# 1 ONS Guidelines ™ to Support Patient Adherence to Oral Anticancer Medications

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## Table 1. Study characteristics of additional studies for PICO 1

Study	Country	Study	N subjects	% female	Age mean	Type of cancer	Tools/methods used	Timing of risk	Findings from the risk assessment	Funding
		Design	(intervention/co		(SD) /	regimen	to assess risk	assessment		Source
			mparator)		Median					
					(IQR)					
Berry,	US	RCT	70 (49/21)	40	Median: 61	Diverse cancers	Measured odds of	Demographic	Symptom distress: OR: SDS-15+1	N/A
2015					Range: 34-	on chemotherapy	low/medium	characteristics	vs SDS-15a 1.1 (1.0–1.2)	
					80	and hormonal	adherence on	at baseline.		
						therapy	Symptom distress:	Unknown when	Depression:	
							SDS-15, Depression:	depression and	Demographic characteristics:	
							PHQ-9; demographic	symptom	Lack of a spouse/	
							characteristics	distress	partner, symptom distress,	
								assessments	younger age, not working at the	
								were taken.	start of therapy, female sex, and	
									oral chemotherapy vs oral	
									hormonal medications	

									NS association with low/medium adherence: cancer stage, working status, education, minority identification, age, married/partner status, time on regimen	
Decke	US	Cohort	30 (23/7)	94	Mean (SD):	Diverse cancers	Depression: CESD-	Baseline and	Functional ability (SF-12): NS btw	N/A
r/200					59.93	on diverse	20;, Functional	end of study (at	adherence and nonadherence	
9					(12.03)	treatments	ability: SF-12	the exit	group	
					Range:			interview)		
					21-71+				Depression (CESD-20): lower	
									scores at baseline (10.91 vs 13.13)	
									and end of study (8.67 vs 11.0) in	
									adherence group (NS)	
DosSa	France	Cohort	129	40%	Median: 70	Renal cell, lung,	Depression: CES-D,	Baseline (before	Significant negative association	N/A
ntos/						prostate,	Anxiety: STAI-Trait	initiation of	between depression and non-	
2019						colorectal, breast	(score range, Global	treatment)	adherence	

						cancers treated	cognitive status:			
						with targeted	MoCA, Digit			
						therapy,	memory: WAIS-III,			
						chemotherapy,	Information			
						and	processing speed:			
						chemoradiothera	TMT, Autonomy:			
						ру	IADL			
Jacob	US	Cohort	90	55.6	Mean (SD):	Diverse cancers	Symptom distress:	Baseline and	- Demographic: Women had	Massac
s/					58.06	on oral	Symptom Distress	post-	greater adherence than men	husetts
2017					(13.08)	chemotherapy	Scale, Anxiety and	assessment (12	(93.48% vs 83.90%) (S)	General
					Range: 28-		depressive	weeks)	- Significant associations with	Hospital
					88		symptoms: Hospital		better adherence: improvements	Cancer
							Anxiety and		in symptom distress (-0.79),	Center
							Depression Scale,		depressive symptoms (-1.57),	
							Cancer-specific		quality of life (0.38),	
							psychological		- Improvements in patient-	
							distress: Cancer		reported symptom distress (23.94	

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3							Worries Inventory		at baseline and -0.22 change from	
							(CWI)		baseline), depressive symptoms	
200									(4.23 at baseline and 0.37 change	
									from baseline), satisfaction with	
									clinician communication and	
, po									treatment (92.68 at baseline and -	
מלום: מרוים: מרוים:									2.84 change from baseline), and	
									perceived burden to others (5.04	
									at baseline and -0.04 change from	
2									baseline) were associated with	
									better adherence. No association	
									between anxiety and adherence	
Krikor	US	RCT	200 (101/99)	77	Interventio	Diverse cancers	Beliefs about	Assessment	Non-adherence was associated	N/A
ian/					n - Mean	on oral	medicines: BMQ	taken at	with forgetfulness, wanting to	
2019					(SD): 61.8	antine oplastic		baseline.	avoid side-effects, being	
					(11.5)	medication		Demographic	depressed or overwhelmed,	
N					Control -			forms were	falling asleep before taking	
				<u> </u>				<u> </u>	<u>l</u>	

					Mean (SD):			updated at later	medication. Numbers not	
5 5 5					61.9 (12)			time points.	provided. Supplement only	
									provides the questions in BMQ.	
									Statistically significant	
									correlations associated with non-	
									adherence were forgetfulness (p =	
									0.009), wanting to avoid side	
									effects (p = 0.02), feeling	
									depressed or overwhelmed (p =	
n 1									0.032), or falling asleep before	
									taking medication (p = 0.048) in	
									both groups	
Krolo	German	Cohort	73	74	N/A	Breast cancer,	N/A	Separated into	Found no associations between	Supple
p/201	у					colorectal cancer,		initially non-	age, gender, any	mentar
3						and esophageal		adherent and	sociodemographic or disease-	y grant
						cancer treated		adherent after	related characteristics to	was

