Can Patients With Controlled Rheumatoid Arthritis Taper Methotrexate From Targeted Therapy and Sustain Remission? A Systematic Review and Metaanalysis

Charis F. Meng¹^(D), Diviya A. Rajesh², Deanna P. Jannat-Khah³^(D), Bridget Jivanelli⁴^(D), and Vivian P. Bykerk¹^(D)

ABSTRACT. Objective. To determine the risk of not being able to sustain remission after tapering methotrexate (MTX) from targeted therapy in patients with controlled rheumatoid arthritis (RA).

Methods. A systematic literature search was conducted in MEDLINE, Embase, and the Cochrane Library for studies reporting remission outcomes after tapering MTX from targeted therapies in RA. Full-text articles and abstracts reported in English were included. Metaanalyses were conducted using random-effects models. Forest and funnel plots were created.

Results. A total of 10 articles were included. Studies evaluated MTX being tapered from combination treatment with tumor necrosis factor inhibitors, tocilizumab, abatacept, and tofacitinib. A total of 9 studies used a randomized design and 1 was observational. Out of 10 studies, 3 focused on early RA (ie, < 1 yr). The MTX-tapering strategy was gradual in 2 studies and rapid in 8 studies. Follow-up ranged from 3 to 18 months in randomized trials and up to 3 years in the observational study. Our metaanalysis, which included 2000 participants with RA from 10 studies, showed that patients who tapered MTX from targeted therapy had a 10% reduction in the ability to sustain remission and an overall pooled risk ratio of 0.90 (95% CI 0.84-0.97). There was no heterogeneity (P = 0%, P = 0.94). Our funnel plot indicated minimal publication bias. **Conclusion.** Patients with controlled RA may taper MTX from targeted therapy with a 10% reduction in the ability to sustain remission for up to 18 months. Longer follow-up studies with attention to radiographic, functional, and patient-reported outcomes are needed. The risk of disease worsening should be discussed with the patient with careful follow-up and prompt retreatment of disease worsening.

Key Indexing Terms: methotrexate, remission, rheumatoid arthritis

Methotrexate (MTX) is recommended to be used in combination with biologic disease-modifying antirheumatic drugs

This research was supported by the Inflammatory Arthritis Center, Hospital for Special Surgery, New York, New York, USA.

¹C.F. Meng MD, V.P. Bykerk, MD, Division of Rheumatology, Hospital for Special Surgery, and Department of Medicine, Weill Cornell Medical College. ²D.A. Rajesh, BA, Division of Rheumatology, Hospital for Special Surgery. ³D.P. Jannat-Khah, DRPH, MSPH, Division of Rheumatology, Epidemiology and Biostatistics CORE, Hospital for Special Surgery, and Department of Medicine, Weill Cornell Medical College. ⁴B. Jivanelli, MLIS, Kim Barrett Memorial Library, HSS Education Institute, Hospital for Special Surgery, New York, NY, USA.

DPJK declares stock ownership in AstraZeneca, Cytodyn, and Walgreens. VPB declares institutional grants from BMS and Amgen; honoraria or consulting fees from Amgen, BMS, Genzyme, Gilead, Janssen, Pfizer, Sanofi-Aventis, and UCB; support by the National Institutes of Health (NIH; National Institute of Allergy and Infectious Diseases/National Institute of Arthritis and Musculoskeletal and Skin; grants 1UH2AR067691-01 and 11652401) and The Cedar Hill Foundation; and participation on the Data Safety Monitoring Board of NIH Cell Therapies. The remaining authors declare no conflicts of interest relevant to this article.

Address correspondence to Dr. C.F. Meng, Hospital for Special Surgery, 535 E 70th Street, New York, NY 10021, USA. Email: mengc@hss.edu. Accepted for publication July 27, 2022. (bDMARDs) in the treatment of rheumatoid arthritis (RA) because of its additive therapeutic benefits and its mitigation of immunogenicity.¹ In clinical practice, however, up to 30% of patients are on bDMARD monotherapy,²⁻⁴ in part because of their intolerance to MTX and other conventional synthetic disease-modifying antirheumatic drugs (csDMARDs). Adverse effects from MTX have been cited to be the most common reason for its discontinuation, particularly from gastrointestinal intolerance, cytopenia, and abnormal liver function tests.⁵ MTX adherence has been observed to be highly variable⁶ and inferior to that of bDMARDs.⁷ In addition, several RA studies have shown the effectiveness of monotherapy with interleukin 6 inhibitors (IL-6i) and Janus kinase inhibitors (JAKi).^{4,8-10} Going forward, we refer to both bDMARDs and JAKi as targeted therapies.

Tapering disease-modifying antirheumatic drugs (DMARD) therapy is a desirable goal for many patients with chronic diseases such as RA. Patients wish to reduce adverse effects, reduce risk of future adverse effects, and maintain control over their own health.¹¹ However, the clinical benefits of tapering treatment in RA are less clear, and many studies have shown a high risk of disease worsening when stopping DMARDs.^{12,13} What is not known is how feasible it is for patients who are taking a combination of targeted therapy with MTX to taper their MTX and continue to be controlled. Observational studies have reported

© 2023 The Journal of Rheumatology

that 34% to 62% of patients with RA using tumor necrosis factor inhibitors (TNFi) later tapered their MTX.^{14,15} The 2021 American College of Rheumatology (ACR) guidelines conditionally recommend (1) continuation of all DMARDs at their current dose over a dose reduction because of a risk of flare, and (2) in patients who are taking both MTX and a targeted therapy, the tapering of MTX before tapering the targeted therapy. However, the ACR acknowledge there is an absence of direct evidence.¹⁶

Prior reviews have focused on tapering of MTX from combination treatment with either csDMARDs or TNFi.¹⁷ A 2015 systematic review of tapering of csDMARDs or bDMARDs reported a flare rate after tapering MTX ranging from 8% at 24 weeks (patients remained on hydroxychloroquine and corticosteroids) to 42% at 32 weeks (patients on infliximab).¹² According to our literature search, there have been no updated reviews addressing MTX tapering from other targeted therapies, such as IL-6i or JAKi, nor has there been a systematic review with a metaanalysis addressing this question. Factors associated with successful tapering, such as disease duration (ie, early vs established RA) or the tapering scheme itself (ie, gradual vs brisk), remain unknown.¹⁸

Therefore, we conducted a systematic literature review to evaluate whether remission can be sustained after the tapering (ie, dose reduction, gradual dose reduction before stopping, or withdrawal) of MTX in patients with RA who are taking MTX in combination with targeted therapy. We also aimed to evaluate the factors associated with successful tapering, such as disease duration and tapering schemes. Our hypothesis was that patients with controlled RA may taper MTX from targeted therapy with low risk of not being able to sustain remission.

METHODS

We searched for tapering studies in which patients received any targeted therapy, including all classes of bDMARDs (ie, abatacept [ABA], certolizumab pegol [CZP], etanercept [ETN], golimumab, infliximab [IFX], rituximab [RTX], tocilizumab [TCZ], and sarilumab) or JAKi (ie, tofacitinib [TOF], baricitinib, and upadacitinib) in combination with MTX, in which the study evaluated the proportion of patients in remission after the dosage of MTX was tapered.

