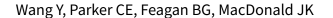


Cochrane Database of Systematic Reviews

Oral 5-aminosalicylic acid for maintenance of remission in ulcerative colitis (Review)



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[Intervention Review]

Oral 5-aminosalicylic acid for maintenance of remission in ulcerative colitis

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ABSTRACT

Background

Oral 5-aminosalicylic (5-ASA) preparations were intended to avoid the adverse effects of sulfasalazine (SASP) while maintaining its therapeutic benefits. Previously, it was found that 5-ASA drugs were more effective than placebo but had a statistically significant therapeutic inferiority relative to SASP. This updated review includes more recent studies and evaluates the effectiveness, doseresponsiveness, and safety of 5-ASA preparations used for maintenance of remission in quiescent ulcerative colitis.

Objectives

The primary objectives were to assess the efficacy, dose-responsiveness and safety of oral 5-ASA compared to placebo, SASP, or 5-ASA comparators for maintenance of remission in quiescent ulcerative colitis. A secondary objective was to compare the efficacy and safety of once daily dosing of oral 5-ASA with conventional (two or three times daily) dosing regimens.

Search methods

A literature search for relevant studies (inception to 9 July 2015) was performed using MEDLINE, EMBASE and the Cochrane Library. Review articles and conference proceedings were also searched to identify additional studies.

Selection criteria

Studies were accepted for analysis if they were randomized controlled trials with a minimum treatment duration of six months. Studies of oral 5-ASA therapy for treatment of patients with quiescent ulcerative colitis compared with placebo, SASP or other 5-ASA formulations were considered for inclusion. Studies that compared once daily 5-ASA treatment with conventional dosing of 5-ASA and 5-ASA dose ranging studies were also considered for inclusion.

Data collection and analysis

The primary outcome was the failure to maintain clinical or endoscopic remission. Secondary outcomes included adherence, adverse events, withdrawals due to adverse events, and withdrawals or exclusions after entry. Trials were separated into five comparison groups: 5-ASA versus placebo, 5-ASA versus sulfasalazine, once daily dosing versus conventional dosing, 5-ASA versus comparator 5-ASA formulation, and 5-ASA dose-ranging. Placebo-controlled trials were subgrouped by dosage. Once daily versus conventional dosing studies were subgrouped by formulation. 5-ASA-controlled trials were subgrouped by common 5-ASA comparators (e.g. Asacol and Salofalk). Doseranging studies were subgrouped by 5-ASA formulation. We calculated the risk ratio (RR) and 95% confidence intervals (95% CI) for each outcome. Data were analyzed on an intention-to-treat basis.



Main results

Forty-one studies (8928 patients) were included. The majority of included studies were rated as low risk of bias. Ten studies were rated at high risk of bias. Seven of these studies were single-blind and three studies were open-label. However, two open-label studies and four of the single-blind studies utilized investigator performed endoscopy as an endpoint, which may protect against bias. 5-ASA was significantly superior to placebo for maintenance of clinical or endoscopic remission. Forty-one per cent of 5-ASA patients relapsed compared to 58% of placebo patients (7 studies, 1298 patients; RR 0.69, 95% CI 0.62 to 0.77). There was a trend towards greater efficacy with higher doses of 5-ASA with a statistically significant benefit for the 1 to 1.9 g/day (RR 0.65; 95% CI 0.56 to 0.76) and the ≥ 2 g/day subgroups (RR 0.73, 95% CI 0.60 to 0.89). SASP was significantly superior to 5-ASA for maintenance of remission. Forty-eight per cent of 5-ASA patients relapsed compared to 43% of SASP patients (12 studies, 1655 patients; RR 1.14, 95% CI 1.03 to 1.27). A GRADE analysis indicated that the overall quality of the evidence for the primary outcome for the placebo and SASP-controlled studies was high. No statistically significant differences in efficacy or adherence were found between once daily and conventionally dosed 5-ASA. Twenty-nine per cent of once daily patients relapsed over 12 months compared to 31% of conventionally dosed patients (8 studies, 3127 patients; RR 0.91, 95% CI 0.82 to 1.01). Eleven per cent of patients in the once daily group failed to adhere to their medication regimen compared to 9% of patients in the conventional dosing group (6 studies, 1462 patients; RR 1.22, 95% CI 0.91 to 1.64). There does not appear to be any difference in efficacy among the various 5-ASA formulations. Forty-four per cent of patients in the 5-ASA group relapsed compared to 41% of patients in the 5-ASA comparator group (6 studies, 707 patients; RR 1.08, 95% CI 0.91 to 1.28). A pooled analysis of two studies showed no statistically significant difference in efficacy between Balsalazide 6 g and 3 g/day. Twenty-three per cent of patients in the 6 g/day group relapsed compared to 33% of patients in the 3 g/day group (216 patients; RR 0.76; 95% CI 0.45 to 2.79). One study found Balsalazide 4 g to be superior to 2 g/ day. Thirty-seven per cent of patients in the 4 g/day Balsalazide group relapsed compared to 55% of patients in the 2 g/day group (133 patients; RR 0.66; 95% CI 0.45 to 0.97). One study found a statistically significant difference between Salofalk granules 3 g and 1.5 g/day. Twenty-five per cent of patients in the Salofalk 3 g/day group relapsed compared to 39% of patients in the 1.5 g/day group (429 patients; RR 0.65; 95% CI 0.49 to 0.86). Common adverse events included flatulence, abdominal pain, nausea, diarrhea, headache, dyspepsia, and nasopharyngitis. There were no statistically significant differences in the incidence of adverse events between 5-ASA and placebo, 5-ASA and SASP, once daily and conventionally dosed 5-ASA, 5-ASA and comparator 5-ASA formulations and 5-ASA dose ranging studies. The trials that compared 5-ASA and SASP may have been biased in favour of SASP because most trials enrolled patients known to be tolerant to SASP which may have minimized SASP-related adverse events.

Authors' conclusions

5-ASA was superior to placebo for maintenance therapy in ulcerative colitis. However, 5-ASA had a statistically significant therapeutic inferiority relative to SASP. Oral 5-ASA administered once daily is as effective and safe as conventional dosing for maintenance of remission in quiescent ulcerative colitis. There does not appear to be any difference in efficacy or safety between the various formulations of 5-ASA. Patients with extensive ulcerative colitis or with frequent relapses may benefit from a higher dose of maintenance therapy. High dose therapy appears to be as safe as low dose and is not associated with a higher incidence of adverse events.

PLAIN LANGUAGE SUMMARY

Oral 5-ASA compounds for maintaining remission in ulcerative colitis

Sulfasalazine (SASP) has been used for treating ulcerative colitis for decades. SASP is made up of 5-aminosalicylic acid (5-ASA) linked to a sulfur molecule. Up to a third of patients treated with SASP have reported side effects, which are thought to be related to the sulfur part of the molecule. Common side effects associated with SASP include nausea, indigestion, headache, vomiting and abdominal pain. 5-ASA drugs were developed to avoid the side effects associated with SASP. This review includes 41 randomized trials with a total of 8928 participants. Oral 5-ASA was found to be more effective than placebo (fake drug) for maintaining remission. Although oral 5-ASA preparations are effective for maintaining remission in ulcerative colitis, they are no more effective than sulfasalazine (SASP) therapy. People who have become well can remain so by continuing to take either medication. There is no evidence that side effects are more frequent with one or the other medication. However, the side effects of 5-ASA may be notably less than those associated with SASP therapy. Common side effects associated with 5-ASA included flatulence, abdominal pain, nausea, diarrhea, headache, dyspepsia (indigestion), and nasopharyngitis (inflammation of the nasal passages). Most of the trials comparing 5-ASA with SASP enrolled patients who were known to tolerate SASP. This may have reduced SASP-related side effects in these trials. Male infertility is associated with SASP and not with 5-ASA, so 5-ASA may be preferred for patients concerned about fertility. 5-ASA therapy is more expensive than SASP, so SASP may be the preferred option where cost is an important factor. Oral 5-ASA administered once daily is as effective and safe as conventional dosing (two or three times daily) for maintaining remission in ulcerative colitis. There does not appear to be any difference in efficacy or safety between the various formulations of 5-ASA. Patients with extensive ulcerative colitis or with frequent relapses may benefit from a higher dose of maintenance therapy. High dose therapy appears to be as safe as low dose and is not associated with a higher incidence of side effects.



Summary of findings for the main comparison. Oral 5-ASA versus placebo for maintenance of remission in ulcerative colitis

Oral 5-ASA versus placebo for maintenance of remission in ulcerative colitis

Patient or population: Patients with quiescent ulcerative colitis

Settings: Outpatients

Intervention: Oral 5-ASA versus placebo

Outcomes Illustrative comparative risks* (95% CI)		ative risks* (95% CI)	Relative effect (95% CI)	No of Participants (studies)	Quality of the evidence	Comments
	Assumed risk	Corresponding risk	(40 % 6.)	(Commiss)	(GRADE)	
	Control	Oral 5-ASA versus placebo				
Failure to maintain clinical or endoscopic remission at study endpoint	584 per 1000 ¹	403 per 1000 (362 to 450)	RR 0.69 (0.62 to 0.77)	1,298 (7 studies)	⊕⊕⊕⊕ high	
Any adverse event	393 per 1000 ¹	369 per 1000 (303 to 452)	RR 0.94 (0.77 to 1.15)	774 (3 studies)	⊕⊕⊕⊝ moderate ²	
Withdrawal due to adverse event	42 per 1000 ¹	36 per 1000 (19 to 68)	RR 0.86 (0.46 to 1.63)	1096 (5 studies)	⊕⊕⊕⊝ moderate ³	

^{*}The basis for the **assumed risk** (e.g. the median control group risk across studies) is provided in footnotes. The **corresponding risk** (and its 95% confidence interval) is based on the assumed risk in the comparison group and the **relative effect** of the intervention (and its 95% CI). **CI:** Confidence interval; **RR:** risk ratio

GRADE Working Group grades of evidence

High quality: Further research is very unlikely to change our confidence in the estimate of effect.

Moderate quality: Further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.

Low quality: Further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.

¹ Control group risk estimates come from control arm of meta-analysis, based on included trials.

² Downgraded one level due to sparse data (320 events).

³ Downgraded one level due to sparse data (41 events).

Oral 5-ASA versus SASP for maintenance of remission in ulcerative colitis

Patient or population: Patients with guiescent ulcerative colitis

Settings: Outpatients

Intervention: Oral 5-ASA versus SASP

Outcomes	Illustrative comparative risks* (95% CI)		Relative effect (95% CI)	No of Participants (studies)	Quality of the evidence	Comments
	Assumed risk	Corresponding risk	(33 % 61)	(Studies)	(GRADE)	
	Control	Oral 5-ASA versus SASP				
Failure to maintain clinical or en- doscopic remission at study end- point	429 per 1000 ¹	489 per 1000 (442 to 545)	RR 1.14 (1.03 to 1.27)	1,655 (12 studies)	⊕⊕⊕⊕ high	
Any adverse event	158 per 1000 ¹	169 per 1000 (130 to 221)	RR 1.07 (0.82 to 1.40)	1,138 (7 studies)	⊕⊕⊕⊝ moderate ²	
Withdrawal due to adverse event	54 per 1000 ¹	69 per 1000 (47 to 101)	RR 1.27 (0.87 to 1.87)	1,585 (10 studies)	⊕⊕⊕⊝ moderate ³	

^{*}The basis for the **assumed risk** (e.g. the median control group risk across studies) is provided in footnotes. The **corresponding risk** (and its 95% confidence interval) is based on the assumed risk in the comparison group and the **relative effect** of the intervention (and its 95% CI). **CI:** Confidence interval; **RR:** risk ratio

GRADE Working Group grades of evidence

High quality: Further research is very unlikely to change our confidence in the estimate of effect.

Moderate quality: Further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.

Low quality: Further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.

 $^{{}^{1}\}operatorname{Control}\operatorname{group}\operatorname{risk}\operatorname{estimates}\operatorname{come}\operatorname{from}\operatorname{control}\operatorname{arm}\operatorname{of}\operatorname{meta-analysis},\operatorname{based}\operatorname{on}\operatorname{included}\operatorname{trials}.$

² Downgraded one level due to sparse data (182 events).

³ Downgraded one level due to sparse data (97 events).

Once daily dosing versus conventional dosing for maintenance of remission in ulcerative colitis

Patient or population: Patients with guiescent ulcerative colitis

Settings: Outpatients

Intervention: Once daily oral 5-ASA versus conventional dosing of 5-ASA

Outcomes	Illustrative comparative risks* (95% CI)		Relative effect (95% CI)	No of Participants (studies)	Quality of the Comments evidence
	Assumed risk	Corresponding risk	(55 % 5.1)	(50.2.55)	(GRADE)
	Control	OD versus conventional			
Failure to maintain clinical or endoscopic remission at 6 months	184 per 1000 ¹	188 per 1000 (157 to 228)	RR 1.02 (0.85 to 1.23)	1,871 (3 studies)	⊕⊕⊙⊝ low ^{2,3}
Failure to maintain clinical or endoscopic remission at 12 months	314 per 1000 ¹	286 per 1000 (258 to 317)	RR 0.91 (0.82 to 1.01)	3,127 (8 studies)	⊕⊕⊕⊝ moderate ⁴
Failure to adhere to study medication regimen	87 per 1000 ¹	106 per 1000 (79 to 143)	RR 1.22 (0.91 to 1.64)	1,462 (6 studies)	⊕⊕⊕⊝ moderate ^{5,6}
Development of any adverse event	453 per 1000 ¹	453 per 1000 (417 to 489)	RR 1.00 (0.92 to 1.08)	2,714 (6 studies)	⊕⊕⊕⊕ high
Withdrawal due to adverse events	15 per 1000 ¹	20 per 1000 (12 to 32)	RR 1.31 (0.80 to 2.13)	3,737 (7 studies)	⊕⊕⊙⊝ low ⁷ ,8

^{*}The basis for the **assumed risk** (e.g. the median control group risk across studies) is provided in footnotes. The **corresponding risk** (and its 95% confidence interval) is based on the assumed risk in the comparison group and the **relative effect** of the intervention (and its 95% CI).

CI: Confidence interval; RR: risk ratio

GRADE Working Group grades of evidence

High quality: Further research is very unlikely to change our confidence in the estimate of effect.

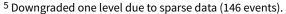
Moderate quality: Further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.

Low quality: Further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.

¹ Control group risk estimates come from control arm of meta-analysis, based on included trials.

² Downgraded one level due to risk of bias. Two studies in the pooled analysis were single-blind (investigator blind).

³ Downgraded one level due to sparse data (349 events).



- ⁶ Adherence was calculated using objective data (pill count or pharmacy data) in 4 of 6 studies in the analysis. One study used patient self-report to calculate adherence and one study did not describe how adherence was assessed.
- ⁷ Downgraded one level due to sparse data (64 events).
- ⁸ Downgraded one level due to risk of bias. One study in the pooled analysis was open-label and was rated as a high risk of bias due to lack of blinding. Three studies in the pooled analysis were single-blind (investigator blinded).

Summary of findings 4. Oral 5-ASA versus comparator 5-ASA formulation for maintenance of remission in ulcerative colitis

Oral 5-ASA versus comparator 5-ASA formulation for maintenance of remission in ulcerative colitis

Patient or population: Patients with quiescent ulcerative colitis

Settings: Outpatients

Intervention: Oral 5-ASA versus 5-ASA (different formulations)

Outcomes			Relative effect (95% CI)	No of Partici-	Quality of the Comm	nents
	Assumed risk		(33 /0 Ci)	(studies)	(GRADE)	
	Control	Oral 5-ASA versus 5-ASA				
Failure to maintain clinical or endoscopic remission at 12 months	407 per 1000 ¹	440 per 1000 (370 to 521)	RR 1.08 (0.91 to 1.28)	707 (6 studies)	⊕⊕⊙⊝ low2,3	
Development of any adverse event	686 per 1000 ¹	645 per 1000 (569 to 734)	RR 0.94 (0.83 to 1.07)	357 (4 studies)	⊕⊕⊙⊝ low ^{4,5}	
Withdrawal due to adverse events	44 per 1000 ¹	55 per 1000 (25 to 122)	RR 1.25 (0.56 to 2.78)	457 (5 studies)	⊕⊝⊝⊝ very low ^{6,7}	

^{*}The basis for the **assumed risk** (e.g. the median control group risk across studies) is provided in footnotes. The **corresponding risk** (and its 95% confidence interval) is based on the assumed risk in the comparison group and the **relative effect** of the intervention (and its 95% CI).

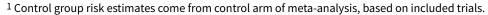
CI: Confidence interval; RR: risk ratio

GRADE Working Group grades of evidence

High quality: Further research is very unlikely to change our confidence in the estimate of effect.

Moderate quality: Further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.

Low quality: Further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.



- ² Downgraded one level due to sparse data (300 events).
- ³ Downgraded one level due to risk of bias. Two studies in pooled analysis were single blind and one was open label.
- ⁴ Downgraded one level due to sparse data (236 events).
- ⁵ Downgraded one level due to risk of bias. One study in the pooled analysis was open label.
- ⁶ Downgraded two levels due to very sparse data (23 events) and very wide confidence intervals.
- ⁷ Downgraded one level due to risk of bias. One study in the pooled analysis due was single blind and another was open label.

Summary of findings 5. High dose oral 5-ASA versus low dose 5-ASA for maintenance of remission in ulcerative colitis

High dose oral 5-ASA versus low dose 5-ASA for maintenance of remission in ulcerative colitis

Patient or population: Patients with guiescent ulcerative colitis

Settings: Outpatients

Intervention: High dose oral 5-ASA versus low dose 5-ASA

Outcomes	Illustrative comparative risks* (95% CI)		Relative effect (95% CI)	No of Partici- pants	Quality of the evidence	Comments
	Assumed risk	Corresponding risk	(33 /0 Ci)	(studies)	(GRADE)	
	Control	High dose 5-ASA versus low dose 5-ASA				
Failure to maintain clinical or endo- scopic remission at 12 months	357 per 1000 ¹	286 per 1000 (164 to 493)	RR 0.80 (0.46 to 1.38)	112 (1 study)	⊕⊝⊝⊝ very low ^{2,3}	Asacol 4.8 g/day versus 2.4 g/day
Failure to maintain clinical or endo- scopic remission at 12 months	330 per 1000 ¹	251 per 1000 (69 to 921)	RR 0.76 (0.21 to 2.79)	216 (2 studies)	⊕⊝⊝⊝ very low ^{34,5}	Balsalazide 6.0 g/day versus 3.0 g/day
Failure to maintain clinical or endo- scopic remission at 12 months	554 per 1000 ¹	366 per 1000 (249 to 537)	RR 0.66 (0.45 to 0.97)	133 (1 study)	⊕⊕⊕⊝ moderate ⁶	Balsalazide 4.0 g/day versus 2.0 g/day
Failure to maintain clinical or endo- scopic remission at 12 months	392 per 1000 ¹	255 per 1000 (192 to 337)	RR 0.65 (0.49 to 0.86)	429 (1 study)	⊕⊕⊕⊝ moderate ⁷	Salofalk granules 3.0 g OD versus 1.5 g OD

^{*}The basis for the **assumed risk** (e.g. the median control group risk across studies) is provided in footnotes. The **corresponding risk** (and its 95% confidence interval) is based on the assumed risk in the comparison group and the **relative effect** of the intervention (and its 95% CI).

CI: Confidence interval; RR: risk ratio; OD: once daily

GRADE Working Group grades of evidence

High quality: Further research is very unlikely to change our confidence in the estimate of effect.

Moderate quality: Further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.

Low quality: Further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.

- ¹ Control group risk estimates come from control arm of meta-analysis, based on included trials.
- ² Downgraded two levels due to very sparse data (36 events).
- ³ Downgraded one due to high risk of bias (open label study).
- ⁴ Downgraded one level due to sparse data (61 events) and very wide confidence intervals.
- ⁵ Downgraded two levels due to very serious inconsistency ($I^2 = 86\%$).
- ⁶ Downgraded one level due to sparse data (61 events) and wide confidence intervals.
- ⁷ Downgraded one level due to sparse data (138 events).



BACKGROUND

The successful management of ulcerative colitis was greatly facilitated after the introduction of sulfasalazine (SASP) by Svartz (Svartz 1942). SASP is composed of 5-aminosalicylic acid (5-ASA) linked to sulfapyridine via a diazo bond. This bond is readily cleaved by bacterial azoreductases in the colon (Peppercorn 1972) to yield the two components. Of these, 5-ASA has been found to be the therapeutically active component, while sulfapyridine, which is primarily absorbed into systemic circulation, is assumed to function solely as a carrier molecule (Azad Khan 1977; Van Hees 1980; Klotz 1980).

Administration of unbound or uncoated 5-ASA revealed that it was readily absorbed in the upper jejunum and was unable to reach the colon in therapeutic concentrations (Schroeder 1972; Nielsen 1983; Myers 1987). Ingested SASP largely resists such premature absorption and thus is able to serve as a delivery system that transports the 5-ASA to the affected regions of the lower intestinal tract (Schroeder 1972). While corticosteroid therapy is more effective in the treatment of severe ulcerative colitis (Truelove 1955; Truelove 1959) the use of SASP in maintaining remission (Misiewitz 1965) has been well established.

Despite its benefits, up to 30% of patients receiving SASP have reported adverse side-effects (Nielsen 1982). It was concluded that many were due to the sulfapyridine moiety, especially those effects found to be dose-dependent (Das 1973; Myers 1987). This discovery spawned more than a decade of research aimed at finding alternative 5-ASA delivery systems.

Asacol® (Proctor and Gamble) consists of a pellet of 5-ASA destined for release in the terminal ileum or colon due to a coating known as Eudragit-S, a resin that dissolves at a pH greater than 7 (Dew 1982a). Claversal® or Mesasal® (GlaxoSmithKline), Salofalk® (Axcan Pharma, Falk Foundation), and Rowasa® (Reid-Rowell) are similar delayed-release preparations of 5-ASA pellets coated with Eudragit L, a resin that dissolves at a pH greater than 6 (the approximate pH of the ileum/colon) (Hardy 1987; Myers 1987). Pentasa® (Marion-Merrell-Dow) is a microsphere formulation that consists of 5-ASA microgranules enclosed within a semi-permeable membrane of ethylcellulose. It is designed for controlled release that begins in the duodenum and continues into the affected regions of the lower bowel (Rasmussen 1982). Olsalazine or Dipentum® (Pharmacia & Upjohn) consists of two 5-ASA molecules linked by a diazo bond (Willoughby 1982; Staerk Laursen 1990). Other formulations, such as benzalazine and balsalazide, are composed of 5-ASA molecules azo-bonded to various benzoic acid derivatives (Chan 1983; Fleig 1988). Like SASP, these compounds are poorly absorbed in the upper digestive tract but are readily metabolized by the intestinal flora in the lower bowel. MMX mesalamine (Lialdaa® or Mezavanta®) uses MMX Multi Matrix System (MMX) technology to delay and extend delivery of active drug throughout the colon (Kamm 2008).

The newer 5-ASA preparations were intended to avoid the adverse effects of SASP while maintaining its therapeutic benefits. These drugs are more costly, however, and have still been shown to cause adverse effects in some patients (Rao 1987). The efficacy and safety of the 5-ASA preparations have been evaluated in numerous clinical trials that have often lacked sufficient statistical power to arrive at definitive conclusions. In an earlier meta-analysis, Sutherland 1993 found that the newer 5-ASA drugs were no more effective than SASP for maintenance of remission in ulcerative colitis. This

systematic review is an update of the Cochrane review published in 2012 (Feagan 2012). We proceeded with this updated review, in accordance with the format of the Cochrane Collaboration, in order to include the more recent studies as well as to evaluate the effectiveness, dose-responsiveness, and safety of the 5-ASA preparations in terms of more precise outcome measures.

Many patients are non-adherent with conventional multi-dose (2 or 3 times daily) treatment regimens which may result in reduced efficacy, and can lead to an increased risk of relapse in patients with quiescent disease (Kane 2001; Kane 2003b), poor long-term prognosis (Kane 2008b) and increased costs of care (Kane 2008b; Beaulieu 2009). Poor adherence may be particularly problematic in quiescent disease (Kane 2001; Kane 2003b), since patients lack continuing symptoms that incentivize them to take medication. Although multiple factors have been shown to influence medication adherence in patients with ulcerative colitis it is commonly believed that a high pill burden and multi-dose regimens are major determinants (Ediger 2007; Kane 2008b). Accordingly, it is reasonable to hypothesize that once daily dosing of mesalamine might improve both adherence with maintenance therapy and outcomes.

The efficacy and safety of once daily oral dosing of mesalamine compared to conventional dosing (two or three times daily) for the treatment of ulcerative colitis has been evaluated in numerous clinical trials. These trials have investigated the efficacy of once daily dosing of various formulations of mesalamine compared to conventional dosing schedules of the same drugs or different formulations. Many of these trials were small in size and lacked sufficient statistical power to arrive at definitive conclusions. A secondary objective of this systematic review was to investigate the efficacy and safety of once daily dosing of mesalamine compared to conventional dosing for the treatment quiescent ulcerative colitis.

OBJECTIVES

The primary objectives were to assess the efficacy, dose-responsiveness, and safety of oral 5-aminosalicylic acid (5-ASA) compared to placebo, sulfasalazine (SASP), or 5-ASA comparators (i.e. other formulations of 5-ASA) for maintenance of remission in quiescent ulcerative colitis. A secondary objective of this systematic review was to compare the efficacy and safety of once daily dosing of oral 5-ASA with conventional dosing regimens.

METHODS

Criteria for considering studies for this review

Types of studies

Prospective, randomized controlled trials of parallel design, with a minimum treatment duration of six months were considered for inclusion.

Types of participants

Patients of any age with mild-to-moderate ulcerative colitis in remission as defined by Truelove and Witts (Truelove 1955) were considered for inclusion.

Types of interventions

Trials of oral 5-ASA therapy for treatment of patients with ulcerative colitis in remission compared with placebo, SASP or



other formulations of 5-ASA were considered for inclusion. Studies that compared once daily 5-ASA treatment with conventional dosing of 5-ASA (two or three times daily) and 5-ASA dose ranging studies were also considered for inclusion.

Types of outcome measures

Outcome measures included endoscopic or clinical relapse, or early withdrawal, as defined by the authors of each study.

