SUPPLEMENTARY APPENDIX 5: ACR/NPF 2018 Psoriatic Arthritis Guideline Evidence Report/Summary

Introduction

Critical outcomes:

- Each table reports the summary of findings (SoF) from randomized trials reporting the critical outcomes. The critical outcomes were chosen at the scoping meeting in September 2016. For nearly all comparisons, the critical outcomes were percentage of patients achieving American College of Rheumatology criteria for 20% improvement (ACR20), Health Assessment Questionnaire Disability Index (HAQ-DI) score, percentage of patients with a 75% reduction in the psoriatic arthritis severity index score (PASI75) and the adverse effect of treatments, including serious infections. If studies did not separately report serious infections, then total infections were included in tables, but not considered a critical outcome. In a few instances, infections were one of the outcomes used in indirect comparisons as described under Evidence Summaries below. A few other critical outcomes were included for individual comparisons.
- It is important to note that serious infections are very rare (infections in general are uncommon/rare), and thus it is quite difficult to achieve a statistically significant difference between groups for this outcome in RCTs powered for efficacy outcomes.
- ACR20 and PASI75 are proportions; The HAQ-DI is typically a continuous outcome but was reported as a binary outcome in some studies (HAQ-DI MCID; % patients who achieved a minimum clinically important difference [0.35 for the HAQ-DI in PsA, but some studies used 0.3 as the minimum change]). Data from studies that reported only continuous HAQ-DI was converted to a binary outcome using the mean change and SD in each group to estimate the proportion of patients achieving a minimal clinically important difference in each group (this method assumes the change scores are normally distributed).
- Not every study identified examined all critical outcomes. Each outcome was analyzed separately.

Therapy groups:

- At our scoping meeting in Sept 2016, we decided upon the following therapy groups:
 - o Oral small molecules: this includes methotrexate, sulfasalazine, cyclosporine, leflunomide and apremilast
 - o TNF inhibitors (TNFi)
 - o IL12/23 inhibitors (IL 12/23i)
 - o IL17 inhibitors (IL17i)
 - o Abatacept
 - o Tofacitinib

Systematic Literature Review

• For most of the outcomes, only randomized controlled trials (RCTs) were included. In some cases, observational studies were examined if relevant RCTs did not exist.

Quality Assessment

- Quality assessment was performed separately for each outcome using the Grading of Recommendations Assessment, Development, and Evaluation (GRADE) system, which results in one of four possible evidence categories that reflect level of confidence in the effect estimate: high, moderate, low, and very low.
- Study design is the starting point for quality assessment: randomized controlled trials (RCTs) start at high quality and observational studies start at low quality.
- Five factors can lower the quality of evidence: risk of bias, inconsistency, indirectness, imprecision, and publication bias.
 - o <u>Risk of bias</u> refers to limitations in study design or execution (e.g. lack of allocation concealment or blinding).
 - o <u>Inconsistency</u> refers to unexplained heterogeneity in results of studies evaluating the same outcome.
 - <u>Indirectness</u> refers to lack of direct comparisons of interventions of interest (e.g. studies comparing drug A vs. placebo and drug B vs. placebo when the comparison of interest is drug A vs. drug B), lack of applicability in the interventions or populations being evaluated, or use of indirect (surrogate) outcome measures.
 - Imprecision refers to uncertainty in the estimate of effect due to very low numbers of patients or events and/or wide 95% confidence intervals that cross a clinical decision threshold (i.e. between recommending and not recommending treatment).
 - <u>Publication bias</u> refers to selective publication of studies that show greater treatment effects (i.e. negative studies are suppressed).
- Three factors can increase the quality of evidence: large effect, dose response, and residual confounding that would reduce the observed treatment effect
 - Large effect refers to an RR >2 or <0.5
 - Dose response gradient refers to an increase in treatment effect or harms observed with increasing medication dose
 - All plausible residual confounding would reduce the demonstrated effect or increase the effect, if no effect was observed
- Quality of evidence can vary from outcome to outcome. The final quality assessment for the PICO question is based on the critical outcome with the lowest quality assessment.

Presentation of effects

- The treatment effects from binary (yes or no) outcomes are presented as relative effects and absolute effects.
- Relative effects capture the difference between intervention and control in relative terms. For example, a 10% event rate in controls

and a 5% event rate in the intervention represents a 50% relative risk reduction [((10- 5)*100)/10]

• The same difference represents a 5% absolute risk reduction (10% - 5% = 5%). In general, for patients, the absolute effect is the most important.

Evidence Summaries including Summary of Findings (= Tables under each PICO question, except some PICO questions for which no evidence was available)

- Indirect comparisons: For PICO questions with a large evidence base, network meta-analysis was used to generate risk ratios (RR) from indirect comparison of different drug classes. Network meta-analysis pools all studies for each comparison of drug A to placebo and studies of drug B to placebo and then divides the pooled odds ratio (OR) for A vs. placebo by the pooled OR for B vs placebo to arrive at an indirect comparison for drug A vs drug B. For PICO questions that had a smaller evidence base (fewer studies), we performed drug drug comparisons using the Bucher adjusted indirect comparison method, which is conceptually similar to the method used in the network meta-analyses.
- Direct comparisons are situations where trials directly compare drug A to drug B. There are very few trials with direct comparisons among patients with PsA but there are studies where direct comparisons are included for psoriasis and the PASI75 outcome.
- In the tables, when RR is specified, the first drug class (e.g. OSM vs. IL17i) is the reference drug class. Therefore a RR >1 for benefits indicates that the medication listed second is more beneficial; similarly a RR>1 for harms indicates the medication listed second is more harmful.

Interpreting the evidence

• It's important to take into account the information presented specifically as it relates to the question of interest. For example, PICO 22 asks whether switching to an OSM improves outcomes, but the trials reviewed address either adding a new OSM or examine a subgroup of patients in the placebo group with continuation of OSM vs. without continuation. Thus, this evidence is indirect and appropriately gets rated down for indirectness, as shown under the column labeled "indirectness." The quality of evidence takes these sorts of things into account, and is appropriately rated as high, moderate, low or very low. This quality of evidence is key to the voting decisions.

Moving from evidence to recommendations

- In GRADE, recommendations can be either strong or conditional. Generally, strong recommendations are restricted to high or moderate quality evidence. Low quality evidence almost invariably mandates a weak recommendation.
- There are, however, situations in which low quality evidence can lead to strong recommendations. For instance, if there is low quality evidence favoring an intervention but high quality evidence of important harm, the voting panel may make a strong recommendation against the intervention.

Bibliography of included studies

• A complete list of studies included as evidence for this report appears at the end of this document, following PICO 78. Shorter lists of studies included for each PICO question with an evidence base appear at the end of the summaries for each question.

Non-pharmacologic Interventions

PICO 1. In adult patients with active PsA, what are the benefits and harms of exercise compared to no exercise?

<u>Summary</u>: The initial literature searches did not identify any studies that addressed this PICO question. However, a systematic review (SR) and meta-analysis[1] of 14 RCTs in patients with RA found that aerobic exercises were significantly more effective than non-aerobic interventions in improving quality of life (Standardized mean difference [SMD] 0.39, p <0.0001), HAQ score (SMD 0.24, p <0.0009), and VAS pain score (SMD 0.31, p=0.02). Reduction in tender or swollen joint count did not show a statistically significant between-group difference (SMD 0.14, p=0.14). The duration of the trials ranged from 2 to 104 weeks. The average quality of the studies as assessed in the SR was moderate, and the indirectness of the population lowers the overall quality of evidence to low.

An updated literature search in March 2018 identified an additional relevant study, an RCT by Roger-Silva et al.[2] that compared resistance exercise to a waiting-list control in 41 patients with PsA. This trial had not been previously reviewed by the guideline panel. The exercise group showed significant improvement in HAQ-S and BASDAI scores compared to the control group at 12 weeks. This study was small, not blinded and at best, moderate quality, so it does not change the overall quality of evidence.

Author, year	Study type	Duration	Population Description	Treatment given to relevant population	Results
Baillet 2010	SR of 14 RCTs	Study duration ranged from 2 to 104 weeks	Patients with RA	Aerobic exercise interventions vs. non-aerobic exercise or usual care	Aerobic exercise significantly more effective than non- aerobic intervention for the following outcomes: Quality of life: SMD 0.39, 95% CI 0.23 to 0.56, p <0.0001 HAQ score: SMD 0.24, 95% CI 0.10 to 0.38, p <0.0009 VAS pain score: SMD 0.31, 95% CI 0.06 to 0.55, p=0.02 No statistically significant between-group difference for tender or swollen joint count: SMD 0.14, 95% CI -0.05 to 0.33 p=0.14

Quality of evidence across all critical outcomes: Low

SMD: standardized mean difference; SR: systematic review

References

- 1. Baillet A, Zeboulon N, Gossec L, Combescure C, Bodin LA, Juvin R, et al. Efficacy of cardiorespiratory aerobic exercise in rheumatoid arthritis: meta-analysis of randomized controlled trials Arthritis Care & Res 2010; 62(7): 984–992.
- 2. Roger-Silva D, Natour J. Moreira E, Jennings F. A resistance exercise program improves functional capacity of patients with psoriatic arthritis: a randomized controlled trial. Clin Rheumatol 2018; 37: 389-395.

PICO 2. In adult patients with active PsA, what are the benefits and harms of low impact exercise (e.g., tai chi, yoga, swimming) compared to high impact exercise (e.g., running)?

<u>Summary</u>: The literature searches did not identify any studies that addressed this PICO question. While there are existing systematic reviews of exercise therapy in other arthritis populations (RA, spondyloarthritis), none of these includes studies with a high-impact exercise (running) arm.

Quality of evidence across all critical outcomes: Very low

PICO 3. In adult patients with active PsA with active peripheral arthritis and/or enthesitis, what are the benefits and harms of physical therapy (PT) compared with no PT?

<u>Summary</u>: The literature searches did not identify any studies that directly addressed this PICO question, or any systematic reviews that specifically evaluated PT in patients with other arthritic conditions (e.g. RA). However, PT interventions often overlap with exercise and occupational therapy interventions, so the evidence from PICO 1 and PICO 4 may be applicable to this PICO question. The evidence from both PICO 1 and 4 is noted as having serious risk of bias and serious indirectness, and since none of those interventions were specifically tailored to PT (they overlap with PT), that increases the indirectness to very serious. Therefore, the overall quality of evidence is very low.

Quality of evidence across all critical outcomes: Very low

PICO 4. In adult patients with active PsA with active peripheral arthritis and/or enthesitis, what are the benefits and harms of occupational therapy (OT) compared with no OT?

<u>Summary</u>: The literature searches did not identify any studies that directly addressed this PICO question. Two systematic reviews evaluated OT interventions in patients with RA. Siegel et al.[1] reviewed evidence from earlier SRs and individual RCTs for a range of interventions associated with OT, including various exercise interventions and psychoeducational interventions. Since exercise interventions were covered under PICO 1 and 2, this PICO focuses on psychoeducational interventions. This SR did not report effect sizes but found that psychological interventions generally had at least small effects on pain and function in patients with RA. Knittle et al.[2] performed meta-analyses of 27 RCTs that compared the efficacy of psychological self-regulatory interventions (CBT, patient education, stress management) to controls in patients with RA. These findings appear in the table below; they indicate a small but significant benefit of psychological interventions for the outcomes disability and physical activity at 2 to 14 months follow-up, but pain at follow-up was not significantly different between groups (although immediately post-treatment there was a significant between-group difference in pain reduction favoring the interventions). The average risk of bias in the studies

in these SRs appears to be serious, and since all studies included patients with RA they are limited by serious indirectness. Therefore, the overall quality of evidence is low.

Quality of evidence across all critical outcomes: Low

Author, year	Study type	Duration	Population Description	Treatment given to relevant population	Results
Knittle	SR of	Study duration	Patients with	Psychological self-regulatory	Outcomes at follow-up (2-14 months post-treatment)
2010	27	ranged from 2	RA	intervention (CBT, patient	Pain: Hedges' $g = 0.127$ (p=0.069), no significant difference
	RCTs	weeks to 14		education, stress management)	Disability: Hedges' g = 0.145 (p=0.047)
		months		vs. control	Physical activity: Hedges' g = 0.361 (p=0.020)

SMD: standardized mean difference; SR: systematic review

References

- 1. Siegel S, Tencza M, Apodaca B, Poole J. Effectiveness of occupational therapy interventions for adults with rheumatoid arthritis: a systematic review. Am J Occ Ther. 2017; 71:1-11.
- 2. Knittle K, Maes S, De Gucht V. Psychological interventions for rheumatoid arthritis: examining the role of self-regulation with a systematic review and meta-analysis of randomized controlled trials. Arth Care & Res. 2010; 62:1460-72.

PICO 5. In adult patients with active PsA who are overweight (e.g., BMI 25 and over), what are the benefits and harms of weight loss compared with no weight loss?

Summary: This PICO question was addressed by direct comparisons in three studies [1-3] but indirect populations in two studies.[2,3] One study compared successful weight loss with unsuccessful weight loss in overweight PsA patients who followed a hypocaloric diet or a free-managed diet. Successful weight loss was defined as ≥ 5% weight loss.[1] Two studies evaluated the effect of weight loss in overweight psoriasis patients following a low-calorie diet.[2,3] Statistically significant differences were reported favoring weight loss over no or unsuccessful weight loss for efficacy outcomes (minimal disease activity [MDA], PASI75).

Quality of evidence across all critical outcomes: Low

V	Weight loss compared to no or unsuccessful weight loss for overweight PsA patients Bibliography: PICO 5: Weight loss versus no weight loss for overweight PsA patients.													
	Quality assessment								Summary of findings					
Nº of participants (studies)	Risk of	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of	Study event ra (%)	ates	Relative effect	Anticipated absolute effects				
Follow-up	Dias					evidence	With no or unsuccessful weight loss	With Weight Ioss	CI)	Risk with no or unsuccessful weight loss	Risk difference with Weight loss			
MDA, 24 v	weeks													
126 (1 RCT)	serious ^a	not serious	not serious	serious ^e	none	⊕⊕⊖⊖ Low	12/52 (23.1%)	37/74 (50.0%)	RR 2.17 (1.26 to 3.74) Favors weight loss	231 per 1,000 (0.231)	270 more per 1,000 (0.270) (60 more to 632 more)			
PASI 75 ,	PASI 75, 24 weeks													

Weight loss compared to no or unsuccessful weight loss for overweight PsA patients Bibliography: PICO 5: Weight loss versus no weight loss for overweight PsA patients.

Quality assessment						Summary of findings					
323 (2 RCTs)	serious ^b	not serious	serious °	not serious	none	⊕⊕⊖⊖ Low	86/162 (53.1%)	132/161 (82.0%)	RR 1.67 (1.09 to 2.54) Favors weight loss	531 per 1,000 (0.531)	299 more per 1,000 (0.299) (212 more to 361 more) ^d

CI: Confidence interval; MDA: Minimal disease activity; RR: Risk ratio

a. Patients/providers not blinded.

b. Randomization and allocation concealment methods not described; no blinding of patients/providers/assessors (1 study, Al-Mutairi 2014)

c. Population is indirect (psoriasis patients)

d. Absolute risk difference and confidence interval calculated based on the odds ratio.

e. Although the effect size is large, the study does not meet optimal information size.

Notes: Limitations described in 1 study[1] included 20% of patients receiving chronic treatment with oral hypoglycemic agents, possible selection bias resulting in a high prevalence of axial involvement in study population and low baseline PASI scores.

Working Group quality of evidence categories:

High quality: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate quality: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low quality: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect Very low quality: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

References:

- 1. Di Minno MN, Peluso R, Iervolino S, Russolillo A, Lupoli R, Scarpa R, et al. Weight loss and achievement of minimal disease activity in patients with psoriatic arthritis starting treatment with tumour necrosis factor alpha blockers. Ann Rheum Dis. 2014;73(6):1157-1162.
- 2. Gisondi P, Del Giglio M, Di Francesco V, Zamboni M, Girolomoni G. Weight loss improves the response of obese patients with moderateto-severe chronic plaque psoriasis to low-dose cyclosporine therapy: a randomized, controlled, investigator-blinded clinical trial. Am J Clin Nutr. 2008;88(5):1242-1247.
- 3. Al-Mutairi N, Nour T. The effect of weight reduction on treatment outcomes in obese patients with psoriasis on biologic therapy: a randomized controlled prospective trial. Expert Opin Biol Ther. 2014;14(6):749-756.

PICO 6. In adult patients with active PsA <u>who smoke</u>, what are the benefits and harms of smoking cessation compared with no smoking cessation?

Summary: The literature searches did not identify any studies that addressed this PICO question. Although specific benefits for PsA or arthritis symptoms are unclear, the general health benefits for smoking cessation are well established. A large RCT in a non-arthritic population found a significant mortality reduction at 14.5 years of follow-up for 5887 patients with asymptomatic airway obstruction who received a smoking cessation intervention compared to a usual care group (hazard ratio [HR] for mortality in usual care group 1.18, 95% CI 1.02 to 1.37).[1] A systematic review of 25 large U.S. and European cohort studies with 503,905 participants ≥60 years of age compared cardiovascular mortality of never smokers, former smokers, and current smokers. With never smokers as the reference group, the individual patient meta-analysis found that former smokers had a lower risk of cardiovascular mortality (HR 1.37, 95% CI 1.25 to 1.49) than current smokers (HR 2.07, 95% CI 1.82 to 2.36).[2] For both studies the only downgrade is for indirectness of the population.

Author, year	Study type	Duration	Population Description	Treatment given to relevant population	Results
Anthonisen 2005	RCT	14.5 years	5887 patients with asymptomatic airway obstruction	10-week smoking cessation intervention (with ipratropium or placebo inhaler) vs. usual care	Mortality: Hazard ratio (HR) 1.18 (95% CI 1.02 to 1.37) for usual care group compared to smoking cessation intervention group.
Mons 2015	SR of 25 cohort studies	Mean follow- up across 25 cohort studies ranged from 1.6 to 15.4 years.	503,905 participants ≥60 years of age from 25 cohort studies	Not applicable. An individual patient meta-analysis compared cardiovascular mortality for never smokers, former smokers and current smokers.	With never smokers as the reference group, the individual patient meta-analysis found that former smokers had a lower risk of cardiovascular mortality (HR 1.37, 95% CI 1.25 to 1.49) than current smokers (HR 2.07, 95% CI 1.82 to 2.36).

Quality of evidence across all critical outcomes: Moderate

SR: systematic review

References

- Anthonisen NR, Skeans MA, Wise RA, Manfreda J, Kanner RE, Connett JE; Lung Health Study Research Group. The effects of a smoking cessation intervention on 14.5-year mortality: a randomized clinical trial. Arthritis Care & Res 2010; 62(7): 984–992. Ann Intern Med. 2005 Feb 15;142(4):233-9.
- 2. Mons U, Müezzinler A, Gellert C, Schöttker B, Abnet CC, Bobak M, et al. Impact of smoking and smoking cessation on cardiovascular events and mortality among older adults: meta-analysis of individual participant data from prospective cohort studies of the CHANCES consortium. BMJ. 2015 Apr 20;350:h1551.

PICO 7. In adult patients with active PsA, what are the benefits and harms of massage therapy compared with no massage therapy?

Summary: The literature searches did not identify any direct studies that addressed this PICO question. One systematic review, involving seven RCTs with total 352 patients, assessed massage therapy (MT) in patients with OA and RA. The outcomes reported are improved range of motion (ROM), WOMAC functional subscale, grip strength in individuals with hand arthritis, walking function among those with OA of the knee, and adverse events. Five studies (310 participants) provided very low-level evidence that MT is superior to non-active therapy for improving range of motion (ROM). Three RCTs involving 233 participants, provided moderate-quality evidence that MT is superior to non-active therapies in improving WOMAC functional subscales. One study (22 participants provided low-quality evidence that MT was superior to a non-active therapy for improving perceived grip strength in individuals with hand arthritis. Two RCTs with a low risk of bias, and one RCT with a high risk of bias, involving 233 participants, provided moderate quality evidence that MT is superior for improving walking function among those with OA of the knee. One study reported significantly faster 50-foot walk times among those receiving MT compared with usual care control participants, whereas another reported decreased time to walk 8 feet in MT recipients compared with participants in a waitlist control group. Adverse effects were reported in two studies: one reported no adverse effects related to the MT intervention, and another reported that one participant experienced an increase in discomfort and subsequently dropped out of the trial.

Author,	Study type	Duration	Population	Treatment given to relevant	Results
year			Description	population	
Nelson L,	Systematic	Seven	352	Massage therapy. The total	Five studies (310 participants) provided very low-level evidence
2017	review	RCTs	participants	minutes of massage exposure	(downgraded because of risk of bias, imprecision, and
		between	with either	for the trial period ranged from	inconsistency) that MT is superior to non-active therapy for
		1997 and	OA or RA	120 to 960 mins.	improving range of motion (ROM). Two trials with a low risk of
		2015			bias and one trial with a high risk of bias RCT involving 233
					participants, provided moderate-quality evidence that MT is
					superior to nonactive therapies in improving WOMAC functional
					subscales. One study (22 participants) with a high risk of bias
					provided low-quality evidence (downgraded because of risk of
					bias and imprecision) that MT was superior to a nonactive
					therapy for improving perceived grip strength in individuals
					with hand arthritis. Two RCTs with a low risk of bias, and one
					RCT with a high risk of bias, involving 233 participants,
					provided moderate guality evidence (downgraded because of
					imprecision) that MT is superior to a nonactive comparator for
					improving walking function among those with OA of the knee.
					One study reported significantly faster 50-foot walk times
					among those receiving MT compared with usual care control
					participants, whereas another reported decreased time to walk
					8 feet in MT recipients compared with participants in a wait list
					control group. Adverse effects were reported in two studies:
					one reported no adverse effects related to the MT intervention

Quality of evidence across all critical outcomes: Very low

Author,	Study type	Duration	Population	Treatment given to relevant	Results
year			Description	population	
					and another reported that one participant experienced an
					increase in discomfort and subsequently dropped out of the
					trial.

References:

1. Nelson L., Churilla J. Massage Therapy for Pain and Function in Patients with Arthritis. A Systematic Review of Randomized Controlled Trials. Am J Phys Med Rehabil 2017;00:00–00

PICO 8. In adult patients with active PsA, what are the benefits and harms of acupuncture compared with no acupuncture?

<u>Summary</u>: The literature searches did not identify any studies that directly addressed this PICO question. One systematic review addressed acupuncture in older osteoarthritis (OA) patients to measure pain intensity (VAS) and functional mobility.[1] This systematic review included 12 trials (1763 participants) comparing acupuncture to sham acupuncture, no treatment or usual care. Most trials have unclear (64%) or high (9%) risk of bias. Acupuncture use was associated with significant reductions in pain intensity (MD –0.29, 95% CI –0.55 to –0.02, I2 0%, 10 trials, 1699 participants), and functional mobility (standardized MD –0.34, 95% CI –0.55 to –0.14, I2 70%, 9 trials, 1543 participants).

Quality of evidence across all critical outcomes: Very low

Acup	Acupuncture Compared to No Acupuncture for Pain Reduction in Osteoarthritis Patients Bibliography: Acupuncture Compared to No acupuncture for Pain Reduction in Osteoarthritis Patients												
		Qu	ality Assess		Summary of Findings								
Number of participants	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence	Study ev rates (%	vent 6)	Relative effect	Anticipat absolute	ed effects		
(studies) Follow-up							With No acupuncture	With Acupuncture	(95% CI)	Risk with No acupuncture	Risk difference with Acupuncture		
VAS pain													
1699 (10 RCTs)	serious ^a	not serious	very serious ^b	not serious	none	⊕○○○ VERY LOW	829	870	-	-	MD 0.29 lower (0.55 lower to 0.22 lower)		
Functional	mobilit	ty											
1543 (9 RCTs)	serious ^a	serious ^c	very serious ^b	not serious	none	⊕⊖⊖⊖ VERY LOW	792	751	-	-	SMD 0.34 SD lower (0.55 lower to 0.14 lower)		

CI: Confidence interval; MD: Mean difference; SMD: Standardized mean difference

Explanations

a. Authors of systematic review assigned the risk of bias as serious

b. Indirect population and outcomes

c. High Chi-squared and I-squared values

References:

1. Manyanga T. et al. Pain management with acupuncture in osteoarthritis: a systematic review and meta-analysis. BMC Complementary and Alternative Medicine 2014, 14:312

Pharmacologic Interventions

PICO 9. In adult patients with active PsA who are treatment-naïve, what are the benefits and harms of MTX vs. NSAID?

Summary: This PICO question was addressed directly by one observational study.[1] This retrospective matched study compared NSAIDS with oral MTX in 38 PsA patients. The intervention group were administered methotrexate (MTX) at a maximum weekly dose of 15-20 mg. Controls, who were matched by damage, actively inflamed joints, gender, and disease duration, received NSAIDS. No statistically significant difference was reported for one efficacy outcome (≥40% improvement in actively inflamed joints) or adverse events (GI side effects, hepatic AEs) at 104 weeks, although there was imprecision in the efficacy estimate and very serious imprecision in the adverse event estimates.

Quality of evidence across all critical outcomes: Very low

	MTX compared to NSAID for treatment-naive PsA patients Bibliography: PICO 9: MTX versus NSAID for treatment-naive PsA patients.													
	Quality assessment								Summary of findings					
Nº of participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality	Study ever rates (%)	ent)	Relative effect	Anticipated absolute effects				
(studies) Follow-up						evidence	With MTX	With NSAI D	(95% CT)	Risk with MTX	Risk difference with NSAID			
>/= 40%	improv	vement in a	ctively infl	amed joint	ts, 104 we	eeks								
38 (1 observational study)	very serious ª	not serious	not serious	serious ^b	none		9/19 (47.4%)	10/19 (52.6%)	RD 0.05 (-0.26 to 0.37)	474 per 1,000 (0.474)	50 more per 1,000 (0.050) (260 fewer to 370 more)			
GI side effects (not described), 104 weeks														

	MTX compared to NSAID for treatment-naive PsA patients Bibliography: PICO 9: MTX versus NSAID for treatment-naive PsA patients.												
Quality assessment								Summary of findings					
42 (1 observational study)	very serious ª	not serious	not serious	serious ^b	none	⊕⊖⊖⊖ VERY LOW	2/23 (8.7%)	0/19 (0.0%)	RR 0.24 (0.01 to 4.71)	87 per 1,000	66 fewer per 1,000 (0.066) (86 fewer to 323 more)		
Hepatic a	dverse	events (no	ot describe	d), 104 we	eks								
42 (1 observational study)	very serious ª	not serious	not serious	serious ^b	none	⊕⊖⊖⊖ VERY LOW	2/23 (8.7%)	0/19 (0.0%)	RR 0.24 (0.01 to 4.71)	87 per 1,000	66 fewer per 1,000 (0.066) (86 fewer to 323 more)		

CI: Confidence interval; RD: Risk difference

a. Retrospective non-randomized design, no blinding, only 60% of MTX arm at follow-up

b. Small study with few patients and events and wide CI.

Working Group quality of evidence categories:

High quality: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate quality: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low quality: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect Very low quality: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

References:

1. Abu-Shakra M, Gladman DD, Thorne JC, Long J, Gough J, Farewell VT. Longterm methotrexate therapy in psoriatic arthritis: clinical and radiological outcome. J Rheumatol. 1995;22(2):241-245.

PICO 10. In adult patients with active PsA who are treatment-naïve, what are the benefits and harms of an Oral Small Molecule (OSM) vs. TNFi?

<u>Summary:</u> This question was addressed by direct comparisons in 5 RCTs[1-5], and by indirect comparisons in 7 RCTs (in 9 publications).[6-14] Indirect populations (psoriasis patients) were evaluated in 2 studies.[4,5] Table 1 shows a comparison of MTX and infliximab (IFX) combination therapy with MTX monotherapy. Table 2 shows an indirect comparison of MTX and TNFi (adalimumab and etanercept) efficacy outcomes (ACR20, HAQ-DI) from placebo-controlled RCTs. Table 3 shows additional efficacy outcomes and adverse events from RCTs comparing OSMs (MTX or sulfasalazine) to placebo. Table 4 shows additional efficacy outcomes and adverse events from RCTs comparing TNFis (adalimumab or etanercept) to placebo. Lastly, Table 5 shows outcomes from two RCTs comparing TNFi (adalimumab or IFX) with MTX in psoriasis patients[4,5] The results from each table are summarized in the following paragraphs.

One RCT comparing MTX and IFX combination therapy (IFX given in 5 mg/kg infusions at weeks 0, 2, 6 and 14) with MTX monotherapy[1] reported statistically significant differences favoring combination therapy over monotherapy for all efficacy outcomes (ACR20, HAQ-DI, PASI75), but no statistically significant difference for adverse events (pulmonary tuberculosis, upper abdominal pain)(Table 1). Two observational studies comparing MTX with TNFi reported statistically significant differences favoring TNFi for some efficacy outcomes (absolute swollen joint count, modified HAQ-DI, absolute PASI score) but no significant difference for other efficacy outcomes (decrease in swollen joint count, absolute tender joint count, absolute HAQ score) and adverse events (liver toxicity, GI intolerance)(data not shown).[2,3]

One RCT of MTX versus placebo and three RCTs of TNFi (adalimumab or etanercept) vs. placebo reported common outcomes (ACR20, HAQ DI) that could be indirectly compared using the Bucher adjusted indirect comparison method (Table 2). TNFi showed significantly greater benefit than MTX for both outcomes. The MTX trial had a higher percentage of OSM-naïve patients (80%) compared to the TNFi trials (50-60%).

Four RCTs comparing OSM with placebo[6-9] reported no statistically significant difference for all efficacy outcomes (absolute tender joint count, absolute swollen joint count, change in tender joint count). Statistically significant differences favoring placebo were reported for one adverse event (GI intolerance). However, no statistically significant difference was reported between OSM and placebo for another adverse event (liver toxicity). A subgroup analyses by drug indicated a statistically significant difference favoring placebo over methotrexate for liver toxicity, but no statistically significant difference on placebo (Table 3).

Three RCTs (5 publications)[10-14] comparing TNFi with placebo reported statistically significant differences favoring TNFi over placebo for all efficacy outcomes (ACR20, HAQ-DI, PASI75), but no statistically significant difference for all adverse events (serious AEs, serious infections, upper respiratory tract infections)(Table 4).

Lastly, two RCTs comparing TNFi (adalimumab or IFX) with MTX in psoriasis patients[4,5] reported a statistically significant difference favoring TNFi over MTX for one skin outcome (measured by PASI75), but no statistically significant difference for two adverse outcomes (liver function test abnormality, serious infection)(Table 5).