Search strategy. The search strategy was initially developed in MEDLINE (ie, PubMed) by a medical librarian. It was then adapted for other databases that were searched: Embase; the Cochrane Library, including the Cochrane Central Register of Controlled Trials; the Health Technology Assessment database; and the NHS (National Health Service) Economic Evaluation Database (for full search strategy and search terms, see Supplementary Data S1, available with the online version of this article). We searched for articles published between January 1, 2014, and August 30, 2021, and ran the last updated search in all databases on August 30, 2021. Additional studies were identified through manually searching reference lists and gray literature references. Studies were excluded if they were not published in English.

Studies were imported into the Covidence platform (Veritas Health Innovation Ltd), allowing duplicates to be removed. The screening process was completed by 2 authors (CFM and DAR). Title and abstract screening were conducted first, followed by full-text screening. Any issues were resolved through consensus with VPB. This review was conducted and reported according to the procedures outlined in the PRISMA (Preferred Reporting Items for Systematic reviews and Meta-Analyses) statement.¹⁹

Study selection. Inclusion criteria for articles comprised the following:

(1) prospective comparative studies, including randomized controlled trials (RCTs), pragmatic trials, and observational studies of patients with RA; (2) subjects were taking MTX and targeted therapy (ie, TNFi, IL-6i, ABA, RTX, or JAKi); (3) the study design included an intervention group who underwent tapering of MTX from a combination with targeted therapy and a comparator group who continued combination therapy; and (4) reporting of subjects who remained in or achieved remission as measured by a composite score. Exclusion criteria comprised the following: (1) retrospective studies and (2) no reporting of the proportion of remission outcomes after tapering treatment.

Data extraction. CFM and DAR selected potential manuscripts for retrieval and, upon retrieval, established study eligibility by applying the selection criteria. Studies in doubt were discussed with VPB until consensus was reached. If trial data relevant to the review were found in a secondary publication or abstract, they were included and noted in the tables. The original publication of the Combination of Methotrexate and Etanercept in Active Early Rheumatoid Arthritis (COMET) trial²⁰ was used to extract study information and baseline data, but remission data were obtained from an updated publication²¹ that was analyzed according to low disease activity (LDA)/remission before tapering, consistent with our inclusion criteria. A standardized data collection form was used to extract the following: study design, patient inclusion/exclusion criteria, prior and baseline treatment, whether patients were MTX-naïve or inadequate responders, and RA duration dichotomized as either early (diagnosis < 1 yr) or established. Included was the implementation information for tapering, including criteria for tapering of therapy, tapering strategy, frequency of assessment, follow-up interval after tapering, as well as the reported outcome measures, including that of remission, disease worsening, duration of remission, retreatment outcomes, radiographic outcomes, patient-reported outcomes, and predictors of either remaining in disease control or losing disease control.

Quality assessment. The methodological quality of each randomized study was assessed using the revised Cochrane risk-of-bias tool for randomized trials²² by CFM and DAR, discussed with DPJK and, where clarification was needed, with VPB. The criteria for evaluation included randomization, deviations from intended interventions, missing outcome data, measurement of outcomes, and selection of reported results. Studies were judged to have an overall low risk of bias if they were found to have a low risk of bias for all domains. Studies were judged to have some concerns overall if they were found to have some concerns in at least 1 domain. Studies were judged to have an overall high risk of bias if they were found to have a high risk of bias in at least 1 domain or some concerns for multiple domains that substantially lowered the confidence in the results.²² Nonrandomized studies were assessed using the Risk Of Bias In Non-randomized Studies - of Interventions (ROBINS-I) tool,²³ which used similar criteria to judge overall risk of bias (Supplementary Table S1, available with the online version of this article).

Statistical analysis. Random-effects models were used to calculate pooled risk ratios (RRs).^{24,25} Heterogeneity was assessed by calculating the I^2 index using the Cochran-Mantel-Haenszel technique.²⁶ Additionally, forest plots were generated for each analysis. A funnel plot was created, and the Egger and Harbord tests were calculated^{27,28} to aid in the assessment of bias. All analyses were performed in Stata (version 14.2; StataCorp).

RESULTS

Literature search. Our search identified 5763 citations using the prespecified search terms. After removal of duplicates and articles not pertaining to the study question using the Covidence platform, 504 full-text articles were reviewed (Figure 1 and Supplementary Data S1, available with the online version of this article). Of these, 10 articles addressed our research question and met our inclusion criteria.

Characteristics of included studies. A total of 10 studies examining

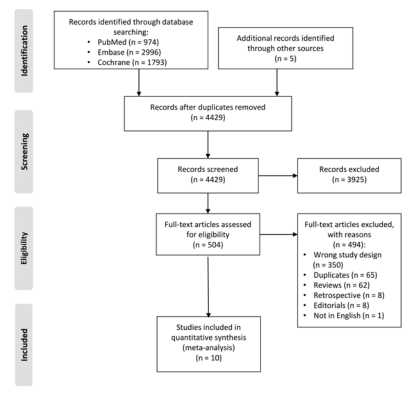


Figure 1. PRISMA flowchart for identification of studies. PRISMA: Preferred Reporting Items for Systematic reviews and Meta-Analyses.

the tapering of MTX from combined treatment with targeted therapy were reviewed. In total, 3 studies tapered MTX from combined treatment with ETN,^{20,21,29-31} 3 studies tapered MTX from TCZ,³²⁻³⁴ and 1 trial each tapered MTX from TOF,^{35,36} CZP,³⁷ adalimumab (ADA),³⁸ and ABA^{39,40} (Table 1). No studies tapering MTX from RTX met our inclusion criteria. In total, 7 articles studied established RA^{29,30,32-35,37} (ie, 6-11 yrs) and 3 studied early RA^{21,38,39} (ie, 1-9 months). Use of prior DMARDs ranged from 11% to 32%^{31,35,37,38} but was not specified in the remaining studies.^{20,29,32,34} Patients who were MTX-naïve were evaluated in the early RA trials, 20,38,39 and the remaining trials^{29,30,32-35,37} studied patients who were MTX-inadequate responders. Seropositivity ranged from 58% to 88% in 7 studies.^{20,29,30,33,35,37,38} There were 9 RCTs,^{20,29,30,32-35,37,39} 7 of which studied withdrawal as the second phase of their study, and 1 was a long-term extension (LTE) study.³⁸ In total, 2 RCTs used a run-in period^{29,35} (Table 1 and Supplementary Table S2, available with the online version of this article). In total, 7 RCTs^{20,29,32-35,39} were placebo-controlled during tapering (Supplementary Table S2).

Criteria for tapering was LDA based on Disease Activity Score in 28 joints (DAS28) in 4 studies,^{31,32,33,38} Clinical Disease Activity Index LDA in 1 study,³⁵ change in DAS28 in 1 study,³⁷ Simplified Disease Activity Index (SDAI) remission in 2 studies,^{29,39} both DAS28 remission and LDA in 1 study,²¹ and based on the European Alliance of Associations for Rheumatology (EULAR) response in 1 study³⁴ (Table 1 and Supplementary Table S2, available with the online version of this article). As their outcome measure, a total of 8 studies^{21,31,32-} ^{35,37,38} used the proportion of patients with DAS28-based remission, with 2 studies^{29,39} using SDAI remission (Table 1 and Supplementary Table S2).}

Follow-up ranged from 28 weeks to 18 months in 9 $RCTs^{20,21,29,30,32\cdot35,37,39}$ but was up to 3 years in the LTE study³⁸ (Table 1). The 3-year LTE study³⁸ did not specify time of with-drawal, so duration of remission after taper was not explicitly reported.