Primary outcomes

The primary outcome was endoscopic or clinical relapse as defined by the authors of each study.

Secondary outcomes

Secondary outcomes included:

- the proportion of patients who failed to adhere with their medication regimen;
- the proportion of patients who experienced at least one adverse event;
- the proportion of patients who withdrew due to adverse events;
- the proportion of patients excluded or withdrawn after entry.

Search methods for identification of studies

MEDLINE (OvidSP), EMBASE (Ovid SP), and the Cochrane Library were searched from inception to January 20, 2012. No language or document type restrictions were applied. The multipurpose search command for the Ovid SP interface (.mp.) was used to search both text and database subject heading fields. Review articles and conference proceedings were also searched to identify additional studies. The search strategies are listed in Appendix 1.

Data collection and analysis

Study Selection

Relevant studies were selected on the basis of the above criteria. When necessary, the original authors were contacted to clarify points regarding trial methodology. The reasons for exclusion are indicated for each ineligible study.

Data Collection

Each study selected for analysis was independently reviewed by two authors. Data were recorded onto standard data extraction forms by each author. Any discrepancies between authors were settled by consensus. Results were recorded on an intention-totreat basis, regardless of whether the original authors had done so.

Outcome Measures

The primary outcome of interest was the failure to maintain clinical or endoscopic remission as evidenced by disease relapse, the criteria for which were defined by the authors of each trial. Data were also extracted, where possible, to investigate the influence of the dose of 5-ASA. As well, the numbers of patients who experienced adverse effects, withdrawal from the study due to adverse events, and exclusion or withdrawal from therapy were recorded where the data were available.

Risk of bias assessment

Two authors independently assessed the risk of bias in the included studies using the Cochrane risk of bias tool (Higgins 2011). Factors assessed included:

- sequence generation (i.e. was the allocation sequence adequately generated?);
- allocation sequence concealment (i.e. was allocation adequately concealed?);
- blinding (i.e. was knowledge of the allocated intervention adequately prevented during the study?);
- 4. incomplete outcome data (i.e. were incomplete outcome data adequately addressed?);
- 5. selective outcome reporting (i.e. are reports of the study free of suggestion of selective outcome reporting?); and
- 6. other potential sources of bias (i.e. was the study apparently free of other problems that could put it at a high risk of bias?).

A judgement of 'Yes' indicates low risk of bias, 'No' indicates high risk of bias, and 'Unclear' indicates unclear or unknown risk of bias. Disagreements were resolved by consensus. Study authors were contacted when insufficient information was provided to determine risk of bias.

We used the GRADE approach for rating the overall quality of evidence for primary outcomes and selected secondary outcomes of interest. Randomized trials start as high quality evidence, but may be downgraded due to: (1) limitations in design and implementation (risk of bias), (2) indirectness of evidence, (3) inconsistency (unexplained heterogeneity), (4) imprecision (sparse data), and (5) reporting bias (publication bias). The overall quality of evidence for each outcome was determined after considering each of these elements, and categorized as high quality (i.e. further research is very unlikely to change our confidence in the estimate of effect); moderate quality (i.e. further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate); low quality (i.e. further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate); and very low quality (i.e. we are very uncertain about the estimate) (Guyatt 2008; Schünemann 2011).

Statistical Methods

Trials were separated into five comparison groups: 5-ASA versus placebo, 5-ASA versus sulfasalazine, once daily dosing versus conventional dosing, 5-ASA versus comparator 5-ASA formulation, and 5-ASA dose-ranging. Within each group, raw data for every measured outcome were extracted and converted into individual two by two tables. The tables for placebo-controlled trials were further subgrouped according to the dose of 5-ASA. The tables for the once daily versus conventional dosing studies were subgrouped by formulation. The tables for 5-ASA-controlled trials were subgrouped by common 5-ASA comparators (e.g. Asacol and Salofalk). The tables for dose-ranging studies were subgrouped by 5-ASA formulation. The risk ratio (RR) and 95% confidence intervals (95% CI) derived from each two by two table were individually calculated and plotted. The results for each comparison group were pooled to determine the RR and 95% CI for each outcome resulting from 5-ASA therapy relative to either placebo, SASP or 5-ASA comparator formulation and once daily 5-ASA therapy relative to conventional dosing. A fixed-effect model was used. Studies were pooled for analysis if patients, outcomes and interventions



were similar (determined by consensus among authors). Studies comparing 5-ASA to comparator 5-ASA formulations were pooled for analysis if they compared equimolar doses of oral 5-ASA.

Dose-responsiveness was analyzed using a Chi² for trend. Trials were also subgrouped according to the specific 5-ASA preparation for those outcomes for which there were two or more studies that used a similar drug. Tests for homogeneity among trials within each comparison group were performed. The presence of heterogeneity among studies was assessed using the Chi² test (a P value of 0.10 was regarded as statistically significant) and the I² statistic (Higgins 2003). If statistically significant heterogeneity was identified, the RR and 95% CI were calculated using a random-effects model. We conducted sensitivity analyses as appropriate to investigate heterogeneity. We also conducted sensitivity analyses excluding studies with a high risk of bias. All statistical analyses were performed using the Cochrane Collaboration RevMan 5 software package.

RESULTS

Description of studies

A literature search conducted on July 9, 2015 identified 2525 studies. Eleven additional studies were identified through searching of references. After duplicates were removed a total of 1619 reports remained for review of titles and abstracts. Two authors independently reviewed the titles and abstracts of these studies and 112 reports of oral 5-ASA maintenance treatment for quiescent ulcerative colitis were selected for full text review (See Figure 1). Thirty reports of 28 studies were excluded (See Characteristics of excluded studies). Eighty-two reports of 41 studies involving a total of 8928 patients were selected for inclusion (Dew 1983a; Sandberg-Gertzen 1986; Andreoli 1987; Ireland 1988; McIntyre 1988; Mulder 1988; Riley 1988; Rutgeerts 1989; Courtney 1992; Giaffer 1992a; Green 1992; Kiilerich 1992; Rijk 1992; Wright 1993; Travis 1994; Ardizzone 1995; Fockens 1995; Kruis 1995; Miner 1995; Nilsson 1995; Hanauer 1996; Hawkey 1997; Green 1998; Ardizzone 1999; Deventer 2001; Kruis 2001; Mahmud 2002; Kane 2003a; Paoluzi 2005; Kamm 2008; Kane 2008a; Dignass 2009; Prantera 2009; Ito 2010; Lichtenstein 2010; Sandborn 2010; Kruis 2011; D'Haens 2012; Hawthorne 2012; Pica 2012; Watanabe 2013) (See Characteristics of included studies).



Figure 1. Study flow diagram.

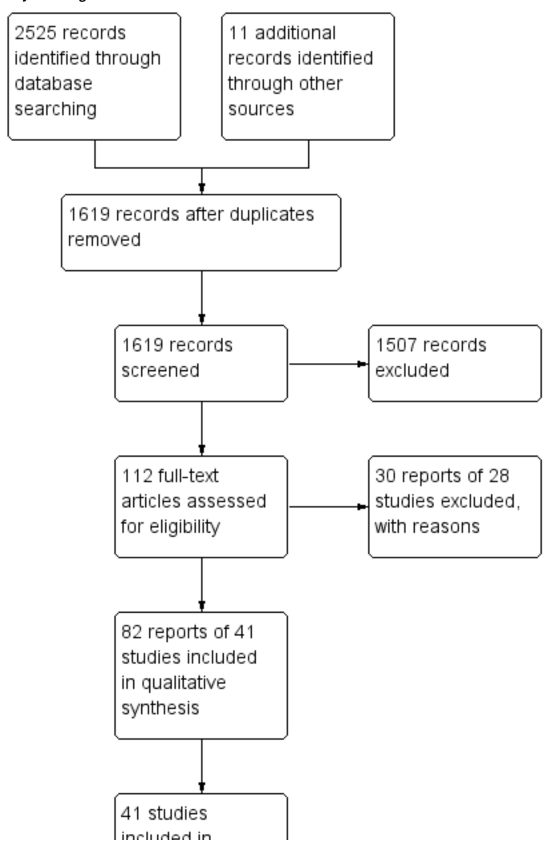




Figure 1. (Continued)

included in quantitative synthesis (meta-analysis)

Seven studies were placebo-controlled (Sandberg-Gertzen 1986; Wright 1993; Miner 1995; Hanauer 1996; Hawkey 1997; Ardizzone 1999; Lichtenstein 2010). Twelve studies compared 5-ASA to SASP (Dew 1983a; Andreoli 1987; Ireland 1988; McIntyre 1988; Mulder 1988; Riley 1988; Rutgeerts 1989; Kiilerich 1992; Rijk 1992; Ardizzone 1995; Kruis 1995; Nilsson 1995). Ten studies were maintenance of remission studies comparing once daily dosing of mesalamine with conventional dosing (Kane 2003a; Kamm 2008; Kane 2008a; Dignass 2009; Prantera 2009; Sandborn 2010; Hawthorne 2012; Kruis 2011; D'Haens 2012; Watanabe 2013). Six studies compared the efficacy and safety of various formulations of oral 5-ASA to other formulations of oral 5-ASA for maintenance treatment (Courtney 1992; Green 1998; Deventer 2001; Kruis 2001; Mahmud 2002; Ito 2010). Ten trials were dose-ranging studies of oral 5-ASA (Giaffer 1992a; Green 1992; Travis 1994; Fockens 1995; Hanauer 1996; Deventer 2001; Kruis 2001; Paoluzi 2005; Kruis 2011; Pica 2012). Six of the included studies were formal non-inferiority studies (Dignass 2009; Ito 2010; Sandborn 2010; Hawthorne 2012; D'Haens 2012; Watanabe 2013).

Risk of bias in included studies

A summary of the risk of bias assessment is provided in Figure 2. Most of the included studies were of high methodological quality. Ten studies were rated as high risk of bias. Seven of these studies were single-blind (Courtney 1992; Deventer 2001; Kane 2003a; Kane 2008a; Dignass 2009; Sandborn 2010; Hawthorne 2012). Outcomes were assessed by a blinded investigator in these studies. Three

studies were open-label and investigators and patients were not blinded to treatment assignment (Mahmud 2002; Kamm 2008; Pica 2012). However, two of the open-label studies (Mahmud 2002; Kamm 2008) and 4 of 7 single-blind studies (Courtney 1992; Kane 2008a; Dignass 2009; Hawthorne 2012) utilized investigator performed endoscopy as an endpoint, which may protect against bias provided the endoscopist is blinded. The methods used for blinding were not described in one study and this study was rated as unclear (Green 1992). Twenty-nine of 41 included studies did not describe the method used for randomization and were rated as unclear for this item (Dew 1983a; Sandberg-Gertzen 1986; Ireland 1988; McIntyre 1988; Mulder 1988; Riley 1988; Rutgeerts 1989; Giaffer 1992a; Green 1992; Rijk 1992; Wright 1993; Travis 1994; Ardizzone 1995; Miner 1995; Nilsson 1995; Hawkey 1997; Green 1998; Ardizzone 1999; Deventer 2001; Kruis 2001; Paoluzi 2005; Kamm 2008; Dignass 2009; Lichtenstein 2010; Sandborn 2010; Hawthorne 2012; D'Haens 2012; Pica 2012; Watanabe 2013). Eighteen studies did not describe methods used for allocation concealment and were rated as unclear for this item (Dew 1983a; Andreoli 1987; Courtney 1992; Giaffer 1992a; Green 1992; Travis 1994; Ardizzone 1995; Fockens 1995; Green 1998; Ardizzone 1999; Deventer 2001; Kruis 2001; Mahmud 2002; Paoluzi 2005; Lichtenstein 2010; Hawthorne 2012; Pica 2012; Watanabe 2013). Thirteen studies were rated as unclear for incomplete outcome data because reasons for withdrawal were not described (Dew 1983a; Sandberg-Gertzen 1986; Andreoli 1987; Rutgeerts 1989; Rijk 1992; Travis 1994; Miner 1995; Green 1998; Hanauer 1996; Deventer 2001; Kruis 2001; Hawthorne 2012; Pica 2012).



Figure 2. Risk of bias summary: review authors' judgements about each risk of bias item for each included study.

	Random sequence generation (selection bias)	Allocation concealment (selection bias)	Blinding (performance bias and detection bias)	Incomplete outcome data (attrition bias)	Selective reporting (reporting bias)	Other bias
Andreoli 1987	•	?	•	?	?	?
Ardizzone 1995	?	?	•	•	•	•
Ardizzone 1999	?	?	•	•	•	•
Courtney 1992	•	?		•	•	•
D'Haens 2012	?	•	•	•	•	•
Deventer 2001	?	?	•	?	?	?
Dew 1983a	?	?	•	?	•	•
Dignass 2009	?	•	•	•	•	•
Dignass 2009 Fockens 1995	?	?	•	•	•	•
	H	_	•	_	_	
Fockens 1995	•	?	_	•	•	•

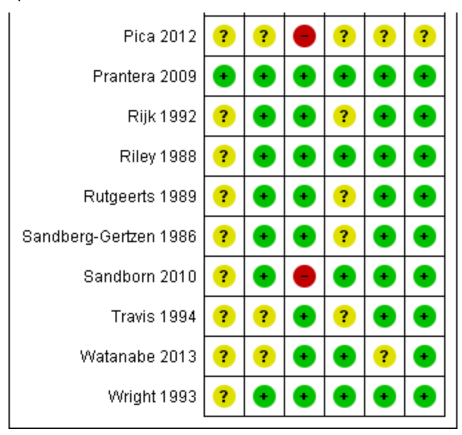


Figure 2. (Continued)

Green 1998	?	?	•	?	•	•
Hanauer 1996	•	•	•	?	•	•
Hawkey 1997	?	•	•	•	•	•
Hawthorne 2012	?	?		?	•	•
Ireland 1988	?	•	•	•	•	•
Ito 2010	•	•	•	•	•	•
Kamm 2008	?	•		•	•	•
Kane 2003a	•	•		•	•	?
Kane 2008a	•	•		•	•	•
Kiilerich 1992	•	•	•	•	•	•
Kruis 1995	•	•	•	•	•	•
Kruis 2001	?	?	•	?	•	•
Kruis 2011	•	•	•	•	•	•
Lichtenstein 2010	?	?	•	•	•	•
Mahmud 2002	•	?		•	•	•
McIntyre 1988	?	•	•	•	•	•
Miner 1995	?	•	•	?	•	•
Mulder 1988	?	•	•	•	•	•
Nilsson 1995	?	•	•	•	•	•
Paoluzi 2005	?	?	•	•	•	•
Pica 2012	?	?		?	?	2



Figure 2. (Continued)



Effects of interventions

See: Summary of findings for the main comparison Oral 5-ASA versus placebo for maintenance of remission in ulcerative colitis; Summary of findings 2 Oral 5-ASA versus SASP for maintenance of remission in ulcerative colitis; Summary of findings 3 Once daily dosing versus conventional dosing for maintenance of remission in ulcerative colitis; Summary of findings 4 Oral 5-ASA versus comparator 5-ASA formulation for maintenance of remission in ulcerative colitis; Summary of findings 5 High dose oral 5-ASA versus low dose 5-ASA for maintenance of remission in ulcerative colitis

EFFICACY

5-ASA versus Placebo

Seven trials (n = 1298 patients) reported treatment outcomes in terms of failure to maintain clinical or endoscopic remission (Sandberg-Gertzen 1986; Wright 1993; Miner 1995; Hanauer 1996; Hawkey 1997; Ardizzone 1999; Lichtenstein 2010). Forty-one per cent of 5-ASA patients relapsed compared to 58% of placebo patients. The pooled RR of failure to maintain clinical or endoscopic remission for all trials was 0.69 (95% CI 0.62 to 0.77; I² = 15%; P < 0.00001) using a fixed-effect model. There was a trend towards greater efficacy with higher doses of 5-ASA with a statistically significant benefit for the 1 to 1.9 g/day (RR 0.65; 95% CI 0.56 to 0.76; I² = 0%; P < 0.00001) and the \geq 2 g/day subgroups (RR 0.73; 95% CI 0.60 to 0.89; I² = 71%; P = 0.002). The GRADE analysis indicated that the overall quality of the evidence for the primary outcome

for the placebo-controlled studies (failure to maintain clinical or endoscopic remission) was high (See Summary of findings for the main comparison). The pooled RR was similar (RR 0.69; 95% CI 0.57 to 0.84) when calculated exclusively with those trials with endpoints at 12 months (Wright 1993; Miner 1995; Ardizzone 1999). Two of the trials involving olsalazine (Sandberg-Gertzen 1986; Wright 1993) had a pooled RR of 0.76 (95% CI 0.58 to 0.99). Two trials involving Asacol had a pooled RR of 0.73 (95% CI 0.60 to 0.88).

5-ASA versus Sulfasalazine

Twelve trials involving a total of 1655 patients compared the efficacy of 5-ASA and SASP (Dew 1983a; Andreoli 1987; Ireland 1988; McIntyre 1988; Mulder 1988; Riley 1988; Rutgeerts 1989; Kiilerich 1992; Rijk 1992; Ardizzone 1995; Kruis 1995; Nilsson 1995). In 8 of the 11 studies the dose of SASP was limited to 2 g/day; in one trial the dose of SASP was 4 g/day; in one trial the mean dose of SASP was 2.7 g/day and ranged from 2.4 to 4.4 g/day (see Table of Included Studies). When the outcome of interest was defined as the failure to maintain clinical or endoscopic remission (withdrawals and relapses), SASP was significantly superior to 5-ASA. Forty-eight per cent of 5-ASA patients relapsed compared to 43% of SASP patients (RR 1.14, 95% CI 1.03 to 1.27; $I^2 = 17\% P = 0.01$). The GRADE analysis indicated that the overall quality of the evidence for the primary outcome for the SASP-controlled studies (failure to maintain clinical or endoscopic remission) was high (See Summary of findings 2). The NNT value was found to be negative (-17), indicating that SASP has a certain degree of therapeutic superiority over the 5-ASA preparations. When the analysis was limited to those studies with endpoints at 12 months (Andreoli 1987; Mulder 1988;



Riley 1988; Rutgeerts 1989; Kiilerich 1992; Rijk 1992; Nilsson 1995; Ardizzone 1995) there was no statistically significant difference between 5-ASA and SASP (RR 1.10; 95% CI 0.98 to 1.23). Similarly, when the analysis was limited to studies that did not use olsalazine (Dew 1983a; Andreoli 1987; McIntyre 1988; Mulder 1988; Riley 1988; Rutgeerts 1989; Ardizzone 1995) there was no statistically significant difference between 5-ASA and SASP (RR 1.08; 95% CI 0.92 to 1.26).

Three trials involving Claversal® (Andreoli 1987; Ardizzone 1995; Rutgeerts 1989) had a pooled RR of 1.15 (95% CI 0.95 to 1.40). When the five trials involving olsalazine (Ireland 1988; Kiilerich 1992; Kruis 1995; Nilsson 1995; Rijk 1992) were pooled, the resulting odds ratio was 1.20 (95% CI 1.04 to 1.38), thus demonstrating that SASP was significantly better than olsalazine for maintenance of remission.

Once Daily Dosing versus Conventional Dosing

Three trials (n = 1871 patients) reported treatment outcomes in terms of failure to maintain clinical or endoscopic remission at six months (Kane 2003a; Sandborn 2010; D'Haens 2012). There was no statistically significant difference in relapse rates at six months. Nineteen per cent of once daily patients relapsed compared to 18% of conventionally dosed patients (RR 1.02; 95% CI 0.85 to 1.23). No statistically significant heterogeneity was detected for this comparison (P = 0.76, I^2 = 0%). None of the subgroup comparisons by formulation showed any differences in efficacy between once daily dosing and conventional dosing. However, only two formulations were evaluated in this pooled analysis. The GRADE analysis indicated that the overall quality of the evidence for the primary outcome (failure to maintain clinical or endoscopic remission at six months) for the studies comparing once daily with conventional dosing was low due to sparse data and a high risk of bias (single blind) in some studies in the pooled analysis (See Summary of findings 3).

Eight trials (n = 3127 patients) reported treatment outcomes in terms of failure to maintain clinical or endoscopic remission at 12 months (Kane 2008a; Kamm 2008; Dignass 2009; Prantera 2009; Sandborn 2010; Hawthorne 2012; Kruis 2011; Watanabe 2013). There was no statistically significant difference in relapse rates at 12 months. Twenty-nine per cent of once daily patients relapsed compared to 31% of conventionally dosed patients (RR 0.91; 95% CI 0.82 to 1.01). No statistically significant heterogeneity was detected for this comparison (P = 0.26, I^2 = 22%). The subgroup comparison for Pentasa showed a statistically significant difference in favour of once daily dosing compared to conventional twice daily dosing (RR 0.75; 95% CI 0.60 to 0.93). None of the other subgroup comparisons (by formulation) showed any differences in efficacy between once daily dosing and conventional dosing. The GRADE analysis indicated that the overall quality of the evidence for the primary outcome (failure to maintain clinical or endoscopic remission at 12 months) for the studies comparing once daily with conventional dosing was moderate due to a high risk of bias in some studies (open label and single blind) in the pooled analysis (See Summary of findings 3).

Eight trials (n = 2126) reported adherence with study medication at study endpoint (Kane 2003a; Kamm 2008; Kane 2008a; Dignass 2009; Prantera 2009; Kruis 2011; Hawthorne 2012; Watanabe 2013). Overall, 9.5% (101/1061) of patients in the once daily group failed to adhere to their medication regimen compared to 7.5% (80/1065) of patients in the conventional dosing group. The pooled RR using

a random-effects model was 1.18 (95% CI 0.69 to 2.03) showing no statistically significant difference in medication adherence between once daily dosing and conventional dosing at study endpoint (6 months for Kane 2003a and 12 months for the other studies in the pooled analysis; P = 0.55). Statistically significant heterogeneity was detected for this comparison (P = 0.009, I^2 = 63%). The heterogeneity appears to be a result of the inclusion of two specific trials (Kamm 2008; Hawthorne 2012). Kamm 2008 reported a significantly higher compliance rate of 99.6% in the twice daily dosing group compared to 93.3% in the once daily group. Hawthorne 2012 reported a significantly higher compliance rate of 97.1% in the once daily dosing group compared to 85.5% in the three times daily dosing group. To investigate if these studies were the source of the heterogeneity the analysis was repeated excluding these trials. The pooled analysis of the ITT population now included 6 studies and 1462 patients (Kane 2003a; Kane 2008a; Dignass 2009; Prantera 2009; Kruis 2011; Watanabe 2013). Overall, 11.2% (83/739) of patients in the once daily group failed to adhere to their medication regimen compared to 8.7% (63/723) of patients in the conventional dosing group. The pooled RR using a fixedeffect model was 1.22 (95% CI 0.91 to 1.64) showing no statistically significant difference in medication adherence between once daily dosing and conventional dosing at study endpoint (P = 0.18). No statistically significant heterogeneity was detected for this comparison (P = 0.55; $I^2 = 0\%$). The GRADE analysis indicated that the overall quality of the evidence for the adherence outcome was moderate due to sparse data (See Summary of findings 3).

5-ASA versus Comparator 5-ASA Formulation

Six studies (n = 707 patients) reported treatment outcomes in terms of the failure to maintain clinical or endoscopic remission at 12 months (Courtney 1992; Green 1998; Deventer 2001; Kruis 2001; Mahmud 2002; Ito 2010). The overall pooled RR showed no statistically significant difference in relapse between various formulations of 5-ASA (including Balsalazide, Pentasa and Olsalazine) and comparator formulations of 5-ASA (including Asacol and Salofalk). Forty-four per cent of patients in the 5-ASA group relapsed compared to 41% of patients in the 5-ASA comparator group. The pooled RR of relapse was 1.08 (95% CI 0.91 to 1.28; $I^2 = 31\%$; P = 0.37) using a fixed-effect model. The GRADE analysis indicated that the overall quality of the evidence for the primary outcome (failure to maintain clinical or endoscopic remission) was low due to sparse data (300 events) and a high risk of bias (lack of blinding) in two studies in the pooled analysis (See Summary of findings 4).

5-ASA Dose Ranging

Several randomized trials have looked at dose-ranging for various formulations of 5-ASA including Asacol, Balsalazide, Olsalazine, Salofalk, and Pentasa. Four studies examined the efficacy of various doses of Asacol for maintenance of clinical or endoscopic remission (Hanauer 1996; Deventer 2001; Paoluzi 2005; Pica 2012). Pica 2012 found no statistically significant difference in efficacy between Asacol 4.8 g/day compared to 2.4 g/day. Twenty-nine per cent of patients in the Asacol 4.8 g/day group relapsed compared to 36% in the 2.4 g/day group (112 patients; RR 0.80; 95% CI 0.46 to 1.38). A GRADE analysis indicated that the overall quality of the evidence for the primary outcome was very low due to very sparse data and risk of bias. Paoluzi 2005 found no statistically significant difference in efficacy between Asacol 2.4 g/day compared to 1.2 g/day.



Seventy per cent of patients in the Asacol 2.4 g/day group relapsed compared to 74% in the 1.2 g/day group (156 patients; RR 0.95; 95% CI 0.78 to 1.16). Deventer 2001 found no statistically significant difference in efficacy between Asacol 3.2 g/day compared to 2 g/day. Fifty-one per cent of patients in the Asacol 3.2 g/day group relapsed compared to 48% of patients in the 2 g/day group (262 patients; RR 1.07; 95% CI 0.83 to 1.37). Hanauer 1996 found no statistically significant difference in efficacy between Asacol 1.6 g/day compared to 0.8 g/day. Fifty-six per cent of patients relapsed in both dose groups (177 patients; RR 1.01; 95% CI 0.78 to 1.32).

