 serious1noneHR 0.87 (0.77 to 0.99)-rent-related death randomis ed trialsno serious inconsisten cyno serious serious2serious2 serious2none1/156 (0.64%)2/155 (1.3% (1.3%)RR fewer per fuon 12 fewer to 57	Design biasRisk ncyInconsiste ncyIndirectn essImprecisi onOther considerati onsOral 5-FU prodrug (capecitabine, ontaining regimeRelati regimeAbsol uteI Survivalrandomis ed trialsno seriousno serious inconsisten cyno serious ssnoneHR 0.87 (0.77 to 0.99)-ODERATErandomis us risk of biasno serious inconsisten cyno serious indirectne ssnoneHR 0.87 (0.77 to 0.99)-ODERATErandomis us risk of biasno serious inconsisten cyno serious indirectne ssnoneHR 0.87 (0.77 to 0.99)-ODERATErandomis us reston-freeno serious inconsisten cyno serious indirectne ssnoneHR 0.87 (0.79 to 1.01)-DERATErent-related CettrWODERAT 0.89 to 1.01)-MODERAT Erendomis us risk of biasno serious indirectne ssserious² serious²none <td< td=""></td<>

Table 39: Summary clinical evidence. Oral 5-FU prodrug (capecitabine) combinations versus intravenous 5-FU combinations

Qualit	y assessme	nt					No of patients		Effect			
No of studi es	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecisi on	Other considerati ons	Oral 5-FU prodrug (capecitabine) containing regime	IV 5- FU conta ining regim es	Relati ve (95% Cl)	Absol ute	Quality	Importanc e
1	randomis ed trials	no serio us risk of bias	no serious inconsisten cy	no serious indirectne ss	very serious ²	none	28/156 (17.9%)	28/15 5 (18.1 %)	RR 0.99 (0.62 to 1.6)	2 fewer per 1000 (from 69 fewer to 108 more)	LOW	CRITICAL
Treatn	nent-related	toxicity	/: Nausea and	vomiting								
1	randomis ed trials	no serio us risk of bias	no serious inconsisten cy	no serious indirectne ss	serious ¹	none	47/494 (9.5%)	60/50 8 (11.8 %)	RR 0.81 (0.56 to 1.16)	22 fewer per 1000 (from 52 fewer to 19 more)	MODERAT E	CRITICAL
Treatn	nent-related	toxicity	/: Diarrhoea									
1	randomis ed trials	no serio us risk of bias	no serious inconsisten cy	no serious indirectne ss	serious ¹	none	42/494 (8.5%)	33/50 8 (6.5%)	RR 1.31 (0.84 to 2.03)	20 more per 1000 (from 10 fewer to 67 more)	MODERAT E	CRITICAL

CI=confidence interval; RR=relative risk; HR=hazard ratio; IV=intravenous; 5-FU=5-fluouracil ¹ 95% CI crosses one default MID ² 95% CI crosses two default MIDs

Table 40: Clinical evidence summary. Cisplatin containing regimes versus oxaliplatin containing regimes

Qualit	y assessmer	nt					No of pat	ients	Effect			
No of studi es	Design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecisi on	Other consideratio ns	Cisplati n containi ng regimes	Oxali platin conta ining regim es	Relat ive (95% Cl)	Absolu te	Quality	Importance
Overa	I Survival											
2	randomise d trials	serious ¹	no serious inconsistenc y	no serious indirectnes s	no serious imprecisio n	none	-	-	HR 0.91 (0.80 to 1.04)	-	MODERAT E	CRITICAL
Progre	ession-free s	urvival										
2	randomise d trials	serious1	no serious inconsistenc y	no serious indirectnes s	serious ²	none	-	-	HR 0.90 (0.79 to 1.02)	-	LOW	CRITICAL
Treatn	nent-related	death										
3	randomise d trials	serious _{3,4}	no serious inconsistenc y	no serious indirectnes s	very serious⁵	none	1/187 (0.53%)	3/176 (1.7%)	RR 0.42 (0.06 to 2.81)	10 fewer per 1000 (from 16 fewer to 31 more)	VERY LOW	IMPORTAN T

Qualit	y assessmer	nt					No of pat	ients	Effect			
No of studi es	Design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecisi on	Other consideratio ns	Cisplati n containi ng regimes	Oxali platin conta ining regim es	Relat ive (95% Cl)	Absolu te	Quality	Importance
Treatn	nent discont	inuation du	ue to toxicity									
1	randomise d trials		no serious inconsistenc y	no serious indirectnes s	very serious⁵	none	12/112 (10.7%)	11/10 2 (10.8 %)	RR 0.99 (0.46 to 2.15)	1 fewer per 1000 (from 60 fewer to 114 more)	VERY LOW	CRITICAL
Treatn	nent-related	toxicity: A	ny grade 3/4 ev	vent								
1	randomise d trials	serious ⁴	no serious inconsistenc y	no serious indirectnes s	very serious⁵	none	26/39 (66.7%)	25/38 (65.8 %)	RR 1.01 (0.74 to 1.39)	7 more per 1000 (from 171 fewer to 257 more)	VERY LOW	CRITICAL
Treatn	nent-related	toxicity: Di	arrhoea									
1	randomise d trials	no serious risk of bias	no serious inconsistenc y	no serious indirectnes s	no serious imprecisio n	none	55/489 (11.2%)	19/51 3 (3.7%)	RR 3.04 (1.83 to 5.04)	76 more per 1000 (from 31 more to 150 more)	HIGH	CRITICAL

	y assessmer						No of pat		Effect			
No of studi es	Design	Risk of bias	Inconsisten cy	Indirectne SS	Imprecisi on	Other consideratio ns	Cisplati n containi ng regimes	Oxali platin conta ining regim es	Relat ive (95% CI)	Absolu te	Quality	Importance
1	randomise d trials	no serious risk of bias	no serious inconsistenc y	no serious indirectnes s	serious ²	none	62/489 (12.7%)	46/51 3 (9%)	RR 1.41 (0.99 to 2.03)	37 more per 1000 (from 1 fewer to 92 more)	MODERAT E	CRITICAL

CI=confidence interval; RR=relative risk; HR=hazard ratio;

¹ Al-Batran 2008: baseline differences between groups in sex and metastatic disease

² 95% Cl crosses one default MID
 ³ Popov 2008: risk of bias in outcome reporting, not ITT
 ⁴ Kim 2014: unclear randomization process, allocation concealment
 ⁵ 95% Cl crosses two default MIDs

Table 41: Clinical evidence summary. 5-FU containing regimes versus non-5FU containing regimes

Qualit	y assessmen	ıt					No of patie	nts	Effect			
No of studi es	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecisi on	Other considerati ons	5FU- containin g regimes	Non- 5FU contai ning regime s	Relati ve (95% Cl)	Absol ute	Quality	Importanc e
Overa	ll survival											
3	randomise d trials	serious ⁴ serious ²	no serious inconsisten cy	no serious indirectne ss	no serious	none	-	-	HR 0.59 (0.39	-	MODERA TE	CRITICAL

Qualit	y assessmer	nt					No of patie	nts	Effect			
No of studi es	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecisi on	Other considerati ons	5FU- containin g regimes	Non- 5FU contai ning regime s	Relati ve (95% Cl)	Absol ute	Quality	Importan e
					imprecisi on				to 0.81)			
Overa	ll survival - D	ocetaxel/p	latinum based	d +/- 5-FU								
1	randomise d trials	no serious risk of bias	no serious inconsisten cy	no serious indirectne ss	serious ²	none	-	-	HR 0.61 (0.45 to 0.84)	-	MODERA TE	CRITICAL
Overa	ll survival – 🤅	5-FU versu	s cisplatin reg	jimen								
1	randomise d trials	serious ⁴ serious ²	no serious inconsisten Cy	no serious indirectne ss	serious ²	none	-	-	HR 0.56 (0.39 to 0.81)	-	LOW	CRITICAL
Two y	ear survival-	5-FU versu	us irinotecan r	egimen								
1	randomise d trials	serious ³	no serious inconsisten cy	no serious indirectne ss	very serious ⁴	none	6/42 (14.3%)	2/43 (4.7%)	RR 3.07 (0.66 to 14.37)	96 more per 1000 (from 16 fewer to 622 more)	VERY LOW	CRITICAL
Progr	ession-free s	urvival										
2	randomise d trials	serious ⁴ serious ²	no serious inconsisten cy	no serious indirectne ss	no serious	none	-	-	HR 0.37 (0.28	-	MODERA TE	CRITICA

Qualit	y assessmer	it					No of patie	nts	Effect			
No of studi es	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecisi on	Other considerati ons	5FU- containin g regimes	Non- 5FU contai ning regime s	Relati ve (95% Cl)	Absol ute	Quality	Importan e
					imprecisi on				to 0.48)			
Progre	ession-free s	urvival - Do	ocetaxel/platir	num based +	/- 5-FU							
1	randomise d trials	no serious risk of bias	no serious inconsisten cy	no serious indirectne ss	no serious imprecisi on	none	-	-	HR 0.34 (0.25 to 0.48)	-	HIGH	CRITICAL
Progre	ession-free s	urvival – 5	-FU versus pla	atinum regim	ien							
1	randomise d trials	serious ⁴ serious ²	no serious inconsisten cy	no serious indirectne ss	no serious imprecisi on	none	-	-	HR 0.41 (0.26 to 0.64)	-	MODERA TE	CRITICAL
Treatn	nent-related	death										
1	randomise d trials	serious ¹ serious ²	no serious inconsisten cy	no serious indirectne ss	very serious ^{4,5}	none	0/72 (0%)	1/74 (1.4%)	RR 0.34 (0.01 to 8.27)	9 fewer per 1000 (from 13 fewer to 98 more)	VERY LOW	IMPORTA NT
Treatn	nent disconti	nuation du	e to toxicity									
2	randomise d trials	serious	no serious inconsisten cy	no serious indirectne ss	very serious ⁴	none	10/114 (8.8%)	16/117 (13.7%	RR 0.64 (0.31	49 fewer per	VERY LOW	CRITICA

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Qualit	y assessmer	nt					No of patie	nts	Effect			
No of studi es	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecisi on	Other considerati ons	5FU- containin g regimes	Non- 5FU contai ning regime s	Relati ve (95% Cl)	Absol ute	Quality	Importance
									to 1.34)	1000 (from 94 fewer to 46 more)		
Treatr	nent disconti	nuation du	e to toxicity -	- 5-FU versus	s irinotecan	regimen						
1	randomise d trials	serious ³	no serious inconsisten cy	no serious indirectne ss	very serious ⁴	none	6/42 (14.3%)	10/43 (23.3%)	RR 0.61 (0.25 to 1.54)	91 fewer per 1000 (from 174 fewer to 126 more)	VERY LOW	CRITICAL
Treatr	nent disconti	nuation du	e to toxicity -	- 5-FU versus	s cisplatin re	egimen						
1	randomise d trials	serious ⁴ serious ²	no serious inconsisten cy	no serious indirectne ss	very serious ⁴	none	4/72 (5.6%)	6/74 (8.1%)	RR 0.69 (0.2 to 2.33)	25 fewer per 1000 (from 65 fewer to 108 more)	VERY LOW	CRITICAL

	y assessmen						No of patie		Effect			
No of studi es	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecisi on	Other considerati ons	5FU- containin g regimes	Non- 5FU contai ning regime s	Relati ve (95% Cl)	Absol ute	Quality	Importanc e
1	randomise d trials	Serious ³	no serious inconsisten cy	no serious indirectne ss	no serious imprecisi on	none	18/42 (42.9%)	7/43 (16.3%)	RR 2.63 (1.23 to 5.64)	265 more per 1000 (from 37 more to 755 more)	MODERA TE	CRITICAL
Treatn	nent-related t	toxicity: Na	usea and von	niting- 5-FU	versus irino	tecan						
1	randomise d trials	serious ³	no serious inconsisten cy	no serious indirectne ss	serious ²	none	7/42 (16.7%)	1/43 (2.3%)	RR 7.17 (0.92 to 55.76)	143 more per 1000 (from 2 fewer to 1000 more)	LOW	CRITICAL

CI=confidence interval; RR=relative risk; HR=hazard ratio; 5-FU=5-fluouracil

1 95% CI crosses one default MID

2 Pozzo 2004: unclear randomization and allocation concealement

3 Roy 2012: unclear randomization and allocation concealment

4 95% CI crosses two default MIDs

5 0 events in one arm

Quality	, assessmen	•	,		- J - J		No of patient		Effect			
No of studi es	Design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecisi on	Other consideratio ns	Platinum containing regimes	Taxa ne conta ining regim es	Relati ve (95% CI)	Absolut e	Quali ty	Importance
Overal	l survival											
1	randomise d trials	seriou s ¹	no serious inconsistenc y	no serious indirectnes s	serious ²	none	-	-	HR 0.75 (0.47 to 1.2)	-	LOW	CRITICAL
Treatm	nent-related	death										
1	randomise d trials	seriou s ¹	no serious inconsistenc y	no serious indirectnes s	very serious ³	none	2/48 (4.2%)	1/46 (2.2%)	RR 1.92 (0.18 to 20.42)	20 more per 1000 (from 18 fewer to 422 more)	VER Y LOW	IMPORTAN T
Treatm	nent disconti	nuation of	due to toxicity									
1	randomise d trials	seriou s ¹	no serious inconsistenc y	no serious indirectnes s	very serious ³	none	6/48 (12.5%)	4/46 (8.7%)	RR 1.44 (0.43 to 4.77)	38 more per 1000 (from 50 fewer to 328 more)	VER Y LOW	CRITICAL
Treatm	nent-related	toxicity: /	Any grade 3/4 e	event								
1	randomise d trials	seriou s ¹	no serious inconsistenc y	no serious indirectnes s	serious ²	none	33/48 (68.8%)	27/46 (58.7 %)	RR 1.17 (0.86 to 1.59)	100 more per 1000 (from 82 fewer to 346 more)	LOW	CRITICAL

Table 42: Clinical evidence summary. Platinum containing regimens versus taxane containing regimens

CI=confidence interval; RR=relative risk; HR=hazard ratio; ¹ Lee 2015: unclear randomization, allocation concealment and blinding