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						with capecitabine		first follow-up	adherence. No numbers	provide
						in combination or			reported.	d by
						monotherapy				Roche,
										Basel
Timm	Netherl	Cohort	62	47	Mean: 63.5	Non small cell	Demographic	Collected at	Relationships with incorrect	Roche,
ers/	ands					lung cancer on	characteristics,	baseline	intake were: older age (OR 1.10,	The
2015						erlotinib	smoking, co-		95 % CI 1.00–1.21), MARS < 25	Netherl
							medications, Quality		(OR 4.83, 95 % CI 1.06–21.99),	ands
							of life: SF-12,		oculair symptoms (OR 3.13, 95 %	
							Attitude(s) towards		CI 1.11–8.82) and stomatitis (OR	
							medication: BMQ,		6.59, 95 % CI 1.77–24.60)	
							Illness perception:			
							Brief IPQ, and		BMQ and Brief IPQ can be found	
							symptoms (likert		in Table 8	
							scale)			
Wicke	US	Cohort	198 (162/36)	100	Mean (SD):	Breast cancer	Sociodemographic	Information on	Depressive symptoms, fatigue,	Nationa
rsham					59.1 (7.5)	treated with	variables: University	predictor	gastrointestinal symptoms,	I

/2013					Range: 39-	Anastrozole,	of Pittsburgh, School	variables was	cognitive symptoms, weight	Institut
					75	Letrozole,	of Nursing Center for	measured pre-	concerns, gynecological	e for
						Examestane,	Research in Chronic	treatment	symptoms, musculoskeletal pain,	Nursing
						Tamoxifen	Disorders		and total BCPT score were	
							Sociodemographic		identified as linear predictors of	
							Questionnaire,		nonadherence. Numbers are not	
							Depressive		reported	
							symptoms: Beck			
							Depression			
							Inventory-II, Anxiety:			
							Profile of Mood			
							States (POMS)			
							Tension-Anxiety			
							subscale, Side effects			
							of hormonal therapy:			
							ВСРТ			
Yusuf	US	Cohort	73 (54/19)	100	Mean (SD):	Breast cancer on	Depression: The	All measured at	Psychological and menopause	N/A

ov/	55 (10.1)	tamoxifen and	Patient Health	baseline	symptoms (depression,
2020		aromatase	Questionnaire (PHQ-		generalized anxiety, insomnia,
		inhibitors	8), Tendency to		somatosensory amplification, hot
			perceive normal		flash frequency, and hot flash-
			visceral or somatic		related interference) were
			sensations as being		assessed pre-AET initiation as
			dangerous,		predictors of subsequent non-
			abnormal, intense,		adherence
			or potentially		Adherent vs non-adherent:
			harmful The		Anxiety: 3.1(4.2) vs 4.1(4.6)
			Somatosensory		Depression: 3.4 (3.3) vs 6.0 ( 3.9)
			Amplification Scale		Insomnia (subthreshold): 7.5 (5.3)
			(SSAS), Anxiety: The		vs 7.7(4.6)
			Generalized Anxiety		Hot flash related interference: 6.2
			Disorder (GAD-7),		(15.2) vs 7.4(14.1)
			Sleep: The Insomnia		Somatosensory Amplification:
			Severity Index (ISI),		22.3(6.5) vs 26.5(8.5)

serves all	Hot flash related	Hot flash frequency: 1.1(2.0) vs
Org. ONS res	interference: The	2.0(3.0)
sions @ ons.	Hot Flash-Related	
i pubpermis:	Daily Interference	
please emai	Scale (HFRDIS)	

- **Table 2. Evidence Profile for PICO 1** 15
- 16 Question: Standardized assessment for risk/barriers compared to standard of care for Patients starting a new oral anti-cancer medication
- 17 regimen
- **Setting**: Outpatient 18

all rights.	5 <b>Ta</b>	able 2. Evidence Profile for PICO 1											
NO Si Constantina de la constantina della consta	6 <b>Q</b> ւ	ı <b>estion</b> : Si	tandardized asse	ssment for ris	k/barriers con	npared to sta	andard of care for	Patients sta	irting a r	new oral anti-ca	ncer medicat	ion	
o.sno@ons.o	7 re	regimen											
il pubpermiss	8 <b>Se</b>	Setting: Outpatient											
e, please ema			Certainty as	sessment			Nº of patie	ents	Effect				
For permission to post conline, reprint, adapt, or reuss		Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerat ions	standardized assessment for risk/barriers	standard of care	Relati ve (95% CI)	Absolute (95% CI)	Certainty	Importance	

### Adherence rate (follow up: 4 months; assessed with: self-report)

1 1	rando	not	not serious	serious <sup>b</sup>	very	none	25 participants who received risk assessment plus	$\oplus$	CRITICAL
	mised	serious			serious <sup>c,d</sup>		tailored intervention had an adherence rate of	VERY LOW	
	trials	а					95.1% vs 20 participants in the control arm with an		
							adherence rate of 82.4%.		
- 16 66	_								

### Self-efficacy to manage medications - not reported

2												
-	-	-	-	-	-	-	-	-	-	-	-	IMPORTANT

Health-related Quality of Life and Patient-reported Outcomes (HRQOL/PROs) - not reported

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reserves all ri	-	-	-	-	-	-	-	-	-	-	-	CRITICAL
Patient s	satisfact	ion - not	reported									
issions @ o	-	-	-	-	-	-	-	-	-	-	-	CRITICAL

- 19 **CI:** Confidence interval
- 20 Explanations
- a. Minimal information provided about randomization and allocation concealment.
- b. Intervention included tailored coaching intervention in addition to risk assessment.
- 23 c. Sample doesn't meet optimal information size. Concerns with fragility.
- d. The possibility of no difference cannot be excluded due to limited information.
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### 28 Table 3. Evidence Profile for PICO 2