Quality assessment. In the RCTs, the overall risk of bias was judged to be low in 1 study,²⁹ have some concerns in 6 studies,^{21,32-35,39} and be high in 2 open-label studies^{30,37} (Figure 2). The LTE study³⁸ was judged to have a serious risk of bias (Supplementary Table S1, available with the online version of this article).

Tapering scheme. Eight studies^{29,30,32,33,35,37-39} stopped MTX in their tapering strategy and 2 studies^{20,21,34} gradually reduced the dose of MTX. The COMET trial tapered MTX from ETN+MTX over 4 weeks and was among the studies reporting a higher remission rate of 70%,²¹ compared to a remission rate ranging from 16% to 76% in the studies that stopped MTX abruptly.^{29,30,32,33,35,37-39} However, the study by Edwards et al (ACT-TAPER) tapered the dose of MTX more slowly over 24 weeks from TCZ+MTX and reported a lower remission rate of 50%.³⁴

Duration of remission and follow-up. Remission outcomes after MTX withdrawal were obtained at varying timepoints, ranging

Study, Author, Year	:	Established RA (duration)	yrs	Number of Naïve Prior DMARDs or IR	Naïve or IR	%	Ireatment	Strategy	REM or LDA Prior to Taper	Outcome Measure	Assessment		Design; WD 2nd Phase or LTE
SEAM ²⁹ , Curtis, 2021	253	estRA (10-11 yrs)	55-56	NR	IR	58-69	ETN+MTX	Stop MTX	SDAI ≤ 3.3 REM; 24-wk run-in	SDAI ≤ 3.3	Every 12 wks	48 wks	RCT; no
AVERT-2 ³⁹ , Emery, 2019 (ABS)	147	eRA (1-1.5 mos)	46-48	DMARD naïve	Naïve	NR	ABA+MTX	Stop MTX	SDAI < 3.3 REM: NR	SDAI ≤ 3.3	Wk 24 (WD), 40,48	48 wks	RCT; 2nd phase
ORAL shift ³⁵ , Cohen, 2019	533	estRA (9 yrs)	56	csDMARD: excluded MTX: 26% Prior TNFi: 30%	IR	62-68	TOF+MTX	Stop MTX	CDAI < 10 LDA; 24-wk run-in	DAS28-CRP < 2.6	Wk 12, 24 (WD), 36, 48	48 wks	RCT; 2nd phase
COMET, Emery, 2010 ²⁰ / 2019 (ABS) ²¹	411	eRA (9 mos)	52	NR	Naïve	68	ETN+MTX	Taper MTX over 4 wks	DAS28 < 2.6 REM or < 3.2 LDA; NR	DAS28 < 2.6	Wk 52 (WD), 104	52 wks	RCT; 2nd phase
IUST-ACT ³² , Pablos, 2019	165	estRA (6 yrs)	50-51	NR	IR	NR	TCZ+MTX	Stop MTX	DAS28 ≤ 3.2; NR	DAS28 < 2.6	Baseline, wk 16, wk 24 (WD), 28	28 wks	RCT; 2nd phase
Pope ³⁷ , 2020	88	estRA (8-10 yrs)	54-58	Prior targeted therapy: 11-14%	IR	60-62	CZP+DMARD (64% on MTX)	Stop DMARDs	Change in DAS28 > 1.2 after adding CZP; NR	DAS28 < 2.6	Baseline, 18 mos (WD time varied)	18 mos	RCT; no
COMP-ACT ³³ , Kremer, 2018	⁵ , 296	cstRA (7 yrs)	54-56	No. of prior csDMARD: 1.2 No. of prior TNFi: 0.2	IR	70-74	TCZ+MTX	Stop MTX	DAS28 ≤ 3.2; NR	DAS28 < 2.6	Wk 24 (WD), 40, 52	52 wks	RCT; 2nd phase
ACT-TAPER ³⁴ , Edwards, 2018	⁴ , 272	estRA (7 yrs)	54-56	NR	IR	NR	TCZ+MTX	Taper MTX over 24 wks	Good/moderate EULAR response ^b ; NR	DAS28 < 2.6	Wk 24 (WD), every 4 wks to wk 72	48 wks	RCT; 2nd phase
CAMEO, Keystone, 2016 ³⁰ / Pope 2014 ³¹	205 º/	estRA (9 yrs)	54	No. prior DMARD (median): 1	IR	65-67	ETN+MTX	Stop MTX I	Subgroup analysis: DAS28-ESR < 3.2 LDA; NR	DAS28 < 2.6	Mo 6 (WD), 12, 18, 24	18 mos	RCT; 2nd phase
PREMIER OLE, 140 Keystone ³⁸ , 2018	E, 140	cRA (0.7-0.8 yrs)	50-51	Prior DMARD: 27-32%	Naïve	84-88	ADA+MTX	Stop MTX	DAS28-CRP < 3.2 LDA; NR	DAS28-CRP < 2.6	Baseline WD2018 (varied), every 12 wks in OLE 1, every 16 wks during OLE yrs 2-3	3 yrs	Pooled post hoc analysis of OLE; LTE

Table 1. Summary characteristics of included studies tapering MTX from targeted therapy.

in 28 joints; DMARD: disease-modifying anticheumatic drug; eRA: early rheumatoid arthritis (≤ 1 yr); ESR: erythrocyte sedimentation rate; estRA: established rheumatoid arthritis; ETN: etanetcept; EULAR: European Alliance of Associations for Rheumatology; IR: inadequate responder; LDA: low disease activity; LTE: long-term extension; MTX: methotrexate; NR: not reported; OLE: open-label extension; RA: rheumatoid arthritis; RCT: randomized controlled trial; REM: remission; SDAI: simplified disease activity index; SEAM: Study of Exanercept and Methotrexate in Combination or as Monotherapy; TCZ: tocilizumab; TNFi: TNF inhibitor; TOF: tofacitinib; WD: withdrawal.

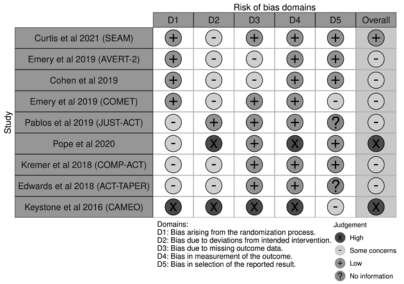


Figure 2. Risk-of-bias assessment of randomized trials using the revised Cochrane risk of bias assessment tool for randomized trials. AVERT-2: Assessing Very Early Rheumatoid Arthritis Treatment-2; COMET: Combination of Methotrexate and Etanercept in Active Early Rheumatoid Arthritis.

from 12 weeks to 18 months in randomized studies^{21,29,30,32-35,37,39} (Table 2). Studies that reported outcomes up to 1 year^{21,29,32-35,37,39} after tapering had remission rates ranging from 48% to 76%, but this dropped to 40% in 1 study that reported 18-month remission outcomes.³⁰ When persistent remission—defined as consistent remission at weeks 12, 24, 36, and 48—after tapering MTX to TOF monotherapy was used, remission rates dropped to 4%.³⁵

Mean disease activity scores after tapering. In total, 8 studies^{29,30,32-} ^{35,37,39} reported on changes in mean disease activity scores after tapering MTX (Table 2). Curtis et al²⁹ found that disease worsening defined as SDAI > 11 was similarly high in those who stopped MTX (75%) compared to those who continued ETN+MTX (78%). In total, 2 studies^{32,33}—COMP-ACT and JUST-ACT-demonstrated noninferiority of change in DAS28 scores in withdrawing MTX from TCZ compared to combination therapy. Pope et al³⁷ did not demonstrate noninferiority of maintaining change in DAS28 scores in the group withdrawing MTX from CZP compared to continuing therapy (Table 2). The Assessing Very Early Rheumatoid Arthritis Treatment-2 (AVERT-2), and Canadian Methotrexate and Etanercept Outcome (CAMEO) trials and study by Edwards et al (ACT-TAPER) did not find a significant difference in mean scores between groups.