² 95% CI cross one default MID
 ³ 95% CI crosses two default MIDs

Table 43: Clinical evidence summary. Epirubicin/cisplatin/capetibacine combinations versus 5-FU/irinotecan combinations

y assessme	ent					No of patients Effect					
Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecis ion	Other considerati ons	Epirubicin/cispl atin/capetibacin e containing regimes	5-FU/Irinotecan containing regimes	Relat ive (95% Cl)	Absol ute	Qua lity	Importanc e
ll survival											
randomis ed trials	no serio us risk of bias	no serious inconsisten cy	no serious indirectne ss	no serious imprecisi on	none	-	-	HR 1.01 (0.82 to 1.24)	-	HIG H	CRITICAL
ession-free	surviva	al									
randomis ed trials	no serio us risk of bias	no serious inconsisten cy	no serious indirectne ss	no serious imprecisi on	none	-	-	HR 0.99 (0.81 to 1.21)	-	HIG H	CRITICAL
nent-related	d death										
randomis ed trials	no serio us risk of bias	no serious inconsisten cy	no serious indirectne ss	very serious ¹	none	7/209 (3.3%)	5/207 (2.4%)	RR 1.39 (0.45 to 4.3)	9 more per 1000 (from 13 fewer to 80	LO W	IMPORTA NT
	Design II survival randomis ed trials ession-free randomis ed trials	Il survival randomis ed trials risk of bias ession-free survive randomis ed trials randomis randomis randomis randomis ed trials serio us risk of bias	DesignRisk of biasInconsiste ncyIl survivalno serio us risk of biasno serious inconsisten cyrandomis ed trialsno serio us risk of biasno serious inconsisten cyrandomis ed trialsno serio us risk of biasno serious inconsisten cyrandomis ed trialsno serio us risk of biasno serious inconsisten cyrandomis ed trialsno serio us risk of biasno serious inconsisten cynent-related death us ed trialsno serio us risk ofno serious inconsisten cy	DesignRisk of biasInconsiste ncyIndirectn essII survivalrandomis ed trialsno serio us risk of biasno serious inconsisten cyno serious indirectne ssrandomis ed trialsno serio us risk of biasno serious inconsisten cyno serious indirectne ssrandomis ed trialsno serio us risk of biasno serious inconsisten cyno serious indirectne serious indirectne serious indirectne ssrandomis ed trialsno serio us risk of biasno serious cyno serious indirectne ssrandomis ed trialsno serio us risk ofno serious inconsisten cyno serious indirectne serious indirectne serious inconsisten cy	DesignRisk of biasInconsiste ncyIndirectn essImprecis ionII survivalrandomis ed trialsno serio us risk of biasno serious inconsisten cyno serious indirectne serious	Design DesignRisk of biasInconsiste ncyIndirectn essImprecis ionOther considerati onsII survivalrandomis ed trialsno serious inconsisten cyno serious indirectne ssno serious indirectne ssno serious imprecisi onnoneed trials ed trialsno serious inconsisten cyno serious indirectne ssno serious imprecisi onnoneed trials ed trialsno serious inconsisten cyno serious indirectne ssno serious imprecisi onnonerandomis ed trialsno serious inconsisten cyno serious indirectne ssno serious imprecisi onnonerandomis ed trialsno serious inconsisten cyno serious indirectne ssno serious imprecisi onnonerandomis ed trialsno serious inconsisten cyno serious indirectne ssvery serious^1none	Design DesignRisk of biasInconsiste ncyIndirectn essImprecis ionOther consideratiEpirubicin/cispl atin/capetibacin e containing regimesII survivalrandomis ed trialsno serious inconsisten cyno serious indirectne ssno serious indirectne serious indirectne ssno serious imprecisi onnone-randomis ed trialsno serious inconsisten cyno serious indirectne ssno serious indirectne seriousnone-randomis ed trials ed trials ofno serious serious inconsisten cyno serious serious indirectne seriousnone serious (3.3%)7/209 (3.3%)	Design biasRisk of biasInconsiste ncyIndirectn essImprecis ionOther considerati onsEpirubicin/cispl atin/capetibacin e containing regimes5-FU/Irinotecan containing regimesII survivalrandomis ed trialsno seriou inconsisten cyno serious serious indirectne ssno serious on onno serious imprecisi onnone e-randomis ed trialsno seriou sinconsisten cyno serious serious indirectne ssno serious onnone serious imprecisi on-randomis ed trialsno seriou seriou seriou srisk of biasno serious serious inconsisten cyno serious serious indirectne ssnone serious imprecisi on-randomis ed trialsno seriou seriou serious risk ofno serious serious serious indirectne ssnone serious serious imprecisi onnone serious serious imprecisi on-randomis ed trialsno serious serious indirectne ssno serious serious indirectne ssnone serious serious serious indirectne ss1000000000000000000000000000000000000	Design biasRisk ncyInconsiste ncyIndirectn essImprecis ionOther considerati onsEpirubicin/cispl atin/capetibacin e containing regimes5-FU/Irinotecan containing regimesRelat ive (95% CI)Il survivalrandomis ed trialsno serious inconsisten us risk of biasno serious inconsisten cyno serious indirectne ssno no serious indirectne ssnone serious imprecisi onHR 1.01 (0.82 to 1.24)ession-free serious risk of biasno serious inconsisten cyno serious indirectne ssno serious imprecisi onnone serious imprecisi onHR 0.92 (0.81 to 1.21)ed trials ed trials ed trialsno serious inconsisten cyno serious indirectne ssno serious imprecisi onnone serious imprecisi onHR 0.99 (0.81 to 1.21)ment-related death randomis ed trials wisk ofno serious inconsisten cyno serious serious indirectne ssnone serious' on7/209 (3.3%)5/207 (2.4%)RR 1.39 (0.45 to 4.3)	Design biasRisk of biasInconsiste ncyIndirectn essImprecis ionOther considerati onsEpirubicin/cispl atin/capetibacin e containing regimes5-FU/Irinotecan containing regimesRelat ive (95% (1)AbsoluteII survivalInconsisten randomis ed trialsno serious inconsisten cyno serious inconsisten cyno serious indirectne ssno serious indirectne ssno serious indirectne ssno serious indirectne ssno serious imprecisi onnone eHR 1.01 (0.82 to to 1.24)-randomis ed trialsno serious inconsisten cyno serious indirectne ssno serious indirectne ssno serious imprecisi onnone serious imprecisi on-HR randomis ed trialsno serious inconsisten cyno serious indirectne ssno serious imprecisi onnone serious imprecisi onHR to <b< td=""><td>Design biasRisk of biasInconsiste ncyIndirectn essImprecis ionOther considerati onsEpirubicin/cisp atin/capetibacin e containing regimes5-FU/Irinotecan containing regimesRelat ive (95%, (21)Absolute Qua Qua ItiII survivalIImprecisionNo serious indirectne</td></b<>	Design biasRisk of biasInconsiste ncyIndirectn essImprecis ionOther considerati onsEpirubicin/cisp atin/capetibacin e containing regimes5-FU/Irinotecan containing regimesRelat ive (95%, (21)Absolute Qua Qua ItiII survivalIImprecisionNo serious indirectne

Qualit	ty assessme	ent					No of patients		Effect			
No of stud ies	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecis ion	Other considerati ons	Epirubicin/cispl atin/capetibacin e containing regimes	5-FU/Irinotecan containing regimes	Relat ive (95% Cl)	Absol ute	Qua lity	Importanc e
1	randomis ed trials	no serio us risk of bias	no serious inconsisten cy	no serious indirectne ss	no serious imprecisi on	none	135/209 (64.6%)	79/207 (38.2%)	RR 1.69 (1.39 to 2.07)	263 more per 1000 (from 149 more to 408 more)	HIG H	CRITICAL

CI=confidence interval; RR=relative risk; HR=hazard ratio; 5-FU=5-Fluouracil ¹ *Downgraded for serious imprecision: 95% CI crosses two default MIDs*

G.16 Second-line palliative chemotherapy

What is the optimal palliative second-line chemotherapy for locally-advanced or metastatic oesophago-gastric cancer?

Table 44: Clinical evidence	profile for 5-FU	versus paclitaxel

Quality	assessment	l					Nº of pa	tients	Effect			
Nº of studie s	Study design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other consideratio ns	5FU	paclitax el	Relativ e (95% Cl)	Absolu te (95% Cl)	Quality	Importanc e
overall	survival											
1	randomise d trials	seriou s ^a	not serious	not serious	serious ^b	none	-/49	-/51	HR 0.89 (0.57	-	LOW	CRITICAL

Quality	assessment						Nº of pa	atients	Effect			
Nº of studie s	Study design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other consideratio ns	5FU	paclitax el	Relativ e (95% CI)	Absolu te (95% Cl)	Quality	Importanc e
									to 1.38)			
progres	ssion free su	rvival										
1	randomise d trials	seriou s ^a	not serious	not serious	not serious	none	-/49	-/51	HR 0.58 (0.38 to 0.88)	-	MODERA TE	IMPORTA NT
nausea												
1	randomise d trials	seriou s ^a	not serious	not serious	very serious ^c	none	3/49 (6.1%)	0/51 (0.0%)	RR 7.28 (0.39 to 137.38)	0 fewer per 1,000 (from 0 fewer to 0 fewer)	VERY LOW	CRITICAL
neutrop	oaenic sepsis	\$										
1	randomise d trials	seriou S ^a	not serious	not serious	very serious ^b	none	2/49 (4.1%)	0/51 (0.0%)	RR 5.20 (0.26 to 105.65)	0 fewer per 1,000 (from 0 fewer to 0 fewer)	VERY LOW	CRITICAL
neutrop	baenia											
1	randomise d trials	seriou s ^a	not serious	not serious	serious ^d	none	14/49 (28.6 %)	6/51 (11.8%)	RR 2.43 (1.02 to 5.81)	168 more per 1,000 (from 2 more to 566 more)	LOW	CRITICAL

Quality	assessment	:					Nº of pa	tients	Effect			
№ of studie s	Study design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other consideratio ns	5FU	paclitax el	Relativ e (95% Cl)	Absolu te (95% Cl)	Quality	Importanc e
diarrho	ea											
1	randomise d trials	seriou s ^a	not serious	not serious	serious ^c	none	5/49 (10.2 %)	0/51 (0.0%)	RR 11.44 (0.65 to 201.55)	0 fewer per 1,000 (from 0 fewer to 0 fewer)	LOW	CRITICAL
treatme	ent related m	ortality										
1	randomise d trials	seriou s ^a	not serious	not serious	serious ^c	none	1/49 (2.0%)	0/51 (0.0%)	RR 3.12 (0.13 to 74.80)	0 fewer per 1,000 (from 0 fewer to 0 fewer)	LOW	IMPORTA NT

CI: Confidence interval; HR: Hazard Ratio; RR: Risk ratio a. No blinding b. 95% CI of the effect includes no effect and clinically important benefit and harm c. 95% CI of the effect includes both default MID thresholds

d. 95% CI of the effect includes one default MID threshold

Table 45: Clinical evidence profile for docetaxel or irinotecan versus BSC

Quality	assessment	t					Nº of patier	nts	Effect			
Nº of studie s	Study design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other consideratio ns	docetaxel or inrinotec an	BSC	Relativ e (95% Cl)	Absolu te (95% Cl)	Quality	Importanc e
overall	overall survival											
1	randomise d trials	seriou s ^a	not serious	serious ^b	not serious	none	-/126	-/62	HR 0.71 (0.54 to 0.97)	-	LOW	CRITICAL

Quality	assessment	:					Nº of patie	nts	Effect			
Nº of studie s	Study design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other consideratio ns	docetaxel or inrinotec an	BSC	Relativ e (95% CI)	Absolu te (95% Cl)	Quality	Importanc e
progres	ssion free su	rvival - n	ot reported									
-	-	-	-	-	-	-	-	-	-	-	-	IMPORTA NT
nausea												
1	randomise d trials	seriou s ^a	not serious	serious ^b	serious °	none	19/126 (15.1%)	20/62 (32.3 %)	RR 0.47 (0.27 to 0.81)	171 fewer per 1,000 (from 61 fewer to 235 fewer)	VERY LOW	CRITICAL
neutrop	baenic sepsis	5										
1	randomise d trials	seriou s ^a	not serious	serious ^b	very serious d	none	6/126 (4.8%)	0/62 (0.0%)	RR 6.45 (0.37 to 112.67)	0 fewer per 1,000 (from 0 fewer to 0 fewer)	VERY LOW	CRITICAL
neutrop	oaenia											
1	randomise d trials	seriou s ^a	not serious	serious ^b	not serious	none	76/126 (60.3%)	8/62 (12.9 %)	RR 4.67 (2.41 to 9.06)	474 more per 1,000 (from 182 more to 1,000 more)	LOW	CRITICAL
diarrho	ea											

Quality	assessment	t					Nº of patier	nts	Effect			
№ of studie s	Study design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other consideratio ns	docetaxel or inrinotec an	BSC	Relativ e (95% CI)	Absolu te (95% CI)	Quality	Importanc e
1	randomise d trials	seriou s ^a	not serious	serious ^b	very serious ^d	none	18/126 (14.3%)	11/62 (17.7 %)	RR 0.81 (0.41 to 1.60)	34 fewer per 1,000 (from 105 fewer to 106 more)	VERY LOW	CRITICAL
treatme	reatment related mortality - not reported											
-	-	-	-	-	-	-	-	-	-	-	-	IMPORTA NT

CI: Confidence interval; HR: Hazard Ratio; RR: Risk ratio; OR: Odds ratio

a. Unclear allocation concealment and blinding b. In the chemotherapy arm choice of drug was at the treating physician's discretion

c. 95% CI of the effect includes one default MID threshold

d. 95% CI of the effect includes both default MID thresholds

Table 46: Clinical evidence profile for docetaxel + cisplatin versus docetaxel + S-1

Quality	assessment	t					Nº of pati	ents	Effect			
Nº of studi es	Study design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other consideratio ns	docetax el + cisplati n	docetax el + S-1	Relativ e (95% Cl)	Absolu te (95% Cl)	Quality	Importanc e
overall	overall survival - not reported											
-	-	-	-	-	-	-	-	-	-	-	-	CRITICAL
progre	ssion free su	rvival - n	ot reported									
-	-	-	-	-	-	-	-	-	-	-	-	IMPORTA NT
nausea - not reported												
-	-	-	-	-	-	-	-	-	-	-	-	CRITICAL

Quality	assessment	t					Nº of pati	ents	Effect			
Nº of studi es	Study design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other consideratio ns	docetax el + cisplati n	docetax el + S-1	Relativ e (95% CI)	Absolu te (95% Cl)	Quality	Importanc e
neutro	paenic sepsis	S										
1	randomise d trials	seriou s ^a	not serious	serious ^b	very serious ^c	none	3/24 (12.5%)	1/23 (4.3%)	RR 2.88 (0.32 to 25.68)	82 more per 1,000 (from 30 fewer to 1,000 more)	VERY LOW	CRITICAL
neutro	paenia											
1	randomise d trials	seriou s ^a	not serious	serious ^b	very serious ^c	none	6/24 (25.0%)	3/23 (13.0%)	RR 1.92 (0.54 to 6.77)	120 more per 1,000 (from 60 fewer to 753 more)	VERY LOW	CRITICAL
diarrho	ea - not repo	orted										
-	-	-	-	-	-	-	-	-	-	-	-	CRITICAL
treatmo	ent related m	ortality -	not reported									
-	-	-	-	-	-	-	-	-	-	-	-	IMPORTA NT

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Table 47: Clinical evidence profile for docetaxel versus BSC

Quality	assessment	:					Nº of pati	ents	Effect			
№ of studie s	Study design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecisi on	Other consideratio ns	docetax el	BSC	Relativ e (95% Cl)	Absolu te (95% Cl)	Quality	Importanc e
overall	survival											
1	randomise d trials	seriou s ^a	not serious	not serious	not serious	none	-/84	-/84	HR 0.67 (0.49 to 0.92)	-	MODERA TE	CRITICAL
progres	ssion free su	rvival										
1	randomise d trials	seriou s ^a	not serious	not serious	not serious	none	-/84	-/84	HR 0.67 (0.48 to 0.93)	-	MODERA TE	IMPORTA NT
nausea	- not reporte	d										
-	-	-	-	-	-	-	-	-	-	-	-	CRITICAL
neutrop	oaenic sepsis	5										
1	randomise d trials	seriou s ^a	not serious	not serious	very serious ^b	none	6/84 (7.1%)	0/84 (0.0%)	RR 13.00 (0.74 to 227.16)	0 fewer per 1,000 (from 0 fewer to 0 fewer)	VERY LOW	CRITICAL
neutrop	oaenia											
1	randomise d trials	seriou s ^a	not serious	not serious	not serious	none	18/84 (21.4%)	0/84 (0.0%)	RR 37.00 (2.27 to 604.13)	0 fewer per 1,000 (from 0 fewer to 0 fewer)	MODERA TE	CRITICAL
diarrho	ea - not repo	rted										

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Quality	assessment	:					Nº of pati	ents	Effect			
№ of studie s	Study design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecisi on	Other consideratio ns	docetax el	BSC	Relativ e (95% Cl)	Absolu te (95% Cl)	Quality	Importanc e
-	-	-	-	-	-	-	-	-	-	-	-	CRITICAL
treatme	ent related m	ortality -	not reported									
-	-	-	-	-	-	-	-	-	-	-	-	IMPORTA NT

CI: Confidence interval; HR: Hazard Ratio; OR: Odds ratio; RR: Risk ratio

a. no blinding
 b. 95% CI of the effect includes both default MID thresholds

Table 48: Clinical evidence profile for docetaxel versus docetaxel + 5'DFUR

Quality	assessment	t					Nº of pati	ents	Effect			
Nº of studi es	Study design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other considerati ons	docetax el	docetax el plus 5'DFUR	Relati ve (95% Cl)	Absolu te (95% CI)	Quality	Importanc e
overall	survival											-
1	randomise d trials	seriou s ^a	not serious	not serious	not serious	none	-/12	-/12	HR 3.11 (1.22 to 7.97)	-	MODERA TE	CRITICAL
progres	ssion free su	i <mark>rvival</mark> - r	ot reported									
-	-	-	-	-	-	-	-	-	-	-	-	IMPORTA NT
nausea	I											
1	randomise d trials	seriou s ^a	not serious	serious ^b	serious ^c	none	1/12 (8.3%)	0/12 (0.0%)	RR 3.00 (0.13 to 67.06)	0 fewer per 1,000 (from 0 fewer to 0 fewer)	VERY LOW	CRITICAL

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Quality	/ assessmen	t					Nº of pat	ents	Effect			
№ of studi es	Study design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other considerati ons	docetax el	docetax el plus 5'DFUR	Relati ve (95% Cl)	Absolu te (95% Cl)	Quality	Importanc e
neutro	paenic sepsi	s - not re	eported									
-	-	-	-	-	-	-	-	-	-	-	-	CRITICAL
neutro	paenia											
1	randomise d trials	seriou S ^a	not serious	serious ^b	serious c	none	4/12 (33.3%)	4/12 (33.3%)	RR 1.00 (0.32 to 3.10)	0 fewer per 1,000 (from 227 fewer to 700 more)	VERY LOW	CRITICAL
diarrho	oea - not repo	orted										
-	-	-	-	-	-	-	-	-	-	-	-	CRITICAL
treatm	ent related m	ortality -	not reported									
-	-	-	-	-	-	-	-	-	-	-	-	IMPORTA NT