**Certainty assessment** 

- 29 **Question**: Educational programs compared to standard of care for patients starting a new oral anticancer medication regimen
- 30 **Setting**: Outpatient

trials

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d												
Nº of studies	Study design	Risk of bias	Inconsistency	Indirectness		Other consid eration	educational programs	standard of care	Relative (95% CI)	Absolute (95% CI)	Certainty	Importance
Adheren	ice rate (fo	ollow up:	3-12 weeks; ass	sessed with: so	elf-report	and pill	count)					
2 <sup>1,2</sup>	randomi	serious	not serious	not serious	very	none	215	156	-	MD <b>0.4 % higher</b>	ФООО	CRITICAL
	sed	а			serious					(1.87 lower to 2.68 higher)	VERY LOW	

Nº of patients

**Effect** 

### Adherence rate (follow up: 2-24 weeks; assessed with: self-report and medication event monitoring system pillboxes)

b,c

4 3,4,5,6	observat	very	not serious	not serious	serious	none	83	100	-	MD <b>10.61</b> % higher	ФООО	CRITICAL
e-user license	ional	serious			b					(7.21 higher to 14.01	VERY LOW	
0-2024. Single	studies	d								higher)		

Proportion with high adherence (follow up: 14-24 weeks; assessed with: MMAS-4 and MMAS-8)

n												
2 <sup>7,8</sup>	randomi	serious	not serious	not serious	not	none	222/391	175/354	RR 1.16	79 more per 1,000	$\Theta\Theta\Theta\Theta$	CRITICAL
	sed	е			serious		(56.8%)	(49.4%)	(1.01 to	(from 5 more to 163 more)	MODERATE	
2 2 2 3 3 4 5 5 6 7	trials								1.33)			
Patient	satisfactio	n (assess	ed with: Helpful	ness of meeti	ng with s	pecialty p	oharmacist an	d medication	on navigator	- % "very")		
<b>1</b> 9	observat	very	not serious	not serious	very	none	30/39	32/37	RR 0.89	95 fewer per 1,000	⊕000	CRITICAL
	ional	serious			serious		(76.9%)	(86.5%)	(0.72 to	(from 242 fewer to 86	VERY LOW	
	studies	f,g			c,h				1.10)	more)		
Patient	satisfactio	n (assess	ed with: Helpful	ness of medic	ation info	sheet - 9	% "very")	ı				
1 <sup>9</sup>	observat	very	not serious	not serious	very	none	25/39	28/37	RR 0.85	114 fewer per 1,000	ФООО	CRITICAL
	ional	serious			serious		(64.1%)	(75.7%)	(0.63 to	(from 280 fewer to 106	VERY LOW	
	studies	f,g			c,h				1.14)	more)		
Patient	satisfactio	n (assess	ed with: Helpful	ness of check	in with n	nedicatio	n navigator -	% very")				
1 <sup>9</sup>	observat	very	not serious	not serious	serious	none	27/39	34/37	RR 0.75	230 fewer per 1,000	⊕○○○	CRITICAL
	ional	serious			b		(69.2%)	(91.9%)	(0.60 to	(from 368 fewer to 46	VERY LOW	
	studies	f,g							0.95)	fewer)		
Patient	knowledge	e of regim	nen (follow up: 2	2 cycles; asses	sed with:	Dosage a	and frequency	y)				
1 <sup>10</sup>	observat	very	not serious	not serious	serious	none	29/29	23/29	RR 1.26	206 more per 1,000	ФООО	CRITICAL
	1				ı			l				

b. Small sample, concerns with fragility.

35

- 36 c. The 95% CI cannot exclude the potential for no difference.
- 37 d. Critical concern with confounding and missing data. Serious concern with bias in the selection of participants.
- e. Some concerns with randomization, effect of assignment to intervention, missing outcome data and measurement of the outcome.
- 39 f. Critical concern with confounding, moderate concern in selection of participants and measurement of outcome.
- 40 g. Not measuring satisfaction before and after intervention, instead looks at satisfaction a little after start of intervention and end of
- 41 intervention.
- 42 h. Few events reported do not meet the optimal information size and suggest fragility of the estimate.
- 43 i. Critical concern with confounding.

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- patients with cancer taking capecitabine: a prospective two-arm cohort study. BMJ Open; 07/2013.
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- 61 with symptom distress, depression, and personal characteristics. Patient Preference and Adherence; 11/2015.
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- Susan. Adherence Measures for Patients with Metastatic Castration-Resistant Prostate Cancer Treated with Abiraterone Acetate plus
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- Angie Mae. Enhancing patients' understanding of and adherence to oral anticancer medication: Results of a longitudinal pilot intervention.
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- anticancer medication management clinic. Journal of Pharmacy Practice and Research; 12/2018.

- Question: Standardized, periodic/ongoing assessment of adherence compared to usual care for patients on an oral anti-cancer medication

70 71 72 73	<b>Ques</b> regin	s <b>tion</b> : Sta			g assessment (	of adhere	ence compared to	o usual care	for patien	ts on an oral anti-cancer medi	ication	
Nº of studies	Study design	Risk of bias	Certainty asso		Imprecision	Other conside rations	ng assessment		Relative (95% CI)	Effect  Absolute  (95% CI)	Certainty	Importance
Adherenc	random	,	: <b>12 weeks; ass</b> not serious	not serious	very serious	none	75	83	-	MD <b>2.34 % higher</b> (5.58 lower to 10.26 higher)	⊕⊕○○ LOW	CRITICAL
0	e rate (fo	•	: 6 months; ass	not serious	elf-report) serious a	none	34	51	-	MD <b>7 % highe</b> r	⊕○○○	CRITICAL