Functional outcomes. In total, 7 studies^{20,30,32,35,37-39} reported on functional or other patient-reported outcomes (Table 2). The AVERT-2 trial found an adjusted mean change in the Health Assessment Questionnaire–Disability Index of 0.16 in those who stopped MTX vs –0.04 in those who continued ABA+MTX.⁴⁰ The physical functioning scale scores from the 36-item Short Form Health Survey were also worse in the group that stopped MTX (–1.45 vs 1.68 in the combination group). Pope et al³⁷ found significantly longer morning stiffness in the CZP monotherapy group (39.9 min) compared to the CZP+MTX group

(21.7 min; P = 0.026). Patient global pain, fatigue, work loss, and tender joint count scores trended worse with CZP monotherapy but did not reach significance (Table 2).

Radiographic outcomes. In total, 2 randomized trials^{30,39} and 1 observational study³⁸ assessed radiographic outcomes after tapering MTX. No significant differences in radiographic progression after tapering MTX to targeted therapy alone was observed.

Predictors of maintaining disease control. In total, 2 RCTs^{29,35,36} and 1 LTE study³⁸ examined predictors of maintaining remission after tapering MTX from targeted therapy. Higher baseline disease activity scores and rheumatoid factor (RF) positivity were found to be associated with lower likelihood of maintaining remission (Table 2). Higher physician global assessment scores were associated with restarting MTX during the open-label LTE³⁸ (P < 0.01; Table 2).

Recapture of remission. In total, 2 studies reported on retreatment outcomes.^{29,38} Curtis et al²⁹ reported that remission was recaptured with retreatment in 75% of patients in the ETN monotherapy group by week 48. The LTE study³⁸ reported that patients who restarted MTX later than 4 weeks after entering the LTE had worse disease activity scores compared to those who restarted MTX earlier (Table 2).

Safety. Safety was reported by all studies and, overall, there were no significant differences between groups in the number of adverse events (AEs), serious AEs, and discontinuations of treatment as a result of AEs. In total, 3 studies—SEAM, COMP-ACT, and Keystone et al³⁶—noted a numerical increase in the frequency of AEs in the patients treated with MTX compared to other arms (Table 2). The study by Pablos et al (JUST-ACT)³² reported higher AEs in the TCZ monotherapy group compared to the MTX-treated group (Table 2).

Metaanalysis. The metaanalysis, conducted among 2000

	Study, Year, Author, Design	Proportion Maintaining REM	Time of REM Assessment After Taper	Mean Disease Activity Scores	Functional Outcomes	Patient-Reported Outcomes	Radiographic Outcomes	Predictors	Proportion Recapture of REM With Retreatment	Proportion Safety AEs/SAEs/ withdrawn because of AEs
7.3.1. Monthly, Monthly, M	SEAM ²⁹ , 2021, Curtis, RCT	SDAI REM: ETN+MTX: 53% ETN mono: 50%	48 wks	Proportion with SDAI > 11: ETN+MTX: 78% ETN mono: 75%	NR	NR	NR	Higher baseline SDAI, RF positivity less likely to maintain RFM	At 12 wks/48wks ETN+MTX: 47%/80% ETN mono: 42%/75%	ETN+MTX: 62%/6%/0% ETN mono: 56%/4%/2%
	WERT-2 ³⁹ , 019 (ABS), .mery, .CT	SDAI REM: ABA+MTX: 74% ABA mono: 57%		Adjusted mean diff in SDAI score from ABA+MTX: Scop MTX: 1.12 (97.5% CI-1.08 to 3.32)	Adjusted mean diff in HAQ-DI: ABA+MTX: -0.04 ABA mono: 0.16 (ABS 2020) ³⁸	Adjusted mean diff in SF-36 PFS: ABA+MTX: 1.68 ABA mono: -1.45 (ABS 2020) ³⁸	Proportion with nonprogression: ABA+MTX: 87% (95% CI77-97) Stop MTX: 87% (95% CI77-98)	NR	NR	ABA+MTX: 44%/6%/0% ABA mono: 51%/0%/0%
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	0.RAL Shift ³⁵ , 019, Sohen, CT		24 wks Persistent REM wks 12, 24, 36, 48	LSM Δ DAS28 4(ESR) ⁶ : 0.30 (95% CI 0.12-0.48) n.s. NI met	Similar LSM A in HAQ-DI	LSM Δ in SF-36 PCS similar	NR	Baseline CDAI OR 0.32 (95% CI 0.24 0.43) P < 0.0001 (multivariable analysis, ABS 2019) ³⁶	NR	TOF+MTX: 41%/2%/2% TOF: 41%/4%/2% n.s.
	COMET, 010 ²⁰ /2019 ABS) ²¹ , Emery CT		52 wks	NR	Proportion normal HAQ-DI: ETN+MTX: 81.5% ETN: 77.8%		NR in 2019 ABS	NR	NR	ETN+MTX: 82%/7%/NR ETN mono: 80%/9%/NR n.s.
7,2020 DAS28 REM: Not specified Maintenance of Δ DHAQ-D12 0.22: CZP+MTX NR NR NR CZP+MTX:41% (WD time NR) DAS28 > 1.2 with CZP+MTX:44% vs CZP: CZP+MTX:41% (WD time NR) DAS28 > 1.2 with CZP+MTX:44% vs CZP: CZP:41% WD time NR) DAS28 > 1.2 with CZP+MTX:44% vs CZP: P = 1.0 2.6% (upper limit of P = 0.377 (VAS mm): 21.7 vs 9.9, (P = 0.026) PGA (VAS mm): 21.7 vs 90% CI: 19%; 1-sided P = 0.377 (VAS mm): 21.7 vs 90% CI: 19%; 1-sided P = 0.377 (VAS mm): 21.7 vs 9.9, (P = 0.026) PGA (VAS mm): 23.3 vs 34.8 Piin (VAS mm): 23.3 vs 34.8	UST-ACT ³² , 019, ablos, CCT	DAS28 REM: TCZ+MTX: 82% TCZ mono: 76%	12 wks		No diff in HAQ score P = 0.674 (9)	 \$\Delta SF-12 PCS: TCZ+MTX: 0.80 \$95% CI -1.1 to 2.7\$; MTX: -2.58 \$5% CI -4.48 to -0.67\$) \$P = 0.015\$ \$P = 0.015\$ \$P eGA, MDGA\$ 	NR	NR	NR	TCZ+MTX: 49%/1%/NR TCZ: 55%/5%/NR
	.CT .CT	DAS28 REM: CZP+MTX: 41% CZP: 41% P = 1.0	Not specified (WD time NR)	Maintenance of Δ DAS28 > 1.2 with absolute risk diff: 2.6% (upper limit of 90% CI: 19%; 1-sided P = 0.402) NI not met		CZP+MTX vs CZP: Morning stiffness (VAS mm): 21.7 vs 39.9 (<i>P</i> = 0.026) PtGA (VAS mm): 32.3 vs 34.8 Pain (VAS mm): 35.0 vs 38.2 Fatigue (VAS mm): 43.7 vs 43.4 & work loss: 7.4 vs 5.4 TJC: 2.1 vs 3.1	NR	NR	NR	CZP+MTX: 72%/5%/0% CZP: 69%/4%/2%