CI: Confidence interval; HR: Hazard Ratio; RR: Risk ratio; OR: Odds ratio

a. Unclear risk of bias due to study limitations - due to poor reporting of study b. Unclear definitions of morbidity outcomes c.95% CI of the effect includes both default MID thresholds

Table 49: Clinical evidence profile for docetaxel versus docetaxel + oxaliplatin

Quality	assessment	t					Nº of pati	ents	Effect			
Nº of studi es	Study design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other considerati ons	docetax el	docetax el plus platinu m	Relati ve (95% Cl)	Absolu te (95% Cl)	Quality	Importanc e
overall	survival											
1	randomise d trials	seriou s ^a	not serious	not serious	serious ^b	none	-/27	-/25	HR 1.17 (0.67	-	LOW	CRITICAL

Quality	assessmen	t					Nº of pati	ients	Effect			
Nº of studi es	Study design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other considerati ons	docetax el	docetax el plus platinu m	Relati ve (95% Cl)	Absolu te (95% Cl)	Quality	Importanc e
									to 2.04)			
progres	ssion free su	irvival										
1	randomise d trials	seriou s ^a	not serious	not serious	not serious	none	-/27	-/25	HR 0.50 (0.27 to 0.91)	-	MODERA TE	IMPORTA NT
nausea	l i i i i i i i i i i i i i i i i i i i											
1	randomise d trials	seriou s ^a	not serious	not serious	serious ^c	none	0/27 (0.0%)	1/25 (4.0%)	RR 0.31 (0.01 to 7.26)	28 fewer per 1,000 (from 40 fewer to 250 more)	LOW	CRITICAL
neutro	paenic sepsi	s										
2	randomise d trials	seriou s ^{c,d}	not serious	serious ^e	serious ^f	none	2/50 (4.0%)	8/49 (16.3%)	RR 0.29 (0.08 to 1.12)	116 fewer per 1,000 (from 20 more to 150 fewer)	VERY LOW	CRITICAL
neutro	paenia											
2	randomise d trials	seriou s ^{a,d}	not serious	serious ^e	serious ^f	none	5/50 (10.0%)	14/49 (28.6%)	RR 0.38 (0.16	177 fewer per	VERY LOW	CRITICAL

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Quality	/ assessmen	t					Nº of pati	ents	Effect			
№ of studi es	Study design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other considerati ons	docetax el	docetax el plus platinu m	Relati ve (95% Cl)	Absolu te (95% Cl)	Quality	Importance
									to 0.93)	1,000 (from 20 fewer to 240 fewer)		
diarrho	bea											
1	randomise d trials	seriou s ^{a,d}	not serious	serious ^e	very serious ^c	none	0/27 (0.0%)	1/25 (4.0%)	RR 0.31 (0.01 to 7.26)	28 fewer per 1,000 (from 40 fewer to 250 more)	VERY LOW	CRITICAL
treatmo	ent related m	ortality -	not reported									
-	-	-	-	-	-	-	-	-	-	-	-	IMPORTA NT

CI: Confidence interval; HR: Hazard Ratio; RR: Risk ratio; OR: Odds ratio

a. unclear risk of bias due to poor reporting of study

b.95% CI of effect includes the possibility of clinically significant benefit and harm

c.95% CI of the effect includes both default MID thresholds d. no blinding

e. unclear definitions of morbidity outcomes

f.95% CI of the effect includes one default MID threshold

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Table 50: Clinical evidence profile for docetaxel versus docetaxel + S-1

Quality	assessment	t					Nº of pati	ents	Effect			
№ of studie s	Study design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other consideratio ns	docetax el	docetac el plus S-1	Relativ e (95% CI)	Absolu te (95% Cl)	Quality	Importanc e
overall	survival - no	t reporte	d									
-	-	-	-	-	-	-	-	-	-	-	-	CRITICAL
progres	ssion free su	<mark>rvival -</mark> n	ot reported									
-	-	-	-	-	-	-	-	-	-	-	-	IMPORTA NT
nausea	- not reporte	ed										
-	-	-	-	-	-	-	-	-	-	-	-	CRITICAL
neutro	paenic sepsi	s										
1	randomise d trials	seriou S ^a	not serious	serious ^b	very serious ^c	none	2/23 (8.7%)	1/23 (4.3%)	RR 2.00 (0.19 to 20.55)	43 more per 1,000 (from 35 fewer to 850 more)	VERY LOW	CRITICAL
neutro	paenia											
1	randomise d trials	seriou s ^a	not serious	serious ^b	very serious ^c	none	5/23 (21.7%)	3/23 (13.0%)	RR 1.67 (0.45 to 6.17)	87 more per 1,000 (from 72 fewer to 674 more)	VERY LOW	CRITICAL
diarrho	ea - not repo	orted										
-	-	-	-	-	-	-	-	-	-	-	-	CRITICAL

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Quality	assessmen	t					Nº of pati	ents	Effect			
Nº of studie s	Study design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other consideratio ns	docetax el	docetac el plus S-1	Relativ e (95% Cl)	Absolu te (95% Cl)	Quality	Importanc e
treatme	ent related m	ortality -	not reported									
-	-	-	-	-	-	-	-	-	-	-	-	IMPORTA NT

CI: Confidence interval; HR: Hazard Ratio; OR: Odds ratio; RR: Risk ratio a. Unclear risk of bias due to poor study reporting b. Unclear definitions of morbidity outcomes c. 95% CI of the effect includes both default MID thresholds

Table 51: Clinical evidence profile for FOLFIRI + sunitinib versus placebo

Quality	assessment	-					Nº of pat	ients	Effect			
№ of studie s	Study design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecisi on	Other consideratio ns	FOLFIR I + sunitini b	placeb o	Relativ e (95% Cl)	Absolut e (95% Cl)	Quality	Importanc e
overall	survival											
1	randomise d trials	seriou s ^a	not serious	not serious	serious ^b	none	-/45	-/46	HR 0.82 (0.50 to 1.34)	-	LOW	CRITICAL
progres	sion free su	rvival										
1	randomise d trials	seriou s ^a	not serious	not serious	serious ^b	none	-/45	-/46	HR 1.11 (0.70 to 1.74)	-	LOW	IMPORTA NT
nausea												
1	randomise d trials	seriou s ^a	not serious	serious c	very serious ^d	none	3/45 (6.7%)	3/46 (6.5%)	RR 1.02 (0.22	1 more per 1,000 (from 51	VERY LOW	CRITICAL

Quality	assessment						Nº of pat	ients	Effect			
№ of studie s	Study design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecisi on	Other consideratio ns	FOLFIR I + sunitini b	placeb o	Relativ e (95% CI)	Absolut e (95% Cl)	Quality	Importanc e
									to 4.80)	fewer to 248 more)		
neutrop	baenic sepsis	s - not re	ported									
-	-	-	-	-	-	-	-	-	-	-	-	CRITICAL
neutrop	baenia											
1	randomise d trials	seriou s ^a	not serious	serious ^c	not serious	none	25/45 (55.6%)	9/46 (19.6%)	RR 2.84 (1.49 to 5.39)	360 more per 1,000 (from 96 more to 859 more)	LOW	CRITICAL
diarrho	ea											
1	randomise d trials	seriou s ^a	not serious	serious ^c	serious ^e	none	1/45 (2.2%)	6/46 (13.0%)	RR 0.17 (0.02 to 1.36)	108 fewer per 1,000 (from 47 more to 128 fewer)	VERY LOW	CRITICAL
treatme	ent related m	ortality -	not reported									
-	-	-	- Ratio: RR: Risk ra	-	-	-	-	-	-	-	-	IMPORTA NT

CI: Confidence interval; HR: Hazard Ratio; RR: Risk ratio; OR: Odds ratio

a. Unclear risk of bias due to poor reporting of methods b. 95% CI of the effect includes both no effect and clinically important benefit c. Unclear definitions of morbidity outcomes d. 95% CI of the effect includes both default MID thresholds e. 95% CI of the effect includes one default MID threshold

Quality	y assessmer	nt					Nº of pati	ients	Effect			
Nº of studi es	Study design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecisi on	Other considerati ons	irinotec an	irinotecan + 5'FU/leucovo rin (mFOLFIRI)	Relati ve (95% CI)	Absolu te (95% CI)	Quality	Importan ce
overal	l survival											
1	randomis ed trials	seriou s ^a	not serious	not serious	serious ^b	none	-/29	-/30	HR 1.04 (0.62 to 1.75)	-	LOW	CRITICAL
progre	ssion free s	urvival										
1	randomis ed trials	seriou s ^a	not serious	not serious	serious ^b	none	-/29	-/30	HR 1.13 (0.68 to 1.89)	-	LOW	IMPORTA NT
nause	a - not repor	ted										
-	-	-	-	-	-	-	-	-	-	-	-	CRITICAL
neutro	paenic seps	is - not r	eported									
-	-	-	-	-	-	-	-	-	-	-	-	CRITICAL
neutro	paenia											
1	randomis ed trials	seriou S ^a	not serious	not serious	very serious ^c	none	8/29 (27.6%)	11/30 (36.7%)	RR 0.75 (0.35 to 1.60)	92 fewer per 1,000 (from 220 more to 238 fewer)	VERY LOW	CRITICAL
diarrh	oea											
1	randomis ed trials	seriou s ^a	not serious	not serious	very serious °	none	1/29 (3.4%)	2/30 (6.7%)	RR 0.52 (0.05	32 fewer per	VERY LOW	CRITICAL

Quality	y assessmen	it					Nº of pati	ients	Effect			
Nº of studi es	Study design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecisi on	Other considerati ons	irinotec an	irinotecan + 5'FU/leucovo rin (mFOLFIRI)	Relati ve (95% CI)	Absolu te (95% Cl)	Quality	Importan ce
									to 5.40)	1,000 (from 63 fewer to 293 more)		
treatm	ent related n	nortality										
1	randomis ed trials	seriou s ^a	not serious	not serious	very serious °	none	1/29 (3.4%)	0/30 (0.0%)	RR 3.10 (0.13 to 73.14)	0 fewer per 1,000 (from 0 fewer to 0 fewer)	VERY LOW	IMPORTA NT

CI: Confidence interval; HR: Hazard Ratio; OR: Odds ratio; RR: Risk ratio

a. no blinding b. 95% CI of the effect includes both no effect and clinically important benefit and harm c. 95% CI of the effect includes both default MID thresholds

Table 53: Clinical evidence profile for irinotecan + cisplatin versus irinotecan

Quality	v assessmen	t					Nº of patie	ents	Effect			
№ of studi es	Study design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other considerati ons	irinotec an + cisplatin	irinotec an	Relati ve (95% Cl)	Absolu te (95% Cl)	Quality	Importanc e
overall	survival											
2	randomise d trials	seriou s ^a	not serious	not serious	not serious	none	-/148	-/150	HR 0.91 (0.71 to 1.16)	-	MODERA TE	CRITICAL
progre	rogression free survival											

Quality	assessment	t					Nº of pati	ents	Effect			
Nº of studi es	Study design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other considerati ons	irinotec an + cisplatin	irinotec an	Relati ve (95% Cl)	Absolu te (95% CI)	Quality	Importanc e
2	randomise d trials	seriou s ^a	not serious	not serious	not serious	none	-/148	-/150	HR 0.77 (0.60 to 0.99)	-	MODERA TE	IMPORTA NT
nausea	l i											
2	randomise d trials	seriou s ^a	not serious	not serious	very serious ^b	none	7/148 (4.7%)	8/150 (5.3%)	RR 0.89 (0.33 to 2.38)	6 fewer per 1,000 (from 36 fewer to 74 more)	VERY LOW	CRITICAL
neutro	paenic sepsi	s										
1	randomise d trials	seriou s ^a	not serious	not serious	very serious ^b	none	0/64 (0.0%)	3/66 (4.5%)	RR 0.15 (0.01 to 2.80)	39 fewer per 1,000 (from 45 fewer to 82 more)	VERY LOW	CRITICAL
neutro	paenia											
2	randomise d trials	seriou s ^a	not serious	not serious	serious ^c	none	60/148 (40.5%)	52/150 (34.7%)	RR 1.17 (0.87 to 1.57)	59 more per 1,000 (from 45 fewer to	LOW	CRITICAL

Quality	/ assessmen	t					Nº of patie	ents	Effect			
№ of studi es	Study design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other considerati ons	irinotec an + cisplatin	irinotec an	Relati ve (95% CI)	Absolu te (95% Cl)	Quality	Importanc e
										198 more)		
diarrho	bea											
2	randomise d trials	seriou s ^a	not serious	not serious	serious ^c	none	1/148 (0.7%)	7/150 (4.7%)	RR 0.20 (0.04 to 1.16)	37 fewer per 1,000 (from 7 more to 45 fewer)	LOW	CRITICAL
treatm	ent related m	ortality -	not reported									
-	-	-	-	-	-	-	-	-	-	-	-	IMPORTA NT

CI: Confidence interval; HR: Hazard Ratio; RR: Risk ratio; OR: Odds ratio

a. high risk due to no (or unclear) blinding

b. 95% CI of the effect includes both default MID thresholds

c. 95% CI of the effect includes one default MID threshold

Table 54: Clinical evidence profile for irinotecan versus BSC

Quality	assessmen	t					Nº of patie	ents	Effect			
№ of studie s	Study design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecisio n	Other consideratio ns	irinoteca n	BS C	Relativ e (95% Cl)	Absolu te (95% Cl)	Quality	Importanc e
overall	survival											
1	randomis ed trials	seriou s ^a	not serious	not serious	not serious	none	-/21	-/19	HR 0.48 (0.25 to 0.92)	-	MODERA TE	CRITICAL

Quality	assessmen	ıt					Nº of patie	ents	Effect			
№ of studie s	Study design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecisio n	Other consideratio ns	irinoteca n	BS C	Relativ e (95% CI)	Absolu te (95% Cl)	Quality	Importanc e
-	-	-	-	-	-	-	-	-	-	-	-	IMPORTA NT
nausea	- not report	ted										
-	-	-	-	-	-	-	-	-	-	-	-	CRITICAL
neutrop	aenic seps	is - not r	eported									
-	-	-	-	-	-	-	-	-	-	-	-	CRITICAL
neutrop	oaenia - not	reported	i									
-	-	-	-	-	-	-	-	-	-	-	-	CRITICAL
diarrho	ea - not rep	orted										
-	-	-	-	-	-	-	-	-	-	-	-	CRITICAL
treatme	nt related n	nortality	- not reported									
-	-	-	-	-	-	-	-	-	-	-	-	IMPORTA NT

CI: Confidence interval; HR: Hazard Ratio; OR: Odds ratio

a. No blinding

Table 55: Clinical evidence profile for olaparib+paclitaxel versus paclitaxel

Quality	/ assessmer	nt					Nº of patients		Effect			
Nº of studi es	Study design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecisi on	Other considerati ons	olaparib+paclit axel	paclita xel	Relati ve (95% Cl)	Absol ute (95% Cl)	Quality	Importan ce
overall	survival											
2	randomis ed trials	not seriou s	not serious	not serious	not serious	none	-/324	-/324	HR 0.74 (0.60 to 0.90)	-	HIGH	CRITICAL
progre	ogression free survival											

Quality	y assessmei	nt					№ of patients		Effect			
Nº of studi es	Study design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecisi on	Other considerati ons	olaparib+paclit axel	paclita xel	Relati ve (95% CI)	Absol ute (95% CI)	Quality	Importan ce
1	randomis ed trials	not seriou s	not serious	not serious	serious ^a	none	-/262	-/263	HR 0.84 (0.67 to 1.05)	-	MODERA TE	IMPORTA NT
nausea	a - not repor	ted										
-	-	-	-	-	-	-	-	-	-	-	-	CRITICAL
neutro	paenic seps	is						-				
1	randomis ed trials	not seriou s	not serious	not serious	very serious ^b	none	1/61 (1.6%)	0/62 (0.0%)	RR 3.05 (0.13 to 73.40)	0 fewer per 1,000 (from 0 fewer to 0 fewer)	LOW	CRITICAL
neutro	paenia											
2	randomis ed trials	not seriou s	not serious	not serious	serious ^c	none	114/323 (35.3%)	84/325 (25.8%)	RR 1.37 (1.08 to 1.72)	96 more per 1,000 (from 21 more to 186 more)	MODERA TE	CRITICAL
diarrho	bea											
1	randomis ed trials	not seriou s	not serious	not serious	very serious ^b	none	2/61 (3.3%)	6/62 (9.7%)	RR 0.34 (0.07 to 1.61)	64 fewer per 1,000 (from 59	LOW	CRITICAL