1 <sup>1</sup>	random	not	not serious	not serious	very serious	none	75	83	-	MD <b>2.34</b> % higher	$\Theta\ThetaOO$	CRITICAL	
	ised	serious			a,b					(5.58 lower to 10.26 higher)	LOW		
	trials												

1 <sup>2</sup>	observa	very	not serious	not serious	serious <sup>a</sup>	none	34	51	1	MD <b>7</b> % higher	ФООО	CRITICAL
	tional	serious								(0.66 higher to 13.34	VERY LOW d	
	studies	С								higher)		

Adherend	ce (Tollow	/ up: 21-	zo days; assess	sea with: reia	tive dose inter	isity)						
1 <sup>3</sup>	random	serious	not serious	not serious	very serious	none	31	37	-	MD <b>0.32</b> % higher	ФООО	CRITICAL
	ised	e			a,b					(0.08 lower to 0.72 higher)	VERY LOW	
	trials											
Quality o	f life (foll	ow up: 1	12 weeks; asse	ssed with: FA	CT-G; higher=l	oetter; N	/IID 5-7; Scale fro	m: 0 to 108	3)			
1 ¹	random	not	not serious	not serious	serious <sup>a</sup>	none	77	85	-	MD <b>2.28 points higher</b>	0000	CRITICAL
	ised	serious								(1.93 higher to 2.63 higher)	MODERATE	
	trials	f										
Quality o	f life (foll	ow up: 3	3 months; asse	ssed with: EO	RTC; higher=b	etter; M	IID 4-11)		<u> </u>			
1 4	observa	serious	not serious	not serious	serious <sup>a</sup>	none	56	56	-	MD <b>15.7 points higher</b>	ФФОО	CRITICAL
	tional	g								(8.84 higher to 22.56	LOW	
	studies									higher)		
Patient sa	atisfactio	n (follow	v up: 3 months	; assessed wi	th: self-report	(single o	question on satis	faction))				
1 <sup>5</sup>	observa	very	not serious	not serious	very serious i	none	20/20 (100.0%)	15/20	RR 1.32	240 more per 1,000	ФООО	CRITICAL
	tional	serious						(75.0%)	(1.02 to	(from 15 more to 540 more)	VERY LOW	
	studies	h							1.72)			
Cancer-re	elated mo	orbidity (	follow up: 24 v	weeks; assess	ed with: globa	l toxicit	y score; higher=w	vorse; Scale	from: 0 to	36)		
											19	

1 6	random	serious	not serious	not serious	very serious	none	92	91	-	MD 1 points higher	ФООО	CRITICAL
	ised	j			a,b					(1.72 lower to 3.72 higher)	VERY LOW	
	trials											
Cancer-re	elated mo	rbidity (	follow up: 21-7	 28 days; asses	ssed with: Sym	nptom Ex	perience Invent	ory; higher	=worse; So	ale from: 0 to 190)		
1 <sup>3</sup>	random	serious	not serious	not serious	very serious	none	31	37	-	MD 1.75 points lower	⊕○○○	CRITICAL
	ised	e			a,b					(9.48 lower to 5.98 higher)	VERY LOW	
	trials											
Cancer-re	elated mo	rbidity (	follow up: 8 w	eeks; assesse	d with: Sympt	om Expe	rience Inventor	 y; higher=w	orse; Scale	e from: 0 to 190)		
1 <sup>7</sup>	observa	very	not serious	not serious	serious <sup>a</sup>	none	24	30	-	MD <b>4.78 points lower</b>	⊕○○○	CRITICAL
	tional	serious								(7.8 lower to 1.76 lower)	VERY LOW	
		k										
;	studies	K										
Self-effic			-28 days; asses	ssed with: MA	ASES-R; higher	=better;	Scale from: 1 to	4)				
Self-effica		w up: 21	-28 days; asses	ssed with: MA	ASES-R; higher	= <b>better</b> ;	Scale from: 1 to	<b>4)</b> 37	-	MD <b>0.51 points lower</b>	⊕○○○	IMPORTANT
	acy (follov	w up: 21	•	1					-	MD <b>0.51 points lower</b> (1.3 lower to 0.28 higher)	⊕○○○ VERY LOW	IMPORTANT
	random	w up: 21	•	1	very serious				-	·		IMPORTANT
1 <sup>3</sup>	random ised trials	w up: 21 serious	•	not serious	very serious	none	31		-	·		IMPORTANT

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serves all rig	tional	serious			a,b					(0.36 lower to 0.34 higher)	VERY LOW	
s.org. ONS res	studies	k										
® Adherence	ce to supp	oortive c	are/lab monito	oring - not rep	oorted							
ail pubper	-	-	-	-	-	-	-	-	-	-	-	IMPORTANT

- 74 CI: Confidence interval; MD: Mean difference; MID: Minimally important difference; RR: Risk ratio; MASES-R: Medication Adherence Self-
- 75 Efficacy Scale Revision
- 76 Explanations
- a. Small sample, concerns with fragility.
- 78 b. 95% CI cannot exclude the possibility of no effect.
- 79 c. Moderate concern with confounding. and measurement of outcome due to subjective measure. Serious concern with missing data.
- d. An additional study reported a risk ratio of 0.92; 95% CI: 0.54, 1.56 comparing on-going assessment to no assessment measured with self-
- reported adherence at 3 months.
- 82 e. Some concerns due to deviations from the intended interventions.
- f. Self-reported outcome measurement could lead to some concerns with risk of bias but not serious.
- 84 g. Critical concern with confounding and serious concern with subjectivity of outcome.
- h. Critical concern for confounding and moderate concern with measurement of outcome due to self-report.
- i. Few events reported do not meet the optimal information size and suggest fragility of the estimate.
- i. Some concerns due to deviations from the intended interventions and self-reported outcome measurement.