Three studies examined the efficacy of various doses of Balsalazide for maintenance of clinical or endoscopic remission (Giaffer 1992a; Green 1992; Kruis 2001). Two of these studies compared Balsalazide 6 g/day to 3 g/day (Green 1992; Kruis 2001). The pooled analysis showed no statistically significant difference in efficacy between Balsalazide 6 g/day and 3 g/day. Twenty-three per cent of patients in the 6 g/day group relapsed (216 patients; RR 0.72; 95% CI 0.46 to 1.13). However, these results should be interpreted with caution as statistically significant heterogeneity was detected for this comparison (P = 0.007; I^2 = 86%). A GRADE analysis indicated that the overall quality of the evidence for the primary outcome was very low due to sparse data, very wide confidence intervals and inconsistency. Giaffer 1992a compared Balsalazide 4 g/day to 2 g/ day and found a statistically significant difference favouring the 4 g/day dose group. Thirty-seven per cent of patients in the 4 g/day Balsalazide group relapsed compared to 55% of patients in the 2 g/day group (133 patients; RR 0.66; 95% CI 0.45 to 0.97). A GRADE analysis indicated that the overall quality of the evidence for the primary outcome was moderate due to sparse data.

Travis 1994 found no statistically significant difference in efficacy between Olsalazine 2 g/day and 1 g/day. Forty per cent of patients in both dose groups relapsed (127 patients; RR 1.01; 95% CI 0,66 to 1.54). Kruis 2011 found a statistically significant difference between Salofalk granules 3 g/day and 1.5 g/day. Twenty-five per cent of patients in the Salofalk 3 g/day group relapsed compared to 39% of patients in the 1.5 g/day group (429 patients; RR 0.65; 95% CI 0.49 to 0.86). Fockens 1995 found no statistically significant difference in efficacy between Pentasa 3.0 g/day and 1.5 g/day. Twenty-eight per cent of patients in the 3.0 g/day group relapsed compared to 38% in the 1.5 g/day group (169 patients; RR 0.74; 95% CI 0.48 to 1.15). A GRADE analysis indicated that the overall quality of the evidence for the primary outcome was moderate due to sparse data.

SAFETY

Three different outcome measures were used to evaluate the safety and clinical utility of 5-ASA relative to placebo and SASP: the number of patients with adverse events, the number of patients withdrawn due to adverse events, and the total number of patients excluded or withdrawn before completion of the study. Since many studies only reported the total number of adverse events rather than the number of patients who experienced an event, we were often unable to include such data in the analysis.

5-ASA versus Placebo

Four studies (n = 962 patients) reported the proportion of patients who experienced at least one adverse event (Wright 1993; Miner 1995; Hanauer 1996; Lichtenstein 2010). There was no statistically significant difference in the incidence of adverse events between 5-ASA and placebo patients. Forty-one per cent of 5-ASA patients

experienced at least one adverse event compared to 34% of placebo patients (RR 0.98; 95% CI 0.69 to 1.39; P = 0.91). Statistically significant heterogeneity was detected for this comparison (P = 0.04; $I^2 = 60\%$). The heterogeneity appears to be a result of the inclusion of one specific trial (Wright 1993). Wright 1993 reported a significantly higher adverse event rate in the 5-ASA group compared to placebo. This was mostly due to a high rate of olsalazinerelated diarrhea in the 5-ASA group. To investigate if this study was the source of the heterogeneity the analysis was repeated excluding this trial. The pooled analysis of the ITT population now included 3 studies and 861 patients (Miner 1995; Hanauer 1996; Lichtenstein 2010). There was no statistically significant difference in the incidence of adverse events between 5-ASA and placebo patients. Overall, 42% of 5-ASA patients experienced at least one adverse event compared to 39% of placebo patients (RR 0.94; 95% CI 0.77 to 1.15). No statistically significant heterogeneity was detected for this comparison (P = 0.31; I^2 = 17%). The GRADE analysis indicated that the overall quality of the evidence for this outcome for the placebo-controlled studies (the proportion of patients who experienced at least one adverse event) was moderate low due to sparse data (See Summary of findings for the main comparison).

Six studies (n = 1197 patients) reported the proportion of patients withdrawn due to adverse events (Wright 1993; Miner 1995; Hanauer 1996; Hawkey 1997; Ardizzone 1999; Lichtenstein 2010). There was no statistically significant difference in withdrawal due to adverse events between 5-ASA and placebo patients. Withdrawals due to adverse events were reported for 5% of 5-ASA patients compared to 4% of placebo patients (RR 1.34; 95% CI 0.78 to 2.30). Statistically significant heterogeneity was detected for this comparison (P = 0.10; $I^2 = 44\%$). Again the heterogeneity appears to be a result of the inclusion of one specific trial (Wright 1993). To investigate if this study was the source of the heterogeneity the analysis was repeated excluding this trial. The pooled analysis of the ITT population now included 5 studies and 1096 patients (Miner 1995; Hanauer 1996; Hawkey 1997; Ardizzone 1999; Lichtenstein 2010). There was no statistically significant difference in withdrawal due to adverse events between 5-ASA and placebo patients. Overall, 3% of 5-ASA patients withdrew due to adverse events compared to 4% of placebo patients in the conventional dosing group (RR 0.86; 95% CI 0.46 to 1.63). No statistically significant heterogeneity was detected for this comparison (P = 0.48; I^2 = 0%). The GRADE analysis indicated that the overall quality of the evidence for this outcome for the placebo-controlled studies (the proportion of patients withdrawn due to adverse events) was moderate due to sparse data (See Summary of findings for the main comparison).

Five studies (n = 1175 patients) reported the proportion of patients excluded or withdrawn after entry (Wright 1993; Miner 1995; Hanauer 1996; Ardizzone 1999; Lichtenstein 2010). There was no statistically significant difference in the proportion of patients withdrawn or excluded after entry. Nineteen per cent of 5-ASA patients were withdrawn or excluded after entry compared to 18% of placebo patients (RR 1.13; 95% CI 0.88 to 1.44; I² = 11%).The GRADE analysis indicated that the overall quality of the evidence for this outcome for the placebo-controlled studies (the proportion of patients withdrawn or excluded after entry) was moderate due to sparse data (202 events; See Summary of findings table 1).



Commonly reported adverse events in the placebo-controlled trials included: headache (Miner 1995; Hanauer 1996; Hawkey 1997; Lichtenstein 2010), nausea (Miner 1995; Hawkey 1997; Lichtenstein 2010), abdominal pain (Miner 1995; Hanauer 1996; Ardizzone 1999; Lichtenstein 2010), dyspepsia (Miner 1995), bloating (Ardizzone 1999), flu syndrome (Hanauer 1996) rhinitis (Hanauer 1996) and nasopharyngitis (Lichtenstein 2010). Diarrhea was reported in one study involving olsalazine (Wright 1993), two Asacol studies (Hanauer 1996; Ardizzone 1999) and one study of Apriso (Lichtenstein 2010).

5-ASA versus Sulfasalazine

Seven studies (n = 1138 patients) reported the proportion of patients who experienced at least one adverse event (Andreoli 1987; Ireland 1988; McIntyre 1988; Mulder 1988; Rutgeerts 1989; Kruis 1995; Nilsson 1995). There was no statistically significant difference in the incidence of adverse events. Sixteen per cent of 5-ASA patients and SASP patients experienced at least one adverse event (RR 1.07; 95% CI 0.82 to 1.40). No statistically significant heterogeneity was detected for this comparison (P = 0.12; I² = 41%). A GRADE analysis indicated that the overall quality of the evidence for this outcome for the SASP-controlled studies (the proportion of patients who experienced at least one adverse event) was moderate low due to sparse data (See Summary of findings 2). Three olsalazine trials including 634 patients (Ireland 1988; Kruis 1995; Nilsson 1995) that were homogeneous (I² = 0%) had a pooled odds ratio of 1.27 (95% CI, 0.92 to 1.76).

Ten studies (n = 1585 patients) reported the proportion of patients withdrawn due to adverse events (Ireland 1988; McIntyre 1988; Mulder 1988; Riley 1988; Rutgeerts 1989; Kiilerich 1992; Rijk 1992; Ardizzone 1995; Kruis 1995; Nilsson 1995). There was no statistically significant difference in withdrawals due to adverse events. Seven per cent of 5-ASA patients were withdrawn due to adverse events compared to 5% of SASP patients (RR 1.27; 95% CI 0.87 to 1.87). No statistically significant heterogeneity was detected for this comparison (P = 0.72; $I^2 = 0\%$). In five olsalazine trials including 906 patients (Ireland 1988, Kruis 1995, Nilsson 1995, Kiilerich 1992, Rijk 1992), 9.2% of those receiving olsalazine and 6.2% of those receiving SASP were withdrawn because of adverse events (RR 1.61; 95% CI, 1.01 to 2.56). A GRADE analysis indicated that the overall quality of the evidence for this outcome for the SASP-controlled studies (the proportion of patients withdrawn due to adverse events) was moderate low due to sparse data (See Summary of findings 2). The results from two Claversal® trials including 422 patients (Ardizzone 1995; Rutgeerts 1989) were not statistically significant (RR 1.10; 95% CI, 0.48 to 2.54).

Nine studies involving 1497 patients (Ireland 1988; McIntyre 1988; Mulder 1988; Riley 1988; Rutgeerts 1989; Kiilerich 1992; Rijk 1992; Kruis 1995; Nilsson 1995) reported the proportion of patients excluded or withdrawn after entry (excluding relapses). Nineteen per cent of 5-ASA patients were excluded or withdrawn after entry compared to 15% of SASP patients (RR 1.30; 95% CI 1.04 to 1.63). No statistically significant heterogeneity was detected for this comparison (P = 0.18; I² = 29%). Withdrawals or exclusions after entry were significantly higher in five olsalazine trials (Ireland 1988; Kiilerich 1992; Rijk 1992; Kruis 1995; Nilsson 1995) involving 906 patients. Seventeen per cent of olsalazine patients were withdrawn or excluded after entry compared to 12% of SASP patients (RR 1.51; 95% CI 1.09 to 2.08).

Commonly reported adverse events in the SASP-controlled trials included: headache (McIntyre 1988; Riley 1988; Kruis 1995), anorexia or appetite loss (Riley 1988; Rutgeerts 1989) nausea (McIntyre 1988; Riley 1988; Rutgeerts 1989; Kiilerich 1992), vomiting (Riley 1988; Rutgeerts 1989; Nilsson 1995), abdominal pain (Ireland 1988; Rutgeerts 1989; Kiilerich 1992; Nilsson 1995), dyspepsia (Ireland 1988; Riley 1988; Kiilerich 1992; Rijk 1992), excessive flatus (McIntyre 1988), bloating (Rutgeerts 1989), urticaria (Kiilerich 1992; Ardizzone 1995) and rash (McIntyre 1988; Mulder 1988; Rijk 1992; Kruis 1995; Nilsson 1995). Diarrhea was reported in five studies involving Olsalazine (Ireland 1988; Kiilerich 1992; Rijk 1992; Kruis 1995; Nilsson 1995), and in two studies involving Claversal (Rutgeerts 1989; Ardizzone 1995).

Once Daily Dosing versus Conventional Dosing

Six studies (n = 2714 patients) reported the proportion of patients who experienced at least one adverse event (Kamm 2008; Dignass 2009; Prantera 2009; Kruis 2011; D'Haens 2012; Watanabe 2013). There was no statistically significant difference in the incidence of adverse events. Approximately 45% of once daily and conventionally dosed patients experienced at least one adverse event (RR 1.00; 95% CI 0.92 to 1.08). No statistically significant heterogeneity was detected for this comparison (P = 0.43; I² = 0%). Seven studies (n = 3737 patients) reported the proportion of patients who were withdrawn due to adverse events (Kamm 2008; Dignass 2009; Prantera 2009; Sandborn 2010; Kruis 2011; D'Haens 2012; Watanabe 2013). There was no statistically significant difference in withdrawal due to adverse events. Withdrawals due to adverse events were 1.9% (36/1858) in the once daily group compared to 1.5% (28/1879) in the conventionally dosed group (RR 1.31; 95% CI 0.80 to 2.13). No statistically significant heterogeneity was detected for this comparison (P = 0.42; $I^2 = 1\%$). Seven studies (n = 3737 patients) reported the proportion of patients who were excluded or withdrawn after entry (Kamm 2008; Dignass 2009; Prantera 2009; Sandborn 2010; Kruis 2011; D'Haens 2012; Watanabe 2013). There was no statistically significant difference in exclusions or withdrawals after entry. Approximately 15% of once daily and conventionally dosed patients were excluded or withdrawn after entry (RR 0.99; 95% CI 0.85 to 1.15). No statistically significant heterogeneity was detected for this comparison (P = 0.48; $I^2 = 0\%$).

The most common adverse events reported in the trials assessing once daily dosing included flatulence (Dignass 2009; Prantera 2009), dyspepsia (D'Haens 2012), abdominal pain (Kamm 2008; Dignass 2009; Prantera 2009; D'Haens 2012; Watanabe 2013), nausea (Prantera 2009), diarrhea (Dignass 2009; Prantera 2009; Watanabe 2013), headache (Kamm 2008; Prantera 2009; D'Haens 2012), nasopharyngitis (Kamm 2008; Dignass 2009; Watanabe 2013), inflammation of the upper respiratory tract (Watanabe 2013), gastroenteritis (Watanabe 2013), dental caries (Watanabe 2013), and worsening ulcerative colitis (Kamm 2008; Prantera 2009; Kruis 2011; D'Haens 2012).

5-ASA versus Comparator 5-ASA Formulation

Four studies (n = 357 patients) reported the proportion of patients who experienced at least one adverse event (Green 1998; Kruis 2001; Mahmud 2002; Ito 2010). There was no statistically significant difference in the incidence of adverse events between various formulations of 5-ASA (including Balsalazide, Pentasa and Olsalazine) and comparator formulations of 5-ASA (including



Asacol and Salofalk). Sixty-four per cent of patients in the 5-ASA group experienced at least one adverse event compared to 69% of patients in the 5-ASA comparator group (RR 0.94; 95% CI 0.83 to 1.07). No statistically significant heterogeneity was detected for this comparison (P = 0.35; $I^2 = 8\%$).

Five studies (n = 457 patients) reported the proportion of patients who were withdrawn due to adverse events (Courtney 1992; Green 1998; Kruis 2001; Mahmud 2002; Ito 2010). There was no statistically significant difference in withdrawal due to adverse events between various formulations of 5-ASA (including Balsalazide, Pentasa and Olsalazine) and comparator formulations of 5-ASA (including Asacol and Salofalk). Six per cent of patients in the 5-ASA group withdrew due to adverse events compared to 4% of patients in the 5-ASA comparator group (RR 1.25; 95% CI 0.56 to 2.78). No statistically significant heterogeneity was detected for this comparison (P = 0.76; I² = 0%).

Five studies (n = 457 patients) reported the proportion of patients who were excluded or withdrawn after entry (Courtney 1992; Green 1998; Kruis 2001; Mahmud 2002; Ito 2010). There was no statistically significant difference in exclusions or withdrawals after entry between various formulations of 5-ASA (including Balsalazide, Pentasa and Olsalazine) and comparator formulations of 5-ASA (including Asacol and Salofalk). Twenty-eight per cent of patients in the 5-ASA group were excluded or withdrawn after entry compared to 22% of patients in the 5-ASA comparator group (RR 1.23; 95% CI 0.90 to 1.70). No statistically significant heterogeneity was detected for this comparison (P = 0.52; $I^2 = 0\%$).

The most common adverse events reported in these trials included dyspepsia (Mahmud 2002), abdominal pain (Courtney 1992; Green 1998; Kruis 2001; Mahmud 2002), nausea (Courtney 1992; Kruis 2001; Mahmud 2002), distension (Mahmud 2002) diarrhea (Courtney 1992; Green 1998; Kruis 2001; Mahmud 2002; Ito 2010), headache (Green 1998; Kruis 2001), nasopharyngitis or respiratory infections (Green 1998; Ito 2010), flu-like disorder (Green 1998) and rash (Courtney 1992).

5-ASA Dose Ranging

Five dose-ranging studies reported the proportion of patients who experienced at least one adverse event (Travis 1994; Hanauer 1996; Kruis 2001; Paoluzi 2005; Kruis 2011). Kruis 2011 found a statistically significant difference in the proportion of patients who experienced at least one adverse event between Salofalk 3 g/day and 1.5 g/day both dosed once daily. Forty-one per cent of patients in the 3 g/day group experienced at least one adverse event compared to 55% of patients in the 1.5 g/day group (429 patients; RR 0.74; 95% CI 0.61 to 0.91). Hanauer 1996 found a statistically significant difference in the proportion of patients who experienced at least one adverse event between Asacol 1.6 g/day and 0.8 g/day. Forty-one per cent of patients in the Asacol 1.6 g/day group experienced at least one adverse event compared to 22% of patients in the 0.8 g/day group (177 patients; RR 1.86; 95% CI 1.18 to 2.95). No statistically significant differences in the incidence of adverse events were found between Asacol 2.4 g/day and 1.2 g/day (RR 2.85; 95% CI 0.12 to 68.95), Balsalazide 6.0 g/day and 3.0 g/day (RR 1.40; 95% CI 0.88 $\,$ to 2.24), and Olsalazine 2.0 g/day and 1.0 g/day (RR 1.37; 95% CI 0.94 to 1.99).

Seven dose-ranging studies reported the proportion of patients who were withdrawn due to adverse events (Giaffer 1992a; Green

1992; Fockens 1995; Hanauer 1996; Kruis 2001; Paoluzi 2005; Kruis 2011). No statistically significant differences in withdrawal due to adverse events were found between Asacol 2.4 g/day and 1.2 g/day (1 study, 156 patients, RR 2.85; 95% CI 0.12 to 68.95); Asacol 1.6 g/day and 0.8 g/day (1 study, 177 patients, RR 0.34; 95% CI 0.04 to 3.25); Balsalazide 6.0 g/day and 3.0 g/day (2 studies, 196 patients, RR 0.59; 95% CI 0.21 to 1.70); Balsalazide 4.0 g/day and 2.0 g/day (1 study, 133 patients, RR 1.43; 95% CI 0.54 to 3.80); Salofalk 3.0 g/day and 1.5 g/day (1 study, 429 patients, RR 0.98; 95% CI 0.29 to 3.33); and Pentasa 3.0 g/day and 1.5 g/day (1 study, 169 patients, RR 1.06; 95% CI 0.07 to 16.69).

Eight dose-ranging studies reported the proportion of patients who were excluded or withdrawn after entry (Giaffer 1992a; Green 1992; Travis 1994; Fockens 1995; Hanauer 1996; Kruis 2001; Paoluzi 2005; Kruis 2011). A statistically significant difference was found between Balsalazide 6.0 g/day and 3.0 g/day (2 studies, 196 patients, RR 0.47; 95% CI 0.26 to 0.84) and between Salofalk 3 g/day and 1.5 g/day (1 study, 429 patients, RR 0.66; 95% CI 0.46 to 0.93). No statistically significant differences were found in exclusions or withdrawals after entry between Asacol 2.4 g/day and 1.2 g/day (1 study, 156 patients, RR 0.95; 95% CI 0.38 to 2.40); Asacol 1.6 g/day and 0.8 g/day (1 study, 177 patients, RR 1.23; 95% CI 0.80 to 1.90); Balsalazide 4.0 g/day and 2.0 g/day (1 study, 133 patients, RR 1.27; 95% CI 0.77 to 2.12); Olsalazine 2.0 g/day and 1.0 g/day (1 study, 127 patients, RR 1.75; 95% CI 0.83 to 3.70); and Pentasa 3.0 g/day and 1.5 g/day (1 study, 169 patients, RR 0.83; 95% CI 0.44 to 1.55).

DISCUSSION

This updated systematic review has largely confirmed the results of previous meta-analyses (Sutherland 1993; Sutherland 1997; Sutherland 2006; Feagan 2012), but differs from the previous work in a variety of aspects. The updated review includes 41 studies and 8928 patients which greatly increases statistical power. Different quality assessment criteria (i.e. Cochrane risk of bias tool) were also used. The current review also utilized the GRADE criteria (Guyatt 2008; Schünemann 2011). to assess the overall quality of the data obtained from the randomized studies included in the review.

Unfortunately, there are limitations to making general conclusions. Almost every study utilized a unique clinical or endoscopic index. Unlike Crohn's disease, the lack of standard indices in ulcerative colitis prevented the collection of consistent treatment efficacy data and makes comparisons across clinical studies difficult. As well, several studies failed to specify the treatment arm to which certain excluded patients were initially randomized. Despite these and other common factors that must be considered when interpreting meta-analyses, the data provided strong evidence that pointed towards a number of conclusions.

The effectiveness of oral 5-ASA preparations for maintenance of remission in quiescent ulcerative colitis was confirmed. Oral 5-ASA is superior to placebo for maintenance of remission in ulcerative colitis. The quality of the placebo-controlled trials was assessed using the Cochrane risk of bias tool and the possibility of bias was rated as low for these studies. The outcome failure to maintain clinical or endoscopic remission was rated as 'high' using the GRADE criteria indicating that further research is very unlikely to change our confidence in the point estimates of effect. There was a trend towards greater efficacy with higher doses of 5-ASA with a statistically significant benefit for the 1 to 1.9 g/day and the \geq 2 g/day dosage groups.



An interesting result was that SASP was found to have a modest, but statistically significant benefit over 5-ASA. The quality of the SASP-controlled trials was assessed using the Cochrane risk of bias tool and the possibility of bias was rated as low for these studies. The outcome failure to maintain clinical or endoscopic remission was rated as 'high' using the GRADE criteria indicating that further research is unlikely to change our confidence in the point estimates of effect. When the pooled analysis was limited to trials with endpoints at 12 months the difference was no longer statistically significant. Nevertheless, certain limitations may have resulted from having combined all trials regardless of whether relapse was defined in terms of clinical or endoscopic criteria. It is possible that the "superiority" of SASP over 5-ASA is a reflection of the intention-to-treat analysis which was employed. This technique considers all patients who received the medication and penalizes medications with high drop-out rates (for example olsalazine). A "per protocol" analysis which includes those patients who are compliant and who tolerate the medication might not support a "superiority" claim.

When data for maintenance therapy were subgrouped according to the specific 5-ASA preparation, olsalazine was observed to be significantly inferior to SASP. Firm conclusions regarding other preparations, which have generally been subject to less rigorous clinical evaluation, could not be surmised. In the case of olsalazine, it appeared that the reduced efficacy was influenced by the significantly higher proportions of withdrawals due to adverse events and total exclusions or withdrawals of patients receiving olsalazine compared to those receiving SASP. In fact, if the analysis of efficacy was restricted to relapses as treatment failures, excluding withdrawals for other reasons, there was no significant difference between olsalazine and SASP (data not shown). The difference may also be related to a misclassification bias in which patients who developed diarrhea were falsely classified as treatment relapses rather than having experienced adverse events.

The overall superiority of SASP over 5-ASA in maintenance therapy may also be attributable to certain pharmacological properties of SASP, including potential therapeutic effects of the sulfapyridine moiety, that are not observed with other 5-ASA delivery systems. The mechanisms of action of SASP and its metabolites have been reviewed by Greenfield 1993.

It was apparent that the newer 5-ASA preparations were not entirely innocent of adverse effects in a number of patients. However, the incidence of adverse events and withdrawals due to the 5-ASA formulations did not significantly differ from that associated with placebo. There was also no apparent difference between the number of adverse events caused by SASP and 5-ASA. It should be noted that there may have been a bias in favour of SASP since many of the studies involved patients who were known to have tolerated SASP in the past. This may have minimized SASP-related adverse events in these trials.

In contrast to these results, olsalazine was associated with a significantly higher proportion of withdrawals due to adverse events relative to SASP. The most common adverse event attributed to olsalazine was diarrhea, an effect previously observed to occur in approximately 10% of patients receiving the drug (Ireland 1988b). It has been suggested that protocol alterations may reduce withdrawal rates in future trials since it has been reported that encouraging patients to take olsalazine with meals appears to reduce the incidence of diarrhea to approximately three per cent of

patients (Jarnerot 1996). However, four of the five olsalazine-SASP trials (Kiilerich 1992, Kruis 1995, Nilsson 1995, Rijk 1992) reported that such recommendations were in fact made.

This meta-analysis indicates that mesalamine administered once daily is as effective as conventional dosing (twice or three times daily) for maintenance of remission over 6 and 12 months periods in patients with quiescent ulcerative colitis. The pooled analyses showed no significant differences between once daily and conventional dosing for maintenance of remission at 6 months (RR 1.02; 95% CI 0.85 to 1.23; P = 0.82) or 12 months (RR 0.91; 95% CI 0.82 to 1.01; P = 0.09). With the exception of Pentasa®, subgroup analyses by drug formulation showed no significant differences in efficacy between once daily and conventional dosing for maintenance of remission. Dignass 2009 found that 2 g of Pentasa® dosed once daily was superior to 1 g Pentasa® dosed twice daily for maintenance of remission at 12 months. The other Pentasa® study found no difference between once daily and conventional dosing for maintenance of remission. A plausible biological explanation for the Dignass 2009 finding is not readily apparent to us.

We believe that the methodological basis for these conclusions is relatively sound. The quality of the trials comparing once daily with conventional dosing was assessed using the Cochrane risk of bias tool and the possibility of bias was judged to be low for most items assessed. However, a concern exists regarding blinding. One open-label study (Kamm 2008) and five studies (Kane 2003a; Kane 2008a; Dignass 2009; Sandborn 2010; Hawthorne 2012) that were single-blind (investigator-blind) were rated as having a high risk of bias. However, the open-label study (Kamm 2008), and 3 of the 5 single-blind studies (Kane 2008a; Dignass 2009; Hawthorne 2012), included endoscopy as an endpoint which may provide some protection against performance and detection bias. The overall quality of the evidence using the GRADE approach was rated as moderate for the selected primary and secondary outcomes of interest due to sparse data or high risk of bias (due to blinding) in the pooled analyses.

The results of this meta-analysis suggest that there is no difference in safety between once daily and conventionally dosed mesalamine. No differences between once daily and conventionally dosed mesalamine were observed for safety outcomes including the overall incidence of adverse events or withdrawal from treatment due to an adverse event. In keeping with the well-established safety profile of mesalamine, most of the adverse events reported in the studies assessing once daily dosing were mild to moderate in intensity. Common adverse events included gastrointestinal symptoms (e.g. flatulence, abdominal pain, nausea, and diarrhea), headache and worsening ulcerative colitis.