Quality	y assessmei	nt					№ of patients		Effect			
№ of studi es	Study design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecisi on	Other considerati ons	olaparib+paclit axel	paclita xel	Relati ve (95% CI)	Absol ute (95% CI)	Quality	Importan ce
										more to 90 fewer)		
treatment related mortality - not reported												
-	-	-	-	-	-	-	-	-	-	-	-	IMPORTA NT

CI: Confidence interval; HR: Hazard Ratio; RR: Risk ratio; OR: Odds ratio a. 95% CI of the effect includes possibility of no effect and clinically important effect b. 95% CI of the effect includes both default MID thresholds

c. 95% CI of the effect includes one default MID threshold

Table 56: Clinical evidence profile for S-1+ irinotecan versus irinotecan

Quality	assessment	t	-				Nº of pati	ents	Effect			
Nº of studi es	Study design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other considerati ons	S-1 + irinotec an	irinotec an	Relati ve (95% Cl)	Absolu te (95% Cl)	Quality	Importanc e
overall	survival											
1	randomise d trials	seriou s ^a	not serious	not serious	not serious	none	-/153	-/151	HR 0.99 (0.78 to 1.25)	-	MODERA TE	CRITICAL
progres	ssion free su	irvival										
1	randomise d trials	seriou s ^a	not serious	not serious	not serious	none	-/153	-/151	HR 0.85 (0.67 to 1.07)	-	MODERA TE	IMPORTA NT
nausea												

Quality	assessment	t					Nº of pati	ents	Effect			
Nº of studi es	Study design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other considerati ons	S-1 + irinotec an	irinotec an	Relati ve (95% CI)	Absolu te (95% CI)	Quality	Importanc e
1	randomise d trials	seriou s ^a	not serious	not serious	very serious ^b	none	7/153 (4.6%)	12/151 (7.9%)	RR 0.58 (0.23 to 1.42)	33 fewer per 1,000 (from 33 more to 61 fewer)	VERY LOW	CRITICAL
neutro	paenic sepsi	s										
1	randomise d trials	seriou s ^a	not serious	not serious	not serious	none	12/153 (7.8%)	1/151 (0.7%)	RR 11.84 (1.56 to 89.96)	72 more per 1,000 (from 4 more to 589 more)	MODERA TE	CRITICAL
neutro	paenia											
1	randomise d trials	seriou s ^a	not serious	not serious	serious ^c	none	57/153 (37.3%)	39/151 (25.8%)	RR 1.44 (1.03 to 2.03)	114 more per 1,000 (from 8 more to 266 more)	LOW	CRITICAL
diarrho	bea											
1	randomise d trials	seriou s ^a	not serious	not serious	very serious ^b	none	7/153 (4.6%)	10/151 (6.6%)	RR 0.69 (0.27	21 fewer per 1,000	VERY LOW	CRITICAL

Quality	assessment	t					Nº of pati	ents	Effect		
№ of studi es	Study design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other considerati ons	S-1 + irinotec an	irinotec an	Relati ve (95% Cl)	Absolu te (95% Cl)	Quality
									to 1.77)	(from 48 fewer to 51 more)	
treatme	ent related m	ortality									
1	randomise d trials	seriou S ^a	not serious	not serious	very serious ^b	none	0/153 (0.0%)	2/151 (1.3%)	RR 0.20 (0.01 to 4.08)	11 fewer per 1,000 (from 13 fewer to 41 more)	VERY LOW

Importanc е

IMPORTA NT

CI:

a. No blinding b. 95% CI of the effect includes both default MID thresholds c. 95% CI of the effect includes one default MID threshold

Table 57: Clinical evidence profile for paclitaxel versus irinotecan

Quality	assessment	t					Nº of pati	ients	Effect			
Nº of studi es	Study design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other considerati ons	paclitax el	irinotec an	Relati ve (95% Cl)	Absolu te (95% Cl)	Quality	Importanc e
Overal	Overall survival											
1	randomise d trials	seriou s ^a	not serious	not serious	not serious	none	-/111	-/112	HR 1.13 (0.86 to 1.49)	-	MODERA TE	CRITICAL

Quality assessment Nº of patients Effect Relati Absolu Nº of Risk Other ve te (95% (95% studi Study of Inconsiste Indirectne Imprecisi considerati paclitax irinotec Importanc el ĊI) CI) Quality es design bias ncy SS on ons an е **Progression free survival** 1 -/111 -/112 HR MODERA IMPORTA randomise seriou not serious not serious not none d trials s a serious 1.14 ΤE NT (0.88 to 1.48) Nausea (assessed with: grade 3 or more) 1 randomise seriou not serious not serious serious b 2/111 5/112 RR 27 LOW CRITICAL none d trials s a (1.8%) (4.5%) 0.40 fewer (0.80 per to 1,000 (from 9 2.04) fewer to 46 more) Neutropaenic sepsis (assessed with: grade 3 or more) 1 randomise seriou not serious not serious serious b 3/111 10/112 RR 63 LOW CRITICAL none 0.30 d trials s a (2.7%) (8.9%) fewer (0.09 per 1,000 to (from 6 1.07) more to 81 fewer) Neutropaenia (assessed with: grade 3 or more) 1 RR 104 LOW CRITICAL randomise seriou not serious not serious serious b none 31/111 43/112 d trials s a (27.9%) (38.4%) 0.73 fewer (0.50 per . 1,000 to 1.06) (from 23 more to

Quality	assessmen	t					Nº of pat	ients	Effect			
Nº of studi es	Study design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other considerati ons	paclitax el	irinotec an	Relati ve (95% CI)	Absolu te (95% Cl)	Quality	Importanc e
										192 fewer)		
Diarrho	oea (assesse	d with: g	rade 3 or mor	e)								
1	randomise d trials	seriou s ^a	not serious	not serious	serious ^b	none	1/111 (0.9%)	1/112 (0.9%)	RR 1.01 (0.06 to 15.93)	0 fewer per 1,000 (from 8 fewer to 133 more)	LOW	CRITICAL
Treatm	ent related n	nortality										
1	randomise d trials	seriou s ^a	not serious	not serious	very serious ^c	none	0/111 (0.0%)	2/112 (1.8%)	RR 0.20 (0.01 to 4.16)	14 fewer per 1,000 (from 18 fewer to 56 more)	VERY LOW	IMPORTA NT

CI: Confidence interval; HR: Hazard Ratio; RR: Risk ratio

a. High risk due to no blinding, moderate risk due to allocation concealment b. 95% CI of the effect includes one default MID threshold

c. 95% CI of the effect includes both default MID thresholds

G.17 Luminal obstruction

What is the optimal management of luminal obstruction for adults with oesophago-gastric cancer not amenable to treatment with curative intent?

			_		-							
Quality	assessmen	t					No of pa	atients	Effect			
No of studie s	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecisi on	Other considerati ons	SEMS	Plasti c tube	Relative (95% CI)	Absol ute	Quality	Importance
Dyspha	igia improve	ment (Be	etter indicated	by lower val	ues)							
2	randomis ed trials	no seriou s risk of bias ¹	serious ²	no serious indirectne ss	no serious imprecisi on	none	141	90	-	MD 0.3 lower (0.69 lower to 0.1 higher)	MODERAT E	CRITICAL
Persist	ent or recuri	rent dysp	hagia									
7	randomis ed trials	seriou s ³	serious ²	no serious indirectne ss	serious ⁴	none	64/241 (26.6%)	95/19 2 (49.5 %)	RR 0.60 (0.39 to 0.91)	198 fewer per 1000 (from 45 fewer to 302 fewer)	VERY LOW	CRITICAL
Proced	ure mortality	y										
7	randomis ed trials	seriou s ³	no serious inconsisten cy	no serious indirectne ss	serious ⁴	none	9/241 (3.7%)	16/19 2 (8.3%)	RR 0.39 (0.17 to 0.88)	51 fewer per 1000 (from 10	LOW	NOT IMPORTAN T

Table 58: Clinical evidence summary. SEMS versus plastic tubes

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Quality	assessmen	t					No of pa	atients	Effect			
No of studie s	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecisi on	Other considerati ons	SEMS	Plasti c tube	Relative (95% CI)	Absol ute	Quality	Importance
										fewer to 69 fewer)		
30-day	mortality											
4	randomis ed trials	no seriou s risk of bias ⁵	no serious inconsisten cy	no serious indirectne ss	serious ⁴	none	33/177 (18.6%)	34/12 7 (26.8 %)	RR 0.74 (0.48 to 1.14)	70 fewer per 1000 (from 139 fewer to 37 more)	MODERAT E	NOT IMPORTAN T
Proced	ure-related i	morbidity	- Perforation									
7	randomis ed trials	seriou s ³	no serious inconsisten cy	no serious indirectne ss	no serious imprecisi on	none	3/241 (1.2%)	14/19 2 (7.3%)	RR 0.24 (0.08 to 0.71)	55 fewer per 1000 (from 21 fewer to 67 fewer)	MODERAT E	CRITICAL
Fistula												
6	randomis ed trials	seriou s ³	no serious inconsisten cy	no serious indirectne ss	very serious ⁶	none	2/137 (1.5%)	3/140 (2.1%)	RR 0.76 (0.17 to 3.28)	5 fewer per 1000 (from 18 fewer	VERY LOW	CRITICAL

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Quality	assessmen	t					No of pa	atients	Effect			
No of studie s	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecisi on	Other considerati ons	SEMS	Plasti c tube	Relative (95% CI)	Absol ute	Quality	Importance
										to 49 more)		
Proced	ure-related i	morbidity	/ - Haemorrha	ge								
7	randomis ed trials	seriou s ³	no serious inconsisten cy	no serious indirectne ss	very serious ⁶	none	28/241 (11.6%)	22/19 2 (11.5 %)	RR 0.83 (0.5 to 1.38)	19 fewer per 1000 (from 57 fewer to 44 more)	VERY LOW	CRITICAL
Chest p	bain											
4	randomis ed trials	seriou s ⁵	no serious inconsisten Cy	no serious indirectne ss	very serious ⁶	none	45/186 (24.2%)	33/14 0 (23.6 %)	RR 1.11 (0.75 to 1.63)	26 more per 1000 (from 59 fewer to 149 more)	VERY LOW	IMPORTAN T
Proced	ure-related i	morbidity	/ - Sepsis									
2	randomis ed trials	seriou s ⁵	no serious inconsisten cy	no serious indirectne ss	very serious ⁶	none	0/41 (0%)	2/41 (4.9%)	RR 0.20 (0.01 to 3.93)	39 fewer per 1000 (from 48 fewer to 143 more)	VERY LOW	CRITICAL

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Quality	uality assessment							atients	Effect			
No of studie s	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecisi on	Other considerati ons	SEMS	Plasti c tube	Relative (95% CI)	Absol ute	Quality	Importance
Reflux												
3	randomis ed trials	seriou s ⁵	no serious inconsisten cy	no serious indirectne ss	very serious ⁶	none	7/63 (11.1%)	5/63 (7.9%)	RR 1.46 (0.43 to 4.92)	32 more per 1000 (from 44 fewer to 218 more)	VERY LOW	IMPORTAN T

RR=relative risk; CI=confidence interval; SEMS=self-expanding metallic stent

¹ Randomisation with appropriate allocation concealment and blinding of participants and personnels

² 12 > 50%

<u>a_cover</u>ere 1998, Sanyika 1999 -2 studies with unclear randomisation and Knyrim 1993, Siersema 1998, Shenfine 2009 - studies with unclear blinding and 3 studies with unclear blinding unclear blinding

⁴ 95%Cl crossed one boundary of default MID
 ⁵ Only one study wasSiersema 1998 conducted in unclear randomisation
 ⁶ 95%Cl crossed 2 boundaries of 95% Cl

Table 59: Clinical evidence summary. SEMS versus laser

Quality	uality assessment							itients	Effect			
No of studie s	Design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other considerati ons	SEMS	Laser	Relativ e (95% Cl)	Absol ute	Quality	Importance
Persist	ent or recuri	rent dysp	hagia									
2	randomis ed trials	seriou s ¹	serious ²	no serious indirectne ss	very serious ³	none	18/73 (24.7%)	16/52 (30.8 %)	RR 0.74 (0.38 to 1.43)	80 fewer per 1000	VERY LOW	CRITICAL

Quality	assessmen	t			-		No of pa	atients	Effect	-		
No of studie s	Design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other considerati ons	SEMS	Laser	Relativ e (95% CI)	Absol ute	Quality	Importanc
										(from 191 fewer to 132 more)		
Need o	f interventio	n for rec	urrent dyspha	gia								
2	randomis ed trials	seriou s ¹	serious ²	no serious indirectne ss	very serious ³	none	25/73 (34.2%)	31/52 (59.6 %)	RR 0.54 (0.23 to 1.26)	274 fewer per 1000 (from 459 fewer to 155 more)	VERY LOW	IMPORTAN T
Proced	ure-related i	morbidity	- Perforation									
2	randomis ed trials	seriou s ¹	no serious inconsisten Cy	no serious indirectne ss	very serious ³	none	0/73 (0%)	3/52 (5.8%)	RR 0.19 (0.02 to 1.64)	47 fewer per 1000 (from 57 fewer to 37 more)	VERY LOW	CRITICAL
Proced	ure-related i	morbidity	- Fistula									
2	randomis ed trials	seriou s ¹	no serious inconsisten cy	no serious indirectne ss	very serious ³	none	0/73 (0%)	4/52 (7.7%)	RR 0.15 (0.02 to 1.35)	65 fewer per 1000 (from	VERY LOW	CRITICAL

Quality	assessmen	t					No of pa	atients	Effect			
No of studie s	Design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other considerati ons	SEMS	Laser	Relativ e (95% Cl)	Absol ute	Quality	Importance
										75 fewer to 27 more)		
Proced	ure-related i	norbidity	- Haemorrhag	le								
2	randomis ed trials	seriou s ¹	no serious inconsisten cy	no serious indirectne ss	very serious ³	none	4/73 (5.5%)	0/52	RR 3.91 (0.53 to 28.66)	-	VERY LOW	CRITICAL
Proced	ure-related r	norbidity	- Sepsis									
2	randomis ed trials	seriou s ¹	no serious inconsisten cy	no serious indirectne ss	very serious ³	none	4/73 (5.5%)	1/52 (1.9%)	RR 2.2 (0.34 to 14.04)	23 more per 1000 (from 13 fewer to 251 more)	VERY LOW	CRITICAL
Proced	ure-related r	norbidity	- All adverse	effects								
2	randomis ed trials	seriou s ¹	no serious inconsisten cy	no serious indirectne ss	serious ⁴	none	28/73 (38.4%)	10/52 (19.2 %)	RR 1.8 (0.93 to 3.47)	154 more per 1000 (from 13 fewer to 475 more)	LOW	CRITICAL

Quality	assessmen	t					No of pa	atients	Effect			
No of studie s	Design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other considerati ons	SEMS	Laser	Relativ e (95% Cl)	Absol ute	Quality	Importance
2	randomis ed trials	seriou s ¹	no serious inconsisten cy	no serious indirectne ss	very serious ³	none	6/73 (8.2%)	2/52 (3.8%)	RR 2.1 (0.46 to 9.57)	42 more per 1000 (from 21 fewer to 330 more)	VERY LOW	NOT IMPORTAN T
Overall	survival (Be	etter indic	ated by highe	r values)								
2	randomis ed trials	seriou S ¹	no serious inconsisten cy	no serious indirectne ss	no serious imprecisio n	none	73	52	-	MD 7.89 higher (24.3 lower to 40.07 higher)	MODERAT E	IMPORTAN T

RR=relative risk; CI=confidence interval; SEMS=self-expanding metallic stent ¹ One study withAdam 1997 unclear allocation concealment ² I2 > 50%

³ 95%Cl crossed 2 boundaries of default MID ⁴ 95%Cl crossed one boundary of default MID