Meng et al

Proportion Safety AEs/SAEs/ withdrawn because of AEs	TCZ+MTX: 68%/6%/NR TCZ mono: 62%/4%/NR Higher in TCZ+MTX 1.5% developed anti-TCZ antibodis (while on MTX)	Proportion A Es/withdrawn: TCZ+stable MTX: 72%/13 TCZ+taper MTX: 72%/12% Proportion S A Es: TCZ-related: 5% MTX-related: 3%	ETN+MTX: 86%/16%/NR ETN mono: 88%/11%/NR n.s.	MTX use group: 93%/29%/9% MTX nonuse: 89%/30%/8% Infectious AEs higher er in MTX use vs nonuse: 73% vs 67%
Proportion Recapture of REM With Retreatment	NR	NR	NR	Median (IQR) time to 1st MTX restart: 5.1 (0.1-31.4) M wks; higher DAS28 if patients restarted later
Predictors	NR	NR	Ŷ	Higher PGA associated with MTX use during OLE P < 0.01
Radiographic Outcomes	NR	NR	Proportion progression in mTSS: ETN+MTX: 13% ETN: 14% n.s.	Proportion with no progression: MTX use: 46% MTX nonuse: 50%
Patient-Reported Outcomes	NR	NR	NR	NR
Functional F Outcomes	NR	NR	HAQ-DI, Δ mean score (SD): ETN+MTX: 0.1 (0.5) ETN: 0.2 (0.4)	Proportion normal function (HAQ-DI < 0.5): MTX use: 45% MTX nonuse: 58%
Mean Disease Activity Scores	Adjusted mean Δ in DAS28-ESR: TCZ+MTX:0.14 (95% CI -0.11 to 0.39) TCZ mono: 0.46 (95% CI 0.22 to 0.70) Adjusted diff: 0.318 (95% CI 0.45 to 0.592) NI met	Mean Δ DAS28 n.s.	DAS28-ESR mean (SD) Δ n.s.	None
Time of REM Assessment After Taper	28 wks	48 wks	18 mos	Up to 3 yrs after MTX WD; MTX restarted at varying timepoints
Proportion Maintaining REM	DAS28 REM: TCZ+MTX: 55% TCZ.48% Between-group diff -7% (95% CI -18% to 5%)	DAS28 REM: TCZ+stable MTX: 51% TCZ+taper MTX: 50% P = 0.902	DAS28 REM: ETN+MTX: 51% (95% CI 37-65) ETN: 40% (95% CI 26-54)	PREMIER OLE, DAS28-CRP 2018, REM: Keystone ³⁸ , MTX use: 48% observational MTX nonuse: 50%
Study, Year, Author, Design	COMP-ACT ³³ , 2018, ' Kremet, B RCT B	ACT-TAPER ³⁴ , 2018, Edwards, RCT	CAMEO, 2016, Keystone ³⁰ / 5 2014, Pope ³¹ RCT	PREMIER OLE, 2018, Keystone ³⁸ , observational ^N

Where available, point estimates are reported.^b Both DAS28-ESR and DAS28-CRP REM defined as score < 2.6; ^c Targeted therapy includes biologic disease-modifying antirheumatic drugs and Janus kinase inhibitor. ABA: abatacept; ABS: abstract; ADA: adalimumab; AE: adverse event; AVERT-2: Assessing Very Early Rheumatoid Arthritis Treatment-2; CAMEO: Canadian Methotrexate and Eranercept Outcome Study; CDAI: Clinical Disease Activity Index; COMET: Combination of Disease Activity Score in 28 joints with four variables including C-reactive protein; DAS28-4(ESR): Disease Activity Score in 28 joints with four variables including ESR; ETN: etanetrept; HAQ: Health Assessment Questionnaire; HAQ-DI: NI: noninferiority; NR: not reported; n.s.: not significant; OLE: open-label extension; OR: odds ratio; PCS: physical component score; PFS: physical functioning scale; PGA: physical assessment; PtGA: patient global assessment; Methotrexate and Etanercept in Active Early Rheumatoid Arthritis; CRP: C-reactive protein; CZP: certolizumab; DAS28: Disease Activity Score in 28 joints; diff: difference; A: change in; ESR: erythrocyte sedimentation rate; DAS28-4(CRP); Health Assessment Questionnaire-Disability Index; LSM: least squares mean; MCS: mental component score; MDGA: physician global assessment of disease; mono: monotherapy; mTSS: modified total Sharp score; MTX: methotrexate; RCT: randomized controlled trial; REM: remission; RF: rheumatoid factor; SAE: serious adverse event;; SDAI: simplified disease activity index; SEAM: Study of Etanercept and Methotrexate in Combination or as Monotherapy; SF-12: 12-12: them Short Form Health Survey; SF-36: 36-Item Short Form Health Survey; TCZ: to cilizumab; TJC: tender joint count; TOF: tofacitinib; WD: withdrawal.

Table 2. Continued.

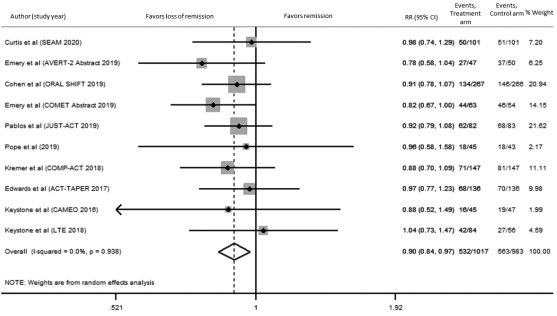


Figure 3. Metaanalysis of studies tapering methotrexate from targeted therapy. AVERT-2: Assessing Very Early Rheumatoid Arthritis Treatment-2; CAMEO: Canadian Methotrexate and Etanercept Outcome; COMET: Combination of Methotrexate and Etanercept in Active Early Rheumatoid Arthritis; LTE: long-term extension; RR: risk ratio; SEAM: Study of Etanercept and Methotrexate in Combination or as Monotherapy

participants with RA from 10 studies, showed a pooled RR for maintaining remission after tapering MTX from targeted therapy of 0.90 (95% CI 0.84-0.97; Figure 3). There was no heterogeneity among the studies in this group $(I^2 = 0\%, P = 0.94)$. Among the studies that enrolled patients with early RA, the RR was 0.84 (95% CI 0.73-0.98) and the heterogeneity was 0% (P =0.39). Among studies with patients with established RA, the RR was 0.92 (95% CI 0.85-1.01) and there was 0% heterogeneity present (P > 0.99; Supplementary Figure S1, available with the online version of this article). We specifically evaluated remission outcomes, rather than LDA, after tapering. Since some studies used LDA in their criteria to taper MTX, we performed a separate metaanalysis on the RRs of maintaining LDA after tapering MTX. We found similar results to the ones reported above (RR 0.92, 95% CI 0.86-0.98; Supplementary Figure S2, available with the online version of this article). Additionally, we performed a sensitivity analysis, where we omitted the 2018 LTE study³⁸ as it had a higher bias. Again, we found similar results (RR 0.90, 95% CI 0.83-0.97; Supplementary Figure S3, available with the online version of this article). Figure 4 shows our funnel plot for all included studies, along with a fitted line representing the Egger test for asymmetry; the results indicate minimal publication bias. Results from both the Egger test and the Harbord modified test for small study effects were found to be not statistically significant, indicating weak evidence of small study effects. Risk differences were calculated with an overall pooled risk difference of -0.05 (95% CI -0.10 to -0.01; Supplementary Figure S4, available with the online version of this article). Using the pooled estimate, if one were to taper MTX from targeted therapy in 20 patients, 2 (10%) patients would not be able to sustain remission.