Quality of evidence across all critical outcomes: Low

	Table 1. MTX/IFX compared to MTX for treatment-naive PsA patients Bibliography: PICO 10: OSM versus TNFi for treatment-naive PsA patients.											
		Qua	ality assessr	nent				Sum	mary of fi	ndings		
№ of participants	Risk of	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of	Study ev (%)	ent rates	Relative effect	Anticipat effects	ed absolute	
Follow-up	Dias					evidence	With MTX	With MTX/IFX	(93% CI)	Risk with MTX	Risk difference with MTX/IFX	
ACR20 response, 16 weeks												
99 (1 RCT)	serious ª	not serious	not serious	not serious	none	⊕⊕⊕⊖ MODERATE	32/48 (66.7%)	44/51 (86.3%)	RR 1.29 (1.03 to 1.63) Favors MTX/IFX	667 per 1,000 (0.667)	196 more per 1,000 (0.196) (32 more to 278 more) ^d	
HAQ-DI ^b ,	16 we	eks								•		
110 (1 RCT)	serious ª	not serious	not serious	not serious	none	⊕⊕⊕⊖ MODERATE	33/54 (61.1%)	45/56 (80.4%)	RR 1.31 (1.03 to 1.69) Favors MTX/IFX	611 per 1,000 (0.611)	192 more per 1,000 (0.192) (24 more to 295 more) ^d	
PASI75, 1	PASI75, 16 weeks											

	Table 1. MTX/IFX compared to MTX for treatment-naive PsA patients Bibliography: PICO 10: OSM versus TNFi for treatment-naive PsA patients.												
	Quality assessment								Summary of findings				
69 (1 RCT)	serious ª	not serious	not serious	not serious	none	⊕⊕⊕⊖ MODERATE	19/35 (54.3%)	33/34 (97.1%)	RR 1.79 (1.31 to 2.44) Favors MTX/IFX	543 per 1,000 (0.543)	428 more per 1,000 (0.428) (259 more to 453 more) ^d		
Pulmonai	ry tube	erculosis ^e	•					1	1				
111 (1 RCT)	serious ª	not serious	not serious	serious ^c	none	⊕⊕⊖⊖ Low	1/57 (1.8%) MTX/IFX	0/54 (0.0%) MTX	RR 0.35 (0.01 to 8.45)	18 per 1,000 MTX/IFX	12 fewer per 1,000 with MTX (0.012) (18 fewer to 134 more)		
Upper ab	Upper abdominal pain												
111 (1 RCT)	serious ª	not serious	not serious	serious ^c	none	⊕⊕⊖⊖ Low	3/54 (5.6%)	0/57 (0.0%)	RR 0.14 (0.01 to 2.56)	56 per 1,000 (0.056)	48 fewer per 1,000 (0.048) (55 fewer to 143 more)		

CI: Confidence interval; RD: Risk difference; RR: Risk ratio

a. Unclear randomization and allocation concealment methods, no blinding

b. HAQ-DI measured as proportion of patients who achieved minimum clinically important improvement (>0.3 or 0.35 in different studies).

c. Single small study with very few events

d. Absolute risk difference and confidence interval calculated based on the odds ratio.

e. For pulmonary tuberculosis there were zero events in the MTX group, so we used MTX/IFX as the denominator in order to calculate the RR. So in this comparison "12 fewer per 1,000" means 12 fewer events in the MTX group compared to MTX/IFX.

	Table 2. MTX compared to TNFi for treatment-naive PsA patients Bibliography: PICO 10: OSM versus TNFi for treatment-naive PsA patients.												
		Qua	ality assessr	ment			Summary of findings						
Nº of participants	Risk of	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of			Relative effect	Anticipated absolute effects			
(studies) Follow-up	DIAS					evidence	With TNFi	With MTX	(95% CI)	Risk with TNFi	Risk difference with MTX		
ACR20 re	spons	e, Bucher ad	djusted ind	lirect com	parison			·					
799 (4 RCTs)	not serious	not serious	serious ^a	not serious	none	⊕⊕⊕⊖ MODERATE	TNFi	МТХ	OR ^b 0.24 (0.07 to 0.72) Favors TNFi	596 per 1,000 (0.596)	335 fewer per 1,000 (0.335) (502 fewer to 58 fewer) ^d		
HAQ-DI [°] ,	Buche	er adjusted	indirect co	mparison					·				
799 (4 RCTs)	not serious	not serious	serious ^a	not serious	none	⊕⊕⊕⊖ MODERATE	TNFi	МТХ	RR 0.63 (0.42 to 0.95) Favors TNFi	596 per 1,000 (0.596)	221 fewer per 1,000 (0.221) (346 fewer to 30 fewer)		

CI: Confidence interval; **RR**: Risk ratio

a. Indirect comparison of placebo-controlled trials
b. Study of MTX reported finding as OR, so indirect comparison had to use OR.
c. HAQ-DI measured as proportion of patients who achieved minimum clinically important improvement (>0.3 or 0.35 in different studies).
d. Absolute risk difference and confidence interval calculated based on the odds ratio.

Table 3. OSM compared to placebo for treatment-naive PsA patients Bibliography: PICO 10: OSM versus TNFi for treatment-naive PsA patients.												
		Qua	ality assessr	nent			Summary of findings					
Nº of participants	Risk of	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of	Study ev (%)	ent rates	Relative effect	Anticipated absolute effects		
Follow-up	DIAS					evidence	With placebo	With OSM	(95% CI)	Risk with placebo	Risk difference with OSM	
Absolute tender joint count (8-12 weeks)												
245 (2 RCTs)	not serious	not serious	serious ^a	serious ^b	none	⊕⊕⊖⊖ Low	126	119	-	-	MD 3.31 lower (9.11 lower to 2.49 higher)	
Absolute	swolle	en joint cou	nt (8-12 w	eeks)								
245 (2 RCTs)	not serious	not serious	serious ^a	serious ^b	none	⊕⊕⊖⊖ Low	126	119	-	-	MD 1.36 lower (5.47 lower to 2.74 higher)	
Change ir	n tend	er joint cou	nt (53 join	ts, week 2	4)							
108 (1 RCT)	not serious	not serious	serious ^a	serious ^b	none	⊕⊕⊖⊖ Low	59	49	-	-	MD 0.8 lower (2.17 lower to 0.57 higher)	
GI intoler	ance	(nausea, vo	miting, abo	dominal pa	ain, diarrh	ea)						

	Table 3. OSM compared to placebo for treatment-naive PsA patients Bibliography: PICO 10: OSM versus TNFi for treatment-naive PsA patients.												
		Qua	ality assessi	ment			Summary of findings						
392 (4 RCTs)	not serious	not serious	serious ^a	not serious	none	⊕⊕⊕⊖ MODERATE	30/205 (14.6%)	67/187 (35.8%)	RR 2.38 (1.64 to 3.46) Favors placebo	146 per 1,000 (0.146)	202 more per 1,000 (0.202) (94 more to 360 more)		
Liver toxi	city ^c -	Methotrexa	ite vs place	ebo									
221 (1 RCT)	not serious	not serious	serious ^a	not serious	none	⊕⊕⊕⊖ MODERATE	2/112 (1.8%)	12/109 (11.0%)	RD 0.09 (0.03 to 0.16) Favors placebo	18 per 1,000 (0.018)	90 more per 1,000 (0.090) (30 more to 160 more)		
Liver toxi	city ^d -	Sulfasalazi	ne vs place	ebo									
117 (1 RCT)	not serious	not serious	serious ^a	serious ^b	none	⊕⊕⊖⊖ Low	1/64 (1.6%)	0/53 (0.0%)	RR 0.40 (0.02 to 9.65)	16 per 1,000 (0.016)	9 fewer per 1,000 (0.009) (15 fewer to 135 more)		

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

a. Indirect comparison

b. Wide 95% CI crosses line of no difference

c. Abnormal liver function tests, but study did not define cutoff.

d. Increased ASAT and ALAT (x 3 compared to baseline) after 1 month. No concomitant NSAID.

Table 4. TNFi compared to Placebo for treatment-naive PsA patients

Bibliography: PICO 10: OSM versus TNFi for treatment-naive PsA patients.

	Quality assessment							Summary of findings				
Nº of participants	Risk of	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of	Study ev (%)	ent rates	Relative effect	Anticipat effects	ed absolute	
(studies) Follow-up	DIAS					evidence	With Placebo	With TNFi	(95% CI)	Risk with Placebo	Risk difference with TNFi	
PASI 75												
326 (3 RCTs)	not serious	not serious	serious ^a	not serious	none	⊕⊕⊕⊖ MODERATE	11/161 (6.8%)	80/165 (48.5%)	RD 0.37 (0.09 to 0.65) Favors TNFi	68 per 1,000 (0.068)	370 more per 1,000 (0.370) (90 more to 650 more)	
Serious I	nfecti	ons										
518 (2 RCTs)	not serious	not serious	serious ^a	not serious	none	⊕⊕⊕⊖ MODERATE	2/266 (0.8%)	1/252 (0.4%)	RD -0.004 (-0.02 to 0.01)	8 per 1,000 (0.008)	4 fewer per 1,000 (0.004) (20 fewer to 10 more)	
Upper res	spirato	bry tract inf	ection 12 v	weeks								
60 (1 RCT)	not serious	not serious	serious ^a	serious ^b	none	⊕⊕⊖⊖ Low	17/30 (56.7%)	17/30 (56.7%)	RR 1.00 (0.64 to 1.56)	567 per 1,000 (0.567)	0 fewer per 1,000 (204 fewer to 317 more)	

CI: Confidence interval; RR: Risk ratio

a. Indirect comparison

b. Wide CIs

	Table 5. TNFi compared to OSM for treatment-naive psoriasis patients Bibliography: PICO 10: OSM versus TNFi for treatment-naive PsA patients.													
		Qua	ality assessi	ment			Summary of findings							
Nº of participants	Risk of	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of	Study even (%)	t rates	Relative effect	Anticipated absolute effects				
(studies) Follow-up	bias					evidence	With OSM (Psoriasis)	With TNFi	(95% CI)	Risk with OSM (Psoriasis)	Risk difference with TNFi			
PASI-75,	16 we	eeks												
1086 (2 RCTs)	serious ^a	not serious	serious ^b	not serious	none	⊕⊕⊖⊖ Low	129/325 (39.7%)	594/761 (78.1%)	RR 1.98 (1.66 to 2.36) Favors TNFi	397 per 1,000 (0.397)	389 more per 1,000 (0.389) (262 more to 540 more)			
Liver fun	ction t	test abnorm	ality ^c - Inf	liximab vs	. MTX									
860 (1 RCT)	serious d	not serious	serious ^b	not serious	none	⊕⊕⊖⊖ Low	3/211 (1.4%)	14/649 (2.2%)	RD 0.01 (-0.01 to 0.03)	14 per 1,000 (0.014)	10 more per 1,000 (0.010) (10 fewer to 30 more)			
Liver fun	ction t	est abnorm	ality ^e - Ada	alimumab	vs. MTX									
217 (1 RCT)	not serious	not serious	serious ^b	not serious	none	⊕⊕⊕⊖ MODERATE	10/110 (9.1%)	2/107 (1.9%)	RR 0.21 (0.05 to 0.92) Favors ADA	91 per 1,000 (0.091)	72 fewer per 1,000 (0.072) (86 fewer to 7 fewer)			

	Table 5. TNFi compared to OSM for treatment-naive psoriasis patients Bibliography: PICO 10: OSM versus TNFi for treatment-naive PsA patients.													
	Quality assessment Summary of findings													
Serious I	nfectio	on												
1077 (2 RCTs)	serious ª	not serious	serious ^b	not serious	none		4/321 (1.2%)	10/756 (1.3%)	RD 0.001 (-0.01 to 0.02)	12 per 1,000 (0.012)	1 more per 1,000 (0.001) (10 fewer to 20 more)			

CI: Confidence interval; RD: Risk difference; RR: Risk ratio

a. 1 RCT (Barker 2011) open label, ITT conducted

b. Indirect population (Psoriasis patients)

c. Elevated liver enzymes (study did not report cutoffs for abnormal test results)

d. Open label, ITT conducted

e. Alanine aminotransferase > 2.5 times the upper normal limit (ULN), aspartate aminotransferase > 2.5 times the ULN, total bilirubin > 1.5 times the ULN, or

ỹ- Gamma-Glutamyltransferase elevation

Working Group quality of evidence categories:

High quality: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate quality: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low quality: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect Very low quality: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

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PICO 11. In adult patients with active PsA who are treatment-naïve, what are the benefits and harms of an OSM vs. IL12/23?

<u>Summary</u>: The literature searches did not identify any studies that addressed this PICO question.

Quality of evidence across all critical outcomes: Very low

PICO 12. In adult patients with active PsA who are treatment-naïve, what are the benefits and harms of an OSM vs. IL17i?

<u>Summary</u>: The literature searches did not identify any studies that addressed this PICO question.

Quality of evidence across all critical outcomes: Very low

PICO 13. In adult patients with active PsA who are treatment-naïve, what are the benefits and harms of a TNFi vs. IL12/23i?

<u>Summary</u>: The literature searches did not identify any studies that addressed this PICO question.

Quality of evidence across all critical outcomes: Very low

PICO 14. In adult patients with active PsA who are treatment-naïve, what are the benefits and harms of TNFi vs. IL17i?

<u>Summary</u>: The literature searches did not identify any studies that addressed this PICO question.

Quality of evidence across all critical outcomes: Very low

PICO 15. In adult patients with active PsA who are treatment-naïve, what are the benefits and harms of IL12/23i vs. IL17i?

<u>Summary</u>: The literature searches did not identify any studies that addressed this PICO question.

Quality of evidence across all critical outcomes: Very low

PICO 16. In adult patients with active PsA despite treatment with OSM, what are the benefits and harms of switching to TNFi compared to switching to IL12/23i?

<u>Summary</u>: Thirteen placebo-controlled RCTs (16 publications) indirectly addressed this PICO question. Nine studies (12 publications) compared TNFi versus placebo in PsA patients.[1-12] Three studies compared IL 12/23 (ustekinumab) with placebo in PsA patients.[13-15] Lastly, one study compared etanercept with ustekinumab in psoriasis patients.[16] Statistically significant differences favoring TNFi over placebo and ustekinumab over placebo were reported for all efficacy outcomes (ACR20, HAQ-DI, PASI-75, data not shown). No statistically significant differences with placebo occurred for any adverse events (see Table 3 for serious infections). In addition, both TNFi-naïve and TNFi-exposed patients responded similarly to TNFi and ustekinumab compared to placebo (data not shown).

A network meta-analysis (method described in the Introduction under Evidence Summaries) was performed based on indirect comparisons of the placebo-controlled RCTs noted above (Table 1). Due to substantial heterogeneity in findings among different TNFis for ACR 20 and PASI75, the individual drugs were separated in the network meta-analysis. Of individual TNFIs, golimumab (GOL) and infliximab were associated with significantly greater proportions of patients who achieved ACR20 compared to ustekinumab, while only infliximab was associated with significantly greater proportions of patients who achieved PASI75 compared to ustekinumab (see table below). The relative risk for golimumab was elevated in part due to a low placebo event rate (8.8%) in the GO-REVEAL trial compared to the ustekinumab trials (average 20.5%). Other individual TNFis did not show superiority over ustekinumab for these outcomes, but imprecision due to wide confidence intervals that overlapped the line of no difference means that the findings were inconclusive (data not shown). The findings for HAQ DI (proportion of patients who achieved a wire associated between TNFIs and ustekinumab due to a wide CI that overlapped the line of no difference. Infection rates (the only relevant adverse event that could be compared between drug classes) did not differ significantly between TNFis and ustekinumab (this outcome was compared using the Bucher adjusted indirect method, see description in the Introduction to this report). For illustrative purposes the network meta-analyses for ACR20 and HAQ-DI are diagrammed in Figures 1 and 2, respectively. An updated literature search in March 2018 identified one additional RCT showing superiority of golimumab over placebo[19]. This trial had not been previously reviewed by the panel, but it was moderate quality (due to indirectness) and the results did not change the overall findings or the overall quality of evidence.

One trial of psoriasis patients found statistically significant differences favoring ustekinumab over etanercept for one efficacy outcome (PASI75), but no significant differences were reported for serious infections (Table 2).[16] However, the event rate for serious infections was so low that the finding is inconclusive. The PASI75 finding was also supported by a published network meta-analysis of RCTs of patients with psoriasis.[17] In this analysis ustekinumab showed superiority over etanercept for PASI75 (OR 1.94, 95% CI 1.31 to 3.01) and did not differ significantly from adalimumab (OR 1.44, 95% CI 0.82 to 2.58), although the latter finding is inconclusive due to imprecision in the CI. However, Infliximab showed superiority over ustekinumab in this network meta-analysis (OR 3.92, 95% CI 1.83 to 9.06), which is consistent with the findings of our

independently performed network meta-analysis of PsA RCTs. A published meta-analysis of psoriasis RCTs found no significant difference between TNFi versus placebo and IL12/23i versus placebo in rates of major adverse cardiovascular events.[18]

Quality of evidence across all critical outcomes: Moderate

	Table 1. TNFi compared to IL12/23i for patients with active PsA despite OSM Bibliography: PICO 16: TNFi compared to IL12/23i for patients with active PsA despite OSM												
Quality assessment Summary of findings													
Nº of participants	Risk of	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of			Relative effect	Anticipated absolute effects			
(studies) Follow-up	bias					evidence	With IL12/23i	With TNFi	(95% CI)	Risk with IL12/23i	Risk difference with TNFi		
ACR20, 1	2-24 v	weeks, netw	ork meta-	analysis	•					•			
1332 (4 RCTs)	Not serious	Not serious	Serious ^a	Not serious	None	⊕⊕⊕⊖ MODERATE	IL12/23i	GOL	RR 2.44 (1.17 to 5.13) Favors GOL	449 per 1,000 (0.449)	289 more per 1000 (0.289) (44 more to 441 more) ^c		
1431 (6 RCTs)	Not serious	Not serious	Seriousª	Not serious	None	⊕⊕⊕⊖ MODERATE	IL12/23i	IFX	RR 3.19 (1.87 to 5.46) Favors IFX	449 per 1,000	417 more per 1000 (0.417) (159 more to 515 more) ^c		
HAQ-DI ^d ,	IAQ-DI ^d , 12-24 weeks, network meta-analysis												

	Table 1. TNFi compared to IL12/23i for patients with active PsA despite OSM Bibliography: PICO 16: TNFi compared to IL12/23i for patients with active PsA despite OSM													
		Qua	ality assessi	ment				Summary of findings						
2477 (10 RCTs)	Not serious	Not serious	Serious ^a	Not serious	None	⊕⊕⊕⊖ MODERATE	IL12/23i	TNFi	RR 0.99 (0.76 to 1.30) No difference	441 per 1,000	4 fewer per 1000 (0.004) (106 fewer to 132 more)			
PASI 75,	12-24	weeks, net	work meta	a-analysis										
1023 (5 RCTs)	Not serious	Not serious	Serious ^a	Not serious	None	⊕⊕⊕⊖ MODERATE	IL12/23i	IFX	RR 3.34 (1.1 to 10.2) Favors IFX	569 per 1,000 (0.569)	306 more per 1000 (0.306) (66 more to 397 more) °			
Infection	, 12-2	4 weeks, Bu	ucher adju	sted indire	ect compa	irison	•	•	•	•	•			
1025 (3 RCTs)	Not serious	Not serious	Serious ^a	Serious ^b	None	⊕⊕⊖⊖ Low	IL12/23i	ADA	RR 0.56 (0.26 to 1.22)	211 per 1,000	93 fewer per 1000 (0.093) (156 fewer to 46 more)			
1199 (3 RCTs)	Not serious	Not serious	Serious ^a	Serious ^b	None	⊕⊕⊖⊖ Low	IL12/23i	CZP	RR 1.19 (0.80 to 1.76)	211 per 1,000	40 more per 1000 (0.040) (42 fewer to 160 more)			

CI: Confidence interval; RR: Risk ratio

a. Indirect comparison, all studies compared drug to placebo, only 50-75% of patients had prior OSM exposure.

b. Wide 95% CI

c. Absolute risk differences calculated from odds ratios obtained using the Bucher method.

d. HAQ-DI measured as proportion of patients who achieved minimum clinically important improvement (>0.3 or 0.35 in different studies).



Figure 1. Network meta-analysis for ACR20. Node (blue circle) size and line thickness varies based on number of studies and patients in each comparison. The largest circle represents the placebo node (because all RCTs had a placebo control) and the branching lines connect to smaller nodes representing each of the treatments. Due to substantial heterogeneity within the TNFi class the individual TNFi drugs were analyzed separately and compared to the other drug classes (OSM, IL12/23i, and IL17i). The figure also identifies the specific trials that provided data for each treatment node.



Figure 2. Network meta-analysis for HAQ-DI. Since there was no substantial heterogeneity within each drug class, a straight drug class comparison was performed for this outcome.

	Table 2. TNFi compared to IL12/23i for patients with Psoriasis despite OSM Bibliography: PICO 16: TNFi versus IL12/23i for PsA patients who failed OSM.												
		Qua	ality assessr	ment			Summary of findings						
Nº of participants	Risk of	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of	Study event rates (%)		Relative effect	Anticipated absolute effects			
(studies) Follow-up	bias					evidence	With TNFi (ETN)	With IL12/23i	(95% CI)	Risk with TNFi	Risk difference with IL12/23i		
PASI-75	at 12 v	weeks											
903 (1 RCT)	not serious	not serious	seriousª	not serious	none	⊕⊕⊕⊖ MODERATE	197/347 (56.8%)	397/556 (71.4%)	RR 1.26 (1.13 to 1.40) Favors IL12/23i	568 per 1,000 (0.568)	148 more per 1,000 (0.148) (74 more to 227 more)		
Serious I	nfectio	on											
903 (1 RCT)	not serious	not serious	seriousª	not serious	none	⊕⊕⊕⊖ MODERATE	1/347 (0.3%)	4/556 (0.7%)	RD 0.004 (-0.002 to 0.010)	3 per 1,000	4 more per 1,000 (0.004) (2 fewer to 10 more)		

CI: Confidence interval; RR: Risk ratio

a. Indirect population (psoriasis)

Table 3. TNFi or IL12/23i compared to placebo for PICO 16: Adverse events Bibliography: PICO 16: TNFi versus IL 12/23i for PsA patients who failed OSM.

Quality assessment

Summary of findings

	Tabl	e 3. TNFi o Biblio	r IL12/23 ography: PICO	i compare 16: TNFi versu	ed to plac Is IL 12/23i fi	e bo for or PsA patie	PICO 1 Ints who fa	I 6: Adv	verse eve	ents	
		Qual	ity assess	ment				Sumn	nary of f	inding	S
№ of participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of	Study event rates (%)		Relative effect	Anticipated absolute effects	
Follow-up						evidence	With placebo	With drug	(7578 61)	Risk with placebo	Risk difference with TNFi or IL12/23i
Serious ir	nfectio	n – TNFi vs.	placebo								
1151 (5 RCTs)	not serious	not serious	serious ^a	not serious	none	⊕⊕⊕⊖ MODERATE	8/564 (1.4%)	4/587 (0.7%)	RR 0.54 (0.17 to 1.77) No difference	14 per 1,000 (0.014)	6 fewer per 1,000 (0.006) (12 fewer to 11 more)
Serious in	fectio	n – Ustekini	umab vs. pl	lacebo		<u>-</u>					
925 (2 RCTs)	not serious	serious	serious ^a	not serious	none	⊕⊕⊕⊖ MODERATE	2/309 (0.6%)	4/616 (0.6%)	RR 0.84 (0.20 to 3.50) No difference	6 per 1,000 (0.006)	1 fewer per 1,000 (5 fewer to 15 more)

CI: Confidence interval; RR: Risk ratio

a. Comparison to placebo

Working Group quality of evidence categories:

High quality: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate quality: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low quality: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect Very low quality: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

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PICO 17. In adult patients with active PsA despite treatment with OSM, what are the benefits and harms of switching to TNFi compared to switching to IL17i?

Summary. This question was addressed indirectly by 19 double-blind RCTs (22 publications). Two reports of the same study directly compared IL17i (secukinumab) to TNFi (etanercept) among patients with psoriasis; one study specifically reported data only for a subgroup of patients with PsA[1], and one report of outcomes for the whole population (patients with psoriasis)[2]. Nine studies (12 publications) involving PsA patients compared TNFis to placebo.[3-14] One study compared ixekizumab to placebo,[15] three studies compared secukinumab to placebo,[16-18] and three studies compared brodalumab to placebo[19-21] (note: brodalumab is currently FDA-approved for psoriasis but not PsA). Three studies (2 publications) compared secukinumab 300mg to secukinumab 150mg. All drugs showed statistically significant improvements in ACR20, PASI75, and HAQ-DI over placebo, while no significant difference was found for infections, but with high imprecision in the effect estimate. Subgroup analysis of previous TNFi exposure did not find differences between exposed and unexposed subgroups, but no studies specified whether TNFi experienced patients had TNFi treatment failure. In the study comparing secukinumab to etanercept, PASI75 was superior in the secukinumab group with statistically significant results (Table 1). A study of patients with psoriasis similarly found that ixekizumab was superior to etanercept for increasing PASI-75 (Table 2).[22] In studies comparing ACR20, PASI-75 and HAQ-DI for patients dosed with 300mg vs. 150mg of secukinumab, all outcomes showed no significant difference between the two doses of secukinumab, with direction of effect slightly favoring 300 mg (RR 1.06-1.15) but slight imprecision in the CIs (data not shown).

A network meta-analysis was performed to indirectly compare TNFis vs IL17is using the placebo-controlled RCTs noted above (see Table 1). Due to substantial heterogeneity in findings among different TNFis for ACR20, the individual TNFis were separated in the network meta-analysis. Of individual TNFis, only golimumab and infliximab were associated with a significantly higher proportion of patients who achieved ACR20 when compared to patients who received IL17is. The remaining TNFis did not show a significant difference compared to IL17is for ACR20, although all of these findings were inconclusive due to imprecise CIs that overlapped with the line of no difference (data not shown). For HAQ-DI minimum improvement, there was no substantial heterogeneity among different TNFis so a straight drug class comparison was performed. No significant difference between TNFis and IL17 is was identified for this outcome, although the findings were inconclusive due to imprecision in the effect estimate. For PASI-75, although there was some heterogeneity among TNFis, none showed a significant difference with IL17is, so a straight drug class comparison was performed (there was also some heterogeneity among trials of secukinumab). The overall comparison found no significant between-class difference, but the findings were inconclusive due to imprecision in the effect estimate. Bucher adjusted indirect comparisons were performed for the outcome of infections (the only relevant adverse event reported by both sets of drug class trials). The two TNFi trials (one adalimumab, one certolizumab) showed heterogeneity and differences in effect direction, so they were separately compared to two trials of secukinumab that reported this outcome. Both comparisons found no significant difference but were inconclusive due to imprecision in the effect estimates. Six TNFi RCTs and 3 IL17i RCTs reported serious infections and found no significant difference between drug and placebo groups, although the finding was imprecise for IL17i due to a wide 95% CI (Table 3). A published meta-analysis of psoriasis RCTs found no significant difference between TNFi versus placebo and IL17i versus placebo in rates of major adverse cardiovascular events. [23] An updated

literature search in March 2018 identified one additional RCT showing superiority of golimumab over placebo[24]. This trial had not been previously reviewed by the panel, but it was moderate quality (due to indirectness) and the results did not change the overall findings or the overall quality of evidence.

Quality of evidence across all critical outcomes: Low

	Tak	ole 1. TNFi Bibliograp	compared bhy: PICO 17: 1	l to IL17i INFi compared	for patie	ents with patients wit	activ	e PsA PsA des	despite	OSM	
		Qua	lity assessn	nent				Su	ummary of	findings	
Nº of participants	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of			Relative effect	Anticipated effects	absolute
(studies) Follow-up						evidence	IL17i	TNFi	- (95% CI)	Risk with IL17i	Risk difference with TNFi
ACR20, 1	2-24 w	veeks, netwo	ork meta-a	nalysis							
1229 (7 RCTs)	Not serious	Not serious	Serious ^a	Not serious	None	⊕⊕⊕⊖ MODERATE	IL17i	GOL	RR 2.21 (1.1 to 4.48) Favors GOL	503 per 1,000 (0.503)	213 more per 1000 (0.213) (33 more to 344 more) ^e
1328 (9 RCTs)	Not serious	Not serious	Serious ^a	Not serious	None	⊕⊕⊕⊖ moderate	IL17i	IFX	RR 2.89 (1.79 to 4.67) Favors IFX	503 per 1,000 (0.503)	351 more per 1000 (0.351) (80 more to 458 more) ^e
HAQ-DI ^f ,	12-24	weeks, net	work meta	-analysis	•	•			·	•	-

	Tab	ble 1. TNFi Bibliograp	compared ohy: PICO 17: 1	to IL17i NFi compared	for patie	nts with patients wit	active	e PsA PsA desp	despite	OSM	
		Qua	lity assessn	nent				Su	immary of	findings	
2258 (11 RCTs)	Not serious	Not serious	Serious ^a	Serious ^b	None	⊕⊕⊖⊖ Low	IL17i	TNFi	RR 1.28 (0.97 to 1.68)	556 per 1,000 (0.556)	156 more per 1000 (0.156) (17 less to 378 more)
PASI 75,	12-24	weeks, netw	vork meta-	analysis							
1837 (14 RCTs)	Not serious	Not serious	Serious ^a	Serious ^b	None	⊕⊕⊖⊖ Low	IL17i	TNFi	OR 0.82 (0.29 to 2.34)	643 per 1,000 (0.643)	47 fewer per 1000 (0.047) (300 fewer to 165 more) ^e
PASI 75 ,	12-24	weeks, dire	ct compari	son (secul	kinumab	versus et	anerce	ept)	-	•	•
1 RCT (143 patients)	Serious ^c	Not serious	Serious ^d	Not serious	None	⊕⊕⊖⊖ Low	IL17i (SEC)	ETN	RR 0.59 (0.40 to 0.88) Favors SEC	657 per 1,000 (0.657)	270 fewer per 1,000 (0.270) (79 fewer to 394 fewer)
Infection	, 12-24	weeks, Bu	cher adjust	ted indired	t compar	ison			-	•	
702 (3 RCTs)	Not serious	Not serious	Serious ^a	Serious ^b	None	⊕⊕⊖⊖ Low	IL17i	ADA	RR 0.45 (0.21 to 1.01)	313 per 1,000 (0.313)	172 fewer per 1000 (0.172) (247 fewer to 0 more)

Table 1. TNFi compared to IL17i for patients with active PsA despite OSM

Bibliography: PICO 17: TNFi compared to IL17i for patients with active PsA despite OSM

1		Qua	lity assessn		Su	mmary of	findings				
876 (3 RCTs)	Not serious	Not serious	Serious ^a	Serious⁵	None	⊕⊕⊖⊖ Low	IL17i	CZP	RR 0.94 (0.59 to 1.50)	313 per 1,000 (0.313)	19 fewer per 1000 (0.019) (128 fewer to 157 more)

CI: Confidence interval; RR: Risk ratio

- a. Indirect comparison, all studies compared drug to placebo, only 50-75% of patients had prior OSM exposure.
- b. Wide CI that is close to or crosses line of no difference
- c. PsA diagnosis based on patient records; PsA activity not measured within the study.
- d. Only 63% of patients had prior OSM exposure
- e. Absolute risk differences calculated from odds ratios obtained using the Bucher method.
- f. HAQ-DI measured as proportion of patients who achieved minimum clinically important improvement (>0.3 or 0.35 in different studies).