Quality	assessmen	+					No of pat	ients	Effect			
No of studie s	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecisi on	Other considerati ons	Covere d Ultrafie x SEMS	Cove red wallst ent SEM S	Relativ e (95% CI)	Absol ute	Quality	Importance
Dyspha	igia improve	ement (Be	etter indicated	by lower va	lues)						·	·
2	randomis ed trials	seriou s ¹	no serious inconsisten cy	no serious indirectne ss	no serious imprecisi on	none	65	55	-	MD 0.15 higher (0.04 lower to 0.33 higher)	MODERAT E	CRITICAL
Persiste	ent or recuri	rent dysp	hagia									
2	randomis ed trials	seriou s ¹	no serious inconsisten cy	no serious indirectne ss	very serious ²	none	13/65 (20%)	10/55 (18.2 %)	RR 1.2 (0.58 to 2.47)	36 more per 1000 (from 76 fewer to 267 more)	VERY LOW	CRITICAL
30-day	mortality											
2	randomis ed trials	seriou s ¹	no serious inconsisten cy	no serious indirectne ss	very serious ²	none	11/65 (16.9%)	8/55 (14.5 %)	RR 1.15 (0.5 to 2.64)	22 more per 1000 (from 73 fewer to 239 more)	VERY LOW	NOT IMPORTAN T

Table 60: Clinical evidence profile. Covered ultraflex SEMS versus covered wallstent SEMS

Quality	assessmen	t					No of pat	ients	Effect			
No of studie s	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecisi on	Other considerati ons	Covere d Ultrafle x SEMS	Cove red wallst ent SEM S	Relativ e (95% CI)	Absol ute	Quality	Importance
All adve	erse effects											
2	randomis ed trials	seriou s ¹	no serious inconsisten cy	no serious indirectne ss	serious ³	none	28/65 (43.1%)	31/55 (56.4 %)	RR 0.82 (0.59 to 1.14)	101 fewer per 1000 (from 231 fewer to 79 more)	LOW	CRITICAL
Advers	e effects - P	erforatio	n						-	-		
2	randomis ed trials	seriou s ¹	no serious inconsisten cy	no serious indirectne ss	very serious ²	none	2/65 (3.1%)	1/55 (1.8%)	RR 1.28 (0.24 to 6.92)	5 more per 1000 (from 14 fewer to 108 more)	VERY LOW	CRITICAL
Advers	e effects - H	aemorrha	age									
2	randomis ed trials	seriou s ¹	no serious inconsisten cy	no serious indirectne ss	very serious ²	none	6/65 (9.2%)	4/55 (7.3%)	RR 1.37 (0.41 to 4.5)	27 more per 1000 (from 43 fewer to 255 more)	VERY LOW	CRITICAL

Quality	assessmen	t					No of pat	ients	Effect			
No of studie s	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecisi on	Other considerati ons	Covere d Ultrafle x SEMS	Cove red wallst ent SEM S	Relativ e (95% CI)	Absol ute	Quality	Importance
Advers	e effects - R	eflux										
2	randomis ed trials	seriou s ¹	no serious inconsisten cy	no serious indirectne ss	very serious ²	none	3/65 (4.6%)	4/55 (7.3%)	RR 0.63 (0.14 to 2.83)	27 fewer per 1000 (from 63 fewer to 133 more)	VERY LOW	IMPORTAN T
Proced	ure related r	nortality										
2	randomis ed trials	seriou s ¹	no serious inconsisten cy	no serious indirectne ss	very serious ²	none	1/65 (1.5%)	1/55 (1.8%)	RR 0.97 (0.06 to 14.88)	1 fewer per 1000 (from 17 fewer to 252 more)	VERY LOW	NOT IMPORTAN T

95%CI = 95% confidence interval; SEMS=self-expanding metal stent; RR=relative risk; ¹ One-study with Subharwal 2003 - unclear randomisation ² 95%CI crossed 2 boundaries of default MID

³ 95%CI crossed one boundary of default MID

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Table 61: Clinical evidence profile. Irradiation SEMS versus conventional SEMS

Quality							No of roti		Effect			
No of studi es	assessmen Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecis ion	Other considerati ons	No of pati Irradiati on SEMS	Conventio nal SEMS	Relativ e (95% CI)	Absol ute	Quality	Importance
Dyspha	igia score (I	Better ir	ndicated by lo	wer values)								
1	randomis ed trials	no serio us risk of bias ¹	no serious inconsisten cy	no serious indirectne ss	serious ²	none	73	75	-	MD 0.26 higher (0.04 lower to 0.56 higher)	MODERAT E	CRITICAL
Overall	survival											
1	randomis ed trials	no serio us risk of bias	no serious inconsisten cy	no serious indirectne ss	serious ²	none	-	-	HR 0.59 (0.41 to 0.86)	-	MODERAT E	IMPORTAN T
Severe	chest pain											
1	randomis ed trials	no serio us risk of bias	no serious inconsisten cy	no serious indirectne ss	very serious ³	none	17/73 (23.3%)	15/75 (20%)	RR 1.16 (0.63 to 2.15)	32 more per 1000 (from 74 fewer to 230 more)	LOW	IMPORTAN T
Fistula	formation											

Quality	v assessmer	nt					No of pati	ents	Effect			
No of studi es	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecis ion	Other considerati ons	Irradiati on SEMS	Conventio nal SEMS	Relativ e (95% CI)	Absol ute	Quality	Importance
1	randomis ed trials	no serio us risk of bias	no serious inconsisten cy	no serious indirectne ss	very serious ³	none	6/73 (8.2%)	5/75 (6.7%)	RR 1.23 (0.39 to 3.86)	15 more per 1000 (from 41 fewer to 191 more)	LOW	CRITICAL
Haemo	rrhage											
1	randomis ed trials	no serio us risk of bias	no serious inconsisten cy	no serious indirectne ss	very serious ³	none	5/73 (6.8%)	5/75 (6.7%)	RR 1.03 (0.31 to 3.4)	2 more per 1000 (from 46 fewer to 160 more)	LOW	CRITICAL

95%CI = 95% confidence interval; SEMS=self-expanding metal stent[÷] RR=relative risk; HR=hazard ratio; ¹ appropriate randomisation with proper allocation concealment ² 95%CI crossed one boundary of default MID ³ 95%CI crossed 2 boundaries of default MID

Table 62: Clinical evidence profile. Polyflex SEMS versus ultraflex SEMS

Quality	/ assessment						No of p	atients	Effect			
No of studi es	Design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecisi on	Other consideratio ns	Polyfi ex SEMS	Ultrafi ex SEMS	Relativ e (95% CI)	Absolu te	Quali ty	Importance
Body v	veight at 4 we	eks in kg	(Better indicate	ed by lower va	alues)							
1	randomise d trials	serious 1	no serious inconsistency	no serious indirectnes s	serious ²	none	47	54	-	MD 1 lower (5.3 lower to 3.3 higher)	LOW	CRITICAL
Dyspha	agia score at	last follov	v-up (Better ind	icated by low	ver values)						-	
1	randomise d trials	serious 1	no serious inconsistency	no serious indirectnes s	serious ²	none	47	54	-	MD 0.2 higher (0.25 lower to 0.65 higher)	LOW	CRITICAL
Major o	complications	s (< 7 days	5)									
1	randomise d trials	serious 1	no serious inconsistency	no serious indirectnes s	very serious ³	none	4/47 (8.5%)	2/54 (3.7%)	RR 2.3 (0.44 to 11.99)	48 more per 1000 (from 21 fewer to 407 more)	VER Y LOW	CRITICAL
Major o	complications	s (> 7 days	s)									
1	randomise d trials	serious 1	no serious inconsistency	no serious indirectnes s	serious ²	none	20/47 (42.6 %)	17/54 (31.5%)	RR 1.35 (0.81	110 more per 1000	LOW	CRITICAL

Quality	/ assessment	t					No of p	atients	Effect			
No of studi es	Design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecisi on	Other consideratio ns	Polyfl ex SEMS	Ultrafi ex SEMS	Relativ e (95% CI)	Absolu te	Quali ty	Importance
									to 2.26)	(from 60 fewer to 397 more)		
Gastro	oesophageal	reflux (wi	thin a week)									
1	randomise d trials	serious 1	no serious inconsistency	no serious indirectnes s	very serious ³	none	0/47 (0%)	2/54 (3.7%)	RR 0.23 (0.01 to 4.66)	29 fewer per 1000 (from 37 fewer to 136 more)	VER Y LOW	IMPORTAN T
Surviv	al days (Bette	er indicate	d by lower valu	ies)								
1	randomise d trials	serious 1	no serious inconsistency	no serious indirectnes s	serious ²	none	47	54	-	MD 12 higher (4.56 to 19.44 higher)	LOW	IMPORTAN T
Days f	rom intervent	tion to rec	urrence of sym	ptoms (Bette	r indicated b	y lower values)						
1	randomise d trials	serious 1	no serious inconsistency	no serious indirectnes s	serious ²	none	47	54	-	MD 12.86 lower (38.49 lower to 12.77 higher)	LOW	CRITICAL

Quality	assessment						No of p	1	Effect			
No of studi es	Design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecisi on	Other consideratio ns	Polyfi ex SEMS	Ultrafi ex SEMS	Relativ e (95% CI)	Absolu te	Quali ty	Importance
1	randomise d trials	serious 1	no serious inconsistency	no serious indirectnes s	very serious ³	none	2/47 (4.3%)	2/54 (3.7%)	RR 1.15 (0.17 to 7.84)	6 more per 1000 (from 31 fewer to 253 more)	VER Y LOW	IMPORTAN T
Retros	ternal pain											
1	randomise d trials	serious 1	no serious inconsistency	no serious indirectnes s	serious ²	none	4/12 (33.3 %)	8/10 (80%)	RR 0.42 (0.18 to 0.98)	464 fewer per 1000 (from 16 fewer to 656 fewer)	LOW	CRITICAL

95%Cl = 95% confidence interval; SEMS=self-expanding metal stent [;] RR=relative risk; HR=hazard ratio; kg=kilograms ¹ appropriate randomisation with unclear allocation concealment ² 95%Cl crossed one boundary of default MID ³ 95%Cl crossed 2 boundaries of default MID

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Quality	assessmen	t					No of pat	ients	Effect			
No of studie s	Design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other considerati ons	Small- diamet er stent	Large- diamet er stent	Relativ e (95% Cl)	Absol ute	Quality	Importance
Dyspha	igia score <	2										
1	randomis ed trials	no serio us risk of bias	no serious inconsisten cy	no serious indirectne ss	no serious imprecisi on	none	47/50 (94%)	47/50 (94%)	RR 1 (0.91 to 1.1)	0 fewer per 1000 (from 85 fewer to 94 more)	HIGH	CRITICAL
immedi	ate adverse	effects	(chest/back p	ain requiring	hospitalisa	tion, persisten	t dysphagi	a, dyspno	ea, GI ha	emorrhag	ge, Arrhythmia	ı)
1	randomis ed trials	no serio us risk of bias	no serious inconsisten cy	no serious indirectne ss	very serious ¹	none	2/50 (4%)	0/50 (0%)	RR 5 (0.25 to 101.58)	-	LOW	CRITICAL
Recurre	ent dysphag	ia										
1	randomis ed trials	no serio us risk of bias	no serious inconsisten cy	no serious indirectne ss	very serious ¹	none	25/50 (50%)	21/50 (42%)	RR 1.19 (0.78 to 1.83)	80 more per 1000 (from 92 fewer to 349 more)	LOW	CRITICAL

Table 63: Clinical evidence profile. Small-diameter stent versus large-diameter stent

Quality	assessmen	t					No of pat	ients	Effect			
No of studie s	Design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other considerati ons	Small- diamet er stent	Large- diamet er stent	Relativ e (95% Cl)	Absol ute	Quality	Importance
1	randomis ed trials	no serio us risk of bias	no serious inconsisten cy	no serious indirectne ss	very serious ¹	none	3/50 (6%)	6/50 (12%)	RR 0.5 (0.13 to 1.89)	60 fewer per 1000 (from 104 fewer to 107 more)	LOW	CRITICAL
ER fistu	ula											
1	randomis ed trials	no serio us risk of bias	no serious inconsisten cy	no serious indirectne ss	very serious ¹	none	2/50 (4%)	5/50 (10%)	RR 0.4 (0.08 to 1.97)	60 fewer per 1000 (from 92 fewer to 97 more)	LOW	CRITICAL
New GE	ERD											
1	randomis ed trials	no serio us risk of bias	no serious inconsisten cy	no serious indirectne ss	very serious ¹	none	13/50 (26%)	12/50 (24%)	RR 1.08 (0.55 to 2.14)	19 more per 1000 (from 108 fewer to 274 more)	LOW	CRITICAL

Quality	assessmen	t					No of pat	tients	Effect			
No of studie s	Design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other considerati ons	Small- diamet er stent	Large- diamet er stent	Relativ e (95% Cl)	Absol ute	Quality	Importance
1	randomis ed trials	no serio us risk of bias	no serious inconsisten cy	no serious indirectne ss	very serious ¹	none	30/50 (60%)	29/50 (58%)	RR 1.03 (0.75 to 1.43)	17 more per 1000 (from 145 fewer to 249 more)	LOW	CRITICAL
Overall	survival at	6 month	S									
1	randomis ed trials	no serio us risk of bias	no serious inconsisten cy	no serious indirectne ss	serious ²	none	25/50 (50%)	15/50 (30%)	RR 1.67 (1 to 2.76)	201 more per 1000 (from 0 more to 528 more)	MODERAT E	IMPORTAN T

95%CI = 95% confidence interval; RR=relative risk; GERD=gastrooesophageal reflux disease; ER fistula = oesophageo-respiratory fistula ¹ 95% CI crossed 2 boundaries of default MID ² 95%CI crossed one boundary of default MID

Quality	assessment	1					No of pat	ients	Effect			
No of studie s	Design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecisi on	Other consideratio ns	Covere d Niti-S stent	Doubl e- layere d Niti- S stent	Relativ e (95% CI)	Absolu te	Qualit y	Importanc e
Dyspha	igia score (Be	etter indic	ated by lower v	/alues)								
1	randomise d trials	very serious 1	no serious inconsistenc y	no serious indirectnes s	very serious ²	none	19	18	-	MD 0.10 higher (0.27 lower to 0.47 higher)	VERY LOW	CRITICAL
Proced	ure-related co	omplicatio	ons									
1	randomise d trials	very serious 1	no serious inconsistenc y	no serious indirectnes s	no serious imprecisio n	none	11/19 (57.9%)	2/17 (11.8 %)	RR 4.92 (1.27 to 19.12)	461 more per 1000 (from 32 more to 1000 more)	LOW	CRITICAL

Table 64: Clinical evidence profile. Covered Niti-S SEMS versus double-layered Niti-S SEMS

95%CI = 95% confidence interval; SEMS=self-expanding metal stent; RR=relative risk; MD=mean difference

¹ Randomisation method was not reported in details ² 95%CI crossed 2 boundaries of default MID

Table 65: Clinical evidence profile. SEMS versus oesophageal bypass

Qual	lity asse	ssment					No of pat	ients	Effect			
No of stu die s	Desig n	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecisi on	Other considerat ions	SEMS	Oesophag eal bypass	Relati ve (95% CI)	Absolute	Quality	Importanc e
Dysp	phagia s	core (Be	tter indicated	by lower val	ues)							
1	rando mised trials	very seriou s ¹	no serious inconsisten cy	no serious indirectne ss	serious ²	none	20	20	-	MD 0.60 higher (0.15 to 1.05 higher)	VERY LOW	CRITICAL

95%CI = 95% confidence interval; SEMS=self-expanding metal stent; MD=mean difference;

¹ Randomisation was not reported in details

² 95%CI crossed one boundary of default MID

Table 66: Clinical evidence profile. SEMS versus External beam RT

Quality	assessment	:					No of p	patients	Effect			
No of studie s	Design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecisi on	Other consideratio ns	SEM S	External beam radiothera py	Relativ e (95% CI)	Absol ute	Qualit y	Importance
Overall	survival day	s (Better	indicated by h	igher values))							
1	randomise d trials	very seriou s ¹	no serious inconsistenc y	no serious indirectnes s	serious ²	none	32	32	-	MD 77.13 lower (116.7 1 to 37.55 lower)	VERY LOW	IMPORTANT

95%CI = 95% confidence interval; SEMS=self-expanding metal stent; MD=mean difference; RT=radiotherapy

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¹ Unclear randomisation and no blinding ² 95%Cl crossed one boundary of default MID