Important patient preference and adherence differences may exist between dosing regimens. In the study that measured patient preference the majority of patients preferred once daily dosing to conventional dosing (Sandborn 2010). Although it is generally believed that administration of fewer tablets and less frequent dosing improves both efficacy and adherence, we could not demonstrate the superiority of once daily dosing for either of these outcomes. This result suggests that patient adherence does not appear to be enhanced by once daily dosing in the clinical trial setting. Several possible explanations exist for these observations, however the most plausible one concerns the unique aspects of



the clinical trial environment. It is noteworthy that adherence with medication was remarkably high in the studies that measured this outcome (Kamm 2008; Dignass 2009; Prantera 2009; Hawthorne 2012; Kruis 2011). The pooled adherence rate for the maintenance of remission studies was 86% for the once daily dosing group compared to 89% for the conventional dosing group. These rates likely reflect the highly supervised environment in which the studies were conducted. Adherence with medication in clinical trials is generally greater than in clinical practice since participants are highly selected volunteers who are more likely, in general, to be adherent with drug regimens (Andrade 1995; Kane 2001; Kane 2006; Kane 2008b). In addition, adherence is continuously reinforced during the clinical trial process. Thus, it may be difficult to detect differences in adherence between once daily and multiple dose regimens in this setting.

Accordingly, a need exists to compare dosing regimens in large scale community-based studies. In this regard reported adherence rates in community based studies range from 40 to 60% and are especially poor among patients in remission (Levy 1999; Kane 2001; Kane 2003b; Shale 2003). However, whether once daily dosing regimens improve adherence in the community remains unknown. Although Kane 2003a demonstrated significantly higher adherence among patients receiving once daily dosing compared to conventional dosing at 3 months, no significant differences were found at 6 months. This time-dependent effect has recently been observed in a larger study (Sandborn 2010). Sandborn 2010 found significantly higher adherence among patients using once daily dosing compared to conventional dosing at 3 months. However no statistically significant difference in adherence was found at 6 and 12 months (Sandborn 2010).

Experience from other indications suggest that factors other than the dosing regimen are important for long-term compliance (Brixner 2007; Kane 2008b). Long-term observations in ulcerative colitis patients as well as in other indications indicate that patients and physicians behaviors play a dominant role in adherence (Magowan 2006; Beaulieu 2009). The patient-physician relationship should reinforce adherence through education, open communication and mutual agreement regarding the value of treatment (Kane 2008b). To ensure continued adherence in a community based setting, Sandborn 2010 have emphasized the importance of health care-providers evaluating and reinforcing compliance with patients after three months of maintenance therapy.

There does not appear to be any difference in efficacy between the various formulations of oral 5-ASA. The overall pooled risk ratio (6 studies, n = 707) showed no statistically significant difference in relapse between various formulations of 5-ASA (including Balsalazide, Pentasa and Olsalazine) and comparator formulations of 5-ASA (including Asacol and Salofalk). However, a GRADE analysis indicated that the overall quality of the evidence for the primary outcome (failure to maintain clinical or endoscopic remission at 12 months) was low due to a high risk of bias (single blind and openlabel) in three studies in the pooled analysis and sparse data (See Summary of findings 4). However, the open-label study (Mahmud 2002) and one of the single-blind studies (Courtney 1992), included endoscopy as an endpoint which may provide some protection against performance and detection bias.

Pharmacokinetic studies suggest that systemic exposure to 5-ASA is similar for all oral 5-ASA formulations and 5-ASA prodrugs

(Sandborn 2002a; Sandborn 2002b; Sandborn 2002c; Sandborn 2003). The excretory function of the kidneys (as measured by the glomerular filtration rate) does not change during maintenance therapy with oral 5-ASA or olsalazine, and nephrotoxicity is rare for Pentasa or Asacol, suggesting that the systemic exposure to 5-ASA that occurs for doses used in clinical practice is safe for all drugs in this class (Sandborn 2002a). With the exception of olsalazinerelated diarrhea, there does not appear to be any difference in safety between the various formulations of oral 5-ASA. The overall pooled risk ratios showed no statistically significant differences in the incidence of adverse events, withdrawal due to adverse events or exclusions or withdrawals after entry. Thus, all of the 5-ASA formulations can be considered safe and effective for the treatment of active ulcerative colitis, and from a practical standpoint, they can be considered therapeutically equivalent at equimolar doses (Sandborn 2002a). Treatment with sulfasalazine and olsalazine may not be preferable due to the high frequency of adverse events. When deciding which 5-ASA formulations to use physicians and patients should consider dose-response data, adherence issues related to dose forms (size of dose form and total number of tablets or capsules per day), and price (Sandborn 2002a).

Few dose ranging maintenance studies were performed which limits the conclusions that can be drawn. Hanauer 1996 compared Asacol at a dosage of 1.6 g/day to 0.8 g/day and found no difference in relapse rates between the dosage groups. Deventer 2001 compared 3.2 g/day to 2 g/day and found no difference in relapse rates between the dosage groups. Paoluzi 2005 compared Asacol at a dosage of 2.4 g/day to 1.2 g/day and found no difference in relapse rates between the dosage groups. However, patients in the higher dosage group remained in remission longer, compared to patients in the low dose group. Paoluzi 2005 recommended a dosage of 2.4 g/day due to the significantly longer time to relapse in the higher dosage group. Pica 2012 compared 4.8 g/day to 2.4 g/day and found no difference in relapse rates between the dosage groups. Further research may be needed to determine the ideal dosage of Asacol for maintenance therapy.

Three studies compared the efficacy of high dose Balsalazide (4.0 to 6.0 g/day) to low dose (2.0 to 3.0 g/day). Giaffer 1992a found a dose of 4.0 g/day to be significantly superior to 2.0 g/day for preventing relapses over a 12 months period. No differences in safety between the dose groups were noted. A pooled analysis of two studies comparing Balsalazide 6.0 g/day to 3.0 g/day found no significant difference in relapse rates. However, these results should be interpreted with caution due to a high degree of heterogeneity and sparse data. The high degree of heterogeneity is due to the fact that the two studies had conflicting results. Green 1992 found no differences in relapse rates between the 6.0 g/day and 3.0 g/day groups at either 6 or 12 months. Green 1992 noted no differences in safety. Kruis 2001 found 6.0 g/day to be superior to 3 g/day for preventing relapse over a 26 week period with no differences in safety. Differences in patient populations may explain these findings. The Green 1992 study included patients with a very distal extent of ulcerative colitis and a large proportion of patients had long term remission (e.g. >1 year) at entry. Kruis 2001 suggested that patients with more extensive ulcerative colitis or with frequent relapses may benefit from a higher dose of maintenance therapy.

Kruis 2011 investigated different doses of once daily Salofalk (3 g or 1.5 g) and found that significantly fewer patients relapsed at 12 months in the group receiving 3 g once daily (25%) compared to



patients in the 1.5 g group (39%; P = 0.002). This analysis involved 439 patients and provides moderate evidence (based on GRADE analysis) that 3 g Salofalk once daily is superior to 1.5 g Salofalk once daily for maintenance treatment of ulcerative colitis. No differences in safety were observed among the dose groups. In a post hoc analysis, Kruis 2011 observed that patients with active inflammation at baseline in the 3 g group continued to maintain a higher rate of remission than patients receiving lower doses (i.e. 1.5 g once daily or 0.5 g three times daily). Kruis 2011 concluded that 3 g once daily is an appropriate dose for maintenance of remission, and may be beneficial for patients with signs of inflammation or in whom endoscopic data are not available. No other maintenance studies looked at dose-ranging for once daily treatment using other 5-ASA formulations.

Travis 1994 compared the efficacy of Olsalazine 2.0 g/day to 1.0 g/day and 0.5 g/day. No significant differences in relapse at 12 months were found between the 2.0 g and 1.0 g/day groups. A dosage of 2.0 g/day was significantly superior to 0.5 g/day for preventing relapse. No differences in safety were noted. Subgroup analysis showed that patients with proctitis and recent relapse may benefit from a dosage of 2.0 g/day (Travis 1994). Fockens 1995 compared the efficacy of Pentasa 3.0 g/day to 1.5 g/day. Although there was a trend favouring the higher dose there was no statistically significant difference in prevention of relapse over a one year period. The higher dosage was not associated with a higher incidence of adverse events (Fockens 1995).

AUTHORS' CONCLUSIONS

Implications for practice

It is clear that oral 5-ASA preparations have yet to be proven to be more clinically beneficial than SASP. Male infertility is associated with SASP and not with 5-ASA (Riley 1987; Kjaergaard 1989), so 5-ASA may be preferred for patients concerned about fertility. 5-ASA therapy is more expensive than SASP, so SASP may be the preferred option where cost is an important factor. 5-ASA could also be offered to patients who are intolerant to SASP. Oral 5-ASA administered once daily is as effective and safe as

conventional dosing (twice or three times daily) for maintenance of remission in quiescent ulcerative colitis. Once daily dosing does not appear to enhance adherence in the clinical trial setting. There does not appear to be any difference in efficacy or safety between the various formulations of 5-ASA. Patients with extensive ulcerative colitis or with frequent relapses may benefit from a higher dose of maintenance therapy. High dose therapy appears to be as safe as low dose and is not associated with a higher incidence of adverse events. When selecting among the various 5-ASA formulations, physicians and patients should consider doseresponse data, adherence issues and price (Sandborn 2002a).

Implications for research

In this time of scarce research dollars, careful thought should be given prior to commissioning new trials of 5-ASA preparations for the treatment of quiescent ulcerative colitis. Future trials comparing 5-ASA with placebo or SASP may not be justified. There does not appear to be any difference in efficacy or safety between the various formulations of 5-ASA. However the overall quality of the evidence from the studies examining differences in efficacy between various 5-ASA formulations is low due to sparse data and risk of bias. Future trials should look at enhancing patient adherence with medication. Adherence to therapy is important for treatment success and may be an important predictor of relapse. Future trials could assess whether once daily dosing regimens improve adherence in the community.

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* Indicates the major publication for the study



CHARACTERISTICS OF STUDIES

Characteristics of included studies [ordered by study ID]

	re			

Methods	Randomized, double-bind trial comparing 5-ASA and SASP. Allocation of drugs was performed using a table of random numbers
Participants	Patients with ulcerative colitis in remission (N = 13)
Interventions	5-ASA, 0.75 g/day (n = 7), or SASP, 1.5 g/day (n = 6) for a duration of 12 months
Outcomes	The development of a new acute phase within 12 months from the beginning of the survey was considered a negative result
Notes	Abstract

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Table of random numbers
Allocation concealment (selection bias)	Unclear risk	Not described
Blinding (performance bias and detection bias) All outcomes	Low risk	Double-blind
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Not described
Selective reporting (reporting bias)	Unclear risk	Not described
Other bias	Unclear risk	Not described

Ardizzone 1995

Methods	Single-center, randomized, double-blind comparison of Claversal and SASP. Clinical, endoscopic and histological examinations were performed at entry, after 6 months, and after 12 months	
Participants	Patients, aged 18 to 61 years, with clinical and endoscopically quiescent ulcerative colitis, who had an episode of active disease requiring systemic corticosteroids at least once in the previous 12 months (N = 88)	
Interventions	5-ASA (Claversal), 1 g daily (n = 44), or SASP, 2 g daily (n = 44), for 1 year. Topical corticosteroids were permitted for short-term treatment of mild distal relapses	
Outcomes	Relapse of disease was defined as the appearance of bloody diarrhea with endoscopic signs of inflammation requiring systemic steroids (major relapse). "Minor relapses" were defined as those requiring a few days of topical corticosteroids. The number of withdrawals due to adverse effects was also reported	



Ardizzone 1995 (Continued)

Notes

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not described
Allocation concealment (selection bias)	Unclear risk	Not described
Blinding (performance bias and detection bias) All outcomes	Low risk	Double-blind
Incomplete outcome data (attrition bias) All outcomes	Low risk	Drop-outs balanced across intervention groups with similar reasons for with-drawal
Selective reporting (reporting bias)	Low risk	Expected outcomes reported
Other bias	Low risk	The study appears to be free of other sources of bias

Ardizzone 1999

Methods	Single-center, randomized, double-blind, placebo-controlled trial	
Participants	Adult patients (aged 18 to 75 years) with confirmed diagnosis of intermittent chronic ulcerative colitis in stable clinical, endoscopic and histological remission for at least one year (N = 112)	
Interventions	Asacol 1.2 g/day (n = 54) or identical placebo (n = 58) for one year	
Outcomes	Primary outcome was maintenance of remission. Relapse was defined as increased stool frequency with blood or mucus and evidence of disease activity on sigmoidoscopy. Secondary outcomes: adverse events, compliance	

Notes

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not described
Allocation concealment (selection bias)	Unclear risk	Not described
Blinding (performance bias and detection bias) All outcomes	Low risk	Double-blind, identical placebo, drugs were provided in identical blister packages



Ardizzone 1999 (Continued)		
Incomplete outcome data (attrition bias) All outcomes	Low risk	Drop-outs balanced across intervention groups with similar reasons for with-drawal
Selective reporting (reporting bias)	Low risk	Expected outcomes were reported
Other bias	Low risk	The study appears to be free of other sources of bias

Courtney 1992

Methods	Randomized single-blind trial
Participants	Adult patients (16 to 75 years) with ulcerative colitis in remission (N = 100)
Interventions	Olsalazine 1.0 g/day (n = 50) or mesalazine (Asacol) 1.2 g/day (n = 50) for 12 months
Outcomes	Remission was defined as absence of symptoms or the presence of only mild stable symptoms of colitis. Relapse was defined as development of new symptoms of colitis sufficiently severe enough to warrant the introduction of systemic steroid therapy (by a blinded investigator). Adverse events

Notes

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Patients were randomly allocated to treatment groups by means of a computer-generated code
Allocation concealment (selection bias)	Unclear risk	Not described
Blinding (performance bias and detection bias) All outcomes	High risk	Single-blind (observer)
Incomplete outcome data (attrition bias) All outcomes	Low risk	Drop-outs balanced across intervention groups with similar reasons for with-drawal
Selective reporting (reporting bias)	Low risk	Expected outcomes were reported
Other bias	Low risk	The study appears to be free of other sources of bias

D'Haens 2012

patients with ulcerative colitis who were in endoscopic remission	Methods	Multi-centre, double-blind, double-dummy, randomized, non-inferiority trial comparing the efficacy and safety of once daily MMX mesalamine (2.4 g/day) with twice daily mesalamine (Asacol, 1.6 g/day) in patients with ulcerative colitis who were in endoscopic remission
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D'Haens 2012 (Continued)

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Adult patients (\geq 18 years) with a diagnosis of UC (confirmed by histology) that was considered to be in remission for \geq 30 days on a stable dose of mesalamine (\leq 2.4 g/ day) or the equivalent dose of sulfasalazine (\leq 6.2 g/ day), with an endoscopy score of \leq 1; and had a combined symptom score (stool frequency and rectal bleeding) of \leq 1. All patients were to have had experienced at least one acute flare of UC (defined as a documented episode of increased bowel frequency with rectal bleeding for which UC therapy was intensified) in the past 12 months, with at least two acute flares in their medical history (N = 826)

Interventions MMX mesalamine (2.4 g/day) dosed once daily (n = 415) versus Asacol (1.6 g/day) dosed twice daily (n = 411) for 6 months

Outcomes

Primary outcome was endoscopic remission at 6 months defined as a modified UC-DAI endoscopy subscore of ≤ 1 point. Secondary outcomes included: maintenance of mucosal healing with no or mild symptoms (combined modified UC-DAI-defined stool frequency and rectal bleeding subscores of ≤ 1 point) at 6 months; time to relapse (relapse was defined as withdrawal due to lack of efficacy), measured from the date of randomization to the date the patient withdrew due to relapse; the modified UC-DAI score and its components (rectal bleeding, stool frequency, endoscopy, and Physician's Global Assessment scores); and safety and tolerability of study treatments

Notes

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not described
Allocation concealment (selection bias)	Low risk	The randomization was performed centrally via an interactive voice response system
Blinding (performance bias and detection bias) All outcomes	Low risk	Double-blind, double-dummy (matched placebos)
Incomplete outcome data (attrition bias) All outcomes	Low risk	Drop-outs balanced across intervention groups with similar reasons for with- drawal
Selective reporting (reporting bias)	Low risk	All expected outcomes reported
Other bias	Low risk	The study appears to be free of other sources of bias

Deventer 2001

Methods	Randomized, multi-center (46), single-blind trial
Participants	Patients with quiescent ulcerative colitis for more than 1 month but less than 3 years (N = 388)
Interventions	Asacol 2 g/day (n = 124), Asacol 3.2 g/day (n = 138), or Salofalk 2 g/day (n = 126) for 18 months
Outcomes	Primary outcome: remission at 18 months
Notes	Abstract publication



Deventer 2001 (Continued)

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not described
Allocation concealment (selection bias)	Unclear risk	Not described
Blinding (performance bias and detection bias) All outcomes	High risk	Single-blind. Sigmoidoscopies were videotaped at study entry and after 9 and 18 months of treatment and blindly scored for disease activity by 2 gastroenterologists
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Drop-outs and reasons for withdrawal were not described
Selective reporting (reporting bias)	Unclear risk	Secondary outcomes were not described
Other bias	Unclear risk	Abstract publication

Dew 1983a

Methods	Randomized, double-blind, double-dummy trial
Participants	Adult patients (N = 67) with ulcerative colitis in remission (3 or less stools daily without blood or slime and normal sigmoidoscopic findings)
Interventions	5-ASA (Asacol mean dose 2.7 g/day; range: 2.4 to 4.4 g/day; n = 32) or SASP (mean dose 2.3 g/day; range 2 to 4 g/day; n = 25) for 6 months
Outcomes	Relapse defined as recurrence of symptoms with sigmoidoscopic changes including contact or spontaneous mucosal hemorrhages, pus or ulceration

Notes

Nich Ci Diuc		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not described
Allocation concealment (selection bias)	Unclear risk	Not described
Blinding (performance bias and detection bias) All outcomes	Low risk	Double-blind, double-dummy (identical placebo tablets)
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	10 of 67 patients withdrew. 4 SASP patients withdrew due to headaches. We do not know which groups the other 6 patients belonged to



Dew 1983a (Continued)		
Selective reporting (reporting bias)	Low risk	Expected outcomes were reported
Other bias	Low risk	The study appears to be free of other sources of bias

Dignass 2009

Methods	Multicenter, randomized, investigator-blinded, non-inferiority trial
Participants	Adult patients (\geq 18 years) with an established diagnosis of ulcerative colitis and in clinical remission (UC-DIA < 2 at entry) (N = 362)
Interventions	Mesalamine (Pentasa) 2 g once daily (n = 169) or 1 g twice daily (n = 184) for 12 months
Outcomes	The primary outcome was maintenance of clinical remission at 12 months. Secondary outcomes: time to relapse, UC-DAI total and subscores, adherence, global acceptability of treatment and adverse events
Notes	Six patients in the OD group and 3 in the BID group were excluded from the ITT analysis for major entry criteria violation. Adherence was self-reported by patients using a VAS scale

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not described
Allocation concealment (selection bias)	Low risk	Patients were randomized centrally using an interactive voice response system
Blinding (performance bias and detection bias) All outcomes	High risk	Single-blind (investigator), outcomes assessed by blinded investigator
Incomplete outcome data (attrition bias) All outcomes	Low risk	Drop-outs balanced across intervention groups with similar reasons for with-drawal
Selective reporting (reporting bias)	Low risk	All expected outcomes reported
Other bias	Low risk	The study appears to be free of other sources of bias

Fockens 1995

Methods	Multicenter, randomized, double-blind trial	
Participants	Adult patients (18 to 75 years) with ulcerative colitis in remission for at least and month and up to 3 years ($N = 169$)	
Interventions	Mesalazine (Pentasa) 1.5 g/day (n = 87) or 3.0 g/day (n = 82) for 12 months	



Fockens 1995 (Continued)

Outcomes Primary outcome: maintenance of remission at 12 months. Secondary outcomes: serious adverse

events

Notes

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer-generated: Randomization was performed with the SAS statistical package in balanced blocks of 10 patients
Allocation concealment (selection bias)	Unclear risk	Not described
Blinding (performance bias and detection bias) All outcomes	Low risk	Double-blind: placebo tablets were identical to Pentasa tablets in appearance, weight and taste
Incomplete outcome data (attrition bias) All outcomes	Low risk	Drop-outs balanced across intervention groups with similar reasons for with- drawal
Selective reporting (reporting bias)	Low risk	Expected outcomes were reported
Other bias	Low risk	The study appears to be free of other sources of bias

Giaffer 1992a

Methods	Multicenter, randomized, double-blind trial		
Participants	Patients with ulcerative colitis in clinical (minimal or no symptoms) and sigmoidoscopic (non-mucosa) remission (N = 133)		
Interventions	Balsalazide 2 g/day (n = 65) or 4 g /day (n = 68)		
Outcomes	Relapse at 12 months and adverse events. Symptomatic relapse was defined as recurrence of previous symptoms with increased stool frequency and associated blood		

Notes

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not described
Allocation concealment (selection bias)	Unclear risk	Not described
Blinding (performance bias and detection bias) All outcomes	Low risk	Double-blind: The placebo and trial drug were in identical hard gelatin capsules



Giaffer 1992a (Continued)		
Incomplete outcome data (attrition bias) All outcomes	Low risk	Drop-outs balanced across intervention groups with similar reasons for withdrawal
Selective reporting (reporting bias)	Low risk	Expected outcomes were reported
Other bias	Low risk	The study appears to be free of other sources of bias

Green 1992

Methods	Multicenter, randomized, double-blind trial	
Participants	Adult patients (aged 19 to 78 years) with ulcerative colitis in clinical and sigmoidoscopic remission (N = 108)	
Interventions	Balsalazide 3 g/day (n = 54) or 6 g/day (n = 54) for 12 months	
Outcomes	Primary outcome: relapse at 12 months. Secondary: adverse events. Relapse was defined on clinical (7 days of increased stool frequency with or without blood and mucus), sigmoidoscopic (friable mucosa or spontaneous hemmorhage) and histological grounds.	

Notes

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not described
Allocation concealment (selection bias)	Unclear risk	Not described
Blinding (performance bias and detection bias) All outcomes	Unclear risk	Double-blind. Dummy capsules were not described in detail
Incomplete outcome data (attrition bias) All outcomes	Low risk	Drop-outs balanced across intervention groups with similar reasons for with-drawal
Selective reporting (reporting bias)	Low risk	Expected outcomes were reported
Other bias	Low risk	The study appears to be free of other sources of bias

Green 1998

Methods	Multicenter, randomized, double-blind, double-dummy, parallel group trial
Participants	Adult patients (aged 18 to 80 years) with ulcerative colitis in remission (N = 99)



Green 1998 (Continued)		
Interventions	Balsalazide 3 g/day (n	= 50) or Asacol 1.2 g/day (n = 49) for 12 months
Outcomes	The primary outcome was the proportion of patients in remission at 3 and 12 months. Seconadry outcomes: symptom free days, symptoms over three months, adverse events (diary cards)	
Notes		
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not described
Allocation concealment (selection bias)	Unclear risk	Not described
Blinding (performance bias and detection bias) All outcomes	Low risk	Double-blind, double-dummy, identical placebos
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	13 patients (26%) withdrew from the balsalazide group compared to 5 patients (11%) in the Asacol group
Selective reporting (reporting bias)	Low risk	Expected outcomes were reported
Other bias	Low risk	The study appears to be free of other sources of bias
Hanauer 1996		
Methods	Multicenter, randomized, double-blind, placebo-controlled trial. Randomization was performed within centers by means of codes using specific patient numbers computer-generated for each study site before the study began. In addition to daily diaries, clinical assessments, endoscopy, and lab analyses were performed at entry, 1, 3 and 6 months	
Participants	Patients, aged 18 to 75 years, with ulcerative colitis in remission for at least 1 month, were recruited from 17 study sites. The presence of symptoms such as loose stools or abdominal cramps were not reasons for exclusion. Corticosteroid or topical rectal therapy was stopped at least 1 month before entry (I = 264)	
Interventions	Mesalamine (Asacol), 0.8 g/day (n = 90) or 1.6 g/day (n = 87) or matched placebo (n = 87) for 6 months. 400 mg tablets were packaged in 4 bottles, each labelled with the time of day at which its contents were to be taken. Compliance was monitored by pill count	
Outcomes	Treatment outcome was rated as either 'success' (maintenance of remission as indicated by endoscopic evaluation) or 'failure' (endoscopic relapse or withdrawal), and was reported on an intention-to-trebasis	
Notes		
Risk of bias		



Hanauer 1996 (Continued)		
Random sequence generation (selection bias)	Low risk	Computer generated
Allocation concealment (selection bias)	Low risk	Adequate
Blinding (performance bias and detection bias) All outcomes	Low risk	The placebo tablets were identical to the mesalamine tablets in odor and appearance but contained no active ingredients
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Seventy-five patients were excluded from the primary efficacy analysis for the following reasons: failure to meet study entry criteria ($n = 36$), noncompliance with study medication ($n = 18$), noncompliance with study procedures ($n = 3$), concomitant medication violation ($n = 10$), loss to follow-up ($n = 4$), and voluntary withdrawal ($n = 4$). The numbers of patients excluded were similar in the three groups
Selective reporting (reporting bias)	Low risk	Expected outcomes were reported
Other bias	Low risk	The study appears to be free of other sources of bias

Hawkey 1997

Methods	International, multicenter, randomized, double-blind study comparing mesalazine and placebo for their ability to maintain remission in UC	
Participants	Patients, age 18 or above, with ulcerative colitis in remission as defined by normal sigmoidoscopic appearances with no rectal bleeding during the week before entry. Patients already receiving salicylates could enter; those receiving oral or rectal steroids could only do so if these were tapered over 2 weeks prior to entry (N = 323)	
Interventions	One 400 mg mesalazine capsule and 2 placebo capsules four times per day ($n = 99$) or 3 placebo capsules four times per day ($n = 111$) for a duration of 6 months. All capsules were identical	
Outcomes	The primary efficacy end point was the maintenance of remission. Relapse was defined as a sigmoido- scopic score of 1 or more or 3 consecutive days of rectal bleeding caused by UC or liquid stools for 1 week. The number of patients withdrawing due to adverse events was also recorded	
Notes	The prime objective of this study was the measurement of the efficacy of zileuton (n = 113) in UC	
Bid office		