Table 2. TNFi compared to IL17i for patients with psoriasis despite OSM

Bibliography: PICO 17 - TNFi compared to IL17i for patients with active PsA despite OSM.

		Qua	lity assessr	nent				Sun	nmary of fi	indings	
№ of participants	Risk of	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of	Study ev (%)	vent rates	Relative effect	Anticipat effects	ed absolute
(studies) Follow-up	bias					evidence	With TNFi (ETN)	With IL17 (IXE)		Risk with TNFi	Risk difference with IL17i
PASI-75	at wee	ek 12			•	•	•	•	•	•	•

	Table 2. TNFi compared to IL17i for patients with psoriasis despite OSM Bibliography: PICO 17 - TNFi compared to IL17i for patients with active PsA despite OSM.												
		Qua	ality assessr	ment				Sun	nmary of fi	ndings			
1476 (1 RCT)	not serious	not serious	serious ^a	not serious	none	⊕⊕⊕⊖ MODERATE	353/740 (47.7%)	651/736 (88.5%)	RR 1.85 (1.71 to 2.01) Favors IL17 (IXE)	477 per 1,000 (0.477)	405 more per 1,000 (0.405) (339 more to 482 more)		
Infection													
1473 (1 RCT)	not serious	not serious	serious ^a	not serious	none	⊕⊕⊕⊖ MODERATE	159/739 (21.5%)	190/734 (25.9%)	RR 1.20 (1.00 to 1.45)	215 per 1,000 (0.215)	43 more per 1,000 (0.043) (0 fewer to 97 more)		

a. Indirect population (psoriasis)

	Table 3. TNFi or IL17i compared to placebo for PICO 17: Adverse events Bibliography: PICO 17: TNFi versus IL17i for PsA patients who failed OSM.												
		Qual	ity assess			Sumn	nary of f	inding	S				
Nº of participants	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of	Study ev (%)	ent rates	Relative effect	Anticipat absolute	ed effects		
Follow-up						evidence	With placebo	With TNFi or IL17i	(93% (1)	Risk with placebo	Risk difference with TNFi or IL17i		
Serious ir	Serious infection – TNFi vs. placebo												

Table 3. TNFi or IL17i compared to placebo for PICO 17: Adverse events

Bibliography: PICO 17: TNFi versus IL17i for PsA patients who failed OSM.

		Qual	ity assess	ment				Sumr	nary of f	inding	S
1151 (5 RCTs)	not serious	not serious	serious ^a	not serious	none	⊕⊕⊕⊖ MODERATE	8/564 4/587 RR 0.54 14 per (1.4%) (0.7%) (0.17 to 1.77) 1,000 (0.014) No difference No				6 fewer per 1,000 (0.006) (12 fewer to 11 more)
Serious ii	nfectio	n – IL17i vs	. placebo								
1189 (3 RCTs)	not serious	not serious	serious ^a	serious ^b	none		5/406 (1.2%)	18/685 (2.6%)	RR 1.79 (0.75 to 4.30)	12 per 1,000 (0.012)	9 more per 1,000 (0.009)

CI: Confidence interval; RR: Risk ratio

a. Comparison to placebo

b. Wide 95% CI that crosses line of no effect

Working Group quality of evidence categories:

High quality: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate quality: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low quality: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect Very low quality: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

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(3 fewer to 52 more)

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PICO 18. In adult patients with active PsA despite treatment with OSM, what are the benefits and harms of switching to IL12/23i compared to switching to IL17i?

Summary: This PICO question was addressed indirectly by 16 double-blind RCTs (13 publications). Three studies (2 publications) involving psoriasis patients compared IL12/23i (ustekinumab) to IL17i.[1,2] Thirteen studies (11 publications) involving PsA patients or psoriasis patients compared either IL12/23i or IL17i to placebo.[3-13] All drugs showed statistically significant improvements in ACR20, PASI-75, and HAQ-DI over placebo (data not shown).

A network meta-analysis was performed to indirectly compare IL12/23i vs IL17is using the placebo-controlled RCTs noted above. The metaanalysis showed no significant between-class differences for the outcomes ACR20, HAQ-DI and PASI-75, but there was imprecision in all effect sizes due to wide Cls (see Table 1). A Bucher adjusted indirect comparison similarly found no significant between-class difference in infection rates, again with imprecision due to a wide Cl.

Studies of patients with psoriasis found no significant difference in PASI-75 when comparing ustekinumab to brodalumab. The psoriasis study comparing ustekinumab to secukinumab reported that PASI-75 was superior in the secukinumab group and there was no significant difference in infection rates between the two groups (Table 2).

Serious infections for drug vs. placebo groups appear in Table 3. Neither IL12/23i nor IL17i showed significantly different serious infection rates compared to placebo groups, but the findings were imprecise for IL17i due to a wide 95% confidence interval.

Quality of evidence across all critical outcomes: Moderate

	Table 1. IL12/23i compared to IL17i for patients with active PsA despite OSM Bibliography: PICO 18: IL12/23i compared to IL17i for patients with active PsA despite OSM												
		Qua	ality assessr	nent				Sur	nmary of f	indings			
№ of participants	Risk of	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of			Relative effect	Anticipated absolute effects			
(studies) Follow-up	bias					evidence	IL12/23i IL17i (95% CI) IL12/23i with IL12/23i with IL1						
ACR20, 1	CP20_12-24 weeks_network_meta-analysis												

	Tab	le 1. IL12/ Bibliograph	23i comp by: PICO 18: IL	ared to II 12/23i compa	L17i for p ared to IL17i	oatients for patients	with active	tive P PsA des	SA despi	te OSM	
		Qua	ality assessr	nent				Sun	nmary of fi	indings	
2043 (9 RCTs)	Not serious	Not serious	Not serious ^a	Serious ^b	None	⊕⊕⊕⊖ MODERATE	IL12/23i	IL17i	RR 1.1 (0.69 to 1.77)	449 per 1,000 (0.449)	45 more per 1000 (0.045) (139 fewer to 346 more)
HAQ-DI [°] ,	12-24	weeks, net	work meta	a-analysis							
1913 (7 RCTs)	Not serious	Not serious	Not serious ^a	Serious ^b	None	⊕⊕⊕⊖ MODERATE	IL12/23i	IL17i	RR 0.81 (0.58 to 1.13)	441 per 1,000 (0.441)	84 fewer per 1000 (0.084) (185 fewer to 57 more)
PASI 75,	12-24	weeks, net	work meta	-analysis	•			<u> </u>			
1630 (9 RCTs)	Not serious	Not serious	Not serious ^a	Serious ^b	None	⊕⊕⊕⊖ MODERATE	IL12/23i	IL17i	RR 1.20 (0.46 to 3.12)	569 per 1,000 (0.569)	112 more per 1000 (0.112) (92 less to 265 more) ^d
Infection	, 12-2	4 weeks, Bu	icher adjus	sted indire	ct compai	rison				•	•
1527 (4 RCTs)	Not serious	Not serious	Not serious ^a	Serious ^b	None	⊕⊕⊕⊖ MODERATE	IL12/23i	IL17i	RR 1.26 (0.81 to 1.97)	211 per 1,000 (0.211)	55 more per 1000 (0.112) (40 less to 205 more)

- a. Indirect comparison, all studies compared drug to placebo, only 50-75% of patients had prior OSM exposure, but patient characteristics and prior drug exposure are similar between drug classes.
- b. Wide 95% CI that includes possibility of no between-group difference or a substantial between-group difference
- c. HAQ-DI measured as proportion of patients who achieved minimum clinically important improvement (>0.3 or 0.35 in different studies).
- d. Absolute risk differences calculated from odds ratios obtained using the Bucher method.

Table 2. IL12/23 (Ustekinumab) compared to IL17 (Brodalumab or Secukinumab) in patientswith psoriasis

Bibliography: PICO 18: IL12/23i versus IL17i for PsA patients who failed OSM.

		Qua	lity assessr	nent		Summary of findings					
№ of participants	Risk of	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of	Study eve (%)	ent rates	Relative effect	Anticipated abs	solute effects
(studies) Follow-up	DIAS					evidence	With IL12/ 23i	With IL17i	(95% CI)	Risk with IL12/23i	Risk difference with IL17i
PASI-75											
1852 (2 RCTs)	not serious	not serious	serious ^a	not serious	none	⊕⊕⊕⊖ MODERATE	427/613 (69.7%)	841/1239 (67.9%) BROD	RR 0.98 (0.91 to 1.04)	697 per 1,000 (0.697)	14 fewer per 1,000 (0.014) (63 fewer to 28 more)
671 (1 RCT)	not serious	not serious	serious ^a	not serious	none	⊕⊕⊕⊖ MODERATE	277/336 (82.4%)	311/335 (92.8%) SEC	RR 1.12 (1.06 to 1.19) Favors SEC	824 per 1,000 (0.824)	99 more per 1,000 (0.099) (49 more to 157 more)
Infection											
671 (1 RCT)	not serious	not serious	serious ^a	not serious	none	⊕⊕⊕⊖ MODERATE	85/336 (25.3%)	98/335 (29.3%) SEC	RR 1.16 (0.90 to 1.49)	253 per 1,000 (0.253)	40 more per 1,000 (0.040) (25 fewer to 124 more)

	Table 3. IL12/23i or IL17i compared to placebo for PICO 18: Adverse events Bibliography: PICO 18: IL12/23i versus IL17i for PsA patients who failed OSM.												
		Qual	ity assess	ment				Summ	ary of fi	ndings	5		
Nº of participants	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of	Study ev (%)	ent rates	Relative effect	Anticipat absolute	ed effects		
Follow-up						evidence	With With placebo IL12/2 or IL17		(73 % 01)	Risk with placebo	Risk difference with IL12/23i or IL17i		
Serious ir	erious infection – Ustekinumab vs. placebo												
925 (2 RCTs)	not serious	serious	serious ^a	not serious	none	⊕⊕⊕⊖ MODERATE	2/309 (0.6%)	4/616 (0.6%)	RR 0.84 (0.20 to 3.50) No difference	6 per 1,000 (0.006)	1 fewer per 1,000 (5 fewer to 15 more)		
Serious ir	nfectio	n – IL17i vs	. placebo										
1189 (3 RCTs)	not serious	not serious	serious ^a	serious ^b	none	⊕⊕ Low	5/406 (1.2%)	18/685 (2.6%)	RR 1.79 (0.75 to 4.30)	12 per 1,000 (0.012)	9 more per 1,000 (0.009) (3 fewer to 52 more)		

a. Comparison to placebo

b. Wide 95% CI that crosses line of no effect

Working Group quality of evidence categories:

High quality: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate quality: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low quality: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect Very low quality: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

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PICO 19. In adult patients with active PsA <u>despite treatment with an OSM</u>, what are the benefits and harms of switching to MTX and TNFi combination therapy compared to switching to TNFi monotherapy?

<u>Summary:</u> This PICO was addressed by direct drug comparisons in 4 studies (2 RCTs, 1 post hoc analysis of 2 RCTs,[1,2,3] 1 observational[4]), but two of these studies had an indirect population (patients with psoriasis).[2,3] Three studies comparing MTX and TNFi (etanercept) combination therapy with TNFi (etanercept) monotherapy reported no statistically significant differences for three efficacy outcomes (ACR20, HAQ-DI, Physician Global). However, results for one efficacy outcome (PASI75) indicated a statistically significant difference in psoriasis patients at 24 weeks in one RCT,[2] but no statistically significant difference in PsA patients at 24 weeks in the post hoc analysis of two RCTs.[1] The psoriasis RCTs reported no statistically significant between-group difference for adverse events (serious infection, infection, hepatic events), but except for total infections the findings were imprecise.

Quality of evidence across all critical outcomes: Low

	MTX + TNFi compared to TNFi monotherapy for PsA patients failed OSM (randomized) Bibliography: PICO 19: MTX + TNFi combination vs. TNFi monotherapy for PsA patients failed OSM.													
		Qua	ality assessn		S	Summary	of findings							
Nº of participants	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of	Study ev (%)	ent rates	Relative effect	Anticipated a	bsolute effects			
Follow-up						evidence	With TNFI mono- therapy	With MTX + TNFi	(95% CI)	Risk with TNFI mono- therapy	Risk difference with MTX + TNFi			
ACR20, 24	ACR20, 24 weeks													
431 (post hoc analysis of 2 RCTs)	$\begin{array}{c c c c c c c c c c c c c c c c c c c $													
HAQ-DI [°] ,	HAQ-DI ^c , 24 weeks													

	MTX -	⊢ TNFi com Bibliogra	pared to T	NFi mono ITX + TNFi con	therapy for the second se	f or PsA p TNFi monoth	oatients berapy for	s failed PsA patier	OSM (rants failed OSI	andomizeo ^{M.}	d)		
		Qu	ality assessm	nent				S	Summary o	of findings			
436 (post hoc analysis of 2 RCTs)	serious ª	not serious	serious ^b	not serious	none	⊕⊕⊖⊖ Low	178/295 (60.3%)	92/141 (65.2%)	RR 1.08 (0.93 to 1.26) No difference	603 per 1,000 (0.603)	48 more per 1,000 (0.048) (42 fewer to 157 more)		
PASI 75, Psoriatic arthritis, 24 weeks													
406 (post hoc analysis of 2 RCTs)	serious ª	not serious	serious ^b	not serious	none	⊕⊕⊖⊖ Low	166/278 (59.7%)	75/128 (58.6%)	RR 0.98 (0.80 to 1.15) No difference	597 per 1,000 (0.597)	12 fewer per 1,000 (0.012) (119 fewer to 90 more)		
PASI 75,	Psorias	sis, 24 weel	<s< td=""><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td></s<>										
537 (2 RCTs)	serious d	not serious	serious ^e	not serious	none	⊕⊕⊖⊖ Low	167/267 (62.5%)	213/270 (78.9%)	RR 1.26 (1.11 to 1.44) Favors MTX + TNFi	625 per 1,000 (0.625)	163 more per 1,000 (0.163) (69 more to 275 more)		
Serious Infection, Psoriasis													
59 (1 RCT)	not serious	not serious	serious ^e	serious ^f	none		1/28 (3.6%)	0/31 (0.0%)	RR 0.30 (0.01 to 7.13)	36 per 1,000 (0.036)	25 fewer per 1,000 (0.025) (35 fewer to 219 more)		

	MTX -	+ TNFi com Bibliogra	pared to T	NFi mono ITX + TNFi con	therapy f nbination vs.	f or PsA p TNFi monoth	patients herapy for	s failed PsA patier	OSM (ra	andomize ^{M.}	d)
		Qu	ality assessn			S	Summary o	of findings			
Infections	s, 24 w	eeks, Psori	asis								
478 (1 RCT)	not serious	not serious	serious ^e	not serious	none	⊕⊕⊕⊖ MODERATE	62/239 (25.9%)	83/239 (34.7%)	RR 1.34 (1.02 to 1.76) Favors TNFi mono	259 per 1,000	88 more per 1,000 (0.088) (5 more to 197 more)
Hepatic a	dverse	event (inci	reased trans	sminases),	24 weeks	s, Psorias	is				
478 (1 RCT)	not serious	not serious	serious ^e	⊕⊕⊖⊖ Low	4/239 (1.7%)	7/239 (2.9%)	RD 0.01 (- 0.01 to 0.04) RR 1.75 (0.52 to 5.90)	17 per 1,000 (0.017)	10 more per 1,000 (0.010) (10 fewer to 40 more)		

a. Both RCTs contributing data to this post hoc analysis did not describe randomization methods or allocation concealment methods, and 1 trial did not report blinding of outcome assessors.

b. Indirect comparison of treatment arms from 2 placebo-controlled RCTs.

c. HAQ-DI measured as proportion of patients who achieved minimum clinically important improvement (>0.3 or 0.35 in different studies).

d. Open label study

e. Indirect population (Psoriasis patients)

f. Low number of events

ГМ	ΓX + 1	NFi comp a Bibliograph	ared to TN by: PICO 19: M	I <mark>Fi monot</mark> TX + TNFi con	herapy for the number of the n	Or PsA TNFi mono	patients fa	ailed O A patients	SM (obs s failed OSM	ervationa	I)		
		Qua	lity assessn	nent				Sum	mary of fi	ndings			
Nº of participants	Risk of	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality	Study event ra	ates (%)	Relative effect	Anticipated at effects	osolute		
(studies) Follow-up	bias					or evidence	With TNFi monotherapy	With MTX + TNFi	(95% CI)	Risk with TNFi monotherapy	Risk difference with MTX + TNFi		
HAQ-DI, 24 weeks													
284 (1 observational study)	serious ª	not serious	not serious	not serious	none	⊕⊕⊖⊖ Low	52/98 (53.1%)	98/186 (52.7%)	RR 0.99 (0.79 to 1.25) No difference	531 per 1,000 (0.531)	5 fewer per 1,000 (0.005) (111 fewer to 133 more)		
Physician	n globa	al, 24 weeks	5										
284 (1 observational study)	serious ª	not serious	not serious	not serious	none	⊕⊕⊖⊖ Low	98 patients MD -24.7 (21.4)	186 patients MD - 22.2 (22.3)	-		MD 2.5 higher (2.81 lower to 7.81 higher)		

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

a. Selection bias and confounding by indication

Working Group quality of evidence categories:

High quality: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate quality: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low quality: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect Very low quality: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

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PICO 20. In adult patients with active PsA despite treatment with an OSM, what are the benefits and harms of switching to MTX and IL12/23i combination therapy compared to switching to IL12/23i monotherapy?

Summary: The literature searches did not identify any studies that addressed this PICO question.

Quality of evidence across all critical outcomes: Very low

PICO 21. In adult patients with active PsA despite treatment with an OSM, what are the benefits and harms of switching to MTX and IL17i combination therapy compared to switching to IL17i monotherapy?

<u>Summary</u>: The literature searches did not identify any studies that addressed this PICO question.

Quality of evidence across all critical outcomes: Very low

PICO 22. In adult patients with active PsA <u>despite treatment with an OSM</u>, what are the benefits and harms of switching to another OSM monotherapy compared to adding another OSM?

<u>Summary</u>: This PICO question was addressed by indirect comparisons in three RCTs.[1-3] All trials had placebo groups that continued to receive whatever prior OSMs they had been receiving, rather than being switched to a new OSM. One RCT comparing MTX and cyclosporine combination therapy with MTX monotherapy plus placebo reported no statistically significant difference in nausea, but there was imprecision in the effect size estimate.[1] Two RCTs comparing apremilast with placebo included a subgroup analysis of patients receiving other OSMs concomitantly with apremilast or placebo. Together these subgroup analyses found a statistically significant difference favoring apremilast over placebo for one efficacy outcome (ACR20) at 16 weeks.[2,3]

Quality of evidence across all critical outcomes: Low

	MTX compared to MTX + Cyclosporine (CSA) for PsA patients who failed OSM Bibliography: PICO 22: OSM monotherapy versus combination OSM for PsA patients failed OSM.												
Quality assessment Summary of findings													
№ of participants	Risk of	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality	Study ev (%)	vent rates	Relative effect	Anticipate effects	ed absolute		
(studies) Follow-up	bias					evidence	With MTX + CSA	With MTX + placebo	(95% CI)	Risk with MTX + CSA	Risk difference with MTX + placebo		
Nausea, 5	52 wee	eks		•	•	•					•		
72 (1 RCT)	not serious	not serious	serious ^a	serious ^b	none	⊕⊕⊖⊖ Low	15/38 (39.5%)	6/34 (17.6%)	RR 0.45 (0.20 to 1.02)	395 per 1,000 (0.395)	217 fewer per 1,000 (0.217) (316 fewer to 8 more)		

CI: Confidence interval; RD: Risk difference

a. All patients had failed to have a partial response to MTX prior to enrollment, so patients in the MTX + placebo group were not "switched" to MTX.

b. Small study that does not meet optimal information size, wide CI

Apremilast plus other OSMs compared to placebo plus other OSMs for PsA patients who failed OSM

Bibliography: PICO 22: OSM monotherapy versus combination OSM for PsA patients failed OSM.

		Qua	lity assessm		Sum	mary of fir	ndings				
№ ofRparticipantso(studies)bFollow-up	Risk of	Inconsistency	Indirectness	Imprecision	sion Publication Overall bias quality	Study event rates (%)		Relative effect	Anticipated absolute effects		
(studies) Follow-up	bias					of evidence	With placebo + OSM	With apremilast + other OSM	(95% CI)	Risk with placebo	Risk difference with apremilast

ACR20, 16 weeks

ous not serious	serious ^b	not serious	none	₽₽○○	45/214	83/214	RR 1.84	210 per	177 more
				LOW	(21.0%)	(38.8%)	(1.35 to 2.51)	1,000	per 1,000 (0.177)
							_		(74 more to
							Favors apremilast		318 more)
							+ other		
							OSM		
2	ous not serious	ous not serious serious ^b	Dus not serious serious b not serious	Dus not serious serious ^b not serious none	Dus not serious serious b not serious none ⊕⊕○○ LOW	Dus not serious serious b not serious none	Dusnot seriousserious $^{\flat}$ not seriousnone $\bigoplus \bigoplus \bigcirc \bigcirc \\ LOW$ 45/214 (21.0%)83/214 (38.8%)	Dusnot seriousserious bnot seriousnone $\bigoplus \bigoplus \bigcirc \bigcirc \\ LOW$ 45/214 (21.0%)83/214 (38.8%)RR 1.84 (1.35 to 	Dusnot seriousseriousnot seriousnone $\bigoplus \bigoplus \bigoplus \bigcup U \otimes W$ $45/214$ (21.0%) $83/214$ (38.8%) RR 1.84 (1.35 to 2.51)210 per 1,000 (0.210)Dus $\bigoplus \bigoplus \bigcup U \otimes W$ $\bigoplus \bigoplus \bigcup U \otimes W$ $45/214$ (21.0%) $83/214$ (38.8%) RR 1.84 (1.35 to 2.51)210 per 1,000 (0.210)Favors apremilast + other OSM $\bigoplus \bigoplus \bigcup U \otimes W$

CI: Confidence interval; RR: Risk ratio

a. Unclear randomization, allocation concealment, and blinding of patients/providers/assessors in 1 RCT[2]. Both studies not consistent with calculating outcomes for ITT populations.

b. Patients in the placebo group were not actually switched to a different OSM, they continued treatment with the same OSM(s).

Working Group quality of evidence categories:

High quality: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate quality: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low quality: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect Very low quality: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

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PICO 23. In adult patients with active PsA despite treatment with an OSM, what are the benefits and harms of switching to a different OSM compared to switching to a TNFi?

Summary. Only one study directly compared a TNFi to an OSM in patients with PsA[1]. This study, which compared cyclosporine and adalimumab, was an unblinded observational study with only partial randomization of enrollees (see footnote in Table 2 for further details). We therefore included two RCTs comparing TNFi to an OSM in patients with psoriasis[2,3] as well as a number of RCTs comparing TNFi or OSM to placebo in patients with PsA. Most of the data presented for this PICO is therefore indirect in nature. Nine RCTs (12 publications) comparing TNFi to placebo were included for this PICO.[4-15] Five RCTs compared OSM to placebo: one compared leflunomide to placebo[16,17], and four compared apremilast to placebo[18-23].

A network meta-analysis was performed to indirectly compare OSMs vs TNFis using the placebo-controlled RCTs noted above (Table 1). Due to considerable heterogeneity in findings among different TNFIs for ACR20, HAQ-DI and PASI-75, these drugs were separated in the network metaanalysis. Of individual TNFIs, golimumab and infliximab were associated with significantly greater numbers of patients who achieved ACR20 and PASI75 compared to the OSM group. Only adalimumab and infliximab were associated with significantly greater numbers of patients who achieved minimum change in HAQ-DI. The remaining individual TNFIs did not show superiority over the OSM group for any outcomes, although imprecision in the findings means that a between-group difference could not be ruled out (data not shown). An updated literature search in March 2018 identified one additional RCT showing superiority of golimumab over placebo[24]. This trial had not been previously reviewed by the panel, but it was moderate quality (due to indirectness) and the results did not change the overall findings or the overall quality of evidence.

The single observational study directly comparing an OSM to a TNFi found cyclosporine to be significantly better than adalimumab for improvement in PASI75. No other outcomes showed significant between-group differences, although there was imprecision in the effect estimates. One RCT of psoriasis patients found a significant benefit of etanercept over 5 mg tofacitinib for improving PASI75 (Table 2).

Adverse events could not be compared using adjusted indirect comparison because the trials of different drug classes did not report similar adverse events. Both apremilast and leflunomide showed a slightly greater risk of causing liver toxicity compared to placebo when combined in a meta-analysis. Apremilast also caused increased GI discomfort compared to placebo (Table 3).

No trial stratified outcomes by history of exposure or failure of OSMs, but a few TNFi vs placebo and apremilast vs placebo RCTs reported outcomes by history of exposure to biologic agents. In the limited data available, TNFi blockers and apremilast seem to work as well in biologic-experienced compared to biologic-naïve patients.

Quality of evidence across all critical outcomes: Moderate

Table 1. Different OSM compared to TNFi for patients with active PsA despite OSM Bibliography: PICO 23: Different OSM compared to TNFi for patients with active PsA despite OSM **Quality assessment** Summary of findings Inconsistency Imprecision Publication Overall Nº of Risk Indirectness Relative Anticipated absolute participants of bias quality of effect effects (studies) evidence bias (95% CI) OSM **Risk with** Risk TNFi Follow-up OSM difference with TNFi ACR20, 12-24 weeks, network meta-analysis GOL 1539 (6 Not Not serious Serious^a Not serious None $\oplus \oplus \oplus \bigcirc$ OSM RR 2.60 380 per 321 more RCTs) serious (1.29 to 1,000 per 1000 MODERATE 5.26) (0.321) (0.380)(139 more Favors to 456 GOL more)^b 1638 (8 Not Not serious Serious^a Not serious None $\oplus \oplus \oplus \bigcirc$ OSM IFX RR 3.40 380 per 464 more (2.10 to 1,000 per 1000 RCTs) serious MODERATE (0.464) 5.51) (0.380) (185 more Favors to 578 IFX more) ^b HAQ-DI^c, 12-24 weeks, network meta-analysis 1580 (6 Not Not serious Serious^a Not serious None $\oplus \oplus \oplus \bigcirc$ OSM ADA RR 1.43 394 per 169 more RCTs) (1.06 to per 1000 serious 1,000 MODERATE 1.94) (0.394)(0.169) (24 more Favors to 370 ADA more)

	Table	1. Differer Bibliography	nt OSM co PICO 23: Diff	mpared to erent OSM cor	O TNFi fO	r patien Fi for patien	ts with ts with ac	activ tive PsA	e PsA de despite OSM	espite OS	SM		
		Qua	ality assessi	ment				Su	immary of	findings			
1367 (5 RCTs)	Not serious	Not serious	Serious ^a	Not serious	None	⊕⊕⊕⊖ MODERATE	OSM	IFX	RR 1.79 (1.21 to 2.64) Favors IFX	394 per 1,000 (0.394)	224 more per 1000 (0.224) (58 more to 367 more) ^b		
PASI 75, 12-24 weeks, network meta-analysis													
1026 (5 RCTs)	Not serious	Not serious	Serious ^a	Not serious	None	⊕⊕⊕⊖ MODERATE	OSM	GOL	RR 4.74 (1.07 to 21.01) Favors GOL	164 per 1,000 (0.164)	405 more per 1000 (0.405) (54 more to 698 more) ^b		
1062 (6 RCTs)	Not serious	Not serious	Serious ^a	Not serious	None	⊕⊕⊕⊖ MODERATE	OSM	IFX	RR 8.23 (2.14 to 31.65) Favors IFX	164 per 1,000 (0.164)	631 more per 1000 (0.631) (320 more to 777 more) ^b		

a. Indirect comparison, all studies compared drug to placebo, only 50-75% of patients had prior OSM exposure.

b. Absolute risk differences calculated from odds ratios obtained using the Bucher method.

c. HAQ-DI measured as proportion of patients who achieved minimum clinically important improvement (>0.3 or 0.35 in different studies).

Table 2. OSM compared to TNFi for patients with active PsA despite OSM

Bibliography: PICO 23: In patients with active PsA despite OSM, benefit/harm of switching to different OSM compared to TNFi.

		Qual			Summ	nary of	finding	gs					
Nº of participants	Risk of	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of	Study ev (%)	ent rates	Relative effect	Anticipa absolute	ted effects		
Follow-up	Dias					evidence	With TNFi	With OSM	CI)	Risk with TNFi	Risk difference with OSM		
ACR20 - 0	yclosp	orine vs TNI	Fi (Adalimu	mab)									
115 (1 study)	serious ª	not serious	not serious	serious ^c	none	⊕⊕⊖⊖ Low	45/58 (77.6%)	36/57 (63.2%)	RR 0.81 (0.64 to 1.04)	776 per 1,000 (0.776)	147 fewer per 1,000 (0.147) (279 fewer to 31 more)		
PASI 75 -	PASI 75 – Apremilast versus TNFi (Etanercept)												
166 (1 RCT)	not serious	not serious	serious ^a	serious ^b	none	⊕⊕⊖⊖ Low	40/83 (48.2%)	33/83 (39.8%)	RR 0.82 (0.58 to 1.17)	482 per 1,000	87 fewer per 1,000 (202 fewer to 82 more)		
PASI-75 -	Tofac	itinib vs TNF	i (Etanerce	ept)									
664 (1 RCT)	not serious	not serious	serious ^b	not serious	none	⊕⊕⊕⊖ MODERATE	197/335 (58.8%)	130/329 (39.5%)	RR 0.67 (0.57 to 0.79) Favors ETN	588 per 1,000 (0.588)	194 fewer per 1,000 (0.194) (253 fewer to 123 fewer)		
PASI-75 -	Cyclo	sporine vs T	NFi (Adalin	numab)									

Bibl	Table 2. OSM compared to TNFi for patients with active PsA despite OSM Bibliography: PICO 23: In patients with active PsA despite OSM, benefit/harm of switching to different OSM compared to TNFi.													
		Qual	ity assess	ment				Summ	ary of	finding	js			
84 (1 study)	serious ^a	not serious	not serious	serious ^c	none	⊕⊕⊖⊖ Low	9/43 (20.9%)	18/41 (43.9%)	RR 2.10 (1.07 to 4.12)	209 per 1,000 (0.209)	230 more per 1,000 (0.230) (15 more to 653 more)			
HAQ-DI ^d - Cyclosporine vs TNFi (Adalimumab)														
115 (1 RCT)	serious ^a	not serious	not serious	serious °	none	⊕⊕⊖⊖ Low	43/58 (74.1%)	33/57 (57.9%)	RR 0.78 (0.60 to 1.02)	741 per 1,000 (0.741)	163 fewer per 1,000 (0.163) (297 fewer to 15 more)			
Serious a	dverse	events - To	facitinib vs	TNFi (Etar	ercept)									
664 (1 RCT)	not serious	not serious	serious ^b	not serious	none	⊕⊕⊕⊖ MODERATE	7/335 (2.1%)	7/329 (2.1%)	RR 1.02 (0.36 to 2.87)	21 per 1,000 (0.021)	0 more per 1,000 (13 fewer to 39 more)			
Liver toxi	city ^e -	Cyclosporine	e vs TNFi (A	dalimuma	b)									
115 (1 study)	serious ª	not serious	not serious	serious ^c	none	⊕⊕⊖⊖ Low	5/58 (8.6%)	3/57 (5.3%)	RR 0.61 (0.15 to 2.44)	86 per 1,000 (0.086)	34 fewer per 1,000 (0.034) (73 fewer to 124 more)			

Table 2. OSM compared to TNFi for patients with active PsA despite OSM

Bibliography: PICO 23: In patients with active PsA despite OSM, benefit/harm of switching to different OSM compared to TNFi.