Table 67: Clinical evidence profile. SEMS versus SEMS plus External beam RT

Quality	/ assess	ment						No of pa	atients	Effect			
No of studi es	Desi gn	Ri sk of bi as	Inco cy	onsisten	Indirectn ess	Imprecisi on	Other considerat ions	SEMS	SEMS plus external beam RT	Relative (95% CI)	Absolute	Qualit y	Importance
Mean o	lysphag	ia free	e sur	vival (Bet	ter indicated	by higher v	alues)						
1	rando mise d trials	no serio risk o bias	of	no serious inconsis tency	no serious indirectne ss	serious ¹	none	37	42	-	MD 21.80 lower (43.63 lower to 0.03 higher)	MODE RATE	CRITICAL
Overal	l surviva	al 👘											
1	rando mise d trials	no seric risk o bias	of	no serious inconsis tency	no serious indirectnes s	serious ¹	none	35/37 (94.6%)	29/42 (69%)	HR 1.94 (1.18 to 3.18)	-	MODE RATE	IMPORTANT

95%CI = 95% confidence interval; SEMS=self-expanding metal stent; MD=mean difference; RT=radiotherapy; HR=hazard ratio ¹ 95%CI crossed one boundary of default MID

Table 68: Clinical evidence profile. SEMS versus Laser plus RT

N CONTRACTOR OF CONTRACTOR OFO		
Relativ Absol e ute (95% CI)		Importanc e
	• • • • • • •	

Quality	assessment						No of p	atients	Effect			
No of studie s	Design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecisi on	Other consideratio ns	SEM S	Laser plus Radiothera py	Relativ e (95% Cl)	Absol ute	Qualit y	Importanc e
1	randomise d trials	very serious 1	no serious inconsistenc y	no serious indirectnes s	serious ²	none	10	21	-	MD 0.08 higher (0.01 lower to 0.17 higher)	VERY LOW	CRITICAL
Recurre	ent dysphagi	a										
1	randomise d trials	very serious 1	no serious inconsistenc y	no serious indirectnes s	very serious ³	none	1/10 (10%)	9/21 (42.9%)	RR 0.23 (0.03 to 1.6)	330 fewer per 1000 (from 416 fewer to 257 more)	VERY LOW	CRITICAL

95%CI = 95% confidence interval; SEMS=self-expanding metal stent ; MD=mean difference; RT=radiotherapy; RR=relative risk; ¹ Unclear randomisation plus no blinding ² 95%CI crossed one boundary of default MID ³ 95%CI crossed 2 boundaries of default MID

Table 69: Clinical evidence profile. SEMS versus laser followed by SEMS

Quality	assessment						No of p	atients	Effect			
No of studie s	Design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecisi on	Other consideratio ns	SEM S	Laser follow ed by SEMS	Relativ e (95% CI)	Absolu te	Qualit y	Importanc e
Recurr	ent dysphagia	a										
1	randomise d trials	very serious 1	no serious inconsistenc y	no serious indirectnes S	very serious ²	none	1/10 (10%)	3/8 (37.5%)	RR 0.27 (0.03 to 2.1)	274 fewer per 1000 (from 364 fewer to 412 more)	VERY LOW	CRITICAL

95%CI = 95% confidence interval; SEMS=self-expanding metal stent[:] RR=relative risk ¹ Unclear randomisation and no blinding ² 95%CI crossed 2 boundaries of default MID

Table 70: Clinical evidence profile. SEMS plus brachytherapy versus brachytherapy alone

Quality	assessmer	nt					No of patients	i	Effect			
No of studi es	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecis ion	Other considerati ons	SEMS plus brachythera py	Brachyther apy	Relativ e (95% Cl)	Absol ute	Qualit y	Importan ce
Numbe	r of patients	s with dy	sphagia impro	ovement								
1	randomis ed trials	seriou s ¹	no serious inconsisten cy	no serious indirectne ss	serious ²	none	12/17 (70.6%)	7/18 (38.9%)	RR 1.82 (1.05 to 3.15)	319 more per 1000 (from 19	LOW	CRITICAL

Quality	assessmer	ıt					No of patients	;	Effect			
No of studi es	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecis ion	Other considerati ons	SEMS plus brachythera py	Brachyther apy	Relativ e (95% CI)	Absol ute	Qualit y	Importan ce
										more to 836 more)		
Proced	lure-related	morbidit	у									
1	randomis ed trials	seriou s ¹	no serious inconsisten cy	no serious indirectne ss	very serious ³	none	4/21 (19%)	0/20 (0%)	RR 8.59 (0.49 to 150)	-	VERY LOW	CRITICAL

95%CI = 95% confidence interval; SEMS=self-expanding metal stent RR=relative risk

Appropriate randomisation with no blinding
 95%Cl crossed one boundary of default MID
 95%Cl crossed 2 boundaries of default MID

Table 71: Clinical evidence profile. Dilatation alone versus dilatation plus laser

Quality	assessmen	t					No of pati	ients	Effect			
No of studie s	Design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other considerati ons	Dilatatio n	Dilatation plus laser	Relativ e (95% Cl)	Absol ute	Qualit y	Importance
Numbe	r of re-interv	ention (E	Setter indicated	d by lower va	alues)							
1	randomis ed trials	very seriou s ¹	no serious inconsisten cy	no serious indirectne ss	very serious ²	none	7	8	-	MD 0.5 higher (0.45 lower to 1.45 higher)	VERY LOW	CRITICAL

	assessmen	1					No of pati		Effect			
No of studie s	Design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other considerati ons	Dilatatio n	Dilatation plus laser	Relativ e (95% Cl)	Absol ute	Qualit y	Importance
Dyspha	igia score at	2 month	s (Better indic	ated by lowe	r values)							
1	randomis ed trials	very seriou s ¹	no serious inconsisten cy	no serious indirectne ss	very serious ²	none	7	8	-	MD 0.1 higher (0.1 lower to 0.3 higher)	VERY LOW	CRITICAL
Surviva	l rate at 30 i	months										
1	randomis ed trials	very seriou s ¹	no serious inconsisten cy	no serious indirectne ss	very serious ²	none	1/7 (14.3%)	2/8 (25%)	RR 0.57 (0.06 to 5.03)	108 fewer per 1000 (from 235 fewer to 1000 more)	VERY LOW	IMPORTAN T

95%CI = 95% confidence interval; SEMS=self-expanding metal stent RR=relative risk; MD=mean difference

¹ RCT with unclear randomisation and blinding ² 95%CI crossed 2 boundaries of MID

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Table 72: Clinical evidence profile. ILRT versus ILRT+5-FU

Quality	assessment						No of	patients	Effect			
No of studie s	Design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecisi on	Other consideratio ns	ILRT	ILRT+5F U	Relativ e (95% Cl)	Absolu te	Qualit y	Importance
Overall 1	survival at 2 randomise d trials	years serious 1	no serious inconsistenc y	no serious indirectnes s	serious ²	none	4/25 (16%)	6/25 (24%)	RR 0.67 (0.21 to 2.08)	79 fewer per 1000 (from 190 fewer to 259 more)	LOW	IMPORTANT
Comple 1	ete regression randomise d trials	n (on bari serious 1	um swallow ar no serious inconsistenc y	nd -ve biopsy no serious indirectnes s) serious ³	none	22/25 (88%)	25/25 (100%)	RR 0.88 (0.75 to 1.04)	120 fewer per 1000 (from 250 fewer to 40 more)	LOW	CRITICAL

95%CI = 95% confidence interval; SEMS=self-expanding metal stent[;] RR=relative risk; ILRT=intraluminal radiotherapy; 5FU=5-Fluouracil; ¹ unclear randomisation with appropriate concealment and unclear outcome of interest ² 95%CI crossed 2 boundaries of default MID ³ 95%CI crossed one default MID

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				-								
Quality	assessment						No of patient	s	Effect			
No of studie s	Design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecisi on	Other consideratio ns	Dilatation plus radiotherap y	Dilata tion alone	Relativ e (95% Cl)	Absol ute	Qualit y	Importanc e
Body w	eight at 6 mo	onths in k	g (Better indic	ated by lowe	r values)							
1	randomise d trials	very seriou s ¹	no serious inconsistenc y	no serious indirectnes s	no serious imprecisio n	none	30	9	-	MD 8.27 higher (3.81 to 12.73 higher)	LOW	CRITICAL
ECOG s	score of 2 or	more at 1	month (lower	, better)								
1	randomise d trials	very seriou s ¹	no serious inconsistenc y	no serious indirectnes s	no serious imprecisio n	none	15/47 (31.9%)	27/41 (65.9 %)	RR 0.48 (0.3 to 0.78)	342 fewer per 1000 (from 145 fewer to 461 fewer)	LOW	CRITICAL
Surviva	I months (Be	etter indic	ated by lower	values)								
1	randomise d trials	seriou s ¹	no serious inconsistenc y	no serious indirectnes s	very serious ²	none	4	10	-	MD 0.34 higher (1.93 lower to 2.61 higher)	VERY LOW	CRITICAL

Table 73: Clinical evidence profile. Dilatation plus radiotherapy versus dilatation alone

95%CI=95%confidence interval; ECOG=Eastern cooperative oncology group; RR=relative risk; MD=mean difference; kg=kilograms ¹ Unclear randomisation and blinding ² 95%CI crossed 2 boundaries of default MID

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No of patients Effect **Quality assessment** No of Design Risk Inconsiste Indirectne Imprecisi Other External Endos Relativ Absol of considerati studie ncy SS on beam recopic ute е dilatati (95% s bias ons irradiation Qualit ĊI) on Importance V Dysphagia grade 2 or more at 4 weeks 1 14/34 32/35 RR 503 LOW CRITICAL randomis verv no serious no serious no none (41.2%) (91.4% 0.45 ed trials seriou inconsistenc indirectnes serious fewer s¹ (0.3 to v s imprecisio per n 0.68) 1000 (from 293 fewer to 640 fewer) Overall survival at the end of study 1 seriou no serious no serious serious² HR -LOW IMPORTANT randomis none _ _ ed trials S^1 inconsistenc indirectnes 0.54 (0.28 to v s 1.03) **Oesophagitis within 4 weeks** 1 randomis very no serious no serious serious² none 20/34 9/35 RR 332 VERY CRITICAL (25.7% 2.29 LOW ed trials inconsistenc indirectnes (58.8%) seriou more S^1 (1.22 to per y s 1000 4.29) (from 57 more to 846 more) Acute chest pain (within 24 hours of dilatation) 1 randomis very no serious no serious no 0/34 35/35 RR 990 LOW IMPORTANT none ed trials inconsistenc indirectnes serious (0%) (100%) 0.01 (0 fewer seriou S¹ y s to 0.23) per

Table 74: Clinical evidence profile. External beam irradiation versus endoscopic dilatation

Quality	assessmen	t					No of patien	its	Effect			
No of studie s	Design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other considerati ons	External beam re- irradiation	Endos copic dilatati on	Relativ e (95% Cl)	Absol ute	Qualit y	Importance
					imprecisio n					1000 (from 770 fewer to 1000 fewer)		
Chest i	nfection wit	hin 4 wee	ks									
1	randomis ed trials	very seriou s ¹	no serious inconsistenc y	no serious indirectnes s	very serious ³	none	4/34 (11.8%)	7/35 (20%)	RR 0.59 (0.19 to 1.83)	82 fewer per 1000 (from 162 fewer to 166 more)	VERY LOW	CRITICAL
Hemete	mesis withi	n 4 weeks	5									
1	randomis ed trials	very seriou s ¹	no serious inconsistenc y	no serious indirectnes s	very serious ³	none	1/34 (2.9%)	0/35 (0%)	RR 3.09 (0.13 to 73.21)	-	VERY LOW	CRITICAL
recurre	nt chest infe	ection after	er 6-10 weeks									
1	randomis ed trials	very seriou s ¹	no serious inconsistenc y	no serious indirectnes s	very serious ³	none	8/34 (23.5%)	3/35 (8.6%)	RR 2.75 (0.79 to 9.49)	150 more per 1000 (from 18 fewer	VERY LOW	CRITICAL

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Quality	assessmen	ıt					No of patier	nts	Effect			
No of studie s	Design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other considerati ons	External beam re- irradiation	Endos copic dilatati on	Relativ e (95% CI)	Absol ute	Qualit y	Importance
										to 728 more)		
Trache	ooesophage	al fistula	after 6-10 wee	ks								
1	randomis ed trials	very seriou s ¹	no serious inconsistenc y	no serious indirectnes s	very serious ³	none	0/34 (0%)	6/35 (17.1%)	RR 0.08 (0 to 1.35)	158 fewer per 1000 (from 171 fewer to 60 more)	VERY LOW	CRITICAL
Tumou	r bleed after	[.] 6-10 wee	ks									
1	randomis ed trials	very seriou s ¹	no serious inconsistenc y	no serious indirectnes s	very serious ³	none	4/34 (11.8%)	5/35 (14.3%)	RR 0.82 (0.24 to 2.81)	26 fewer per 1000 (from 109 fewer to 259 more)	VERY LOW	CRITICAL

95%CI=95%confidence interval; RR=relative risk; MD=mean difference; ¹ Randomisation method was not reported in details ² 95%CI crossed one boundary of default MID ³ 95%CI crossed 2 boundaries of default MID

Table 75: Clinical evidence profile. 8Gy per fraction 2 times radiotherapy within 3 days versus 6 Gy per fraction 3 times radiotherapy within 5 days

Quality	assessment						No of pa	tients	Effect			
No of studie s	Design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecisi on	Other consideratio ns	8 Gy per fractio n	6 Gy per fracti on	Relativ e (95% Cl)	Absolu te	Qualit y	Importance
Trachee	ooesophagea	al fistula										
1	randomise d trials	very serious 1	no serious inconsistenc y	no serious indirectnes s	very serious ²	none	11/118 (9.3%)	12/10 4 (11.5 %)	RR 0.81 (0.37 to 1.75)	22 fewer per 1000 (from 73 fewer to 87 more)	VERY LOW	CRITICAL
Fibrous	strictures											
1	randomise d trials	very serious 1	no serious inconsistenc y	no serious indirectnes s	very serious ²	none	12/118 (10.2%)	13/10 4 (12.5 %)	RR 0.81 (0.39 to 1.7)	24 fewer per 1000 (from 76 fewer to 88 more)	VERY LOW	CRITICAL
Patients	s necessitatio	on additic	onal treatment									
1	randomise d trials	very serious	no serious inconsistenc y	no serious indirectnes s	serious ³	none	37/50 (74%)	45/50 (90%)	RR 0.82 (0.68 to 0.99)	162 fewer per 1000 (from 9 fewer	VERY LOW	IMPORTANT

Quality	assessment						No of pa	tients	Effect			
No of studie s	Design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecisi on	Other consideratio ns	8 Gy per fractio n	6 Gy per fracti on	Relativ e (95% Cl)	Absolu te	Qualit y	Importance
										to 288 fewer)		

95%CI=95%confidence interval; RR=relative risk; ¹ inappropriate randomisation with unclear allocation concealment and blinding

² 95%CI crossed two boundaries of default MID
 ³ 95%CI crossed one boundary of default MID

Table 76: Clinical evidence profile. 16 Gy/2 fractions weekly versus 18Gy/3 fractions weekly

Quality	assessmen	t					No of patier	nts	Effect			
No of studie s	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecisi on	Other considerati ons	16Gy/2fra ct weekly	18Gy/3fr act weekly	Relativ e (95% CI)	Absol ute	Qualit y	Importance
Overall	survival rat	e at 12 m	onths									
1	randomis ed trials	very seriou s ¹	no serious inconsisten cy	no serious indirectne ss	serious ²	none	14/60 (23.3%)	19/55 (34.5%)	RR 0.68 (0.38 to 1.21)	111 fewer per 1000 (from 214 fewer to 73 more)	VERY LOW	IMPORTAN T
Dyspha	agia free sur	vival rate	i.									
1	randomis ed trials	very seriou s ¹	no serious inconsisten cy	no serious indirectne ss	serious ²	none	15/60 (25%)	21/55 (38.2%)	RR 0.65 (0.38 to 1.14)	134 fewer per 1000 (from	VERY LOW	CRITICAL

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Quality	assessmen	ıt					No of patie	nts	Effect			
No of studie s	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecisi on	Other considerati ons	16Gy/2fra ct weekly	18Gy/3fr act weekly	Relativ e (95% CI)	Absol ute	Qualit y	Importance
										237 fewer to 53 more)		
Strictu	res											
1	randomis ed trials	very seriou s ¹	no serious inconsisten cy	no serious indirectne ss	serious ²	none	15/60 (25%)	23/55 (41.8%)	RR 0.6 (0.35 to 1.02)	167 fewer per 1000 (from 272 fewer to 8 more)	VERY LOW	CRITICAL
Persist	ent disease											
1	randomis ed trials	very seriou s ¹	no serious inconsisten cy	no serious indirectne ss	very serious ³	none	4/60 (6.7%)	4/55 (7.3%)	RR 0.92 (0.24 to 3.49)	6 fewer per 1000 (from 55 fewer to 181 more)	VERY LOW	CRITICAL
Fistula												
1	randomis ed trials	very seriou s ¹	no serious inconsisten cy	no serious indirectne ss	very serious ³	none	2/60 (3.3%)	6/55 (10.9%)	RR 0.31 (0.06 to 1.45)	75 fewer per 1000 (from 103	VERY LOW	CRITICAL