- k. Serious concern with confounding, bias in selection of participants, missing data and measurement of outcome. Moderate concern withdeviations from intervention.
  - References

99

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#### **Table 5. Evidence Profile for PICO 4** 112

#### **Setting**: Outpatient 114

eserves all rights.	112	2 Tab	le 5. Evid	ence Profile for	PICO 4								
org. ONS reser	113	3 Que	e <b>stion</b> : Ac	tive follow-up co	ompared to us	ualcare for pat	tients on ar	oral antica	ancer medicatio	n regimen who ha	ve additional risk fac	tors	
issions@ons.c	114	1 Sett	i <b>ng</b> : Outp	patient									
il pubperm				Certainty ass	sessment			Nº of	fpatients	Ef	fect		
or reus	of dies	Study design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other consider	active follow-	standard of	Relative (95% CI)	Absolute (95% CI)	Certainty	Importance
reprint, ada	aics	acsign	Dias				ations	up	care	(3370 Ci)	(33% C.)		
to bost online,	neren	ce rate (	follow up	o: 6 cycles; asses	sed with: MEN	MS (medicatio	n event mo	onitoring sy	rstem) pillboxes	s)			
ermission 1	1	observ	very	not serious	not serious	very serious	none	10	10	-	MD <b>17.8</b> % higher	ФОО	CRITICAL
ciety. For pr		ational	serious			b					(6.43 higher to	0	
Nursing So		studies	a								29.17 higher)	VERY	
the Oncology												LOW	
ght 202 by I	ncer-r	elated m	orbidity	- not reported		<u> </u>	<u> </u>	<u> </u>			l	<u>I</u>	
only. Copyri	-	-	-	-	-	-	-	-	-	-	-	-	CRITICAL
Qua	ality c	of life - n	ot report	ed		I			1		1	ı	l .
4. Single-t	-	-	-	-	-	-	-	-	-	-	-	-	CRITICAL

reserves all rig	-	-	-	-	-	-	-	-	-	-	-	CRITICAL
Patient	self-effica	cy about	treatment - no	t reported								
nissions @ o	-	1	-	-	-	-	-	-	-	-	-	IMPORTANT

- 115 **CI:** Confidence interval; **MD:** Mean difference
- 116 Explanations
- 117 a. Critical concern with confounding.
- b. Small sample, concerns with fragility.
- 119 References
- 120 1. Vacher, Laure, Thivat, Emilie, Poirier, Camille, Mouret-Reynier, Marie-Ange, Chollet, Philippe, Devaud, Hervé, Dubray-Longeras, Pascale,
- 121 Kwiatkowski, Fabrice, Durando, Xavier, van Praagh-Doreau, Isabelle, Chevrier, Régine. Improvement in adherence to Capecitabine and Lapatinib
- by way of a therapeutic education program. Supportive Care in Cancer; 07/2020.

#### **Table 6. Evidence Profile for PICO 5** 123

#### 125 **Setting**: Outpatient

erves all rights.	.23 <b>Tal</b>	ole 6. Evi	dence Profile fo	r PICO 5								
g. ONS reserves	.24 <b>Q</b> u	<b>estion</b> : C	oaching compar	ed to usual ca	are for patien	ts on an ora	l anti-can	cer medica	tion regimen v	who have additional risk factors		
ons.ons @ ons.or	.25 <b>Se</b> t	t <b>ting</b> : Out	tpatient									
ail pubpermi			Certainty ass	sessment			Nº of ∣	patients		Effect		
t, or reuse, please ema	Study	Risk of	Inconsistency	Indirectness	Imprecision	Other considerati		standard	Relative	Absolute	Certainty	Importance
studies	design	bias				ons	g	of care	(95% CI)	(95% CI)		
Adhere	ence rate	(follow u	p: 3-4 weeks; as	sessed with:	pill count)							
noissim 1 1	random	serious	not serious	not serious	very serious	none	101	99	-	MD <b>0.8</b> % higher	ФООО	CRITICAL
g Society. For	ised	a			b,c					(2.24 lower to 3.84 higher)	VERY LOW	
oav Nursin	trials											
Adhere	ence rate	(follow u	p: 2 educational	sessions eve	ry three cycl	es; assessed	with: M	EMS pillbo	kes) <sup>d</sup>			
1 <sup>2</sup>	observa	very	not serious	not serious	serious <sup>c</sup>	none	10	10	-	MD <b>17.8 % higher</b>	ФООО	CRITICAL
e only. Copy	tional	serious								(6.43 higher to 29.17 higher)	VERY LOW	
gle-user license	studies	е										
Adhere	ence (follo	w up: 3 i	months; assesse	d with: MPR	greater than	or equal to	90%)	<u>.                                      </u>		ı	ı	
0. 1 3 aged on 05-2-2	random	serious <sup>f</sup>	not serious	serious <sup>g</sup>	very serious	none	59/64	54/59	RR 1.01	9 more per 1,000	⊕○○○	CRITICAL
Down	1	1	I	1	ı		1	<u> </u>		1	26	