Quality assessment

Summary of findings

Serious infection - OSM vs TNFi

945 (3 RCTs)	Not serious	not serious	serious ^b	not serious	none	⊕⊕⊕⊖ MODERATE	4/476 (0.8%)	3/469 (0.6%)	RR 0.81 (0.20 to 3.38)	8 per 1,000 (0.008)	2 fewer per 1,000 (0.002) (6 fewer to 19 more)

CI: Confidence interval; RR: Risk ratio

a. The Karanikolas et al 2011 cyclosporine vs adalimumab study[1] is an unblinded clinical trial in which most but not all patients were randomized to treatment groups. Specifically, 76 patients were randomly assigned to cyclosporine, adalimumab, or combination therapy. The remaining 94 patients were not randomly assigned: 32 patients could not receive adalimumab due to insurance restrictions, so they were assigned to cyclosporine. The remaining 62 patients were randomly assigned to adalimumab (32 patients) and combination therapy (30 patients).

b. The Bachelez 2015 tofacitinib vs etanercept RCT[2] and the Reich apremilast vs. etanercept RCT[3] focused on patients with plaque psoriasis, not psoriatic arthritis.

c. Wide CI that crosses line of no effect

d. HAQ-DI measured as proportion of patients who achieved minimum clinically important improvement (>0.3 or 0.35 in different studies).

e. Alanine aminotransferase or aspartate aminotransferase value \geq 3 times upper limit of normal.

Table 3. OSM or TNFi compared to placebo for PICO 23: Adverse events

Bibliography: PICO 23: In patients with active PsA despite OSM, benefit/harm of switching to different OSM compared to TNFi.

		Qual	ity assess	ment				Sumn	nary of f	inding	S
Nº of participants	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of	Study ev (%)	ent rates	Relative effect	Anticipat absolute	ed effects
Follow-up						evidence	With placebo	With TNFi or OSM	(7378 01)	Risk with placebo	Risk difference with TNFi or OSM

Bibli	Table 3. OSM or TNFi compared to placebo for PICO 23: Adverse events Bibliography: PICO 23: In patients with active PsA despite OSM, benefit/harm of switching to different OSM compared to TNFi.												
		Qual	ity assess	ment				Sumn	nary of f	inding	S		
Serious in	nfectio	ns – TNFi vs											
1151 (5 RCTs)	not serious	not serious	serious ^a	not serious	none	⊕⊕⊕⊖ MODERATE	8/564 (1.4%)	4/587 (0.7%)	RR 0.54 (0.17 to 1.77) No difference	14 per 1,000 (0.014)	6 fewer per 1,000 (0.006) (12 fewer to 11 more)		
Liver toxicity ^d – OSM vs. placebo													
1309 (5 RCTs)	not serious	not serious	serious ^a	not serious	none	⊕⊕⊕⊖ MODERATE	6/652 (0.9%)	19/657 (2.9%)	RR 2.58 (1.13 to 5.85) Favors placebo	9 per 1,000 (0.009)	15 more per 1,000 (0.015) (1 more to 45 more)		
GI intoler	ance (diarrhea an	d nausea)	– OSM vs.	placebo								
1308 (5 RCTs)	not serious	serious ^c	serious ^a	not serious	none	⊕⊕℃ LOW	76/646 (11.8%)	220/662 (33.2%)	RR 2.71 (1.62 to 4.52) Favors placebo	118 per 1,000 (0.118)	202 more per 1,000 (0.169) (73 more to 415 more)		
GI intoler	ance (diarrhea an	d nausea)	- Apremila	ist vs plac	ebo							

Table 3. OSM or TNFi compared to placebo for PICO 23: Adverse events

Bibliography: PICO 23: In patients with active PsA despite OSM, benefit/harm of switching to different OSM compared to TNFi.

Quality assessment								Summary of findings			
1120 (4 RCTs)	not serious	serious	serious ^a	not serious	none	⊕⊕⊕⊖ MODERATE	56/554 (10.1%)	188/566 (33.2%)	RR 3.15 (1.79 to 5.54) Favors placebo	101 per 1,000 (0.101)	217 more per 1,000 (0.217) (80 more to 459 more)

CI: Confidence interval; RR: Risk ratio

a. Comparison to placebo

b. Wide confidence intervals due to low event rates

c. The study of leflunomide had a RR of 1.53 (0.95-2.48) whereas the studies of apremilast showed a higher risk, with an overall RR of 3.15 (1.79 to 5.54)

d. Studies had slightly different cutoffs for abnormality: $ALT \ge 1.5$ times the upper normal limit (1 study), ALT > 150 u/l (2 studies), ALT > upper normal limit (1 study)

Working Group quality of evidence categories:

High quality: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate quality: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low quality: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect Very low quality: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

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PICO 24. In adult patients with active PsA despite treatment with an OSM, what are the benefits and harms of switching to a different OSM compared to switching to an IL12/23i?

Summary. No studies directly compared IL12/23i with an OSM. Eight placebo-controlled RCTs provided indirect evidence for this question: three RCTs of ustekinumab[1-3], and 5 RCTs of OSMs (apremilast, 4 RCTs[4-9], and leflunomide, 1 RCT[10,11]). A majority of patients in most studies had a history of OSM use. None of the studies stratified outcomes by history of OSM use, which also contributes to indirectness of the data relative to the PICO question. For all studies we used data for drug dosages that were closest to standard clinical practice.

A network meta-analysis was performed to indirectly compare OSMs vs IL12/23i using the placebo-controlled RCTs noted above. The metaanalysis found a significant difference favoring IL12/23i for HAQ-DI, but no significant between-class differences in ACR20 or PASI 75 outcomes (these findings were inconclusive due to imprecision in the wide CIs that cross the line of no effect) (Table 1).

Adverse events could not be compared using adjusted indirect comparison because the trials of different drug classes did not report similar adverse events. Ustekinumab trials reported rates of infection while OSM trials reported liver toxicity and gastrointestinal (GI) discomfort. Comparing adverse events in drug versus placebo groups, ustekinumab did not demonstrate any increase in severe infection rates compared to placebo. Both apremilast and leflunomide showed a slightly greater risk of causing liver toxicity compared to placebo when combined in a meta-analysis. Apremilast also caused increased GI discomfort compared to placebo (Table 2).

Quality of evidence across all critical outcomes: Low

Table 1. Different OSM compared to IL12/23i for patients with active PsA despite OSM Bibliography: PICO 24: Different OSM compared to IL12/23i for patients with active PsA despite OSM												
Quality assessment								Summary of findings				
№ of participants	Risk of	Inconsistency	Indirectness I	Imprecision	Publication bias	Overall quality of evidence			Relative effect	Anticipated absolute effects		
(studies) Follow-up	bias						OSM	IL12/23i	(95% CI)	Risk with OSM	Risk difference with IL12/23i	
ACR20, 12-24 weeks, network meta-analysis												

Table 1. Different OSM compared to IL12/23i for patients with active PsA despite OSM Bibliography: PICO 24: Different OSM compared to IL12/23i for patients with active PsA despite OSM													
Quality assessment								Summary of findings					
2353 (8 RCTs)	Not serious	Not serious	Serious ^a	Serious⁵	None	⊕⊕⊖⊖ Low	OSM	IL12/23i	RR 1.06 (0.67 to 1.70)	380 per 1,000 (0.380)	23 more per 1000 (0.023) (125 fewer to 266 more)		
HAQ-DI ^c , 12-24 weeks, network meta-analysis													
2233 (7 RCTs)	Not serious	Not serious	Serious ^a	Not serious	None	⊕⊕⊕⊖ MODERATE	OSM	IL12/23i	RR 1.37 (1.00 to 1.88) Favors IL12/23i	394 per 1,000 (0.394)	146 more per 1,000 (0.146) (0 more to 347 more)		
PASI 75, 12-24 weeks, network meta-analysis													
1637 (7 RCTs)	Not serious	Not serious	Seriousª	Serious⁵	None	⊕⊕⊖⊖ Low	OSM	IL12/23i	RR 2.01 (0.70 to 5.76)	164 per 1,000 (0.164)	166 more per 1000 (0.166) (49 fewer to 781 more)		

a. Indirect comparison, all studies compared drug to placebo, in most studies only 50-75% of patients had prior OSM exposure.

b. Wide CI that overlaps line of no difference

c. HAQ-DI measured as proportion of patients who achieved minimum clinically important improvement (>0.3 or 0.35 in different studies).

Table 2. OSM (apremilast or leflunomide) or IL12/23i compared to placebo for PICO 24: Adverse events

Bibliography: PICO 24: In patients with active PsA despite OSM, benefit/harm of switching to different OSM compared to IL-12/23i.

Quality assessment							Summary of findings					
№ of participants	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence	Study event rates (%)		Relative effect	Anticipated absolute effects		
Follow-up							With placebo	With drug	(95% CI)	Risk with placebo	Risk difference with OSM	
Serious infections – Ustekinumab vs. placebo												
925 (2 RCTs)	not serious	serious	serious ^a	not serious	none	⊕⊕⊕⊖ MODERATE	2/309 (0.6%)	4/616 (0.6%)	RR 0.84 (0.20 to 3.50) No difference	6 per 1,000 (0.006)	1 fewer per 1,000 (5 fewer to 15 more)	
Liver toxicity ^d – OSM vs. placebo												
1309 (5 RCTs)	not serious	not serious	serious ^a	not serious	none	⊕⊕⊕⊖ MODERATE	6/652 (0.9%)	19/657 (2.9%)	RR 2.58 (1.13 to 5.85) Favors placebo	9 per 1,000 (0.009)	15 more per 1,000 (0.015) (1 more to 45 more)	
GI intoler	GI intolerance (diarrhea and nausea) – OSM vs. placebo											

Table 2. OSM (apremilast or leflunomide) or IL12/23i compared to placebo for PICO 24:Adverse events

Bibliography: PICO 24: In patients with active PsA despite OSM, benefit/harm of switching to different OSM compared to IL-12/23i.

Quality assessment							Summary of findings				S
1308 (5 RCTs)	not serious	serious °	serious ^a	not serious	none	⊕⊕ Low	76/646 (11.8%)	220/662 (33.2%)	RR 2.71 (1.62 to 4.52) Favors placebo	118 per 1,000 (0.118)	202 more per 1,000 (0.169) (73 more to 415 more)
GL intolor	Clintoloranea (diarrhaa and naucaa) Anromilast va plaasha										

GI intolerance (diarrnea and nausea) - Apremilast vs placebo

1120 (4 RCTs)not seriousnot seriousseriousnot seriousnone $\bigoplus \bigoplus \bigoplus \bigoplus$ MODERATE	56/554 188/566 (10.1%) (33.2%)	56/554 188/566 RR 3.15 10 (10.1%) (33.2%) (1.79 to 1, 5.54) Favors (0	101 per 217 more 1,000 per 1,000 (0.101) (0.217) (80 more to 459 more)
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CI: Confidence interval; RR: Risk ratio

a. Comparison to placebo

b. Wide confidence intervals due to low event rate in both groups

c. The study of leflunomide had a RR of 1.53 (0.95-2.48) whereas the studies on apremilast showed a higher risk, with an overall of 3.15 (1.79-5.54)

d. Studies had slightly different cutoffs for abnormality: ALT \geq 1.5 times the upper normal limit (1 study), ALT > 150 u/l (2 studies), ALT > upper normal limit (1 study)

Working Group quality of evidence categories:

High quality: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate quality: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low quality: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect Very low quality: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

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PICO 25. In adult patients with active PsA despite treatment with an OSM, what are the benefits and harms of switching to a different OSM monotherapy compared to switching to an IL17i?

Summary: This PICO was addressed indirectly by 16 double-blind RCTs. Eight studies involving PsA patients or psoriasis patients compared IL17i to placebo.[1-8] One study compared IL17i to TNFi.[8] The outcomes measured when available were ACR20, PASI-75, HAQ-DI, and adverse events. All drugs showed statistically significant improvements in ACR20, PASI-75, and HAQ-DI over placebo (data not shown), with no significant between-group difference in adverse events. Eight studies involving PsA patients compared OSMs (apremilast and leflunomide) to placebo.[9-16] Both OSMs showed statistically significant improvements in ACR20, PASI-75, and HAQ-DI over placebo (data not shown). Liver toxicity was significantly greater in patients receiving OSM compared to placebo. GI intolerance did not differ significantly between leflunomide and placebo patients, but a significantly increased risk appeared among patients receiving apremilast compared to patients receiving placebo (Table 2).

A network meta-analysis was performed to indirectly compare OSMs vs IL17is using the placebo-controlled RCTs noted above (Table 1). Although the meta-analysis found no significant difference between drug classes for ACR20 and HAQ DI, the findings are inconclusive due to imprecision in the effect estimates. For PASI75, although heterogeneity was detected among studies of secukinumab, the findings showed a significant benefit of IL17is over OSMs for improvement in this outcome. Although there was no substantial heterogeneity among IL17i studies for ACR20, the effect sizes were larger in trials of secukinumab compared to trials of ixekizumab and brodalumab. Therefore, we used the Bucher adjusted indirect comparison method to compare OSMs to secukinumab alone. This analysis found a significant benefit of secukinumab over OSMs for improvement in ACR20.

Ta	Table 1. Different OSM compared to IL17i for patients with active PsA despite OSM Bibliography: PICO 25: Different OSM compared to IL17i for patients with active PsA despite OSM												
Quality assessment Summary of findings													
№ of participants (studies)	Risk of	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of			Relative effect	Anticipated effects	absolute		
(studies) Follow-up	bias					evidence	Ince OSM IL17i (95% CI) Risk with R OSM IL17i CI) Risk with R OSM d						
ACR20, 1	CR20, 12-24 weeks, network meta-analysis												

Ta	able 1	. Different Bibliography:	OSM com PICO 25: Diffe	pared to	L17i for	patients 7i for patien	s with ts with	active PsA	PsA des despite OS	spite OSI ^M	V
		Qua	ality assessr	nent				Su	mmary of	findings	
2162 (11 RCTs)	Not serious	Not serious	Serious ^a	Serious ^b	None	⊕⊕⊖⊖ Low	OSM	IL17i	RR 1.19 (0.71 to 2.00)	386 per 1,000 (0.386)	73 more per 1000 (0.073) (112 fewer to 386 more)
ACR20, 1	2-24 v	veeks, Buch	er adjuste	d indirect	compariso	on – OSM	lvers	us Secul	kinumab		
1828 (8 RCTs)	Not serious	Not serious	Serious ^a	Not serious	None	⊕⊕⊕⊖ MODERATE	OSM	SEC	RR 1.49 (1.07 to 2.09) Favors SEC	386 per 1,000 (0.386)	189 more per 1000 (0.189) (27 more to 421 more)
HAQ-DI [°] ,	12-24	weeks, net	work meta	a-analysis							
2014 (8 RCTs)	Not serious	Not serious	Serious ^a	Serious ^b	None	⊕⊕⊖⊖ Low	OSM	IL17i	RR 1.11 (0.82 to 1.52)	394 per 1,000 (0.394)	43 more per 1000 (0.043) (71 fewer to 205 more)
PASI 75,	12-24	weeks, net	work meta	-analysis							
1576 (8 RCTs)	Not serious	Not serious	Serious ^a	Not serious	None	⊕⊕⊕⊖ MODERATE	OSM	IL17i	RR 2.41 (1.02 to 5.74) Favors IL17i	164 per 1,000 (0.164)	231 more per 1000 (0.231) (3 more to 777 more)

CI: Confidence interval; RR: Risk ratio

- a. Indirect comparison, all studies compared drug to placebo, only 50-75% of patients had prior OSM exposure.
- b. Wide CI that overlaps line of no difference
- c. HAQ-DI measured as proportion of patients who achieved minimum clinically important improvement (>0.3 or 0.35 in different studies).

Table 2. OSM (apremilast or leflunomide) or IL17i compared to placebo for PICO 25: Adverseevents

Bibliography: PICO 25: In patients with active PsA despite OSM, benefit/harm of switching to different OSM compared to IL-17i.

		Qual	ity assess	ment				Summ	ary of	finding	S
Nº of participants	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of	Study ev (%)	ent rates	Relative effect	Anticipat effects	ed absolute
Follow-up						evidence	With placebo	With IL17i or OSM	CI)	Risk with placebo	Risk difference with IL17i or OSM
Serious in	fectio	n – IL17i vs.	placebo								
1189 (3 RCTs)	not serious	not serious	serious ^b	serious ^d	none	⊕⊕℃ LOW	5/406 (1.2%)	18/685 (2.6%)	RR 1.79 (0.75 to 4.30)	12 per 1,000 (0.012)	9 more per 1,000 (0.009) (3 fewer to 52 more)
Liver toxi	city ^d –	OSM vs. pla	cebo								
1309 (5 RCTs)	not serious	not serious	serious ^a	not serious	none	⊕⊕⊕⊖ MODERATE	6/652 (0.9%)	19/657 (2.9%)	RR 2.58 (1.13 to 5.85) Favors placebo	9 per 1,000 (0.009)	15 more per 1,000 (0.015) (1 more to 45 more)

Table 2. OSM (apremilast or leflunomide) or IL17i compared to placebo for PICO 25: Adverseevents

Bibliography: PICO 25: In patients with active PsA despite OSM, benefit/harm of switching to different OSM compared to IL-17i.

Quality assessment

Summary of findings

GI intolerance (diarrhea and nausea) - OSM vs. placebo

1308 (5 RCTs)	not serious	serious ^c	serious ^a	not serious	none	⊕⊕Ĵ) Low	76/646 (11.8%)	220/662 (33.2%)	RR 2.71 (1.62 to 4.52) Favors placebo	118 per 1,000 (0.118)	202 more per 1,000 (0.169) (73 more to 415 more)
GI intoler	ance (diarrhea an	d nausea) -	Apremilas	st vs place	bo					

1120 (4 RCTs)	not serious	serious	serious ^a	not serious	none	⊕⊕⊕⊖ MODERATE	56/554 (10.1%)	188/566 (33.2%)	RR 3.15 (1.79 to 5.54) Favors placebo	101 per 1,000 (0.101)	217 more per 1,000 (0.217) (80 more to 459 more)
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CI: Confidence interval; RR: Risk ratio

a. Comparison to placebo

b. Wide CI that overlaps the line of no effect

c. The study of leflunomide had a RR of 1.53 (0.95-2.48) whereas the studies on apremilast showed a higher risk, with an overall of 3.15 (1.79-5.54)

d. Studies had slightly different cutoffs for abnormality: ALT \geq 1.5 times the upper normal limit (1 study), ALT > 150 u/l (2 studies), ALT > upper normal limit (1 study))

Working Group quality of evidence categories:

High quality: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate quality: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

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PICO 26. In adult patients with active PsA despite treatment with a TNFi monotherapy, what are the benefits and harms of switching to a second TNFi + MTX compared to adding MTX to the same TNFi monotherapy?

<u>Summary</u>: The literature searches did not identify any studies that addressed this PICO question.

Quality of evidence across all critical outcomes: Very low

PICO 26 (alternate). In adult patients with active PsA despite treatment with a TNFi monotherapy, what are the benefits and harms of switching to a second TNFi monotherapy compared to adding MTX to the same TNFi monotherapy?

<u>Summary</u>: The literature searches did not identify any studies that addressed this PICO question.

<u>Quality of evidence across all critical outcomes</u>: Very low

PICO 27. In adult patients with active PsA despite treatment with a TNFi monotherapy, what are the benefits and harms of switching to a second TNFi compared to switching to IL12/23i?

<u>Summary</u>: This PICO question was addressed indirectly by three double-blind RCTs (4 publications). In addition to lack of direct drug comparisons, only 19% to 50% of patients in each study had prior TNFi exposure. One study (2 publications) involving PsA patients compared TNFi (certolizumab pegol [CZP]) to placebo.[1,2] Two studies involving PsA patients compared IL12/23i to placebo.[3,4] The outcomes measured when available were ACR20, PASI-75, HAQ-DI, serious infections and infections. All drugs showed statistically significant improvements in ACR20, PASI-75, and HAQ-DI over placebo; for serious infections there were no significant between-group differences, but with very high imprecision due to the low number of events (data not shown).

The adjusted indirect comparison method was used to calculate RRs for CZP vs. ustekinumab. For ACR20 we also performed this calculation using data only from patients with prior TNFi exposure (a few studies reported separate data for these patients). Ustekinumab showed a significant benefit over CZP for HAQ-DI and PASI-75, but not for ACR20 or rate of infection (see table below).

	TNFi compared to IL12/23i for patients with active PsA despite prior TNFi Bibliography: PICO 27: TNFi compared to IL12/23i for patients with active PsA despite prior TNFi												
Quality assessment Summary of findings													
Nº of participants	Risk of	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of	Study eve (%)	nt rates	Relative effect	Anticipate effects	d absolute		
(studies) Follow-up	bias					evidence	Risk with IL12/23i	Risk difference with TNFi					
ACR20, 1	ACR20, 12-24 weeks, Bucher adjusted indirect comparison												
732 (3 RCTs) Not serious Not serious Serious ^a Serious ^b None \bigoplus_{LOW} IL12/23i CZP RR 1.00 (0.62 to 1.00) (0.433) Omore per 1000 (165 fewer to 260 more) (0.433)													
ACR20, 1	2-24 v	veeks, Buch	er adjuste	d indirect	comparis	on, TNFi-	exposed	d patie	nts				

	TNFi compared to IL12/23i for patients with active PsA despite prior TNFi Bibliography: PICO 27: TNFi compared to IL12/23i for patients with active PsA despite prior TNFi												
		Qua	ality assessi	nent				Sum	mary of fi	ndings			
260 (2 RCTs)	Not serious	Not serious	Serious ^a	Serious⁵	None	⊕⊕⊖⊖ Low	CZP	IL12/23i	RR 0.32 (0.01 to 2.52)	296 per 1,000 (0.296)	201 fewer per 1000 (293 fewer to 450 more)		
HAQ-DI [°] ,	12-24	weeks, Bu	cher adjus	ted indired	ct compar	ison							
725 (3 RCTs)	Not serious	Not serious	Serious ^a	Not serious	None	⊕⊕⊕⊖ MODERATE	IL12/23i	CZP	RR 0.63 (0.41 to 0.97) Favors IL12/23i	389 per 1,000 (0.389)	144 fewer per 1000 (0.144) (230 fewer to 12 fewer)		
PASI 75,	12-24	weeks, Bud	cher adjust	ted indired	t compar	ison							
535 (3 RCTs)	Not serious	Not serious	Serious ^a	Not serious	None	⊕⊕⊕⊖ MODERATE	IL12/23i	CZP	RR 0.32 (0.13 to 0.83) Favors IL12/23i	531 per 1,000 (0.531)	361 fewer per 1000 (0.361) (462 fewer to 90 fewer)		
Infection	, 12-2	4 weeks, Bu	icher adjus	sted indire	ct compa	rison							
(3 RCTs)	Not serious	Not serious	Serious ^a	Serious⁵	None	⊕⊕⊖⊖ Low	IL12/23i	CZP	RR 1.01 (0.61 to 1.67)	271 per 1,000 (0.271)	3 more per 1000 (0.003) (106 fewer to 182 more)		

CI: Confidence interval; RR: Risk ratio

- a. Indirect comparison, all studies compared drug to placebo, only 50-75% of patients had prior OSM exposure.
- b. Wide 95% CI
- c. HAQ-DI measured as proportion of patients who achieved minimum clinically important improvement (>0.3 or 0.35 in different studies).

Working Group quality of evidence categories:

High quality: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate quality: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low quality: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect Very low quality: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

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PICO 28. In adult patients with active PsA despite treatment with a TNFi monotherapy, what are the benefits and harms of switching to a second TNFi compared to switching to IL17i?

Summary: This PICO was addressed indirectly by six double-blind RCTs (in 7 publications). In addition to lack of direct drug comparisons, only 28% to 50% of patients in each study had prior TNFi exposure. One study (in 2 publications) involving PsA patients compared TNFi (CZP) to placebo.[1,2] Two studies involving PsA patients compared Brodalumab to placebo.[3,4] Three studies compared Secukinumab to Placebo.[5-7] All drugs showed statistically significant improvements in ACR20, PASI-75, and HAQ-DI over placebo (data not shown).

The adjusted indirect comparison method was used to calculate RRs for CZP vs. IL17i (see table below). For ACR20 we also performed this calculation using data only from patients with prior TNFi exposure (a few studies reported separate data for these patients). The analyses found no significant differences in ACR20, HAQ-DI, PASI-75, or total infections, but almost all effect sizes were imprecise due to wide confidence intervals.

	TNFi compared to IL17i for patients with active PsA despite prior TNFi Bibliography: PICO 28: TNFi compared to IL17i for patients with active PsA despite prior TNFi												
		Qua	ality assessr		Sun	nmary of fi	ndings						
Nº of participants (studies) Follow-up	Risk of	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of			Relative effect	Anticipate effects	ed absolute		
(studies) Follow-up	bias					evidence	IL17i	TNFi	(95% CI)	Risk with IL17i	Risk difference with TNFi		
ACR20, 1	2-24 v	veeks, Buch	er adjuste	d indirect	compariso	on							
1022 (5 RCTs)	Not serious	Not serious	Serious ^a	Serious ^b	None	⊕⊕⊖⊖ Low	IL17i	CZP	RR 0.85 (0.56 to 1.28)	479 per 1,000 (0.479)	72 fewer per 1000 (0.072) (211 fewer to 134 more)		

	Т	NFi comp Bibliograp	ared to IL hy: PICO 28: 1	.17i for pa	tients N to IL17i fo	vith active	e PsA c active Ps	despite	e prior TI	NFi			
		Qu	ality asses	sment				Su	mmary of	findings			
ACR20, 12-24 weeks, Bucher adjusted indirect comparison, TNFi-exposed patients													
270 (3 RCTs)	Not serious	Not serious	Serious ^a	Serious ^b	None		CZP	IL17i	RR 0.33 (0.04 to 2.55)	296 per 1,000 (0.296)	198 fewer per 1000 (284 fewer to 459 more)		
HAQ-DI ^c , 12-24 weeks, Bucher adjusted indirect comparison													
910 (4 RCTs)	Not serious	Not serious	Serious ^a	Not serious	None	⊕⊕⊕⊖ MODERATE	IL17i	CZP	RR 1.05 (0.78 to 1.40)	545 per 1,000 (0.545)	27 more per 1000 (0.027) (120 fewer to 218 more)		
PASI 75,	12-24	weeks, Bu	cher adjus	sted indired	t compa	arison		1					
494 (3 RCTs)	'4 (3 RCTs) Not serious Serious ^a Serious ^b None $\bigoplus \bigoplus \bigcup \ LOW$ IL17i CZP RR 0.70 (0.24 to 2.05) 566 per 1,000 (0.140)												
Infection	, 12-2	4 weeks, B	ucher adju	usted indire	ect comp	oarison	<u> </u>	1	1				

TNFi compared to IL17i for patients with active PsA despite prior TNFi Bibliography: PICO 28: TNFi compared to IL17i for patients with active PsA despite prior TNFi Quality assessment Summary of findings 876 (3 RCTs) Not Not serious Serious^a Serious^b IL17i C7P RR 0.94 321 per 19 fewer None $\Theta \Theta \odot \odot$ (0.59 to 1,000 per 1000 serious LOW 1.50) (0.321) (0.019) (132 fewer to 161 more)

CI: Confidence interval; RR: Risk ratio

a. Indirect comparison, all studies compared drug to placebo, only 50-75% of patients had prior OSM exposure.

b. Wide 95% CI

c. HAQ-DI measured as proportion of patients who achieved minimum clinically important improvement (>0.3 or 0.35 in different studies).

d. Absolute risk difference and confidence interval calculated based on the odds ratio.

Working Group quality of evidence categories:

High quality: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate quality: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low quality: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect Very low quality: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

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PICO 29. In adult patients with active PsA despite treatment with a TNFi monotherapy, what are the benefits and harms of switching to an IL12/23i compared to switching to IL17i?

<u>Summary:</u> This PICO was addressed indirectly by 8 double-blind RCTs involving PsA patients or psoriasis patients compared either IL12/23i or IL17i to placebo.[1-8] In addition to lack of direct drug comparisons, only one study[8] included 100% of patients with prior TNFi exposure; the remaining studies included only 28% to 50% of patients with prior TNFi exposure. All drugs showed statistically significant improvements in ACR20, PASI-75, and HAQ-DI over placebo, while adverse event rates did not differ significantly between the two groups (data not shown).

The adjusted indirect comparison method was used to calculate RRs for IL12/23i vs. IL17i (see table below). For ACR20 we also performed this calculation using data only from patients with prior TNFi exposure (a few studies reported separate data for these patients). No significant between-class difference was observed for the outcomes ACR20, HAQ-DI, PASI75, and total infections, but all of these findings were inconclusive due to imprecision in the effect estimates.

	IL12/23i compared to IL17i for patients with active PsA despite prior TNFi Bibliography: PICO 29: IL12/23i compared to IL17i for patients with active PsA despite prior TNFi													
		Qua	lity assessr		Su	mmary of f	findings							
Nº of participants	Risk of	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of			Relative effect	Anticipate effects	d absolute			
(studies) Follow-up	DIAS					evidence IL12/23i IL17i (95% CI) Risk Risk differen IL12/23i with IL								
ACR20, 1	ACR20, 12-24 weeks, Bucher adjusted indirect comparison													
662 (5 RCTs)	62 (5 RCTs) Not serious Serious ^a Serious ^b None \bigoplus_{LOW} IL12/23i IL17i RR 1.19 (0.79 to 1.80) 381 per 1000 (0.072) (80 fewer to 305 more)													
ACR20, 1	ACR20, 12-24 weeks, Bucher adjusted indirect comparison, TNFi-exposed patients													

	IL12/23i compared to IL17i for patients with active PsA despite prior TNFi Bibliography: PICO 29: IL12/23i compared to IL17i for patients with active PsA despite prior TNFi											
		Qua	ality assessi	ment				Su	mmary of f	findings		
611 (4 RCTs)	Not serious	Not serious	Serious ^a	Serious ^b	None	⊕⊕⊖⊖ Low	IL12/23i	IL17i	RR 0.97 (0.47 to 2.00)	356 per 1,000 (0.356)	11 fewer per 1000 (0.011) (189 fewer to 356 more)	
HAQ-DI [°] ,	12-24	weeks, Bu	cher adjus	ted indired	ct compar	ison						
1089 (5 RCTs)	Not serious	Not serious	Serious ^a	Serious ^b	None	⊕⊕⊖⊖ Low	IL12/23i	IL17i	RR 0.68 (0.44 to 1.07)	389 per 1,000 (0.389)	124 fewer per 1000 (0.124) (218 fewer to 27 more)	
PASI 75 ,	12-24	weeks, Bud	cher adjus	ted indired	t compar	ison	•		•	•		
639 (5 RCTs)	Not serious	Not serious	Serious ^a	Serious ^b	None	⊕⊕⊖⊖ Low	IL12/23i	IL17i	RR 0.52 (0.22 to 1.22)	493 per 1,000 (0.493)	237 fewer per 1000 (0.237) (385 fewer to 108 more)	
Infection	, 12-2	4 weeks, Bu	ucher adju	sted indire	ect compa	rison						
913 (3 RCTs)	Not serious	Not serious	Serious ^a	Serious ^b	None	⊕⊕⊖⊖ Low	IL12/23i	IL17i	RR 1.07 (0.62 to 1.85)	271 per 1,000 (0.271)	19 more per 1000 (0.019) (103 fewer to 230 more)	

CI: Confidence interval; RR: Risk ratio

- a. Indirect comparison, all studies compared drug to placebo, in some studies the majority of patients did not have prior TNFi exposure and data was not reported separately for those with TNFi exposure.
- b. Wide 95% CI
- c. HAQ-DI measured as proportion of patients who achieved minimum clinically important improvement (>0.3 or 0.35 in different studies).