Bias	Authors' judgement	Support for judgement
Dias	Authors judgement	Support for Judgement
Random sequence generation (selection bias)	Unclear risk	Not described
Allocation concealment (selection bias)	Low risk	Centralized randomization: Concealed randomization schedules were held at each participating hospital for code break in the event of serious adverse events
Blinding (performance bias and detection bias) All outcomes	Low risk	Each capsule was identical in appearance



Hawkey 1997 (Continued)		
Incomplete outcome data (attrition bias) All outcomes	Low risk	Drop-outs balanced across intervention groups with similar reasons for withdrawal
Selective reporting (reporting bias)	Low risk	Expected outcomes were reported
Other bias	Low risk	The study appears to be free of other sources of bias

Hawthorne 2012

Methods	Multicenter, single-blind, randomized, non-inferiority study comparing once daily mesalazine to conventionally dosed mesalazine (TID) for maintenance of remission in UC	
Participants	Adult UC patients taking mesalazine or sulphasalazine in remission for > 4 weeks and < 2 years (N = 213)	
Interventions	Asacol 2.4 g/day given once daily (3 x 800 mg tablets; $n = 103$) versus Asacol 2.4 g/day given three times daily (1 X 800 mg tablet 3 times per day, $n = 110$) for one year	
Outcomes	The primary end-point was the difference between groups in relapse rates over one year. Relapse was defined as typical symptoms of relapse with a Baron sigmoidoscopy score of 2 or 3. Secondary outcomes: adherence	

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not described
Allocation concealment (selection bias)	Unclear risk	Not described
Blinding (performance bias and detection bias) All outcomes	High risk	Investigator-blinded, outcomes assessed by blinded investigator
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Not described
Selective reporting (reporting bias)	Low risk	Expected outcomes were reported
Other bias	Low risk	The study appears to be free of other sources of bias



Ireland 1988	
Methods	Randomized, double-blind, double-dummy trial comparing olsalazine and SASP. Randomization was in blocks of 10, and the drugs were dispensed by the hospital pharmacy. At entry, 3, and 6 months, a history, clinical exam, sigmoidoscopy, and rectal biopsy were performed
Participants	Out-patients, aged 18 to 75 years, with ulcerative colitis in remission (without a relapse for previous 6 months) taking an average dose of 2 g/day SASP, were recruited over an 18 month period (N = 164)
Interventions	SASP, 2 g/day, plus olsalazine-matched placebo (n = 82) or olsalazine, 1 g/day, plus SASP-matched placebo (n = 82) for 6 months
Outcomes	Relapse was defined as increased stool frequency with or without blood or mucus and with evidence of inflammation on sigmoidoscopy. Withdrawals and the occurrence of side-effects were also reported
Notes	

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not described
Allocation concealment (selection bias)	Low risk	Adequate
Blinding (performance bias and detection bias) All outcomes	Low risk	Double-blind, double-dummy
Incomplete outcome data (attrition bias) All outcomes	Low risk	Drop-outs generally balanced across intervention groups with similar reasons for withdrawal with the exception of a greater number of patients withdrawing from the olsalazine group ($n = 10$) due to diarrhea ($n = 3$ in SASP group)
Selective reporting (reporting bias)	Low risk	Expected outcomes were reported
Other bias	Low risk	The study appears to be free of other sources of bias

Ito 2010

Methods	Multicenter, randomized, double-blind, double-dummy, non-inferiority trial		
Participants	Adult patients (aged \geq 16 to < 65 years) with quiescent (UC-DAI of 2 or less and bloody stool score of 0) ulcerative colitis (N = 131)		
Interventions	Pentasa 2.25 g/day (n = 66) or Asacol 2.4 g/day (n = 65) for 48 weeks		
Outcomes	Primary outcome: the proportion of patients without bloody stools. Secondary outcomes were time to bloody stools, proportion of patients without relapse, time to relapse, decrease in UC-DAI and adverse events. Relapse was defined as a bloody stool score of 1 or more and UC-DAI of 3 or more		
Notes	One patient from the Pentasa group was excluded from the ITT analysis due to a GCP violation		
Risk of bias			



Ito 2010 (Continued)

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer generated. Treatment assignment was balanced using a biased-coin minimization algorithm
Allocation concealment (selection bias)	Low risk	Centralized randomization: A person independent from the study was in charge of the random allocation. The randomization code was sealed and stored until the blind was removed
Blinding (performance bias and detection bias) All outcomes	Low risk	Double-blind, double-dummy
Incomplete outcome data (attrition bias) All outcomes	Low risk	Drop-outs balanced across intervention groups with similar reasons for with-drawal
Selective reporting (reporting bias)	Low risk	Expected outcomes were reported
Other bias	Low risk	The study appears to be free of other sources of bias

Kamm 2008

Methods	Multicenter, randomized, open-label trial		
Participants	Adult patients were entered following induction of remission after an acute flare of mild to moderate ulcerative colitis (N = 451) in clinical and endoscopic remission (modified UC-DAI score of \leq 1, with rectal bleeding and stool frequency scores of 0, a combined PGA and sigmoidoscopy score of \leq 1, no mucosal friability and a \geq 1 point reduction from baseline in sigmoidoscopy score)		
Interventions	MMX mesalazine 2.4 g/day given once daily (n = 225) or 2.4 g/day given in two divided doses of 1.2 g (n = 234) for 12 months		
Outcomes	Primary outcome: safety and tolerability of the two dosage regimens over 12 months. Secondary out comes: proportion of patients in remission at 12 months, changes from baseline in components of the modified UC-DAI, and adherence		
Notes	Compliance with study medication was calculated by pill count at study visits		

Nisk of Dias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	Unclear risk	Not described	
Allocation concealment (selection bias)	Low risk	Centralized randomization:Patients entering this 12-month maintenance study were randomized via an interactive voice recognition system	
Blinding (performance bias and detection bias) All outcomes	High risk	Open-label	
Incomplete outcome data (attrition bias)	Low risk	Drop-outs balanced across intervention groups with similar reasons for with-drawal	



Kamm 2008	(Continued)
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All outcomes

Selective reporting (reporting bias)	Low risk	All expected outcomes reported
Other bias	Low risk	The study appears to be free of other sources of bias

Kane 2003a

Methods	Single-center, investigator-blinded, randomized trial		
Participants	Adult patients with ulcerative colitis in remission (defined as absence of blood in stools, urgency or cramping) for at least 4 months prior to entry (N = 22)		
Interventions	Mesalamine (Asacol) dosed once daily (n = 12) or twice or three times daily (n = 10) for 6 months		
Outcomes	The primary outcome was the presence of absence of disease activity at the end of 6 months treatment. Relapse was defined as a HBI > 3. Secondary outcomes: adherence and patient satisfaction with treatment		
Notes	The mean dose of Asacol in the once daily group was 2.5 g/day. The mean dose of Asacol in the conventionally dosed group was 2.7 g/day. Adherence was calculated using pharmacy data		

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Random-numbers table
Allocation concealment (selection bias)	Low risk	Opaque sealed envelopes
Blinding (performance bias and detection bias) All outcomes	High risk	Investigator-blinded
Incomplete outcome data (attrition bias) All outcomes	Low risk	Drop-outs balanced across intervention groups with similar reasons for with-drawal
Selective reporting (reporting bias)	Low risk	All expected outcomes reported
Other bias	Unclear risk	The study appears to be free of other sources of bias

Kane 2008a

Methods	Two center, investigator-blinded, randomized trial	
Participants	Adult patients with ulcerative colitis in remission (defined as absence of blood in stools, urgency or cramping) for at least 4 months prior to entry (N = 20)	



ane 2008a (Continued)		
Interventions	Mesalamine (Asacol) dosed once daily (n = 12) or twice or three times daily (n = 8) for 12 months	
Outcomes	The primary outcome was the percentage of patients with quiescent disease at 12 months. Remission was defined as a modified UC-DAI \leq 3. Relapse was defined as a UC-DAI $>$ 3. Secondary outcomes: adherence and adverse events	
Notes	Adherence was calcula	ited using pharmacy data
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer-generated
Allocation concealment (selection bias)	Low risk	Opaque sealed envelopes
Blinding (performance bias and detection bias) All outcomes	High risk	Investigator-blinded, patient assessed by blinded physician at 6 and 12 month assessment
Incomplete outcome data (attrition bias) All outcomes	Low risk	One patient in the once daily mesalamine group died
Selective reporting (reporting bias)	Low risk	All expected outcomes reported
Other bias	Low risk	The study appears to be free of other sources of bias
Gillerich 1992		
Methods	ization scheme was co secutive patients withi	ed, double-blind, double-dummy comparison of olsalazine and SASP. Random- mputer-generated, stratified for each center and performed in blocks of 4 con- n each center. Clinical exam, sigmoidoscopy, and blood tests were performed at 2 months (or upon relapse)
Participants	Outpatients, aged 18 to 80 years, with ulcerative colitis in remission, but who had at least 2 attacks of ulcerative colitis, were recruited from March 1988 to March 1989 (N = 226)	
Interventions	Olsalazine, $1g/day$ (n = 114), or SASP, $2g/day$ (n = 112), in a double-dummy method for 12 months. Patients were instructed to take medications with meals. Patients were questioned upon clinical examination to verify their compliance	
Outcomes	Relapse was defined as inflammation of the rectal mucosa grade 3-4 on sigmoidoscopy (no distinct ves sels in the mucosa, spontaneous bleeding or bleeding by contact with the sigmoidoscope). The total failure rate was also reported	
Notes		
Risk of bias		
Bias	Authors' judgement	Support for judgement



Kiilerich 1992 (Continued)		
Random sequence generation (selection bias)	Low risk	Computer generated
Allocation concealment (selection bias)	Low risk	Adequate
Blinding (performance bias and detection bias) All outcomes	Low risk	Double-blind, double-dummy
Incomplete outcome data (attrition bias) All outcomes	Low risk	Drop-outs balanced across intervention groups with similar reasons for with-drawal
Selective reporting (reporting bias)	Low risk	Expected outcomes were reported
Other bias	Low risk	The study appears to be free of other sources of bias

Kruis 1995

Multicenter, randomized, double-blind comparison of 3 doses of olsalazine and a standard dose of SASP. Randomization scheme was computer-generated and performed in blocks of 8 and stratified for each center. Clinical assessments were performed at entry, 2, 4 and 6 months. Endoscopic exam was performed at entry and after 6 months (or at relapse)
Patients with proven ulcerative colitis in remission were recruited at 15 centers in 3 countries. No concomitant medications were permitted (N = 148)
Olsalazine 0.5 g/day, 1.25 g/day, 2.0 g/day, or SASP 2 g/day for 6 months. Patients were instructed to take medications with meals. Medications were similar in color, size and weight. Full dosage was reached after 5 days. Compliance was monitored
Relapse was defined as a change in the endoscopic grading, and corresponding histologic assessment, to at least moderate activity. Withdrawals were also reported
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Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	Low risk	Randomization scheme was computer-generated and performed in blocks of 8 and stratified for each center	
Allocation concealment (selection bias)	Low risk	Adequate	
Blinding (performance bias and detection bias) All outcomes	Low risk	The study was double-blind and the medication was similar in colour, size and weight	
Incomplete outcome data (attrition bias) All outcomes	Low risk	Drop-outs balanced across intervention groups with similar reasons for with- drawal	



Kruis 1995 (Continued)				
Selective reporting (reporting bias)	Low risk	Expected outcomes were reported		
Other bias	Low risk	The study appears to be free of other sources of bias		

Kruis 2001

Methods	Multicenter, randomized, double blind, double dummy, three armed, parallel group trial		
Participants	Adult patients (aged 18 to 70 years) with ulcerative colitis in remission for less than one year (N = 133)		
Interventions	Balsalazide 3 g twice daily (n = 40), Balsalazide 1.5 g twice daily (n = 48) or Salofalk 0.5 g three times daily (n = 44) for 26 weeks		
Outcomes	Primary: proportion of patients in clinical remission at 26 weeks. Secondary: CAI, endoscopic score, urine analysis, adverse events. Clinical remission was defined as a CAI < 6. Endoscopic findings were also classified according to Rachmilewitz and an EI score < 4 was considered to indicate remission. Relapse was defined as CAI > 6 and EI > 4		
Notes	1 patient randomized to the Balsalazide 1.5 g group did not receive any study medication and was excluded from the ITT analysis		

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not described
Allocation concealment (selection bias)	Unclear risk	Not described
Blinding (performance bias and detection bias) All outcomes	Low risk	Double-blind, double-dummy, identical placebos
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Patients were more likely to drop-out from the low dose balsalazide group (26.5%) due to lack of efficacy than the high dose balsalazide (7.5%) or the Salofalk (13.6%) groups
Selective reporting (reporting bias)	Low risk	Expected outcomes were reported
Other bias	Low risk	The study appears to be free of other sources of bias

Kruis 2011

Methods	Randomized, multicenter, double-blind, double-dummy, parallel group, phase III trial	
Participants	Adult patients (aged 18 to 75 years) with endoscopically and histologically confirmed ulcerative colitis (N = 648) in remission (CAI \leq 4 and EI \leq 3)	



Kruis 2011 (Continued)		
Interventions	Mesalazine (Salofalk granules) 3.0 g/day dosed once daily (n = 217), 1.5 g/day dosed once daily (n = 212) or 0.5 g dosed three times daily (1.5 g/day n = 218) for one year	
Outcomes	The primary outcome was clinical remission at the final visit (relapse was defined as CAI > 4 and an increase of \geq 3 from baseline. Secondary outcomes included: clinical remission at month 12 in patients with mucosal inflammation at baseline (EI > 1), endoscopic remission at month 12, change from baseline to month 12 in number of stools and number of bloody stools per week, adherence and adverse events	
Notes	One patient from the 3.0 g/day OD group was excluded from the ITT analysis for not receiving any study medication. Adherence was calculated by pill count at study visits and by patient diary	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer-generated randomisation list
Allocation concealment (selection bias)	Low risk	Randomisation was performed centrally in blocks of three by means of a computer-generated randomisation list
Blinding (performance bias and detection bias) All outcomes	Low risk	Double-blind, double-dummy
Incomplete outcome data (attrition bias) All outcomes	Low risk	Drop-outs balanced across intervention groups with similar reasons for with-drawal
Selective reporting (reporting bias)	Low risk	All expected outcomes reported

Lichtenstein 2010

Other bias

Methods	Multicenter, randomized, double-blind, placebo-controlled trial
Participants	Adult patients (≥ 18 years) with ulcerative colitis (N = 305) in remission (revised Sutherland Disease Activity Index (SDAI) rectal bleeding = 0 and mucosal appearance < 2)). Relapse was defined as SDAI rectal bleeding score ≥ 1 and a mucosal appearance score ≥ 2, a UC flare, or initiation of medication to treat a UC flare
Interventions	Mesalamine granules (Apriso) 1.5 g/day dosed once daily (n = 209) or placebo (n = 96) for 6 months
Outcomes	Primary: percentage of patients who were relapse free at 6 months. Secondary: percentages of patients with each level of change from baseline in rectal bleeding score, mucosal appearance score, physician's rating of disease activity and stool frequency on SDAI at months 1, 3, and 6 and end of treatment; percentage of patients classified as treatment success, relapse-free duration, and adverse events
Notes	
Risk of bias	

The study appears to be free of other sources of bias

Low risk



Lichtenstein 2010 (Continued)

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not described
Allocation concealment (selection bias)	Unclear risk	Not described
Blinding (performance bias and detection bias) All outcomes	Low risk	Double-blind, matching placebo
Incomplete outcome data (attrition bias) All outcomes	Low risk	Drop-outs balanced across intervention groups with similar reasons for with-drawal
Selective reporting (reporting bias)	Low risk	All expected outcomes reported
Other bias	Low risk	The study appears to be free of other sources of bias

Mahmud 2002

Methods	Single center, randomized, parallel group trial	
Participants	Adult patients (aged \geq 18 to 70 years) with a confirmed diagnosis of ulcerative colitis in remission (N = 40)	
Interventions	Asacol 1.2 g/day (n = 20) or Olsalazine 1 g/day (n = 20) for 9 months	
Outcomes	Clinical remission, laboratory assessments and adverse events. Remission was defined as HBI $<$ 3 and a sigmoidoscopy score of 0 to 1	
Notes	The primary objective was to evaluate the effects of 9 months of therapy with Asacol or Olsalazine on renal function	

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer-generated code
Allocation concealment (selection bias)	Unclear risk	Not described
Blinding (performance bias and detection bias) All outcomes	High risk	Open study
Incomplete outcome data (attrition bias) All outcomes	Low risk	Drop-outs balanced across intervention groups with similar reasons for with-drawal



Mahmud 2002 (Continued)		
Selective reporting (reporting bias)	Low risk	Expected outcomes were reported
Other bias	Low risk	The study appears to be free of other sources of bias

McIntyre 1988

Methods	Randomized, double-blind comparison of balsalazide and SASP. Extent of disease was initially classified by most recent barium enema, sigmoidoscopy and rectal biopsy as: extensive, substantial, left-sided, proctosigmoiditis, or proctitis. Evaluation continued throughout study at 4, 8, 16 and 24 weeks by clinical exam, sigmoidoscopy, and rectal biopsy (if deterioration occurred)
Participants	Adult patients (aged 18 to 79 years) with ulcerative colitis with clinical and sigmoidoscopic (non-friable mucosa) remission, and who were not currently on maintenance therapy, or who agreed to substitute trial capsules for maintenance SASP (N = 79)
Interventions	Balsalazide (n =41) or SASP, 2 g/day (n =38) in identical 500 mg capsules, for 6 months
Outcomes	Relapse was defined as the recurrence of previous symptoms. Patients completing 6 months without recurrence were classified as maintaining remission; the rest were considered to be treatment failures. Exclusions, withdrawals, and the occurrence of adverse side-effects were also recorded

Notes

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not described
Allocation concealment (selection bias)	Low risk	Adequate
Blinding (performance bias and detection bias) All outcomes	Low risk	Double-blind: identical capsules
Incomplete outcome data (attrition bias) All outcomes	Low risk	Drop-outs balanced across intervention groups with similar reasons for with-drawal
Selective reporting (reporting bias)	Low risk	Expected outcomes reported
Other bias	Low risk	The study appears to be free of other sources of bias

Miner 1995

Methods

Multicenter, randomized, double-blind, placebo-controlled and parallel design trial. Patients were stratified according to location of the disease. Clinical assessments were performed at entry, 1, 2, 3, 6, 9 and 12 months. At entry and final visit, sigmoidoscopy and histological assessments were performed. Patients were also required to keep daily diaries



Miner 1995 (Continued)			
Participants	Patients, 18 years or older, previously diagnosed with ulcerative colitis and in remission. Use of immunosuppressives or steroids required 90-day and 60-day wash-out periods, respectively (N =205)		
Interventions	Controlled-release mesalamine, in 250 mg capsules (Pentasa) 4 g/day (n = 103) or identical-appearing placebo (n = 102) for 48 weeks, distributed in blister packaging. Compliance was verified by pill count		
Outcomes	Recurrence of disease was defined as one of the following: 1) a sigmoidoscopic index of greater or equal to 5, and 5 or more trips to the toilet per day or rectal bleeding for 3 of 7 continuous days; 2) sigmoidoscopic index of 5 or more, with missing data for toilet visits or rectal bleeding; 3) missing data for final sigmoidoscopic index and early termination due to insufficient therapeutic effect		
Notes	Failure rate was calcula	ated as "1 - reported remission rate"	
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	Unclear risk	Not described	
Allocation concealment (selection bias)	Low risk	Adequate	
Blinding (performance bias and detection bias) All outcomes	Low risk	Identical placebo	
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	34 patients in the placebo dropped out due to adverse reaction compared to 14 in the mesalamine group	
Selective reporting (reporting bias)	Low risk	Expected outcomes were reported	
Other bias	Low risk	The study appears to be free of other sources of bias	
Mulder 1988			
Methods	Multicenter, randomized, double-blinded, double-dummy comparison of Pentasa and SASP. Evaluation at entry involved clinical, endoscopic, and histologic examinations. Follow-ups occurred at 3, 6, 9 and 12 months after entry		
Participants	Male and female outpatients, 18 years or older, with ulcerative colitis in remission for between 1 month and 5 years, who had not taken steroids or azathioprine during the month before entry (N = 78)		
Interventions	Six tablets (1.5 g) of Pentasa (5-ASA) plus 6 SASP-matched placebo tablets (n = 42) or 6 Pentasa-matched placebo tablets plus 6 tablets (3 g) of SASP daily (n = 36) for 12 months. Patients were to take medication in 3 doses before or after main meals. Placebo tablets were matched in appearance, weight and taste		
Outcomes	If the data obtained at each examination were assessed as 'normal' or 'in remission', the patient was		

considered to have remained in remission. If, at any time, the patient had not been considered 'normal' or 'in remission', then the patient was considered to have relapsed. Outcomes were reported in terms

of remission rates. Side effects and withdrawals were also recorded



Mulder 1988 (Continued)

Notes

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not described
Allocation concealment (selection bias)	Low risk	Adequate
Blinding (performance bias and detection bias) All outcomes	Low risk	Double-blind, double-dummy: placebo tablets were identical in appearance, weight and taste
Incomplete outcome data (attrition bias) All outcomes	Low risk	3 patients dropped out (2 from Pentasa group and 1 from SASP)
Selective reporting (reporting bias)	Low risk	Expected outcomes were reported
Other bias	Low risk	The study appears to be free of other sources of bias

Nilsson 1995

Methods	Multicenter, randomized, double-blind, double-dummy comparison of SASP and olsalazine. Clinical assessments were performed after every 3 months or more frequently if relapse was suspected. Rigid sigmoidoscopy was performed at entry and after 6 and 18 months	
Participants	Patients with ulcerative colitis in remission, with at least 2 episodes of active disease during the last 5 years. No other concomitant medications were permitted (N = 322)	
Interventions	Olsalazine, 1 g/day, with SASP-matched placebo (n =161) or active SASP, 2 g/day, with olsalazine-matched placebo (n = 161) for either 6, 9, 12, 15 or 18 months. Patients were instructed to medication with meals. Full treatment regimen was reached after 7 days. Compliance was verified pill counts	
Outcomes	Relapse was defined as macroscopic changes in the rectum of grade 3 or 4 on a 4-point scale. Other measures of efficacy were changes in number of stools per day, blood in stools, and consistency of stools. The failure rates were reported on an intention-to-treat basis	

Notes

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not described
Allocation concealment (selection bias)	Low risk	A - Adequate



Nilsson 1995 (Continued)		
Blinding (performance bias and detection bias) All outcomes	Low risk	Double-blind, double-dummy (matched placebos)
Incomplete outcome data (attrition bias) All outcomes	Low risk	Drop-outs balanced across intervention groups with similar reasons for with-drawal
Selective reporting (reporting bias)	Low risk	Expected outcomes were reported
Other bias	Low risk	The study appears to be free of other sources of bias

Paoluzi 2005

Methods	Single center, randomized, investigator-blind trial		
Participants	Adult patients (> 18 years) with confirmed ulcerative colitis in clinical, endoscopic and histological remission (N = 156)		
Interventions	Asacol 2.4 g/day (n = 80) or 1.2 g/day (n = 76) for 12 months		
Outcomes	Primary: maintenance of remission (Truelove and Witts). Secondary: time to relapse, adverse events		
Notes			

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not described
Allocation concealment (selection bias)	Unclear risk	Not described
Blinding (performance bias and detection bias) All outcomes	Low risk	Investigator-blinded
Incomplete outcome data (attrition bias) All outcomes	Low risk	Drop-outs balanced across intervention groups with similar reasons for with-drawal
Selective reporting (reporting bias)	Low risk	Expected outcomes were reported
Other bias	Low risk	The study appears to be free of other sources of bias

Pica 2012

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Pica 2012 (Continued)		
Participants	Patients with ulcerative colitis in remission (N = 112)	
Interventions	Asacol 4.8 g/day (n = 56) or 2.4 g/day (n = 56) for 12 months	
Outcomes	Remission at 12 months	
Notes	Abstract publication	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not described
Allocation concealment (selection bias)	Unclear risk	Not described
Blinding (performance bias and detection bias) All outcomes	High risk	Open label
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Drop-outs and reasons for withdrawal were not described
Selective reporting (reporting bias)	Unclear risk	Secondary outcomes were not described
Other bias	Unclear risk	Abstract publication
Prantera 2009		
Methods	Double-blind, double-dummy, parallel-group, randomized trial evaluating the efficacy and safety of MMX mesalazine dosed once daily (2.4 g/day) compared with Asacol (1.2 g twice daily) for the maintenance of clinical remission in patients with quiescent ulcerative colitis	
Participants	Adult patients (aged 18 to 75 years) in remission (modified UC-DAI score of \leq 1 supported by a rectal sigmoidoscopy in the preceding 3 months or colonoscopy in the preceding 6 months) for \geq 1 month prior to entry and had experienced at least one clinical or endoscopic relapse within the previous 12 months were recruited from centers in Italy, Poland and the Ukraine (N = 331)	
Interventions	MMX mesalazine 2.4 g/day (n = 162) given once daily or Asacol 2.4 g/day (n = 169) given in two divided doses of 1.2 g for 12 months	
Outcomes	The primary outcomes were the proportion of patients in clinical remission and the proportion of patients in clinical and endoscopic remission at 12 months. Secondary outcomes included the time to relapse (UC-DAI > 1) and assessment of the safety and tolerability of MMX mesalazine 2.4 g/day. Patients were assessed at screening, entry and at 3, 6, 9 and 12 months (or early withdrawal)	
Notes	Adherence was checked by tablet counts at each visit. Adherence was defined as taking ≥ 80% of the	

study medication



Prantera	2009	(Continued)
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Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer generated
Allocation concealment (selection bias)	Low risk	Centralized randomization
Blinding (performance bias and detection bias) All outcomes	Low risk	Double-blind, double-dummy
Incomplete outcome data (attrition bias) All outcomes	Low risk	Drop-outs balanced across intervention groups with similar reasons for withdrawal
Selective reporting (reporting bias)	Low risk	All expected outcomes reported
Other bias	Low risk	The study appears to be free of other sources of bias