DISCUSSION

To our knowledge, this is the first study and systematic review with metaanalysis to examine the effects of tapering MTX in patients with RA who combined MTX with a broad range of targeted therapies. Our metaanalysis showed that patients who tapered MTX from targeted therapy had a 10% reduction in the ability to sustain remission compared to not tapering therapy (RR 0.90, 95% CI 0.84-0.97) for up to 18 months. There was no heterogeneity, and our CIs were narrow.

These data extend those from Subesinghe et al¹⁷ who published a narrative review on tapering MTX, which included 2 trials of MTX, one with IFX in Infliximab Rheumatoid Arthritis Methotrexate Tapering (iRAMT) trial and ETN (COMET trial,²¹ included in the present review). In the 2005 iRAMT trial, MTX was tapered in patients who had achieved a 40% reduction in tender and swollen joint counts from baseline with combination IFX/MTX therapy. In total, 75% of patients were able to taper MTX to a minimum dose of 5 mg/week without loss of efficacy, suggesting low doses of MTX may help protect against loss of efficacy of IFX. Other classes of targeted therapies were not evaluated. To our knowledge, this is the first systematic review and metaanalysis to address tapering of MTX from a range of targeted therapies, including IL-6i and JAKi. Both of these targeted therapies have also been shown to be effective as monotherapies in RA.^{4,8-10,41,42} Several of our reviewed studies showed numerically increased AEs in patients treated with

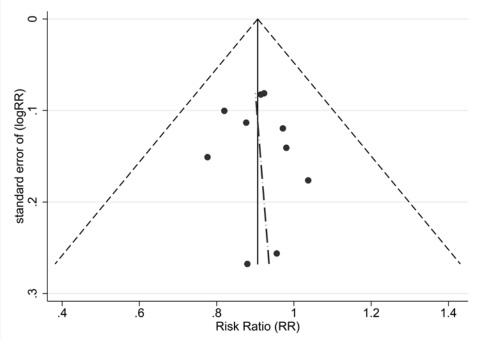


Figure 4. Funnel plot with pseudo 95% CIs for studies tapering methotrexate from targeted therapy. The fitted line represents the Egger test for asymmetry.

MTX compared to those on targeted therapies alone.^{29,33,38} Our patients who may now be taking any of a wide range of targeted therapies often wish to taper their MTX because of intolerance. This review helps inform patients and their physicians as to whether this is a good decision.

Longer follow-up times were associated with lower remission rates, underscoring the importance of including longer follow-up times in tapering studies in RA. Loss of remission over time is common even without changes in treatment.⁴³ Three^{21,29,30} of the 5 TNFi studies examined ETN, which is not associated with antidrug antibodies and may not benefit as much from concomitant treatment with MTX. It is possible that if the other TNFi drugs were more broadly represented, the data may have been different. Although the development of antidrug antibodies could occur if patients remain on monotherapy with bDMARDs and specifically TNFi after MTX tapering, there is little evidence to support this. An observational study found that the long-term drug survival of TNFi was not significantly different between those who discontinued MTX and those who continued it (hazard ratio 1.046, 95% CI 0.76-1.44), though how long patients remained off MTX was not explicitly reported.14

We expected that patients who tapered MTX gradually or allowed a dose reduction without stopping would maintain remission more so than with abrupt withdrawal. There was no clear association of tapering schemes with remission outcomes; however, only 2 studies^{21,34} performed a gradual dose reduction, one of which tapered MTX off within 4 weeks.²¹ The other study tapered MTX over 24 weeks and stopped tapering in the event of flare,³⁴ allowing subjects to remain in the taper group if retreatment recaptured disease control at a dose that was not higher than at randomization. This was the only study we reviewed that allowed dose reduction of MTX without stopping in their protocol; the mean dose of MTX in the tapering group was not reported.

We analyzed both early and established RA and found that both groups had an increased risk of not being able to sustain remission, but it did not reach significance in established RA. Prior studies have shown that those with early RA may be more successful in tapering bDMARDs.^{18,44,45} Only 3 studies on early RA^{21,38,39} were analyzed in our review, and more studies are needed to address this question.

Patient-reported physical function was reported to worsen when MTX was tapered; although it was not statistically significant, it could become significant with longer follow-up periods. Only 3 studies reported on predictors of maintaining disease control after tapering MTX.^{29,35,38} Higher baseline disease activity and RF seropositivity were associated with reduced likelihood of maintaining remission, similar to prior studies.⁴⁶ A higher physician global score was associated with restarting MTX in the LTE study. A systematic review of biomarkers for successful tapering of bDMARDs found that shorter symptom duration, lower erosion scores, and higher ADA drug levels were significant predictors for successful tapering, but evidence was limited by low-quality studies and reporting bias.¹⁸ Understanding the subset of patients who can successfully taper RA therapies will help prevent disease worsening and avoid the undesirable scenario of not being able to recapture disease control with retreatment.

Only 1 study looked at retreatment after tapering MTX and reported a 75% rate of recapture of remission,²⁹ similar to that reported by prior studies tapering bDMARDs.^{47,48} However, these results should be interpreted with caution, as only 1 study reported on retreatment outcomes. Only 1 trial, not eligible for the metaanalysis, evaluated outcomes of patients tapering both csDMARDs and bDMARDs and reported recapture of DAS28 remission by 65% of patients tapering csDMARDs.⁴⁹ More research on the recapture of remission after tapering MTX from targeted therapy is needed.

Several limitations of our review should be considered. Studies differed with respect to whether patients had early or established RA, whether patients were MTX-naïve or inadequate responders, the tapering strategy used, and the criteria used to taper (Table 1). Patients who were MTX-naïve were studied, not surprisingly, in the 3 early RA studies^{20,38,39} that we analyzed separately as previously mentioned (Supplementary Figure S1, available with the online version of this article). The current guidelines recommend gradual tapering of MTX if this is necessary for the care of a given patient; however, most available evidence for MTX tapering is based on studies in which MTX is either abruptly or rapidly withdrawn. It is possible that more gradual tapering of MTX, as examined in some studies, may have allowed each patient to determine the optimal dose of MTX needed to maintain remission after tapering. Overall, our studies had no heterogeneity, with an I^2 of 0% (P = 0.94). This could limit the external validity of this study, but it more likely reflects the similarity of the populations being studied.

We included pragmatic studies to increase generalizability to patients seen in routine practice; however, because of their open-label design, they scored higher on the risk-of-bias assessment tool. These studies also provided longer follow-up data, which we felt was important in addressing our study question. Our 1 observational LTE study was judged to have serious risk of bias.³⁸ We included it because it met our inclusion criteria; reported remission outcomes, including an adjusted analysis with propensity scoring; and evaluated MTX tapering in a realworld setting. Our sensitivity analysis excluding the LTE study showed similar results.