Working Group quality of evidence categories:

High quality: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate quality: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low quality: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect Very low quality: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

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PICO 30. In adult patients with active PsA despite treatment with a TNFi monotherapy, what are the benefits and harms of switching to a second TNFi and MTX combination therapy compared to a second TNFi monotherapy?

<u>Summary</u>: The literature searches did not identify any studies that addressed this PICO question.

Quality of evidence across all critical outcomes: Very low

PICO 31. In adult patients with active PsA despite treatment with a TNFi monotherapy, what are the benefits and harms of switching to MTX and IL12/23i combination therapy compared to switching to IL12/23i monotherapy?

Summary: The literature searches did not identify any studies that addressed this PICO question.

Quality of evidence across all critical outcomes: Very low

PICO 32. In adult patients with active PsA despite treatment with a TNFi monotherapy, what are the benefits and harms of switching to MTX and IL17i combination therapy compared to switching to IL17i monotherapy?

Summary: The literature searches did not identify any studies that addressed this PICO question.

Quality of evidence across all critical outcomes: Very low

PICO 33. In adult patients with active PsA despite treatment with a TNFi and MTX combination therapy,, what are the benefits and harms of switching to a second TNFi and MTX combination therapy compared to switching to a second TNFi monotherapy?

Summary: The literature searches did not identify any studies that addressed this PICO question.

Quality of evidence across all critical outcomes: Very low

PICO 34. In adult patients with active PsA despite treatment with a TNFi and MTX combination therapy,, what are the benefits and harms of switching to MTX and IL12/23i combination therapy compared to switching to IL12/23i monotherapy?

<u>Summary</u>: The literature searches did not identify any studies that addressed this PICO question.

PICO 35. In adult patients with active PsA despite treatment with a TNFi and MTX combination therapy, what are the benefits and harms of switching to MTX and IL17i combination therapy compared to switching to IL17i monotherapy?

<u>Summary</u>: The literature searches did not identify any studies that addressed this PICO question.

Quality of evidence across all critical outcomes: Very low

PICO 36. In adult patients with active PsA despite treatment with an IL23/23i, what are the benefits and harms of adding MTX to the IL12/23i compared to switching to TNFi?

Summary: The literature searches did not identify any studies that addressed this PICO question.

Quality of evidence across all critical outcomes: Very low

PICO 37. In adult patients with active PsA despite treatment with an IL23/23i, what are the benefits and harms of adding MTX to the IL12/23i compared to switching to IL17i?

Summary: The literature searches did not identify any studies that addressed this PICO question.

Quality of evidence across all critical outcomes: Very low

PICO 38. In adult patients with active PsA despite treatment with an IL23/23i, what are the benefits and harms of switching to a TNFi compared to switching to IL17i?

Summary: The literature searches did not identify any studies that addressed this PICO question.

Quality of evidence across all critical outcomes: Very low

PICO 39. In adult patients with active PsA despite treatment with an IL17i, what are the benefits and harms of switching to a TNFi compared to switching to IL12/23i?

Summary: The literature searches did not identify any studies that addressed this PICO question.

Quality of evidence across all critical outcomes: Very low

PICO 40. In adult patients with active PsA despite treatment with an IL17i, what are the benefits and harms of adding MTX to the IL17i compared to switching to IL12/23i?

Summary: The literature searches did not identify any studies that addressed this PICO question.

Quality of evidence across all critical outcomes: Very low

PICO 41. In adult patients with active PsA despite treatment with an IL17i, what are the benefits and harms of adding MTX to the IL17i compared to switching to TNFi?

Summary: The literature searches did not identify any studies that addressed this PICO question.

Quality of evidence across all critical outcomes: Very low

PICO 42. In adult patients with active PsA despite treatment with an IL17i, what are the benefits and harms of switching to a different IL17i compared to switching to TNFi?

<u>Summary</u>: The literature searches did not identify any studies that addressed this PICO question.

Quality of evidence across all critical outcomes: Very low

PICO 43. In adult patients with active PsA despite treatment with an IL17i, what are the benefits and harms of switching to a different IL17i compared to switching to IL12/23i?

Summary: The literature searches did not identify any studies that addressed this PICO question.

PICO 44. Among adults with active PsA, what are the benefits and harms of treat to target (or intensive therapy) compared to a not treat to target strategy (include liver toxicity, zoster, malignancy, infection, cardiovascular, IBD, uveitis)?

Summary: This PICO was directly addressed by one open label, multicenter RCT that compared tight control to standard therapy.[1] Methotrexate was the initial therapy for the tight control arm and followed by a specific treatment algorithm. Patients who did not achieve minimal disease activity (MDA) following 12 weeks of MTX (starting at 15 mg/week and ending at 25 mg/week) received combination therapy (MTX + sulfasalazine) for 8 weeks. Patients who still did not achieve MDA then received either combination treatment (MTX + cyclosporine or MTX + leflunomide) for 12 weeks if they had <3 swollen joints or first-line anti-TNFi therapy (usually etanercept unless contraindicated) for 12 weeks if they had ≥3 swollen joints. Patients who still did not achieve MDA and had ≥3 swollen joints then received second-line anti TNFi therapy for 12 weeks. Patients were seen every 4 weeks. At 48 weeks follow-up, statistically significant differences favoring tight control over standard therapy were reported for disease activity (measured by ACR 20 response) and skin (measured by PASI-75) in the evaluable patient population. Abdominal/GI discomfort occurred significantly more often in the MTX tight control group. Nausea was also more frequent in this group but the difference did not reach statistical significance. Liver enzyme abnormalities did not differ significantly between groups, but the effect size was imprecise due to a wide CI.

		Tight con Bibliog	trol compa raphy: PICO 44	ared to st	andard of versus Sta	ndard Care	Adults for Adults	with Activ	ctive PsA ve PsA.	L			
	Quality assessment Summary of findings												
№ of participants	Risk of	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of	Study eve (%)	ent rates	Relative effect	Anticipate effects	ed absolute		
(studies) Follow-up	bias					evidence	With standard care	With MTX tight control	(95% CT)	Risk with standard care	Risk difference with MTX tight control		
ACR20, 4	R20, 48 weeks												

	Tight control compared to standard care for Adults with Active PsA Bibliography: PICO 44: Tight Control versus Standard Care for Adults with Active PsA.												
		Qua	ality assessi	ment			Summary of findings						
173 (1 RCT)	serious ^a	not serious	not serious	not serious	none	⊕⊕⊕⊖ MODERATE	37/84 (44.0%)	55/89 (61.8%)	RR 1.40 (1.05 to 1.88) Favors tight control	440 per 1,000 (0.440)	176 more per 1,000 (0.176) (22 more to 388 more)		
PASI 75, 4	48 we	eks											
156 (1 RCT)	serious ^a	not serious	not serious	not serious	none	⊕⊕⊕⊖ MODERATE	27/81 (33.3%)	44/75 (58.7%)	RR 1.76 (1.23 to 2.53) Favors tight control	333 per 1,000 (0.333)	253 more per 1,000 (0.253) (77 more to 510 more)		
Liver enz	yme a	bnormalitie	es ^c	1		1	1			1	1		
206 (1 RCT)	a a	not serious	not serious	serious ⁶	none	⊕⊕⊖⊖ Low	28/105 (26.7%)	23/101 (22.8%)	RR 0.85 (0.53 to 1.38)	267 per 1,000 (0.267)	40 fewer per 1,000 (0.040) (125 fewer to 101 more)		
Abdomin	al/GI	upset ^d											

	Tight control compared to standard care for Adults with Active PsA Bibliography: PICO 44: Tight Control versus Standard Care for Adults with Active PsA.											
		Qua	ality assessr	ment				Sun	nmary of fi	ndings		
206 (1 study)	serious ^a	not serious	not serious	not serious	none	⊕⊕⊕⊖ MODERATE	12/105 (11.4%)	31/101 (30.7%)	RR 2.69 (1.46 to 4.93) Favors standard care	114 per 1,000 (0.114)	193 more per 1,000 (0.193) (53 more to 449 more)	
Nausea												
206 (1 RCT)	serious ª	not serious	not serious	serious ^b	none	⊕⊕⊖⊖ Low	27/105 (25.7%)	36/101 (35.6%)	RR 1.39 (0.91 to 2.10)	257 per 1,000 (0.257)	100 more per 1,000 (0.100) (23 fewer to 283 more)	

CI: Confidence interval; RR: Risk ratio

- a. No blinding of patients/physicians. No patient data provided for primary outcome (ITT population); only OR presented in narrative. Recall bias for AEs described by authors as a limitation.
- b. Wide CI that overlaps with line of no effect.
- c. Cutoff for abnormal enzyme level was not defined in the study
- d. Not defined in the study

Note: TICOPA trial. Additional limitations noted by authors included inability to test the efficacy of masked assessors; "blunting of the efficacy of tight control" since standard arm being treated by consultant rheumatologists at teaching hospitals and may already be following a more aggressive approach to treatment; possible dilution of intended treatment effect due to "deviations from the treatment escalation protocol."

Working Group quality of evidence categories:

High quality: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate quality: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low quality: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect Very low quality: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

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PICO 45. In adult patients with active axial PsA despite treatment with NSAIDs, what are the benefits and harms of switching to IL12/23i compared to switching to TNFi?

<u>Summary</u>: The literature searches did not identify any studies that addressed this PICO question.

Quality of evidence across all critical outcomes: Very low

PICO 46. In adult patients with active axial PsA despite treatment with NSAIDs, what are the benefits and harms of switching to IL17i compared to switching to TNFi?

<u>Summary</u>: The literature searches did not identify any studies that addressed this PICO question.

Quality of evidence across all critical outcomes: Very low

PICO 47. In adult patients with active axial PsA despite treatment with NSAIDs, what are the benefits and harms of switching to IL17i compared to switching to IL12/23i?

Summary: The literature searches did not identify any studies that addressed this PICO question.

PICO 48. In adult patients with active PsA and predominant enthesitis who are both OSM and biologic treatment-naïve, what are the benefits and harms of starting OSM compared to starting NSAIDs?

Summary: This PICO was addressed indirectly by three double-blind RCTs comparing OSM to Placebo, all using apremilast.[1-3] No studies included a treatment arm of patients starting NSAIDs. The relevant outcomes measured included enthesitis, serious infection, liver toxicity, nausea and diarrhea. Because all trials had a placebo comparison and all (or almost all) patients had prior treatment with OSM and/or biologics, the indirectness of the evidence is very serious.

Only one of the apremilast studies [3] provided sufficient information to import into Table 1. The other [2] reported a non-significant trend toward greater improvement in MASES score in patients receiving apremilast compared with placebo; the magnitude of difference was smaller in this study. The apremilast groups had higher liver toxicity (non-significant difference with imprecision in the CI), but a statistically significant increase in GI intolerance compared to placebo.

Quality of evidence across all critical outcomes: Very low

Table 1. OSM or NSAIDs compared to placebo in patients with enthesitis who are treatmentnaive

Bibliography: PICO 48: In patients with PsA and enthesitis who are treatment-naive, benefit/harm of starting OSM vs. starting NSAIDs.

	Quality assessment								Summary of findings					
№ of participants	Risk of	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of	Study ev rates (%	vent 5)	Relative effect	Anticipate effects	ed absolute			
(studies) Follow-up	bias					evidence	With With placebo OSM		(95% CI)	Risk with placebo	Risk difference with OSM			

Enthesitis score (MASES) (LS mean change) – Apremilast vs. placebo

326 (1 RCT)	not serious	serious ^a	very serious ^b	not serious	none	⊕ VERY LOW	165	161	-	The mean enthesitis score (MASES) (LS mean change)	MD 0.9 lower (1.73 lower to 0.07 lower) Favors
										change) was 0	apremilast

Table 1. OSM or NSAIDs compared to placebo in patients with enthesitis who are treatmentnaive

Bibliography: PICO 48: In patients with PsA and enthesitis who are treatment-naive, benefit/harm of starting OSM vs. starting NSAIDs.

		Qua	ality assessr		Summary of findings						
Nausea –	TNFi	vs. placebo									
259 (1 RCT)	not serious	not serious	very serious ^b	serious ^c	none	⊕ VERY LOW	5/113 (4.4%)	4/146 (2.7%)	RR 0.62 (0.17 to 2.25)	44 per 1,000 (0.044)	17 fewer per 1,000 (0.017) (37 fewer to 55 more)
GI intoler	ance	(diarrhea ar	nd nausea)	– Apremil	ast vs. pl	acebo					
983 (3 RCTs)	not serious	not serious	very serious ^b	not serious	none	⊕⊕⊖⊖ Low	38/486 (7.8%)	162/497 (32.6%)	RR 4.17 (3.00 to 5.80) Favors placebo	78 per 1,000 (0.078)	248 more per 1,000 (0.248) (156 more to 375 more)
Liver toxi	city ^d -	- Apremilast	t vs. placeb	00			<u> </u>	<u> </u>			
984 (3 RCTs)	not serious	not serious	very serious ^b	not serious	none	⊕⊕⊖⊖ Low	1/492 (0.2%)	6/492 (1.2%)	RD 0.01 (-0.001 to 0.02) Favors placebo	2 per 1,000 (0.002)	10 more per 1,000 (0.005) (1 fewer to 20 more)

CI: Confidence interval; RD: Risk difference RR: Risk ratio

a. A second study, by Edwards et al 2016, reported a smaller improvement in enthesitis score but could not be added to RevMan. See discussion in summary paragraph.

b. Indirect comparison to placebo and all or almost all patients had prior exposure to OSMs or biologics.

c. Wide CI crossing significant effect and no-effect lines

d. Studies had slightly different cutoffs for abnormality: ALT > 150 u/l (2 studies), ALT > upper normal limit (1 study)

Working Group quality of evidence categories:

High quality: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate quality: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low quality: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect Very low quality: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

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PICO 49. In adult patients with active PsA and predominant enthesitis despite treatment with OSM, what are the benefits and harms of switching to TNFi compared to switching to OSM?

Summary: This PICO was addressed indirectly by five double-blind RCTs (6 publications). Two studies (in three publications) involving PsA patients compared TNFi (golimumab and certolizumab pegol) to placebo.[1-3] Three studies compared OSM to Placebo, all using apremilast.[4-6] The relevant outcomes measured included enthesitis, serious infection, liver toxicity, nausea and diarrhea.

Enthesitis and adverse events could not be compared using adjusted indirect comparisons, so these findings are presented for drug versus placebo in Table 1. Only one of the apremilast studies [6] provided sufficient information to import into the comparison tables. The other [5] reported a non-significant trend toward greater improvement in MASES score in patients receiving apremilast compared with placebo; the magnitude of difference was smaller in this study. TNFis were superior to placebo in improving enthesitis. There was no significant difference between TNFi and placebo in serious infections, nausea and diarrhea, but all findings were imprecise. The apremilast groups had higher liver toxicity (non-significant difference with imprecision in the CI), but a statistically significant increase in GI intolerance.

Ta Bibl	Table 1. TNFi or OSM compared to placebo in patients with enthesitis despite OSM Bibliography: PICO 49: In patients with PsA and enthesitis despite OSM, benefit/harm of switching to TNFi compared to OSM.											
		Qua	ality assessr	nent				Sur	nmary of f	indings		
№ of participants	Risk of	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of	Study event rates (%)		Relative effect	Anticipated absolute effects		
(studies) Follow-up	bias					evidence	With placebo	With TNFi or OSM	(95% CI)	Risk with placebo	Risk difference with TNFi or OSM	
Enthesitis	sscore	e (MASES) (LS mean c	hange) – <i>I</i>	Apremilas	t vs. plac	ebo	•	1			

T Bik	able 1	I. TNFi or C y: PICO 49: In p	DSM comp patients with P	ared to p sA and enthes	lacebo in sitis despite C	patient SM, benefit/	S with ^{/harm of s}	enthes switching	sitis desp to TNFi com	Dite OSI	И sм.
		Qu	ality assess	ment				Sun	nmary of f	indings	
326 (1 RCT)	not serious	serious ^a	serious ^b	Not serious	none	⊕⊕ Low	165	161	-	The mean enthesitis score (MASES) (LS mean change) was 0	MD 0.9 lower (1.73 lower to 0.07 lower) Favors apremilast
Enthesiti	s, 14 v	veeks – Gol	imumab vs	s. placebo	•	1			•	1	•
247 (1 RCT)	not serious	not serious	serious ^b	not serious	none	⊕⊕⊕⊖ MODERATE	75/105 (71.4%)	78/142 (54.9%)	RR 0.77 (0.63 to 0.93) Favors GOL	714 per 1,000 (0.714)	164 fewer per 1,000 (0.164) (264 fewer to 50 fewer)
Serious i	nfectio	on – TNFi vs	. placebo	1		•					ł
533 (2 RCTs)	not serious	not serious	serious ^b	not serious	none	⊕⊕⊕⊖ MODERATE	5/249 (2.0%)	3/284 (1.1%)	RD -0.009 (-0.02 to 0.02)	20 per 1,000 (0.020)	9 fewer per 1,000 (0.009) (20 fewer to 20 more)
Diarrhea	– TNF	i vs. placeb	0								

-

Bi	Table 1	I. TNFi or C y: PICO 49: In p	SM comp	ared to p	lacebo in itis despite O	patients SM, benefit	s with /harm of s	enthes switching	sitis desp to TNFi comp	Dite OS	И 5м.
		Qua	ality assess	ment				Sur	nmary of f	indings	
259 (1 RCT)	not serious	not serious	serious ^b	serious ^c	none	⊕⊕⊖⊖ Low	4/113 (3.5%)	5/146 (3.4%)	RR 0.97 (0.27 to 3.52)	35 per 1,000 (0.035)	1 fewer per 1,000 (0.001) (26 fewer to 89 more)
Nausea	– TNFi	vs. placebo	•	•	•	•	•	•	•	•	•
259 (1 RCT)	not serious	not serious	serious ^b	serious ^c	none		5/113 (4.4%)	4/146 (2.7%)	RR 0.62 (0.17 to 2.25)	44 per 1,000 (0.044)	17 fewer per 1,000 (0.017) (37 fewer to 55 more)
GI intole	erance	(diarrhea ai	nd nausea)) – Apremi	last vs. pl	acebo	-	•	•	•	•
983 (3 RCTs)	not serious	not serious	serious ^b	not serious	none	⊕⊕⊕⊖ MODERATE	38/486 (7.8%)	162/497 (32.6%)	RR 4.17 (3.00 to 5.80) Favors placebo	78 per 1,000 (0.078)	248 more per 1,000 (0.248) (156 more to 375 more)
Liver to	kicity ^d -	- Apremilas	t vs. placel	oo							
984 (3 RCTs)	not serious	not serious	serious ^b	not serious	none	⊕⊕⊕⊖ MODERATE	1/492 (0.2%)	6/492 (1.2%)	RD 0.01 (-0.001 to 0.02) Favors placebo	2 per 1,000 (0.002)	10 more per 1,000 (0.005) (1 fewer to 20 more)

CI: Confidence interval; RD: Risk difference RR: Risk ratio

a. A second study, by Edwards et al 2016, reported a smaller improvement in enthesitis score but could not be added to RevMan. See discussion in summary paragraph.

- b. Indirect comparison to placebo
- c. Wide CI crossing significant effect and no-effect lines
- d. Studies had slightly different cutoffs for abnormality: ALT > 150 u/l (2 studies), ALT > upper normal limit (1 study)

Working Group quality of evidence categories:

High quality: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate quality: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low quality: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect Very low quality: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

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PICO 50. In adult patients with active PsA and predominant enthesitis despite treatment with OSM, what are the benefits and harms of switching to OSM compared to switching to IL12/23i?

Summary: This question was indirectly addressed using 5 placebo-controlled RCTs (6 publications).[1-6] Apremilast was the only OSM with data suitable for this question. Two RCTs comparing ustekinumab versus placebo[1,2] and three studies comparing apremilast versus placebo[3-6] were included. None of the studies stratified outcomes by history of OSM use, which adds another layer of indirectness. However, the majority of patients in these studies had a history of OSM use, usually methotrexate.

Enthesitis and adverse events could not be compared using adjusted indirect comparisons, so these findings are presented for drug versus placebo in Table 1. Enthesitis scores (MASES) were provided by both ustekinumab studies and two of the three apremilast studies. Only one of the apremilast studies[4] provided sufficient information to import into the comparison tables. The other[6] reported a non-significant trend toward greater improvement in MASES score in patients receiving apremilast compared with placebo; the magnitude of difference was smaller than was reported by Kavanaugh et al.[4] Ustekinumab was superior to placebo for reducing the number of patients with MASES score >1. The apremilast groups had higher liver toxicity (non-significant difference with imprecision in the CI), but a statistically significant increase in GI intolerance. Serious infection rates did not differ significantly between ustekinumab and placebo groups.

Tab Biblio	le 1. (graphy:	DSM or IL1 PICO 50: In pati	2/23i con ients with PsA	npared to and enthesitis	placebo s despite OSM	in patie 1, benefit/ha	nts wit arm of sw	tching to C	e sitis de DSM compar	spite O ed to IL12	SM /23i.
		Qua	ality assessr	nent				Sum	mary of fi	ndings	
№ of participants	Risk of	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of	Study event rates (%)		Relative effect	Anticipated absolute effects	
(studies) Follow-up	bias					evidence	With placebo	With OSM or IL12/23i	(95% CI)	Risk with placebo	Risk difference with OSM or IL12/23i
Enthesitis	s score	e (MASES) (LS mean c	hange) – A	Apremilas	t vs. plac	ebo				

Tal ^{Bibli}	ble 1. (lography:	OSM or IL1 PICO 50: In pat	2/23i cor tients with PsA	npared to	s despite OS	in patie M, benefit/ha	nts wi arm of sw	th enth	esitis de OSM compar	spite O ed to IL12	SM /23i.
		Qu	ality assess	ment				Sum	mary of fi	ndings	
326 (1 RCT)	not serious	serious ^a	serious ^b	Not serious	none		165	161	-	The mean enthesitis score (MASES) (LS mean change) was 0	MD 0.9 lower (1.73 lower to 0.07 lower)
Enthesit	is (MA	SES score >	1) – Ustek	inumab vs	. placebo	1					<u>I</u>
633 (2 RCTs)	not serious	not serious	serious ^b	not serious	none	⊕⊕⊕⊖ MODERATE	171/205 (83.4%)	288/428 (67.3%)	RR 0.81 (0.74 to 0.88) Favors IL12/23i	834 per 1,000 (0.834)	158 fewer per 1,000 (0.158) (217 fewer to 100 fewer)
Serious	infectio	on – Ustekir	numab vs.	placebo	1	1					<u>I</u>
925 (2 RCTs)	not serious	serious	serious ^a	not serious	none	⊕⊕⊕⊖ MODERATE	2/309 (0.6%)	4/616 (0.6%)	RR 0.84 (0.20 to 3.50) No difference	6 per 1,000 (0.006)	1 fewer per 1,000 (5 fewer to 15 more)
GI intole	erance	(diarrhea a	nd nausea) – Apremi	last vs. p	lacebo					
Table 1. OSM or IL12/23i compared to placebo in patients with enthesitis despite OSM

Bibliography: PICO 50: In patients with PsA and enthesitis despite OSM, benefit/harm of switching to OSM compared to IL12/23i.

		Qua	ality assessi	ment			Summary of findings					
983 (3 RCTs)	not serious	not serious	serious ^b	not serious	none	⊕⊕⊕⊖ MODERATE	38/486 (7.8%)	162/497 (32.6%)	RR 4.17 (3.00 to 5.80) Favors placebo	78 per 1,000 (0.078)	248 more per 1,000 (0.248) (156 more to 375 more)	
Liver tox	Liver toxicity ^d – Apremilast vs. placebo											
984 (3 RCTs)	not serious	not serious	serious ^b	serious ^c	none	⊕⊕⊖⊖ Low	1/492 (0.2%)	6/492 (1.2%)	RR 3.32 (0.67 to 16.35)	2 per 1,000 (0.002)	5 more per 1,000 (0.005) (1 fewer to	

CI: Confidence interval; RR: Risk ratio

a. A second study, by Edwards et al 2016, reported a smaller improvement in enthesitis score but could not be imported to RevMan. See discussion in summary paragraph.

b. Indirect comparison to placebo

c. Wide CI crossing significant effect and no-effect lines

d. Studies had slightly different cutoffs for abnormality: ALT > 150 u/l (2 studies), ALT > upper normal limit (1 study)

Working Group quality of evidence categories:

High quality: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate quality: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low quality: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect Very low quality: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

References:

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PICO 51. In adult patients with active PsA and predominant enthesitis despite treatment with OSM, what are the benefits and harms of switching to OSM compared to switching to IL-17i?

<u>Summary:</u> This question was indirectly addressed using 7 placebo-controlled RCTs (in 8 publications).[1-8] Apremilast was the only OSM with data suitable for this question. In all, four RCTs comparing IL17i (ixekizumab, secukinumab and brodalumab)[1-4] versus placebo and three studies comparing apremilast versus placebo[5-8] were included. None of the studies stratified outcomes by history of OSM use, which adds another layer of indirectness. However, the majority of patients in these studies had a history of OSM use, usually methotrexate.

Enthesitis and adverse events could not be compared using adjusted indirect comparisons, so these findings are presented for drug versus placebo in Table 1. Enthesitis scores (MASES) were provided by two of the three apremilast studies. Only one of the apremilast studies[6] provided sufficient information to import into the comparison tables. The other[8] reported a non-significant trend toward greater improvement in MASES score in patients receiving apremilast compared with placebo; the magnitude of difference was smaller than was reported by Kavanaugh et al.[6] IL17i was superior to placebo for increasing the number of patients with enthesitis resolution. Two IL17i studies compared mean change in Leeds Enthesitis Index score in drug vs. placebo. One trial found a significant benefit of ixekizumab over placebo, while the other trial found no significant difference in change score between brodalumab and placebo (the estimate was imprecise). The apremilast groups had higher liver toxicity (non-significant difference with imprecision in the CI), but a statistically significant increase in GI intolerance. Rates of serious infection did not differ significantly between IL17i and placebo groups, but the effect size was imprecise.

Ta	Table 1. OSM or IL17i compared to placebo in patients with enthesitis despite OSM Bibliography: In patients with PsA and enthesitis despite OSM, benefit/harm of switching to IL17i compared to OSM.													
	Quality assessment Summary of findings													
№ of participants (studies)	Risk of	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of	Study ev rates (%	vent >)	Relative effect	Anticipate effects	ed absolute			
Follow-up	DIAS					evidence	With placebo	With OSM or IL17i	(95% CI)	Risk with placebo	Risk difference with OSM or IL17i			
Enthesitis	Enthesitis score (MASES) (LS mean change) – Apremilast vs. placebo													

Quality of evidence across all critical outcomes: Low

Τa	able 1 Bibliogi	. OSM or II raphy: In patien	_17i comp ts with PsA an	ared to p	lacebo in espite OSM, b	patient	s with of switch	enthes ning to IL	sitis des 17i compared	pite OS d to OSM.	Μ
		Qua	ality assessi	ment				Sun	nmary of f	indings	
326 (1 RCT)	not serious	serious ^a	serious ^b	not serious	none	⊕⊕℃ LOW	165	161	-	-	MD 0.9 lower (1.73 lower to 0.07 lower)
Enthesiti	s Reso	olution – IL1	7i vs. plac	ebo	•	•	•	•	•	-	•
822 (3 RCTs)	not serious	not serious	serious ^b	not serious	none	⊕⊕⊕⊖ MODERATE	58/284 (20.4%)	242/538 (45.0%)	RR 2.33 (1.80 to 3.02) Favors IL17i	204 per 1,000 (0.204)	272 more per 1,000 (0.272) (163 more to 413 more)
Enthesiti	s score	e (Leeds Ent	thesitis Ind	dex) – I xe	kizumab v	/s. placel	00				
197 (1 RCT)	not serious	not serious	serious ^b	not serious	none	⊕⊕⊕⊖ MODERATE	102	95	-	-	MD 0.7 lower (1.37 lower to 0.03 lower)
Enthesiti	s score	e (Leeds En	thesitis Ind	dex) – Bro	dalumab	vs. place	bo	•	•		
112 (1 RCT)	not serious	not serious	serious ^b	serious ^d	none	⊕⊕ Low	55	57	-	-	MD 0.1 lower (1.06 lower to 0.86 higher)

Ta	able 1 Bibliogr	. OSM or II	_17i comp ts with PsA and	ared to p d enthesitis de	lacebo in espite OSM, b	patient	s with of switch	enthes	sitis desp 17i compared	bite OS to OSM.	М		
		Qua	ality assessr	nent				Sun	nmary of f	indings			
Serious ir	nfectio	on – IL17i vs	s. placebo										
1189 (3 RCTs)	not serious	not serious	serious ^b	serious ^d	none	⊕⊕ Low	5/406 (1.2%)	18/685 (2.6%)	RR 1.79 (0.75 to 4.30)	12 per 1,000 (0.012)	9 more per 1,000 (0.009) (3 fewer to 52 more)		
GI intoler	31 intolerance (diarrhea and nausea) – Apremilast vs. placebo												
983 (3 RCTs)	not serious	not serious	serious ^b	not serious	none	⊕⊕⊕⊖ MODERATE	38/486 (7.8%)	162/497 (32.6%)	RR 4.17 (3.00 to 5.80) Favors placebo	78 per 1,000 (0.078)	248 more per 1,000 (0.248) (156 more to 375 more)		
Liver toxi	city ^e –	- Apremilast	t vs. placeb	00	<u></u>	1	ł	ł	<u> </u>	1	1		
984 (3 RCTs)	not serious	not serious	serious ^b	serious ^d	none	⊕⊕⊖⊖ Low	1/492 (0.2%)	6/492 (1.2%)	RR 3.32 (0.67 to 16.35)	2 per 1,000 (0.002)	5 more per 1,000 (0.005) (1 fewer to 31 more)		

a. A second study, by Edwards et al 2016, reported a smaller improvement in enthesitis score but could not be imported to RevMan. See discussion in summary paragraph.

b. Indirect comparison to placebo

c. Substantial heterogeneity between secukinumab studies.

d. Wide CI crossing significant effect and no-effect lines

e. Studies had slightly different cutoffs for abnormality: ALT > 150 u/l (2 studies), ALT > upper normal limit (1 study)

Working Group quality of evidence categories:

High quality: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate quality: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low quality: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect Very low quality: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

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PICO 52. In adult patients with active PsA and predominant enthesitis despite treatment with OSM, what are the benefits and harms of switching to TNFi compared to switching to IL12/23i?