	ised				b,h		(92.2%)	(91.5%)	(0.91 to 1.12)	(from 82 fewer to 110 more)	VERY LOW	
	trials											
Adhere	ence (folio	ow up: 6-3	31.9 months; as	ssessed with:	MPR)							
2 <sup>4,5</sup>	observa	very	serious <sup>j</sup>	serious <sup>g</sup>	serious <sup>c</sup>	none	84	281	-	MD <b>2.98 % higher</b>	⊕○○○	CRITICAL
	tional	serious <sup>i</sup>								(2.95 higher to 3.01 higher)	VERY LOW	
	studies											
Cancer	related r	norbidity	-Symptom seve	erity (follow u	ıp: 3 months;	assessed w	/ith: 13 ite	em M.D. A	nderson Sympt	om Inventory; higher=worse; N	1ID 1.0 per 10	) point scale
Scale f	rom: 0 to	130)										
. 2		f		not corious	very serious	none	64	62	_	MD 0 points	ФООО	CRITICAL
1 <sup>3</sup>	random	serious <sup>f</sup>	not serious	not serious	very serious	Hone	04	02	-	WID <b>o points</b>	10000	CRITICAL
1 ³	ised	serious	not serious	not serious	b,c	none	04	02	-	(0.55 lower to 0.55 higher)	VERY LOW	CRITICAL
1°		serious	not serious	not serious	,	none	04	02	-	·		CRITICAL
	ised trials				b,c				er; Scale from: 1	(0.55 lower to 0.55 higher)		CRITICAL
	ised trials t self-effic			s; assessed wi	b,c				er; Scale from: 1	(0.55 lower to 0.55 higher)	VERY LOW	
Patient	ised trials t self-effic	cacy (follo	w up: 3 month	s; assessed wi	b,c ith: General s	elf-efficacy	scale; hig	her=bette	er; Scale from: 1	(0.55 lower to 0.55 higher)	VERY LOW	
Patient	ised trials t self-effice random	cacy (follo	w up: 3 month	s; assessed wi	b,c ith: General so	elf-efficacy	scale; hig	her=bette	er; Scale from: 1	(0.55 lower to 0.55 higher)  to 40)  MD 1.8 points higher	VERY LOW	
Patient	ised trials t self-effice random ised trials	serious <sup>f</sup>	w up: 3 month	s; assessed wi	b,c  ith: General so  very serious  b,c,h	e <b>lf-efficacy</b> none	scale; hig	her=bette	-	(0.55 lower to 0.55 higher)  to 40)  MD 1.8 points higher	VERY LOW	IMPORTAN <sup>-</sup>

erves all	ised		b,c			(6.18 lower to 6.58 higher)	VERY LOW	
ns. org. ONS rese	trials							

gratient satisfaction (follow up: 3 months; assessed with: self-designed scale; higher=better; Scale from: 0 to 5)

1 <sup>3</sup>	random	serious <sup>f</sup>	not serious	not serious	very serious	none	64	62	-	MD <b>0.1 points higher</b>	ФООО	CRITICAL
	ised				b,c					(0.9 lower to 1.1 higher)	VERY LOW	
	trials											

- 126 CI: Confidence interval; MD: Mean difference; MEMS: Medication event monitoring system; MPR: Medication possession ratio; RR: Risk ratio;
- 127 MID: Minimally important difference
- 128 Explanations
- a. Serious concern with missing outcome data and selection of the reported result.
- b. The 95% CI cannot exclude the potential for no difference.
- 131 c. Small sample, concerns with fragility.
- d. Reflects the mean of the daily adherence scores which correspond to the proportion of pills actually taken (recorded opening by MEMS) in
- 133 comparison with prescribed amounts (expected openings).
- e. Critical concern with confounding and missing outcome data.
- 135 f. Serious concerns with missing outcome data.
- g. MPR is surrogate for adherence.

- 137 h. Few events reported do not meet the optimal information size and suggest fragility of the estimate.
- i. Critical concern with confounding.
- j. Concerns with heterogeneity due to I2 value of 100%.
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153 Table 7. Evidence Profile for PICO 6

Question: Motivational interviewing compared to usual care for patients on an oral anti-cancer medication regimen who have additional risk

155 factors

154

156 **Setting**: Outpatient

se, please e				Certainty a	assessment			Nº of pati	ents		Effect		
eprint, a	№ of studies		Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations		standard of care	Relative (95% CI)		Certainty	Importance
SSic	۸dharan	co rato l	follow u	n. 12 wooks. 20	caccad with	colf_ranart)							

#### Adherence rate (follow up: 12 weeks; assessed with: self-report)

1 <sup>1</sup>	random	not	not serious	not serious	very serious	none	57	114	-	MD <b>3.23</b> % higher	$\Theta\ThetaOO$	CRITICAL	
	ised	seriou			a,b					(0.45 higher to 6.02	LOW		
	trials	S								higher)			

### Cancer-related morbidity - Summed symptom severity (follow up: 8 weeks; assessed with: Symptom Experience Inventory; Higher=worse; Scale from: 0 to 190)

1 <sup>2</sup>	observa	very	not serious	not serious	serious <sup>a</sup>	none	24	30	-	MD <b>4.78 points lower</b>	ФООО	CRITICAL
	tional	seriou								(7.8 lower to 1.76	VERY LOW	
-Z0Z4. SIIIgir	studies	s <sup>c</sup>								lower)		

Patient-self efficacy about treatment (follow up: 12 weeks; assessed with: MASES; higher=better; Scale from: 1 to 96)

 $\Theta\ThetaOO$ 

LOW

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**VERY LOW** 

**IMPORTAN** 

**IMPORTAN** 

Т

- d. Some concerns with bias due to subjectivity of outcome measurement and limited information provided about analysis used to estimate theeffect of assignment to intervention.
- e. Scale used to measure outcome not specified.
- 166 f. CI does not have meaningful difference thus not docked down for CI.