Rijk 1992

Methods	Multicenter, randomized, double-blind comparison of SASP and olsalazine. Randomization used a standard variance allocation method to ensure equal distribution of prognostic factors among both treatments. In addition to patient diaries, clinical exam, blood counts and urinalysis were performed after 4, 12, 24, 36 and 48 weeks. Sigmoidoscopy and biopsy was performed at entry and 48 weeks to verify remission
Participants	Patients with ulcerative colitis in remission for no longer than 2 years, and with active colitis in the past (proven by endoscopy/biopsies) were recruited between Dec. 1985 and Jan. 1989. Remission was defined as the absence of clinical signs of inflammation (3 or fewer stools/day) and a normal mucosa on sigmoidoscopic assessment. Some patients had participated in a trial comparing olsalazine and SASP in active ulcerative colitis and had achieved remission after 6 to 12 weeks of treatment (N = 46)
Interventions	SASP, 4 g/day (n = 23), or olsalazine, 2 g/day (n = 23), for 48 weeks, full dosage being reached by the 5th day. Dose reduction to 3 g and 1.5 g, respectively, was allowed in case of adverse effects. Drugs were given in physically indistinguishable capsules in sealed plastic containers
Outcomes	Relapse was defined as a recurrence of symptoms (blood in stools, with or without diarrhea) and signs of inflammation at endoscopy, even without the presence of patient complaints. Adverse effects and withdrawals were also reported
Notes	

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not described
Allocation concealment (selection bias)	Low risk	Centralized randomization



Rijk 1992 (Continued)		
Blinding (performance bias and detection bias) All outcomes	Low risk	Double-blind: SASP and olsalazine were given in externally indistinguishable capsules
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	8 patients dropped out from the olsalazine group (3 due to diarrhea) compared to 4 in the SASP group
Selective reporting (reporting bias)	Low risk	Expected outcomes were reported
Other bias	Low risk	The study appears to be free of other sources of bias

Riley 1988

Methods	Randomized, double-blind, double-dummy, parallel group design comparing mesalazine (Asacol) and SASP. Randomization was governed by a centrally-held pharmacy code and medication was pre-packaged and equally and randomly allocated to each center. In addition to daily diary records, patients were evaluated at entry and at 4, 12, 24, 36 and 48 weeks for clinical exam, urine specimen, blood tests. At entry and at completion (or relapse), sigmoidoscopy and biopsy were performed	
Participants	Adult outpatients with chronic ulcerative colitis in clinical remission for a minimum of 1 month before entry, recruited from three hospitals in close geographic proximity. All had appearance of normal mucosa or only erythema upon entry. All had previously taken SASP maintenance therapy (N = 100)	
Interventions	SASP, min 2 g/day, plus mesalazine-matched placebo (n = 50) or mesalazine, min. 0.8 g/day plus SASP-matched placebo (n = 50) for 48 weeks. Compliance was verified by pill counts	
Outcomes	Relapse was indicated by symptomatic disease recurrence. Confirmation of worsened appearance of rectal mucosa was verified by sigmoidoscopy. Withdrawals and occurrence of side-effects were also reported	

Notes

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not described
Allocation concealment (selection bias)	Low risk	Randomization was governed by a centrally-held pharmacy code
Blinding (performance bias and detection bias) All outcomes	Low risk	Double-blind, double-dummy (matched placebo)
Incomplete outcome data (attrition bias) All outcomes	Low risk	12% of the SASP group withdrew (n = 6) compared to 4% of the 5-ASA group (n = 2)
Selective reporting (reporting bias)	Low risk	Expected outcomes were reported



Ril	ey 1	988	(Continued)
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Other bias	Low risk	The study appears to be free of other sources of bias	
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Rutgeerts 1989

Methods	Randomized, double-blind, double-dummy comparison of Claversal (5-ASA) and SASP. Clinical exam, including blood work and urinalysis was performed at entry, 1, 2, 3, 6, 9, and 12 months (or earlier for relapse). Colonoscopy was performed if symptoms recurred
Participants	Adult patients with inactive ulcerative colitis controlled (score of 4 or lower on a standard index) for 1 month pre-trial on a stable sulfasalazine dose of 1.5 to 2 g/day ($N = 334$)
Interventions	SASP, 1.5 to 2 g/day (n = 167), or coated 5-ASA (Claversal), 0.75 g/day (n = 167), dispensed according to a double-dummy protocol for 12 months
Outcomes	Relapse was defined as a clinical index of 6 or greater. Withdrawals and adverse events were reported
Notes	

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Drug supplies were labelled and randomized in blocks according to a predetermined computer-generated list
Allocation concealment (selection bias)	Low risk	Adequate
Blinding (performance bias and detection bias) All outcomes	Low risk	Double-blind, double-dummy
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	60 patients in the 5-ASA did not complete the study compared to 41 SASP patients. The reasons for withdrawal are not adequately described
Selective reporting (reporting bias)	Low risk	Expected outcomes were reported
Other bias	Low risk	The study appears to be free of other sources of bias

Sandberg-Gertzen 1986

Methods	Randomized, double-blind, placebo-controlled trial. Patients were evaluated (clinically, sigmoidoscopically, and histologically) upon entry and completion, plus by regular telephone contact
Participants	Patients, of any age, with ulcerative colitis who, after 6 months of ADS (Dipentum) therapy, were in remission (N = 102)
Interventions	Azodisal sodium (ADS, Dipentum), 1 g/day (n = 52), or identical-appearing placebo (16 mg riboflavin; n = 49) for 6 months. Compliance was tested by urine and serum analyses and pill count



Sandberg	-Gertzen	1986	(Continued)
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Outcomes	Relapse was defined as the occurrence of diarrhea with macroscopic blood, together with the finding
	of active inflammation on sigmoidoscopy

Notes One placebo patient was excluded from the ITT analysis for entry criteria violation

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not described
Allocation concealment (selection bias)	Low risk	Adequate
Blinding (performance bias and detection bias) All outcomes	Low risk	Double-blind: identical placebo
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	No drop-outs were reported
Selective reporting (reporting bias)	Low risk	Expected outcomes were reported
Other bias	Low risk	The study appears to be free of other sources of bias

Sandborn 2010

Methods	12-month, multi-centre, investigator-blinded, randomized, non-inferiority trial comparing the efficacy and safety of mesalamine (Asacol) 1.6 to 2.4 g/day administered once daily with mesalamine (Asacol) 1.6 to 2.4 g/day administered as a divided twice daily dose for the maintenance of clinical remission in patients with ulcerative colitis who have experienced a mild to moderate disease course
Participants	Adult patients (aged ≥ 18 years) with quiescent ulcerative colitis (Simple Clinical Colitis Activity Index ≤

2) of at least 3 months duration on a stable dose of Asacol ranging from 1.6 to 2.4 g/day were recruited from 193 sites in the USA, Puerto Rico and Canada for the study (n = 1023). Patients must have experienced at least one flare of ulcerative colitis within 18 months of entry (N = 1023)

Interventions Mesalamine (Asacol) 1.6 to 2.4 g/day dosed either once daily (n = 512) or twice daily (n = 511) for 12 months

Primary outcome was maintenance of remission at 6 months (SSCAI ≥ 5 defined relapse). Secondary outcomes included: patient defined remission index, adherence (MARS scale), patient satisfaction and preference with dosing regimen, serious adverse events, adverse events leading to withdrawal

Notes

Risk of bias

Outcomes

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not described



Sandborn 2010 (Continued)		
Allocation concealment (selection bias)	Low risk	The randomization was performed centrally via an interactive voice response system
Blinding (performance bias and detection bias) All outcomes	High risk	Investigator-blinded, blinded investigator assessed outcome
Incomplete outcome data (attrition bias) All outcomes	Low risk	Drop-outs balanced across intervention groups with similar reasons for with-drawal
Selective reporting (reporting bias)	Low risk	All expected outcomes reported
Other bias	Low risk	The study appears to be free of other sources of bias

Travis 1994

Methods	Two-center, randomized, parallel group dose-ranging study
Participants	Adult patients with ulcerative colitis in remission (N = 198) for three or more months. Remission was defined as no clinical symptoms of active disease and no signs of active inflammation on sigmoidoscopy (grade 0 or 1: normal or pink mucosa of quiescent colitis without visible vessels)
Interventions	Olsalazine at a dose of 0.5 g/day (n = 67), 1.0 g/day (n = 65) or 2.0 g/day (n = 62) for 12 months
Outcomes	Primary: relapse defined as an increase in bowel frequency with blood or mucus and evidence of active disease on sigmoidoscopy. Secondary: time in remission from start of treatment, laboratory measurements, adverse events
Notes	4 patients did not receive any study drugs and were excluded from the ITT analysis

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not described
Allocation concealment (selection bias)	Unclear risk	Not described
Blinding (performance bias and detection bias) All outcomes	Low risk	Double-blind: patients in the lower dose group took a physically indistinguishable placebo
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	19% of the 0.5 g/day group withdrew before the end of the study compared to 14% of the 1.0 g/day group and 24% of the 2.0 g/day group. Reasons for withdrawal are described but are not attributed to individual treatment groups
Selective reporting (reporting bias)	Low risk	Expected outcomes are reported
Other bias	Low risk	The study appears to be free of other sources of bias



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Methods	Phase III, double-blind, double-dummy, randomized, multicenter non-inferiority study
	Patients were assessed at entry and once every 4 weeks until week 52
Participants	Patients (15 to 64 years) with documented diagnosis of ulcerative colitis. All patients were in remission (defined as UC-DAI score ≤ 2 and rectal bleeding score of 0) at study entry, had experienced clinical relapse during the previous year, and were receiving ≤ 2.25 g of mesalazine or 4.5 g of salazosulfapyridine per day as maintenance therapy (N = 301)
Interventions	Prolonged-release oral mesalazine (Pentasa; Kyorin Pharmaceutical Co., Ltd., Tokyo, Japan) at doses of 1.5 to 2.25 g/d once daily (n = 152) or 3 times daily (n = 149) for 52 weeks
Outcomes	Primary outcome: remission at 52 weeks
	Secondary outcomes: duration of remission, UC-DAI at final assessment, adherence, adverse events

Notes

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not described
Allocation concealment (selection bias)	Unclear risk	Not described
Blinding (performance bias and detection bias) All outcomes	Low risk	Double-blind, double dummy design
Incomplete outcome data (attrition bias) All outcomes	Low risk	Drop-outs balanced across intervention groups with similar reasons for with-drawal
Selective reporting (reporting bias)	Unclear risk	Most expected outcomes were reported 12 serious adverse events were reported and a causal relationship with the study drug was ruled out - but the authors did not report to which group these patients belonged
Other bias	Low risk	The study appears to be free of other sources of bias

Wright 1993

Methods	Randomized, double-blind, placebo-controlled trial. Patients were assessed at entry, after 3 weeks, and after 3, 6, 9 and 12 months. Sigmoidoscopy was performed at each visit. Biopsy was taken at entry, 6 months and 1 year
Participants	Patients, ages 18 to 75 years, with inactive ulcerative colitis who were asymptomatic for not less than one week and not more than one month prior to entry (N = 101). Steroid or SASP therapy was stopped at least one week prior to entry



Wright 1993 (Continued)								
Interventions	Olsalazine 2 g/day (500	Olsalazine 2 g/day (500 mg capsules 4 times per day; $n = 49$) or matching placebo ($n = 52$) for 1 year						
Outcomes		Clinical relapse was defined as a relapse of diarrhea (with or without blood and mucous) thought by the attending physician to warrant introduction of rectal or oral corticosteroids						
Notes								
Risk of bias								
Bias	Authors' judgement	Support for judgement						
Random sequence generation (selection bias)	Unclear risk	Not described						
Allocation concealment (selection bias)	Low risk	Adequate						
Blinding (performance bias and detection bias) All outcomes	Low risk	Double-blind: matching placebo						
Incomplete outcome data (attrition bias) All outcomes	Low risk	3 patients from the placebo group did not complete the study (1 lost top follow up, 1 lost to compliance, 1 initial protocol exclusion)						
Selective reporting (reporting bias)	Low risk	Expected outcomes were reported						
Other bias	Low risk	The study appears to be free of other sources of bias						

UC-DAI = Ulcerative Colitis Disease Activity Index

Characteristics of excluded studies [ordered by study ID]

Study	Reason for exclusion
Bardazzi 1994	Trial compares intermittent dosing of 5-ASA (2.4 g/day for first week of the month) with continuous dosing of 5-ASA (1.6 g/day) for maintenance treatment
d'Albasio 1997	Trial compares the combination of oral 5-ASA and 5-ASA enemas with oral 5-ASA alone
Dew 1982b	Treatment duration of less than 6 months (16 weeks)
Eliakim 1990	Single-center report of patients who were abstracted from a larger multi-center trial (Rutgeerts 1989)
Ewe 1996	Treatment duration of less than 6 months (10 days)
Fernández-Bañares 1999	Trial compares oral mesalamine to Plantago ovata seeds
Frieri 2005	Not an RCT
Giaffer 1992b	Treatment duration of less than 6 months (3 months)
Gionchetti 1990	Treatment duration of less than 6 months



Study	Reason for exclusion						
Gionchetti 1996	Treatment duration of less than 6 months (2 weeks)						
Green 2004	Not a randomized controlled trial. No comparison group						
Hanauer 2009	Not a RCT - open-label extension study						
Karamanolis 1996	Treatment duration of less than 6 months (2 weeks)						
Kruis 1997	Trial compares oral mesalazine (Salofalk) to probiotic (Esherichia coli Nissle 1917)						
Kruis 2004	Trial compares oral mesalazine to probiotic (Esherichia coli Nissle 1917)						
Mani 1994	Treatment duration of less than 6 months (4 weeks)						
Mantzaris 2004	Trial compares azathioprine monotherapy to combination of azathioprine and olsalazine for maintenance of remission in patients with steroid-dependent ulcerative colitis						
Odes 1997	Not a RCT						
Osterman 2014	Treatment duration of less than 6 months (6 weeks)						
Paoluzi 2002	Trial looks at 4 weeks of combined oral and topical 5-ASA (mesalazine) versus 8 weeks of combined oral and topical 5-ASA (mesalazine)						
Papatheodoridis 1995	Treatment duration of less than 6 months (14 days)						
Pelech 1998	Trial compares oral mesalazine to probiotic (Esherichia coli Nissle 1917)						
Staerk Laursen 1990	Treatment duration of less than 6 months (28 days)						
Stoa-Birketvedt 1999	Treatment duration of less than 6 months (14 days)						
Tragnone 1996	Trial compares 800 mg 5-ASA tablets (total dose 1.6 g/day) with 400 mg 5-ASA tablets (total dose 1.6 g/day). Treatment duration of less than 6 months (3 months)						
Tzivras 1997	Treatment duration of less than 6 months (14 days)						
Yokoyama 2007	Trial compares oral 5-ASA with 5-ASA enemas						
Zocco 2006	Trial compares oral mesalazine to probiotic (Lactobacillus GG) to combination of oral mesalazine + probiotic						

DATA AND ANALYSES



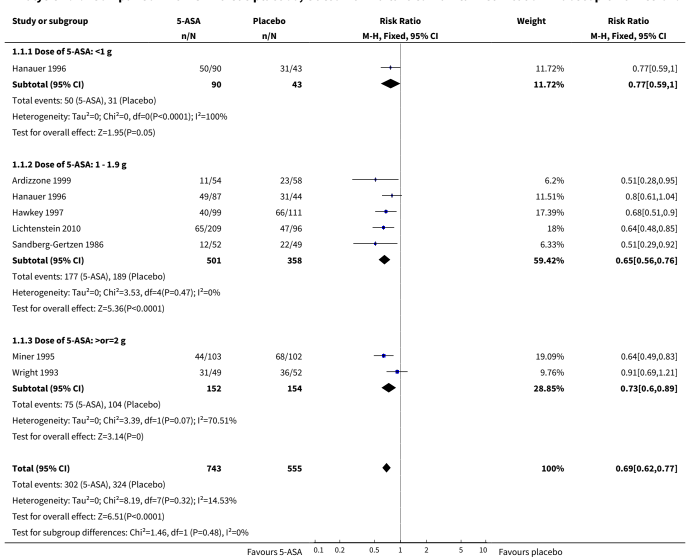
Comparison 1. 5-ASA versus placebo

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	0.69 [0.62, 0.77]	
1 Failure to Maintain Clinical or Endoscopic Remission	7	1298	Risk Ratio (M-H, Fixed, 95% CI)		
1.1 Dose of 5-ASA: <1 g	1	133	Risk Ratio (M-H, Fixed, 95% CI)	0.77 [0.59, 1.00]	
1.2 Dose of 5-ASA: 1 - 1.9 g	5	859	Risk Ratio (M-H, Fixed, 95% CI)	0.65 [0.56, 0.76]	
1.3 Dose of 5-ASA: >or=2 g	2	306	Risk Ratio (M-H, Fixed, 95% CI)	0.73 [0.60, 0.89]	
2 Development of Any Adverse Event	4	875	Risk Ratio (M-H, Random, 95% CI)	0.98 [0.69, 1.39]	
2.1 Dose of 5-ASA: <1 g	1	133	Risk Ratio (M-H, Random, 95% CI)	0.82 [0.51, 1.31]	
2.2 Dose of 5-ASA: 1 - 1.9 g	2	436	Risk Ratio (M-H, Random, 95% CI)	1.02 [0.86, 1.20]	
2.3 Dose of 5-ASA: >or=2 g	2	306	Risk Ratio (M-H, Random, 95% CI)	1.69 [0.14, 20.58]	
3 Development of Any Adverse Event (Sensitivity analysis)	3	774	Risk Ratio (M-H, Random, 95% CI)	0.94 [0.77, 1.15]	
3.1 Dose of 5-ASA: <1 g	1	133	Risk Ratio (M-H, Random, 95% CI)	0.82 [0.51, 1.31]	
3.2 Dose of 5-ASA: 1 - 1.9 g	2	436	Risk Ratio (M-H, Random, 95% CI)	1.02 [0.86, 1.20]	
3.3 Dose of 5-ASA: >or=2 g	1	205	Risk Ratio (M-H, Random, 95% CI)	0.52 [0.25, 1.12]	
4 Withdrawal from Study due to Adverse Event	6	1197	Risk Ratio (M-H, Fixed, 95% CI)	1.34 [0.78, 2.30]	
4.1 Dose of 5-ASA: <1 g	1	133	Risk Ratio (M-H, Fixed, 95% CI)	1.43 [0.15, 13.38]	
4.2 Dose of 5-ASA: 1 - 1.9 g	4	758	Risk Ratio (M-H, Fixed, 95% CI)	1.05 [0.50, 2.23]	
4.3 Dose of 5-ASA: >or=2 g	2	306	Risk Ratio (M-H, Fixed, 95% CI)	1.80 [0.78, 4.15]	
5 Withdrawal from Study due to Adverse Event (Sensitivity analysis)	5	1096	Risk Ratio (M-H, Fixed, 95% CI)	0.86 [0.46, 1.63]	
5.1 Dose of 5-ASA: <1 g	1	133	Risk Ratio (M-H, Fixed, 95% CI)	1.43 [0.15, 13.38]	
5.2 Dose of 5-ASA: 1 - 1.9 g	4	758	Risk Ratio (M-H, Fixed, 95% CI)	1.05 [0.50, 2.23]	
5.3 Dose of 5-ASA: >or=2 g	1	205	Risk Ratio (M-H, Fixed, 95% CI)	0.33 [0.07, 1.60]	
6 Exclusion/Withdrawal after Entry (not due to relapse)	5	1074	Risk Ratio (M-H, Fixed, 95% CI) 1.13 [0.88, 1.4		
6.1 Dose of 5-ASA: <1 g	1	177	Risk Ratio (M-H, Fixed, 95% CI)	0.90 [0.58, 1.40]	



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
6.2 Dose of 5-ASA: 1 - 1.9 g	3	591	Risk Ratio (M-H, Fixed, 95% CI)	1.22 [0.87, 1.71]
6.3 Dose of 5-ASA: >or=2 g	2	306	Risk Ratio (M-H, Fixed, 95% CI)	1.26 [0.69, 2.29]

Analysis 1.1. Comparison 1 5-ASA versus placebo, Outcome 1 Failure to Maintain Clinical or Endoscopic Remission.





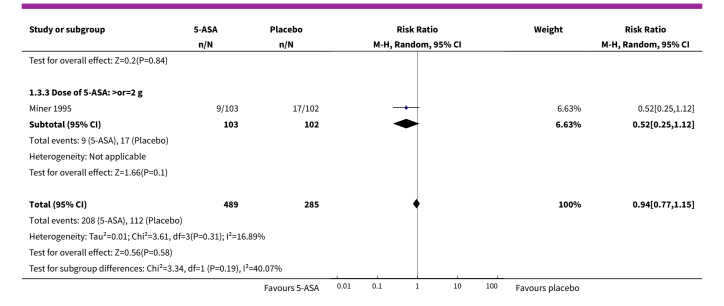
Analysis 1.2. Comparison 1 5-ASA versus placebo, Outcome 2 Development of Any Adverse Event.

Study or subgroup	5-ASA	Placebo	Risk Ratio	Weight	Risk Ratio	
	n/N	n/N	M-H, Random, 95% CI		M-H, Random, 95% CI	
1.2.1 Dose of 5-ASA: <1 g						
Hanauer 1996	29/90	17/43		22.54%	0.82[0.51,1.31]	
Subtotal (95% CI)	90	43	*	22.54%	0.82[0.51,1.31]	
Total events: 29 (5-ASA), 17 (Placebo)						
Heterogeneity: Not applicable						
Test for overall effect: Z=0.84(P=0.4)						
1.2.2 Dose of 5-ASA: 1 - 1.9 g						
Hanauer 1996	36/87	17/44	<u> </u>	23.64%	1.07[0.68,1.68]	
Lichtenstein 2010	134/209	61/96	•	35.11%	1.01[0.84,1.21]	
Subtotal (95% CI)	296	140	\oddsymbol{\displaysim}	58.75%	1.02[0.86,1.2]	
Total events: 170 (5-ASA), 78 (Placebo)						
Heterogeneity: Tau ² =0; Chi ² =0.06, df=1(F	P=0.8); I ² =0%					
Test for overall effect: Z=0.2(P=0.84)						
1.2.3 Dose of 5-ASA: >or=2 g						
Miner 1995	9/103	17/102		13.65%	0.52[0.25,1.12]	
Wright 1993	12/49	2/52		5.06%	6.37[1.5,27.01]	
Subtotal (95% CI)	152	154		18.71%	1.69[0.14,20.58]	
Total events: 21 (5-ASA), 19 (Placebo)						
Heterogeneity: Tau ² =2.91; Chi ² =9.39, df=	:1(P=0); I ² =89.35%					
Test for overall effect: Z=0.41(P=0.68)						
Total (95% CI)	538	337	•	100%	0.98[0.69,1.39]	
Total events: 220 (5-ASA), 114 (Placebo)						
Heterogeneity: Tau ² =0.08; Chi ² =9.9, df=4	(P=0.04); I ² =59.58 ^o	%				
Test for overall effect: Z=0.12(P=0.91)						
Test for subgroup differences: Chi ² =0.92	, df=1 (P=0.63), I ² =0	0%				
		Favours 5-ASA 0.	01 0.1 1 10 100) Favours placebo		

Analysis 1.3. Comparison 1 5-ASA versus placebo, Outcome 3 Development of Any Adverse Event (Sensitivity analysis).

Study or subgroup	5-ASA	Placebo			Risk Ratio			Weight	Risk Ratio	
	n/N	n/N	M-H, Random,			95% CI			M-H, Random, 95% CI	
1.3.1 Dose of 5-ASA: <1 g										
Hanauer 1996	29/90	17/43			+			15.58%	0.82[0.51,1.31]	
Subtotal (95% CI)	90	43			•			15.58%	0.82[0.51,1.31]	
Total events: 29 (5-ASA), 17 (Placebo)										
Heterogeneity: Not applicable										
Test for overall effect: Z=0.84(P=0.4)										
1.3.2 Dose of 5-ASA: 1 - 1.9 g										
Hanauer 1996	36/87	17/44			+			17.25%	1.07[0.68,1.68]	
Lichtenstein 2010	134/209	61/96			•			60.54%	1.01[0.84,1.21]	
Subtotal (95% CI)	296	140			•			77.79%	1.02[0.86,1.2]	
Total events: 170 (5-ASA), 78 (Placebo)										
Heterogeneity: Tau ² =0; Chi ² =0.06, df=1(P=0.8); I ² =0%		_							
		Favours 5-ASA	0.01	0.1	1	10	100	Favours placebo		





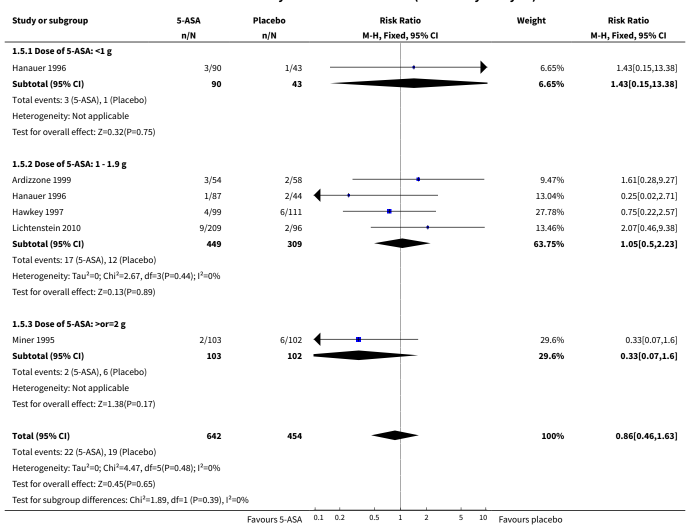
Analysis 1.4. Comparison 1 5-ASA versus placebo, Outcome 4 Withdrawal from Study due to Adverse Event.