Summary: This PICO was indirectly addressed by 4 studies (5 publications).[1-5] Two RCTs (3 publications) compared TNFis with placebo,[1-3] while two RCTs compared ustekinumab with placebo.[4,5]

Enthesitis and adverse events could not be compared using adjusted indirect comparisons, so these findings are presented for drug versus placebo in Table 1. Both ustekinumab and golimumab were superior to placebo in reducing the number of patients with enthesitis or the number of patients with severe enthesitis. There was no significant difference between TNFi and placebo in serious infections, nausea and diarrhea, but all findings were imprecise. Ustekinumab showed no significant difference with placebo in rate of serious infections.

Quality of evidence across all critical outcomes: Low

Tab	Table 1. TNFi or IL12/23i compared to placebo in patients with enthesitis despite OSM Bibliography: PICO 52: TNFi compared to IL12/23i for patients with active PsA and enthesitis despite OSM													
Quality assessment Summary of findings														
Nº of participants	Risk of	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of	Study ev (%)	ent rates	Relative effect	Anticipat effects	ed absolute			
Follow-up	DIAS					evidence	With placebo	With TNFi or IL12/23i	(95% CI)	Risk with placebo	Risk difference with TNFi or IL12/23i			
Enthesitis	Enthesitis (MASES) score >1 – Ustekinumab vs. placebo													
633 (2 RCTs)	$\begin{array}{c} 333\\ (2 \text{ RCTs}) \end{array} \begin{array}{c} \text{not} \\ \text{serious} \end{array} \begin{array}{c} \text{not} \\ \text{serious} \end{array} \begin{array}{c} \text{not} \\ \text{serious} \end{array} \begin{array}{c} \text{serious} \\ \text{serious} \end{array} \begin{array}{c} \text{serious} \\ \text{serious} \end{array} \begin{array}{c} \text{not} \end{array} \begin{array}{c} \text{not} \end{array} \begin{array}{c} \text{not} \end{array} \begin{array}{c} \text{not} \end{array} \begin{array}{$													
Enthesitis	Enthesitis, 14 weeks – Golimumab vs. placebo													

Tab	Table 1. TNFi or IL12/23i compared to placebo in patients with enthesitis despite OSM Bibliography: PICO 52: TNFi compared to IL12/23i for patients with active PsA and enthesitis despite OSM													
		Qua	ality assessr	ment				Sum	mary of fi	ndings				
247 (1 RCT)	not serious	not serious	serious ^a	not serious	none	⊕⊕⊕⊖ MODERATE	75/105 (71.4%)	78/142 (54.9%)	RR 0.77 (0.63 to 0.93) Favors GOL	714 per 1,000 (0.714)	164 fewer per 1,000 (0.164) (264 fewer to 50 fewer)			
Serious ir	Serious infection – Ustekinumab vs. placebo													
925 (2 RCTs)	not serious	serious	serious ^a	not serious	none	⊕⊕⊕⊖ MODERATE	2/309 (0.6%)	4/616 (0.6%)	RR 0.84 (0.20 to 3.50) No difference	6 per 1,000 (0.006)	1 fewer per 1,000 (5 fewer to 15 more)			
Serious ir	nfectio	on – TNFi vs	. placebo											
533 (2 RCTs)	not serious	not serious	serious ^a	serious ^b	none	⊕⊕⊖⊖ Low	5/249 (2.0%)	3/284 (1.1%)	RR 0.55 (0.11 to 2.77)	20 per 1,000 (0.020)	9 fewer per 1,000 (0.009) (18 fewer to 36 more)			
Diarrhea	– TNF	i vs. placebo	D											

Table 1. TNFi or IL12/23i compared to placebo in patients with enthesitis despite OSM

Bibliography: PICO 52: TNFi compared to IL12/23i for patients with active PsA and enthesitis despite OSM

		Qua	ality assessr	ment			Summary of findings					
259 (1 RCT)	not serious	not serious	serious ^a	serious ^b	none	⊕⊕⊖⊖ Low	4/113 (3.5%)	5/146 (3.4%)	RR 0.97 (0.27 to 3.52)	35 per 1,000 (0.035)	1 fewer per 1,000 (0.001) (26 fewer to 89 more)	
Nausea –	TNFi	vs. placebo										
259 (1 RCT)	not serious	not serious	serious ^a	serious ^b	none	⊕⊕⊖⊖ Low	5/113 (4.4%)	4/146 (2.7%)	RR 0.62 (0.17 to 2.25)	44 per 1,000 (0.044)	17 fewer per 1,000 (0.017) (37 fewer to 55 more)	

CI: Confidence interval; RR: Risk ratio

a. Indirect comparison to placebo

b. Wide CI crossing significant effect and no-effect lines

Working Group quality of evidence categories:

High quality: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate quality: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low quality: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect Very low quality: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

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PICO 53. In adult patients with active PsA and predominant enthesitis despite treatment with OSM, what are the benefits and harms of switching to TNFi compared to switching to IL17i?

<u>Summary</u>: This PICO was addressed indirectly by 5 double-blind RCTs (6 publications) involving PsA patients or psoriasis patients comparing either TNFi or IL17i to placebo.[1-6] The relevant outcomes included enthesitis resolution, Leeds enthesitis index change from baseline, and adverse events. All drugs showed statistically significant improvements in enthesitis resolution over placebo.

Since both TNFi and IL17i studies reported enthesitis resolution, we compared the findings for this outcome between drug classes using adjusted indirect comparisons (Table 1). Golimumab was the only TNFi with a study that reported this outcome. The comparison found no significant difference between golimumab and IL17i (secukinumab and ixekizumab) but was imprecise due to a wide CI that overlapped the line of no effect.

Adverse events are presented for drug versus placebo in Table 2. Both IL17i and golimumab were superior to placebo in reducing the number of patients with enthesitis. Two IL17i studies compared mean change in Leeds Enthesitis Index score in drug vs. placebo. One trial found a significant benefit of ixekizumab over placebo, while the other trial found no significant difference in change score between brodalumab and placebo (the estimate was imprecise). There was no significant difference between TNFi and placebo in serious infections, nausea and diarrhea, but all findings were imprecise. IL17i showed no significant difference with placebo in rate of serious infections.

Quality of evidence across all critical outcomes: Low

Table	Table 1. TNFi compared to IL17i for patients with active PsA and predominant enthesitis despite OSM Bibliography: PICO 53: TNFi compared to IL17i for patients with active PsA and enthesitis despite OSM											
Quality assessment Summary of findings												
Nº of participants	Risk of	sk Inconsistency as	Indirectness	Imprecision	Publication bias	Overall quality of			Relative effect	Anticipat absolute	ed effects	
Follow-up	DIAS					evidence	IL17i	TNFi	(95% CI)	Risk with IL17i	Risk difference with TNFi	
Enthesitis	Inthesitis resolution, Bucher adjusted indirect comparison											

Table 1. TNFi compared to IL17i for patients with active PsA and predominant enthesitisdespite OSM

Bibliography: PICO 53: TNFi compared to IL17i for patients with active PsA and enthesitis despite OSM

		Qua	ality assessr	ment				Sum	mary of fi	ndings	
1069 (4 RCTs)	Not serious	Not serious	Serious ^a	Serious⁵	None	⊕⊕⊖⊖ Low	IL17i	GOL	RR 0.71 (0.39 to 1.30)	450 per 1,000 (0.450)	130 fewer per 1000 (0.130) (274 fewer to 135 more)

CI: Confidence interval; RR: Risk ratio

a. Indirect comparison, all studies compared drug to placebo, only one trial had 100% patients with prior OSM exposure, most had 50-83% of patients with prior OSM exposure.

b. Wide 95% CI

Ta	Table 2. TNFi or IL17i compared to placebo in patients with enthesitis despite OSM Bibliography: PICO 53: TNFi compared to IL17i for patients with active PsA and enthesitis despite OSM												
Quality assessment Summary of findings													
Nº of participants (studies)	Risk of	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of	Study ev rates (%	ent)	Relative effect	Anticipate effects	ed absolute		
(studies) Follow-up	bias					evidence	With placebo	With TNFi or IL17i	(95% CI)	Risk with placebo	Risk difference with TNFi or IL17i		
Enthesitis Resolution – IL17i vs. placebo													

Ta	able 2	. TNFi or II Bibliography: PI	_17i comp co 53: TNFi co	ared to p	lacebo in 17i for patien	patient ts with activ	s with re PsA and	enthes denthesit	s itis des is despite OS	oite OS ™	М			
		Qua	ality assessr	nent				Sun	nmary of f	indings				
822 (3 RCTs)	not serious	not serious	serious ¹	not serious	none	⊕⊕⊕⊖ MODERATE	58/284 (20.4%)	242/538 (45.0%)	RR 2.23 (1.36 to 3.66) Favors IL17i	204 per 1,000 (0.204)	251 more per 1,000 (0.251) (74 more to 543 more)			
Enthesitis	nthesitis, 14 weeks – Golimumab vs. placebo													
247 (1 RCT)	not serious	not serious	serious ^a	not serious	none	⊕⊕⊕⊖ MODERATE	75/105 (71.4%)	78/142 (54.9%)	RR 0.77 (0.63 to 0.93) Favors GOL	714 per 1,000 (0.714)	164 fewer per 1,000 (0.164) (264 fewer to 50 fewer)			
Enthesitis	s score	e (Leeds En	thesitis Ind	dex) – I xe	kizumab v	/s. placek	00							
197 (1 RCT)	not serious	not serious	serious ^a	not serious	none	⊕⊕⊕⊖ MODERATE	102	95	-	-	MD 0.7 lower (1.37 lower to 0.03 lower)			
Enthesitis	Enthesitis score (Leeds Enthesitis Index) – Brodalumab vs. placebo													
112 (1 RCT)	not serious	not serious	serious ^a	serious ^b	none	⊕⊕℃ LOW	55	57	-	-	MD 0.1 lower (1.06 lower to 0.86 higher)			

Ta	Table 2. TNFi or IL17i compared to placebo in patients with enthesitis despite OSM Bibliography: PICO 53: TNFi compared to IL17i for patients with active PsA and enthesitis despite OSM												
		Qua	ality assessr	nent				Sur	nmary of f	indings			
Serious in	nfectio	on – IL17i ve	s. placebo										
1189 (3 RCTs)	not serious	not serious	serious ^a	serious ^b	none	⊕⊕ Low	5/406 (1.2%)	18/685 (2.6%)	RR 1.79 (0.75 to 4.30)	12 per 1,000 (0.012)	9 more per 1,000 (0.009) (3 fewer to 52 more)		
Serious infection – TNFi vs. placebo													
533 (2 RCTs)	not serious	not serious	serious ^a	serious ^b	none	⊕⊕⊖⊖ Low	5/249 (2.0%)	3/284 (1.1%)	RR 0.55 (0.11 to 2.77)	20 per 1,000 (0.020)	9 fewer per 1,000 (0.009) (18 fewer to 36 more)		
Diarrhea	– TNF	i vs. placebo	D	•		•				1			
259 (1 RCT)	not serious	not serious	serious ^a	serious ^b	none	⊕⊕⊖⊖ Low	4/113 (3.5%)	5/146 (3.4%)	RR 0.97 (0.27 to 3.52)	35 per 1,000 (0.035)	1 fewer per 1,000 (0.001) (26 fewer to 89 more)		
Nausea –	TNFi	vs. placebo	I	<u>I</u>	<u>-</u>	1		<u>.</u>		1			
259 (1 RCT)	not serious	not serious	serious ^a	serious ^b	none	⊕⊕⊖⊖ Low	5/113 (4.4%)	4/146 (2.7%)	RR 0.62 (0.17 to 2.25)	44 per 1,000 (0.44)	17 fewer per 1,000 (0.017) (37 fewer to 55 more)		

Cl: Confidence interval; RR: Risk ratioa. Indirect comparison to placebob. Wide Cl crossing significant effect and no-effect lines

Working Group quality of evidence categories:

High quality: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate quality: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low quality: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect Very low quality: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

References:

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PICO 54. In adult patients with active PsA and predominant enthesitis despite treatment with OSM, what are the benefits and harms of switching to IL12/23i compared to switching to IL17i?

Summary: This PICO was addressed indirectly by 7 double-blind RCTs comparing either IL12/23i or IL17i to placebo in patients with PsA.[1-7] The relevant outcomes included enthesitis resolution, enthesitis (MASES) score >1, Leeds Enthesitis Index change from baseline, and adverse events.

Infection was the only relevant outcome that was comparable between IL12/23i studies and IL17i studies, so we compared infection rates between drug classes using adjusted indirect comparisons for various outcomes (Table 1). No significant between-class difference was identified, but the effect estimate had serious imprecision due to a wide CI that overlapped the line of no effect.

Enthesitis findings could not be compared using adjusted indirect comparisons because of differences in measurement between the drug classes, so these findings are presented for drug versus placebo in Table 2. IL17i and ustekinumab were superior to placebo in enthesitis resolution and reducing the number of patients with enthesitis and a MASES score >1, respectively. Two IL17i studies also compared mean change in Leeds Enthesitis Index score in drug vs. placebo. One trial found a significant benefit of ixekizumab over placebo, while the other trial found no significant difference in change score between brodalumab and placebo (the estimate was imprecise). Serious infection rates were so low for IL17i and ustekinumab that adjusted indirect comparison was not performed. Neither drug class showed significantly different serious infection rates compared to placebo groups, but the findings were imprecise for IL17i due to wide 95% confidence intervals.

Quality of evidence across all critical outcomes: Moderate

Table 1. IL12/23i compared to IL17i for patients with active PsA and predominant enthesitisdespite OSM

Bibliography: PICO 54: IL12/23i compared to IL17i for patients with active PsA and enthesitis despite OSM

	Quality assessment							Summary of findings					
Nº of participants (studies)	Risk of	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of			Relative effect	Anticipate effects	ed absolute		
(studies) Follow-up	bias					evidence	IL12/23i	IL17i	(95% CI)	Risk with IL12/23i	Risk difference with IL17i		
Infection	*	•		•	•	•	•	•	•	•	•		

Table 1. IL12/23i compared to IL17i for patients with active PsA and predominant enthesitisdespite OSM

Bibliography: PICO 54: IL12/23i compared to IL17i for patients with active PsA and enthesitis despite OSM

	Quality assessment							Summary of findings				
1527 (4 RCTs)	Not serious	Not serious	Not serious ^a	Serious ^b	None	⊕⊕⊕⊖ MODERATE	IL12/23i	IL17i	RR 1.26 (0.81 to 1.97)	211 per 1,000 (0.211)	55 more per 1000 (0.055) (40 fewer to 205 more)	

CI: Confidence interval; RR: Risk ratio

a. Indirect comparison, all studies compared drug to placebo, only 50-83% of patients had prior OSM exposure in most trials, but patient characteristics and prior drug exposure are similar between drug classes.

b. Wide 95% CI

Tabl	Table 2. IL12/23i or IL17i compared to placebo in patients with enthesitis despite OSM Bibliography: PICO 54: IL12/23i compared to IL17i for patients with active PsA and enthesitis despite OSM										
		Qua	ality assessr	nent				Sum	mary of fi	ndings	
Nº of participants (studies)	Risk of	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of	Study event rates (%)		Relative effect	Anticipated absolute effects	
(studies) Follow-up	bias					evidence	With placebo	With IL12/23i or IL17i	(95% CI)	Risk with placebo	Risk difference with IL12/23i or IL17i
Enthesitis	s Reso	olution – IL1	7i vs. plac	ebo	•	•	•	•	•		

Table 2. IL12/23i or IL17i compared to placebo in patients with enthesitis despite OSM Bibliography: PICO 54: IL12/23i compared to IL17i for patients with active PsA and enthesitis despite OSM													
		Qua	ality assessr	nent			Summary of findings						
822 (3 RCTs)	not serious	not serious	serious ^a	not serious	none	⊕⊕⊕⊖ MODERATE	58/284 (20.4%)	242/538 (45.0%)	RR 2.23 (1.36 to 3.66) Favors IL17i	204 per 1,000 (0.204)	251 more per 1,000 (0.251) (74 more to 543 more)		
Enthesitis	s score	e (Leeds Ent	hesitis Ind	dex) – I xel	kizumab v	/s. placeb	oo						
197 (1 RCT)	not serious	not serious	serious ^a	not serious	none	⊕⊕⊕⊖ MODERATE	102	95	-	-	MD 0.7 lower (1.37 lower to 0.03 lower)		
Enthesitis	s score	e (Leeds Ent	hesitis Ind	lex) – Bro	dalumab	vs. place	bo	•	•	•	•		
112 (1 RCT)	not serious	not serious	serious ^a	serious ^b	none	⊕⊕℃ LOW	55	57	-	-	MD 0.1 lower (1.06 lower to 0.86 higher)		
Enthesitis	s (MAS	SES) score >	1 – Usteki	numab vs	. placebo	•	•						
633 (2 RCTs)	not serious	not serious	serious ^a	not serious	none	⊕⊕⊕⊖ MODERATE	171/205 (83.4%)	288/428 (67.3%)	RR 0.81 (0.74 to 0.88) Favors IL12/23i	834 per 1,000 (0.834)	158 fewer per 1,000 (0.158) (217 fewer to 100 fewer)		

Table 2. IL12/23i or IL17i compared to placebo in patients with enthesitis despite OSM Bibliography: PICO 54: IL12/23i compared to IL17i for patients with active PsA and enthesitis despite OSM											
		Qua	ality assessr		Summary of findings						
Serious ir	on – IL17i v										
1189 (3 RCTs)	not serious	not serious	serious ^a	serious ^b	none	⊕⊕ Low	5/406 (1.2%)	18/685 (2.6%)	RR 1.79 (0.75 to 4.30)	12 per 1,000 (0.012)	9 more per 1,000 (0.009) (3 fewer to 52 more)
Serious ir	nfectio	on – Ustekin	umab vs. p	olacebo	•	•	•	•	•	•	•
925 (2 RCTs)	not serious	serious	serious ^a	not serious	none	⊕⊕⊕⊖ MODERATE	2/309 (0.6%)	4/616 (0.6%)	RR 0.84 (0.20 to 3.50) No difference	6 per 1,000 (0.006)	1 fewer per 1,000 (5 fewer to 15 more)

CI: Confidence interval; RR: Risk ratio

a. Indirect comparison to placebo

b. Wide 95% CI that crosses line of no effect

Working Group quality of evidence categories:

High quality: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate quality: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low quality: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect Very low quality: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

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PICO 55. In adult patients with active PsA and predominant enthesitis despite treatment with NSAIDs, what are the benefits and harms of switching to tofacitinib compared to switching to OSM?

Summary: The literature searches did not identify any studies that addressed this PICO question.

Quality of evidence across all critical outcomes: Very low

Special Populations

PICO 56: In patients with active PsA, what are the benefits and harms of vaccination with killed vaccines prior to starting biologic compared to vaccination while using a biologic?

<u>Summary</u>: Only one study directly addressed this PICO question[1] This was an RCT comparing the antibody response of patients with PsA randomized to etanercept or placebo and subsequently vaccinated with a 23-valent pneumococcal vaccine. Table 1 shows that the percentage of patients with 2-fold or 4-fold increase in antibody titers to five different antigens did not differ significantly between groups, but the 95% CIs showed serious imprecision in all effect estimates.

Adverse events were indirectly addressed by six studies evaluating vaccination safety in patients (mostly with rheumatoid arthritis) treated with TNFi.[2-7] All of the studies evaluated adverse events and serious adverse events in patients vaccinated during treatment with TNFi or Placebo. No significant difference was found in all studies between TNFi and Placebo patients. In one study, even though there were no cases reported in the category of tuberculosis (TB) in either group during the RCT, 2 cases (0.9%) were reported during the open-label extension (OLE) after CZP treatment, among which one was considered related to study drug by the investigator. In the same study, serious adverse events occurred at a rate of 1.8% in the CZP group during the RCT period, and in 6.8% of CZP-treated patients overall (combined RCT and OLE), which might be attributable to both longer duration and the OLE period. In another study, the rate of infectious adverse events was statistically significantly higher in the placebo treatment group [23.5% (27/115)] versus the adalimumab group [12.6% (14/111)] (p = 0.039).

Table 1. Antibody Response in Patients Who Received Etanercept or Placebo Prior to Vaccination with Pneumococcal Vaccine Bibliography: Etanercept versus Placebo for PV vaccinated patients. Summary of findings Quality assessment Nº of Study event rates Risk Inconsistency Indirectness Imprecision Publication Overall Relative Anticipated absolute participants of bias quality (%) effect effects (95% CI) (studies) bias of With With Risk Risk evidence Follow-up Placebo ETN with difference Placebo with ETN 2-fold increase in titer 9V

Quality of evidence across all critical outcomes: Low

Tak	Table 1. Antibody Response in Patients Who Received Etanercept or Placebo Prior to Vaccination with Pneumococcal Vaccine Bibliography: Etanercept versus Placebo for PV vaccinated patients.												
		Qua	lity assessr	nent			Summary of findings						
184 (1 RCT)	not serious	not serious	serious ^a	serious ^b	none		53/90 (58.9%)	47/94 (50.0%)	OR 0.70 (0.39 to 1.25)	589 per 1,000	88 fewer per 1,000 (230 fewer to 53 more)		
2-fold ind	rease	in titer 14											
184 (1 RCT)	not serious	not serious	serious ^a	serious ^b	none		50/90 (55.6%)	55/94 (58.5%)	OR 1.13 (0.63 to 2.02)	556 per 1,000	30 more per 1,000 (115 fewer to 161 more)		
2-fold ind	rease	in titer 180	2										
184 (1 RCT)	not serious	not serious	serious ^a	serious ^b	none		56/90 (62.2%)	58/94 (61.7%)	OR 0.98 (0.54 to 1.77)	622 per 1,000	5 fewer per 1,000 (151 fewer to 122 more)		
2-fold ind	rease	in titer 19F	-				•				•		
184 (1 RCT)	not serious	not serious	serious ^a	serious ^b	none		36/90 (40.0%)	33/94 (35.1%)	OR 0.81 (0.45 to 1.48)	400 per 1,000	49 fewer per 1,000 (169 fewer to 97 more)		
2-fold ind	rease	in titer 23F	:	•	·	·	·	·	·	·	·		

Table 1. Antibody Response in Patients Who Received Etanercept or Placebo Prior to Vaccination with Pneumococcal Vaccine Bibliography: Etanercept versus Placebo for PV vaccinated patients.													
	Quality assessment								Summary of findings				
184 (1 RCT)	not serious	not serious	serious ^a	serious ^b	none		52/90 (57.8%)	48/94 (51.1%)	OR 0.76 (0.43 to 1.36)	578 per 1,000	68 fewer per 1,000 (207 fewer to 73 more)		
4-fold inc	rease	of titer 9V											
184 (1 RCT)	not serious	not serious	serious ^a	serious ^b	none		41/90 (45.6%)	32/94 (34.0%)	OR 0.62 (0.34 to 1.12)	456 per 1,000	114 fewer per 1,000 (234 fewer to 28 more)		
4-fold inc	rease	in titer 14	•				•	•		•	•		
184 (1 RCT)	not serious	not serious	serious ^a	serious ^b	none		41/90 (45.6%)	40/94 (42.6%)	OR 0.89 (0.49 to 1.59)	456 per 1,000	29 fewer per 1,000 (165 fewer to 115 more)		
4-fold inc	rease	in titer 180)				<u> </u>						
184 (1 RCT)	34 RCT) not serious not serious serious a serious a serious b none $\bigoplus_{LOW} 42/90$ $42/90$ (40.4%) (40.4%) (40.4%) (40.4%) (40.4%) (40.4%) (40.4%) (40.4%) $(1000$ (1000) (100) (1000) $(10$												
4-fold inc	rease	in titer 19F	:	·	·		·	·	·	·	·		

Tat	Table 1. Antibody Response in Patients Who Received Etanercept or Placebo Prior to Vaccination with Pneumococcal Vaccine Bibliography: Etanercept versus Placebo for PV vaccinated patients.											
	Quality assessment Summary of findings											
184 (1 RCT)	not serious	not serious	serious ^a	serious ^b	none	⊕⊕⊖⊖ Low	20/90 (22.2%)	18/94 (19.1%)	OR 0.83 (0.41 to 1.69)	222 per 1,000	31 fewer per 1,000 (117 fewer to 103 more)	
4-fold inc	crease	in titer 23F		•	•		•	•	•		•	
184 (1 RCT)not seriousserious a serious b none $\bigoplus \bigoplus \bigoplus \bigoplus B = 1000$ $31/90$ (34.4%) $25/94$ (26.6%)OR 0.69 (0.37 to 1.30) 344 per per 1,000 (182 fewer to 61 more)												

CI: Confidence interval; OR: Odds ratio

Explanations

a. Outcomes not direct and only a single vaccine is used, which may not be representative of the immune response to other vaccines

b. Wide CI crossing significant effect and no-effect lines

Author, year	Study type	Duratio n	Population Description	Treatment given to relevant population	Results
Ribeiro, 2013	Cohort study	21 days	99 Patients with RA	Influenza A/H1N1 killed virus vaccine given to 11 pts with Abatacept (RA-ABA), 33 MTX (RA-MTX) and 55 healthy controls	The rates of minor side effects were comparable: 55% in RA- ABA patients, 39% in RA-MTX patients, and 40% in control groups (P=0.64). Severe side effects were not reported during the followup period.
Migita, 2015	Randomized, double-blind placebo- controlled study	6 weeks	703 patients with RA	PPSV23 administered to 353 patients on MTX, ABA, other biologics, or controls	There were no reported adverse events associated with PPSV23 vaccination.
Kivitz A, 2014	Single-blind Randomized placebo- controlled	6-week	224 patients with RA	Pneumococcal (polysaccharide 23) and influenza vaccines administered at Week 2 to 110 patients with CZP and 114 with	AE occurred 62.3% in placebo and 63.6% CZP groups. Serious AE occurred in 1 patient (0.9%) in the placebo group, and in 2 patients (1.8%) in the CZP group. Serious AE occurred in 6.8% of

Author,	Study type	Duratio	Population	Treatment given to relevant	Results
year		n	Description	population	
	trial			Placebo	CZP-treated patients overall (combined RCT and OLE). Even though there were no cases reported in the category of tuberculosis (TB) in either group during the RCT, 2 cases (0.9%) were reported during the OLE after CZP treatment, among which one was considered related to study drug by the investigator
Franca I, 2012	Cohort study	21 days	236 pts with RA, SpA and PsA, and 117 healthy controls	anti-influenza A H1N1/2009 vaccine administered to 120 pts on anti-TNF agents, 116 inflammatory arthritis patients on DMARDs, and 117 healthy controls.	Only mild systemic reactions were more often observed in patients on anti-TNF compared with healthy controls: fever (8.3% vs 0.9%, P = 0.01), arthralgia (12.5% vs 4.3%, P = 0.03), and nasal congestion (13.3% vs 4.3%, P = 0.014). No severe adverse event was reported in any group.
Elkayam O, 2008	Cohort study	4 to 6 weeks	43 RA patients and 18 AS patients	Split-virion inactivated vaccine containing15g hemagglutinin/dose of each of A/New Caledionan/20/1999 (H1N1), A/Wisconsin/67/2005 (H3N2), and B/Malaysia/2506/2004 (M) was given to 22 pts on the day of administration of infliximab, while 16 received the vaccine 3 weeks after infliximab.	No adverse effects other than injection site pain were recorded.
Kaine J, 2007	Double- blind, randomized trial	4 weeks	208 adult patients with RA	Pneumococcal and influenza vaccines were administered on Day 8 to patients who received adalimumab (99) or placebo (109).	During the blinded period of the study no deaths were reported, and one patient receiving placebo reported a serious AE. A slightly greater percentage of patients in the placebo group reported an AE than did patients in the adalimumab group [54.8% (63/115) vs 45.9% (51/111), respectively]. The most frequently reported treatment-emergent AE occurring during the blinded period of the study were upper respiratory tract infection and injection site reaction; both were reported more frequently by placebo-treated patients. There were no serious infectious AE, malignancies, or opportunistic infections, including tuberculosis, reported during the double-blind period. The rate of infectious AE was statistically significantly higher in the placebo treatment group [23.5% (27/115)] versus the adalimumab group [12.6% (14/111)] (p = 0.039).

References

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- 6. Ribeiro AC, Laurindo IM, Guedes LK, Saad CG, Moraes JC, Silva CA, et al. Abatacept and reduced immune response to pandemic 2009 influenza A/H1N1 vaccination in patients with rheumatoid arthritis. Arthritis Care Res (Hoboken). 2013;65(3):476-480.
- 7. Migita K, Akeda Y, Akazawa M, Tohma S, Hirano F, Ideguchi H, et al. Effect of abatacept on the immunogenicity of 23-valent pneumococcal polysaccharide vaccination (PPSV23) in rheumatoid arthritis patients. Arthritis Res Ther. 2015;17:357.

PICO 57. In patients with active PsA, what are the benefits and harms of vaccination with live attenuated vaccines prior to starting biologic compared to vaccination while using a biologic?

<u>Summary</u>: The literature searches identified one large retrospective cohort study of patients with immune-mediated diseases who received the Herpes Zoster vaccine that addressed this PICO question.[1] As a retrospective observational study the GRADE rating started at low; since only 3% of patients had PsA (the rest had psoriasis, ankylosing spondylitis, rheumatoid arthritis, and inflammatory bowel disease); the rating was further downgraded to very low due to indirectness of the patient population. The Herpes Zoster incidence rate was similar in patients who received biologics and patients who did not receive biologics (see table below).

Author,	Study type	Duration	Population	Treatment given to		Results	
year			Description	relevant population			
Zhang et al.	Retrospective cohort study	2 years	463,541 patients over	Herpes Zoster vaccine given to pts with TNFi		HZ cases	HZ incidence rate
2012			oo with RA, psoriasis, psoriatic arthritis, ankylosing spondylitis,	and non-TNFT biologics, DMARDs, and oral glucocorticoids	Overall Medications (mutually exclusive groups): Biologics (regardless of concomitant DMARDs or oral glucocorticoids)	138 14	6.7 (5.7-7.9) 8.5(5.1-14.4)
			and/or IBD		Anti-TNF therapies DMARDs (without biologics but regardless of oral glucocorticoids)	12 25	8.5(4.8-15.0) 7.0(4.7-10.3)
					Oral glucocorticoids alone	21	10.3(6.7-15.8)

Quality of evidence across all critical outcomes: Very low

References

1. Zhang J, Xie F, Delzell E, Chen L, Winthrop K, Lewis JD, et al.. Association between Vaccination for Herpes Zoster and Risk of Herpes Zoster Infection among Older Patients with Selected Immune-mediated Diseases. JAMA. 2012 July 4; 308(1): 43–49.