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167

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ves all rights.	75 <b>Ta</b> l	ble 8. Ev	idence Profile 1	or PICO 7								
ons reserved.	76 <b>Q</b> u	estion: -	Гесhnology con	npared to usu	ıal care for pa	itients on an o	ral anti-cancer n	nedication regir	nen			
ons @ ours @ ours @ ours @ ours	77 <b>Se</b> t	tting: Ou	itpatient									
ail pubpermis			Certainty a	ssessment			Nº of pa	tients		Effect		
Nº of studies	Study design		Inconsistency	Indirectness	Imprecision	Other consideratio	technology	standard of	Relative (95% CI)	Absolute (95% CI)	Certainty	Importance
Adherei	_	•					ottle openings)	00		MD 0 22 0/ bish siz		CDITICAL
sing Society. For permis	mised	serious a	serious <sup>s</sup>	not serious	serious <sup>c</sup>	none	91	99	-	MD <b>8.23</b> % higher (2.9 higher to 13.55	VERY	CRITICAL
A by the Oncology Nur Adherei	trials	(follow u	up: 6 months; a	ssessed with	: MPR)					higher)	LOW	
1 3 3	observ ational	very serious	not serious	not serious	serious <sup>c</sup>	none	50	51	-	MD <b>4.7</b> % higher (1.19 higher to 8.21	⊕○○○ VERY	CRITICAL
4. Single-user licer	studies		se intensity (fo	llow un: 3-13	weeks: asses	ssed with: nill	counts)			higher)	LOW	
7 4,5		serious			very serious	none	149	152		MD <b>0.01</b> % lower	<b>⊕</b> ○○○	CRITICAL
Down <u>loaded r</u>		50.7000			12., 20.1043					3.32 / 3.03.61	22	

	mised	е			c,g					(0.04 lower to 0.02	VERY	
	trials									higher)	LOW	
Cancer r	related n	norbidity	- Summed syı	mptom sever	ty (follow up	: 21 days; ass	essed with: Symp	otom Experien	ce Inventory; hi	gher=worse; Scale fro	om: 0 to 190	))
1 <sup>6</sup>	rando	not	not serious	not serious	very serious	none	49	26	-	MD 3.5 points	<b>ФФОО</b>	CRITICAL
	mised	serious			c,g					lower	LOW	
	trials									(12.48 lower to 5.48		
										higher)		
Quality	of Life (f	ollow up	: 3-12 weeks;	assessed with	: FACT-G and	WHO Quality	of Life-BREF Sca	ale; higher=bet	tter)			
2 1,7	rando	serious	serious <sup>h</sup>	not serious	serious <sup>c</sup>	none	77	85	-	SMD <b>1.44 SD higher</b>	<del>Ф</del> ООО	CRITICAL
	mised	a								(1.15 higher to 1.74	VERY	
	trials									higher)	LOW	
Quality	of Life (f	ollow up	: 6 months; as	sessed with:	assessed usin	g the EuroQo	  -5D (EQ-5D); MI	D 0.061; highe	r=better)			
1 <sup>3</sup>	observ	very	not serious	not serious	serious <sup>c</sup>	none	50	51	-	MD 0.13 points	⊕OOO	CRITICAL
	ational	serious								higher	VERY	
	studies	d								(0.07 lower to 0.2	LOW	
										higher)		
Patient :	satisfact	ion (follo	ow up: 6 cycles	(ranging fron	n 21 day to 90	0 day cycles);	assessed with: F	ACIT-TS-PS; hi	gher=better; Sc	ale from: 0 to 73)	<u>                                       </u>	
											34	

18	rando	serious	not serious	not serious	very serious	none	56	33	-	MD <b>0 points</b>	ФООО	CRITICAL
	mised	i			c,g					(1.31 lower to 1.31	VERY	
	trials									higher)	LOW	

178 CI: Confidence interval; MD: Mean difference; MPR: Medication possession ratio; SMD: Standardised mean difference

### Explanations

179

- a. Limited information on effect of assignment to intervention and some concerns with measurement of the outcome.
- 181 b. Rated down due to I2 value of 74%.
- 182 c. Small sample, concerns with fragility.
- d. Critical concerns with confounding. Serious concerns with missing data.
- e. Some concerns with bias due to deviations from the intended interventions.
- 185 f. I2 value is 61%; however, rating down for imprecision accounts for the variability between study findings.
- 186 g. 95% CI cannot exclude the possibility of no effect.
- h. Rated down due to the I2 value of 95%.
- i. Some concerns with effect of assignment to intervention and measurement of outcome.

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### 213 Table 9. Evidence Profile for PICO 8

214 **Question**: Interactive technology compared to non-interactive technology for patients on an oral anti-cancer medication regimen

### 215 **Setting**: Outpatient

trials

mail pubpe	Certainty assessment							f patients	Efi			
line, reprint, adapt, or reuse, please e	Study s design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations		non-interactive technology	Relative (95% CI)	Absolute (95% CI)	Certainty	Importance
Adher	ence (follo	ow up: 8	weeks; assesse	ed with: only a	adherence ra	te ≥80%)						
1 <sup>1</sup>	rando	very	not serious	not serious	very serious	none	56/79	33/40 (82.5%)	RR 0.86	116 fewer per	ФОО	CRITICAL
ng Society. For	mised	seriou			b,c		(70.9%)		(0.70 to 1.05)	1,000	0	

### Cancer related morbidity - Exit symptom severity (follow up: 8 weeks; assessed with: Symptom Experience Inventory range 0-150; higher = worse)