Study or subgroup	5-ASA	Placebo	Risk Ratio	Weight	Risk Ratio M-H, Fixed, 95% CI
	n/N	n/N	M-H, Fixed, 95% CI		
1.4.1 Dose of 5-ASA: <1 g					
Hanauer 1996	3/90	1/43	+	6.07%	1.43[0.15,13.38]
Subtotal (95% CI)	90	43		6.07%	1.43[0.15,13.38]
Total events: 3 (5-ASA), 1 (Placebo)					
Heterogeneity: Not applicable					
Test for overall effect: Z=0.32(P=0.75)					
1.4.2 Dose of 5-ASA: 1 - 1.9 g					
Ardizzone 1999	3/54	2/58	-	8.65%	1.61[0.28,9.27]
Hanauer 1996	1/87	2/44	+	11.91%	0.25[0.02,2.71]
Hawkey 1997	4/99	6/111		25.36%	0.75[0.22,2.57]
Lichtenstein 2010	9/209	2/96	+	12.29%	2.07[0.46,9.38]
Subtotal (95% CI)	449	309		58.2%	1.05[0.5,2.23]
Total events: 17 (5-ASA), 12 (Placebo)					
Heterogeneity: Tau ² =0; Chi ² =2.67, df=3	(P=0.44); I ² =0%				
Test for overall effect: Z=0.13(P=0.89)					
1.4.3 Dose of 5-ASA: >or=2 g					
Miner 1995	2/103	6/102	-	27.03%	0.33[0.07,1.6]
Wright 1993	12/49	2/52	· · · · · · · · · · · · · · · · · · ·	8.7%	6.37[1.5,27.01]
Subtotal (95% CI)	152	154		35.73%	1.8[0.78,4.15]
Total events: 14 (5-ASA), 8 (Placebo)					
Heterogeneity: Tau ² =0; Chi ² =7.38, df=1	(P=0.01); I ² =86.45%				
Test for overall effect: Z=1.38(P=0.17)					
Total (95% CI)	691	506		100%	1.34[0.78,2.3]
Total events: 34 (5-ASA), 21 (Placebo)					
Heterogeneity: Tau ² =0; Chi ² =10.62, df=	6(P=0.1); I ² =43.5%				
Test for overall effect: Z=1.07(P=0.28)			İ		



Study or subgroup	5-ASA n/N	Placebo n/N				sk Rat	tio 95% CI			Weight	Risk Ratio M-H, Fixed, 95% CI
Test for subgroup differences: Chi²=0.88, df=1 (P=0.65), I²=0%											
		Favours 5-ASA	0.1	0.2	0.5	1	2	5	10	Favours placebo	

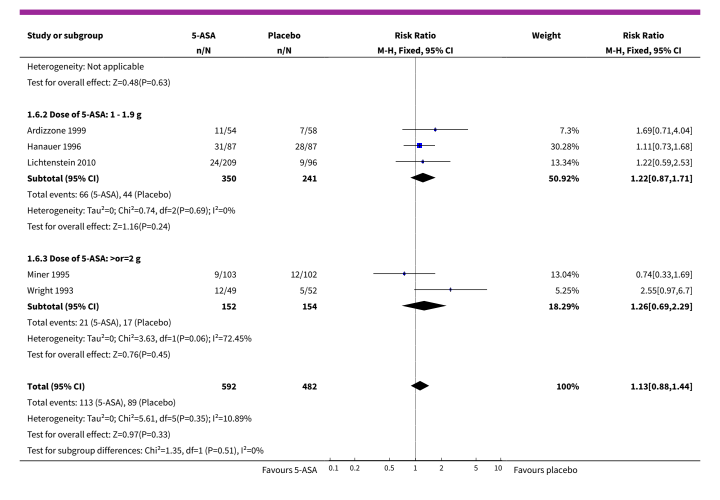
Analysis 1.5. Comparison 1 5-ASA versus placebo, Outcome 5 Withdrawal from Study due to Adverse Event (Sensitivity analysis).



Analysis 1.6. Comparison 15-ASA versus placebo, Outcome 6 Exclusion/Withdrawal after Entry (not due to relapse).

Study or subgroup	5-ASA	Placebo	Risk Ratio				Weight	Risk Ratio
	n/N	n/N		M-H, Fixed, 95%	CI			M-H, Fixed, 95% CI
1.6.1 Dose of 5-ASA: <1 g								
Hanauer 1996	26/90	28/87					30.79%	0.9[0.58,1.4]
Subtotal (95% CI)	90	87		*			30.79%	0.9[0.58,1.4]
Total events: 26 (5-ASA), 28 (Placebo)								
		Favours 5-ASA	0.1 0.2	0.5 1 2	5	10	Favours placebo	



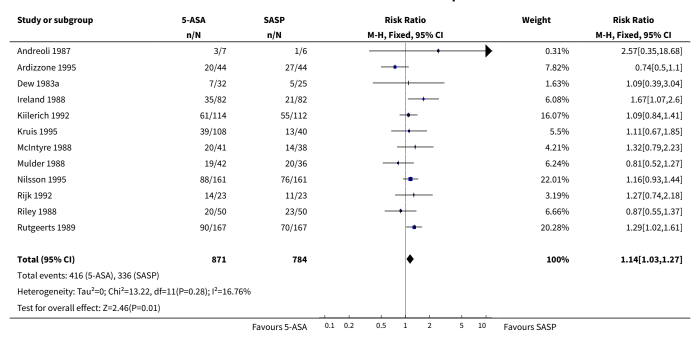


Comparison 2. 5-ASA versus sulfasalazine

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Failure to Maintain Clinical or Endoscopic Remission	12	1655	Risk Ratio (M-H, Fixed, 95% CI)	1.14 [1.03, 1.27]
2 Failure to Maintain Remission (trials without olsalazine)	7	749	Risk Ratio (M-H, Fixed, 95% CI)	1.08 [0.92, 1.26]
3 Development of Any Adverse Event	7	1138	Risk Ratio (M-H, Fixed, 95% CI)	1.07 [0.82, 1.40]
4 Withdrawal from Study due to Adverse Event	10	1585	Risk Ratio (M-H, Fixed, 95% CI)	1.27 [0.87, 1.87]
5 Exclusion/Withdrawal after Entry (not due to relapse)	9	1497	Risk Ratio (M-H, Fixed, 95% CI)	1.30 [1.04, 1.63]



Analysis 2.1. Comparison 2 5-ASA versus sulfasalazine, Outcome 1 Failure to Maintain Clinical or Endoscopic Remission.



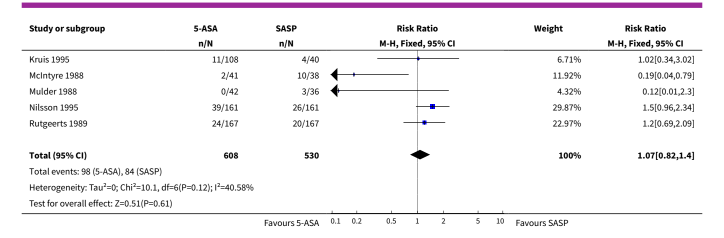
Analysis 2.2. Comparison 2 5-ASA versus sulfasalazine, Outcome 2 Failure to Maintain Remission (trials without olsalazine).

Study or subgroup	5-ASA	SASP			Ri	sk Ratio			Weight	Risk Ratio
	n/N	n/N			M-H, F	ixed, 95%	6 CI		M-H, Fixed, 95%	M-H, Fixed, 95% CI
Andreoli 1987	3/7	1/6					•	→	0.66%	2.57[0.35,18.68]
Ardizzone 1995	20/44	27/44			_	•+			16.59%	0.74[0.5,1.1]
Dew 1983a	7/32	5/25				+			3.45%	1.09[0.39,3.04]
McIntyre 1988	20/41	14/38				+	_		8.93%	1.32[0.79,2.23]
Mulder 1988	19/42	20/36			_	+			13.23%	0.81[0.52,1.27]
Riley 1988	20/50	23/50				+			14.13%	0.87[0.55,1.37]
Rutgeerts 1989	90/167	70/167							43.01%	1.29[1.02,1.61]
Total (95% CI)	383	366				•			100%	1.08[0.92,1.26]
Total events: 179 (5-ASA), 160 (SASP)										
Heterogeneity: Tau ² =0; Chi ² =9.45, df=6	(P=0.15); I ² =36.48%									
Test for overall effect: Z=0.96(P=0.34)										
		Favours 5-ASA	0.1	0.2	0.5	1 :	2 5	10	Favours SASP	

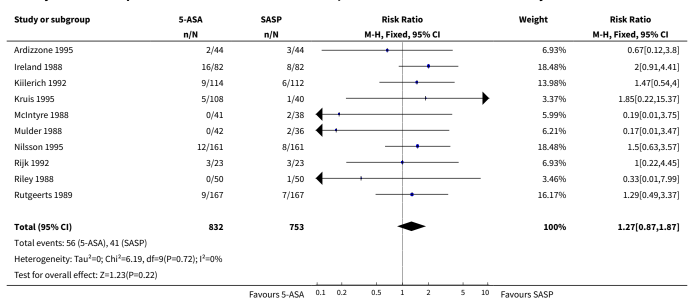
Analysis 2.3. Comparison 2 5-ASA versus sulfasalazine, Outcome 3 Development of Any Adverse Event.

Study or subgroup	5-ASA	SASP			Ri	sk Rat	io			Weight	Risk Ratio
	n/N	n/N			M-H, F	ixed, 9	95% CI				M-H, Fixed, 95% CI
Andreoli 1987	1/7	1/6	+						→	1.24%	0.86[0.07,10.96]
Ireland 1988	21/82	20/82			_	+	_			22.97%	1.05[0.62,1.78]
		Favours 5-ASA	0.1	0.2	0.5	1	2	5	10	Favours SASP	





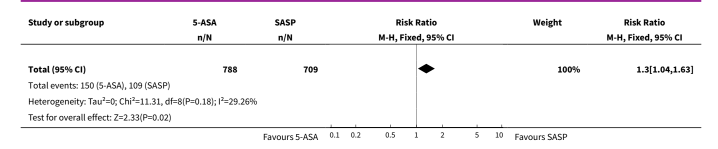
Analysis 2.4. Comparison 2 5-ASA versus sulfasalazine, Outcome 4 Withdrawal from Study due to Adverse Event.



Analysis 2.5. Comparison 2 5-ASA versus sulfasalazine, Outcome 5 Exclusion/Withdrawal after Entry (not due to relapse).

Study or subgroup	5-ASA	5-ASA SASP Risk Ratio		Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
Ireland 1988	19/82	11/82		9.94%	1.73[0.88,3.4]
Kiilerich 1992	16/114	13/112		11.85%	1.21[0.61,2.4]
Kruis 1995	12/108	2/40	- 	2.64%	2.22[0.52,9.5]
McIntyre 1988	3/41	6/38		5.63%	0.46[0.12,1.72]
Mulder 1988	1/42	5/36	—	4.87%	0.17[0.02,1.4]
Nilsson 1995	29/161	21/161	+-	18.98%	1.38[0.82,2.32]
Rijk 1992	8/23	4/23		3.62%	2[0.7,5.73]
Riley 1988	2/50	6/50	—	5.42%	0.33[0.07,1.57]
Rutgeerts 1989	60/167	41/167		37.05%	1.46[1.05,2.04]
		Favours 5-ASA	0.1 0.2 0.5 1 2 5 1	0 Favours SASP	



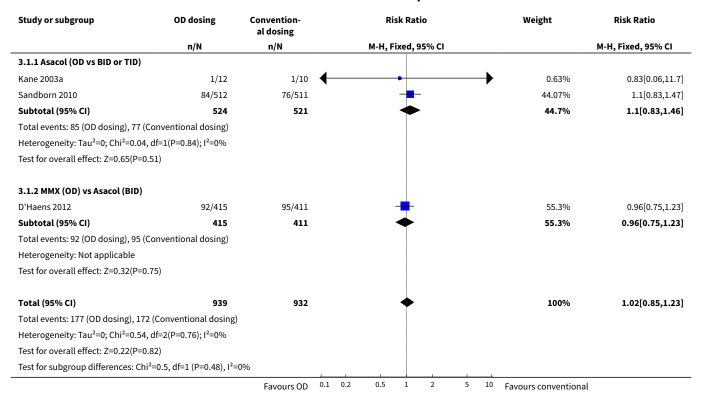


Comparison 3. Once daily versus conventional dosing

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Failure to Maintain Clinical or Endo- scopic Remission at 6 months	3	1871	Risk Ratio (M-H, Fixed, 95% CI)	1.02 [0.85, 1.23]
1.1 Asacol (OD vs BID or TID)	2	1045	Risk Ratio (M-H, Fixed, 95% CI)	1.10 [0.83, 1.46]
1.2 MMX (OD) vs Asacol (BID)	1	826	Risk Ratio (M-H, Fixed, 95% CI)	0.96 [0.75, 1.23]
2 Failure to Maintain Clinical or Endo- scopic Remission at 12 months	8	3127	Risk Ratio (M-H, Fixed, 95% CI)	0.91 [0.82, 1.01]
2.1 Asacol (OD vs BID or TID)	3	1256	Risk Ratio (M-H, Fixed, 95% CI)	0.95 [0.79, 1.15]
2.2 MMX (OD) vs Asacol (BID)	1	331	Risk Ratio (M-H, Fixed, 95% CI)	0.99 [0.74, 1.33]
2.3 Pentasa (OD vs BID)	2	654	Risk Ratio (M-H, Fixed, 95% CI)	0.75 [0.60, 0.93]
2.4 MMX (OD vs BID)	1	451	Risk Ratio (M-H, Fixed, 95% CI)	1.13 [0.87, 1.47]
2.5 Salofalk granules (OD vs TID)	1	435	Risk Ratio (M-H, Fixed, 95% CI)	0.81 [0.60, 1.10]
3 Failure to adhere to study medication regimen at study endpoint	8	2126	Risk Ratio (M-H, Random, 95% CI)	1.18 [0.69, 2.03]
4 Failure to adhere to study medication regimen (Sensitivity analysis - excluding outliers)	6	1462	Risk Ratio (M-H, Fixed, 95% CI)	1.22 [0.91, 1.64]
5 Development of Any Adverse Event	6	2714	Risk Ratio (M-H, Fixed, 95% CI)	1.00 [0.92, 1.08]
6 Withdrawal due to adverse event	7	3737	Risk Ratio (M-H, Fixed, 95% CI)	1.31 [0.80, 2.13]
7 Exclusion/Withdrawal after Entry (not due to relapse)	7	3737	Risk Ratio (M-H, Fixed, 95% CI)	0.99 [0.85, 1.15]



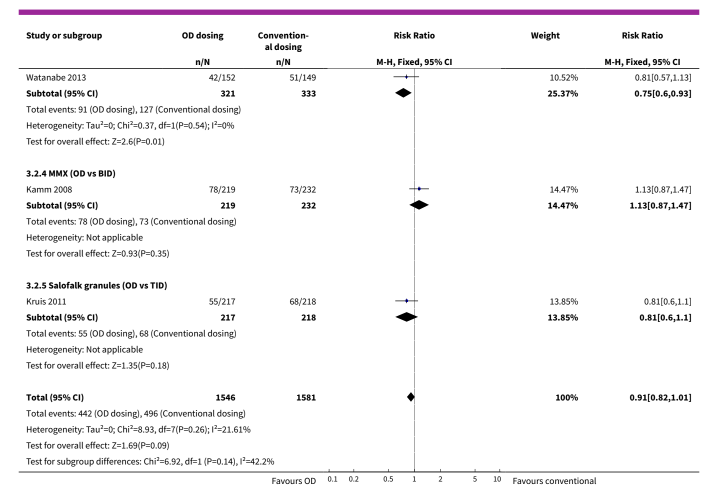
Analysis 3.1. Comparison 3 Once daily versus conventional dosing, Outcome 1 Failure to Maintain Clinical or Endoscopic Remission at 6 months.



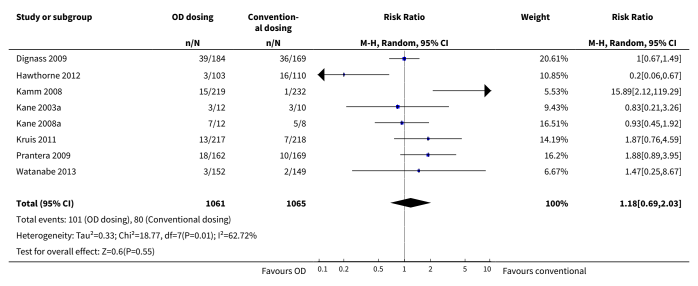
Analysis 3.2. Comparison 3 Once daily versus conventional dosing, Outcome 2 Failure to Maintain Clinical or Endoscopic Remission at 12 months.

Study or subgroup	OD dosing	Convention- al dosing	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
3.2.1 Asacol (OD vs BID or TID)					
Hawthorne 2012	23/103	33/110		6.52%	0.74[0.47,1.18]
Kane 2008a	6/12	5/8		1.22%	0.8[0.37,1.74]
Sandborn 2010	133/512	131/511	+	26.77%	1.01[0.82,1.25]
Subtotal (95% CI)	627	629	*	34.51%	0.95[0.79,1.15]
Total events: 162 (OD dosing), 16	9 (Conventional dosing)				
Heterogeneity: Tau ² =0; Chi ² =1.64	I, df=2(P=0.44); I ² =0%				
Test for overall effect: Z=0.49(P=0	0.62)				
3.2.2 MMX (OD) vs Asacol (BID)					
Prantera 2009	56/162	59/169	-	11.79%	0.99[0.74,1.33]
Subtotal (95% CI)	162	169	*	11.79%	0.99[0.74,1.33]
Total events: 56 (OD dosing), 59 (Conventional dosing)				
Heterogeneity: Not applicable					
Test for overall effect: Z=0.07(P=0	0.95)				
3.2.3 Pentasa (OD vs BID)					
Dignass 2009	49/169	76/184		14.86%	0.7[0.52,0.94]
		Favours OD 0.1	0.2 0.5 1 2 5	10 Favours convention	al





Analysis 3.3. Comparison 3 Once daily versus conventional dosing, Outcome 3 Failure to adhere to study medication regimen at study endpoint.





Analysis 3.4. Comparison 3 Once daily versus conventional dosing, Outcome 4 Failure to adhere to study medication regimen (Sensitivity analysis - excluding outliers).

Study or subgroup	OD dosing	Convention- al dosing	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
Dignass 2009	39/184	36/169		57.21%	1[0.67,1.49]
Kane 2003a	3/12	3/10		4.99%	0.83[0.21,3.26]
Kane 2008a	7/12	5/8		9.15%	0.93[0.45,1.92]
Kruis 2011	13/217	7/218		10.65%	1.87[0.76,4.59]
Prantera 2009	18/162	10/169	 	14.92%	1.88[0.89,3.95]
Watanabe 2013	3/152	2/149		3.08%	1.47[0.25,8.67]
Total (95% CI)	739	723	•	100%	1.22[0.91,1.64]
Total events: 83 (OD dosing),	63 (Conventional dosing)				
Heterogeneity: Tau ² =0; Chi ² =4	1.02, df=5(P=0.55); I ² =0%				
Test for overall effect: Z=1.33(P=0.18)				
		Favours OD	0.1 0.2 0.5 1 2 5	Favours conventional	

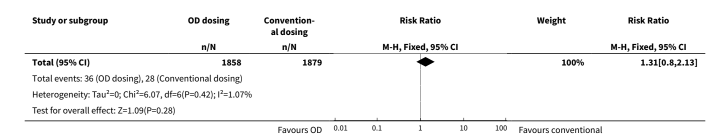
Analysis 3.5. Comparison 3 Once daily versus conventional dosing, Outcome 5 Development of Any Adverse Event.

Study or subgroup	OD dosing	Convention- al dosing		Risk Ratio	Weight	Risk Ratio
	n/N	n/N		M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
D'Haens 2012	154/415	148/411		+	24.16%	1.03[0.86,1.23]
Dignass 2009	75/175	68/187		+-	10.68%	1.18[0.91,1.52]
Kamm 2008	88/225	86/234		+	13.7%	1.06[0.84,1.34]
Kruis 2011	89/217	105/218		-+-	17.02%	0.85[0.69,1.05]
Prantera 2009	92/162	99/169		+	15.74%	0.97[0.81,1.17]
Watanabe 2013	110/152	114/149		+	18.7%	0.95[0.83,1.08]
Total (95% CI)	1346	1368		•	100%	1[0.92,1.08]
Total events: 608 (OD dosing)	, 620 (Conventional dosing)					
Heterogeneity: Tau ² =0; Chi ² =4	1.89, df=5(P=0.43); I ² =0%					
Test for overall effect: Z=0.12(P=0.9)					
		Favours OD	0.1 0.2	0.5 1 2	5 10 Favours conventiona	nl

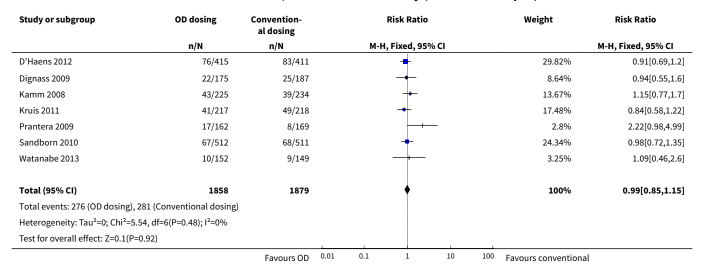
Analysis 3.6. Comparison 3 Once daily versus conventional dosing, Outcome 6 Withdrawal due to adverse event.

Study or subgroup	OD dosing	Convention- al dosing		Risk Ratio			Weight	Risk Ratio	
	n/N	n/N		M-H	, Fixed, 95 ⁹	% CI			M-H, Fixed, 95% CI
D'Haens 2012	6/415	3/411			-			10.86%	1.98[0.5,7.87]
Dignass 2009	5/175	1/187			-	+	_	3.48%	5.34[0.63,45.28]
Kamm 2008	11/225	9/234			-			31.78%	1.27[0.54,3.01]
Kruis 2011	5/217	3/218			+			10.78%	1.67[0.41,6.92]
Prantera 2009	3/162	3/169		_		_		10.58%	1.04[0.21,5.09]
Sandborn 2010	2/512	7/511						25.24%	0.29[0.06,1.37]
Watanabe 2013	4/152	2/149			+			7.28%	1.96[0.36,10.54]
		Favours OD	0.01	0.1	1	10	100	Favours conventional	





Analysis 3.7. Comparison 3 Once daily versus conventional dosing, Outcome 7 Exclusion/Withdrawal after Entry (not due to relapse).



Comparison 4. 5-ASA versus comparator 5-ASA

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Failure to Maintain Clinical or Endoscopic Remission at 12 months	6	707	Risk Ratio (M-H, Fixed, 95% CI)	1.08 [0.91, 1.28]
1.1 Asacol comparator	5	615	Risk Ratio (M-H, Fixed, 95% CI)	1.05 [0.87, 1.26]
1.2 Salofalk comparator	1	92	Risk Ratio (M-H, Fixed, 95% CI)	1.30 [0.86, 1.98]
2 Development of Any Adverse Event	4	357	Risk Ratio (M-H, Fixed, 95% CI)	0.94 [0.83, 1.07]
2.1 Asacol comparator	3	265	Risk Ratio (M-H, Fixed, 95% CI)	0.96 [0.85, 1.08]
2.2 Salofalk comparator	1	92	Risk Ratio (M-H, Fixed, 95% CI)	0.83 [0.51, 1.34]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
3 Withdrawal from Study due to Adverse Event	5	457	Risk Ratio (M-H, Fixed, 95% CI)	1.25 [0.56, 2.78]
3.1 Asacol comparator	4	365	Risk Ratio (M-H, Fixed, 95% CI)	1.64 [0.61, 4.42]
3.2 Salofalk comparator	1	92	Risk Ratio (M-H, Fixed, 95% CI)	0.69 [0.16, 2.90]
4 Exclusion/Withdrawal after Entry (not due to relapse)	5	457	Risk Ratio (M-H, Fixed, 95% CI)	1.23 [0.90, 1.70]
4.1 Asacol comparator	4	365	Risk Ratio (M-H, Fixed, 95% CI)	1.21 [0.81, 1.80]
4.2 Salofalk comparator	1	92	Risk Ratio (M-H, Fixed, 95% CI)	1.28 [0.76, 2.16]

Analysis 4.1. Comparison 4 5-ASA versus comparator 5-ASA, Outcome 1 Failure to Maintain Clinical or Endoscopic Remission at 12 months.

Study or subgroup	5-ASA for- mulations	5-ASA com- parator	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
4.1.1 Asacol comparator					
Courtney 1992	13/50	23/50		15.98%	0.57[0.32,0.99]
Deventer 2001	71/126	59/124	+=-	41.32%	1.18[0.93,1.51]
Green 1998	21/49	20/46		14.33%	0.99[0.62,1.56]
Ito 2010	20/65	15/65	+-	10.42%	1.33[0.75,2.37]
Mahmud 2002	6/20	6/20		4.17%	1[0.39,2.58]
Subtotal (95% CI)	310	305	*	86.23%	1.05[0.87,1.26]
Total events: 131 (5-ASA formulat	ions), 123 (5-ASA compa	arator)			
Heterogeneity: Tau ² =0; Chi ² =6.5, o	df=4(P=0.16); I ² =38.44%				
Test for overall effect: Z=0.47(P=0.	.64)				
4.1.2 Salofalk comparator					
Kruis 2001	27/48	19/44	++-	13.77%	1.3[0.86,1.98]
Subtotal (95% CI)	48	44	*	13.77%	1.3[0.86,1.98]
Total events: 27 (5-ASA formulation	ons), 19 (5-ASA compara	tor)			
Heterogeneity: Not applicable					
Test for overall effect: Z=1.23(P=0.	.22)				
Total (95% CI)	358	349	•	100%	1.08[0.91,1.28]
Total events: 158 (5-ASA formulat	ions), 142 (5-ASA compa	arator)			
Heterogeneity: Tau ² =0; Chi ² =7.23,	, df=5(P=0.2); I ² =30.84%				
Test for overall effect: Z=0.9(P=0.3	37)				
Test for subgroup differences: Chi	i ² =0.88, df=1 (P=0.35), I ²	=0%			
Test for subgroup differences: Chi		-ASA formulations 0.1	0.2 0.5 1 2 5	10 Comparator formula	ation



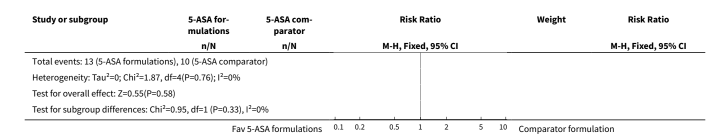
Analysis 4.2. Comparison 4 5-ASA versus comparator 5-ASA, Outcome 2 Development of Any Adverse Event.