Comorbidities

PICO 58. In adult patients with active PsA and IBD despite treatment with an OSM, what are the benefits and harms of switching to TNFi (monoclonal antibodies [MABs]) vs. switching to TNFi soluble receptor biologic (i.e. etanercept)?

<u>Summary</u>: The literature searches did not identify any direct studies that addressed this PICO question. Two systematic reviews compared TNFi (monoclonal antibodies [MABs]) to placebo, one in Crohn's Disease patients,[1] and another in Ulcerative Colitis patients.[2] One RCT compared etanercept with Placebo in patients with Crohn's Disease.[3] The systematic reviews revealed that anti-TNF MABs result in a 1.66-fold higher likelihood of induction of remission (95% CI: 1.17–2.36) and 1.43-fold higher likelihood of induction of response (95% CI: 1.17–1.73) compared to placebo in patients with Crohn's Disease. For patients with ulcerative colitis, meta-analyses found a 2.45-fold higher likelihood of induction of remission and 1.65-fold higher likelihood of induction of response compared to placebo (RR: 2.45, 95% CI: 1.72–3.47 and RR: 1.65, 95% CI: 1.37–1.99 respectively). In the RCT the rates of clinical response at week 2, 4 and 8 were similar in patients treated with etanercept compared with placebo, RR=1.91 (0.8-4.75), 0.87 (0.43-1.76), and 1.01 (0.41-2.52) respectively.

Quality of evidence across all critical outcomes: Moderate

TNFi (MAB) compared to Placebo for Crohn's disease Bibliography: TNFi (MAB) compared to Placebo for Crohn's disease												
		Qua	ality assess	ment			Summary of findings					
Nº of participants	Risk of	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of	Study ev rates (%	ent)	Relative effect	Anticipate effects	ed absolute	
(studies) Follow-up	DIAS					evidence	With Placebo	With TNFi (MAB)	(95% CT)	Risk with Placebo	Risk difference with TNFi (MAB)	
Induction	of re	mission end	lpoint									
1771 (6 RCTs)	not serious	not serious	serious ^a	not serious	none	⊕⊕⊕⊖ MODERATE	150/882 (17.0%)	227/889 (25.5%)	RR 1.66 (1.17 to 2.36)	170 per 1,000	112 more per 1,000 (29 more to 231 more)	
Induction	of re	sponse end	point									
1771 (6 RCTs)	not serious	not serious	serious ^a	not serious	none	⊕⊕⊕⊖ MODERATE	246/882 (27.9%)	346/889 (38.9%)	RR 1.43 (1.17 to 1.73)	279 per 1,000	120 more per 1,000 (47 more to 204 more)	
Maintena	nce of	f remission	endpoint									
1690 (5 RCTs)	$\begin{array}{c} \begin{array}{c} \begin{array}{c} \begin{array}{c} 0 \\ 0 \\ 0 \\ \end{array} \end{array} \\ \begin{array}{c} 0 \\ 0 \\ \end{array} \end{array} \\ \begin{array}{c} 0 \\ 0 \\ \end{array} \\ \begin{array}{c} 0 \\ 0 \\ \end{array} \end{array} \\ \begin{array}{c} 0 \\ 0 \\ 0 \\ \end{array} \\ \begin{array}{c} 0 \\ 0 \\ 0 \\ \end{array} \\ \begin{array}{c} 0 \\ 0 \\ 0 \\ 0 \\ \end{array} \\ \begin{array}{c} 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ \end{array} \\ \begin{array}{c} 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ \end{array} \\ \begin{array}{c} 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ 0 $											
Maintena	nce of	f response e	endpoint									

1467 (4 RCTs)	not serious	not serious	serious ^a	not serious	none	⊕⊕⊕⊖ MODERATE	184/729 (25.2%)	315/738 (42.7%)	RR 1.68 (1.46 to 1.93)	252 per 1,000	172 more per 1,000 (116 more to 235 more)
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CI: Confidence interval; RR: Risk ratio

Explanations a. Indirect comparison to placebo

	TNFi (MAB) compared to Placebo for Ulcerative colitis Bibliography: TNFi (MAB) compared to Placebo for Ulcerative colitis													
Quality assessment								Summary of findings						
Nº of participants	Risk of	Inconsistency	ency Indirectness Imprecision Publication Overall quality evidence	Imprecision	Publication bias	Overall quality of	Study event rates (%)		Relative effect	Anticipated absolute effects				
(studies) Follow-up	DIAS			evidence	With Placebo	With TNFi (MAB)	(95% CI)	Risk with Placebo	Risk difference with TNFi (MAB)					
Induction	Induction of remission endpoints													
1823 (6 RCTs)	not serious	not serious	serious ^a	not serious	none	⊕⊕⊕⊖ MODERATE	82/912 (9.0%)	210/911 (23.1%)	RR 2.45 (1.72 to 3.47)	90 per 1,000	130 more per 1,000 (65 more to 222 more)			
Induction	n of re	sponse end	point											
1780 (5 RCTs)	not serious	not serious	serious ^a	not serious	none	⊕⊕⊕⊖ MODERATE	300/892 (33.6%)	491/888 (55.3%)	RR 1.65 (1.37 to 1.99)	336 per 1,000	219 more per 1,000 (124 more to 333 more)			

	TNFi (MAB) compared to Placebo for Ulcerative colitis Bibliography: TNFi (MAB) compared to Placebo for Ulcerative colitis													
		Qu	ality assess		Sun	nmary of f	indings							
Maintenance of remission endpoint														
1070 (3 RCTs)	not serious	not serious	serious ^a	not serious	none	⊕⊕⊕⊖ MODERATE	65/537 (12.1%)	129/533 (24.2%)	RR 2.00 (1.52 to 2.62)	121 per 1,000	121 more per 1,000 (63 more to 196 more)			
Maintena	ince of	f response e	endpoint											
1070 (3 RCTs)	not serious	not serious	serious ^a	not serious	none	⊕⊕⊕⊖ MODERATE	118/537 (22.0%)	208/533 (39.0%)	RR 1.76 (1.46 to 2.14)	220 per 1,000	167 more per 1,000 (101 more to 251 more)			

Explanations

a. Indirect comparison to placebo

	Etanercept compared to Placebo for Crohn's Disease Bibliography: Etanercept compared to Placebo for Crohn's Disease												
Quality assessment								Summary of findings					
№ of participants	Risk of bias	k Inconsistency bias	Indirectness Imprecis	Imprecision	Publication bias	Overall quality of evidence	Study eve (%)	ent rates	Relative effect	Anticipated absolute effects			
(studies) Follow-up							With Placebo	With Etanercept	(95% CI)	Risk with Placebo	Risk difference with Etanercept		

		I	Etanercep Bibliogra	ot compa aphy: Etanei	ared to F	Placebo fo	or Croh	n's Dise n's Disease	ease				
		Qu	ality asses		Su	mmary of f	indings						
Clinical	linical Response at 4 weeks (Primary Study Endpoint)												
43 (1 RCT)	not serious	not serious	serious ^a	serious ^b	none		9/20 (45.0%)	9/23 (39.1%)	RR 0.87 (0.43 to 1.76)	450 per 1,000	59 fewer per 1,000 (257 fewer to 342 more)		
Clinical	Respo	nse at 2 w	eeks	-			•	-					
43 (1 RCT)	not serious	not serious	serious ^a	serious ^b	none		5/20 (25.0%)	11/23 (47.8%)	RR 1.91 (0.80 to 4.57)	250 per 1,000	227 more per 1,000 (50 fewer to 893 more)		
Clinical	Respo	nse at 8 w	eeks				1				•		
43 (1 RCT)	not serious	not serious	serious ^a	serious ^b	none		6/20 (30.0%)	7/23 (30.4%)	RR 1.01 (0.41 to 2.52)	300 per 1,000	3 more per 1,000 (177 fewer to 456 more)		

CI: Confidence interval; RR: Risk ratio

Explanations

a. Indirect comparison to placebo

b. C.I. crosses no effect line

References

- 1. Stidham RW, Lee TC, Higgins PD, Deshpande AR, Sussman DA, Singal AG, et al.. Systematic review with network meta-analysis: the efficacy of anti-TNF agents for the treatment of Crohn's disease. Aliment Pharmacol Ther. 2014 Jun;39(12):1349-62.
- Stidham RW, Lee TC, Higgins PD, Deshpande AR, Sussman DA, Singal AG, et al. Systematic review with network meta-analysis: the efficacy of anti-tumour necrosis factor-alpha agents for the treatment of ulcerative colitis. Aliment Pharmacol Ther. 2014 Apr;39(7):660-71.

3. Sandborn WJ, Hanauer SB, Katz S, Safdi M, Wolf DG, Baerg RD, et al. Etanercept for active Crohn's disease: a randomized, double-blind, placebo-controlled trial. Gastroenterology. 2001 Nov;121(5):1088-94.

PICO 59. In adult patients with active PsA and IBD despite treatment with an OSM, what are the benefits and harms of switching to TNFi (MABs) vs. switching to IL17i?

<u>Summary</u>: The literature searches did not identify any studies that addressed this PICO question. However, IL17i was not approved for IBD based on evidence from an RCT of harms in this patient population (the trial was discontinued early due to excess harms and lack of efficacy in the secukinumab arm).[1] This resulted in a warning on the IL17i package insert regarding harms in this population.

Quality of evidence across all critical outcomes: Moderate

Author,	Study	Duration	Population	Treatment given to	Results
year	type		Description	relevant population	
Hueber,	RCT	6 weeks,	59 patients with	Secukinumab (2X10	Primary endpoint was change in Crohn's Disease Activity Index
2012		with	moderate to	mg/kg intravenous, 39	(CDAI) at 6 weeks. The difference between groups was not
		follow-up	severe Crohn's	patients) vs. placebo	significant (change in CDAI score 33.9, 95% Bayesian
		of	disease	(20 patients)	credible interval -4.9 to 72.9), and area under the curve analysis
		secondary		-	at 4 to 10 weeks showed a significant difference (mean change in
		endpoints			CDAI=49; 95% CI (2 to 96), p=0.043) favoring placebo.
		to 10			
		weeks			Adverse events were more common in the secukinumab group
					(74% vs. 50%) and the infection rate was much higher in the
					secukinumab group (43% vs. 0%).

References

1. Hueber W, Sands BE, Lewitzky S, Vandemeulebroecke M, Reinisch W, Higgins PDR, et al. Secukinumab, a human anti-IL-17A monoclonal antibody, for moderate to severe Crohn's disease: unexpected results of a randomised, double-blind, placebo-controlled trial. Gut 2012;61:1693–1700.

PICO 60. In adult patients with active PsA and IBD despite treatment with an OSM, what are the benefits and harms of switching to IL12/23i vs. switching to IL17i?

<u>Summary</u>: The literature searches did not identify any studies that addressed this PICO question. However, IL12/23i is approved for IBD and IL17i was not approved for IBD based on evidence of harms in this patient population (see PICO 59). This resulted in a warning on the IL17i package insert regarding harms in this population.

Quality of evidence across all critical outcomes: Moderate

References

 Hueber W, Sands BE, Lewitzky S, Vandemeulebroecke M, Reinisch W, Higgins PDR, et al. Secukinumab, a human anti-IL-17A monoclonal antibody, for moderate to severe Crohn's disease: unexpected results of a randomised, double-blind, placebo-controlled trial. Gut 2012;61:1693–1700.

PICO 61. In adult patients with active PsA and IBD despite treatment with an OSM, what are the benefits and harms of switching to TNFi (MABs) vs. switching to IL12/23i?

Summary: The literature searches did not identify any studies that addressed this PICO question.

Quality of evidence across all critical outcomes: Very low

PICO 62. In adult patients with active PsA and IBD who are <u>both OSM and biologic treatment-naïve</u>, what are the benefits and harms of starting OSMs vs. starting TNFi (MABs)?

Summary. This PICO was addressed indirectly by one systematic review that included 3 RCTs. The studies looked at patients with acute ulcerative colitis naïve to OSM or biologics comparing treatment with infliximab vs. cyclosporine [1]. The only relevant outcome reported was serious adverse events, which showed no significant difference between treatment groups. However, the findings are inconclusive due to imprecision in the effect estimate.

<u>Quality of evidence across all critical outcomes</u>: Very low

Infliximab compared to Cyclosporine for PsA and IBD patients who are OSM and Biologic-naive Bibliography: PICO 62 - PsA and IBD patients who are OSM and Biologic naive: OSM vs. TNFi.

Quality assessment

Summary of findings

Infliximab compared to Cyclosporine for PsA and IBD patients who are OSM and Biologic-naive Bibliography: PICO 62 - PsA and IBD patients who are OSM and Biologic naive: OSM vs. TNFi.

Quality assessment								Summary of findings					
№ of participants (studies) Follow-up	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of	Study event rates (%)		Relative effect	Anticipated absolute effects			
						evidence	With CYC	With IFX	(95% CI)	Risk with CYC	Risk difference with IFX		

Serious Adverse Events

-			1					1			
415 (1 systematic review with 3 RCTs)	a a	not serious	very serious ^b	serious ^c	none	⊕⊖⊖⊖ VERY LOW	14/206 (6.8%)	20/209 (9.6%)	RR 1.41 (0.73 to 2.71)	68 per 1,000 (0.068)	28 more per 1,000 (0.028) (18 fewer to 116 more)

CI: Confidence interval; CYC: Cyclosporine; IFX: Infliximab; RR: Risk ratio

a. No mention in the review of randomization, allocation concealment, or blinding

b. Entire patient population are those with acute UC

c. Wide CI that crosses no effect line

Working Group quality of evidence categories:

High quality: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate quality: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low quality: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect Very low quality: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

References:

 Narula N, Marshall JK, Colombel JF, Leontiadis GI, Williams JG, Muqtadir Z, et al. Systematic Review and Meta-Analysis: Infliximab or Cyclosporine as Rescue Therapy in Patients With Severe Ulcerative Colitis Refractory to Steroids. Am J Gastroenterol. 2016;111(4):477-491.
PICO 63. In adult patients with active PsA and diabetes who are both OSM and biologic treatment-naïve, what are the benefits and harms of starting OSM vs. starting TNFi?

<u>Summary</u>: The literature searches did not identify any studies that directly addressed this PICO question. Two case series studies evaluated effects of diabetes on liver in patients with psoriasis on MTX treatment. One study [1] compared the impact of diabetes cumulatively with other risk factors such as obesity, alcohol consumption, chronic hepatitis B and C, while another study [2] studied impact of diabetes separately from other risk factors. Results show that diabetes has more impact on developing liver pathologies, such as fibrosis, in patients taking MTX at least 1000mg of cumulative dose.

Author,	Study	Duration	Population	Treatment given to	Results
year	type		Description	relevant population	
Rosenberg, 2007	Case series	Between 1975 and 2003	169 Patients with psoriasis and with/without diabetes mellitus type 2	Median cumulative dose MTX 1500-2100 mg	26 patients had one or more of the risk factors (diabetes mellitus type 2, overweight, alcohol over-consumption, and chronic hepatitis B or C) and 25 (96%) of these (median cumulative dose methotrexate 1500 mg) developed liver fibrosis. Of those without risk factor, 26 (58%) ($p = 0.012$) developed fibrosis (median cumulative dose methotrexate 2100 mg). Ten (38%) of the patients with risk factor(s) had severe fibrosis (stage 3–4) (mean cumulative dose methotrexate 1600 mg), while four (9%) ($p =$ 0.0012) of those without risk factors had severe fibrosis (median cumulative dose methotrexate 1900 mg). Seven (100%) of the patients with diabetes mellitus developed liver fibrosis compared to 37 (52%) of those without. Four (57%) of the seven patients with diabetes developed severe fibrosis
					compared to nine (14%) of those without $(p = 0.003)$
Malatjalian, 1996	Case series	Mean treatment duration 3.38 yrs	104 patients with psoriasis and diabetes	MTX cumulative dose 1000 to 1500 mg	Progression of liver pathology to higher grade in patients with diabetes Odds Ratio 2.07 (0.35-12.35) p=0.42. Progression of liver pathology to grades IIIB and IV Odds Ratio 5.68 (1.34-24.39), p= 0.02

Quality of evidence across all critical outcomes: Very low

- 1. Rosenberg P et al. Psoriasis patients with diabetes type 2 are at high risk of developing liver fibrosis during methotrexate treatment. Journal of Hepatology 46 (2007) 1111–1118
- 2. Malatjalian D et al. Methotrexate hepatotoxicity in psoriatics: Report of 104 patients from Nova Scotia, with analysis of risks from obesity, diabetes and alcohol consumption during long term follow-up. Can J Gastroenterol 1996;10(6):369-375.

PICO 64. In adult patients with active PsA and frequent serious infections who are both OSM and biologic treatment-naïve, what are the benefits and harms of starting OSMs vs. starting TNFi?

<u>Summary</u>: The literature searches did not identify any studies that addressed this PICO question. However, evidence concerning serious infections appears in the evidence base of earlier questions. Although the serious infection rate in most RCTs was so low that the findings were always imprecise, there is a black box warning against the use of TNFi in patients with recurrent serious infections.

Quality of evidence across all critical outcomes: Moderate

PICO 65. In adult patients with active PsA and frequent serious infections despite treatment with an OSM, what are the benefits and harms of switching to TNFi vs. switching to IL12/23i?

<u>Summary</u>: The literature searches did not identify any studies that directly addressed this PICO question. However, evidence concerning serious infections appears in the evidence base of earlier questions. The serious infection rate in most RCTs was so low that the findings were always imprecise. However, there is indirect evidence from a large retrospective cohort study (Yun et al. 2016) using Medicare data from RA patients.[1] The final cohort had 31,801 new courses of biologic therapy. The study compared hospitalized infection rates associated with the following biologic therapies: adalimumab, certolizumab, etanercept, golimumab, infliximab, rituximab, tocilizumab, and abatacept. Since abatacept had the lowest crude incidence rate of hospitalized infection (13.1/100 patients-years) it was used as the reference comparison. Biologic therapies with significantly higher adjusted hazard ratios (HR) for hospitalized infection compared to abatacept include etanercept (adjusted HR 1.24, 95% CI 1.07-1.45), infliximab (adjusted HR 1.39, 95% CI 1.21-1.60), and rituximab (adjusted HR 1.36, 95% CI 1.21-1.53).[1]

Quality of evidence across all critical outcomes: Very low

References:

1. Yun H et al. Comparative risk of hospitalized infection associated with biologic agents in rheumatoid arthritis patients enrolled in Medicare. Arth Rheumatol 2016; 68:56-66.

PICO 66. In adult patients with active PsA and frequent serious infections despite treatment with an OSM, what are the benefits and harms of switching to TNFi vs. switching to IL17i?

<u>Summary</u>: The literature searches did not identify any studies that addressed this PICO question. However, evidence concerning serious infections appears in the evidence base of earlier questions. However, the serious infection rate in most RCTs was so low that the findings were always imprecise.

Quality of evidence across all critical outcomes: Very low

Update: Additional PICO Questions

PICO 67. In adult patients with active PsA despite treatment with an OSM, what are the benefits and harms of switching to a TNFi compared to switching to abatacept?

<u>Summary</u>: Eleven placebo controlled RCTs (14 publications) indirectly addressed this PICO question. Nine studies (12 publications) compared TNFi versus placebo in PsA patients.[1-12] Two studies compared abatacept with placebo in PsA patients.[13,14] Statistically significant differences favoring TNFi over placebo were reported for all efficacy outcomes (ACR20, HAQ-DI, PASI-75, data not shown), while abatacept showed a significant benefit over placebo only for ACR20 (data not shown). No statistically significant differences with placebo occurred for any adverse events (see Table 2 for serious infections).

The adjusted indirect comparison method was used to calculate RRs for abatacept versus TNFi. The two TNFi studies (one using adalimumab, one using certolizumab) that reported infection rates had effect sizes in different directions, so these were separately compared to the abatacept trials. TNFi showed a significant benefit over abatacept for ACR20 and PASI-75, but not for HAQ-DI or rate of infection due to imprecision in effect estimates (Table 1).

	Table 1. TNFi compared to abatacept for patients with active PsA despite OSM Bibliography: PICO 67: TNFi compared to abatacept for patients with active PsA despite OSM											
	Quality assessment Summary of findings											
№ of participants	P of articipantsRisk ofInconsistencyIndirectnessImprecisionPublication biasOverall 											

Table 1. TNFi compared to abatacept for patients with active PsA despite OSM Bibliography: PICO 67: TNFi compared to abatacept for patients with active PsA despite OSM													
		Qua	ality assessi			Sur	nmary of	findings	5				
(studies) Follow-up	bias					evidence	With TNFi	With ABT	(95% CI)	Risk with TNFi	Risk difference with ABT		
ACR20, 1	2-24 v	veeks, Buch	er adjuste	d indirect	comparis	on							
2075 (11 RCTs)	Not serious	Not serious	Serious ^a	Not serious	None	⊕⊕⊕⊖ MODERATE	TNFi	ABT	RR 0.46 (0.30 to 0.69) Favors TNFi	583 per 1,000 (0.583)	315 fewer per 1,000 (0.315) (408 fewer to 181 fewer)		
HAQ-DI [°] ,	12-24	weeks, Bu	cher adjus [.]	ted indired	ct compar	ison	•						
1715 (9 RCTs)	Not serious	Not serious	Serious ^a	Serious ^b	None	⊕⊕⊖⊖ Low	TNFi	ABT	RR 0.85 (0.43 to 1.66)	547 per 1,000 (0.547)	82 fewer per 1,000 (0.082)(312 fewer to 361 more)		
PASI 75,	12-24	weeks, Bud	her adjust:	ed indirec	t compari	ison	<u>.</u>						
1342 (10 RCTs)	Not serious	Not serious	Seriousª	Not serious	None	⊕⊕⊕⊖ MODERATE	TNFi	ABT	RR 0.19 (0.08 to 0.43) Favors TNFi	494 per 1,000 (0.494)	400 fewer per 1,000 (0.4) (454 fewer to 282 fewer)		
Infection	, 12-2	4 weeks, Bu	icher adjus	sted indire	ct compa	rison							

Table 1. TNFi compared to abatacept for patients with active PsA despite OSM

Bibliography: PICO 67: TNFi compared to abatacept for patients with active PsA despite OSM

	_										
		Qua	ality assessr	nent				Sum	mary of	findings	
524 (2 RCTs)	Not serious	Not serious	Serious ^a	Serious ^b	None	⊕⊕⊖⊖ Low	ADA	ABT	RR 1.67 (0.75 to 3.69)	176 per 1,000 (0.176)	118 more per 1,000 (0.118) (44 fewer to 473 more)
698 (2 RCTs)	Not serious	Not serious	Serious ^a	Serious⁵	None	⊕⊕⊖⊖ Low	CZP	ABT	RR 0.79 (0.52 to 1.21)	435 per 1,000 (0.435)	91 fewer per 1,000 (0.091) (209 fewer to 91 more)

CI: Confidence interval; RR: Risk ratio

a. Indirect comparison, all studies compared drug to placebo, only 50-75% of patients had prior OSM exposure.

b. Wide 95% CI

c. HAQ-DI measured as proportion of patients who achieved minimum clinically important improvement (>0.3 or 0.35 in different studies).

Table 2. TNFi or abatacept compared to placebo for PICO 67: Adverse events Bibliography: PICO 67: TNFi versus abatacept for PsA patients who failed OSM.

		Qual	ity assess	ment				Summ	nary of f	inding	S
Nº of participants	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of	Study ev (%)	vent rates	Relative effect	Anticipat absolute	ted effects
Follow-up						evidence	With ABT or TNFi	With placebo	(43 % 61)	Risk with ABT or TNFi	Risk difference with placebo
Serious ir	nfectio	n – Abatace	pt vs. plac	ebo							

	Table 2. TNFi or abatacept compared to placebo for PICO 67: Adverse events Bibliography: PICO 67: TNFi versus abatacept for PsA patients who failed OSM.														
Quality assessment Summary of findings															
506 (2 RCTs)	not serious	not serious	serious ^a	serious ^b	none	⊕⊕⊖⊖ Low	2/253 (0.8%)	0/253 (0%)	RR 0.33 (0.03 to 3.13)	8 per 1,000 (0.008)	5 fewer per 1,000 (0.005) (0 fewer to 25 more)				
Serious ir	nfectio	n – TNFi vs.	placebo												
1151 (5 RCTs)	not serious	not serious	serious ^a	serious ^b	none	⊕⊕⊖⊖ Low	4/587 (0.7%)	8/564 (1.4%)	RR 1.85 (0.56 to 5.88)	7 per 1,000 (0.007)	6 more per 1,000 (0.006) (3 fewer to 34 more)				

a. Comparison to placebo

b. Wide 95% CI

Working Group quality of evidence categories:

High quality: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate quality: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low quality: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect Very low quality: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

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PICO 68. In adult patients with active PsA despite treatment with an OSM, what are the benefits and harms of switching to an IL12/23i compared to switching to abatacept?

<u>Summary</u>: Five placebo-controlled RCTs indirectly addressed this PICO question. Three studies compared IL 12/23 (ustekinumab) with placebo in PsA patients.[1-3] Two studies compared abatacept with placebo in PsA patients.[4,5] Statistically significant differences favoring ustekinumab over placebo were reported for all efficacy outcomes (ACR20, HAQ-DI, PASI-75, data not shown), while abatacept showed a significant benefit over placebo only for ACR20 (data not shown). No statistically significant differences with placebo occurred for any adverse events (see Table 2 for serious infections).

The adjusted indirect comparison method was used to calculate RRs for abatacept versus ustekinumab. Ustekinumab showed a significant benefit over abatacept for PASI-75, but not for ACR20, HAQ-DI and rate of infections (there was imprecision in the effect estimates for these outcomes, see Table 1)).

-	Fable	1. IL12/23 Bibliography:	Bi compare PICO 68: IL12	ed to abai	tacept fo d to abatacep	r patien ot for patien	ts with ts with acti	active ve PsA de	PsA de espite OSM	spite OS	SM
		Qua	ality assessr	nent				Sum	mary of 1	findings	
Nº of participants	Risk of	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of			Relative effect	Anticipate effects	d absolute
(studies) Follow-up	DIAS					evidence	With IL12/23i	With ABT	(95% CI)	Risk with IL12/23i	Risk difference with ABT
ACR20, 1	2-24 v	veeks, Buch	er adjuste	d indirect	compariso	on					
1579 (5 RCTs)	Not serious	Not serious	Serious ^a	Serious⁵	None	⊕⊕⊖⊖ Low	IL12/23i	ABT	RR 0.87 (0.61 to 1.24)	449 per 1,000 (0.449)	58 fewer per 1,000 (175 fewer to 108 more)
HAQ-DI [°] ,	12-24	weeks, Bud	cher adjust	ed indirec	t compari	ison	•		•	•	·

-	Table	1. IL12/23 Bibliography:	Bi compare PICO 68: IL12	ed to aba 2/23i compare	tacept fo	r patien ot for patien	ts with ts with act	active ive PsA de	PsA de espite OSM	spite O	SM
	Quality assessment Summary of findings										
1370 (5 RCTs)	Not serious	Not serious	Serious ^a	Serious ^b	None	⊕⊕⊖⊖ Low	IL12/23i	ABT	RR 0.84 (0.43 to 1.65)	441 per 1,000 (0.441)	71 fewer per 1,000 (251 fewer to 287 more)
PASI 75,	12-24	weeks, Bud	her adjust	ed indirec	t compari	son					
1135 (5 RCTs)	Not serious	Not serious	Serious ^a	Not serious	None	⊕⊕⊕⊖ MODERATE	IL12/23i	ABT	RR 0.26 (0.14 to 0.49) Favors IL12/23i	569 per 1,000 (0.569)	421 fewer per 1,000 (290 fewer to 489 fewer)
Infection	, 12-2	4 weeks, Bu	icher adjus	sted indire	ct compai	rison					
1349 (3 RCTs)	Not serious	Not serious	Serious ^a	Serious ^b	None	⊕⊕⊖⊖ Low	IL12/23i	ABT	RR 0.94 (0.63 to 1.41)	211 per 1,000 (0.211)	13 fewer per 1,000 (78 fewer to 87 more)

a. Indirect comparison, all studies compared drug to placebo, only 50-75% of patients had prior OSM exposure.

b. Wide 95% CI

c. HAQ-DI measured as proportion of patients who achieved minimum clinically important improvement (>0.3 or 0.35 in different studies).

Table 2. IL12/23i or abatacept compared to placebo for PICO 67: Adverse events Bibliography: PICO 68: IL 12/23i versus abatacept for PsA patients who failed OSM.

Quality assessment

Summary of findings

т	able	2. IL12/23 Bibliog	i or abata _{Iraphy:} PICO 6	cept com 8: IL 12/23i v	pared to ersus abatac	placebo ept for PsA	for PIC	CO 67: A	Adverse	events	
		Qual	ity assess	ment				Sumn	hary of f	indings	
Nº of participants	Risk of	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of	Study eve (%)	nt rates	Relative effect	Anticipate effects	ed absolute
Follow-up	DId3					evidence	With ABT or IL12/23i	With placebo	(7378 01)	Risk with ABT or IL12/23i	Risk difference with placebo
Serious ir	nfectio	on – Abatace	ept vs. plac	cebo							
506 (2 RCTs)	not serious	not serious	serious ^a	serious ^b	none	⊕⊕⊖⊖ Low	2/253 (0.8%)	0/253 (0%)	RR 0.33 (0.03 to 3.13)	8 per 1,000	5 fewer per 1,000 (0 fewer to 25 more)
Serious ir	nfectio	on – Ustekin	umab vs. p	olacebo	•	•	•		•		•
925 (2 RCTs)	not serious	serious	serious ^a	serious ^b	none	⊕⊕⊖⊖ Low	4/616 (0.6%)	2/309 (0.6%)	RR 1.19 (0.29 to 4.98)	6 per 1,000 (0.006)	1 more per 1,000 (4 fewer to 24 more)

a. Comparison to placebo

b. Wide 95% CI

Working Group quality of evidence categories:

High quality: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate quality: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low quality: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect Very low quality: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

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PICO 69. In adult patients with active PsA despite treatment with an OSM, what are the benefits and harms of switching to an IL17i compared to switching to abatacept?

<u>Summary</u>: Nine placebo-controlled RCTs indirectly addressed this PICO question. Seven studies compared IL17i to placebo: One study compared ixekizumab to placebo,[1] three studies compared secukinumab to placebo,[2-4] and three studies compared brodalumab to placebo[5-7] (note: brodalumab is currently FDA-approved for psoriasis but not PsA). Two studies compared abatacept with placebo in PsA patients.[8,9] Statistically significant differences favoring IL17i over placebo were reported for all efficacy outcomes (ACR20, HAQ-DI, PASI-75, data not shown), while abatacept showed a significant benefit over placebo only for ACR20 (data not shown). No statistically significant differences with placebo occurred for any adverse events (see Table 2 for serious infections).