Quality	assessmen	t					No of patier	nts	Effect			
No of studie s	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecisi on	Other considerati ons	16Gy/2fra ct weekly	18Gy/3fr act weekly	Relativ e (95% CI)	Absol ute	Qualit y	Importance
										fewer to 49 more)		

95%CI=95%confidence interval; RR=relative risk; ¹ Inappropriate randomisation and no blinding ² 95%CI crossed one boundary of default MID ³ 95%CI crossed 2 boundaries of default MID

Table 77: Clinical evidence profile. Brachytherapy versus brachytherapy plus radiotherapy

Quality	/ assessmen	t					No of patie	ents	Effect			
No of studi es	Design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecisi on	Other consideratio ns	Brachyth erapy	Brachyth erapy plus radiother apy	Relati ve (95% CI)	Absolu te	Quali ty	Importan ce
Advers	se effects - S	tricture										
2	randomise d trials	very serious 1	serious2	no serious indirectnes s	very serious ³	none	9/138 (6.5%)	8/139 (5.8%)	RR 1.43 (0.18 to 11.34)	25 more per 1000 (from 47 fewer to 595 more)	VER Y LOW	CRITICAL

Quality	/ assessmen	t					No of patie	ents	Effect			
No of studi es	Design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecisi on	Other consideratio ns	Brachyth erapy	Brachyth erapy plus radiother apy	Relati ve (95% CI)	Absolu te	Quali ty	Importan ce
2	randomise d trials	very serious 1	no serious inconsistenc y	no serious indirectnes s	very serious ³	none	13/138 (9.4%)	10/139 (7.2%)	RR 1.09 (0.27 to 4.35)	6 more per 1000 (from 53 fewer to 241 more)	VER Y LOW	CRITICAL

95%CI=95%confidence interval; RR=relative risk;

¹ Both studies with Rosenblatt 2010 and Sur 2004 - no clear randomisation and no blinding

² 12> 50%
 ³ 95%CI crossed 2 boundaries of default MID

Table 78: Clinical evidence profile. Covered stent versus uncovered stent

	assessmen						No of pa		Effect			
No of studie s	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecis ion	Other considerati ons	Covere d stent	Uncover ed stent	Relativ e (95% Cl)	Absol ute	Quality	Importance
Clinical	success											
3	randomis ed trials	very seriou s ¹	no serious inconsisten cy	no serious indirectne ss	no serious imprecisi on	none	96/104 (92.3%)	95/103 (92.2%)	RR 1 (0.92 to 1.08)	0 fewer per 1000 (from 74 fewer	LOW	CRITICAL

Quality	assessmen	ıt					No of pa	tients	Effect			
No of studie s	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecis ion	Other considerati ons	Covere d stent	Uncover ed stent	Relativ e (95% CI)	Absol ute	Quality	Importance
										to 74 more)		
Clinica	l success - (GOO-tail	ored stent vs	Standard un	covered ste	nt						
1	randomis ed trials	very seriou s ²	no serious inconsisten cy	no serious indirectne ss	no serious imprecisi on	none	31/33 (93.9%)	30/32 (93.8%)	RR 1 (0.88 to 1.13)	0 fewer per 1000 (from 113 fewer to 122 more)	LOW	CRITICAL
Clinica	l success - (Covered	pyloric stent v	/s uncovered	d pyloric ste	ent						
2	randomis ed trials	seriou s ³	no serious inconsisten cy	no serious indirectne ss	no serious imprecisi on	none	65/71 (91.5%)	65/71 (91.5%)	RR 1 (0.9 to 1.11)	0 fewer per 1000 (from 92 fewer to 101 more)	MODERAT E	CRITICAL
Patenc	y at final fol	low-up										
1	randomis ed trials	seriou s ⁴	no serious inconsisten cy	no serious indirectne ss	very serious ⁵	none	14/31 (45.2%)	13/36 (36.1%)	RR 1.25 (0.7 to 2.24)	90 more per 1000 (from 108 fewer	VERY LOW	CRITICAL

Quality	assessmen	t					No of pa	tients	Effect			
No of studie s	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecis ion	Other considerati ons	Covere d stent	Uncover ed stent	Relativ e (95% Cl)	Absol ute	Quality	Importance
										to 448 more)		
Major c	omplication	l i										
3	randomis ed trials	very seriou s ¹	no serious inconsisten cy	no serious indirectne ss	no serious imprecisi on	none	14/104 (13.5%)	3/103 (2.9%)	RR 4.06 (1.32 to 12.44)	89 more per 1000 (from 9 more to 333 more)	LOW	CRITICAL
Major c	omplication	- GOO-t	ailored covere	ed stent vs S	tandard un	covered stent						
1	randomis ed trials	very seriou s ²	no serious inconsisten cy	no serious indirectne ss	no serious imprecisi on	none	11/33 (33.3%)	2/32 (6.3%)	RR 5.33 (1.28 to 22.2)	271 more per 1000 (from 17 more to 1000 more)	LOW	CRITICAL
	omplication	- Covere	ed pyloric ste	nt vs Uncove	ered pyloric	stent						
2	randomis ed trials	seriou s ³	no serious inconsisten cy	no serious indirectne ss	very serious ⁵	none	3/71 (4.2%)	1/71 (1.4%)	RR 2.33 (0.35 to 15.42)	19 more per 1000 (from 9 fewer	VERY LOW	CRITICAL

Quality	assessmen	t					No of pa	tients	Effect			
No of studie s	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecis ion	Other considerati ons	Covere d stent	Uncover ed stent	Relativ e (95% Cl)	Absol ute	Quality	Importance
										to 203 more)		
Reinter	vention rate)										
2	randomis ed trials	very seriou s ⁶	no serious inconsisten cy	no serious indirectne ss	no serious imprecisi on	none	9/75 (12%)	21/69 (30.4%)	RR 0.39 (0.19 to 0.79)	186 fewer per 1000 (from 64 fewer to 247 fewer)	LOW	IMPORTAN T
Reinter	vention rate	- WAVE	-covered SEM	S vs Uncove	ered SEMS							
1	randomis ed trials	seriou s ⁴	no serious inconsisten Cy	no serious indirectne ss		none	6/42 (14.3%)	14/37 (37.8%)	RR 0.38 (0.16 to 0.88)	235 fewer per 1000 (from 45 fewer to 318 fewer)	LOW	IMPORTAN T
Reinter	vention rate	- GOO-t	ailored stent v	/s uncovered	d stent							
1	randomis ed trials	very seriou s ⁸	no serious inconsisten cy	no serious indirectne ss	very serious ⁵	none	3/33 (9.1%)	7/32 (21.9%)	RR 0.42 (0.12 to 1.47)	127 fewer per 1000 (from 192 fewer	VERY LOW	IMPORTAN T

Quality	assessmen	ıt					No of pa	tients	Effect			
No of studie s	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecis ion	Other considerati ons	Covere d stent	Uncover ed stent	Relativ e (95% CI)	Absol ute	Quality	Importance
										to 103 more)		
Advers	e events											
1	randomis ed trials	seriou s ⁹	no serious inconsisten cy	no serious indirectne ss	very serious⁵	none	6/31 (19.4%)	10/31 (32.3%)	RR 0.6 (0.25 to 1.45)	129 fewer per 1000 (from 242 fewer to 145 more)	VERY LOW	CRITICAL
Overall	survival											
1	randomis ed trials	seriou s ⁴	no serious inconsisten cy	no serious indirectne ss	serious ⁷	none	-	-	HR 0.62 (0.34 to 1.14)	-	LOW	IMPORTAN T
Recurre	ent obstruct	tive symp	otoms									
1	randomis ed trials	seriou s ⁹	no serious inconsisten cy	no serious indirectne ss	serious ⁷	none	1/31 (3.2%)	9/31 (29%)	RR 0.11 (0.01 to 0.83)	258 fewer per 1000 (from 49 fewer to 287 fewer)	LOW	CRITICAL

Quality	assessmen	it			-		No of pa	tients	Effect			
No of studie s	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecis ion	Other considerati ons	Covere d stent	Uncover ed stent	Relativ e (95% Cl)	Absol ute	Quality	Importance
1	randomis ed trials	very seriou s ⁸	no serious inconsisten cy	no serious indirectne ss	serious ⁷	none	33	32	-	MD 19 higher (8.06 to 29.94 higher)	VERY LOW	IMPORTAN T
Gastric	outlet obst	ruction s	core (GOOS)	change (Bett	ter indicate	d by lower valu	ies)					
1	randomis ed trials	very seriou s ⁸	no serious inconsisten cy	no serious indirectne ss	serious ⁷	none	33	32	-	MD 0.1 higher (0.12 lower to 0.32 higher)	VERY LOW	CRITICAL

95%CI=95%confidence interval; RR=relative risk; MD=mean difference; GOO=gastric outlet obstruction; HR=hazard ratio ¹ <u>All 3 studies Shi 2014, Kim 2010, Maetani 2014-</u> unclear or inappropriate randomization and unclear blinding

² RCT with inappropriate randomisation and unclear blinding
 ³ One studyKim 2010, unclear randomisation and another study withMaetani 2014, unclear allocation concealment
 ⁴ One study with unclear allocation concealment and unclear blinding

⁵ 95%CI crossed 2 boundaries of default MID

⁶ one study with unclear randomization, one study with inappropriatre randomisation and unclear blinding

⁷ 95%CI crossed one boundary of MID

⁸ one study with inappropriate randomisation

⁹ One study with unclear randomisation and blinding

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Table 79: Clinical evidence profile. Stent versus gastroenterostomy

Quality	assessmer /	nt					No of pa	atients	Effect			
No of studi es	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecis ion	Other considerati ons	Stent	Gastroenterost omy	Relativ e (95% CI)	Absol ute	Qualit y	Importance
Mortali	ty											
1	randomis ed trials	very seriou s ¹	no serious inconsisten cy	very serious	no serious imprecisi on	none	0/9 (0%)	0/9 (0%)	No event in either arm	-	VERY LOW	NOT IMPORTAN T
Minor o	complication	าร										
2	randomis ed trials	seriou S ²	no serious inconsisten cy	very serious3	very serious ⁴	none	5/30 (16.7%)	6/27 (22.2%)	RR 0.73 (0.26 to 2.11)	60 fewer per 1000 (from 164 fewer to 247 more)	VERY LOW	CRITICAL
Major o	complication	1 I										
2	randomis ed trials	seriou s ²	no serious inconsisten cy	very serious ³	very serious ⁴	none	5/30 (16.7%)	1/27 (3.7%)	RR 3.37 (0.57 to 19.9)	88 more per 1000 (from 16 fewer to 700 more)	VERY LOW	CRITICAL

Quality	assessmer	nt					No of pa	atients	Effect			
No of studi es	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecis ion	Other considerati ons	Stent	Gastroenterost omy	Relativ e (95% Cl)	Absol ute	Qualit y	Importanc
1	randomis ed trials	very seriou s ¹	no serious inconsisten cy	very serious ³	very serious ⁴	none	8/9 (88.9%)	6/9 (66.7%)	RR 1.33 (0.8 to 2.23)	220 more per 1000 (from 133 fewer to 820 more)	VERY LOW	CRITICAL
Persist	ent obstruc	tive sym	ptoms									
1	randomis ed trials	seriou s ⁵	no serious inconsisten cy	very serious ³	very serious⁴	none	3/21 (14.3%)	3/18 (16.7%)	RR 0.86 (0.2 to 3.73)	23 fewer per 1000 (from 133 fewer to 455 more)	VERY LOW	CRITICAL
Recurr	ent obstruct	tive symp	otom									
1	randomis ed trials	seriou s ⁵	no serious inconsisten cy	very serious ³	very serious⁴	none	5/21 (23.8%)	1/18 (5.6%)	RR 4.29 (0.55 to 33.38)	183 more per 1000 (from 25 fewer to 1000 more)	VERY LOW	CRITICAL

Quality	assessmer	π					No of pa	atients	Effect			
No of studi es	Design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecis ion	Other considerati ons	Stent	Gastroenterost omy	Relativ e (95% CI)	Absol ute	Qualit y	Importance
1	randomis ed trials	seriou s ⁵	no serious inconsisten cy	very serious ³	very serious ⁴	none	7/21 (33.3%)	2/18 (11.1%)	RR 3 (0.71 to 12.66)	222 more per 1000 (from 32 fewer to 1000 more)	VERY LOW	CRITICAL
Mean ti	ime for oral	intake (B	etter indicate	d by lower v	/alues)							
1	randomis ed trials	very seriou s ¹	no serious inconsisten cy	very serious ³	no serious imprecisi on	none	9	9	-	MD 4.20 lower (5.53 to 2.87 lower)	VERY LOW	CRITICAL

95%CI=95%confidence interval; RR=relative risk

³ Inappropriate randomisation and no blinding
 ² Only one study.Jeurnink 2010 with inappropriate randomisation; Fiori 2004, Jeurnink 2010 - but no blinding in both studies
 ³ Majority people with gastric outlet obstruction from non-gastric origin
 ⁴ 95%CI crossed 2 boundaries of default MID
 ⁵ Appropriate randomisation but no blinding

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G.18 Curative treatment

What is the effectiveness of nutritional support interventions for adults undergoing curative treatment for oesophago-gastric cancer?

Quality	assessment	t			-		Nº of pat	tients	Effect	-		
Nº of studi es	Study design	Risk of bias	Inconsiste ncy	Indirectnes s	Impreci sion	Other considerati ons	Enteral nutritio n	parenter al nutrition or IV fluids	Relati ve (95% Cl)	Absolu te (95% Cl)	Quality	Importanc e
			cally during h		· •		17/047	00/004				
6	randomise d trials	seriou s ^a	no serious inconsisten cy	no serious indirectness	serious ^b	none	17/217 (7.8%)	33/224 (14.7%)	RR 0.52 (0.30 to 0.91)	71 fewer per 1,000 (from 13 fewer to 103 fewer)	LOW	CRITICAL
Surgic	al site infecti	ons (follo	ow up: Typica	lly during hos	pital stay)							
7	randomise d trials	seriou S ^a	no serious inconsisten cy	no serious indirectness	very serious °	none	26/217 (2.4%)	34/224 (15.2%)	RR 0.81 (0.46 to 1.42)	29 fewer per 1,000 (from 64 more to 82 fewer)	VERY LOW	CRITICAL
Anasta	motic leaks ((follow u	p: Typically d	uring hospital	stay)							
6	randomise d trials	seriou s ^a	no serious inconsisten cy	no serious indirectness	serious ^b	none	10/193 (5.2 %)	27/197 (13.7%)	RR 0.43 (0.22	78 fewer per 1,000	LOW	CRITICAL

Table 80: Clinical evidence profile. Early enteral feeding versus parenteral nutrition or IV support immediately after surgery

Quality	assessment	t					Nº of pat	tients	Effect			
№ of studi es	Study design	Risk of bias	Inconsiste ncy	Indirectnes s	Impreci sion	Other considerati ons	Enteral nutritio n	parenter al nutrition or IV fluids	Relati ve (95% Cl)	Absolu te (95% Cl)	Quality	Importanc e
									to 0.85)	(from 21 fewer to 107 fewer)		
Short t	erm mortality	y (follow	up: Typically	during hospita	al stay)							
6	randomise d trials	seriou S ^a	no serious inconsisten cy	no serious indirectness	very serious ^c	none	5/206 (2.4%)	4/213 (1.9%)	RR 1.08 (0.29 to 4.00)	2 more per 1,000 (from 13 fewer to 56more)	VERY LOW	IMPORTA NT
Length	of hospital s	stay (day	rs)									
4	randomise d trials	seriou s ^a	no serious inconsisten Cy	no serious indirectness	serious ^d	none	121	110	-	MD 0.96 days lower (2.54 lower to 0.61 higher)	LOW	IMPORTA NT
Weight	change (%)	(follow ເ	ıp: 14 days; as	sessed with:	Percentage	change from	baseline w	eight)				
1	randomise d trials	seriou s ^a	no serious inconsisten cy	no serious indirectness	no serious imprecis on	none	24	23	-	MD 2.11 % higher (0.15 higher to 4.07 higher)	MODERA TE	IMPORTA NT