One strength of this study is that we were able to estimate the proportion of people who could sustain remission when withdrawing MTX from therapy combined with multiple classes of advanced therapies. Although there were too few studies to draw conclusions about specific classes of drugs, the pooled data were consistent and could inform a broader group of RA patients needing to stop MTX, regardless of which targeted therapy was currently in use, be it a TNFi, IL-6i, or JAKi. Of note, we found no randomized MTX-tapering study for patients using it in combination with RTX that met our criteria.

We specifically evaluated remission outcomes rather than LDA after tapering. Only 2 studies^{29,39} in our review used remission alone as their tapering criteria, with the other studies^{21,30,32-35,37,38} using less stringent criteria to taper. It is possible that if we looked at LDA as our outcome after tapering, our results may have shown higher proportions of maintaining disease control. We further evaluated this by performing a metaanalysis of the RR of maintaining LDA after tapering MTX, and we found similar results (RR 0.92, 95% CI 0.86-0.98; Supplementary Figure S2, available with the online version of this article).

Current guidelines recommend achieving sustained remission prior to tapering therapy; thus, this stricter criterion was applied to inform tapering of MTX from targeted therapy.

In summary, the results of our systematic review and metaanalysis supported our hypothesis that patients with controlled RA have a low risk of not being able to sustain remission when tapering MTX from targeted therapy up to 18 months. This review adds to the body of evidence to help inform ACR guidelines regarding tapering of MTX from combination therapy. It can also help inform discussions with patients who have controlled RA and who are struggling with common MTX-related intolerances, such as hair loss, stomatitis, nausea, diarrhea, and elevated liver enzymes, and wish to taper it. Our data may aid in the discussion among female patients of childbearing age, who are concerned about the teratogenicity of MTX. The authors of this review advocate for the continuation of MTX with targeted therapy when it is well tolerated, as the long-term effects of tapering beyond 18 months requires further study and there were indicators of potential worsening of functional outcomes in some studies. Patients need to be informed that disease control may be lost over time if they remain on targeted monotherapy and that the recapture of remission may not be possible with retreatment. Most importantly, patients need to continue careful follow-up over time, as prompt retreatment to recapture disease control is essential.

Further research is needed that includes studies with longer follow-up periods that also address predictors of successful tapering and long-term consequences of treatment withdrawal, including worsening of function, measures of joint damage, whether drug immunogenicity develops, and whether there is an advantage to gradual tapering regimens. Whether targeted therapy used as monotherapy in RA can also be tapered is an important sequitur to this study.

ACKNOWLEDGMENT

The authors wish to thank Omar J. Bruce for his assistance with some of the analyses performed for this review.

ONLINE SUPPLEMENT

Supplementary material accompanies the online version of this article.

REFERENCES

- Smolen JS, Landewé RBM, Bijlsma JWJ, et al. EULAR recommendations for the management of rheumatoid arthritis with synthetic and biological disease-modifying antirheumatic drugs: 2019 update. Ann Rheum Dis 2020;79:685-99.
- Choy E, Aletaha D, Behrens F, et al. Monotherapy with biologic disease-modifying anti-rheumatic drugs in rheumatoid arthritis. Rheumatology 2017;56:689-97.
- Emery P, Sebba A, Huizinga TWJ. Biologic and oral diseasemodifying antirheumatic drug monotherapy in rheumatoid arthritis. Ann Rheum Dis 2013;72:1897-904.
- Detert J, Klaus P. Biologic monotherapy in the treatment of rheumatoid arthritis. Biologics 2015;9:35-43.
- Nikiphorou E, Negoescu A, Fitzpatrick JD, et al. Indispensable or intolerable? Methotrexate in patients with rheumatoid and psoriatic arthritis: a retrospective review of discontinuation rates from a large UK cohort. Clin Rheumatol 2014;33:609-14.

- Curtis JR, Bykerk VP, Aassi M, Schiff M. Adherence and persistence with methotrexate in rheumatoid arthritis: a systematic review. J Rheumatol 2016;43:1997-2009.
- Harley CR, Frytak JR, Tandon N. Treatment compliance and dosage administration among rheumatoid arthritis patients receiving infliximab, etanercept, or methotrexate. Am J Manag Care 2003;9:S136-43.
- 8. Emery P, Pope JE, Kruger K, et al. Efficacy of monotherapy with biologics and JAK inhibitors for the treatment of rheumatoid arthritis: a systematic review. Adv Ther 2018;35:1535-63.
- 9. Yamanaka H, Tanaka Y, Takeuchi T, et al. Tofacitinib, an oral Janus kinase inhibitor, as monotherapy or with background methotrexate, in Japanese patients with rheumatoid arthritis: an open-label, long-term extension study. Arthritis Res Ther 2016;18:34.
- 10. Fleischmann R, Takeuchi T, Schiff M, et al. Efficacy and safety of long-term baricitinib with and without methotrexate for the treatment of rheumatoid arthritis: experience with baricitinib monotherapy continuation or after switching from methotrexate monotherapy or baricitinib plus methotrexate. Arthritis Care Res 2020;72:1112-21.
- Shaw Y, Metes ID, Michaud K, et al. Rheumatoid arthritis patients' motivations for accepting or resisting disease-modifying antirheumatic drug treatment regimens. Arthritis Care Res 2018;70:533-41.
- Kuijper TM, Lamers-Karnebeek FBG, Jacobs JWG, Hazes JMW, Luime JJ. Flare rate in patients with rheumatoid arthritis in low disease activity or remission when tapering or stopping synthetic or biologic DMARD: a systematic review. J Rheumatol 2015; 42:2012-22.
- Mangoni AA, Al Okaily F, Almoallim H, Al Rashidi S, Mohammed RHA, Barbary A. Relapse rates after elective discontinuation of anti-TNF therapy in rheumatoid arthritis: a meta-analysis and review of literature. BMC Rheumatol 2019;3:10.
- 14. Manders SHM, van de Laar MAFJ, Rongen-van Dartel SAA, et al. Tapering and discontinuation of methotrexate in patients with RA treated with TNF inhibitors: data from the DREAM registry. RMD Open 2015;1:e000147.
- 15. Kremer JM, Weinblatt ME, Bankhurst AD, et al. Etanercept added to background methotrexate therapy in patients with rheumatoid arthritis: continued observations. Arthritis Rheum 2003;48:1493-9.
- Fraenkel L, Bathon JM, England BR, et al. 2021 American College of Rheumatology Guideline for the Treatment of Rheumatoid Arthritis. Arthritis Care Res 2021;73:924-39.
- Subesinghe S, Scott IC. Key findings from studies of methotrexate tapering and withdrawal in rheumatoid arthritis. Expert Rev Clin Pharmacol 2015;8:751-60.
- Tweehuysen L, van den Ende CH, Beeren FMM, Been EMJ, van den Hoogen FHJ den Broeder AA. Little evidence for usefulness of biomarkers for predicting successful dose reduction or discontinuation of a biologic agent in rheumatoid arthritis: a systematic review. Arthritis Rheumatol 2017;69:301-8.
- Liberati A, Altman DG, Tetzlaff J, et al. The PRISMA statement for reporting systematic reviews and meta-analyses of studies that evaluate health care interventions: explanation and elaboration. PLoS Med 2009;6:e1000100.
- 20. Emery P, Breedveld F, van der Heijde D, et al. Two-year clinical and radiographic results with combination etanercept-methotrexate therapy versus monotherapy in early rheumatoid arthritis: a two-year, double-blind, randomized study. Arthritis Rheum 2010;62:674-82.
- 21. Emery P, Breedveld F, Pedersen R, et al. Relationships between DAS28 response and clinical, functional and radiographic outcomes

in year 2 of the COMET study of etanercept in patients with rheumatoid arthritis [abstract]. Arthritis Rheumatol 2019;71:2382.