The adjusted indirect comparison method was used to calculate RRs for abatacept versus IL17i. IL17i showed a significant benefit over abatacept for PASI-75, but not for ACR20, HAQ-DI and rate of infections (there was imprecision in the effect estimates for these outcomes).

	Tabl	le 1. IL17i Bibliograph	compared y: PICO 69: IL	to abata	cept for to abatacept	patients for patients	with activ	ctive P e PsA des	SA desp pite OSM	oite OSN	Λ
		Qua	ality assessr	nent				Sum	mary of t	findings	
Nº of participants	Risk of	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of			Relative effect	Anticipate effects	d absolute
(studies) Follow-up	DIAS					evidence	With IL17i	With ABT	(95% CI)	Risk with IL17i	Risk difference with ABT
ACR20, 1	2-24 v	veeks, Buch	er adjuste	d indirect	compariso	on		•			•
1463 (7 RCTs)	Not serious	Not serious	Serious ^a	Serious ^b	None	⊕⊕⊖⊖ Low	IL17i	ABT	RR 0.76 (0.53 to 1.09)	505 per 1,000 (0.505)	121 fewer per 1,000 (0.121)(237 fewer to 45 more)
HAQ-DI ^c ,	12-24	weeks, Bud	cher adjust	ted indired	t compari	ison	•	•	•	•	

	Tab	le 1. IL17i Bibliograph	compared y: PICO 69: IL	l to abata 17i compared	cept for to abatacept	Datients for patients	with active	ctive P PsA des	sA desp ^{pite OSM}	oite OSN	1
		Qua	ality assessr	nent				Sum	mary of f	findings	
1151 (6 RCTs)	Not serious	Not serious	Serious ^a	Serious ^b	None	⊕⊕⊖⊖ Low	IL17i	ABT	OR 0.98 (0.39 to 2.50)	556 per 1,000 (0.556)	5 fewer per 1,000 (0.005) (228 fewer to 202 more) ^d
PASI 75 ,	12-24	weeks, Bud	her adjust:	ed indirec	t compari	son					
1167 (9 RCTs)	Not serious	Not serious	Serious ^a	Not serious	None	⊕⊕⊕⊖ MODERATE	IL17i	ABT	RR 0.21 (0.10 to 0.42) Favors IL17i	643 per 1,000 (0.643)	508 fewer per 1,000 (0.508)(373 fewer to 579 fewer)
Infection	, 12-2	4 weeks, Bu	icher adjus	sted indire	ct compar	ison					
1026 (3 RCTs)	Not serious	Not serious	Serious ^a	Serious ^b	None	⊕⊕⊖⊖ Low	IL17i	ABT	RR 0.74 (0.46 to 1.20)	321 per 1,000 (0.321)	83 fewer per 1,000 (0.083)(173 fewer to 64 more)

a. Indirect comparison, all studies compared drug to placebo, only 50-75% of patients had prior OSM exposure.

b. Wide 95% CI

c. HAQ-DI measured as proportion of patients who achieved minimum clinically important improvement (>0.3 or 0.35 in different studies).

d. Absolute risk differences calculated from odds ratios obtained using the Bucher method.

	Table	e 2. IL17i o Biblio	r abatace graphy: PICO o	pt compai 69: IL 17i vers	red to pla	acebo fo for PsA pat	r PICO ients who	69: Ad	verse e\ ^{1.}	vents	
		Qual	ity assess	ment				Summ	hary of f	inding	5
Nº of participants	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of	Study ev (%)	ent rates	Relative effect	Anticipat absolute	ed effects
Follow-up						evidence	With ABT or IL17i	With placebo	(45 % 61)	Risk with ABT or IL17i	Risk difference with placebo
Serious ir	nfectio	n – Abatace	pt vs. plac	ebo							
506 (2 RCTs)	not serious	not serious	serious ^a	serious ^b	none	⊕⊕⊖⊖ Low	2/253 (0.8%)	0/253 (0%)	RR 0.33 (0.03 to 3.13)	8 per 1,000 (0.008)	5 fewer per 1,000 (0.005) (8 fewer to 25 more)
Serious ir	nfectio	n – IL17i vs	. placebo							<u> </u>	
1189 (3 RCTs)	not serious	not serious	serious ^a	serious ^b	none	⊕⊕℃ LOW	18/685 (2.6%)	5/406 (1.2%)	RR 0.56 (0.23 to 1.33)	26 per 1,000 (0.026)	11 fewer per 1,000 (0.011) (20 fewer to 9 more)

a. Comparison to placebo

b. Wide 95% CI

Working Group quality of evidence categories:

High quality: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate quality: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low quality: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect Very low quality: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

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PICO 70. In adult patients with active PsA despite treatment with a TNFi monotherapy, what are the benefits and harms of switching to a second TNFi compared to switching to abatacept?

<u>Summary</u>: This PICO question was addressed indirectly by three double-blind RCTs (4 publications). In addition to lack of direct drug comparisons, only 19% to 39% of patients in each study had prior TNFi exposure. One study (2 publications) involving PsA patients compared TNFi (CZP) to placebo.[1,2] Two studies compared abatacept with placebo in PsA patients.[3,4] CZP showed statistically significant benefit over placebo for all effectiveness outcomes (ACR20, HAQ-DI, and PASI-75, data not shown), while abatacept showed significant benefit over placebo only for ACR20 (data not shown). For serious infections there were no significant between-group differences, but with very high imprecision due to the low number of events (data not shown).

The adjusted indirect comparison method was used to calculate RRs for CZP vs. abatacept. For ACR20 we also performed this calculation using data only from patients with prior TNFi exposure (two studies reported separate data for these patients). CZP showed a significant benefit over abatacept for PASI75, but not for other outcomes (all were imprecise, see Table 1).

	Table 1. TNFi compared to abatacept for patients with active PsA despite TNFi Bibliography: PICO 70: TNFi compared to abatacept for patients with active PsA despite TNFi													
		Qua	ality assessr	nent				Sun	nmary of	findings				
Nº of participants	Risk of	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of			Relative effect	Anticipate effects	ed absolute			
(studies) Follow-up	DIAS					evidence	With TNFi	With ABT	(95% CI)	Risk with TNFi	Risk difference with ABT			
ACR20, 1	2-24 v	veeks, Buch	er adjuste	d indirect	compariso	on		·	·					
780 (3 RCTs)	Not serious	Not serious	Serious ^a	Serious ^b	None	⊕⊕⊖⊖ Low	CZP	ABT	RR 0.69 (0.45 to 1.06)	638 per 1,000 (0.638)	198 fewer per 1,000 (0.198) (351 fewer to 38 more)			
ACR20, 1	2-24 v	veeks, Buch	er adjuste	d indirect	compariso	on, TNFi-	expose	d	•					

	Table 1. TNFi compared to abatacept for patients with active PsA despite TNFi Bibliography: PICO 70: TNFi compared to abatacept for patients with active PsA despite TNFi													
		Qua	ality assessr	nent			Summary of findings							
339 (2 RCTs)	Not serious	Not serious	Serious ^a	Serious⁵	None	⊕⊕⊖⊖ Low	CZP	ABT	RR 0.32 (0.10 to 1.01)	593 per 1,000 (0.593)	403 fewer per 1,000 (0.403) (534 fewer to 6 more)			
HAQ-DI [°] ,	12-24	weeks, Bud	cher adjust	ed indirec	t compari	son								
578 (3 RCTs)	Not serious	Not serious	Serious ^a	Serious⁵	None	⊕⊕⊖⊖ Low	СZР	ABT	RR 1.14 (0.59 to 2.21)	580 per 1,000 (0.580)	16 more per 1,000 (0.016) (213 fewer to 210 more) ^d			
PASI 75,	12-24	weeks, Bud	her adjust	ed indirec	t compari	son								
512 (3 RCTs)	Not serious	Not serious	Serious ^a	Not serious	None	⊕⊕⊕⊖ MODERATE	CZP	ABT	RR 0.41 (0.19 to 0.90) Favors CZP	622 per 1,000 (0.622)	367 fewer per 1,000 (0.367)			
Infection	, 12-2	4 weeks, Bu	icher adjus	ted indire	ct compar	ison								
698 (2 RCTs)	Not serious	Not serious	Serious ^a	Serious ^b	None	⊕⊕⊖⊖ Low	CZP	ABT	RR 0.79 (0.52 to 1.21)	435 per 1,000 (0.435)	91 fewer per 1,000 (0.091)(209 fewer to 91 more)			

- a. Indirect comparison, all studies compared drug to placebo, only 50-75% of patients had prior OSM exposure.
- b. Wide 95% CI
- c. HAQ-DI measured as proportion of patients who achieved minimum clinically important improvement (>0.3 or 0.35 in different studies).
- d. Absolute risk differences calculated from odds ratios obtained using the Bucher method.

Working Group quality of evidence categories:

High quality: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate quality: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low quality: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect Very low quality: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

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- 4. Mease PJ, Gottlieb AB, van der Heijde D, FitzGerald O, Johnsen A, Nys M, et al. Efficacy and safety of abatacept, a T-cell modulator, in a randomised, double-blind, placebo-controlled, phase III study in psoriatic arthritis. Ann Rheum Dis. 2017.

PICO 71. In adult patients with active PsA despite treatment with a TNFi monotherapy, what are the benefits and harms of switching to an IL12/23i compared to switching to abatacept?

<u>Summary</u>: This PICO question was addressed indirectly by four double-blind RCTs. In addition to lack of direct drug comparisons, only 28% to 50% of patients in each study had prior TNFi exposure. Two studies involving PsA patients compared IL12/23i (ustekinumab) to placebo.[1,2] Two studies compared abatacept with placebo in PsA patients.[3,4] Ustekinumab showed statistically significant benefit over placebo for all effectiveness outcomes (ACR20, HAQ-DI, and PASI-75, data not shown), while abatacept showed significant benefit over placebo only for ACR20 (data not shown). For serious infections there were no significant between-group differences, but with very high imprecision due to the low number of events (data not shown).

The adjusted indirect comparison method was used to calculate RRs for IL12/23i vs. abatacept. For ACR20 we also performed this calculation using data only from patients with prior TNFi exposure (two studies reported separate data for these patients). Ustekinumab showed a significant benefit over abatacept for PASI75, but not for other outcomes (all were imprecise, see Table 1).

-	Table 1. IL12/23i compared to abatacept for patients with active PsA despite TNFi Bibliography: PICO 71: IL12/23i compared to abatacept for patients with active PsA despite TNFi													
		Qua	ality assessr	Summary of findings										
Nº of participants	Risk of	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of			Relative effect	Anticipate effects	d absolute			
(studies) Follow-up	bias					evidence	With IL12/23i	With ABT	(95% CI)	Risk with IL12/23i	Risk difference with ABT			
ACR20, 1	2-24 v	veeks, Buch	er adjuste	d indirect	comparis	on								
964 (4 RCTs)	Not serious	Not serious	Serious ^a	Serious ^b	None	⊕⊕⊖⊖ Low	IL12/23i	ABT	RR 0.79 (0.50 to 1.22)	433 per 1,000 (0.433)	91 fewer per 1,000 (0.091)(216 fewer to 95 more)			
ACR20, 1	2-24 v	veeks, Buch	er adjuste	d indirect	comparis	on, TNFi	-exposed	d patie	ents					

-	Table	1. IL12/2: Bibliography:	3i compar PICO 71: IL12	ed to aba 2/23i compare	tacept for ed to abatace	pr patien	ts with act	active	e PsA de espite TNF	espite T	NFi
		Qua	ality assessr	nent				Sum	mary of	findings	
394 (2 RCTs)	Not serious	Not serious	Serious ^a	Serious⁵	None	⊕⊕⊖⊖ Low	IL12/23i	ABT	RR 0.67 (0.31 to 1.42)	356 per 1,000 (0.356)	117 fewer per 1,000 (0.117) (246 fewer to 150 more)
HAQ-DI ^c ,	12-24	weeks, Bu	cher adjus	ted indired	ct compar	ison					
755 (4 RCTs)	Not serious	Not serious	Serious ^a	Serious ^b	None	⊕⊕⊖⊖ Low	IL12/23i	ABT	RR 0.72 (0.36 to 1.47)	389 per 1,000 (0.389)	109 fewer per 1,000 (0.109) (249 fewer to 183 more)
PASI 75,	12-24	weeks, Bud	cher adjust	ted indired	t compar	ison					
695 (4 RCTs)	Not serious	Not serious	Serious ^a	Not serious	None	⊕⊕⊕⊖ MODERATE	IL12/23i	ABT	RR 0.16 (0.06 to 0.42) Favors IL12/23i	531 per 1,000 (0.531)	446 fewer per 1,000 (0.446) (499 fewer to 308 fewer)
Infection	, 12-2	4 weeks, Bu	ucher adjus	sted indire	ect compa	rison					
735 (2 RCTs)	Not serious	Not serious	Serious ^a	Serious ^b	None	⊕⊕⊖⊖ Low	IL12/23i	ABT	RR 0.80 (0.48 to 1.33)	271 per 1,000 (0.271)	54 fewer per 1,000 (141 fewer to 89 more)

a. Indirect comparison, all studies compared drug to placebo, only 50-75% of patients had prior OSM exposure.

b. Wide 95% CI

c. HAQ-DI measured as proportion of patients who achieved minimum clinically important improvement (>0.3 or 0.35 in different studies).

Working Group quality of evidence categories:

High quality: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate quality: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low quality: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect Very low quality: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

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- 4. Mease PJ, Gottlieb AB, van der Heijde D, FitzGerald O, Johnsen A, Nys M, et al. Efficacy and safety of abatacept, a T-cell modulator, in a randomised, double-blind, placebo-controlled, phase III study in psoriatic arthritis. Ann Rheum Dis. 2017.

PICO 72. In adult patients with active PsA despite treatment with a TNFi monotherapy, what are the benefits and harms of switching to an IL17i compared to switching to abatacept?

<u>Summary</u>: This PICO was addressed indirectly by seven double-blind RCTs. In addition to lack of direct drug comparisons, only 29% to 50% of patients in each study had prior TNFi exposure. Two studies involving PsA patients compared Brodalumab to placebo.[1,2] Three studies compared Secukinumab to Placebo in patients with PsA.[3-5] One study compared Ixekizumab to placebo in patients with PsA.[6] Two studies compared abatacept with placebo in PsA patients.[7, 8] IL17i showed statistically significant benefit over placebo for all effectiveness outcomes (ACR20, HAQ-DI, and PASI-75, data not shown), while abatacept showed significant benefit over placebo only for ACR20 (data not shown). For serious infections there were no significant between-group differences, but with very high imprecision due to the low number of events (data not shown).

The adjusted indirect comparison method was used to calculate RRs for IL17i vs. abatacept. For ACR20 we also performed this calculation using data only from patients with prior TNFi exposure (two studies reported separate data for these patients). IL17i showed significant benefit over abatacept for PASI-75 and ACR20, but the additional analysis of ACR20 using only patients with prior TNFi exposure did not show a significant difference between drugs (there was imprecision due to a wide 95% CI). The findings for HAQ-DI and infections were inconclusive due to imprecision in effect estimates.

	Table 1. IL17i compared to abatacept for patients with active PsA despite TNFi Bibliography: PICO 72: IL17i compared to abatacept for patients with active PsA despite TNFi											
		Qua	ality assessr	ment				Sun	nmary of	findings		
№ of participants	Risk of	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of	Rela		Relative effect	Anticipate effects	d absolute	
follow-up	bias					evidence	With IL17i	With ABT	(95% CI)	Risk with IL17i	Risk difference with ABT	
ACR20, 1	2-24 v	veeks, Buch	er adjuste	d indirect	compariso	on						

	Table 1. IL17i compared to abatacept for patients with active PsA despite TNFi Bibliography: PICO 72: IL17i compared to abatacept for patients with active PsA despite TNFi													
		Qua	ality assessr	nent			Summary of findings							
1254 (6 RCTs)	Not serious	Not serious	Serious ^a	Not serious	None	⊕⊕⊕⊖ MODERATE	IL17i	ABT	RR 0.66 (0.46 to 0.97) Favors IL17i	479 per 1,000 (0.479)	163 fewer per 1,000 (0.163)(259 fewer to 15 fewer)			
ACR20, 1	2-24 v	veeks, Buch	er adjuste	d indirect	comparis	on, TNFi-	exposed	d						
293 (2 RCTs)	Not serious	Not serious	Seriousª	Serious ^b	None	⊕⊕⊖⊖ Low	IL17i	ABT	RR 0.73 (0.37 to 1.43)	354 per 1,000 (0.354)	96 fewer per 1,000 (0.096) (223 fewer to 152 more)			
HAQ-DI ^c ,	12-24	weeks, Buo	cher adjust	ted indired	t compari	ison								
942 (5 RCTs)	Not serious	Not serious	Serious ^a	Serious ^b	None	⊕⊕⊖⊖ Low	IL17i	ABT	RR 1.20 (0.64 to 2.25)	545 per 1,000 (0.545)	36 more per 1,000 (0.036) (175 fewer to 367 more) ^d			
PASI 75 ,	12-24	weeks, Bud	her adjust	ed indirec	t compari	son	•	•			•			
865 (6 RCTs)	Not serious	Not serious	Serious ^a	Not serious	None	⊕⊕⊕⊖ MODERATE	IL17i	ABT	RR 0.21 (0.07 to 0.60) Favors IL17i	594 per 1,000 (0.594)	469 fewer per 1,000 (0.469)(552 fewer to 238 fewer)			

Table 1. IL17i compared to abatacept for patients with active PsA despite TNFi

Bibliography: PICO 72: IL17i compared to abatacept for patients with active PsA despite TNFi

Quality assessment

Summary of findings

Infection, 12-24 weeks, Bucher adjusted indirect comparison

1026 (3 RCTs)	Not serious	Not serious	Serious ^a	Serious ^b	None	⊕⊕⊖⊖ Low	IL17i	ABT	RR 0.74 (0.46 to 1.20)	321 per 1,000 (0.321)	83 fewer per 1,000 (0.083)(173 fewer to 64 more)
											more)

CI: Confidence interval; RR: Risk ratio

a. Indirect comparison, all studies compared drug to placebo, only 50-75% of patients had prior OSM exposure.

- b. Wide 95% CI
- c. HAQ-DI measured as proportion of patients who achieved minimum clinically important improvement (>0.3 or 0.35 in different studies).
- d. Absolute risk differences calculated from odds ratios obtained using the Bucher method.

Working Group quality of evidence categories:

High quality: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate quality: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low quality: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect Very low quality: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

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- 8. Mease PJ, Gottlieb AB, van der Heijde D, FitzGerald O, Johnsen A, Nys M, et al. Efficacy and safety of abatacept, a T-cell modulator, in a randomised, double-blind, placebo-controlled, phase III study in psoriatic arthritis. Ann Rheum Dis. 2017.

PICO 73. In adult patients with active PsA despite treatment with an OSM, what are the benefits and harms of switching to a TNFi compared to switching to tofacitinib?

<u>Summary</u>: Ten placebo controlled RCTs (13 publications) indirectly addressed this PICO question. Nine studies (12 publications) compared TNFi versus placebo in PsA patients.[1-12] One study that pooled data from two psoriasis RCTs compared tofacitinib (5 mg twice daily) to placebo in a subgroup of patients with PsA.[13] The only relevant outcome presented in this study was PASI-75, so this is the only outcome that was compared between TNFi and tofacitinib. Statistically significant differences favored TNFi over placebo and tofacitinib over placebo for this outcome (data not shown). An adjusted indirect comparison found no significant difference in PASI-75 between TNFi and tofacitinib, but the finding was imprecise due to a wide CI that overlapped with the line of no effect (Table 1).

A search update in March 2018 identified two new RCTs comparing tofacitinib to placebo in patients with PsA.[14,15] The guideline panel had previously reviewed these studies in abstract form when formulating recommendations. Only one of the trials (Mease et al.[14]) enrolled a population of patients with prior OSM treatment but no prior TNFi exposure. This study also had a comparison arm of patients receiving adalimumab. At 3 months, tofacitinib showed significant benefit over placebo for ACR 20, HAQ-DI, and PASI 75, and no significant difference between tofacitinib and adalimumab for these outcomes. The tofacitinib versus placebo comparison is moderate quality due to indirectness, and the tofacitinib versus adalimumab comparison is moderate quality due to imprecision. These trials do not change the imprecision in the adjusted indirect comparisons and do not alter the overall quality of evidence.

Quality	/ of	evidence	across	all	critical	outcomes:	Low

	Table 1. TNFi compared to tofacitinib for patients with active PsA despite OSM Bibliography: PICO 73: TNFi compared to tofacitinib for patients with active PsA despite OSM												
		Qua	ality assessr	nent				Sun	nmary of	findings			
№ of participantsR of (studies)Follow-up	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence			Relative effect	Anticipated absolute effects			
	DIAS						With TNFi	With TOF	(95% CI)	Risk with TNFi	Risk difference with TOF		
PASI 75,	12-24	weeks, Buc	her adjust:	ed indirec	t compari	son	•	-					

Table 1. TNFi compared to tofacitinib for patients with active PsA despite OSM

Bibliography: PICO 73: TNFi compared to tofacitinib for patients with active PsA despite OSM

	Quality assessment								nmary of	findings	
10 RCTs	Not serious	Not serious	Serious ^a	Serious⁵	None	⊕⊕⊖⊖ Low	TNFi	TOF	RR 0.90 (0.28 to 2.92)	494 per 1,000 (0.494)	113 fewer per 1,000 (0.113) (352 fewer to 199 more) ^c

CI: Confidence interval; RR: Risk ratio

- a. Indirect comparison, all studies compared drug to placebo, only 50-75% of patients had prior OSM exposure.
- b. Wide 95% CI
- c. Absolute risk differences calculated from odds ratios obtained using the Bucher method.

Working Group quality of evidence categories:

High quality: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate quality: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low quality: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect Very low quality: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

- 1. Mease PJ, Fleischmann R, Deodhar AA, Wollenhaupt J, Khraishi M, Kielar D, et al. Effect of certolizumab pegol on signs and symptoms in patients with psoriatic arthritis: 24-week results of a Phase 3 double-blind randomised placebo-controlled study (RAPID-PsA). Ann Rheum Dis. 2014;73(1):48-55.
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PICO 74. In adult patients with active PsA despite treatment with an OSM, what are the benefits and harms of switching to an IL12/23i compared to switching to tofacitinib?

<u>Summary</u>: Four placebo-controlled RCTs indirectly addressed this PICO question. Three studies compared IL 12/23 (ustekinumab) with placebo in PsA patients.[1-3] One study that pooled data from two psoriasis RCTs compared tofacitinib (5 mg twice daily) to placebo in a subgroup of patients with PsA.[4] The only relevant outcome presented in this study was PASI-75, so this is the only outcome that was compared between ustekinumab and tofacitinib. Statistically significant differences favored ustekinumab over placebo and tofacitinib over placebo for this outcome (data not shown). An adjusted indirect comparison found no significant difference in PASI-75 between IL12/23i and tofacitinib, but the finding was imprecise due to a wide CI that overlapped with the line of no effect (Table 1).

A search update in March 2018 identified two new RCTs comparing tofacitinib to placebo in patients with PsA.[5,6] The guideline panel had previously reviewed these studies in abstract form when formulating recommendations. Only one of the trials (Mease et al.[5]) enrolled a population of patients with prior OSM treatment but no prior TNFi exposure. Tofacitinib showed significant benefit over placebo for ACR 20, HAQ-DI, and PASI 75. The tofacitinib versus placebo comparison is moderate quality due to indirectness. These trials do not change the imprecision in the adjusted indirect comparisons and do not alter the overall quality of evidence.

	Iable 1. IL12/231 compared to tofacitinib for patients with active PsA despite OSM Bibliography: PICO 74: IL12/23i compared to tofacitinib for patients with active PsA despite OSM												
		Qua	ality assessr	nent				Sum	nmary of	findings			
Nº of participants	Risk of	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of			Relative effect	Anticipate effects	d absolute		
Follow-up	DIAS					evidence	With IL12/23i	With TOF	(95% CI)	Risk with IL12/23i	Risk difference with TOF		
PASI 75,	12-24	weeks, Bud	her adjust	ed indirec	t compari	son				•			
4 RCTs	Not serious	Not serious	Serious ^a	Serious⁵	None	⊕⊕⊖⊖ Low	IL12/23i	TOF	OR 0.84 (0.27 to 2.65)	569 per 1,000	43 fewer per 1,000 (0.043) (306 fewer to 209 more) ^c		

CI: Confidence interval; OR: Odds ratio

- a. Indirect comparison, all studies compared drug to placebo, only 50-75% of patients had prior OSM exposure.
- b. Wide 95% CI
- c. Absolute risk differences calculated from odds ratios obtained using the Bucher method.

Working Group quality of evidence categories:

High quality: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate quality: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low quality: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect Very low quality: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

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- 6. Gladman D, Rigby W, Azevedo VF, Behrens F, Blanco R, Kaszuba A, et al. Tofacitinib for psoriatic arthritis in patients with an inadequate response to TNF inhibitors. NEJM. 2017;377(16):1525-1536.

PICO 75. In adult patients with active PsA despite treatment with an OSM, what are the benefits and harms of switching to an IL17i compared to switching to tofacitinib?

<u>Summary</u>: Nine placebo-controlled RCTs indirectly addressed this PICO question. Seven studies compared IL17i to placebo: One study compared ixekizumab to placebo,[1] three studies compared secukinumab to placebo,[2-4] and three studies compared brodalumab to placebo[5-7] (note: brodalumab is currently FDA-approved for psoriasis but not PsA). One study that pooled data from two psoriasis RCTs compared tofacitinib (5 mg twice daily) to placebo in a subgroup of patients with PsA.[8] The only relevant outcome presented in this study was PASI-75, so this is the only outcome that was compared between IL17i and tofacitinib. Statistically significant differences favored IL17i over placebo and tofacitinib over placebo for this outcome (data not shown). An adjusted indirect comparison found no significant difference in PASI-75 between IL17i and tofacitinib, but the finding was imprecise due to a wide CI that overlapped with the line of no effect (Table 1).

A search update in March 2018 identified two new RCTs comparing tofacitinib to placebo in patients with PsA.[9,10] The guideline panel had previously reviewed these studies in abstract form when formulating recommendations. Only one of the trials (Mease et al.[9]) enrolled a population of patients with prior OSM treatment but no prior TNFi exposure. Tofacitinib showed significant benefit over placebo for ACR 20, HAQ-DI, and PASI 75. The tofacitinib versus placebo comparison is moderate quality due to indirectness. These trials do not change the imprecision in the adjusted indirect comparisons and do not alter the overall quality of evidence.

Table 1. IL17i compared to tofacitinib for patients with active PsA despite OSM Bibliography: PICO 75: IL17i compared to tofacitinib for patients with active PsA despite OSM											
Quality assessment							Summary of findings				
№ of participants (studies) Follow-up	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence			Relative effect	Anticipated absolute effects	
							With IL17i	With TOF	(95% CI)	Risk with IL17i	Risk difference with TOF
PASI 75, 12-24 weeks, Bucher adjusted indirect comparison											

Table 1. IL17i compared to tofacitinib for patients with active PsA despite OSM

Bibliography: PICO 75: IL17i compared to tofacitinib for patients with active PsA despite OSM

Quality assessment							Summary of findings					
9 RCTs	Not serious	Not serious	Serious ^a	Serious ^b	None	⊕⊕⊖⊖ Low	IL17i	TOF	OR 0.52 (0.15 to 1.85)	643 per 1,000 (0.643)	159 fewer per 1,000 (0.159) (430 fewer to 126 more) ^c	

CI: Confidence interval; OR: Odds ratio

- a. Indirect comparison, all studies compared drug to placebo, only 50-75% of patients had prior OSM exposure.
- b. Wide 95% CI
- c. Absolute risk differences calculated from odds ratios obtained using the Bucher method.

Working Group quality of evidence categories:

High quality: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate quality: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low quality: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect Very low quality: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

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- 2. McInnes IB, Mease PJ, Kirkham B, Kavanaugh A, Ritchlin CT, Rahman P, et al. Secukinumab, a human anti-interleukin-17A monoclonal antibody, in patients with psoriatic arthritis (FUTURE 2): a randomised, double-blind, placebo-controlled, phase 3 trial. Lancet. 2015;386(9999):1137-1146.
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- 9. Mease P, Hall S, Fitzgerald O, van der Heijde D, Merola JF, Avila-Zapata F, et al. Tofacitinib or adalimumab versus placebo for psoriatic arthritis. NEJM. 2017;377(16):1537-1550.
- 10. Gladman D, Rigby W, Azevedo VF, Behrens F, Blanco R, Kaszuba A, et al. Tofacitinib for psoriatic arthritis in patients with an inadequate response to TNF inhibitors. NEJM. 2017;377(16):1525-1536.

PICO 76. In adult patients with active PsA despite treatment with a TNFi monotherapy, what are the benefits and harms of switching to a second TNFi compared to switching to tofacitinib?

<u>Summary</u>: See summary for PICO 73. The available tofacitinib study did not separately report data for patients with prior TNFi exposure, and did not report the percentage of patients with PsA who had prior TNFi exposure (the percentage with prior biologic exposure was 26% for the overall population of patients with psoriasis), so the comparison in PICO 73 can be used as indirect evidence. In the main population of patients with psoriasis the findings did not differ for biologic-naïve and biologic-exposed patients.

A search update in March 2018 identified two new RCTs comparing tofacitinib to placebo in patients with PsA.[1,2] The guideline panel had previously reviewed these studies in abstract form when formulating recommendations. Only one of the trials (Gladman et al.[2]) enrolled a population of patients with prior TNFi exposure. At 3 months, tofacitinib showed significant benefit over placebo for ACR 20 and HAQ-DI; only the 10 mg dose showed significant benefit over placebo for PASI 75. This new evidence is moderate quality due to indirectness, does not change the imprecision in the adjusted indirect comparisons and therefore does not alter the overall quality of evidence.

Quality of evidence across all critical outcomes: Low

- 1. Mease P, Hall S, Fitzgerald O, van der Heijde D, Merola JF, Avila-Zapata F, et al. Tofacitinib or adalimumab versus placebo for psoriatic arthritis. NEJM. 2017;377(16):1537-1550.
- 2. Gladman D, Rigby W, Azevedo VF, Behrens F, Blanco R, Kaszuba A, et al. Tofacitinib for psoriatic arthritis in patients with an inadequate response to TNF inhibitors. NEJM. 2017;377(16):1525-1536.

PICO 77. In adult patients with active PsA despite treatment with a TNFi monotherapy, what are the benefits and harms of switching to an IL12/23i compared to switching to tofacitinib?

<u>Summary</u>: See summary for PICO 74, with explanation in PICO 76.

Quality of evidence across all critical outcomes: Low

PICO 78. In adult patients with active PsA despite treatment with a TNFi monotherapy, what are the benefits and harms of switching to an IL17i compared to switching to tofacitinib?

<u>Summary</u>: : See summary for PICO 75, with explanation in PICO 76.

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