Study or subgroup	5-ASA for- mulations	5-ASA com- parator	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
4.2.1 Asacol comparator					
Green 1998	30/49	30/46	_	25.4%	0.94[0.69,1.28]
Ito 2010	62/65	62/65	•	50.9%	1[0.93,1.08]
Mahmud 2002	6/20	8/20		6.57%	0.75[0.32,1.77]
Subtotal (95% CI)	134	131	+	82.87%	0.96[0.85,1.08]
Total events: 98 (5-ASA formulations)	, 100 (5-ASA compa	rator)			
Heterogeneity: Tau ² =0; Chi ² =1.38, df=	=2(P=0.5); I ² =0%				
Test for overall effect: Z=0.64(P=0.52)					
4.2.2 Salofalk comparator					
Kruis 2001	18/48	20/44		17.13%	0.83[0.51,1.34]
Subtotal (95% CI)	48	44		17.13%	0.83[0.51,1.34]
Total events: 18 (5-ASA formulations)	, 20 (5-ASA compara	ator)			
Heterogeneity: Not applicable					
Test for overall effect: Z=0.77(P=0.44)					
Total (95% CI)	182	175	•	100%	0.94[0.83,1.07]
Total events: 116 (5-ASA formulations	s), 120 (5-ASA comp	arator)			
Heterogeneity: Tau ² =0; Chi ² =3.27, df=	=3(P=0.35); I ² =8.38%				
Test for overall effect: Z=0.98(P=0.33)					
Test for subgroup differences: Chi ² =0	.36, df=1 (P=0.55), I ²	=0%			
	Fav 5	-ASA formulations 0.1	0.2 0.5 1 2 5	10 Comparator formula	ition

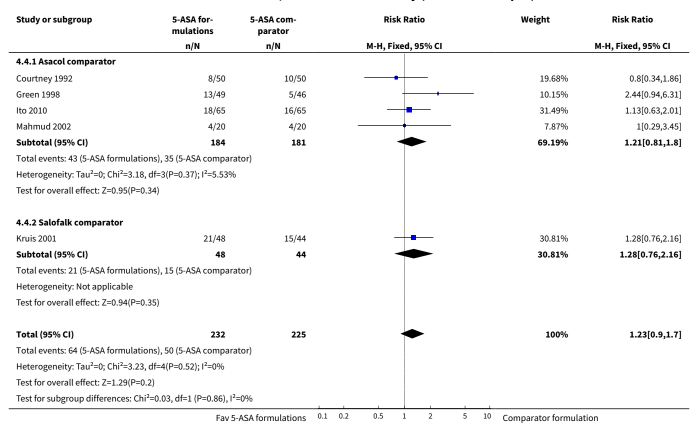
Analysis 4.3. Comparison 4 5-ASA versus comparator 5-ASA, Outcome 3 Withdrawal from Study due to Adverse Event.

5-ASA for- 5-ASA com- Risk Ratio mulations parator		Weight	Risk Ratio	
n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
2/50	2/50		19.6%	1[0.15,6.82]
3/49	1/46	+	10.11%	2.82[0.3,26.12]
3/65	1/65	-	9.8%	3[0.32,28.09]
2/20	2/20		19.6%	1[0.16,6.42]
184	181		59.1%	1.64[0.61,4.42]
s), 6 (5-ASA comparato	or)			
f=3(P=0.79); I ² =0%				
3)				
3/48	4/44		40.9%	0.69[0.16,2.9]
48	44		40.9%	0.69[0.16,2.9]
), 4 (5-ASA comparator	·)			
1)				
232	225		100%	1.25[0.56,2.78]
	mulations n/N 2/50 3/49 3/65 2/20 184 s), 6 (5-ASA comparator f=3(P=0.79); I²=0% 3) 3/48 48 48), 4 (5-ASA comparator	mulations parator n/N	mulations parator n/N n/N M-H, Fixed, 95% CI 2/50 2/50 3/49 1/46 3/65 1/65 2/20 2/20 184 181 s), 6 (5-ASA comparator) f=3(P=0.79); 1²=0% 3) 3/48 4/44 48 44 48 44 48 44	mulations parator n/N n/N M-H, Fixed, 95% CI 2/50 2/50 19.6% 3/49 1/46 10.11% 3/65 1/65 9.8% 2/20 2/20 19.6% 184 181 59.1% s), 6 (5-ASA comparator) f=3(P=0.79); I²=0% 3) 3/48 4/44 40.9% 48 44 40.9%





Analysis 4.4. Comparison 4 5-ASA versus comparator 5-ASA, Outcome 4 Exclusion/Withdrawal after Entry (not due to relapse).



Comparison 5. 5-ASA (dose ranging)

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Failure to Maintain Clinical or Endoscopic Remission	10		Risk Ratio (M-H, Random, 95% CI)	Subtotals only
1.1 Asacol 4.8 g versus 2.4 g/day	1	112	Risk Ratio (M-H, Random, 95% CI)	0.8 [0.46, 1.38]
1.2 Asacol 3.2 g versus 2 g/day	1	262	Risk Ratio (M-H, Random, 95% CI)	1.07 [0.83, 1.37]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1.3 Asacol 2.4 g versus 1.2 g/day	1	156	Risk Ratio (M-H, Random, 95% CI)	0.95 [0.78, 1.16]
1.4 Asacol 1.6 g versus 0.8 g/day	1	177	Risk Ratio (M-H, Random, 95% CI)	1.01 [0.78, 1.32]
1.5 Balsalazide 6.0 g versus 3.0 g/day	2	216	Risk Ratio (M-H, Random, 95% CI)	0.76 [0.21, 2.79]
1.6 Balsalazide 4.0 g versus 2.0 g/day	1	133	Risk Ratio (M-H, Random, 95% CI)	0.66 [0.45, 0.97]
1.7 Olsalazine 2.0 g versus 1.0 g/day	1	127	Risk Ratio (M-H, Random, 95% CI)	1.01 [0.66, 1.54]
1.8 Salofalk granules 3 g versus 1.5 g OD	1	429	Risk Ratio (M-H, Random, 95% CI)	0.65 [0.49, 0.86]
1.9 Pentasa 3.0 g versus 1.5 g/ day	1	169	Risk Ratio (M-H, Random, 95% CI)	0.74 [0.48, 1.15]
2 Development of Any Adverse Event	5		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
2.1 Asacol 2.4 g versus 1.2 g/day	1	156	Risk Ratio (M-H, Fixed, 95% CI)	2.85 [0.12, 68.95]
2.2 Asacol 1.6 g versus 0.8 g/day	1	177	Risk Ratio (M-H, Fixed, 95% CI)	1.86 [1.18, 2.95]
2.3 Balsalazide 6.0 g versus 3.0 g/day	1	88	Risk Ratio (M-H, Fixed, 95% CI)	1.4 [0.88, 2.24]
2.4 Olsalazine 2.0 g versus 1.0 g/ day	1	127	Risk Ratio (M-H, Fixed, 95% CI)	1.37 [0.94, 1.99]
2.5 Salofalk granules 3 g versus 1.5 g OD	1	429	Risk Ratio (M-H, Fixed, 95% CI)	0.74 [0.61, 0.91]
3 Withdrawal from Study due to Adverse Event	7		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
3.1 Asacol 2.4 g versus 1.2 g/day	1	156	Risk Ratio (M-H, Fixed, 95% CI)	2.85 [0.12, 68.95]
3.2 Asacol 1.6 g versus 0.8 g/day	1	177	Risk Ratio (M-H, Fixed, 95% CI)	0.34 [0.04, 3.25]
3.3 Balsalazide 6.0 g versus 3.0 g/day	2	196	Risk Ratio (M-H, Fixed, 95% CI)	0.59 [0.21, 1.70]
3.4 Balsalazide 4.0 g versus 2.0 g/day	1	133	Risk Ratio (M-H, Fixed, 95% CI)	1.43 [0.54, 3.80]
3.5 Salofalk granules 3 g versus 1.5 g OD	1	429	Risk Ratio (M-H, Fixed, 95% CI)	0.98 [0.29, 3.33]
3.6 Pentasa 3.0 g versus 1.5 g/ day	1	169	Risk Ratio (M-H, Fixed, 95% CI)	1.06 [0.07, 16.69]

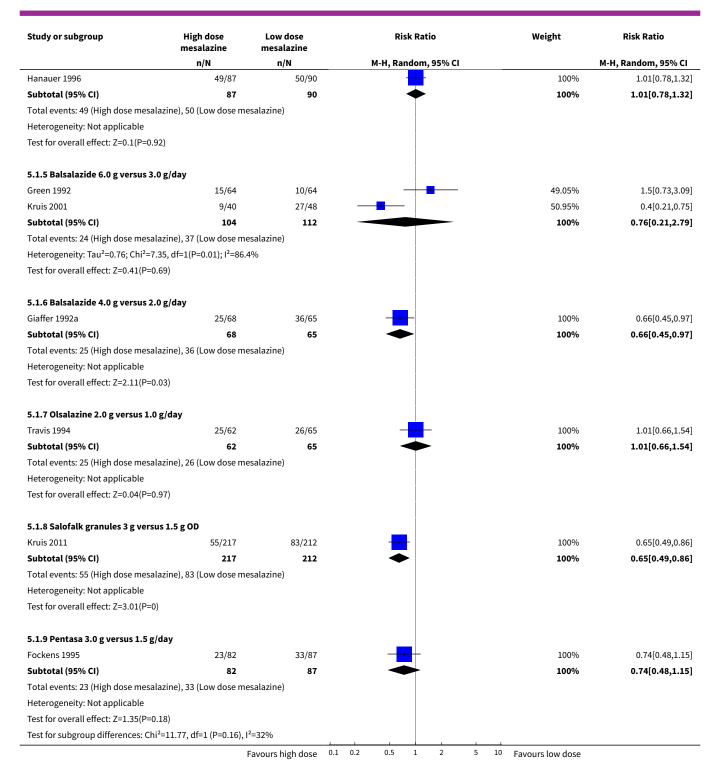


Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
4 Exculsion/Withdrawal after Entry (not due to relapse)	8		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
4.1 Asacol 2.4 g versus 1.2 g/day	1	156	Risk Ratio (M-H, Fixed, 95% CI)	0.95 [0.38, 2.40]
4.2 Asacol 1.6 g versus 0.8 g/day	1	177	Risk Ratio (M-H, Fixed, 95% CI)	1.23 [0.80, 1.90]
4.3 Balsalazide 6.0 g versus 3.0 g/day	2	196	Risk Ratio (M-H, Fixed, 95% CI)	0.47 [0.26, 0.84]
4.4 Balsalazide 4.0 g versus 2.0 g/day	1	133	Risk Ratio (M-H, Fixed, 95% CI)	1.27 [0.77, 2.12]
4.5 Olsalazine 2.0 g versus 1.0 g/day	1	127	Risk Ratio (M-H, Fixed, 95% CI)	1.75 [0.83, 3.70]
4.6 Salofalk granules 3 g versus 1.5 g OD	1	429	Risk Ratio (M-H, Fixed, 95% CI)	0.66 [0.46, 0.93]
4.7 Pentasa 3.0 g versus 1.5 g/ day	1	169	Risk Ratio (M-H, Fixed, 95% CI)	0.83 [0.44, 1.55]

Analysis 5.1. Comparison 5 5-ASA (dose ranging), Outcome 1 Failure to Maintain Clinical or Endoscopic Remission.

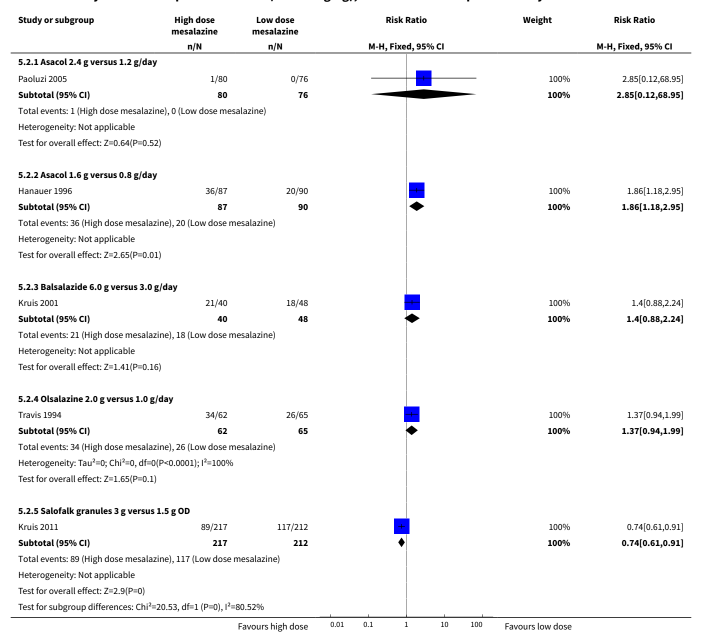
Study or subgroup	roup High dose Low dose Risk Ratio mesalazine mesalazine		Weight	Risk Ratio		
	n/N	n/N	M-H, Random, 95% CI		M-H, Random, 95% CI	
5.1.1 Asacol 4.8 g versus 2.4 g/day						
Pica 2012	16/56	20/56	- 1	100%	0.8[0.46,1.38]	
Subtotal (95% CI)	56	56		100%	0.8[0.46,1.38]	
Total events: 16 (High dose mesalazir	ne), 20 (Low dose m	esalazine)				
Heterogeneity: Not applicable						
Test for overall effect: Z=0.81(P=0.42)						
5.1.2 Asacol 3.2 g versus 2 g/day						
Deventer 2001	70/138	59/124	-	100%	1.07[0.83,1.37]	
Subtotal (95% CI)	138	124	*	100%	1.07[0.83,1.37]	
Total events: 70 (High dose mesalazir	ne), 59 (Low dose m	esalazine)				
Heterogeneity: Not applicable						
Test for overall effect: Z=0.51(P=0.61)						
5.1.3 Asacol 2.4 g versus 1.2 g/day						
Paoluzi 2005	56/80	56/76	<u> </u>	100%	0.95[0.78,1.16]	
Subtotal (95% CI)	80	76	→	100%	0.95[0.78,1.16]	
Total events: 56 (High dose mesalazir	ne), 56 (Low dose m	esalazine)				
Heterogeneity: Not applicable						
Test for overall effect: Z=0.51(P=0.61)						
5.1.4 Asacol 1.6 g versus 0.8 g/day						
		Favours high dose 0.1	0.2 0.5 1 2 5	¹⁰ Favours low dose		







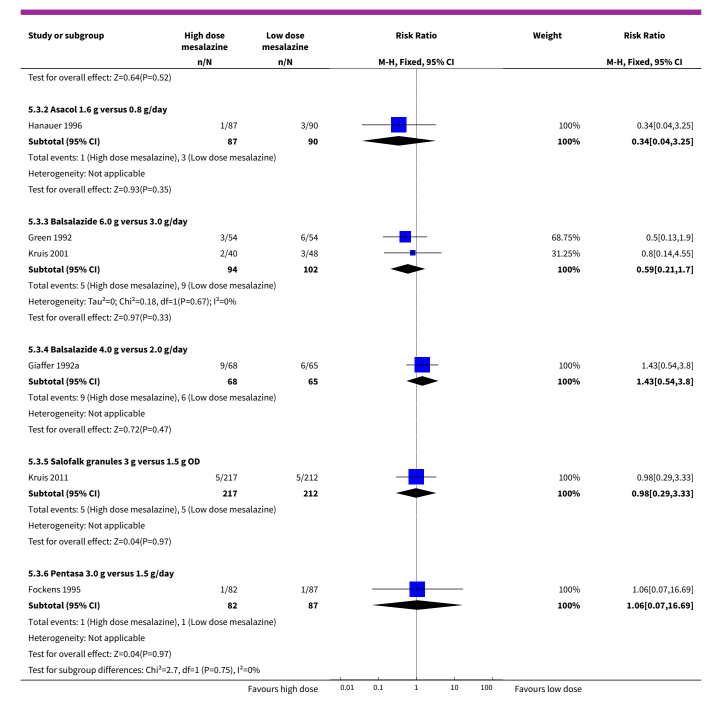
Analysis 5.2. Comparison 5 5-ASA (dose ranging), Outcome 2 Development of Any Adverse Event.



Analysis 5.3. Comparison 5 5-ASA (dose ranging), Outcome 3 Withdrawal from Study due to Adverse Event.

Study or subgroup	High dose mesalazine	Low dose mesalazine		Risk Ratio		Weight	Risk Ratio		
	n/N	n/N		М-Н,	Fixed, 95	% CI			M-H, Fixed, 95% CI
5.3.1 Asacol 2.4 g versus 1.2 g	/day								
Paoluzi 2005	1/80	0/76		-				100%	2.85[0.12,68.95]
Subtotal (95% CI)	80	76						100%	2.85[0.12,68.95]
Total events: 1 (High dose mesa	alazine), 0 (Low dose mesa	alazine)							
Heterogeneity: Not applicable									
		Favours high dose	0.01	0.1	1	10	100	Favours low dose	

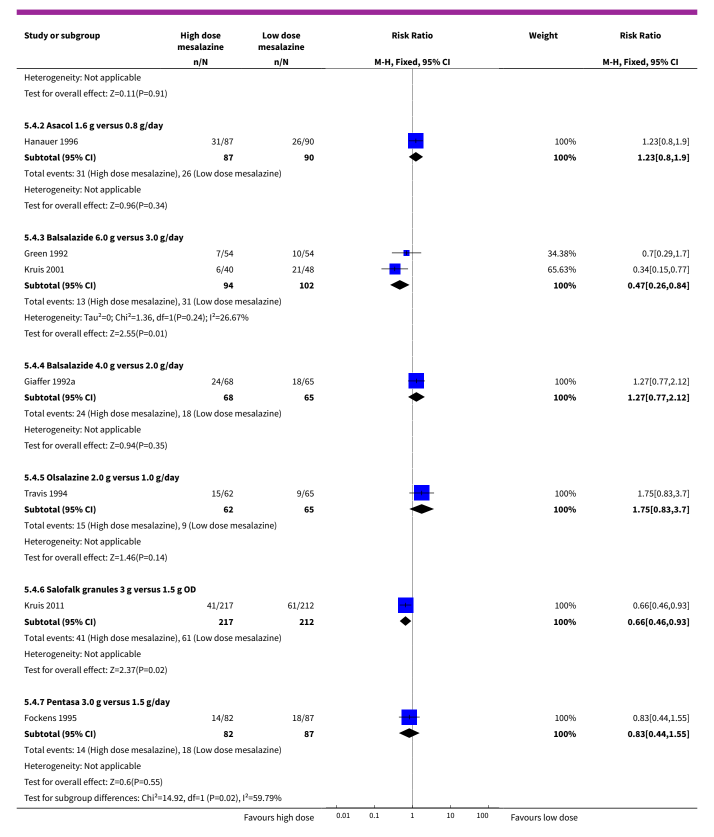




Analysis 5.4. Comparison 5 5-ASA (dose ranging), Outcome 4 Exculsion/Withdrawal after Entry (not due to relapse).

Study or subgroup	High dose mesalazine	Low dose mesalazine			•	Weight		Risk Ratio	
	n/N	n/N		М-Н	Fixed, 95	5% CI			M-H, Fixed, 95% CI
5.4.1 Asacol 2.4 g versus 1.2 g	g/day								
Paoluzi 2005	8/80	8/76			_			100%	0.95[0.38,2.4]
Subtotal (95% CI)	80	76						100%	0.95[0.38,2.4]
Total events: 8 (High dose mes	alazine), 8 (Low dose mesa	alazine)							
		Favours high dose	0.01	0.1	1	10	100	Favours low dose	







APPENDICES

Appendix 1. Search strategies

MEDLINE Search Strategy:

- 1. random\$.tw.
- 2. factorial\$.tw.
- 3. (crossover\$ or cross over\$ or cross-over\$).tw.
- 4. placebo\$.tw.
- 5. single blind.mp.
- 6. double blind.mp.
- 7. triple blind.mp.
- 8. (singl\$ adj blind\$).tw.
- 9. (double\$ adj blind\$).tw.
- 10. (tripl\$ adj blind\$).tw.
- 11. assign\$.tw.
- 12. allocat\$.tw.
- 13. crossover procedure/
- 14. double blind procedure/
- 15. single blind procedure/
- 16. triple blind procedure/
- 17. randomized controlled trial/
- 18. or/1-17
- 19. (exp animal/ or animal.hw. or nonhuman/) not (exp human/ or human cell/ or (human or humans).ti.)
- 20. 18 not 19
- 21. (colitis and ulcerat*).mp.
- 22. ulcerative colitis.mp. or exp ulcerative colitis/
- 23. (inflammatory bowel disease* or IBD).mp.
- 24. 21 or 22 or 23
- 25. 20 and 24
- 26 5-aminosalicylic acid.mp. or exp Mesalamine/
- 27. Mesalazine.mp. or exp Mesalamine/
- 28. Sulfasalazine.mp. or exp Sulfasalazine/
- 29. sulphasalazine.mp. or exp Sulfasalazine/
- 30. 26 or 27 or 28 or 29
- 8.25 and 30

EMBASE Search Strategy:



- random\$.tw.
 factorial\$.tw.
- 3. (crossover\$ or cross over\$ or cross-over\$).tw.
- 4. placebo\$.tw.
- 5. single blind.mp.
- 6. double blind.mp.
- 7. triple blind.mp.
- 8. (singl\$ adj blind\$).tw.
- 9. (double\$ adj blind\$).tw.
- 10. (tripl\$ adj blind\$).tw.
- 11. assign\$.tw.
- 12. allocat\$.tw.
- 13. crossover procedure/
- 14. double blind procedure/
- 15. single blind procedure/
- 16. triple blind procedure/
- 17. randomized controlled trial/
- 18. or/1-17
- 19. (exp animal/ or animal.hw. or nonhuman/) not (exp human/ or human cell/ or (human or humans).ti.)
- 20.18 not 19
- 21. (colitis and ulcerat*).mp.
- 22. ulcerative colitis.mp. or exp ulcerative colitis/
- 23. (inflammatory bowel disease* or IBD).mp.
- 24. 21 or 22 or 23
- 25. 20 and 24
- 26 5-aminosalicylic acid.mp. or exp Mesalamine/
- 27. Mesalazine.mp. or exp Mesalamine/
- 28. Sulfasalazine.mp. or exp Sulfasalazine/
- 29. sulphasalazine.mp. or exp Sulfasalazine/
- 30. 26 or 27 or 28 or 29
- 8. 25 and 30

Cochrane Library Search Strategy:

- 1. MeSH descriptor: [Colitis, Ulcerative] explode all trees
- 2. colitis
- 3. #1 or #2



- 4.5-ASA
- 5. 5-aminosalicylic acid
- 6. Mesalamine
- 7. Sulfasalazine
- 8. Salazosulfapyridine
- 9. Sulphasalazine
- 10. #4 or #5 or #6 or #7 or #8 or #9
- 11. #3 and #10

Cochrane IBD Specialized Register:

- 1. 5-ASA (ab/ti)
- 2.5-Amino* (ab/ti)
- 3. Mesala* (ab/ti)
- 4. Sulfa* (ab/ti)
- 5. Sulpha* (ab/ti)
- 6.1 or 2 or 3 or 4 or 5
- 7. Colitis (ab/ti)
- 8.6 and 7

WHAT'S NEW

Date	Event	Description
9 July 2015	New citation required but conclusions have not changed	Updated review with new authors
9 July 2015	New search has been performed	A new literature search was conducted on 9 July 2015. New studies added

DECLARATIONS OF INTEREST

Yongjun Wang: None known

Claire E Parker: None known

Brian G Feagan has received fees from Abbott/AbbVie, Amgen, Astra Zeneca, Avaxia Biologics Inc., Bristol-Myers Squibb, Celgene, Centocor Inc., Elan/Biogen, Ferring, JnJ/Janssen, Merck, Nestles, Novartis, Novonordisk, Pfizer, Prometheus Laboratories, Protagonist, Salix Pharma, Takeda, Teva, TiGenix, Tillotts Pharma AG and UCB Pharma for Scientific Advisory Board membership; fees from Abbott/AbbVie, Actogenix, Akros, Albireo Pharma, Amgen, Astra Zeneca, Avaxia Biologics Inc., Avir Pharma, Axcan, Baxter Healthcare Corp., Biogen Idec, Boehringer-Ingelheim, Bristol-Myers Squibb, Calypso Biotech, Celgene, Elan/Biogen, EnGene, Ferring Pharma, Roche/Genentech, GiCare Pharma, Gilead, Given Imaging Inc., GSK, Ironwood Pharma, Janssen Biotech (Centocor), JnJ/Janssen, Kyowa Kakko Kirin Co Ltd., Lexicon, Lilly, Lycera BioTech, Merck, Mesoblast Pharma, Millennium, Nektar, Nestles, Novonordisk, Pfizer, Prometheus Therapeutics and Diagnostics, Protagonist, Receptos, Salix Pharma, Serono, Shire, Sigmoid Pharma, Synergy Pharma Inc., Takeda, Teva Pharma, TiGenix, Tillotts, UCB Pharma, Vertex Pharma, VHsquared Ltd., Warner-Chilcott, Wyeth, Zealand, and Zyngenia for consultancy; payment for lectures from Abbott/AbbVie, JnJ/Janssen, Takeda, Warner-Chilcott, UCB Pharma; his institution has received grants/grants pending from Abbott/AbbVie, Amgen, Astra Zeneca, Bristol-Myers Squibb (BMS), Janssen Biotech (Centocor), JnJ/Janssen, Roche/Genentech, Millennium, Pfizer, Receptos, Santarus, Sanofi, Tillotts, and UCB Pharma



John K MacDonald: None known

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Internal sources

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External sources

• Searle Mucosal Defense Unit, Oakville, Ontario, Canada.

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INDEX TERMS

Medical Subject Headings (MeSH)

Administration, Oral; Aminosalicylic Acids [*administration & dosage]; Anti-Inflammatory Agents, Non-Steroidal [*administration & dosage]; Colitis, Ulcerative [*drug therapy] [prevention & control]; Maintenance Chemotherapy [*methods]; Medication Adherence [statistics & numerical data]; Mesalamine [*administration & dosage]; Randomized Controlled Trials as Topic; Recurrence; Remission Induction [methods]; Sulfasalazine [administration & dosage]

MeSH check words

Humans