- 22. Higgins JPT, Savovic J, Page MJ, Sterne JAC. Revised Cochrane risk-of-bias tool for randomized Trials (RoB 2). [Internet. Accessed September 27, 2022.] Available from: https://www.riskofbias.info/ welcome/rob-2-0-tool/current-version-of-rob-2
- 23. Jonathan AC, Sterne MAH, Barnaby C Reeves, et al. Current version of ROBINS-I. [Internet. Accessed September 27, 2022.] Available from: https://www.riskofbias.info/welcome/home/ current-version-of-robins-i
- 24. Ranganathan P, Aggarwal R, Pramesh CS. Common pitfalls in statistical analysis: Odds versus risk. Perspect Clin Res 2015; 6:222-4.
- Tufanaru C, Munn A, Stephenson M, Aromotaris E. Common pitfalls in statistical analysis: odds versus risk. Int J Evid Based Healthc 2015;13:196-207.
- 26. von Hipple PT. The heterogeneity statistic I(2) can be biased in small meta-analyses. BMC Med Res Methodol 2015;15:35.
- 27. Harbord RM, Egger M, Sterne JAC. A modified test for small-study effects in meta-analyses of controlled trials with binary endpoints. Stat Med 2006;25:3443-57.
- 28. Harbord RM, Harris RJ, Sterne JAC. Updated tests for small-study effects in meta-analyses. Stata J 2009;9:197-210.
- 29. Curtis JR, Emery P, Karis E, et al. Etanercept or methotrexate withdrawal in rheumatoid arthritis patients in sustained remission. Arthritis Rheumatol 2021;73:759-68.
- Keystone EC, Pope JE, Thorne JC, et al. Two-year radiographic and clinical outcomes from the Canadian Methotrexate and Etanercept Outcome study in patients with rheumatoid arthritis. Rheumatology 2016;55:327-34.
- 31. Pope JE, Haraoui B, Thorne JC, et al. The Canadian methotrexate and etanercept outcome study: a randomised trial of discontinuing versus continuing methotrexate after 6 months of etanercept and methotrexate therapy in rheumatoid arthritis. Ann Rheum Dis 2014;73:2144-51.
- 32. Pablos JL, Navarro F, Blanco FJ, et al. Efficacy of tocilizumab monotherapy after response to combined tocilizumab and methotrexate in patients with rheumatoid arthritis: the randomised JUST-ACT study. Clin Exp Rheumatol 2019;37:437-44.
- 33. Kremer JM, Rigby W, Singer NG, et al. Sustained response following discontinuation of methotrexate in patients with rheumatoid arthritis treated with subcutaneous tocilizumab: results from a randomized, controlled trial. Arthritis Rheumatol 2018;70:1200-8.
- Edwards CJ, Östör AJK, Naisbett-Groet B, et al. Tapering versus steady-state methotrexate in combination with tocilizumab for rheumatoid arthritis: a randomized, double-blind trial. Rheumatology 2018;57:84-91.
- 35. Cohen S, Pope J, Haraoui B, et al. Methotrexate withdrawal in patients with rheumatoid arthritis who achieve low disease activity with tofacitinib modified-release 11 mg once daily plus methotrexate (ORAL Shift): a randomised, phase 3b/4, non-inferiority trial. The Lancet Rheumatology, 2019;1:e23-34.
- 36. Yamaoka K, Cohen SB, Sugiyama N, et al. Predictors of durable clinical response to tofacitinib 11 mg once daily with or without methotrexate in patients with rheumatoid arthritis: post hoc analysis of data from a phase 3b/4 methotrexate withdrawal study [abstract]. Arthritis Rheumatol 2020;72:0200.
- Pope J, Rampakakis E, Vaillancourt J, et al. An open-label randomized controlled trial of DMARD withdrawal in RA patients achieving therapeutic response with certolizumab pegol combined with DMARDs. Rheumatology 2020;59:1522-8.
- 38. Keystone EC, Breedveld FC, Kupper H, Li Y, Florentinus S,

Sainsbury I. Long-term use of adalimumab as monotherapy after attainment of low disease activity with adalimumab plus methotrexate in patients with rheumatoid arthritis. RMD Open 2018;4:e000637.

- Emery P, Tanaka Y, Bykerk V, et al. Maintenance of remission following dose de-escalation of abatacept in early, MTX-naïve, ACPA-positive patients with RA: results from a randomized phase IIIb study [abstract]. Arthritis Rheumatol 2019;71:L11.
- 40. Emery P, Tanaka Y, Bykerk V, et al. Maintenance of SDAI remission and patient-reported outcomes (PROs) following dose de-escalation of abatacept in MTX-naïve anti-citrullinated protein antibody (ACPA)+ patients with early RA: results from AVERT-2, a randomised phase IIIB study [abstract]. Ann Rheum Dis 2020;79:985-6.
- Fleischmann R, Mysler E, Hall S, et al. Efficacy and safety of tofacitinib monotherapy, tofacitinib with methotrexate, and adalimumab with methotrexate in patients with rheumatoid arthritis (ORAL Strategy): a phase 3b/4, double-blind, head-to-head, randomised controlled trial. Lancet 2017;390:457-68.
- 42. Smolen JS, Pangan AL, Emery P, et al. Upadacitinib as monotherapy in patients with active rheumatoid arthritis and inadequate response to methotrexate (SELECT-MONOTHERAPY): a randomised, placebo-controlled, double-blind phase 3 study. Lancet 2019;393:2303-11.
- 43. Prince FHM, Bykerk VP, Shadick NA, et al. Sustained rheumatoid arthritis remission is uncommon in clinical practice. Arthritis Res Ther 2012;14:R68.

- 44. Emery P, Burmester GR, Bykerk VP, et al. Evaluating drug-free remission with abatacept in early rheumatoid arthritis: results from the phase 3b, multicentre, randomised, active-controlled AVERT study of 24 months, with a 12-month, double-blind treatment period. Ann Rheum Dis 2015;74:19-26.
- Vodencarevic A, Tascilar K, Hartmann F, et al. Advanced machine learning for predicting individual risk of flares in rheumatoid arthritis patients tapering biologic drugs. Arthritis Res Ther 2021;23:67.
- 46. Schlager L, Loiskandl M, Aletaha D, Radner H. Predictors of successful discontinuation of biologic and targeted synthetic DMARDs in patients with rheumatoid arthritis in remission or low disease activity: a systematic literature review. Rheumatology 2020;59:324-34.
- 47. Ghiti Moghadam M, Vonkeman HE, Ten Klooster PM, et al. Stopping tumor necrosis factor inhibitor treatment in patients with established rheumatoid arthritis in remission or with stable low disease activity: a pragmatic multicenter, open-label randomized controlled trial. Arthritis Rheumatol 2016;68:1810-7.
- 48. Tanaka Y, Takeuchi T, Mimori T, et al. Discontinuation of infliximab after attaining low disease activity in patients with rheumatoid arthritis: RRR (remission induction by Remicade in RA) study. Ann Rheum Dis 2010;69:1286-91.
- Kuijper TM, Luime JJ, de Jong PHP, et al. Tapering conventional synthetic DMARDs in patients with early arthritis in sustained remission: 2-year follow-up of the tREACH trial. Ann Rheum Dis 2016;75:2119-23.