$1^1$ rand	do seriou	not serious	not serious	very serious	none	79	40	-	MD 4.12 points	$\oplus$	CRITICAL
mise	ed s <sup>d</sup>			b,e					higher	0	
trial	ls								(0.4 lower to 8.64	VERY	
יישייניט חס ס									higher)	LOW	

**VERY** 

LOW

(from 248 fewer

to 41 more)

**CRITICAL** 

**CRITICAL** 

### **Table 10. Evidence Profile for PICO 9**

Question: Structured oral anti-cancer medication program compared to no structured oral anti-cancer medication program for institutions providing care to patients on an oral anti-cancer medication regimen

**Setting**: Outpatient

observat very

not serious

serious <sup>d</sup>

not serious

none

e, please e			Certainty as	ssessment			Nº of p	patients	E	ffect				
No of studies	Study design	bias				ions	structured oral anti-cancer medication program	no structured oral anti-cancer medication program	Relative (95% CI)	Absolute (95% CI)	Certainty	Importance		
Adheren	dherence rate (follow up: 6 cycles; assessed with: medication event monitoring system)													
2 1,2	observat	very	not serious	not serious	serious <sup>b</sup>	none	18	29	-	MD <b>12.22</b> %	$\oplus$	CRITICAL		
y the Oncold	ional	serio								higher	0			
Copyright 2024 by the Or	studies	us <sup>a</sup>								(9.19 higher	VERY			
e only. Copy										to 15.24	LOW			
gle-user licens										higher)				
Adheren	ce rate (fo	llow u	p: 6 months - e	nd of treatme	nt; assessed	with: medica	ation possession rati	o)		1				

12536

31123

**CRITICAL** 

 $\Theta$ 

MD 6 %

rights.														
erves all r	ional	serio								higher	0			
org. ONS res	studies	us <sup>c</sup>								(4 higher to	VERY			
ilssions@ons.										8 higher)	LOW			
Adheren	Adherence (follow up: end of treatment; assessed with: pill counting)													
1 <sup>7</sup>	observat	very	not serious	serious <sup>d</sup>	very serious	none	87/100 (87.0%)	38/50 (76.0%)	RR 1.14	106 more	ФОО	CRITICAL		
apt, or reuse,	ional	serio			b,f				(0.96 to	per 1,000	0			
e, reprint, ada	studies	us <sup>e</sup>							1.36)	(from 30	VERY			
to post online										fewer to 274	LOW			
r permission t										more)				
Cancer-r	elated mo	rbidity	- Physical func	tioning (follow	w up: 1 year;	assessed wi	th: EORTC QoL physic	cal function; higher = I	better; MID	6 points; Scal	e from: 0 to	o 100)		
1 8	observat	very	not serious	serious <sup>g</sup>	serious <sup>b</sup>	none	56	56	-	MD <b>11.1</b>	ФОО	CRITICAL		
/ the Oncolog	ional	serio								points	0			
right 2024 by	studies	us <sup>e</sup>								higher	VERY			
se only. Copy										(7.45 higher	LOW			
e-user licens										to 14.75				
7-2024. Singl										higher)				

rights.													
erves all rights.	18	observat	very	not serious	not serious	serious <sup>b</sup>	none	56	56	-	MD <b>15.7</b>	$\oplus$	CRITICAL
org. ONS res		ional	serio								points	0	
sions@ons.c		studies	us <sup>e</sup>								higher	VERY	
I pubpermis											(12.7 higher	LOW	
please ema											to 18.7		
apt, or reuse,											higher)		
Page sgr	tient s	atisfactio	n (follo	w up: once dur	ing or after tr	eatment; asse	essed with:	telephone survey)					
oost on ne	1 <sup>9</sup>	observat	very	not serious	not serious	serious <sup>b</sup>	none	20/20 (100.0%)	15/20 (75.0%)	RR 1.32	240 more	ФОО	CRITICAL
rmission to p		ional	serio							(1.02 to	per 1,000	0	
ciety. For pe		studies	us <sup>h</sup>							1.72)	(from 15	VERY	
y Nursing So											more to 540	LOW	
the Oncolog											more)		
right 2024 by	tient f	inancial to	oxicity	(follow up: 1 ye	ear; assessed v	with: EORTC f	inancial diff	iculties; higher = wor	rse; Scale from: 0 to 10	00)			
nly. Copyr	18	observat	very	not serious	not serious	very serious	none	56	56	-	MD <b>0</b>	ФОО	CRITICAL
user license o		ional	serio			b,f					(1.57 lower	0	
024. Single-t		studies	us <sup>e</sup>								to 1.57	VERY	
d on 05-20-2											higher)	LOW	
nloade													

Time to	obtain me	dicatio	n - not reporte	d										
org. ONS	-	-	-	-	-	-	-	-	-	-	-	CRITICAL		
OCM mo	©CM model/value-based care - not reported													
mail pubper	-	-	-	-	-	-	-	-	1	-	1	CRITICAL		

CI: Confidence interval; MD: Mean difference; RR: Risk ratio

### **Explanations**

- a. Critical concerns with confounding and missing data. Moderate concern with measurement of outcome.
- b. Small sample, concerns with fragility.
- c. Critical concerns with confounding. Moderate concerns with selection of participants.
- d. Indirect measure of adherence.
- e. Critical concerns with confounding.
- f. The 95% CI cannot exclude the potential for no difference.
- g. Indirect measure of morbidity.
- h. Critical concerns with confounding. Serious concerns with selection of participants.

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