CI=confidence interval; RR=relative risk; MD=mean difference;

a. Randomisation and allocation concealment unclear in most cases. Blinding either unclear or not present.
 b. 95% CI of the effect estimate includes one MID threshold [0.80, 1.25]
 c. 95% CI of the effect estimate includes both MID thresholds [0.80, 1.25]
 d. 95% CI of the effect estimate includes both the MID (1 day) and no effect

Table 81: Clinical evidence profile: immunonutrition versus standard nutrition during the perioperative period

	assessmen						Nº of patients	· ·	Effect			
№ of studi es	Study design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecisi on	Other considerati ons	Immunonutrit ion	standa rd nutritio n	Relati ve (95% CI)	Absolu te (95% CI)	Quality	Importan ce
Pneum	onia (follow	up: duri	ing hospital s	tay)								
12	randomis ed trials	seriou s ^a	no serious inconsisten cy	no serious indirectne ss	very serious ^b	none	74/550 (13.5%)	75/523 (14.3%)	RR 0.95 (0.71 to 1.26)	7 fewer per 1,000 (from 37 more to 42 fewer)	VERY LOW	CRITICAL
Surgic	al site infect	ions (fol	low up: durin	g hospital st	ay)							
12	randomis ed trials	seriou s ^a	no serious inconsisten cy	no serious indirectne ss	very serious ^b	none	43/550 (7.8%)	51/523 (9.8%)	RR 0.84 (0.56 to 1.25)	16 fewer per 1,000 (from 24 more to 43 fewer)	VERY LOW	CRITICAL
Anasta	motic leaks	(follow u	up: during ho	spital stay)								
8	randomis ed trials	seriou s ^a	no serious inconsisten cy	no serious indirectne ss	very serious ^b	none	20/442 (4.5%)	29/416 (7.0%)	RR 0.71 (0.41 to 1.22)	20 fewer per 1,000 (from 15	VERY LOW	CRITICAL

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Quality	/ assessmer	nt					№ of patients		Effect			
№ of studi es	Study design	Risk of bias	Inconsiste ncy	Indirectn ess	Imprecisi on	Other considerati ons	Immunonutrit ion	standa rd nutritio n	Relati ve (95% CI)	Absolu te (95% Cl)	Quality	Importan ce
										more to 41 fewer)		
Short t	erm mortalit	ty (follow	v up: Typically	y during hos	pital stay)							
9	randomis ed trials	seriou s ^a	no serious inconsisten cy	no serious indirectne ss	very serious ^b	none	14/476 (2.9%)	15/455 (3.3%)	RR 0.93 (0.46 to 1.90)	2 fewer per 1,000 (from 18 fewer to 30 more)	VERY LOW	IMPORTA NT
Overal	l survival - n	ot repor	ted									
1	randomis ed trials	very seriou s ^a	no serious inconsisten cy	no serious indirectne ss	no serious impreciso n	none	-	-	HR 0.93 (0.57 to 1.45)	-	LOW	CRITICAL
Length	of hospital	stay (da	ys)									
9	randomis ed trials	seriou s ^a	no serious inconsisten cy	no serious indirectne ss	no serious impreciso n	none	475	458	-	MD 2.7 days lower (3.19 lower to 2.21 lower)	MODERA TE	IMPORTA NT

CI=confidence interval; RR=relative risk; HR=Hazard ratio;

a. Allocation concealment unclear in most cases.
 b.95% CI of the effect estimate includes both MID thresholds [0.80, 1.25]
 c 32% not included in survival analysis but no ITT analysis

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Table 82: Clinical evidence profile. Oral nutritional supplements

Quality	assessment	t _					Nº of patien	ts	Effect			
№ of studi es	Study design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other considerati ons	Oral nutritional suppleme nts	placeb o	Relati ve (95% Cl)	Absolu te (95% Cl)	Quality	Importanc e
Advers	e events (gra	ade 2 or	more) (follow	up: range 4 v	weeks to 6 w	veeks)						
1	randomise d trials	seriou s ^a	no serious inconsisten cy	no serious indirectnes s	very serious ^b	none	15/58 (25.9%)	10/53 (18.9%)	RR 1.37 (0.68 to 2.78)	70 more per 1,000 (from 60 fewer to 336 more)	VERY LOW	CRITICAL
Short t	erm mortality	y (follow	up: range 4 w	eeks to 6 we	eks)							
1	randomise d trials	seriou s ^a	no serious inconsisten cy	no serious indirectnes s	serious ^c	none	1/58 (1.7%)	0/53 (0.0%)	RR 2.75 (0.11 to 65.98)	0 fewer per 1,000 (from 0 fewer to 0 fewer)	LOW	IMPORTA NT
Weight	change (%)	(follow u	p: range 4 we	eks to 6 wee	ks; assesse	d with: change	from baselin	e)				
2	randomise d trials	seriou s ^d	no serious inconsisten Cy	no serious indirectnes s	no serious impreciso n	none	77	69	-	MD 1.03 % higher (0.23 higher to 1.82 higher)	MODERA TE	IMPORTA NT

CI=confidence interval; RR=relative risk; MD=mean difference;

a. No blinding, unclear allocation concealment b. 95%Cl includes both MID thresholds [0.80, 1.25] c. 95%Cl includes both MID thresholds [0.80, 1.25], but the absolute risk difference is small d. No blinding in one trial, unclear allocation concealment in both

Quality	assessment	t					Nº of patie	ents	Effect			
№ of studi es	Study design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other considerati ons	Extra nutrition al support during CRT	placeb o	Relati ve (95% Cl)	Absolu te (95% Cl)	Quality	Importanc e
Treatm	ent related a	dverse e	ffects - Oral m	ucositis (gra	de 3 or more	e) (follow up: d	uring chem	o(radio)th	erapy)			
4	randomise d trials	seriou s ^a	no serious inconsisten cy	no serious indirectnes s	very serious ^b	none	10/123 (8.1%)	16/119 (13.4%)	RR 0.59 (0.17 to 2.03)	55 fewer per 1,000 (from 112 fewer to 138 more)	VERY LOW	CRITICAL
Treatm	ent related a	dverse e	ffects - Oesop	hagitis (grad	e 3 or more)	(follow up: du	ring chemo	(radio)the	rapy)			
1	randomise d trials	seriou s ^a	no serious inconsisten cy	no serious indirectnes s	very serious ^b	none	1/35 (2.9%)	1/36 (2.8%)	RR 1.03 (0.07 to 15.81)	1 more per 1,000 (from 26 fewer to 411 more)	VERY LOW	CRITICAL
Treatm	ent related a	dverse e	ffects - Diarrh	ea (grade 3 o	r more) (foll	ow up: during o	chemo(radi	o)therapy				
3	randomise d trials	seriou s ^a	no serious inconsisten cy	no serious indirectnes s	very serious ^b	none	10/113 (8.8%)	17/110 (15.5%)	RR 0.55 (0.26 to 1.14)	70 fewer per 1,000 (from 22 more to 114 fewer)	VERY LOW	CRITICAL

Table 83: Clinical evidence profile. Additional nutritional support during chemotherapy or chemoradiotherapy

Quality	assessment	t					Nº of patie	ents	Effect			
№ of studi es	Study design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other considerati ons	Extra nutrition al support during CRT	placeb o	Relati ve (95% Cl)	Absolu te (95% Cl)	Quality	Importanc e
Treatm	ent related a	dverse e	ffects - Nause	a (grade 3 or	[•] more) (follo	w up: during c	hemo(radio)therapy)				
3	randomise d trials	seriou s ^a	no serious inconsisten cy	no serious indirectnes s	serious °	none	35/113 (31.0%)	43/110 (39.1%)	RR 0.76 (0.56 to 1.04)	94 fewer per 1,000 (from 16 more to 172 fewer)	LOW	CRITICAL
Treatm	ent related a	dverse e	ffects - Vomit	ing (grade 3 d	or more) (fol	low up: during	chemo(radi	io)therapy	r)			
3	randomise d trials	seriou s ^a	no serious inconsisten cy	no serious indirectnes s	very serious ^b	none	3/113 (2.7%)	3/110 (2.7%)	RR 0.98 (0.19 to 5.22)	1 fewer per 1,000 (from 22 fewer to 115 more)	VERY LOW	CRITICAL
Treatm	ent related a	dverse e	ffects - compl	ication relate	ed infection (follow up: duri	ng chemo(r	adio)thera	apy)			
1	randomise d trials	seriou s ^a	no serious inconsisten cy	no serious indirectnes s	serious ^b	none	3/25 (12.0%)	11/25 (44.0%)	RR 0.27 (0.09 to 0.86)	321 fewer per 1,000 (from 62 fewer to 400 fewer)	LOW	CRITICAL

Quality	/ assessment	t					Nº of patie	ents	Effect			
№ of studi es	Study design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other considerati ons	Extra nutrition al support during CRT	placeb o	Relati ve (95% Cl)	Absolu te (95% Cl)	Quality	Importance
4	randomise d trials	seriou s ^a	no serious inconsisten cy	no serious indirectnes s	serious ^b	none	128/138 (92.8%)	120/13 5 (88.9%)	RR 1.03 (0.95 to 1.12)	27 more per 1,000 (from 44 fewer to 107 more)	LOW	IMPORTA NT
Short t	erm mortality	y (follow	up: during ch	emo(radio)th	erapy)							
1	randomise d trials	seriou S ^a	no serious inconsisten cy	no serious indirectnes s	very serious ^b	none	2/35 (5.7%)	3/36 (8.3%)	RR 0.69 (0.12 to 3.86)	26 fewer per 1,000 (from 73 fewer to 238 more)	VERY LOW	IMPORTA NT
Length	of hospital s	stay (day	s)									
1	randomise d trials	seriou s ^a	no serious inconsisten cy	no serious indirectnes s	no serious impreciso n	none	25	25	-	MD 4.48 days lower (7.08 lower to 1.88 lower)	MODERA TE	IMPORTA NT

Quality	assessment	t					Nº of patie	ents	Effect			
Nº of studi es	Study design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other considerati ons	Extra nutrition al support during CRT	placeb o	Relati ve (95% Cl)	Absolu te (95% Cl)	Quality	Importanc e
4	randomise d trials	seriou s ^a	no serious inconsisten cy	no serious indirectnes s	no serious impreciso n	none	138	138	-	MD 0.11 % higher (0.78 lower to 1 higher)	MODERA TE	IMPORTA NT

CI=confidence interval; RR=relative risk; MD=mean difference;

^a. No blinding or blinding unclear. Allocation concealment unclear

b. 95% CI of the effect estimate includes both MID thresholds [0.8, 1.25]
 c. 95% CI of the effect estimate includes one MID threshold [0.8, 1.25]

Table 84: Clinical evidence profile. Ccontinued routine nutritional support after discharge from hospital versus standard care

Quality	assessment	:					Nº of pati	ents	Effect			
Nº of studi es Jeiuno	Study design	Risk of bias	Inconsiste ncy - In hospital c	Indirectne ss	Imprecisi on	Other consideratio ns during hospital	Post dischar ge nutritio n support stav)	placeb o	Relativ e (95% CI)	Absolu te (95% Cl)	Quality	Importanc e
1	randomise d trials	seriou s ^a	no serious inconsistenc y	no serious indirectnes s	very serious ^b	none	11/22 (50.0%)	7/23 (30.4%)	RR 1.64 (0.78 to 3.46)	195 more per 1,000 (from 67 fewer to 749 more)	VERY LOW	CRITICAL

Quality	assessment						Nº of pati	ents	Effect			
Nº of studi es	Study design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other consideratio ns	Post dischar ge nutritio n support	placeb o	Relativ e (95% CI)	Absolu te (95% Cl)	Quality	Importanc e
Jejuno	stomy compl	ications	- Post dischar	ge (out of ho	spital) comp	lications (follow	w up: rang	e 6 weeks	s to 6 mor	nths)		
2	randomise d trials	seriou s ^a	no serious inconsistenc y	no serious indirectnes s	very serious ^b	none	12/43 (27.9%)	15/42 (35.7%)	RR 0.83 (0.51 to 1.35)	61 fewer per 1,000 (from 125 more to 175 fewer)	VERY LOW	CRITICAL
Pneum	onia											
1	randomise d trials	seriou S ^a	no serious inconsistenc y	no serious indirectnes s	very serious ^b	none	5/22 (22.7%)	7/23 (30.4%)	RR 0.75 (0.28 to 2.00)	76 fewer per 1,000 (from 219 fewer to 304 more)	VERY LOW	CRITICAL
Surgica	al site infection	ons										
1	randomise d trials	seriou s ^a	no serious inconsistenc y	no serious indirectnes s	very serious ^b	none	7/22 (31.8%)	6/23 (26.1%)	RR 1.22 (0.49 to 3.06)	57 more per 1,000 (from 133 fewer to 537 more)	VERY LOW	CRITICAL

Quality	assessment						Nº of pati	ents	Effect			
Nº of studi es	Study design motic leak	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other consideratio ns	Post dischar ge nutritio n support	placeb o	Relativ e (95% CI)	Absolu te (95% Cl)	Quality	Importanc e
							3/22	6/23	RR	125	VERY	CRITICAL
1	randomise d trials	seriou s ^a	no serious inconsistenc y	no serious indirectnes s	very serious ^b	none	3722 (13.6%)	(26.1%)	0.52 (0.15 to 1.84)	fewer per 1,000 (from 219 more to 222 fewer)	LOW	URITICAL
Sarcop	enia (follow i	up: range	e 6 weeks to 6	months; ass	essed with:	change in grip	strength fr	om basel	ine)			
3	randomise d trials	seriou s ^a	no serious inconsistenc y	not serious	no serious impreciso n	none	68	75	-	MD 1.02 kg (0.11 lower to 1.93 kg higher)	MODERA TE	IMPORTA NT
Short to	erm mortality	,										
1	randomise d trials	seriou s ^a	no serious inconsistenc y	no serious indirectnes s	very serious ^c	none	1/22 (4.5%)	0/23 (0.0%)	RR 3.13 (0.13 to 72.99)	0 fewer per 1,000 (from 0 fewer to 0 fewer)	LOW	IMPORTA NT
	Change in QC 100 to 100)	DL from b	paseline to 6 n	nonths (follow	v up: mean 6	6 months; asse	ssed with:	change ii	1 EORTC	QLQ-C30	from baselin	e; Scale
1	randomise d trials	seriou s ^a	no serious inconsistenc y	no serious indirectnes s	very serious ^d	none	16	20	-	MD 2 higher (12.57 lower to	VERY LOW	CRITICAL

Quality	/ assessment	t					Nº of pati	ents	Effect			
Nº of studi es	Study design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisi on	Other consideratio ns	Post dischar ge nutritio n support	placeb o	Relativ e (95% Cl)	Absolu te (95% Cl)	Quality	Importanc e
										16.57 higher)		
QOL -	QOL at the er	nd of foll	ow up (follow	up: range 6 v	veeks to 6 m	onths; assesse	d with: EO	RTC QLC	Q-C30; Sca	ale from: 0	to 100)	
2	randomise d trials	seriou s ^a	no serious inconsistenc y	no serious indirectnes s	serious ^e	none	30	33	-	MD 4.81 lower (15.52 lower to 5.89 higher)	LOW	CRITICAL
Weight	t change (kg)	assesse	d with: change	e from baseli	ne follow up	: range 6 week	s to 6 mont	ths				
3	randomise d trials	seriou s ^a	no serious inconsistenc y	no serious indirectnes s	serious ^f	none	30	75	-	MD 2.37 kg higher (0.48 to 4.27 higher)	LOW	IMPORTA NT

^a. No blinding

^a. No billinging
 ^b. 95% CI of the effect estimate includes both MID thresholds [0.80, 1.25]
 ^c. 95% CI of the effect estimate includes both MID thresholds [0.80, 1.25] - but absolute risk difference is small – so only downgraded one level
 ^d. 95% CI of the effect estimate includes both MID thresholds [-9, +9] - based on 0.5 SD of the control group
 ^e. 95% CI of the effect estimate includes one MID thresholds [-9, +9] - based on 0.5 SD of the control group
 ^f. 95% CI of the effect estimate includes one MID thresholds [-4, +4] - based on 0.5 SD of the control group

G.19 Palliative care

What is the effectiveness of nutritional interventions in adults with oesophago-gastric cancer receiving palliative care?

No evidence was identified for this review.

G.20 Routine follow-up

In adults who have undergone treatment for oesophago-gastric cancer with curative intent, with no symptoms or evidence of residual disease, what is the optimal method(s), frequency, and duration of routine follow-up for the detection of concurrent disease?

GRADE was not used for this review. See modified clinical evidence profile for evidence tables.

DRAFT FOR CONSULTATION Contents

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GRADE Profiles

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